Edited by Karl Jousten

Handbook of Vacuum Technology

Second Edition



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Second, Completely Revised and Updated Edition



Editor

Dr. Karl Jousten

Physikalisch-Technische Bundesanstalt (PTB) AG Vakuummetrologie Abbestr. 2 -12 10587 Berlin Germany

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Preface

In 2008, Wiley-VCH published a translation of the ninth edition of the German handbook "Wutz – Handbook of Vacuum Technology," named after the author of the first edition Max Wutz. This book has been a great success for five decades and the object of many requests for a translation. Since its second edition, the "Wutz – Handbook of Vacuum Technology" has become a multi-author book covering the field of vacuum science, vacuum technology, and vacuum techniques comprehensively. Since 2008, the German handbook underwent significant changes and when it could be foreseen that the English edition would run out of print, Wiley-VCH suggested to issue a second English edition "Handbook of Vacuum Technology," which is a translation of the 11th German edition of the "Wutz – Handbook of Vacuum Technology," published by Springer Vieweg. Chapter 17, however, received a new author and was newly written for this second English edition. Compared with the first English edition, also Chapters 10 and 12 were written by new authors while improvements were made in most of the other chapters according to the changes in techniques.

Although multi-author, the book aims to be read as a single-author work, a goal to which the present editor who himself has revised almost half of the content has stringently adhered to. The style is as uniform as possible, there are only recurrences where necessary, and the same symbols and notation are used throughout. Hence, the book has taken on textbook character, though it was originally intended to be used as a technical handbook.

The main idea of the book is to cover all aspects of vacuum science and technology in order to enable engineers, technicians, and scientists to develop and work successfully with the equipment and "environment" of vacuum. Beginners in the field of vacuum shall be able to start and experts shall be able to deepen their knowledge and find the necessary information and data to continue their work.

Despite the fact that the applications of vacuum technology are steadily increasing both quantitatively and qualitatively – note, for instance, that the next chip generation will be illuminated under vacuum by extreme ultraviolet (EUV) lithography – the number of scientists researching and teaching in the field is on a steady decline. Thus, another task for a book like this is to both preserve the knowledge of vacuum science and technology and enable self-studying

in the field. For this reason, the book may be at times too introductive and simple for experts and sometimes too specialized for beginners. The reader should not be discouraged when experiencing this, but rather choose the information as his/her personal level requires. Short explanations following the title of each chapter describe the contents and may help the reader to choose the right chapter for his/her needs.

We hope that also this second edition will be helpful to all readers of English interested in a comprehensive and up-to-date overview in the field of vacuum technology including its underlying science.

Even after many people read drafts and proofs, there will always be mistakes in a book of this size. If you discover such or if you have any suggestions for improvements, please send an email to the editor (karl.jousten@ptb.de). I will be glad to consider your suggestions in future editions.

June 2015

Karl Iousten Berlin, Germany

1

The History of Vacuum Science and Vacuum Technology

Dr. Karl Jousten

Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

In old Greece, before the time of Socrates, the philosophers searched for the constancy in the world, that is, what is behind the daily experience. The Greek philosopher *Democritus* (circa 460 to 375 BC) (Figure 1.1) assumed that the world was made up of many small and undividable particles that he called *atoms* (atomos, Greek: undividable). In between the atoms, *Democritus* presumed empty space (a kind of microvacuum) through which the atoms moved according to the general laws of mechanics. Variations in shape, orientation, and arrangement of the atoms would cause variations of macroscopic objects. Acknowledging this philosophy, *Democritus*, together with his teacher *Leucippus*, may be considered as the inventor of the concept of vacuum. For them, the empty space was the precondition for the variety of our world, since it allowed the atoms to move about and arrange themselves freely. Our modern view of physics corresponds very closely to this idea of *Democritus*. However, his philosophy did not dominate the way of thinking until the sixteenth century.

It was *Aristotle's* (384 to 322 BC) philosophy that prevailed throughout the Middle Ages and until the beginning of modern times. In his book *Physica* [1], around 330 BC, *Aristotle* denied the existence of an empty space. Where there is nothing, space could not be defined. For this reason, no vacuum (Latin: empty space, emptiness) could exist in nature. According to his philosophy, nature consisted of water, earth, air, and fire. The lightest of these four elements, fire, is directed upward, whereas the heaviest, earth, downward. Additionally, nature would forbid vacuum since neither *up* nor *down* could be defined within it. Around 1300, the medieval scholastics began to speak of a *horror vacui*, meaning *nature's fear of vacuum*. Nature would abhor vacuum and wherever such a vacuum may be on the verge to develop, nature would fill it immediately.

Around 1600, however, the possibility or impossibility of an evacuated volume without any matter was a much-debated issue within the scientific–philosophical community of Italy, and later in France and Germany as well. This happened at the time when the first scientists were burnt at the stake (*Bruno* in 1600).



Figure 1.1 Democritus. Bronze statue around 250 BC, National Museum in Naples.

In 1613, *Galileo Galilei* in Florence attempted to measure the weight and density of air. He determined the weight of a glass flask containing either compressed air, air at atmospheric pressure, or water. He found a value of $2.2 \,\mathrm{g} \,\ell^{-1}$ for the density of air (the modern value is $1.2 \,\mathrm{g} \,\ell^{-1}$). This was a big step forward: air could now be considered as a substance with weight. Therefore, it could be assumed that air, in some way, could also be removed from a volume.

In 1630, *Galilei* was in correspondence with the Genoese scientist *Baliani* discussing the water supply system of Genoa. *Galilei* argued that, for a long time, he had been aware of the fact that the maximum height of a water column in a vertical pipe produced by a suction pump device was about 34 feet. *Baliani* replied that he thought this was due to the limited pressure of the atmosphere!

One can see from these examples that in Italy in the first half of the seventeenth century the ground was prepared for an experiment, which was performed in 1640 by *Gasparo Berti* and in 1644 by *Evangelista Torricelli*, a professor in Florence. The *Torricelli* experiment was bound to be one of the key experiments of natural sciences.

Torricelli filled a glass tube of about 1 m in length with mercury. The open end was sealed with a fingertip. The tube was then brought to an upright position with the end pointing downward sealed by the fingertip. This end was immersed in a mercury reservoir and the fingertip removed so that the mercury inside the tube was in free contact with the reservoir. The mercury column in the tube sank to a height of 76 cm, measured from the liquid surface of the reservoir. Figure 1.2 shows a drawing of the *Torricellian* apparatus.

The experiment demonstrated that the space left above the mercury after turning the tube upside down was in fact a vacuum: the mercury level was independent of the volume above, and it could be filled completely with water



Figure 1.2 Torricelli's vacuum experiment in 1644. The level AB of mercury in both tubes C and D was equal, independent of the size of the additional volume E in tube D. (From Ref. [2].)

admitted from below. This experiment was the first successful attempt to produce vacuum and subsequently convinced the scientific community. An earlier attempt by Berti who used water was less successful.

In 1646, the mathematician Pierre Petit in France informed Blaise Pascal (Figure 1.3) about Torricelli's experiment. Pascal repeated the experiment and, in addition, tried other types of liquid. He found that the maximum height was exactly inversely proportional to the used liquid's density. Pascal knew the equally famous philosopher Descartes. During a discussion in 1647, they developed the idea of air pressure measurements at different altitudes using a *Torricellian* tube.

Pascal wrote a letter to his brother-in-law Périer and asked him to carry out the experiment on the very steep mountain Puy de Dôme, close to Périer's home.



Figure 1.3 Portrait of Blaise Pascal.

Périer agreed and on September 19, 1648 [3], he climbed the *Puy de Dôme* (1500 m) accompanied by several men who served to testify the results, which was common practice at the time. They recorded the height of the mercury column at various altitudes. From the foot to the top of the mountain, the difference of the mercury column's height was almost 8 cm and *Pascal* was very pleased: the first successful pressure measurement had been carried out! *Torricelli*, however, never enjoyed the triumph of the experiment based on his invention: he had died a year before.

Despite these experiments, the discussion between the *plenists* (no vacuum is possible in nature) and the *vacuists* (vacuum is possible) continued. One of the leading vacuists was *Otto von Guericke*, burgomaster of Magdeburg in Germany from 1645 to 1676 (Figure 1.4).

He was the first German scientist who gave experiments a clear priority over merely intellectual considerations when attempting to solve problems about nature.

Around 1650, Guericke tried to produce a vacuum in a water-filled, wooden cask by pumping out the water with a pump used by the fire brigade in Magdeburg. Although the cask was specially sealed, the experiment failed: the air rushed into the empty space above the water through the wood, developing a chattering noise. Consequently, Guericke ordered to build a large copper sphere, but when the air was pumped out, the sphere was suddenly crushed. Guericke correctly recognized atmospheric pressure as the cause and ascribed the weakness of the sphere to the loss of sphericity. The problem was solved by constructing a thicker and more precisely shaped sphere. After evacuating this sphere and leaving it untouched for several days, Guericke found that the air was seeping into the sphere, mainly through the pistons of the pump and the seals of



Figure 1.4 Portrait of Otto von Guericke in 1672. Engraving after a master of Cornelius Galle the Younger. (From Ref. [4].)

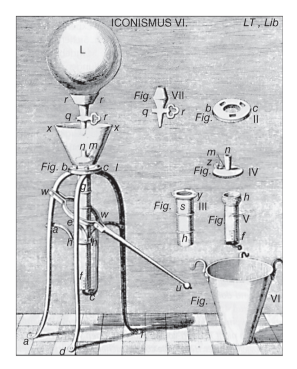


Figure 1.5 Guericke's air pump no. 3. Design for Elector Friedrich Wilhelm, 1663. (From Ref. [4].)

the valves. To avoid this, he constructed a new pump where these parts were sealed by water, an idea still used in today's vacuum pumps, but with oil instead of water.

Guericke's third version (Figure 1.5) was an air pump, which pumped air directly out of a vessel. These pumps were capable of producing vacua in much larger volumes than Torricellian tubes.

The word pump is still used for today's vacuum pumps, although they are actually rarefied gas compressors. This is due to the origin of the vacuum pump: the water pump used by the fire brigade in Magdeburg.

Guericke was also a very successful promoter of his own knowledge and experiments, which he used to catch attention for political purposes. In 1654, he performed several spectacular experiments for the German Reichstag in Regensburg. The most famous experiment demonstrating the new vacuum technique was displayed in Magdeburg in 1657.

Guericke used two hemispheres with a diameter of 40 cm, known as the Magdeburg hemispheres (Figure 1.6). One of the hemispheres had a valve for evacuation, and between the hemispheres, Guericke placed a leather ring soaked with wax and turpentine as seal. Teams of eight horses on either side were just barely able to separate the two hemispheres after the enclosed volume had been evacuated.

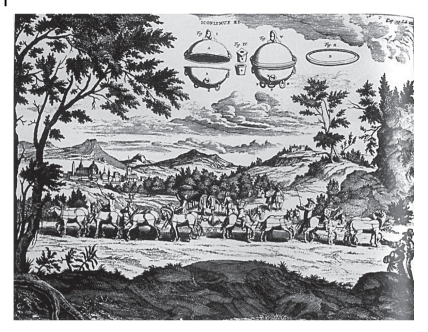


Figure 1.6 Painting of *Guericke* showing his experiment with the hemispheres to the German emperor, *Kaiser Ferdinand III*. (From Ref. [4].)

News of *Guericke's* experiment spread throughout Europe and his air pump can be considered as one of the greatest technical inventions of the seventeenth century, the others being the telescope, the microscope, and the pendulum clock.

The new vacuum technology brought up many interesting experiments. Most of them were performed by *Guericke* and *Schott* in Germany, by *Huygens* in the Netherlands, and by *Boyle* and *Hooke* in England.

Guericke showed that a bell positioned in a vacuum could not be heard; a magnetic force, however, was not influenced by the vacuum. Instead of metal, he often used glass vessels in order to make the processes in vacuum visible. For this, he used glass flasks from the pharmacist. These were called *recipients*, a word still used today for vacuum vessels. Guericke put a candle in a glass vessel and found that the candle extinguished slowly as evacuation proceeded. Huygens suspended a lump of butter in the center of a vacuum jar and, after evacuation, he placed a hot iron cap over the jar. In spite of the hot jar, the butter did not melt. Animals set into vacuum chambers died in a cruel manner. Guericke even put fish in a glass vessel, half filled with water. After evacuating the air above and from the water, most of the fish swelled and died.

Noble societies of the seventeenth and eighteenth century enjoyed watching experiments of this kind for amusement (Figure 1.7).

However, scientific experiments were performed as well during the early days of vacuum. *Huygens* verified that the free fall of a feather in a vacuum tube was



Figure 1.7 "Experiment on a bird in the air pump," 1768, by *Joseph Wright*, National Gallery, London. A pet cockatoo (top center) was placed in a glass vessel and the vessel was evacuated. The lecturer's left hand controls

the plug at the top of the glass globe. By opening it, he saves the life of the already dazed bird. The man below the "experimenter" stops the time until the possible death of the bird.

exactly equal to that of a piece of lead. *Boyle* found that the product of volume and pressure was constant, while *Amontons* in France showed that this constant was temperature-dependent (1699).

In 1673, *Huygens* attempted to build an internal combustion engine using the pressure difference between the atmosphere and a vacuum to lift heavy weights (Figure 1.8). Gunpowder, together with a burning wick, is placed in container C, arranged at the lower end of cylinder AB. The violent reaction of the gunpowder drives the air out of the cylinder through the wetted leather tubes EF. Cylinder AB cools down and produces a vacuum. The tubes EF then flatten and seal, and the atmospheric pressure drives down piston D, thus lifting weight G.

During the experiments, the importance of carefully cleaned materials became obvious and it was realized that the quality of pumps would have to be improved. Engineering improvements by *Hooke, Hauksbee* (1670–1713), and others followed. Somewhat later, the Englishman *H. A. Fleuss* developed a piston pump that he named *Geryk* in honor of *Otto von Guericke*.

However, it was not until 1855 that significantly better vacua could be produced using a pump designed by *Geissler* in Germany. *Sprengel* improved this pump in 1865 and 1873 (Figures 1.9 and 1.10), which used *Torricelli's* principle. Ten kilograms of mercury had to be lifted up and down by hand for a pump speed of about $0.004 \, \ell \, \rm s^{-1}$. About 6 h of pumping action was required to

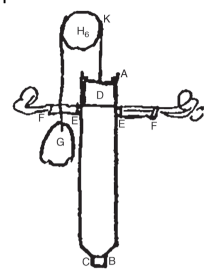


Figure 1.8 *Huygens'* explosion motor. After the explosion of gunpowder in container C, the temperature drops creating vacuum that lifts weight G. (From Ref. [3].)

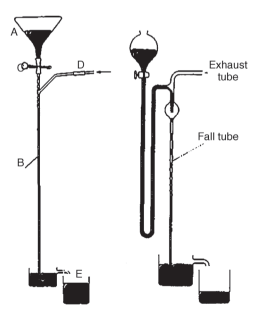


Figure 1.9 *Sprengel's* first mercury pumps of 1865. A falling mercury droplet formed a piston that drove the air downward (suction ports at D and "exhaust tube"). Later, *Sprengel* improved the pump by adding a mechanism to recover the mercury. (From Ref. [5].)

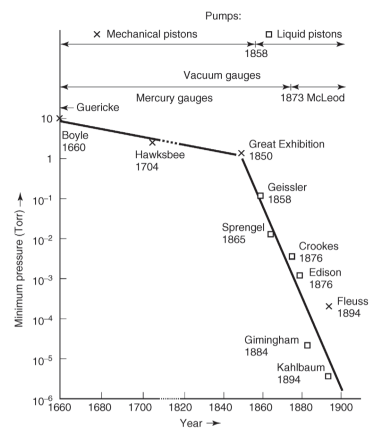


Figure 1.10 Progress in lowest generated and measured pressures in vacuum from 1660 to 1900. (Data from Ref. [6].)

evacuate a vessel of $6\,\ell$ from $0.1\,\mathrm{mmHg}$ (13 Pa) to about $2\times10^{-5}\,\mathrm{mmHg}$ (2.7 \times 10⁻³ Pa)! With these pumps, for the first time, the high-vacuum regime became available. In 1879, *Edison* used them in his *Menlo Park* to evacuate the first incandescent lamps (Figure 1.11).

The early scientists who produced vacuum still had no clear definition of a vacuum. They had no idea that air could consist of atoms and molecules, which in part are removed to produce a vacuum. Until 1874, the *Torricellian* tube was the only instrument available for measuring vacuum, and limited to about 0.5 mmHg (67 Pa). The idea of vacuum was still quite an absolute (present or not) as in the *Aristotelian* philosophy but it was not accepted as a measurable quantity. The gas kinetic theory by *Clausing*, *Maxwell*, *Boltzmann*, and others as well as the invention of the gauge by *McLeod* (1874), however, showed that vacuum indeed was a measurable physical quantity.

The *McLeod* gauge (Figure 1.12), still applied in a few laboratories today, uses *Boyle's* law. By compressing a known volume of gas by a known ratio to a higher

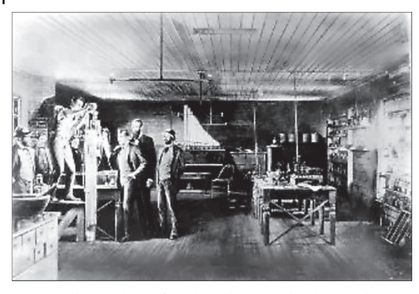


Figure 1.11 *Edison's* production of incandescent lamps in *Menlo Park* in 1879. The man standing elevated pours mercury into a *Sprengel* pump (Figure 1.9) to evacuate an incandescent lamp.

pressure, which can be measured using a mercury column, the original pressure can be calculated.

Huygens' idea of using the pressure difference between the atmosphere and a vacuum to build an engine was continued by *Thomas Newcomen* in the eighteenth century. He used condensed steam to create vacuum. Newcomen's engines were broadly used in England to pump water from deep mine shafts, to pump domestic water supplies, and to supply water for industrial water wheels in times of drought. His machines predate rotary steam engines by 70 years.

Another exciting development in the history of vacuum technology took place when *atmospheric railways* were constructed in England during the mid-nine-teenth century. Since steam locomotives at the time were rather unreliable, dirty, noisy, heavy, and not able to face steep gradients, a group of imaginative engineers conceived a plan to build clean, silent, and light trains driven by the force between the atmosphere and a vacuum on the surface of a piston placed between the rails.

In 1846, *Brunel* built such a system on the South Devon coast of England (Figure 1.13).

A continuous line of a cast iron tube was arranged centrally between the rails. The pressure difference of the external atmosphere on its rear and the rough vacuum on its front surface propelled a tightly fitted piston inside the tube. Huge stationary pumps placed in about 5 km intervals along the track generated the vacuum. The underside of the first railway coach was connected to a frame forming the rear end of the piston. Along the top of the tube was a slot closed by

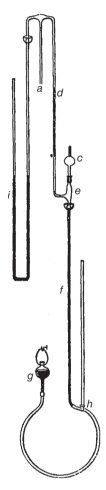


Figure 1.12 Original McLeod vacuum gauge [7]. (a) Measuring port; (b) simple siphon barometer; (c) glass bulb with a volume of 48 ml and a volume tube at the upper end having identical diameter as the measuring tube (d); (f) vertical 80 cm long

tube; and (g) reservoir of mercury. As soon as the mercury is lifted to the level of (e), the gas in (c) is compressed developing a height difference between (d) and the tube above (c) according to the volume ratio.

a longitudinal airtight valve, consisting of a continuous leather flap reinforced with iron framing.

An average speed of 103 km h⁻¹ over 6 km was reported for these trains, which was breathtaking at the time. However, atmospheric railways did not prevail. Accidents with starting trains, the lack of control by the engineer on board, and the inefficiency of the longitudinal valve (e.g., rats ate through the leather sealing), among other reasons, contributed to their demise.

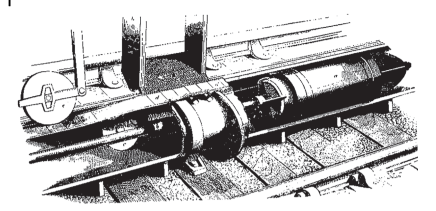


Figure 1.13 Drawing of the vacuum traction tube to propel an atmospheric railway. Piston (a) slides forward due to the action of a vacuum pump positioned in front of (to the right

of) the piston. (b) connects the piston with the leading wagon of the train. Wheel (c) lifts and opens the longitudinal valve (d) while wheel (e) closes it. (From Ref. [8].)

The large advances in physics in the second half of the nineteenth century are almost unthinkable without the aid of vacuum technology. *Hauksbee* already discovered gas discharges at the beginning of the eighteenth century. Significant progress, however, was only possible after the invention of the *Geissler* pump in 1855. Three years later, *Plücker* found that the glow of the glass wall during a gas discharge shifts when a magnetic field is applied. In 1860, *Hittorf* discovered that the rays from a cathode produce a very sharp shadow if an object is placed in between the cathode and a glass. Many scientists continued research on cathode rays, which finally led to the discovery of the electron as a component of the cathode rays by *J. Thomson* in 1898.

In 1895, *Röntgen* reported that when a discharge is pumped to less than 1 Pa, a highly penetrating radiation is produced capable of passing through air, flesh, and even thin sheets of metal. He named the beams X-rays.

In 1887, *Hertz* discovered the photoelectric effect under vacuum. In 1890, *Ramsay* and *Rayleigh* discovered the noble gases. All these experiments helped to understand the nature of vacuum: the increasing rarefaction of gas atoms and molecules. At the time, it became clear that any matter in nature consists of atoms.

In 1909, *Knudsen* [9] published a comprehensive investigation on the flow of gases through long and narrow tubes. He divided this flow into three regimes: the molecular regime at very low pressures, where the particles are so dilute that they do not interact with each other but only with the surrounding walls, the viscous regime at higher pressures, where the motion of particles is greatly influenced by mutual collisions, and an intermediate regime. This publication can be considered as the beginning of vacuum physics.

For his experiments, *Knudsen* used the so-called *Gaede* pump. *Gaede*, a professor at the University of Freiburg in Germany, was the most important

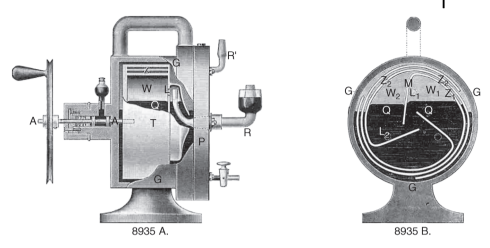


Figure 1.14 *Gaede's* mercury rotation pump. R indicates the position of the suction port. (With kind permission of the *Gaede* Foundation at *Oerlikon Leybold GmbH*, Cologne, Germany.)

inventor of vacuum pumps since *Guericke*. *Gaede's* pump was a rotary mercury pump (Figure 1.14), in which the *Torricellian* tube was wound up so that it allowed continuous pumping by rotary action. The pump was driven by an electromotor. Its pumping speed was 10 times faster than the *Sprengel*-type pump and produced vacua down to 1 mPa. However, it required an additional pump in series because it was able to compress the gas only up to 1/100 of atmospheric pressure.

The sliding vane rotary vacuum pump was developed between 1904 and 1910, based on an idea of aristocrat *Prince Rupprecht*, which dated back to 1657 [10]. *Gaede* optimized these pumps in 1935 by inventing the gas ballast, which allowed pumping condensable gases as well.

Gaede carefully studied Knudsen's work, and at a meeting of the German Physical Society in 1912 introduced his first molecular pump (Figure 1.15) [11]. Gaede used the finding that any gas molecule hitting a wall stays at its location for a while and accommodates to the wall before it leaves the same. If therefore a gas particle hits a fast moving wall, it will adopt the velocity of the wall and is transported in the direction of the motion during its sojourn time. The pumps based on this principle require very high rotor speeds and low clearances of about $20\,\mu m$ between the moving wall and the fixed wall. The pump floundered on these requirements, which were too stringent for the technology available at the time. In 1958, however, Becker utilized the principle and invented the turbomolecular pump [12], which eased the clearance problem.

In the years 1915 and 1916, *Gaede* and *Langmuir* developed the mercury diffusion pump [13,14]. Twelve years later, the oil diffusion pump followed, which was the most widespread pump until the turbomolecular pump was developed.

In addition, vacuum measurement also developed further (Figure 1.16) using other pressure-dependent properties of gases: *Sutherland* suggested to use the

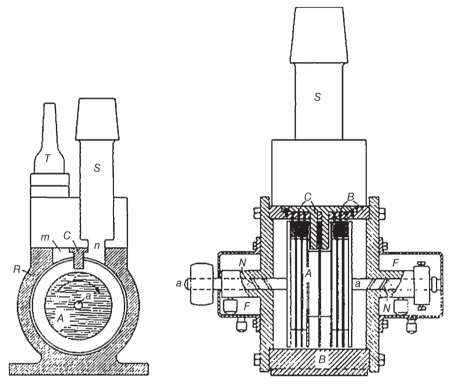


Figure 1.15 Gaede's molecular pump of 1912.

viscosity of gases in 1897. Langmuir put this principle into practice in 1913 using an oscillating quartz fiber. The decrement in amplitude of the oscillations gave a measure of gas pressure. In 1960, J. W. Beams demonstrated that the deceleration in rotational frequency of a magnetically suspended steel ball rotating at about 1 MHz under vacuum could be used as a measure of pressure. Fremerey optimized this device during the 1970s and 1980s. Pirani [15] used the pressure dependence of thermal conductivity and built the first thermal conductivity gauge in 1906. In 1909, von Baeyer showed that a triode vacuum tube could be used as a vacuum gauge. Penning invented the cold cathode gauge in 1937 in which a gas discharge is established by crossed electric and magnetic fields. During the Second World War, mass spectrometers were developed, and they became crucial parts of the weapons industry.

After World War II, it was generally believed that diffusion pumps would not be able to generate pressures below 10⁻⁸ Torr although the underlying effect was unknown. All manufacturers' pumping speed curves showed a value of zero at this point. The pressure was measured using triode gauges. During the *Physical Electronics Conference* in 1947, *Nottingham* suggested that the impingement of X-ray photons on the collector of the triode causing secondary electrons might

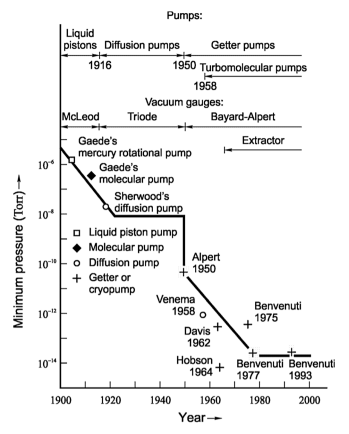


Figure 1.16 Progress in lowest pressures generated and measured in the twentieth century. (Data from Ref. [6].)

be the reason for this lower pressure limit. This was a breakthrough. A competition for a significant improvement of the ion gauge started, which *Nottingham's* own group did not win, to his regret. Instead, in 1950, *Bayard* and *Alpert* [16] succeeded with an idea as simple as ingenious (Figure 13.48).

Since all vacuum gauges except for the *McLeod* and the *Torricellian* tube had to be calibrated, and because, at the same time, vacuum industry grew to an important branch (see Chapter 2), independent metrological laboratories were set up in state-owned institutes in the late 1950s. The first were established at the *National Physical Laboratory (NPL)* in England. The *Laboratory for Vacuum Physics* (today: *Vacuum Metrology*) at the *Physikalisch-Technische Bundesanstalt*¹⁾ (*PTB*) in Germany followed in 1966, and in the 1970s the *Vacuum Laboratory* at the *National Bureau of Standards (NBS*; today: *NIST*) in the United States.

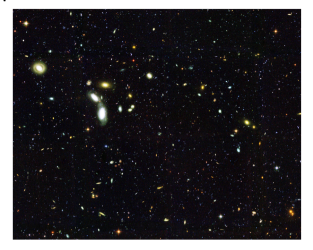


Figure 1.17 In between galaxies, there are small volumes of a few dm³ without any massive particle (absolute or ideal vacuum). Oldest known galaxies pictured by the Hubble Space Telescope. (Courtesy of NASA.)

Coming back to the philosophical considerations at the beginning of this chapter, let us make a concluding remark on the nature of vacuum from the point of view of today's physics [17,18]: without any doubt, there are macroscopic areas, for example, small volumes between galaxies, where there is no single molecule (Figure 1.17). For such a volume, the term *absolute vacuum* was introduced. We know today, however, that even absolute vacuum is not empty (in terms of energy). Otherwise, it would not be in accordance with the laws of nature. A vacuum energy with still unknown nature, which may be related to the cosmological constant introduced by *Einstein*, permits particles to be generated spontaneously by fluctuating quantum fields for short time intervals, even in *absolute vacuum*. In this sense, there is no space in the world, which is truly empty.

References

- 1 Aristotle (1997) *Physica*, Akademie Verlag,
- 2 Middleton, W.E.K. (1964) The History of the Barometer, John Hopkins University Press, Baltimore, MD.
- 3 Sparnaay, M.J. (1992) *Adventures in Vacuum*, Elsevier Science, North-Holland, p. 78.
- 4 de Guericke, O. (1672) Experimenta Nova Magdeburgica de Vacuo Spatio, Amstelodami.
- 5 Zur Geschichte der Vakuumtechnik, six reports in *Vak.-Tech.*, **35** (4/5), 99–157 (1986).
- **6** Redhead, P.A. (1999) The ultimate vacuum. *Vacuum*, **53**, 137–149.
- 7 McLeod, H.G. (1874) Apparatus for measurement of low pressures of gas. *Philos. Mag.*, 48, 110 ff.; *Proc. Phys. Soc.*, 1, 30–34 (1874).

- 8 Madey, T.E. (1984) Early applications of vacuum, from Aristotle to Langmuir. J. Vac. Sci. Technol. A, 2, 100-117.
- 9 Knudsen, M. (1909) Die Gesetze der Molekularströmung und der inneren Reibungsströmung der Gase durch Röhren. Ann. Phys. Lpz., 28, 75 - 130.
- 10 Redhead, P.A. (2005) Vacuum and the electron tube industry. J. Vac. Sci. Technol. A, 23, 1252-1259.
- 11 Gaede, W. (1912) The external friction of gases and a new principle for vacuum pumps: The molecular pump. Z. Phys., 13,
- 12 Henning, J. (1991) Die Entwicklung der Turbomolekularpumpe. Vak. Prax., 1, 28 - 30.
- 13 Gaede, W. (1915) Die Diffusion der Gase durch Quecksilberdampf bei niedrigen Drücken und die Diffusionsluftpumpe. Ann. Phys., 6, 357-392.
- 14 Langmuir, I. (1916) Gen. Electr. Rev., 19,
- 15 Pirani, M. (1906) Selbstanzeigendes Vakuum-Meßinstrument. Verh. Dtsch. Phys. Ges., 8, 686-694.
- 16 Bayard, R.T. and Alpert, D. (1950) Extension of low pressure range of the ionization gauge. Rev. Sci. Instrum., 2, 571-572.
- 17 Genz, H. (2004) Nichts als das Nichts -Die Physik des Vakuums, Wiley-VCH Verlag GmbH, p. 266.
- 18 Bojowald, M. (2011) Alles aus dem Nichts. Phys. J., 10 (3), 37.

Further Reading

- Auwaerter, M. (1983) Das Vakuum und Wolfgang Gaede. Vak.-Tech., 32 (8), 234-246.
- Dunkel, M. (1978) Gedenken an Wolfgang Gaede. Vak.-Tech., 27, 99-101.
- Gaede, H. (1954) Wolfgang Gaede, Der Schöpfer des Hochvakuums, A. Braun, Karlsruhe, p. 127.
- Hablanian, M.H. (1984) Comments on the history of vacuum pumps. J. Vac. Sci. Technol. A, 2, 118-125.
- Hoppe, E. (1936/1965) Geschichte der Physik, Vieweg, Braunschweig.
- Jahrreiß, H. (1987) Otto von Guericke (1602-1686) in memoriam. J. Vac. Sci. Technol. A, 5, 2466-2471.
- Madey, T.E. (1984) History of Vacuum Science and Technology, AIP, New York.
- Mulder, Th. (1987) Otto von Guericke. Vak.-Tech., 35 (4/5), 101-110.
- Redhead, P. (1984) The measurement of vacuum pressures. J. Vac. Sci. Technol. A, 2, 132-138.
- Singleton, J.H. (1984) The development of valves, connectors, and traps for vacuum systems during the 20th century. J. Vac. Sci. Technol. A, 2, 126-131.
- von Guericke, O. (1968) Neue Magdeburger Versuche über den leeren Raum (German by H. Schimank), VDI-Verlag, Düsseldorf, p. 291.

2

Applications and Scope of Vacuum Technology

Dr. Karl Jousten

Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

By the time *Edison* produced incandescent light bulbs, vacuum technology left the niche of a pure laboratory technique. Five hundred automatic *Sprengel* pumps were utilized in *Edison's* first production site. Since then, many other important industries, for example, microelectronics to name only one, developed that required vacuum technology. From an economic point of view, these industrial applications are much more important for vacuum industry than its applications in physics research. According to a survey of engineering federations in Europe, the United States, and Japan [1], about 40% of the sales of vacuum-related equipment of companies in these regions go into semiconductor industry. This is by far the largest segment of the vacuum technology market. The significance of vacuum technique for physical research, however, has not dropped at all. Hardly any physical experiment is conducted outside of a vacuum environment.

Figure 2.1 provides a selection of industrial applications of vacuum technique under different vacuum regimes; Figure 2.2 shows applications in physical research methods. Both lists are not intended to be complete. They rather show the variety and diversity in the field of applications of vacuum technology. Even for products in everyday use, vacuum technology plays an important role, for example, freeze drying or vacuum packaging of food, coating of PET lemonade bottles to reduce the loss of carbon dioxide gas, coating of architectural glass to reduce the loss of thermal energy, or recycling of mercury from batteries or electronic waste.

Applications of vacuum technology require a pressure range of about 15 orders of 10, from 10^{-10} Pa in the extreme vacuum to 10^5 Pa at atmospheric pressure. This is an enormous challenge for vacuum measuring technique as well as the applied vacuum pumps and materials. Table A.20 lists a selection of pumping systems used in different areas.

Some important new research areas such as life science, analysis of polymers, and catalytic research require vacuum for analytical tools but higher (atmospheric) pressure for the targets under investigation. This is accomplished with pressure stages. Additionally, in electron beam welding, the target is at

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Industrial applications o	f vacuum technology
Vacuum metallurgy Annealing, melting, casting of metals Melting by electron beam Degassing of metal or steel	10⁻³ Pa 10³ Pa
Crystal growth Zone melting of silicon	10 ⁻⁴ Pa 10 ⁻¹ Pa
Electron beam melting Physical vapor deposition	10 ^{-₃} Pa 10⁵ Pa 10 ^{-₃} Pa 1 Pa
Architectural glass coating Coating/hardening of tools Wear protection Coating of PET bottles Coating of optical devices, glasses, lenses Decorative coating Metallization of plastic material and foils Magnetic memories	s, mirrors
Chemical vapor deposition	1 Pa 10³ Pa
Medicine and medical engineering X-ray tubes, cancer therapy Sterilisation of instruments, wound healing	10 ⁻⁷ Pa 10⁴ Pa
Drying and degassing Degassing of liquids Casting of resin and lacquer Casting and drying of plastic materials Drying of insulating materials	10 ⁻² Pa 10⁴ Pa
Recycling	10 Pa 10⁴ Pa
Chemical industry Laboratory technique Processes	10² Pa 10³ Pa
Eroozo dincina	1 Do 10 ⁴ Do



Food processing 10² Pa ... 10⁴ Pa Food packaging 10⁻⁷ Pa ... 10⁻¹ Pa **Electrical engineering**

TV picture tubes Monitors, oscilloscopes Light bulbs, fluorescent tubes Transmitter tubes, receiver tubes Vacuum high power switches

Pharmaceutical industry

10⁻⁵ Pa ... 10² Pa Semiconductor technology

Wafers: oxidation, plasma etching, ion implantation Mobile phones Flat panel displays CD metallization

EUV lithography











Figure 2.1 Selected industrial applications of vacuum technology.

Vacuum technology in the research area

Biotechnology	10 ⁻⁸ Pa–10 ⁵ Pa	
Elementary particle physics	10 ⁻ Pa–10 ⁻⁶ Pa	CERN accelerator
Gravitational wave detectors	10 ⁻⁸ Pa−10 ⁻⁶ Pa	SPS
Growth of thin films Molecular beam epitaxy Nuclear fusion	10 ⁻⁶ Pa–1 Pa 10 ⁻¹⁰ Pa–10 ⁻⁸ Pa 10 ⁻⁶ Pa–10 ⁻¹ Pa	
Mass spectrometers	10⁻⁵ Pa–10⁻³ Pa	
Materials research Neutron scattering	10 ⁻⁴ Pa–10 ⁻¹ Pa	Fusion research in the ASDEX, IPP,
Metrology Mass determination Radiometry	10 ⁻⁸ Pa–10 ⁻¹ Pa	Germany
Nanotechnology (STM, AFM) Surface analytics SIMS SEM, TEM ESCA/XPS, UPS AES, LEED Ion sources	10 ⁻⁸ Pa−10 ⁵ Pa 10 ⁻⁹ Pa−10 ⁻¹ Pa	Neutron scattering on
Plasma research	10 ⁻⁷ Pa –10 ³ Pa	silicon
Synchrotron radiation Soft to hard X-radiation	10 ⁻⁸ Pa−10 ⁻⁵ Pa	
Low-temperature research	10 ⁻⁸ Pa−10 ⁻¹ Pa	
Space simulation	10⁻⁵ Pa−10⁻³ Pa	Surface analytics by Omikron

Figure 2.2 Selected physical research methods operated under vacuum.

environmental pressure whereas electrons are accelerated under high-vacuum conditions. A new branch of cancer therapy using ion beams represents another example where the target (patient) is at atmospheric pressure, but beams are accelerated in ultrahigh and high vacuum.

In the microelectronics industry (Figure 2.3), vacuum is mainly necessary for producing thin oxide layers, plasma etching, chemical and physical vapor deposition, and ion implantation. A good portion of the investment for a new DRAM factory is spent on vacuum technology. As the integration on chips increased, the purity of vacuum as well as of process gases gained importance [2–5]. Also, exhaust management for health protection of staff and for protecting the environment is an important issue [6]. The service intervals of pumps in the semiconductor industry were extended greatly by utilizing dry (oil-free) pumps.

The increased corrosion resistance of dry pumps is also of great value in the fluorination of plastics [7]. Surfaces of synthetic materials require activation by fluorine as a pretreatment for coating and gluing.

Since around the year 2000, sustainable energy plants have become a strong market for vacuum technology. The production of solar cells needs vacuum for the manufacturing of silicon crystals, thin film coating (also for thin film solar cells), and final laminating. Solar thermal power stations need glass tubes coated in vacuum. Rotors in wind turbines are manufactured from fiber-reinforced materials that have been fabricated by vacuum infusion. They are mounted on their posts by vacuum lifts able to carry weights of 32 t. In the inner part of the wind turbine, vacuum contactors limit and switch electrical currents of more than 2000 A.



Figure 2.3 Vacuum technology is critical for microelectronics industry. (Photo by the Max-Planck Society.)

A less known application of vacuum technology, not listed in Figure 2.1, is water treatment, for example, of long-distance heating water or high-purity water, used mainly for reducing oxygen content [8] but also for wastewater treatment [9]. Problematic wastewater is evaporated and the distillate can be recovered.

In the automotive industry, rough- and fine-vacuum pump systems are used before the filling of brake systems, servo-steering systems, and air-conditioning systems [10]. Vacuum methods are employed to check the tightness of such systems and engines.

Vacuum is of great importance for heat insulation purposes, be it in cryotechnology, refrigerators, or even buildings [11]. Evacuated, nanoporous foams yield 10 times higher heat flow resistance than conventional foams.

To mention more of a curiosity, acoustic characteristics of wood for musical instruments can be improved by applying vacuum heat treatment [12].

Surface analysis is certainly the main field of application in research. In this area, the transition between industrial and other applications is smooth. An example is vacuum metallization of forensic traces by vaporization sources [13]: fingerprints become visible and identifiable by metallization with zinc or gold.

The longest vacuum system in the world is the 27 km long vacuum tube of the LEP accelerator (Large Electron Positron Collider) of CERN, placed in a subterranean tunnel near Geneva. It is not in operation currently because the vacuum tube of the LHC (Large Hadron Collider) is built into the same tube. However, the latter will soon be completed. These facilities provide a basis for investigating the elementary particles of all matter and of the processes that occurred shortly after the Big Bang gave birth to our universe.

Even larger in terms of volume are the vacuum systems of gravitational wave detectors constructed at several sites across the globe [14]. The LIGO detectors (Laser Interferometer Gravitational Wave Observatory) in Washington and Louisiana feature two vacuum tubes each with a length of 4km and a diameter of 1.2 m (Figure 2.4).

Space simulation chambers require even larger volumes. The biggest vacuum space chamber, used by NASA, is located in Sandusky, Ohio, USA (Figure 2.5).

The probably largest single compact ultrahigh vacuum chamber of the world, with 1400 m³, was installed in 2006 in the Research Center Karlsruhe (now Karlsruhe Institute of Technology). The huge chamber contains the electron energy spectrometer by which means the mass of the neutrino shall be determined (Figure 2.6).

Sufficiently efficient, economical, and easy-to-operate vacuum pumps are available for all vacuum ranges.

• For the rough-vacuum regime $(10^5 \text{ to } 10^2 \text{ Pa})$

- Side channel blowers
- Rotary lobe blowers
- Dry rotary pumps
- Claw vacuum pumps
- Diaphragm pumps
- Liquid ring pumps (combined with vapor jet pumps, if applicable)

• For the rough- and fine-vacuum regime - Oil-sealed vacuum pumps (gas ballast pumps) $(10^2 \text{ to } 10^{-1} \text{ Pa})$ - Screw pumps - Spiral pumps - Scroll pumps - Roots pumps - Vapor jet pumps • For the high-vacuum regime - Diffusion pumps $(10^{-1} \text{ to } 10^{-5} \text{ Pa})$ - Turbomolecular pumps - Multistage roots pumps • For the high- and ultrahigh-vacuum - Turbomolecular pumps regime ($< 10^{-5} \text{ Pa}$) - Ion getter pumps - Titanium sublimation pumps - NEG pumps - Cryopumps

To connect the various components (pumps, valves, vacuum gauges, pipes, chambers, vapor traps, and other accessories), a range of standardized flanges are available up to large diameters (DN 1000), as well as carefully developed and thoroughly tested welded and brazed joints. For leak testing, the helium leak detector has become the standard test instrument. Its sensitivity is sufficient to detect and localize even the smallest leaks that could affect the performance of an apparatus or plant. National and international standardization activities

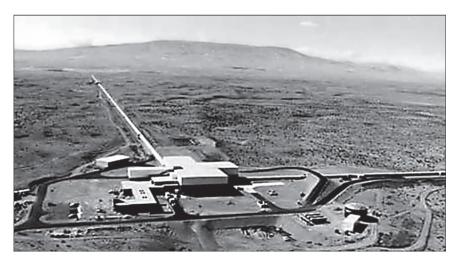


Figure 2.4 The gravitational wave detector LIGO near Hanford in the desert of the State of Washington in the United States. Each arm of the Michelson interferometer is under ultrahigh vacuum and 4 km long.



Figure 2.5 Largest space simulation chamber in the world in Sandusky, Ohio, USA. The Plum Brook Station Space Power Facility is part of NASA's Glenn Research Centre. The vacuum chamber has a diameter of about 30 m and a height of about 40 m. (Photo by NASA.)



Figure 2.6 Transportation of the main spectrometer chamber of the Karlsruhe Neutrino Project (KATRIN) through Leopoldshafen-Eggenstein on November 25, 2006. (Picture courtesy of KIT.)

covering the area of vacuum technology (Table A.22) have helped much to simplify design, operation, and maintenance of vacuum plants, as well as to increase their flexibility and make them economical. Standardized measuring methods are available for a series of vacuum technological quantities, such as the pumping speed of pumps or the critical backing pressure. Vacuum plants can be constructed in virtually any size.

The considerable advances in vacuum technology of the past decades were triggered mainly by increasing technological requirements arising from the chemical and process engineering side [15]. Most recent developments were concerned particularly with the following problems:

- Improved purity of vacuum by dry rough pumps.
- · Development of single-stage pumps.
- A comprehensive calibration system for vacuum gauges.
- Reduction of outgassing rates for bakeable and non-bakeable vacuum systems.

The range of problems concerning pure vacuum is associated particularly with the production of hydrocarbon-free vacuum. In the past, it was created preferably with sorption and condensation pumps. Research on their behavior used the methods of surface and boundary layer physics, which focuses on the interaction of gases and solid surfaces. Thus, a close association between this discipline and that of vacuum physics and technology evolved. Of the numerous research methods and procedures, those of electron spectroscopy and secondary ion mass spectrometry have found wide-scale applications. These processes, carried out under ultrahigh-vacuum conditions, are applied routinely in analytics.

Today, a large number of mechanical pump varieties such as mechanical pumps, piston pumps, screw pumps, claw pumps for generating dry, that is, carbon-free, vacuum, are available. They serve as roughing pumps for highvacuum pumps as well as for generating rough and fine vacuum. In addition to the advantage of an oil-free vacuum, these types of pumps are robust against aggressive media and dust and feature higher vapor tolerance. Due to their prolonged service intervals and their improved environmental compatibility compared with oil-sealed pumps, their cost of ownership is low.

Generating vacuum and maintaining it under various operating conditions has become a routine task. Since the 1970s, the number of physicists dealing with vacuum science is in steady decline. Today, advances in vacuum technology are mainly engineering developments. The withdrawal of science from the field bears the risk that the basics of vacuum science and technology play a less important role in the educational work of universities. Thus, today's state of the art in vacuum technology might face deterioration in practical applications. Since vacuum science and technology is a basic science for many areas of industry and other sciences, this development would have many undesired effects. Continuing the education in the field and keeping it up to date is one of the tasks of this book.

References

- 1 EVTA (European Vacuum Technology Association), AVEM International (Association of Vacuum Equipment Manufacturers), and IVIA (Japanese Vacuum Industry Association) (2014) Press release of the year 2004. EVTA secretary with VDMA, Fachverband Kompressoren, Druckluft- und Vakuumtechnik, Frankfurt, Germany.
- 2 Arthur, J.R. (1987) Vacuum gauging in the electronics industry. J. Vac. Sci. Technol. A, 5, 3230–3231.
- 3 O'Hanlon, J.F. and Parks, H.G. (1992) Impact of vacuum equipment contamination on semiconductor yield. J. Vac. Sci. Technol. A, 10, 1863-1868.
- 4 Waits, R.K. (1994) Controlling your vacuum process: effective use of a QMA. Semicond. Int., 79-84.
- 5 Buckley, M.E. (1993) Process control in the semiconductor manufacturing environment using a high pressure quadrupole mass spectrometer. Vacuum, 44, 665-668.
- 6 Gennermann, D. (2000) Abgasmanagement von Vakuumprozessen moderner Halbleiter-Produktionsanlagen. Vak. Forsch. Prax., 1, 29-33.

- 7 Fischer, S. (2000) Fluorvorbehandlungsanlagen mit trockenlaufenden Vakuumpumpen. Vak. Forsch. Prax., 2, 112-113.
- 8 Zilly, G. (2000) Entgasung von Wasser unter Vakuum. Vak. Forsch. Prax., 3, 180-183.
- 9 Schneider, W. (1996) Vakuum in der Abwasseraufbereitung. Vak. Forsch. Prax., **2**, 92–96.
- 10 Nikutta, K.-H. (1993) Vakuum-Befüllverfahren beim Automobil. Vak. Forsch. Prax., 3, 159-164.
- 11 Fricke, J., Heinemann, U., and Ebert, H.P. (2008) Vacuum insulation panels - from research to market. Vacuum, 82, 680-690.
- 12 Hix, P. (1992) Vakuum und Musik. Vak. Forsch. Prax., 4, 271-272.
- 13 Herrmann, R. and Rustler, P. (2002) Vakuum-Metallisierung als daktyloskopische Spurensicherungsmethode bei der deutschen Polizei. Vak. Forsch. Prax., 1, 30-32.
- 14 Faber, S. (1994) Gravity's secret. New Scientist, November, 40-44.
- 15 Hobson, J.P. (1984) The future of vacuum technology. J. Vac. Sci. Technol. A, 2, 144-149.

3

Gas Laws and Kinetic Theory of Gases

Prof. Dr. Wolfgang Jitschin

University of Applied Sciences, Vakuumlabor, Wiesenstr. 14, 35390, Giessen, Germany

This chapter explains the most important fundamentals of vacuum physics, focusing on the macroscopic equation of state, the kinetic theory of gases, and the description of transport phenomena.

3.1 Description of the Gas State

3.1.1

State Variables

Due to the bond between its molecular particles, a solid or liquid substance occupies a certain volume hardly influenced by ambient conditions (temperature, pressure, etc.). Therefore, this volume is an inherent property of the substance. A gas behaves differently: when a container holds a certain amount of gas, the gas spreads across the complete inner volume of the container and fills it homogeneously. The larger the container, the thinner the gas. The container's $volume\ V$ determines volume as well as state of the gas.

The gas in the container exerts a force on the walls of the container (Figure 3.1). A larger wall area is subject to a larger force than a smaller wall area. Therefore, it is convenient to introduce the term *pressure* p. The quantity *pressure* is defined as the ratio of the force F, exerted perpendicularly to a surface element of the container's wall, to the area of this surface element A:

$$p \coloneqq \frac{F}{A}.\tag{3.1}$$

In vacuum technology, the term *pressure* usually refers to *absolute* pressure (based on an ideal vacuum). Pressure is an important quantity when describing the gas state.

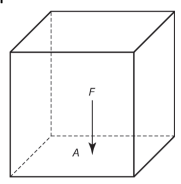


Figure 3.1 Pressure exerted to the walls of a container by a gas.

The word vacuum typically means a dilute gas or the corresponding state at which the pressure or density is lower than that in the surrounding atmosphere (ISO 3529/1). DIN 28400/1, on the other hand, defines vacuum as a state where pressure is smaller than 30 kPa (300 mbar), which is the lowest pressure that may exist on the surface of the earth.

The defining equation (3.1) shows that pressure is a derived quantity. The unit of pressure in the International System of Units (SI) is given by

$$[p] = \frac{[F]}{[A]} = \frac{\text{newton}}{\text{meter}^2} = \text{pascal} = \text{Pa}.$$
 (3.2)

One pascal (unit symbol Pa), therefore, is the pressure at which a force of 1 N $(= 1 \text{ kg m s}^{-2})$ is exerted perpendicularly to a flat surface of 1 m^2 .

A number of additional pressure units are in use; the most important are listed in Table 3.1.

According to SI, the only additional unit accepted besides Pa is bar (and mbar). The unit mmHg is often used in medicine (for blood pressure, internal eye pressure). In the United States, the units torr and, for higher pressures, psi are common in vacuum technology.

A certain pressure value, corresponding approximately to the pressure of atmospheric air at sea level, that is, normal atmosphere, has been defined as standard pressure p_n (ISO 554, ISO 3529/1, DIN 1343):

$$p_{\rm n} \coloneqq 101\,325\,\text{Pa} = 1013.25\,\text{mbar}$$
 (3.3)

In vacuum-technology applications, where the value of negative pressure with respect to ambient pressure is of interest (e.g., lifting devices with a vacuum sucker), the term relative vacuum is also used. At normal pressure, the relative vacuum is 0%, whereas it is 100% in an ideal vacuum. For any pressure p, relative vacuum is calculated according to

Relative vacuum :=
$$\frac{p_n - p}{p_n} \times 100\%$$
. (3.4)

Table 3.1 Pressure units according to ISO 3529/1, Appendix A.

Unit symbol	Unit, definition	Conversion
bar	Bar	1 bar = 10 ⁵ Pa
Mbar	Millibar	1 mbar = 100 Pa
Torr	Torr = $1/760$ of standard pressure p_n	1 Torr = $101 \ 325/760 \ Pa \approx 133.322 \ Pa \approx 4/3 \ mbar$
mmHg	Millimeters of mercury = the pressure exerted at the bottom of a vertical column of mercury, 1 mm deep, at standard acceleration due to gravity and at 0 °C	1 mmHg ≈ 133.322 Pa
μ	Micron = micrometers of mercury = the pressure exerted at the bottom of a vertical column of mercury, 1 µm deep, at standard acceleration due to gravity and at 0°C	1 μmHg ≈ 0.133322 Pa
psi	Pound-force per square inch = pressure due to weight (at standard acceleration due to gravity) of one American pound to an area of one square inch	1 psi ≈ 6894.76 Pa

Standard acceleration due to gravity is $g_n = 9.80665 \,\mathrm{m\,s^{-2}}$.

Example 3.1

A lifting device for aluminum sheet metal (or glass panes) establishes a pressure of $p = 120 \,\mathrm{mbar}$ in its vacuum suckers. The relative vacuum calculates to $[(1013 - 120) \text{ mbar}/1013 \text{ mbar}] \times 100\% = 88\%.$

The pressure of a gas in a container changes when the temperature changes. Therefore, also temperature is an important quantity, characterizing the gas state. In everyday life, temperature is usually given in Celsius (centigrade). Here, we will use ϑ (theta) as a symbol for Celsius temperature, $[\vartheta] = C$ (°C).

For characterizing the gas state, it is convenient to use the thermodynamic temperature (referring to absolute zero) rather than using Celsius. This temperature is referred to as Kelvin temperature, the symbol is T, and the unit [T] is K

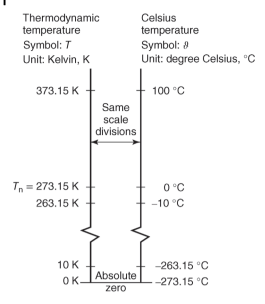


Figure 3.2 Comparison of Celsius and Kelvin temperature scales.

(Kelvin). 1 K is defined as the 1/273.16 fraction of the temperature at the triple point (θ = 0.01 °C) of pure water.

Then, the normal freezing point of water is $T_{\rm n}=273.15$ K. Thus, the relationship between the thermodynamic temperature T and the Celsius temperature ϑ is (Figure 3.2)

$$\frac{T}{K} := \frac{\theta}{C} + 273.15. \tag{3.5}$$

The different temperature scales are compared in Figure 3.2.

International agreements promoted introduction of the thermodynamic-temperature scale [1]: the International Temperature Scale of 1990 (ITS-90) published by the Consultative Committee for Thermometry (CCT) of the International Committee for Weights and Measures (CIPM) is an equipment calibration standard for making measurements on the Kelvin and Celsius temperature scales. ITS-90 is an approximation of the thermodynamic temperature scale that facilitates the comparability and compatibility of temperature measurements internationally by defining calibration points (Table 3.2), that is, triple and freezing temperatures of pure substances. Temperature values related to ITS-90 are denoted by the subscript 90. As reference gauges, ³He- and ⁴He-vapor-pressure thermometers are used in the temperature range of 0.65–5 K, gas thermometers for 3–24.6 K, platinum resistance thermometers above 13.8 K, and spectral pyrometers above 1235 K.

1084.62

Equilibrium condition T_{90} (K) t₉₀ (°C) Helium vapor pressure 3 to 5 -270.15 to -268.15 Triple point of equilibrium hydrogen^{a)} 13.8033 -259.3467 Vapor pressure of equilibrium hydrogen^{a)} \approx 17 to \approx 20.3 \approx -256.15 to \approx -252.85 Triple point of neon 24.5561 -248.5939 Triple point of oxygen 54.3484 -218.7916Triple point of argon 83.8058 -189.3442Triple point of mercury 234.3156 -38.8344Triple point of water 273.16 0.01 Melting point of gallium 302.9146 29.7646 Freezing point of indiumb) 429.7485 156.5985 Freezing point of tin^{b)} 505.078 231.928 Freezing point of zincb) 692.677 419.527 Freezing point of aluminum^{b)} 933.473 660.323 Freezing point of silverb) 1234.93 961.78 Freezing point of gold^{b)} 1337.33 1064.18

Table 3.2 Fixed reference points of ITS-90 [1].

1357.77

Freezing point of copper^{b)}

The temperature of the melting point of water is referred to as standard temperature (ISO 554, ISO 3529/1, DIN 1343):

$$T_n := 273.15 \text{ K}, \quad \vartheta_n := 0.00 \,^{\circ}\text{C}.$$
 (3.6)

If the pressure and temperature of a gas each are of standard values, the gas is in standard condition, that is, normal atmosphere.

To summarize, three state variables characterize the condition of a gas in a closed container:

- volume V;
- pressure p;
- temperature T or ϑ .

3.1.2

Extensive Quantities

The amount of a gas (or of a liquid or solid) can be specified in different ways:

- mass *m*:
- particle number *N*;
- amount of substance ν .

a) At room temperature, equilibrium hydrogen (e-H₂) is a mixture in which the hydrogen nuclei in 75% of the H₂ molecules spin in the same direction (ortho-H₂), and the nuclei in 25% of the H₂ molecules spin in opposite directions (para-H₂).

b) Freezing points at pressure 101 325 Pa.

The unit of mass *m* in the International System of Units (SI) is kg (kilogram). However, mass often is a rather inconvenient unit to describe the amount of a gas. On one hand, the mass is usually small and therefore difficult to measure, and on the other hand, the particle number, or the amount of the substance, is a quantity better suitable for characterizing the physical behavior of gases.

In this book, the term *particle* refers to both simple atoms and composite molecules. In many languages, the terms atoms (monatomic particles, e.g., noble gases) and molecules (polyatomic particles, e.g., nitrogen) are differentiated when considering gas particles. In English, however, the word molecule often refers to a small particle, without discriminating between atoms and molecules. A gas consists of many individual gas particles. Therefore, the number of individual particles is a quantity that describes the amount of a gas. This dimensionless quantity is referred to as the particle number N. The particle number is a comprehensible quantity and appears in calculations frequently. For common amounts of gas, the particle number is very large and therefore practically impossible to measure.

A practical method to describe the amount of a gas is to specify the amount of the substance ν . It is obtained by scaling the actual amount of a substance to a certain reference amount. In the International System of Units (SI), this reference is one mole (symbol: mol). Following Avogadro's constant N_A ,

$$N_{\rm A} = 6.022142 \times 10^{23} \,\text{mol}^{-1},\tag{3.7}$$

the amount of one mole of a substance corresponds to 6.022142×10^{23} particles. The amount of the substance ν of any amount of a gas can be calculated from the number N of its gas particles:

$$\nu = \frac{N}{N_{\rm A}}.\tag{3.8}$$

If the temperature T of a gas is known, the pV value at T may also be used since the product $p \cdot V$ is proportional to the amount of gas, according to Eqs. (3.18)– (3.20).

For a gas that is distributed evenly within a volume (filling the volume homogeneously), its density, that is, the ratio of its amount and volume, can be calculated:

Mass density (density)
$$\rho := \frac{m}{V}, \quad [\rho] = \text{kg m}^{-3}.$$
 (3.9)

Number density of molecules
$$n = \frac{N}{V}$$
, $[n] = m^{-3}$. (3.10)

For calculations, it is useful to introduce the mass m_P of an individual gas particle. The mass m of an amount of gas with N particles is obtained by multiplying the number of particles with the particle mass:

$$m = Nm_{\rm P}, \quad [m] = [m_{\rm P}] = \text{kg}.$$
 (3.11)

Particle	Relative particle mass $M_{\rm r}$	Particle mass m _P	Molar mass M
He (atom)	4.003	$04.003 \mathrm{u} = 6.647 \times 10^{-27} \mathrm{kg}$	$04.003 \times 10^{-3} \text{ kg/mol}$
H ₂ (molecule)	2.016	$02.016 \mathrm{u} = 3.348 \times 10^{-27} \mathrm{kg}$	$02.016\times10^{-3}~\mathrm{kg/mol}$
N ₂ (molecule)	28.013	$28.013 \mathrm{u} = 4.652 \times 10^{-26} \mathrm{kg}$	$28.013\times10^{-3}~kg/mol$

Table 3.3 Selected relative particle masses, particle masses, and molar masses.

Additionally, the *molar mass M* of a substance is introduced by

$$M := \frac{m}{\nu} = N_{\rm A} m_{\rm P}, \quad [M] = \text{kg mol}^{-1}.$$
 (3.12)

The mass of a gas particle is very small. Therefore, mass is often not given in the SI unit kg but in the atomic mass unit u. It is defined as 1/12 of the mass of carbon-12 isotope and has the value

$$u = 1.660538 \times 10^{-27} \text{ kg.}$$
 (3.13)

Following the definitions, a simple relation connects Avogadro's constant $N_{\rm A}$ and the atomic mass unit *u*:

$$N_{\rm A}u \equiv \frac{\rm kg}{10^3 \,\rm mol}.\tag{3.14}$$

In addition, the relative particle mass M_r is also used. It is obtained by scaling the mass of a particle to the atomic mass unit (see also Table 3.3):

$$M_{\rm r} = \frac{m_{\rm P}}{u}, \quad [M_{\rm r}] = 1.$$
 (3.15)

 $A_{\rm r}$ (relative atomic mass) is also used as a symbol for the particle mass of an atom.

3.1.3

Equation of State of an Ideal Gas

In the seventeenth century, Boyle and Mariotte conducted experiments in England and France, respectively, to investigate the relationship between the pressure and volume of fixed amounts of gases. As the experiments showed, the volume V decreases when the pressure p rises. The two found a quantitative relation, indicating that, for constant temperature, the product of pressure and volume is constant (Boyle-Mariotte law):

$$pV = \text{constant}$$
, fixed amount of gas at $T = \text{constant}$. (3.16)

In 1704, Amontons discovered that a change in gas temperature leads to a change in pressure. Toward the end of the eighteenth century, experiments conducted by Charles and Gay-Lussac revealed that, for a fixed amount of gas, the

product pV increases linearly with a rise in temperature. Conversely, the product pV decreases linearly when the temperature drops and, for sufficiently low temperatures, approaches zero. This indicated the existence of a lowest possible temperature referred to as absolute zero.

For characterizing gas properties, it is appropriate to use the thermodynamic temperature scale, based on absolute zero. Kelvin introduced it around 1900 (Section 3.1.1).

This leads to the equation of state of an ideal gas:

$$\frac{pV}{T}$$
 = constant, for a fixed amount of gas.

The constant is proportional to the amount of gas. The three types of the equation of state of an ideal gas are obtained by expressing the amount of gas with mass m, particle number N, or the amount of the substance ν :

$$pV = mR_{\rm s}T,\tag{3.18}$$

$$pV = NkT$$
 or $p = nkT$, (3.19)

$$pV = \nu RT. \tag{3.20}$$

The above equations include the following fundamental constants:

Boltzmann's constant
$$k = 1.380650 \times 10^{-23} \text{ J K}^{-1}$$
. (3.21)

Molar gas constant
$$R = 8.314472 \,\mathrm{J} \,\mathrm{mol}^{-1} \,\mathrm{K}^{-1}$$

= $8.314472 \,\mathrm{Pa} \,\mathrm{m}^{3} \,\mathrm{mol}^{-1} \,\mathrm{K}^{-1}$
= $83.14472 \,\mathrm{mbar} \,\mathrm{l} \,\mathrm{mol}^{-1} \,\mathrm{K}^{-1}$. (3.22)

The constant R_s , however, depends on the gas species:

Specific gas constant
$$R_{\rm s} = \frac{k}{m_{\rm P}} = \frac{R}{M}$$
, $[R_{\rm s}] = {\rm J \ kg^{-1} \ K^{-1}}$. (3.23)

The equation of state of an ideal gas allows calculating the volume of the gas per amount of substance under standard conditions ($p_n = 101325 \, \text{Pa}, \ T_n = 273.15 \, \text{K}$). This volume is referred to as the *molar volume under standard conditions* $V_{\text{molar.n}}$:

$$V_{\text{molar,n}} = \frac{RT_{\text{n}}}{p_{\text{n}}} = 22.413996 \times 10^{-3} \text{ m}^3 \text{ mol}^{-1}.$$
 (3.24)

Therefore, under standard conditions, one mole of an ideal gas requires a volume of approximately 22.4 ℓ .

Additionally, the number density of particles n_n under standard conditions, known as the Loschmidt constant, can be calculated:

$$n_{\rm n} = \frac{p_{\rm n}}{kT_{\rm n}} = 2.686777 \times 10^{25} \,\mathrm{m}^{-3}.$$
 (3.25)

The equation of state of an ideal gas can be plotted. In a p-V diagram, the curves for constant temperatures appear as an array of hyperbolic curves, called isotherms (Figure 3.3).

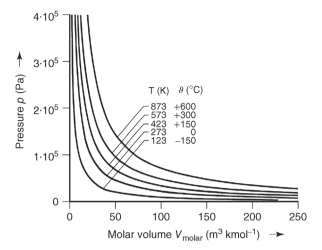


Figure 3.3 Isotherms of an ideal gas.

The behavior of real gases differs more or less from ideal-gas behavior, depending on the conditions of state. Section 3.4.1 focuses on this in more detail. For example, the value of the product pV for air under standard conditions is approximately 0.02% smaller than for an ideal gas. The deviation is higher under higher pressure and at lower temperature.

3.1.4 Mixtures of Different Gas Species

To this point, we assumed that the investigated gas consists solely of particles that are all of the same mass. In practice, however, one often deals with mixtures of gases. According to *Dalton*, the total pressure p_{tot} caused by the gas mixture is equal to the sum of the partial pressures p_i of the individual gases (each marked with the subscript *i*); thus (Dalton's law of partial pressures),

$$p_{\text{tot}} = \sum_{i} p_{i} = \sum_{i} \frac{N_{i}kT}{V} = \frac{NkT}{V} \sum_{i} \frac{N_{i}}{N}.$$
 (3.26)

This equation introduces the number of gas particles N_i of gas species i as well as the total number $N = \sum N_i$ of all gas particles. The ratio N_i/N , the number of particles of gas species i relative to the total number, corresponds to the relative volume fraction of gas *i*, and also to the ratio p_i/p_{tot} .

For many calculations, it is practical to treat the gas as if it was made up of hypothetical gas particles with a mean molar mass \overline{M} . This quantity is obtained from the weighted average

$$\overline{M} = \frac{\sum_{i} \left(M_{i} \frac{N_{i}}{N} \right)}{\sum_{i} \frac{N_{i}}{N}}.$$
(3.27)

Example 3.2

Dry air is a mixture with the main components (Table A.6):

Gas species	Molar mass (10 ⁻³ kg mol ⁻¹)	Relative volume function
Nitrogen	28.013	0.7809
Oxygen	31.999	0.2095
Argon	39.948	0.0093
Carbon dioxide	44.010	0.0003

The composition of air can be thought of as 780.9 ℓ nitrogen, 209.5 ℓ oxygen, 9.3 ℓ argon, 0.3 ℓ carbon dioxide, brought together and mixed to yield 1000 ℓ of dry air. Using Eq. (3.27), the mean molar mass of the gas mixture

$$\overline{M} = \frac{\begin{pmatrix} 28.013 \cdot 0.7809 + 31.999 \cdot 0.2095 \\ +39.948 \cdot 0.0093 + 44.010 \cdot 0.0003 \end{pmatrix} \times 10^{-3} \text{ kg mol}^{-1}}{0.7809 + 0.2095 + 0.0093 + 0.0003}$$

$$= \frac{28.964 \times 10^{-3} \text{ kg mol}^{-1}}{1.0000} = 28.964 \times 10^{-3} \text{ kg mol}^{-1}.$$
(3.28)

Solving for the density ρ of an ideal gas yields

$$\rho = \frac{m}{V} = \frac{pM}{RT} \,. \tag{3.29}$$

Example 3.3

The density of dry air under standard conditions (101 325 Pa, 273.15 K) is

$$\rho = \frac{101\ 325\ \text{Pa} \cdot 28.964 \times 10^{-3}\ \text{kg mol}^{-1}}{8.314472\ \text{J mol}^{-1}\ \text{K}^{-1} \cdot 273.15\ \text{K}} = 1.2922\ \text{kg m}^{-3}. \tag{3.30}$$

3.2 **Kinetic Theory of Gases**

3.2.1

Model Conceptions

A gas completely fills an available volume and shows a number of macroscopic properties: it has a temperature and exerts a temperature-dependent pressure to the walls. An equation of state, Eqs. (3.18)–(3.20), connects the state quantities pressure, volume, and temperature. Additionally, a gas is capable of conducting frictional force between surfaces in motion (viscosity), transferring thermal energy between surfaces with unequal temperatures (thermal conductivity), and can influence spreading of molecular particles (diffusion).

These different properties of a gas can be explained easily by considering the microscopic behavior of individual gas particles (atoms, molecules), by means of the kinetic theory of gases. This theory is based on the conception that a gas consists of a very large number of tiny particles that move thermally (kinetics). The moving particles hit the walls of the container and one another. All collisions are assumed elastic, that is, the total energy is conserved. During a collision, however, velocities of the colliding particles change with respect to value and direction, following the mechanical laws of collisions. The kinetic theory of gases derives the macroscopic properties of a gas from the microscopic motion of individual particles.

Krönig developed the kinetic theory of gases as a model in Berlin during the mid-nineteenth century. Later, it was verified in experiments and has proven very successful. Using the model, the pressure on a wall can be calculated from the molecular impacts of many individual particles. It therefore permits developing the equation of state of a gas. Furthermore, the transport properties viscosity, thermal conductivity, and diffusion can be derived easily. This is briefly discussed in the following sections.

In its simplest form, the kinetic theory of gases assumes that gas particles are small, hard spheres with a fixed diameter, and which remain practically unaltered during a collision, such as billiard balls. This conception often already yields good understanding of reality and is used in this chapter. When further developing the model, soft spheres can be assumed that deform like rubber balls during a collision and additionally attract one another mutually when they come close.

3.2.2

Wall Pressure due to Impacting Particles

The hard-sphere model of gas particles is used to calculate wall pressure. In the calculation, N gas particles, each having the mass m_P , being in a volume V, that is, a cube with an edge length d_i are considered (Figure 3.4). To simplify matters, the particle size is assumed to be negligibly small. The particles are evenly (homogeneously) distributed in the volume and move about randomly (kinetic motion). The directions of motion are distributed isotropically in three dimensions.

Of all gas particles, one third, 1/3N, each move along or reversely to the x-, y-, or z-axis. The movement is described by the terms velocity c (vector c or vector component c_x) or speed c (absolute value). We will now consider an individual particle moving back and forth horizontally between the confining walls of the cube in the x-direction. The velocity c_x of the particle is constant. Before impacting a wall, the momentum of the particle is $m_P c_x$; after colliding, it leaves

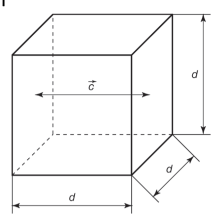


Figure 3.4 Particle motion in a cube and resulting wall pressure.

the wall with a momentum $-m_Pc_x$. The value of the momentum therefore changes by $2m_Pc_x$ during the collision.

If the particle moves at constant velocity c_x , it hits the walls periodically. The collision frequency, that is, the number of collisions per time interval, is the ratio of velocity c_x and traveling distance (2d for back and forth distance), that is, $c_x/(2d)$.

According to the laws of mechanics, the force that is exerted to the wall is the product of the change in momentum per collision and the collision frequency:

Wall force caused by particle =
$$2m_P c_x \frac{c_x}{2d} = \frac{m_P c_x^2}{d}$$
. (3.31)

The pressure is calculated from the force by dividing the wall force by the wall's surface area (d^2) :

Wall pressure caused by particle =
$$\frac{m_P c_x^2}{d} \cdot \frac{1}{d^2} = \frac{m_P c_x^2}{d^3}$$
. (3.32)

 d^3 can be written as volume V. The pressure applied to the wall by the total gas is obtained by multiplying the wall pressure caused by a single particle with the number of particles hitting the wall (1/3N):

$$p = \frac{m_{\rm P}c_x^2}{d^3} \cdot \frac{N}{3} = \frac{N}{3} \cdot \frac{m_{\rm P}c_x^2}{V}.$$
 (3.33)

Rearranging the equation finally yields

$$pV = N \frac{m_{\rm P} c_{\rm x}^2}{3}. (3.34)$$

When comparing Eq. (3.34), obtained from the kinetic theory of gases, with the experimentally found equation of state (3.19), the two equations correspond, if the velocity c_x complies with the following relation:

$$c_x = \sqrt{\frac{3kT}{m_{\rm P}}}. (3.35)$$

In our simple model, the speed c of a particle is just the absolute value of its velocity c_x . Then, by rewriting using Eq. (3.23):

$$c = \sqrt{\frac{3RT}{M}} = \sqrt{3R_sT}. ag{3.36}$$

Example 3.4

The speed of a hypothetical air particle (Table A.6) at 20 °C amounts to

$$c(air) = \sqrt{\frac{3 \cdot 8.314472 \,\text{J mol}^{-1} \,\text{K}^{-1} \cdot 293.15 \,\text{K}}{0.028964 \,\text{kg mol}^{-1}}} = 502 \,\text{m s}^{-1}. \tag{3.37}$$

Thus, the speed of a hypothetical air particle is very high, higher than the speed of sound (343 m s⁻¹ at 20 °C). This result is understandable when we consider that the sound is transmitted as pressure variations by the gas particles.

Additionally, Eq. (3.36) yields that the speed of gas particles rises when the temperature increases and that heavy gas particles are slower than lighter particles.

3.2.3

Maxwell-Boltzmann Velocity Distribution

In the previous section, we assumed that all gas particles travel with the same velocity c_x . Collisions between particles were neglected.

In fact, however, particles do collide mutually due to their finite particle size. Depending on the type of collision between two particles (head-on or rather grazing), velocity values and directions change. Similar considerations apply to collisions with a wall. A real vessel wall is not a static, flat surface but shows microscopic roughness and vibrates thermally. Thus, the collision with a wall is not a simple reflection.

Overall, a large number of collisions occur within a gas, and the sheer number makes it impossible to consider them individually. This may initially create the impression that quantitative relations of gas properties cannot be derived from microscopic behavior. However, this is not the case. On closer inspection, we find that just the large number of particles allows deriving accurate mean values of motion quantities.

To begin with particle velocity, general symmetry considerations suggest that all directions of motion (arbitrary orientation in three dimensions) appear equally often. Considering the component velocity of all gas particles in any given direction, for example, the x-direction, particles possess different values that can be positive (along the considered direction) or negative (opposite to the

considered direction). This behavior can be expressed mathematically using a distribution function, for example, the function F_1 for the normalized x-component velocity c_x . Normalization is performed by dividing by the most probable velocity $c_{\rm mp}$ (see Eq. (3.42) below):

$$F_1\left(\frac{c_x}{c_{\rm mp}}\right) := \frac{1}{N} \cdot \frac{dN}{d(c_x/c_{\rm mp})}.$$
(3.38)

Here, dN is the fraction of particles from the total number N with a velocity component in the x-direction in the interval from c_x/c_{mp} to $(c_x + dc_x)/c_{mp}$ when the components in the y- and z-directions are of arbitrary values. As all particles are considered, the normalizing condition reads

$$\int_{-\infty}^{\infty} F_1 \left(\frac{c_x}{c_{\rm mp}} \right) d\left(\frac{c_x}{c_{\rm mp}} \right) = 1.$$
 (3.39)

Now what does the velocity distribution F_1 look like? Around 1860, Maxwell presumed a Gaussian bell-like distribution curve. Boltzmann determined the absolute value of the velocity about one decade later. The velocity distribution is therefore referred to as the Maxwell-Boltzmann velocity distribution (Figure 3.5). Later, it was derived precisely from statistical mechanics [2]. Modern computer simulations that calculate the motion and collisions of large numbers of particles, as well as many experiments, have verified this distribution. It reads

$$F_1\left(\frac{c_x}{c_{\rm mp}}\right) = \frac{1}{\sqrt{\pi}} \exp\left(-\frac{c_x^2}{c_{\rm mp}^2}\right). \tag{3.40}$$

This one-dimensional velocity distribution is symmetric to the axis of the ordinate because positive and negative velocity values appear equally often.

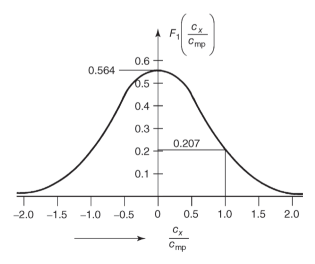


Figure 3.5 Distribution function of the normalized one-dimensional particle velocity c_x according to Maxwell and Boltzmann.

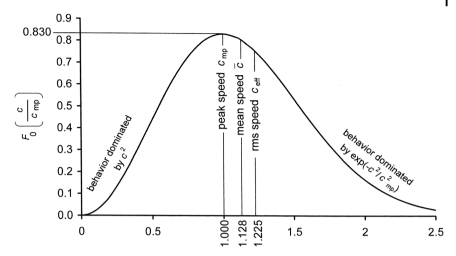


Figure 3.6 Normalized distribution function of the absolute values of particle velocity c according to Maxwell and Boltzmann.

The distribution F_0 of speed values c, that is, the absolute value of the velocity vector, can be obtained by integration over the F_1 distributions of the three directions with boundary condition of fixed c. Calculation leads to the function given below, represented in Figure 3.6:

$$F_0\left(\frac{c}{c_{\rm mp}}\right) = \frac{4}{\sqrt{\pi}} \cdot \frac{c^2}{c_{\rm mp}^2} exp\left(-\frac{c^2}{c_{\rm mp}^2}\right). \tag{3.41}$$

For normalization, the most probable speed $c_{\rm mp}$, that is, the peak speed, has already been used, indicating the speed value at the peak of the distribution function F_0 . In order to describe macroscopic phenomena, it can be advantageous to use other speed values. The mean speed \bar{c} is obtained by calculating the weighted average of the gas particles' speed values. The effective speed $c_{\rm eff}$ is determined by calculating the square root of the weighted average of the gas particles' squared speed values. Calculation yields the following values:

Most probable speed c_{mp} = argument value where F_0 obtains its maximum

$$= \sqrt{\frac{2kT}{m_{\rm P}}} = \sqrt{\frac{2RT}{M}} = \sqrt{2R_{\rm S}T} = \sqrt{\frac{2p}{\rho}}.$$
 (3.42)

Arithmetic mean thermal speed $\overline{c} = \int_{0}^{\infty} cF_0 \left(\frac{c}{c_{max}}\right) dc$

$$= \sqrt{\frac{8kT}{\pi m_{\rm P}}} = \sqrt{\frac{8RT}{\pi M}} = \sqrt{\frac{8}{\pi}R_{\rm s}T} = \sqrt{\frac{8p}{\pi \rho}}.$$
 (3.43)

Effective speed = root-mean-square speed
$$c_{\text{rms}} = \sqrt{\int_0^\infty c^2 F_0 \left(\frac{c}{c_{\text{mp}}}\right) dc}$$

$$= \sqrt{\frac{3kT}{m_{\rm P}}} = \sqrt{\frac{3RT}{M}} = \sqrt{3R_{\rm s}T} = \sqrt{\frac{3p}{\rho}}.$$
 (3.44)

Example 3.5

Velocity calculation for hypothetical air particles (Table A.6) at 20 °C. The molar mass $M = 0.028964 \text{ kg mol}^{-1}$ (Eq. (3.28)):

$$c_{\text{mp}} = 410 \text{ ms}^{-1},$$
 (3.45)
 $\overline{c} = 463 \text{ ms}^{-1},$ (3.45)
 $c_{\text{rms}} = 502 \text{ ms}^{-1}.$ (3.47)

$$\overline{c} = 463 \,\mathrm{ms}^{-1},$$
 (3.45)

$$c_{\rm rms} = 502 \,{\rm ms}^{-1}$$
. (3.47)

3.2.4

Collision Rate and Effusion

Many macroscopic properties of a gas, for example, pressure, are determined by the impingement rate at which gas particles collide with a surface. Here, the term collision rate j_N is introduced, also referred to as the rate of incidence. It is defined as the number of collisions with a surface per unit area and time. The kinetic theory of gases allows calculating the collision rate, when assuming a Maxwell-Boltzmann velocity distribution:

Collision rate
$$j_N := \frac{\text{Number of collisions with wall}}{\text{Area of wall} \times \text{time}} = \frac{N}{At} = \frac{n\overline{c}}{4}$$

$$= \frac{p\overline{c}}{4kT}.$$
(3.48)

An application example for using the collision rate is a gas flow through an opening in a wall, referred to as effusion (gas escape) (Figure 3.7). A thin wall that has a small hole with the area A separates one vessel from the other.

If the pressure in vessel 1, to the left of the wall, is p_1 , the temperature is T_1 , and the pressure in vessel 2 is negligible, the particle flow (= number of particles per unit time) leaving the vessel 1 is

Effusion-particle flow
$$q_N:=\frac{\text{Number of emanating particles}}{\text{Time}}$$

$$=j_NA=\frac{p\overline{c}}{4kT}A=\frac{n\overline{c}}{4}A. \tag{3.49}$$

A precondition for this equation is that the pressure is small enough so that disturbing collisions between gas particles do not occur in the area of the

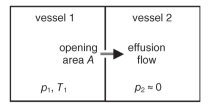


Figure 3.7 Effusion from a vessel.

opening (molecular flow). For the volumetric flow of the escaping gas, the above equation yields

Effusion-volumetric flow
$$q_V := \frac{\text{Emanating gas volume}}{\text{Time}}$$

$$= \frac{\Delta V}{\Delta t} = \frac{\Delta N/n}{\Delta t} = \frac{\overline{c}}{4}A. \tag{3.50}$$

Example 3.6

The vessel to the left (Figure 3.7) contains air at pressure p = 1 mPa and a temperature of 20 °C; the opening has an area $A = 1 \text{ cm}^2$. With $\bar{c} = 463 \text{ m s}^{-1}$ and T = 293 K, we calculate

$$q_N = \frac{10^{-3} \text{ Pa} \times 463 \text{ m s}^{-1}}{4 \times 1.381 \times 10^{-23} \text{ J K}^{-1} \cdot 293 \text{ K}} \times 1 \times 10^{-4} \text{ m}^2$$
$$= 2.86 \times 10^{15} \text{ s}^{-1}, \tag{3.51}$$

$$q_V = \frac{463 \,\mathrm{m \, s^{-1}}}{4} \cdot 1 \times 10^{-4} \,\mathrm{m}^2 = 0.0116 \,\mathrm{m}^2 \,\mathrm{s}^{-1} = 11.6 \,\ell \,\mathrm{s}^{-1}. \tag{3.52}$$

3.2.5

Size of Gas Particles and Free Path

So far, the size of the gas particles remained unconsidered. Their sizes play a crucial role for transport phenomena. Different methods are available to determine the size, as shall be discussed next.

When a gas is cooled far enough, it initially liquefies and finally freezes. A certain amount of gas then forms a liquid or solid with a certain volume. The assumption is plausible that the individual atoms and molecules in a solid are arranged as small, closely packed spheres. Using this model, the volume filled by an individual particle can be calculated by dividing the particle mass by the density of the solid. Based on the used volume, the diameter of the sphere is obtained after specifying the structure in which the spheres are arranged in the solid.

Example 3.7

At 4 K, nitrogen is a solid with a density of 1035 kg m⁻³. The volume required by a nitrogen molecule amounts to $28u/1035 \text{ kg m}^{-3} = 4.5 \times 10^{-29} \text{ m}^3$, corresponding to a cube with an edge length of 3.6×10^{-10} m = 0.36 nm.

Modern experimental methods, such as structure analysis by X-ray diffraction or surface scanning with an atomic force microscope, allow direct measurement of the distance between two particles, and therefore, of their size. Results reveal that the diameter of simple gas particles (i.e., individual gas atoms) amounts to approximately 3×10^{-10} m = 0.3 nm, quite independent of the gas species. As gas particles, in fact, are not hard spheres, their size is not well defined but depends on the type of phenomenon observed, as will be discussed in Section 3.4.2.

During their kinetic motion, gas particles come into contact when the distance between their centers drops below their diameter. The collision changes the particles' directions and speeds. Due to multiple particle collisions, the path of an individual particle follows a zigzag route (Figure 3.8).

The path lengths that a particle travels between two successive collisions vary due to the statistical motion of the particles. An average value of this path length can be defined, referred to as the *mean free path l*.

We will now calculate this mean free path, while assuming that the gas particles are small hard spheres with diameter d. Furthermore, we shall presume that no force is transferred between particles except during elastic collisions. First, we will consider a simplified case in which a gas particle travels through a virtual gas volume V (cross-sectional area A, thickness s), containing static particles of the same species (Figure 3.9).

A moving gas particle collides with a stationary gas particle inside the volume if the distance between their centers drops below the particle diameter d. Thus, the effective collision area (perpendicular to the particle's trajectory) for this



Figure 3.8 Zigzag path of a gas particle.

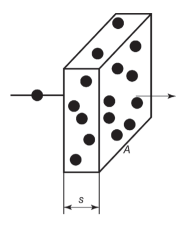


Figure 3.9 A gas particle traveling through a volume of gas.

collision amounts to πd^2 . The total effective collision area for all possible collisions is obtained by multiplying the individual areas with the number N of atoms in the volume:

Total collision area =
$$N\pi d^2 = nV\pi d^2 = nAs \pi d^2$$
. (3.53)

The larger the thickness s of the layer, the more probable a collision. In the case of a statistical (irregular) arrangement of the stationary gas particles in the volume, the thickness of the layer amounts to the mean free path, thus $s = \bar{l}$, if the total effective collision area (Eq. (3.53)) is equal to the geometrical area A. This leads to the condition

$$\bar{l} = \frac{1}{\pi d^2 n}$$
 (gas particles in the volume assumed stationary). (3.54)

Due to the statistical arrangement of the gas particles in the volume, an incoming particle passes the distance $s = \bar{l}$ without collisions with a probability of 37%, and passes the distance $s = 4\bar{l}$ with a probability of just under 2%.

In reality, all gas particles travel with a statistic velocity distribution according to Maxwell-Boltzmann. Therefore, more collisions occur and the mean free path drops. Maxwell investigated this problem in 1860 and added a factor of $1/\sqrt{2}$ to Eq. (3.54):

$$\bar{l} = \frac{1}{\sqrt{2\pi d^2 n}}$$
 (all gas particles in motion). (3.55)

This formula is valid when the mean free path is defined as the total distance traveled by molecules in a time period divided by the total number of their collisions in this period. Other definitions of the mean free path may be used, for example, the mean distance moved by a molecule between a given instant and its next collision. These definitions lead to slightly different numerical values, see [2]. Nearly all literature uses Eq. (3.55) and so will we.

By replacing the number density of particles n with the term p/(kT) in Eq. (3.55), and by moving p to the left-hand side, we find

$$\bar{l}p = \frac{kT}{\sqrt{2\pi}d^2} \quad . \tag{3.56}$$

Thus, for a particular gas (with particles of diameter d), the product of mean free path and pressure depends only on the temperature.

Example 3.8

At a temperature of 20 °C, a hypothetical air particle (Tables A.7 and A.9) has a diameter of d = 0.37 nm. The product $\bar{l}p$ amounts to

$$\bar{l}p = \frac{1.38 \times 10^{-23} \text{ J K}^{-1} \cdot 293 \text{ K}}{\sqrt{2} \pi (0.37 \times 10^{-9} \text{ m})^2} = 0.0066 \text{ m Pa}.$$
 (3.57)

Under atmospheric pressure (10⁵ Pa), the mean free path is only 6.6×10^{-8} m = 66 nm. However, under high vacuum of 10^{-4} Pa, it reaches 66 m and therefore exceeds the size of common vacuum systems.

Mean free path is an important term, for both a descriptive characterization of gas behavior and a quantitative calculation of macroscopic gas properties. The larger a gas particle, the more often collisions occur, the lower the free path, and the poorer the transport properties for frictional force (viscosity) and heat energy (thermal conductivity).

When comparing the transport properties calculated using the kinetic theory of gases with experimental data at different temperatures, the gas particles seem to grow when the temperature drops. Sutherland gave an empirical description of this behavior in 1894. He formulated the following relation between the diameter d of a gas particle and the temperature T:

$$d(T) = d_{\infty} \sqrt{1 + T_{\rm D}/T}.$$
 (3.58)

Here, d_{∞} is the particle diameter at very high temperature and $T_{\rm D}$ is Sutherland's constant, with the dimension of a temperature. At temperature $T_{\rm D}$, the effective particle diameter is twice as high as at very high temperature.

The value of Sutherland's constant for hypothetical air amounts to approximately 102 K (Table A.9). Therefore, the diameter of an air particle at 20 °C is larger by a factor of $\sqrt{1 + 102 \text{ K}/293 \text{ K}} = 1.16$ than the diameter at very high temperature.

As is known today, Sutherland's approach describes the fact that real gas particles attract one another due to electrostatic polarization when they come close. When the temperature drops, the particle velocity decreases. The mutual attraction then increasingly affects the particle paths and the particles seem to grow.

The ratio of mean free path to mean particle velocity is referred to as mean time τ between individual collisions:

$$\tau = \frac{\overline{I}}{\overline{c}}.\tag{3.59}$$

Example 3.10

Hypothetical air at standard conditions shows $\bar{l} = 6.6 \times 10^{-8} \, \text{m}$ and $\bar{c} =$ 463 m s⁻¹ so that the mean time between two collisions is only 1.4×10^{-10} s.

The volume collision rate χ , that is, the temporal average of the number of collisions between two gas particles in a volume per unit time and volume, is calculated from

$$\chi = \frac{n}{2\tau} = \frac{n\overline{c}}{2\overline{l}} = \frac{\pi}{\sqrt{2}}\overline{c}d^2n^2 = \frac{\pi}{\sqrt{2}}\overline{c}d^2\left(\frac{p}{kT}\right)^2. \tag{3.60}$$

The factor 1/2 in this equation takes into account that two particles are involved in each particle-particle collision.

Example 3.11

The volume collision rate for hypothetical air under standard conditions is

$$\chi = \frac{\pi}{\sqrt{2}} 463 \text{ m s}^{-1} (3.7 \times 10^{-10} \text{ m})^2 \left(\frac{10^5 \text{ Pa}}{1.38 \times 10^{-23} \text{ J K}^{-1} \cdot 293 \text{ K}} \right)^2$$

$$= 8.6 \times 10^{34} \text{ m}^{-3} \text{ s}^{-1}.$$
(3.61)

The concepts of time between two collisions and volume collision rate are intuitively understandable. However, they are not required for a precise quantitative calculation of observable quantities. Thus, whether the given definitions in fact represent statistically correct average values is irrelevant in this context. Figure 3.10 compiles various gas properties as a function of pressure.

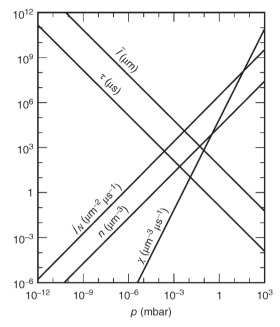


Figure 3.10 Gas-kinetic diagram for air at 20 °C: pressure dependence of the mean free path \bar{l} , average time τ between two collisions, collision rate j_N , particle number density n, and volume collision rate χ .

3.3 Transport Properties of Gases

3.3.1

Pressure Dependence

Transport properties of a gas include the following macroscopic properties:

- Transmission of frictional force through the gas shear stress (viscosity).
- Transfer of thermal energy through the gas heat flux (thermal conductivity).
- Influence on spreading of particular individual particles through the gas (diffusion).

In order to understand the concepts of transport of frictional forces and thermal energy by the gas, the geometry of two plates at a distance x is considered. The transport properties depend crucially on the ratio of the free path \bar{l} to the distance between the two plates x (Figure 3.10).

If the ratio \overline{l}/x is far above 1, the system is in the so-called *molecular* regime. Here, gas particles nearly travel freely from one plate to the other. When the pressure, that is, the particle number density, increases, more and more particles are

available for transport. Thus, the transport ability increases linearly with the pressure. In the molecular regime, the transport properties are proportional to pressure.

In the case of \overline{l}/x being far below 1, the condition is referred to as viscous regime. An individual gas particle now only travels a small fraction of the distance between one plate and the other before it encounters a collision. By the collision, only part of the transported quantity (momentum or energy) is transferred in the forward direction while the remaining portion moves backward. Thus, collisions impede transport processes from one plate to the other. If the pressure rises, the number of particles available for transport increases but the mean free path decreases. This means that more and more collisions hinder the transport of momentum and energy. The result is that, in the viscous regime, transport properties of a gas (viscosity and thermal conductivity) are independent of pressure.

3.3.2

Transport of Frictional Forces in Gases and Viscosity

In order to understand internal friction, two plane parallel plates are considered, with area A and distance x. The bottom plate is stationary while the above plate moves at velocity ν (Figure 3.11).

Initially, we consider the *low-pressure case* (molecular regime) in which the gas particles travel back and forth between the plates (nearly) without any mutual collisions. The number of gas particles hitting the upper plate per unit time is calculated by multiplying the collision rate j_N (Eq. (3.48)) with the area A of the plate:

$$\frac{\mathrm{d}N}{\mathrm{d}t} = \frac{1}{4} \cdot \frac{p\overline{c}}{kT}A. \tag{3.62}$$

We will first presume that the particles that hit the moving plate, on average, do not carry any velocity component in the direction in which the plate is moving. During reflection at the plate, the particles, on average, take on a part of the plate's velocity. This fraction is described by the so-called tangential-momentum accommodation coefficient:

Tangential-momentum accommodation coefficient
$$\sigma_t$$

$$= \frac{\text{mean tangential velocity of reflected particles}}{\text{velocity of plate}}$$
(3.63)

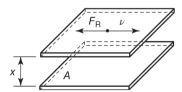


Figure 3.11 Frictional force between two moving plates.

Because the moving plate gives the gas particles a tangential velocity, a decelerating force occurs at the plate, that is, the frictional force. This force is calculated from the number of particles hitting per unit time (Eq. (3.48)) area and the mean change in momentum of a particle:

$$F_{\rm R} = \frac{1}{4} \cdot \frac{p\overline{c}}{kT} A \sigma_{\rm t} m_{\rm P} \nu$$
 in the molecular regime. (3.64)

Using the definition of the mean speed, Eq. (3.43), the equation can be rewritten

$$F_{\rm R} = \frac{2}{\pi} p A \sigma_{\rm t} \frac{\nu}{\bar{c}}$$
 in the molecular regime. (3.64b)

Actually, it is necessary to take into account the momentum accommodation of the gas particles at both plates. After leaving the upper plate, the gas particles, on average, carry a tangential velocity component. This is reduced (but not to zero) during the reflection at the bottom plate, so that in effect the particles, on average, already travel at a tangential velocity when hitting the above plate. Let σ_{t1} and σ_{t2} denote the momentum-accommodation coefficients at plates 1 and 2, respectively. For the effective total momentum-accommodation coefficient needed in Eq. (3.64), calculation yields

$$\sigma_{t} = \frac{\sigma_{t1}\sigma_{t2}}{\sigma_{t1} + \sigma_{t2} - \sigma_{t1}\sigma_{t2}}.$$
(3.65)

In the case of equal momentum-accommodation coefficients at both plates $(\sigma_{t2} = \sigma_{t1})$ we find

$$\sigma_{\rm t} = \frac{\sigma_{\rm tl}}{2 - \sigma_{\rm tl}}.\tag{3.66}$$

We will now consider the high-pressure case (viscous regime), that is, the mean free path is small compared to the distance between the plates $(\bar{l} \ll x)$. In this case, the accommodation behavior of the gas at the plates only has an influence in the immediate vicinity of the plates or, more precisely, in a boundary layer with a thickness of several free-path lengths. For calculation of the frictional force, this boundary layer is negligible. Thus, for approximation, full accommodation may be assumed in the high-pressure case.

Calculating the frictional force via the kinetic theory of gases requires taking into account and averaging the momentum transfer from one gas particle to another during collisions, a tedious and complicated task. In order to understand the process, a descriptive, simplified approach is helpful. For this, we divide the volume (thickness x) between the plates into numerous thin sheets separated by imaginary parallel plates at distances of twice the free path 2l (layer model) (Figure 3.12). The factor 2 appears to be chosen arbitrarily at the moment. Just this factor will yield the exact expression of the viscosity, which will be calculated later.

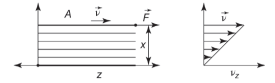


Figure 3.12 Velocity profile of layers between two moving plates.

When the layer thickness is $2\bar{l}$, there are $x/(2\bar{l})$ layers between the real outer plates. The relative velocity between the upper and lower side of a layer amounts to $2\nu \bar{l}/x$.

The frictional force between the two plates is the same as that between the upper and the lower side of a layer. The latter can be taken from Eq. (3.64), setting $\sigma_t = 1$ and replacing ν by $2\nu \bar{l}/x$. Thus, we get

$$F_{R} = \frac{2}{\pi} p A \frac{2\nu \bar{l}}{\bar{c}x}$$

$$= \frac{4}{\pi} A \frac{p\bar{l}}{\bar{c}} \cdot \frac{\nu}{x}$$
 in the viscous regime. (3.67)

Experimental investigations have revealed that, in laminar flow, the frictional force is proportional to the plate area A and to the velocity ν , and is inversely proportional to the distance x between the plates. Therefore, Newton's formulation is used for the frictional force:

$$F_{\rm R} = \eta A \frac{v}{x}$$
 and $F_{\rm R} = \eta A \frac{\mathrm{d}v}{\mathrm{d}x}$. (3.68)

This is the definition of the (dynamic) viscosity η , a property of the fluid between the plates. Comparing the frictional force, Eq. (3.67), with Newton's formulation, we obtain for the viscosity

$$\eta = \frac{4}{\pi} \cdot \frac{p\bar{l}}{\bar{c}}.\tag{3.69}$$

As $p\bar{l}$ is independent of pressure, so is η . Frequently, Eq. (3.69) is rewritten by taking the pressure from Eq. (3.43):

$$p = \frac{\pi}{8}\rho \overline{c}^2. \tag{3.70}$$

This produces the following result for viscosity:

$$\eta = \frac{1}{2}\rho\bar{c}\bar{l}.\tag{3.71}$$

This equation gives a prefactor of 1/2, which agrees well with experimental data. As early as 1860, Maxwell performed qualitative assessments and had already obtained an equation such as Eq. (3.71) for viscosity, but including a prefactor of 1/3. The numerically incorrect prefactor first derived by Maxwell still appears in a number of textbooks today.

With considerable mathematical effort, a correct calculation of the viscosity of a gas from the individual particle-particle collisions is possible. In 1915, Chapman used an analytical calculation in the hard-sphere model of gas particles that, as one would expect, again yielded an equation of the type (3.71), but included a prefactor of 0.491 in first approximation, and a prefactor of 0.499 in second approximation. Later, statistical calculations confirmed this result. For practical applications, the prefactor 0.499 may be replaced by 1/2, as written in Eq. (3.71). Remember that this result was derived here by proper choice of the layer thickness $2\bar{l}$ in the simple layer model.

The viscosity of a gas in the viscous regime can be measured precisely in experiments (e.g., by assessing the frictional force on moving plates or laminar flow through a pipe). Figure 3.13 surveys viscosity data for various gas species.

As the density ρ of a gas and the mean particle velocity \overline{c} can be calculated reliably, Eq. (3.71) provides the straightest method to obtain the mean free path from experimental viscosity values:

$$\bar{l} = \frac{2\eta}{\rho \bar{c}} \tag{3.72}$$

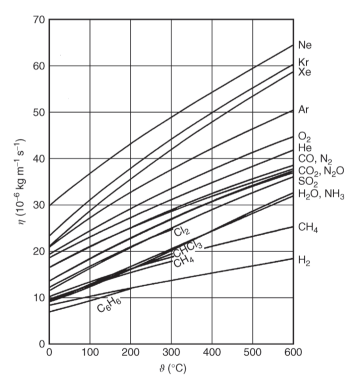


Figure 3.13 Viscosity of selected gas species in the viscous range versus temperature. (Data taken from Ref. [3].)

and the equivalent free path ℓ , which is defined as

$$\ell = \frac{2}{\sqrt{\pi}}\bar{l} = \frac{4}{\sqrt{\pi}} \cdot \frac{\eta}{\rho \bar{c}} = \frac{\eta c_{\rm mp}}{p}.$$
(3.73)

Furthermore, one calculates the product of mean free path and pressure,

$$\bar{l}p = \frac{\pi}{4}\bar{c}\eta,\tag{3.74}$$

and the particle diameter (Eq. (3.56) rearranged)

$$d = \frac{2}{\pi} \sqrt{\frac{1}{\sqrt{2}} \cdot \frac{kT}{\overline{c}\eta}}. (3.75)$$

Example 3.12

At 20 °C, the viscosity of air is 18.2×10^{-6} Pa s. Using this, calculation yields

$$\bar{l}p = \frac{\pi}{4}463 \text{ m s}^{-1} 18.2 \times 10^{-6} \text{ Pa s} = 6.6 \times 10^{-3} \text{ m Pa},$$
 (3.76)

$$d = \frac{2}{\pi} \sqrt{\frac{1}{\sqrt{2}}} \times \frac{1.38 \times 10^{-23} \text{ J K}^{-1} \cdot 293 \text{ K}}{463 \text{ m s}^{-1} \cdot 18.2 \times 10^{-6} \text{ Pa s}} = 3.7 \times 10^{-10} \text{ m}.$$
 (3.77)

In some applications, the frictional force over the whole pressure regime is required. A simple formula, which describes the limiting cases of the molecular regime and the viscous regime correctly, is obtained by combining the corresponding results given in Eqs. (3.64) and (3.67):

$$\frac{1}{F_{R}} = \frac{1}{F_{R}(\text{molecular})} + \frac{1}{F_{R}(\text{viscous})}$$
(3.78)

giving

$$F_{\rm R} = \frac{pA\nu}{\frac{\pi}{2} \cdot \frac{\bar{c}}{a} + \frac{x}{n} p}.$$
 (3.79)

The transitional region is treated in more detail in Section 5.4.1.

3.3.3

Transport of Heat in Gases and Thermal Conductivity

We will consider two plane plates with area A and distance x. The temperatures T_1 and T_2 of the plates differ (Figure 3.14).

If the space between the plates contains matter, heat transport occurs from the warmer to the colder plate. Transport of heat through a gas is very similar to friction behavior. From a macroscopic point of view, thermal energy is transferred by

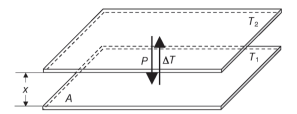


Figure 3.14 Heat transfer between two plates at different temperatures.

heat transport, while a force is transferred by friction. On the microscopic scale, gas particles absorb energy at the warmer plate and release it at the cold plate during heat transport. In force transport, they pick up momentum at the fast-moving plate and release it at the slow-moving plate. Calculating the thermal transport of a gas in the kinetic theory of gases is analogous to calculating the frictional force in the previous section, and therefore, will be presented in brief.

Clearly, the transport of heat is proportional to the amount of heat that an individual gas particle can absorb and carry. The thermal energy of a particle is stored in its forward motion (translatory energy). In molecules, internal motion (vibration \times vibration and rotation) adds to the thermal energy. This concept is described using the degree of freedom f of a particle.

Atoms (noble gases and metal vapors) have a degree of freedom f=3 because they can perform translatory motion in three dimensions. At room temperature, the degree of freedom for *diatomic molecules* (e.g., air) amounts to f=5 because three translatory motions (as in atoms) and, in addition, two directions of rotation may occur. Although the molecule may rotate around all three axes (x,y,z), the rotation of a diatomic molecule around the axis that connects the two nuclei hardly contains any energy. This is because the particle mass concentrates nearly completely in the extremely small atomic nucleus. Therefore, the angular momentum of rotation around the axis connecting the nuclei is very small.

Additional degrees of freedom arise from the vibrating motion of the individual atoms within a molecule. For air at room temperature, this motion is hardly excited, as the necessary quantum-mechanical energy barrier is higher than the thermal energy. In the case of polyatomic hydrocarbon molecules with weak C–H bonds, room temperature is sufficient to excite many vibrations, and thus, to create a correspondingly high degree of freedom. While the temperature rises, the thermal energy increases and more states of motion are excited. Thus, the degree of freedom and the heat capacity increase with temperature.

The *heat capacity* C_V of an individual gas particle located in a vessel with fixed volume (indicated by the subscript V) amounts to

$$C_V = \frac{f}{2}k. (3.80)$$

Thus, the *specific* heat capacity c_V (ratio of heat capacity and mass) and the *molar* heat capacity c_{mV} (ratio of heat capacity to the amount of substance)

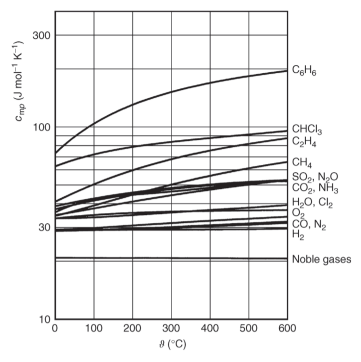


Figure 3.15 Molar heat capacity at constant pressure of selected gas species versus temperature. (Data taken from Ref. [3].)

of a gas are

$$c_V = \frac{f}{2} \cdot \frac{k}{m_P}, \quad c_{mV} = \frac{f}{2} k N_A = \frac{f}{2} R.$$
 (3.81)

Data collections often list the heat capacity at constant pressure (indicated by subscript p) (Figure 3.15). This has a higher value than the heat capacity at constant volume because, for constant pressure, the volume increases with temperature and, therefore, additional work is spent for the volume change. For ideal gases, the heat capacities at constant pressure and constant volume are easy to convert:

$$c_p = \frac{f+2}{2} \cdot \frac{k}{m_p} = c_V + \frac{k}{m_p}, \quad c_{mp} = c_{mV} + R = \left(1 + \frac{f}{2}\right)R.$$
 (3.82)

As an example, for noble gases f = 3; then $c_{mp} = (1 + 3/2)R = 20.8 \,\text{J mol}^{-1} \,\text{K}^{-1}$. Furthermore, the quantities can be converted using the isentropic exponent κ :

$$\frac{c_p}{c_V} = \frac{c_{\rm mp}}{c_{\rm mV}} = \kappa. \tag{3.83}$$

Carbon dioxide

For calculating the heat transport of a gas, we will first consider the *low-pressure* case (molecular regime). The number of gas particles that hit one of the plates per unit time is given by the collision rate j_N .

For an initial investigation, we will presume that the mean energy of the particles hitting Plate 2 at temperature T_2 averagely corresponds to the energy of the temperature T_1 of Plate 1 ($T_2 > T_1$). During reflection at Plate 2, the particles pick up part of the higher thermal energy. This fraction is referred to as the energy-accommodation coefficient a_E :

Energy-accommodation coefficient a_E

$$= \frac{\text{Real heat flux}}{\text{Theoretical heat flux at complete accommodation}}.$$
 (3.84)

Table 3.4 contains a selection of experimental data on the energy-accommodation coefficient.

Plate 2 is cooled by the fact that the gas particles, on average, pick up thermal energy during the reflection at the plate. The heat transferred here is calculated by multiplying the collision rate j_N , area A, energy-accommodation coefficient a_E , heat capacity C_V (Eq. (3.81)) of an individual particle, and the temperature difference. Incorporating the expressions for the variables yields

$$P = j_N A a_E \frac{f}{2} k (T_2 - T_1). \tag{3.85}$$

This equation is not quite correct because deriving it included an incorrect average across the statistical velocity distribution of the gas particles. Not only does a particle with higher velocity have more energy, but also it moves faster and therefore transfers this energy in less time, and thus, better. When

Gas species		Clean surface	Technical surface
Helium	He	0.03	0.38
Neon	Ne	0.07	0.74
Argon	Ar	0.55	0.86
Krypton	Kr		0.84
Xenon	Xe		0.86
Mercury	Hg	1.00	1.00
Hydrogen	H_2	0.15	0.29
Nitrogen	N_2		0.77
Oxygen	O_2	0.42	0.79
Carbon monoxide	CO		0.78

 CO_2

0.77

Table 3.4 Energy accommodation coefficients for selected gas species on platinum surfaces.

calculating this effect, we find that the factor f in Eq. (3.86) must be replaced with f + 1:

$$P = j_N A a_E \frac{f+1}{2} k (T_2 - T_1). \tag{3.86}$$

Data tables rarely list the degree of freedom f of a gas, but the isentropic exponent κ instead. Using the relation $f = 2/(\kappa - 1)$, we finally obtain the following equation for the heat transport:

$$P = \frac{1}{8} p \, \overline{c} A \, a_E \, \frac{\kappa + 1}{\kappa - 1} \frac{T_2 - T_1}{T} \quad \text{in the molecular regime.}$$
 (3.87)

To this point, the energy accommodation of the gas particles was only considered at the upper plate. In fact, it has to be taken into account at both plates. If we consider the energy-accommodation coefficients a_{E1} and a_{E2} for plates 1 and 2, calculation for the total effective energy-accommodation coefficient yields

$$a_E = \frac{a_{E1}a_{E2}}{a_{E1} + a_{E2} - a_{E1}a_{E2}}. (3.88)$$

If the energy-accommodation coefficients are equal at both plates ($a_{E2} = a_{E1}$), then

$$a_E = \frac{a_{E1}}{2 - a_{E1}}. (3.89)$$

We will now consider the high-pressure case (viscous regime), that is, the mean free path is small compared to the distance between the plates $(\bar{l} \ll x)$. In this case, the accommodation behavior of the gas is of interest only in the immediate vicinity of the plates or, more precisely, in a boundary layer with a thickness of several free-path lengths. For calculation of the amount of energy transferred, this layer is negligible. Thus, as an approximation, complete energy accommodation can be assumed.

Calculating heat transfer using the kinetic theory of gases requires taking into account and averaging the energy transfer from one gas particle to another during individual collisions, a tedious task. In order to understand the process qualitatively, we can think of the volume (thickness x) between the plates as separated by thin sheets, arranged at a distance twice the free path \bar{l} (layer model). This yields $x/(2\bar{l})$ layers, each with thickness $2\bar{l}$. The temperature difference between the two plates of a single layer is $2(T_2 - T_1)\overline{l}/x$. As in each layer the regime, by approximation, may be considered molecular, Eq. (3.88), as derived above, may be used. Now, the distance between the plates is equal to the thickness of the layer, and accommodation is set to 1:

$$P = \frac{1}{4} A p \bar{l} \bar{c} \frac{\kappa + 1}{\kappa - 1} \frac{1}{T} \frac{T_2 - T_1}{\kappa}, \text{ approximation in layer model.}$$
 (3.90)

Experimental investigations reveal that, in many cases, the conducted heat is proportional to the area A, to the difference in temperatures $T_2 - T_1$, and inversely proportional to the distance x between the plates, formulated in the following equation:

$$P = \lambda A \, \frac{T_2 - T_1}{x} \,. \tag{3.91}$$

This formulation defines the thermal conductivity λ , a property of the fluid between the plates.

Comparing the two previous equations, one obtains for the thermal conductivity of the gas

$$\lambda = \frac{1p\bar{l}\bar{c}\kappa + 1}{4T\kappa - 1}, \quad \text{approximation in layer model.}$$
 (3.92)

This equation may be rewritten by introducing the viscosity, Eq. (3.69), the molar heat capacity, Eq. (3.82), and the mean speed, Eq. (3.43):

$$\lambda = \frac{\kappa + 1}{2} \eta c_V$$
, approximation in layer model. (3.93)

For noble gases (isentropic exponent $\kappa = 5/3$), the numerical prefactor $(\kappa + 1)/2$ amounts to 4/ern - 0.15em3 = 1.33. In 1860, Maxwell determined the thermal conductivity in a qualitative approach and found an equation of the type 3.94 but, however, with a numerical prefactor of 1 instead of 4/3. With considerable mathematical effort, a correct calculation of the macroscopic thermal conductivity of a gas from microscopic particle-particle collisions is possible. As a result, an equation is obtained with a prefactor of 5/2 for noble gases. In 1913, Eucken adopted this equation empirically for other gases:

$$\lambda = \frac{9\kappa - 5}{4}\eta c_V \qquad (3.94)$$

Figure 3.16 shows experimental data of thermal conductivities for a selection of gas species.

The heat transport is then given by

$$P = \frac{9\kappa - 5}{4} \eta c_V A \frac{T_2 - T_1}{x} \quad \text{in the viscous regime.}$$
 (3.95)

Thermal conductivity λ is useful when describing steady-state heat transport. In dynamic processes with variable temperature, behavior is determined by heat transport (thermal conductivity) as well as by the ability to store heat. The ability to store heat is proportional to the density and to the specific heat capacity. The thermal diffusivity a is a convenient quantity to describe this type of process. It is

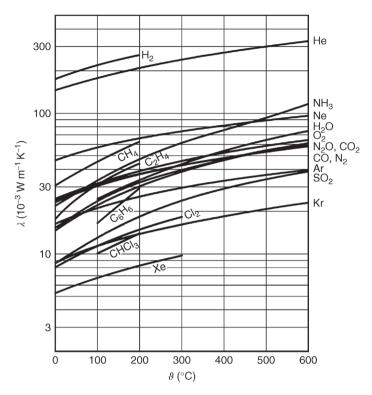


Figure 3.16 Thermal conductivities of selected gas species in the viscous range versus temperature. (Data taken from Ref. [3].)

defined as

$$a = \frac{\lambda}{\rho c_p}. (3.96)$$

In some applications, the heat transport over the whole pressure range is required. A rough approximation, which describes the limits of molecular and viscous regions correctly, is the expression

$$\frac{1}{P} = \frac{1}{P_{\text{molecular}}} + \frac{1}{P_{\text{viscous}}}$$
 (3.97a)

giving

$$P = \frac{pA(T_2 - T_1)}{8\frac{\kappa - 1}{\kappa + 1}\frac{T}{\bar{r}_{g_1}} + \frac{\kappa}{2}p}.$$
(3.97b)

The heat transfer over the whole pressure range is treated exactly for two parallel plates in Section 5.4.3 and for two coaxial cylinders in Section 5.4.4.

A Pirani vacuum gauge makes use of the pressure dependence of thermal conductivity. Here, the aim is to calculate the operational data of a typical gauge.

Typically, the pressure sensor is designed with cylinder symmetry (see Figures 13.26 and 13.27). On the axis, a thin wire is arranged, which is heated directly by means of an electrical current. A tube at ambient temperature is placed around the wire concentrically. Thus, a heat flux develops from the wire, through the surrounding gas, and toward the cylinder. The transported thermal power for this cylinder geometry can be calculated using the above equations for planar geometries. For this, the volume between the wire and the tube is divided into many virtual hollow cylinders with finite wall thickness, slid into one another telescopically. The arrangement of cylinders completely fills the volume. Imagining the cylinders to be unrolled gives the previously calculated geometry of parallel plates.

Usually, the wire diameter is small compared to the diameter of the tube. The gas particles therefore collide frequently with the tube but rarely with the wire. Due to the large number of collisions with the tube, the value of the energyaccommodation coefficient at the wall of the tube is negligible because practically all particles show a thermal energy corresponding to the temperature of the tube, after only few collisions. In contrast, the energy accommodation at the wire is relevant.

The length of the tube is assumed large compared to its diameter. By determining the relations for the radii r_1 , r_2 for the wire and tube, as well as the temperatures T_1 , T_2 for the wire and tube, the thermal power transported by the gas is calculated using Eq. (3.87) as well as Eqs. (3.48) and (3.80) by integration:

$$P = a_{E1} 2\pi r I \frac{T_1 - T_2}{T_2} \frac{f + 1}{8} \overline{c} p \quad \text{in the molecular regime,}$$
 (3.98)

$$P = \pi l \left[\lambda(T_1) + \lambda(T_2) \right] \frac{T_1 - T_2}{\ln(r_2/r_1)} \quad \text{in the viscous regime.}$$
 (3.99)

For better understanding, the power appearing in such a gauge is now calculated numerically. Specified data: wire diameter $d = 10 \,\mu\text{m}$, wire length 5 cm, and tube diameter 16 mm. The temperature of the wire and tube shall be 120 °C and 20 °C, respectively. The gas is air and the accommodation coefficient is 0.8. It follows that

For $p = 10 \, \text{Pa}$:

$$P = 0.8 \times 2\pi \times 5 \times 10^{-6} \text{ m} \cdot 0.05 \text{ m} \frac{393 \text{ K} - 293 \text{ K}}{293 \text{ K}} \cdot \frac{5+1}{8} 463 \text{ m s}^{-1} \cdot 10 \text{ Pa}$$

= 0.0015 W.

(3.100)

For
$$p = 10^5$$
 Pa:

$$P = \pi \times 0.05 \text{ m} \left[0.0322 \text{ W m}^{-1} \text{ K}^{-1} + 0.0256 \text{ W m}^{-1} \text{ K}^{-1} \right] \times \frac{393 \text{ K} - 293 \text{ K}}{\ln(0.008 \text{ m/5} \times 10^{-6} \text{ m})}$$

$$= 0.12 \text{ Pa}.$$
(3.101)

Transport of heat in the viscous regime is higher than in the molecular regime and independent of the pressure.

3.3.4 Diffusion

The motion of particles in a medium is referred to as diffusion. For example, if a bottle of perfume is opened in a room, after a while, the scent of the perfume is perceived at some distance. In spite of the high velocity of individual molecules, spreading of the perfume requires a considerable amount of time. This is because the free path of the perfume molecules in air is short (see Section 3.2.5), and because the directions change in collisions with air molecules. Thus, the path of a perfume molecule is a random zigzag route, and the distance between a molecule and its place of origin only increases slowly.

Now, the diffusion of a gas (species 1) in another gas (species 2) is assessed quantitatively. Gas 2 fills a volume homogeneously (constant particle number density). Gas 1 is added at a certain location; thus, initially, it is distributed inhomogeneously within the volume. Due to the inhomogeneous particle number density n_1 of the added gas, the thermal motion of all gas particles leads to a net particle flux, of species 1 directed opposite to the gradient of the number density. To simplify matters, the particle number density is assumed to change only in one dimension, the *z*-direction. *Fick's first law* describes the flux j_{NI} of molecule species 1 through an imaginary plane at position *z* (Figure 3.17):

$$j_{N1} = -D_{12} \frac{\mathrm{d}n_1}{\mathrm{d}z} \,. \tag{3.102}$$

This equation defines the diffusion coefficient D_{12} of gas 1 in gas 2.

For the *low-pressure case* (molecular regime), particle–particle collisions are negligible. Here, no real diffusion occurs; instead, the system rather features flow, which is covered in Section 4.4.

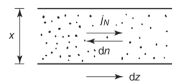


Figure 3.17 The diffusion flow is directed toward lower particle density.

In the high-pressure case (viscous regime), it is understandable that an increase in velocity c and in the free path \bar{l} of gas particles promotes the diffusion motion of particles. Accordingly, the qualitative behavior follows the formulation $D\bar{c}\bar{l}$. Eq. (3.55) previously described the mean free path for a single particle species. Assuming that the particles are hard spheres with diameter d, mathematically solving for diffusion in the model of the kinetic theory of gases yields

$$D_{12} \approx \frac{4}{3\pi} \cdot \frac{\sqrt{\overline{c}_2^1 + \overline{c}_2^2}}{(n_1 + n_2)(d_1 + d_2)^2}.$$
 (3.103)

An interesting special case is self-diffusion, featuring only one type of particles, that is, both gas species are of the same kind. Experiments aimed at investigating this phenomenon use (e.g., radioactively) marked individual particles that are observed while they spread. In the case of self-diffusion,

$$\overline{c}_1 = \overline{c}_2 = \overline{c}, \quad n_1 + n_2 = n, \quad d_1 = d_2 = d.$$
 (3.104)

Using Eq. (3.104), Eq. (3.103) simplifies to

$$D_{11} = \frac{\sqrt{2}}{3\pi} \cdot \frac{\overline{c}_1}{n_1 d_1^2}. (3.105)$$

Introducing the mean free path \bar{l} according to Eq. (3.55) and viscosity as in Eq. (3.72), in order to eliminate the particle diameter d, finally yields

$$D_{11} = \frac{2}{3}\overline{c}_1\overline{l}_1 = \frac{4}{3} \cdot \frac{\eta}{\rho} \quad . \tag{3.106}$$

In 1860, Maxwell had already found a similar expression by qualitative assessments, though with a prefactor only half as high. Eq. (3.103) produces values consistent with experimental data (Table 3.5).

Table 3.5 Diffusion constants for selected gas species in air at 20 °C and 1 bar.

Gas species	Diffusion constant D_{12} (10 ⁻⁵ m ² s ⁻¹) experimental values	Diffusion constant D_{12} (10 ⁻⁵ m ² s ⁻¹) calculated values			
H_2	7.2	7.4			
Не	7.1	6.5			
H_2O	2.5	1.9			
Ne	3.2	3.1			
N_2	2.2	2.0			
O_2	2.0	2.0			
Ar	1.9	1.9			
CO_2	1.5	1.5			
Kr	1.5	1.5			
Xe	1.2	1.2			

Calculated values were determined using Eq. (3.103).

3.4 **Real Gases**

3.4.1

Equations of State

Section 3.1.3 introduced the equation of state for an ideal gas as a relation between pressure, volume, and temperature. Solving for pressure yields

$$p = \frac{\nu RT}{V} = \frac{NkT}{V}. ag{3.107}$$

In 1840, Regnault conducted precise measurements revealing that real gases behave differently from what this equation of state describes, especially at low temperature and for high particle number densities. We will now investigate these deviations more precisely.

First, the deviations occurring at high particle number densities are examined. As described earlier, the gas particles are considered small spheres with diameter d. Thus, an individual gas particle, as well as a certain portion of gas, has an inherent volume. For high particle number density, the inherent volume of a gas is not negligibly small compared to the total volume available to the gas. Today, statistical mechanics and computer simulations can be used to solve the problem of the inherent volume in the hard-sphere model, yielding a modified equation of state,

$$p = \frac{NkT}{V} \cdot \frac{1 + y + y^2 - y^3}{(1 - y)^3},$$
(3.108)

where y represents the dimensionless ratio of the total inherent volume of all gas particles to the volume of the vessel, that is,

$$y = \frac{\pi d^3}{6} \cdot \frac{N}{V}.\tag{3.109}$$

Vacuum technology deals with dilute gases, featuring low particle densities and, therefore, $\gamma \ll 1$. Thus, the above equation of state, Eq. (3.108), may be approximated to first order in γ and gives

$$p = \frac{NkT}{V} \cdot \frac{1}{1 - 4\gamma}.\tag{3.110}$$

We will now discuss the deviations at low temperature. The reason for the different behavior of real gases, compared to the equation of state for an ideal gas, is that the gas particles attract one another when they come close (before actually touching in a collision). As a model conception, the gas particles can be thought of as being surrounded by an attractive field of force. This type of attractive force (cohesive force) is observed in liquids as well, and is known to create an excess pressure (inner pressure) in small drops of liquid. As for gases, a similar effect occurs when mutual attractive forces between particles reduce the pressure exerted on a confining wall. The value of the inner pressure (inherent pressure) is proportional to the volume collision rate χ (Eq. (3.60)) and, thus, proportional to the squared particle number density N/V. Therefore, the appropriate correction of the equation of state, Eq. (3.107), for the inner pressure will be proportional to $(N/V)^2$.

In 1873, van der Waals formulated an equation of state, named after him, taking into account the inherent volume and inner pressure of a gas:

$$p = \frac{\nu RT}{V - \nu b_{\rm m}} - \frac{\nu^2}{V^2} a_{\rm m} \qquad (3.111)$$

This equation includes two empirically found parameters $a_{
m m}$ and $b_{
m m}$, referred to as van der Waals constants. A comparison of the first term on the right-hand side of Eq. (3.111), describing the inherent volume, with the right-hand side of Eq. (3.110) shows that the parameter $b_{\rm m}/N_{\rm A}$ corresponds to the fourfold inherent volume of a gas particle:

$$b_{\rm m} = 4\frac{\pi}{6}d^3N_A. {(3.112)}$$

The parameter $a_{\rm m}$ gives the value of the inner pressure resulting from the attractive force.

Both parameters $a_{\rm m}$ and $b_{\rm m}$ can be obtained from precise measurements of the behavior of real gases. As experimental data show, $b_{\rm m}$ is nearly temperature independent, which is to be expected for fixed particle size, whereas a_m shows a significant dependence on temperature (of approximately $T^{-1/2}$). This is explained by the fact that the velocity of gas particles decreases when the temperature drops (as the velocity is proportional to $T^{1/2}$). Thus, the attractive force appears over a longer period during a collision and therefore has a greater effect.

From the physical point of view, van der Waals' approach (Eq. (3.111)) is unfavorable since the parameter $a_{\rm m}$ is assumed to be constant (= not temperature dependent) although, in fact, it is not. A far better way of describing the behavior of real gases is to formulate the temperature dependence of the attractive force's influence explicitly, as was done in the equation of state formulated 1949 by Redlich and Kwong [4]. However, for calculation of thermodynamic state variables, the involved mathematics is uncomfortable.

An alternative means of describing the behavior of a real gas is to introduce a formal power series in density or pressure (virial series) in the equation of state. Terminating at the first-order term and solving for pressure yields

$$p = \frac{\nu RT}{V} (1 + B''(T)p). \tag{3.113}$$

This equation introduces the so-called second virial coefficient B" that is temperature dependent.

We will now assume that the difference in behavior of real gas compared to ideal gas, described by $a_{\rm m}$, $b_{\rm m}$, and B'', is small, and a first-order approximation is adequate. By comparing van der Waals' equation, Eq. (3.111), with the virial equation, Eq. (3.113), the relationship between the virial coefficient and van der Waals constants is identified as

$$B''(T) = \frac{b_{\rm m}}{RT} - \frac{a_{\rm m}}{R^2 T^2}.$$
 (3.114)

Table 3.6 lists experimental data for van der Waals constants as well as viscosity and particle diameters calculated from these quantities. The diameter calculated from $b_{\rm m}$ should correspond to the diameter of the hard-sphere particle as the attractive force between particles.

Table 3.6 Properties of selected gas species at 20 °C, sorted according to relative particle mass M_r .

Gas species		M _r (-)	a _m m ^{6 Pa} mol ²	$b_{\rm m}~(\times 10^{-6}~{\rm m^3~mol}^{-1})$	d from b _m (nm)	$\eta~(imes 10^{-6}~{ m Pa~s})$	d from η (N m)
Hydrogen	H ₂	2.016	0.0244	26.6	0.276	8.82	0.274
Helium	He	4.003	0.0034	23.7	0.266	19.65	0.218
Methane	CH_4	16.043	0.2253	42.8	0.324	11.08	0.410
Ammonia	NH_3	17.031	0.4170	37.1	0.309	10.05	0.437
Water vapor	H_2O	18.015	0.5464	30.5	0.289	9.7	0.451
Neon	Ne	20.180	0.0211	17.1	0.238	31.50	0.258
Acetylene	C_2H_2	26.038	0.4390	51.4	0.344	10.08	0.486
Carbon monoxide	CO	28.010	0.1485	39.9	0.316	17.64	0.374
Nitrogen	N_2	28.013	0.1390	39.1	0.314	17.59	0.374
Ethylene	C_2H_4	28.054	0.4471	57.1	0.356	10.15	0.493
Ethane	C_2H_6	30.07	0.5489	63.8	0.370	9.29	0.524
Oxygen	O_2	31.999	0.1360	31.8	0.293	20.39	0.359
Hydrogen chloride	HCl	36.461	0.3667	40.8	0.319	14.08	0.447
Argon	Ar	39.948	0.1345	32.2	0.294	22.3	0.363
Carbon dioxide	CO_2	44.010	0.3592	42.7	0.324	14.88	0.456
Dinitrogen monoxide	N_2O	44.013	0.3782	44.2	0.327	14.52	0.461
Propane	C_3H_8	44.097	0.8664	84.5	0.406	8.18	0.615
n-Butane	C_4H_{10}	58.113	1.447	122.6	0.460	7.60	0.683
Sulfur dioxide	SO_2	64.065	0.6714	56.4	0.355	12.97	0.536
Chlorine	Cl_2	70.905	0.6493	56.2	0.355	12.87	0.552
Benzole	C_6H_6	78.114	1.800	115.4	0.451	7.5	0.741
Krypton	Kr	83.80	0.2318	39.8	0.316	25.07	0.412
Xenon	Xe	131.29	0.4194	51.1	0.343	22.79	0.484
Tetrachloromethane	CCl_4	153.822	2.039	138.3	0.479	11.9	0.697
Mercury	Hg	200.59	0.8093	17.0	0.238	22.6	0.540

 a_{m} and b_{m} are the van der Waals constants, and η is the viscosity. As indicated, the particle diameters d are calculated from b_{m} using Eqs. (3.112) and (3.75). Data taken from Refs. [3,5,6].

According to Eq. (3.114), the virial coefficient of nitrogen at 20 °C is calculated from van der Waals constants as

$$\begin{split} B''(T) &= \frac{39.1 \times 10^{-6} \text{ m}^3 \text{ mol}^{-1}}{8.314 \text{ J mol}^{-1} \text{ K}^{-1} \cdot 293 \text{ K}} \\ &- \frac{0.139 \text{ Pa m}^6 \text{ mol}^{-2}}{(8.314 \text{ J mol}^{-1} \text{ K}^{-1} \cdot 293 \text{ K})^2} \\ &= 1.61 \times 10^{-8} \text{ Pa}^{-1} - 2.34 \times 10^{-8} \text{ Pa}^{-1} = -0.73 \times 10^{-8} \text{ Pa}^{-1}. \end{split}$$

For comparison, the experimental value for $B'' = -0.24 \times 10^{-8} \, \text{Pa}^{-1}$ (Figure 3.18). The large deviation between calculated (Eg. (3.115)) and experimental values is understandable as the calculation subtracts two terms of nearly the same value. Thus, the result strongly depends on the values of the terms, which are uncertain due to the van der Waals constants.

As a result, reliable calculations of the gas state should use the virial equation, Eq. (3.113), with accurate virial coefficients (Figure 3.18 and Table A.8), instead of the van der Waals equation, Eq. (3.111).

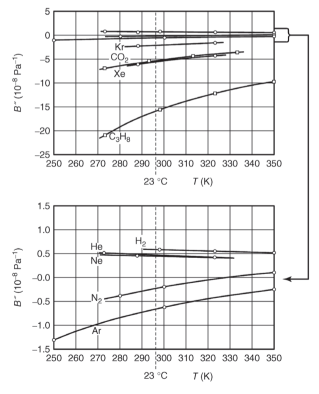


Figure 3.18 Second virial coefficient of selected gas species versus temperature. The lower plot shows a magnification of the upper, with the ordinate stretched by a factor of 10.

To this point, gas particles were described as small hard spheres that attract one another when they come close. This conception corresponds to the state of knowledge at the beginning of the twentieth century. Today, structures of atoms and molecules as well as their interactions during approach have been investigated thoroughly [3,8,9]. Calculations take into account the microscopic properties of individual gas particles in order to obtain the macroscopic properties of gases, as well as substances that, in fact, are made up of many individual particles.

As is known today, an atom consists of a nearly point-particle-like nucleus and a surrounding cloud of electrons. The density of the electron cloud is very high near the nucleus and drops gradually with increasing distance from the nucleus. By using quantum mechanics, the density can be calculated (Figure 3.19). A diatomic or polyatomic molecule is made up of two or more bound atoms with overlapping electron clouds. Thus, in contrast to a hard sphere, a gas particle does not have a definite diameter.

In complicated molecules such as water, the electron cloud, on one hand, can feature a permanent electrical dipole moment. On the other hand, particles in the vicinity can shift the electron cloud of simple atoms (e.g., noble gases) relative to the nucleus and, thus, induce an electrical dipole moment.

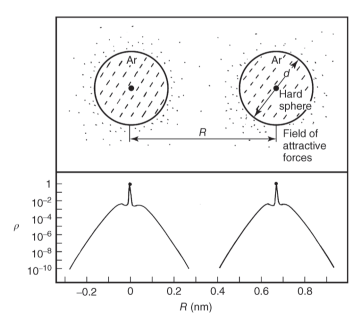


Figure 3.19 Model of gas particles, argon taken as an example. (a) Traditional conception of hard spheres with diameter d, attracting each other at short distance R. (b) Modern

conception of electron clouds with charge distribution ρ (normalized to maximum value). (From Ref. [7].)

When two gas particles approach, an electrical force develops due to the electrical dipole moments, even when the particles are still far apart. This force can be attractive or repellent, depending on the type of interacting particles and the symmetry properties of the electron cloud as a whole. In noble gases, the force is weakly attractive and the resulting potential energy behaves, with respect to internuclear distance R, as a function of R^{-6} . Thus, it drops rapidly with increasing distance.

When two colliding particles approach, their outer electron clouds overlap increasingly. The repellent electrostatic force between the electrons causes a rapidly increasing repellent force that finally exceeds the attractive force caused by dipole moments. As a rough approximation, the size of a particle can be described by defining the diameter as the internuclear distance at which the interaction shifts from far-range attractive to low-range repellent.

We have now formulated the attractive-soft-sphere model of a gas particle and have described the interaction force between two particles qualitatively. The microscopic interaction force determines collision behavior, and thus, macroscopic gas properties.

The interaction force is a result of the change in potential interaction energy that occurs when particle distance changes. Today, quantum mechanics provides the means to calculate the potential energy of two particles theoretically, and advanced experiments allow us to obtain detailed information on atomic interaction in targeted collision experiments. Figure 3.20 shows the corresponding data

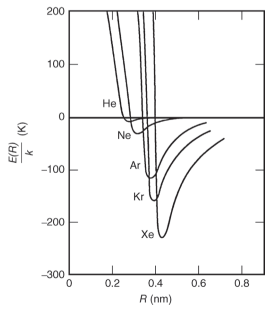


Figure 3.20 Potential energy E(R) between two noble gas atoms of the same species versus internuclear distance. The y-axis gives the energy divided by Boltzmann's constant k in Kelvin.

of noble gases. The energy's zero crossing appears at an internuclear distance of 0.3-0.4 nm, which is approximately in compliance with the particle diameter in the hard-sphere model.

As indicated in Figure 3.20, the size of atoms increases gradually with growing atomic number; the attractive force, however, intensifies considerably. When complex particles collide, the interaction energy also depends on the mutual orientation during the collision.

A relatively simple analytical function, referred to as the Lennard-Jones potential, approximates the actual potential energy E(R) between two particles fairly accurately. It contains only two parameters, the minimum energy ε and the particle distance σ at which the energy is zero (Figure 3.21):

$$E(R) = 4\varepsilon \left[\left(\frac{\sigma}{R} \right)^{12} - \left(\frac{\sigma}{R} \right)^{6} \right]. \tag{3.116}$$

The Lennard-Jones potential has a minimum of the potential energy at a distance $R_{\rm m}$ that is obtained by differentiating Eq. (3.116):

$$R_{\rm m} = 2^{1/6}\sigma = 1.12\sigma. \tag{3.117}$$

Figure 3.20 shows the potential energy between two noble gas atoms of the same species. Table 3.7 lists values of adopted parameters for Lennard-Jones potentials of different gases.

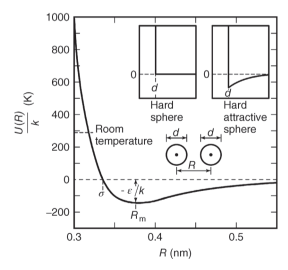


Figure 3.21 Potential energy of two argon atoms versus internuclear distance. In the plotted range, the adopted 12-6-Lennard-Jones potentials and precise potentials do not differ any more than approximately the line

widths used in the plot. For comparison, the inserted small plots show the potential energies in the hard-sphere and in the attractivehard-sphere models. (From Ref. [7].)

			$\sigma(\mathbf{nm})$	
Gas	A _r (–)	ϵ/\mathbf{k} (K)		
Не	4.00	11	0.27	
Ne	20.18	42	0.28	
Ar	39.94	142	0.34	
Kr	83.7	195	0.36	
Xe	131.3	270	0.39	
H_2	2.02	107	0.28	
N_2	28.02	103	0.36	
O_2	32.00	129	0.34	
CO_2	44.01	246	0.38	
CH ₄	16.04	152	0.37	
CF ₄	88.01	152	0.47	

Table 3.7 Mass number A_r , minimum ε , and root σ of the potential energy for selected gas

Literature values for ε scatter considerably. The reason is that the values are often derived from experimental data of the temperature-dependent viscosity. When adopting the values, the quality of adoption hardly changes if a greater change in ε is compensated by a slight change in σ . From Ref. [8].

A defined internuclear potential E(R) allows deriving the macroscopic values of viscosity, thermal conductivity, diffusion, and second virial coefficient. This is a time-consuming task as, initially, momentum and energy transfer for different collision geometries (head-on and grazing collisions) of two gas particles, and subsequently, the correct mean values across a large number of individual collisions have to be calculated. However, the problem is solvable, as was shown first by Chapman and Enskog in 1916.

The macroscopic quantities depend on the temperature: at elevated temperature, gas particles have more thermal kinetic energy. The interaction force between particles then less affects their paths and they can come closer to each other in spite of the repellent short-range force. Solving for viscosity η , thermal conductivity λ , self-diffusion D, and second virial coefficient B'' of a monatomic gas yields

$$\eta(T) = \frac{5\sqrt{2}kT}{8\pi} \frac{1}{\bar{c}} \frac{1}{\sigma^2 \Omega^{(2,2)^*}(kT/\epsilon)},\tag{3.118}$$

$$\lambda(T) = \frac{75}{128\sqrt{2}} k\overline{c} \frac{1}{\sigma^2} \frac{1}{\Omega^{(2,2)^*} (kT/\varepsilon)},$$
(3.119)

$$D_{11}(T) = \frac{3}{16\sqrt{2}} \,\overline{c} \, \frac{1}{n} \frac{1}{\sigma^2} \frac{1}{\Omega^{(1,1)^*} (kT/\varepsilon)},\tag{3.120}$$

$$B''(T) = \frac{2\pi}{3} \frac{\sigma^3}{kT} B^*(kT/\varepsilon). \tag{3.121}$$

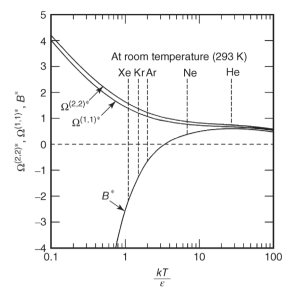


Figure 3.22 Reduced collision integrals and reduced second virial coefficient versus the parameter kT/ϵ (reduced temperature). For a representation of practical conditions, reduced

temperatures are marked by dashed lines for the indicated noble gases at room temperature. (From Ref. [7].)

These equations use the reduced collision integrals $\Omega^{(2,2)*}$ and $\Omega^{(1,1)*}$ as well as the reduced virial coefficient B^* . These standardized dimensionless quantities ultimately contain the interatomic potential via the ratio of thermal energy kTand energy ε at the minimum potential. This dimensionless ratio is referred to as reduced temperature. Assuming a Lennard-Jones potential, these quantities can be calculated numerically (Figure 3.22).

An interpretation of the reduced collision integral Ω^* is that a real gas behaves just as a gas of hard spheres with $\sigma\sqrt{\Omega^*}$ as the effective diameter, σ being the particle distance where the Lennard-Jones potential is zero. The temperature dependence of the reduced collision integrals $\Omega^{(2,2)^*}$ and $\Omega^{(1,1)^*}$ as well as the reduced virial coefficient B^* (Figure 3.22) is understandable: at elevated temperature ($kT/\varepsilon > 30$), particles behave as hard spheres with a diameter slightly below σ . At even higher temperatures, during collisions, the particles approach more and more, and thus, seem to shrink. At low temperatures (kT/silon < 1), the mutual attraction between particles gains in importance. The effective range of interaction forces increases to larger distances and the particle route is disturbed, corresponding to a collision. Thus, toward lower temperatures, particles seem to increase in size.

The introduction of an atomic potential allows predicting precisely different properties of real gases fundamentally.

Calculating viscosity, thermal conductivity, self-diffusion (at 1 bar), and second virial coefficient of argon at 20 °C, using Eqs. (3.118)–(3.121).

According to Table 3.7, $\sigma = 0.34 \times 10^{-9}$ m and $\epsilon/k = 142$ K.

As follows, the argument of the collision integrals $kT/\epsilon = 293 \text{ K}/142 \text{ K} = 2.06$.

The reduced collision integrals and the reduced virial coefficient read (data from Figure 3.22)

$$\Omega^{(2,2)^*}(2.06) = 1.16, \ \Omega^{(1,1)^*}(2.06) = 1.06, \ \text{and} \ \textit{B}^*(2.06) = -0.6.$$

This calculates to

$$\eta = \frac{5\sqrt{2}}{8\pi} \times \frac{1.38 \times 10^{-23} \text{ J K}^{-1} \cdot 293 \text{ K}}{394 \text{ m s}^{-1}} \times \frac{1}{(0.34 \times 10^{-9} \text{ m})^2} \times \frac{1}{1.16}$$

$$= 22 \times 10^{-6} \text{ Pa s.}$$
(3.122)

$$\lambda = \frac{75}{128\sqrt{2}} 1.38 \times 10^{-23} \,\text{J K}^{-1} \cdot 394 \,\text{m s}^{-1} \,\frac{1}{(0.34 \times 10^{-9} \,\text{m})^2} \cdot \frac{1}{1.16}$$

$$= 17 \times 10^{-3} \,\text{W m}^{-1} \,\text{K}^{-1}. \tag{3.123}$$

$$D_{11} = \frac{3}{16\sqrt{2}} 394 \,\mathrm{m \, s^{-1}} \, \frac{\mathrm{m}^3}{2.47 \times 10^{25}} \cdot \frac{1}{(0.34 \times 10^{-9} \,\mathrm{m})^2} \cdot \frac{1}{1.06}$$

$$= 1.7 \times 10^{-5} \,\mathrm{m}^2 \,\mathrm{s}^{-1}, \qquad (3.124)$$

$$B'' = \frac{2\pi}{3} \cdot \frac{(0.34 \times 10^{-9} \text{ m})^3}{1.38 \times 10^{-23} \text{ J K}^{-1} \times 293 \text{ K}} \cdot (-0.6)$$
$$= -1.2 \times 10^{-8} \text{ Pa}^{-1}.$$
 (3.125)

The calculated values can be compared to experimental data:

$$\eta = 22.3 \times 10^{-6} \text{ Pa s},$$

 $\lambda = 17.3 \times 10^{-3} \text{ W m}^{-1} \text{ K}^{-1},$
 $D_{11} = 1.8 \times 10^{-5} \text{ m}^2 \text{ s}^{-1},$
 $B'' = -0.74 \times 10^{-8} \text{ Pa}^{-1}.$

Theoretical and experimental values are consistent, except for the virial coefficient B". The supposed reason for this deviation is that the calculated value B'' is determined directly by the reduced virial coefficient B^* , which is very sensitive to the abscissa kT/ϵ , that is, to the minimum in potential ϵ .

3.5 **Vapors**

3.5.1

Saturation Vapor Pressure

The state of a certain amount of substance inside a vessel depends on the prevailing conditions and the volume. Figure 3.23 shows a vessel whose volume can be varied by moving a piston.

At sufficiently high pressure, only liquid (or solid) is present inside the vessel. If the vessel's volume is increased by pulling the piston upward, additional volume is created. This volume does not remain empty. Instead, the free surface of the substance releases particles due to thermal motion, thus filling the additional volume. This process is referred to as evaporation (or sublimation) (Figure 3.24).

We will assume that the walls of the vessel reflect the released gas particles diffusely so that they do not stick to the wall. Thus, a particle emitted by the surface will eventually revisit the surface after a series of collisions with other particles or the walls. With a certain probability, it will condense here, that is, will be reintegrated into the liquid (or solid), or may again be reflected diffusely.

After a sufficient amount of time, a closed system establishes a steady-state equilibrium in terms of the amount of evaporating (sublimating) and condensing particles. This state is referred to as saturation, and the resulting pressure is called saturation vapor pressure (Figure 3.23b). If the piston is fairly close to the bottom end, the main part of the substance is in the liquid (or solid) phase

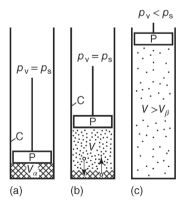


Figure 3.23 Phase transition. P piston; V volume; checkered area: liquid or solid; dotted: gas; p_V : pressure in volume V_i p_s : saturation pressure of substance. (a) Liquid only, (b) coexisting liquid and gas, (c) gas only.

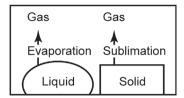


Figure 3.24 Phase transitions.

and only a small proportion is in gaseous condition. If the piston's position is further toward the top, most of the substance is gaseous and only a minor part is in the liquid (or solid) state. The saturation vapor pressure always has the same value, independent of the piston's position. The liquid (or solid) substance can be interpreted as a kind of reservoir. If the piston is pulled up even further, at some point, the reservoir will be depleted and the pressure drops (Figure 3.23c).

Saturation vapor pressure $p_{\rm s}$ depends only on the kind of substance and the temperature. Thermodynamics provides a simple model to describe the temperature dependence of saturation vapor pressure. The predominant quantity is the energy necessary to release a particle from the liquid or solid. This energy is given by the specific enthalpy of vaporization Δh , that is, the enthalpy of vaporization per unit mass. Thermodynamic considerations show that a simple function describes the dependence of saturation vapor pressure and temperature T as well as specific enthalpy of vaporization Δh . This relationship is expressed in the *Clausius–Clapeyron equation*,

$$\frac{\mathrm{d}p_{\mathrm{s}}}{\mathrm{d}T} = \frac{\Delta h}{\Delta \nu} \cdot \frac{1}{T}.\tag{3.126}$$

This equation introduces $\Delta \nu$, a quantity describing the increase in specific volume (= volume/mass) for the transition from the liquid (or solid) state to the gas phase.

The saturation vapor pressure is obtained by integrating the *Clausius–Clapeyron* equation over temperature. To simplify, we will assume the specific volume of substance in gaseous condition $\nu=R_{\rm s}T/p_{\rm s}$, and negligible (zero) in the liquid or solid state. Furthermore, the specific enthalpy of vaporization Δh shall be temperature independent. The (arbitrary) starting point of the integration is the boiling point $T_{\rm B.P.}$, that is, the temperature at which the saturation vapor pressure is equal to standard pressure $p_{\rm n}=101~325~{\rm Pa}$. Integration of Eq. (3.126) then yields

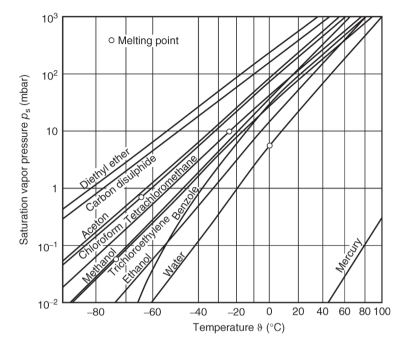


Figure 3.25 Saturation vapor pressures of selected solvents.

$$\ln \frac{p_{\rm s}}{p_{\rm n}} = \frac{\Delta h}{R_{\rm s}} \left(\frac{1}{T_{\rm B.P.}} - \frac{1}{T} \right).$$
(3.127)

A straight line should be the result when plotting the saturation vapor pressure in a diagram with a logarithmic-scale ordinate, against the inverse temperature T^{-1} on a linear abscissa, according to this equation. This chart is named the Arrhenius plot.

Experimental data (Figures 3.25 and 3.26), indeed, appear nearly as straight lines across several orders of magnitude. The fact that the enthalpy of vaporization is not, as assumed, constant, but drops with an increase in temperature, causes the slight bend. This is understandable as the thermal motion of liquids and solids increases with temperature, the inner cohesion weakens, and, thus, releasing particles (evaporation) requires less energy. At the socalled *critical point*, the enthalpy of vaporization actually drops to zero. Because the Clausius-Clapeyron equation tends to describe the strong change in saturation vapor pressure with temperature well across many orders of magnitude, it is used even in accurate analyses of experimental data. However, correction values for temperature dependence are then added to the righthand side of the equation.

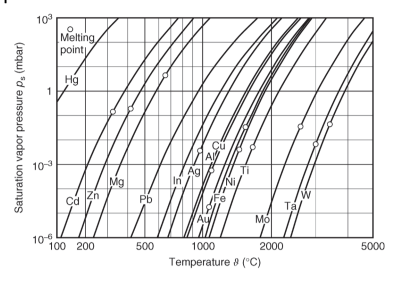


Figure 3.26 Saturation vapor pressures of selected metals.

Calculating saturation vapor pressure of water vapor at 20 °C, using Eq. (3.127).

The boiling point of water (according to the International Temperature Scale ITS-90), is $T = 373.124 \, \text{K}$. The specific gas constant can be calculated from the particle mass, $R_{\rm s}=461.5\,{\rm J\,kg^{-1}\,K^{-1}}$. At 20 °C, the specific heat of vaporization is 2.454 MJ/kg, and drops to 2.257 MJ/kg at 100 °C. For rough approximation, we will use the average 2.36 MJ/kg for this temperature interval. With these values, we can calculate the saturation vapor pressure at 20 °C, using Eq. (3.127):

$$\begin{split} p_{\rm s} &= p_{\rm n} \exp \left[\frac{\Delta h}{R_{\rm s}} \left(\frac{1}{T_{\rm B.P.}} - \frac{1}{T} \right) \right] \\ &= 101\,325\,{\rm Pa}\,\exp \left[\frac{2.36\times10^6\,{\rm J\,kg^{-1}}}{461.5{\rm J\,kg^{-1}\,K^{-1}}} \left(\frac{1}{373.12\,{\rm K}} - \frac{1}{293.15\,{\rm K}} \right) \right] \\ &= 101\,325\,{\rm Pa}\,\exp \left[5114\,{\rm K} \frac{-7.31\times10^{-4}}{{\rm K}} \right] = 2411\,{\rm Pa}. \end{split}$$

For comparison, the experimental value of the saturation vapor pressure of water at 20 °C is $p_{\rm s} = 2338 \, {\rm Pa}$.

Temperature (K)	Mass evaporation rate with respect to surface area $(kg m^{-2} s^{-1})$	Saturation vapor pressure (Pa)	Mean particle velocity (m s ⁻¹)
2000	1.76×10^{-12}	1.33×10^{-9}	480
2400	4.26×10^{-9}	3.52×10^{-6}	526
2800	1.10×10^{-6}	9.84×10^{-4}	568
3200	6.38×10^{-5}	6.13×10^{-2}	607
3600	1.51×10^{-3}	1.53	644

Table 3.8 Vacuum evaporation of tungsten.

From Ref. [6].

3.5.2 **Evaporation Rate**

At saturation vapor pressure, a system is in steady-state equilibrium of particles released by the surface of a liquid or solid, and of particles impinging and condensing from the gas phase. Surface evaporation rate and surface condensation rate are equal. The latter is calculated from the probability of condensation σ_c and the collision rate (Eq. (3.48)):

$$\frac{\text{Evaporating particles}}{\text{Area} \cdot \text{Time}} = \frac{\text{Condensating particles}}{\text{Area} \cdot \text{Time}}$$

$$= \sigma_c \frac{n_s \overline{c}}{4} = \sigma_c \frac{p_s \overline{c}}{4kT}.$$
(3.129)

Here, n_s denominates the particle number density and p_s the pressure of the saturated vapor.

The situation changes when released particles do not return because, for example, they are pumped out, drawn away by a gas flow, or simply condensate on the chamber walls. In these cases, the amount of liquid or solid continuously decreases due to evaporation.

Assuming that none of the evaporating gas particles return to the surface and condensate there, the particle loss per unit area and time is given by the surface evaporation rate, Eq. (3.129).

From this, the mass loss per unit area and time (specific mass flow rate) is obtained by multiplying with the particle mass m_P :

$$\frac{\text{Evaporating mass}}{\text{Area} \cdot \text{Time}} = \sigma_{c} \frac{p_{s} \overline{c} m_{P}}{4kT} = \sigma_{c} \frac{2p_{s}}{\pi \overline{c}}.$$
 (3.130)

Experimental data of very low vapor pressures, as for refractory metals such as tungsten, molybdenum, and tantalum (Table 3.8), are obtained by measuring the mass evaporation rate and subsequently calculating the saturation vapor pressure using Eq. (3.130). Here, the condensation coefficient is practically $\sigma_c = 1.00$.

According to Eq. (3.130), the mass loss per unit time of a tungsten wire (diameter 0.1 mm, length 100 mm) glowing in a vacuum at 3200 K is

$$\frac{\text{Mass}}{\text{Time}} = \sigma_c \frac{2p_s}{\pi \overline{c}} A = 1 \frac{2 \times 6.13 \times 10^{-2} \text{ Pa}}{\pi 607 \text{ m s}^{-1}} \pi 10^{-4} \text{ m} \cdot 0.1 \text{ m}$$

$$= 2 \times 10^{-9} \text{ kg s}^{-1} = 7.2 \times 10^{-6} \text{ kg h}^{-1}.$$
(3.131)

Since tungsten has a density of $19\,254\,\mathrm{kg}\,\mathrm{m}^{-3}$, the mass of the wire is $1.5\times10^{-5}\,\mathrm{kg}$. At such high operating temperature, nearly 1% of the wire's mass evaporates per minute.

If the volume around the wire contains an additional gas with which the evaporating gas particles can collide, a portion of the particles may be reflected and can return to the surface. This reduces the net evaporation rate. A region, mostly saturated with evaporated particles, thus builds up in the immediate vicinity of the wire. The ratio of the predominant partial vapor pressure here and the saturation vapor pressure is referred to as the *saturation ratio*. For water vapor in air, this corresponds to the *relative humidity*.

The reduction of evaporation rate by additional gas is used in gas-filled lamps. By filling the bulb with a gas, the evaporation of the wire drops by several orders of magnitude, and thus, increases lifetime. Heavy gas particles are particularly appropriate. Kr ($M_{\rm r}=80$) is more favorable than argon ($M_{\rm r}=40$), but also more expensive.

Example 3.18							
The tungsten filament in a gas-filled bulb has an operating temperature of 2870 K. The bulb is filled with an $\rm Ar-N_2$ gas mixture. The surface evaporation rate was measured.							
Pressure of filling gas	0	1×10^3	5×10^3	2×10^4	1×10^5	3×10^5	
86% Ar, 14% N ₂ (Pa)							
Surface mass evaporation	230	66	31	14	4.1	1.5	
Rate $(10^{-8} \text{ kg m}^{-2} \text{ s}^{-1})$							
Vapor pressure/saturation	0	0.73	0.87	0.94	0.982	0.9935	
vapor pressure							

During evaporation, the evaporating substance loses the heat of evaporation. Thus, if it is not heated, evaporation cools the substance. With falling temperature, the evaporation rate drops.

Calculation of the evaporation of ethanol (ethyl alcohol, C_2H_5OH , $M_r = 46$). A container holding ethyl alcohol at 27 °C is placed in a vacuum chamber. Surface $A = 100 \text{ cm}^2$, the filling height is 10 cm, and the initial volume is 1 liter.

According to Figure 3.25, the saturation vapor pressure $p_s = 1 \times 10^4$ Pa. The condensation coefficient $\sigma_c = 0.024$. Following Eq. (3.130), the evaporating mass per time

$$\frac{\text{Mass}}{\text{Time}} = \frac{m}{t} = \sigma_c \frac{2p_s}{\pi \overline{c}} A = 0.024 \frac{2 \times 1 \times 10^4 \text{ Pa}}{\pi \cdot 372 \text{ m s}^{-1}} 1 \times 10^{-2} \text{ m}^2$$

$$= 4.1 \times 10^{-3} \text{ kg s}^{-1}.$$
(3.132)

If this amount of ethanol vapor is to be pumped out, the pump must have a pumping speed

$$\frac{\text{Volume}}{\text{Time}} = \frac{m}{t} \cdot \frac{R_s T}{p} = 4.1 \times 10^{-3} \frac{\text{kg}}{\text{s}} \cdot \frac{181 \text{ J kg}^{-1} \text{ K}^{-1} \cdot 300 \text{ K}}{1 \times 10^4 \text{ Pa}}$$

$$= 0.022 \text{ m}^3 \text{ s}^{-1} = 80 \text{ m}^3 \text{ h}^{-1}.$$
(3.133)

The specific heat of evaporation of ethanol is 840 kJ kg⁻¹. Thus, the thermal power removed by evaporation is

$$P = q_{\rm s} \frac{\rm dm}{\rm dt} = 840 \times 10^3 \,\mathrm{J \, kg^{-1}} \,4.1 \times 10^{-3} \,\mathrm{kg \, s^{-1}} = 3.44 \times 10^3 \,\mathrm{J \, s^{-1}}$$

$$= 3.44 \,\mathrm{kW}$$
(3.134)

If this heat is not replaced in some way, the liquid will cool rapidly. The initial amount of 1ℓ has a mass of 0.79 kg. For a specific heat capacity $c = 2.43 \text{ kJ kg}^{-1} \text{ K}^{-1}$, the temperature drop per unit time

$$\frac{dT}{dt} = \frac{1}{mc}P = \frac{1}{0.79 \text{ kg} \cdot 2430 \text{ J kg}^{-1} \text{ K}^{-1}} 3.44 \times 10^3 \text{ J s}^{-1} = 1.8 \text{ K s}^{-1}. (3.135)$$

Without an external heating, the liquid would cool from 27 °C to 9 °C in only 10 s, and the saturation vapor pressure would drop to 3×10^3 Pa.

References

- 1 Preston-Thomas, H. (1990) The International Temperature Scale of 1990 (ITS-90). Metrologia, 27, 3-10.
- 2 Chapman, S. and Cowling, T.G. (1970) The Mathematical Theory of Non-Uniform Gases, 3rd edn, Cambridge University Press, Cambridge, UK.
- 3 Kaye, G.W.C. and Laby, T.H. (1986) Tables of Physical and Chemical Constants, Longman, Harlow.
- 4 Redlich, O. and Kwong, J.N.S. (1949) On the thermodynamics of solutions - an equation of state, fugacities of gaseous solutions. Chem. Rev., 44, 233-245.

- 5 Blanke, W. (ed.) (1989) Thermophysikalische Stoffgrößen, Springer, Berlin.
- 6 Lide, D.R. (1990) CRC Handbook of Chemistry and Physics, 71th edn, CRC Press, Boca Raton, FL.
- 7 Jitschin, W. (1999) Vakuum-Lexikon, Wiley-VCH Verlag GmbH, Weinheim.
- 8 Reid, R.C., Prausnitz, J.M., and Poling, B.E. (1987) The Properties of Gases and Liquids, McGraw-Hill, New York.
- 9 Maitland, G.C., Rigby, M., Smith, E.B., and Wakeham, W.A. (1981) Intermolecular Forces, Clarendon Press, Oxford.

Comprehensive general treatments of the subject

Edelmann, C. (1998) Vakuumphysik, Spektrum Akademischer Verlag, Heidelberg.

Lafferty, J.M. (ed.) (1998) Foundations of Vacuum Science and Technology, John Wiley & Sons, Inc., New York.

4

Gas Flow

Prof. Dr. Wolfgang Jitschin

University of Applied Sciences, Vakuumlabor, Wiesenstr. 14, 35391, Giessen, Germany

This chapter covers various types of gas flows, each developing in specific geometry and in characteristic pressure range.

4.1

Types of Flows and Definitions

4.1.1

Characterizing Flow, Knudsen Number, and Reynolds Number

Gas flow patterns play an important role in vacuum technology. When a vessel is evacuated, the gas that initially filled the vessel flows to the pump through tubes. During operation of the vessel, gas released by components (desorption) or supplied to the process flows from high-pressure to low-pressure regions. Knowledge of flow patterns is vital for designing vacuum systems intelligently and understanding their performance characteristics.

Flow (or flux) is a three-dimensional movement of substance. In a gas, the thermal motion of individual gas particles, as well as macroscopic forces due to local pressure deviations, causes flow. Pressure forces, inertial forces, and frictional forces determine flow behavior. Gravity, however, is usually negligible for gas flow. Usually, the total gas flow through a tube is of interest, but, in certain cases, knowledge of local flow densities in an apparatus is required.

Depending on the prevailing conditions, different types of flows arise. In order to understand flow patterns, it is favorable to consider the different types of flows individually in their pure form. Figure 4.1 shows the types of flows that occur in a tube of arbitrary length.

Depending on pressure and the cross dimensions of a tube, three types of flows can be differentiated:

 For sufficiently low pressure, the mean free path of gas particles is high, compared with the cross dimensions of the tube. Any mutual particle collision

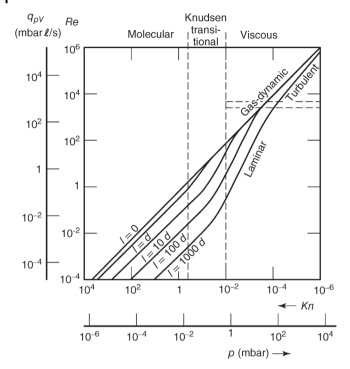


Figure 4.1 Flow types in tubes with circular cross section, diameter d=1 cm, and length l as indicated taken as an example. The gas is air at 20 °C. Inlet pressure is taken as abscissa and the outlet pressure is assumed negligible.

hardly occurs. Each gas particle travels through the tube due to its thermal motion, independent of other particles. However, frequent collisions with the tube walls cause a zigzag route. On average, the paths of many individual particles combine to form the macroscopic flow behavior. This situation is referred to as single-particle motion or *molecular flow*.

- 2) Under high pressure, the mean free path of gas particles is much lower than the cross dimensions of the tube. The particles experience frequent mutual collisions, thereby exchanging momentum and energy continuously. Even a small volume contains many frequently colliding particles. Thus, the gas behaves as a continuum. A flow is the result of local pressure gradients. This situation is referred to as continuum flow or *viscous flow*.
- 3) The medium-pressure range is characterized by a transition between molecular and viscous flows. In this transition, collisions of gas particles with the wall occur just about as often as mutual collisions among gas particles. This situation is referred to as transitional flow or Knudsen flow.

Thus, for a particular type of flow to occur, two main criteria can be identified: one criterion is the mean free path of gas particles in relation to the cross dimensions of the tube (for circular cross sections, the diameter). The second

criterion is the velocity of flow for given cross dimensions of the tube and internal friction of the gas. Thus, two dimensionless characteristic numbers may be defined to describe these criteria quantitatively.

The Knudsen number Kn is the ratio of the mean free path \bar{l} of the gas particles between two particle-particle collisions and the characteristic geometrical dimension d of the tube's cross section (for circular tube cross sections, the diameter):

$$Kn := \frac{\bar{l}}{d}.\tag{4.1}$$

As shown in Chapter 3, the mean free path can be obtained from viscosity n(Eq. (3.72)). Thus, for practical reasons, Eq. (4.1) can be rewritten as

$$Kn = \frac{\pi}{4} \cdot \frac{\overline{c}\eta}{pd},\tag{4.2}$$

denoting that the Knudsen number is inversely proportional to the pressure. A high Knudsen number indicates low pressure, and thus molecular flow, whereas a low value of the Knudsen number suggests viscous flow. Transition between the two types of flows is smooth and leads to a change in gas flow through the tube. The limiting cases of molecular or viscous flow are approximately reached when roughly 90% of this change in flow has established. The quantitative investigations described below show that this assumption leads to the following conditions:

$$Kn > 0.5$$
, molecular flow,
 $0.5 > Kn > 0.01$, transitional flow,
 $Kn < 0.01$, viscous flow. (4.3)

We will now investigate the second criterion for the type of flow: the velocity of flow. The velocity ν of a gas flow is the mean velocity component of the gas particles in the direction of the tube. Usually, the velocity's mean value is given as an average across the tube's cross section.

In the case of molecular flow, the individual gas particles travel back and forth between the walls of the tube with thermal velocity. A particle's direction after hitting the wall is (nearly) independent of its direction prior to the collision. Thus, a zigzag route develops (Figure 4.2a). The geometry of the tube determines the resulting velocity of flow.

The situation is different in the case of viscous flow. Here, three types of flows in a tube are differentiated. The length of the tube determines the type of flow (Figure 4.2b-d).

1) Initially, the gas has to leave a reservoir (vessel) to reach the entrance of the tube. Subsequently, it streams into the tube (Figure 4.2b). Here, the gas accelerates from a quiescent state (velocity of flow equals zero) to a finite velocity of flow. This process requires acceleration energy that is taken from pressure energy (pressure drops) and thermal energy (temperature drops).

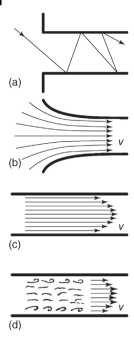


Figure 4.2 Different types of gas flows. (a) Molecular flow. (b–d) Different types of viscous flows: gas-dynamic (intake flow), laminar, and turbulent.

Thus, as a volume element of gas travels along a path, velocity rises, and simultaneously temperature and pressure drop. For short distances, wall friction is usually negligible. This so-called *intake flow* is a particular type of *gasdynamic flow*.

- 2) Now, the gas flows through the tube. The velocity of flow at the inlet is approximately constant across the complete cross section. As the gas continues its way through the tube, the gas layers near the walls decelerate, and the velocity of flow drops to zero in the boundary layer at the wall. The thickness of the boundary layer increases along the way. The velocity of flow, the friction behavior of the gas, and the dimensions of the cross section determine the type of flow that develops after a certain intake stretch. For low velocities, all individual volume elements move in the direction of the tube. Now, the volume elements in the center of the tube move quicker than the volume elements at the boundary of the tube. Thus, a velocity profile develops across the cross section of the tube (Figure 4.2c). This type of flow is referred to as *laminar flow*.
- 3) If, however, flow velocity is high, frictional forces are high as well because they are determined by flow velocity. A volume element, traveling at higher velocity and some distance from the wall of the tube, is deflected toward the wall by the decelerating action of the slower moving layers near the wall. The deflecting effects increase with friction and thus velocity, whereas the inertia

of mass, which tends to preserve the direction of flow, remains unchanged by a change in velocity. Thus, for sufficiently high velocities, deflecting forces dominate and the flow shows turbulences and eddies (Figure 4.2d). The criterion for turbulences to develop is the ratio of frictional force (proportional to gas viscosity η) and inertia of mass (proportional to gas density ρ) for a specified velocity of flow ν (cross-section average) and specified cross section. Typically, the *Reynolds number Re* is used to describe this criterion:

$$Re := -\frac{\rho}{\eta} vd. \tag{4.4}$$

The quantity d characterizes the cross section of the tube. For a downpipe, this corresponds to the diameter d.

$$Re < 2300$$
, laminar flow,
 $Re > 4000$, turbulent flow. (4.5)

4.1.2

Gas Flow, Throughput, and Pumping Speed

The flow rate q of a gas flowing through a duct is defined as transported gas per time. Several approaches of describing the amount of gas yield several different types of flow rates:

Volume flow rate:
$$q_V = \frac{\Delta V}{\Delta t} = \dot{V}, \quad [q_V] = \text{m}^3 \text{ s}^{-1}.$$
 (4.6)

Mass flow rate:
$$q_m = \frac{\Delta m}{\Delta t} = \dot{m}, \quad [q_m] = \text{kg s}^{-1}.$$
 (4.7)

Molar flow rate :
$$q_{\nu} = \frac{\Delta \nu}{\Delta t} = \dot{\nu}, \quad [q_{\nu}] = \text{mol s}^{-1}.$$
 (4.8)

Particle flow rate:
$$q_N = \frac{\Delta N}{\Delta t} = \dot{N}, \quad [q_N] = s^{-1}.$$
 (4.9)

At times, the terms rate of flow and flux rate are used synonymously when referring to *flow rate*. It should be considered that the flow rate can change along the length of the tube. For example, the volume flow rate at the end of a tube is higher than that at the beginning because the pressure drops along the tube and, correspondingly, the volume of the gas increases.

Furthermore, the concept of pV flow or throughput is used frequently:

$$pV$$
 flow : $q_{pV} = p\dot{V}$, $[q_{pV}] = \text{Pam}^3 \text{ s}^{-1} = 10 \text{ mbar } \ell \text{ s}^{-1}$. (4.10)

Using the equation of state of an ideal gas (Eqs. (3.18)–(3.20)),

$$pV = NkT = mR_sT = \nu RT, \tag{4.11}$$

and the definition of the mean particle speed (Eq. (3.43)),

$$\overline{c} = \sqrt{\frac{8}{\pi} \cdot \frac{kT}{m_{\rm P}}} = \sqrt{\frac{8}{\pi} R_{\rm s} T} = \sqrt{\frac{8}{\pi} \cdot \frac{p}{\rho}},\tag{4.12}$$

the different types of flows can be converted:

$$q_V = \frac{q_{pV}}{p},\tag{4.13}$$

$$q_{m} = \frac{q_{pV}}{R_{c}T} = \frac{8}{\pi} \cdot \frac{q_{pV}}{\overline{c}^{2}} = Mq_{\nu}, \tag{4.14}$$

$$q_{\nu} = \frac{q_{pV}}{RT},\tag{4.15}$$

$$q_N = \frac{q_{pV}}{kT}. (4.16)$$

Table 4.1 lists a number of common units for gas flow.

Table 4.1 Conversion of selected common units for gas flow.

Unit	Conversion		Definition
Pa m ³ s ⁻¹	= 1	Pa $\mathrm{m^3~s^{-1}}$	pV flow of 1 Pa m 3 s $^{-1}$
$\operatorname{mbar} \ell \operatorname{s}^{-1}$	= 0.1	$\mathrm{Pa}\ \mathrm{m}^3\ \mathrm{s}^{-1}$	pV flow of 1 mbar ℓ s ⁻¹
$\operatorname{Torr} \mathscr{C} \operatorname{s}^{-1}$	= 0.133322	$\mathrm{Pa}\ \mathrm{m}^3\ \mathrm{s}^{-1}$	pV flow of 1 Torr ℓ s ⁻¹
$\rm atm~cm^3~s^{-1}$	= 0.101325	$\mathrm{Pa}\ \mathrm{m}^3\ \mathrm{s}^{-1}$	pV flow of 1 atm cm 3 s $^{-1}$
lusec	= 0.000133322	Pa $\mathrm{m}^3~\mathrm{s}^{-1}$	pV flow of 1ℓ $\mu \mathrm{mHg}\mathrm{s}^{-1}$
sccm	⇔0.0018124	Pa m 3 s $^{-1}$ at 20 $^{\circ}$ C	Flow of standard cm ³ min ⁻¹
slm	⇔1.8124	Pa m 3 s $^{-1}$ at 20 $^{\circ}$ C	Flow of standard ℓ min ⁻¹ = 10^3 sccm
$mols^{-1}$	⇔2437.4	Pa $\rm m^3~s^{-1}$ at 20 $^{\circ}C$	Molar flow per second

The prefix "standard" refers to gas volume under standard conditions (101 325 Pa, 0 °C).

Example 4.1

Flowmeters often measure in "sccm" (standard cubic centimeters per minute). 1 sccm means a gas flow of 1 cm³ min⁻¹, referring to standard conditions ($p_{\rm n}=$ 101 325 Pa, $\theta_{\rm n}=$ 0 °C). Conversion to pV flow at 20 °C is as follows:

$$1 \, \text{sccm} \Leftrightarrow \frac{101\,325\,\text{Pa} \times 1\,\text{cm}_{273\,\text{K}}^3}{\text{min}} \times \frac{293.15\,\text{K}}{273.15\,\text{K}}$$

$$= 1.8124 \times 10^{-3}\,\text{Pa}\,\text{m}^3\,\text{s}^{-1} = 1.8124 \times 10^{-2}\,\text{mbar}\,\ell\,\text{s}^{-1}\,\text{at}\,20\,^\circ\text{C}. \tag{4.17}$$

Example 4.2

The permissible leakage of an air conditioner is 3 g per year. What is the pV value (at 20 °C) for the leakage? Tetrafluoroethylene R134a (CH₂F-CF₃), with a total mass number of 102, is the refrigerating medium. With Eq. (4.14), it follows that

$$q_{pV} = q_m \frac{R}{M} T = \frac{0.003 \text{ kg}}{3.156 \times 10^7 \text{ s}} \times \frac{8.314 \text{ J mol}^{-1} \text{ K}^{-1}}{0.102 \text{ kg mol}^{-1}} \times 293 \text{ K}$$
$$= 2.27 \times 10^{-6} \text{ Pa m}^3 \text{ s}^{-1} = 2.27 \times 10^{-5} \text{ mbar } \ell \text{ s}^{-1}. \tag{4.18}$$

When a vacuum vessel is evacuated by a vacuum pump, the gas volume flowing through the pump inlet per unit time (volume flow rate at the pump inlet) is the *pumping speed S* of the pump:

$$S = \dot{V}_{\text{inlet}} = q_{V \text{inlet}}, \quad [S] = \text{m}^3 \text{ s}^{-1} = 1000 \, \ell \, \text{s}^{-1} = 3600 \, \text{m}^3 \, \text{h}^{-1}.$$
 (4.19)

The pV flow at the inlet of the pump is referred to as throughput \dot{Q} of the pump:

$$\dot{Q} = q_{pV,\text{inlet}}, \quad [\dot{Q}] = \text{Pa m}^3 \text{ s}^{-1} = W = 10 \text{ mbar } \ell \text{ s}^{-1}.$$
 (4.20)

The previous two equations indicate that pumping speed and throughput of a vacuum pump are related according to (p is the pressure at the inlet)

$$\dot{Q} = q_{pV,\text{inlet}} = pS. \tag{4.21}$$

For many vacuum pumps, pumping speed S is (nearly) pressure independent. Then, throughput \hat{Q} is proportional to pressure as indicated in Eq. (4.21). Especially, throughput is low at low pressure. This is a comprehensible behavior because a volume element, at low pressure, contains less gas particles (and thus less mass). Figure 4.3 shows the pressure-independent pumping speed S (top) and the pressure-dependent throughput Q (bottom) of a pump as a function of inlet pressure.

Example 4.3

A mechanical displacement pump has a pumping speed $S = 360 \text{ m}^3 \text{ h}^{-1} =$ $100 \,\ell\,\mathrm{s}^{-1}$ and pumps air with a temperature of $20\,^{\circ}\mathrm{C}$ at an inlet pressure p = 10 mbar. This calculates to

$$pV$$
 flow: $q_{pV} = pS = 10 \text{ mbar} \times 100 \,\ell \text{ s}^{-1} = 1000 \text{ mbar} \,\ell \text{ s}^{-1}$. (4.22)

Throughput:
$$\dot{Q} = q_{pV} = 1000 \text{ mbar } \ell \text{ s}^{-1} = 100 \text{ Pa m}^3 \text{ s}^{-1} = 100 \text{ W}.$$
 (4.23)

Mass flow rate :
$$q_m = \frac{q_{pV}}{R_s T} = \frac{q_{pV} M}{RT}$$

= $\frac{100 \text{ W} \times 0.029 \text{ kg mol}^{-1}}{8.3 \text{ J mol}^{-1} \text{ K}^{-1} \times 293 \text{ K}} = 0.0012 \text{ kg s}^{-1}$. (4.24)

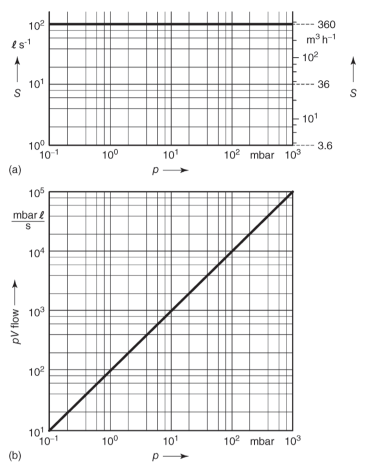


Figure 4.3 Pumping speed (a) and throughput (b) of a pump with a pressure-independent pumping speed of $S = 100 \, \ell \, s^{-1}$ versus inlet pressure.

4.1.3 Flow Resistance and Flow Conductance

The terms flow resistance and flow conductance are exemplified by considering a tube, connecting a vacuum chamber with a vacuum pump, as shown in Figure 4.4.

During evacuation, gas flows from the chamber, through the tube, and to the pump. This requires the pressure difference

$$\Delta p = p_{\text{chamber}} - p_{\text{pump inlet}} = p_{\text{c}} - p_{\text{in}}. \tag{4.25}$$

The behavior of a gas flow q is analogous to that of an electrical current, and the pressure of a gas corresponds to an electrical potential (see also

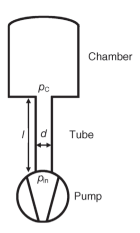


Figure 4.4 Vacuum system with pump line between vacuum chamber (pressure p_c) and pump (pressure p_{in} at the pump inlet).

Section 18.2.2). As an analogy to the ohmic resistance of an electrical component, the flow resistance R and conductance C of a tube are defined as

$$R = \frac{\text{pressure difference}}{\text{flux}} = \frac{\Delta p}{q},$$
 (4.26)

$$C := \frac{1}{R} = \frac{q}{\Delta p} \,. \tag{4.27}$$

Depending on the unit used for the flow, the quantities flow resistance and conductance are obtained with the corresponding units. Typically, pV flow is used, which leads to $[R] = s \text{ m}^{-3} \text{ or } s \ell^{-1} \text{ and } [C] = \text{m}^3 \text{ s}^{-1} \text{ or } \ell \text{ s}^{-1}$. If the particle flow rate is given, then [R] = Pa s and $[C] = Pa^{-1} s^{-1}$. Unless otherwise stated, pV flow will be used here.

For *electrical* conductors connected in series and parallel, the individual resistances and conductances add up to the total resistance R and total conductance C, respectively:

Series connection:

$$R = R_1 + R_2 + R_3 + \cdots$$
 and $\frac{1}{C} = \frac{1}{C_1} + \frac{1}{C_2} + \frac{1}{C_3} + \cdots$ (4.28)

Parallel connection:

$$C = C_1 + C_2 + C_3 + \cdots$$
 and $\frac{1}{R} = \frac{1}{R_1} + \frac{1}{R_2} + \frac{1}{R_3} + \cdots$ (4.29)

The validity of Eqs. (4.28) and (4.29) for gases is quite limited: an inlet flow develops at the inlet of a two-tube-series connection or in regions where a change occurs in the cross section. An additional flow resistance accompanies this type of flow. Thus, the overall assembly determines the flow resistance of a tube section. Flow resistance is higher if the tube is mounted

directly at the vessel, as opposed to being mounted in a position beyond an additional tube. Series connections of components will be discussed in detail for molecular flow (Sections 4.4.7 and 4.4.8). Equation (4.29) is only applicable for parallel connections if the tube inlets are separated far enough, so that the inlet flows do not interfere.

Practical calculations of multicomponent tube assemblies subdivide the systems into individual segments, according to geometrical dimensions and types of flow. Nonstationary gas flow can be treated analogously to an electrical current as well: the tube volume corresponds to the capacity of a capacitor in the same way as the inertia of the flowing gas mass corresponds to the inductance of a coil.

4.1.4

Effective Pumping Speed of a Vacuum Pump

We will consider a vacuum pump, attached to a vessel via a long tube (Figure 4.4), and will investigate the influence of the connecting tube on the pumping action.

For the (quasi-)stationary case, gas flow (given as, e.g., particle flow) is constant; that is, particle flows at the inlet and outlet of the pump are the same. The temperature of the gas can change due to the flow in the connecting tube (expansion), and due to the subsequent pumping process (compression). However, due to heat exchange with the environment, the change in temperature remains moderate. As an approximation, gas temperature is assumed to be constant, and thus, pV flow q_{pV} remains constant as well.

At the inlet of the connecting tube (chamber flange CF), the pressure is p_c , and at the outlet of the tube (inlet flange PF of the pump), the pressure is p_{in} . Assuming constant pV flow, it follows that

$$q_{pV} = p_{c} \dot{V}_{c} = p_{in} \dot{V}_{in}.$$
 (4.30)

The volume flow rate at the inlet flange of the pump is just the pumping speed of the pump; thus, $\dot{V}_{\rm in} = S$. The volume flow rate at the vessel flange is the pumping speed available for evacuating the vessel; thus, $\dot{V}_{\rm c} = S_{\rm eff}$ and is referred to as effective pumping speed (net pumping speed). Using these quantities, Eq. (4.30) can be rewritten as

$$S_{\text{eff}} = \frac{p_{\text{in}}}{p_c} S < S. \tag{4.31}$$

The effective pumping speed S_{eff} is lower than the pumping speed S because $p_{\rm c} > p_{\rm in}$, in order to maintain flow. However, due to the condition of continuity, the throughputs are the same at the inlet and the outlet of the tube.

By rewriting Eq. (4.27), the conductance C of the tube

$$C = \frac{q_{pV}}{p_{c} - p_{in}} = \frac{p_{in}S}{p_{c} - p_{in}} = \frac{p_{c}S_{eff}}{p_{c} - p_{in}}$$
(4.32)

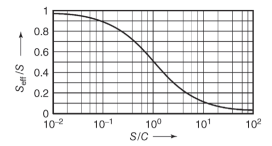


Figure 4.5 Dependence of pump efficiency $S_{\rm eff}/S$ on the ratio of pumping speed S of the pump and the conductance C of the tube.

yields the pressure ratio $p_c/p_{\rm in}$:

$$\frac{p_{\rm c}}{p_{\rm in}} = 1 + \frac{S}{C}.\tag{4.33}$$

The series connection of pump (pumping speed S) and tube (conductance C) determines the effective pumping speed S_{eff} available at the vessel:

$$\frac{1}{S_{\text{eff}}} = \frac{1}{S} + \frac{1}{C}$$
 and $S_{\text{eff}} = \frac{S}{1 + S/C}$. (4.34)

Figure 4.5 shows a plot of Eq. (4.34). Obviously, a pumping efficiency $S_{\rm eff}/S =$ 0.9 = 90% requires at least a 10-fold line conductance, compared with the pumping speed of the pump. If the conductance is just equal to the pumping speed, the effective pumping speed is only 50% of the pump's pumping speed. If the conductance is considerably below the pumping speed, the effective pumping speed is determined largely by the conductance and hardly by the pumping speed of the pump. Thus, any larger pump would not increase the effective pumping speed significantly. Consequentially, when installations are planned, tubes with maximum possible conductance should be selected (short tubes with large cross sections).

4.2 **Inviscid Viscous Flow and Gas Dynamics**

4.2.1

Conservation Laws

We will now consider viscous flow of gas through a tube. To simplify, flow shall be stationary (constant in time). In the stationary case, mass flow, that is, the mass flowing through a cross section of the tube, remains constant along the

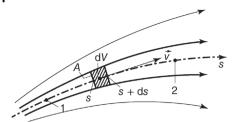


Figure 4.6 Flow field with flow filament.

line. Thus, the product of cross-sectional area A, density ρ , and velocity ν of gas flow remains constant as well:

$$q_m = \frac{\mathrm{d}m}{\mathrm{d}t} = \rho v A = \text{constant}$$
, conservation of mass. (4.35)

For a short tube, interactions of the gas with the walls are often negligible; that is, momentum exchange due to friction as well as energy transfer due to heat exchange does not occur. Assuming this, additional conservation laws apply. These are derived by considering a small mass element (face area A, length ds), and by investigating its motion along the direction s of flow (Figure 4.6).

If the static pressure changes along the path, the forces on the two face areas of the volume element are not balanced, and thus, a decelerating or accelerating force develops that affects the mass element. The inertial force of the mass element opposes this force. Thus, when frictional force is neglected,

$$dm\frac{dv}{dt} + dpA = 0. (4.36)$$

Since $dm = \rho A ds$, we obtain

$$\rho A s \frac{v}{t} + pA = 0. \tag{4.37}$$

Dividing by ρA and integrating across the path from location 1 to location 2 returns

$$\int_{1}^{2} \frac{\mathrm{d}\nu}{\mathrm{d}t} \, \mathrm{d}s + \int_{1}^{2} \frac{1}{\rho} \, \mathrm{d}p = 0. \tag{4.38}$$

Solving the integrals finally gives

$$\frac{1}{2}(v_2^2 - v_1^2) + \int_1^2 \frac{\mathrm{d}p}{\rho} = 0 \quad , \quad \text{conservation of momentum.}$$
 (4.39)

This is referred to as Bernoulli's equation for gases. It describes the relationship between static pressure and the velocity of flow along the path. If, for example, the pressure drops, then the gas accelerates and thus velocity increases.

If energy transfer with the wall is neglected, the total energy of a flowing mass element remains constant. This is made up of three components: pressure energy $(p \, dV)$, kinetic energy (motion with velocity of flow), and thermal energy (random particle movement):

$$\frac{pm}{\rho} + \frac{1}{2}mv^2 + c_V mT = \text{constant}. \tag{4.40}$$

Because mass is constant, dm can be canceled out, yielding

$$\frac{p}{\rho} + \frac{1}{2}v^2 + c_V T = \text{constant}$$
 , conservation of energy. (4.41)

The three conservation laws for mass, momentum, and energy do not yet define flow behavior clearly. Now, two cases are discussed:

Case 1: The cross-sectional area changes gradually along the line. The flowing gas adjusts to this change. Density, velocity of flow, pressure, and temperature alter gradually. Furthermore, no energy transfer between a mass element and its environment shall occur, giving an isentropic change of state. This case is discussed in Section 4.2.2.

Case 2: For supersonic flow, the flow can change abruptly at a certain area (shock surface). Density, velocity of flow, pressure, and temperature change abruptly. Because the cross-sectional area is unchanged at the shock surface, mass flow density $j_m = \rho v$ must be constant due to conservation of mass (Eq. (4.35)). This case is discussed in Sections 4.2.6 and 4.2.7.

4.2.2

Gradual Change of Cross-Sectional Area: Isentropic Change of State

We will examine the flow of a mass element in a tube, with negligible friction and energy transfer at the wall. The cross-sectional area A of the mass element $dm = \rho A ds$ is equal to the cross section of the tube. If the crosssectional area gradually decreases or increases along the direction of flow, the gas volume increases (expansion) or decreases (compression) accordingly (Figure 4.7).

A volume change in a gas is accompanied by a change in temperature. Without heat exchange between the mass element and its environment, its entropy will remain constant along the path. This behavior is referred to as isentropic or

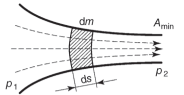


Figure 4.7 Flow through a tube with changing cross section (nozzle).

adiabatic. For this, thermodynamics derives the following relationships between pressure, volume, temperature, and density (Poisson's equations):

$$\frac{p_2}{p_1} = \left(\frac{V_2}{V_1}\right)^{-\kappa} = \left(\frac{T_2}{T_1}\right)^{\kappa/(\kappa - 1)} = \left(\frac{\rho_2}{\rho_1}\right)^{\kappa}.$$
 (4.42)

Here, κ is the isentropic exponent, that is, the ratio of the heat capacities at constant pressure and constant volume. Values for κ are as follows:

 $\kappa = \frac{5}{3} \approx 1.667$ for monatomic gases (noble gases, metal vapors such as Hg),

 $\kappa = \frac{7}{5} = 1.400$ for diatomic gases (e.g., nitrogen),

 $\kappa = \frac{4}{3} \approx 1.333$ for buckled three-atom molecules (e.g., water vapor),

 $\kappa \approx 1.1$ for more complicated molecules (e.g., oil vapors).

If we use the relation between density and pressure (Eq. (4.42)) for the isentropic change of state in Eq. (4.39), the integral can be solved and we obtain

$$v_2^2 - v_1^2 = \frac{p_1}{\rho_1} \cdot \frac{2\kappa}{\kappa - 1} \left[1 - \left(\frac{p_2}{p_1} \right)^{(\kappa - 1)/\kappa} \right] = \frac{\pi}{4} \overline{c}_1^2 \frac{\kappa}{\kappa - 1} \left[1 - \left(\frac{p_2}{p_1} \right)^{(\kappa - 1)/\kappa} \right]. \tag{4.43}$$

We will now assume that the gas flows from a vessel, where it is nearly motionless, into a tube. Thus, initially, velocity is negligible ($v_1 = 0$) and we find

$$\nu_{2} = \sqrt{\frac{p_{1}}{\rho_{1}} \cdot \frac{2\kappa}{\kappa - 1} \left[1 - \left(\frac{p_{2}}{p_{1}} \right)^{(\kappa - 1)/\kappa} \right]}$$

$$= \overline{c}_{1} \sqrt{\frac{\pi}{4} \cdot \frac{\kappa}{\kappa - 1} \left[1 - \left(\frac{p_{2}}{p_{1}} \right)^{(\kappa - 1)/\kappa} \right]}.$$
(4.44)

This relationship describes in which way the velocity increases, from initially zero to v_2 along the path, while at the same time the pressure decreases, from initially p_1 to p_2 (Figure 4.8). In addition, the temperature drops.

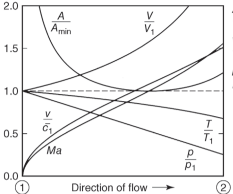


Figure 4.8 Changes in selected state quantities in gas-dynamic flow of a gas with an isentropic exponent $\kappa = 1.4$ from a vessel into a

Temperature

Velocity of flow

Area of cross section Volume of mass element

opening 1 (infinite cross-sectional area) to the

point 2. The gradient in cross-sectional area is

Pressure

Ma: Mach number

adjusted so that the pressure drops linearly nozzle, along the path from the vessel from p_1 at 1 to $p_2 = \frac{1}{4} p_1$ at 2.

In the extreme case, the pressure at the end of the tube is (nearly) zero; that is, $p_2 = 0$. Here, the velocity of flow v_2 reaches the maximum possible value:

$$\nu_{2,\text{max}} = \sqrt{\frac{p_1}{\rho_1} \cdot \frac{2\kappa}{\kappa - 1}} = \overline{c}_1 \sqrt{\frac{\pi}{4} \cdot \frac{\kappa}{\kappa - 1}}.$$
(4.45)

In a stationary flow, mass flow is constant along the path (Eq. (4.35)). After rewriting using Eq. (4.36), the mass flow density j_m , that is, the ratio of mass flow q_m and cross-sectional area A, now calculates to

$$j_m = \frac{q_m}{A} = -\frac{\mathrm{d}p_2}{\mathrm{d}v_2} = -\frac{1}{\mathrm{d}v_2/\mathrm{d}p_2}.$$
 (4.46)

 dv_2/dp_2 can be obtained by differentiating Eq. (4.44), which yields

$$j_m = p_1 \sqrt{2 \frac{\rho_1}{p_1}} \psi\left(\frac{p_2}{p_1}\right) = \frac{4}{\sqrt{\pi}} \cdot \frac{p_1}{\overline{c}_1} \psi\left(\frac{p_2}{p_1}\right). \tag{4.47}$$

This equation introduces the dimensionless *flow function* ψ , which is determined only by the ratio of the inlet and outlet pressures of the tube (Figure 4.9):

$$\psi\left(\frac{p_2}{p_1}\right) = \sqrt{\frac{\kappa}{\kappa - 1} \left[\left(\frac{p_2}{p_1}\right)^{2/\kappa} - \left(\frac{p_2}{p_1}\right)^{(1+\kappa)/\kappa} \right]}.$$
 (4.48)

According to Eqs. (4.47) and (4.48), mass flow density depends only on the inlet pressure p_1 , the ratio of inlet and outlet pressures, the particle velocity \overline{c}_1 (at inlet temperature), and the isentropic exponent κ of the gas.

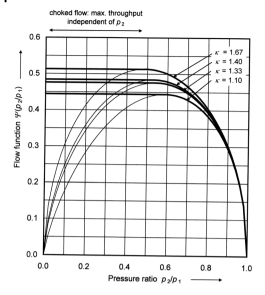


Figure 4.9 Flow function curves calculated for gas-dynamic flow by Eq. (4.48). The thinner drawn parts of the curves to the left do not describe real flow.

4.2.3 Critical Flow

As in the previous section, we will examine a gas, flowing out of a vessel through a tube with gradually narrowing cross section. In the vessel (at the inlet to the tube), the pressure is p_1 and the velocity is zero, $v_1 = 0$. At a distance (at the end of the tube), the pressure is p_2 .

If the outlet pressure p_2 varies, while all other conditions (inlet pressure, p_1 , in particular) remain unchanged, it follows from Eq. (4.47) that the mass flow of the gas is proportional to the flow function ψ (Eq. (4.48), Figure 4.9). Thus, the following behavior is observed for different ranges of outlet pressure:

- 1) If outlet pressure is equal to inlet pressure, gas flow is zero.
- 2) If the outlet pressure drops with respect to the inlet pressure $(p_2/p_1 = 1 \text{ to } 0.6)$, gas flow rises.
- 3) If the outlet pressure is lowered further, compared with inlet pressure $(p_2/p_1 \approx 0.6 \text{ to } 0.5, \text{ depending on the gas species})$, mass flow approaches a maximum.
- 4) If the outlet pressure is reduced even more, gas flow does not increase but, in fact, drops according to the flow function ψ .

The formation of a maximum in mass flow is easy to understand. In pressure range 2 $(p_2/p_1 = 1 \text{ to } 0.6)$, the velocity of flow, and thus mass flow, increases with decreasing outlet pressure p_2 . When the outlet pressure decreases, the volume of the gas increases. Therefore, if outlet pressure approaches the zero limit,

(4.53)

the volume expands to infinity. However, for energy reasons, the velocity of flow approaches a finite value. Thus, for low outlet pressure, mass flow approaches zero (pressure range 4). The maximum lies between ranges 2 and 4. Section 4.2.8 discusses pressure range 4 in more detail. Now, we will investigate the maximum in range 3.

Mass flow reaches its maximum at the so-called *critical point*. Correspondingly, values of quantities, at this point, are referred to as critical values, denoted by a superscripted asterisk (*). The critical point features the following values:

$$\frac{p^*}{p_1} = \left(\frac{2}{\kappa + 1}\right)^{\kappa/(\kappa - 1)}, \quad \text{pressure ratio}, \tag{4.49}$$

$$\psi\left(\frac{p^*}{p_1}\right) = \sqrt{\frac{\kappa}{2} \left(\frac{2}{\kappa+1}\right)^{(\kappa+1)/(\kappa-1)}}, \quad \text{flow function}, \tag{4.50}$$

$$\frac{T^*}{T_1} = \frac{2}{\kappa + 1}, \quad \text{temperature ratio}, \tag{4.51}$$

$$\frac{\rho^*}{\rho_1} = \left(\frac{2}{\kappa + 1}\right)^{1/(\kappa - 1)}, \quad \text{density ratio}, \tag{4.52}$$

$$\nu^* = \sqrt{\frac{p_1}{\rho_1} \cdot \frac{2\kappa}{\kappa + 1}} = \overline{c}_1 \sqrt{\frac{\pi}{4} \cdot \frac{\kappa}{\kappa + 1}} = \sqrt{R_{\rm s} T_1 \frac{2\kappa}{\kappa + 1}}, \quad \text{velocity of flow}.$$

From this, it follows for the mass flow density at the critical point:

$$j_{m}^{*} = \rho^{*} v^{*} = \rho_{1} \left(\frac{2}{\kappa+1}\right)^{1/(\kappa-1)} \sqrt{\frac{p_{1}}{\rho_{1}} \cdot \frac{2\kappa}{\kappa+1}}$$

$$= \frac{p_{1}}{\overline{c}_{1}} \left(\frac{2}{\kappa+1}\right)^{1/(\kappa-1)} \sqrt{\frac{16}{\pi} \cdot \frac{\kappa}{\kappa+1}}.$$
(4.54)

An additional, important value for gas dynamics is the speed of sound or acoustic velocity a. For an ideal gas with temperature-independent heat capacity,

$$a = \sqrt{\frac{p}{\rho}\kappa} = \overline{c}\sqrt{\frac{\pi}{8}\kappa} = \sqrt{\kappa R_{\rm s}T}$$
, speed of sound. (4.55)

Because the gas cools during expansion, the speed of sound decreases. If the temperature at the critical point is used in Eq. (4.55), the local speed of sound at the critical point is obtained:

$$a^* = \sqrt{\kappa R_s T^*} = \sqrt{\kappa R_s \frac{2T_1}{\kappa + 1}}.$$
(4.56)

Comparing velocity of flow (Eq. (4.53)) and speed of sound (Eq. (4.56)) shows that, at the critical point, the local velocity of flow is equal to the local speed of sound.

Introducing the Mach number Ma is useful for a simple description of flow velocity. Ma is defined as the ratio of local velocity of flow (Eq. (4.44)) and local speed of sound (Eq. (4.55)):

$$Ma := \frac{v}{a} = \sqrt{\frac{2}{\kappa - 1} \left[\left(\frac{p}{p_1} \right)^{(1 - \kappa)/\kappa} - 1 \right]}.$$
 (4.57)

Before reaching the critical point, the velocity of flow is less than the velocity of sound; thus, Ma < 1. This condition is referred to as subsonic flow. At the critical point, Ma = 1. Behind the critical point, Ma > 1, and the condition is termed supersonic flow (see Figure 4.8).

4.2.4

Choked Flow at Low Outlet Pressure

As an application example of gas-dynamic flow, we will consider flow through a nozzle (Figure 4.10) in which the cross-sectional surface A gradually narrows along the direction of flow, until reaching the minimum cross section A_{\min} . The inlet zone is designed short in order to reduce the friction between gas and walls, which increases with the length. In the following, friction in the inlet zone and in the attached tube is neglected as well.

At the inlet to the nozzle (marked with subscript 1), the cross-sectional area $A_1 \approx \infty$, bulk velocity $v_1 = 0$, and the state quantities of the gas are p_1 and T_1 . In the nozzle inlet, the cross-sectional area reduces to A_{\min} at its narrowest part (subscript 2). The gas has the state quantities p_2 , T_2 , and v_2 . The ratio p_2/p_1 of inlet and outlet pressures determines the type of flow. Three cases are differentiated:

Case 1: Outlet pressure p_2 is higher than the critical pressure p^* ; thus, $p^* < p_2 < p_1$. In this case, the gas accelerates along the path, so pressure, temperature, and density drop. The velocity of flow remains below the speed of sound and reaches the value as given by Eq. (4.44). According to Eq. (4.47), mass flow

$$q_{m} = A_{\min} j_{m} = A_{\min} p_{1} \sqrt{2 \frac{\rho_{1}}{p_{1}}} \psi \left(\frac{p_{2}}{p_{1}}\right) = A_{\min} \frac{4}{\sqrt{\pi}} \cdot \frac{p_{1}}{\overline{c}_{1}} \psi \left(\frac{p_{2}}{p_{1}}\right). \tag{4.58}$$

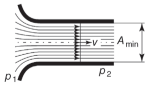


Figure 4.10 Flow through a nozzle.

Quantity	Monatomic gases: noble gases, metal vapors	Diatomic gases, for example, nitrogen	Buckled three-atom gases, for example, water vapor	Polyatomic gases, for example, oil vapor
κ	1.667	1.400	1.333	1.100
$(\kappa-1)/\kappa$	0.400	0.286	0.250	0.091
$(\kappa + 1)/\kappa$	1.600	1.714	1.750	1.909
p^*/p_1	0.487	0.528	0.540	0.585
T^*/T_1	0.750	0.833	0.857	0.952
$ ho^*/ ho_1$	0.650	0.634	0.630	0.614
v^*/\overline{c}_1	0.701	0.677	0.670	0.641
$\psi(p^*/p_1)$	0.513	0.484	0.476	0.444

Table 4.2 Values for calculating critical flow for selected gas species.

Thus, q_{pV} flow (with respect to the temperature T_1 at the outlet of the nozzle) is

$$q_{pV} = A_{\min} \sqrt{\frac{\pi}{4}} p_1 \overline{c}_1 \psi\left(\frac{p_2}{p_1}\right). \tag{4.59}$$

Case 2: Outlet pressure p_2 is equal to the critical pressure p^* ; that is, $p_2 = p^* \approx$ $\frac{1}{2} p_1$ (Eq. (4.49)). In this case, the gas accelerates so rapidly while flowing through the nozzle that it reaches the critical velocity, which is equal to the local speed of sound. Mass flow and mass flow density reach a maximum at the maximum of the flow function $\psi(p^*/p_1)$ (Eq. (4.50), Table 4.2). Mass flow and q_{nV} flow are calculated according to Eqs. (4.58) and (4.59), respectively. Here, the flow function reaches its maximum value (p^*/p_1) (Eq. (4.50)).

Case 3: Outlet pressure p_2 is lower than the critical pressure p^* ; that is, $p_2 < p^* < p_1$. In this case, the inlet pressure of the gas is identical to the inlet pressure in case 2. Again, behind the narrowing of the cross section to the minimum, the velocity increases to the highest possible value, that is, critical velocity. As in case 2, mass flow reaches a maximum, which is independent of the pressure at the outlet, but is determined only by the critical values in the narrowest cross section. When the velocity of the gas reaches its maximum, the condition is referred to as choked flow. Here, the flow behavior of the gas is no longer influenced by the flow behavior beyond the narrowing (e.g., outlet pressure) because in supersonic flow no effect can spread against the direction of flow.

The pressure drop from critical pressure p^* at the narrowing to outlet pressure p_2 occurs abruptly at the outlet of the nozzle (see Section 4.2.8).

4.2.5

Contraction of Flow into Aperture and Tube

Flow through an aperture or into a tube differs from flow through a nozzle in the sense that apertures and tubes feature an abrupt change of cross section instead

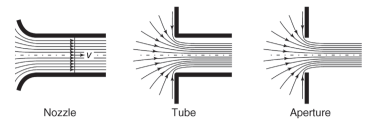


Figure 4.11 Intake flow into selected components with equal inlet cross sections.

of a smooth transition. In an abrupt change, the flow is not guided. The gas volume elements, approaching from various directions, tend to conserve their direction due to their inertia. Thus, the cross section contracts (*vena contracta*) (see Figure 4.11).

The flow passing through the components can be characterized by using the equations for nozzles presented in the previous section. However, the minimum area A_{\min} to use then is not the geometric opening A_0 but the area of the contracted flow. For a sharp-edged circular aperture,

$$A_{\min} = \begin{cases} 0.60A_0 & \text{if} \quad p_2 \approx p_1 \text{ (low drop in pressure)}, \\ 0.86A_0 & \text{if} \quad p_2 < p^* \text{ (choked flow)}. \end{cases} \tag{4.60}$$

4.2.6 Examples of Nozzle Flow

Often in practice, the gas to consider is air. Figure 4.12 shows the conductances of nozzles and apertures with circular cross sections for choked flow of air at 20 $^{\circ}$ C.

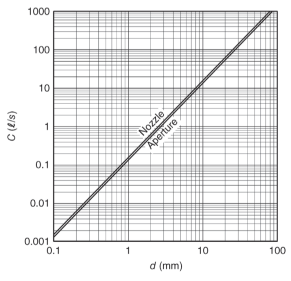


Figure 4.12 Flow conductances of nozzles and apertures with circular cross section (diameter d) for choked flow of air at 20 °C.

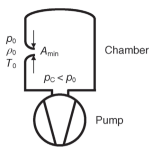


Figure 4.13 Flow into a chamber through a nozzle.

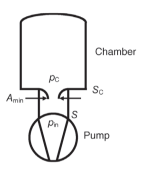


Figure 4.14 Evacuating a chamber through a nozzle.

We will now calculate two practical examples of gas-dynamic flow: first, venting a vacuum chamber through a nozzle (Figure 4.13), and second, evacuating a vacuum chamber through a nozzle (Figure 4.14).

In both cases, the ratio p_2/p_1 of the pressures at opposite ends of the nozzle plays a significant role. If this ratio is below the critical value (0.53 for air, see Table 4.2), then gas flow through the nozzle is determined only by the inlet pressure and is independent of the pressure at the outlet. However, if the pressure ratio p_2/p_1 is greater, the flow is determined by the pressure ratio: as the pressure ratio increases, flow drops and finally reaches zero for a pressure ratio of 1.

Example 4.4

Flow of air through a nozzle into an evacuated vessel (Figure 4.13).

This situation arises, for example, if the chamber is vented. The nozzle shall be circular with diameter d=2 mm. The ambient air has a pressure of 1000 mbar and a temperature of 20 °C. Using Table 4.2, we find

Critical pressure : $p^* = 0.528 \times 1000 \text{ mbar} = 528 \text{ mbar}.$

 $v^* = 0.677 \times 463 \text{ m s}^{-1} = 313 \text{ m s}^{-1}$. Critical velocity of flow:

Critical temperature : $T^* = 0.833 \times 293 \text{ K} = 244 \text{ K}$, that is, $-29 \,^{\circ}\text{C}$.

Initially, the pressure in the chamber is below critical pressure, leading to choked flow. Figure 4.12 is used to determine the conductance, $C = 0.62 \,\ell\,\mathrm{s}^{-1}$, and thus, for an inlet pressure of $p_1 = 1000 \,\mathrm{mbar}$, a pV flow of $p_1 C =$ 620 mbar ℓ s⁻¹ is obtained. In addition, the gas flow can be calculated. According to Table 4.2, the flow function ψ reaches the critical value 0.484. Thus, the airflow into the chamber amounts to

Critical mass flow (Eq. (4.58)) : $q_m^* = 0.74 \text{ g s}^{-1}$. Critical pV flow (Eq. (4.59)) : $q_{nV}^* = 623 \text{ mbar } \ell \text{ s.}^{-1}$

A problem that can occur in practice during venting through a nozzle is that ice may cover the nozzle due to the humidity of the air. Air, when traveling through the nozzle, cools to -29 °C, and then, the saturation vapor pressure of water/ice is only 0.47 mbar. Calculating back the isentropic expansion, this value corresponds to a water vapor pressure for the air inflow of 0.87 mbar at 20 °C. If the partial pressure of the water vapor is above this value, the vapor condenses to ice when cooling in the nozzle. After all, ambient air, typically, has a relative humidity of 50%, and thus a partial pressure of water vapor of 11.7 mbar.

After awhile, the pressure in the chamber is above the critical pressure. We will assume the pressure in the chamber to be $p_c = 800$ mbar. The pressure ratio at the nozzle then amounts to $p_2/p_1 = 0.8$. Considering Figure 4.9, the flow function reads $\psi \approx 0.40$. Solving according to Eq. (4.48) yields the accurate value of $\psi = 0.396$. As the maximum value was $\psi = 0.484$, mass flow is now 82% of the maximum mass flow $(q_m^* = 0.74 \text{ g s}^{-1})$; thus, $q_m = 0.61 \text{ g s}^{-1}$.

Example 4.5

Nozzle in the inlet line of a pump (Figure 4.14).

A pump (pumping speed S) is used to evacuate a chamber (Figure 4.14) through a tube with a large cross section. A nozzle placed inside the tube acts as flow resistance and affects the flow rate as well as the pressure ratio. At chamber pressure p_c and pump inlet pressure p_{in} , Eq. (4.31) gives the effective pumping speed S_c available at the chamber:

$$S_{c} = \frac{p_{in}}{p_{c}} S. \tag{4.61}$$

For the following investigations, the critical pumping speed *S** is introduced. This is the maximum pumping speed, obtained when the pump is powerful enough to keep the pressure p_{in} sufficiently low to produce choked flow in the nozzle with critical gas flow q_{nV}^* through the nozzle. For this situation,

$$\frac{q_{pV}^*(\text{nozzle})}{q_{pV}(\text{pump inlet})} = \frac{p_c S^*}{p_{\text{in}} S},$$
(4.62)

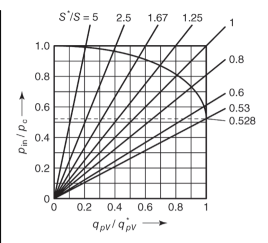


Figure 4.15 Nomogram for determining the gas flow through a nozzle for inviscid flow of a diatomic gas ($\kappa = 1.4$). The straight lines represent Eq. (4.63). The bended curve represents Eq. (4.59), that is, $q_{pV}(p_2/p_1)$ normalized to $q_{pV}(p_2/p^*)$.

and thus,

$$\frac{S^*}{S} = \frac{p_{\text{in}}}{p_c} / \frac{q_{pV}(p_c/p_{\text{in}})}{q_{pV}^*}.$$
 (4.63)

The functional dependence between the gas flow through the nozzle and the pressure ratio is complicated (Eq. (4.59)), so a nomogram is useful for practical calculations (Figure 4.15).

For the following calculation, we need to know the critical pumping speed

$$S^* = \frac{q_{pV}^*}{p_c} = A_{\min} \sqrt{\frac{\pi}{4}} \overline{c}_1 \psi \left(\frac{p^*}{p_1} \right). \tag{4.64}$$

Let us give an example. If the diameter of the nozzle $d_{min} = 1$ cm, the critical pumping speed for air at 20 °C calculates to

$$S^* = \frac{\pi}{4} (0.01 \text{ m})^2 \sqrt{\frac{\pi}{4}} 463 \text{ m s}^{-1} \times 0.484 = 15.6 \,\ell \text{ s}^{-1}. \tag{4.65}$$

If the pump has a pumping speed $S = 72 \text{ m}^3 \text{ h}^{-1} = 20 \text{ } \ell \text{ s}^{-1}$, this is just above the critical pumping speed S^* of the nozzle because $S^*/S = 0.78$. The straight line through the origin of the nomogram (Figure 4.15) corresponding to this value crosses the curve at an abscissa value $p_{\rm in}/p_{\rm c}=0.72$. Following Eq. (4.61), an effective pumping speed $S_c = 0.72 \times 20 \,\ell\, s^{-1} = 14.4 \,\ell\, s^{-1}$ is available at the chamber, that is, 72% of the pumping speed of the pump.

If a pump is used, featuring half of the pumping speed assumed above, that is, $S = 36 \,\mathrm{m}^3 \,\mathrm{h}^{-1} = 10 \,\ell \,\mathrm{s}^{-1}$, the corresponding quantities calculate to $S^*/S = 1.56$ and $p_{\rm in}/p_{\rm c}=0.9.$ At the chamber, the available effective pumping speed $S_c = 0.9 \times 10 \,\ell \,\mathrm{s}^{-1} = 9 \,\ell \,\mathrm{s}^{-1}$, that is, 90% of the pumping speed of the pump.

If the pump has twice the pumping speed of the value assumed initially, that is, $S = 144 \,\mathrm{m}^3 \,\mathrm{h}^{-1} = 40 \,\ell \,\mathrm{s}^{-1}$, then the ratio of critical pumping speed of the nozzle and pumping speed of the pump is $S^*/S = 0.39$. Thus, the pressure ratio $p_{\rm in}/p_{\rm c} < 0.523$, and accordingly, flow is choked. The gas flow reaches its maximum value, determined by the nozzle. The pumping speed available at the chamber is equal to the critical pumping speed $S^* = 15.6 \,\ell\,\mathrm{s}^{-1}$, and even a more powerful pump cannot increase it.

It should be mentioned that the above calculations apply only to viscous flow, that is, to a Knudsen number $Kn = \overline{I}/d$ below 10^{-2} . In the example, this condition prevails down to a chamber pressure p of approximately 0.65 mbar.

4.2.7

Straight and Oblique Compression Shocks

We will now consider stationary flow through a tube. The velocity of flow shall be greater than or equal to the speed of sound. Supersonic velocities can occur, for example, in the outlet of a Laval nozzle (Section 4.2.8). Then, changes at a certain point have no backward effect on successive gas flow.

In this type of flow, it may happen that flow suddenly changes, for example, due to a minor disturbance: at the shock front, the state variables such as velocity of flow, density, temperature, and pressure change abruptly. Thus, the supersonic flow changes to subsonic flow. Density then rises abruptly, and the effect is termed a compression shock, or simply, shock.

First, we will investigate the straight or perpendicular shock. In order to calculate the state variables for the shock, we will again use the conservation laws for mass, momentum, and energy introduced in Section 4.2.1 (Eqs. (4.35), (4.39), and (4.41)). In the shock, the cross-sectional area A in Eq. (4.35) (conservation of mass) remains unchanged; however, the quantities ν , p, and ρ in Eq. (4.59) (conservation of momentum) change abruptly. Thus, the conservation laws can be formulated as

$$\rho v = \text{constant}, \quad \text{conservation of mass},$$
 (4.66a)

$$p + \rho v^2 = \text{constant}, \quad \text{conservation of momentum},$$
 (4.66b)

$$\frac{p}{\rho} + \frac{1}{2}v^2 + c_V T = \text{constant}, \quad \text{conservation of energy.}$$
 (4.66c)

Calculating the values of state variables behind the shock requires assumptions regarding temperature behavior T. For the isentropic flow in Section 4.2.2, the temperature was obtained using *Poisson's* equations (Eq. (4.42)). For the shock considered here, the term $c_V T$ is calculated from the characteristics of an ideal gas:

$$c_V T = \frac{1}{\kappa - 1} R_{\rm s} T = \frac{1}{\kappa - 1} \cdot \frac{p}{\rho}. \tag{4.67}$$

Thus, by putting in Eq. (4.67), the temperature is eliminated from the conservation laws. Furthermore, when rewriting the three conservation laws (Eqs. (4.66a)-(4.66c)) for the gas state in front of and behind the shock, the flow velocities v_1 (in front of the shock) and v_2 (behind the shock) can be eliminated as well. The result describes the relationship between the ratio p_2/p_1 of the pressures in front of and behind the shock on the one hand and the density ratio ρ_2/ρ_1 on the other hand. This relation is referred to as the *Hugoniot equation*:

$$\frac{p_2}{p_1} = \frac{(\kappa+1)(\rho_2/\rho_1) - (\kappa-1)}{(\kappa+1) - (\kappa-1)(\rho_2/\rho_1)}.$$
(4.68)

Figure 4.16 shows a plot of the Hugoniot curve (Eq. (4.68)) for an abrupt compression shock, and, for comparison, includes a corresponding plot for a gradual isentropic change of state, Poisson's equation (Eq. (4.42)).

The *Hugoniot* curve has a number of interesting characteristics. Density ρ_2 behind the shock can reach a maximum value $(\kappa + 1)/(\kappa - 1)$ -fold of ρ_1 in front of the shock as, for this, pressure would approach infinity. However, Poisson's equation does not show such a limit. At $\rho_2/\rho_1 = 1$, not only both curves have the same ordinate value, but their first and second derivatives also correspond. Thus, for low compression in the shock, that is, $\rho_2/\rho_1 \approx 1$, *Poisson's* curve represents an acceptable approximation for the Hugoniot curve. The deviation of the Hugoniot curve from Poisson's isentropic curve indicates a change in entropy caused by the shock.

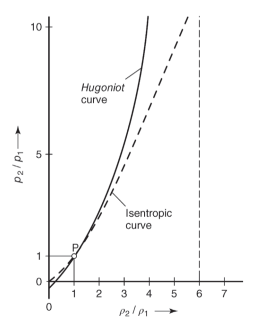


Figure 4.16 Hugoniot curve, Eq. (4.68), and *Poisson* curve (isentropic), Eq. 4.42, for $\kappa = 1.4$, that is, $(\kappa + 1)/(\kappa - 1) = 6$.

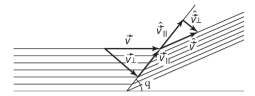


Figure 4.17 Oblique compression shock. The velocities of flow in front of and behind the shock front are resolved into components perpendicular and parallel to the shock surface.

Due to the second law of thermodynamics, the entropy of the gas cannot drop while it passes through the shock surface. Thus, the only branch of the Hugoniot curve with physical significance lies above *Poisson's* curve (Figure 4.16). This branch describes a compression shock (increase of mass density). A diluting shock (drop of mass density) is impossible. Compression of gas flow can occur in a discontinuous shock, with a rise in entropy following the *Hugoniot* equation, as well as, gradually, with conservation of entropy (isentropic) according to Poisson's equation. In contrast, dilution always happens gradually and isentropically.

In technical compressors, for example, vapor-iet pumps or diffusion pumps, compression occurs in one or more compression shocks until the medium is at rest ($\nu = 0$). Behind the shock, the pressure rises to static pressure p_0 , the maximum possible compression pressure of the pump. The pressure ratio p_2/p_1 = p_0/p_1 increases until it reaches the *static pressure ratio*.

We will now investigate the *oblique compression shock*. The previous considerations for the straight shock can be applied to the oblique shock. This requires separating the vector components of the flow velocity, in front of and behind the shock, into the components parallel and perpendicular to the shock surface (see Figure 4.17).

The equations derived above may be used if we substitute

$$\nu$$
 by ν_{\perp} ,
$$Ma \text{ by } Ma \sin \theta,$$

$$(4.69)$$

in which θ is the angle between the direction of flow and the surface of the compression shock.

As Ma > 1 is one necessary condition for a straight shock, and Ma is replaced by $Ma \sin \theta$ in the oblique shock, the corresponding condition for an oblique shock reads $Ma > Ma \sin \theta > 1$. Thus, the angle of the shock surface cannot be lower than the *Mach* angle α , defined by $\sin \alpha = 1/Ma$. The angle θ represents an additional geometric controlling factor for the oblique shock, compared with the straight shock. For small angles, the change in the values of state quantities is low (weak shocks).

4.2.8

Laval Nozzle and Effluent Flow against Counterpressure

A Laval nozzle is a nozzle with an inlet zone of decreasing cross section and an outlet zone of increasing cross section (Figure 4.18). The inlet zone is designed short in

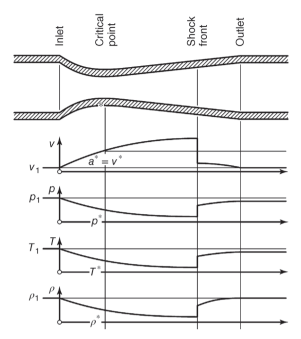


Figure 4.18 Laval nozzle. Top: cross section. Bottom: state quantities along the path (velocity of flow, pressure, temperature, density). (After Ref. [1].)

order to minimize friction between the walls and the gas. The concept of the outlet zone affords a compromise: the outlet zone should be short to avoid frictional loss. This can be obtained by using a wide aperture angle. If, however, the aperture angle is too wide, the flow, depending on its velocity, might not follow the rapidly increasing cross section, and thus, undesired stall from the walls develops.

The cross section gradually decreases in the inlet zone of a Laval nozzle, down to the narrowing (throat), and the velocity of flow increases as pressure and temperature fall. In contrast, the outlet zone features different types of flow. If only the amount of gas passing through a Laval nozzle is of interest, two cases are differentiated:

Case 1: If the exit pressure p_2 is just barely below inlet pressure p_1 , the flow remains subcritical; that is, across the complete nozzle, the velocity of flow is below the critical value. After passing the nozzle throat, the gas decelerates to the outlet while pressure and temperature rise. Without any frictional loss, the gas condition at the outlet is equal to the inlet state. Depending on flow conditions and the geometry of the nozzle, real flow can be very close to the frictionless ideal case. In such a case, flow can be calculated very well. Thus, Laval nozzles are used frequently for flow monitoring. Here, the static pressures at the inlet and throat are measured and, for known cross-sectional areas, the amount of the passing substance is calculated. Deviations from the ideal case have been investigated thoroughly in experiments and theory, and are tabulated in DIN 1952 and ISO 5167-1.

Case 2: If the exit pressure p_2 is below the critical pressure p^* , the flow accelerates to the speed of sound in the throat, and becomes choked. The gas flow develops according to Eqs. (4.58) and (4.59) if the cross-sectional area A_{min} is set to the area of the nozzle's throat, and the critical value is used for the flow function ψ (Table 4.2). Here also, depending on flow conditions, real flow can be very close to the frictionless ideal case. ISO 9300 lists the deviations from the ideal case. Thus, for known inlet pressure and nozzle geometry, gas flow can be measured precisely.

The flow in the outlet zone of a Laval nozzle depends on nozzle geometry and exit pressure.

Figure 4.18 shows an interesting flow condition: the gas passing through the nozzle initially accelerates to the speed of sound at the nozzle throat. Thus, choked flow develops and the gas flow is independent of the exit pressure. In the outlet of the nozzle, the gas continues to accelerate to supersonic velocity while the pressure drops further. As, however, the pressure against which the gas escapes at the exit is not sufficiently small, a compression shock with abrupt pressure rise develops still inside the nozzle.

A nice application example of Laval nozzles are jet pumps (Section 9.3). In these pumps, a working fluid (or pump fluid), which can be a gas or vapor, enters a volume referred to as mixing chamber via a Laval nozzle. The mixing chamber contains the gas to be pumped at a certain pressure that can vary considerably according to the operating state. Depending on working-fluid pressure and mixing-chamber pressure, different operating states of the nozzle are obtained.

Operating state 1 (Figure 4.19, left-hand image): The pressure p_{out} at the exit of the nozzle is just barely below the inlet pressure. In the complete nozzle, the pressure remains above critical pressure p^* . In this case, the flow accelerates in

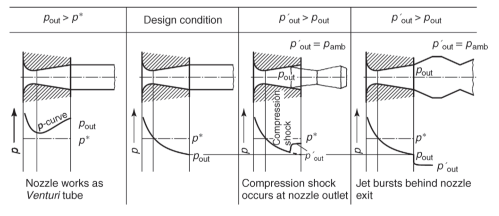


Figure 4.19 Different operating conditions of a Laval nozzle under counterpressure. (After Ref. [2].)

the narrowing part of the nozzle and decelerates in the widening part. The velocity of flow remains below the speed of sound at all times.

Operating state 2 (Figure 4.19, second image): The pressure p_{out} at the outlet of the nozzle is just equal to the designed exit pressure. In this case, the flow accelerates in the narrowing part, and reaches the critical value in the throat. Then, the flow accelerates further in the widening part where the cross-sectional area increases. Thus, the flow accelerates along the complete path through the nozzle, and thus, the pressure drops continuously. The resulting exit pressure is determined by the cross-sectional area of the outlet, which is designed to produce just the pressure in the exit volume. This is the optimal case.

Operating state 3 (Figure 4.19, third image): The pressure $p'_{\rm out}$ in the exit volume is below critical pressure p^* but above the designed exit pressure p_{out} of the nozzle. At the throat, the velocity of flow is equal to the speed of sound. Thus, the flow is choked, and is independent of the exit pressure. In the outlet of the nozzle, the gas accelerates further and reaches supersonic flow, and thus, the pressure continues to drop. As, however, the counterpressure at the outlet is significant, a straight compression shock develops inside the Laval nozzle. Additional straight and oblique shocks can arise in the nozzle outlet and in the emerging jet. Possibly, stalling flow may occur.

Operating state 4 (Figure 4.19, fourth image): The pressure p'_{out} in the outlet volume is below critical pressure p^* and below the designed outlet pressure p_{out} of the nozzle. After the jet leaves the nozzle, oblique shocks develop in the jet, which expand the jet more than the extension of the nozzle geometry would suggest. Further along the jet, compression shocks and diluting waves succeed each other. As a result, oscillation can develop in the jet.

4.2.9

Flow Around a Corner (Prandtl-Meyer Flow)

The flow filament theory usually does not explain the free supersonic jet behind a nozzle. In vacuum technology, however, calculation of supersonic-jet behavior near the outlet of the nozzle is often sufficient. For this, the *Prandtl–Meyer* procedure is used. We will calculate a parallel flow, restricted by a wall at one side. Furthermore, at the end of the wall (corner), the flow enters a region of lower counterpressure p, compared with the pressure within the parallel flow (p_1) (Figure 4.20). Thus, the flow is deflected from its original direction and the directions of the streamlines (velocity) now are at an angle θ to the parallel flow.

The two-dimensional (plane) approach to the problem reveals that the absolute value and direction of velocity ν , as well as the state quantities pressure, density, and temperature, are constant along each polar vector starting from the corner. Furthermore, the polar vectors obviously are *Mach* lines; that is, they cross the flow filaments (in the direction of the velocity) at the *Mach* angle α , with $Ma = 1/\sin \alpha$. The angle ϑ is the difference

$$\vartheta = \lambda - \lambda_1. \tag{4.70}$$

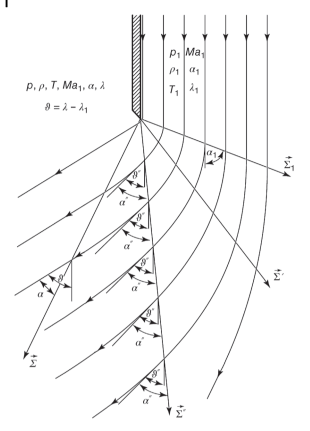


Figure 4.20 Supersonic flow around the corner of a wall. The polar vectors Σ are *Mach* lines on which the state quantities p, ρ , T, and the deflection angle $\vartheta = \lambda - \lambda_1$ are constant. The angle α between the *Mach* lines and the stream lines follows $Ma = 1/\sin \alpha$.

Here, λ and λ_1 denote a flow parameter at the considered point of the flow field at a location past and in front of the deflection, respectively. The flow parameter is defined by

$$\lambda = \frac{1}{2} \arccos\left(\kappa - \frac{\kappa - 1}{1 - (p/p_0)^{(\kappa - 1)/\kappa}}\right) + \frac{1}{2} \sqrt{\frac{\kappa + 1}{\kappa - 1}} \arccos\left(\kappa - (\kappa + 1) \left[1 - \left(\frac{p}{p_0}\right)^{(\kappa - 1)/\kappa}\right]\right) - \frac{\pi}{2}.$$
(4.71)

If the parallel flow in front of the bend is just critical, that is, Ma = 1, so that the pressure ratio p/p_0 complies with the relation in Eq. (4.49), both arccos terms in Eq. (4.71) are just zero, and it follows that

$$\lambda_1 = \lambda^* = 0. \tag{4.72}$$

In this case, the deflection angle θ is equal to the flow parameter λ that, by this, can be perceived. If the gas showing critical flow in front of the bend expands to the pressure p = 0 behind the bend, it follows from Eq. (4.71) that the maximum deflection angle

$$\vartheta_{\text{max}} = \frac{\pi}{2} \left[\sqrt{\frac{\kappa + 1}{\kappa - 1}} - 1 \right]. \tag{4.73}$$

For gases with $\kappa < 1.25$, this angle is greater than 180°. As this is physically impossible, the gas can only expand to a finite value of the pressure p_{\min} . Equation (4.71) may be used to calculate p_{\min} if $\lambda = \pi$. Similar concepts apply to supercritical flow.

Example 4.6

This example investigates the aperture angle of a free supersonic jet, emerging from a nozzle to a region of low counterpressure (fourth operating condition of Laval nozzle in the previous section). The jet shall be composed of oil vapor with $\kappa = 1.1$ and a relative particle mass of 435. At the nozzle exit, the expansion ratio $p_1/p_0 = 0.03$, and the counterpressure p behind the nozzle corresponds to an expansion ratio of $p/p_0 = 0.001$. Equation (4.71) yields $\lambda_1 = 65^{\circ}$ and $\lambda = 113^{\circ}$, so that the deflection angle, according to Eq. (4.70), calculates to

$$\vartheta = \lambda - \lambda_1 = 113^{\circ} - 65^{\circ} = 48^{\circ}. \tag{4.74}$$

For the state variables of the gas, it follows:

• At nozzle exit $(p_1/p_0 = 0.33)$:

$$Ma_1=2.8, \quad Ma_1^*=2.4, \quad \frac{T_1}{T_0}=0.72, \quad \frac{\rho_1}{\rho_0}=0.04.$$
• Behind nozzle ($p/p_0=0.001$):

$$Ma = 4.2$$
, $Ma^* = 3.2$, $\frac{T_1}{T_0} = 0.53$, $\frac{\rho_1}{\rho_0} = 0.002$.

The quantities p_0 , ρ_0 , and T_0 are not independent, and thus cannot be chosen arbitrarily. Their relationship is given by the equation of state (Eq. (4.11)) and by the vapor pressure diagram. For the oil used ($\kappa = 1.1$), for example, the vapor pressure curve suggests a boiling temperature of $T_0 = 520 \,\mathrm{K}$ for a boiling pressure of p = 1333 Pa. Thus, the thermal velocity at the inlet (Eq. (4.12)) is

$$\overline{c}_1 = \sqrt{\frac{8}{\pi} \cdot \frac{8.314 \,\mathrm{J \, mol^{-1} \, K^{-1} \times 520 \, K}}{0.435 \,\mathrm{kg \, mol^{-1}}}} = 159 \,\mathrm{m \, s^{-1}}.$$
(4.75)

The critical velocity (Eq. (4.53)) is

$$\nu^* = \sqrt{\frac{8.314 \,\mathrm{J \, mol}^{-1} \,\mathrm{K}^{-1} \times 520 \,\mathrm{K}}{0.435 \,\mathrm{kg \, mol}^{-1}} \cdot \frac{2 \times 1.1}{1.1 + 1}} = 102 \,\mathrm{m \, s}^{-1}, \tag{4.76}$$

and the critical flow density (Eq. (4.54)) is

$$j_m^* = \frac{1333 \,\text{Pa}}{159 \,\text{m s}^{-1}} \left(\frac{2}{1.1+1}\right)^{1/(1.1-1)} \sqrt{\frac{16}{\pi} \cdot \frac{1.1}{1.1+1}} = 8.4 \,\text{kg m}^{-2} \,\text{s}^{-1}. \tag{4.77}$$

4.3 Frictional–Viscous Flow through a Tube

4.3.1

Laminar and Turbulent Flows through a Tube

This section deals with viscous flow in long tubes and covers its fundamental phenomena. Figure 4.21 shows the various phenomena of tube flow. The phenomena are investigated in the order in which they appear along the course of the gas flow.

The inflow of the fluid into the tube with flow contraction has already been treated in Section 4.2. In the subsequent region of the tube, boundary layers near the wall are slowed down by friction. In nozzles and short tubes, the boundary layer is so thin that the total flow is hardly influenced by friction.

The thickness of the boundary layer increases gradually along the tube and eventually reaches considerable size. Thus, it affects the flow increasingly and, ultimately, determines the flow considerably. Due to wall friction, a velocity distribution develops across the cross section of the tube. This velocity profile changes gradually within the intake range until it eventually reaches its final shape (Figure 4.22).

Wall friction decelerates the flowing fluid, and thus, momentum is no longer a conserved quantity. Furthermore, the moving fluid's wall interaction leads to an exchange of thermal energy, which results in (nearly) isothermal flow. Consequentially, energy also is no longer a conserved quantity.

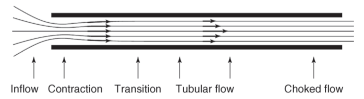


Figure 4.21 Gas flow through a long tube.

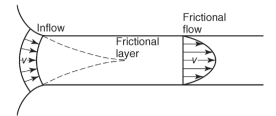


Figure 4.22 Flow pattern developing across the cross section of a tube behind inflow into the

The following investigations consider flow through a tube. We will assume a long tube and will examine a section of the tube, far beyond the inlet of the tube, where the final velocity profile has established.

Equations given here are true for stationary isothermal flow. In the case of pulsating flow, flow resistance is usually higher (compare electrical analogy: additional resistance caused by impedances).

As described in Section 4.1, flow in long tubes can be laminar as well as turbulent. The Reynolds number Re is the criterion used to differentiate between the two types of flows:

$$Re = \frac{\rho vd}{n},\tag{4.4}$$

where ρ is the density of gas flow, ν is the velocity of gas flow (mean value across tube's cross section), d is the cross section (for circular tubes, diameter), and η is the dynamic viscosity of the gas.

For a tube with circular cross section (*circular tube*),

$$Re \begin{cases} < 2300 \text{ characterizes laminar flow,} \\ > 4000 \text{ characterizes turbulent flow.} \end{cases}$$
 (4.5)

The Reynolds number is not appropriate for precisely characterizing the accurate transition between the two types of flows. This is because the flow behavior is determined by the condition of flow at the inlet into the considered stretch, and is very sensitive to wall roughness.

Usually, flow velocity ν is unknown but it can be calculated from pV flow q_{pV} :

$$\nu = \frac{1}{A} \cdot \frac{\mathrm{d}V}{\mathrm{d}t} = \frac{4}{\pi d^2} \cdot \frac{q_{pV}}{p}.\tag{4.78}$$

Putting in the velocity of flow into the term for the *Reynolds* number (Eq. (4.4)) yields

$$Re = \frac{4}{\pi} \cdot \frac{q_m}{\eta d} = \frac{4}{\pi} \cdot \frac{\rho q_{pV}}{p \eta d} = \frac{32}{\pi^2} \cdot \frac{q_{pV}}{\eta \overline{c}^2 d}.$$
 (4.79)

Thus, the condition given by Eq. (4.5) for the type of flow can be rewritten as

$$\frac{q_{pV}}{d} \begin{cases} < 709 \, \eta \overline{c}^2 \text{ for laminar flow,} \\ > 1234 \, \eta \overline{c}^2 \text{ for turbulent flow.} \end{cases}$$
 (4.80)

In laminar flow, the individual layers slide along one another. The frictional force is proportional to the velocity and viscosity of the flowing liquid (Newtonian approach). As derived elementally in textbooks on fluid mechanics, this type of arrangement yields the following Hagen-Poiseuille equation for volumetric flow:

$$\frac{\mathrm{d}V}{\mathrm{d}t} = \frac{\pi}{128} \frac{1}{n} \frac{\mathrm{d}p}{\mathrm{d}l} d^4. \tag{4.81}$$

This equation applies to fluid elements with constant volume. In gas flow, the pressure drops along the tube, and thus, the volume increases accordingly. If the temperature remains constant along the flow path due to heat exchange with the walls of the tube, pV flow q_{nV} , however, is constant. In this case, Eq. (4.81) that applies to a short piece of tube can be integrated over the length l of the tube, from point 1 to point 2. Thus, pV flow

$$q_{pV} = \frac{\pi}{256} \frac{1}{\eta} \frac{d^4}{l} (p_1^2 - p_2^2)$$
, circular tube, laminar flow, (4.82)

and conductance

$$C = \frac{q_{pV}}{p_1 - p_2} = \frac{\pi}{256} \frac{1}{\eta} \frac{d^4}{l} (p_1 + p_2)$$
, circular tube, laminar flow. (4.83)

For turbulent flow, a semi-empirical method is used to calculate throughput and conductance. As above, the temperature is assumed to be constant. The following semi-empirical formulation describes the pressure loss dp in the tube per length dl:

$$\frac{\mathrm{d}p}{\mathrm{d}l} = \lambda \frac{1}{2d} \rho v^2. \tag{4.84}$$

Here, λ is the so-called dimensionless friction coefficient of the tube and ρ is the gas density. The mean longitudinal flow velocity ν is simply the ratio of volume flow and cross-sectional area (Eq. (4.78)).

The Blasius equation is commonly used to calculate the coefficient of friction λ in a circular tube with smooth inner surface:

$$\lambda = \frac{0.3164}{\sqrt[4]{Re}} = 0.3164 \sqrt[4]{\frac{\eta}{\rho v d}}.$$
(4.85)

Equations (4.84) and (4.85) yield the pressure loss per distance as a function of flow velocity. The velocity of flow can be substituted by the flow (Eq. (4.78)), and density may be eliminated using pressure (Eq. (3.43)):

$$\frac{\mathrm{d}p}{\mathrm{d}l} = \frac{0.3164}{2} \frac{1}{p} \sqrt[4]{\frac{8^3 \times 4^7}{\pi^{10}}} \frac{1}{d^{19}} \frac{\eta}{\overline{c}^6} q_{pV}^7. \tag{4.86}$$

Integrating the pressure loss over the length of the tube and rearranging yields the pV flow through a circular tube with diameter d and length l:

$$q_{pV} = 1.015 d^{19/7} \left(\frac{\overline{c}^6}{\eta}\right)^{1/7} \left(\frac{p_1^2 - p_2^2}{l}\right)^{4/7}$$
 for turbulent flow, (4.87)

and conductance

$$C = 1.015d^{19/7} \left(\frac{\overline{c}^6}{\eta}\right)^{1/7} \left(\frac{p_1 + p_2}{l}\right)^{4/7} (p_1 - p_2)^{-3/7}$$
 for turbulent flow. (4.88)

A tube is considered smooth if wall roughness is less than 1% of the diameter, if the Reynolds number is not too high. In rough tubes and tubes with sharp bends, gas flow and conductance are below the values calculated with the preceding equations.

Careful attention should be paid to the outlet pressure p_2 of the tube. In laminar as well as turbulent flow, the pressure drops along the tube due to wall friction. Thus, at constant temperature, the volume, volume flow, and velocity of flow increase. If the outlet pressure p_2 of the tube is sufficiently low, the velocity of flow approaches a maximum, namely, the speed of sound a. At this critical pressure p^* , flow becomes choked and the throughput attains its maximum value q_{nV}^* . The pressure at the exit of the tube cannot drop below the critical pressure. If, however, the vessel pressure is reduced further, the emanating gas suffers an abrupt pressure drop when escaping from the tube into the vessel. Reducing vessel pressure below the critical pressure does not increase gas flow. We will now calculate the critical outlet pressure p^* at which chocked flow

The maximum pV flow at a cross section A (a = speed of sound, see Eq. (4.55)) is

$$q_{pV}^* = p^* A \, a. \tag{4.89}$$

Equation (4.82) specifies the pV flow through a circular tube for laminar flow, and Eq. (4.87) for turbulent flow. For the approximation $p_1^2 - p_2^2 \approx p_1^2$, the critical pressure is obtained by putting in the pV flow into Eq. (4.89) and rearranging:

$$p^* = \frac{d^2 p_1^2}{64 \, \eta l a}$$
, critical pressure for laminar flow, (4.90)

$$p^* = 1.92 \frac{1}{ad} \left(\frac{\overline{c}^6}{\eta}\right)^{1/7} \left(\frac{d^3 p_1^2}{2l}\right)^{4/7}, \quad \text{critical pressure for turbulent flow.}$$
(4.91)

Maximum gas flows q_{nV}^* for laminar and turbulent flows through a circular tube are calculated from Eqs. (4.82) and (4.87), respectively, if the critical pressure p^* is used as exit pressure p_2 , according to Eqs. (4.90) and (4.91), respectively.

4.3.2

Airflow through a Tube

We will now study airflow through a circular tube (diameter d) at 20 °C (Figure 4.23) and will derive matched numeric-value equations. Numeric-value equations will be marked with a bar to the left in order to differentiate them from primarily used physical-quantity equations. A tube with pressures p_1 at the inlet and p_2 at the outlet is examined. To obtain numeric-value equations, the following properties of air will be used:

- Mean thermal particle speed, $\overline{c} = 463 \,\mathrm{m \, s^{-1}}$.
- Viscosity, $\eta = 18.2 \times 10^{-6} \text{ Pa s.}$
- Flow function at critical point, $\Psi(p^*/p_1) = 0.484$.
- Speed of sound, $a = 343 \,\mathrm{m \, s^{-1}}$.

A short tube behaves just like a nozzle. Here, pV flow is given by Eq. (4.59). Supposing outlet pressure is less than, approximately, half of the inlet pressure $(p_2 < \frac{1}{2} p_1)$, choked flow develops. For this, the numeric-value equation $(q_{pV} \text{ in mbar } \ell \text{ s}^{-1}, d \text{ in cm, } p_1 \text{ in mbar}) \text{ reads}$

$$q_{pV} = 15.6 d^2 p_1$$
, choked nozzle airflow. (4.92)

For long tubes, it is necessary to verify whether flow is laminar or turbulent. The criterion for this is expressed in Eq. (4.80). With the values, the numeric-

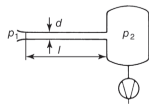


Figure 4.23 Flow inside a tube. Flow at the inlet is nearly inviscid, whereas the flow is viscid inside the long tube.

value equation is obtained (q_{vV} in mbar ℓ s⁻¹, d in cm):

$$\frac{q_{pV}}{d}$$
 $\begin{cases} < 277, & \text{laminar flow,} \\ > 481, & \text{turbulent flow.} \end{cases}$

Equation (4.82) is used for laminar flow. With the values for air, the following numeric-value equation is found $(q_{pV}$ in mbar ℓ s⁻¹, p_1 and p_2 in mbar, d and lin cm):

$$\frac{q_{pV}}{d} = 135 \frac{d^3}{l} \cdot \frac{p_1^2 - p_2^2}{2}, \quad \text{laminar airflow.}$$
 (4.94)

Equation (4.87) is applicable to turbulent flow. With the values for air, the following numeric-value equation is found (q_{pV} in mbar ℓ s⁻¹, p_1 and p_2 in mbar, dand l in cm):

$$\frac{q_{pV}}{d} = 136 \left(\frac{d^3}{l} \frac{p_1^2 - p_2^2}{2}\right)^{4/7}, \quad \text{turbulent airflow.}$$
 (4.95)

Finally, we will consider the critical pressure by introducing the values to Eqs. (4.90) and (4.91) (p^* and p_1 in mbar, d and l in cm):

$$p^* = 2.5 \frac{d^2 p_1^2}{I}$$
, critical pressure for laminar airflow, (4.96)

$$p^* = 5.1 \frac{1}{d} \left(\frac{d^3 p_1^2}{2l} \right)^{4/7}$$
, critical pressure for turbulent airflow. (4.97)

For convenient practical use, the previous equations are plotted in Figures 4.24 and 4.25.

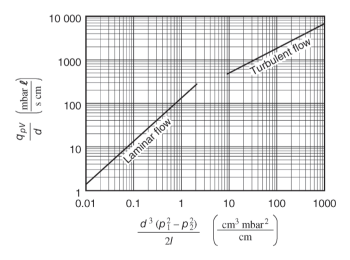


Figure 4.24 Nomogram for obtaining the airflow (20 °C) according to the equations for laminar airflow, Eq. (4.94), and turbulent airflow, Eq. (4.95), through a long tube with circular cross section for fixed inlet and outlet pressures.

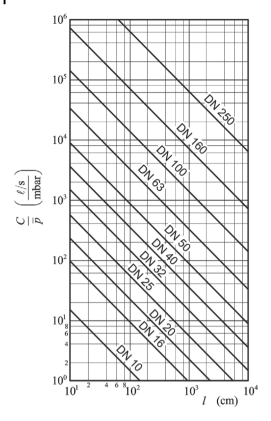


Figure 4.25 Conductance C according to Eq. (4.83) divided by the mean pressure $\overline{p} = \frac{1}{5}(p_1 + p_2)$ for laminar airflow (20 °C) through tubes with circular cross section of given diameters (in mm) versus tube length / in cm.

4.3.3 Air Inflow to a Vessel: Examples

Example 4.7

A vessel is connected to ambient air by means of a capillary tube (inside diameter d = 1 mm, length l = 10 m). Ambient pressure $p_1 = 1000$ mbar and vessel pressure $p_2 = 5$ mbar. The gas flow is the missing quantity.

A prompt solution is found using Figure 4.26. It contains the considered case in the curve DN 1.0 at the right border of the figure. The annotation at the curve indicates laminar flow, and the ordinate value specifies a pV flow close to 7 mbar ℓ s⁻¹ (Figure 4.27).

In general, a plot containing the desired solution is usually unavailable. Therefore, we will also solve the problem by calculation using the nomogram in Figure 4.24. First, the abscissa parameter for the nomogram is calculated

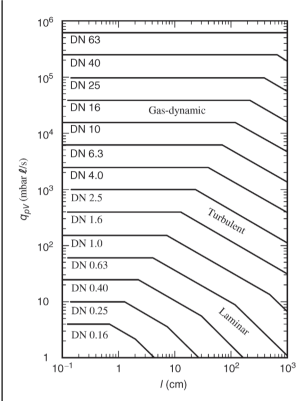


Figure 4.26 pV flow of air (20 °C) through tubes with circular cross section at inlet pressure $p_1 = 1000$ mbar and negligible outlet pressure $p_2 \ll p_1$ versus tube

length I. The gas-dynamic (choked) flow is calculated from Eq. (4.92), the laminar flow from Eq. (4.94), and the turbulent flow from Eq. (4.95).

(numeric-value equation, d and l in cm, p_1 and p_2 in mbar):

$$\frac{d^3}{2l}(p_1^2 - p_2^2) = \frac{0.1^3}{2 \times 1000}(1000^2 - 5^2) = 0.5. \tag{4.98}$$

For this abscissa value, according to Figure 4.24, flow is laminar. The ordinate and Eq. (4.94) reveal that $(q_{pV} \text{ in mbar } \ell \text{ s}^{-1}, d \text{ in cm})$

$$\frac{q_{pV}}{d} = 68. (4.99)$$

Thus, pV flow

$$q_{pV} = \frac{q_{pV}}{d}d = 68 \times 0.1 \text{ mbar } \ell \text{ s}^{-1} = 6.8 \text{ mbar } \ell \text{ s}^{-1}.$$
 (4.100)

When using the nomogram, it is necessary to verify whether the entrance effect at the tube has a significant influence on the amount of gas flow. For

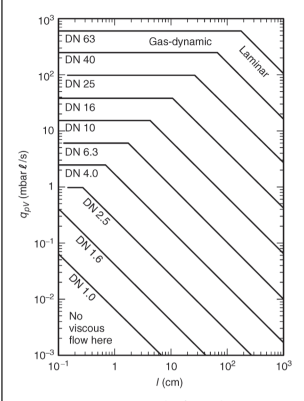


Figure 4.27 Same as Figure 4.26 but for an inlet pressure $p_1 = 1$ mbar.

choked nozzle flow, according to Eq. (4.92),

$$q_{pV} = 15.6 \times 0.1^2 \times 1000 \text{ mbar } \ell \text{ s}^{-1} = 156 \text{ mbar } \ell \text{ s}^{-1}.$$
 (4.101)

For the given opening of the tube, the flow through a nozzle (Eq. (4.101)) is considerably higher than flow through a tube (Eq. (4.100)). Thus, the entrance effect has practically no influence on the amount of gas flow.

Furthermore, we can examine whether choked flow develops in the tube. Following Eq. (4.96), the critical pressure for laminar flow is

$$p^* = 2.5 \frac{0.1^2 \times 1000^2}{1000} \text{mbar} = 25 \text{ mbar}.$$
 (4.102)

The critical pressure (25 mbar) is higher than the vessel pressure (5 mbar). Thus, choked flow in fact occurs. Here, the pressure in the outlet is 25 mbar, and after the gas leaves the tube, it expands to vessel pressure in the vessel. Therefore, when calculating gas flow, the pressure in the outlet must be used for p_2 in Eq. (4.98), instead of the vessel pressure (see Section 4.3.1). However, because the pressure at the outlet of the tube is low compared with the inlet, the effect on pV flow through the tube is minute.

Finally, we will check whether flow through the complete tube is viscous. Equations (4.2) and (4.3) require that

$$Kn = \frac{\pi}{4} \cdot \frac{\overline{c} \,\eta}{p \,d} < 0.01. \tag{4.103}$$

Thus, the pressure must comply with the following condition:

$$p > 25\pi \frac{\overline{c} \, \eta}{d} = 25\pi \frac{463 \,\mathrm{m \, s^{-1}} \times 18.2 \times 10^{-6} \,\mathrm{Pa \, s}}{10^{-3} \,\mathrm{m}} = 662 \,\mathrm{Pa}.$$
 (4.104)

Here, the condition reads p > 7 mbar, which is valid for the complete length of the tube.

Example 4.8

The air in the environment of a chemical reactor is monitored continuously to identify certain pollutants with a laser-optical detector, operating at a pressure of a few millibars. For this, the so-called sniffing operation is used to suck ambient air ($p_1 = 1000 \text{ mbar}$) into the detector via a tube (inside diameter d = 2 mm, length I = 12 m). A displacement pump with pumping speed S = 18 m³ h⁻¹ = $5 \ell s^{-1}$ evacuates the detector (Figure 4.23). Missing quantity is the stationary pressure that develops inside the detector.

In a stationary condition, the flow of air into the detector is equal to the flow of air drawn by the pump. The airflow is usually given as pumping speed, that is, as volume flow rate q_{V} , which is pressure dependent (see Figure 4.28).

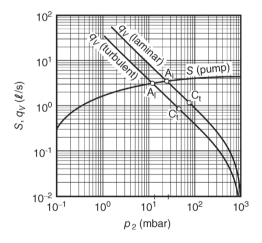


Figure 4.28 Pumping speed and incoming volume flow versus detector pressure in Example 4.8. "A" indicates the stationary operating point, and C the critical point for laminar and turbulent flows (indicated by subscripts "I" and "t," respectively).

To solve the problem, it is helpful to provide the air inflow as a volume flow rate as well. If the volumetric flow through the tube is plotted into Figure 4.28 as a function of detector pressure, the crossing between the curves marks the missing stationary working point. As it is unknown whether the flow through the tube is laminar or turbulent, calculation of both conditions is necessary.

a. Laminar flow of air pV flow is given by Eq. (4.94). Maximum flow occurs for low detector pressure $p_2^2 \ll p_1^2$ (q_{pV} in mbar ℓ s⁻¹, p_1 in mbar, d and l in cm):

$$q_{pV,max} = 135 \frac{d^4}{I} \cdot \frac{p_1^2}{2} = 135 \frac{0.2^4}{1200} \cdot \frac{(10^3)^2}{2}$$

= 90 mbar ℓ s⁻¹. (4.105)

The corresponding volume flow rate $q_V = q_{pV,max}/p_2$ appears as a falling 45° diagonal in the logarithmic representation of Figure 4.28. As detector pressure p_2 increases, q_{pV} drops below the maximum value. Equation (4.94) shows that q_{pV} is 1% lower for $p_2 = 100$ mbar, compared with $p_2 = 0$. If the detector pressure p_2 approaches ambient pressure p_1 , the curve approaches a vertical line at $p_2 = p_1 = 1000$ mbar asymptotically. In Figure 4.28, this part of the curve was drawn freehand.

According to Eq. (4.96), critical detector pressure p_2 at which choked flow develops is $(p^* \text{ and } p_1 \text{ in mbar, } d \text{ and } l \text{ in cm})$

$$p^* = 2.5 \frac{d^2}{I} p_1^2 = 2.5 \frac{0.2^2}{1200} (10^3)^2 \text{ mbar}$$

= 83 mbar. (4.106)

In Figure 4.28, this pressure value is marked by point C_I. Thus, choked flow develops at pressures $p_2 < 83$ mbar. However, as $p_1^2 - p^{*2} \approx p_1^2$, the effective gas flow, according to Eq. (4.94), is practically equal to the maximum gas flow calculated above.

The intersecting point of the curves for pumping speed S and volume flow rate q_V entering the detector through the tube indicates a stationary detector pressure $p_2 = 26$ mbar (working point A_I).

b. Turbulent flow of air pV flow is calculated using Eq. (4.95). Maximum flow occurs for low detector pressure $p_2^2 \ll p_1^2$ (q_{pV} in mbar ℓ s⁻¹, p_1 in mbar, d

$$q_{pV,\text{max}} = 136 d \left(\frac{d^3}{I} \cdot \frac{p_1^2}{2}\right)^{4/7}$$

$$= 136 \times 0.2 \left(\frac{0.2^3}{1200} \cdot \frac{1000^2}{2}\right)^{4/7} = 54 \text{ mbar } \ell \text{ s}^{-1}.$$
(4.107)

The corresponding volume flow rate is plotted in Figure 4.28. According to Eq. (4.97), the critical detector pressure p_2 at which choked flow develops is $(p^* \text{ and } p_1 \text{ in mbar, } d \text{ and } l \text{ in cm})$

$$p^* = 5.1 \frac{1}{d} \left(\frac{d^3}{l} \cdot \frac{p_1^2}{2} \right)^{4/7} = 5.1 \frac{1}{0.2} \left(\frac{0.2^3}{1200} \cdot \frac{1000^2}{2} \right)^{4/7} = 51 \text{ mbar.}$$
 (4.108)

The corresponding point is marked with Ct.

The intersecting point of the curves for S and q_V gives a stationary detector pressure $p_2 = 13$ mbar (working point A_t). Here also, choked flow develops, which has hardly any impact on the gas flow.

c. In order to assess a possible effect of the entrance flow, we will calculate the gas flow through a nozzle of the same diameter (Eq. (4.92)) $(q_{pV}$ in mbar ℓ s⁻¹, p_1 in mbar, d in cm):

$$q_{pV} = 15.6 d^2 p_1 = 15.6 \times 0.2^2 \times 1000$$

= 624 mbar ℓ s⁻¹. (4.109)

The value is far above laminar or turbulent gas flow, and thus, entrance effects can be neglected in this example.

With the Reynolds number, or the abscissa in the nomogram of Figure 4.24, we can determine whether the flow is laminar or turbulent. The abscissa parameter calculates to $(p_1 \text{ and } p_2 \text{ in mbar, } d \text{ and } l \text{ in cm})$

$$\frac{d^3}{l} \cdot \frac{p_1^2 - p_2^2}{2} = \frac{d^3}{l} \cdot \frac{p_1^2}{2} = \frac{0.2^3}{1200} \cdot \frac{1000^2}{2} = 3.3.$$
 (4.110)

This shows that the condition is located in the transition region between laminar and turbulent flows where behavior can only be approximated. Thus, the detector pressure is in the range of 13–26 mbar.

4.3.4

Tube at the Inlet of a Pump: Examples

A chamber with pressure p_c is evacuated via a tube with length l and diameter d, using a pump with pumping speed S. At the inlet flange of the pump, the pressure is p_{in} (Figure 4.29). Due to flow resistance (conductance C) in the tube, the effective pumping speed S_{eff} at the chamber remains below the pumping speed Sof the pump. As shown in Section 4.1.4,

$$\frac{S_{\text{eff}}}{S} = \frac{p_{\text{in}}}{p_c} = \frac{C}{C+S} = \frac{1}{1+S/C}.$$
(4.111)

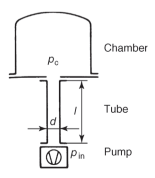


Figure 4.29 Evacuating a vacuum chamber through a tube.

The pressures p_{in} and p_c determine the conductance C of the tube. We will now calculate the pressure ratio p_c/p_{in} for the laminar and turbulent cases in a circular tube.

Assuming laminar and turbulent flows through a circular tube, conductance C is given by Eqs. (4.83) and (4.88), respectively. By using the formula for conductance, Eq. (4.111) can be converted so that the pressure ratio is

$$\frac{p_{\rm c}}{p_{\rm in}} = \sqrt{1 + \frac{256 \,\eta \,l}{\pi \,d^4} \cdot \frac{S}{p_{\rm in}}} \quad \text{for laminar flow}, \tag{4.112}$$

$$\frac{p_{\rm c}}{p_{\rm in}} = \sqrt{1 + 0.974 \, l \left(\frac{\eta \, S^7}{d^{19} \, \overline{c}^6 \, p_{\rm in}}\right)^{1/4}} \quad \text{for turbulent flow.}$$
 (4.113)

For air at $20 \,^{\circ}$ C ($\eta = 18.2 \times 10^{-6} \, \text{Pa s}$, $\bar{c} = 463 \, \text{m s}^{-1}$), these two equations, rewritten as fitted numeric-value equations, read ($p_{
m c}$ and $p_{
m in}$ in mbar, S in ℓ s⁻¹, d and l in cm)

$$\frac{p_{\rm c}}{p_{\rm in}} = \sqrt{1 + 0.0148 \frac{l}{d^4} \cdot \frac{S}{p_{\rm in}}} \quad \text{for laminar airflow}, \tag{4.114}$$

$$\frac{p_{\rm c}}{p_{\rm in}} = \sqrt{1 + 0.00036l \left(\frac{S^7}{d^{19}p_{\rm in}}\right)^{1/4}}$$
 for turbulent airflow. (4.115)

Example 4.9

A pump with the pumping speed $S = 18 \text{ m}^3 \text{ h}^{-1}$ sucks in air from a chamber at a temperature $\vartheta = 20\,^{\circ}\text{C}$ with a pressure $p_{\text{in}} = 3\,\text{mbar}$, through a tube with the diameter d=25 mm (equal to the pump's inlet flange) and length l=2 m. How high is the effective pumping speed S_{eff} at the chamber flange?

First, we will determine whether the entrance flow into the tube is relevant, that is, whether choked flow occurs. The maximum pV flow sucked by the pump is obtained by attaching it directly to the vessel:

$$q_{\rm pV} = p_{\rm in} S = 3 \,\text{mbar} \times 5 \,\ell \,\text{s}^{-1} = 15 \,\text{mbar} \,\ell \,\text{s}^{-1}.$$
 (4.116)

According to Eq. (4.92), the choked aperture flow ($q_{\rm DV}$ in mbar ℓ s⁻¹, p_1 in mbar, d in cm) is

$$q_{pV} = 15.6d^2p_1 = 15.6 \times 2.5^2 \times 3 = 293 \text{ mbar } \ell \text{ s}^{-1}.$$
 (4.117)

Flow through the pump is considerably below this maximum aperture flow. As a result, the effect of entrance flow on the overall flow is negligible and the equations for frictional flow are applicable. We will now analyze whether the flow is laminar or turbulent. For this, the Reynolds number is assessed according to

Eq. (4.93). To calculate pV flow, we will use the maximum pV flow sucked by the pump (q_{pV} in mbar ℓ s⁻¹, d in cm):

$$\frac{q_{pV}}{d} = \frac{15}{2.5} = 6. {(4.118)}$$

As this value is below 277, the flow is laminar according to Eq. (4.93). Then Eq. (4.114) allows calculating the pressure ratio:

$$\frac{p_{\rm in}}{p_c} = \sqrt{1 + 0.0148 \times \frac{200}{2.5^4} \cdot \frac{5}{3}} = 1.061. \tag{4.119}$$

The effective pumping speed at the vessel is calculated with Eq. (4.111):

$$S_{\text{eff}} = \frac{p_{\text{in}}}{p_{\text{c}}} S = \frac{1}{1.061} \times 18 \text{ m}^3 \text{ h}^{-1} = 17.0 \text{ m}^3 \text{ h}^{-1}.$$
 (4.120)

As the example shows, flow resistance is negligible in the rough vacuum range (>1 mbar) if appropriate tubing is used. In a certain pressure range, the pumping speed of many pumps hardly depends on the pressure. In contrast, the conductance of tubes depends on the pressure under laminar as well as turbulent flow conditions: flow resistance increases with falling pressure. As a result, a tube placed between vessel and pump reduces pumping speed marginally under atmospheric pressure but causes a significant reduction of pumping speed under low-pressure conditions.

If the tolerated loss in pumping speed of a pump, caused by the inlet tubing, is 10% (90% utilization of pump), then the ratio $S_{\rm eff}/S$ must not be lower than 0.9, and the ratio p_c/p_{in} not any higher than 1.11. From Eqs. (4.114) and (4.115), it follows that the maximum permissible length of the suction line (fitted numericvalue equation, S in ℓ s⁻¹, p_{in} in mbar, l and d in cm) is

$$l > 16 \frac{d^4}{S} p_{\text{in}}$$
 for laminar airflow, i.e., $\frac{p_{\text{in}}S}{d} < 277$, (4.121)

$$l < 650 \frac{d^5}{S^2} \sqrt[4]{\frac{p_{\rm in}S}{d}}$$
 for turbulent airflow, i.e., $\frac{p_{\rm in}S}{d} > 481$. (4.122)

Example 4.10

A displacement pump with a pumping speed $S = 72 \text{ m}^3 \text{ h}^{-1}$ is to be connected to a vacuum vessel via an inlet tube with diameter d = 40 mm. What is the maximum length of the tube if the utilization of the pump is 90%, and the lowest inlet pressure at the pump is (a) 700 mbar and (b) 1 mbar?

Case (a): pV flow for low inlet pressure is

$$q_{pV} = p_{in}S = 700 \text{ mbar} \times 20 \,\ell \,\text{s}^{-1} = 14\,000 \,\text{mbar} \,\ell \,\text{s}^{-1}.$$
 (4.123)

Thus, assessing the type of flow according to Eq. (4.93) leads to (q_{pV}) in mbar ℓ s⁻¹, d in cm)

$$\frac{q_{pV}}{d} = \frac{14\,000}{4} = 3500 > 481. \tag{4.124}$$

According to this, the flow appears to be turbulent. Equation (4.122) is used to calculate the length, and thus,

$$I < 650 \times \frac{4^5}{20^2} \sqrt[4]{\frac{700 \times 20}{4}} \approx 12\,800 \,\text{cm}, \quad \text{i.e.}, \quad I_{\text{max}} = 128 \,\text{m}. \quad (4.125)$$

Case (b): Calculation is analogous to case (a).

The flow is laminar and the maximum acceptable length is $I_{\text{max}} = 2 \text{ m}$.

4.3.5

Flow through Ducts with Noncircular Cross Sections

So far, flow conditions in tubes with circular cross sections were investigated. In these devices, the cross section is characterized by the diameter d. If, however, the cross section has a different shape, the hydraulic diameter d_h can be used as a characteristic value to describe the cross section. The hydraulic diameter is defined by

$$d_{\rm h} = 4 \times \frac{\text{cross-sectional area of duct}}{\text{circumference of duct}}$$
 (4.126)

Now, ducts with circular, ring-shaped (annular), and rectangular cross sections are investigated (Figure 4.30).

For a circular cross section, the hydraulic diameter corresponds to the geometrical diameter, that is, $d_h = d$. When considering an annular cross section, the hydraulic diameter is

$$d_{\rm h} = 4 \frac{(\pi/4)d_{\rm o}^2 - (\pi/4)d_{\rm i}^2}{\pi d_{\rm o} + \pi d_{\rm i}} = d_{\rm o} - d_{\rm i},$$
 annular cross section. (4.127)

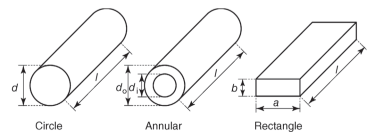


Figure 4.30 Ducts with selected cross sections.

In a rectangular cross section (area $a \times b$), the hydraulic diameter is

$$d_{\rm h} = 4 \frac{a \, b}{2(a+b)} = \frac{2 \, a \, b}{a+b},$$
 rectangular cross section. (4.128)

The Reynolds number, calculated with Eq. (4.4), helps differentiating between laminar and turbulent flows. Here, the hydraulic diameter d_h is used for the quantity d.

In the case of laminar flow, flow conductance values of certain tubes can be derived elementally from the definition of viscosity η . Section 4.3.1 introduced the conductance of a tube with circular cross section (diameter *d*):

$$C = \frac{\pi}{256} \cdot \frac{1}{\eta} \cdot \frac{d^4}{l} (p_1 + p_2) \quad ,$$

tube with circular cross section, laminar flow.

(4.83)

For the annular gap formed by two concentric tubes with circular cross sections (diameters d_0 and d_i), we find that

$$C = \frac{\pi}{256} \cdot \frac{1}{\eta l} \left[d_o^4 - d_i^4 - \frac{(d_o^2 - d_i^2)^2}{ln(d_o/d_i)} \right] (p_1 + p_2) \quad ,$$

tube with annular cross section, laminar flow.

(4.129)

The conductance of a narrow slot with rectangular cross section (area $a \times b$, with $a \gg b$) is calculated elementally as well:

$$C = \frac{1}{24} \cdot \frac{1}{\eta} \cdot \frac{a b^3}{l} (p_1 + p_2) \quad , \quad \text{narrow slot, laminar flow.}$$
 (4.130)

This equation can be generalized to apply to an arbitrary tube with rectangular *cross section* (area $a \times b$, with $a \ge b$):

$$C = \frac{1}{24} \cdot \frac{1}{\eta} \cdot \frac{a b^3}{l} (p_1 + p_2) \left[1 - \frac{192 b}{\pi^5 a} \sum_{n=1,3,5,\dots}^{\infty} \frac{1}{n^5} \tanh \left(\frac{\pi n}{2} \cdot \frac{a}{b} \right) \right]$$

rectangular cross section, laminar flow.

(4.131)

The series converges rapidly but calculation of the terms is tedious. However, the following equation represents a close approximation that is accurate for

a = b and $a \gg b$, and in other cases produces an error of less than 3%:

$$C \approx \frac{1}{24} \cdot \frac{1}{\eta} \cdot \frac{a^3 b^3 (p_1 + p_2)}{l(a^2 + b^2 + 0.371ab)}, \quad \text{rectangular cross section,}$$

$$\text{laminar flow.} \quad (4.132)$$

The conductance of a tube with circular cross section under turbulent flow conditions was already given in Section 4.3.1:

$$C \approx d^{19/7} \left(\frac{\overline{c}^6}{\eta}\right)^{1/7} \left(\frac{p_1 + p_2}{l}\right)^{4/7} (p_1 - p_2)^{-3/7} \quad , \quad \text{circular cross section,}$$
turbulant flow. (4.88)

The conductance C_x of a tube with noncircular cross section can be estimated by substituting the tube with an appropriately picked circular tube, for example, with the same hydraulic diameter, and by subsequently calculating the conductance of the latter.

4.3.6 Influence of Gas Species on Flow

In the previous section, equations were often written out in two different ways: as physical-quantity equations for any type of fluid and as fitted numeric-value equations for air at 20 °C. If equations are required for other gas species, the equations for air can be used if appropriate scaling is taken into account. The following properties of a gas determine the developing flow.

Viscosity η determines frictional behavior, and thus the velocity of flow under frictional flow conditions. In the viscous regime, viscosity is almost pressure independent and increases proportionally to T^{ω} , where $0.66 \le \omega \le 1.1$, according to Table A.1 in Ref. [3].

The velocity \bar{c} of the thermal particle motion directly determines the velocity of flow in gas-dynamic flow. This velocity is proportional to the square root of the thermodynamic temperature T and inversely proportional to the square root of particle mass m_P .

The heat capacity of a gas enters into calculation of the isentropic exponent κ , which affects flow behavior in gas-dynamic flow: during expansion, a gas with low heat capacity cools more rapidly than a gas with high heat capacity. Cooling changes the gas volume and particle velocity, and thus overall flow behavior. The dependence of flow rate and conductance from the isentropic exponent is formulated in the flow function ψ .

Table 4.3 lists the scaling behavior for different types of flows.

Under laminar flow conditions, for example, for a component of arbitrary geometry and different gas species under equal conditions (line dimensions,

Type of flow	Scaling behavior of throughput $q_{ m pV}$ and conductance C	
Gas-dynamic	Proportional to $\overline{c}\psi_{\kappa}$	
Laminar	Proportional to $1/\eta$	
Turbulent	Proportional to $\overline{c}^{6/7}/\eta^{1/7}$	

Table 4.3 Dependence of viscous flow on the properties of the gas species.

pressures at inlet and outlet),

$$q_{pV} \propto \frac{1}{\eta}$$
 and $q_{pV}(\text{any gas}) = q_{pV}(\text{air}) \frac{\eta(\text{air})}{\eta(\text{any gas})}$. (4.133)

If we compare the pV flow through a capillary featuring laminar flow for the gases helium and air at 20 °C, we find

$$q_{pV}(\text{helium}) = q_{pV}(\text{air}) \frac{18.2 \times 10^{-6} \text{ Pa s}}{19.6 \times 10^{-6} \text{ Pa s}} = q_{pV}(\text{air}) \times 0.93, \tag{4.134}$$

that is, pV flow for helium is 7% lower than that of air.

4.4 Molecular Flow under High-Vacuum and Ultrahigh-Vacuum Conditions

4.4.1

Flow Pattern, Definitions, and Transmission Probability

Under high-vacuum and ultrahigh-vacuum conditions, the mean free path l of gas particles is large compared with the cross dimension d of a tube, and thus, the *Knudsen* number $Kn = \overline{l}/d \gg 0.5$. This indicates that gas particles traveling through an aperture pass the aperture without suffering mutual collisions, and that, considering flow through a tube, an individual particle hits the walls of the tube much more often than it hits other particles.

For calculation of molecular flow through a component (aperture, tube), the concept of transmission probability P is useful, because the motion of large numbers of gas particles through a component is statistical. Here, the assumption is used that an approaching gas particle initially passes through the entrance plane of the component and then travels through the component. There, the particle makes numerous wall collisions and finally leaves the component, either through the exit plane or through the initial entrance plane. The probability of a gas particle, which entered through the entrance plane, leaving the component at the exit plane is referred to as the transmission probability P. In real situations, the number of particles is always large, and thus, averages over many particles and over particle flow (= particles/time) can be taken:

$$\left\{ \begin{array}{l} \text{particle flow at} \\ \text{exit plane } q_{N2} \end{array} \right\} = \left\{ \begin{array}{l} \text{particle flow at} \\ \text{entrance plane } q_{N1} \end{array} \right\} \cdot \left\{ \begin{array}{l} \text{transmission} \\ \text{probability } P \end{array} \right\}.$$
 (4.135)

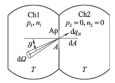


Figure 4.31 Molecular flow through a thin aperture Ap placed in between two vacuum chambers Ch1 and Ch2.

First, the direction distribution of gas particles for molecular flow through a thin aperture with opening area *A*, placed between two chambers, is investigated. The size and shape of the chambers shall suggest that the velocity distribution of the gas particles is completely isotropic (Figure 4.31).

A stream of gas particles entering from the left-hand side, and hitting the wall on the right side of chamber Ch1 (Figure 4.31) at a right angle, passes through the complete area A of the aperture. For a stream of gas particles entering at an angle ϑ to the perpendicular direction, only an area $A\cos\vartheta$ is available for passing through the aperture. Thus, considering angular distribution, the particles leaving the aperture show a *cosine distribution* (Figure 4.32).

Inside chamber Ch1, the pressure is p_1 and the particle number density is n_1 . Chamber Ch2, however, shall contain perfect vacuum. Under these conditions, the number of particles approaching the opening area A per unit time, that is, particle flow q_N , is

$$q_N = j_N A = \frac{1}{4} n_1 \bar{c}_1 A. \tag{4.136}$$

Here, j_N and \overline{c}_1 denote the collision rate (Eq. (3.48)) and the thermal particle velocity (Eq. (3.43)) in chamber C_1 , respectively.

We will now substitute the aperture with a *tube* of finite length. Once more, we will investigate the angular distribution of the particles passing through. Initially, the length l of the tube shall be short compared with its cross dimensions (= diameter, for circular cross section) (see Figure 4.33).

All gas particles hitting the entrance plane at normal direction travel through the tube and leave it at the exit plane. Therefore, the transmission probability for these particles is P = 1. A certain portion of the particles passing through the entrance plane at an angle collides with the wall of the tube and is reflected. Experimental investigations on wall collisions have shown that, with good

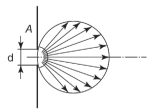


Figure 4.32 Angular distribution of particles passing through a thin aperture.

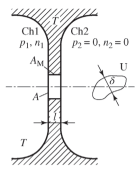


Figure 4.33 Molecular flow through a short tube in between two chambers.

approximation, emission is diffuse for technical surfaces. Thus, a particle's direction after being emitted by the wall is practically independent of the initial angle of incidence and shows a cosine distribution. A particle reflected by the wall moves in or opposite to the direction of flow with the same probability. As a result, the particle leaves the tube either at the entrance plane or at the exit plane.

Now we will consider a tube that is longer than its cross dimensions (Figure 4.34). Here, many of the particles colliding with the wall continue to hit the wall and will undergo further emissions. Especially in long tubes, particles can suffer many wall collisions.

For each individual collision, the probability of the particle to move forward or backward, with respect to the direction of flow, is equal. If the tube is long enough, only few particles make it to the exit plane; most of the particles suffer many wall collisions and return to the entrance plane.

The better a particle's direction of motion is aligned with the direction of flow, the longer the path that a particle travels within the tube without colliding. As a result, particles leaving the tube at the exit plane feature a direction of motion that is mostly perpendicular to this plane. In contrast, particles returning are practically never directed perpendicularly to the entrance plane.

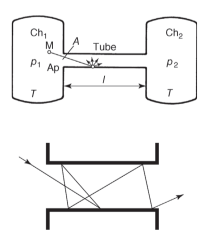


Figure 4.34 Molecular flow through a long tube in between two chambers.

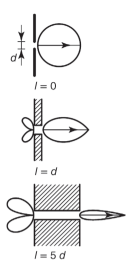


Figure 4.35 Molecular flow through a tube with circular cross section. Particles reach the opening from the left-hand side. The figure shows the angular distribution of particles

leaving the tube in forward and backward directions, for selected values of the ratio of tube length I to diameter d of the tube.

P. Clausing (1930) conducted quantitative calculations of the angular distribution of particles emanating from a tube with circular cross section under the assumption of diffuse wall reflection. Results are shown in Figure 4.35.

We will now introduce several terms. The particle flow moving from chamber Ch1 to chamber Ch2 is already given in Eq. (4.136). If the pressure in chamber C_2 is finite as well, an opposed particle flow develops, directed from chamber Ch2 to chamber Ch1. The net particle flow is

$$q_N = \frac{1}{4} n_1 \overline{c}_1 A P_{12} - \frac{1}{4} n_2 \overline{c}_2 A P_{21} \tag{4.137}$$

if the interflowing particle streams do not interfere, which is true for molecular flow. In the case considered here, that is, a tube with constant cross-sectional area, the transmission probability P_{12} from chamber Ch1 to chamber Ch2 is the same as P_{21} from chamber Ch2 to chamber Ch1, as suggested by symmetry. Thus, the subscript for P can be omitted.

Furthermore, we will assume that both vessels contain the same type of gas at the same temperature, and thus, $\overline{c}_1 = \overline{c}_2 = \overline{c}$. Using this, the net flow through the tube is

$$q_N = \frac{\overline{c}}{4}AP(n_1 - n_2), \quad \text{particle flow}, \tag{4.138}$$

$$q_{pV} = \frac{\overline{c}}{4}AP(p_1 - p_2), \quad \text{throughput}, pV \text{ flow}, \tag{4.139}$$

$$q_m = \frac{2}{\pi} \cdot 1\overline{c}AP(p_1 - p_2), \quad \text{mass flow.}$$
 (4.140)

From this and according to the defining equation, conductance *C* is given by

$$C = \frac{q_N}{n_1 - n_2} = \frac{q_{pV}}{p_1 - p_2} = \frac{\overline{c}}{4}AP, \quad [C] = \text{m}^3 \text{ s}^{-1}.$$
 (4.141)

Flow resistance *R* is just the reciprocal value of conductance *C*:

$$R = \frac{1}{C} = 4\overline{c} \cdot \frac{1}{AP}, \quad [R] = \text{s m}^{-3}.$$
 (4.142)

The assembly influences the conductance of a component. If the component, as assumed above, is arranged between two very large vessels, the gas flow into the component shows isotropic distribution of the direction of flow. The conductance developing in this case is termed inherent conductance or intrinsic conductance (DIN 28400, Part 1, and ISO 3529-1).

However, if the component is built into a tube, the particle flow at the entrance plane of the component is directional, and the direction perpendicular to the plane is privileged (beam formation) (see also Figure 4.35). In this case, gas particles travel through the part more easily than under isotropic inflow. Here, the developing conductance is termed assembly conductance or reduced conductance.

4.4.2

Molecular Flow through an Aperture

In a thin aperture (tube of length zero), the transmission probability is P = 1, independent of the geometry of the opening (cross section). Thus, according to Eq. (4.141), intrinsic conductance is

$$C_{\mathcal{A}} = \frac{\overline{c}}{4}A. \tag{4.143}$$

As example, for air at 20 °C (particle velocity $\bar{c} = 463 \text{ m s}^{-1}$), intrinsic conductance is

$$C_{\rm A} = 11.6 \, \ell \, \rm s^{-1}$$
, for air at 20 °C, aperture area $A = 1 \, \rm cm^2$, (4.144)

$$C_{\rm A}=9.1\,\ell~{\rm s}^{-1},~{\rm for~air~at~20^{\circ}C,~circular~aperture~with~diameter}=1~{\rm cm}.$$
 (4.145)

Table 4.4 lists conductance values of thin circular orifices having the diameter of standardized flanges.

In order to calculate conductance values for other gas species and different temperatures, \bar{c} is taken from Table A.10, or calculated according to Eq. (3.43). Conductance is proportional to the square root of the thermodynamic temperature and inversely proportional to the square root of the particle mass (Table 4.5):

$$C_{\rm A} \propto \sqrt{\frac{T}{m_{\rm P}}}$$
 (4.146)

Table 4.4 Flow conductances of apertures with diameters corresponding to the clear opening widths of standard flanges.

Nominal sizes given as DN (dia- mètre nominale) without unit according to DIN ENISO 6708/ ISO 3445	Real inside diameter (mm) according to DIN 28403/ISO 2861, DIN 28404/ISO 1609	Intrinsic conductance C_A for molecular flow of air at 20 °C (\mathscr{C} s ⁻¹)
10	10	9.1
16	16	23.3
20	21	40.1
25	24	52.4
32	34	105
40	41	153
50	51	237
63	70	446
80	83	627
100	102	947
125	127	1468
160	153	2130
200	213	4129
250	261	6199
320	318	9202
400	400	14 560
500	501	22 840
630	630 or 651	38 570
800	800	58 240
1000	1000	91 000

This behavior is observed not only in apertures but, generally, in the case of molecular flow because the transmission probability P depends only on the geometry of the component (assuming diffuse emission after collisions with wall).

Table 4.5 Flow conductances for various gases with respect to the flow conductance of air under molecular flow conditions.

Gas	Relative particle mass	C(gas)/C(air)		
H_2	2	3.8		
He	4	2.7		
H_2O	18	1.27		
Air	29	1.00		
Ar	40	0.85		
CO_2	44	0.81		

Example 4.11

A thin aperture with diameter d = 2 mm separates two vessels (see also Figure 4.31). The vessels contain argon at 20 °C, and pressures in the vessels $p_1 =$ 0.1 Pa and $p_2 = 0.01$ Pa. Net pV flow through the aperture is the missing quantity. The problem is solved using Eq. (4.139):

$$\begin{split} q_{pV} &= \frac{\overline{c}}{4} A(p_1 - p_2) \\ &= \frac{394 \text{ m s}^{-1}}{4} \cdot \frac{\pi}{4} (0.02 \text{ m})^2 (0.1 \text{ Pa} - 0.01 \text{ Pa}) \\ &= 2.8 \times 10^{-3} \text{ Pa m}^3 \text{ s}^{-1} = 0.028 \text{ mbar } \ell \text{ s}^{-1}. \end{split}$$

$$(4.147)$$

4.4.3

Molecular Flow through a Tube with Constant Cross-Sectional Area

In the previous section, we calculated the conductance C_A of a thin aperture (open area A, length l = 0). Now, we will investigate a straight tube with arbitrary cross section and will consider the two particular cases of short and very long lengths.

The number of gas particles approaching the intake area A per unit time is

$$q_{N,A} = j_N A = \frac{1}{4} n_1 \overline{c}_1 A. \tag{4.148}$$

We start with the case of a short tube (length small compared with cross dimensions). Here, some of the gas particles that cross the inlet hit the interior surface $A_{\rm S}$ of the tube (Figure 4.33).

Their number per time is

$$q_{N,S} = \frac{1}{2} \cdot \frac{1}{4} n_1 \overline{c}_1 A_S. \tag{4.149}$$

The factor $\frac{1}{2}$ in the equation takes into account that the collision rate at the interior surface of the tube is just half as high as that at the interior surface of the vessel because particles only impact the interior surface from one half-space. Under diffuse reflection conditions, half of the particles hitting the interior surface are emitted in the backward direction and leave through the entrance plane. Thus, the net particle flow through a thin aperture is

$$q_{N,A} = q_{N,A} - \frac{1}{2}q_{N,S} = \frac{1}{4}n_1\bar{c}_1A - \frac{1}{2}\cdot\frac{1}{2}\cdot\frac{1}{4}n_1\bar{c}_1A_S$$

$$= \frac{1}{4}n_1\bar{c}_1A\left(1 - \frac{A_S}{4A}\right). \tag{4.150}$$

The bracketed term in Eq. (4.150) corresponds to the transmission probability

$$P_{\text{short tube}} = 1 - \frac{A_{\text{S}}}{4A}, \quad \text{short tube.}$$
 (4.151)

For long tubes, that is, the length is large compared with cross dimensions, M. Smoluchowski (1910) derived an equation to calculate the transmission probability:

$$P_{\text{long tube}} = \frac{1}{4lA} \int_{s} ds \int_{-\pi/2}^{+\pi/2} d\theta \ b^{2} \cos \theta, \quad \text{long tube.}$$
 (4.152)

Here, l and A are the length and cross-sectional area of the tube, respectively. The integration over s leads along the border (circumference) of the crosssectional area. $b(\theta)$ is the distance that a particle emitted from the wall travels until it hits the wall the next time. θ is the angle between the particle path b and the differential circumference ds. Equation (4.152) is universally valid and allows calculation of the transmission probabilities for tubes with any cross section. Calculation of the integrals is tedious. For a rough estimate, we will assume a circular cross section and equate path b and the (hydraulic) diameter d of the tube. Then, integration simplifies to

$$P_{\text{long tube}} \approx \frac{1}{4l(\pi/4)d^2} \int_{\sigma} ds \int_{-\pi/2}^{+\pi/2} d\theta \, d^2 \cos \theta. \tag{4.153}$$

Therefore, in a long tube of arbitrary, constant cross-sectional area, the transmission probability is proportional to the ratio of hydraulic diameter and length, whereby the proportional factor depends on the geometry of the cross section.

Calculating the transmission probability for a tube with medium-sized length is more challenging. For a rough approximation, we can suppose a series connection of an aperture and a long tube:

$$\frac{1}{P} \approx \frac{1}{P_{\rm A}} + \frac{1}{P_{\rm long \; tube}} \approx 1 + \frac{1}{P_{\rm long \; tube}},$$
 tube with arbitrary length. (4.154)

This equation correctly describes the two limiting cases of a very short and a very long tube. However, for medium-sized lengths, depending on crosssectional geometry, where flow develops from inlet flow (isotropic angular distribution of incoming particles) to pure tube flow (particle velocities mainly in the direction of flow), an error of 10% and more must be accepted.

Knudsen (1909), Smoluchowski (1910), Clausing (1932), and others conducted precise calculations of flow in a tube of arbitrary length. Analytical investigation yields integrals that are evaluated numerically. Today, numerical calculation of the transmission probability is feasible: in the so-called *Monte Carlo simulation*, a computer calculates the paths of individual gas particles entering a tube. These are distributed statistically, and, after each collision with the wall, they reflect statistically in different directions according to a cosine distribution. A random number generator provides data for the statistical distribution. State-of-the-art desktop computers can calculate the paths of numerous gas particles in a tube (one million and more) within minutes, depending on the geometry of the investigated duct. The transmission probability can thus be calculated with high statistical accuracy. Usually, analytical calculations as well as computer simulations rely on the following assumptions:

- · Stationary flow.
- No mutual particle collisions.
- Isotropic inflow.
- Diffuse angular distribution after collision with wall (cosine distribution).

Tubes with simple geometries are appropriate to verify analytical and statistical calculations of the transmission probability. Both approaches lead to the same results. Deviations appearing in some papers are caused by miscalculations or due to differing assumptions. Calculation of more complicated ducts (e.g., valves) can be carried out only by means of statistical methods.

The intrinsic conductance of a component is the product of the aperture conductance of the entrance plane C_A and the transmission probability P:

$$C = C_{\mathcal{A}}P. \tag{4.155}$$

4.4.4

Molecular Flow through a Tube with Circular Cross Section

We will investigate the transmission probability P_C for a tube with circular cross section (diameter d, length l).

For a *short* tube with circular cross section, the transmission probability is obtained from Eq. (4.151):

$$P_{C,\text{short tube}} = 1 - \frac{A_{S}}{4A} = 1 - \frac{l}{d}.$$
 (4.156)

For a long tube with circular cross section, Eq. (4.152) must be solved accurately. As a result,

$$P_{\text{C,long tube}} = \frac{4}{3} \cdot \frac{d}{l}. \tag{4.157}$$

For a tube of medium-sized length, the approximation formula in Eq. (4.154) produces a maximum error of approximately +13% for $l/d \approx 2$. The numerically found results of analytical and statistical calculations can be approximated by an analytical, cut-and-try type function. Choosing the following function, the relative error remains below 0.6%:

$$P_{\rm C} = \frac{14 + 4(l/d)}{14 + 18(l/d) + 3(l/d)^2}$$
,

tube with circular cross section, arbitrary length.

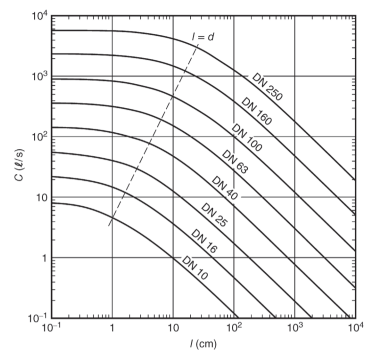


Figure 4.36 Characteristic flow conductances of tubes with circular cross sections and nominal sizes (in mm) for molecular flow of air at 20 °C, calculated from Eq. (4.159).

From the transmission probability P, the intrinsic conductance is obtained by multiplying with the aperture conductance C_A (Eq. (4.143)):

$$C_{\rm C} = C_{\rm A} P_{\rm C} = \frac{\pi}{16} \bar{c} d^2 \frac{14 + 4(l/d)}{14 + 18(l/d) + 3(l/d)^2},$$

tube with circular cross section, arbitrary length. (4.159)

Once more, it should be noted that the probabilities of passage given here assume isotropic inflow. This is the case if large vessels are connected to the inlet and outlet of the tube.

Figure 4.36 shows the intrinsic conductances of tubes with circular cross sections and standardized nominal diameters, for air at 20 °C. Figure 4.37 illustrates the transmission probabilities $P_{\rm C}$ of tubes with circular cross sections.

4.4.5

Molecular Flow through Tubes with Simple Cross-Sectional Geometry

Technical literature lists analytical formulas, lengthy in part, as well as extensive data tables for the transmission probabilities in tubes with simple cross-sectional geometry (see, e.g., Ref. [4]). For practical applications where accuracy in the

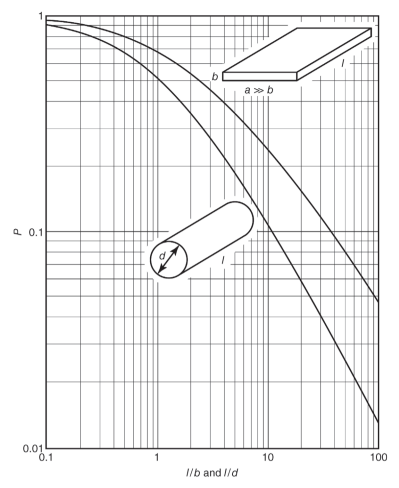


Figure 4.37 Transmission probabilities for molecular flow in a thin gap (top curve) according to Eq. (4.160) and in a tube with circular cross section according to Eq. (4.158) versus length.

percent range is sufficient, values are conveniently read from figures. In the following, several simple analytical formulas as well as figures are listed.

Figures 4.36 and 4.37 show intrinsic conductances and transmission probabilities, respectively, for tubes with circular cross sections.

The transmission probability for a *narrow slot* (height b very small compared with width a) is approximated (error < 1%) by

$$P = \frac{1 + \ln(0.433(l/b) + 1)}{(l/b) + 1} \tag{4.160}$$

where b and l denote the height and length of the slot, respectively. The equation is plotted by the uppermost curves in Figures 4.37 and 4.38.

Figure 4.38 displays the transmission probability for a tube with rectangular *cross section* (or *quadratic* for a = b).

The transmission probability of an annular slot (clearance between two coaxial tubes with radii r_0 and r_i) is shown in Figure 4.39. In the special case with $r_{\rm i}/r_{\rm o}
ightarrow 0$, the annular slot approaches a tube with circular cross section with $d=2r_{\rm o}$, and in the special case $r_{\rm i}/r_{\rm o} \rightarrow 1$ it approaches a narrow slot with b= $r_{\rm o} - r_{\rm i}$ and $a = \pi(r_{\rm o} + r_{\rm i})$.

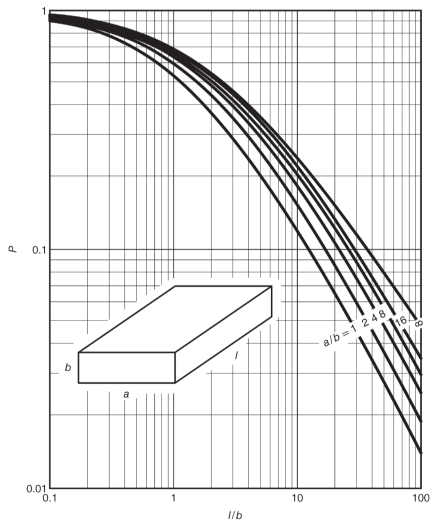


Figure 4.38 Transmission probabilities in tubes with rectangular cross sections (selected aspect ratios) for molecular flow versus length.

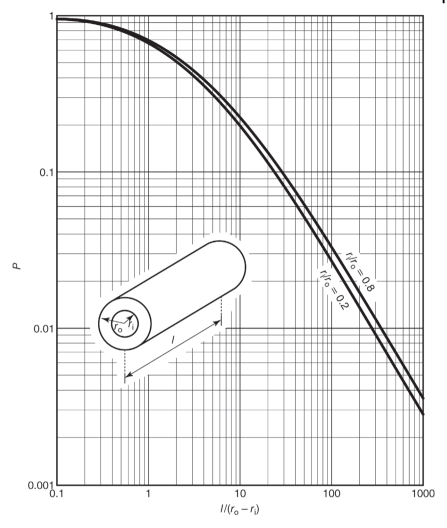


Figure 4.39 Transmission probabilities in tubes with annular cross section (coaxial double tube) with selected radii ratios for molecular flow versus length.

4.4.6 **Tube Bend and Tube Elbow**

Connecting ducts often contain tube bends and elbows (Figure 4.40). Such components feature approximately the same transmission probability as a straight tube with equal cross section and axis length.

For simple geometries, for example, a 90° bend with circular cross-section tubes, calculations for the transmission probability by means of Monte Carlo simulation are available (Figure 4.41).

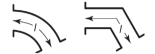


Figure 4.40 Axis length in a tube bend and a tube elbow.

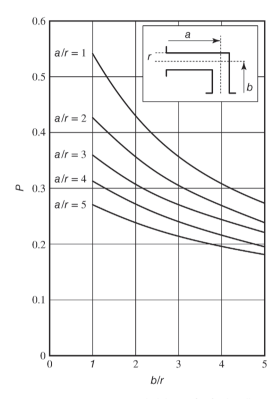


Figure 4.41 Transmission probabilities of 90° tube elbows with circular cross sections and selected dimensions. As in the previous figures, the transmission probabilities apply to a mounting position between two chambers but not to a position between tubes.

Example 4.12

Two vacuum vessels are connected via a 90° elbow consisting of a tube with nominal diameter DN 25 (d = 25 mm). The mounting dimensions of the elbow are the common metric ones, that is, 50 mm displacement in both x- and y-directions.

Case 1: A radius elbow is used. The effective length is $\frac{1}{4}$ of a full circle with 50 mm radius of curvature, and thus, $I = \frac{1}{2}\pi r_{\rm C} = 79 \, {\rm mm}$. The I/d ratio is 79/25 = 3.16. Using Eq. (4.158) or Figure 4.37, the transmission probability P = 26.5%.

Case 2: A mitered elbow is used. The effective length of the tube is the sum of the side lengths, 100 mm. The I/d ratio is 100/25 = 4. As calculated with Eq. (4.158) or obtained from Figure 4.37, the transmission probability P = 22.4%. The precise result of Monte Carlo simulation (Figure 4.41) in the considered case (a/r = b/r = 50/12.5 = 4) is $P = 24 \pm 1\%$ (reading error of figure).

Considering conductance, the radius elbow is superior to the mitered elbow. Better than both of these components is a large sphere with two attached short tubes for the connections. It should be noted that the transmission probabilities given in the example correspond to the intrinsic conductance, that is, are valid for a component mounted directly between the two vessels where the gas particles approach isotropically. If the components are attached via stubs, as commonly used, other transmission probabilities result due to the developing tubular flow at the inlet to the components. This type of arrangement is covered in Section 4.4.8.

4.4.7

Series Connection of Tube and Aperture

In practice, tubes with different cross sections are often combined in a serial arrangement. A simple example is a series connection between two chambers, made up of a tube (cross-sectional area A_1) and an aperture (cross-sectional area A_2) (Figure 4.42). First, we will investigate molecular flow from the left-hand chamber Ch1 into the right-hand chamber Ch2.

The particle flow hitting the entrance plane A_1 from chamber Ch1 is q_N . The transmission probability for the tube shall be P. Then, a particle flow $q_N P$ reaches the area A_2 at the aperture. Here, a portion A_2/A_1 passes through the aperture and enters into chamber Ch2. The other portion, $1 - A_2/A_1$, is reflected backward, in the direction opposite to the inflow. Again, from the reflected gas particles, a portion P reaches A_1 and thus finds its way back into chamber Ch1 while the remaining portion 1 - P reaches A_2 . From this portion, a fraction (corresponding to the surface-area ratio A_2/A_1) enters into chamber Ch2 while the rest is reflected backward just as in the first reflection. The total particle flow leaving the aperture toward the right-hand side is the sum of the contributing particle flows, traveling the tube once, with an extra double passage, with two

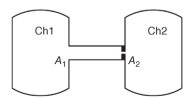


Figure 4.42 Two chambers connected by a series connection of a tube (cross-sectional area A_1) and an aperture (cross-sectional area A_2).

extra double passages, and so on. Adding up the geometric series yields the transmission probability P_{12} of the arrangement according to Figure 4.42 in the direction oriented from surface A_1 to surface A_2 :

$$\frac{1}{P_{12}} = \frac{1}{P} + \frac{A_1}{A_2} - 1. \tag{4.161}$$

Analogous treatment of a flow from chamber Ch2 to chamber Ch1 yields the transmission probability in the opposite direction, from 2 to 1:

$$\frac{1}{P_{21}} = \frac{A_2}{A_1} \cdot \frac{1}{P} - \frac{A_2}{A_1} + 1. \tag{4.162}$$

By eliminating the quantity P from the previous two equations, it follows that

$$A_2 P_{21} = A_1 P_{12}. (4.163)$$

The product of the area of the exit plane and the transmission probability is just proportional to the conductance. Thus, we find that the conductance from chamber Ch1 to chamber Ch2 is identical to the conductance in the opposite direction. This relationship is universally valid for any arrangement of passive line elements. It indicates that, under stationary conditions, the pressure among connected chambers with the same temperature is balanced.

4.4.8

Series Connection of Components

We will consider a series connection of several components with dissimilar cross sections and lengths. The components include tubes, apertures (tube of length zero), inserted chambers (tube with very large diameter), and other types of components such as bent tubes (Figure 4.43).

The net conductance cannot be obtained by treating the arrangement as a simple series connection of the individual components. The intrinsic conductance of a component applies to the case where the component lies between two large chambers. This quantity has two constituents: the inflow to the component and the tubular flow through the component. If the cross section of the line at the transition between two subsequent components remains unchanged or even

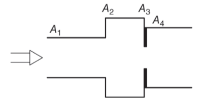


Figure 4.43 Series connection of four tubes with different cross-sectional areas A.

increases, the inflow loss disappears for the succeeding component. However, if the cross-sectional area of the line decreases at the transition from one component to the next, inflow loss occurs to an extent determined by the drop in the cross-sectional area. In the previous section, a descriptive example showed that, in a series connection of components, the transmission probability depends on the direction.

The sum of resistances due to inflow and tubular flow is a good approximation for the intrinsic flow resistance of an individual component. In a series connection of components, the total resistance is then given by the sum of individual resistances. However, if the cross-sectional area of the line at the intake of the component does not decrease, the resistance due to inflow is set to zero (Oatley's approach). Thus, the following addition theorem is obtained for a series connection of components:

$$\frac{1}{A_1} \left(\frac{1}{P_{1n}} - 1 \right) = \sum_{i=1}^n \frac{1}{A_i} \left(\frac{1}{P_i} - 1 \right) + \sum_{i=1}^{n-1} \left(\frac{1}{A_{i+1}} - \frac{1}{A_i} \right) \delta_{i,i+1}. \tag{4.164}$$

where i denotes the component number, A_1 is the cross-sectional area of intake to the arrangement, A_i is the cross-sectional area of component i, P_{1n} is the total intrinsic transmission probability for the arrangement, P_i is the intrinsic transmission probability for component i, $\delta_{i,i+1} = 1$ if $A_{i+1} < A_i$ (decreasing crosssectional area), and $\delta_{i,i+1} = 0$ if $A_{i+1} \ge A_i$ (without reduction in cross-sectional area).

Special case: Series connection of two components with cross-sectional areas A_1 and A_2 , and transmission probabilities P_1 and P_2 . From Eq. (4.164), it follows for the general case with arbitrary cross sections:

$$\frac{1}{P_{12}} = \frac{1}{P_1} + \frac{A_1}{A_2} \left(\frac{1}{P_2} - 1\right) + \left(\frac{A_1}{A_2} - 1\right) \delta_{1,2}. \tag{4.165}$$

Case 1: The cross-sectional areas of both tubes are the same, $A_1 = A_2$. Thus, $\delta_{1,2} = 0$ and

$$\frac{1}{p_{12}} = \frac{1}{p_1} + \frac{1}{p_2} - 1. \tag{4.166}$$

Case 2: The cross-sectional area increases, $A_2 > A_1$. Thus, $\delta_{1,2} = 0$ and

$$\frac{1}{P_{12}} = \frac{1}{P_1} + \frac{A_1}{A_2} \left(\frac{1}{P_2} - 1 \right). \tag{4.167}$$

Case 3: The cross-sectional area decreases, $A_2 < A_1$. Thus, $\delta_{1,2} = 1$ and

$$\frac{1}{P_{12}} = \frac{1}{P_1} + \frac{A_1}{A_2} \cdot \frac{1}{P_2} - 1. \tag{4.168}$$

Example 4.13

Two tubes with circular cross sections, equal diameter d, and lengths $l_1 = 2d$ and $I_2 = 3d$ are connected and thereby form a tube with length $I_{12} = 5d$.

With Eq. (4.158), the transmission probabilities of the two tubes are P_1 0.355 and $P_2 = 0.274$. If the transmission probability for the assembled tube is calculated by considering a simple series connection of conductances, as in electrical engineering, with the equation $1/P_{12} = 1/P_1 + 1/P_2$, the result is $P_{12} = 0.155$. If the transmission probability is calculated with the approximation formula for vacuum ducts in Eq. (4.166), $1/P_{12} = 1/P_1 + 1/P_2 - 1$, then $P_{12} = 0.183$.

For the assembled tube, the effective transmission probability, according to Eq. (4.158), is $P_{12} = 0.190$.

The given approximation formula for a series connection of flow conductances thus yields a considerably more accurate result compared with the simple equation for serially connected electrical conductances. The reason is that the flow at the outlet of the first tube is no longer isotropic, as it is when it enters the first tube. In fact, it is directed forward in the tube (beaming effect) and thus passes through the second tube more easily.

Example 4.14

Series connection of elements in a line.

Certain components such as valves feature a more complex geometry and their transmission probabilities are usually unknown. Manufacturers list the intrinsic conductance C in technical specifications from which the transmission probability can be calculated using $C = \frac{1}{4}\overline{C}AP$. If, furthermore, isothermal conditions are assumed (constant thermal velocity \bar{c} within the line), the conductance of the series connection can be calculated by rewriting Eq. (4.164):

$$\left(\frac{1}{C_{1n}} - \frac{4}{\overline{c}A_1}\right) = \sum_{i=1}^{n} \left(\frac{1}{C_i} - \frac{4}{\overline{c}A_i}\right) + 4\overline{c} \sum_{i=1}^{n-1} \left(\frac{1}{A_{i+1}} - \frac{1}{A_i}\right) \delta_{i,i+1}. \tag{4.169}$$

Molecular Flow through Conical Tube with Circular Cross Section (Funnel)

Figure 4.44 shows a conical tube (circular cross section) as is used, for example, for reducing adaptors between pump flanges and vacuum lines.

If the transition in cross section is smooth, the cone can be thought of as a series connection of short circular-cylindrical tube elements with changing

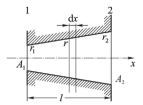


Figure 4.44 Conical tube.

diameters. According to Eq. (4.156), the inverse transmission probability $1/P_{\rm dx}$ of a short element of length dx is given by

$$\frac{1}{P_{\rm dx}} = \frac{1}{1 - (dx/2r)} \approx 1 + \frac{dx}{2r}.$$
 (4.170)

The latter approximation is correct, since $dx/2r \ll 1$. Analogous to Eq. (4.164), the transmission probability of the conical tube piece can be calculated by treating it as a series connection of tube elements and by integrating over the length of the tube (rather than summing up the contributions of the tube elements).

This yields the transmission probability in the direction of increasing diameter:

$$\frac{1}{P_{12}} = 1 + A_1 \int_{x_1}^{x_2} \frac{1}{A(x)} \cdot \frac{\mathrm{d}x}{2r} = 1 + \frac{r_1 + r_2}{4r_2^2} l, \tag{4.171}$$

and facing toward the direction of decreasing diameter,

$$\frac{1}{P_{21}} = 1 + A_2 \int_{x_2}^{x_1} \left(\frac{1}{A(x)} \cdot \frac{1}{2r} - \frac{1}{A^2(x)} \cdot \frac{dA}{dx} \right) dx = \frac{r_2^2}{r_1^2} + \frac{r_1 + r_2}{4r_1^2} l.$$
 (4.172)

By calculation, it is easy to verify Eq. (4.163) for a funnel; therefore, conductances for both directions of flow are the same as it has to be.

The given equations can be tested by equating the two radii of the conical tube, which then leads to the known case of a cylindrical tube with circular cross section. The obtained equation corresponds to the approximation formula in Eq. (4.154). This is because both equations are based on the approach that the transmission probability for an arbitrarily long tube can be calculated with the approximation of inversely summing up the probabilities of passage for an entry aperture and a very long tube. This approximation is correct for the special cases of very short and very long tubes. However, for $l/d \approx 2$, it produces a positive error of 13%.

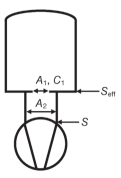


Figure 4.45 Pump evacuating a chamber through an aperture.

4.4.10

Component in the Inlet Line of a Pump

At low pressures (molecular regime), pumping speed S = dV/dt of many vacuum pumps is pressure independent. Therefore, as a model conception, a pump can be idealized as a series connection of a tube and an ideal pump with zero inlet pressure. The tube has the same intake area as the pump and its conductance is equal to the pumping speed of the pump (C = S).

If this type of pump connects to a chamber via a component with finite conductance, the effective pumping speed available at the chamber S_{eff} is lower than that of the pump (Figure 4.45). The effective pumping speed can be calculated roughly as a series connection between the component and the tube (pump model) according to Eq. (4.169).

Example 4.15

A turbomolecular pump with pumping speed $S = 100 \,\ell\,s^{-1}$ for air (flange diameter 70 mm) is mounted directly to the vacuum flange of a chamber. The effective pumping speed at the chamber shall be reduced to $S_{\rm eff} = 40 \, \ell \, \rm s^{-1}$ by insertion of an aperture into the chamber flange. What diameter does the aperture need to have?

Figure 4.45 shows the arrangement of chamber, aperture, and pump. Equation (4.169), rewritten to the given case (subscript 1 for the aperture, subscript 2 for the pump), reads

$$\left(\frac{1}{S_{\text{eff}}} - \frac{4}{\overline{c}A_1}\right) = \left(\frac{1}{C_1} - \frac{4}{\overline{c}A_1}\right) + \left(\frac{1}{S} - \frac{4}{\overline{c}A_2}\right). \tag{4.173}$$

Solving for the intrinsic conductance C_1 of the aperture yields

$$\frac{1}{C_1} = \frac{1}{S_{\text{eff}}} - \frac{1}{S} + \frac{4}{\overline{c}A_2}$$

$$= \frac{1}{40 \,\ell \, s^{-1}} - \frac{1}{100 \,\ell \, s^{-1}} + \frac{4}{4630 \,\text{dm} \, s^{-1} \times 0.385 \,\text{dm}^2}$$

$$= \frac{1}{58.0 \,\ell \, s^{-1}}.$$
(4.174)

From this, the aperture diameter is calculated:

$$d = \sqrt{\frac{16}{\pi} \cdot Cc} = \sqrt{\frac{16}{\pi} \cdot \frac{58.0 \,\ell \,\mathrm{s}^{-1}}{4630 \,\mathrm{dm} \,\mathrm{s}^{-1}}} = 0.253 \,\mathrm{dm} = 25.3 \,\mathrm{mm}. \tag{4.175}$$

If a simple series connection of aperture and pump would have been calculated as is common in electrical engineering, the following equation would have been found, rather than Eq. (4.174):

$$\frac{1}{C_1} = \frac{1}{S_{\text{eff}}} - \frac{1}{S},\tag{4.176}$$

which yields $C_1 = 66.7 \, \ell \, \mathrm{s}^{-1}$ and $d = 27.1 \, \mathrm{mm}$. The simple calculation neglects the preferential forward gas motion caused by the entrance aperture (beaming effect, Figure 4.35).

4.5 Flow throughout the Entire Pressure Range

4.5.1

Flow Ranges

In practice, one often requires the flow conductance of a component throughout the entire range from molecular to viscous gas flow. Also, it may happen that the pressure is high enough for flow to be viscous at the inlet of a tube while the pressure drops toward the end of the tube to an extent that flow here becomes molecular. The previous sections cover the two special cases of molecular and viscous flows. In the transition range, flow behavior is complicated and defies precise calculation. However, instructive experimental data and approximation formulas are available.

4.5.2

Flow through a Thin Aperture with Circular Cross Section

We will consider the ideal case of a very thin circular aperture (tube of length $l \ll \text{diameter } d$). The aperture is arranged between two chambers, with pressure

Table 4.6 Flow function ψ^* for choked flow, Eq. (4.50), and ratio of conductances in the viscous, Eq. (4.177), and molecular regimes, Eq. (4.143), for flow through an aperture with exit pressure low compared with inlet pressure.

	Monatomic gases: noble gases, metal vapors	Diatomic gases: air and so on	Buckled three- atom gases, for example, water	Polyatomic gases, for example, oil vapor
ψ^*	0.513	0.484	0.476	0.444
$C_{ m viscous}/C_{ m molecular}$	1.56	1.48	1.45	1.35

 p_1 in one chamber and negligible pressure p_2 ($p_2 \ll p_1$) in the other chamber. In the case of molecular flow, conductance

$$C_{\text{molecular}} = \frac{1}{4} \overline{c} A. \tag{4.143}$$

Under viscous flow conditions, flow is determined by gas dynamics. Choked flow develops if the outlet pressure is low. Conductance is derived from Eq. (4.27) with Eq. (4.59), taking into account the contraction of flow (Eq. (4.60)):

$$C_{\text{viscous}} = 0.86 \sqrt{\frac{\pi}{4}} \overline{c} A \Psi^*. \tag{4.177}$$

Table 4.6 lists values of the flow function Ψ^* for choked flow as well as the ratio of conductances in the viscous and molecular ranges for selected gas species.

The conductance of the thin aperture shows a change of only approximately 50% across the entire range of flow. Experimental data of aperture conductances for different gas species are compiled in Figure 4.46. In Figure 4.47, the same

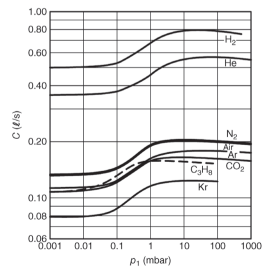


Figure 4.46 Measured flow conductances of a thin aperture with a diameter of 1.22 mm for selected gas species at room temperature. Inlet pressure is taken as abscissa and the outlet pressure is negligible.

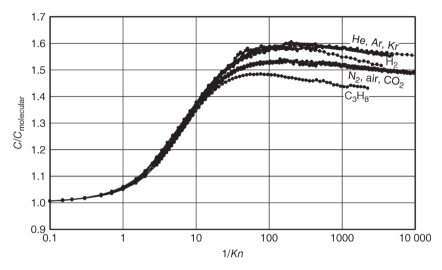


Figure 4.47 Flow conductances of a thin aperture. Same data as in Figure 4.46 but plotted at normalized abscissa and normalized ordinate.

data are plotted in normalized form. For normalizing, conductance is divided by the conductance under molecular flow conditions. In addition, rather than pressure, the inverse Knudsen number was used for the abscissa. This was calculated according to Eq. (4.2), with pressure being the inlet pressure p_1 (rather than average pressure).

Comparison shows that theoretical predictions (Table 4.6) are in good compliance with experimental data in the viscous and molecular ranges (Figure 4.47). Remarkably, conductance passes through a maximum for Knudsen number $Kn^{-1} \approx 100$, that is, at the upper end of the transition range, rather than increasing monotonically from molecular to viscous values. This is because flow is mostly viscous at this point but the density of the gas is not sufficient for full contraction (assumed value 0.86) to develop.

Figure 4.47 confirms the assumption in Eq. (4.3) that the flow in an aperture is molecular for $Kn^{-1} < 2$ and viscous for $Kn^{-1} > 100$.

Sharipov [5,6] calculated conductances for thin apertures using the DSCM (direct simulation Monte Carlo) method in the range $Kn^{-1} = 0-1000$, which agree reasonably with experiment.

Example 4.16

We will investigate airflow at 20 °C through a thin aperture. According to Eq. (4.144), the volumetric flow density under molecular flow conditions is

$$j_V = 11.6 \,\ell\,\text{s}^{-1}\,\text{cm}^{-2},$$
 (4.178)

and under gas-dynamic conditions (choked flow) is

$$j_V = \frac{C_{\text{viscous}}}{A} = 0.86 \sqrt{\frac{\pi}{4}} 463 \text{ m s}^{-1} \times 0.484 = 17.1 \,\ell\text{ s}^{-1} \text{ cm}^{-2}.$$
 (4.179)

The change in conductance of the aperture is low throughout the entire range of flow regimes. Under molecular flow conditions, a gas particle passes through the aperture only if it strikes the inlet by chance. Here, conductance (for air) is approximately 73% of the value under viscous flow conditions where gas particles move through the aperture collectively from the space in front of the aperture.

4.5.3

Flow through a Long Tube with Circular Cross Section

In a long tube, that is, length $l \gg$ (hydraulic) diameter d, the conductance $C_{\text{molecular}}$ in the molecular range is independent of pressure whereas conductance C_{viscous} in the viscous range increases with rising pressure. Thus, for sufficiently low pressures (molecular regime), $C_{\text{viscous}} \ll C_{\text{molecular}}$, and for sufficiently high pressures (viscous regime), $C_{\text{molecular}} \ll C_{\text{viscous}}$. Two tubes placed in parallel, one with molecular conductance and the other with viscous conductance, represent a good approximation for the conductance of a long tube. This can be expressed as

$$C \approx C_{\text{molecular}} + C_{\text{viscous}}.$$
 (4.180)

This equation correctly describes the two special cases of molecular and viscous flows, and represents a good approximation for the transition range.

We will now consider a long tube with circular cross section. The conductance $C_{\text{molecular}}$ in the molecular flow range is given by Eq. (4.141) with Eq. (4.157). If a tube is sufficiently long, flow in the viscous range is laminar. The corresponding conductance C_{viscous} was also given previously (Eq. (4.83)). Summing up the two equations according to Eq. (4.180) yields

$$C = \frac{\pi}{12} \left(\frac{3}{32} \cdot \frac{\overline{p}d}{\eta \overline{c}} + Z \right) \frac{d^3}{l} \overline{c}. \tag{4.181}$$

Here, η is the viscosity of the gas, \bar{c} is the mean thermal speed of the particles, and $\overline{p} = (p_1 + p_2)/2$, the arithmetical mean of the pressures at the inlet and outlet. The dimensionless, bracketed term in Eq. (4.181) is referred to as the conductance function *f* :

$$f = \frac{3}{32} \cdot \frac{\overline{p}d}{\eta \overline{c}} + Z = \frac{3\pi}{128} \cdot \frac{1}{Kn} + Z. \tag{4.182}$$

The Knudsen number, Eq. (4.2), used here has to be calculated for mean pressure \overline{p} .

Calculation shows that the term Z is just equal to 1. Then, the conduction function f has the value 1 in the molecular regime and increases with increasing pressure (see curve $f_{Z=1}$ in Figure 4.48). Knudsen (1909) performed experimental

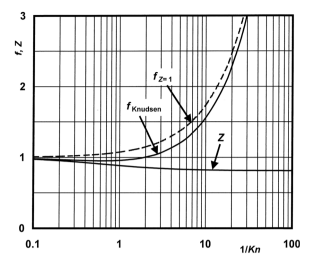


Figure 4.48 Calculated conductance function f, Eq. (4.182), and term Z, Eq. (4.183), versus inverse Knudsen number.

investigations and observed a dip of the conductance function f. It has the value of 1 in the molecular regime, then drops slightly with rising pressure until it reaches a minimum of 0.952 at $Kn^{-1} \approx 0.6$, and then rises rapidly (see curve $f_{\rm Knudsen}$ in Figure 4.48). Thus, flow conductance shows a slight minimum as well.

In order to describe this tube, he introduced a semi-empirical expression for the dimensionless term *Z*:

$$Z = \frac{1 + \sqrt{8/\pi(\bar{p}d/\bar{c}\eta)}}{1 + (21/17)\sqrt{8/\pi(\bar{p}d/\bar{c}\eta)}} = \frac{1 + (1.28/Kn)}{1 + (1.58/Kn)}.$$
 (4.183)

Both term Z and the conductance function f only depend on the Knudsen number *Kn* and thus apply to all gas species.

For a tube, Figure 4.48 verifies the assumption made in Eq. (4.3) that flow is molecular for $Kn^{-1} < 2$ and viscous for $Kn^{-1} > 100$. pV flow through the tube is calculated using conductance, Eq. (4.181):

$$q_{pV} = \frac{\pi}{128} \cdot \frac{1}{\eta} \cdot \frac{d^4}{l} \cdot \frac{p_1^2 - p_2^2}{2} + Z \frac{\pi}{12} \overline{c} \frac{d^3}{l} (p_1 - p_2). \tag{4.184}$$

Numeric-value variants of the previous equations are listed here for air at 20 °C. Units used are mbar for \overline{p} , p_1 , and p_2 , cm for d and l, and mbar ℓ s⁻¹ for q_{nV} :

$$f(\overline{p}d) = 11.1\overline{p}d + \frac{1 + 189\overline{p}d}{1 + 234\overline{p}d},\tag{4.185}$$

$$q_{pV} = 12.1 \frac{d^3}{l} f(\overline{p}d)(p_1 - p_2)$$

$$= \left(135 \frac{d^4}{l} \overline{p} + 12.1 \frac{d^3}{l} \cdot \frac{1 + 189 \overline{p}d}{1 + 234 \overline{p}d}\right) (p_1 - p_2).$$
(4.186)

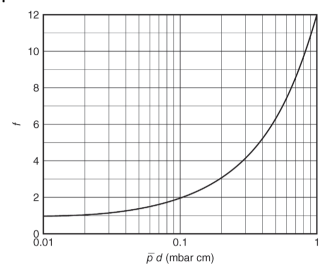


Figure 4.49 Conductance function $f(\overline{p}d)$ according to Eq. (4.185) for air at 20 °C.

The large bracketed term is the conductance of the tube. The conductance function for air at 20 °C is plotted in Figure 4.49. It features the following limiting values (\overline{p} in mbar, d in cm):

$$f(\overline{p} d) = 1$$
 for molecular flow, i.e., $\overline{p} d < 0.01$ mbar cm, (4.187)

$$f(\overline{p}d) = 0.86 + 11.1\overline{p}d$$
 for laminar viscous flow, i.e., $\overline{p}d$
> 0.66 mbar cm. (4.188)

These equations contain the mean pressure $\overline{p} = (p_1 + p_2)/2$. In certain cases, one of the two pressures p_1 and p_2 is unknown, and thus impedes calculation of \overline{p} . Here, an iterative method is applied: in the first step, we approximate $\overline{p} = p_1$ or $\overline{p} = p_2$, or we choose a reasonable value for \overline{p} . Then, calculation yields a value for p_1 or p_2 . With this, a new value for \overline{p} can be obtained, and so forth, until the desired accuracy is obtained.

Equation (4.186) includes the previously derived formulas for the pV flow of air at 20 °C in the special cases of laminar and molecular flows.

Example 4.17

A small Roots pump with a pumping speed $S = 40 \,\ell\,s^{-1}$, pressure independent throughout the operating range, is connected to a chamber via a tube of length $l=0.3\,\mathrm{m}$ and diameter $d=40\,\mathrm{mm}$. The pressure in the chamber is $p_\mathrm{C}=$ 0.02 mbar (see also Figure 4.29). The missing quantity is the effective pumping speed $S_{\rm eff}$.

 $p_{\rm C}$ is the chamber pressure and $p_{\rm in}$ is the pressure at the inlet flange of the pump. Equation (4.186) is used to solve the problem (q_{pV} in mbar ℓ s⁻¹, $p_{\rm in}$, $p_{\rm C}$, and \overline{p} in mbar, d and l in cm):

$$q_{pV} = p_{in}S = 12.1 \frac{d^3}{I} f(\overline{p}d)(p_C - p_{in}).$$
 (4.189)

Rewriting yields ($S = 40 \ \ell \ s^{-1}$, $I = 30 \ cm$, $d = 4 \ cm$)

$$\frac{p_{\rm C}}{p_{\rm in}} = 1 + \frac{SI}{12.1d^3 f(\overline{p}d)} = 1 + \frac{1.55}{f(\overline{p}d)}.$$
 (4.190)

The rest of the calculation is carried out iteratively. In the first step, we will estimate $p_{in} = 0.015$ mbar as the initial value.

This leads to $p_{\rm C}/p_{\rm in} = 0.02/0.015 = 1.33$ and $\overline{p} = \frac{1}{2}(0.02 + 0.015)\,{\rm mbar} =$ 0.0175 mbar.

Thus, $\overline{p}d = 0.07$ mbar cm. From this, Eq. (4.185) or Figure 4.49 yields f = 1.60. Using this value, $p_{\rm C}/p_{\rm in}=1.97$ is obtained from Eq. (4.190), a considerable deviation from the initial assumption $p_{\rm C}/p_{\rm in}=1.33$.

In the second step, we use the result obtained in step 1 to calculate p_{in} : $p_{\rm in} = p_{\rm C}/(p_{\rm C}/p_{\rm in}) = 0.02 \, {\rm mbar}/1.97 = 0.0102 \, {\rm mbar}.$

It follows that $\bar{p} = \frac{1}{2}(0.02 + 0.0102) \text{ mbar} = 0.0151 \text{ mbar}, \ \bar{p}d = 0.0604 \text{ mbar cm},$ and f = 1.50.

With this value, the result $p_{\rm C}/p_{\rm in}=2.03$ is obtained, which is already close to the initial value of the pressure ratio in the second step.

We can use an additional third step: $p_{in} = p_C/(p_C/p_{in}) = 0.02 \,\text{mbar}/2.03 =$ 0.0099 mbar.

Thus, $\overline{p} = \frac{1}{2}(0.02 + 0.0099)$ mbar = 0.0150 mbar, $\overline{p}d = 0.060$ mbar cm, f = 1.49, and $p_{\rm C}/p_{\rm in} = 2.04$.

It follows that the effective pumping speed according to Eq. (4.31) is $S_C = S_{in}p_C/p_{in} = 40 \,\ell \, s^{-1}/2.04 = 19.6 \,\ell \, s^{-1}$.

In the boundary case of viscous laminar flow, the quantity $\overline{p}d$ reaches high values and the previously stated result is obtained (numeric-quantity equation, \overline{p} , p_1 , and p_2 in mbar, d and l in cm, q_{pV} in mbar ℓ s⁻¹):

$$q_{pV} = 135 \frac{d^4}{l} \overline{p}(p_1 - p_2)$$
, viscous laminar airflow. (4.94)

For the special case of molecular flow, the quantity \overline{p} d in Eq. (4.186) is very low and the result is equal to that of Eqs. (4.145) and (4.157):

$$q_{pV} = 12.1 \frac{d^3}{I} (p_1 - p_2),$$
 molecular airflow. (4.91)

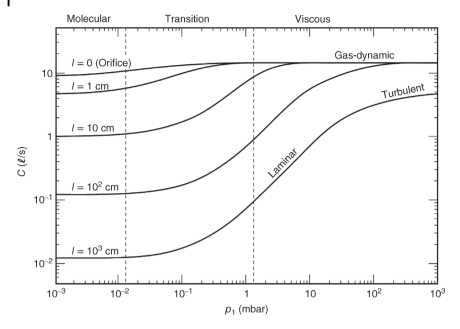


Figure 4.50 Flow conductance of tubes with circular cross sections, diameter d = 1 cm, and selected lengths for air at 20 °C. Inlet pressure p_1 is taken as abscissa and outlet

pressure p_2 is assumed negligible. Molecular and viscous regions were calculated, and in the transition region, freehand interpolation was used.

Finally, Figure 4.50 shows the flow conductance of tubes with circular cross section for air.

4.6 Flow with Temperature Difference, Thermal Effusion, and Transpiration

The previous sections covered isothermal systems with temperature T. Analogous to Figure 4.34, we will now consider two chambers C1 and C2 connected by a tube T (diameter d, length l) or an aperture (tube of length zero). The wall temperatures T_1 and T_2 of the chambers are unequal. Chambers C_1 and C_2 contain gas with the state quantities p_1 , n_1 , T_1 and p_2 , n_2 , T_2 , respectively (Figure 4.51).

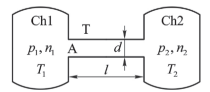


Figure 4.51 Two connected chambers showing unequal temperatures T_1 and T_2 .

For the following, we will assume a stationary condition of equilibrium in which the net gas flow between the chambers is zero. Different states of equilibrium develop, determined by the pressures.

For high pressures, the gas particles' mean free path \bar{l} is short compared with the diameter of the aperture or the tube; thus, $Kn = \bar{l}/d \ll 1$. The gas particles collide frequently and therefore a pressure difference would directly cause a net particle flow. Thus, in the stationary case,

$$p_2 = p_1$$
 , in the viscous range. (4.192)

For *small* pressures, $Kn = \overline{l}/d > 1$. The particle flow from chamber Ch1 to chamber Ch2 is just the product of the collision rate j_{N1} in chamber Ch1, area A of the tube, and the transmission probability P_{12} from chamber Ch1 to chamber Ch2. The particle flow in the reverse direction is calculated analogously. Thus, as previously stated, the total net particle flow (effusion rate) is

$$q_N = \frac{1}{4}n_1\bar{c}_1AP_{12} - \frac{1}{4}n_2\bar{c}_2AP_{21}. \tag{4.137}$$

In the stationary case, $q_N = 0$. In addition, the probabilities of passage from chamber Ch1 to chamber Ch2 and vice versa are equal $(P_{12} = P_{21})$ as long as wall reflections of gas particles are temperature independent, which is assumed here. As the particle velocity \bar{c} is proportional to the square root of the thermodynamic temperature T, the particle number densities

$$n_1\sqrt{T_1} = n_2\sqrt{T_2}$$
, in the molecular range. (4.193)

Using the equation of state for an ideal gas, p = nkT, the ratio of the pressures in both chambers amounts to

$$\frac{p_2}{p_1} = \sqrt{\frac{T_2}{T_1}} \quad , \quad \text{in the molecular range.} \tag{4.194}$$

Thus, under molecular flow conditions, the particle density is lower in the warmer chamber than it is in the cooler chamber (because warmer gas particles escape more easily). However, pressure is higher in the warmer chamber (because the gas particles move faster, and thus collide with the walls more frequently and more violently).

The phenomenon of unequal pressures in connected chambers at different temperatures is referred to as thermal effusion, or more commonly, thermal transpiration.

Example 4.18

Capacitance diaphragm manometers are used as sensitive vacuum gauges that directly measure the pressure acting upon a diaphragm. The temperature of the diaphragm sensor can be raised and controlled thermostatically in order to improve measuring accuracy or to prevent gas condensation. Under viscous flow conditions, pressures in the sensor (chamber C₁) and the vacuum chamber (chamber C₂) are equal. However, in the molecular range, the pressure in the sensor (chamber C_1) is higher than that in the vacuum chamber (chamber C_2) as described by Eq. (4.194). Thus, the pressure registered by the gauge is too high.

At an ambient temperature of 23 °C and a sensor temperature of 45 °C, the pressure ratio

$$\frac{p_2}{p_1} = \sqrt{318.2/296.2} = 1.036$$
, in the molecular range.

Figure 4.52 shows the measured characteristics of a heated (temperature-controlled) diaphragm manometer for different gas species. Helium has the smallest gas particles of all gas species. A molecule of the refrigerant R12 (CCl₂F₂) has a gas-kinetic impact area 8.3 times as large as helium. The data of different gas species coincide fairly well if the pressure axis is scaled to the inverse Knudsen number (Figure 4.53).

An empirical function found by Takaishi and Sensui [7] describes the measured characteristics. Setina [8] determined the coefficients universally for all gas species and formulated

$$\frac{p_2}{p_1} = 1 + \frac{\sqrt{T_2/T_1} - 1}{0.0181 \, Kn^{-2} + 0.229 \, Kn^{-1} + 0.211 \, Kn^{-1/2} + 1},\tag{4.195}$$

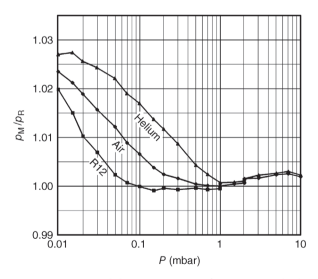


Figure 4.52 Measured characteristic curve of a capacitance diaphragm vacuum gauge heated to approximately 45 °C at an ambient temperature of 23 °C. The plot shows the measured ratio of pressures at the warm internal sensor and at the cold connecting flange.

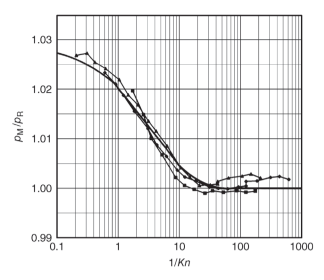


Figure 4.53 Characteristic curves of a thermostatically controlled capacitance diaphragm vacuum gauge. Same measured data as in Figure 4.52 but with normalized abscissa. The thick continuous line is the empirical curve according to Eq. (4.195).

with the pressure dependence being in the Knudsen number:

$$Kn = \frac{\pi}{4} \cdot \frac{\eta \, \overline{c}}{p \, d}.\tag{4.2}$$

 η and \overline{c} denote the viscosity and particle velocity of the gas, respectively, and d is the diameter of the diaphragm manometer's joining pipe (typically, $d = 3/16'' = 4.76 \,\mathrm{mm}$).

The thermal effusion is treated theoretically for various conditions in Section 5.5.7.

Example 4.19

In cryotechnology, pressures are often not measured directly at the low-temperature sample but rather by using a measuring instrument at ambient temperature, which is connected to the sample via a line. The measured pressure is a multiple of the actual pressure if the sample is at the temperature of liquid helium ($T_1 = 4.2 \text{ K}$) and the measuring equipment is at ambient temperature $(T_2 = 300 \text{ K})$:

$$\frac{p_2}{p_1} = \sqrt{\frac{300 \text{ K}}{4.2 \text{ K}}}$$
= 8.5. (4.196)

Example 4.20

At different chamber temperatures, gas particles have unequal velocities and are reflected slightly differently from the walls of a tube. As a result, the probabilities of passage P_{12} and P_{21} in the two directions of a tube are different to some extent, depending on the surface of the tube walls. Thus, if several chambers with, in turns, ambient and cryo temperature are connected in a series via tubes with alternating surface characteristics, a large pressure ratio between the first and the last chamber can be obtained under molecular flow conditions [9].

4.7

Measuring Flow Conductances

4.7.1

Necessity of Measurement

As shown in previous sections, flow conductances of vacuum components with simple geometries can be calculated reliably. The flow conductance of complicated components, especially in the transition range of different types of flow, can only be estimated. In these cases, measurements are indispensable.

The flow conductance of a component is defined by

$$C = \frac{q_{pV}}{p_1 - p_2}. (4.27)$$

 q_{pV} denotes the throughput (or pV flow) of the component and p_1-p_2 is the pressure difference between both ends of the component. In vacuum technology, the mounting position is crucial because entrance effects occur throughout the range from molecular to viscous flow. Therefore, intrinsic and reduced conductances are differentiated. The intrinsic conductance arises when the component is mounted between two large chambers so that the gas streams into the component from a chamber. In contrast, the reduced conductance is effective when the component is placed into a tube having the same cross section as the component.

The effects due to entrance under molecular flow conditions are discussed above. For this, a tube was treated as a series connection of an aperture and a long tube (Section 4.4.7). Intrinsic conductance C_{int} is obtained from the conductance of the aperture C_A (describing the inflow effect) and the reduced conductance C_{red} of the long tube (pure tubular flow):

$$\frac{1}{C_{\rm int}} \approx \frac{1}{C_{\rm A}} + \frac{1}{C_{\rm red}}.\tag{4.197}$$

4.7.2

Measurement of Intrinsic Conductances (Inherent Conductances)

In order to measure the intrinsic conductance, the investigated component is placed between two large chambers. Then, a known flow of gas q_{pV} is supplied and the pressure difference between the two chambers is measured.

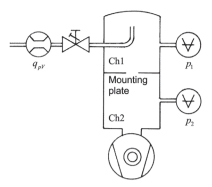


Figure 4.54 Measuring setup for determining the characteristic flow conductance of a component.

As apparent when studying technical literature, arrangements for measuring intrinsic conductances vary considerably. Direct mounting an inspected component between two chambers would require time-consuming welding work on every single component. Mounting the component with adapters on the chambers would disturb the flow. More suitable is an arrangement where the part is fixed to a mounting plate on one side. This assembly is then embedded in a divided measurement dome, similar to those used for pumping speed measurements on ultrahigh-vacuum pumps (DIN 28428, DIN 28429, ISO 5302) (Figure 4.54).

The gas flows through a flowmeter (e.g., thermoelectric flowmeter) and through an inlet against the lid of the upper chamber Ch1, thus featuring a practically isotropic distribution of particle velocities. The center of the measurement chamber contains a (replaceable) mounting plate carrying the investigated vacuum component that is fixed to the plate vacuum tightly. A turbomolecular pump evacuates the lower chamber C2. Pressures in both chambers are recorded. Calibrated vacuum gauges (viscosity vacuum gauges, diaphragm manometers) yield measuring uncertainties below 1%. The systematic uncertainty of conductance measurements depends on the size of the component's inlet opening compared with the inner surface of the chamber. This type of arrangement can be tested by measuring conductance values for simple components (e.g., tubes with circular cross sections), for which theoretical conductance values are available.

Measuring conductances of larger components under molecular flow conditions (e.g., valves with connecting flanges of 100 mm diameter and more) requires measurement domes of considerable size, and thus leads to an uneconomical measuring routine. The problem is solved by measuring the conductance of a scaled-down model. In the range of molecular flow, the conductance C of a component scales with the cross-sectional area of the line because the transmission probability P of the scaled-down model is equal to the original part's transmission probability.

4.7.3

Calculation of Reduced Conductance (Assembly Conductance)

Valves are important components in vacuum systems. Therefore, knowing their precise conductance as a function of pressure is crucial for system design. Open ball valves and gate valves can be treated as short tubes. However, valves with a more complex structure (shape of valve housing, elements inside the housing for valve actuation, gap-type opening between sealing plate and housing), for example, right-angle valves, call for conductance measurements. As valves are usually mounted to a line or flanges, the relevant quantity here is the assembly conductance (reduced conductance).

As an example, we will consider a commercial right-angle valve of nominal diameter DN 40 (actual diameter 41 mm). The apparatus described in the previous section yielded a measured intrinsic conductance of 28 ℓ s⁻¹ for molecular flow and ambient air. The conductance of an aperture with the diameter of the flange is $153 \, \ell \, \mathrm{s}^{-1}$. For the reduced conductance (assembly conductance), Eq. (4.197) yields

$$\frac{1}{C_{\text{red}}} \approx 1 C_{\text{int}} - \frac{1}{C_{\text{A}}} = \frac{1}{28 \,\ell \, \text{s}^{-1}} - \frac{1}{153 \,\ell \, \text{s}^{-1}} = \frac{1}{34 \,\ell \, \text{s}^{-1}}.$$
 (4.198)

This value is close to the reduced conductance $32 \ell s^{-1}$, measured directly at the valve.

4.7.4

Measuring Reduced Conductances

The setup shown in Figure 4.55 is appropriate for measuring the reduced conductances of components throughout the entire range of flow.

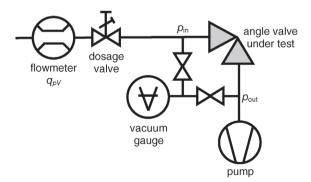


Figure 4.55 Setup for measuring the reduced flow conductance of right-angle valve (shaded) under any flow regime.

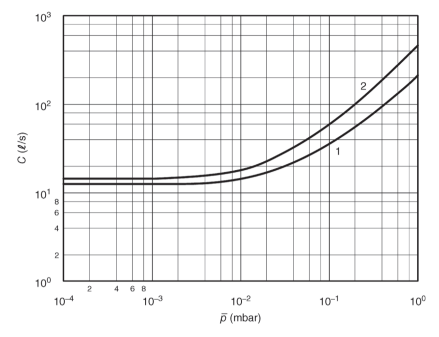


Figure 4.56 Reduced (assembly) conductance of a right-angle valve of nominal diameter 25 mm with length of buckled axis: 100 mm. Curve 1 shows the measured values for the

valve, and curve 2 shows the calculated conductance according to Eq. (4.197) for a tube of the same axis length.

The test gas flows into the apparatus through a flowmeter (pV flow). Downstream, a regulating valve is used to adjust the flow. The subsequent tube moderates the gas flow (to avoid entrance effects) and is made up of a tube with the same nominal diameter as the measured component. Here, the test object is a right-angle valve. A single vacuum gauge is used to measure the pressures p_1 and p_2 in front of and behind the test object. For this, it is equipped with valves that allow alternate connections to the two metering points. The usage of a single gauge compared with two gauges has the advantage of avoiding errors caused by differences between two gauges in addition to that of saving one gauge.

As an example, Figure 4.56 illustrates a measuring curve for the reduced conductance of a right-angle valve of nominal diameter DN 25 as a function of mean pressure $\overline{p} = \frac{1}{2}(p_1 + p_2)$.

The effective length of the axis in the corner valve is 100 mm. For comparison, the figure contains the reduced conductance of a tube of 100 mm length (calculated using Eq. (4.181)). Obviously, the experimental results are entirely below the calculated values because of the small cross section inside the valve.

References

- 1 Schade, H. and Kunz, E. (1989) Strömungslehre, Walter de Gruyter, Berlin.
- 2 Bohl, W. (1991) Technische Strömungslehre, Vogel, Würzburg.
- 3 Bird, G.A. (1994) Molecular Gas Dynamics and the Direct Simulation of Gas Flows, Oxford University Press, Oxford.
- 4 Lafferty, J.M. (1998) Foundations of Vacuum Science and Technology, John Wiley & Sons, Inc., New York, Chapter 2.
- 5 Sharipov, F. and Seleznev, V. (1998) J. Phys. Chem. Ref. Data, 27, 657-705.
- 6 Sharipov, F. (2004) J. Fluid Mech., 518, 35-60.
- 7 Takaishi, T. and Sensui, Y. (1963) Trans. Faraday Soc., 59, 2503-2515.
- 8 Setina, J. (1999) Metrologia, 36, 623-626.
- 9 Hobson, J.P. (1969) J. Vac. Sci. Technol., 7, 351-357.

Further Reading

Comprehensive description of gas flow

Jitschin, W. (1999) Vakuum-Lexikon, Wiley-VCH Verlag GmbH, Weinheim.

O'Hanlon, J.F. (2003) A User's Guide to Vacuum Technology, 3rd edn, John Wiley & Sons, Inc., New York, Chapter 3 and Appendix B4.

Gas-dynamic flow

Becker, E. (1965) Gasdynamik, Teubner, Stuttgart.

Sauer, R. (1960) Einführung in die theoretische Gasdynamik, Springer, Berlin.

Molecular flow

Berman, A. (1992) Vacuum Engineering Calculations, Formulas, and Solved Exercises, Academic Press, San Diego, CA.

Saksaganskii, G.L. (1988) Molecular Flow in Complex Vacuum Systems, Gordon and Breach Science Publishers.

Steckelmacher, W. (1986) Rep. Prog. Phys., 49, 1083-1107.

5

Analytical and Numerical Calculations of Rarefied Gas Flows

Prof. Felix Sharipov

Universidade Federal do Parana, Departamento de Fisica, Caixa Postal 19044, 81531-990, Curitiba. Brasil

The aim of this chapter is to detail the concepts described in Chapters 3 and 4 and to describe the main results of rarefied gas flow calculations based on the kinetic Boltzmann equation. A simple and accessible presentation of results without a hard mathematical formalism shall enable physicists and engineers to understand and simulate rarefied gas flows over the whole range of the Knudsen number.

5.1

Main Concepts

5.1.1

Knudsen Number and Gas Rarefaction

The mean free path \overline{l} introduced previously for hard-sphere molecules depends on the molecular diameter, which is usually unknown and calculated via the shear viscosity η . Moreover, the mean free path expression will be different if one assumes another potential of the intermolecular interaction. To avoid such an ambiguity, here the equivalent free path ℓ is introduced, which is related directly to the shear viscosity as

$$\ell := \frac{\eta c_{\rm mp}}{p},\tag{5.1}$$

where $c_{\rm mp}$ is the most probable molecular speed defined by Eq. (3.42) and p is the gaseous pressure.

The gas rarefaction parameter defined as

$$\delta := \frac{a}{\ell} = \frac{ap}{\eta c_{\rm mp}} \tag{5.2}$$

is frequently used instead of the Knudsen number. Here a is a characteristic size of a duct in which the gas flows. In the literature, most numerical results are given in terms of this parameter. The limit of large values, that is, $\delta \gg 1$, corresponds to the viscous regime, while the opposite limit, that is, $\delta \ll 1$, represents the free molecular condition. Thus, the rarefaction parameter is inversely proportional to the Knudsen number.

5.1.2

Macroscopic Quantities

Besides the previously introduced quantities such as number density n, pressure p, temperature T, and bulk velocity \mathbf{v} , here some additional macroscopic quantities will be used. Note that the term "bulk velocity" is used in order to distinguish it from the velocity of gaseous molecules.

Consider an area segment A_x with a normal directed along the x-axis. Let F_z be a force acting in the z-direction. Then the quantity defined as

$$P_{xz} := \frac{F_z}{A_x} \tag{5.3}$$

is called the shear stress. It has the same dimension as a pressure.

If \dot{Q}_x is a magnitude of heat crossing the area segment A_x per unit time, then the quantity

$$q_x = \frac{\dot{Q}_x}{A_x} \tag{5.4}$$

is the x component of the heat flux vector q. The dimension of this quantity is W m^{-2} .

5.1.3

Velocity Distribution Function

The velocity distribution function $f(\mathbf{r}, \mathbf{c})$ used in this chapter is assumed to be dependent on the spatial position vector r and on the molecular velocity vector c. Generally speaking, it also depends on the time, but here only time-independent flows will be considered so that the dependence on the time will be omitted. The distribution function is defined as

$$f(\mathbf{r}, \mathbf{c}) \coloneqq \frac{\mathrm{d}N}{\mathrm{d}^3 \mathbf{r} \mathrm{d}^3 \mathbf{c}},\tag{5.5}$$

where dN is the number of particles in the phase volume $d^3\mathbf{r}d^3\mathbf{c}$ near the point (\mathbf{r}, \mathbf{c}) . All macro-characteristics of gas flow can be calculated via the distribution function:

Number density

$$n(\mathbf{r}) = \int f(\mathbf{r}, \mathbf{c}) d^3 \mathbf{c}, \tag{5.6}$$

Bulk velocity

$$\mathbf{v}(\mathbf{r}) = \frac{1}{n} \int \mathbf{c} f(\mathbf{r}, \mathbf{c}) d^3 \mathbf{c}, \tag{5.7}$$

Shear stress

$$P_{xz}(\mathbf{r}) = m_{\rm P} \int (c_x - \nu_x)(c_z - \nu_z) f(\mathbf{r}, \mathbf{c}) \mathrm{d}^3 \mathbf{c}, \qquad (5.8)$$

Temperature

$$T(\mathbf{r}) = \frac{m_{\rm P}}{3nk} \left[(\mathbf{c} - \mathbf{v})^2 f(\mathbf{r}, \mathbf{c}) d^3 \mathbf{c}, \right]$$
 (5.9)

Heat flux

$$q_x(\mathbf{r}) = \frac{m_P}{2} \left[(\mathbf{c} - \mathbf{v})^2 (c_x - \nu_x) f(\mathbf{r}, \mathbf{c}) d^3 \mathbf{c}. \right]$$
 (5.10)

The integration in Eqs. (5.6)–(5.10) with respect to the molecular velocity means the threefold integral over the whole velocity space, that is,

$$\int \dots d^3 \mathbf{c} = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \dots dc_x dc_y dc_z.$$
 (5.11)

5.1.4

Global Equilibrium

Global equilibrium means that no macroscopic motion of one part of the system is relative to another, there is no heat exchange between different parts of the system, and no chemical reactions occur. In such a state, all macroscopic quantities (pressure, temperature, and concentrations) are constant over time and space. Under these conditions, the velocity distribution function is given by the absolute Maxwellian:

$$f^{\rm M}(n,T) = n \left(\frac{m_{\rm P}}{2\pi kT}\right)^{3/2} \exp\left(-\frac{m_{\rm P}c^2}{2kT}\right).$$
 (5.12)

Note that the function F_0 given by Eq. (3.41) characterizes the distribution of the molecular speed c, while the function $f^{\rm M}$ describes the distributions of the velocity components according to Eq. (5.5).

The mean (or thermal) molecular speed is calculated as

$$\bar{c} := \frac{1}{n} \int c f^{M} d^{3} \mathbf{c} = \sqrt{\frac{8kT}{\pi m_{P}}} = \frac{2}{\sqrt{\pi}} c_{mp}.$$
 (5.13)

To calculate the wall flux density or the number of particles striking a unit surface per unit time, we consider a plane surface fixed at x = 0 and a gas occupying the space $x \le 0$. Then, the flux of particles to the surface is calculated as

$$j_{\rm N} = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \int_{0}^{\infty} c_x f^{\rm M} dc_x dc_y dc_z = \frac{1}{2} n \sqrt{\frac{2kT}{\pi m_{\rm P}}} = \frac{1}{4} n \overline{c}, \tag{5.14}$$

where Eq. (5.12) with $c^2 = c_x^2 + c_y^2 + c_z^2$ and Eq. (5.13) have been used.

5.1.5

Local Equilibrium

Let us divide a system into many small subsystems so that each of them still contains a large number of molecules. Each subsystem can stay in equilibrium, but the concentration n and temperature T can slowly vary from one subsystem to another. Moreover, each subsystem can move with its one bulk velocity v. Such a state is called local equilibrium. It occurs in the viscous regime, that is, when the Knudsen number is small. In this case, we have the local Maxwellian distribution function determined by the local values of n, T, and \mathbf{v} and denoted as

$$f^{\mathrm{M}}(n,T,\mathbf{v}) = n(\mathbf{r}) \left[\frac{m_{\mathrm{P}}}{2\pi k T(\mathbf{r})} \right]^{3/2} \exp\left\{ -\frac{m_{\mathrm{P}}[\mathbf{c} - \mathbf{v}(\mathbf{r})]^{2}}{2k T(\mathbf{r})} \right\}.$$
(5.15)

However, in the transitional and free molecular regimes, the local equilibrium is broken, and then the distribution function is calculated from the kinetic Boltzmann equation.

5.1.6

Boltzmann Equation

The Boltzmann equation determines the evolution of the distribution function. The main idea of its derivations consists of the fact that at a fixed point of the physical space, the distribution function varies due to the intermolecular collisions, that is, its total derivative is given as

$$\frac{\mathrm{d}f}{\mathrm{d}t} = J^+ - J^-,\tag{5.16}$$

where J^+ is the rate of gain of particles in the velocity space near the point c due to the collisions and J^- is the rate of the loss of particles at the same point. The total derivative can be written as

$$\frac{\mathrm{d}f}{\mathrm{d}t} = \frac{\partial f}{\partial t} + \mathbf{c} \cdot \frac{\partial f}{\partial \mathbf{r}} + \frac{\partial \mathbf{c}}{\partial t} \cdot \frac{\partial f}{\partial \mathbf{c}}.$$
(5.17)

Note that the second term on the right-hand side means the scalar product, that is,

$$\mathbf{c} \cdot \frac{\partial f}{\partial \mathbf{r}} = c_x \frac{\partial f}{\partial x} + c_y \frac{\partial f}{\partial y} + c_z \frac{\partial f}{\partial z}.$$
 (5.18)

The third term on the right-hand side of Eq. (5.17) is calculated similarly.

The rates J^+ and J^- are calculated regarding the details of binary collisions. For our purpose, it is enough to consider the Boltzmann equation in the absence of external forces, that is, $\partial \mathbf{c}/\partial t = 0$, and under the stationary conditions, that is, $\partial f/\partial t=0.$

The stationary flow means that all macroscopic quantities (pressure, temperature, concentrations, bulk velocity, shear stress, and heat flow vector) are time independent, but they are functions of the space coordinates.

Finally, the stationary Boltzmann equation reads

$$\mathbf{c} \cdot \frac{\partial f}{\partial \mathbf{r}} = J^{+} + J^{-} = J(ff_{*}), \tag{5.19}$$

where the collision integral $I(ff_*)$ takes the form

$$J(ff_*) = \iiint (f'f'_* - ff_*)wd^3\mathbf{c}'d^3\mathbf{c}'_*d^3\mathbf{c}_*.$$
(5.20)

Here, the affixes to f correspond to those of their arguments \mathbf{c} : $f' = f(\mathbf{c}')$, $f_* = f(\mathbf{c}_*)$. The quantity $w = w(\mathbf{c}, \mathbf{c}_*; \mathbf{c}', \mathbf{c}'_*)$ is the probability density that two molecules having the velocities \mathbf{c}' and \mathbf{c}'_* before a collision will have the velocities c and c*, respectively, after they collide. Its specific expression depends on the potential of the intermolecular interaction. The main properties of the collision integral $I(ff_*)$ can be found in Refs [1–6]. Note that Eq. (5.19) was obtained considering only binary intermolecular collisions, which is not valid at high pressures. However, for any pressure lower than or equal to atmospheric pressure, this assumption is well fulfilled.

In the case of weak nonequilibrium, the Boltzmann equation can be linearized representing the distribution function as

$$f(\mathbf{r}, \mathbf{c}) = f_{R}^{M} [1 + \xi h(\mathbf{r}, \mathbf{c})], \quad |\xi| \ll 1,$$
 (5.21)

where ξ is a small parameter, h is the perturbation function, and f_R^M is the reference Maxwellian given by Eq. (5.15) with the reference number density n_R , temperature T_R , and bulk velocity \mathbf{v}_R , that is,

$$f_{\mathbf{R}}^{\mathbf{M}} = f^{\mathbf{M}}(n_{\mathbf{R}}, T_{\mathbf{R}}, \mathbf{v}_{\mathbf{R}}). \tag{5.22}$$

The quantities n_R , T_R , and \mathbf{v}_R are chosen such that computational effort is reduced. Particularly, they can be constant and equal to their equilibrium values. Substituting (5.21) into (5.19), the linearized Boltzmann equation is obtained as

$$\mathbf{c} \cdot \frac{\partial h}{\partial \mathbf{r}} = \hat{L}h + g(\mathbf{r}, \mathbf{c}), \tag{5.23}$$

where \hat{L} is the linearized collision operator:

$$\hat{L}h = \iiint f_{R}^{M}(\mathbf{c}_{*})(h' + h'_{*} - h - h_{*})wd^{3}\mathbf{c}'d^{3}\mathbf{c}'_{*}d^{3}\mathbf{c}_{*}.$$
 (5.24)

Here h'_* , h', h_* , and h represent the perturbations of f'_* , f', f_* , and f, respectively. The nonequilibrium source function g is given as

$$g(\mathbf{r}, \mathbf{c}) = -\frac{\mathbf{c}}{\xi} \cdot \frac{\partial \ln f_{R}^{M}}{\partial \mathbf{r}}.$$
 (5.25)

If the reference quantities n_R , T_R , and \mathbf{v}_R are constant, then $g(\mathbf{r}, \mathbf{c}) = 0$.

Substituting Eq. (5.21) into Eqs. (5.7), (5.8), and (5.10), the linearized expressions of the moments are obtained:

$$\mathbf{v}(\mathbf{r}) - \mathbf{v}_{R} = \frac{\xi}{n_{R}} \int \mathbf{c} f_{R}^{M} h(\mathbf{r}, \mathbf{c}) d^{3} \mathbf{c},$$
 (5.26)

$$P_{xz}(\mathbf{r}) = \xi m_{\rm P} \int c_x c_z f_{\rm R}^{\rm M} h(\mathbf{r}, \mathbf{c}) \mathrm{d}^3 \mathbf{c}, \qquad (5.27)$$

$$q_x(\mathbf{r}) = \frac{\xi}{2} \int (m_P c^2 - 5kT_R) c_x f_R^M h(\mathbf{r}, \mathbf{c}) d^3 \mathbf{c}.$$
 (5.28)

5.1.7

Transport Coefficients

In this section, the definitions of the transport coefficients, that is, shear viscosity η and thermal conductivity λ , and main ideas of how to calculate them from the Boltzmann equation are described. It is necessary to emphasize that both shear viscosity n and thermal conductivity λ are defined so that they do not depend on the pressure of gas, but they depend only on the gas species and on its temperature. The explicit expressions of the coefficients will be given only for the hard-sphere model of intermolecular potential. For other kinds of potential, the transport coefficient expressions can be found in Refs [2,3].

Viscosity Coefficient

Let us consider a gas flow in a boundless region with the bulk velocity v given as

$$\mathbf{v} = [0, 0, \nu_z], \quad \nu_z \propto x, \tag{5.29}$$

that is, the bulk velocity has only the z component linearly depending on x, while the pressure p and temperature T of the gas are constant over the whole space. According to the Newton law, the shear stress P_{xz} is proportional to the velocity gradient, that is,

$$P_{xz} = -\eta \frac{\mathrm{d}\nu_z}{\mathrm{d}x}.\tag{5.30}$$

This relation defines the viscosity coefficient η . Under such conditions, the Boltzmann equation is linearized using the quantity $\xi = (\ell/c_{mp})(d\nu_z/dx)$ as the small parameter. The reference density n_R and the temperature T_R are assumed to be constant in Eq. (5.22), while the reference velocity \mathbf{v}_{R} to be given by Eq. (5.29). Then Eq. (5.23) takes the following form:

$$\hat{L}h - 2\frac{c_x c_z}{\ell c_{\rm mp}} = 0. \tag{5.31}$$

This is a complicated integral equation where h is an unknown quantity depending only on the molecular velocity c. Once the perturbation function h is obtained from Eq. (5.31), it is substituted into Eq. (5.27) and the viscosity coefficient η is calculated with the help of Eq. (5.30).

A numerical solution of Eq. (5.31) is a very difficult task, which requires knowledge of the intermolecular interaction law. For the hard-sphere molecules, such calculations were carried out in Ref. [7] where the following expression for the viscosity was obtained:

$$\eta = 1.016034 \frac{5\pi}{32} m_{\rm P} n \bar{c} \bar{l} \approx \frac{1}{2} m_{\rm P} n \bar{c} \bar{l}.$$
 (5.32)

Here, the mean free path \bar{l} is given by Eq. (3.55).

Thermal Conductivity Coefficient

Now, let us consider a gas being at rest, that is, v = 0, and occupying an infinite region. However, the gas temperature is not constant, but its deviation from the equilibrium value T_0 linearly depends on the x-coordinate, that is,

$$(T(x) - T_0) \propto x. \tag{5.33}$$

According to the Fourier law, the heat flux is determined as

$$q_x = -\lambda \frac{\mathrm{d}T}{\mathrm{d}x},\tag{5.34}$$

where λ is the heat conductivity. To calculate it, the Boltzmann equation is linearized using the quantity $\xi = (\ell/T)(dT/dx)$ as the small parameter and assuming the reference temperature T_R to be given by Eq. (5.33), the pressure to be constant, and the bulk velocity to be equal to zero. Then Eq. (5.23) is reduced to

$$\hat{L}h - \frac{c_x}{\ell} \left(\frac{c^2}{c_{\rm mp}^2} - \frac{5}{2} \right) = 0. \tag{5.35}$$

This is practically the same integral equation as Eq. (5.31) with the different free term. Here h also depends only on the molecular velocity c. When the perturbation function h is obtained from Eq. (5.35), it is substituted into Eq. (5.28). Then the heat conductivity is obtained with the help of Eq. (5.34).

The numerical solution of Eq. (5.35) for the hard-sphere potential is reported in Ref. [7], where the following expression is given:

$$\lambda = 1.025218 \frac{75\pi}{128} k n \overline{c} \overline{l} \approx \frac{15}{8} k n \overline{c} \overline{l}, \tag{5.36}$$

where \overline{l} is given by Eq. (3.55).

Prandtl Number

The transport coefficients η and λ are related via the Prandtl number defined as

$$\Pr = c_p \frac{\eta}{\lambda},\tag{5.37}$$

where c_p is the specific heat at constant pressure. If one substitutes the expressions (5.32) and (5.36) into this definition and takes into account $c_p = 5k/(2m)$, then one obtains

$$\Pr = 0.66069 \approx \frac{2}{3}.\tag{5.38}$$

It can be verified that the Prandtl number calculated on the basis of experimental data for the transport coefficient is very close to 2/3 for all monoatomic gases.

5.1.8

Model Equations

A numerical solution of Eqs. (5.19) and (5.23) with the exact expression of the collision integral requires a great computational effort; that is why some simplified expressions of $I(ff_*)$ are used. The kinetic equations with such expressions maintain the main properties of the exact Boltzmann equation, but they allow us to reduce significantly the computational effort to calculate rarefied gas flows. The simplified kinetic equations are called model equations.

The most usual model equation was proposed by Bhatnagar, Gross, and Krook (BGK) [8] and by Welander [9]. They presented the collision integral as

$$J_{\text{BGK}}(ff_*) = \nu [f^{\text{M}} - f(\mathbf{r}, \mathbf{c})]. \tag{5.39}$$

Here $f^{\mathrm{M}} = f^{\mathrm{M}}(n, T, \mathbf{v})$ is the local Maxwellian given by Eq. (5.15), where the local values of the number density $n(\mathbf{r})$, bulk velocity $\mathbf{v}(\mathbf{r})$, and temperature $T(\mathbf{r})$ are unknown and calculated via the distribution function $f(\mathbf{r}, \mathbf{c})$ in accordance with the definitions (5.6), (5.7), and (5.9), respectively. Thus, the kinetic equation (5.19) with the model collision integral (5.39) continues to be nonlinear. The parameter ν is chosen so as to provide the correct expression of one transport coefficient, that is, η or λ . However, no choice of ν provides the correct Prandtl number (5.38). Thus, it is impossible to obtain correctly both viscosity and heat conductivity using a unique expression of ν . If one uses the expression

$$\nu(\mathbf{r}) = \frac{p(\mathbf{r})}{\eta},\tag{5.40}$$

then one obtains the correct viscosity coefficient and hence a correct description of the momentum and mass transfer. However, if one is interested in a correct description of heat transfer, one should use the expression

$$\nu(\mathbf{r}) = \frac{5}{2} \frac{pk}{m\lambda} = \Pr \frac{p(\mathbf{r})}{n},\tag{5.41}$$

which provides the correct heat conductivity. Here, Eq. (5.37) has been used.

The S model proposed by Shakhov [10] is a modification of the BGK model giving the correct Prandtl number. The collision integral of this model reads as

$$J_{S}(ff_{*}) = \frac{p}{\eta} \left\{ f^{M} \left[1 + \frac{4}{15} \frac{(\mathbf{q} \cdot \mathbf{C})}{pc_{mp}^{2}} \left(\frac{C^{2}}{c_{mp}^{2}} - \frac{5}{2} \right) \right] - f(\mathbf{r}, \mathbf{c}) \right\}, \quad \mathbf{C} = \mathbf{c} - \mathbf{v}.$$

$$(5.42)$$

Since the model equations (5.39) and (5.42) significantly reduce the computational effort, they are widely used in practical calculations. However, to obtain reliable results, one should apply an appropriate model equation. If a gas flow is isothermal and the heat transfer is not important, the BGK equation is the most suitable model equation. If a gas flow is nonisothermal, it is better to apply the S model.

5.1.9

Gas-Surface Interaction

On a boundary surface, the velocity distribution function of incident particles $f^$ is related to that of reflected molecules f^+ as

$$c_n f^+(\mathbf{c}) = -\int_{c_n < 0} c'_n f^-(\mathbf{c}') R(\mathbf{c}', \mathbf{c}) d^3 \mathbf{c}', \quad c_n \ge 0,$$

$$(5.43)$$

where \mathbf{c}' is the velocity of incident particles, \mathbf{c} is the velocity of reflected particles, $c_n = \mathbf{c} \cdot \mathbf{n}$ is a normal velocity component, and \mathbf{n} is the unit vector normal to the surface directed to the gas. The explicit expression of the scattering kernel $R(\mathbf{c}', \mathbf{c})$ depends on the gas-surface interaction law.

In practice, the concept of accommodation coefficient α is frequently used, which is defined as

$$\alpha(\psi) = \frac{j^{-}(\psi) - j^{+}(\psi)}{j^{-}(\psi) - j^{+}_{\text{diff}}(\psi)}, \quad j^{\pm}(\psi) = \int_{c_n > 0} |c_n| f(\mathbf{c}) \psi(\mathbf{c}) d\mathbf{c},$$

$$c_n < 0$$
(5.44)

where $j^{\pm}(\psi)$ is the flux of the property $\psi(\mathbf{c})$ for the reflected/incident particles and j_{diff}^+ is the flux corresponding to the diffuse scattering. For instance, when $\psi = (1/2)mc^2$, then $\alpha(\psi)$ will be the energy accommodation coefficient.

-			
Gas	α		
	a	b	с
Не	0.71	0.71	0.96
Ne	0.80		
Ar	0.88	0.87	0.98
Kr	0.92	0.92	1.00

Table 5.1 Accommodation coefficient α extracted from the experimental data [11] applying the diffuse-specular kernel (5.46).

The well-known diffuse scattering (cosine law) corresponds to the following kernel:

$$R_{\rm d}(\mathbf{c}', \mathbf{c}) = \frac{m_{\rm P}^2 c_n}{2\pi (kT_{\rm w})^2} \exp\left(-\frac{m_{\rm P}c^2}{2kT_{\rm w}}\right),\tag{5.45}$$

where $T_{\rm w}$ is the surface temperature. Physically, it means that a particle can be reflected to any direction independent of its velocity before the collision with a surface. Such an interaction is called complete accommodation because it provides $\alpha = 1$ calculated by Eq. (5.44). In many practical applications, the diffuse scattering is well justified and provides reliable results. It usually happens for technical surfaces, which are rough and uncleaned. However, the interaction of gas with a smooth and atomically clean surface can differ from the diffuse scattering.

To take into account a noncomplete accommodation, it is assumed that a part of incident particles are scattered diffusely, while the rest of the particles are reflected specularly. Such a model of the gas-surface interaction is called diffuse-specular. If one calculates the accommodation coefficient α defined by Eq. (5.44) for this model, one obtains that the part reflected diffusely is exactly equal to the accommodation coefficient α for any function $\psi(\mathbf{c})$. Thus, the corresponding kernel is written down as

$$R_{ds}(\mathbf{c}', \mathbf{c}) = \alpha R_{d}(\mathbf{c}', \mathbf{c}) + (1 - \alpha)\delta(\mathbf{c}'_{t} - \mathbf{c}_{t})\delta(c'_{n} + c_{n}), \tag{5.46}$$

where \mathbf{c}_t is the two-dimensional tangential velocity.

Numerical values of the coefficient α obtained from the experimental results reported in Ref. [11] are given in Table 5.1. The corresponding experimental results were obtained for an atomically clean surface, that is, surface prepared in vacuum conditions by vapor deposition.

However, some experimental data contradict theoretical results based on the kernel (5.46) (see Section 5.5.7). The main reason of such a contradiction is that the diffuse-specular kernel contains just one parameter and cannot describe the complexity of the gas-surface interaction. Thus, another kernel containing more parameters should be used, for example, that proposed by Cercignani and

a: atomically clean silver; b: atomically clean titanium; c: titanium covered by oxygen.

Gas	σ_{t}	a _n
Не	0.90	0.10
Ne	0.89	0.75
Ar	0.96	1.00
Kr	1.00	1.00
Xe	1.00	1.00
H_2	0.95	
N_2	0.91	
CO_2	1.00	

Table 5.2 Accommodation coefficients σ_t and a_n extracted from the experimental data [13,14] applying the CL kernel (5.47): surface is typically technical.

Lampis (CL) [12]:

$$R_{\rm CL}(\mathbf{c}', \mathbf{c}) = \frac{m_{\rm P}^2 c_n}{\pi^2 a_n \sigma_{\rm t} (2 - \sigma_{\rm t}) (2kT_{\rm w})^2} \times \exp\left\{-\frac{m_{\rm P} [\mathbf{c}_{\rm t} - (1 - \sigma_{\rm t}) \mathbf{c}_{\rm t}']^2}{2kT_{\rm w} \sigma_{\rm t} (2 - \sigma_{\rm t})}\right\}$$

$$\times \exp\left\{-\frac{m_{\rm P} \left[c_n^2 + (1 - a_n)c_n'^2\right]}{2kT_{\rm w} a_n}\right\}$$

$$\times \int_0^{2\pi} \exp\left\{\frac{m_{\rm P} \sqrt{1 - a_n} c_n c_n' \cos \phi}{kT_{\rm w} a_n}\right\} d\phi,$$
(5.47)

where the coefficient a_n is the accommodation coefficient of energy corresponding to the normal velocity c_n , which varies from 0 to 1, while σ_t is the accommodation coefficient of the tangential momentum, which varies from 0 to 2. In other words, if one substitutes the kernel (5.47) into Eq. (5.44) using $\psi = \frac{1}{2}mc_n^2$ one obtains $\alpha(\psi) = a_n$. Using the function $\psi = m\mathbf{c}_t$ in Eq. (5.44), one obtains $\alpha(\psi) = \sigma_t$. In the particular case $a_n = \sigma_t = 1$, the kernel (5.47) coincides with the diffuse one (5.45). The combination $a_n = 0$ and $\sigma_t = 2$ corresponds to the back scattering, which can occur on a rough surface. Numerical values of the accommodation coefficients σ_t and a_n extracted from the experimental data on the slip coefficients [13] and on the heat transfer [14] are presented in Table 5.2. The corresponding experimental measurements [13,14] were carried out for a technical surface, that is, without any special treatment.

From Tables 5.1 and 5.2, it can be seen that heavy gases such as Kr, Xe, and CO₂ are characterized by the complete accommodation on a technical surface, while light gases such as He and Ne do not undergo diffuse scattering. The accommodation coefficients also depend on the chemical composition of surfaces. If a surface is atomically clean, the scattering is less diffuse.

5.2

Methods of Calculations of Gas Flows

5.2.1

General Remarks

Methods of calculations depend on the gas flow regime. In the viscous regime (Kn < 0.01), the continuum mechanics equations are successfully used. Main results based on these equations are given in Chapter 4. A moderate gas rarefaction, say Kn < 0.1, can also be considered on the continuum mechanics level if the velocity slip and temperature jump boundary conditions are applied. The explicit form of such conditions and recommended values of the slip and jump coefficients are given in Section 5.3. The free molecular regime (Kn > 100) – when all molecules move independent of each other - is relatively easy for analytical and numerical calculations. Some results for this regime are also given in Chapter 4.

In this chapter, more details will be given on the methods used in the transitional regime when the Navier-Stokes equation is not valid any more, but the intermolecular collisions cannot be neglected. All methods used in this regime can be divided into two large groups: deterministic approach based on solving of the kinetic equation (5.19) and probabilistic approach representing a Monte Carlo simulation of a large amount of model particles considering collisions between them and their interaction with a solid surface. The main ideas, advantages, and disadvantages of both approaches are described in the following sections.

5.2.2

Deterministic Methods

The deterministic methods are based on analytical or numerical solution of the kinetic equation (5.19) or its linearized form (5.23). Usually, the collision integral J is substituted by its model, for example, (5.39) or (5.42). Here, the method is illustrated for the BGK model, that is, Eq. (5.19) with (5.39), but it remains the same for any other model and for the Boltzmann equation itself.

If a set of values of the molecular velocity \mathbf{c}_i is chosen, then the kinetic equation (5.19) is replaced by a system of differential equations for the functions $f_i(\mathbf{r}) = f(\mathbf{r}, \mathbf{c}_i)$ coupled with the collision integral, that is,

$$\mathbf{c}_{i} \cdot \frac{\partial f_{i}(\mathbf{r})}{\partial \mathbf{r}} + \nu(\mathbf{r})f_{i}(\mathbf{r}) = \nu(\mathbf{r})f_{i}^{\mathbf{M}}(\mathbf{r}), \tag{5.48}$$

where Eq. (5.39) has been used. The Maxwellian $f_i^{\rm M}({\bf r})$ defined by Eq. (5.15) depends on the coordinates \mathbf{r} via the moments $n(\mathbf{r})$, $\mathbf{v}(\mathbf{r})$, and $T(\mathbf{r})$, which are calculated by some integration rule, that is,

$$\begin{bmatrix} n(\mathbf{r}) \\ \mathbf{v}(\mathbf{r}) \\ T(\mathbf{r}) \end{bmatrix} = \sum_{i} \begin{bmatrix} \frac{1}{n} \mathbf{c}_{i} \\ \frac{m_{\mathrm{p}}}{3nk} (\mathbf{c}_{i} - \mathbf{v})^{2} \end{bmatrix} f_{i}(\mathbf{r}) W_{i}, \tag{5.49}$$

according to Eqs. (5.6), (5.7) and (5.9), respectively. Here, W_i is the weight of the node \mathbf{c}_i .

The system (5.48) with (5.49) is solved by an iteration procedure. First, some values of the moments $n(\mathbf{r})$, $\mathbf{v}(\mathbf{r})$, and $T(\mathbf{r})$ are assumed in all points of the physical space r. Then, the following steps are executed:

- i) The differential equations (5.48) are solved for each molecular velocity c_i applying a finite difference scheme.
- ii) New values of the moments are calculated at all points of the physical space using Eq. (5.49).
- iii) The convergence is verified comparing the moments obtained in two successive iterations. If the convergence is reached, all moments of practical interest (density, bulk velocity, pressure tensor, temperature, etc.) are calculated using the rule (5.49), and the calculations are stopped. If the convergence is not reached, all steps are repeated.

The main advantages of the discrete velocity method are as follows: (i) Since the method is deterministic, it is completely free from any kinds of statistical noise; therefore, this method is indispensable in case of low Mach number flows. (ii) It requires a modest computational memory because the calculations are carried out so that it is not necessary to store the distribution function at all points of the physical and velocity spaces. However, a realization of this method needs a careful choice of numerical grids in both physical and velocity spaces. It is not so easy to adapt a physical grid to a complicated geometrical configuration. In many practical problems, the distribution function is discontinuous, requiring a special modification of the method. The iteration convergence is very slow at small values of the Knudsen number, that is, a special methodology of acceleration [15] must be used. Details of the method are given in Section 3.13 of Ref. [4] and in Refs [5,16].

If one applies a linearized model equation, then a system of integral equations can be obtained for distribution function moments. These equations can be solved by a variational method with a quite modest computational effort. Applying this method, care must be taken regarding the velocity space grid and distribution function discontinuity. However, the method needs a large computational memory. Details about the method can be found in Chapter IV, Section 12 of Ref. [1].

5.2.3

Probabilistic Methods

The probabilistic methods consist of numerical simulations of molecule motion, interaction between them, and their interaction with a solid surface. Since both gas-gas and gas-surface interactions are stochastic processes, random numbers are used in their simulations. Therefore, this approach is called the direct simulation Monte Carlo (DSMC) method.

To realize the DSMC method, the region of the gas flow is divided into a network of cells with dimensions such that the change in flow properties across each cell is small. Then a huge number (about 10⁷) of molecules are distributed over the gas flow region, that is, their positions \mathbf{r}_i and velocities \mathbf{c}_i are stored in a computer memory. The time is advanced in discrete steps of magnitude Δt , such that it is small compared with the mean time between two successive collisions. The particle motion and intermolecular collisions are uncoupled over the time increment Δt by the repetition of the following procedures:

i) The particles are moved through the distance determined by their velocities \mathbf{c}_i and Δt and new positions are calculated as

$$\mathbf{r}_{i. \text{ new}} = \mathbf{r}_{i. \text{ old}} + \mathbf{c}_i \Delta t. \tag{5.50}$$

If the straight trajectory crosses a solid surface, a simulation of the gassurface interaction is performed according to a given law, that is, a new velocity \mathbf{c}_i is generated and the particle continues to move with the new velocity. If the new position $\mathbf{r}_{i,\text{ new}}$ is out of the computational region, then the information about the corresponding particle is removed. It happens if the gas flow region is not closed, but some surfaces allow influx and outflux of the gas.

- ii) New particles are generated at boundaries when there is an inward flux. This step is necessary in the case of a nonclosed region of the gas flow. The generation is done according to a boundary condition. Usually, a local Maxwellian (5.15) with given values of the density n, bulk velocity \mathbf{v} , and temperature T is generated.
- iii) Following the no time counter (NTC) method [17], a representative number of collisions appropriate to Δt and number of particles in every cell is calculated as

$$N_{\text{coll}} = \frac{1}{2} N \overline{N} F_{\text{N}} (\sigma c_{\text{r}})_{\text{max}} \frac{\Delta t}{V_{C}}, \tag{5.51}$$

where N is the number of model particles in the cell at the current time interval, \overline{N} is its average magnitude during all previous intervals, $F_{\rm N}$ is the number of real particles represented by one model particle, σ is the molecule cross section that can be dependent on the relative molecular velocity $c_{\rm r}$, $(\sigma c_{\rm r})_{\rm max}$ represents a maximum value of the product (σc_r) , and V_C is the cell volume. Then a random pair of particles being in the same cell is chosen and its product (σc_r) is calculated. The pair is accepted for collision if the ratio $(\sigma c_r)/(\sigma c_r)_{max}$ is larger than a random number. Otherwise, it is rejected. Such a procedure provides more collisions between fast particles and less collisions for low-speed ones. If the pair is accepted, the precollision velocities of the particles are replaced by the postcollision values in accordance with the given potential of the intermolecular interaction. Recently, a technique allowing application of any intermolecular potential has been proposed [18].

iv) The moments are calculated according to Eqs. (5.6)-(5.10). For instance, the bulk velocity in every cell is calculated as the average velocity of all molecules in the cell, that is,

$$\mathbf{v} = \frac{1}{N} \sum_{i=1}^{N} \mathbf{c}_i. \tag{5.52}$$

Steps (i)-(iv) must be repeated many times in order to establish a stationary flow. Then the simulations must be continued in order to calculate the average values of the moments over many iterations (samples).

In the free molecular regime ($Kn \gg 1$), the particles do not suffer any mutual collisions and it is not necessary to simulate their motion simultaneously, but the motion of every particle can be simulated independent of the others. First, a particle is generated on a boundary of gas influx, that is, its position and velocity are generated according to a boundary condition. Then, using the generated position and velocity, the point of it hitting a solid surface is calculated. A new particle velocity is generated according to the gas-surface interaction law and the point of the next hit is calculated. In this way, the whole trajectory of the particle is simulated up to its exit from the computational region. Repeating this procedure with many particles, statistical information is accumulated, which allows us to calculate macroscopic quantities such as the mass flow rate. This technique is called the test particle Monte Carlo method.

Thus, the idea of the probabilistic method is very clear. To use it, neither grids in the velocity space nor finite difference scheme is necessary. The physical cells can be easily adapted to any geometrical configuration. It is not difficult to simulate nonelastic collisions occurring in polyatomic gases. Even more complicated phenomena such as dissociation, ionization, and so on are considered without a great effort. The book by Bird [17] contains numerical codes that can be modified and used in practical calculations. Because of these advantages, the direct simulation and test particle Monte Carlo methods are so widely used in practice that many researchers think that any gas dynamic problem can be solved by these methods.

However, the probabilistic methods have their own shortcomings and cannot be considered as a universal remedy. The main defect of the method is the statistical scattering (or statistical noise), which is reduced by increasing the number of samples. Initially, the DSMC method was elaborated for aerothermodynamic problems, where the Mach number is extremely high. Under such conditions, the statistical noise is very low and a small number of samples provide reliable results. When the Mach number is small, which is usually the case in vacuum systems, then one needs such large numbers of samples to reduce the noise that the computational time becomes inadmissibly long. In this case, an application of the deterministic approach is considerably more advantageous.

5.3 Velocity Slip and Temperature Jump Phenomena

A moderate gas rarefaction can be taken into account by solving the continuum mechanics equations with the velocity slip and temperature jump boundary conditions. In some applications, these solutions can be applied practically up to the transition regime. Analytical expressions based on the slip and jump boundary conditions represent an asymptotic behavior of a numerical solution based on the kinetic equation in the limit $Kn \to 0$. Thus, the slip and jump solutions can be used as a criterion to verify an uncertainty of numerical results.

In this section, the velocity slip and temperature jump conditions are formulated. Recommended data on the corresponding coefficients are provided. Some examples of their applications are given in the subsequent sections.

5.3.1

Viscous Slip Coefficient

Consider a gas flowing in the z-direction near a solid surface being at rest. Let the x-axis be directed normally to the surface toward the gas, as is shown in Figure 5.1. According to the slip condition, the bulk velocity of a gas is not equal to zero on the surface, but its tangential component v_z is proportional to its normal gradient:

$$v_z = \beta_P \ell \frac{\partial v_z}{\partial x}, \quad \text{at} \quad x = 0,$$
 (5.53)

where the equivalent free path ℓ is defined by Eq. (5.1) and β_P is the viscous slip coefficient calculated from the kinetic equation applying to the Knudsen layer. Such a layer is adjacent to the solid surface and has the thickness on the order of the molecular free path. The condition (5.43) is assumed on the solid surface, that is, on the lower boundary of the Knudsen layer, while a local Maxwellian distribution function is assumed on the upper boundary of the Knudsen layer. The quantity $\xi = (\ell/c_{mp})(\partial v_z/\partial x)$ is used as the linearization parameter. Then

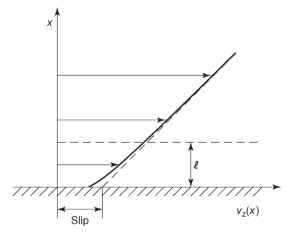


Figure 5.1 Scheme of viscous velocity slip, Eq. (5.53): the solid line represents the real velocity profile; the dashed line represents the extrapolation of the linear profile up to the surface.

the linearized kinetic equation (5.23) is solved numerically. The detailed technique of calculations of the slip coefficient $\beta_{\rm P}$ and their numerical values can be found in Refs [19-21].

The velocity profile shown in Figure 5.1 by the solid line represents a numerical solution of the S model. It can be seen that outside of the Knudsen layer $(x \ge \ell)$, the velocity linearly depends on the *x*-coordinate, that is, $\nu_z \propto (\beta_p \ell + x)$. However, near the surface, $0 \le x \le \ell$, the profile is not linear, but has a small defect, which does not contribute to the first-order slip correction. Thus, the viscous slip coefficient β_P is calculated via the extrapolation of the linear velocity profile up to the surface.

The value of the slip coefficients β_P recommended in practical calculations is as follows:

$$\beta_{\rm P} = 1.018,$$
 (5.54)

which was obtained under an assumption of complete accommodation on the surface. The BGK model provides practically the same velocity profile and the same value of the slip coefficient.

In the case of noncomplete accommodation, the kinetic equation was solved with the boundary condition (5.47) in Ref. [21]. It was observed that the slip coefficient $\beta_{\rm P}$ is very weakly affected by the energy accommodation coefficient a_n , but it strongly depends on the momentum accommodation coefficient σ_t . The numerical data based on the CL scattering law can be interpolated by the formula

$$\beta_{\rm P}(\sigma_{\rm t}) = \frac{1.772}{\sigma_{\rm t}} - 0.7540. \tag{5.55}$$

The data on the viscous slip coefficients for gaseous mixtures can be found in Ref. [22].

5.3.2

Thermal Slip Coefficient

If the solid surface is nonisothermal, but its temperature varies along the z-axis, $T_{\rm w} = T_{\rm w}(z)$, as is depicted in Figure 5.2, then the gas begins to flow near such a surface from the cold to the hot region. In this case, the tangential velocity v_z of the gas is proportional to the temperature gradient:

$$v_z = \beta_{\rm T} \frac{\eta}{T_{\rm HMD}} \frac{\partial T}{\partial z}, \quad \text{at} \quad x = 0,$$
 (5.56)

where $\beta_{\rm T}$ is the thermal slip coefficient, which is calculated from the linearized kinetic equation applied to the Knudsen layer. The quantity $\xi = (\ell/T)(\partial T/\partial z)$ is used as the small parameter. The detailed technique of calculations of the thermal slip coefficient on the basis of the kinetic equation (5.23) and their numerical values can be found in Refs [21,23].

The velocity profile shown in Figure 5.2 by the solid line represents a numerical solution of the S model. It can be seen that outside of the Knudsen layer,

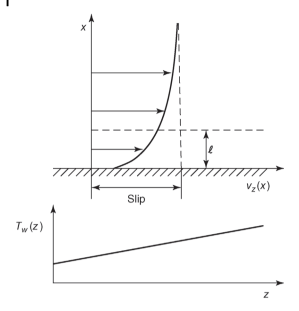


Figure 5.2 Scheme of thermal velocity slip, Eq. (5.56): the solid line represents the real velocity profile; the dashed line represents the velocity magnitude far from the surface.

 $x>2\ell$, the velocity is practically constant, but inside of the Knudsen layer, $0< x<2\ell$, it sharply decreases. The variation of the velocity profile near the surface does not contribute to the first-order slip coefficient. So, the thermal slip coefficient $\beta_{\rm T}$ is calculated via the value of the velocity far from the surface, that is, at $x>2\ell$.

In practice, it is recommended to use the following value:

$$\beta_{\rm T} = 1.175,$$
 (5.57)

which was obtained for the diffuse gas–surface interaction. As was shown in Ref. [21], the thermal slip coefficient $\beta_{\rm T}$ is significantly affected by both accommodation coefficients a_n and $\sigma_{\rm t}$, so it is difficult to propose a formula interpolating the numerical data reported in Ref. [21].

If one deals with a gaseous mixture, the corresponding data on the coefficient β_T are reported in Ref. [24].

5.3.3

Temperature Jump Coefficient

Let us assume that the temperature varies in the direction normal to a wall. Then the temperature of gas $T_{\rm g}$ near the wall is not equal to the wall temperature $T_{\rm w}$, but there is a jump proportional to the normal temperature gradient, that is,

$$T_{\rm g} - T_{\rm w} = \zeta_{\rm T} \ell \frac{\partial T_{\rm g}}{\partial x}, \quad \text{at} \quad x = 0,$$
 (5.58)

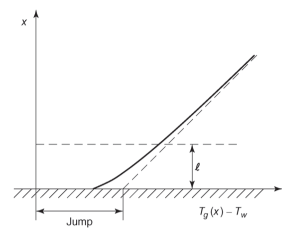


Figure 5.3 Scheme of temperature jump, Eq. (5.58): the solid line represents the real temperature profile; the dashed line represents the extrapolation of the linear profile up to the surface.

where ζ_T is the temperature jump coefficient. The scheme of the jump is shown in Figure 5.3. To calculate the coefficient ζ_T , the kinetic equation (5.23) is solved in the Knudsen layer using the small parameter $\xi = (\ell/T)(\partial T/\partial x)$. The detailed technique of the calculations and numerical values of ζ_T can be found in Ref. [21].

The temperature profile shown in Figure 5.3 by the solid line represents a numerical solution of the S model. It can be seen that outside of the Knudsen layer, $x > 2\ell$, the temperature linearly depends on the x-coordinate, while inside of the Knudsen layer, $0 < x < 2\ell$, there is a small deviation from the linear dependence. Such a deviation is of the second order with respect to the Knudsen number and is neglected in calculations of the temperature jump coefficient.

It is recommended to use the value

$$\zeta_{\rm T} = 1.954$$
 (5.59)

in practice, which was obtained for the gas-surface interaction corresponding to the complete accommodation.

The values of ζ_T for the nondiffuse scattering kernel (5.47) can be found in Ref. [21]. Like the thermal slip coefficient, it is also significantly affected by both accommodation coefficients a_n and σ_t , so no interpolating formula was obtained.

In case of gaseous mixtures, the temperature jump coefficient ζ_{T} was calculated in the work [25].

5.4 Momentum and Heat Transfer through Rarefied Gases

In this section, two classical problems of fluid mechanics will be considered, namely, Couette flow, that is, gas confined between two surfaces moving relative to one another, and heat transfer through a gas between two surfaces having different temperatures. The results are given over the whole range of the gas rarefaction, including the velocity slip and temperature jump analytical solutions.

5.4.1

Planar Couette Flow

Consider two parallel plates placed at x = 0 and x = d, as is shown in Figure 5.4. The lower plate (x = 0) is fixed, while the upper plate moves with a speed ν_w to the right. To apply the linearized kinetic equation, it is assumed that $\nu_{\rm w} \ll c_{\rm mp}$. Then, the ratio $\xi = v_{\rm w}/c_{\rm mp}$ is used as the small parameter of the linearization. The distance d is adopted as the characteristic size, so that the rarefaction parameter is given as

$$\delta = \frac{dp}{\eta c_{\rm mp}} \,. \tag{5.60}$$

Our aim is to calculate the velocity profile $v_z(x)$ and the shear stress P_{xz} as functions of the rarefaction parameter δ . Note that the quantity P_{xz} does not vary between the plates due to the momentum conservation law.

In the free molecular regime ($\delta \to 0$), the solution of the kinetic equation (5.23) can be obtained analytically, see Section 4.2 of Ref. [4]. For the diffuse gassurface interaction, the shear stress and the bulk velocity read

$$P_{xz}^{\text{fm}} = -\frac{p}{\sqrt{\pi}} \frac{\nu_{\text{w}}}{c_{\text{mp}}}, \quad \nu_{z}(x) = \frac{\nu_{\text{w}}}{2}, \quad \text{for} \quad \delta \to 0.$$
 (5.61)

So, in this regime, the velocity $v_z(x)$ is constant over the gap and equal to the mean value of the speeds of the two plates.

Using the method of successive approximations based on the kinetic equation (5.23), it is possible to obtain the first correction for small values of δ :

$$P_{xz} = P_{xz}^{\text{fm}} \left(1 - \frac{\sqrt{\pi}}{2} \delta \right), \quad \text{for} \quad \delta \ll 1.$$
 (5.62)

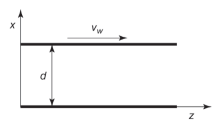


Figure 5.4 Scheme of planar Couette flow.

In the viscous regime $(\delta \to \infty)$, the Navier-Stokes equation with the slip boundary condition (5.53) is applied, which for the problem in question reads

$$v_z = \begin{cases} v_w - \beta_P \,\ell \, \frac{\mathrm{d}v_z}{\mathrm{d}x}, & \text{at} \quad x = d, \\ \beta_P \ell \, \frac{\mathrm{d}v_z}{\mathrm{d}x}, & \text{at} \quad x = 0. \end{cases}$$
 (5.63)

At the upper plate (x = d), the slip condition determines the difference between the surface speed $v_{\rm w}$ and that of the gas $v_{\rm z}$. Then, the velocity profile is obtained analytically

$$v_z(x) = v_w \left(\frac{x}{d} + \frac{\beta_P}{\delta}\right) \left(1 + 2\frac{\beta_P}{\delta}\right)^{-1}, \quad \text{for} \quad \delta \gg 1.$$
 (5.64)

The shear stress is obtained from Eq. (5.30) as

$$P_{xz} = -\eta \frac{v_{\rm w}}{d} \left(1 + \frac{2\beta_{\rm p}}{\delta} \right)^{-1} = P_{xz}^{\rm fm} \frac{\sqrt{\pi}}{\delta + 2\beta_{\rm p}}, \quad \text{for} \quad \delta \gg 1.$$
 (5.65)

As expected, the velocity slip on the surfaces reduces the shear stress.

In the transition regime ($\delta \sim 1$), the kinetic equation (5.23) is solved numerically. Once the perturbation function h is known, then the bulk velocity $v_z(z)$ and shear stress P_{xz} are calculated with the help of Eqs. (5.26) and (5.27), respectively. The corresponding numerical data can be found in Refs [26-28]. The velocity profile $v_z(x)$ obtained from the BGK model assuming the complete accommodation is plotted in Figure 5.5 for four values of the rarefaction parameter. At $\delta = 10$, the solution of the kinetic equation is very close to the analytical slip solution (5.64). In the free molecular regime ($\delta \to 0$), the expression (5.64) and the free molecular solution (5.61) coincide with each other and provide the velocity equal to $v_{\rm w}/2$. The numerical solution for $\delta = 0.01$ is very close to this

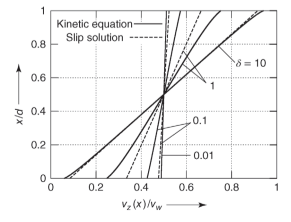


Figure 5.5 Velocity profile $v_z(x)$ in planar Couette flow for different values of the rarefaction δ .

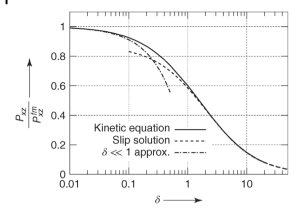


Figure 5.6 Shear stress P_{xz} in planar Couette flow versus rarefaction parameter δ . Solid line – numerical solution of BGK model [26]; dashed line – slip solution Eq. (5.65); point-dashed line – near free molecular solution Eq. (5.62).

value. However, in the transition ($\delta = 1$) and near free molecular ($\delta = 0.1$) regimes, the slip solution (5.64) does not provide reliable results.

The shear stress is shown in Figure 5.6 and in Table 5.A.1 as a function of the rarefaction parameter δ . The slip solution (5.65) works well up to the transition regime, that is, in the range of $\delta \geq 1$. However, this is a peculiarity of the planar Couette flow. For other situations, the slip solution works in a smaller interval of δ .

Exercise 5.1 Consider nitrogen confined between two parallel plates as is shown in Figure 5.4. Calculate its velocity $v_z(x)$ at x=0 and x=d and the shear stress P_{xz} under the following conditions: distance between the plates d=2 mm, speed of the moving plate $v_{\rm w}=300\,{\rm m~s^{-1}}$, and gas temperature $T=293\,{\rm K}$. Use $M = 0.028 \text{ kg mol}^{-1}$, $\eta = 17.5 \times 10^{-6} \text{ Pa s}$, $\beta_p = 1.018$. Consider two values of the gas pressure: (a) p = 0.05 Pa and (b) p = 50 Pa.

Calculation of the most probable speed by Eq. (3.42):

$$c_{\rm mp} = \sqrt{\frac{2 \times 8.31 \,\mathrm{J \, mol}^{-1} \,\mathrm{K}^{-1} \times 293 \,\mathrm{K}}{0.028 \,\mathrm{kg \, mol}^{-1}}} = 417 \,\mathrm{m \, s}^{-1}.$$

a) Calculation of the rarefaction parameter for p = 0.05 Pa using (5.60):

$$\delta = \frac{0.002 \text{ m} \times 0.05 \text{ Pa}}{17.5 \times 10^{-6} \text{ Pa s} \times 417 \text{ m s}^{-1}} = 0.0137.$$

Since $\delta \ll 1$, Eq. (5.61) is used. Then,

$$v_z(0) = v_z(d) = \frac{300 \text{ m s}}{2} = 150 \text{ m s}^{-1}.$$

$$P_{xz} = -\frac{0.05 \text{ Pa}}{1.77} \frac{300 \text{ m s}^{-1}}{417 \text{ m s}^{-1}} = -0.0203 \text{ Pa}.$$

b) Calculation of the rarefaction parameter for p = 50 Pa using (5.60):

$$\delta = \frac{0.002 \text{ m} \times 50 \text{ Pa}}{17.5 \times 10^{-6} \text{ Pa s} \times 417 \text{ m s}^{-1}} = 13.7.$$

Since $\delta \approx 10$, Eqs. (5.64) and (5.65) can be used:

$$v_z(0) = 300 \text{ m s}^{-1} \frac{1.018/13.7}{1 + 2 \times 1.018/13.7} = 19.4 \text{ m s}^{-1}.$$

$$\label{eq:vz} \nu_z(d) = 300~{\rm m~s^{-1}} \frac{1 + 1.018/13.7}{1 + 2 \times 1.018/13.7} = 281~{\rm m~s^{-1}}.$$

$$P_{xz} = -\frac{17.5 \times 10^{-6} \text{ Pa s} \times 300 \text{ m s}^{-1}}{0.002 \text{ m} \times (1 + 2 \times 1.018/13.7)} = -2.29 \text{ Pa}.$$

5.4.2 Cylindrical Couette Flow

Consider a gas flow between two coaxial cylinders of radii R_1 and R_2 , as is depicted in Figure 5.7. The internal cylinder R_1 rotates with an angular speed ω , while the external one R_2 is fixed. Since this flow is considered in the cylindrical coordinate, the notation $P_{r\varphi}$ is used instead of P_{xz} . It is assumed that the surface speed of the internal cylinder, that is, the quantity ωR_1 , is small compared with the most probable speed $c_{\rm mp}$. Thus, the small parameter $\xi = \omega R_1/c_{\rm mp}$ can be used to linearize the kinetic equation. The internal cylinder radius R_1 is assumed as the characteristic size, so the rarefaction parameter is given as

$$\delta = \frac{R_1 p}{\eta c_{\rm mp}}.\tag{5.66}$$

We will calculate the shear stress $P_{r\varphi}$ and the azimuth bulk velocity ν_{φ} , which are determined by the rarefaction parameter δ and by the radius ratio R_2/R_1 . For

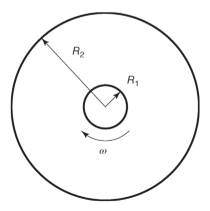


Figure 5.7 Scheme of cylindrical Couette flow.

the cylindrical Couette flow, the quantity $P_{r\varphi} r^2$ is constant because of the momentum conservation law. Here, r is the radial coordinate, that is, the distance from the cylinder axis.

In the free molecular regime, the kinetic equation (5.23) is solved analytically, and the shear stress $P_{r\omega}$ thus reads

$$P_{r\varphi}^{\text{fm}}(r) = \frac{p}{\sqrt{\pi}} \frac{\omega R_1^3}{c_{\text{mp}} r^2}, \quad \text{for} \quad \delta \to 0.$$
 (5.67)

Note that at the internal cylinder surface $r = R_1$, this expression coincides with that for the planar Couette flow given by Eq. (5.61). The velocity profile is given as

$$\nu_{\varphi}(r) = \frac{\omega R_1}{\pi} \left[\frac{r}{R_1} \arcsin\left(\frac{R_1}{r}\right) - \sqrt{1 - \left(\frac{R_1}{r}\right)^2} \right], \quad \text{for} \quad \delta \to 0.$$
 (5.68)

It is interesting that the free molecular solution, Eqs. (5.67) and (5.68), does not depend on the external cylinder radius R_2 , but it is determined only by the internal cylinder radius R_1 .

In the viscous regime, the Navier-Stokes equation in the cylindrical coordinates is solved. The slip boundary condition (5.53) must also be written in the cylindrical variables (r, φ) as

$$\nu_{\varphi} = \begin{cases} \omega R_1 + \beta_{p} \ell \left(\frac{\mathrm{d}\nu_{\varphi}}{\mathrm{d}r} - \frac{\nu_{\varphi}}{r} \right), & \text{at} \quad r = R_1, \\ -\beta_{p} \ell \left(\frac{\mathrm{d}\nu_{\varphi}}{\mathrm{d}r} - \frac{\nu_{\varphi}}{r} \right), & \text{at} \quad r = R_2. \end{cases}$$
(5.69)

Then the velocity profile reads

$$\nu_{\varphi}(r) = \omega R_1^2 \left[\frac{1}{r} - \frac{r}{R_2^2} \left(1 - 2 \frac{R_1}{R_2} \frac{\beta_{\rm P}}{\delta} \right) \right] \mathcal{D}, \quad \text{for} \quad \delta \gg 1,$$
 (5.70)

where

$$\mathcal{D} = \left\{ 1 - \left(\frac{R_1}{R_2}\right)^2 + 2\frac{\beta_P}{\delta} \left[\left(\frac{R_1}{R_2}\right)^3 + 1 \right] \right\}^{-1}.$$
 (5.71)

Equation (5.30) could be written down in the cylindrical coordinate as

$$P_{r\varphi}(r) = -\eta \left(\frac{\mathrm{d}\nu_{\varphi}}{\mathrm{d}r} - \frac{\nu_{\varphi}}{r}\right). \tag{5.72}$$

Then, with the help of Eq. (5.70), the shear stress takes the following form:

$$P_{r\varphi}(r) = 2\eta\omega \frac{R_1^2}{r^2} \mathcal{D} = P_{r\varphi}^{\text{fm}}(r) \frac{2\sqrt{\pi}}{\delta} \mathcal{D}, \quad \text{for} \quad \delta \gg 1.$$
 (5.73)

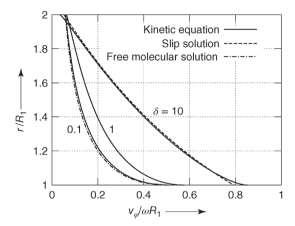


Figure 5.8 Velocity profiles $v_{\omega}(r)$ in cylindrical Couette flow at $R_2/R_1=2$.

In the limit of high radius ratio, $R_2/R_1 \rightarrow \infty$, this solution yields

$$P_{r\varphi}(r) = 2\eta\omega \frac{R_1^2}{r^2} \left[1 + 2\frac{\beta_p}{\delta} \right]^{-1}, \quad \text{for} \quad \delta \gg 1 \quad \text{and} \quad R_2 \gg R_1.$$
 (5.74)

If $R_2/R_1 \ge 5$, the limit expression provides the shear stress within the uncertainty of 4%.

In the transition regime ($\delta \sim 1$), the kinetic equation (5.23) is solved numerically. The technique and corresponding numerical data can be found in Refs. [29,30]. The velocity profiles $\nu_{\varphi}(r)$ calculated from the BGK equation assuming the diffuse gas-surface interaction are plotted in Figure 5.8 for some values of the rarefaction parameter δ at $R_2/R_1=2$. For $\delta=10$, the numerical solution is close to that of the slip (5.70). For the small value of the rarefaction parameter, that is, $\delta = 0.1$, the analytical free molecular solution (5.68) practically coincides with the numerical solution. In the transition regime ($\delta = 1$), the numerical solution cannot be presented by the slip expression (5.70) and differs significantly from the free molecular solution (5.68).

The shear stress $P_{r\varphi}$ is plotted in Figure 5.9 and presented in Table 5.A.1. In the cylindrical Couette flow, the slip solution (5.73) works well only for moderately large values of the rarefaction parameter, say up to $\delta = 5$. The difference between the shear stress $P_{r\varphi}$ at $R_2/R_1=3$ and that at $R_2/R_1=5$ is very small. In practice, it means that the results corresponding to the radius ratio $R_2/R_1=5$ can be successfully applied for larger values of this ratio.

It is important to note that in the transition and free molecular regimes, the relation of the shear stress to the bulk velocity, that is, Eqs. (5.30) and (5.72), is not valid.

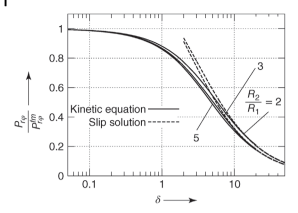


Figure 5.9 Shear stress $P_{r\omega}$ in cylindrical Couette flow versus rarefaction parameter δ and radius ratio R_2/R_1 . Solid line – numerical solution of BGK model; dashed line – slip solution, Eq. (5.73).

Exercise 5.2 Consider nitrogen confined between two coaxial cylinders, as is shown in Figure 5.7. Calculate its velocity $v_r(r)$ and the shear stress $P_{r\omega}(r)$ at r= R_1 and $r = R_2$ under the following conditions: radii $R_1 = 0.5$ cm and $R_2 = 2R_1$, rotation speed $\omega = 10^4$ Hz, gas temperature T = 293 K. Use M = 0.028 kg mol⁻¹, $\eta=17.5\times 10^{-6}$ Pa s, $\beta_{\rm p}=1.018$. Consider two values of the gas pressure: (a) p=1.0180.015 Pa and (b) p = 15 Pa.

The most probable speed is the same as in Exercise 5.1, that is, $c_{\rm mp} = 417 \; {\rm m \; s^{-1}}.$

a) Calculation of the rarefaction parameter for p = 0.015 Pa using (5.66):

$$\delta = \frac{0.005 \text{ m} \times 0.015 \text{ Pa}}{17.5 \times 10^{-6} \text{ Pa s} \times 417 \text{ m s}^{-1}} = 0.0103.$$

Since $\delta \ll 1$, Eqs. (5.67) and (5.68) are used:

$$v_{\varphi}(R_1) = \frac{\omega R_1}{2} = \frac{10^4 \text{ Hz} \times 0.005 \text{ m}}{2} = 25 \text{ m s}^{-1}.$$

$$\nu_{\varphi}(R_2) = \frac{10^4 \ \mathrm{Hz} \times 0.005 \ \mathrm{m}}{3.14} \left[2 \arcsin \left(\frac{1}{2} \right) - \sqrt{1 - 0.25} \right] = 2.88 \ \mathrm{m \ s^{-1}}.$$

$$P_{r\varphi}(R_1) = \frac{p}{\sqrt{\pi}} \frac{\omega R_1}{c_{\rm mp}} = \frac{0.015 \text{ Pa}}{1.77} \times \frac{10^4 \text{ Hz} \times 0.005 \text{ m}}{417 \text{ m s}^{-1}} = 1.02 \times 10^{-3} \text{ Pa}.$$

$$P_{r\varphi}(R_2) = \left(\frac{R_1}{R_2}\right)^2 P_{r\varphi}(R_1) = \left(\frac{1}{2}\right)^2 \times 1.01 \times 10^{-3} \text{ Pa} = 2.54 \times 10^{-4} \text{ Pa}.$$

b) Calculation of the rarefaction parameter for p = 15 Pa using (5.66):

$$\delta = \frac{0.005 \text{ m} \times 15 \text{ Pa}}{17.5 \cdot 10^{-6} \text{ Pa s} \times 417 \text{ m s}^{-1}} = 10.3.$$

Since $\delta > 10$, Eqs. (5.70) and (5.73) are used. First, the factor \mathcal{D} is calculated using (5.71):

$$\begin{split} \mathcal{D} &= \left\{1 - \left(\frac{1}{2}\right)^2 + 2\frac{1.018}{10.3} \left[\left(\frac{1}{2}\right)^3 + 1\right]\right\}^{-1} = 1.03. \\ \nu_{\varphi}(R_1) &= \omega R_1 \left[1 - \left(\frac{R_1}{R_2}\right)^2 \left(1 - 2\frac{R_1}{R_2}\frac{\beta_p}{\delta}\right)\right] \mathcal{D} \\ &= 10^4 \, \mathrm{Hz} \times 0.005 \, \mathrm{m} \left[1 - \left(\frac{1}{2}\right)^2 \left(1 - 2\frac{1}{2}\frac{1.018}{10.3}\right)\right] \times 1.03 = 39.8 \, \mathrm{m \, s^{-1}}. \\ \nu_{\varphi}(R_2) &= 2\omega \left(\frac{R_1}{R_2}\right)^2 \frac{\beta_p}{\delta} \mathcal{D} = 2 \times 10^4 \, \mathrm{Hz} \times 0.005 \, \mathrm{m} \left(\frac{1}{2}\right)^2 \frac{1.018}{10.3} \times 1.03 \\ &= 2.54 \, \mathrm{m \, s^{-1}}. \\ P_{r\varphi}(R_1) &= 2\eta\omega\mathcal{D} = 2 \times 17.5 \times 10^{-6} \, \mathrm{Pa \, s} \times 10^4 \, \mathrm{Hz} \times 1.03 = 0.361 \, \mathrm{Pa}. \end{split}$$

$$P_{r\varphi}(R_2) = \left(\frac{R_1}{R_2}\right)^2 P_{r\varphi}(R_1) = \left(\frac{1}{2}\right)^2 \times 0.360 \text{ Pa} = 0.0901 \text{ Pa}.$$

5.4.3

Heat Transfer between Two Plates

Let us again consider two parallel plates fixed at x = 0 and x = d. The upper is maintained at temperature T_0 , while the lower plate has a different temperature $T_0 + \Delta T$ as is depicted in Figure 5.10. To apply the linearized kinetic equation, it is assumed that $\Delta T \ll T_0$. Then, the quantity $\xi = \Delta T/T_0$ is used as the linearization small parameter. The distance d is assumed to be the characteristic size, so that the rarefaction parameter is given by Eq. (5.60). In this problem, the heat flux q_x and temperature distribution T(x) in the gap between the plates are calculated over the whole range of the gas rarefaction δ . According to the energy conservation law, the heat flux q_x does not depend on the coordinate x.

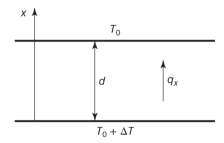


Figure 5.10 Scheme of heat transfer between two plates.

In the free molecular regime ($\delta \rightarrow 0$), the kinetic equation (5.23) can be solved analytically, see Section 4.2 of Ref. [4]. In case of the diffuse gas-surface scattering, this solution provides the following expressions for the heat flux and temperature profile:

$$q_x^{\text{fm}} = \frac{pc_{\text{mp}}}{\sqrt{\pi}} \frac{\Delta T}{T_0}, \quad T = T_0 + \frac{1}{2} \Delta T, \quad \text{for} \quad \delta \to 0,$$
 (5.75)

that is, the temperature is constant over the gap and equal to the mean value of the temperatures of the plates.

In the viscous regime ($\delta \gg 1$), the Fourier equation with the temperature jump boundary condition (5.58) is solved. For the planar heat transfer, this condition reads

$$T - T_0 = \begin{cases} -\zeta_T \ell \frac{\mathrm{d}T}{\mathrm{d}x} & \text{at } x = d, \\ \Delta T + \zeta_T \ell \frac{\mathrm{d}T}{\mathrm{d}x} & \text{at } x = 0. \end{cases}$$
 (5.76)

As a result, the temperature distribution is obtained as

$$T(x) = T_0 + \Delta T \left[1 - \left(\frac{x}{d} + \frac{\zeta_T}{\delta} \right) \left(1 + \frac{2\zeta_T}{\delta} \right)^{-1} \right], \quad \text{for} \quad \delta \gg 1.$$
 (5.77)

The heat flux is calculated from Eq. (5.34) as

$$q_x = \lambda \frac{\Delta T}{d} \left(1 + \frac{2\zeta_T}{\delta} \right)^{-1} = q_x^{\text{fm}} \frac{15\sqrt{\pi}}{8\delta} \left(1 + \frac{2\zeta_T}{\delta} \right)^{-1}, \quad \text{for} \quad \delta \gg 1.$$
 (5.78)

It is evident that the heat flux decreases when the temperature jump condition is applied.

In the transition regime ($\delta \sim 1$), the kinetic equation (5.23) is solved numerically, see, for example, Refs [31–33]. The temperature profile obtained from the S model assuming the complete accommodation is plotted in Figure 5.11. At $\delta = 10$, the solution of the kinetic equation is close to the analytical expression (5.77) obtained on the basis of the Fourier law with the jump boundary condition. In the free molecular regime, the temperature jump solution (5.77) provides the temperature value equal to that given by the free molecular solution (5.75). The numerical solution at $\delta = 0.01$ yields practically the same value. However, in the transition ($\delta = 1$) and near free molecular regimes, the jump solution (5.77) does not work well.

The heat flux q_x is shown in Figure 5.12 and given in Table 5.A.2 as a function of the rarefaction parameter δ . Like for the planar Couette flow, the temperature jump solution (5.78) describes well the numerical data up to the transition regime, $\delta \sim 1$. However, for cylindrical geometry, the jump solution works for a smaller range of the rarefaction parameter.

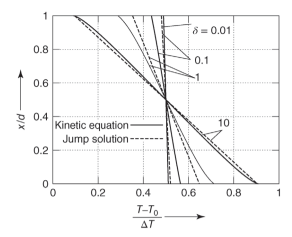


Figure 5.11 Temperature profile T(x) between two plates for various values of the rarefaction δ .

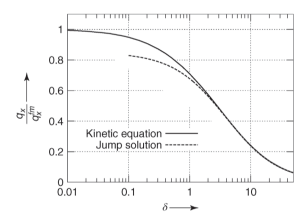


Figure 5.12 Heat flux q_y between two plates versus rarefaction parameter δ . Solid line – numerical solution of S model; dashed line - jump solution, Eq. (5.78).

5.4.4 Heat Transfer between Two Coaxial Cylinders

Consider two coaxial cylinders with radii R_1 and R_2 , as is drawn in Figure 5.13. The external cylinder is maintained at temperature T_0 , while the internal one has a different temperature $T_0 + \Delta T$. Like in the previous case, the parameter $\xi = \Delta T/T_0$ is used to linearize the kinetic equation. The internal cylinder radius R_1 is assumed to be the characteristic size, so that the rarefaction parameter is given by Eq. (5.66). The quantities of our interest are the temperature distribution T(r) and the radial heat flux q_r determined by the rarefaction parameter δ and by the radius ratio R_2/R_1 . Applying the energy conservation law, it is concluded that the quantity $q_r r$ is constant over the gap between the cylinders.

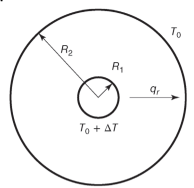


Figure 5.13 Scheme of heat transfer between two cylinders.

In the free molecular regime $(\delta \to 0)$, the temperature distribution and the radial heat flux q_r are calculated analytically as

$$T(r) = T_0 + \frac{\Delta T}{\pi} \arcsin\left(\frac{R_1}{r}\right), \quad \text{for} \quad \delta \to 0,$$
 (5.79)

$$q_{\rm r}^{\rm fm}(r) = \frac{pc_{\rm mp}}{\sqrt{\pi}} \frac{R_1}{r} \frac{\Delta T}{T_0}, \quad \text{for} \quad \delta \to 0.$$
 (5.80)

Note that the free molecular solution is not affected by the external cylinder radius R_2 . At the internal cylinder $r = R_1$, the expression of the radial heat flux (5.80) coincides with the planar heat flux given by Eq. (5.75).

In the viscous regime, the Fourier law is applied with the temperature jump boundary condition (5.58), which for the cylindrical heat flux is quite similar to that of the planar heat transfer, that is,

$$T - T_0 = \begin{cases} -\zeta_T \ell \frac{\mathrm{d}T}{\mathrm{d}r}, & \text{at } r = R_2, \\ \Delta T + \zeta_T \ell \frac{\mathrm{d}T}{\mathrm{d}r}, & \text{at } r = R_1. \end{cases}$$
 (5.81)

Then the temperature profile is obtained as

$$T(r) = T_0 + \Delta T \left[1 - \left(\ln \frac{r}{R_1} + \frac{\zeta_T}{\delta} \right) \mathcal{B} \right], \quad \text{for} \quad \delta \gg 1,$$
 (5.82)

$$\mathcal{B} = \left[\ln \frac{R_2}{R_1} + \frac{\zeta_T}{\delta} \left(1 + \frac{R_1}{R_2} \right) \right]^{-1}.$$
 (5.83)

The heat flux is calculated from Eq. (5.34):

$$q_{\rm r} = \lambda \frac{\Delta T}{r} \mathcal{B} = q_r^{fm} \frac{15\sqrt{\pi}}{8\delta} \mathcal{B}, \quad \text{for} \quad \delta \gg 1.$$
 (5.84)

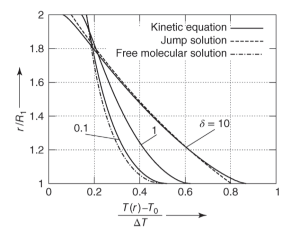


Figure 5.14 Temperature profile T(r) between two cylinders at $R_2/R_1=2$.

When the external cylinder radius is significantly larger than that of the internal cylinder, $R_2 \gg R_1$, then $\mathcal{B} \sim \left| \ln \frac{R_2}{R_1} \right|^{-1}$, this means that, the influence of the external cylinder does not vanish as one could expect.

In the transition regime ($\delta \sim 1$), the kinetic equation (5.23) is solved numerically. The temperature distribution obtained from the S model assuming the complete accommodation [34] is shown in Figure 5.14 for some values of the rarefaction parameter δ and at $R_2/R_1=2$. For $\delta=10$, the numerical results are in a good agreement with the jump solution (5.82). The profile at $\delta = 0.1$ is close to the free molecular solution (5.79). In the transition regime ($\delta = 1$) the numerical solution differs significantly from both free molecular and jump solutions.

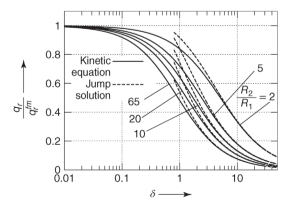


Figure 5.15 Heat flux q_r between two cylinders versus rarefaction parameter δ and radius ratio R_2/R_1 . Solid line – numerical solution of S model [34]; dashed line – jump solution, Eq. (5.84).

Table 5.3 Function $Q(\delta)$.

δ	1	2	5	10	20
$\mathcal Q$	2.72	1.25	0.450	0.212	0.0986

The radial heat flux q_r is presented in Figure 5.15 and in Table 5.A.2. In this case, the temperature jump solution does not provide a good approximation up to the transition regime. It works well up to reasonably large values of the rarefaction parameter, that is, for $\delta > 5$. Unlike the cylindrical Couette flow, the dependence of the radial heat flux q_r on the radius ratio R_2/R_1 is strong. In Ref. [34], the following asymptotic behavior of q_r under the condition $\delta_{R_1}^{R_2} \gg 1$ was obtained:

$$q_{\rm r} = \lambda \frac{\Delta T}{r} \left[\mathcal{Q}(\delta) + \ln \frac{R_2}{R_1} \right]^{-1}, \quad \text{for} \quad \delta \frac{R_2}{R_1} \gg 1,$$
 (5.85)

where the function $\mathcal{Q}(\delta)$ presented in Table 5.3 was calculated numerically. Comparing Eq. (5.85) with Eqs. (5.83) and (5.84), one derives the asymptotic behavior of $\mathcal{Q}(\delta)$ when $\delta \to \infty$, that is, $\mathcal{Q}(\delta) \to \zeta_{\mathrm{T}}/\delta$.

A comparison of the numerical results based on the CL scattering kernel (5.47) with experimental data reported in Ref. [14] is performed in Figure 5.16. The comparison shows that heavy gases such as argon, krypton, and xenon interact diffusely, that is, $\sigma_{\rm t}=1$ and $a_n=1$, with a surface, while light gases such as helium and neon represent a significant deviation from the complete accommodation. The corresponding values of the accommodation coefficients $\sigma_{\rm t}$ and a_n are given in Table 5.2.

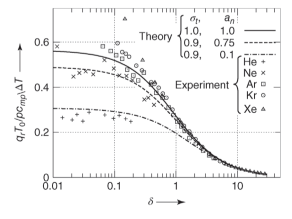


Figure 5.16 Heat flux q_r between two cylinders versus rarefaction parameter δ for $R_2/R_1 = 65$. Curves – theoretical results [34] based on S model and CL scattering law (5.47); symbols – experimental data [14].

Exercise 5.3 Consider helium confined between two coaxial cylinders, as is shown in Figure 5.13. Calculate the radial heat flux q_r on the internal cylinder $(r = R_1)$ under the following conditions: radii $R_1 = 0.1$ mm and $R_2 = 100R_1$, temperature $T_0 = 293$ K, and temperature difference $\Delta T = 5$ K. Use M = 0.004kg mol⁻¹, $\eta = 19.7 \times 10^{-6}$ Pa s, $\lambda = 0.154$ W m⁻¹ K⁻¹, and $\zeta_T = 1.175$. Consider two values of the gas pressure: (a) p = 3 Pa and (b) $p = 3 \times 10^3$ Pa.

Calculation of the most probable speed by Eq. (3.42):

$$c_{\rm mp} = \sqrt{\frac{2 \times 8.31 \,\mathrm{J \, mol^{-1} \, K^{-1} \times 293 \, K}}{0.004 \,\mathrm{kg \, mol^{-1}}}} = 1103 \,\mathrm{m \, s^{-1}}.$$

a) Calculation of the rarefaction parameter for p = 3 Pa using (5.66):

$$\delta = \frac{10^{-4} \text{ m} \times 3 \text{ Pa}}{19.7 \times 10^{-6} \text{ Pa s} \times 1103 \text{ m s}^{-1}} = 0.0138.$$

Since $\delta \ll 1$, Eq. (5.80) is used:

$$q_{\rm r} = \frac{pc_{\rm mp}}{\sqrt{\pi}} \frac{\Delta T}{T_0} = \frac{3 \text{ Pa} \times 1103 \text{ m s}^{-1}}{1.77} \frac{5 \text{ K}}{293 \text{ K}} = 31.9 \text{ W m}^{-2}.$$

b) Calculation of the rarefaction parameter for $p = 3 \times 10^3$ Pa using (5.66):

$$\delta = \frac{10^{-4} \text{ m} \times 3 \times 10^{3} \text{ Pa}}{19.7 \times 10^{-6} \text{ Pa s} \times 1103 \text{ m s}^{-1}} = 13.8.$$

Calculation of \mathcal{B} by Eq. (5.83):

$$\mathcal{B} = \left[\ln 100 + \frac{1.175}{13.8} \times 101 \right]^{-1} = 0.213.$$

Since $\delta \gg 1$, Eq. (5.84) is used:

$$q_{\rm r} = 0.154 \,\mathrm{W m^{-1} K^{-1}} \frac{5 \,\mathrm{K}}{10^{-4} \,\mathrm{m}} \times 0.213 = 1.64 \,\mathrm{kW m^{-2}}.$$

5.5

Flows Through Long Pipes

A gas flow through pipes is a most usual problem that one deals with in vacuum technology. This section contains analytical and numerical data on flow fields and mass flow rates for various shapes of pipes over the whole range of the gas rarefaction and for various conditions at the pipe ends.

5.5.1

Definitions

Consider long pipes with two types of cross sections: (i) rectangular with a height a and width b, as is depicted in Figure 5.17 and (ii) cylindrical with the

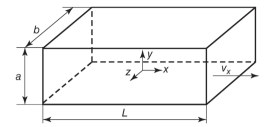


Figure 5.17 Scheme of flow through a channel.

radius a, as is drawn in Figure 5.18. In both cases, a is adopted as the characteristic size so that the rarefaction parameter is defined as

$$\delta = \frac{ap}{\eta c_{\rm mp}}.\tag{5.86}$$

Furthermore, the word "pipe" will be used for all kinds of cross-sectional shapes. However, the rectangular cross-sectional pipe will be referred to as "channel," while the pipe with the cylindrical cross section will be called "tube."

We assume the pipe length L to be significantly larger than its cross-sectional size, that is, $L \gg a, b$. This assumption allows us to neglect end effects and to consider only the *x*-component of the bulk velocity.

A rarefied gas can flow along the pipe due to small longitudinal gradients of pressure p and temperature T denoted as

$$\xi_{\rm P} = \frac{a}{\rho} \frac{\mathrm{d}p}{\mathrm{d}x}, \quad \xi_{\rm T} = \frac{a}{T} \frac{\mathrm{d}T}{\mathrm{d}x},$$
 (5.87)

respectively. Since the gradients $\xi_{\rm P}$ and $\xi_{\rm T}$ are small, the mass flow rate depends linearly on them, that is,

$$\dot{m} = \frac{Ap}{c_{\rm mp}} (-G_{\rm P} \xi_{\rm P} + G_{\rm T} \xi_{\rm T}),$$
 (5.88)

where *A* is the cross-sectional area, that is,

$$A^{\text{ch}} = ab, \quad A^{\text{tb}} = \pi a^2$$
 (5.89)

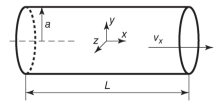


Figure 5.18 Scheme of flow through a tube.

for channel and tube, respectively. Note that the superscripts ch or tb mean that the quantity corresponds to channel or tube, respectively. If the superscript is omitted, the quantity is referred to both channel and tube. The coefficient G_P describes the gas flow induced by a pressure gradient and is called the Poiseuille coefficient. A temperature gradient can also cause a flow of rarefied gas. If the pressure is constant along a pipe, that is, $\xi_{\rm P}=0$, then the gas flows in the direction of the temperature gradient, that is, from a cold to a hot region. This phenomenon is called thermal creep and the quantity G_T is called the thermal creep coefficient. The coefficients G_P and G_T are introduced so as to be always positive. They are calculated from the linearized kinetic equation (5.23) using the gradients ξ_P and ξ_T as the small parameters of the linearization. The coefficients $G_{\rm P}$ and $G_{\rm T}$ are determined by the rarefaction parameter δ . The details of such calculations can be found in Ref. [5]. In the following sections, some recommended data on the coefficients G_P and G_T are given.

5.5.2

Free Molecular Regime

In the free molecular regime ($\delta \rightarrow 0$), the kinetic equation (5.23) is integrated analytically, then the perturbation function is substituted into Eq. (5.26) to calculate the velocity profile. In case of the channel flow it reads [35,36]

$$\nu_{x}(y,z) = \frac{c_{\text{mp}}}{8\sqrt{\pi}} \left(-\xi_{P} + \frac{1}{2}\xi_{T} \right) \sum_{i=1}^{2} \sum_{j=1}^{2} \left(\zeta_{j} \ln \frac{C_{ij} + \eta_{i}}{C_{ij} - \eta_{i}} + \eta_{i} \ln \frac{C_{ij} + \zeta_{j}}{C_{ij} - \zeta_{j}} \right), \tag{5.90}$$

where

$$\eta_i = \frac{1}{2} + (-1)^i \frac{y}{a}, \quad \zeta_j = \frac{b}{2a} + (-1)^j \frac{z}{a}, \quad C_{ij} = \sqrt{\eta_i^2 + \zeta_j^2}.$$
(5.91)

The velocity profile for the tube flow is expressed as

$$\nu_x(r) = \frac{c_{\rm mp}}{\sqrt{\pi}} \left(-\xi_{\rm P} + \frac{1}{2} \xi_{\rm T} \right) \int_0^{\pi/2} \sqrt{1 - (r \sin \phi)^2} \, d\phi. \tag{5.92}$$

Integrating the velocity over the cross section, the coefficients G_P and G_T are obtained. As a result, the Poiseuille coefficient G_p^{ch} for the channel flow reads [36]

$$G_{\rm P} = \frac{1}{\sqrt{\pi}} \left[\frac{b}{a} \left(\ln \mathcal{F} \left(\frac{a}{b} \right) - \frac{1}{3\mathcal{F}(b/a)} \right) + \ln \mathcal{F} \left(\frac{b}{a} \right) - \frac{1}{3\mathcal{F}(a/b)} \right],\tag{5.93}$$

where $\mathcal{F}(x) = x + \sqrt{1 + x^2}$. The numerical values of G_p^{ch} based on this expression are given in Table 5.4 for some aspect ratios b/a. If the channel is

	b/a = 1	2	5	10	50	100	∞
$G_{\rm P}^{\rm ch}(\delta=0)$	0.839	1.152	1.618	1.991	2.884	3.273	∞
\mathcal{H}	0.422	0.686	0.874	0.937	0.989	0.994	1.0
$\mathcal S$	0.562	0.749	0.899	0.949	0.990	0.994	1.0

Table 5.4 Coefficients $G_p^{ch}(\delta = 0)$, H, and S versus aspect ratio b/a.

sufficiently wide, that is, $b \gg a$, then the expression (5.93) is simplified to

$$G_{\rm P}^{\rm ch} = \frac{1}{\sqrt{\pi}} \left(\ln \frac{2b}{a} + \frac{1}{2} \right), \quad \text{for} \quad b \gg a.$$
 (5.94)

In the case of a tube, the expression of the Poiseuille coefficient $G_{\mathrm{P}}^{\mathrm{tb}}$ is quite simple

$$G_{\rm p}^{\rm tb} = \frac{8}{3\sqrt{\pi}}.$$
 (5.95)

The thermal creep coefficient G_T for any kind of pipe is given by

$$G_{\rm T} = \frac{G_{\rm P}}{2}.\tag{5.96}$$

5.5.3

Slip Flow Regime

To calculate the velocity profiles and the coefficients G_P and G_T in the viscous regime ($\delta \gg 1$), the Navier–Stokes equation is solved with the velocity slip boundary conditions (5.53) and (5.56). For the channel flow, the velocity profile reads

$$\nu_{x}(y,z) = -\frac{ap}{2\eta} \left[\frac{1}{4} - \left(\frac{y}{a} \right)^{2} - \sum_{i=0}^{\infty} \frac{(-1)^{i}}{\mu_{i}^{3}} \frac{\cosh(2\mu_{i}z/a)\cos(2\mu_{i}y/a)}{\cosh(\mu_{i}b/a)} + \frac{\beta_{P}}{\delta} s(y,z) \right] \xi_{P}$$

$$+ \frac{c_{mp}\beta_{T}}{2\delta} \xi_{T}, \quad \mu_{i} = \left(i + \frac{1}{2} \right) \pi,$$
(5.97)

where the function s(y, z) is calculated numerically [35] and analytically [37] from the slip boundary condition. In the case of a wide channel, $b \gg a$, the velocity profile is simplified:

$$\nu_x(y) = -\frac{ap}{2\mu} \left[\frac{1}{4} - \left(\frac{y}{a} \right)^2 + \frac{\beta_P}{\delta} \right] \xi_P + \frac{c_{mp}\beta_T}{2\delta} \xi_T.$$
 (5.98)

For tube flow, the profile is also simple:

$$\nu_x(r) = -\frac{ap}{2\mu} \left[1 - \left(\frac{r}{a}\right)^2 + \frac{\beta_P}{\delta} \right] \xi_P + \frac{c_{\rm mp}\beta_T}{2\delta} \xi_T.$$
 (5.99)

Integrating the velocity over the cross section of the pipe yields the coefficients G_P and G_T . The Poiseuille coefficient for the channel flow reads [35,37]

$$G_{\rm P}^{\rm ch} = \frac{\delta}{6} \mathcal{H} + \beta_{\rm P} S,\tag{5.100}$$

where

$$\mathcal{H} = 1 - 6\frac{a}{b} \sum_{i=0}^{\infty} \frac{\tanh(\mu_i b/a)}{\mu_i^5}, \quad \mathcal{S} = \frac{4}{3} \mathcal{H} - 2\left(1 - \frac{a}{b}\right) \sum_{n=0}^{\infty} \frac{\tanh(\mu_i b/a)}{\mu_i^4}.$$
(5.101)

Some numerical values of \mathcal{H} and \mathcal{S} are given in Table 5.4. If the channel is wide, that is, $b \gg a$, then the expression (5.100) is reduced to

$$\lim_{b/a \to \infty} G_{\rm P}^{\rm ch} = \frac{\delta}{6} + \beta_{\rm P}, \quad \text{for} \quad b \gg a. \tag{5.102}$$

For the tube flow, the Poiseuille coefficient is calculated as

$$G_{\rm P}^{\rm tb} = \frac{\delta}{4} + \beta_{\rm P}.\tag{5.103}$$

The thermal creep coefficient G_T does not depend on the cross section, and for both channel and tube, it takes the form

$$G_{\rm T} = \frac{\beta_{\rm T}}{\delta}.\tag{5.104}$$

Note that expressions (5.100), (5.103), and (5.104) are valid for any kind of slip coefficients $\beta_{\rm P}$ and $\beta_{\rm T}$, including those obtained for the nondiffuse gas–surface interaction in Ref. [21] and for gaseous mixtures in Refs [22,24].

5.5.4

Transitional Regime

To calculate the velocity profiles and the coefficients G_P and G_T in the transitional regime ($\delta \sim 1$), the linearized kinetic equation (5.23) is applied using the gradients ξ_P and ξ_T as the small parameters. The details of such calculations can be found in Refs [5,38-40].

The velocity profile $v_x(y)$ for the flow through a wide channel, $b \gg a$, due to the pressure gradient ξ_P is plotted in Figure 5.19 by solid lines. For the large value of the rarefaction parameter ($\delta = 10$), the numerical solution is very close to that of the slip (5.99), plotted by a dashed line. The profile becomes flatter by decreasing the rarefaction parameter δ . However, the magnitude of the bulk velocity increases when δ tends to zero. The similar behavior is observed for the velocity profile $v_x(r)$ in the tube plotted in Figure 5.20 with the difference that for the small values of the rarefaction parameter $\delta < 1$, the speed magnitude practically does not change and is close to that corresponding to the free molecular profile given by Eq. (5.92).

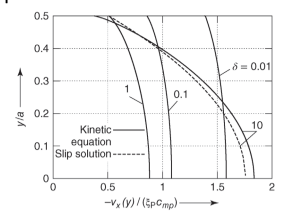


Figure 5.19 Velocity profile $v_x(y)$ for planar Poiseuille flow.

The velocity profiles due to the temperature gradient ξ_{T} are shown in Figures 5.21 and 5.22 for the flows in channel and tube, respectively. The behaviors of both profiles are similar to each other. They are more or less flat and their magnitudes decrease by increasing the rarefaction parameter δ . For $\delta = 10$, the velocity magnitude obtained numerically is close to that obtained from the slip solutions (5.98) and (5.99) only in the pipe axis. Near the pipe wall, the numerical profiles differ from the corresponding slip solution.

The Poiseuille coefficient $G_{\rm P}^{\rm ch}$ for the channel flow obtained from the linearized BGK model in Ref. [35] is presented in Figure 5.23 and in Table 5.A.3. For all values of the aspect ratio b/a, the coefficient G_P^{ch} has the Knudsen minimum near the point $\delta \approx 1$. For the square channel (b/a = 1), the minimum is rather shallow, while for large values of the aspect ratio, $b/a \ge 10$, the Knudsen

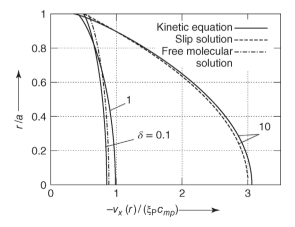


Figure 5.20 Velocity profile $v_x(r)$ for cylindrical Poiseuille flow.

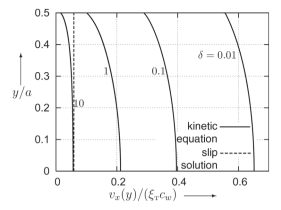


Figure 5.21 Velocity profile $v_x(y)$ for planar thermal creep.

minimum is deep. The existence of the minimum is explained by the fact that in the free molecular regime ($\delta \rightarrow 0$), there are many particles moving long distances parallel to the channel walls without undergoing any strikes. They contribute significantly to the mass flow rate. However, when the intermolecular collisions occur with a small frequency, that is, when $\delta < 1$, then the particles moving along the wall are scattered and they cannot travel a long distance. Thus, their contribution to the mass flow rate decreases. If the intermolecular collisions are quite frequent, that is, $\delta > 1$, the particles begin to drag each other and the mass flow rate increases by increasing the rarefaction parameter. Thus, in the transition regime, the flow rate has a minimum because the scattering effect is still significant, but the drag phenomenon is not so strong.

The slip solution (5.100) presented in Figure 5.23 by dashed lines works well in the range of $\delta \geq 10$. For the small values of the aspect ratio $b/a \leq 10$, the numerical solution at $\delta = 0.001$ is close to the corresponding free

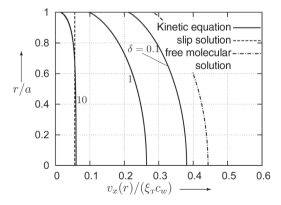


Figure 5.22 Velocity profile $v_x(r)$ for cylindrical thermal creep.

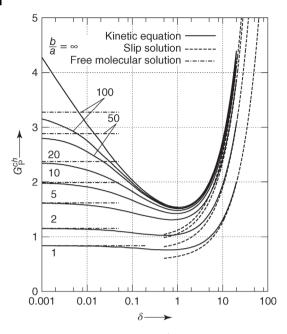


Figure 5.23 Poiseuille coefficient G_p^{ch} versus rarefaction parameter δ and aspect ratio b/a. Solid line – numerical solution of BGK model [35]; dashed line – slip solution, Eq. (5.100); pointdashed line – free molecular solution, Eq. (5.93).

molecular value of $G_{\rm P}^{\rm ch}$ given by Eq. (5.93); while for the large values b/a>20, the numerical solution is still far from the free molecular value.

The thermal creep coefficient for the channel $G_{\rm T}^{\rm ch}$ is shown in Figure 5.24 and in Table 5.A.4. It vanishes in the viscous regime $(\delta \to \infty)$ in accordance with

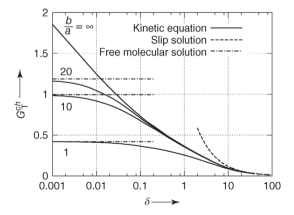


Figure 5.24 Thermal creep coefficient G_T^{ch} versus rarefaction parameter δ and aspect ratio b/a. Solid line – numerical solution of S model [42]; dashed line – slip solution, Eq. (5.104); point line – free molecular solution, Eqs. (5.93) and (5.96).

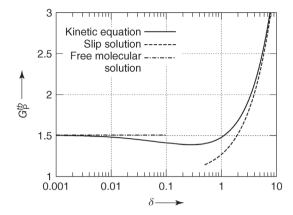


Figure 5.25 Poiseuille coefficient G_P^{tb} versus rarefaction parameter δ. Solid line – numerical solution of S model [41]; dashed line – slip solution, Eq. (5.103); point line – free molecular solution, Eq. (5.95).

Eq. (5.104), and in the free molecular regime it tends to a constant value given by Eqs. (5.93) and (5.96). Note that both coefficients $G_{\rm P}^{\rm ch}$ and $G_{\rm T}^{\rm ch}$ have a singularity, that is, they tend to infinity, at $\delta \to 0$ and $b/a \to \infty$, which is related to the degenerated geometry. However, in practice, the aspect ratio b/a is always finite, so the coefficients $G_{\rm P}^{\rm ch}$ are never infinite.

so the coefficients $G_{\rm p}^{\rm ch}$ and $G_{\rm p}^{\rm ch}$ are never infinite. The coefficients $G_{\rm p}^{\rm th}$ and $G_{\rm T}^{\rm ch}$ for the tube flow obtained from the linearized S model in Ref. [41] are presented in Figures 5.25 and 5.26, respectively. They are also given in Tables 5.A.3 and 5.A.4, respectively. As for the channel, the Poiseuille coefficient $G_{\rm p}^{\rm th}$ also has a small minimum in the transition regime ($\delta \approx 1$). Its variation near the free molecular regime is very small. For large values of the

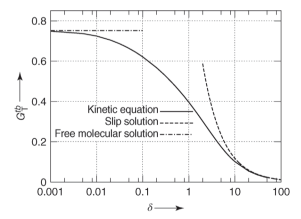


Figure 5.26 Thermal creep coefficient G_T^{tb} versus rarefaction parameter δ. Solid line – numerical solution of S model [41]; dashed line – slip solution, Eq. (5.104); point line – free molecular solution, Eqs. (5.95) and (5.96).

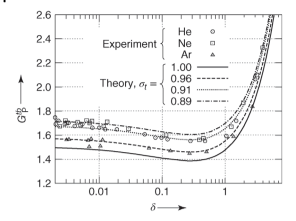


Figure 5.27 Poiseuille coefficient G_p^{tb} versus rarefaction parameter δ . Curves – theoretical results [40] based on S model and CL scattering law (5.47) assuming $a_n = 1$; symbols – experimental data [13].

rarefaction parameter δ , the numerical solution tends to the analytical expression (5.103). The thermal creep coefficient G_T^{tb} vanishes in the viscous limit $(\delta \to \infty)$ according to Eq. (5.104) and it tends to the constant value given by Eqs. (5.95) and (5.96) in the free molecular regime ($\delta \to 0$). By combining the limit solutions (5.95) and (5.103) with the numerical results, the following interpolated formula was obtained:

$$G_{\rm p}^{\rm tb} = \frac{8}{3\sqrt{\pi}} \frac{1 + 0.04\delta^{0.7} \ln \delta}{1 + 0.78\delta^{0.8}} + \left(\frac{\delta}{4} + 1.018\right) \frac{\delta}{1 + \delta},\tag{5.105}$$

by the least-squares method. It represents the numerical data plotted in Figure 5.25 and given in Table 5.A.3 with an accuracy less than 0.4%.

Numerical data for a nondiffuse gas-surface interaction can be found in Refs [39,40] where the CL scattering kernel was applied. A comparison of these results with experimental data reported in Ref. [13] is given in Figure 5.27. It can be seen that the experimental values of the flow rate for the light gases (He and Ne) are in good agreement with the theoretical results corresponding to $\sigma_t \approx 0.9$, while the heavier gas (Ar) undergoes practically diffuse scattering, that is, $\sigma_{\rm t} \approx 1$.

Flows of gaseous mixtures through long pipes of different cross sections are considered in Refs [43-45].

5.5.5

Arbitrary Pressure and Temperature Drops

In the previous sections, the flow rate was calculated as a function of the local gradients of pressure ξ_P and temperature ξ_T . However, in practice, these gradients are unknown, but the pressures and temperatures on the pipe ends are

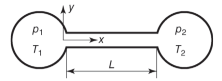


Figure 5.28 Scheme of flows at arbitrary pressure and temperature drops.

measured. In this section, a methodology of flow rate calculations as a function of these pressures and temperatures is described.

Consider two chambers containing a gas and connected by a pipe of length L, as is depicted in Figure 5.28. The gas in the left chamber is maintained at a pressure p_1 and temperature T_1 , while in the right chamber the pressure is p_2 and the temperature is T_2 . The temperature distribution along the channel/tube is denoted as $T_w(x)$ and satisfies the conditions $T_w(0) = T_1$ and $T_w(L) = T_2$.

To calculate the flow rate between the chambers as a function of the pressures p_1 and p_2 and the temperatures T_1 and T_2 , two rarefaction parameters are introduced:

$$\delta_1 = \frac{ap_1}{\eta_1 c_{\text{mp1}}}, \quad \delta_2 = \frac{ap_2}{\eta_2 c_{\text{mp2}}}, \tag{5.106}$$

where the viscosities η_1 , η_2 and most probable speeds c_{mp1} , c_{mp2} correspond to the temperatures T_1 and T_2 , respectively. The results are expressed via the reduced flow rate G related to the mass flow rate as

$$\dot{m} = \frac{ap_1 A}{Lc_{\rm mpl}}G,\tag{5.107}$$

where A is the cross-sectional area.

Since we assume the pipe to be long, $L \gg a, b$, the pressure and temperature gradients are small in each cross section and Eq. (5.88) is valid locally. Combining this equation with Eq. (5.107) and considering that the gas temperature in each section is equal to the pipe wall temperature $T_{\rm w}$, the differential equation for the local pressure p(x) is obtained:

$$G = \frac{p(x)}{p_1} \sqrt{\frac{T_1}{T_{\mathrm{w}}(x)}} \left[-G_{\mathrm{P}}(\delta) \frac{L}{p} \frac{\mathrm{d}p}{\mathrm{d}x} + G_{\mathrm{T}}(\delta) \frac{L}{T_{\mathrm{w}}} \frac{\mathrm{d}T_{\mathrm{w}}}{\mathrm{d}x} \right],\tag{5.108}$$

where $G_{\rm P}(\delta)$ and $G_{\rm T}(\delta)$ are functions of the local rarefaction parameter δ expressed via the local pressure p(x) and temperature T(x). In general, this equation is solved numerically by a finite difference method.

If the flow rate G and the lower pressure p_2 are known, the integration of Eq. (5.108) is realized from x = L to x = 0 with the boundary condition $p(L) = p_2$. As a result of the integration, the pressure p_1 is obtained. If the pressures p_1 and p_2 are known, the quantity G is fitted to satisfy the boundary conditions $p(0) = p_1$ and $p(L) = p_2$. Some particular examples of application of Eq. (5.108) are given in the following section.

Isothermal Flows

First, let us consider an isothermal flow, that is, $T_w = T_1 = T_2$. Then the integral equation (5.108) is simplified and the reduced flow rate G is calculated directly via G_P as

$$G(\delta_1, \delta_2) = \frac{1}{\delta_1} \int_{\delta_1}^{\delta_1} G_{\mathcal{P}}(\delta) d\delta.$$
 (5.109)

Once the function $G_P = G_P(\delta)$ is known, the integration (5.109) is easily performed. Some examples of such integrations can be found in Ref. [46] for tubes and in Ref. [35] for channels. Substituting (5.109) into Eq. (5.107) and using Eq. (5.106), the mass flow rate is related directly to the Poiseuille coefficient G_P as

$$\dot{m} = \frac{A\eta}{L} \int_{\delta_2}^{\delta_1} G_{\rm P}(\delta) \mathrm{d}\delta. \tag{5.110}$$

In the case of a square channel b/a = 1 and a cylindrical tube, the approximation

$$G(\delta_1, \delta_2) = \frac{\delta_1 - \delta_2}{\delta_1} G_P\left(\frac{\delta_1 + \delta_2}{2}\right)$$
 (5.111)

provides a good accuracy, that is, the disagreement between the exact integration (5.109) and approximate formula (5.111) does not exceed 2%. For a channel with a large aspect ratio, say b/a = 100, Eq. (5.111) provides an accuracy of about 6%. If the approximation (5.111) is used, the mass flow rate is calculated as

$$\dot{m} = \frac{A\mu}{L}(\delta_1 - \delta_2)G_P(\overline{\delta}) = \frac{aA(p_1 - p_2)}{c_{mn1}L}G_P(\overline{\delta}), \quad \overline{\delta} = \frac{\delta_1 + \delta_2}{2}.$$
 (5.112)

If the pressure drop is small, $p_1 - p_2 \ll p_1$, then the pressure gradient ξ_P is constant and the mass flow rate is calculated directly from Eq. (5.88):

$$\dot{m} = \frac{aA(p_1 - p_2)}{c_{\text{mpl}}L}G_P(\delta_1), \quad \text{for} \quad p_1 - p_2 \ll p_1.$$
 (5.113)

When the flow rate G is known, the pressure distribution along a pipe can be calculated integrating Eq. (5.108) from any intermediate value of δ , that is,

$$x = \frac{L}{G(\delta_1, \delta_2)} \frac{1}{\delta_1} \int_{\delta}^{\delta_1} G_{\mathbf{P}}(\delta) d\delta, \tag{5.114}$$

where $\delta_2 \le \delta \le \delta_1$. This equation provides the function $x = x(\delta)$, which is inverted into $\delta = \delta(x)$. Since $p(x)/p_1 = \delta(x)/\delta_1$, the pressure distribution p(x) is known. Some typical distributions corresponding to the case $\delta_2 = 0$ are shown in Figure 5.29. In the transition regime ($\delta_1 = 1$), the density distribution is linear. The same distribution is observed in the free molecular regime ($\delta_1 \ll 1$). For high values of rarefaction ($\delta > 10$), the density linearly depends on the coordinate x in the most part of the tube and then it sharply decreases up to zero near the tube exit.

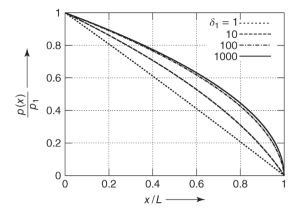


Figure 5.29 Pressure distribution along tube at $\delta_2 = 0$.

Nonisothermal Flows

If the temperatures T_1 and T_2 are different, Eq. (5.108) should be solved numerically. To calculate the local rarefaction parameter $\delta(x)$, its dependence on the temperature should be known, which is determined by the viscosity $\eta = \eta(T)$. To calculate $\delta(x)$, a theoretical expression of the viscosity $\eta(T)$ can be used, for example, that obtained on the basis of the hard-sphere potential. Then the local rarefaction parameter is related to δ_1 as

$$\delta(x) = \delta_1 \frac{T_1}{p_1} \frac{p(x)}{T_w(x)}. (5.115)$$

Usually, a chamber with a lower pressure also has a lower temperature, that is, $T_2 < T_1$. Two examples of the tube flow corresponding to such a situation, namely, $T_2/T_1 = 0.5$ and $T_2/T_1 = 0.25$, under the condition $\delta_2 = 0$ are shown in Figure 5.30. At large values of δ_1 , the flow rate G increases by decreasing the

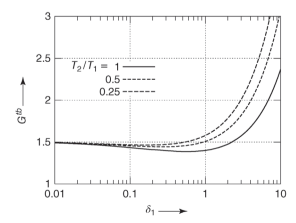


Figure 5.30 Reduced flow rate G^{tb} versus rarefaction parameter δ_1 at $\delta_2 = 0$.

temperature ratio T_2/T_1 , while near the free molecular regime $\delta \ll 1$ the temperature variation does not affect the flow rate G significantly.

If the pressure and temperature drops are small, that is, $p_1 - p_2 \ll p_1$ and $T_1 - T_2 \ll T_1$, then the mass flow rate is calculated directly from Eq. (5.88) as

$$\dot{m} = \frac{aAp_1}{c_{mp1}L} \left[\frac{p_1 - p_2}{p_1} G_P(\delta_1) - \frac{T_1 - T_2}{T_1} G_T(\delta_1) \right],$$
for $p_1 - p_2 \ll p_1$ and $T_1 - T_2 \ll T_1$. (5.116)

Exercise 5.4 Consider helium flowing through a long tube shown in Figure 5.28. Calculate the mass flow rate \dot{m} of helium under the following conditions: tube radius a = 0.5 mm, tube length L = 10 cm, temperatures $T_1 = T_2 = 293 \,\mathrm{K}$, and pressure ratio $p_1/p_2 = 10$. Use $M = 0.004 \,\mathrm{kg/mol}$, $\eta = 19.7 \times 10^{-6}$ Pa s, $\sigma_p = 1.018$. Consider three values of the upflow pressure: (a) $p_1 = 1$ Pa, (b) $p_1 = 200$ Pa, and (c) $p_1 = 1000$ Pa.

Since $p_1/p_2 \gg 1$, Eq. (5.112) must be used. The most probable speed is the same as in Exercise 5.3, that is, $c_{\rm mp} = 1103 \,\mathrm{m \, s^{-1}}$.

a) Calculation of the rarefaction parameters δ_1 and δ_2 for $p_1=1\,\mathrm{Pa}$ using (5.106):

$$\delta_1 = \frac{5 \times 10^{-4} \text{ m} \times 1 \text{ Pa}}{19.7 \times 10^{-6} \text{ Pa s} \times 1103 \text{ m s}^{-1}} = 0.0230,$$

$$\delta_2 = \delta_1 \frac{p_2}{p_1} = 0.00230.$$

Calculation of the average rarefaction parameter:

$$\overline{\delta} = \frac{1}{2}(\delta_1 + \delta_2) = 0.0127.$$

Since $\overline{\delta} \ll 1$, Eq. (5.95) is used to calculate $G_{\rm p}$, that is,

$$G_{\rm p}(\overline{\delta}) = 1.5045.$$

Then, the mass flow rate \dot{m} is calculated using Eq. (5.112):

$$\dot{m} = \frac{3.14 \times (5 \times 10^{-4} \text{ m})^3 \times (1 \text{ Pa} - 0.1 \text{ Pa})}{1103 \text{ m s}^{-1} \times 0.1 \text{ m}} \times 1.5045 = 4.82 \times 10^{-12} \text{ kg s}^{-1}.$$

Using Eq. (4.14), the molar flow rate is calculated:

$$q_{\nu} = \frac{\dot{m}}{M} = \frac{4.82 \times 10^{-12} \ \mathrm{kg \ s^{-1}}}{0.004 \ \mathrm{kg \ mol^{-1}}} = 1.21 \times 10^{-9} \ \mathrm{mol \ s^{-1}}.$$

Using Table 4.1, the flow rate is expressed in other units, for example,

$$q_{pV} = 2437.4 \,\mathrm{J \, mol^{-1}} \times q_{\nu} = 2437.4 \,\mathrm{J \, mol^{-1}} \times 1.21 \times 10^{-9} \,\mathrm{mol \, s^{-1}}$$

= $2.95 \times 10^{-6} \,\mathrm{Pa \, m^3 \, s^{-1}}$ at 20 °C.

b) Calculation of the rarefaction parameters δ_1 and δ_2 for $p_1 = 200 \, \text{Pa}$ using

$$\delta_1 = \frac{5 \times 10^{-4} \text{ m} \times 200 \text{ Pa}}{19.7 \times 10^{-6} \text{ Pa s} \times 1103 \text{ m s}^{-1}} = 4.60.$$
$$\delta_2 = \delta_1 \frac{p_2}{p_1} = 0.460.$$

Calculation of the average rarefaction parameter:

$$\overline{\delta} = \frac{1}{2}(\delta_1 + \delta_2) = 2.53.$$

Since $\overline{\delta} \sim 1$, Eq. (5.105) is used to calculate $G_p(\overline{\delta})$:

$$G_{\rm p} = 1.5045 \frac{1 + 0.04 \times 2.53^{0.7} \ln{(2.53)}}{1 + 0.78 \times 2.53^{0.8}} + \left(\frac{2.53}{4} + 1.018\right) \frac{2.53}{1 + 2.53} = 1.79.$$

Substituting G_p into Eq. (5.112), we obtain

$$\dot{m} = \frac{3.14 \times (5 \times 10^{-4} \text{ m})^3 \times (200 \text{ Pa} - 20 \text{ Pa})}{1103 \text{ m s}^{-1} \times 0.1 \text{ m}} \times 1.79 = 1.15 \times 10^{-9} \text{ kg s}^{-1}.$$

Using Eq. (4.14), the molar flow rate is calculated:

$$q_{\nu} = \frac{\dot{m}}{M} = \frac{1.15 \times 10^{-9} \; \mathrm{kg \; s^{-1}}}{0.004 \; \mathrm{kg \; mol^{-1}}} = 2.87 \times 10^{-7} \; \mathrm{mol \; s^{-1}}.$$

Using Table 4.1, the flow rate is expressed in other units:

$$\begin{split} q_{pV} &= 2437.4\,\mathrm{J\,mol^{-1}} \times q_{\nu} = 2437.4\,\mathrm{J\,mol^{-1}} \times 2.87 \times 10^{-7}\,\mathrm{mol\,s^{-1}} \\ &= 7.00 \times 10^{-4}\,\mathrm{Pa\,m^3\,s^{-1}}\,\mathrm{at\,20\,^{\circ}C}. \end{split}$$

c) Calculation of the rarefaction parameters δ_1 and δ_2 for $p_1 = 1000 \, \text{Pa}$ using (5.106):

$$\delta_1 = \frac{5 \times 10^{-4} \text{ m} \times 1000 \text{ Pa}}{19.7 \times 10^{-6} \text{ Pa s} \times 1103 \text{ m s}^{-1}} = 23.0.$$

$$\delta_2 = \delta_1 \frac{p_2}{p_1} = 2.30.$$

Calculation of the average rarefaction parameter:

$$\overline{\delta} = \frac{1}{2}(\delta_1 + \delta_2) = 12.7.$$

Since $\overline{\delta} > 10$, Eq. (5.103) can be used to calculate $G_p(\overline{\delta})$:

$$G_{\rm p}(\overline{\delta}) = \frac{12.7}{4} + 1.018 = 4.19.$$

Substituting $G_{\rm p}$ into Eq. (5.112), we obtain

$$\dot{m} = \frac{3.14 \times (5 \times 10^{-4} \text{ m})^3 \times (1000 \text{ Pa} - 100 \text{ Pa})}{1103 \text{ m s}^{-1} \times 0.1 \text{ m}} \times 4.19 = 1.34 \times 10^{-8} \text{ kg s}^{-1}.$$

Using Eq. (4.14), the molar flow rate is calculated:

$$q_{\nu} = \frac{\dot{m}}{M} = \frac{1.34 \times 10^{-8} \text{ kg s}^{-1}}{0.004 \text{ kg mol}^{-1}} = 3.35 \times 10^{-6} \text{ mol s}^{-1}.$$

Using Table 4.1, the flow rate is expressed in other units:

$$\begin{split} q_{pV} &= 2437.4\,\mathrm{J\,mol^{-1}} \times q_{\nu} = 2437.4\,\mathrm{J\,mol^{-1}} \times 3.35 \times 10^{-6}\,\mathrm{mol\,s^{-1}} \\ &= 8.16 \times 10^{-3}\,\mathrm{Pa\,m^3\,s^{-1}}\,\mathrm{at\,20\,^{\circ}C}. \end{split}$$

5.5.6

Variable Cross Section

Equation (5.108) can be generalized for a pipe of variable cross section. For the sake of simplicity, only the tube flow is considered here. It is assumed that the tube radius gradually depends on the x-coordinate, that is, the derivative da(x)/dx is sufficiently small. In this case, the reduced flow rate G^{tb} is related to the mass flow rate as

$$\dot{m} = \frac{\pi a_1^3 p_1}{L c_{\rm mpl}} G^{\rm tb},\tag{5.117}$$

where $a_1 = a(0)$ is the tube radius at its entrance. Then, with the help of Eq. (5.88), we obtain [47]

$$G^{\text{tb}} = \frac{p(x)}{p_1} \sqrt{\frac{T_1}{T_{\text{w}}(x)}} \left[\frac{a(x)}{a_1} \right]^3 \left(-G_p^{\text{tb}} \frac{L}{p} \frac{dp}{dx} + G_T^{\text{tb}} \frac{L}{T_{\text{w}}} \frac{dT_{\text{w}}}{dx} \right).$$
 (5.118)

Some numerical examples calculated in Ref. [47] on the basis of Eq. (5.118) are given in Table 5.A.5, where a conical tube is considered, that is,

$$a(x) = a_1 + \frac{x}{I}(a_2 - a_1), \tag{5.119}$$

 $a_2 = a(L)$ is the tube radius at its exit. In the case of isothermal flow, that is, $T_{\rm w} = T_1 = T_2$, the integration (5.118) can be performed analytically for such a tube in the free molecular $(\delta_1, \delta_2 \ll 1)$ and viscous regimes $(\delta_1, \delta_2 \gg 1)$, that is,

$$G^{\text{tb}} = \frac{16}{3\sqrt{\pi}} \frac{(a_2/a_1)^2}{1 + a_2/a_1} \left(1 - \frac{p_2}{p_1}\right), \text{ for } \delta_1 \ll 1 \quad \text{and} \quad \delta_2 \ll 1, \tag{5.120}$$

$$G^{\text{tb}} = \frac{3\delta_1}{8} \frac{(a_2/a_1)^3}{1 + a_2/a_1 + (a_2/a_1)^2} \left[1 - \left(\frac{p_2}{p_1}\right)^2 \right], \quad \text{for} \quad \delta_1 \gg 1 \quad \text{and} \quad \delta_2 \gg 1.$$
(5.121)

These expressions give an idea about the influence of the radius variation. Substituting (5.120) and (5.121) into Eq. (5.117), the mass flow rate is obtained as

$$\dot{m} = \frac{16\sqrt{\pi} (a_1 a_2)^2 (p_1 - p_2)}{3 c_{\text{mpl}}(a_1 + a_2)L}, \quad \text{for} \quad \delta_1 \ll 1 \quad \text{and} \quad \delta_2 \ll 1.$$
 (5.122)

$$\dot{m} = \frac{3\pi}{8} \frac{(a_1 a_2)^3 (p_1^2 - p_2^2)}{\eta c_{\text{mpl}}^2 (a_1^2 + a_1 a_2 + a_2^2) L}, \quad \text{for} \quad \delta_1 \gg 1 \quad \text{and} \quad \delta_2 \gg 1,$$
 (5.123)

in the free molecular and viscous regimes, respectively.

In practice, one deals frequently with the situation when $\delta_1 \gg 1$, $\delta_2 \ll 1$, and $a_2 \gg a_1$. The dependence of G^{tb} on the rarefaction parameter δ_1 and radius ratio a_2/a_1 at $\delta_2 = 0$ is shown in Figure 5.31 and Table 5.A.6. It can be seen that at the large values of the ratio a_2/a_1 , the flow rate G^{tb} is proportional to this ratio,

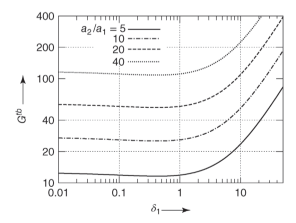


Figure 5.31 Reduced flow rate G^{tb} for a tube with variable cross section versus rarefaction parameter δ_1 at $\delta_2 = 0$.

that is.

$$G^{\text{tb}} \propto \frac{a_2}{a_1}, \quad \text{for} \quad \frac{a_2}{a_1} \gg 1.$$
 (5.124)

Such a proportionality is confirmed by Eqs. (5.120) and (5.121). Thus, for large values of a_2/a_1 , the quantity G^{tb} can be calculated from the data corresponding to $a_2/a_1 = 40$, using the correction factor $\frac{1}{40} \frac{a_2}{a}$.

5.5.7

Thermomolecular Pressure Ratio

The thermal creep causes another interesting phenomenon called the thermomolecular pressure ratio (TPR). Let us assume the system (pipe + chambers) shown in Figure 5.28 to be closed and the temperatures T_1 and T_2 are maintained different. Then the pressures p_1 and p_2 are established so that the net flow rate through the pipe is zero. It happens when the thermal creep caused by the temperature difference is compensated by the Poiseuille flow driven by the pressure drop. The established pressure ratio p_1/p_2 in this state can be related to the maintained temperature ratio as

$$\frac{p_1}{p_2} = \left(\frac{T_1}{T_2}\right)^{\gamma},\tag{5.125}$$

where γ is the exponent of the TPR. The knowledge of this quantity is important if a pressure is measured not directly in a vacuum chamber, but a gauge is connected with the chamber by a pipe. Frequently, in such situations, the gauge temperature can be different from that of the chamber.

If the temperature drop is small, then the TPR exponent is given as

$$\gamma = \frac{G_{\rm T}}{G_{\rm P}}, \quad \text{for} \quad |T_1 - T_2| \ll T_1.$$
 (5.126)

For an arbitrary temperature drop, Eq. (5.108) is used assuming G = 0, that is,

$$\frac{\mathrm{d}p}{\mathrm{d}T_{\mathrm{w}}} = \frac{p}{T_{\mathrm{w}}} \frac{G_{\mathrm{T}}(\delta)}{G_{\mathrm{P}}(\delta)},\tag{5.127}$$

where the local rarefaction parameter δ is determined by the local pressure p and temperature $T_{\rm w}$. In general, this differential equation is solved numerically. Once p_1 and p_2 are known, the TPR exponent is calculated using Eq. (5.125). It is interesting that the TPR exponent γ does not depend on the temperature distribution $T_{\rm w}$, but it is determined only by the rarefaction parameter δ_1 and temperature ratio T_2/T_1 .

In the free molecular limit, $\delta_1 \ll 1$ and $\delta_2 \ll 1$, the integration of Eq. (5.127) can be performed analytically. Since for the diffuse gas-surface interaction G_T = $G_{\rm P}/2$ for $\delta \to 0$,

$$\gamma = \frac{1}{2}$$
, for $\delta_1 \ll 1$ and $\delta_2 \ll 1$ (5.128)

for any temperature ratio T_1/T_2 and for any kind of pipe. If one calculates G_P and $G_{\rm T}$ through a long pipe in the free molecular regime assuming the diffusespecular gas-surface interaction (5.46), one also obtains the relation $G_T = G_P/2$ at any value of the coefficient α . Thus, the value given by Eq. (5.128) is valid for the diffuse-specular reflection too.

According to Ref. [48], the experimental value of γ for krypton is exactly 1/2. However, for helium, the experimental value of γ is lower than 0.5, namely, $\gamma = 0.4$ and 0.464, according to Refs [49] and [48], respectively. Such a deviation from the theoretical value is explained by the nondiffuse interaction of helium with a pipe wall. However, an application of the diffuse-specular kernel (5.128) provides the value $\gamma = 1/2$ for any value of the accommodation coefficient α , while applying the CL kernel (5.47) one can obtain a TPR exponent γ lower than 1/2. This fact indicates that the last kernel (5.47) provides a more physical gas—surface interaction.

In the viscous regime, $\delta_1 \gg 1$ and $\delta_2 \gg 1$, the ratio G_T/G_P has the order δ^{-2} and then

$$\gamma \propto \frac{1}{\delta_1^2}$$
, for $\delta_1 \gg 1$ and $\delta_2 \gg 1$, (5.129)

where the proportionality coefficient is determined by the thermal slip coefficient β_T and by the cross-sectional shape of the pipe.

An empirical formula of the TPR for tube based on experiment data for several species of gases was proposed in Ref. [50]. Considering that the experimental temperature difference was sufficiently small, namely, $(T_2 - T_1)/T_1 = 0.07$, the empirical formula representing all gases used in the measurements [50] can be written in our notations as

$$\gamma = 0.5 \left(1 + 0.0920 \delta^2 + 0.518 \delta + 0.317 \delta^{1/2} \right)^{-1},$$
 for tube at $\frac{|T_2 - T_1|}{T_1} \ll 1.$ (5.130)

The theoretical results [41,42] on the exponent γ corresponding to the diffuse scattering law are plotted in Figure 5.32 for both tube (solid line) and channel (dashed line). It can be seen that in the transitional and viscous regimes, the TPR exponent is affected by the pipe shape and by the value of the temperature ratio T_2/T_1 . The empirical formula (5.130) is plotted in Figure 5.32 by the pointed line and represents good agreement with the theoretical results [41]. For the large temperature ratio $(T_2/T_1 = 3.8)$, the theoretical results [41] agree well with the experimental data [49] for the gas helium presented in Figure 5.32 by circles.

The theoretical values of the exponent γ for tube based on the S model and CL scattering law obtained in Ref. [40] are shown in Figure 5.33 together with the experimental data reported in Ref. [51]. It is evident that the heavy gas (Xe) undergoes the complete accommodation on the tube wall because its experimental values of γ coincide with the theoretical results for $\sigma_t = 1$. The experimental values of γ for the lighter gas (Ar) are in agreement with the theoretical results for $\sigma_t = 0.94$, that is, the deviation from the diffuse scattering is weak.

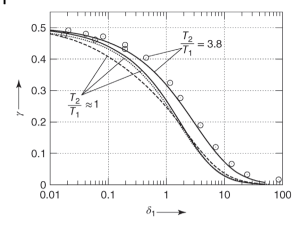


Figure 5.32 TPR exponent γ versus rarefaction parameter δ_1 . Solid line – theoretical results [41] for tube with diffuse scattering law; dashed line – theoretical results [42] for channel at b/a = 10

with diffuse scattering; pointed line - empirical formula (5.130); circles - experimental data for tube [49].

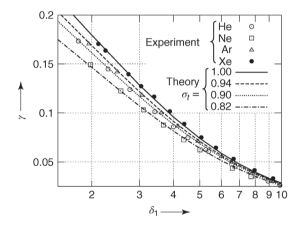


Figure 5.33 TPR exponent γ versus rarefaction parameter δ_1 for tube at $T_2/T_1 \approx 1$. Curves – theoretical results [40] based on CL scattering law (5.47) assuming $a_n = 1$; symbols – experimental data [51].

The light gases He and Ne represent a stronger deviation from the diffuse scattering, that is, $\sigma_t = 0.9$ and 0.82, respectively.

5.6 Flow Through an Orifice

An orifice flow represents a limit opposite to the long pipe, that is, here an infinitesimally thin partition separating chambers is considered. The chambers

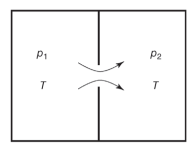


Figure 5.34 Scheme of orifice flow.

contain a gas at the same temperature T, but at different pressures p_1 and p_2 . Without loss of generality, it is assumed that $p_2 < p_1$. The chambers are connected by a circular orifice, which allows the gas to flow as is shown in Figure 5.34. In practice, one is interested in the mass flow rate \dot{m} through the orifice determined by the pressure ratio p_2/p_1 and by the rarefaction parameter δ_1 calculated via the pressure p_1 and via the orifice radius a as

$$\delta = \frac{ap_1}{\eta c_{\rm mp}}.\tag{5.131}$$

In the free molecular regime ($\delta_1 = 0$) at $p_2/p_1 = 0$, the mass flow rate is calculated via the molecular fluxes given by Eq. (5.14):

$$\dot{m}_0 = \pi a^2 m_{\rm P} j_1 = \frac{\sqrt{\pi} a^2}{c_{\rm mp}} p_1, \quad \text{for} \quad \delta_1 \to 0 \quad \text{and} \quad \frac{p_2}{p_1} \to 0,$$
 (5.132)

where j_1 corresponds to the pressure p_1 . In case of arbitrary pressure ratio p_2/p_1 , the mass flow rate is calculated as the difference of two opposite fluxes and reads

$$\dot{m} = \pi a^2 m_{\rm P} (j_1 - j_2) = \dot{m}_0 \left(1 - \frac{p_2}{p_1} \right), \quad \text{for} \quad \delta_1 \to 0,$$
 (5.133)

where j_2 corresponds to the pressure p_2 .

References [52,53] report numerical data on the flow rate through an orifice obtained by the DSMC method based on the hard-sphere potential. Simulations of the same flow based on ab initio potentials [54] showed that the influence of the molecule model does not exceed 1.5% so that the hard-sphere model provides reliable results for orifice flows. The numbers of particles and samples were sufficient to reduce the statistical scattering of the flow rate up to 1%. The dependence of the reduced flow rate \dot{m}/\dot{m}_0 on the rarefaction parameter δ_1 for $p_2/p_1 = 0, 0.1, 0.5, \text{ and } 0.9$ are represented in Figure 5.35. The numerical results are in good agreement with the corresponding experimental data [55,56]. In the free molecular regime ($\delta_1 = 0$), the numerical value of \dot{m}/\dot{m}_0 tends to its theoretical expression (5.133). For $\delta_1 > 100$, the variations of the flow rate are within the numerical accuracy for all pressure ratios considered here. Therefore, the data presented here cover the whole range of the gas rarefaction δ_1 .

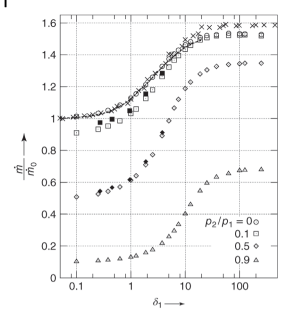


Figure 5.35 Flow rate \dot{m} through an orifice versus rarefaction parameter δ_1 . Open symbols – DSMC simulation [52,53]; filled symbols – experimental data [56]; crosses – experimental data [55]; dashed line – empirical formula, Eq. (5.134) [57].

The following empirical formula was proposed in Ref. [57] for outflow into vacuum, $p_2/p_1 = 0$:

$$\dot{m} = \dot{m}_0 \left(1 + \frac{0.4733 + 0.6005 / \sqrt{\delta_1}}{1 + 4.559 / \delta_1 + 3.094 / \delta_1^2} \right), \quad \text{for} \quad \frac{p_2}{p_1} \to 0$$
and $0 \le \delta_1 \le 20$, (5.134)

which works well in the range of $0 \le \delta_1 \le 20$. This formula is plotted in Figure 5.35 by the dashed line.

In many practical applications, the flow rate is needed only for small values of the gas rarefaction, $\delta_1 \ll 1$. Under such condition, Eq. (5.133) is corrected by a linear term, that is, the flow rate can be written as

$$\dot{m} = \dot{m}_0 \left(1 - \frac{p_2}{p_1} \right) (1 + A\delta_1) \quad \text{for} \quad \delta_1 \ll 1.$$
 (5.135)

The values of the constant \mathcal{A} are given in Table 5.5 for some pressure ratios p_2/p_1 . These values were obtained by the least-squares method on the basis of the numerical results for $0 \le p_2/p_1 \le 0.9$ and on the basis of the experimental data [58] for $p_2/p_1 \approx 1$.

Table 5.5	Coefficient .	$\mathcal A$ in Eq.	(5.135)	versus p_2	p_1 .
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p_2/p_1	0	0.1	0.5	0.9	1
\mathcal{A}	0.13	0.15	0.23	0.31	0.34

5.7 Modeling of Holweck Pump

In this section, the main ideas of how complex flows that usually occur in diverse kinds of pumps are modeled. Such flows can be calculated employing a superposition of several solutions of the kinetic equation. The Holweck pump considered as an example is composed of two coaxial cylinders. One of them has grooves in a spiral form and the other is smooth. A rotation of the smooth cylinder causes a gas flow from a chamber of low pressure to that of high pressure, that is, the pumping effect is induced. Generally, the gas flow through such a pump is three dimensional and requires a lot of computational effort. To reduce the effort, a two-dimensional flow is considered, that is, the groove curvature and the end effects are neglected. More exactly, a plane with regularly distributed grooves is considered. Another surface, which is smooth, moves to the left over the grooved surface and causes an upward gas flow, as shown in Figure 5.36a. The cross section of one groove, that is, AA, is depicted in Figure 5.36b.

The problem solution includes two stages. In the first stage, four independent problems are solved over the whole range of the gas rarefaction: (i) Longitudinal Couette flow, that is, the gas flow due to a surface motion along the z-axis. The coordinate system (x, y, z) is shown in Figure 5.36b. (ii) Longitudinal Poiseuille flow, that is, the gas flow caused by a pressure gradient along the z-axis. (iii) Transversal Couette flow, that is, the gas flow due to a surface motion along the x-axis. (iv) Transversal Poiseuille flow, that is, the gas flow caused by a pressure drop in the x-direction through a pair of groove and ridge. The solution to these four problems is determined by the grove and ridge sizes and by the local rarefaction parameter δ . Usually, this stage takes a long computational time.

In the second stage, a linear superposition of the four solutions obtained previously is realized in accordance with the methodology described in

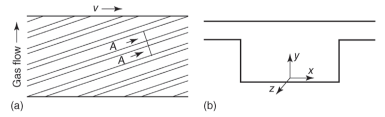


Figure 5.36 Scheme of pump and cross section of groove AA.

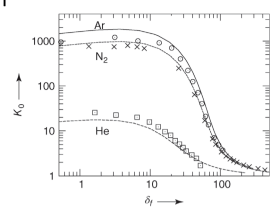


Figure 5.37 Limit compression pressure ratio K_0 versus fore-vacuum rarefaction δ_f [60]. Lines – theoretical results; symbols – experimental data.

Refs [35,41,42,46,59,60]. This stage does not require much computational effort and allows us to easily change many parameters such as groove inclination, forevacuum and high-vacuum pressures, angular velocity of rotating cylinder, species of gas, temperature of the gas, and so on.

Applying the present approach, the compression ratio and pumping speed were calculated. The results related to the limit compression ratio are shown in Figure 5.37, from which it can be seen that the numerical results are in fine agreement with the experimental data. The results for the pumping speed are given in Figure 5.38 in terms of the dimensionless pumping speed defined as

$$G = \frac{S}{a^2 c_{\rm mp} N_{\rm gr}},\tag{5.136}$$

where S is the pumping speed, a is the height of the groove, and $N_{\rm gr}$ is the number of the groove. Physically, G is the dimensionless flow rate through one

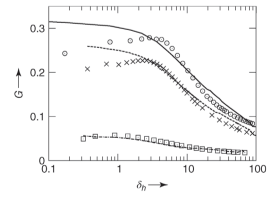


Figure 5.38 Dimensionless pumping speed G versus high-vacuum rarefaction δ_h [60]. Lines – theoretical results; symbols – experimental data.

groove in the vertical direction, see Figure 5.36a. A comparison of the numerical results on G with experimental data shows the efficiency of the methodology based on the superposition of several solutions obtained from the linearized kinetic equation. The details of the numerical calculations and measurements can be found in Ref. [60].

5.8 Appendix A

5.8.1 **Tables**

Table 5.A.1 Shear stress in Couette flow versus rarefaction parameter δ [26,30].

	Planar P_{xz}/P_{xz}^{fm}	Cylindrical $P_{rarphi}/P_{rarphi}^{ m fm}$			
δ		$\frac{R_2}{R_1} = 2$	3	5	
0.01	0.9914	0.9988	0.9987	0.9987	
0.1	0.9258	0.9883	0.9871	0.9866	
1.0	0.6008	0.8811	0.8669	0.8601	
2.0	0.4437	0.7725	0.7465	0.7338	
5.0	0.2523	0.5458	0.5076	0.4874	
10.0	0.1473	0.3540	0.3180	0.3057	
20.0	0.0805	0.2080	0.1816	0.1750	

Table 5.A.2 Heat flux versus rarefaction parameter δ [33,34].

	Planar		Cylindrical $q_{ m r}/q_{ m r}^{ m fm}$					
δ	$\overline{q_x/q_x^{ m fm}}$	$\overline{R_2/R_1=2}$	5	10	20	65		
0.01	0.9939	0.9982	0.9965	0.9954	0.9942	0.9920		
0.1	0.9485	0.9818	0.9653	0.9532	0.9410	0.9190		
1.0	0.7092	0.8393	0.7191	0.6429	0.5753	0.4823		
2.0	0.5736	0.7219	0.5493	0.4576	0.3885	0.3065		
5.0	0.3740	0.5050	0.3121	0.2387	0.1921	0.1437		
10.0	0.2390	0.3334	0.1788	0.1313	0.1034	0.0758		
20.0	0.1390	0.1973	0.0960	0.0688	0.0536	0.0389		

Table 5.A.3 Poiseuille coefficient G_P versus rarefaction parameter δ [35,41].

	Channel G _P ^{ch}							Tube G _P ^{tb}	
δ	b/a = 1	2	5	10	20	50	100	∞	
0	0.8387	1.152	1.618	1.991	2.373	2.884	3.273	∞	1.505
0.001	0.8373	1.150	1.612	1.978	2.344	2.798	3.015		1.501
0.01	0.8315	1.137	1.577	1.910	2.217	2.551	2.695	3.050	1.480
0.02	0.8261	1.125	1.549	1.858	2.130	2.400	2.510	2.711	1.464
0.05	0.8124	1.099	1.492	1.759	1.971	2.149	2.214	2.302	1.434
0.1	0.7958	1.073	1.437	1.665	1.826	1.943	1.983	2.033	1.410
0.2	0.7766	1.046	1.379	1.563	1.678	1.752	1.776	1.808	1.391
0.5	0.7607	1.026	1.319	1.454	1.526	1.569	1.580	1.602	1.401
1.0	0.7660	1.041	1.315	1.424	1.480	1.513	1.520	1.539	1.476
2.0	0.8076	1.115	1.391	1.491	1.541	1.571	1.577	1.595	1.680
5.0	0.9846	1.413	1.753	1.870	1.929	1.962	1.973	1.991	2.367
10.0	1.314	1.955	2.437	2.599	2.683	2.729	2.753	2.769	3.575
20.0	2.000	3.077	3.864	4.121	4.267	4.341	4.368	4.397	6.049

Table 5.A.4 Thermal creep coefficient $G_{\rm T}$ versus rarefaction parameter δ [41,42].

		Channel G_{T}^{ch}					
δ	b/a = 1	10	20	∞			
0	0.4193	0.9955	1.186	∞	0.7523		
0.001	0.4181	0.9839	1.162	1.855	0.7486		
0.01	0.4110	0.9165	1.044	1.246	0.7243		
0.02	0.4037	0.8658	0.9662	1.078	0.7042		
0.05	0.3857	0.7695	0.8291	0.8719	0.6637		
0.1	0.3637	0.6763	0.7089	0.7320	0.6210		
0.2	0.3390	0.5814	0.5968	0.6105	0.5675		
0.5	0.2953	0.4490	0.4553	0.4620	0.4779		
1.0	0.2545	0.3553	0.3593	0.3633	0.3959		
2.0	0.2070	0.2667	0.2693	0.2719	0.3016		
5.0	0.1366	0.1598	0.1609	0.1621	0.1752		
10.0	0.0868	0.0956	0.0961	0.0966	0.1014		
20.0	0.0495	0.0522	0.0524	0.0526	0.0543		

66.21

δ_1	G ^{tb}							
	$p_2/p_1 = 0$	0.01	0.1	0.5	0.9			
0.0	27.35	27.08	24.62	13.68	2.735			
0.01	27.04	26.76	24.29	13.42	2.672			
0.1	25.95	25.67	23.21	12.78	2.547			
1.0	25.99	25.73	23.57	13.80	2.910			
10.0	52.29	52.17	50.66	35.56	8.531			

Table 5.A.5 Reduced flow rate G^{tb} for conical tube versus rarefaction parameter δ_1 and pressure ratio p_2/p_1 at $a_2/a_1 = 10$ and T = const. [47].

Table 5.A.6 Reduced flow rate G^{tb} for conical tube versus rarefaction parameter δ_1 and radius ratio a_2/a_1 at $\delta_2=0$ and T= const. [47].

350.1

263.1

354.2

		$G^{ m tb}$		
δ_1	$a_2/a_1 = 5$	10	20	40
0.01	12.39	27.02	56.54	115.6
0.02	12.29	26.79	56.07	114.7
0.05	12.09	26.35	55.13	112.7
0.1	11.90	25.93	54.25	110.9
0.2	11.72	25.54	53.45	109.3
0.5	11.63	25.34	53.03	108.4
1.0	11.90	25.96	54.35	111.1
2.0	12.89	28.18	59.03	120.7
5.0	16.64	36.59	76.79	157.1
10.0	23.63	52.25	109.8	224.8
20.0	38.34	85.15	179.2	367.0
50.0	83.32	185.6	391.0	800.9

References

100.0

354.5

- 1 Cercignani, C. (1988) The Boltzmann Equation and Its Application, Springer, New York.
- 2 Chapman, S. and Cowling, T.G. (1952) The Mathematical Theory of Non-Uniform Gases, University Press, Cambridge.
- 3 Ferziger, J.H. and Kaper, H.G. (1972) Mathematical Theory of Transport Processes in Gases, North-Holland Publishing Company, Amsterdam.
- 4 Kogan, M.N. (1969) Rarefied Gas Dynamics, Plenum, New York.
- 5 Sharipov, F. and Seleznev, V. (1998) Data on internal rarefied gas flows. J. Phys. Chem. Ref. Data, 27, 657-706.
- 6 Sone, Y. (2002) Kinetic Theory and Fluid Mechanics, Birkhäuser, Boston.
- 7 Pekeris, C.L. and Alterman, Z. (1957) Solution of the Boltzmann-Hilbert integral equation: II. The coefficients of

- viscosity and heat conduction. Proc. Natl. Acad. Sci. USA, 43, 998-1007.
- 8 Bhatnagar, P.L., Gross, E.P., and Krook, M.A. (1954) A model for collision processes in gases. Phys. Rev., 94, 511-525.
- 9 Welander, P. (1954) On the temperature jump in a rarefied gas. Ark. Fys., 7, 507-553.
- 10 Shakhov, E.M. (1968) Generalization of the Krook kinetic relaxation equation. Fluid Dyn., 3, 95-96.
- 11 Sazhin, O.V., Borisov, S.F., and Sharipov, F. (2001) Accommodation coefficient of tangential momentum on atomically clean and contaminated surfaces. J. Vac. Sci. Technol. A, 19, 2499-2503 (erratum: 20 (3), 957 (2002)).
- 12 Cercignani, C. and Lampis, M. (1971) Kinetic model for gas-surface interaction. Transp. Theory Stat. Phys., 1, 101-114.
- 13 Porodnov, B.T., Suetin, P.E., Borisov, S.F., and Akinshin, V.D. (1974) Experimental investigation of rarefied gas flow in different channels. J. Fluid Mech., 64, 417-437.
- 14 Semyonov, Y.G., Borisov, S.F., and Suetin, P.E. (1984) Investigation of heat transfer in rarefied gases over a wide range of Knudsen numbers. Int. J. Heat Mass Transf., 27, 1789-1799.
- 15 Valougeorgis, D. and Naris, S. (2003) Acceleration schemes of the discrete velocity method: Gaseous flows in rectangular microchannels. SIAM J. Sci. Comp., 25, 534-552.
- 16 Sharipov, F.M. and Subbotin, E.A. (1993) On optimization of the discrete velocity method used in rarefied gas dynamics. Z. Angew. Math. Phys., 44, 572-577.
- 17 Bird, G.A. (1994) Molecular Gas Dynamics and the Direct Simulation of Gas Flows, Oxford University Press, Oxford.
- 18 Sharipov, F. and Strapasson, J.L. (2012) Direct simulation Monte Carlo method for an arbitrary intermolecular potential. Phys. Fluids, 24, 011703.
- 19 Albertoni, S., Cercignani, C., and Gotusso, L. (1963) Numerical evaluation of the slip coefficient. Phys. Fluids, 6, 993-996.
- 20 Ohwada, T., Sone, Y., and Aoki, K. (1989) Numerical analysis of the shear

- and thermal creep flows of a rarefied gas over a plane wall on the basis of the linearized Boltzmann equation for hardsphere molecules. Phys. Fluids A, 1, 1588-1599.
- 21 Sharipov, F. (2003) Application of the Cercignani-Lampis scattering kernel to calculations of rarefied gas flows: II. Slip and jump coefficients. Eur. J. Mech. B Fluids, 22, 133-143.
- 22 Sharipov, F. and Kalempa, D. (2003) Velocity slip and temperature jump coefficients for gaseous mixtures: I. Viscous slip coefficient. Phys. Fluids, 15, 1800-1806.
- 23 Loyalka, S.K. (1989) Temperature jump and thermal creep slip: rigid sphere gas. Phys. Fluids A, 1, 403-408.
- 24 Sharipov, F. and Kalempa, D. (2004) Velocity slip and temperature jump coefficients for gaseous mixtures: II. Thermal slip coefficient. Phys. Fluids, 16, 759-764.
- 25 Sharipov, F. and Kalempa, D. (2005) Velocity slip and temperature jump coefficients for gaseous mixtures: IV. Temperature jump coefficient. Int. J. Heat Mass Transf., 48, 1076-1083.
- 26 Cercignani, C. and Pagani, C.D. (1966) Variational approach to boundary value problems in kinetic theory. Phys. Fluids, 9, 1167-1173.
- 27 Sharipov, F., Cumin, L.M.G., and Kalempa, D. (2004) Plane Couette flow of binary gaseous mixture in the whole range of the Knudsen number. Eur. J. Mech. B Fluids, 23, 899-906.
- 28 Sone, Y., Takata, S., and Ohwada, T. (1990) Numerical analysis of the plane Couette flow of a rarefied gas on the basis of the linearized Boltzmann equation for hard-sphere molecules. Eur. J. Mech. B Fluids, 9, 273-288.
- 29 Cercignani, C. and Sernagiotto, F. (1966) Cylindrical Poiseuille flow of a rarefied gas. Phys. Fluids, 9, 40-44.
- 30 Sharipov, F.M. and Kremer, G.M. (1996) Linear Couette flow between two rotating cylinders. Eur. J. Mech. B Fluids, 15, 493-505.
- 31 Bassanini, P., Cercignani, C., and Pagani, C. (1968) Influence of the accommodation coefficient on the heat transfer in a

- rarefied gas. Int. J. Heat Mass Transf., 11, 1359-1369.
- 32 Valougeorgis, D. and Thomas, J.R. (1985) The F_N-method in kinetic theory: II. Heat transfer between parallel plates. Transport Theory Stat. Phys., 14, 497-512.
- 33 Sharipov, F., Cumin, L.M.G., and Kalempa, D. (2007) Heat flux through a binary gaseous mixture over the whole range of the Knudsen number. Physica A, 378, 183-193.
- 34 Sharipov, F. and Bertoldo, G. (2006) Heat transfer through a rarefied gas confined between two coaxial cylinders with high radius ratio. J. Vac. Sci. Technol. A, 24, 2087-2093.
- 35 Sharipov, F. (1999) Rarefied gas flow through a long rectangular channel. J. Vac. Sci. Technol. A, 17, 3062-3066.
- 36 Titarev, V.A. and Shakhov, E.M. (2010) Kinetic analysis of the isothermal flow in a long rectangular microchannel. J. Comput. Math. Math. Phys., 50, 1221-1237.
- 37 Méolans, J., Nacer, M., Rojas, M., Perrier, P., and Graur, I. (2012) Effects of two transversal finite dimensions in long microchannel: analytical approach in slip regime. Phys. Fluids, 24, 112005.
- 38 Valougeorgis, D. and Thomas, J.R. (1986) Exact numerical results for Poiseuille and thermal creep flow in a cylindrical tube. Phys. Fluids, 29, 423-429.
- 39 Sharipov, F. (2002) Application of the Cercignani-Lampis scattering kernel to calculations of rarefied gas flows: I. Plane flow between two parallel plates. Eur. J. Mech. B Fluids, 21, 113-123.
- 40 Sharipov, F. (2003) Application of the Cercignani-Lampis scattering kernel to calculations of rarefied gas flows: III. Poiseuille flow and thermal creep through a long tube. Eur. J. Mech. B Fluids, 22, 145-154.
- 41 Sharipov, F. (1996) Rarefied gas flow through a long tube at any temperature difference. J. Vac. Sci. Technol. A, 14, 2627-2635.
- 42 Sharipov, F. (1999) Non-isothermal gas flow through rectangular microchannels. J. Micromech. Microeng., 9, 394-401.
- 43 Sharipov, F. and Kalempa, D. (2002) Gaseous mixture flow through a long tube

- at arbitrary Knudsen number. J. Vac. Sci. Technol. A. 20, 814-822.
- 44 Naris, S., Valougeorgis, D., Kalempa, D., and Sharipov, F. (2004) Gaseous mixture flow between two parallel plates in the whole range of the gas rarefaction. Physica A, 336, 294-318.
- 45 Naris, S., Valougeorgis, D., Sharipov, F., and Kalempa, D. (2004) Discrete velocity modelling of gaseous mixture flows in MEMS. Superlattices Microstruct., **35**, 629–643.
- 46 Sharipov, F. and Seleznev, V. (1994) Rarefied gas flow through a long tube at any pressure ratio. J. Vac. Sci. Technol. A, 12, 2933-2935.
- 47 Sharipov, F. and Bertoldo, G. (2005) Rarefied gas flow through a long tube of variable radius. J. Vac. Sci. Technol. A, 23, 531-533.
- 48 Seleznev, V.D., Porodnov, B.T., Kulev, A.N., Flyagin, A.G., Kudertzev, A.N., and Obraz, S.P. (1988) Knudsen molecular flow through a channel for small temperature difference at its ends. J. Eng. Phys., 54, 475-480.
- 49 Edmonds, T. and Hobson, J.P. (1965) A study of thermal transpiration using ultrahigh-vacuum techniques. J. Vac. Sci. Technol., 2, 182-197.
- 50 Setina, J. (1999) New approach to corrections for thermal transpiration effects in capacitance diaphragm gauges. Metrologia, 36, 623-626.
- 51 Porodnov, B.T., Kulev, A.N., and Tukhvetov, F.T. (1978) Thermal transpiration in a circular capillary with a small temperature difference. J. Fluid Mech., 88, 609-622.
- 52 Sharipov, F. (2002) Rarefied gas flow into vacuum through a thin orifice: influence of the boundary conditions. AIAA J., 40, 2006-2008.
- 53 Sharipov, F. (2004) Numerical simulation of rarefied gas flow through a thin orifice. J. Fluid Mech., 518, 35-60.
- 54 Sharipov, F. and Strapasson, J.L. (2014) Ab initio simulation of rarefied gas flow through a thin orifice. Vacuum, 109, 246-252.
- 55 Jitschin, W., Ronzheimer, M., and Khodabakhshi, S. (1999) Gas flow measurement by means of orifices

- and Venturi tubes. Vacuum, 53, 181-185.
- 56 Sreekanth, A.K. (1965) Transition flow through short circular tubes. Phys. Fluids, 8, 1951-1956.
- 57 Fujimoto, T. and Usami, M. (1984) Rarefied gas flow through a circular orifice and short tubes. J. Fluids Eng., 106, 367-373.
- 58 Borisov, S.F., Neudachin, I.G., Porodnov, B.T., and Suetin, P.E. (1973) Flow of
- rarefied gases through an orifice for small pressure drop. Zh. Tekh. Fiz., 43, 1735-1739. in Russian].
- 59 Sharipov, F. (1997) Rarefied gas flow through a long tube at arbitrary pressure and temperature drops. J. Vac. Sci. Technol. A, 15, 2434-2436.
- 60 Sharipov, F., Fahrenbach, P., and Zipp, A. (2005) Numerical modeling of Holweck pump. J. Vac. Sci. Technol. A, 23, 1331-1339.

6

Sorption and Diffusion

Dr. Karl Jousten

Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

This chapter explains how molecules stick to a surface and travel through a solid (the material of a vacuum vessel), and furthermore, what consequences this has for the pressure inside a vacuum chamber.

6.1 Sorption Phenomena and the Consequences, Definitions, and Terminology

Atoms or molecules from the gas or vapor phase impinging on a solid surface, referred to as *adsorbent* (Figure 6.1), remain at the surface with a sticking probability $s \le 1$, in other words, are reflected with a probability (1-s). The sticking *adparticles*, termed the *adsorbate*, adhere to the surface either due to *dipole forces* or *van der Waals forces* (*physisorption*), or due to *covalent linkage* (*atomic bonds, chemisorption*). The binding energy resulting from these forces is referred to as adsorption energy $E_{\rm ad}$ (or adsorption heat).

Figure 6.2 shows the potential curves of a molecule A_2 and of two molecules 2A with respect to the distance from the surface. For physisorption, $E_{\rm ad}=E_{\rm p}$. In certain cases of chemisorption of molecular adsorbates (e.g., for H_2 , O_2 , N_2 , but not CO, CO_2), an energy barrier termed activation energy $E_{\rm act}$ must be surpassed [1]. At this point, the considered molecules dissociate into atoms $(xA_2 \to 2A)$, which, subsequently, are chemisorbed. Here, the energy released corresponds to the sum of the activation energy and twice the adsorption energy $(E_{\rm act}+E_{\rm c})$ (Figure 6.2). This type of activated adsorption occurs only for molecules carrying sufficient kinetic energy in the direction perpendicular to the surface when they are still distant, $E_{\rm kin, \ \perp} > E_{\rm act}$. For adparticles to desorb, expenditure of the desorption energy $E_{\rm des}$ is required, which is equal to the adsorption energy $E_{\rm ad}$. Molar desorption energies for physisorption are below $E_{\rm des} \approx 40~{\rm kJ~mol}^{-1}$ (0.4 eV per particle), and for chemisorption, in the range of $E_{\rm des} \approx 80~{\rm to~800~kJ~mol}^{-1}$ (0.8–8 eV per particle). Thus, bonds due to chemisorption are approximately 10 times stronger than bonds produced by physisorption. Table 6.1 lists a number of

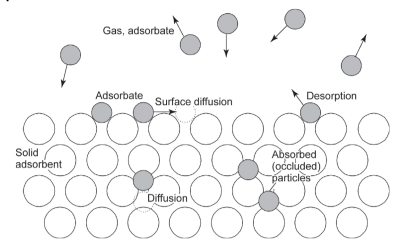


Figure 6.1 Concepts of sorption processes. White circles represent atoms in the solid. Gas atoms or molecules (gray circles) impinge on the surface and subsequently adsorb, diffuse (become absorbed), or desorb.

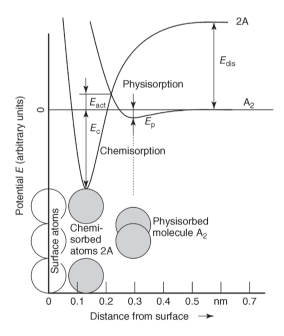


Figure 6.2 Potential of a molecule A₂ or two atoms 2 A at a distance from the surface of an adsorbent. The surface holds a physisorbed molecule A_2 with the energy $E_{ad} = E_p$ at a distance of approximately 0.3 nm (varying between 0.2 nm and 0.4 nm depending on the combination of adsorbate and adsorbent

species). Near the surface, the energy E_{dis} required for the dissociation $A_2 \rightarrow 2 \mbox{A}$ is reduced considerably to $E_{\rm act}$. If this activation energy E_{act} is overcome, the molecule dissociates and both atoms A are chemisorbed with the energy E_c to the surface at a distance of 0.15 nm.

Table 6.1 Adsorption energies E_{ad} equal to desorption energies E_{des} of adsorbate for selected gas species on some vacuum technologically relevant substances in eV (1 eV \triangleq 96.2 kJ mol⁻¹) for θ = 0.

Adsorbates	Adsorbents									
	Ti	Fe	Ni	Pd	Ta	w	Au	Pt	Al6063	Stainless steel
H ₂ /H		1.4	1.3	0.9-1.1	2.0	1.9	1.7			
O_2/O	10.8	5.5-6.2	5.5	2.4-2.9	9.5	8.4-9.6				
N ₂ /N		0.32/0.29								
CO	6.7-6.9	2.0	1.3-1.8	1.8	5.8	3.6		1.5-2.2		
CO_2	7.1	2.6	1.9		7.3	4.7				
H_2O							1.1		0.82 - 1.05	0.89-1.08

Due to measuring uncertainty and the specific condition of the surface (e.g., crystal plane, steps), values are limited to two decimals. Ranges are listed for values scattering any greater.

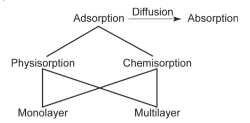


Figure 6.3 Taxonomy of sorption processes.

values for E_{des} . If the *adparticles* react chemically with *surface particles* and create a chemical, stoichiometric bond, values of adsorption energy rise to those of chemical reaction energies, which in fact are slightly above chemisorption values. This type of reaction requires surface particles to rearrange.

Also, adparticles can diffuse into the adsorbent: they dissolve in the solid. This process is named absorption or occlusion. The term sorption is used when information regarding the relative portions of both effects, adsorption and absorption, is unavailable or concealed (see also diagram in Figure 6.3).

The time constant with which occluded gas emerges to vacuum from a solid is considerably higher than the time constant with which surface adsorbed gas of the same type emerges. This process is referred to as outgassing, and is covered separately in Section 6.3. To simplify matters, the physically distinguished processes of desorption (from the surface) and outgassing (from the solid) are often subsumed under the term desorption. This is because a particle dissolved in the solid initially has to travel to the surface, quasi adsorb from within, to subsequently desorb. However, we will distinguish desorption and outgassing. The superordinate degassing for both concepts should be avoided. It usually refers to a controlled removal of dissolved gas from solids or liquids.

In the adsorption phase, adparticles can be close packed in a single layer (Figure 6.4). In this monomolecular layer, the number density per unit area

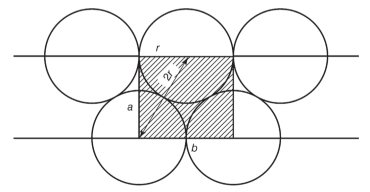


Figure 6.4 Close-packed atoms at a surface. The shaded area indicates the smallest possible rectangular cell with surface area A and the two sides a and b. b = 2r and according to the Pythagorean theorem, $a = \sqrt{3}r^2$, and thus, $A = 2\sqrt{3}r^2$.

 \tilde{n}_{mono} is

$$\tilde{n}_{\text{mono}} := \frac{N}{A} \,. \tag{6.1}$$

N denotes the number of adjacent particles on the required area A. If less adparticles adhere to the surface than in the monomolecular case, that is, $\tilde{n} < \tilde{n}_{\text{mono}}$, we define surface coverage

$$\theta = \frac{\tilde{n}}{\tilde{n}_{\text{mono}}}.$$
(6.2)

Example 6.1

According to Table A.9, the radius of a typical adsorbed molecule or atom $r = 1.6 \times 10^{-10}$ m. In closest packing, a particle (see Figure 6.4) requires the area $A = 2\sqrt{3} r^2$. Thus, the number density of the monolayer (monomolecular, monatomic) is

$$\tilde{n}_{\text{mono}} = 1/A = [2\sqrt{3}(1.6 \times 10^{-10})^2 \text{m}^2]^{-1} = 1.13 \times 10^{19} \text{ m}^{-2} \approx 10^{15} \text{ cm}^2.$$

For monatomic coverage, approximately 10¹⁵ particles lie on a geometric surface area of 1 cm². This important characteristic value of monomolecular coverage should be noted. However, it should be perceived that a technical surface can have a significantly larger area than a geometric surface because it features many steps and tips on the microscopic scale (from only one up to many interatomic distances). This issue is significant in ultrahigh-vacuum technology.

If an additional adlayer forms on top of the monolayer, the adsorption forces on this succeeding layer also include forces between identical adsorbate particles. For subsequent layers, these forces are nearly exclusively relevant. Thus, the desorption energy of particles in these layers approaches the evaporation heat Δh of the solid adsorbate for static adsorption layers, or of the liquid adsorbate for adsorption layers in motion (e.g., Δh (water, 0 °C) = 45.00 kJ mol⁻¹ \triangleq 0.47 eV/molecule, $\Delta h(\text{ice}, 0^{\circ}\text{C}) = 50.86 \text{ kJ} \text{ mol}^{-1} \triangleq 0.53 \text{ eV/molecule}.$

If the number density n, that is, pressure, of the adsorbate species is high in the gas phase, many superimposed adsorption layers can form. This process is called condensation (see also Section 3.5).

The phenomena of sorption and condensation are important throughout the entire pressure range in vacuum technology. Sorption pumps (Chapter 11), sputter ion pumps (Section 11.4), condensers (Chapter 8), and cryopumps (Section 12.4) utilize adsorption and condensation of gas molecules in order to pump. In non-evaporable getter (NEG) pumps (Section 11.3.2), diffusion of adsorbed particles is used to activate the pumps.

When a vented high-vacuum vessel is evacuated, adsorbed gas particles can delay or even practically prevent a certain pressure to be obtained. Gases that are adsorbed or absorbed (e.g., oxygen, nitrogen, water vapor) at higher pressure (e.g., when exposed to air or during ventilation) are released at different rates in a vacuum, depending on the value of desorption energy. Such desorbed molecules are potential sources of contaminants and impurities, an important issue in semiconductor industry. For example, the effect of oxygen atoms in metallization processes is devastating when they influence electric conductivity. The smaller the structures in integrated circuits (IC), the fewer harming particles are necessary to cause an IC to fail.

In particle accelerators, residual gas particles limit the lifetime and quality of the particle beam. In gravitational-wave detectors, they decrease the resolution with which changes of length are detected in Michelson interferometers.

In vacuum measurement, adsorption can influence results if a measuring device is sensitive to surface effects (e.g., ionization vacuum gauges).

Example 6.2

A monolayer of nitrogen is adsorbed at the interior surface A_S of a spherical vessel, radius r, volume $V = 1 \ell$.

According to Example 6.1, the area $A_S = 4\pi r^2 = 4\pi \left(\frac{3}{4\pi}v\right)^{2/3} = 485 \text{ cm}^2 \text{ carries}$

$$N_{\rm ad} = \tilde{n}_{\rm mono} A_{\rm S} = 5 \times 10^{17}$$
 molecules of nitrogen.

If they desorb entirely (e.g., due to rising temperature), the partial pressure of nitrogen in the vessel, according to Eq. (3.19), at ambient temperature is

$$p_{\rm N_2} = \frac{N_{\rm ad}}{V} kT = \frac{5 \times 10^{17} \cdot 1.4 \times 10^{-23} \text{ J} \cdot 300 \text{ K}}{10^{-3} \text{ m}^3 \text{ K}} \approx 2 \text{ N m}^{-2} = 2 \text{ Pa}.$$

The reactions of sorption and outgassing of gas and vapor at solid surfaces are very complex. Here, they are discussed only with the simplest models in order to provide basic understanding of vacuum technological processes.

6.2 Adsorption and Desorption Kinetics

6.2.1

Adsorption Rate

Equation (3.48) describes the particle flow density of gas or vapor (temperature T) onto the surface of an adsorbent (temperature T_{W}) where the particles adhere with the sticking probability s. The sticking probability of a particle depends on whether the particle reaches a free adsorption site at the surface,

		•
Gas species	<i>s</i> ₀	$ heta_{c}$
H ₂	0.08-0.30	0.26-0.50
CO	0.18-0.97	0.30-0.66
N_2	0.11-0.55	0.14-0.50
O_2	0.14-0.15	00.4-0.70

Table 6.2 Sticking probabilities s₀ of selected gas species on tungsten at 300 K. Ranges represent the values on different types of single- and polycrystalline tungsten surfaces as listed by a number of references. s is independent of θ in the range $\theta = 0$ to $\theta = \theta_c$ (see Eq. (6.3)).

From [2].

and, if it does, how high the probability of resting is at the site. This probability is denominated s_0 . It is temperature dependent, $s_0(T, T_W)$, particularly when the adsorption requires activation energy. However, it was shown that this is rarely the case, except for dissociating adsorption, or that the temperature dependence is only weak. Thus, to simplify, s_0 is assumed to be temperature independent, that is, constant:

$$s = s_0 f(\theta). \tag{6.3}$$

Table 6.2 lists a number of values for s_0 on tungsten at 300 K.

Langmuir formulated the simplest coverage dependence of s by assuming that particles are adsorbed only if they strike a vacant site, that is, if

$$f(\theta) = 1 - \theta,\tag{6.4}$$

as long as no dissociation occurs. If dissociation does take place, all n dissociation fragments require a vacant site and the probability drops with the *n*th power of $(1 - \theta)$. However, experiments show that s, in contrast to *Langmuir's* assumption, is practically independent of coverage in the range of $\theta = 0$ to approximately $\theta_c = 0.3-0.4$ (Table 6.2). In spite of this, *Langmuir's* coverage expression is often a very useful approximation. With Langmuir's assumption, the adsorption rate or adsorption flux density per unit area is

$$j_{\rm ad} = s_0 (1 - \theta) \frac{n c}{4}. \tag{6.5}$$

6.2.2

Desorption Rate

Adparticles adsorbed at the surface oscillate with a frequency in an order of magnitude $\nu_0 \approx 10^{13} \, \mathrm{s}^{-1}$, that is, with an oscillation period of $\tau_0 \approx 10^{-13} \, \mathrm{s}$. For desorption, they must carry kinetic energy $E_{kin} > E_{des}$. If the activation energy for adsorption is zero, then $E_{\text{des}} = E_{\text{ad}}$. According to *Boltzmann*, of \tilde{n} particles, only a portion $\Delta \tilde{n} = \tilde{n} \exp(-E_{\rm des}/(RT_{\rm W}))$ meets this requirement so that the

surface-desorption rate or desorption flux density,

$$j_{\text{des}} = \frac{d\tilde{n}}{dt} = -\nu_0 \,\Delta \tilde{n} = -\nu_0 \,\tilde{n} \exp\left(-\frac{E_{\text{des}}}{RT_{\text{W}}}\right),\tag{6.6}$$

incorporates the product of the particle number $\Delta \tilde{n}$ with sufficient energy $E_{\rm des}$ and the frequency ν_0 at which they oscillate to the outside, that is, away from the surface. Equation (6.6) is known as the Polanyi-Wigner equation. The frequency ν_0 is estimated by equating the energy $h\nu_0$ of the oscillator with the thermal energy kT:

$$\nu_0 = \frac{kT}{h} \tag{6.7}$$

At T = 296 K, it is $\nu_0 = 6.2 \times 10^{12}$ s⁻¹ $\approx 10^{13}$ s⁻¹.

Among the adparticles, some remain at the surface for a longer period of time before they desorb, whereas the residence time for others is low. According to Eq. (6.6), the average of all residence times, the mean residence time τ , is

$$\tau = \tau_0 \exp\left(\frac{E_{\text{des}}}{RT_{\text{W}}}\right). \tag{6.8}$$

Example 6.3

The molar desorption energy E_{des} of H_2O on stainless steel or aluminum is approximately 96 kJ mol⁻¹ (see also Table 6.1). Thus, at room temperature $(T_{\rm W}=300~{\rm K})$, the exponential factor in Eq. (6.8) $\varepsilon=\exp(\frac{96~000}{8.314\times300})=5.196\times10^{16}$, and at a temperature of 500 K, $\varepsilon = 1.070 \times 10^{10}$.

If the prefactor is assumed to be $\tau_0\approx 10^{-13}\,\text{s},$ the mean residence time of water molecules on a stainless steel or aluminum surface at $T_{\rm W}$ = 300 K amounts to $\tau \approx 10^{-13} \text{ s} \cdot 5.2 \times 10^{16} = 5200 \text{ s} \approx 1.4 \text{ h}$, and at $T_W = 500 \text{ K}$, $\tau \approx 10^{-13} \text{ s} \cdot 1.07 \times 10^{-13} \text{ s}$ $10^{10} = 1.07 \, \text{ms}.$

The desorption rate according to Eq. (6.6) is

$$j_{\text{des}} = -v_0 \tilde{n} \exp\left(-\frac{E_{\text{des}}}{RT_w}\right) = -10^{13} \text{ s}^{-1} \times 10^{15} \text{ cm}^2 \times \exp\left(-\frac{96000}{8.314 \times 300}\right)$$

= 1.9 × 10¹¹ s⁻¹ cm².

In more common units as pressure unit times volume unit this is

$$q_{pV} = q_N kT = 1.9 \times 10^{-11} \text{ s}^{-1} \text{cm}^{-2} \times 1.38 \times 10^{23} \text{ J/K}$$

 $\times 300 \text{ K} = 7.9 \times 10^{-10} \text{ Pa m}^3 \text{ s}^{-1} \text{ cm}^{-2}$
= $7.9 \times 10^{-9} \text{ mbar } \ell \text{ s}^{-1} \text{ cm}^{-2}$.

Hence, the typical desorption rate of a nonbaked inner surface of vacuum is about 10^{-8} mbar ℓ s⁻¹ cm⁻².

If, at the time t = 0, the surface is covered with \tilde{n}_0 particles, integration of Eq. (6.6) yields the particle number $\tilde{n}(t)$ at the time t:

$$\tilde{n}(t) = \tilde{n}_0 \exp\left(-\frac{t}{\tau}\right),\tag{6.9}$$

analogous to the law of radioactive decay. Thus, the time necessary for an adsorption layer to desorb into a vacuum with only a fraction $f = \tilde{n}(t)/\tilde{n}_0$ remaining is

$$t = \tau \ln\left(\frac{1}{f}\right),\tag{6.10}$$

for example, $t = 13.8 \tau$ for $f = 10^{-6}$.

Equations (6.9) and (6.10) are valid only if none of the desorbed particles readsorb at the surface. This condition is obtained, for example, by facing the desorbing surface to a pump with nearly infinite pumping speed, for example, a cryocondensation pump. In the practice of sealed vacuum chambers with a pump flange, however, readsorption is common. Hence, the reduction over time of adsorbed particles in a system slows down considerably as is discussed in the following section.

Example 6.4

A monolayer of H₂O molecules is adsorbed on a stainless steel surface. How long does it take the layer to desorb to 1% (f = 0.01), assuming that readsorption does not occur (a) at room temperature at $T_{\rm W} = 300\,\rm K$ and (b) at elevated temperature at T_W = 500 K? A monolayer of H₂O molecules accommodates approximately $\tilde{n}_0 = 10^{15}$ molecules/cm² (see Example 6.1). The desorption energy of H_2O on stainless steel shall be $E_{des} = 96 \text{ kJ mol}^{-1}$. According to Eqs. (6.10) and (6.8), and with $\tau_0 = 10^{-13}$ s,

a.
$$T = 300 \text{ K}$$

$$t = \tau \ln 100 = \tau_0 \exp\left(\frac{E_{\text{des}}}{RT_{\text{W}}}\right) \ln 100 = 10^{-13} \text{ s}$$

$$\times \exp\left(\frac{96 \text{ kJ mol}^{-1}}{8.314 \text{ J mol}^{-1} \text{ K}^{-1} \times 300 \text{ K}}\right) \ln 100$$

$$= 2.4 \times 10^4 \text{ s} = 6.6 \text{ h}.$$

b. T = 500 K. Analogous calculation yields

$$t = 4.9 \times 10^{-3} \text{ s} \approx 5 \text{ ms}.$$

As the example shows, a rise in temperature accelerates desorption remarkably. This is also valid for outgassing (Section 6.3). For this reason, the term bake-out was chosen.

Table 6.3 lists values of the exponential factor and residence time for τ_0 = 10^{-13} s and for selected desorption energies $E_{
m des}$ as well as surface (wall) temperatures T_W. However, it should be mentioned that experimental investigations also yielded prefactors τ_0 in the range $10^{-4} \text{ s} \ge \tau_0 \ge 10^{-15} \text{ s}$. Measured values of $E_{\rm des}$ vary between 0.08 kJ mol⁻¹ (\triangleq 0.8 meV, evaporation heat of liquid helium) and 1040 kJ mol^{-1} ($\triangleq 10.8 \text{ eV}$, adsorption of O_2 on Ti) (see also Table 6.1).

6.2.3

Hobson Model of a Pump-down Curve

Calculating a p(t) curve of a vacuum chamber, that is, the pressure change over time in the chamber, requires a description of the physical processes at the interior surface of the chamber and in its volume.

We will consider an isothermal chamber with the interior surface A_S and volume V, flanged to a vacuum pump. The pumping speed of the pump is expressed in terms of an effective pumping area A_p . All particles passing through this area are pumped out. Therefore, the number of particles that are removed from the volume by the pump is $n \, \overline{c} 4 A_p$, where n denotes the particle number density in the volume and \bar{c} is their mean thermal velocity. Per unit time, $\tilde{n}\tau A_{\rm S}$ particles desorb from the surface. $n \bar{c} 4 s A_S$ particles become readsorbed. The following equation may be formulated:

$$V\frac{\mathrm{d}n}{\mathrm{d}t} = \frac{\tilde{n}}{\tau}A_{\mathrm{S}} - \frac{n\,\bar{c}}{4}A_{\mathrm{p}} - \frac{n\,\bar{c}}{4}s\,A_{\mathrm{S}}.\tag{6.11}$$

The change in the number of particles in the volume per unit time V dn/dt is equal to the number of particles desorbing from the surface to the vacuum minus the number of particles that are pumped out or that readsorb.

At the surface,

$$A_{\rm S} \frac{\mathrm{d}\tilde{n}}{\mathrm{d}t} = \frac{n\,\bar{c}}{4} s\,A_{\rm S} - \frac{\tilde{n}}{\tau} A_{\rm S}.\tag{6.12}$$

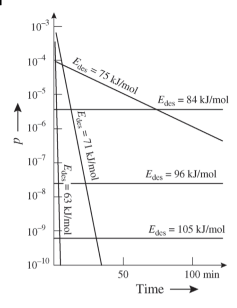
The change in particle number at the surface is equal to the number of particles that readsorb per unit time minus the number of particles that desorb. The previous equation shows that the effective net desorption rate $-A_{\rm S}\left[rac{d ilde{n}}{dt}
ight]$ generally depends on *n* and therefore on the pressure.

Combining Eqs. (6.11) and (6.12) yields a differential equation:

$$\frac{\mathrm{d}^2 n}{\mathrm{d}t^2} + \left(\frac{\overline{c}}{4V}(sA_\mathrm{S} + A_\mathrm{p}) + \frac{1}{\tau}\right)\frac{\mathrm{d}n}{\mathrm{d}t} + \frac{\overline{c}}{4V}\frac{A_\mathrm{p}}{\tau}n = 0. \tag{6.13}$$

Table 6.3 Exponential factor $\epsilon = \exp(E_{\text{des}}/(RT_{\text{w}}))$ and mean residence times τ according to Eq. (6.7) with $\tau_0 = 10^{-13}$ s versus desorption energy E_{des} and temperature T_{w} of the solid (1 year is equal to 3.15×10^7 s).

E_{des}	E _{des}		$T_{\rm w}=77{\rm K}\;(-196^{\circ}{\rm C})$		298 K (25 °C)		523 K (250 °C)		1273 K (1000 °C)		2273 K (2000 °C)	
eV	kJ mol ⁻¹	kcal mol ⁻¹	ε	τ/s	ε	τ/s	ε	τ/s	ε	τ/s	ε	τ/s
0.004	0.42	0.1	1.93	2×10^{-13}	1.18	1.2×10^{-13}	1.10	1.1×10^{-13}	1.04	1×10^{-13}	1.02	1×10^{-13}
0.0436	4.19	1	698	7×10^{-11}	5.43	5.4×10^{-13}	2.62	2.6×10^{-13}	1.48	1.5×10^{-13}	1.25	1.3×10^{-13}
0.436	41.9	10	2.7×10^{28}	3×10^{15}	2.2×10^{7}	2.2×10^{-6}	1.5×10^4	1.5×10^{-9}	52.5	5.3×10^{-12}	9.19	9×10^{-13}
2.18	210	50			6.7×10^{36}	6.7×10^{23}	9.4×10^{20}	9.4×10^{7}	4.2×10^8	4.2×10^{-5}	6.7×10^4	6.7×10^{-9}
4.36	419	100					7.1×10^{41}	7×10^{28}	1.6×10^{17}	1.6×10^4	4.3×10^{9}	4×10^{-4}
6.54	629	150							6.6×10^{25}	6.6×10^{12}	2.9×10^{14}	29
13.1	1257	300									8×10^{29}	8×10^{15}



of a vacuum system covered with adsorbate layers showing the indicated desorption energies E_{des} . At t = 0, the thickness of the layers

Figure 6.5 Pressure versus time for evacuation is one monolayer. $V = 1 \ell$, $A_S = 100 \text{ cm}^2$, $S = 100 \text{ cm}^2$ $1 \ell s^{-1}$ ($A_p = 0.09 \text{ cm}^2 \text{ for air}$), T = 295 K, andfirst-order desorption, that is, without any dissociation.

Using certain simplifications, *Hobson* [3] solved these equations for the important range of desorption energies from 63 to 105 kJ mol⁻¹. Figure 6.5 shows the pressure plotted against time in a vacuum chamber of $V = 1 \ell$, $A_S =$ 100 cm^2 , $S = 1 \ell \text{ s}^{-1}$ ($A_p = 0.09 \text{ cm}^2$ for air), T = 295 K, for first-order desorption (i.e., no dissociation). *Hobson* assumed pressure-independent sticking probability; thus, applications of his model are limited.

Yet, an important conclusion can be drawn from this calculation: molecules with low desorption $\leq 71 \text{ kJ mol}^{-1}$ leave the chamber quickly, within approximately 30 minutes, and subsequently, are irrelevant. Very high desorption $\geq 105 \,\mathrm{kJ} \,\mathrm{mol}^{-1}$ impede desorption so effectively that the molecules can hardly be pumped out at room temperature but, however, they hardly cause the pressure to rise. Only in the transition region of 75–105 kJ mol⁻¹, molecules contribute considerably to pressure, in the long term.

Unfortunately for vacuum technology, the desorption energy of water lies precisely in this range. Values measured for H2O on stainless steel and aluminum range between 80 kJ mol⁻¹ and 104 kJ mol⁻¹ (Table 6.1). This means that H₂O is the main component in evacuating a previously vented vacuum chamber, and this can prolong for days, even weeks, if the temperature is not adjusted (for baking).

In a pump-down process, the net desorption rate (desorption minus readsorption) drops, following an exponential law with time:

$$j_{\rm des} = K t^{-n}. (6.14)$$

Thus, if pumping speed is constant, the pressure in the system follows a power law as well:

$$p = k t^{-n}. (6.15)$$

Experiments yielded values for n from 0.7 to 2.0, mostly in the range of 0.9–1.3 [4]. This span indicates that not only one single type of H₂O adsorption site, that is, a corresponding desorption energy value, exists. This is understandable because a technical surface features a very complex structure and it is unlikely that only a single desorption energy level is realized. Indeed, Redhead explained this nearly 1/t-behavior with a distribution of desorption energy levels [5].

It should be noted, however, that there is an additional model that explains the 1/t-behavior: Dayton [6] assumed that water molecules gather in small pores and capillaries in the oxide layer on the metal surface. During pumpdown, these pores would gradually empty due to diffusion of H₂O molecules. Later, in Section 6.3, we will see that when degassing is diffusion controlled, the rate of gas evolution, and therefore pressure, drops with $t^{-1/2}$ although an explanation for 1/t-behavior is needed. Dayton suggested that diameters and lengths of the pores and capillaries in the oxide vary and lead to differing diffusion coefficients in calculations. He introduced a time constant τ , describing the type of pore. If the pumping time $> \tau$, gas evolution follows an exponential law. A broader distribution of τ causes the cumulative curve to show 1/t-behavior (Figure 6.6).

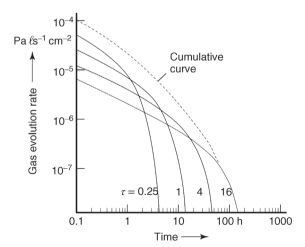


Figure 6.6 Specific water evolution rate of a vacuum chamber wall according to *Dayton* [6] based on four selected effective time constants τ (0.25 h, 1 h, 4 h, 16 h) as in Eq. (6.7). In this case, these constants do not represent

four different desorption energies (which would yield the same results) but combinations of a single desorption energy and diffusion coefficients of four capillaries with different dimensions.

6.2.4

Monolayer Adsorption Isotherms

The so-called adsorption isotherm indicates the relationship between pressure and surface coverage θ . Application of this isotherm is reasonable if adsorption and desorption rates are in equilibrium, that is, neither rapid changes in pressure nor in temperature occur. Kanazawa [7] showed that, for relative pressure changes in time.

$$\left| \frac{1}{p} \cdot \frac{\mathrm{d}p}{\mathrm{d}t} \right| \ll s \frac{A}{V} \cdot \frac{\overline{c}}{4}. \tag{6.15}$$

Here, V and A denote the volume and the interior surface of the system, respectively. If the system fulfills the condition in this inequation, changes are quasistationary and the surface coverage can follow pressure changes according to the adsorption isotherm.

Example 6.5

What is the maximum value for dp/dt in a vacuum chamber of $V = 16.7 \, \text{l}$, $A = 4750 \text{ cm}^2$, and T = 300 K, assuming that the pressure $p = 10^{-2} \text{ Pa}$ is caused only by water vapor (pump-down process) and s = 0.1? The condition is

$$\frac{dp}{dt} \ll p \, s \frac{A}{V} \cdot \frac{\overline{c}}{4} = 10^{-2} \, \text{Pa} \cdot 0.1 \frac{4750}{16700} \, \text{cm}^{-1} \frac{59400}{4} \, \text{cm s}^{-1} = 4.2 \, \text{Pa s}^{-1}.$$

In pump-down processes under high- or fine-vacuum conditions, the rate at which the pressure changes is far below this value, that is, the process is quasistationary.

When using Langmuir's assumption for the coverage dependence of s, the isotherms are calculated by equating the equations for adsorption rate (6.5) and desorption rate (6.6). Using the relationship between the pressure p and number density n of the adsorbate in Eq. (3.19), this equalization yields the Langmuir adsorption isotherm:

$$\theta = \frac{\tilde{n}}{\tilde{n}_{\text{mono}}} = \frac{a p}{1 + a p} \tag{6.16}$$

with

$$a = \frac{s_0 \, \tau_0 \exp\left(\frac{E_{\text{des}}}{RT_{\text{W}}}\right)}{\tilde{n}_{\text{mono}} \sqrt{2\pi \, M_{\text{molar}} \, RT}},\tag{6.17}$$

in which τ_0 as well as E_{des} can depend on coverage and T_{W} .

Figure 6.7 shows the characteristic plot of this isotherm. If p is low $(p \ll a^{-1})$, then $\theta \propto p$. This proportionality is referred to as *Henry's* law or *Henry adsorption*

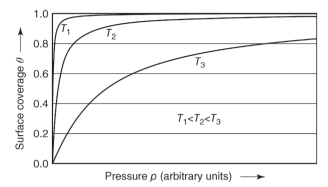


Figure 6.7 Langmuir adsorption isotherm $\theta(p)$ according to Eq. (6.16) for three different temperatures.

isotherm. If $p \to \infty$, then $\theta \to 1$, but θ cannot rise above 1. This means that under *Langmuir's* assumption, $s \propto (1 - \theta)$, at most, a monatomic layer can develop.

Additionally, Langmuir supposed that the adsorption energy (= E_{des}) was independent of θ . In contrast, Freundlich assumed an exponential relation between adsorption energy and θ :

$$E_{\rm des} = -E' \ln \theta. \tag{6.18}$$

This assumption uses the concept of a heterogeneous surface with exponential energy dependence among the adsorption sites, leading to the Freundlich adsorption isotherm in the form

$$\theta = k \, p^{\beta},\tag{6.19}$$

where k and β are constants. A disadvantage of the assumption underlying the *Freundlich* isotherm is that $E_{\text{des}} = 0$ for $\theta = 1$, and $E_{\text{des}} \to \infty$ for $\theta \to 0$. Thus, for both these boundary cases, it does not realistically model the physical behavior. Therefore, the *Freundlich isotherm* is useful only for medium-high values of coverage.

For the Temkin isotherm, a linear change of adsorption energy with surface coverage θ is assumed:

$$E_{\text{des}} = E_{\text{des},\theta=0}(1 - \alpha \theta). \tag{6.20}$$

 $E_{\mathrm{des},\theta=0}$ is the adsorption/desorption energy for $\theta=0$. α is a constant. For $\theta=1$, $E_{\rm des}$ drops to a value $E_{{\rm des},\theta=1}>0$. The drop in $E_{\rm des}$ with increasing coverage can be explained with repellant forces among adsorbed molecules. The calculation of the adsorption isotherm yields [8]

$$\theta = \frac{RT}{E_{\text{des},\theta=0} - E_{\text{des},\theta=1}} \ln \left(\frac{1 + \frac{p}{p^*} \exp\left(\frac{E_{\text{des},\theta=0}}{RT}\right)}{1 + \frac{p}{p^*} \exp\left(\frac{E_{\text{des},\theta=1}}{RT}\right)} \right), \tag{6.21}$$

where
$$p^* = \frac{\tilde{n}_{\mathrm{mono}}}{s_0 \, \tau_0 \sqrt{2\pi \, M \, RT}}$$
.

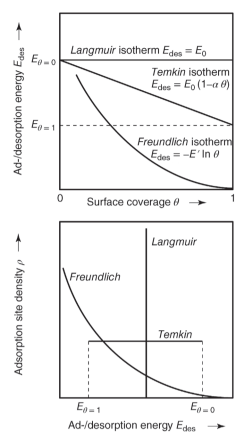


Figure 6.8 Assumptions made for selected adsorption isotherms regarding the dependence of desorption energy on coverage and adsorption site distribution on desorption energy.

For the three simple adsorption isotherms introduced here, Figure 6.8 shows the energy dependence of ad-/desorption energy $E_{\rm des}$ from θ , and the corresponding density ρ of adsorption sites versus E_{des} . All of these adsorption isotherms are applicable only if $\theta \leq 1$.

6.2.5 Multilayer Adsorption and Brunauer-Emmett-Teller (BET) Isotherm

It was observed in many cases that surface coverage increases to $\theta > 1$ when the pressure rises. This indicates that additional physisorbed layers grow on top of the chemi- or physisorbed monolayer. The simplest model for this phenomenon assumes that each layer is completed to full surface coverage, that is, layer

(n+1) does not start to grow on an incomplete partial layer n. Brunauer, Emmett, and Teller formulated these conceptions. As experiments show, they are applicable to many cases as long as the gas pressure p of the adsorbate is well below the vapor pressure p_s of the condensed adsorbate.

The parameters in Section 6.2.4 apply to the first layer. For all succeeding layers, desorption energy $E_{\rm des}$ in Eq. (6.7) is replaced by the heat of evaporation Δh and a different prefactor τ' . Summing up yields the BET isotherm

$$\theta_{\text{BET}} = \frac{\frac{p}{p_{\text{S}}} C_{\text{BET}}}{\left(1 - \frac{p}{p_{\text{S}}}\right) \left(1 + (C_{\text{BET}} - 1)\frac{p}{p_{\text{S}}}\right)},\tag{6.22}$$

where

$$C_{\text{BET}} = \frac{\tau_0}{\tau'} \exp\left(\frac{E_{\text{des}} - \Delta h}{RT_{\text{W}}}\right) \tag{6.23}$$

is the ratio of the residence times on the first adlayer and the condensing layer.

Figure 6.9 gives a schematic representation of Eq. (6.22). If $p \rightarrow p_s$, then $\theta \to \infty$, and for $p \ll p_{\rm S}, \theta_{\rm BET} \propto p$ (Henry isotherm). A type of Langmuir saturation characterizes the transition range. However, as described previously, if $p \rightarrow p_S$, the BET isotherm differs from data found in experiments. In practice, the amount of gas condensing for $p \rightarrow p_S$ is limited. Instead, continuously pumped systems develop an equilibrium pressure p_{eq} [9] and a corresponding $\theta_{\rm eq}$ because the adsorbate is pumped away ($p_{\rm S}$ is the saturation pressure in a sealed, unpumped system). The amount of gas that can condense in a sealed system is limited as well because the enclosed amount of gas is finite. The BET isotherm features a θ_{eq} -value of several 1000, whereas θ_{eq} for other adsorption isotherms such as the Frankel-Halsey-Hill (FHH) isotherm [10] or the McMillan-Teller (MT) isotherm [11] is in the range of 10–100. Between $0.5 < \theta < 2$, Eq. (6.22) also serves to calculate the true surface of adsorbents.

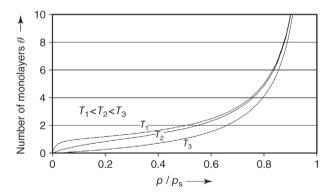


Figure 6.9 BET adsorption isotherms $\theta(p/p_S)$ for three selected temperature levels. p_S is the saturation vapor pressure of the condensed adsorbate.

6.2.6

Monolayer Time

The term *monolayer time* (monolayer formation time) was introduced in order to estimate the period of time available for surface analysis of a clean, pure surface at given pressure. Monolayer time is the period in which a monolayer develops, assuming that all atoms and molecules striking the analyzed surface from the gas phase adhere to the surface permanently (s = 1). The equation for monolayer time,

$$j_{\rm ad} t_{\rm mono} = \tilde{n}_{\rm mono}, \tag{6.24}$$

can be rewritten using Eq. (6.5), the equation of state, Eq. (3.19), p = n k T, and Eq. (3.43) for the mean velocity of gas molecules to yield

$$t_{\text{mono}} = \frac{\tilde{n}_{\text{mono}}}{p N_{\text{A}}} \sqrt{2\pi M_{\text{molar}} RT}, \tag{6.25}$$

or as an abbreviated numerical value equation,

$$t_{\rm mono}/{\rm s} = 3.18 \times 10^{-25} \, {\rm s} \, \frac{\tilde{n}_{\rm mono}/{\rm m}^2}{p/{\rm Pa}} \, \sqrt{M_{\rm r} \, T/{\rm K}}.$$

If the residual gas is air ($M_{\rm r} \approx 29$) at $T \approx 300$ K, a useful approximation formula is obtained ($\tilde{n}_{\text{mono}} \approx 10^{15} \text{ cm}^{-2}$):

$$t_{\text{mono}} = \frac{3.6 \times 10^{-4} \,\text{Pa}}{p} \,\text{s}$$
 for air. (6.26)

Table 6.4 lists a few values. In investigations of surface properties, for example, measurements of electron work functions, the monolayer time must fulfill the condition $t_{\rm ex} \ll t_{\rm mono}$ if nearly constant surface coverage θ for the duration of the experiment $t_{\rm ex}$ is desired. Thus, the pressure must be sufficiently low (Table 6.4), that is, ultrahigh-vacuum technology is usually indispensable.

Table 6.4 Monolayer time t_{mono} versus gas pressure p for air, water, and hydrogen (at 300 K, $\tilde{n}_{\text{mono}} \approx 10^{15} \text{ cm}^{-2}$).

p	Pa	100	0.1	10 ⁻⁵	10 ⁻⁷	10 ⁻⁹
	mbar	1	10 ⁻³	10 ⁻⁷	10 ⁻⁹	10 ⁻¹¹
t _{mono} (air)		$3.6 \times 10^{-6} \text{ s}$	$3.6 \times 10^{-3} \text{ s}$	36 s.3	01 h	100 h
$t_{\rm mono}$ (H ₂ O)		$2.8 \times 10^{-6} \text{ s}$	$2.8 \times 10^{-3} \text{ s}$	$28 \mathrm{s.3}$	47 min	078 h
$t_{\mathrm{mono}} \left(\mathrm{H}_{2} \right)$		$9.3 \times 10^{-7} \text{ s}$	$9.3 \times 10^{-4} \text{ s}$	09.3 s	16 min	026 h

Absorption, Diffusion, and Outgassing

Adsorbed particles can migrate into a solid by skipping to interstitial sites or lattice defects or by moving along grain boundaries of crystallites (practically all technical substances are polycrystalline). They are absorbed.

As every jump from one site to the next requires an activation energy E_{dif} , the process is temperature dependent. The totality of events is called diffusion. According to Eq. (3.102), it is caused by a concentration gradient and features a particle flow rate (Fick's first law):

$$j_{\rm dif} = -D \frac{\mathrm{d}n_{\rm dis}}{\mathrm{d}x},\tag{6.27}$$

where $n_{\rm dis}$ denotes the density of dissolved (occluded, absorbed) particles and the coefficient of diffusion

$$D = D_0 \exp\left(-\frac{E_{\text{dif}}}{RT}\right). \tag{6.28}$$

The amount of absorbed gas can be far greater than the amount of adsorbed gas because the number of sites available to dissolved, occluded particles in the bulk of a solid is large compared with eligible sites at the surface.

Diffusion into a solid is technically relevant as, for example, in the case of H₂ and O2: Ta and Nb can serve as hydrogen reservoirs. However, too high concentrations cause embrittlement in these metals. In steel production, considerable amounts of hydrogen from the atmosphere dissolve in the steel. In stainless-steel vacuum systems, this hydrogen diffuses out and is the main source of residual gas in baked-out, ultrahigh-vacuum systems.

To exemplify, we will consider outgassing of a *thin* piece of sheet metal, that is, a sheet metal with thickness 2d lying in the direction of the x-coordinate, low compared with length l and width b (y- and z-coordinates, area A = l b) (Figure 6.10). This one-dimensional diffusion problem is fairly simple to treat mathematically. The change in the particle density $n_{\rm dis}$ of absorbed particles over time is (Fick's second law)

$$\frac{\partial n_{\rm dis}}{\partial t} = D \frac{\partial^2 n_{\rm dis}}{\partial r^2}.$$
 (6.29)

From fabrication, the number density $n_{\text{dis},0}$ shall be evenly distributed throughout the plate at the begin of diffusion, that is, at time t = 0. Diffusion starts at t = 0 due to an increase in temperature (alternatively, prior to t = 0, the gas is in an equilibrium condition with the surrounding, and subsequently, the outer space is evacuated). A diffusion current j_{dif} A, Eq. (6.27), emanates symmetrically to both sides of the sheet metal and a density distribution $n_{dis}(x, t)$, falling with time, develops. The boundary condition

$$n_{\rm dis}(\pm d) = 0 \quad \text{for} \quad t > 0, \tag{6.30}$$

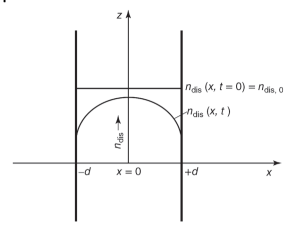


Figure 6.10 Gas evolution on a thin plate of thickness 2d. At t = 0, the adsorbate (gas) is homogeneously dissolved in the solid (particle symmetric sinusoidal half-wave profile density $n_{dis,0}$) and in equilibrium with the

surrounding. Subsequently, the volume on both sides of the plate is evacuated and a develops.

indicates that desorption is far more rapid than diffusion and the surface is essentially free of the molecules diffusing to the surface. Assuming this, and with the boundary condition

$$n_{\rm dis}(x) = n_{\rm dis,0}$$
 for $t = 0$, (6.31)

the result to the differential equation, Eq. (6.29), at the surfaces reads [12]

$$j_{\text{dif}}(x = \pm d) = \frac{2D}{d} n_{\text{dis},0} \sum_{i=0}^{\infty} \exp\left(-\frac{(2i+1)^2 \pi^2 D t}{4d^2}\right). \tag{6.32}$$

The value

$$t_{\rm out} = \frac{4d^2}{\pi^2 D} \tag{6.33}$$

in the argument of the exponential function is a time constant characterizing outgassing.

Example 6.6

Calculate t_{out} for hydrogen and a stainless steel plate with thickness 2d = 2 mmat 296 and 550 K (typical baking temperature). The diffusion coefficient is given in Eq. (6.28). D_0 for stainless steel is 0.012 cm²s⁻¹, and $E_{dif} = 56 \text{ kJ mol}^{-1}$ according to Table 6.5. At room temperature,

$$D = 0.012 \text{ cm}^2 \text{ s}^{-1} \exp\left(-\frac{56 \text{ kJ mol}^{-1}}{8.314 \text{ kJ kmol}^{-1} \text{ K}^{-1} 296 \text{ K}}\right)$$
$$= 1.57 \times 10^{-12} \text{ cm}^2 \text{ s}^{-1},$$

and thus.

$$t_{\text{out}} = \frac{4(0.1 \text{ cm})^2}{\pi^2 \cdot 1.57 \times 10^{-12} \text{ cm}^2 \text{s}^{-1}} = 2.58 \times 10^9 \text{ s} = 81.9\text{a}.$$

Analogous calculation for 550 K yields $D = 5.77 \times 10^{-8} \text{ cm}^2 \text{ s}^{-1}$ and

$$t_{\text{out}} = 7.024 \times 10^4 \text{ s} \approx 19.5 \text{ h}$$

The time constant t_{out} is temperature dependent because D is temperature dependent. For $t \gg 0.5 \, t_{\rm out}$, the summation terms with $i \ge 1$ amount to less than 2% so that, in this case,

$$j_{\text{dif}}(x = \pm d) = \frac{2D}{d} n_{\text{dis},0} \exp\left(-\frac{\pi^2 D}{4d^2} t\right) = j_0 \exp\left(-\frac{t}{t_{\text{out}}}\right).$$
 (6.34)

For $t \ll 0.5 t_{\text{out}}$, instead of Eq. (6.32), the following equation can be used for approximation:

$$j_{\text{dif}} = \frac{2D}{d} n_{\text{dis, 0}} \sqrt{\frac{\pi}{16} \cdot \frac{t_{\text{out}}}{t}} = j_0 \sqrt{\frac{\pi}{16} \cdot \frac{t_{\text{out}}}{t}}.$$
 (6.35)

Values in the two latter equations are determined by the constant

$$j_0 = \frac{2D}{d} n_{\text{dis}, 0}. \tag{6.36}$$

Table 6.5 Diffusion constant D, outgassing time constant t_{out} , and outgassing time t for selected outgassing levels according to the pure diffusion model for hydrogen in a stainless steel sheet of thickness 2d = 2 mm, $D_0 = 0.012 \text{ cm}^2 \text{ s}^{-1}$, $E_{\text{dif}} = 56 \text{ kJ mol}^{-1}$, Eq. (6.28).

<i>T</i> (K)	$D(T) \text{ (cm}^2 \text{ s}^{-1}\text{)}$	t _{out} (s)	f = 0.1			$f = 10^{-6}$
			t (s ⁻¹)	t	t (s ⁻¹)	t
0296	1.6×10^{-12}	02.6×10^9	005.4×10^9	170 a	3.5×10^{10}	1100 a
0500	1.7×10^{-8}	02.4×10^5	005.0×10^5	006 d	3.3×10^6	0038 d
0823	3.4×10^{-6}	01.2×10^3	002.5×10^{3}	042 min	1.6×10^4	0004.4 h
1223	4.9×10^{-5}	83	174	003 min	1.1×10^3	0019 min
Eq.	(6.28)	(6.33)	(6.40)	(6.40)	(6.40)	(6.40)

Example 6.7

Calculate the gas evolution of hydrogen in Pa ℓ s⁻¹ for a stainless steel plate of thickness 2d = 2 mm at 296 K after $t = 10^7 \text{ s}$ (116 days). Following Example 6.6, $t \ll t_{\rm out}$, and thus, Eq. (6.35) is applicable. Typically, $n_{\rm dis,\,0}$ amounts to 40 Pa ℓ cm⁻³. From Eq. (6.36), it follows that

$$j_0(296 \text{ K}) = \frac{2 \times 1.57 \times 10^{-12} \text{ cm}^2 \text{s}^{-1}}{0.1 \text{ cm}} 40 \text{ Pa } \ell \text{ cm}^{-3}$$
$$= 1.26 \times 10^{-9} \text{ Pa } \ell \text{ s}^{-1} \text{ cm}^{-2}.$$

Thus, gas evolution

$$j_{\text{dif}} = 1.26 \times 10^{-9} \text{ Pa } \ell \text{ s}^{-1} \text{ cm}^{-2} \sqrt{\frac{\pi}{16} \cdot \frac{2.58 \times 10^9}{10^7}}$$

= 8.9 × 10⁻⁹ Pa ℓ s⁻¹ cm⁻².

Example 6.8

Calculate the pressure in a vacuum chamber made of stainless steel as in Example 6.7. The chamber has an interior surface of 10 000 cm² and is pumped by a pump with $100 \, \ell \, \mathrm{s}^{-1}$. The total outgassing rate is $q_{pV} = j_{\mathrm{dif}} \, A = 8.9 \times 10^{-1}$ $10^{-5} \text{ Pa } \ell \text{ s}^{-1}$. The equilibrium pressure

$$p = \frac{q_{pV}}{S} = \frac{8.9 \times 10^{-5} \text{ Pa } \ell \text{ s}^{-1}}{100 \ell \text{ s}^{-1}} = 8.9 \times 10^{-7} \text{ Pa}.$$

Thus, a hydrogen partial pressure of about 10^{-6} Pa $(10^{-8}$ mbar) can be expected for a stainless steel vacuum system of thickness 2 mm that has never been baked out.

As a result, under the considered initial and boundary conditions, the diffusion current first drops following a complicated, Eq. (6.32), and later, according to a simple, Eq. (6.34), exponential distribution law.

Vacuum chambers are often baked, that is, the material is exposed to high temperature temporarily, in order to desorb molecules that were previously adsorbed. The diffusion coefficient increases rapidly with rising temperature, and thus, the increased temperature determines the concentration profile of the dissolved substance, which then is preserved at low temperature. If t_1 denotes the baking period, then, for $t_1 \gg t_{\text{out}}$ (which is usually the case, see Example 6.6),

$$j_{\text{dif}} = \frac{2D_{\text{r}}}{d} n_{\text{dis, 0}} \exp\left(-\frac{\pi^2 D_1 t_1}{4d^2}\right)$$
 (6.37)

if $D_{\rm r}$ and $D_{\rm 1}$ denote the diffusion coefficients at room temperature and baking temperature, respectively.

Example 6.9

Calculate $j_{\rm dif}$ at 296 K for a stainless steel plate of thickness $2d=2\,{\rm mm}$, which was previously baked out at 550 K for $t_1 = 24 \,\mathrm{h}$. Considering Example 6.6, $t_1 \gg t_{\rm out}$, and thus, Eq. (6.37) is applicable. The prefactor in Eq. (6.37) corresponds to j_0 in Example 6.7. Therefore (see also Example 6.6 for D_1),

$$\begin{split} j_{\mathrm{dif}} &= 1.26 \times 10^{-9} \, \mathrm{Pa} \, \ell \, \mathrm{s}^{-1} \, \mathrm{cm}^{-2} \, \mathrm{exp} \bigg(-\frac{\pi^2 \, 5.76 \times 10^{-8} \, \mathrm{cm}^2 \, \mathrm{s}^{-1} \, 8.64 \times 10^4 \, \mathrm{s}}{4 (0.1 \, \mathrm{cm})^2} \bigg) \\ &= 3.69 \times 10^{-10} \, \, \, \mathrm{Pa} \, \ell \, \mathrm{s}^{-1} \, \mathrm{cm}^{-2}. \end{split}$$

Comparison of this value shows that this type of baking already reduces gas evolution by a factor of 25.

Equation (6.32) allows calculating the relative number of dissolved particles in the sheet metal with respect to the beginning of outgassing. A piece of sheet metal of area A accommodates $N_0 = 2dA n_{\text{dis}, 0}$ particles at the time outgassing begins. After a time t,

$$\Delta N = 2 \int_{0}^{t} A j_{\text{dif}} \, \mathrm{d}t \tag{6.38}$$

particles have diffused out to both sides. Integration of Eq. (6.38) from t to ∞ yields the residual gas content N(t) after an outgassing period t, referred to as the outgassing ratio (or exhalation ratio)

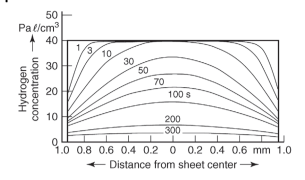
$$f = \frac{N(t)}{N_0} = \frac{8}{\pi^2} \left(e^{-\zeta} + \frac{e^{-9\zeta}}{9} + \frac{e^{-25\zeta}}{25} + \cdots \right), \tag{6.39}$$

where $\zeta = t/t_a$; for $\zeta = 1$, that is, $t = t_a$, we find f = 0.3; here, higher terms of the progression already are negligible. Thus, for f < 0.3, the simpler relation

$$t/t_{\rm a} = \ln\left(\frac{8}{\pi^2 f}\right) \tag{6.40}$$

can be used to calculate outgassing time t. Table 6.5 lists values for the case of hydrogen in a stainless steel sheet metal of thickness 2 mm.

It must be pointed out that the previous equations apply only to the given idealized conditions. In particular, the assumption that diffusion is far quicker than desorption from the surface can be wrong for outgassing of hydrogen. Hydrogen diffuses as individual atoms within a solid but desorbs only as molecules. A hydrogen atom reaching the surface requires a second hydrogen atom to form a molecule and desorb. Thus, the assumption $n_{\text{dis}}(\pm d) = 0$, Eq. (6.30), must be wrong because then the probability that two H atoms meet would be



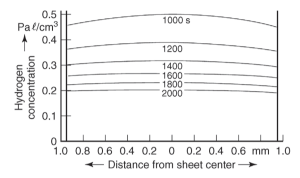


Figure 6.11 Calculated hydrogen concentration (starting value for t=0 is 40 Pa ℓ cm⁻³) in a plate of 1.9 mm thickness versus time according to *Moore's* recombination

model [13]. The volume on both sides of the plate is evacuated at 950 °C. Note that the lower ordinate is stretched by a factor of 100. (Data from [13].)

zero. In fact, the number of atoms at the surface will accumulate until the recombination rate of hydrogen atoms equals the diffusion rate to the surface. *Moore* [13] numerically calculated this process and found the following.

Figure 6.11 shows a concentration distribution of H atoms in a cross section of an infinitely extended, 1.9-mm-thick stainless steel plate during degassing at 950 °C in a vacuum furnace (vacuum firing). The numbers denote the period of baking in s. A considerable deviation from the purely diffusion-controlled concentration distribution develops due to the concentration at the surface, which is different from zero $(n_{\rm dis}(\pm d) \neq 0)$. With time, the plot of $n_{\rm dis}$ approaches a sine half-wave plus a constant instead of a simple sine half-wave as for the diffusion-controlled model. The sine component drops with time until the concentration is nearly homogeneous after 2000s. After this period, the recombination rate limits outgassing while diffusion is rapid enough to replace two recombined hydrogen atoms spontaneously. The rate of recombination $j_{\rm rec}$ is proportional to the square of the density of surface atoms $n_{\rm S}$. The symbol for the proportionality constant is $K_{\rm rec}$.

$$j_{\rm rec} = K_{\rm rec} n_{\rm S}^2. \tag{6.41}$$

Moore adopted his model to experimental data and found [13] K_{rec} = 3×10^{-22} cm² s⁻¹ for 950 °C, and $K_{\text{rec}} = 1.14 \times 10^{-27}$ cm² s⁻¹ for 25 °C.

After high-temperature degassing under vacuum (vacuum firing) of 1-2 h, the calculated outgassing at room temperature in the recombination-controlled model of outgassing is many orders of magnitude higher than in the diffusioncontrolled model. Therefore, much longer degassing periods are required to produce equivalent outgassing rates.

If outgassing of hydrogen is controlled by the rate of recombination, one would expect surface roughness, which influences surface diffusion, to also influence the effective hydrogen desorption energy. Indeed, Chun [14] observed this phenomenon in experimental investigations.

It is still openly discussed in the literature, if the outgassing rate of hydrogen from stainless steel is controlled by diffusion or recombination. Investigations of surfaces of stainless steel after vacuum firing showed [15,16] that grain boundaries are more pronounced and deepened compared with the state before vacuum firing while the surfaces of the grains are smoother (Figure 6.12). With deeper grain boundaries there are more active sites for adsorption and recombination available. On the smoother surface, terraces are formed, which, at their steps, also contain many adsorption sites. Surface diffusion, on the other hand, is greatly helped by smooth terraces, so that the atoms can easily diffuse across them and recombine. In addition, the enrichment of nickel compared with iron on the surface after vacuum firing favors recombination. Leisch [15,16] estimates that 50 recombination sites are available on 100 nm² of such a surface. These facts at least qualitatively



Figure 6.12 STM image of a vacuum fired stainless steel surface with (111) terraces and steps in between. The detail shows an area of 1 μm × 1 μm. (By courtesy of M. Leisch, University Graz, Austria.)

support the assumption that for the case of vacuum fired stainless steel desorption is the faster process than diffusion in the bulk.

6.4 Permeation

In the previous section, we assumed $n_{dis}(d) = n_{dis}(-d)$ as the boundary condition for calculating outgassing, that is, equal conditions on both surfaces of the vacuum wall. If, however, the outer surface is exposed to air while the interior surface is exposed to vacuum, it should be considered whether components of the air or other external gas species penetrate the vessel walls and propagate into the vacuum. This process is referred to as permeation and features three steps (Figure 6.13):

- adsorption of a molecule to the outer surface (generally, high-pressure side);
- diffusion through bulk material;
- desorption from the interior surface (low-pressure side).

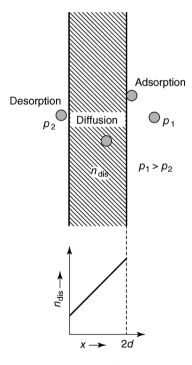


Figure 6.13 Processes during permeation. On the right-hand high-pressure (p_1) side, (net) adsorption of gas develops. The gas diffuses through the solid and (net) desorption is observed on the left-hand

low-pressure (p_2) side. At equilibrium, a linear concentration gradient establishes in the solid due to a homogeneous gas flow density throughout the entire system.

 p_1 , p_2 , and 2d denote the pressure of permeating gas on the high-pressure side, the pressure on the low-pressure side, and the thickness of the bulk material, respectively. If permeation is diffusion controlled, then the permeation rate j_{perm} depends on the kinetics of the involved molecules. Under stationary conditions,

$$j_{\text{perm}} = K_{\text{perm}} \frac{1}{2d} (p_1 - p_2),$$
 (6.42)

if the gas molecules do not dissociate on the considered material (e.g., N2, O2, H₂O on many different materials), that is,

$$j_{\text{perm}} = K'_{\text{perm}} \frac{1}{2d} (p_1^{0.5} - p_2^{0.5}),$$
 (6.43)

if a diatomic gas dissociates (e.g., H2 on stainless steel). In a stationary condition, a linear concentration profile develops between both surfaces (Figure 6.12). The constant K_{perm} is a characteristic for each molecule/material combination and heavily temperature dependent. K_{perm} is the product of solubility K_S and the diffusion constant D,

$$K_{\text{perm}} = K_{S} D. \tag{6.44}$$

Solubility K_S describes the ability of a substance to take up a certain type of (gas) particle (atom or molecule) from the surrounding. A concentration discontinuity of the corresponding particle species is observed at the boundary between the substance and the environment. As well as D, K_S is exponentially temperature dependent:

$$K_{\rm S} = K_{\rm S0} \exp\left(-\frac{E_{\rm S}}{RT}\right),\tag{6.45}$$

where K_{SO} is the solubility at $T \to \infty$ and E_S is the enthalpy of solution of the molecule. At higher density (pressure) and thermal equilibrium, K_S describes the concentration discontinuity

$$\frac{K_{\rm S}}{K_{\rm S0}} = \frac{n_{\rm i}}{n_{\rm o}} \quad \text{(with } n_{\rm i} \text{ the particle density inside, and } n_{\rm o}$$
the particle density outside) (6.46)

at the interface of the substance.

Example 6.10

Under the assumption that diffusion is rate determining, how high is the stationary permeation rate of air-born hydrogen ($p_{H_2} = 0.01 \, \text{Pa}$) into a vacuum vessel made of stainless steel with thickness 2d = 2 mm at 23 °C? According to Louthan [17],

$$\begin{split} K'_{\text{perm}} &= 6 \times 10^{-2} \, \text{Pa} \, \ell \, \text{s}^{-1} \, \text{cm}^{-1} \, \text{Pa}^{-1/2} \text{exp} \bigg(-\frac{59.9 \, \text{kJ mol}^{-1}}{\textit{RT}} \bigg) \\ &= 1.61 \times 10^{-12} \, \text{Pa} \, \ell \, \text{s}^{-1} \, \text{cm}^{-1} \, \text{Pa}^{-1/2}. \end{split}$$

The pressure p_2 in the vacuum chamber is negligible $p_2 \ll 0.01 \, \text{Pa}$.

According to Eq. (6.43),

$$j_{\text{perm}} = 1.61 \times 10^{-12} \text{Pa} \, \ell \, \text{s}^{-1} \, \text{cm}^{-1} \, \text{Pa}^{-1/2} \cdot \frac{1}{0.2 \, \text{cm}} \sqrt{10^{-2} \, \text{Pa}}$$

= $8.06 \times 10^{-13} \text{Pa} \, \ell \, \text{s}^{-1} \, \text{cm}^{-2}$.

Compared with typical outgassing rates, this gas flow is extremely low (see also the result in Example 6.9). Thus, the contribution of permeation to the residual pressure is important only when specially prepared or very thin stainless steel sheet metal is used. Furthermore, it should be taken into account that the oxide layer at the surface often acts as a hydrogen-diffusion barrier and additionally restrains permeation rates.

If permeation is controlled by sorption, then [18]

$$j_{\text{perm}} = \frac{s}{\sqrt{2\pi \, m \, k \, T}} (p_1 - p_2). \tag{6.47}$$

References

- 1 Venema, A. (1961) Processes limiting the ultimate pressure in ultra high vacuum systems, Trans 8th AVS Vac. Symp. 1-7.
- 2 Redhead, P.A., Hobson, J.P., and Kornelson, E.V. (1968) The Physical Basis of Ultrahigh Vacuum, Chapman and Hall Ltd, London, p. 68.
- 3 Hobson, J.P. (1961) Trans. 8th AVS Vac. Symp. 26.
- 4 Dylla, H.F., Manos, D.M., and LaMarche, P.H. (1993) J. Vac. Sci. Technol. A, 11, 2623-2633.
- 5 Redhead, P.A. (1995) Modeling the pumpdown of a reversibly adsorbed phase I. Monolayer and submonolayer initial coverage. J. Vac. Sci. Technol. A, 13, 467-475.
- 6 Dayton, B.B. (1961) Outgassing rate of contaminated metal surfaces, Trans. 8th AVS Vac. Symp. 42-57.
- 7 Kanazawa, K. (1989) J. Vac. Sci. Technol. A, 7, 3361.
- 8 Hayward, D.O. and Trapnell, B.M.W. (1964) Chemisorption, Butterworth, London.
- 9 Redhead, P.A. (1995) Modeling the pumpdown of a reversibly adsorbed phase II. Multilayer coverage. J. Vac. Sci. Technol. A, 136, 2791-2796.
- 10 Halsey, G.D. Jr. (1948) J. Chem. Phys., 16,

- 11 McMillan, W.G. and Teller, E. (1951) J. Chem. Phys., 19, 25.
- 12 Calder, R. and Lewin, G. (1967) Reduction of stainless steel outgassing in ultra-high vacuum. Brit. J. Appl. Phys., 18, 1459-1472.
- 13 Moore, B.C. (1995) Recombination limited outgassing of stainless steel. J. Vac. Sci. Technol. A, 13, 545-548.
- 14 Chun, I., Cho, B., and Chung, S. (1996) J. Vac. Sci. Technol. A, 14, 2636.
- 15 Leisch, M. (2008) Surface morphology, surface composition and outgassing behaviour of vacuum fired stainless steel. J. Phys.: Conf. Ser., 100, 092003.
- 16 Stupnik, A., Frank, P., and Leisch, M. (2009) Atom probe, AFM and STM studies on vacuum fired stainless steels. *Ultramicroscopy*, **109**, 563–567.
- 17 Louthan, M.R. and Derrick, R.G. (1975) Hydrogen transport in austenitic stainless steel. Corros. Sci., 15, 565-577.
- 18 Nemanic, V. and Bogataj, T. (1998) Outgassing of thin stainless steel chamber. *Vacuum*, **50**, 431–437.

Further Reading

Benard, J. (ed.) (1983) Adsorption on Metal Surfaces, Elsevier Scientific Publishing Comp., Amsterdam, Oxford, New York.

- Hauffe, K. and Morrison, S.R. (1974) Adsorption, Walter de Gruyter, Berlin-New York.
- Mikchail, R.Sh. and Robens, E. (1983) Microstructure and Thermal Analysis of Solid Surfaces, John Wiley & Sons, Ltd, Chichester.
- Prutton, M. (1983) Surface Physics, Oxford Physics Series 11, 2nd edn, Clarendon Press, Oxford.
- Redhead, P.A., Hobson, J.P., and Kornelsen, E.V. (1968) The Physical Basis of Ultrahigh Vacuum, Chapman and Hall Ltd, London (This book was reprinted by the American Vacuum Society in the 2000s).
- Wedler, G. (1970) Adsorption, Chem. Taschenbuch No. 9, Chemie, Weinheim. Zangwill, A. (1988) Physics at Surfaces, Cambridge University Press, Cambridge.

7

Positive Displacement Pumps

Dr. Karl Jousten,¹ Dr. Jürgen Dirscherl,² Dr. Rudolf Lachenmann,³ Dr. Alfons Jünemann,⁴ Dr.-Ing. Friedrichsen,⁵ Dr. Erik Lippelt,⁵ and Dr. Ing. Boris Kossek⁶

This chapter explains vacuum pumps based on the principle of positive displacement. These pumps are used as stand-alone pumps in the low and medium vacuum range or serve as backing pumps for high-vacuum pumps.

7.1

Introduction and Overview

Positive displacement pumps are the most important and most commonly used pumps in vacuum technology. According to DIN 28400, ¹⁾ Section 2 (1980), a positive displacement pump is defined as a "vacuum pump that aspirates, compresses, and discharges a gas to be pumped, using valves if necessary, by means of pistons, rotors, sliders, etc., which are sealed from another either with or without liquids."

Oscillating pumps are the simplest positive displacement pumps (Section 7.2). Historically, they are among the first pumps used to generate vacuum (Chapter 1). Either a diaphragm with a connecting rod or a piston aspirates gas through an inlet valve during one half-cycle of the motion and ejects the gas during the other half-cycle via a discharge valve.

In the beginning of the twentieth century, liquid sealed rotating positive displacement pumps were developed to yield lower pressures and higher pumping speeds. They can be used in the low vacuum range, certain types of pumps also far into medium vacuum. These pumps use sickle-shaped suction and compression

- Translator's note: corresponds to ISO 3529.
- 2) Translator's note: translated from the German.

¹Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

²Vacuubrand GmbH & Co, Alfred-Zippe-Str. 4, 97877, Wertheim, Germany

³(retired from) Vacuubrand GmbH & Co, Alfred-Zippe-Str. 4, 97877, Wertheim, Germany

⁴Sterling SIHI GmbH, Lindenstr. 170, 25524, Itzehoe, Germany

⁵Dr.-Ing. K. Busch GmbH, Schauinslandstr. 1, 79689, Maulburg, Germany

⁶Busch Produktions GmbH, Schauinslandstr. 1, 79689, Maulburg, Germany

chambers. In sliding vane rotary pumps, the chambers are sealed by vanes arranged in a rotor. Rotating plunger pumps use a piston and an eccentrically guided rotary plunger arranged inside a housing to seal the chambers. Liquid ring pumps feature working liquid and impeller blades in order to seal the chambers. In operation, the sickle-shaped suction chamber forms anew in every cycle, starting with zero volume each time. As there is no dead space, pumping speed is high, far into the medium vacuum range.

In the mid-1950s, and particularly towards the end of the twentieth century, dry-running rotating positive displacement pumps, that is, without liquids such as oil or water for lubrication or sealing, were developed. Listed in the order of their introduction to the market, these include Roots pumps, multistage pumps (dry sliding vane rotary pumps), tooth-type rotary pumps (claw pumps), scroll pumps, and screw-type pumps. They combine the advantages of being oil-free such as oscillating positive displacement pumps and of having high pumping speeds such as rotating displacers.

Distinct PNEUROP acceptance guidelines [1] and DIN standards for rating assessments demonstrate the significance of positive displacement pumps. ISO standardization includes these regulations as well (see also Table A.22).

Pumping speed is the most important characteristic of a positive displacement pump. According to the acceptance criteria and standards [1], pumping speed S is the volume flowing from a given measurement dome at given pressure through the inlet cross section of a vacuum pump per unit time. The units used for S are m^3h^{-1} and ℓ s⁻¹.

Table 7.1 provides an overview of positive displacement pumps introduced in this chapter. Here, oscillating and single- or twin-shaft rotating displacers are differentiated according to the driving principle. Screw-type pumps transport gas axially along the screws, as opposed to twin-shaft claw or Roots pumps that deliver radially. However, all three types of pumps include two drive shafts, spindles, or screws. The basic operating principle of scroll pumps uses two geared spirals to transport the gas. The principle is very similar to that of twin-shaft

Table 7.1 Taxonomy of positive displacement pumps according to the driving principle. The bracketed numbers indicate the corresponding sections that cover the specified pump type.

Positive displacement pumps							
Oscillating positive displacement Rotary positive displacement pumps pumps (7.2)							
Piston pumps (7.2.1)	Single-spool (7.3)	Twin-spool (7.4)					
Diaphragm pumps (7.2.2)	Liquid ring pumps (7.3.1)	Screw-type pumps (7.4.1)					
	Sliding vane rotary pumps (7.3.2)	Claw pumps (7.4.2)					
	Multistage pumps (7.3.2.2)	Roots pumps (7.4.3)					
	Rotary plunger pumps (7.3.3)						
	Trochoidal pumps (7.3.4)						
	Scroll pumps (7.3.5)						

pumps but, in scroll pumps, one spiral is generally fixed, for why this type of pump is categorized as a single-shaft drive.

In contrast to other types of pumps that are usually used for low vacuum, Roots pumps, named after the first person to use them in air compression, are used mainly in the medium vacuum range. They feature two symmetrically designed pistons moving on rolling contact and synchronized by a gear pair, so that only a small gap remains between both pistons as well as between the pistons and the pump housing. Due to noncontact operation, the pistons can rotate at high speed, and thus, high pumping speed is obtained with small pumps. At high pressures, however, high gas backflow through the gaps develops, leading to poor compression. For operation, a Roots vacuum pump, therefore, usually requires one of the above-mentioned pumps as a backing pump, which compresses to atmospheric pressure.

All types of positive displacement pumps are often used as main pumps in vacuum systems. However, they may also serve as fore pumps to steam-jet or vapor pumps, fluid entrainment pumps, and turbomolecular pumps, or as auxiliary pumps to ion pumps, getter pumps, and cryopumps. Choosing the appropriate, most efficient combination of pumps is an important criterion for a particular application.

Many processes in industry and research use oil-sealed rotating vacuum pumps for generating low and medium vacuum. They are favorable due to their relatively high pumping speeds, wide operating ranges of pressures (see Table A.15), high compression ratios, and because they allow continuous operation. However, two basic disadvantages of using oil are:

- backward migration of oil (vapor and eventually liquid) to the inlet with possible contamination of vacuum (process) chamber (see Section 7.5.2), and
- contaminated and, therefore, deteriorated pump oil due to pumped-down process gas and vapors, often with accumulated solid particles (dust) (see Section 7.8).

More or less costly measures are necessary in order to avoid or at least reduce these disadvantages, depending on the particular application. These include additionally attached filters of all sorts as well as usage of expensive specialpurpose oils (see Table A.17) and systems for their purification and reuse.

Facing these problems, semiconductor industry, in particular, called for a lubricant-free rotating positive replacement pump. Vacuum systems in this branch of industry use processed gases that are often toxic and/or corrosive. When these gases are pumped down, liquid or solid particles can form along the path through the pump, especially in contact with humidity. All of this reduces the lubricating properties in the suction chamber and makes environmentally acceptable disposal of the oil considerably more difficult. Since approximately 1987 [2], such dry systems are available on request.

After introducing the pumps listed in Table 7.1 section wise, Section 7.5 covers specific characteristics of oil-sealed rotating pumps in general. Principles of gas ballast and power requirements, common to all types of positive displacement pumps, are discussed in Section 7.6. General operating and safety instructions for positive displacement pumps are discussed in Section 7.7. Section 7.8 describes accessories that are required for many processes and are connected directly to mostly oil-sealed positive displacement pumps.

7.2 Oscillating Positive Displacement Pumps

This type of pump includes piston and diaphragm pumps. Both types are dry pumps in which the transferred gas or the gas vapor mixture is not exposed to any sealing or lubricating agents. Either an oscillating piston, connected via a shaft and connecting rod, or a diaphragm with connecting rod aspirates the gas during one half-cycle of motion and ejects the gas during the second half-cycle (through a valve).

Even in the most precisely manufactured systems, the basic design of these pumps leads to a so-called dead space after ejecting at the dead center of the piston or connecting rod. From this space, the gas to be pumped is not pushed into the exhaust line. Therefore, piston displacement pumps with a high ratio of suction chamber volume to dead space develop typical ultimate pressures in the range of 0.1-1 Pa, while diaphragm pumps with lower compression ratios due to limited motion freedom of the diaphragm usually develop ultimate pressures around 100 Pa in single-stage operation.

7.2.1

Piston Pumps

A large variety of small piston pumps is available for technical applications. In vacuum technology, they have gained in importance recently in the realm of oilfree vacuum pump development. Initially, single- and two-stage designs of these pumps developed ultimate pressures of approximately 500 Pa and 10 Pa, respectively. Figure 7.1 shows a model of a simple reciprocating piston pump. During the downward movement with open right-hand inlet valve, the pump aspirates gas. For the period of upward motion, the outlet valve is open, the inlet valve is closed, and the pumped gas is compressed and ejected. A dead space remains at the top dead center (TDC) from which no gas is ejected. The gas remaining in this dead space at exhaust pressure expands during the subsequent inlet stroke and partially or fully fills the active volume, thus preventing new gas from being aspirated. The compression ratio of a positive displacement pump with dead space is therefore limited to the ratio of its maximum working volume to the dead space.

Recent developments, for example, four-stage pumps, have reduced the ultimate pressure considerably (down to approximately 2 Pa) in order to allow for direct connection to a high-vacuum pump [3]. Figure 7.2 shows a schematic arrangement of a four-stage dry-running pump, a further development of the reciprocating piston pump [4,5]. The diameter of the four identical pistons is 100 mm; the cylinder height is 25.4 mm (1 inch). Due to the relatively low stroke

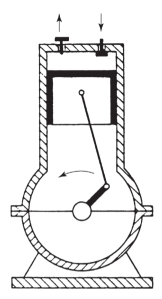


Figure 7.1 Diagram of a reciprocating piston pump.

frequency of 1200 min⁻¹, the pressure-dependent, maximum (net) pumping speed is low, $34 \text{ m}^3\text{h}^{-1}$. The ultimate pressure is in the range of 1.5–3 Pa.

After some time of operation, in addition to the dead space, the gas leaking in at the cylinder from the outside due to wear also leads to an increase of ultimate pressure.

Modern surface treatments as well as materials such as fluoroplastic coatings and pistons with special sliding surfaces have led to considerable advances [6].

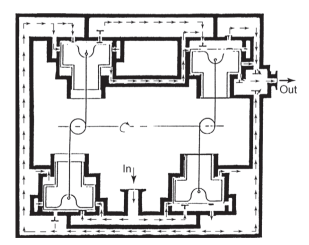


Figure 7.2 Diagram of a four-stage reciprocating piston pump with a compression of 50 000 [4,5].

Nevertheless, the disadvantage of the gap remains, as opposed to a hermetically sealed, fixed diaphragm (see the following section). Small amounts of pumped media can always leak from the drive system through the gap and cause corrosion of bearings, etc. Also, pumped particles can cause wear on the sealing surfaces of the gap.

7.2.2

Diaphragm Pumps

The origin of modern diaphragm pumps is unknown. In the 1940s, development of elastomer materials promoted widespread use of diaphragm pumps and compressors. Further advances in elastomer materials in the early 1960s led to diaphragm pumps with mechanical diaphragm drives featuring higher pumping speed, lower ultimate pressure, and above all, longer service life of the diaphragms [7]. Additional progress in materials technology as well as new design and fabrication methods such as CAD and CNC contributed to the development of today's diaphragm vacuum pumps as standard laboratory pumps.

Towards the end of the last century, two main steps characterize the development: the introduction of chemical diaphragm pumps that use chemically resistant fluoroplastics for chemical laboratories in the early 1980s, and, around 1990, the emerging multistage diaphragm pumps as oil-free backing pumps for the newly developed turbo molecular pumps with molecular stage. Additionally, materials and manufacturing optimizations, three- and fourstage diaphragm pumps, and kinetic improvements due to the introduction of speed-controlled drive systems provided process-controlled vacuum systems with pumping speeds and ultimate pressures found only in rotating vane pumps, at the time.

7.2.2.1 Design and Principle of Operation

Diaphragm vacuum pumps are oscillating positive displacement pumps according to the definition in ISO 3529-2. Today, diaphragm vacuum pumps are built in a large variety of different sizes with individual vacuum technological properties. For reasons of physics and mechanics, they cover a range of low pumping speeds only. Diaphragm pumps are manufactured with pumping speeds up to approximately 20 m³h⁻¹; however, the pumping speed of most commercially available pumps is below approximately 12 m³h⁻¹. As for ultimate vacuum, they are limited to >10 Pa (0.1 mbar) for physical reasons.

Figure 7.3 shows the design of a diaphragm pump: the head cover and the diaphragm with the diaphragm clamping disk define the pumping chamber. The diaphragm is fixed between the head cover and the housing at the outer perimeter. The upward and downward motion of the connecting rod results in a

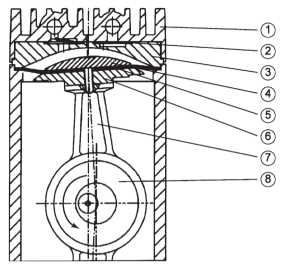


Figure 7.3 Diagram of a diaphragm pump stage: ① housing, ② valves, ③ head cover, ④ diaphragm clamping disk, 3 diaphragm, 6 diaphragm supporting disk, 2 connecting rod, 8 eccentric rotor (crank shaft).

periodic change of the suction volume in aspiration and compression phases. Gas-flow-controlled inlet and outlet valves are mounted between the housing lid and the head cover.

The gas in the dead space remaining at the top of stroke is not ejected. During the following inlet stroke, this volume expands anew and fills the working volume partially, thus limiting ultimate vacuum. The compression ratio of a positive displacement pump is limited to the ratio of maximum working volume and dead space.

7.2.2.2 Pumping Speed and Ultimate Pressure

The pumping speed of a diaphragm pump is a function of the suction chamber volume $V_{\rm S}$, rotational speed n, and dead space $V_{\rm D.S.}$.

The effective pV flow q_{pV} then is

$$q_{pV} = n(V_{\rm S} p_{\rm in} - V_{\rm D.S.} p_{\rm out})$$
 (7.1)

($p_{\rm in}$ inlet pressure; $p_{\rm out}$ outlet pressure, generally, atmospheric pressure), and the effective pumping speed S of a single-stage pump

$$S = \frac{q_{pV}}{p_{\text{in}}}. (7.2)$$

Here, backflow is neglected. The pumping speed of a multistage pump is calculated from a system of equations incorporating analogous expressions for the individual stages. The equations are linked by the condition of gas-flow conservation [8].

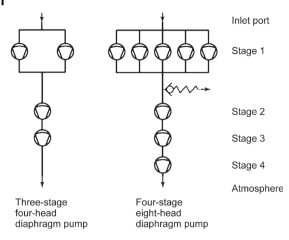


Figure 7.4 Examples of heads connected in parallel and in series. High pumping capacity by parallel connection of intake heads and with a pressure relief valve (if appropriate).

A more accurate calculation would have to take into account the backflow through gaps and the influence of flow dissipation in the inlet. Leakage, for example, through valves, if unavoidable, has negative effect on the theoretically achievable ultimate pressure of the pump.

Typical values of compression ratios $p_{\rm out}/p_{\rm in}$ range from 10 to 30. Vacuum technical connections of pump heads in series lead to lower ultimate pressure. Single- to four-stage arrangements are common (Figure 7.4).

An increase in pump chamber volume of a diaphragm pump causes structural and fabricating problems. Therefore, parallel connections of pump heads are favorable when higher pumping speed of up to approximately $20\,\mathrm{m}^3h^{-1}$ is requested. Use of diaphragm pumps is limited and restricted mainly to laboratory and pilot installations due to the limited pumping speed.

Figure 7.4 shows diaphragm pumps in different three- to four-stage arrangements. A four-cylinder pump, for example, can be connected single-, two-, three-, or four-stage-wise, all featuring the same external dimensions. This leads to the following vacuum technological data (see also plots in Figure 7.5):

- single-stage, 8 m³h⁻¹ at atmospheric pressure (A), 70–80 mbar (hPa) ultimate pressure;
- two-stage, 4 m³h⁻¹ (B₁), approximately 7–9 mbar (hPa) ultimate pressure;
- three-stage, 2.8 m³h⁻¹ (B₂), approximately 2 mbar (hPa) ultimate pressure;
- four-stage, 2 m³h⁻¹ (C), approximately 0.6 mbar (hPa) ultimate pressure.

7.2.2.3 Gas Ballast

Greater amounts of condensate in a pump cause mechanical pressure peaks due to the limited dead space, and therefore, decrease the service life of valves and diaphragms. Chemical diaphragm pumps, in particular, are often equipped with

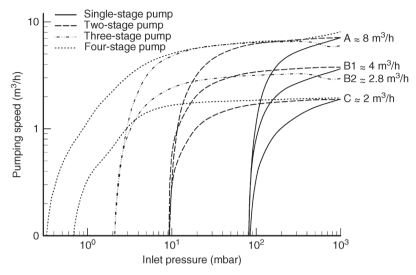


Figure 7.5 Pumping capacities and ultimate pressures of geometrically same-sized multistage diaphragm pumps with the same number of heads but different connection layouts for the heads.

a gas ballast device, which either prevents condensation or discharges condensate produced by the pump [9]. As in sliding vane rotary pumps, electromagnetic gas valves with process-determined control are used here as well [10].

7.2.2.4 Drive Concepts

Diaphragm pumps for laboratory use are usually driven by monophase motors with a nominal rotational speed of 1500 revolutions per minute (at 50 Hz power frequency). Increasingly, variable-speed diaphragm pumps are used that feature better characteristics in terms of pumping speed, ultimate pressure, and controllability.

Two types of speed controlled motors are applied (see example in Figure 7.6):

- 1) Frequency converter-controlled three-phase AC motors, preferably for large pumps.
- 2) Electronically commutated, brushless DC motors, being far more compact than three-phase AC motors with frequency converters.

Table 7.2 shows the increase in maximum pumping speed by up to 40%, and the lowering of the ultimate pressure by a factor of 2, with the same mechanical vacuum unit by using a speed-controlled drive.

7.2.2.5 Ultimate Pressure

Typical ultimate pressures for diaphragm pumps with fixed speed and flat diaphragm are approximately 70–80 mbar (hPa) for single-stage designs, approximately 9 mbar (hPa) for two-stage designs, approximately 2 mbar (hPa) for



Figure 7.6 Two diaphragm pumps by VACUUBRAND: (a) MV 10 NT VARIO: $12.1 \text{ m}^3 \text{ h}^{-1}$, 30 Pa; (b) MD 1 VARIO-SP: $1.8 \text{ m}^3 \text{ h}^{-1}$, 100 Pa (compare Table 7.2).

three-stage designs, and approximately 0.6 mbar (hPa) for four-stage designs. The dead space can be reduced and the ultimate pressure improved by tangential fixing and/or by using preformed diaphragms that move to direct contact with the wall of the compression volume in the top of stroke [7,11].

A speed-controlled, four-stage diaphragm pump reaches 30 Pa ultimate pressure, the lowest of all commercially available diaphragm pumps [12].

Table 7.2 Selected diaphragm pumps with a single-phase AC motor (fixed frequency), frequency converter-controlled three-phase AC motor (VARIO), and brushless DC motor (VARIO-SP) (VACUUBRAND).

Model	Length (mm)	Weight (kg)	Maximum pumping speed (m ³ h ⁻¹)	Ultimate pressure (hPa)
MD 1 230 V/50 Hz AC	303	6.5	1.2	1.5
MD 1 24 V DC Vario-sp	223	4.1	1.8	1
MD 4 230 V/50 Hz AC	325	16.4	3.3	2
MD 4 VARIO	325	18.6	4.3	1
MD 4 24 V DC Vario-sp	260	12.2	4	1
MV 10 NT (AC)	554	30.6	10.4	0.5
MV 10 NT VARIO	554	31.2	12.1	0.3

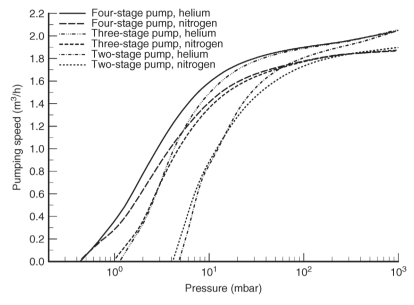


Figure 7.7 Pumping capacities for helium and nitrogen versus inlet pressure for two-, three-, and four-stage diaphragm pumps.

7.2.2.6 Influence of Gas Species on Pumping Speed and Ultimate Pressure

The displaced volume and the suction volume geometry mainly determine pumping speed and ultimate pressure of diaphragm pumps. In practice, the gas composition has a minor influence on both parameters (Figure 7.7). A 10% increase in pumping speed for light gases, compared to nitrogen, is probably due to better filling of the suction chambers because of different gas viscosities [13,14]. The influence of memory effects on the diaphragm material is below measuring sensitivity for plane diaphragms [15].

7.2.2.7 Influence of Rotational Speed on Ultimate Pressure

At lower pressure ($<10\,\text{mbar}$ (hPa)), gas forces on the valves are very low. This also contributes to a speed-dependent ultimate pressure (Table 7.2, Figure 7.8). Because of lower driving forces, the valves require more time to follow the gas flow at lower pressures. The lowest ultimate pressure is not obtained at higher, but at lower rotational speed than the typical nominal revolutions of 1500 min⁻¹. This behavior is important for the use of diaphragm pumps as backing pumps for turbomolecular pumps (see Section 7.2.2.10).

7.2.2.8 Design Principles

Pump Heads and Diaphragm

The geometric design of pump heads is of particular importance. Modern, computer aided design and CNC-fabricating techniques allow adapting the geometry of pump heads, and if necessary clamping disks, optimally to the motion and to

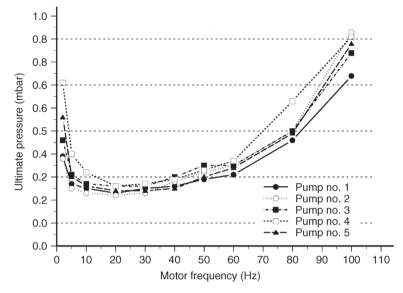


Figure 7.8 Speed dependence of ultimate pressure for several four-stage diaphragm pumps.

the surface of the diaphragm. This applies likewise to preformed or structured as well as flat diaphragms. Flat diaphragms are punched out of a flat elastomer plate and are fixed between a clamping disk and a supporting plate. Preformed and structured diaphragms are vulcanized onto a usually metallic core inside a mold resulting in a higher amount of elastomer. Aging after vulcanizing as well as embrittlement of elastomers are discussed in [12]. A flat diaphragm requires higher precision mechanical fabrication of head components. Then, it shows excellent results with respect to pumping speed, ultimate pressure, and service life, which, today, usually exceeds 15 000 h. Manufacturers' ratings of different diaphragm designs vary in terms of advantages and disadvantages.

Valves

As valves, gas-flow-controlled flapper valves are common. The desired maximum conductance stands in contrast to geometrical restrictions of valve shape and reliable sealing at the inlet and outlet seats. Standard materials for valves are FKM (e.g., Viton®), PTFE (e.g., Teflon®), PEEK, or FFKM (e.g., Kalrez®, Chemraz®), depending on the application. Single-stage diaphragm pumps generate reasonable ultimate pressures, even with relatively stiff valves of PTFE or PEEK. Diaphragm pumps with very low ultimate pressures require lightweight, elastic, and very well sealing valves as well as nonwetting and adhesion-reduced valve seats.

Materials

For gas-wetted parts, metallic materials as well as fluoroplastics are used. The latter have significant advantages in corrosive applications or in presence of condensing media as opposed to liquid-sealed and lubricated rotating pumps, which,

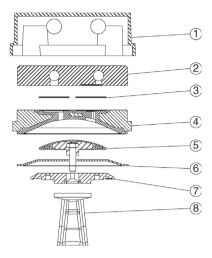


Figure 7.9 Exploded view of a pump head for chemical applications. 1: housing cover; 2: housing cover insert (carbon-fiber-reinforced PTFE); 3: valves (PTFE or FFKM); 4: head cover (ceramic-reinforced PTFE or ETFE with

reinforcing core); 5: diaphragm clamping disk (ETFE with reinforcing core); 6: flat diaphragm (PTFE) with fabric-reinforced FKM support; 7: diaphragm-supporting disk; 8: connecting rod.

for design and tolerance aspects, require metallic materials that are subject to corrosion.

Figure 7.9 illustrates the design of a pump head for chemical applications. In spite of the used reinforced PTFE, creeping under load can lead to undesired changes in geometry. If the PTFE is completely housed in (2) by an appropriate casing cover (1), mechanical stresses are carried by the metallic components, and the pumped-down gas is still not exposed to the metal.

Requirements on the head cover (4) and the clamping disk (5) are high in terms of chemical resistance and mechanical stability. Mechanical deformation of only 0.1 mm is tolerable for silent operation and constantly high-vacuum performance of chemical application diaphragm pumps, and this, throughout a service life of many years and under high mechanical and thermal load. Solutions combining mechanical and material advantages use fiber-reinforced thermoplastic fluoroplastics with wall thicknesses $> 0.5\,\mathrm{mm}$ molded around stabilizing inlets made of high-strength plastics or metals.

7.2.2.9 Diaphragm Pumps in Chemical Laboratories

During the last two decades, diaphragm pumps have become the dominating pump type in chemical laboratories. Other than oil-sealed sliding vane rotary pumps, they do not require a cooling trap on the vacuum side to protect the pump. In a diaphragm pump, pumped-down solvents can be retained effectively, thermodynamically favorable, on the exhaust side at atmospheric pressure and room temperature. Specially designed, appropriate emission condensers are combined with diaphragm pumps to compact pumping units (Figure 7.10) [15–17].



Figure 7.10 Chemistry diaphragm pump PC 3001 VARIOpro by VACUUBRAND: $2\,\mathrm{m}^3\,\mathrm{h}^{-1}$, 200 Pa, with solvent recovery.

For such units, vacuum control is of particular importance [18]. Modern electronics and new algorithms allow simple control of different processes as well as automatic operation without any necessary programming [19].

7.2.2.10 Diaphragm Pumps as Backing Pumps to Turbomolecular Pumps

Development of modern turbomolecular pumps with improved critical backing pressure at outlet pressures of up to 30 mbar (hPa) allows utilization of oil-free diaphragm pumps as backing pumps. The required size of the fore-vacuum pump depends on a number of factors, for example, the desired pump-down time (Figure 7.11).

If a wide-range turbomolecular pump is used in a pressure range higher than the ultimate vacuum, that is, at high gas load, the pumping speed of the diaphragm pump must be rated for maximum gas throughput at the optimal operating point of the wide-range turbomolecular pump [20].

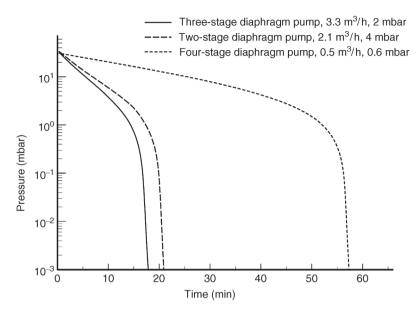


Figure 7.11 Pump-down time for a 100 ℓ vacuum chamber using a wide-range turbomolecular pump (20 ℓ s⁻¹) with three different backing pumps.

Low ultimate pressures of fore-vacuum pumps reduce the power consumption and cooling demand of wide-range turbomolecular pumps due to lower gas drag [21]. Also, low ultimate pressure of a backing pump contributes to lower ultimate pressure performance of a turbomolecular pump in the high vacuum range and increased compression values for light gases. Therefore, a diaphragm pump should provide an ultimate pressure <5 mbar (hPa) although wide-range turbomolecular pumps tolerate a fore vacuum of up to 30 mbar (hPa).

Service life and maintenance rates are determined primarily by the service life of diaphragm and valves. Flat diaphragms and optimized geometries yield diaphragm service lives of more than 15 000 h. Speed-controlled diaphragm pumps (e.g., VACUUBRAND® "VARIO®") reach considerably higher diaphragm service lives and thus reach the maintenance intervals of turbomolecular pumps.

For corrosive gas applications, only chemical-type diaphragm pumps are used, in which gas-wetted parts are made of highly corrosion proof and chemically resistant materials such as fluoroplastics.

The influence of diaphragm pumps as backing pumps on the residual gas spectrum, particularly on hydrogen partial pressure, is discussed in [22] and illustrated in Figure 7.12. As indicated by the diagram in the lower image in Figure 7.12, a variation in fore-vacuum pressure influences hydrogen partial pressure. Shifting the working pressure to an area with better compression ratio in the turbomolecular pump reduces hydrogen partial pressure substantially. An influence on hydrogen partial pressure due to memory effects in the diaphragm could not be detected within measurement accuracy [18].

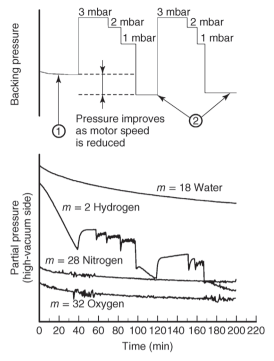


Figure 7.12 Residual gas composition of a tur- 1500 min⁻¹: ultimate pressure 0.3 mbar (hPa) bomolecular pump (lower image) versus forevacuum pressure (upper image). The forevacuum pressure is varied solely by changing the speed of the four-stage diaphragm pump (without any gas intake): ① operation at

(30 Pa), ② operation at self-optimized speed for minimum ultimate pressure (approximately 700 min⁻¹), ultimate pressure approximately 0.1 mbar (hPa) (10 Pa).

Compared to constant-speed pumps, speed-controlled VARIO® diaphragm pumps considerably improve the capacity of the overall system when combined with wide-range turbomolecular pumps. During pump-down of a vacuum system or in processes with high gas throughput, the speed of the diaphragm pump is increased as required, up to 2400 min⁻¹. After the ultimate pressure of the pump is reached, the speed of the VARIO® diaphragm pump is lowered considerably. VARIO[®] pumps reduce pump-down time to 10 mbar (hPa) by approximately 20%. A special control algorithm automatically adjusts the speed to deliver the best ultimate pressure in the diaphragm pump (VACUUBRAND "Turbo Mode").

When operating at high and ultrahigh vacuum, a wide-range turbomolecular pump delivers low mass flow only. In many cases, a speed of $< 500 \, \rm min^{-1}$ is sufficient for the VARIO® diaphragm pump to cope with this gas flow. This controlled reduction of speed reduces the average number of strokes and thus yields noticeably higher diaphragm service life. Therefore, service life and maintenance intervals of VARIO® diaphragm pumps reach the high service life of over

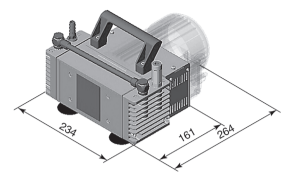


Figure 7.13 Very compact diaphragm pump as a backing pump to turbomolecular pumps: MZ 2 VARIO-SP: $2.5 \text{ m}^3\text{h}^{-1}$, 9 mbar (hPa), 6.5 kg. For comparison: MZ 2 with a single-phase

AC motor (shaded in the image): $1.9 \text{ m}^3 \text{h}^{-1}$, 9 mbar (hPa), 10.1 kg, approximately 100 mm longer than the DC model.

40 000 h of wide-range turbomolecular pumps. Furthermore, diaphragm pumps in this operating mode run very silently and smoothly. Additionally, the forevacuum pressure at the wide-range turbomolecular pump is improved (see also Figure 7.8).

Diaphragm pumps are often used in oil-free leak detectors. While scroll pumps feature higher pumping speed and lower ultimate vacuum, not required, however, for wide-range turbomolecular pumps, diaphragm pumps have the advantage of better sealing and considerably higher service lives of wearing parts. This is true, even more, for speed-controlled diaphragm pumps, which also reduce noise levels and power consumption.

Figure 7.13 shows a remarkably compact solution for a mobile, dry leak detector: a two-stage diaphragm pump with electronically controlled DC drive.

7.2.2.11 Diaphragm Pumps Combined with Other Types of Vacuum Pumps

In certain applications, combinations of diaphragm pumps and pumps with different types of operating principles are favorable.

Combining diaphragm pumps and Roots pumps [23] (Section 7.4.3) yields an increase in pumping speed and lower ultimate pressure while generally maintaining the advantages of the diaphragm pump, namely oil-free suction chambers. Depending on the size of the Roots pump and the diaphragm pump, pumping speeds vary between 6 and 35 m³h⁻¹ and ultimate pressures from 10 to 30 Pa. Gas-load controlled, frequency converter-operated Roots pumps are available as very compact units without bypass and yield high pumping speeds even at comparably high pressures.

Very common as a chemical hybrid pump in certain medium-vacuum laboratory applications is a combination of a sliding vane rotary pump and a diaphragm pump (Figure 7.14). This corrosion-optimized combination of a twostage sliding vane rotary pump and a diaphragm pump is made of widely chemically resistant materials. The diaphragm pump continuously evacuates the

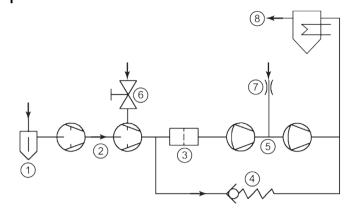


Figure 7.14 Chemical HYBRID pump system with ① (optional) separator at the inlet and ® emission condenser. The four-stage pump includes ② two sliding vane rotary pump stages with ⑤ two succeeding diaphragm

pump stages, and in between, ③ an oil separator with overpressure relief valve ④. Gas ballast for the sliding vane rotary pump is operated manually as required ⑥ and continuously for the diaphragm pump ⑦.

oil reservoir of the sliding vane rotary pump module. This prevents condensation problems in oil-sealed components and corrosion problems in nearly all possible cases [24,25].

Only special applications feature combinations of diaphragm pumps and scroll pumps (Section 7.3.5). By this, pumping speed and, in particular, ultimate pressure of small, single-stage scroll pumps, limited in terms of ultimate pressure due to internal leakage, can be improved considerably.

Combinations of claw pumps or screw-type pumps (Sections 7.4.2 and 7.4.1) with small diaphragm pumps, operated as auxiliary pumps if required, can reduce power consumption of large dry-runners by up to 70%, depending on design, which then do not need to compress to atmospheric pressure.

7.3 Single-Shaft Rotating Positive Displacement Pumps

7.3.1

Liquid Ring Vacuum Pumps [26-32]

As early as 1890, the liquid ring vacuum pump was invented as a so-called water ring pump, still comparable to basic designs of today. Due to the robust operating behavior and characteristics, its compressor design is one of the most prominent in vacuum pumps of chemical engineering. But also in other branches of industry, it is widespread as a main pump or fore pump, combined with other types of vacuum pumps, for low- and medium-high vacuum production. Established applications include, for example, energy production, plastics industry, medical technology, food and beverage industry, paper production, and building-materials industry.

Liquid ring vacuum pumps are capable of pumping nearly any type of gas and vapor. These machines feature an oil-free operating principle, low temperature levels, and the possibility to deliver liquids alongside the gas flow. Increasingly, applications in process technology emerge in which heat and material exchange or chemical reactions are initiated inside the pump.

Liquid ring compressors are built for suction volume flows below $10 \text{ m}^3\text{h}^{-1}$, and up to above $10\,000 \text{ m}^3\text{h}^{-1}$.

7.3.1.1 Design and Principle of Operation

In liquid ring pumps, a rotating ring of liquid transmits momentum and energy to the delivered medium (Figure 7.15). An impeller drives the ring. Nearly isothermal compression is obtained due to the intense contact between the delivered gas and the operating liquid.

The impeller is suspended eccentrically in a cylindrical housing (central body). During operation, the rotating impeller carries along the pump fluid, which is driven outward by centrifugal forces thus establishing a uniform fluid ring. The impeller rises from the liquid ring on one side and submerges into it on the other. Together with the liquid ring, the impeller blades form separate chambers, and thus, the liquid enters into and emerges from the impeller cells in a piston-like motion during rotation.

The lateral limits of the impeller cells are formed by guide disks equipped with aspirating and pressure holes. The suction channel is arranged in the area where

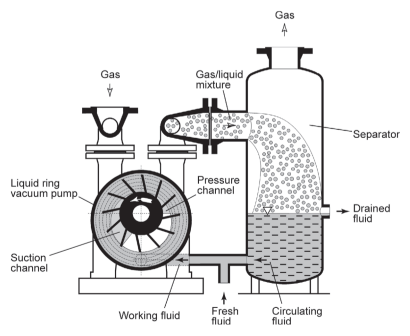


Figure 7.15 Operating principle of a liquid ring vacuum pump.

the blades rise from the liquid. The pressure channel is at the opposite side where the blades submerge.

Besides compressing the pumped gas, the working fluid has additional functions. These include sealing the axial gap between the impeller and the guide disk as well as transporting compression heat from the pumped gas. Due to the intense contact between gas and liquid, the working liquid completely absorbs the heat produced during compression. Thus, the temperature of the pumped gas hardly rises and the compression can be considered quasi-isothermal. Compared to other types of vacuum pumps, exhaust temperatures are low.

During operation, part of the liquid forming the ring is ejected continuously on the outlet side, together with the pumped gas. A connection is provided in order to allow supply of working liquid.

Water is the standard working fluid in most applications. However, in chemical engineering and pharmaceutical industry, for example, where reactive gases and vapors are pumped, chemical properties of the delivered medium must be taken into account when selecting the working fluid.

7.3.1.2 Operating Properties and Dimensioning

Physical properties of the working fluid, particularly vapor pressure (Section 3.5.1), influence the pumping behavior of a liquid ring vacuum pump.

When dry gases are sucked in, a small amount of working fluid evaporates at the inlet of the vacuum pump. A state of saturation establishes in the impeller cell, and thus, only part of the cell's volume is available for gas transport.

In a vapor-saturated gas mixture, the partial pressure of the vapor is equal to its vapor pressure. According to Dalton's law (Section 3.1.4), the proportion of vapor is proportional to the partial pressure:

$$\frac{V_{\text{vapor}}}{V_{\text{tot}}} = \frac{p_{\text{vapor}}}{p_{\text{tot}}}.$$
(7.3)

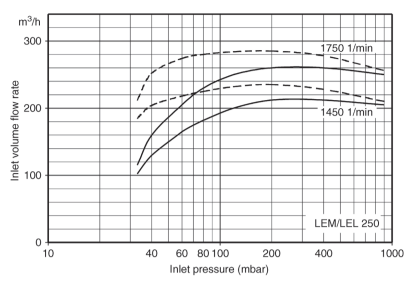
This shows that the effective amount of displaced volume provided for gas transport by the impeller cells depends on the temperature, that is, vapor pressure, of the working fluid and on the inlet pressure.

When vapor-saturated gas is aspirated, no working fluid evaporates in the impeller cells on the inlet side. Thus, the complete cell volume is available for gas transport and the pumping speed is not reduced. If vapor from the transported gas mixture condenses in the inlet of the pump due to a temperature drop, this results in an additional increase in pumping speed. This condensation effect is particularly apparent for hot, vapor-saturated gas mixtures.

Figure 7.16 shows a plot of pumping speeds and power consumptions over inlet pressure for transport of dry and water-vapor saturated air.

Generally, modern liquid ring vacuum pumps work down to inlet pressures of approximately 30 mbar (hPa).

The dashed plot in Figure 7.16 represents the increased pumping speed for water-vapor saturated air due to the condensation effect.



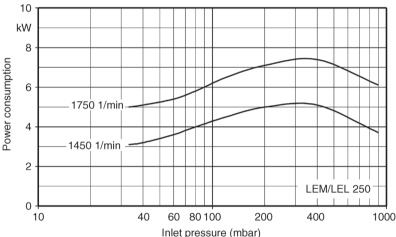


Figure 7.16 Characteristic curves for the liquid ring vacuum pump LEM/LEL 250 (Sterling SIHI): pumping capacity (inlet volume flow) and power consumption versus inlet pressure at selected speeds.

Manufacturers' catalogs plot the pumping speed (inlet volume flows) over inlet pressure. They apply to dry or water-vapor saturated air as aspirated medium at 20 °C and water as working fluid at 15 °C. Here, compression pressure is atmospheric pressure of 1013 mbar (hPa). Assessment of the characteristic curves is performed in accordance with the acceptance criteria defined in DIN 28431 [32].

For selecting liquid ring vacuum pumps, catalog data is converted to fit the actual operating conditions in the considered application. Power consumption and pumping speed, in particular, can be influenced considerably under different

operating conditions. In addition to the temperatures of the pumped-down medium and the working fluid, the state of the pumped gas, in terms of dry or vapor-saturated condition, is essential.

Using Dalton's law, Eq. (7.3), and assuming that the temperature ϑ_A of the pumped air adjusts to the temperature θ_B of the working fluid by the time it enters into the impeller cells, pumping speed S calculates to:

$$S_{A,dry} = S_K \left(\frac{\theta_A + 273}{\theta_B + 273} \times \frac{288}{293} \times \frac{p_A - p_{\text{vapor,B}}}{p_A - 17.04 \text{ hPa}} \right).$$
 (7.4)

Here, S_{K} is the pumping speed at inlet pressure p_{A} (in hPa) according to catalog conditions (DIN 28431). The bracketed term takes into account the air temperature deviating from 20 °C (293 K) and the working fluid temperature deviating from 15 °C (288 K). It should be noted that vapor pressure $p_{\rm vapor,B}$ is a function of working fluid temperature θ_B . The value 17.04 hPa represents the vapor pressure of water at 15 °C.

In order to determine the transported volume flow when saturated air is pumped down, Dalton's law is applied to pumping speed as well:

$$S_{A,\text{sat}} = S_{A,\text{dry}} \left(1 + \frac{p_{\text{vapor,A}}}{p_A - p_{\text{vapor,A}}} \right). \tag{7.5}$$

Here, vapor pressure $p_{\text{vapor.A}}$ is a function of inlet temperature ϑ_{A} .

In practice, the change in pumping speed differs slightly from these idealized equations. Therefore, extensive series of measurements were conducted to find empirical equations. Figure 7.17 shows, for example, the influence of workingfluid temperature for two-stage vacuum pumps.

Thus, for compressing dry air and using water as working fluid, the pumping speed of a liquid ring vacuum pump

$$S_{A} = S_{K} \lambda. \tag{7.6}$$

The required λ -value can be acquired from the diagram (Figure 7.17), for known working fluid temperature and inlet pressure.

If the liquid ring vacuum pump operates close to the vapor pressure of the working fluid, cavitation might occur in the pump. In this case, vapor bubbles develop in the liquid ring that abruptly collapse in the compression phase and thus produce considerable pressure and sound waves. As a result, components may be damaged, particularly impellers and guide disks. The line referred to as cavitation limit in Figure 7.17 provides guiding values for inlet pressure, which should be considered as minimum tolerable values for a given working-fluid temperature under cavitation-free operation.

Additional parameters that should be considered when dimensioning a liquid ring vacuum pump for complex applications are gas species, type of working fluid, deviations in compression pressure, and possible liquid carried along in the gas flow.

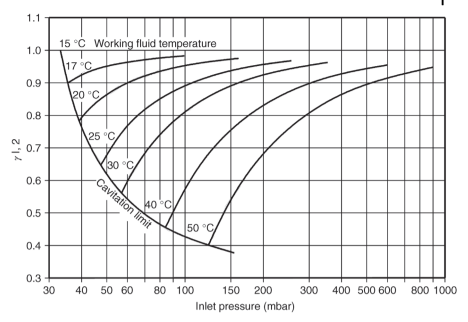


Figure 7.17 Influencing factor λ on the pumping capacity of a two-stage liquid ring vacuum pump versus inlet pressure at selected temperature levels of working fluid.

While traveling through the liquid ring vacuum pump, the temperature of the working fluid rises. The heat flow, which has to be removed by the working fluid, is mainly determined by compression heat and condensation heat; and the compression heat corresponds nearly to the power consumption of the vacuum pump. The condensation heat is important when the inlet flow contains vapor. The condensation effect is particularly eminent for low inlet pressures and high inlet temperatures.

The thermal balance is based on an equilibrium of heat fluxes into and out of the system. It allows calculating the temperature of the compressed gas and the working fluid at the outlet of the pressure joint.

7.3.1.3 **Designs**

The different types of liquid ring vacuum pumps feature different designs and are supplemented by application-oriented material variants. High operational reliability is common to all types, a result of the noncontact operation of impeller and housing in the compression chamber.

Liquid ring vacuum pumps are available in both single- and two-stage designs (Figure 7.18).

Whether single-stage vacuum pumps are applicable at inlet pressures down to 120 mbar (hPa) or even 33 mbar (hPa), is determined by the design of the impeller and the guide disks. Single-stage pumps that accommodate self-acting valves in the guide disks, in addition to the pressure outlet, can be used down to 33 mbar (hPa), just as two-stage pumps.

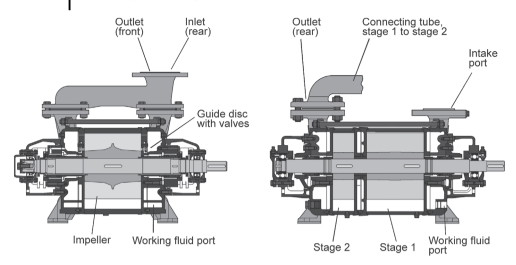


Figure 7.18 Single- and two-stage liquid ring vacuum pumps.

The advantages of two-stage vacuum pumps are a lower susceptance to failure and higher pumping speed at low inlet pressure, in particular, when pumping saturated gas mixtures and for high working-fluid temperatures.

For low and medium-high pumping speeds, single-stage liquid ring vacuum pumps are available as compact units in a block assembly (Figure 7.19). Here, the impeller is mounted directly onto the motor shaft. The bearing of the motor shaft thus additionally carries the pump's impeller.

The lowest achievable inlet pressure of a liquid ring vacuum pump depends on the vapor pressure of the working fluid. If lower inlet pressure is desired, a liquid ring vacuum pump can be combined with a gas jet pump (Section 9.3)

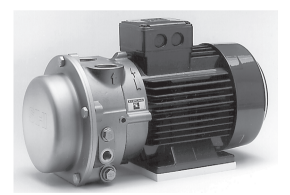


Figure 7.19 Motor unit vacuum pump with modern sheet-metal design (Sterling SIHI: LEM 26). Inlet and pressure ports are located at the top; the working fluid port is arranged laterally.

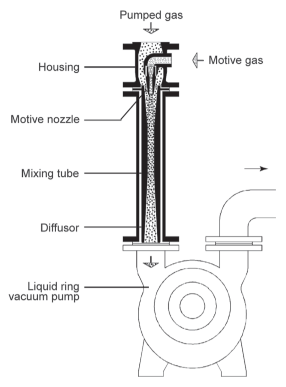


Figure 7.20 Liquid ring vacuum pump combined with a gas jet pump.

(Figure 7.20), boosting the transported gas to the appropriate inlet pressure of the liquid ring vacuum pump.

Gas jet vacuum pumps (Section 9.3) use the ejector principle. A driving gas (motive gas) relaxes in a motive nozzle and accelerates whereby the pumped gas is sucked in. In a diffuser, the kinetic energy of the mixture gas flow converts to static pressure energy due to higher pressure.

Figure 7.21 shows plots of pumping speeds for a liquid ring vacuum pump with and without gas jet.

Obviously, a combination of a liquid ring vacuum pump and a gas jet pump is capable of producing lower inlet pressures. At high inlet pressures, however, the volume flow is lower than in a stand-alone liquid ring vacuum pump. Therefore, the gas jet is initially bypassed in evacuating processes and subsequently connected at lower pressure.

Gas jet pumps provide ideal cavitation protection for liquid ring vacuum pumps because the driving gas is a gas species, which is noncondensing at operating temperatures. Even if the inlet amount of the gas jet approaches zero or the inlet of the gas jet is closed, the liquid ring vacuum pump operates outside the unsafe cavitation range.

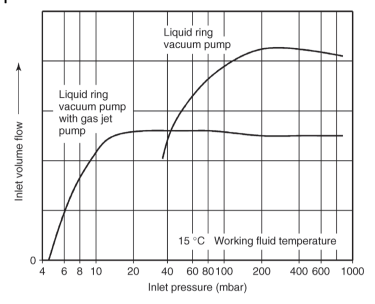


Figure 7.21 Pumping speeds (inlet volume flow) of a liquid ring vacuum pump with and without a gas jet pump.

7.3.1.4 Pump Units with Liquid Ring Vacuum Pumps

Figure 7.22 shows a connection diagram of a vacuum system with a liquid ring vacuum pump operating in circulating liquid mode. Typical suction pressures range from 50 mbar (hPa) to 500 mbar (hPa).

A downstream condenser is favorable when vapor-saturated gases are pumped. Thereby, pumping speed increases, that is, smaller vacuum pumps are sufficient. The developing condensate can be removed via the gas flow in the liquid ring vacuum pump (Figure 7.23).

For ultimate pressures between 5 mbar (hPa) and 50 mbar (hPa), a combination of a liquid ring vacuum pump and a gas jet provides an economical, robust solution. The gas jet compresses the pumping gas to inlet pressure of the liquid ring vacuum pump. The carrier gas is supplied by the separator or the highpressure line.

Combinations of Roots pumps (Section 7.4.3) and liquid ring vacuum pumps provide even lower suction pressures and higher pumping speeds.

In the simplest arrangement, a preceding Roots pump is connected to the inlet of the gas jet, adding to the structure in Figure 7.24.

However, Roots pumps can also be connected in a multistage series, directly preceding the liquid ring vacuum pump. This is appropriate in applications where a gas jet should be avoided. Then, the liquid ring vacuum pump can be reduced in size, as it does not need to additionally transport the motive gas of the gas jet.

Vacuum systems with Roots pumps yield ultimate pressures far below 1 mbar (hPa).

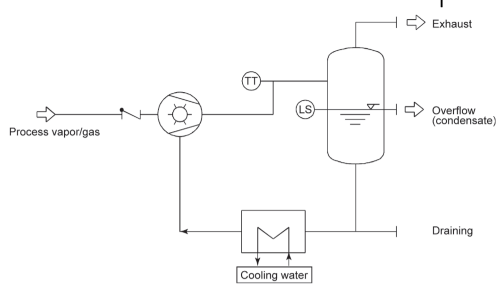


Figure 7.22 Vacuum system with a closed working-fluid cycle, equipped with monitoring equipment for maximum tolerable compression temperature (TT) and liquid level (LS) in the separator.

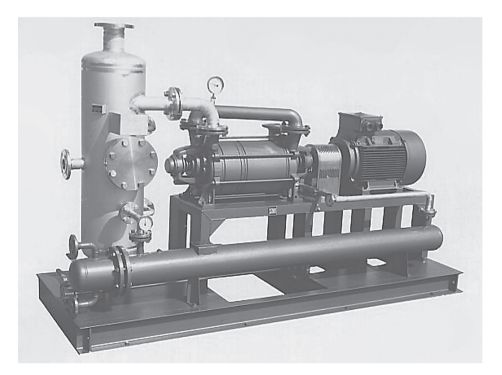


Figure 7.23 Vacuum system including a liquid ring vacuum pump (Sterling SIHI: LPH 65327) with a liquid separator and heat exchanger. The heat exchanger is mounted at the lowest part of the system to prevent gas pockets.

Figure 7.24 Vacuum system with a liquid ring vacuum pump and gas jet pump: working-fluid level (LS) and compression temperature (TT) are monitored.

Cooling water

7.3.1.5 Suggestions for Economical Operation

Liquid ring vacuum pumps are robust units. They are very tolerant towards pollutants in the pumping gas. If installation and operating mode are within specifications, maintenance of the pump is hardly ever necessary, even after long periods of operation.

The working fluid can be connected in one of the following modes:

- The connection diagram in Figure 7.24 shows circulating liquid mode, particularly common to the chemical and pharmaceutical industries. Here, other working fluids than water are often used, for example, oil, alkaline solutions, and acids. Tightly sealed working-fluid circulation is mandatory if corrosive, wastewater polluting, or toxic media are pumped. However, this setup is also used to recover condensate. The liquid flowing back from the vacuum pump's separator is re-cooled in a heat exchanger.
- The simplest operating mode is fresh-fluid mode. Here, the working fluid is not fed back from the separator to the vacuum pump but disposed off on the high-pressure side. Fresh-fluid mode requires a sufficient supply of working fluid (water). In practice, it is usually found only in applications with small-sized liquid ring vacuum pumps.
- Combined fluid mode is the most common type of operating arrangement. Here, a part of the fluid from the separator is fed back to serve as working

fluid. Fresh fluid, for example, water, from a fresh-fluid supply is used to supplements this flow. The inlet temperature of the working fluid into the pump corresponds to the mixture temperature of the two partial flows. It can be adjusted by regulating the amount of added fresh fluid.

Real operating conditions of a liquid ring vacuum pump often differ from basic design data. For overdimensioned vacuum pumps, in particular, adjusting pumping speed, that is, reducing pumping speed, is advisable. For this, the following approaches are suggested:

- Adjusting working-fluid temperature: The temperature of the working fluid is the simplest controllable parameter and the most effective way of controlling the volume flow of a liquid ring vacuum pump. In circulating and combined fluid mode, raising/lowering the pump's working fluid inlet temperature allows reducing/increasing pumping speed, respectively. If the working fluid is water, the temperature should be held above 10 °C to prevent icing inside the pump.
- Speed control: A transmission or frequency converter serves to adjust rotational speed. Frequency converters have the advantage that the working point can be adjusted instantly after a change in operating conditions. However, it must be considered that the speed of liquid ring vacuum pumps is restricted to a maximum of 30% deviation from nominal velocity, that is, the speed cannot be adjusted throughout the total velocity range. At too low speeds, the liquid ring does not form evenly, and the pump runs unsteadily. Too high velocities overstress components and can lead to impeller fracture;
- Bypass control: A portion of the compressed gas, or atmospheric air, is recirculated to the suction channel. This prevents unacceptably low inlet pressures.

Regulating pumping speed by throttling the inlet should be avoided at all times due to the risk of cavitation. The same applies to throttling on the outlet as this overstresses components and may lead to a destroyed impeller.

For cavitation-free operation, liquid ring vacuum pumps always require a certain amount of noncondensing gas in the flow. This is particularly important when the inlet flow contains a high proportion of vapor. Many pumps are equipped with a special cavitation-preventing connection providing a channel for feeding inert gas directly to the impeller cells. Thus, cavitation is avoided and, at the same time, the lowest tolerable inlet pressure is guaranteed.

When the pump is started, the fluid level should approximately reach the shaft. Thus, the complete output is obtained instantaneously and overload of the driving motor is prevented.

Liquid ring machines do not act as ignition sources when explosive mixtures are pumped.

7.3.2

Sliding Vane Rotary Pumps

Sliding vane rotary pumps are the most used vacuum pumps. The sliding vane rotary pump was developed between 1904 and 1910. A nobleman named Prince

Rupprecht gave the basic idea for this kind of operating principle as early as 1657. The capsule pump, developed by Gaede in 1909, is considered to be the origin of the sliding vane rotary pump [33]. Oil-lubricated sliding vane rotary pumps are economically relevant since applications in food vacuum packaging. Largest growth developed in the 1970s and 1980s. At the same time, printing and general packaging industry prospered, demanding many dry-running sliding vane rotary pumps.

7.3.2.1 Operating Principle and Design

The operating principle of a sliding vane rotary pump is defined in ISO 3529/2 (1981):

A sliding vane rotary pump is a rotary positive displacement pump in which an eccentrically placed rotor is turning tangentially to the fixed surface of the stator (housing). Two or more vanes sliding in slots of the rotor (usually radial) and rubbing on the internal wall of the stator, divide the stator chamber into several parts of varying volume.

Figure 7.25 shows the general design and the operating principle of a sliding vane rotary pump.

The sliding vane rotary pump has a housing, cylindrical on the inside, in which an eccentric, slotted rotor revolves. In the slots, vanes glide that move outward due to centrifugal forces, and slide along the casing wall. This divides the space between rotor and housing into chambers. The pumped gas enters through the inlet into the sickle-shaped suction (expansion) chamber that increases its volume during rotation. In Figure 7.25a, a newly formed sickle-shaped suction chamber reaches the inlet. Ideally, the chamber would start with zero volume. As the rotor turns the volume of the expansion chamber increases and causes suction. After reaching maximum suction volume, the volume starts to decrease while rotation continues. Figure 7.25d indicates the position of the vanes at the time the maximum expansion volume is obtained. At this point, the expansion chamber and the inlet are no longer connected, that is, the vane has passed by the inlet.

While the rotation continues, the sucked in gas is compressed until the compression chamber opens to the outlet (Figure 7.25e). The internal compression is determined by the geometrical position of the outlet. The compressed gas is driven out through the outlet until, ideally, the chamber volume reaches zero volume, Figure 7.25f. As the rotor continues its rotation, the considered working chamber sweeps by the inlet anew and the next compression cycle starts.

As the suction volume, ideally, starts with zero volume, there is no dead space. In practice, however, gaps are present due to manufacturing tolerances, and to allow for thermal expansion. Gaps appear radially as clearances between the rotor and the housing near the TDC, axially to both sides of the

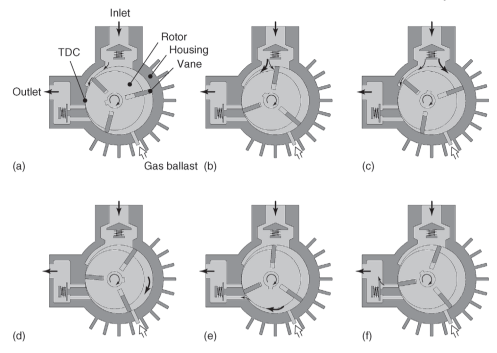


Figure 7.25 Design and operating principle of sliding vane rotary pumps.

rotor, as well as around the vanes in the slots and here, axially as well, toward the case lids.

7.3.2.2 Dry-Running Sliding Vane Rotary Pumps

Dry-running sliding vane rotary pumps reach ultimate pressures of 80-200 hPa. Generally, they feature a fairly large number of vanes. Figure 7.26 shows an example of a dry runner. Here, the vanes not only seal the chambers but also serve to fulfill the delicate task of lubricating the running surface. The materials for vanes are often chosen from composite materials containing mainly graphite.

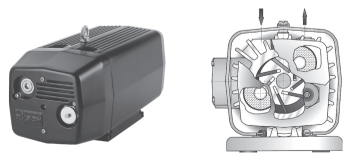


Figure 7.26 Dry sliding vane rotary pump.

The exact composition and the manufacturing are strictly confidential, since the materials selection for vanes as well as the running surfaces (case) is crucial.

Today's materials allow maintenance rates in between vane replacements of approximately 2000 to above 10000 operating hours. Usually, designs feature seven vanes in order to reduce the load on each individual vane. Dry-running sliding vane rotary pumps are also referred to as multiple vane pumps or lamella pumps (the vanes are also called lamella).

Often, a filter is used on the inlet side to prevent pump damage due to contaminants. During operation, the vanes of a dry runner are subject to wear. Therefore, a filter is also included in the outlet side. It serves solely to protect the environment against particles produced by vane abrasion and is pre-integrated into the pump. Dry-running sliding vane rotary pumps are available for volume flow rates of $1.5-500 \,\mathrm{m}^3 \,\mathrm{h}^{-1}$.

7.3.2.3 Oil-Lubricated Sliding Vane Rotary Pumps

In order to generate low ultimate pressures, an oil film is used that improves gap sealing. Bore holes and channels maintain an overall supply with oil between case lids and the fore-parts of rotor and vanes, and help the vanes to push forward a small wave of oil continuously between the suction and compression volumes. This leads to sufficient sealing between the compression and suction volumes. An oil-lubricated sliding vane rotary pump usually has a spring-loaded valve in the outlet channel of the compression chamber. This does not open until the internal compression pressure rises far enough above ambient pressure in front of the valve to overcome the spring resistance of the valve. At low gas throughput, that is, when inlet pressure is low, the oil pushed ahead by the vanes, just before ejection, nearly completely fills the exhaust channel. This quickly leads to a strong increase in pressure inside the chamber. Now the vanes attempt to compress incompressible oil until the oil flows out through the outlet valves. This pressure produces a blow of oil, and well into the 1970s, this was considered a signal that ultimate pressure had been attained, and thus, as a kind of gauge. Today, this oil blow is regarded as nuisance, and thus, a small amount of gas is artificially fed to the pump. As long as gas is still pumped, the valve opens smoothly due to the gas cushion, and oil blow is prevented. However, the obtainable ultimate pressure is affected negatively by this measure. For a variety of applications, single-stage sliding vane rotary pumps are available with ultimate pressures in the range of 0.1–20 hPa.

The working fluid, usually a high-grade mineral oil, seals and lubricates the pump. A relatively large amount of working fluid is injected to the compression volume. Therefore, this fluid also takes up a considerable proportion of the compression heat. By this, the temperature in the pump can be adjusted between 70 °C and 90 °C. In practice, this temperature range has proven to be a fair compromise between a long service life of the oil and avoiding condensation inside the pump. In addition to high temperature in the pump, gas ballast (see also Section 7.6.1) also prevents condensation in vapor-generating processes. The gas ballast opens into the compression chamber, just after this is no longer in contact with the inlet; illustrated in Figure 7.25d. Ambient air is used as gas ballast in the sealed compression chamber. The mixture of ambient air and humid process gas cannot condensate inside the pump. However, in dry processes, the service life of the operating fluid is higher as long as the pump is cooled sufficiently.

Additional functions of the working fluid include corrosion protection and cleaning. Particularly in contaminated applications, the oil cleans all moving parts. Critical parts are mainly the vanes in the slits and the valves. The valves increase the efficiency of the sliding vane rotary pump and are required for obtaining sufficient ultimate pressure. Due to the cleaning effect of the oil, valves in oil-lubricated pumps hardly ever fail. Inside the compression chamber, the working fluid is exposed to the pumped media. In many cases, not only clean, dry air is pumped, but also, for example, fat and dust can enter the pump and then be absorbed by the operating fluid. Sufficiently large oil reservoirs as well as oil filters provide working fluids that perform well for long periods of time without harming the pump. A specially designed oil-mist separator divides the oil from the pumped gas. It requires particular technology to induce the oil mist to form larger oil droplets for recycling (see Section 7.8.4).

Figure 7.27 shows the design of an oil-lubricated sliding vane rotary pump. Individual functions are spatially separated. The gas enters the pump case through a nonreturn valve. The check valve separates the recipient from the

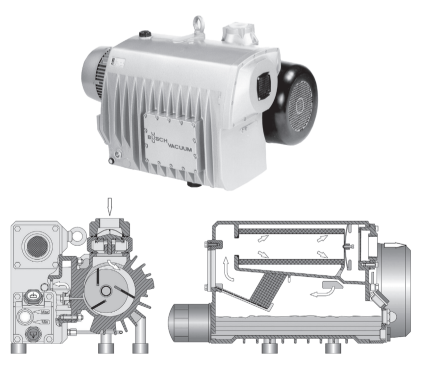


Figure 7.27 Oil-lubricated sliding vane rotary pump.

pump when it stands still. It prevents backflow of gas or working fluid. After gas transport and compression, the gas leaves the pump case through outlet valves and enters into the oil separator housing. This serves as an oil reservoir and separates the working fluid from the gas. In a first chamber, as transport velocity drops, larger drops of oil leave the flow. Then, the finest droplets agglomerate to large drops in specially designed oil-mist separators, which are still subject to continuous improvement. The oil filter cleans the working fluid and feeds it back to the pump. Depending on the pump's power, the available surface area, a simple fan, an oil-air heat exchanger, or an oil-water heat exchanger is required for re-cooling the working fluid.

Oil-lubricated sliding vane rotary pumps are available for $3-1600 \,\mathrm{m}^3\mathrm{h}^{-1}$. They are widespread for producing low and medium vacuum because they are robust, reliable, economically priced, and provide long service lives.

7.3.2.4 Once-Through Lubricated Sliding Vane Rotary Pumps

In applications where the operating fluid degrades or is destroyed due to the particular process, the usually common closed-loop oil cycle is not applicable. In this case, pumps operate with so-called once-through lubrication. Here, the lubricating oil is not fed back to the pump, as in circular lubrication, but is only used once and then drained. Thus, fresh oil continuously lubricates the pump. Figure 7.28 describes the principle.

In this arrangement, the oil lubricates the suction chambers and creates a protective film between functional surfaces in order to prevent corrosion and for rinsing out condensed vapor from the pump. For transport of aggressive/corrosive vapors, the operating reliability of this type of pump is high. The permanent slope from the inlet to the outlet and continuous rinsing of condensate prevents damage to the vanes due to accumulating liquid as well as a possible fall off in

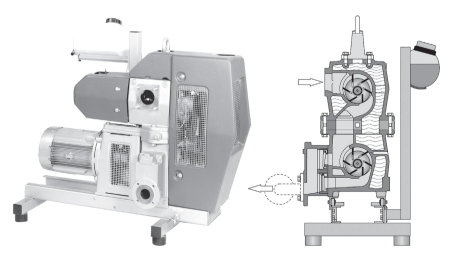


Figure 7.28 Once-through lubricated sliding vane rotary pump in two-stage piggyback design.

vacuum quality due to lubricating oil that is polluted with condensed vapor. Because the oil stays inside the pump only for a very limited amount of time, cheap oil is applicable. An oil pump feeds the oil. Pumps, available between 160 and 780 m³h⁻¹, consume approximately 84–188 cm³h⁻¹ of oil.

7.3.2.5 Operating Behavior and Recommendations

Individual descriptions are given here because the operating behaviors of the three described sliding vane rotary pumps - dry-running, oil-lubricated, and once-through lubricated - differ considerably.

Dry-Running Sliding Vane Rotary Pumps

Regular inspections of possible vane wear in a dry-running sliding vane rotary pump are necessary in order to prevent vane fracturing during operation, and thus, pump failure. Scheduled filter cleaning avoids possible power drops. Intervals for this kind of maintenance work depend heavily on the particular application. In dusty environments, more frequent filter cleaning is necessary than when clean and dry gas is pumped. Additionally, vane wear increases when pressure differences are high and dust is aspirated. If condensable vapors appear in the pump, second running with flush gas can prevent corrosion inside the pump. Appropriate measures are required to prevent downtime corrosion if the pump is turned off for longer periods. In typical applications, sliding vane rotary pumps suck in ambient air and require pressure of only approximately 400 hPa. The two largest market shares in this context are packaging and printing industry. In both cases, pumps lift and transport goods and corresponding packaging material. Here, pumps are usually placed locally, in the immediate vicinity of the application. Pump failure would interrupt operation of the complete system; therefore, high reliability and simple maintenance of the pump is mandatory. This type of pump meets these requirements and, at the same time, is cheap. In spite of simple maintenance, disadvantages are high maintenance rates and frequent inspections of vane wear (check every 1000-2000 operating hours).

Oil-Lubricated Sliding Vane Rotary Pumps

During operation, oil-lubricated sliding vane rotary pumps depend on the working fluid, usually a high-grade mineral oil. In clean applications where the oil is not polluted, this type of pump operates for long periods without requiring maintenance work. However, oil level and oil draining components should be checked every 2000 operating hours. If other substances apart from dry and clean gas enter the pump, the contaminated oil pollutes oil filters and oil mist separators. In this case, oil filter, oil-mist separator, and the oil require replacement, which is easy and swift. Generally, the oil is sensitive to aging that also leads to a change in operating conditions and reduces lubricating properties. Then, an oil change is necessary. For pump operation, the oil fulfills an additional important task: cold oil has higher viscosity than oil at operating temperature. Therefore, the drive must deliver a higher driving torque when the pump starts in the cold state. Depending on the temperature, this can mean that the drive may not be able to deliver enough torque and start the pump. A workaround would be to use thinner oil. General use of thinner oil, however, is not recommended due to volatile components, service life, and with respect to oil mist separation. Thus, manufacturers offer different oil viscosities for different ambient temperatures.

If condensable vapors, particularly water, are sucked in, oil and condensate join and emulsify. This leads to an immediate increase of the obtainable ultimate pressure and also lower pumping speed, and eventually, to lowered operating behavior of the working fluid, particularly lubrication and corrosion protection. Gaede prevented this by inventing the gas ballast. Gas ballast lowers the obtainable ultimate pressure. If condensable vapors are present, the vapor pressure of the liquid defines the operational limit. In the application, the pressure in the vessel cannot drop lower, and thus, a pump with gas ballast is always superior to a pump without any gas ballast. The amount of gas ballast is adjusted according to experimental values for each individual application. Section 7.6.1 explains the function of a gas ballast.

In contrast to dry-running sliding vane rotary pumps, vanes in oil-lubricated pumps experience practically no wear at all. In addition, the running surface on the internal side of the case remains essentially unaltered, which leads to very high service life of the pump. Epoxy bind fiberglass covered by carbon fiber is employed as vane material, but also aluminum. Oil-lubricated sliding vane rotary pumps operate in applications with pressures of approximately 1–800 hPa. This wide range is differentiated into an area <200 hPa, out of reach for dry-running pumps, and into an area >200 hPa for applications in which humidity and pollution impede use of other types of pumps. The largest market share for this type of pump is vacuum packaging of food.

Small oil-lubricated sliding vane rotary pumps are available in single- and twostage designs. For two-stage pumps, series connections produce better ultimate pressures, and, in conjunction with this, still good pumping speeds even at low pressures. In two-stage pumps, the stages are connected by attaching the outlet of the first stage directly to the inlet of the second stage without any interconnected valve. Thus, the second stage serves, so to speak, as a backing pump to the first stage. A two-stage sliding vane rotary pump can therefore be thought of as a pump unit (see also Sections 7.4.3.6 and 7.4.3.7).

Once-Through Lubricated Sliding Vane Rotary Pumps

In applications of once-through lubricated sliding vane rotary pumps, the working fluid, usually mineral oil, protects the pump against aggressive pumping media or large amounts of condensate. The application must therefore consider pump protection at all times. For example, after shutdown, no corrosive substances should remain inside the pump. Usually, rinsing or second running is recommendable. The largest market share for this type of pump is in the chemical and pharmaceutical industries. Two-stage designs with good pumping speeds at operating pressure are common, which distribute the compression heat among two stages. The amount of oil put to use only lubricates and protects the

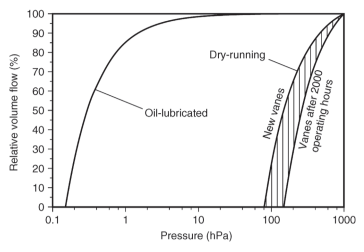


Figure 7.29 Typical pumping capacities versus inlet pressure of oil-lubricated and dry-running sliding vane rotary pumps, also considering operating hours for the latter.

surfaces. It cannot take up the compression heat as in circular lubrication systems. Temperatures are comparable to dry runners. Too high temperatures harm the oil. Therefore, the compression heat is divided between the two stages. Many applications that rely on limited process temperatures benefit from this setup.

7.3.2.6 Characteristic Curves and Ratings

Figure 7.29 shows characteristic curves of a dry-running and an oil-lubricated sliding vane rotary pump. The characteristic curve of a two-stage once-through lubricated sliding vane rotary pump depends on the internal staging, explained in more detail for a Roots pump in Section 7.4.3.6. Here, the relative volume flow with respect to inlet pressure is plotted versus absolute pressure. Standards for measurement of such characteristic curves are defined in *PNEUROP* [1] and DIN 28400.³⁾ It should be noted that, for dry-running sliding vane rotary pumps, the characteristic curve changes noticeably due to wear on the vanes. If the operating point is close to ultimate pressure with worn vanes, the difference of volume flow is considerable. This has to be taken into account when dimensioning the pump.

Oil-lubricated sliding vane rotary pumps, in contrast, do not suffer any changes due to wear. If the working fluid is not exposed to condensate, the characteristic curve remains unaltered. The pump practically provides full pumping speed throughout a wide pressure range. Only below approximately 2 hPa, the volume flow drops due to internal leakage.

For a pump with internal leakage, known ultimate pressure, and nominal pumping speed that operates at low vacuum, the characteristic curve follows a

³⁾ Translator's note: corresponds to ISO 3529.

simple relationship:

$$S(p) = S_1 \left(1 - \frac{\frac{p_1}{p} - 1}{\frac{p_1}{p_{\text{odd}}} - 1} \right), \tag{7.7}$$

with pumping speed S_1 at pressure p_1 (usually, $p_1 = p_n$), and the ultimate pressure $p_{\rm ult}$ of the pump. This simple relationship corresponds very well to measured characteristics under low-vacuum conditions and applies to other types of displacement pumps as well. Real characteristics of oil-lubricated sliding vane rotary pumps usually show a slight deviation because injected oil and therefore internal sealing increases when the working point approaches ultimate pressure. In this sense, available pumps behave quite differently. High-quality pumps operate very close to the theoretical curve whereas less sophisticated pumps show considerably flatter characteristics. For example, certain pumps feature the same nominal volume flow and ultimate pressure but produce different results in practical applications. For users, the time required to obtain a certain pressure in a vessel is of interest. This pump-down time is determined mainly by the integral of the volume-flow–pressure curve.

7.3.3 Rotary Plunger Pumps

7.3.3.1 Principle of Operation and Technical Design

Figures 7.30 and 7.31 show the principle of operation and the working cycle of a rotary plunger pump.

The piston of a rotary plunger pump runs in a piston bearing. As the rotor moves eccentrically and along the cylindrical case walls in noncontact motion,

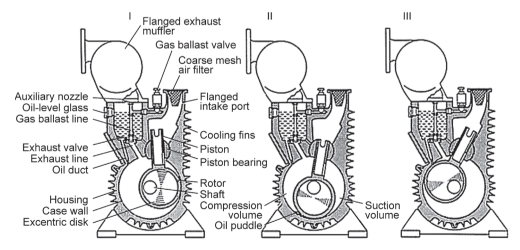


Figure 7.30 Pumping phases I-III of a rotary plunger pump.

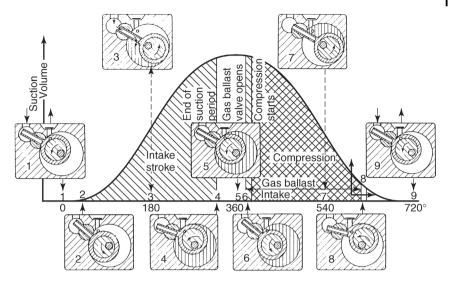


Figure 7.31 Working cycle of a rotary plunger pump. Position 1: top dead center. Position 2: the groove at the inlet channel of the piston is enabled, begin of intake period. Position 3: bottom dead center. The groove at the inlet channel is free. The pumped gas (arrows) travels into the suction chamber (shaded area). Position 4: the groove at the inlet channel is

sealed by lamellae. End of intake period. Position 5: top dead center, maximum volume for the suction chamber. Position 6: just before the compression period starts, the face of the rotor opens the gas ballast port. Begin of gas ballast intake. Position 7: gas ballast port is completely open. Position 8: end of gas ballast intake. Position 9: end of pumping period.

the piston reciprocates. The pumped-down gas enters into the sickle-shaped suction chamber through the inlet and a peripheral aperture in the piston. The suction chamber expands during rotation and causes the suction effect.

When the rotor reaches the TDC (just after position III, Figure 7.30, and position 5, Figure 7.31), the suction chamber has reached its maximum volume. At the same time, the piston is far enough to the top for the peripheral aperture to seal. As rotation continues, a new suction chamber forms that begins with zero volume (therefore, there is no dead space). The pumped down gas from the previous cycle enters the compression chamber, which shrinks continuously and thus compresses the gas until the outlet valve opens and ejects the gas at approximately 105 kPa (1050 mbar (hPa)).

The oil filling is adjusted to a level so that, for low-pressure operation, the outlet valve is covered with oil whereas it is free of oil at full throughput. Largely, the ejected air is separated mechanically from entrained oil in an exhaust chamber that succeeds the valve chamber. Boreholes and channels guarantee that sufficient amounts of oil are available at the fore parts of the rotor and the piston. Additionally, an oil puddle develops between the suction chamber and the compression chamber, which follows the rotation thus sealing and separating both chambers from another.

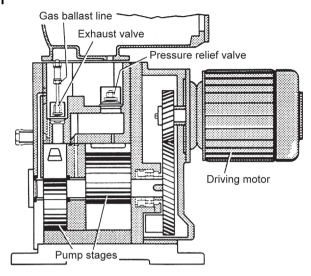


Figure 7.32 Section of a rotary plunger pump (schematic).

At low gas throughput and just before the TDC is reached, the oil puddle between the suction and compression chambers fills the valve channel. This guarantees that there is no dead space in the compression chamber as well. However, as the rotation continues, the valve opens instantaneously due to the oil blow-out, which causes clattering of valves in rotary plunger pumps as well, thus indicating ultimate pressure operation. The rotating frequency of the eccentric rotor lies in the interval of 400–600 min⁻¹, and thus, is far below common rotating speeds of oil-sealed sliding vane rotary pumps (Section 7.3.2.3).

Rotary plunger pumps also pump oil in addition to gas, which makes oil circulation an important issue, just as the gas path. In Figures 7.32 and 7.33, a rotary plunger pump is illustrated in a cross section and in an additional view. The cross section shows a pump with two rotary pistons, with lengths in the ratio 2:1 and an offset of 180°, driven by a motor via a gear pair. A characteristic feature in this design is that the air of the motor fan travels over the pump and thus simultaneously cools the pump. Fins, added to the housing of larger pumps, increase the surface available for heat transmission. The larger the size of a rotary plunger pump, the less sufficient is air cooling. Therefore, all larger rotary plunger pumps are water-cooled.

The two rotors serve as pump stages. In single-stage pumps, both stages are placed in parallel. In two-stage pumps, the exhaust of the first stage is connected to the inlet of the second stage. Due to the difference in length of the rotors, here necessary for mechanical balancing, the pump stages feature different pumping speeds. The longer rotor is connected to serve as high-vacuum stage. At high inlet pressures, this causes overpressure between the two stages, which is let off to the exhaust through a pressure relief valve. (Due to the 2:1 ratio,



Figure 7.33 Rotary plunger pump with a nominal pumping capacity of 250 m³ h⁻¹. Total length: 1190 mm, total height: 740 m. With courtesy of Oerlikon Leybold Vacuum GmbH.

overpressure develops between the stages for an inlet pressure range from ambient pressure down to 50 kPa (500 mbar).)

Figure 7.34 shows a schematic illustration of the gas path in single- and twostage pumps. The oil path in a two-stage pump is designed to first feed oil that was exposed to air at ambient pressure to the fore-vacuum stage for degassing.

Series and parallel connections of pump stages are provided by incorporating flow channels in the pump. This is why single- and two-stage designs look alike in the considered examples. A single-stage pump of the same size, however, has an about 50% higher pumping speed. The only difference in pumps for higher

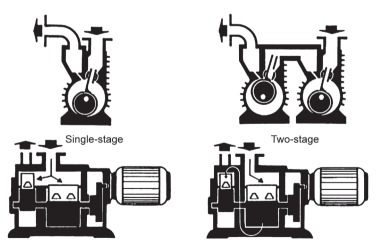


Figure 7.34 Gas paths through single- and two-stage rotary plunger pumps.

pumping speeds is a longer rotor. Pump diameters remain the same. Steps from one pump to the next higher are designed in such a way that the smaller rotor of the next larger pump is equal to the larger rotor in the next smaller pump. This reduces the variety of necessary rotor types.

7.3.3.2 A Comparison of Sliding Vane Rotary Pumps and Rotary Plunger Pumps

Sliding vane rotary pumps and rotary plunger pumps produce approximately the same ultimate pressures so that this criterion is irrelevant for selection purposes.

The motion of the rotary plunger pump is constrained. The rotor runs along the case walls in noncontact operation, and thus, experiences no wear. However, wear can occur in the bearings between the eccentric guidance and the rotor as well as in the guiding surfaces of the piston. Mass balancing improves smoothness of running considerably, which usually is relatively poor.

The motion of the sliding vane rotary pump is not constrained. Its vanes press against the internal wall of the housing due to spring power or centrifugal force (or both). A sliding vane rotary pump operates in nonpositive motion, that is, is actuated by forces. Sliding vane rotary pumps run extremely smoothly due to very well balanced masses. Modern sliding vane rotary pumps use bearings, separated from the rest of the lubricating cycle, so that the vanes are the only parts in the pump chamber that suffer minor wear. This is due to continuously improved materials. By separating the challenging bearing lubrication from the uncritical lubrication in the pump chamber, the latter can be adjusted to fit miscellaneous industrial applications, that is, a large variety of different lubricants is utilizable.

Thus, in large-sized chemical and other industrial installations, large sliding vane rotary pumps with nominal pumping speeds of several 100 m³h⁻¹ are applied in addition to rotary plunger pumps.

7.3.4

Trochoidal Pumps

Adequate valve control provided, any conventional piston engine can be used as a vacuum pump. Therefore, transferring the principle of a planetary piston engine to a vacuum pump suggested itself. This idea led to a vacuum pump known as the trochoidal⁴⁾ pump.

Due to economical reasons associated with relatively high manufacturing costs of the ellipse and cardioid (heart-shaped) geometry of the piston and the case, this type of pump did not prevail. However, it shall be discussed here in brief because the general principle is very interesting.

4) A cycloid, epicycloid, or hypocycloid is a curve described by a point on the periphery of a circle as the circle rolls, without sliding, along a straight line, along the outside of a static circle, or along the inside of a static circle. Trochoids, epitrochoids, or hypotrochoides are analogous curves represented by a point on the plane of the rolling circle, and inside or outside the periphery (Brieskorn, E. and Körner, H. (1981) Ebene algebraische Kurven, Birkhäuser, Basel).

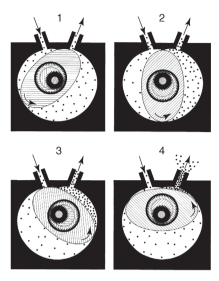


Figure 7.35 Pump phases in a trochoidal pump.

Figure 7.35 shows the operating principle and pump phases of a trochoidal pump.

An elliptical piston moves around a shaft eccentrically in a toothed case. The case is in continuous, sealing point-contact with the piston. The suction chamber expands from this point (starting at zero volume), and thus, sucks in the gas to be pumped down from the recipient. Both tips of the piston form the other sealing points, which move along the housing in noncontact motion. Oil is fed to the pump chamber for sealing. Gas continues to enter the suction chamber as long as there is a connection between the suction chamber and the inlet. After the tip of the piston has passed by the inlet, the gas is compressed in the sealed chamber until it reaches the pressure required to open the exhaust valve (approximately 1100 mbar (hPa)). The outlet valve is connected in such a way that oil overlay is guaranteed.

Considering the variety of available trochoid geometries for housing and rotor, only the rotor is suitable for a hypotrochoid because this is the only geometry meeting the requirements of a vacuum pump (static spot between inlet and outlet, suction volume starting with zero volume). Calculation of the hypotrochoid used in this pump is described in [34].

Considering motion aspects, a trochoidal pump can be understood as a combination of a sliding vane rotary pump and a rotary plunger pump. The pistons have positive fit whereas the seal, similar to vanes in a sliding vane rotary pump, is actuated by force, pressed to the piston, and slides along the same. In trochoidal pumps, short and large inlet cross sections provide stable pumping speeds down to low inlet pressures.

As all mass motion is rotary, a trochoidal pump, as a sliding vane rotary pump, can be perfectly balanced. Only the small amounts of transported oil represent minor unbalanced masses. Therefore, a trochoidal pump can operate at equally high speeds as a sliding vane rotary pump; and this, although, when perceived as a rotary plunger pump, it actually should only allow half to a third of the speed of sliding vane rotary pumps. This leads to pump units small in size considering the nominal pumping speed.

7.3.5

Scroll Pumps

The scroll pump is a recent development featuring a dry positive displacement pump that compresses against atmospheric pressure. The operating principle of a scroll pump goes back to an idea patented by the Frenchman Leon Creux in 1905. Technology at the time, however, did not allow producing sufficiently strong and precise components. Around 1970, this principle of compression was picked anew in the United States, and utilized in compressors for refrigeration. In response to the energy crisis, scroll compressors became very popular due to their efficiency, which is considerably higher than that of piston compressors.

In the mid-1990s, the Japanese company *Iwata* built the first scroll vacuum pumps. Other suppliers followed, and since, the use of scroll pumps continues to spread rapidly.

7.3.5.1 Principle of Compression

A scroll compressor features two nested Archimedean screws. Each screw contains an equidistant spiral wall, built onto a circular base plate. If the screws, arranged at a 180° offset, are joined, the walls sectionally enclose sickle-shaped volumes. Centrally-symmetric oscillation (orbiting) of one hull against the other causes the volume to move along the spirals (Figure 7.36). Thus, positive displacement occurs. The inlet can be arranged at the outside; an axial borehole provides the outlet.

Following the historical development, Sawada [35] provided the initial theoretical basics. During gas transport, all types of flow occur, from turbulent to molecular, and thus, calculation is difficult. The authors considered an individual transported gas volume V, assumed isothermal conditions, and calculated the transported amount of gas iteratively along a complete spiral passage. The conclusions were compared with results obtained in experimental investigations with a real pump. It turned out that, for velocities below nominal speed (1500 min⁻¹), the produced ultimate pressure is lower than predicted. This effect was believed to be caused by heat released from the air-cooled shells, leading to













Figure 7.36 Gas path in a scroll spiral. Courtesy of E. Reuschling, Edwards GmbH.

reduced clearance to the inner component, and thus, creating increased backflow. In the experiments with a swift transition to low speeds, the authors proved that theoretical values could be obtained.

This is an important aspect in dimensioning scroll pumps. The power output reaches an optimum after an even distribution of operating temperature is obtained throughout the pump.

7.3.5.2 **Design**

The basic principle of scroll pumps does not require any inlet or outlet valves. Usually, CNC machines are used to manufacture the aluminum scroll shells. The height of the spiral walls (in the centimeter range), the clearance between them, and the rotational speed determine the throughput of the pump. The length of the spiral path defines the ultimate pressure (due to dynamic sealing). Coating the spiral walls with Teflon reduces friction and improves chemical resistance. The faces, that is, the top edges of the spiral walls, require sealing to the opposite surfaces. For this, a flexibly mounted seal with rectangular cross section, usually made of plastic compound, is applied. The backside of the oscillating shell is attached to an eccentric gearing.

Considering possible lock-up of the shells, tolerances in the gearing and in shell manufacturing are critical. Sufficiently accurate tolerances provided (<1/10 mm), sealing agent is not necessary. The pump self-seals dynamically at low backflow. Tolerances must be matched to the thermal equilibrium in the pump chamber. A directly coupled electric motor drives the gearing, that is, the pump can run at speeds above 1000 min⁻¹.

The gas path influences the ultimate pressure. Therefore, early developments already featured multistage scrolls. A compact design incorporates the moving component in scroll configuration at the front and back as well as symmetrical sealing with scroll walls (Figure 7.37). Here, the gas initially passes through the spiral at the front, and subsequently through the one at the back, which improves ultimate pressure considerably. However, this arrangement has certain disadvantages; for example, moving components require complex guidance and a hermetical variant is not available (see below).

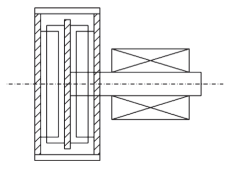


Figure 7.37 Diagram of a two-stage scroll pump with drive unit on the right-hand side. Courtesy of Edwards GmbH, Wiesbaden.

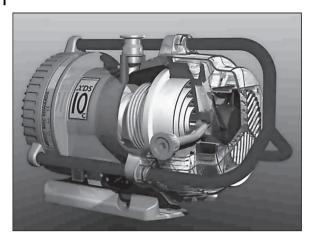


Figure 7.38 Section of a scroll pump. Far left: disk armature motor. Center: eccentric drive, fully encapsulated by stainless-steel bellows. Right: pump chamber. Courtesy of *Edwards GmbH*, Wiesbaden.

In new developments, the inlet of the scroll spiral is divided into several channels that merge on their way to the inside. This increases compression and thus allows ultimate vacua in the area of 0.1 Pa, even in single-stage pumps.

Recent advances use a principle patented in nuclear technology in which the complete eccentric drive including all ball bearings lies beneath stainless-steel bellows connected to the moving shell (Figure 7.38). Theoretically, the bellows follow the tumbling motion with infinite service life. Thus, pumped media cannot damage the bearings and lubricating grease does not contaminate the vacuum.

7.3.5.3 Applications and Advantages

Oil-free scroll pumps can replace oil-sealed sliding vane rotary pumps in the medium-high pumping-speed range $(5-30\,\mathrm{m}^3\mathrm{h}^{-1})$. Ultimate pressures reach several Pa, slightly above ultimate pressures of two-stage sliding vane rotary pumps. However, scroll pumps increasingly serve as components of oil-free high-vacuum pumping systems, particularly in combinations with turbomolecular pumps with *Holweck* or *Gaede* stages where the ultimate pressure of scroll pumps is sufficient. In addition, use of scroll pumps is spreading in many areas of research and analytics. Typical users are, for instance, manufacturers of analytical instruments, electron microscopes, and inert glass glove boxes.

In the area of very low pump speeds, diaphragm pumps are more economical because the basic price of a scroll pump is more or less independent of its size. Scroll pumps are not available for high pump speeds since large eccentric gearings would be too expensive.

Main advantages of the scroll pump include oil-free operation (depending on separation and sealing between bearings and evacuated volume), that is, high

purity of the vacuum, no disposal costs for working fluids (pump oil), primarily only one moving part, no valves, hardly any wearing parts, and high efficiency due to high speed at low required driving energy.

7.4 Twin-Spool Rotating Positive Displacement Pumps

7.4.1

Screw-Type Pumps

Screw-type vacuum pumps are dry-running positive displacement pumps for inlet pressures in the low and medium vacuum range. The operating principle relies on two noncontacting, screw-type rotors, rotating in opposite direction. They form several chambers along their axes, moving continuously from the inlet to the outlet.

Screw-type vacuum pumps can be differentiated according to rotor types (symmetric and asymmetric tooth profile), or according to the principle of gas compression in the working chamber. An end plate on the high-pressure side with a defined, controllable outlet aperture determines the compression of conventional screw-type vacuum pumps. Modern screw-type vacuum pumps, in contrast, control the compression of the pumped medium with a gradient in tooth profile and open gas outlet from the screw-type rotors.

Symmetric as well as asymmetric tooth profiles were initially developed for screw-type compressors. The earliest patents published can be traced back to the year 1917. In 1990, industrial use of screw-type rotors as dry-running vacuum pumps began. As opposed to worm compressors, used industrially since circa 1950, most commonly containing oil as working fluid for sealing and heat removal, screw-type vacuum pumps only in very rare cases require working fluid in the working chamber.

For this reason, and due to the wide operating range, that is, a nominal pumping speed of 70-2500 m³ h⁻¹ at inlet pressures from ambient pressure to 0.001 mbar, this design has prevailed in chemical and pharmaceutical processing technology as well as in semiconductor production and other branches of industry. Many applications also feature scroll pumps as fore pumps combined with Roots pumps or other types of vacuum pumps for the medium and high vacuum range because of their low inlet pressures.

7.4.1.1 Operating Principle and Technical Design

Screw-type vacuum pumps are twin-spool machines with screw-type rotors, arranged in parallel. During operation, the rotors mesh in noncontact motion and rotate in opposite directions. One or more chambers on the inlet side enclose the transported medium and deliver it to the high-pressure side.

Figure 7.39 shows a conventional screw-type vacuum pump with evaporative cooling on the pressure side of the working chamber's outer housing.

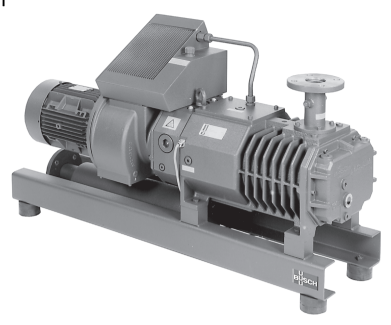


Figure 7.39 Screw-type vacuum pump with nominal pumping capacity 250 m³ h⁻¹; sealed circuit cooling with a air-cooled condenser on the pump.

Evaporative cooling with closed cooling circuit and direct flow-through cooling with fluid are the most common methods for limiting operating temperatures.

Figure 7.40a—d shows the transport process in this screw-type vacuum pump with an asymmetric tooth profile of constant gradient, in which compression takes place against an end plate with a controllable aperture on the pressure side. The inlet lies radially above the vacuum pump between the screw-type rotors. Beginning here, the first chamber fills with transported medium. The

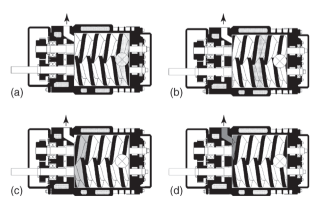


Figure 7.40 Transport in a standard screw-type vacuum pump: (a) intake, (b) transport, (c) compression, (d) ejection.

images do not reveal the total axial length of the individual chambers because they extend, below the visible rotor contours, to an angle of wrap of 720° around the considered tooth profile.

Therefore, throughout the length of the screw rotors, 2.5 working chambers develop around both rotors transporting medium continuously from the suction to the high-pressure side. In the shown asymmetric tooth profile, the chambers form in a way that the height of the clearance along the complete width of the sealing gaps is nearly constant. 2.5 working chambers provide that gas backflow remains sufficiently low. Compression at the controllable high-pressure aperture leads to a local rise in temperature and pulsating exhaust gas flow.

Cooling liquid is necessary to carry off the compression heat produced during operation. Large-sized pumps increasingly use cooling gas, referred to as gas ballast, fed to the working chamber close to the high-pressure end thus limiting exhaust temperatures.

After operation starts, high exhaust temperature causes the screw-type rotors to expand until a thermal equilibrium establishes. As in other dry-running vacuum pumps (e.g., Roots pumps, claw pumps), cooled parts of the housing more or less maintain their original dimensions due to lower temperature and thus reduce radial clearance. Stable inlet pressure develops only after thermal equilibrium has established.

Figure 7.41 shows an overview of today's screw-type rotors with the top-level distinction between screw-type rotors with symmetrical and asymmetrical tooth profiles.

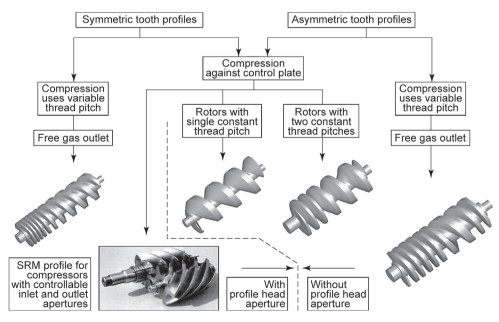


Figure 7.41 Classification of screw-type rotors into symmetric and asymmetric tooth profiles.

The right-hand side shows the different types of screw rotors with asymmetric tooth profile. Of these, the two on the left, in the center of the image, maintain nearly constant clearance height throughout the total width of the chamber's sealing gaps. Furthermore, they produce chamber geometries, which each extend around both rotors. Next to the screw rotor with constant thread pitch, the figure shows a profile with two stages of constant pitch each. In this rotor, compression starts further to the inlet side of the working chamber. Nevertheless, a limiting end plate with controllable aperture remains indispensable. To the far right, the image shows a novel rotor geometry with a continuous gradient in thread pitch. This rotor geometry also features nearly constant gap height around the pumping chambers. Additionally, final compression against an end plate with controllable aperture is unnecessary. Figure 7.42 shows a simplified representation of this screw-type vacuum pump. When introduced, this rotor geometry fulfilled a number of requirements in problematical applications, for example, chemical engineering. Noteworthy characteristics are free drainage of liquid components in transported media, comparably high internal compression for reduced power consumption, as well as low rotor velocities for simple and variable shaft sealing.

A conventional asymmetric SRM profile (Svenska Rotor Maskiner) [36] is shown at the bottom of Figure 7.41. These rotors, used with different kinds of asymmetric and symmetric tooth profiles in screw-type compressors, consist of one rotor with only tooth tips, and a second rotor, with only tooth roots. They often have different tooth numbers and thus different driving torques and rotational speeds [37]. As opposed to all other types of rotor geometries, a controllable inlet aperture spanning the complete rotor length is necessary due to the lower angle of wrap of each tooth profile. A characteristic common to all screwtype vacuum pumps with variable thread pitch and symmetric tooth profile

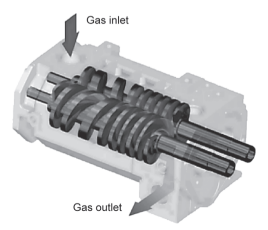


Figure 7.42 Simplified diagram of a modern screw-type vacuum pump with variable thread pitch and asymmetric tooth profile.

(Figure 7.41, left) is a profile head aperture. Minimizing the profile head aperture, and thus, the leakage backflow, is a prime goal of rotor design [38].

A peculiarity of rotors with the described symmetrical tooth profile is that the individual chambers do not extend around both rotors. The gearing teeth separate the chambers of the two rotors. A large number of transporting chambers develop, which require high velocities due to the inevitable profile head apertures [39,40]. Rotors with symmetric tooth profiles are advantageous in conducting compression heat in large-sized vacuum pumps. For example, if pumping speeds exceed 300 m³h⁻¹, the comparably large rotor surface and the leakage rates through the profile head apertures, combined with direct internal liquid cooling of the rotors, effectively lower exhaust temperatures.

Today, different cooling methods are used for large-sized systems in order to limit exhaust temperatures in addition to fluid case cooling:

- gas cooling by way of feeding foreign gas into the working chamber (gas ballast);
- gas cooling by means of recirculating a proportion of the cooled exhaust gas into the working chamber;
- gas cooling using direct internal cooling of screw rotors, for example, with oil.

It appears that direct internal screw rotor cooling is establishing in the modern screw-type vacuum pump market. Here, the mass flow of cooling medium carries off compression heat from the working chamber without requiring any additional sealing elements.

Optimizing the course of compression by increasing the rate of change in the thread pitch, that is, by raising the internal compression, is limited. Power consumption would then decrease significantly only at low inlet pressures. For high inlet pressures, in contrast, it would increase rapidly and thus require unreasonably high driving power.

Figure 7.43 shows typical characteristic curves of dry-running screw-type vacuum pumps. For a comparison of various types of screw rotors, a set of differently sized pumps is considered:

- pump 1 with 160 m³h⁻¹ pumping speed, asymmetric tooth profile, constant thread pitch and compression against an end plate;
- pump 2 with 250 m³h⁻¹ pumping speed, asymmetric tooth profile, variable thread pitch without compression against an end plate;
- pump 3 with 400 m³h⁻¹ pumping speed, symmetric tooth profile, variable thread pitch with comparably high compression ratio and without compression against an end plate.

Figure 7.44 shows the corresponding power-consumption curves. A further increase of compression ratio than in the 400 size would reduce power consumption even further for low inlet pressure but would lead to considerably increased power consumption at high inlet pressure. The power consumption at high inlet pressure would then demand an unreasonably large driving motor.

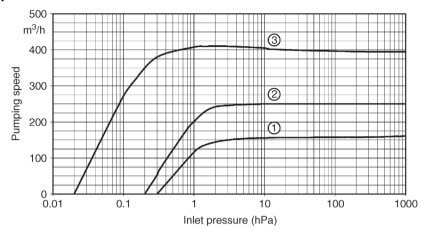


Figure 7.43 Pumping capacity curves for three different types of screw-type vacuum pumps.

All considered screw-type vacuum pumps require sealing between the rotors and the case at both the suction and high-pressure sides. In recent years, cantilevered rotors are also used in screw rotor machines. These screw-type vacuum pumps have bearings on the high-pressure side, and therefore, require sealing only on the gearbox side. The advantage of this type of arrangement is that sealing is dispensable on the suction side. However, there is a risk of rotors touching each other during operation due to a lack of parallelism.

Independent of rotor setup, touching as well as nontouching seals are used. Furthermore, different combinations of both sealing variants are common on individual seal surfaces.

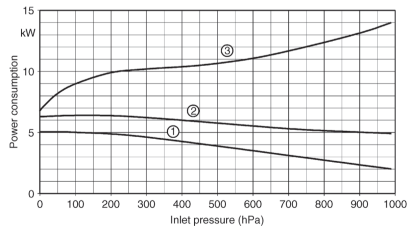


Figure 7.44 Power consumption curves for the screw-type vacuum pumps considered in Figure 7.43.

Often, noncontacting conventional seals are applied, frequently supplemented with seal gas in labyrinth seals made of several piston rings. In contrast, noncontact seals include:

- rotary shaft seals or combinations of rotary shaft seals;
- liquid-lubricated floating ring seals using the gear oil as cooling and lubricating fluid on the high-pressure side, and with their own oil reservoir on the suction side that also serves to lubricate the bearings on that side;
- gas-lubricated floating ring seals used only for sealing at the high-pressure side to the gearbox. They require continuous supply of foreign gas. Due to high costs, however, they are only used for special media to be pumped, or, if exceptional requirements exist in terms of closeness.

7.4.1.2 Heat Behavior and Technical Notes

An important characteristic of dry-running screw-type vacuum pumps are the gas temperatures developing under polytropic compression of transported media. Neglecting friction (bearings, seals, and synchronizing gears), the temperature increase can be estimated by balancing the powers entering and leaving the pump during operation:

$$P_{\text{drive}} = \dot{Q}_{\text{gas}} + \dot{Q}_{\text{L.C.}} + \dot{Q}_{\text{A.C.}}. \tag{7.8}$$

Here, P_{drive} is the driving power of the pump, Q_{gas} the heat flow fed to the transported gas, $\dot{Q}_{\text{L.C.}}$ the heat flow carried off by the cooling liquid, and $\dot{Q}_{\text{A.C.}}$ the heat flow carried off by air cooling. Using $\dot{Q}_{gas} = \Delta T \, \dot{m} \, c_p$, the temperature increase ΔT of the transported medium can be calculated using mass flow \dot{m} and the thermal capacity c_n :

$$\Delta T = \frac{P_{\text{drive}} - \dot{Q}_{\text{L.C.}} - \dot{Q}_{\text{A.C.}}}{\dot{m} c_p}.$$
(7.9)

The hottest areas in the working chamber of a conventional dry-running screw-type vacuum pump for pumping speeds above 500 m³h⁻¹ reach temperature peaks up to 500 °C. In spite of the polytropic compression of the transported medium, a theoretical relationship for the temperature increase due to a pressure change can be derived on the strength of past evidence using Poisson's equation (see Section 4.2.2) for an adiabatic change of state of an ideal gas. This attempt requires considering operating states in which only a comparatively low proportion of the compression heat is transferred to the environment. Then,

$$T_2 = T_1 \left(\frac{p_2}{p_1}\right)^{\kappa - 1\kappa}. (7.10)$$

The rise in outlet temperature T_2 compared to the inlet temperature T_1 is determined by the pressure ratio of outlet pressure p_2 and inlet pressure p_1 as well as by the isentropic exponent κ of the transported medium.

For example, theoretically, a gas inlet temperature of 30 °C, inlet pressure of 50 hPa, outlet pressure of 1013 hPa, and an isentropic exponent κ of 1.4 yield an

outlet temperature of 443 °C. In fact, similarly high temperatures can be measured in individual areas just behind the working chamber of pumps with high pumping speeds and reduced cooling.

During pump operation, outlet temperature T_2 drops as the mass flow \dot{m} of the transported medium increases. On the other hand, the driving power required for compression rises for higher inlet pressures. Analogous to the calculation of compressor power, presented in Section 7.6.2.4, the compression work in a dry-running screw-type vacuum pump approaches zero at low inlet pressure. The driving power is consumed almost exclusively for frictional losses in bearings, seals, and the synchronizing gear. At zero delivery, temperatures measured in the exhaust of a pump start approximately at ambient temperature, show a relative maximum between 100 hPa and 400 hPa, and ultimately drop at higher inlet pressures as a consequence of the increasing mass flow *m*.

Considering pumps with higher pumping speeds, the ratio of cooled housing surface area and pumping speed drops (see Section 7.6.2.3). Gas cooling with supplied foreign gas or by recycling cooled exhaust gas is limited in its utilization for certain applications. Therefore, modern screw-type vacuum pumps with pumping speeds above 160 m³h⁻¹ also use direct internal liquid cooling of the screw rotors. This method employs the large surface area of the screw rotors for an additional heat removal and lowers the maximum temperatures appearing inside the pump on surfaces and in the gas.

Limiting exhaust temperature using very cold cooling water and high cooling water flow is problematic if the transported medium contains vapor. This vapor can condense at cold housing surfaces and develop corrosive condensates, or can destroy sealing gaps and thus the pump rapidly due to cavitation.

Although damage caused by corrosive liquids also occurs during operation, it is particularly likely to develop when the pump stands still. In applications where corrosive liquids are anticipated, a drying cycle with sealed inlet shut-off device is initiated for screw-type vacuum pumps immediately after working cycles. Usually, the speed is lowered and the increased temperature in the working chamber is utilized to support the drying process.

As opposed to liquid ring vacuum pumps or sliding vane rotary pumps that require a certain minimum velocity to build up the working chambers, such restrictions do not exist for operating speed-controlled screw-type vacuum pumps. For practical use of speed control, it should be noted that reducing speed does not only lead to a drop in flow output, but also, the ultimate vacuum has higher values. Figure 7.45 shows the effects of different velocities for a conventional screw-type vacuum pump.

The use of speed control to meet temperature limits, for example, for pumping explosive gases or vapors [41], usually lowers pumping speed and limits the inlet pressure of a pump. In addition, flame arresters at the inlet and outlet, often required for delivering explosive media, reduce inlet pressures and pumping speeds.

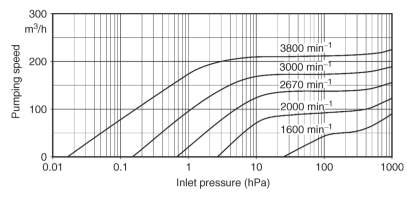


Figure 7.45 Pumping capacity curves of a screw-type vacuum pump for selected drive speeds.

7.4.2 Claw Pumps

Claw pumps, named after their characteristic rotor shapes, are twin-shaft rotating positive displacement pumps with a basic design similar to Roots pumps. They feature dry compression, that is, without any sealing liquids such as oil and water, and operate in noncontact motion between rotors as well as between the rotors and the stage walls. Sealing gaps provide internal sealing of the compression chambers and between the suction and high-pressure sides. As opposed to Roots pumps (Section 7.4.3), claw pumps operate on internal compression. This means that the volume of the positive displacement chamber decreases prior to opening the outlet and compresses the enclosed gas before it is ejected. This process is controlled inherently by the rotor profiles, and thus, no separate outlet valves are required.

Northey was the first company to introduce claw pumps [42]. This type of claw pump, operating with two identical rotors with one rotating claw each, is therefore also referred to as the *Northey* principle. This principle still is utilized, most commonly in two- or multistage vacuum pumps.

In the early 1970s, technical literature first described claw pumps containing two different types of rotors, a main and a control rotor [43]. Their special rotor geometry guarantees that the gas is released completely, and thus, lifts a restriction of the *Northey* type. Here, these pumps will also be referred to as *exact* rotating claw profiles. Theoretically, the ejecting compression chamber shrinks to zero volume. In practice, the exact profiles were initially used in single- or two-stage compressors.

Exact rotating claw profiles, as opposed to the *Northey* type, allow multiclaw rotor geometries. The most common profiles have one or two teeth per rotor because efficiency drops when more teeth are used.

7.4.2.1 Compression Principle

Northey Type

Figure 7.46 shows the working principle of claw pumps of the *Northey* type. Inlet and outlet apertures lie at the faces of the pump stage, that is, in the axial direction. Rotation of the rotors is synchronized, inverse, and noncontacting. Sealing gaps of approximately $1/10 \, \mathrm{mm}$ (when cold) provide sealing between the rotors and towards the stage walls in the axial and radial directions. Indentations in the rotors seal or open the inlet and outlet depending on the rotors' position and thus control suction, compression, and ejection. The rotating teeth divide the pump volume into chambers that aspirate, compress, and discharge the gas by increasing or decreasing their volume.

A full pump cycle spans two revolutions (Figure 7.46). In Pos. 1, the inlet aperture (1) just opens, the suction chamber expands as shown in Pos. 2 until, in Pos. 3, maximum volume is established and the inlet shuts. The rotors mesh as shown in Pos. 4. At this point, inlet and outlet are sealed and there is no sealing gap between the rotors; thus, aspirated gas fills the entire compression chamber. Geometrically, Pos. 5 corresponds to Pos. 1. The gas sucked in during the

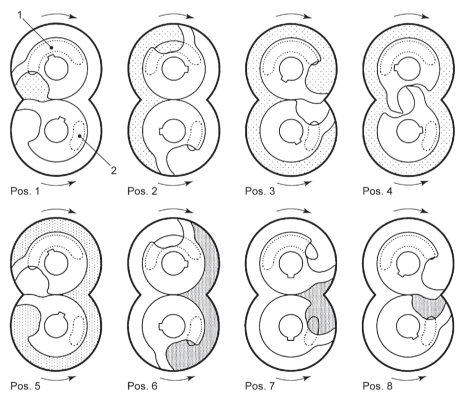


Figure 7.46 Operating principle of a tooth-type rotary pump of the Northey type.

previous revolution is now captured inside the compression chamber. The volume of this chamber reduces thus compressing the gas (internal compression) until, in Pos. 6, the outlet aperture (2) opens. Pos. 7 shows continuing ejection and, in Pos. 8, ejection terminates and the outlet closes. The rotors mesh and return to Pos. 1.

In Pos. 7, obviously, the compressed gas is not ejected entirely. A dead space (or dead volume) remains inside the pump, amounting to approximately 7% of the pump-chamber volume. The enclosed gas, which is under outlet/atmospheric pressure, expands anew while the rotors mesh. Now, the inlet is sealed, as opposed to the condition in other pumps where the dead space can re-expand on the suction side and thus limit the obtainable ultimate pressure. Therefore, expanding of the dead space slightly increases the pressure of the sucked in gas in the total volume, and thus cuts efficiency, but only indirectly influences the producible ultimate pressure. However, the pressure in the suction chamber in Pos. 1, spanning 7% of the total volume as well, is higher than inlet pressure, which leads to a backflow towards the suction side. Roughly estimated, this adds up to 7% of 7% of the pump volume, thus, approximately 0.5%.

Exact Profile

In exact rotating claw profiles, as opposed to the *Northey* profile, the compressed gas is ejected entirely. Theoretically, the dead space is of zero volume and a permanent sealing gap exists between the rotors, even at times when they mesh. Therefore, designing a radial inlet (6) as shown in Figure 7.47 is possible and flow restriction in the inlet can be reduced. The figure shows the most common profile: a double-claw-type design with a main rotor (1) and a control rotor (2). The rotors divide the pump volume into four chambers: the inlet chamber (3), two transport chambers (4), and the compression chamber (5). The control rotor opens and closes the outlet.

Aspiration begins in Pos. 1 (Figure 7.47). The suction chamber increases its volume (Pos. 2) until, in Pos. 3, the inlet is separated from the suction chamber, initially, by the main rotor, and subsequently, by the control rotor. The rotors divide the suction chamber into two transport chambers. Ideally, the pressure in the transport chambers is equal to the inlet pressure. At constant pressure, the enclosed gas is simply moved, that is, pushed, until the transport chambers unite and form the compression chamber (Pos. 4). The gas is compressed via Pos. 5; the outlet (7) opens in Pos. 6, and ejection starts. Pos. 7 shows the continuing ejection, in Pos. 8, the gas is ejected entirely and the outlet closes. The rotors mesh and return to their initial position.

Obviously, in Pos. 4, and during the succeeding compression cycle in Pos. 8, the proportion of slightly precompressed gas located between the rotating teeth is separated from the rest. This proportion amounts to approximately 10% of the chamber volume, is not ejected, and returns to the inlet side as dead space. The effectively transported volume thus decreases by approximately 12% because the pressure in the dead space is not equal to the outlet pressure but only approximately 16% above inlet pressure. Therefore, in spite of reduced

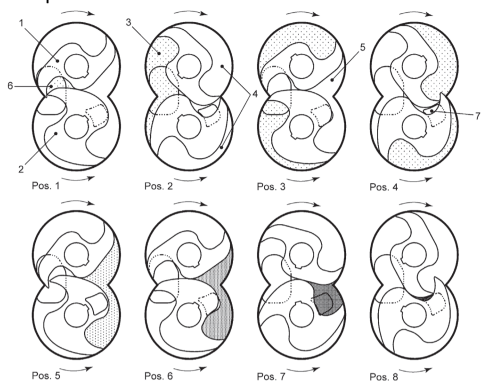


Figure 7.47 Operating principle of a tooth-type rotary pump with true rotor profile.

efficiency, there is no negative impact on ultimate pressure because the dead space, in an exact profile, does not deliver outlet-side air backward but only inlet-side air. Apart from other aspects, the number of teeth in multiclaw-type designs is limited by the fact that the dead space appears correspondingly more often during one revolution.

Comparison

In the investigated exact profile, the sucked-in gas remains inside the pump stage for about half a revolution before it is ejected. However, in the *Northey* profile, it remains in the stage for a period of two revolutions. Assuming equally large gaps, the exact profile suffers less from disturbing internal backflow through these gaps. In the exact profile, the rotors smoothly control the ejection of the compressed gas down to zero volume whereas the *Northey* profile causes dynamic overcompressions due to abruptly terminated, incomplete ejection [44]. Thus, for single-stage pumps, the specific power consumption (shaft power per unit volume flow) of an exact-profile claw pump is approximately 30% lower than for the *Northey* type [45].

On principle, the *Northey* type is less sensitive to extra liquid or solid transport than an exact rotating claw profile because the *Northey* type, at all times, maintains

a minimum volume between the rotors, even during ejection. Thus, in spite of the energetic disadvantage, it has prevailed in applications where this issue is critical.

7.4.2.2 Comparison with Roots Pumps

The basic design of a claw pump and a Roots pump (Section 7.4.3) is very similar. The main distinction with respect to the compression process of the two working principles is that claw pumps operate on internal compression. In order to be able to compress gas internally prior to ejection, the outlet must be sealed temporarily. Thus, the outlet is arranged axially on the sidewalls, and is opened and closed by indentations in the control rotor (see, e.g., Figure 7.47). In contrast, gas passage through a Roots pump (see Section 7.4.3) is much more streamlined.

The aperture angle of the outlet determines the value of the internal compression. For example, lengthening the outlet towards the direction of rotation reduces internal compression. In single-stage claw pumps, internal compression is usually adjusted to values between 1:1.4 and 1:2.

Figure 7.48 shows typical values of the relative pumping speed and specific power with respect to inlet pressure of a two-claw pump as well as a three-clawtype Roots pump, operating against atmospheric pressure (i.e., not designed as a fore or booster pump). Both pumps have a nominal pumping speed of approximately 250 m³h⁻¹. The pressure range of the Roots pump is limited to approximately 400 hPa in order to avoid overheating at lower pressures. The pumping-speed curve of the claw pump is flatter. Due to the internal compression, it can produce a deeper vacuum. Energetically, the Roots pump is very advantageous when the pressure difference is low. Inlet pressures above 500-600 hPa lead to undesired overcompression in the claw pump due to the internal compression. For high pressure differences, the claw-type rotary pump has a lower specific power consumption, as can be anticipated from the pV diagrams with/without internal compression. Overall, claw pumps feature the energetically most favorable working principle throughout a wide range of low vacuum.

7.4.2.3 Multistage Claw Pumps and Pump Combinations

A single-stage c law pump (Figure 7.49) is capable of producing an ultimate pressure of 30-50 hPa. If an application requires lower working pressure, twoor multistage pumps are utilized. Pumps with up to four stages are available, some of which are designed as combinations of claw and Roots pumps [46]. A particular variant is referred to as twin-flow design consisting of a vacuum stage and an independent pressure stage [47].

For multistage pumps, the internal compression in the individual stages is a parameter used in pump selection for a particular application. The steps, that is, ratios of sizes, between individual stages represent additional parameters used for the selection process. For example, if the low-pressure stage is larger than the high-pressure stage in a two-stage pump, power consumption at low inlet pressures is reduced. For high inlet pressures, however, overcompression would occur between the stages. This shows how important application details are for the selection process.

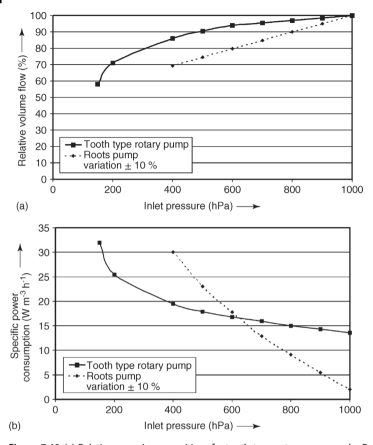


Figure 7.48 (a) Relative pumping capacities of a tooth-type rotary pump and a Roots pump. (b) Specific power consumptions of a tooth-type rotary pump and a Roots pump.

7.4.2.4 Speed Control

Centrifugal forces in claw pumps, as opposed those in sliding vane rotary pumps and liquid ring pumps, are not part of the working principle. Therefore, it appears appropriate to operate claw pumps at varying velocities in order to



Figure 7.49 Single-stage tooth-type rotary pump.

adopt them to the demands of particular applications. Obviously, this can be achieved by alternatively equipping the pump either with a two-pole (3000 rpm) or with a four-pole (1500 rpm) rotary-current asynchronous motor. However, the sealing gaps must be manufactured precisely in order to prevent excessive drop of the characteristic curves at low speeds.

If an application demands variable pumping speeds, speed-controlled claw pumps are used. Usually, a frequency converter controls the rotary-current asynchronous motor in such applications. Here, the speed can be adjusted from approximately 1000 to about 4500 rpm. Motor cooling determines the lower limit and the thermal load tolerable by the pump controls the upper limit.

7.4.2.5 Fields of Application

Single-stage claw pumps are used primarily in pneumatic transport, printing industry, vacuum networks, and central vacuum systems in hospitals.

Two-stage claw pumps are appropriate for delivering working pressures below 100 mbar.

Semiconductor industry and chemical/pharmaceutical processes use multistage claw pumps.

7.4.3

Roots Pumps

Roots pumps are based on a principle invented by the Englishman Isaiah Davies in 1848. 20 years later, the Americans Francis M. and Philander H. Roots took over the basic design, which subsequently was known as the *Roots blower* concept. Since, Roots pumps are used in technical applications (mainly as superchargers with compression ratios 1.5–2), and, with reversed operating mode, as gas meters (volume measurement). If a Roots blower operates as a Roots pump against atmospheric pressure, ultimate pressure is limited to approximately 15 kPa. It was not until circa 1954 that vacuum technology rediscovered Roots pumps [48].

7.4.3.1 Principle of Operation

Vacuum technology uses Roots pumps with two-lobe impellers. Figure 7.50 explains the basic working principle. Two eight-shaped impellers rotate in opposite direction inside a case. Forced coupling via a gear pair with equal tooth numbers guarantees that the impellers match during rotation but touch neither each other nor the case walls. The resulting clearance between both lobes as well as between the lobes and the case walls is kept as small as possible and determined by the size of the pump, the desired efficiency, and the expected operating conditions. The actually obtained clearance is the result of a compromise and amounts to about 0.1 mm.

For a simplified description of the operating principle, we will consider only the right-hand side of the pump in Figure 7.50. In the impeller positions I and II, the pump volume facing the recipient increases. In impeller position III, the sickle-shaped volume V_2 is separated from the inlet. As rotation continues, the volume opens towards the pressure side (fore-vacuum side) and gas at

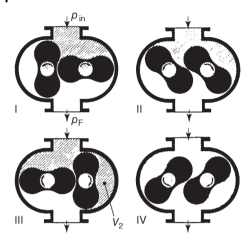


Figure 7.50 Pump phases I–IV in a Roots pump. Right impeller turns clockwise, and the left counterclockwise.

fore-vacuum pressure $p_{\rm F}$ flows into the previously sealed volume (impeller position IV). The gas inflow compresses the gas inside the volume and is ejected together with the gas previously transported from the suction side when the impeller rotates further. Thus, if loss is neglected, the transported gas volume is equal to the volume V_2 of the sickle-shaped chamber in position III. This volume develops twice during each revolution and the system contains two impellers. Therefore, the suction chamber volume (corresponding to the piston displacement in a displacement piston pump) of a Roots pump

$$V_{S} = 4V_{2}. (7.11)$$

At high rotating frequencies (e.g., $\nu = 3000\,\mathrm{min}^{-1} = 50\,\mathrm{Hz}$) high pumping speeds $S_{\mathrm{th}} = \nu V_{\mathrm{S}}$ can be produced with small pumps. Here, the impeller material determines the tolerable speed due to the developing centrifugal forces.

Pump operation inside the pump chamber is completely dry. Solely the bearings and gearwheels require oil lubricant. Piston rings, labyrinths, etc. on the shaft ducts seal the pump chamber from the bearing volumes and generally prevent oil from penetrating the dry pump chamber. This leads to absolute pressures below 0.1 Pa. The only sealing between the suction side and the forevacuum side is provided by tight clearances (dynamic sealing). Particularly in the low and medium vacuum range, compression leads to different warming of the impeller and the case. This, combined with the resistance of flow, limits the pressure difference that can be produced by the pump, which, therefore, requires particular measures (re-cooling, outlet cooling) if operation against atmospheric pressure is desired. Usually, a fore pump is required for operation.

7.4.3.2 Technical Setup

Figure 7.51 shows a simplified design for a Roots pump. The axes of the two impellers extend beyond the pump chamber. A gear pair with equal tooth

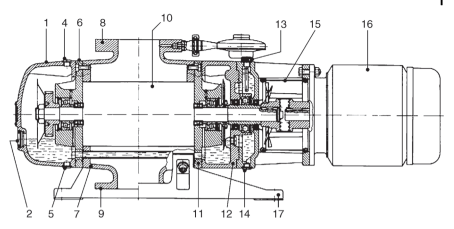


Figure 7.51 Longitudinal section of a Roots pump. 1: front cover; 2: oil-level glass; 4: oil filling screw; 5: oil drain plug; 6: bearing flange on the impeller side; 7: pump housing; 8: inlet flange; 9: outlet flange (connects to roughing pump); 10: driven impeller; 11: bearing flange

on the motor side; 12: intermediate flange; 13: oil filling screw with oil dip stick (shaft lead through); 14: oil drain plug (shaft lead through); 15: lantern with protecting cage; 16: driving motor; 17: foot. Horizontal shading: oil.

number, lying outside the actual pump chamber, couples the impellers. Rotating disks (oil-thrower lubrication) supply the gearwheels and bearings with oil from the oil reservoir. Piston rings, labyrinth seals, etc. on the shaft ducts seal the oil volumes from the pump chamber.

Larger pressure differences between the pump chamber and the oil chambers can cause oil to leak through the seals and into the pump chamber. Therefore, the oil volumes are connected either to the suction side or to the fore-vacuum side of the Roots pump in order to ease sealing. The entire inside volume of the Roots pump is tightly sealed against ambient pressure (usually atmospheric pressure). Pumps from series production easily remain below leak rates of $0.1 \, \mathrm{Pa} \, \ell \, \mathrm{s}^{-1}$. O-ring seals guarantee simple dismounting and assembly for cleaning and, if necessary, repairs.

For high rotating frequencies, vacuum-tight rotary transmission leadthroughs, exposed to atmospheric pressure on one side, are problematic and can cause malfunctions. They are designed as oil-covered chambers in which the oil reservoir simultaneously cools the rotary shaft seals. Furthermore, the sealing rings tend to run in into the shaft, which is at least troublesome for (usually necessary) replacement work. Therefore, modern Roots pump designs for medium vacuum feature a canned motor. Here, a thin tube made of nonmagnetic material, e.g., austenitic stainless steel, assures vacuum-tight separation of stator and rotor (Fig. 7.52). Thus, the rotor is under vacuum. Pumps operating under low-vacuum conditions must handle higher pressure differences. Here, and if special motors are used, e.g., explosion proof or with special voltages and frequencies, the disadvantages of rotary shaft sealing rings are tolerated, and the motor is coupled from the outside as a flange or conventional motor. If the rotary shaft



Figure 7.52 Partial section of a single-stage, hermetically sealed Roots pump RUVAC WH 7000 by *Oerlikon Leybold* Vacuum. Inlet flange size 320 mm, nominal pump speed

10.000 m³ h⁻¹, motor power 11 kW, 4200 rev min⁻¹. With courtesy of *Oerlikon Leybold Vacuum GmbH*.

sealing rings are replaced, a bush placed onto the shaft, sealed with O-ring seals, is removed as well. This, to a certain extent, prevents the run-in problem associated with the rotary shaft seal.

Many pump types are prepared to connect to a manometric switch (usually, a diaphragm pressure switch), which automatically starts the pump as soon as the inlet pressure reaches appropriate values.

7.4.3.3 Theoretical Basics

Particularly when considering Roots pumps, in-depth knowledge of the basic theoretical principles introduced here is of practical significance because it allows calculation of pumping-speed curves for a large variety of pump combinations. These can be highly complex and experimental simulations are often too expensive – alone due to the size of the systems [49,50].

7.4.3.4 Effective Gas Flow

A Roots pump delivers an effective gas flow $q_{pV,\rm eff}$, which is calculated from the theoretical gas flow $q_{pV,\rm th}=V_{\rm S}\,\nu\,p_{\rm in}$ of a lossless Roots pump diminished by the loss in gas flow $q_{pV,\rm loss,\rm gaps}=C(p_{\rm F}-p_{\rm in})$ caused by clearances (conductance C) or by the transition from position II to position III (Figure 7.50), just before position III is reached. Here, the wedge-shaped space between the impellers shrinks so rapidly that the gas molecules do not completely travel to the fore-vacuum side. As the rotation continues this amount of residual gas at higher pressure (higher than fore-vacuum pressure $p_{\rm F}$) expands back into the lower-pressure chamber on the suction side. Additional backflow occurs due to the fact that the impeller surfaces and their hollows, following the conditions of equilibrium, charge with gas, which is released when the pressure drops on the suction side. These adsorption and desorption effects become more and more crucial as the operating pressure of the Roots pump drops. The losses described here are summarized as $S_{\rm back}\,p_{\rm F}$. The effectively delivered gas flow

$$q_{pV,\text{eff}} = p_{\text{in}} S = V_S \nu p_{\text{in}} - C(p_F - p_{\text{in}}) - S_{\text{back}} p_F.$$
 (7.12)

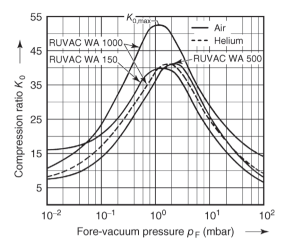


Figure 7.53 Measured compression ratios at zero delivery K_0 versus fore-vacuum pressure p_F (pressure at the outlet of the Roots pump) for differently sized Roots pumps.

7.4.3.5 Compression Ratio K₀ at Zero Delivery

The compression ratio at zero delivery $q_{pV,\text{eff}} = 0$ is measured by sealing the inlet with a blind flange. Rewriting Eq. (7.12) then yields the compression ratio at zero delivery:

$$\left(\frac{p_{\rm F}}{p_{\rm in}}\right)_0 = \frac{S_{\rm th} + C}{S_{\rm back} + C} = \frac{S_{\rm th}}{S_{\rm back} + C} + \frac{C}{C + S_{\rm back}} = K_0. \tag{7.13}$$

This value is one of the most important characteristic parameters of a Roots pump.

DIN 28426,5 part 2, and [9] provide details concerning measurement setup and method for obtaining K_0 with respect to fore-vacuum pressure p.

Figure 7.53 shows curves for the compression ratio K_0 of selected Roots pumps. As K_0 is usually above 10, and the second term in Eq. (7.13) is less than 1, it follows that, approximately,

$$K_0 = \frac{S_{\text{th}}}{S_{\text{back}} + C}.\tag{7.14}$$

In the high-pressure range ($p_F > 1.5 \text{ kPa}$), conductance C of the clearances is high (see Eq. (4.188) and Figure 4.49). Then, the disturbing backflow S_{back} is negligible compared to conductance C, and thus,

$$K_0 = \frac{S_{\text{th}}}{C}$$
 for $p_{\text{F}} > 1.5 \text{ kPa}$. (7.15)

Pure molecular flow occurs in the low-pressure range $p_{\rm F} < 10 \, {\rm Pa}$. This leads to conductance values that are small (Eq. (4.188) and Figure 4.49) compared to the

⁵⁾ Translator's note: corresponds to ISO 1607.

unwanted backflow, and thus.

$$K_0 = \frac{S_{\text{th}}}{S_{\text{back}}} \quad \text{for } p_{\text{F}} < 10 \text{ Pa.}$$
 (7.16)

For viscous flow, the conductance increases with rising pressure. This causes the compression ratio K_0 to drop at higher pressure. Conductance drops towards lower pressure and ultimately becomes pressure-independent, and thus constant, in the molecular-flow regime. At the same time, however, the harmful backflow increases at lower pressure so that the compression ratio K_0 drops at low pressure as well. As indicated in Figure 7.53, the developing maximum in the compression ratio $K_{0,\text{max}}$ appears at a fore-vacuum pressure $p_{\text{F}} \approx 100 \, \text{Pa}$ (1 mbar (hPa)). As pump size increases, the theoretical pumping speed S_{th} increases more rapidly than conductance C, which thus also increases the compression ratio K_0 (see Figure 7.53).

Using the ultimate pressure $p_{F,\text{ult}}$ of the fore pump and the corresponding compression ratio $K_{0,\text{ult}}$, the producible ultimate pressure of the Roots pump (combination of Roots pump and fore pump) calculates to

$$p_{\rm in,ult} = \frac{p_{\rm F,ult}}{K_{\rm 0.ult}}.\tag{7.17}$$

7.4.3.6 Effective Compression Ratio and Volumetric Efficiency [51]

In a system where a Roots pump (pumping speed S) and an attached fore pump (pumping speed S_F) deliver a gas flow (developing in a process or due to leakage), both pumps feature the same pV gas flow (throughput), according to the principle of continuity. In a series connection, the pressure p_F at the outlet of the Roots pump can be assumed to be equal to the inlet pressure of the fore pump:

$$p_{\rm in} S = p_{\rm F} S_{\rm F}.$$
 (7.18)

The effective (K_{eff}) and theoretical (K_{th}) compression ratios are defined as

$$K_{\text{eff}} =: \frac{p_{\text{F}}}{p_{\text{in}}} = \frac{S}{S_{\text{F}}}, \quad K_{\text{th}} =: \frac{S_{\text{th}}}{S_{\text{F}}},$$
 (7.19)

with $S_{\rm th} = \nu V_{\rm S}$. Putting Eq. (7.18) into Eq. (7.12) for the effective pV gas flow, and considering the definitions in Eq. (7.19),

$$\frac{1}{K_{\text{eff}}} = \frac{p_{\text{in}}}{p_{\text{F}}} = \frac{S_{\text{F}}}{S_{\text{th}} + C} + \frac{S_{\text{back}} + C}{S_{\text{th}} + C}.$$
 (7.20)

Generally, the conductance C can be neglected with respect to the theoretical pumping speed S_{th} , so that

$$\frac{1}{K_{\text{eff}}} = \frac{S_{\text{F}}}{S_{\text{th}}} + \frac{S_{\text{back}} + C}{S_{\text{th}}} = \frac{1}{K_{\text{th}}} + \frac{1}{K_{0}},\tag{7.21}$$

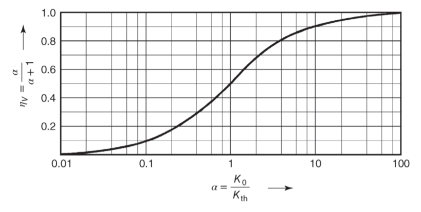


Figure 7.54 For determining the volumetric efficiency η_V of Roots pumps according to Eq. (7.22), $\alpha = K_0/K_{th}$.

or using Eq. (7.19),

$$\frac{K_{\text{eff}}}{K_{\text{th}}} = \frac{S}{S_{\text{th}}} =: \eta_{\text{V}} = \frac{K_0/K_{\text{th}}}{1 + K_0/K_{\text{th}}}.$$
 (7.22)

 $\eta_{\rm V}$ is termed volumetric efficiency. It can be calculated from the pumping speed of the fore pump $S_F = f(p_F)$, the theoretical pumping speed of the Roots pump S_{th} , and the measured compression ratio K_0 of the Roots pump (see Figure 7.53).

Then, η_V allows determining the pumping speed S of Roots pump combinations (Roots pump plus fore pump) according to

$$S = \eta_{V} S_{th}. \tag{7.23}$$

Figure 7.54 shows a plot of η_V using the abbreviation $\alpha = K_0/K_{\rm th}$.

For determining the volumetric efficiency η_V of Roots pumps according to Eq. (7.22), $\alpha = K_0/K_{\text{th}}$.

Initially, Eq. (7.23) gives the pumping speed S with respect to fore-vacuum pressure $p_{\rm F}$. However, the latter is related to inlet pressure $p_{\rm in}$ following the equation of continuity, Eq. (7.18). Thus, pumping speed is given as a function of inlet pressure p_{in} as well (pumping-speed curve).

If multistage combinations are considered, a stepwise approach is convenient, considering the low-stage combination as a fore pump to the subsequent Roots pump.

Gradation of Pumping Speed between Fore Pump and Roots Pump

Two main aspects determine the gradation of the pumping speed from the Roots pump to the fore pump:

- a) High volumetric efficiency η_V is desired.
- b) The maximum tolerable pressure difference $\Delta p_{\text{max}} = p_{\text{F}} p_{\text{in}}$ of the Roots pump shall not be exceeded.

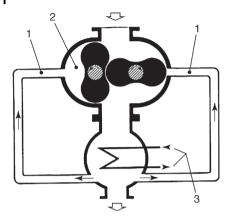


Figure 7.55 Principle of pre-inlet gas cooling for Roots pumps. 1: pre-inlet channel accepting part of the incoming gas flow; 2: suction chamber; 3: gas cooler.

If volumetric efficiency is low, the effective pumping speed of the Roots pump drops considerably. Exceeding the maximum pressure difference causes the compression to heat the pump to an extent, which might cause the impellers to jam due to thermal expansion. This effect is promoted by the intensively cooled pump housing that hardly expands at all [52].

For short pumps, the maximum tolerable pressure difference $\Delta p_{\rm max}$ is higher than for long pumps. It amounts to approximately 4–10 kPa. In the higher-pressure range (above 15 kPa), the tolerable pressure difference is slightly higher due to the transported mass and the associated cooling effect of the impellers.

Special gas coolers attached to the fore-vacuum connection (Figure 7.55) allow higher tolerable pressure differences due to a reduced heat input, which is obtained by recirculating cooled gas. Roots pumps with this type of equipment can handle high levels of contaminants, for example, from steel degassing.

In order to investigate pump gradation according to the above conditions, the equation of continuity, Eq. (7.18), is rewritten as

$$\frac{p_{\rm F}}{p_{\rm in}} = \frac{S}{S_{\rm F}} \quad \text{or} \quad \frac{p_{\rm F} - p_{\rm in}}{p_{\rm in}} = \frac{S}{S_{\rm F}} - 1.$$
(7.24)

Assuming that the pressure difference $p_{\rm F}-p_{\rm in}$ at the Roots pump is below $\Delta p_{\rm max}$ (maximum tolerable pressure difference), then

$$\frac{S}{S_{\rm F}} \le \frac{\Delta p_{\rm max}}{p_{\rm in}} + 1. \tag{7.25}$$

Under medium vacuum ($p_{\rm in} < 100~{\rm Pa}$), with a maximum tolerable pressure difference $\Delta p_{\rm max}$ of 5 kPa, and at an inlet pressure of $p_{\rm in} = 100~{\rm Pa}$ and 10 Pa,

according to Eq. (7.25), $S/S_{\rm F}=51$ and 510, respectively. Thus, gradation is independent of the maximum tolerable pressure difference. The only aspect to be taken into account is that volumetric efficiency is sufficient. If the average maximum compression ratio is around 30, then the volumetric efficiency $\eta_{\rm V}=0.75$ for the theoretical compression ratio $K_{\rm th}$. This value is sufficiently high, and thus, as a rule of thumb:

In the range of medium to high vacuum, the gradation of pumping speed from the fore pump S_F to S of the Roots pump should comply with $S_F/S = 1/10$.

It is important that these considerations apply to a stationary process. For strongly changing working pressure or a fore-vacuum pressure so low that the compression ratio K_0 is small, reduced gradation should be selected (e.g., 1/5).

In the low vacuum range ($p_{\rm in} > 100$ Pa), the maximum tolerable pressure difference is significant. For example, if the maximum tolerable pressure difference $\Delta p_{\rm max}$ is 5 kPa and the inlet pressure $p_{\rm in} = 1$ kPa and 5 kPa, then $S/S_{\rm F} = 6$ and 2, respectively, according to Eq. (7.25).

This means that, according to the maximum tolerable pressure difference and the given inlet pressures, the gradation ratio between the Roots pump and the fore pump is limited to only 6:1 and 2:1, respectively.

The circumstances are best conceived if the *throughputs* of the Roots pump and the fore pump are plotted against inlet pressure p_{in} . Figure 7.56 shows the

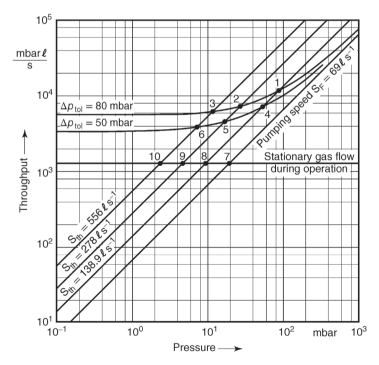


Figure 7.56 Throughput diagram for determining differential pressure Δp .

pV gas flow through the pumps versus p_{in} for selected gradations ($K_{th} = 2, 4$, and 8) of the pumping speed between the Roots pump and the fore pump with constant, pressure-independent pumping speed $S_F = 250 \,\mathrm{m}^3 \mathrm{h}^{-1} \,(69 \,\ell \,\mathrm{s}^{-1})$. Additionally, the diagram takes into account two different maximum tolerable pressure differences of 5 kPa and 8 kPa.

The points 1, 2, 3 and 4, 5, 6 in the diagram give the starting pressures for the three Roots pumps at a maximum tolerable pressure difference of 8 kPa and 5 kPa, respectively. This approximation is pessimistic because it uses the theoretical rather than the effective throughputs of the Roots pumps. Generally, starting pressures are higher without overloading the pumps.

However, for a given throughput, the diagram furthermore serves for determining the producible pressures, the transfer pressure to the fore pump, and the pressure differences. Figure 7.56, for instance, highlights a throughput of $q_{nV} = 1.3 \times 10^{-3}$ mbar(hPa) ℓ s⁻¹(= 0.13 Pa ℓ s⁻¹). Here, the fore pump produces a pressure of $p_{\rm F} = 1.9 \, \text{kPa}$ (point 7). If, subsequently, a Roots pump with $1000 \,\mathrm{m}^3\mathrm{h}^{-1}$ (278 $\ell\,\mathrm{s}^{-1}$) is connected in series, it would theoretically reach a pressure of 4.7 mbar (hPa) (470 Pa) (point 9). In reality, the pressure is higher because $S_{\rm eff} < S_{\rm th}$. The pressure difference $\Delta p_{\rm max} = p_{\rm F} - p_{\rm in} = 19 \, {\rm mbar} \, ({\rm hPa})(1.43 \, {\rm kPa})$ (hPa) - 4.7 mbar (hPa) = 14.3 mbar.

If the pumping speed of the fore pump is pressure-independent, which is common, for example, in oil-sealed positive displacement pumps (see Figure 7.61), the pump gradation and the maximum tolerable inlet pressure can be obtained from Figure 7.57. This diagram, plotted using Eq. (7.25), shows that, for example,

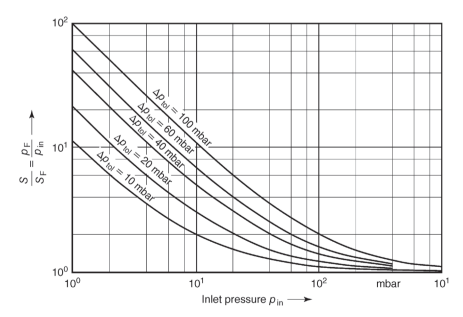


Figure 7.57 Diagram for determining pump gradation $S_{Rootspump}/S_{F(forepump)}$.

for a gradation ratio of $S/S_{\rm F}=10/1$ and $\Delta p_{\rm max}=6\,{\rm kPa}$ (60 mbar (hPa)), the Roots pump can operate continuously at an inlet pressure $p_{in} = 0.66 \text{ kPa}$ (6.6 mbar (hPa)).

7.4.3.8 Pumping Speed and Ultimate Pressure

Roots pumps are positive displacement pumps. Therefore, their pumping speeds and ultimate pressures are nearly constant for all gas species. One exception, however, is the pumping speed for gases with a molecular mass that is below that of nitrogen, for example, hydrogen and helium. Here, pumping speed is slightly lower due to the loss caused by clearances. This leads to lower compression values K_0 as well (see Figure 7.53).

Pumping Speed and Ultimate Pressure in Combinations with Oil-Sealed **Fore Pumps**

Figure 7.58 shows curves of the pumping speed of Roots pumps with oil-sealed sliding vane rotary pumps. The maximum pumping speed is reached at inlet pressures p_{in} of approximately 10 Pa (0.1 mbar (hPa)) to 100 Pa (1 mbar (hPa)). In the actual suction chamber, the Roots pump operates dry. Thus, the pumping-speed curve relates to the total pressure (for oil-sealed pumps, the curves depend on the pressure of the permanent gases). The selected fore pump affects the obtainable ultimate total pressure of $p_{in} = 0.1 \, \text{Pa}$. The producible ultimate partial pressure is lower.

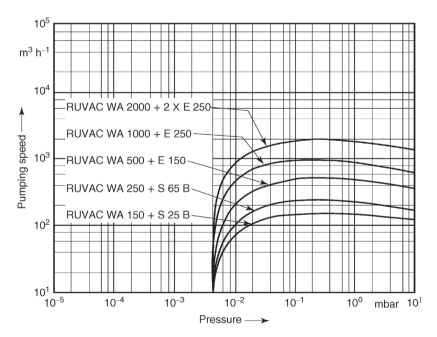


Figure 7.58 Pumping-speed curves of Roots pumps with oil-sealed rotary vacuum pumps as fore pumps.

Pumping Speed and Ultimate Pressure in Combinations with Liquid Ring Vacuum Pumps as Fore Pumps

If liquid ring pumps are used, pumping speeds and ultimate pressures depend on the type of liquid used. To date, water is the most commonly applied liquid, and thus, we will focus on water ring pumps here. Depending on the temperature of the water, the ultimate pressure of a water ring pump generally lies between 2 and 3 kPa. Therefore, the producible ultimate pressure is given by the compression ratio of the Roots pump and amounts to approximately 100 Pa. The ultimate pressure is a result of the water vapor, which develops in the water ring pump and passes through the Roots pump. At higher pressures, that is, if higher gas flows are delivered, the vapor pressure of the water on the inlet side of the Roots pump is lower than would be expected from the compression ratio. This is because the gas flow through the pump represents an additional barrier for water vapor.

Multistage Pump Combinations

Today pairs of impellers may be arranged one behind the other on a common axis of rotation to form multiple stages (Figure 7.59). The outlet of the previous stage is guided in a screw-type manner to the inlet of the following stage. Since the compression ratios of each stage are multiplied, it is even possible to build Roots pumps with an exhaust to atmosphere with a still acceptable pump volume speed. In this way an additional rough pump may be unnecessary. For higher pump speeds at high inlet pressures, however, the Roots pump will be

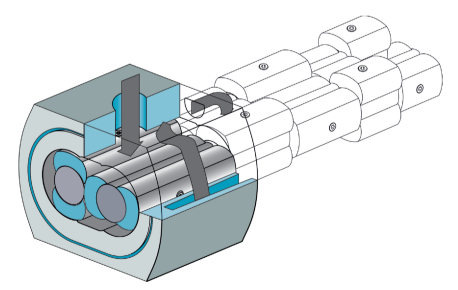


Figure 7.59 Scheme of a five-stage roots pump. All stages are on a common axis. The gray arrows show the direction of gas flow in the first and to the second stage. With courtesy of Pfeiffer Vacuum GmbH.

combined with a suitable rough pump into a pump unit. Rotary vane and screw pumps are chosen quite often for this application [53].

7.4.3.9 Installation and Operating Suggestions

In installation, the fore-vacuum joint of the Roots pump is connected to the inlet joint of the fore pump. In order to prevent overload on the Roots pump at higher pressures due to the pressure difference $p_{\rm F} - p_{\rm in}$, the fore pump initially runs alone during pump down. In this phase, the impellers of the Roots pump follow the flow and rotate without any load as in a gas meter. The throttling effect of the Roots pump is comparably low. It causes a pressure loss between 100 Pa and 500 Pa, depending on the pump type. For pressures $p_{in} > 2 \text{ kPa}$, throttling is so low that it hardly influences the pumping speed of the fore pump. Operation of the Roots pump is not initiated until the pressure has dropped far enough to guarantee that the startup pressure difference at the Roots pump is below the maximum tolerable pressure difference Δp_{max} . Depending on pump gradation, this is the case for inlet pressures between 1 and 10 kPa.

During electrical hookup, attention must be paid to the correct direction of the rotary-current motor's rotation. After the pump is assembled, the correct direction of rotation should be verified by checking the pressure: if the sense of rotation is correct, the pressure drops in the vessel, and if not, it rises after the Roots pump is started. Short periods of operation with incorrect sense of rotation do not damage a Roots pump.

7.5 Specific Properties of Oil-Sealed Positive Displacement Pumps

7.5.1

Pumping Speed and Producible Ultimate Pressure

The volume of the suction chamber and the rotational speed determine the pumping speed of a positive displacement pump. For the same pump size, higher speed increases the pumping speed while production costs remain low. However, rotational speed is subject to mechanical and thermodynamic limitations. Today, positive displacement pumps rotate at speeds of 300-1500 min⁻¹, occasionally up to 3000 min⁻¹, depending on size and type of the considered pump.

According to DIN 28426, part 1 (acceptance specifications for oil-sealed sliding vane rotary pumps and rotary piston pumps), the ultimate pressure of a vacuum pump is the lowest, asymptotically obtainable pressure produced in a measurement dome by the pump. Here, ultimate total pressure (including oilvapor pressure) and ultimate partial pressure are differentiated.

⁶⁾ Translator's note: corresponds to ISO 1607, positive displacement vacuum pumps; measurement of performance characteristics.

7.5.1.1 Pumping Speed and Ultimate Partial Pressure

If a condensation plane of sufficiently low temperature (e.g., a liquid nitrogen (LN₂)-cooling trap) lies between a pressure gauge and a pump, oil vapors flowing back from the pump operating at ultimate pressure condense at the cooled surface, do not reach the gauge, and therefore, are not measured. The pressure measured in this way is referred to as partial pressure, produced as a cumulative pressure only by the permanent gases. Pumping speed measurements according to the acceptance guidelines of it *PNEUROP* and DIN 28426⁷⁾ rely on this value.

Although, in practice, an LN2-cooled condensation plane is rarely used between the pump and the gauge as well as between the pump and the recipient, the acquired pumping-speed curves express the limit of what is producible with the pump. Furthermore, they represent characteristic values of the pump's quality widely independent of the sealing liquid.

In single-stage positive displacement pumps, pumping speed (Figure 7.60) is constant from 100 kPa (1000 mbar) to approximately 1 hPa (1 mbar). At lower pressures, the pumping speed initially drops gradually until ultimate pressure is reached at values of several Pa and zero pumping speed. Values of producible ultimate pressure are about ten times higher if gas ballast (see Section 7.6.1) is used in operation and air is additionally fed to the compression chamber.

Two-stage pumps show a flatter drop in pumping speed below 1 mbar (1 hPa). These pumps reach ultimate pressures around 10⁻² Pa. Generally, larger pumps produce lower ultimate pressures because the ratio of sealing area to suction volume decreases, that is, improves. Gas ballast operation yields ultimate pressures of about 1 Pa.

For other gas species and superheated steam (if no condensation occurs inside the pump), pumping-speed curves are nearly equal to those of air and nitrogen because the pumps work as mechanical scoop pumps.

7.5.1.2 Ultimate Pressure and Oil Selection

The minimal obtainable total ultimate pressure of an oil-sealed positive displacement pump (partial pressures plus oil-vapor pressure) depends on the vapor pressure, the temperature, and the dissolved gas amount in the oil or sealing medium. The temperature is given by the operating temperature of the pump. Using high-quality mineral oils without additives (closely cut fractions), ultimate total pressures of approximately 0.5 Pa are possible in two-stage pumps after longer periods of pump operation when the oil is well-degassed and clean.

If heavy silicone oil (e.g., CR 200, DC 705) or diffusion-pump oil with very low vapor pressure is used, the initial ultimate pressure reaches 0.05 Pa. After a while, however, the ultimate total pressure increases to nearly the value obtained with an unalloyed mineral oil (Figure 7.61).

Diffusion-pump oils are designed for low vapor pressures. Most of them are inappropriate for continuous operation in oil-sealed pumps because, generally,

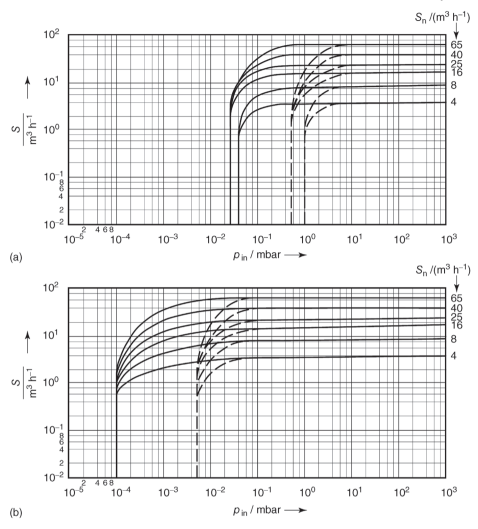


Figure 7.60 Pumping capacities S versus inlet pressure p_{in} and values for nominal pumping capacity S_n in m^3h^{-1} . (a) Single-stage sliding vane rotary pump, (b) two-stage sliding vane rotary pump. Dashed lines: operation with gas ballast (see Section 7.6.1).

the lubricating properties of these oils, particularly silicone oils, are so bad that heavy wear would develop after short periods of operation.

As opposed to mineral oils, silicone oils provide hardly any corrosion resistance at all. Thus, plain iron parts of the pumps corrode rapidly if exposed to oxygen (e.g., from ambient air).

If any other type of alloyed mineral oil, for example, simply picked up from a nearby gas station, is utilized in a positive displacement pump, ultimate

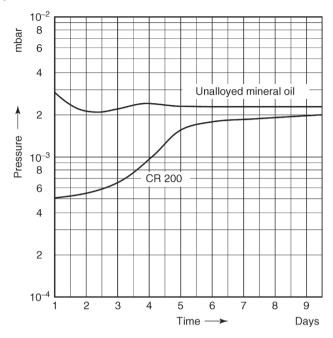


Figure 7.61 Pressure at the inlet port of a two-stage rotary plunger pump for operation with silicone oil CR 200 and an unalloyed mineral oil.

pressures of a two-stage pump around 1 Pa can be expected. In practice, the difference to closely cut, unalloyed mineral oil is often negligible. It follows:

Nearly any mineral oil with appropriate lubricating properties and proper viscosity (kinematic viscosity at 50 °C approximately $\nu = 60~\text{cSt} = 60~\text{mm}^2~\text{s}^{-1}$) is suitable as pump oil. Unsuitable, highly alloyed mineral oils have high vapor pressures, are highly emulsifiable, and are subject to considerable saponification as well as slight oxidation.

Usually, positive displacement pumps run on pure mineral oils with a slight corrosion protection, anti-emulsifying additives (practical for pumping down vapors), and oxidation inhibitors. By this, an ultimate pressure of 0.5 Pa develops in a two-stage pump if the operating temperature remains within certain limits.

Oil with higher corrosion protection, referred to as corrosion-inhibition oil or slushing oil, usually contains alkaline additives that are capable of neutralizing certain amounts of acid depending on the type of additive. Their vapor pressure is slightly higher than that of regular mineral oils and often they are hygroscopic.

After a pump is supplied with fresh oil, it takes some time before the oil is sufficiently degassed and ultimate pressure is actually reached. If degassed oil is used in the pump, initially, the produced ultimate total pressure is lower. A

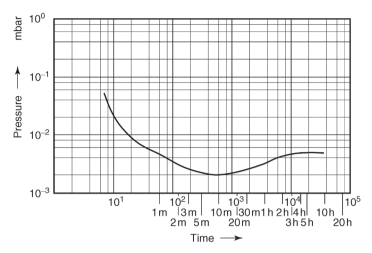


Figure 7.62 Pressure at the (sealed) inlet port in a two-stage rotary plunger pump for operation with pre-degassed oil and open gas ballast valve, m = minutes.

stationary ultimate pressure does not develop until the temperature of the oil has risen (temperature-dependence of vapor pressure), which takes some time when the inlet is shut (no compression work) (Figure 7.62). The ultimate total pressure in single-stage pumps is about ten times higher than in two-stage pumps because of the missing oil degassing in the primary stage.

During the past years, requirements on pump oils have increased considerably. The specific power of the pumps continues to rise, the velocities of sliding motion in bearings, at vanes, etc. increase, and chemical as well as thermal demands rise. Often, regular mineral oils, whether alloyed or unalloyed, cannot cope with these requirements. Thus, certain applications use special lubricating oils and sealing fluids of appropriate viscosity.

For example, lubricating fluids based on phosphoric esters are recommended when pure oxygen is to be pumped down because mineral oil ages particularly rapidly if exposed to pure oxygen. Vacuum and lubricating properties of such liquids are comparable to those of unalloyed mineral oils.

Furthermore, the use of fluorinated hydrocarbons has increased during recent years. One of the main reasons is requirements of uranium separation. Such fluorinated hydrocarbons are practically scentless, noncombustible, and noncorrosive. By now, they are also used in other applications (pumping down oxygen, hydrogen, and other gases). Lubricating properties compare to those of unalloyed mineral oils. However, they are expensive and thus restricted to certain applications.

Sporadically, inhibitors have improved lubricating properties of silicone oils as well. However, here again, high costs impede widespread utilization.

The pump oil is an important design feature of pumps [54–56].

Table A.17 lists additional technical data of oils used in oil-sealed positive displacement pumps and their corresponding fields of application.

In this context, it should be noted that using a special (often expensive) oil does not always represent the correct technical solution to a problem, for example, in chemical [57–59] and the semiconductor industry [60–62]. Moreover, often an additional modification of a standard positive displacement pump using appropriate supplementary devices (e.g., chemical and/or mechanical filters, oil filters, oil circulation devices, etc.), leads to a chemically or corrosion resistant pump [63-67]. Standard pumps are equipped with pressure-proof housings and flame barriers in the inlet and outlet (see Section 7.7.4) for pumping down flammable and explosive substances [68,69].

In these and other special cases, for example, recovery of expensive pump oils [70], it is often advisable to consult the pump manufacturer.

7.5.2 Oil Backflow

During operation of oil-sealed positive displacement pumps, oil vapor flows in the direction opposite to the flow of pumped gases and vapors. It reaches the inlet of the pump and the suction pipe, and if no high-vacuum pump is present, arrives at the vacuum chamber [71,72]. This oil-vapor backflow, shortly referred to as oil backflow, increases with lessening gas counter flow. Thus, it reaches maximum values when the positive displacement pump operates at ultimate pressure (Figure 7.63). The main proportion of the oil backflow is made up of light fractions of the pump oil [73].

However, in production of hydrocarbon-free high and ultrahigh vacua, for example, using Penning type pumps or turbomolecular pumps, a mostly

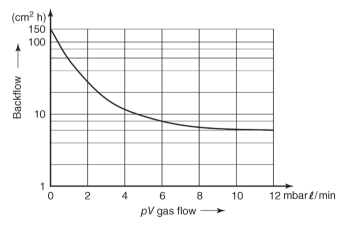


Figure 7.63 Oil-vapor backflow versus pumped-down gas flow [74].

oil-free vacuum is required at the fore-vacuum side of these pumps as well. In practice, several methods are successful for preventing oil backflow [4,74]:

- a) An artificial leakage in the suction line that prevents the pressure from dropping below 10 Pa reduces backflow by approximately 98%. The gas flow prevents back diffusion and permits transport of the pump oil's light fractions through and out of the pump.
- b) In single-stage pumps, it is often sufficient to open the gas ballast valve.
- c) Mounting of a medium vacuum sorption trap (see Section 7.8.1).
- d) Other measures of reducing oil backflow include using special, but often very expensive pump oils, or installing deep-cooled vapor traps (see also Section 9.4.3). However, these reduce pumping speed considerably, and, for this and other reasons have not prevailed in combinations with oilsealed positive displacement pumps.

7.6 **Basics of Positive Displacement Pumps**

7.6.1

Pumping Down Vapors - Gas Ballast

Positive displacement pumps, which compress from inlet pressure to atmospheric pressure p_{atm} or higher, are inappropriate for pumping gases or vapors that condense at the operating temperatures T of 60 °C to 90 °C prevailing in the pump. If during compression the saturation pressure $p_S(T)$ is reached, the gas or vapor liquefies. This has the consequence that the necessary exhaust pressure $\alpha p_{\rm atm}$ ($\alpha \approx 1.3$ for an efficient exhaust) cannot be reached. In particular, this applies to water vapor that is most relevant for vacuum technology. Its saturation pressure (Table A.13) amounts to 20 kPa at 60 °C and 70 kPa at 90 °C. Water accumulates in the pump and is carried over to the inlet side and considerably reduces the ultimate pressure of the pump due to re-evaporation. Additionally, the oil film on lubricating points may be interrupted, and thus, the pump may jam. Furthermore, if larger amounts of water are present, the danger of corrosion arises.

In a first step, we want to make clear when condensation will occur. The suction pressure p_A is composed of partial pressure p_1 containing noncondensing gas and partial pressure $p_{D,1}$ containing the condensable vapor (or gas): $p_A = p_1$ $+p_{\rm D.1}$. To avoid condensation, $p_{\rm D.2}$ of the vapor at the outlet of the pump must not exceed $p_S(T)$:

$$p_{\rm D,2} < p_{\rm S}(T).$$
 (7.26)

The compression ratio K, that is, the ratio of sucked volume to compressed (exhaust) volume, is given by

$$K = \frac{\alpha p_{\text{atm}}}{p_{\text{A}}} = \frac{\alpha p_{\text{atm}}}{p_1 + p_{\text{D,1}}}.$$
(7.27)

As long as condensation does not occur, both the condensable part and the noncondensable (permanent) part of the gas are compressed in the same way. Since *K* is given by the pump, for no condensation the following condition must hold:

$$p_{D,2} = Kp_{D,1} < p_{S}(T). (7.28)$$

Insertion of (7.27) in (7.28) delivers the condition for $p_{D,1}$ where no condensation occurs:

$$p_{\mathrm{D},1} < \frac{p_{\mathrm{S}} p_{\mathrm{1}}}{\alpha p_{\mathrm{atm}} - p_{\mathrm{S}}}.\tag{7.29}$$

It is important that p_S is evaluated for the coldest spot of the suction flow in the pump. Even in the space behind the exhaust valve condensation should not occur.

Example 7.1

A rotary vane pump pumps a permanent gas flow at a pressure of $p_1 = 1$ kPa at the inlet. The exhaust pressure is 30% above atmospheric pressure (101.3 kPa). The operating pressure of the pump is 70 °C. Calculate the maximum allowable, additional water vapor pressure at the inlet so that no condensation occurs.

It is $\alpha = 1.3$ and according to Table A.19 p_s (70 °C) = 31.2 kPa. Applying Eq. (7.29), we obtain

$$p_{\mathrm{D},1} < \frac{31.2 \, \mathrm{kPa} \times 1 \, \mathrm{kPa}}{1.3 \times 101.3 \, \mathrm{kPa} - 31.2 \, \mathrm{kPa}} = 0.31 \, \mathrm{kPa}.$$

This corresponds to a maximum relative water vapor content at the inlet of 0.31/ (1+0.31)=24%.

Raising the pump temperature to approximately 110 °C, for example, by using a heater or by installing thermal insulation, prevents water vapor condensation inside the pump. Also, continuous oil regeneration or exchange can be used. Gaede presented the most elegant method to prevent condensation inside the pump: the gas ballast [33]. Here, a carefully controlled amount of fresh gas (referred to as gas ballast) is fed continuously to the suction chamber of the pump. Thus, the opening pressure of the outlet valve is reached before the water vapor is compressed to the saturation vapor pressure corresponding to the pump temperature, and therefore, before condensation occurs. The fresh gas inflow (usually atmospheric air) starts immediately after the suction volume is separated from the inlet. Hence, ultimate pressure impairment is limited.

The amount of required fresh gas or the amount of gas ballast q_{nV}^{ballast} is derived from the following considerations: the gas ballast reduces the compression ratio (7.27), since, in addition to the sucked gas flow, the gas ballast flow has to be pumped away. With the gas ballast valve open, the exhaust valve opens at a smaller compression of the sucked gas compared to gas ballast valve closed. With gas ballast we obtain the compression ratio K to

$$K = \frac{\alpha p_{\text{atm}}}{p_1 + p_{\text{D,1}} + \frac{q_{pV}^{\text{ballast}}}{S}}.$$
(7.30)

This equation is based on the consideration that the partial pressure of the fresh gas develops according to the ratio of gas ballast flow rate in $q_{pV}^{\mathrm{ballast}}$ and pumping speed S of the pump. Use of S makes sense, since the gas ballast valve immediately opens after isolation of the sucked volume from the inlet port (see Figure 7.25). At this very moment the sucked volume is not yet reduced, and therefore the same pumping speed S can be assumed. Putting (7.30) in (7.28) we obtain the condition for $p_{D,1}$ with gas ballast flow:

$$p_{\mathrm{D},1} < \frac{p_{\mathrm{S}}p_{1} + p_{\mathrm{S}} \frac{q_{pV}^{\mathrm{ballast}}}{S}}{\alpha p_{\mathrm{atm}} - p_{\mathrm{S}}}.$$
 (7.31)

The other way round, for given $p_{D,1}$, we obtain the necessary gas ballast flow for no condensation:

$$q_{pV}^{\text{ballast}} > S\left(\frac{\alpha p_{\text{atm}} p_{\text{D},1}}{p_{\text{S}}} - p_{\text{D},1} - p_1\right). \tag{7.32}$$

Example 7.2

Consider the conditions of Example 7.1 Determine the gas ballast flow necessary so that no condensation occurs, when the water vapor pressure at the inlet amounts to 0.4 kPa. The pumping speed be $S = 6 \text{ m}^3 \text{ h}^{-1}$.

It is
$$6 \text{ m}^3 \text{ h}^{-1} = 1.67 \times 10^{-3} \text{ m}^3 \text{ s}^{-1}$$
.

$$q_{pV}^{\text{ballast}} > 1.67 \times 10^{-3} \text{ m}^3 \text{ s}^{-1} \left(\frac{1.3 \times 101.3 \text{ kPa} \times 0.4 \text{ kPa}}{31.2 \text{ kPa}} - 0.4 \text{ kPa} - 1 \text{ kPa} \right)$$
$$= 0.481 \text{ Pa m}^3 \text{ s}^{-1}.$$

Comparing this value with the pump throughput

$$\dot{Q} = 1.67 \times 10^{-3} \text{ m}^3 \text{ s}^{-1} \times 1.6 \times 10^3 \text{ Pa} = 2.5 \text{ Pa m}^3 \text{ s}^{-1}$$

we see that the gas ballast throughput has to be nearly 20% of the pump throughput. By this percentage the effective compression ratio is reduced by the gas ballast.

In deriving Eqs. (7.31) and (7.32,) we have not yet considered two facts: if, as common, fresh air is used for gas ballast, this also contains condensable water vapor. Second, the fresh environmental air at temperature T_0 is heated up by the pump. Let us denote the throughput of water vapor in the gas ballast by q_{pV}^{W} . Here exists the additional condition that this water vapor of the fresh air does not condense:

$$K \frac{q_{pV}^{W}}{S} \frac{T}{T_0} < p_{S}(T).$$
 (7.33)

If water vapor is also the condensable gas at the inlet pump port, we can combine Eqs. (7.28) and (7.30) as

$$K\left(p_{D,1} + \frac{q_{pV}^{W}}{S} \frac{T}{T_0}\right) < p_{S}(T).$$
 (7.34)

Equation (7.30) must be modified if the gas ballast throughput contains water vapor throughput q_{pV}^{W} in addition to the noncondensable part q_{pV}^{0} . It is $q_{pV}^{\text{ballast}} =$ $q_{pV}^{W} + q_{pV}^{0}$ and we obtain as replacement for Eq. (7.30)

$$K = \frac{\alpha p_{\text{atm}}}{p_1 + p_{\text{D},1} + \frac{q_{pV}^{\text{W}}}{S} + \frac{q_{pV}^{0}}{S}}.$$
 (7.35)

Putting Eqs. (7.35) into (7.34), we obtain after some calculation the condition for noncondensation of water vapor

$$p_{\mathrm{D,1}} < \frac{p_{\mathrm{S}}}{S} \left[\frac{p_{1}S + q_{pV}^{0} \left(1 + \frac{q_{pV}^{\mathrm{W}}}{q_{pV}^{0}} - \frac{\alpha p_{\mathrm{atm}}}{p_{\mathrm{S}}} \frac{q_{pV}^{\mathrm{W}}}{q_{pV}^{0}} \frac{T}{T_{0}} \right)}{\alpha p_{\mathrm{atm}} - p_{\mathrm{S}}} \right]. \tag{7.36}$$

Example 7.3

We consider the conditions of Examples 7.1 and 7.2. Let $q_{\rho V}^{\text{ballast}} = 0.482 \, \text{Pa m}^3 \text{s}^{-1}$, $S = 6 \text{ m}^3 \text{ h}^{-1}$, T = (273 + 70) K, $T_0 = 296 \text{ K}$, $\alpha = 1,3$, and $q_{pV}^W/q_{pV}^0 = 0.014$, according to a relative humidity of 50%. Determine the maximum value of $p_{D,1}$, which is the water vapor tolerance, as accurate as possible.

It is $q_{pV}^{W} = 0.0067 \text{ Pa m}^3 \text{s}^{-1}$ and $q_{pV}^{0} = 0.475 \text{ Pa m}^3 \text{s}^{-1}$. Application of Eq. (7.36) yields

$$p_{\text{D,1}} < \frac{31.2 \times 10^{3} \text{ Pa}}{1.67 \times 10^{-3} \text{ Pa m}^{-3} \text{ s}^{-1}} \times \left[\frac{1 \times 10^{3} \text{ Pa} \times 1.67 \times 10^{-3} \text{ Pa m}^{-3} \text{ s}^{-1}}{1.3 \times 1.013 \times 10^{5} \text{ Pa}} \times \frac{1.3 \times 1.013 \times 10^{5} \text{ Pa} \times 1.67 \times 10^{-3} \text{ Pa m}^{-3} \text{ s}^{-1} + 0.475 \times \left(1 + 0.014 - \frac{1.3 \times 1.013 \times 10^{5}}{31.2 \times 10^{3} \text{ Pa}} 0.014 \frac{343 \text{ K}}{296 \text{ K}}\right)}{1.3 \times 1.013 \times 10^{5} \text{ Pa} - 31.2 \times 10^{3} \text{ Pa}}$$

In Example 7.2, we assumed $p_{\rm D,1} = 400 \, \rm Pa$ for the calculation of $q_{\rm pV}^{\rm ballast}$. Compared to this result, we see that the water vapor tolerance is practically equal to the approximations made for Eq. (7.32).

The maximum gas ballast throughput is needed when pure water vapor is pumped. In this case $p_1 = 0$. We can simplify Eq. (7.36) with $T/T_0 \approx 1$:

$$p_{\rm D,1} < \frac{q_{pV}^0}{S} \frac{p_{\rm S} \left(1 + \frac{q_{pV}^{\rm W}}{q_{pV}^0} \right) - \alpha p_{\rm atm} \frac{q_{pV}^{\rm W}}{q_{pV}^0}}{\alpha p_{\rm atm} - p_{\rm S}}.$$
 (7.37)

Setting $p_{D,1}$ equal to the right term in Eq. (7.37), we obtain the so defined water vapor tolerance $p_{\rm W}$ of a pump (ISO 21360-2, DIN 28426). This is the pressure that must not be reached at the pump inlet when no condensation occurs. Multiplying p_{W} with S one obtains the maximum water vapor throughput for no condensation.

For given $p_{D,1}$ we can determine the necessary gas ballast throughput so that no condensation occurs:

$$q_{pV}^{\text{ballast}} \cong q_{pV}^{0} > S \frac{p_{\text{D,1}}}{p_{\text{S}}} \frac{\alpha p_{\text{atm}} - p_{\text{S}}}{1 + \frac{q_{pV}^{\text{W}}}{q_{pV}^{0}} \left(1 - \frac{\alpha p_{\text{atm}}}{p_{\text{S}}}\right)}.$$
 (7.38)

Finally, it shall be mentioned that usage of gas ballast was also found to be beneficial when gases pumped previously have to be removed from the pump system, for example, helium in rough pumps of leak detectors [75].

7.6.2

Power Requirements

The amount of energy required to operate pumps of the types investigated in Sections 7.2 and 7.3 depends on the compression work, the frictional loss, and the exhaust stroke work. Compression work is best described by considering a piston pump, Figure 7.1. If the piston travels a distance dl and the effective force F = pA (with A = piston area), then the work

$$dW = -p A dl = p dV. (7.40)$$

The negative sign is due to the fact that work is done upon the system if the volume decreases (negative dW). Equation (7.40) depends on the type of pump used. It applies to any kind of positive displacement pump (as described above).

We will now investigate the course of pressure using a pV diagram (Figure 7.64) of a piston pump as shown in Figure 7.1:

- **4–1** the intake stroke (filling of the cylinder) at inlet pressure p_1 in the inlet of
- **1–2** compression in the cylinder to exhaust pressure p_2 , usually atmospheric pressure;
- 2-3 the exhaust stroke that removes the compressed gas from the cylinder after the outlet valve has opened;
- 3-4 the exhaust valve closes, the inlet valve opens; Figure 7.64 idealizes the slight increase in volume to zero.

The work required for a single process cycle is obtained by integrating Eq. (7.40):

$$W = -\oint p \, \mathrm{d}V. \tag{7.41}$$

Geometrically, this integral corresponds to the shaded area in Figure 7.64, which can be written as

$$W = \int_{p_1}^{p_2} V \, \mathrm{d}p. \tag{7.42}$$

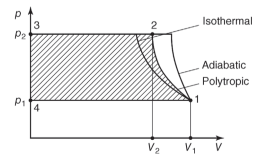


Figure 7.64 pV-diagram of a compression process.

In order to calculate the area, we need to know the shape of the 1–2 curve, which depends on whether the compression is isothermal, adiabatic, or polytropic.

7.6.2.1 Isothermal Compression

If appropriate measures are taken to effectively remove the compression heat, compression is nearly isothermal (compression at constant temperature). This is the case, for example, in a liquid ring pump. As then the 1-2 curve is an isothermal line, the *Boyle –Mariotte* law, Eq. (3.16), applies and $V = p_1 V_1/p$. Thus, the isothermal compression work

$$W_{\text{isothermal}} = p_1 V_1 \int_{p_1}^{p_2} \frac{\mathrm{d}p}{p} = p_1 V_1 \ln \frac{p_2}{p_1}. \tag{7.43}$$

7.6.2.2 Adiabatic Compression

During adiabatic (isentropic) compression, no heat is released during compression. The 1-2 curve is an adiabatic line, Eq. (4.42), and because of $p V^{\kappa} = \text{constant}.$

$$V = \frac{V_1 \, p_1^{1/\kappa}}{p^{1/\kappa}},\tag{7.44}$$

and the adiabatic compression work

$$W_{\text{ad}} = \int_{p_1}^{p_2} V \, \mathrm{d}p = V_1 \, p_1^{1/\kappa} \int_{p_1}^{p_2} \frac{\mathrm{d}p}{p^{1/\kappa}} = V_1 \, p_1^{1/\kappa} \frac{\kappa}{\kappa - 1} \, \left(p_2^{\kappa - 1\kappa} - p_1^{\kappa - 1\kappa} \right)$$
$$= \frac{\kappa}{\kappa - 1} p_1 V_1 \left[\left(\frac{p_2}{p_1} \right)^{\kappa - 1\kappa} - 1 \right]$$
(7.45)

7.6.2.3 Polytropic Compression

The compression in positive displacement pumps does not correspond to the special cases of isothermal or adiabatic compression, but is rather polytropic. This means that part of the compression heat is released and the other part remains with the gas. Then, the 1-2 curve is nearly a polytropic line, which can be described as

$$pV^{\zeta} = \text{constant} = p_1 V_1^{\zeta} = p_2 V_2^{\zeta}, \tag{7.46}$$

in which $1 < \zeta < \kappa$. The resulting compression work for a polytropic compression is:

$$W_{\text{pol}} = \frac{\zeta}{\zeta - 1} p_1 V_1 \left[\left(\frac{p_2}{p_1} \right)^{\zeta - 1\zeta} - 1 \right]. \tag{7.47}$$

The less heat is released by the pump, that is, the higher rotating speed and pumping speed, the closer the polytropic exponent ζ approaches κ (for high pumping speeds, the ratio of surface and pumping speed decreases, and thus, less heat is released).

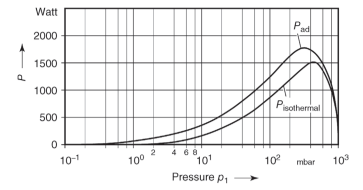


Figure 7.65 Calculated power consumptions, Eqs. (7.43) and (7.45) with right side multiplied by ν , of a single-stage rotary plunger pump with a suction chamber volume $V_1=5\,\mathrm{I}$ and a rotary frequency of $\nu=450\,\mathrm{min}^{-1}$. Abscissa:

pressure p_1 at the inlet port of the pump; ordinate: adiabatic (P_{ad}) and isothermal ($P_{isothermal}$) power consumptions. Exhaust pressure $p_2 = 110$ kPa.

7.6.2.4 Compression Power

In order to calculate compression power P, the work-terms W_{isotherm} , W_{ad} , and W_{pol} are simply multiplied with the rotating frequency ν .

Figure 7.65 shows plots of power consumptions versus pressure p_1 in the inlet. Here, the ejection pressure (pressure at which the valve opens) was set to $p_2=110\,\mathrm{kPa}$. Obviously, the compression power shows a maximum at $p_1=405\,\mathrm{mbar}$ (40.5 kPa) for the isothermal case and at $p_1=339\,\mathrm{mbar}$ (33.9 kPa) for the adiabatic case. The maximum value is obtained by differentiating the above equations. For the isothermal case, $P_{\mathrm{isothermal,max}}=1517\,\mathrm{W}$, and for the adiabatic case, $P_{\mathrm{ad,max}}=1780\,\mathrm{W}$, considering air ($\kappa=1.4$).

The total power of the pump includes the compression power, and in addition, the power required to overcome friction (bearings, seals, vanes, etc.). In liquid ring pumps, the latter is mainly caused by fluid friction in the liquid ring, in sliding vane rotary pumps and rotary plunger pumps, by oil friction. Because oil viscosity is highly temperature-dependent, so is the power required to overcome friction (Figure 7.66).

For low pressures, hardly any compression work is done. The power consumed levels the frictional losses. If the pump operates with gas ballast, the drop in power consumption for lower pressures is flatter due to the continuously required compression work of the gas ballast (Figure 7.67).

The larger the pump, the larger cross sections are required for the outlet, and therefore, for the exhaust valves as well. Sufficient sealing of these valves calls for higher closing forces, and thus, higher opening pressure for the valve. This applies not only to exhaust valves but to other downstream components as well, such as separators that require higher counter pressure due to the throttling effect on the gas flow. Here, additional power is needed, which absolutely can be in the range of the plain compression power. This fact must be considered in selection processes for driving motors, particularly for larger positive displacement pumps.

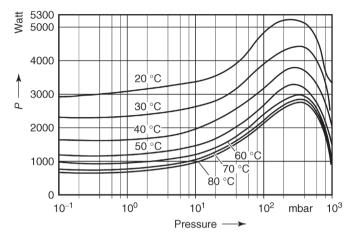


Figure 7.66 Power consumptions for operation without gas ballast at selected oil temperatures (same pump as in Figure 7.65).

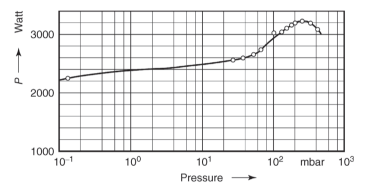


Figure 7.67 Power consumptions for operation with gas ballast at normal operating temperature (same pump as in Figure 7.65).

7.7 **Operating and Safety Recommendations**

7.7.1 Installation

Except for large piston pumps, positive displacement pumps do not require a special foundation because, usually, they run smoothly. However, they should be secured to the floor. At recipients, even the smallest vibrations generally are disturbing. Therefore, a damping element (e.g., a cushioning device) that absorbs such vibrations is often placed between the pump and the recipient.

If pumped gases are harmful to the environment, the pump is connected to a central exhaust line, which connects to necessary cleaning facilities. Also, the exhaust line can contain filters that separate the oil in the oil mist ejected from the pumps at higher pressure. When the exhaust line is hooked up, it should be considered whether harmful pump vibrations occur and whether a damping or elastic element is again required between the pump and the exhaust line.

7.7.2

Starting and Shut Down, Inlet Valves

Large positive displacement pumps are equipped with rotary-current motors; smaller types have AC motors. The latter usually include power switches and the motor determines the sense of rotation. For pumps with rotary-current motors, the direction of rotation must be checked each time after hookup.

If the pump rotates in the wrong direction, it transports pump oil into the recipient. Therefore, such pumps are frequently equipped with a reverse-run safety device that prevents rotation in the wrong direction. If the rotary-current motor is then hooked up the wrong way, the pump does not start. The motor would overheat and therefore features a protective motor switch (usually, a bimetal excess-current circuit breaker), which should be used in any case. AC motors are protected by excess-temperature switches (*Klixon*).

Power consumption of the motors in oil-sealed vacuum pumps is completely different from standard machines. Shortly after starting up, a considerable amount of power is required due to the compression work at high pressure and because of the high viscosity of the cold pump oil. In contrast, a warm pump, operating close to ultimate pressure, only consumes a fraction of the initially required power (see Figure 7.66). This must be considered when adjusting the protective motor switch. If the protection is designed to work during ultimate pressure operation, the current is adjusted to correspondingly low values. In this case, however, the protective switch must be bypassed (sometimes using a time lag relay) during the starting phase.

A pump's power consumption is high when it starts and the suction and compression chambers contain cold oil. Starting is particularly difficult if the recipient is evacuated because the pressure difference between the suction and compression chambers is 1 bar (100 kPa).

Usually, motors are designed to start even at 5% under voltage, which is accepted by the VDE, 80 and under heavy starting conditions (vacuum at inlet, oil completely inside the compression and suction chambers, open gas-ballast valve, and cold pump). PNEUROP acceptance guidelines [1] and DIN 28426⁹⁾ specify the lowest tolerable starting temperatures in order to keep pump sizes within reasonable limits.

⁸⁾ Translator's note: Association for Electrical, Electronic and Information Technologies.

⁹⁾ Translator's note: corresponds to ISO 1607.

The lowest starting temperature is the mean temperature of the pump at which the pump starts flawlessly with ventilated inlet using a driving motor supplied or recommended by the manufacturer after operating was interrupted for a minimum of one hour. If no temperature is specified, 12 °C is the lowest starting temperature. 100

During operation, the pump temperature rises to 70-90 °C, which is not alarming at all and rather beneficial for pumping down vapors. Usually, not the pump but the pump oil limits the tolerable temperature.

If the pump is stopped and no particular measures are taken, it will rotate backwards under the influence of the ambient air pressure. Thus, the oil is pressed into the inlet from the pump and the recipient is vented. Pumps with a reverse-run safety device avoid backward rotation but not venting of the recipient.

Venting is prevented by sealing the recipient using special inlet valves (Section 7.8.2).

7.7.3

Pump Selection and Operating Recommendations

Single-stage oil-sealed pumps are ideal for low and the beginning medium vacuum range. They operate with or without gas ballast without showing a noteworthy drop in pumping speed. The critical pressure for the operating range of these pumps is approximately 10 Pa.

If reliable operating pressure is intended to be lower than this value, or if operation below 100 Pa and with gas ballast is desired, then a two-stage pump is preferable. (Of course, for pressures in the area of 100 Pa, not the total gas ballast would be required. However, closing the gas ballast valve during pump down would require additional operator action, and therefore, remains undone in practice.) For longer periods of operation at high inlet pressures, care must be taken that lubrication, often relying on gravity and the pressure difference between suction chamber and exhaust, is still sufficient. If necessary, inlet lubrication must be applied.

A number of accessory components (separator, filter devices, exhaust filters, and others, see Section 7.8) are available that protect pumps under rough and contaminated operating conditions.

Modern oil-sealed pumps require little maintenance work. Attention should be paid mainly to the amount and quality of the oil. Therefore, the oil level must be checked at least once a week. The oil level is checked during operation after the system has reached ultimate pressure because most of the oil can gather in the suction chamber when the pump stands still. From

time to time, oil levels should also be checked in pumps with separate gearboxes.

If the pump mostly operates at high inlet pressure or with gas ballast, oil levels should be checked on a daily basis because here oil loss is particularly high. For a discharge of one cubic meter of standard air, the oil loss amounts to approximately 2-3 cm³.

Example 7.4

A pump with a nominal pumping speed of 250 m³ h⁻¹ operating at an inlet pressure of $p_{in} = 100 \text{ Pa}$, delivers approximately $250/1013 \text{ m}^3 \text{h}^{-1} = 0.25 \text{ m}^3 \text{h}^{-1}$ of air at standard pressure. This value increases to $25 \,\mathrm{m}^3 \mathrm{h}^{-1}$ for $p_{\mathrm{in}} = 10 \,\mathrm{kPa}$. The corresponding oil consumption is approximately $0.5-0.75 \text{ cm}^3\text{h}^{-1}$ at $p_{\text{in}} = 100 \text{ Pa}$ and approximately $50-75 \text{ cm}^3\text{h}^{-1}$ at $p_{\text{in}}=10 \text{ kPa}$. If the pump operates with 10% gas ballast, an additional 25 m³h⁻¹ gas ballast air is delivered at standard pressure. Thus, the oil consumption for the gas ballast amounts to $50-75 \,\mathrm{cm^3h^{-1}}$ as well. During a 24 h shift with an inlet pressure of $p_{\mathrm{in}}=100\,\mathrm{Pa}$ and additional gas ballast, the pump consumes $1.25-2 \ell$ of oil.

Oil-change intervals depend on the degree of contamination. The oil is changed either if decomposition products in the oil prevent obtaining ultimate pressure or if dirt (mechanical contaminants) or insufficient lubricating properties of the oil threaten to damage the pump. Generally, changing the oil once to often is better than neglecting an oil change. Under smooth ultimate-pressure operation, a yearly oil change is usually sufficient.

These recommendations regarding oil, oil levels, and oil change apply to sliding vane rotary pumps and rotary plunger pumps using oil circulation. Lately, in certain special applications, such pumps operate on once-through lubrication [76,77] (see Section 7.3.2.4).

7.7.4

Technical Safety Recommendations

Since July 1, 2003, in the European Union (EU), operation of vacuum pumps under specified intended operation in potentially explosive environments, that is, for installation in potentially explosive atmospheres and/or pumping of potentially explosive gases, vapors, or dusts, is regulated by the 94/9/EC (ATEX 95) Directive. It guarantees a mutual legal law for explosion protection of nonelectrical systems for all member states, in addition to the recognized national regulations on explosion protection for electrical appliances. The 94/9/EC Directive aims at reducing trade barriers and establishing collective safety standards. The legally valid national implementation of the 94/9/EC Directive for Germany incorporates a new approach. This means that the directive defines minimum standards for appliances and protecting devices for specified intended operation in potentially explosive environments rather than containing detailed test specifications and/or legal rules. Unfortunately, interpretation is sometimes ambiguous due to this concept.

Two new EU guidelines divide the responsibility among system operators and manufacturers (suppliers) of vacuum pumps for specified intended operation in potentially explosive environments.

Directive 1999/92/EC (ATEX 137) regulates responsibilities of system operators. It describes the minimum standards for protecting persons working in potentially explosive environments. A key issue is the zone division based on a risk analysis performed by the system operator [78]. The zone division determines the required equipment category of a vacuum pump. For example, the inside of a drying plant is defined as Zone 0 (regularly potentially explosive atmosphere) and the vicinity of the drying plant is part of Zone 1 (rarely but potentially explosive atmosphere even under standard operation). This translates into the required equipment Category 1 (inside) and equipment Category 2 (outside) of a vacuum pump.

Directive 94/9/EC, described above, defines the required approach that manufacturers of vacuum pumps use to test their products and document the results in order to declare explicitly the authorized areas of application with a CE symbol and an explosion proof designation [41,79].

Particularly dry-running vacuum pumps, for example, screw-type vacuum pumps, claw vacuum pumps, or Roots pumps, must be regarded as potential ignition sources on the inside of the pump. For any dry-running vacuum pump operating in potentially explosive low vacuum, it is likely that the rotors expand and touch the cooled housing when they heat up during operation. Therefore, Category 1 appliances usually feature flame arresters on the inlet and outlet to the pump. This measure prevents flame propagation from the pump. In contrast, Category 2 appliances nowadays generally do without flame arresters due to a number of electronic monitoring systems. These safety measures prevent ignition sources from developing.

In the realm of initial experimental tests for EC type approvals of vacuum pumps according to Directive 94/9/EC, it was found that typical flame arresters used until then, which were tested in conventional testing stands [80], in part, showed flame propagation. Obviously, the effect was produced by a change in geometrical and physical conditions. For this reason, any experimental EC type approval nowadays includes triggering of controlled explosions under different operating conditions with mounted flame arresters and an observation to determine whether flame propagation occurs.

Figure 7.68 shows a dry-running screw-type vacuum pump, type COBRA TC 2250. This device is a Category 1 appliance in accordance with Directive 94/9/ EC and includes flame arresters.

When flame arresters are used, potential contamination and clogging caused by solid particles that are carried along must be considered. In certain applications, the pumping speed can decrease dramatically within a short period due to



Figure 7.68 Screw-type vacuum pump COBRA TC 2250, Category 1 (inside), according to 94/9/EC; top left: flame trap on the inlet with endurance burning flame arrester and

shutting flap; bottom right: flame trap on the pressure side and temperature switch as well as pressure switch between the pump and the flame trap.

low clearance in the flame arresters. Additionally, the pressure loss in the flame arrester on the inlet side can be considerable. Below approximately $50\,hPa$, the pressure loss in this area can cause progressively lowered pumping speed in the vacuum pump, and for $10\,hPa$, it can decrease to 25% of the initial pumping speed.

7.8 Specific Accessories for Positive Displacement Pumps

7.8.1

Sorption Traps

To prevent oil from emanating out of an oil-sealed positive displacement pump and reaching the high-vacuum part of a vacuum system, a sorption trap is placed in between the positive displacement pump and the high-vacuum pump (usually, a turbomolecular pump).

A sorption trap (Figure 7.69) contains an appropriate sorption medium. Besides zeolite, activated, granular alumina is the most effective of all investigated media. The efficiency is 99% and the throttling effect on the pumping speed is

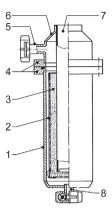


Figure 7.69 Partial section of a fine-vacuum sorption trap. 1: trap housing; 2: strainer basket; 3: molecular sieve (filling); 4: sealing flanges; 5: connecting flange with small flange; 6: trap top; 7: heating or cooling medium vessel; 8: connecting port with small flange.

low when designed correctly. However, it should be taken into account that the sorption medium is saturated after a certain period and must be replaced or regenerated. Saturation does not occur solely due to sorption of back flowing oil vapors but also because of the sorption of pumped down gases and vapors. Therefore, it is often advisable to connect two traps in parallel using valves to alternately operate and maintain the traps (see also Section 11.2.3.2). If this approach is impractical for design or economical reasons, an alternate line that bypasses the sorption trap can be used. Initially, large amounts of gas, for example, when evacuating starts at atmospheric pressure, are pumped through this line when the trap is shut off. Subsequently, the trap is activated after the pressure has dropped to several 100 Pa.

7.8.2 Safety Valves

A safety valve (Figure 7.70) attached to the inlet of a pump separates the inlet from the recipient and vents the pump in case of a power failure or when the pump is shut down. Sealing and venting are subsequent steps, and thus, air backflow is practically prevented when the valve closes. The electromagnetic control valve is connected electrically with the motor and controls the air inlet, which closes the actual valve by means of the pressure difference between the inlet line and ambient pressure, and subsequently vents the pump. Thus, no oil flows into the suction chamber. When the pump starts, the control valve is shut and the pump evacuates the safety valve until the integrated spring reopens it. Another method incorporates a so-called inlet valve which is built into modern sliding vane rotary pumps (Figure 7.27). Here, different mechanisms (centrifugal switches, magnetic switches, hydraulic switches, and others) provide that the inlet of the pump is closed and the pump is vented. If an inlet valve is used,

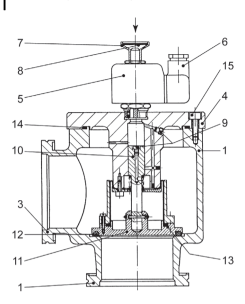


Figure 7.70 Section of safety valve SECUVAC. 1: housing; 2: connecting flange to vacuum equipment; 3: connecting flange for pump; 4: top; 5: air intake valve (electromagnetic); 6:

cable connection; 7: air intake, with 8: filter disk; 9: nozzle; 10: piston; 11: valve plate; 12: valve plate seal; 13: roll diaphragm; 14: seal; 15: connecting bolt.

safety valves as described above are not required. The leak rate of these valves is very low and comparable to the leak rates of standard valves.

7.8.3 Oil Filter and Oil Cleaning

Often, at some phase in vacuum pump operation, larger amounts of solid particles may appear, which remain inside the pump, accumulate, and cause disturbance or excessive wear. In these cases, using a device that provides continuous oil filtering during operation is advisable. This can be realized in different ways. Some filter units bypass the vacuum pump with an independent oil pump. Others are integrated into the oil supply to the pump as full-flow filters.

The way such a filter works shall be exemplified with an oil filter unit. In an oil filter unit, a filter pot attaches to the vacuum pump with a concentric adapter tube on one side, at the point usually holding the oil drain plug. The filter pot holds a filter cartridge. During operation, oil penetrates from the pump reservoir above the exhaust valves through the outer concentric tube (Figure 7.71) and into the filter pot where it moves on to the filter cartridge (e.g., a standard truck filter cartridge). After passing through the filter cartridge where mechanical contaminants are held back, the cleaned oil reenters the pump's oil circuit through the inner tube of the two concentric tubes. Thus, an independent oil circulation develops between the filter, the oil reservoir on the exhaust side, and the inlet

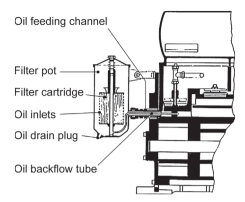


Figure 7.71 Oil filter, schematic.

chamber of the vacuum pump. Here, the pressure difference is used to transport the oil whereas separate oil pumps are required for this task in other types of systems.

Generally, a significant amount of oil is exposed to high-pressure gas (usually, standard air at atmospheric pressure) inside the filter. Therefore, the ultimate pressure is slightly lower in systems with oil filters than in systems without oil filters. In two-stage pumps, the oil supply can be arranged in a way that the oil initially degases before it enters the stage on the suction side (high-vacuum stage). A filter unit increases the amount of oil required for oil circulation. In air-cooled pumps, this is advantageous because the oil is cooled more effectively. A particular advantage is observed in applications where corrosive gases or vapors are pumped down. If no particular measures have been taken, the amount of oil present (including appropriate inhibitors) is the only variable determining what amount of critical media can be stored or neutralized in the oil. Additionally, the filter unit allows feeding chemically active substances (e.g., sodium carbonate under acid conditions) to the oil circuit, which reduces oil loading.

Cool tubes and filters are a sign of faulty oil circulation because hot oil from the vacuum pump should pass through the filter unit. In this case, usually, the filter is saturated and has built up a high pressure difference, which prevents the oil from passing through. However, special monitoring elements (e.g., manometers) are often present, which indicate a saturated filter unit. Then, filter replacement or cleaning is necessary. In the course of this work, it is also advisable to change the pump oil and clean the connecting tubes.

7.8.4 Exhaust Filter (Oil-Mist Separator)

Due to the intense mixing of gas and pump oil during compression and ejection phases, a vacuum pump not only delivers pumped gas at the exhaust but also oil

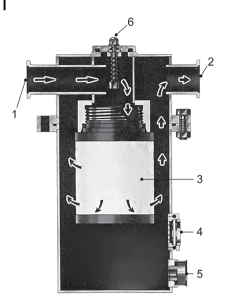


Figure 7.72 Oil mist separator. 1: connecting flange to pump; 2: oil-free exhaust; 3: filter; 4: oil-level glass; 5: oil drain; 6: overpressure valve.

particles. These are carried over by the gas (aerosol) and are visible as smoke or oil mist in the exhaust (additionally, small amounts of oil vapor corresponding to the vapor pressure of the oil occur). This oil mist is disturbing in the atmosphere of the operating area. In the past, oil mist was collected in the exhaust line and discharged to the outside together with the pumped gas. Filter elements represent an environmentally friendly solution.

The droplet size of the oil aerosol in the exhaust is in the range of $0.01-0.8\,\mu m$. Conventional fibrous or ceramic filters cannot separate these fine droplets. Therefore, a filter material was developed that is made of very fine borosilicate fibers, formed into a compound providing sufficient porosity to absorb contaminants. This filter material was pressed into cylindrical filter elements and supported appropriately for it to withstand pressure differences. Exhaust filters as the one shown in Figure 7.72 contain such filter elements in cylindrical vessels.

The exhaust gas of the pump enters into the exhaust filter, is cleaned from oil mist in the filter element, and leaves the filter after passing through the filter element. The filter element collects the oil, which drips out at the bottom of the element. Using the oil-sight glass, the separated oil can be checked and, if necessary, drained. When the filter element is filled with contaminants, the pressure loss increases. At a pressure of 150 kPa (50 kPa or 0.5 bar overpressure), the integrated pressure control valve opens, which indicates that filter element exchange is required.

The size of the exhaust filters is adopted to the pumping speed of the corresponding pump. However, it can be advisable to connect a series of smaller pumps to a large separator, particularly when the pumps operate intermittently.

7.8.5 **Dust Filters**

In certain processes (e.g., steel degassing), larger amounts of dust develop that are carried away by the gas, enter the vacuum pump, and produce abrasive mixtures with the pump oil that can be harmful to the pump. If the amount of dust is low, filtering of the oil circuit is sufficient. Oil filters are inappropriate for higher amounts of dust, particularly, if the dust can settle inside the tubes and form agglomerates that reach the pump. In order to prevent damage to the pump, different types of dust filters are used depending on the specific application and pump size.

Smaller pumps often contain cotton filters in the tubes. They feature large areas in order to keep throttling loss low. Sieving filters (approximately 25 µm mesh size) or paper filters as used in air filters for large motors can be applied in pumping processes with lowest working pressures in the area of approximately 1 kPa (e.g., in packaging industry). In this pressure range, throttling is not a key issue and the tightness of the filters is sufficient.

Below 1 kPa, filters have prevailed that work according to a double principle (Figure 7.73). At 1, the air enters tangentially whereby the outer cylindrical housing 2 works as a cyclone separator. After the coarser contaminants

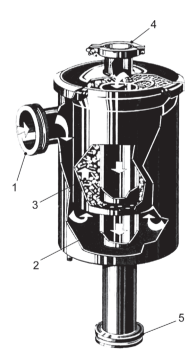


Figure 7.73 Dust filter connecting flange to the vacuum system. 1: intake flange; 2: cyclone separator; 3: oil-drenched fine separator. Optional pump connectors. 4: small flange; 5: integral flange; overpressure valve below.

are separated here, the air enters into the inner housing, which contains oilwetted Raschig rings 3. The separation rate amounts to approximately 99.9% for particle sizes up to 10 um, and approximately 99.8% for particle sizes of 2 μm. Throttling caused by the filter can be neglected down to 1 kPa, at 0.1 kPa it reaches 10%, and 50% of pumping speed of the corresponding pump at 10 Pa. A problem that frequently arises here is that vapors are pumped down that can harden inside the pump (e.g., plastic vapors). Then, frequent (often daily or even hourly) oil exchange is required because hardening in the inside of the pump is a serious threat. The pump might even stall or fail to start the next day (after downtime). Activated carbon filters can keep these vapors away from the pump very effectively. The activated carbon holds off the vapor by adsorption.

References

- 1 Pneurop (1983) Vakuumpumpen, Abnahmeregeln, Teil 1, Maschinenbau-Verlag GmbH, Ffm. 1979, and DIN 28426, part 1.
- 2 Troup, A.P. and Turell, D. (1989) Dry pump operating under harsh conditions in the semiconductor industry. J. Vac. Sci. Technol., A7, 2381-2386.
- 3 Bez, E. and Guarnaccia, D.G. (1990) Operational experience with totally oil-free rough vacuum pumps. Vacuum, 41 (7-9), 1819-1821.
- 4 Hablanian, M., Bez, E., and Farrand, J.L. (1987) Elimination of backstreaming from mechanical vacuum pumps. J. Vac. Sci. Technol., A5, 2612-2615.
- 5 Hablanian, M.H. (1988) The emerging technologies of oil-free vacuum pumps. J. Vac. Sci. Technol., A6, 1177-1182.
- 6 Bahnen, R. (1998) Völlig ölfreie neue Vorvakuumpumpe für saubere Anwendungen. Vakuum in Forschung und Praxis, 4, 279-283.
- 7 Becker, E. (1997) Membranpumpen Mit Mechanischem Membranantrieb für Gase. Vulkan Verlag, Essen, ISBN 3 8027-2184-5.
- 8 Bickert, P., Eckle, F.J., Lachenmann, R., and Rüster, G. (1993) Die Membranpumpe -Entwicklung und technischer Stand. Vakuum in der Praxis, 3, 165-171.
- 9 Lachenmann, R. (1988) Oil reservoir evacuation for pumping condensable vapours through oil sealed rotary vane pumps. Vacuum, 38 (8-10), 659-663.

- 10 Hauser, E. (2002) Diaphragm pumps and systems for gases, vapours, and liquids. GIT Laboratory Journal, 4, 184-185.
- 11 Möckel, H. (2001) Membranpumpen mit hoher Kompression durch tangentiale Membraneinspannung. Vakuum in Forschung und Praxis, 5, 273-275.
- 12 VACUUBRAND, Neue Membranpumpen MD 12, MV 10 und MV 10 VARIO, Company information, 7/2002.
- 13 Eckle, F.J. (1998) Diaphragm Pumps, in Handbook of Vacuum Science and Technology, Academic Press, London, ISBN 0-12-325065-7 pp. 84-96.
- 14 Eckle, F.J., Bickert, P., Lachenmann, R., and Wortmann, B. (1996) Pumping speed of diaphragm pumps for various gases. Vacuum, 47 (6-8), 799-801.
- 15 Lachenmann, R. (2001) The influence of backing turbomolecular pumps on the hydrogen partial pressure in high vacuum systems. Zprarodaj CVS (Bulletin of the Czech Vacuum Society), 10 (3/4), 2-7.
- 16 Lachenmann, R. (1989) Vakuumerzeugung im chemischen Labor. Labor, 8, 20-25 and 9 (1989), 108-115.
- 17 Eckle, F.J., Jorisch, W., and Lachenmann, R. (1991) Vakuumtechnik im Chemielabor. Vakuum in der Praxis, 2, 126-133.
- 18 Dirscherl, J. (2000) Vakuumregelung im Chemielabor. GIT Labor-Fachzeitschrift, 6, 750-752.
- 19 Dirscherl, J., Lachenmann, R., and Dunin von Przychowski, M. (2002) Schnellere

- Verdampfung mit geringerem Bedienaufwand durch neue Algorithmen. GIT Labor-Fachzeitschrift, 6, 711-713.
- 20 Breitenbach, J. and Lachenmann, R. (2000) Diaphragm pumps clean up high vacuums. Vacuum Solutions, 7/8, 41-46.
- 21 Breitenbach, J.B., Bickert, P., Eckle, F.J., and Lachenmann, R. (1998) Weitbereichturbomolekularpumpen und Membranvorvakuumpumpen als trockene HV- und UHV-pumpsysteme. Vakuum in Forschung und Praxis, 4, 307-312.
- 22 Lachenmann, R. and Dirscherl, J. (2004) Advanced performance of small diaphragm vacuum pumps through the use of mechatronics. Appl. Phys., A78, 671 - 673.
- 23 Eckle, F.J., Blösl, S., Lachenmann, R., and Rüster, G. Oil-free vacuum by combining diaphragm and dry rotary pumps, Le Vide, Supplement au no. 252, 5/6/7 (1990), 146-148.
- 24 Eckle, F.J., Bickert, P., and Lachenmann, R. (1995) Rotary vane and roots pumps backed by diaphragm pumps - progress in corrosive applications and clean vacuum requirements. Vacuum, 46 (8-10), 793-796.
- 25 Dirscherl, J. (2000) Chemie-HYBRID-Pumpen setzen sich durch. GIT Labor-Fachzeitschrift, 10, 1166-1168.
- 26 Faragallah, W.H. (1989) Flüssigkeitsring-Vakuumpumpen und -Kompressoren, Vulkan-Verlag, Essen.
- 27 Bannwarth, H. (1991) Flüssigkeitsring-Vakuumpumpen-Kompressoren und -Anlagen: Konventionell und Hermetisch, 2nd edn, VCH Verlagsgesellschaft mbH, Weinheim, Weinheim 1994.
- 28 Segebrecht, U. (1993) Flüssigkeitsringvakuumpumpen und Flüssigkeitsringkompressoren: Technik und Anwendung, Verlag Moderne Industrie, Landsberg/Lech.
- 29 Jünemann, A. (1994) Grundlagen zur Auslegung von Flüssigkeitsring-Vakuumpumpen und -Kompressoren: Technische Information, Sterling SIHI GmbH, Itzehoe.
- 30 Teifke, J. (1995) Flüssigkeitsring-Vakuumpumpen als verfahrenstechnische Maschine, Fachhochschule Westküste, Heide.

- 31 Faragullah, W.H. (1985) Flüssigkeitsring-Vakuumpumpen und -Kompressoren, 2nd edn (Self-published Pf 2502 D-6231 Schwalbach/Ts).
- 32 DIN 28431: Vacuum technology; acceptance specifications for liquid ring vacuum pumps.
- 33 Gaede, W. (1947) Gasballastpumpen. Zeitschrift für Naturforschung, 2a, 233-238.
- 34 Wutz, M. (1975) Vakuumpumpen nach dem Kreiskolbenprinzip. VDI-Z., 117 (6), 271 - 281.
- 35 Sawada et al. (1999) Experimental verification of theory for the pumping mechanism of a dry-scroll vacuum pump. Vacuum, 53, 233-237.
- 36 Rinder, L. (1979) Schraubenverdichter, Springer, Vienna, New York.
- 37 Rinder, L. and Grafinger, M. (2002) Entwicklung, Beurteilung und Optimierung von Rotorprofilen mir der Profilsteigungsfunktion, Schraubenmaschinen 2002, VDI-Berichte 1715, VDI-Verlag, Düsseldorf.
- 38 Kauder, K. and Wenderott, D. (1998) Spaltproblematik in Schraubenspindel-Vakuumpumpen, Schraubenmaschinen 1998, VDI-Berichte 1391 VDI-Verlag, Düsseldorf.
- 39 Sachs, R. (2002) Experimentelle Untersuchungen von Gasströmungen in Schraubenmaschinen, doctoral thesis, Dortmund.
- 40 Wenderott, D. (2001) Spaltströmungen im Vakuum, doctoral thesis, Dortmund.
- 41 Friedrichsen, U. (2001) Explosive Medien fördern, Chemie Anlagen + Verfahrenstechnik, No. 3.
- 42 Northey, A.J. (1929) patent specifications CAN 286 637 and CAN 353 007 (1935).
- 43 Arnegard, B. et al. (1970) patent specifications DE 20 29 831-833.
- 44 Kriehn, H. et al. (1996) patent specification DE 196 29 174.
- 45 Busch GmbH, Dr.-Ing.K. product brochures Mink 1080-1250 AV (Northey profile), Mink 1122-1502 BV (true profile). http://www.buschvacuum.com/ de/en/products/mink, 2016.
- 46 Wycliffe, H. (1981) patent specification DE 31 47 824.

- 47 Busch GmbH, Dr.-Ing.K. (1990) patent specification DE 40 38 704.
- 48 C. M., vanAtta. (1956) Theory and performance characteristics of a positive displacement rotary compressor as a mechanical booster vacuum pump, Trans. Nat. Sym. Vacuum Technology.
- 49 Hamacher, H. (1970) Beitrag zur Berechnung des Saugvermögens von Rootspumpen. Vakuum-Technik, 19, 215 - 221.
- 50 Hamacher, H. (1969) Kennfeld-Berechnung für Rootspumpen. DLR FB
- 51 Lorenz, A. and Armbruster, I.V. (1958) Das maximale Kompressionsverhältnis und der volumetrische Wirkungsgrad von Vakuumpumpen nach dem Rootsprinzip. Vakuumtechnik, 7, 81–85.
- 52 Bormuth, Ph. (1961) Ermittlung der temperaturerhöhung in roots-gebläsen. Konstruktion, 13, 21-23.
- 53 Janicki, M. and Schiller, D. (2011) Abpumpverhalten von Roots- und Schraubenvakuumpumpen. Vakuum in Forschung und Praxis (VIP), 23 (4), 6-11.
- 54 Laurenson, L. (1982) Technology and application of pumping fluids. J. Vac. Sci. Technol., 20 (4), 989-995.
- 55 O'Hanlon, J.E. (1984) Vacuum pump fluids. J. Vac. Sci. Technol., A2, 174-181.
- 56 Lang, H.G. (1990) Betriebsmittel für Vakuumpumpen in der chemischen Industrie und in der Kunststoffindustrie, in Handbuch Verdichter, part II Vakuumpumpen, Vulkan-Verlag, Essen, pp. 418-423.
- **57** Jorisch, W. and Oswald, D. (1987) Mechanische Vakuumpumpen und -Pumpsysteme als maßgeschneiderte Vakuumeinheiten in der Chemie (3 pages). Chemie-Technik Issue 12.
- 58 Jorisch, W. and Moll, J. (1990) Vakuum-Pumpen und Vakuum-Pumpsätze für die industrielle Vakuumtechnik, in Handbuch Verdichter, part II; Vakuumpumpen, Vulkan-Verlag, Essen, pp. 402-409.
- 59 Pujol, E. (1990) Vakuumtechnik im Einsatz bei chemischen Prozessen. Handbuch Verdichter, part II Vakuumpumpen, Vulkan-Verlag, Essen, pp. 424-430.

- 60 Duval, P. (1980) Using mechanical vacuum pumps for L.P., CVD, plasma etching and reactive ion etching, Proc. 8th Intern. Vac. Congr. Cannes, Suppl. Rev. Le Vide No. 201, Vol. 2, pp. 26-29.
- 61 Duval, P. (1983) Pumping chlorinated gases in plasma etching. J. Vac. Sci. Technol., A1 (2), 233-236.
- 62 Valente, M. (1985) Perfluoropolyether vacuum fluids for safety in semiconductor processing. *Vacuum*, **35**, 511–512 (abstract).
- 63 Connock, P., Devaney, A., and Ctirrington, L. (1981) Vacuum pumping of aggressive and dust laden vapors. J. Vac. Sci. Technol., 18 (3), 1033-1036.
- 64 Dennis, N.T.M., Budgen, L.L., and Laurenson, L. (1981) Mechanical boosters on clean or corrosive applications. J. Vac. Sci. Technol., 18 (3), 1030-1032.
- 65 Fischer, K., Henning, J., Abbel, K., and Lotz, H. (1981) Pumping of corrosive or hazardous gases with turbomolecular and oil-filled rotary vane backing pumps. J. Vac. Sci. Technol., 18 (3), 1026-1029.
- 66 Carrington, L. et al. (1982) Mechanical vacuum pumping equipment involving corrosive and aggressive materials. J. Vac. Sci. Technol., 20 (4), 1019-1082.
- 67 Berges, H.P. et al. (1980) Increased life and reliability of rotary vane pumps by using process fitting accessories for pumping aggressive gases. Proc. 8th Intern. Vac. Congress, Cannes, Suppl. Rev. Le Vide. No. 201. Vol. 2, pp. 30-33.
- 68 Fischer, K. et al. (1982) Pumping of corrosive or hazardous gases with turbomolecular and oil-filled rotary vane backing pumps. Vacuum, 32 (10/11), 619-621.
- 69 Bachmann, P. and Berges, H.-P. (1987) Sicherheitsaspekte beim Einsatz ölgedichteter drehschiebervakuumpumpen in CVD-Anwendungen. Vakuum-Technik, 36, 41-47.
- 70 Whikman, Chr.B. (1987) Reclamation of vacuum pump fluids. J. Vac. Sci. Technol., A5, 255-261.
- 71 Fulker, M. 1 (1968) Backstreaming from rotary pumps. Vacuum, 18 (8), 445-449.

- 72 Harris, N.S. (1978) Rotary pump backmigration. Vacuum, 28 (6/7), 261-268.
- 73 Laurenson, L. et al. (1988) Rotary pump backstreaming; an analytical appraisal of practical results and the factors affecting them. J. Vac. Sci. Technol., A6, 238-242.
- 74 Baker, M.A., Holland, L., and Stanton, D.A.G. (1972) The design of rotary pumps and systems to provide clean Vacua. J. Vac. Sci. Technol., 9 (1), 412-415.
- 75 Chew, A.D. (1999) The measurement of helium retention in forepumps. Vacuum, **53**, 243-246.
- 76 Baratti, A. (1990) Absaugen von feuchten und korrosiven Gasen mit mechanischen Vakuumpumpen, in Handbuch "Verdichter" part II Vakuumpumpen, Vulkan-Verlag, Essen, pp. 379-383.
- 77 van Atta, C.M. (1965) Vacuum Science and Engineering, McGraw-Hill Bock Company, New York, pp. 184ff.
- 78 DIN EN 13463-1 (April 2002).
- 79 Redeker, T. (2001) Die neuen europäischen Richtlinien 94/9/EG (ATEX 95) and 99/92/EG (ATEX 137) zum Explosionsschutz – Bedeutung für Hersteller und Betreiber. PTB-Bericht ThEx-20, Braunschweig, August.
- 80 DIN EN 12874 (April 2001).

Further Reading on Positive Displacement Pumps

- Adam, R.W. and DahmIos, C. (1980) Vakuumpumpen in der chemischen Industrie. Flüssigkeitsring-Vakuumpumpen/B. Vakuum-Technik, 29 (5), 141-148.
- Bartels, D. (1980) Vakuumpumpen in der chemischen Industrie. Flüssigkeitsring-Vakuumpumpen/A. VakuumTechnik, 29 (5), 131-140.
- Berges, H.P. and Goetz, D. (1988) Oil-free vacuum pumps of compact design. Vacuum, 38, 761-763.
- Berger, H.P. and Kuhn, M. Handling of particles in fore vacuum.
- DIN 28400 Vacuum technology; terms and definitions. part 1: general terms, part 2: vacuum pumps, Beuth, Berlin.

- Duval, P. and Long, J. (1983) Water vapor pumping with vane pumps. A critic of the PN-F-UROP method. Proc. 9. Intern. Vac. Congr., Madrid p. 89.
- Duval, P. (1989) Selection criteria for oil-free vacuum pumps. J. Vac. Sci. Tech., A7, 2369-2372.
- Grabow, G. (2002) Optimalbereiche von Fluidenergiemaschinen – Pumpen und Verdichter. Forschung im Ingenieurwesen, 7, 100-106.
- Jorisch, W. (1999) Vakuumtechnik in der Chemischen Industrie, Wiley-VCH, Weinheim, pp. 82–88.
- Knobloch, D. (1984) Verdrängerpumpen I. VDI-Bildungswerk, Handbuch Vakuumtechnik BW41-01-36, Stuttgart.
- Knobloch, D. (1989) Wirkungsweise und Eigenschaften von Verdrängerpumpen – Verschiedene Bauarten für eine Aufgabe Technische Rundschau No. 36/1989, pp. 148-156.
- Laurenson, L. and Turell, D. (1988) The performance of a multistage dry pump operating under non-standard conditions. Vacuum, 38, 665-668.
- Mathy, C. (1980) Energy saving in industrial vacuum by the use of liquid ring machines. Proc. 8th International Vac. Congress, Cannes Suppl. Rev. Le Vide, No. 295, Vol. 2, pp. 38-48.
- Mirgel, K.H. (1980) Vane-type pump with fresh oil lubrication and 100°C-technique saves energy and avoids pollution. Proc 8th Internat. Vac. Congr., Cannes Suppl. Rev. Le Vide, No. 201, Vol. 2, pp. 49-52.
- Powle, U.W. and Kar, S. (1983) Investigations on pumping speed and compression work of liquid ring vacuum pumps. Vacuum, 33 (5), 255-263.
- Reyländer, H. (1958) Über die wasserdampfverträglichkeit von Gasballastpumpen. Vakuum-Technik, 7, 78-81.
- Segebrecht, U. (1990) Förderung von trockener Luft etc. in Handbuch Verdichter, part II Vakuumpumpen, Vulkan-Verlag, Essen, pp. 356-363.
- Teifke, W. and Bohnet, M. (1990) Vakuumpumpen in der Verfahrenstechnik. in Handbuch Verdichter, part II Vakuumpumpen, Vulkan-Verlag, Essen, pp. 250-260.

Thees, R. (1957) Vakuumpumpen und ihr Einsatz zum Absaugen von Dämpfen. Vakuum-Technik, 6, 160-170.

Wycliffe, H. (1987) Mechanical high-vacuum pumps with an oil-free swept volume. J. Vac. Sci. Technol., A5, 2608-2611.

Woreg, W. et al. (1988) An evaluation of the composition of the residual gas atmosphere above a commercial dry pump. J. Vac. Sci. Technol., A6, 1183-1186.

8

Condensers

Dr. Harald Grave

GEA Jet Pumps GmbH, Andreas-Hofer-Str. 3, 76718, Karlsruhe, Germany

This chapter explains how condensation is used for pumping in the low-vacuum range, particularly in drying processes.

8.1 Condensation Processes Under Vacuum

8.1.1

Fundamentals

In drying and condensation processes under vacuum, the main task of the vacuum pumping system is to pump down developing gases and vapors. For vapors, condensers serve as particularly simple and economical vacuum pumps. Pumping with a condenser is limited to vapors. Therefore, a combination with vacuum pumps that initially pump down the air from the process container and, subsequently, pump the process and leakage gases is required. Figure 8.1 illustrates the basic setup: a condensation surface 2, kept at low temperature by means of a coolant flow from in to out in the x direction, is arranged in the condenser housing 1. At the entrance port 3, vapor enters and releases its condensation heat when hitting condenser plate 2 if the temperature of the condenser plate $T_{\rm K}$ is considerably below the saturation temperature T (temperature of dew point, condensation temperature) of the vapor. The coolant absorbs the condensation heat released by the liquefying vapor, and thus heats up and carries off the heat. The condensate drains off through discharge 5. The noncondensed part is removed by a vacuum pump connected to outlet 4.

The vapor pressure curve characterizes the relationship between the saturation temperature $T_{\rm s}$ and vapor pressure $p_{\rm s}$ of a substance (see Section 3.5.1). Figure 8.2 shows vapor pressure curves for water and selected solvents in the range of $-50\,^{\circ}{\rm C}$ to $+290\,^{\circ}{\rm C}$. To the right of the curve, the substance is in gaseous form. To the left of the curve, it is liquid or solid. On the curve, liquid and saturated

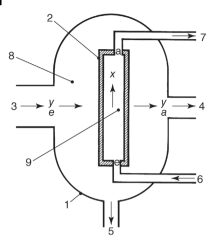


Figure 8.1 Diagram of a condenser. 1: housing; 2: condensation surface = wall of coolant channel; 3(e): incoming vapor/gas mixture; 4(a): discharge of noncondensable components of

mixture; 5: condensate drain; 6(e) and 7(a): coolant inlet and outlet, respectively; 8: vapor/gas chamber; 9: coolant channel; x: direction of coolant flow; y: direction of vapor/gas mixture flow.

vapor coexist in equilibrium. If the temperature of a nonsaturated (superheated) vapor drops, it condenses as soon as it reaches the saturation temperature (temperature of dew point) corresponding to the considered vapor pressure.

Condensation requires removal of the condensation heat, which is equal to the evaporation heat. The specific condensation heat r is temperature dependent. Water at $\theta = 25$ °C, for example, shows a specific condensation heat approximately 10% higher than at $\theta = 100$ °C. Table 8.1 lists values for selected substances at 101.3 kPa condensation pressure.

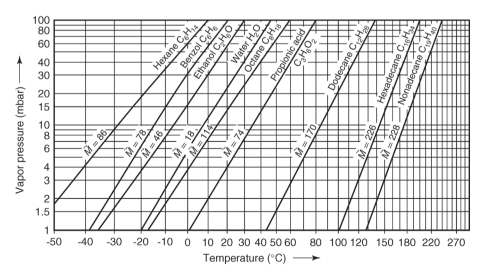


Figure 8.2 Vapor pressure curves $p_s(\vartheta)$ for selected substances.

Substance	Molar mass M (kmol kg ⁻¹)	Condensation enthalpy Δh (kJ kg ⁻¹)	
Acetone	58.08	523	
Benzole	78.11	394	
Ethanol	46.07	846	
Hexane	86.18	335	
Octane	114.23	301	
Water	18.02	2257	

Table 8.1 Molar mass M and specific condensation enthalpy Δh at pressure $p_{\rm n}=101.3$ kPa for selected substances.

If, according to the condition given by the vapor pressure curve, a vapor is not saturated but hotter, that is, superheated, removal of the additional superheat is required as well. The superheat can be calculated from the vapor's specific heat capacity (see Section 3.3.3).

Precise values of the specific heat capacity and the specific condensation heat at different temperatures can be obtained from steam tables [1,2].

8.1.2 **Condensation of Pure Vapors**

We will consider a container with a certain amount of liquid to be evaporated. A zero-mass, movable piston lies on the liquid surface as shown in Figure 8.3 (see also Figure 3.23). If, for example, the liquid is water and it is heated as illustrated in Figure 8.4, the water will evaporate at a temperature of 100 °C and the piston will move upward. Thus, the process of evaporation takes place at constant pressure (1000 mbar) and constant temperature (100 °C).

The volume between the piston and the water surface gets filled with water vapor at atmospheric pressure. Now, heat input is interrupted, the piston is fixed, and the evaporation container is placed into a cooling bath as shown in Figure 8.5 until the whole setup cools down to 20 °C. Most of the water vapor

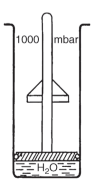


Figure 8.3 Beaker glass with zero-mass piston.

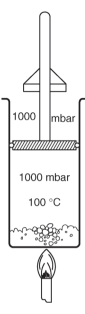


Figure 8.4 Boiling water and water vapor at atmospheric pressure.

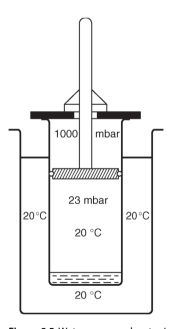


Figure 8.5 Water vapor and water in phase equilibrium.

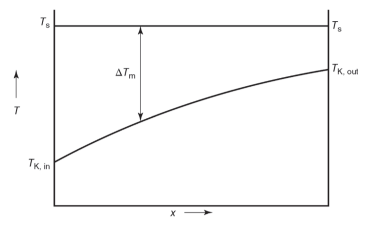


Figure 8.6 Course of temperature during condensation of pure vapor (temperature of dew point $T_s = \text{constant}; T_K : \text{coolant temperature}).$

will condense and, assuming thoroughly sealed equipment, a vacuum pressure of 23 mbar will develop. Thus, under the prevailing conditions, vacuum is produced by means of condensation (see also Section 3.5.1).

In condensers, the condensation heat released by the vapor is transferred to the coolant. Thereby, the heat flow \dot{Q} is proportional to the exchange area A, the mean temperature difference $\Delta T_{\rm m}$, and the heat transfer coefficient k:

$$\dot{Q} = A\Delta T_{\rm m}k. \tag{8.1}$$

For a representation of the mean temperature difference $\Delta T_{\rm m}$, Figure 8.6 shows the temperature distributions of vapor and coolant across condensation plate 2 in the x direction of flow (Figure 8.1). Constant vapor pressure p_s is assumed, which applies only to the condensation of pure vapor without any noncondensable fraction. Additionally, the condenser must have large cross sections in order to prevent pressure loss in the vapor flow. Under real conditions (see Section 8.1.3), the saturation temperature T_s drops along the condensation plate.

Coolant temperature $T_{K,in}$ at the entrance rises to outlet temperature $T_{K,out}$. The mass flow $\dot{m}_{\rm K}$ and the specific heat capacity $c_{\rm K}$ of the coolant determine the temperature increase, as given by

$$T_{\text{K,out}} - T_{\text{K,in}} = \frac{\dot{Q}}{\dot{m}_{\text{K}} c_{\text{K}}}.$$
 (8.2)

 $T_{\rm K}(x)$ approximately follows an exponential law. Thus, for precise calculations, the mean temperature difference in Eq. (8.1) should be calculated from the logarithmic average

$$\Delta T_{\rm m} = \frac{\Delta T_{\rm in} - \Delta T_{\rm out}}{\ln \frac{\Delta T_{\rm in}}{\Delta T_{\rm out}}}$$
(8.3)

of the temperature difference $\Delta T_{\rm in} = T_{\rm s} - T_{\rm K, in}$ at the entrance and the temperature difference $\Delta T_{\text{out}} = T_{\text{s}} - T_{\text{K,out}}$ at the outlet of the condenser.

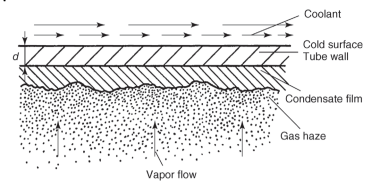


Figure 8.7 Diagram of heat transferred from condensing vapor to coolant.

For a ratio $\Delta T_{\rm in}/\Delta T_{\rm out} < 3$, the error stays below 10% if the mean temperature is calculated from the algebraic mean:

$$\Delta T_{\rm m} = \frac{\Delta T_{\rm in} - \Delta T_{\rm out}}{2}.\tag{8.4}$$

The heat transfer coefficient k is determined by the heat transmission coefficient $\alpha_{\rm C}$ on the condensation side, thickness d and thermal conductivity λ of the cooling surface walls, and the heat transmission coefficient α_K on the coolant side (Figure 8.7):

$$\frac{1}{k} = \frac{1}{\alpha_C} + \frac{d}{\lambda} + \frac{1}{\alpha_K}.\tag{8.5}$$

For typical thin sheet-metal designs, the heat transmission resistance d/λ of the cooling surface material is negligible. The heat transmission resistance $1/\alpha_K$ on the coolant side is favorably low if the velocity of flow is sufficiently high to produce turbulent flow and, additionally, to prevent fouling. For condensation of pure vapors without any noncondensable proportion, the heat transmission resistance $1/\alpha_{\rm C}$ on the condensation side is determined nearly exclusively by the thermal conductivity of the condensation film and usually shows very favorable values. Established methods for reliably calculating the heat transfer coefficient k for such conditions are available [1].

8.1.3

Condensation of Gas-Vapor Mixtures

Considering the condition illustrated in Figure 8.4, we will assume that the sealed volume contains a certain amount of air (i.e., a noncondensable gas termed inert gas in the following) in addition to water vapor prior to cooling. At a total pressure of 1000 mbar, for example, the water vapor partial pressure shall be 950 mbar. If the system is cooled down to 20 °C at constant volume, the mass of air remains unchanged. The air partial pressure drops from 50 mbar to 40 mbar due to the temperature drop from 100 °C to 20 °C. After the water vapor condenses, this air pressure adds to the water vapor partial pressure of 24 mbar so that, in this case, a vacuum pressure of 64 mbar establishes. The remaining vapor-air mixture is saturated with water vapor because the water vapor partial pressure is equal to the saturated vapor pressure of water at the considered temperature of 20 °C. Instead of water, any liquid can be investigated. The position of the vapor pressure curve determines the resulting vacuum.

Often, vacuum processes produce condensable vapors continuously. Such a vapor flow usually contains a certain proportion of inert gas, for example, the airflow penetrating the vacuum equipment due to leakage. Obviously, it should be kept as low as possible. If this mixed flow passes by an appropriately cooled surface at constant pressure, that is vacuum in the considered case, vapor condenses and the proportion of inert gas increases. A suitable vacuum pump is used to remove the inert gas-vapor mixture remaining at the end of the condensation plate. Naturally, the lower the temperature reached, the lower the vapor content in the saturated inert gas at the end of the condensation plate. Therefore, it is beneficial to pump down at the coolest part of the condenser where the coolant is supplied. For this reason, coolant flow is usually designed following the countercurrent principle with respect to the direction of the condensing vapor-inert gas mixture.

If a noncondensable gas is present, the vapor partial pressure falls short of the total pressure by the inert gas partial pressure. This means that initially the vapor cannot condense at saturation vapor temperature corresponding to the total pressure but at lower temperature (vapor pressure reduced by inert gas partial pressure means lower condensation temperature). If vapor condenses due to heat removal, the proportion of inert gas rises as the vapor condenses. Then, however, the condensation temperature drops continuously as the proportion of inert gas increases. The thermal behavior is illustrated in Figure 8.8.

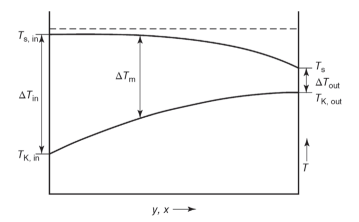


Figure 8.8 Course of temperature in a condenser for presence of inert gas. $T_{s,in}$ and $T_{s,out}$: temperatures of due points at the inlet and outlet, respectively (course of

temperature in the y direction, see Figure 8.1). $T_{K,in}$ and $T_{K,out}$: coolant temperatures at the inlet and outlet, respectively (course of temperature in the x direction).

Comparison with Figure 8.6 shows that the temperature regime here is far more unfavorable. Due to the inert gas proportion, the condensation temperature is already reduced at the entrance to the condenser. The condensation temperature drops continuously in the course of condensation and reaches its lowest value at the end of the condenser because here the proportion of inert gas in the remaining total amount reaches a maximum. As indicated, the natural aim is to reduce the vapor content as far as possible, therefore the lowest possible temperatures are desired but limited to the available coolant temperature.

Across the condenser, the temperature gradient for inert gas containing vapor is lower than that for the condensation of inert-gas-free vapor. Particularly, towards the end of the condenser (after the major proportion of heat is removed), the condensation temperature drops increasingly quickly. Therefore, it is easily accepted that the concept of countercurrent with respect to the direction of the condensing vapor flow is crucial for coolant flow. Condenser designs, thus, sometimes incorporate a special supercooling section, which forces the remaining inert gas-vapor mixture to pass through the coolest area of the condenser. Components of different geometry are used that all serve the same purpose. However, if these parts are not completely immersed in the vapor, dead zones can develop. Here, the inert gas collects and blocks the flow of vapor to the heat-exchanging plate so that parts of the surface remain unused.

However, the presence of inert gases not only reduces the mean temperature difference $\Delta T_{\rm m}$ but heat transmission as well because the heat transmission resistance $1/\alpha_C$ on the condensation side drops. The cool walls of the cooling surfaces represent vapor sinks. If vapor condenses here, the inert gas remains in front of the heat exchange surface. Vapor following must diffuse through the layer enriched with inert gas in order to reach the cold wall. Thus, the higher the proportion of inert gas, the more it inhibits heat transfer. Therefore, the lowest heat transfer occurs at the condenser outlet.

A high velocity of flow in the condensation chamber can rinse the inert gas from the heat exchange surface, which improves heat transfer, while reduced velocity impedes heat transfer. However, for constant cross-sectional area, the volume flow and, thus, the velocity of flow decrease as condensation continues. At the same time, the temperature of condensation drops. All these factors cause increasingly disadvantageous conditions towards the condenser outlet.

In order to calculate condensation to a certain extent, such condensers are modeled in succeeding steps. In some cases, a series connection of several condensers is considered, for example, a main condensation apparatus and an aftercondenser for undercooling inert gas.

Having said this, many factors such as inert gas proportion and flow velocity in the condensation volume, in particular, determine heat transition coefficients in vacuum condensers (see Table 8.2). Turbine condensers, some of which condense enormous amounts of turbine exhaust steam under vacuum, show relatively high heat transition coefficients. Here, transition coefficients reach $2500-5000 \,\mathrm{W}\,\mathrm{m}^{-2}\,\mathrm{K}^{-1}$, depending on the design and inert gas fraction. Here, of course, pure water vapor condenses so that no contamination (fouling) is to be

Condenser Type	<i>k</i> (W m ⁻² K ⁻¹)
Large condenser for pure vapor (turbine condenser)	3500
Small coiled-tube condenser for pure vapor	1200
Coiled-tube condenser with low gas portion (approximately 5% at the outlet)	800
Coiled-tube condenser with high gas portion (approximately 30% at the outlet)	400

Table 8.2 Standard values for heat transmission coefficients k in W m⁻² K⁻¹ for water vapor condensation using coolant.

expected from the vapor side. On the water side, contamination is determined by the quality of cooling water, and the corresponding fouling factors must be considered.

In condensers for vapor mixtures, for example, hydrocarbons from distillation columns in oil refineries, heat transition coefficients are in the range of 200-2500 W m⁻² K⁻¹. Apparently, precise calculations here are very difficult, keeping in mind that different components condense in different areas under varying inert gas fractions, changing velocities and directions of flow, as well as nonconstant condensation temperatures. Anticipated heavy contaminations on the vapor and water sides often compensate for an uncertain calculation because refineries, for example, specify fouling factors in the range of 0.0004 m² KW⁻¹ for the vapor side and about 0.0002 m² KW⁻¹ for the cooling water side. This corresponds to heat transition coefficients of 2500 and 5000 W m⁻² K⁻¹, and can mean a condensation area twice or three times the size of a completely clean condenser. Thus, faulty calculation of heat transition coefficients on the vapor side only changes cleaning intervals. However, systems, as delivered, always provide sufficient reserve for smooth operation. When purchasing a system, offers should be examined with respect to the calculated heat exchange surface as this, naturally, determines the price.

8.1.4 Coolants

The predominantly used cooling fluid for condensers is water. Due to fresh water costs, using recooled water is becoming increasingly popular. While seasonal variations in fresh water temperature usually are low, the inlet temperature of recooled water can rise to $\vartheta_{\text{C.in}} > 25\,^{\circ}\text{C}$ during the summer. Too high cooling water temperature, however, might cause excess of the vapor tolerance in the downstream vacuum pump (see Section 8.3.1).

Certain special applications use other types of coolants. For condensing substances with high boiling points and for separating vapor mixtures by partial condensation, coolant temperatures are increased, for example, by using warm water. In order to obtain particularly low vapor pressure or recover solvents, refrigerated water just above the melting point (e.g., $\theta_{\rm C} \approx 1$ °C) is used. Lowtemperature cooled brine (e.g., $\vartheta_{\rm C} \approx -35\,^{\circ}{\rm C}$) is used for cooling ice condensers, condensing solvents, and for operating protective condensers in front of vacuum pumps. Ice condensers, condensers for solvents and for complete condensation of high-value substances as well as condensers for protecting vacuum pumps also use directly injected and evaporating coolants ($\theta_{\rm C} \approx -20$ °C to -100 °C).

All of the special tasks that differ from standard applications in terms of condensation conditions on the vapor and coolant side require separate calculations taking into account these special conditions.

8.2 Condenser Designs

8.2.1 Surface Condensers for Liquid Condensation

Vacuum technology usually uses surface condensers for liquid condensation. Often, tubes provide the condensation surface and coolant (commonly, cooling water) flows through the tubes at velocities of 0.4-2 m s⁻¹. Generally, the vapor flows around the pipes because here wide cross sections for the large volumes of low-pressure vapor are simple to design. As an example of such a condenser, Figure 8.9 shows a coiled-tube condenser, which is widely used in vacuum technology for condensation surfaces up to several square meters. It not only provides high cooling water velocities but also high vapor cross sections and is very air-tight even for longer periods of operation because of the low number of connections.

Manufacturing tube bundle condensers with large condensation areas is simple. They utilize the condenser volume very well, permit high coolant flow rates

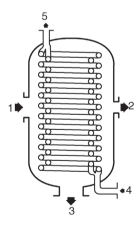


Figure 8.9 Coiled-tube condenser. 1: vapor inlet; 2: residual gas discharge; 3: condensate discharge; 4: coolant inlet; 5: coolant outlet.



Figure 8.10 Tube bundle surface condenser with fixed tube bundle. Coolant flows through the tubes, while vapor flows inside the housing in between the tubes.

due to tubes connected in parallel, and provide easy accessibility for cleaning the inside of the tubes after the heads are removed. Figure 8.10 shows a horizontal tube bundle condenser with a fixed tube bundle in which condensation takes place around the tubes and the coolant flows through the tubes. One disadvantage of this design is that it is not possible to clean the inside of the condensation chamber.

Figure 8.11 also illustrates a surface condenser with a fixed vertical tube bundle. Here, however, the vapor condenses on the inside of the tubes to allow for easier cleaning. The cooling water flows around the tubes. Obviously, cleaning is difficult on the cooling water side. In this design, vapor enters the tubes at high velocity, which guarantees good distribution and a thin film condensate. Yet, it must be paid attention not to reach the sonic velocity at the tube entrance. Additionally, deflectors increase the velocity of coolant flow to increase heat transfer.

Also, floating head designs and pull-out tube bundles (Figure 8.12) are available with cleanable outer tube surfaces. They are common to the chemical and petrochemical industry. The design provides easy cleaning on the vapor and coolant sides.

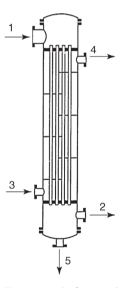


Figure 8.11 Surface condenser with fixed tube bundle. Condensation inside the tubes and cooling water around the tubes. 1: vapor inlet; 2: residual gas outlet; 3: coolant inlet; 4: coolant outlet; 5: condensate discharge.

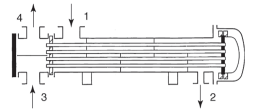


Figure 8.12 Floating head condenser with pull-out tube bundle. 1: vapor inlet; 2: residual gas and condensate discharge; 3: coolant inlet; 4: coolant outlet.

A special design of surface condensers is found in thin film evaporators for molecular distillation (Figure 8.13). Here, vapors of high molecular weight come into contact with the tube bundle from all sides and displace noncondensable parts through the tubes. A vacuum pump attached to the inside of the tube bundle then removes this proportion.

A large variety of additional surface condenser designs is available including U-tube condensers, block condensers, and air condensers, all of which are also used as vacuum condensers.

8.2.2 Direct Contact Condensers

Coolant and the vapor to be condensed mix thoroughly in direct contact condensers (see also Section 9.3.3). Figure 8.14 shows different designs of direct contact condensers. Frequently, they include cascades for coolant distribution,

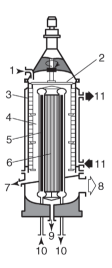


Figure 8.13 Molecular distillation column. 1: product intake; 2: valve equipment; 3: evaporator surface; 4: vapor chamber; 5: tubular condenser; 6: residual gas chamber; 7: residue

discharge; 8: residual gas discharge; 9: distillate discharge; 10: coolant inlet and outlet; 11: warm-liquid cycle heating.

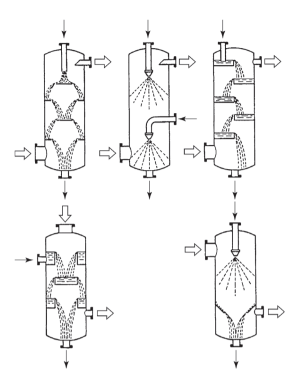


Figure 8.14 Selected designs of direct contact condensers.

that is, for providing a sufficiently large liquid surface. In the so-called spray condensers, nozzles distribute the water.

Direct contact condensers are usually designed as counterflow condensers. In certain cases, however, cocurrent flow principles are used. Cocurrent flow principles can be advantageous if, for example, intense undercooling of the inert gas-vapor mixture is not tolerable because components of the pumped-down product could precipitate. In cocurrent condensers, the vapor flows from top to bottom under gravity and in the same direction as the coolant. Thus, higher velocities of flow are acceptable and smaller cross sections are sufficient.

In cocurrent condensers, the pumped-down mixture is exposed to warmed coolant towards the end of the condenser. Therefore, the outlet temperature is higher than in counterflow condensers and a larger proportion of vapor must be removed. Occasionally, a cocurrent condenser used for the main condensation is combined with a downstream countercurrent, direct contact condenser for undercooling the exhaust mixture.

Advantages of direct contact condensers are low purchase costs and best possible utilization of the coolant. In direct contact condensers, the coolant can be heated nearly to the boiling point of the condensing vapor without requiring a particularly large condenser. This is because heat transmission coefficients are very high in the direct heat transmission occurring in such condensers. This also means less cooling water consumption compared to surface condensers. The greatest disadvantage of direct contact condensers, however, is that the vapor condensate mixes with the coolant, which is permitted only if the condensate is harmless and further use is not intended.

A main advantage of direct contact condensers is the insensitivity to fouling. Sometimes mixing of vapor condensate and cooling water is prohibited, and at the same time, surface condensation is not applicable due to the danger of fouling. Then, occasionally, direct contact condensers use the vapor condensate itself as coolant (which often dissolves the contaminants).

A surface heat exchanger cools this condensate using cooling water and circulates it back to the direct contact condenser. The overflow from this circulation contains the dissolved contaminants and can be fed back to the process after undergoing appropriate treatment.

The insensitivity of direct contact condensers to fouling is due to relatively large amounts of liquid flowing around and rinsing the walls and other components. Appropriate designs of direct contact condensers improve this property by preventing dry areas on the inside. Experience shows that fouling and incrustation arise mainly in areas not wetted by liquids.

8.2.3

Condensate Discharge

In vacuum systems, condensate discharge describes the removal of condensate from the vacuum system produced during vacuum condensation. For small amounts of condensate and in batch processing, condensate discharge is discontinuous. A large collecting tank is the simplest type of condensate collector, which takes up a certain amount of condensate in a batch and subsequently discharges it. The volume of condensate can be measured. A float switch is used for safety reasons. It triggers an alarm as soon as the condensate reaches its maximum level. Discharging the condensate during vacuum operation requires three valves: a stop valve between the condenser and the collecting tank, a ventilation valve, and a drain valve. Figure 8.15 shows a reliable design. The collecting tank is re-evacuated after draining the condensate. Here, care should be taken that the amount of air flowing back into the vacuum system is kept low.

In continuous processes developing larger amounts of condensate, appropriate pumps are used for discharging. If the flow of condensate is constant, the pump is adjusted to the required pumping level and an intermediate buffering tank is inserted into the intake line. For discontinuously developing condensate, the pump must be designed for the peak demand. A control valve, which is inserted into a bypass line and actuated by a liquid level indicator in the buffering tank, compensates for fluctuations in condensate flow.

Barometric discharge is a very common type of condensate discharge. If water is the condensate, a difference in elevation of at least 10 m is required (see Figure 9.9). This column of water corresponds to 1000 mbar atmospheric pressure so that the condensate can drain into a collecting tank from the vacuum

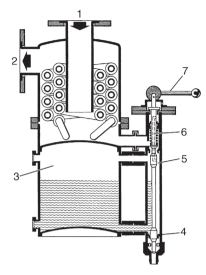


Figure 8.15 Rapid condensate discharge. 1: vapor inlet; 2: residual gas discharge; 3: condensate collector; 4: straight-way condensate valve; 6: ventilation bore; 7: actuation of valve combination.

system without requiring a pump. Especially in direct contact condensers, which require discharge not only of condensate but also of approximately 60 times as much cooling water in addition, this type of discharge has proven successful.

8.2.4 **Surface Condensers for Solid Condensation**

If vapors develop below the triple point in a vacuum process, condensation of these vapors without previous pressure increase produces solids. For such desublimation of vapors, alternating condensers are used, one of which is loaded with the product while the other is sealed.

Obviously, this procedure requires appropriate coolant, usually brine or evaporating refrigerant from a refrigerating plant. In freeze drying, for example, ice condensers (Figure 8.16) are used because of water vapor developing from frozen

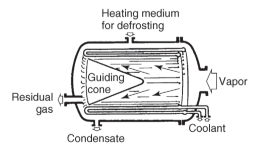


Figure 8.16 Ice condenser.

goods, which is removed at temperatures of approximately -20 °C and condenses as ice at a pressure of about 1 mbar.

8.3

Integrating Condensers into Vacuum Systems

8.3.1

Condensers Combined with Vacuum Pumps

As explained in Section 8.1.3, establishing and maintaining vacuum in a condenser requires a downstream vacuum pump matched to the mass flow and pressure delivered by the condenser. Regardless of condenser design, an inert gas-vapor mixture saturated with condensed vapor must be pumped down. If only a single vapor species is present, the saturation flow rate can be calculated by

$$\dot{m}_{\rm v} = \dot{m}_{\rm i} \frac{M_{\rm v}}{M_{\rm i}} \times \frac{p_{\rm v}}{p_{\rm i}},\tag{8.6}$$

where M is the molecular weight, p is the partial pressure, and \dot{m} is the mass flow rate of the vapor phase (subscript v) and inert gas phase (subscript i), respectively. An example illustrates the impact of condensation conditions on the specifications of the vacuum pump.

Example 8.1

In a process, water vapor is to be condensed under a vacuum of 60 mbar. The corresponding saturated steam temperature is 36 °C. The vapor flow that is to be condensed contains 10 kg h⁻¹ of air. The condenser is capable of cooling down the air including the vapor content to a temperature of 30 °C. What amount of vapor-air mixture must be removed by the vacuum pump? The water vapor partial pressure of a saturated mixture at 30 °C is $p_s = 42.4$ mbar (which is equal to the saturated water vapor pressure at 30 °C). The partial pressure of the inert gas then is the difference between the total pressure and the water vapor partial pressure, that is, $p_i = 60 \text{ mbar} - 42.4 \text{ mbar} = 17.6 \text{ mbar}$.

Thus, the saturation flow rate of water vapor, calculated according to Eq. (8.6), is

$$\dot{m}_{\rm v} = 10 \,{\rm kg}\,{\rm h}^{-1} \frac{18}{29} \times \frac{42.4}{17.6} = 15 \,{\rm kg}\,{\rm h}^{-1}.$$

This means that the vacuum pump has to pump down 15 kg h^{-1} of water vapor from the container at 60 mbar, in addition to 10 kg h^{-1} of air.

Apparently, saturation flow rates under vacuum are considerable. How important sufficient cooling is for the flow emanating from a condenser becomes obvious when considering a variation of the previous example. A drop in the outlet temperature from 30 °C to 25 °C reduces the water vapor content to a mere 7 kg h^{-1} .

Naturally, not in any case conditions are as simple as in the example, considering that usually mixtures of vapors are condensed in which the individual components follow different vapor pressure curves. Components may be completely soluble, partially soluble, or insoluble. The previous example of a mixture of water and air is particularly simple also because the liquid phase contains only a single component. In a mixture including several components, the equilibrium condition, that is, the molar or mass fraction of the individual components in the gas and liquid, must be calculated from appropriate systems of equations. For the gas phase, the condition remains valid that the sum of all partial pressures is equal to the total pressure and, thus,

$$p_i = y_i p, \tag{8.7}$$

where p_i is the partial pressure of component i, y_i is the mole fraction of component i in the gas phase, and p is the total pressure. For an ideal solution, the connection to the liquid phase is given by

$$p_i = x_i \, p_{si}, \tag{8.8}$$

where x_i is the mole fraction of component i in the liquid phase and p_{si} is the vapor pressure of the pure component i. For the water vapor-air system, Eq. (8.6) is easily derived from Eqs. (8.7) and (8.8) if solubility of air in water is neglected and the molar fraction of water in the liquid phase $x_{H_2O} = 1$. If the liquid phase contains more than one component, an iterative approach is required for solving Eqs. (8.7) and (8.8).

Figure 8.17 shows an example for condenser use in an intermittently operating vacuum drying system. A vacuum pump in such a batch processing system must meet three requirements: (1) the time required for the initial evacuation

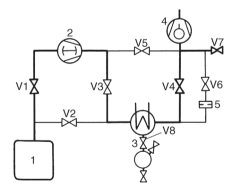


Figure 8.17 Diagram of a vacuum pumping system for batch operation. 1: vacuum chamber (e.g., drying chamber); 2: Roots pump; 3: condenser with condensate discharge; 4:

rotary plunger pump; 5: throttling aperture; V1-V8: shut-off valves; heavy lines: standard pumping system equipment.

(primary drying) has to be in reasonable relation to the total batch time, (2) the gas partial pressure during condensation must be kept sufficiently low, and (3) the pumping speed of the vacuum must be high enough for usually intended ultimate drying at low pressure, if necessary, in combination with an upstream Roots pump. Generally, the size of the pump is determined by the first or third condition.

This leads to a pump that is too large for pumping down gases from the condenser. It would additionally suck in large amounts of vapor. Therefore, throttling is advisable if recovery of condensate is desired. The condenser should be matched so that the vapor partial pressure $p_{s,out}$ at the condenser outlet does not exceed the vapor tolerance of the vacuum pump. For a vapor partial pressure $p_{\rm s,out}$ at the condenser outlet that exceeds the vapor tolerance of the vacuum pump, valve V6 can activate a throttling aperture (valve V4, connected in parallel, is then shut). The aperture increases the gas partial pressure $p_{s,out}$ at the condenser outlet.

As drying continues (ultimate drying), vapor partial pressure p_s drops below the saturation pressure corresponding to the coolant temperature $T_{K,out}$. The condensate collector is then sealed using valve V8 in order to avoid re-evaporation of the condensate. Even with the condensate collector sealed, it takes the vacuum pump approximately 1 h to pump down the liquid residue from the condenser. Therefore, the complete condenser should be sealed using valves V3 and V4, and bypassed using valve V5, for rapid drying processes.

In rare cases, vapor developing during initial drying stages (primary drying) can be too high for the downstream Roots pump, which is matched to the ultimate drying phase and usually operates continuously throughout the process. Then, an additional bypass including valve V2 is required. During this phase, valves V1 and V3 lock off the Roots pump.

In continuous processes, vacuum pumps downstream to the condenser are matched to the peaks in gas flow rates. The gases have to be pumped down at such condenser outlet gas partial pressures $p_{\rm i,out}$ that smooth condenser operation is possible. Depending on the application, gas partial pressure should be in the range of 5-50% of the total pressure. Low gas partial pressures are desired in main condensers that deposit process vapors. Intermediate condensers are intended to compress noncondensable components as far as possible to take load off downstream vacuum pumps and, thus, ask for high gas partial pressures.

Here, the condenser deposits the greater amount of the vapor, in certain cases, after precompressing the vapor. The condenser should be designed to deliver at its outlet a vapor partial pressure $p_{s,out}$ that is below the vapor tolerance of the vacuum pump under all circumstances.

Figure 8.18 shows another example of a combination of vacuum pumps and condensers: a multistage steam jet vacuum pump as used for generating vacuum in distillation columns for the chemical industry. The intake pressure is approximately 1 mbar and the gases are compressed to atmospheric pressure. However, the maximum compression ratio of a jet pump stage is insufficient for this task,

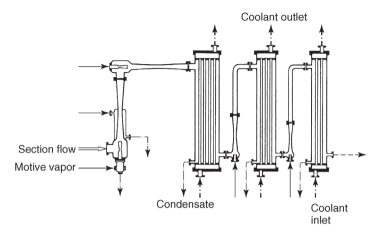


Figure 8.18 Multistage vapor jet vacuum pump with intermediate condensers (from left to right: two prestages with succeeding main condenser and two venting stages with one succeeding condenser each).

and, thus, multistage systems are required. As far as possible, condensers are arranged in the outlet of each stage in order to take load off succeeding stages, and for downstream stages to be designed smaller and to use less motive steam.

At the outlet of the first stage, the pressure is approximately 10 mbar. Therefore, common cooling water temperatures do not yet allow condensation at this point. Behind the second stage, at approximately 60 mbar, however, condensation is possible. Noncondensable substances are then exhausted together with accompanying vapors by the third jet pump stage. Succeeding condensers operate at approximately 300 mbar and atmospheric pressure.

8.3.2

Control

Several factors determine the vacuum generated in a condenser system. These include the vapor mass flow rate, inert gas fraction, condenser fouling, composition of the vapor (if more than one component is present), cooling water mass flow, and cooling water temperature. These input values are available to control the operation of a given system. Parameters such as the area of the heat exchange surface, condenser design, and the size of the vacuum pump are usually predefined and cannot be changed easily.

If control is required, for example, to guarantee constant vacuum pressure, the first step is to decide which values will change and could influence the vacuum. Then, it should be considered as to how these values can be kept constant. However, if this is impossible, an appropriate parameter is to be identified that balances variations in the input values. Often, cooling water mass flow is suitable. Generally, vacuum depends on condensation conditions. Obviously, the variable that mostly influences condensation conditions should be considered. Also, adding ballast air (gas ballast, see Section 7.6.1) changes the conditions of condensation. However, gas ballast variation is limited by the capability of the vacuum pump to pump down this additional amount of air.

8.4 Calculation Examples

Example 8.2

A drying process characterized by a pressure p = 35 mbar (which corresponds to saturation temperature $T_s = 27$ °C) develops a water vapor mass flow rate of $\dot{m}_{\rm v} = 50 \, {\rm kg \, h^{-1}}$ which is to be condensed. Considering a condensation enthalpy of 2439 kJ kg⁻¹ [2], the resulting condensation heat amounts to 33.8 kW. Fresh water with $T_{K,in} = 12$ °C is available as coolant and can be heated to $T_{K,out} = 22$ °C. Thus, according to Eq. (8.4), the mean temperature difference is

$$\Delta T_m = \frac{(27-12) + (27-22)}{2} \, \text{K} = 10 \, \text{K}.$$

For a heat transition coefficient $k = 1200 \,\mathrm{W}\,\mathrm{m}^{-2}\mathrm{K}^{-1}$ and with Eq. (8.1), it follows that the required condensation surface is

$$A = \frac{33\,800\,\text{W}}{10\,\text{K} \times 1200\,\text{W}\,\text{m}^{-2}\,\text{K}^{-1}} = 2.8\,\text{m}^2.$$

Example 8.3

Turbine condenser evacuation:

Steam turbines are equipped with large condensers that condense the working vapor expanded to pressures of 10-100 mbar in the low-pressure stage. Attached to the large condensers are vacuum systems that pump down leakage air from the equipment. These pumps are equipped with smaller condensers that initially separate a part of the vapor content in the pumped-down leakage air. We will assume a leakage airflow of $\dot{m}_i = 5 \text{ kg h}^{-1}$, pumped down at a total pressure of p = 100 mbar, and a large condenser outlet temperature of 45 °C, that is, a water vapor partial pressure of $p_s = 95.8$ mbar (according to Table A.13). Thus, inert gas

partial pressure $p_i = 100 \,\text{mbar} - 95.8 \,\text{mbar} = 4.2 \,\text{mbar}$. Then, the vapor mass flow accompanying the pumped-down airflow is given by Eq. (8.6):

$$\dot{m}_{\rm v} = 5 \,{\rm kg}\,{\rm h}^{-1} \frac{18}{29} \times \frac{95.8}{4.2} = 70.8 \,{\rm kg}\,{\rm h}^{-1}.$$

The volume flow rate at condenser input (pumping speed S) is (for a specific volume of 15.28 m³ kg⁻¹, according to Table A.13)

$$S = \dot{V} = 70.8 \text{ kg h}^{-1} \times 15.28 \text{ m}^3 \text{ kg}^{-1} = 1082 \text{ m}^3 \text{ h}^{-1}$$

The condenser uses fresh water of 12 °C that warms to 22 °C. Therefore, a condenser with a saturation temperature of $T_{s,out} = 29$ °C at the outlet can be utilized. The vapor partial pressure at the condenser outlet corresponding to this temperature is $p_{s,out} = 40$ mbar (Table A.13). With an estimated pressure loss in the condenser of $\Delta p = 10$ mbar, the air partial pressure at the vacuum pump is

$$p_{\text{i.out}} = p - \Delta p - p_{\text{s.out}} = 100 \text{ mbar} - 10 \text{ mbar} - 40 \text{ mbar} = 50 \text{ mbar}.$$

The vapor mass flow accompanying the air that is pumped down by the vacuum pump is calculated from Eq. (8.6):

$$\dot{m}_{\rm v} = 5 \text{ kg h}^{-1} \frac{18}{29} \times \frac{40}{50} = 2.5 \text{ kg h}^{-1}.$$

The volume flow rate (for a specific volume of 34.74 m³ kg⁻¹) at the condenser outlet is

$$\dot{V} = 2.5 \text{ kg h}^{-1} \times 34.74 \text{ m}^3 \text{ kg}^{-1} = 87 \text{ m}^3 \text{ h}^{-1}.$$

Of course, the installed vacuum pump will not have exactly this pumping speed because a pump will be picked from the vacuum pump manufacturer's delivery program that has a pumping speed slightly above the required speed. Thus, the pumped down vapor mass flow will increase. For a pumping speed of 140 m³ h⁻¹, for example, the air partial pressure $p_{i,out}$ will drop from 50 mbar to

$$p_{i,\text{out}} = 50 \text{ mbar} \frac{87 \text{ m}^3 \text{ h}^{-1}}{140 \text{ m}^3 \text{ h}^{-1}} = 31 \text{ mbar}.$$

The vapor partial pressure then rises to

$$p_{\text{sout}} = p - \Delta p - p_{\text{iout}} = 100 \text{ mbar} - 10 \text{ mbar} - 31 \text{ mbar} = 59 \text{ mbar}.$$

Here, the water vapor tolerance of the applied pump must be considered.

The vapor mass flow pumped down from the condenser is calculated from the volume flow rate, 140 m³ h⁻¹, and the specific volume, 24.1 m³ kg⁻¹, corresponding to the vapor partial pressure (Table A.13):

$$\dot{m}_{v,\text{out}} = \frac{140 \text{ m}^3 \text{ h}^{-1}}{24.1 \text{ m}^3 \text{ kg}^{-1}} = 5.8 \text{ kg h}^{-1}.$$

The difference between the incoming and emerging vapor mass flow rates

$$\dot{m}_{\rm v} = (70.8 - 5.8) \, {\rm kg \, h^{-1}} = 65 \, {\rm kg \, h^{-1}} = 0.01806 \, {\rm kg \, s^{-1}}$$

must be deposited in the condenser. The heat to be removed is

$$\dot{Q} = \dot{m}_{\rm v} r = 0.01806 \,\rm kg \, s^{-1} \times 2417 \,\rm kJ \, kg^{-1} = 43.7 \,\rm kW.$$

Using the above-mentioned cooling water temperatures as well as the saturation temperatures $T_{s,in} = 45$ °C (corresponding to the vapor partial pressure $p_{\rm s.in} = 95$ mbar) and $T_{\rm s.out} = 36$ °C (corresponding to $p_{\rm s.out} = 59$ mbar), Eq. (8.3) yields the mean temperature difference:

$$\Delta T_{m} = \frac{(45 - 12) - (35 - 22)}{\ln \frac{45 - 12}{35 - 22}} = 21.5 \text{ K}.$$

If a heat transition coefficient of $k = 500 \,\mathrm{Wm}^{-2} \,\mathrm{K}^{-1}$ is assumed in this example with a high air content, the condensation surface is given by Eq. (8.1):

$$A = \frac{43700 \text{ W}}{21.5 \text{ K} \times 500 \text{ W} \text{ m}^{-2} \text{ K}^{-1}} = 4.1 \text{ m}^2.$$

Furthermore, it should be considered that the leakage air mass flow of 5 kg h⁻¹ is a maximum value and the actual value in standard operation can be considerably lower. This then leads to a lower gas partial pressure at the intake to the vacuum pump and the water vapor partial pressure increases correspondingly. If neither the pumping speed of the vacuum pump is reduced according to the gas flow nor the total pressure is lowered by throttling, the water vapor tolerance of the vacuum pump might be exceeded.

References

- 1 (1997) VDI-Wärmeatlas, 8th edn, Springer, Berlin.
- 2 Schmidt, E. (1968) VDI-Wasserdampftafeln, 7th edn, Oldenbourg-Verlag, Munich.

9

Jet and Diffusion Pumps

Dr. Karl Jousten¹ and Klaus Galda²

¹Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany ²Körting Hannover AG, Badenstedter Str. 56, 30453, Hannover, Germany

This chapter describes the functionality and operating characteristics of liquid jet, vapor (steam) jet, gas jet, and diffusion pumps.

9.1 Introduction and Overview

Jet pumps (or fluid entrainment pumps) are characterized (DIN 28400, part 2) by a liquid, gas, or vapor medium (motive fluid) that travels through the pump, thereby transporting the gas to be pumped down. Figure 9.1 shows a general schematic illustration of a jet pump. The jet with velocity v_2 is produced by releasing the motive fluid from pressure p_0 in pressure chamber 1 to pressure p_2 in the jet. In the mixing chamber 3, at suction pressure p_s , the pumped-down gas mixes with the motive fluid. This causes a transmission of momentum to the pumped-down gas particles that accelerate in the expanding direction of the motive fluid until they reach the prevacuum chamber 4. Pressure p_3 here is higher than p_2 in the expanding jet so that the gas transported by the motive fluid can be released to the ambient atmosphere either directly from the prevacuum side or by means of an additional vacuum pump (prevacuum pump).

The expanding working fluid in the jet nozzle 5 as well as in the mixing chamber 3, and the pressure increase in the diffuser nozzle 6 follow *Bernoulli's* equation, Eq. (4.39), derived in Section 4.2.1 for frictionless media. Here, ν is the jet velocity, p is the static pressure in the jet, ρ is the density of the working fluid in the jet, and p_0 is the pressure in the pressure chamber ($\nu_0 = 0$):

$$\nu^2 = 2 \int_p^{p_0} \frac{\mathrm{d}p}{\rho} \,. \tag{9.1}$$

1) Translator's note: corresponds to ISO 3529.

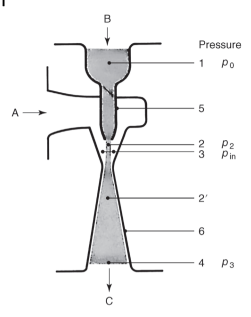


Figure 9.1 Diagram of a jet vacuum pump. 1: pressure chamber (pressure p_0); 2 and 2': motive medium jet; 3: mixing chamber (pressure p_2 in the jet); 4: compression chamber

(pressure p_3); 5: jet nozzle; 6: diffuser nozzle; B: motive medium inlet; A: vacuum connection (intake pressure $p_{\rm in}$); C: fore-vacuum connection (pressure $p_{\rm F}$).

Technical designs of fluid entrainment pumps and the detailed functionality vary considerably. A first classification is obtained by considering the physical types of motive fluids:

- liquid jet vacuum pumps;
- · gas jet vacuum pumps;
- vapor (steam) jet vacuum pumps.

An additional characteristic is working pressure range. The working pressure influences spreading of the jet in the expansion chamber and the mixing process involving pumped-down gas and motive fluid. Due to fundamentally different operating principles, we differentiate:

- 1) *jet vacuum pumps*, where intake pressure p_{in} in the mixing chamber is about equal to the static pressure p_2 in the jet, and
- 2) diffusion pumps with an intake pressure $p_{\rm in}$ far below the static pressure p_2 in the jet.

In jet pumps, the working fluid and the pumped gas preferably mix in a turbulent boundary layer of the working fluid jet. In diffusion pumps, mixing takes place due to diffusion of the pumped-down gas into the motive fluid jet.

A further important distinctive feature is the mean free path \bar{l} of gas molecules at the inlet flange to the pump. In jet pumps, \bar{l} is lower than the annular passage

between the jet nozzle and the pump wall (Figure 9.1), and thus, flow is predominantly viscous. For diffusion pumps, \bar{l} is higher than the opening passage, thus exhibiting molecular flow.

The following considerations focus on the most important pump types for vacuum technology, particularly for high-vacuum applications.

Processes in fluid entrainment pumps are complicated. Therefore, theories that usually are very simplified comply quantitatively with experimental results only in certain individual cases. The most important results that are helpful for practical use are presented.

9.2 Liquid Jet Vacuum Pumps

Many industrial applications use jet pumps with liquid working media for mixing and transporting fluids and even solids. For producing vacuum, these types of jet pumps, such as the most widely known water jet pumps, are used to deliver and compress gases and vapors by means of a liquid working fluid. Here, depending on pressure and temperature conditions, vapors can be condensed partially or completely.

On principle, any liquid can serve as working fluid. However, dimensioning requires thorough knowledge of physical properties such as density, viscosity, and boiling behavior.

Example 9.1

Calculate the jet velocity in a water jet pump (Figure 9.1). Tap water with a pressure of $p_0 = 5$ bar is released to $p_2 = 0.03$ bar in the mixing chamber. Considering the density of water to be pressure independent is a close approximation. Therefore, Bernoulli's equation (9.1) can be written as

$$v^2 = \frac{2(p_0 - p_2)}{\rho}.$$

Using the density of water, $\rho = 1000 \text{ kg m}^{-3}$, jet velocity $v_2 = 32 \text{ m s}^{-1}$.

In many cases, a closed working fluid cycle is used to prevent wasting working fluid at the outlet of a liquid jet vacuum pump and to recover it for further use. Behind the jet pump, a separator divides the working fluid from the gas, and a circulation pump feeds the liquid back as working fluid. The temperature of the circulating fluid inevitably increases in this operating mode due to the input of pump power and possible condensation of suction stream components. Therefore, a heat exchanger for recooling the liquid is required in the cycle.

The vapor pressure of the working fluid at operating temperature limits the obtainable minimum suction pressures. A water jet vacuum pump with a

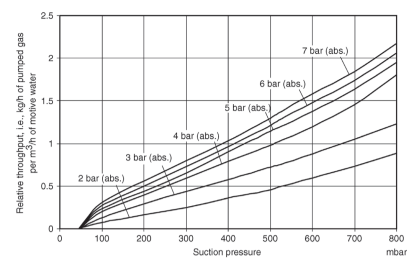


Figure 9.2 Relative suction flows of water jet vacuum ejectors for a motive water temperature of 25 °C, atmospheric discharge pressure, and motive pressures between 2 bar and 7 bar (abs.). Condensation of suction flow constituents unconsidered.

working fluid temperature of 20 °C, for example, cannot deliver an ultimate suction pressure below 23 hPa. In contrast, ultimate pressures down to 4 hPa can be obtained by using a motive fluid with negligible vapor pressure such as oil.

Counterpressure (outlet pressure) is usually atmospheric pressure although pumping against increased counterpressure is possible. The working fluid's fore pressure essentially determines the tolerable counterpressure and the corresponding motive fluid flow. The higher the motive pressure the lower the motive fluid consumption. Figure 9.2 gives motive fluid consumptions for selected motive pressures.

Intense mixing of the liquid jet and the pumped-down gas is necessary in order to obtain optimal pumping speed. This is achieved by integrating a torsion body into the jet nozzle that breaks the liquid jet at the outlet of the jet nozzle. Flow in a water jet pump created in this way is an extremely complex phenomenon. Thus, dimensioning and design until today are based solely on empirical investigations.

Because pumping speed in a liquid jet vacuum pump is much higher at high suction pressure than at ultimate pressure, this type of pump is particularly useful for startup evacuation of vacuum systems. Typical applications are:

- startup evacuation of suction lines in large centrifugal pumps;
- a variety of vacuum processes in chemical industry;
- evacuation of turbine condensers in power plants.

Liquid jet vacuum pumps are especially suitable for applications where process characteristics require mixing of pumped-down gases and a liquid because the high velocities of flow in the mixing zone of the jet pump present ideal conditions for heat and mass transfer. Due to this property, wastewater technology and drinking water conditioning increasingly utilize water jet pumps for intensively mixing air, oxygen, and ozone with the water to be treated.

Their simple design allows manufacturing jet pumps from a variety of materials. Apart from carbon steel, cast iron, bronze, and stainless steel used in standard designs, other materials such as plastics, porcelain, graphite, or glass are also applicable. Due to this circumstance, jet pumps are utilizable in applications processing aggressive and extremely corrosive media.

Due to relatively low prime costs and easy installation, standard applications use liquid jet vacuum pumps when investment costs are particularly important.

9.3 Steam Jet Vacuum Pumps

The physical principle of jet pumps has been presented in Section 4.2.8, using the Laval nozzle as example.

Jet pumps using water vapor as motive fluid (motive steam) are very important for vacuum technology and are applicable down to suction pressures of 1 Pa (0.01 mbar). The typical pressure range for the motive steam is in the range of 2-20 bar. Due to relatively simple basic geometry of the jet pump and the absence of moving parts, hardly any restrictions arise in terms of materials selection and utilization. Standard materials such as carbon steel, cast iron, bronze, stainless steel, and a variety of plastics already cover a large number of applications. However, using special materials, for example, Hastelloy, titanium, graphite, porcelain, or even glass, is possible and in fact common in jet pump fabrication.

9.3.1

Design and Function

Figure 9.3 shows the principle design of a steam jet vacuum pump.

Motive steam accelerates in jet nozzle 2 and the pressure drops, that is, pressure is transformed into velocity following Eq. (9.1). In the jet pump head 3, the suction stream B enters at the lowest pressure within the entire flow channel. In succeeding diffusers 4 and 5, the joined stream of suction flow and motive flow decelerates while the pressure rises. The joined stream finally reaches the counterpressure level at C.

Figure 9.4 illustrates pressure and velocity distributions in a steam jet vacuum pump at an overcritical pressure drop. Under this condition, the pressure of the motive medium is at least twice as high as the suction pressure. The motive medium already reaches the velocity of sound in the narrowest cross section of

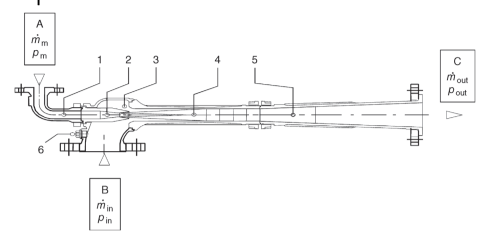


Figure 9.3 Design of a steam jet vacuum pump. 1: steam chest with steam filter; 2: motive nozzle; 3: head; 4: inlet diffuser; 5: outlet diffuser; 6: measuring port. A: motive

steam; B: suction flow; C: mixed flow; $\dot{m}_{\rm m}$: motive flow; $p_{\rm m}$: motive steam pressure; $\dot{m}_{\rm in}$: suction flow; $p_{\rm in}$: suction pressure; $\dot{m}_{\rm out}$: discharge flow; $p_{\rm out}$: discharge pressure.

the *Laval* nozzle, which widens beyond this point. The flow then continues accelerating, expands further behind the nozzle mouth, and can here reach several times the speed of sound. At this point of highest velocity and lowest pressure, mixing with the suction stream starts. In the succeeding inlet or supersonic

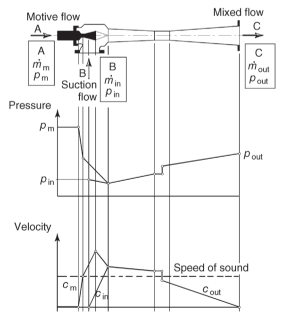


Figure 9.4 Pressure and velocity distributions in a steam jet pump.

diffuser 4, supersonic mixing proceeds as deceleration and pressure increase continue simultaneously. Velocity approaches the speed of sound and, in the narrowest diffuser cross section, changes to subsonic velocity in a compression shock. Subsequent continuing compression to counterpressure in the outlet or subsonic diffuser 5 then occurs at subsonic velocity. Stable operation of a steam jet pump is assured if the compression shock wave occurs in the narrowest diffuser cross section or further downstream. If counterpressure increases, the compression shock wave moves upstream and leads to a sudden rise in the suction pressure and, ultimately, to instable operating conditions if it reaches the inlet diffuser 4. This occurs on exceeding the so-called maximum backpressure on the discharge side. Thus, any pressure change below the maximum backpressure occurring on the mixing side has no effect on the pumping speed of the jet pump.

9.3.2 Performance Data, Operating Behavior, and Control

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The compression ratio of a single-stage steam jet ejector pump is limited to approximately 1:10 (1:20, at higher vacuum). Therefore, multistage steam jet pumps are required for suction pressures below 100 mbar. Table 9.1 lists reference values regarding appropriate stage numbers for desired suction pressures.

Section 9.3.3 covers multistage systems. First, however, the behavior of a single stage is considered. Figure 9.5 shows performance characteristics of a single stage for suction pressures between 25 hPa (25 mbar) and 40 hPa (40 mbar). Obviously, the suction flow remains unchanged for constant suction pressure and increasing backpressure as long as the maximum backpressure is not reached. If the backpressure exceeds the maximum backpressure, the suction flow suddenly decreases heavily down to zero flow. The slope of the descending characteristic curves depends on the compression ratio. For high compressions, they are very steep and cannot be used as operating points. However, this range can be used for operating (and jet pump control) in cases of lower compression ratios $(p_{\text{out}}/p_{\text{in}} < 3)$ because then the descending slope is rather gradual.

In order to illustrate the operating behavior of a single stage across a wide range of suction pressures, the maximum backpressures and the suction

pressure $7 < p_{\rm m} < 15$ in bar (abs.).	
Stage number	Intake pressure $p_{\rm in}$ in mbar (hPa)
1	100
2	30

4

0.2 0.05

Table 9.1 Stage numbers for selected suction pressures p_{in} for motive

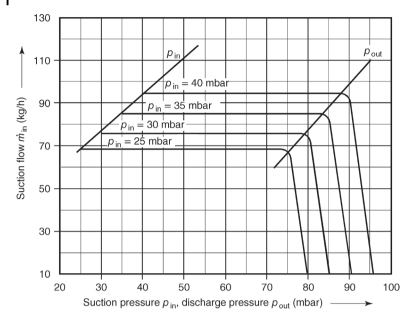


Figure 9.5 Characteristics of a single-stage ejector for selected suction pressures $p_{\rm in}$. $p_{\rm out}$: discharge pressure.

pressures are drawn as individual lines leading to the diagram in Figure 9.6. The suction flow, independent of the backpressure as long as this is below the maximum backpressure, can now be read from the suction-pressure curve for any desired suction pressure.

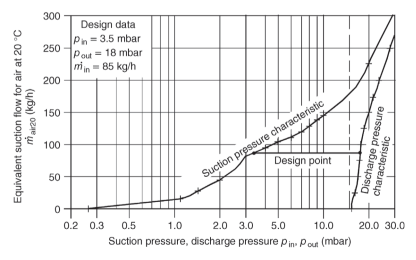


Figure 9.6 Typical characteristics of a single-stage ejector showing suction and discharge pressure curves.

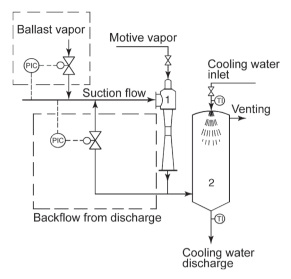


Figure 9.7 Suction-pressure control of steam jet pumps.

For determining motive steam consumption, it is necessary to define the suction flow composition, the temperature of the suction flow, the suction pressure, the maximum backpressure and the motive steam pressure. The pressures are given by the considered application. The suction stream arises at a given temperature and is usually made up of different fractions (air, vapor, hydrocarbons, etc.). A standard basis for calculation is obtained by scaling the real suction flow to a normalized suction flow, which is air at 20 °C. This conversion is performed according to VDMA standard 24924, sheet 2 [1]:

For gases,

$$\dot{m}_{\text{air }20} = \frac{\dot{m}_{\text{G}}}{\tau \vartheta_{\text{G}}},\tag{9.2}$$

and for water vapor,

$$\dot{m}_{\text{air }20} = \frac{\dot{m}_{\text{v}}}{\tau \vartheta_{\text{v}}},\tag{9.3}$$

using the conversion factors for relative molar mass M_r ,

$$\tau = \exp\{-2.44731 + (\ln M_{\rm r} \times 1.028) - [(\ln M_{\rm r})^2 \times 0.0894]\},\tag{9.4}$$

the temperature of gases

$$\vartheta_{G} = \frac{(\vartheta - 20)(0.8 - 1)}{460} + 1,\tag{9.5}$$

and the temperature of water vapor

$$\vartheta_{\rm v} = \frac{(\vartheta - 20)(0.725 - 1)}{460} + 1. \tag{9.6}$$

Example 9.2

A suction stream of 35 kg h^{-1} CO₂ ($M_r = 44$) with a temperature of 120 °C is converted according to Eq. (9.2) to

$$\dot{m}_{\text{air }20} = \frac{35}{1.176 \times 0.956} = 31.1 \text{ kg h}^{-1}$$

of air at 20 °C.

The required motive flow $\dot{m}_{\rm m}$ is calculated from the equation for the relative suction throughput:

$$\frac{1}{\mu} = \frac{\dot{m}_{\text{air }20}}{\dot{m}_{\text{m}}} = \frac{1}{c_L \frac{p_{\text{out}}}{p_{\text{in}}}}.$$
(9.7)

The dimensionless factor $c_{\rm L}$ is an empirical value that depends on suction and motive pressures. Table 9.2 lists values for compressions up to atmospheric pressure or compression ratios $p_{\rm out}/p_{\rm in} < 10$.

Options for performance control of jet pumps are very limited. Pumping capacity can be varied only in jet pumps with small compression ratios $(p_{\rm out}/p_{\rm in} < 3)$ by utilizing the descending slopes of the characteristic curves. Otherwise, matching to different pumping speeds can be obtained by connecting several pumps in parallel. For small compression ratios $(p_{\rm out}/p_{\rm in} < 3)$, suction pressure is controlled by variation of motive pressure, while for higher compression rates, it is done only by using suction-side throttling, by adding ballast suction flow, or by recirculating part of the total mixture stream back to the suction side (Figure 9.7).

Suction pressure (abs.) (mbar)	Motive steam pressure in bar (abs.)				
	4	8	12		
500	0.71	0.41	0.36		
200	1.20	0.54	0.48		
100	5.00	0.92	0.50		
010	0.29	0.26	0.22		
005	0.27	0.22	0.21		
001	0.25	0.21	0.21		
000.5	0.25	0.20	0.20		

Multistage Steam Jet Vacuum Pumps

At low suction pressures (1 hPa), very high compression ratios must be overcome in order to pump against atmospheric pressure. Here, multistage systems with intermediate condensation are used. As far as pressure and temperature conditions allow, water vapor and other condensable components are condensed from the suction and motive streams in direct contact condensers or surface condensers after each jet pump stage. Each succeeding jet pump stage then further compresses only the water vapor saturated noncondensable fractions and not the motive steam itself. Figure 9.8 shows the basic design of a five-stage system with direct contact condensers, matched to a suction pressure of 1 hPa (1 mbar, compression ratio 1000).

The first condenser operates at the lowest possible pressure, depending on the available cooling water temperature (using cooling water of 25 °C, water vapor can be condensed at approximately 40 hPa (40 mbar)). Two jet pump stages precede the first condenser (main condenser) because the necessary 40-fold compression (from 1 hPa to 40 hPa) cannot be obtained in a single stage. Behind the main condenser, three additional jet pump stages are arranged including intermediate condensation (three-stage air evacuation unit) to ultimately deliver the noncondensable gases against atmospheric pressure. A two-stage air evacuation unit could also do this work; however, motive steam consumption would be

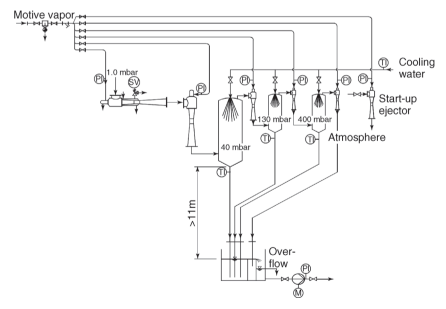


Figure 9.8 Five-stage steam jet vacuum pump with intermediate direct contact condensers and startup ejector pump (for quick evacuation of processing system during startup). Tl: thermometer; PI: pressure gauge; SV: safety valve on heating jacket; M: motor.



Figure 9.9 First two stages of a five-stage steam jet vacuum system with direct contact condenser for the deodorizing column in an edible oil refinery. The first stage (pumping top-down diagonally, nominal pumping diameter DN 500, length approximately 6 m) sucks from the deodorizing column at 3 hPa (pumping speed 66 000 m³ h⁻¹). The second stage (pumping upward from below, nominal

pumping diameter DN 400, length approximately 5 m) features counterpressure-controlled motive vapor for reducing motive vapor consumption (803–485 kg h⁻¹) at low cooling water temperatures (33 °C-23 °C). The first direct contact condenser (main condenser \emptyset 700 \times 2800 mm) is visible at the outlet of the second stage.

higher. Utilizing simple and cheap direct contact condensers (Figure 9.9) is possible only when mixing of process media and cooling water is tolerable. If this is not acceptable, surface condensers are used that feature separated process medium and cooling water.

The cooling water draining from direct contact condensers or the condensate draining from surface condensers can be realized either barometrically by means of a downpipe that is at least 11 m in length (i.e., 11 m level difference) or at lower elevation by using appropriate centrifugal pumps (e.g., side channel pumps). This is necessary because the condensers operate under vacuum. In practice, barometrical assemblies (Figure 9.9) are preferred because the complete vacuum system then operates trouble-free without any moving parts.

Typical applications for systems with direct contact (mixing) condensation:

- Edible oil refining (Figure 9.9). Process steps include bleaching, drying, neutralizing, and deodorizing or physical refining under vacuum. Nearly exclusively, multistage steam jet vacuum pumps are used here for producing vacuum.
- Steam jet refrigerating systems.

Typical applications in systems with surface condensation:

- vacuum production in mineral oil distillation (oil refineries);
- vacuum production in urea synthesis;
- turbine condenser evacuation in power plants;
- vacuum generation in evaporation plants;
- vacuum production in seawater desalinization;
- vacuum production in extruder processes.

Hybrid systems containing multistage steam jet vacuum pumps, surface condensers, and a liquid ring vacuum pump as atmospheric stage are used frequently for vacuum production in processes of

- chemical industry;
- pharmaceutical industry;
- drying technology;
- petrochemical industry.

9.3.4

Organic Vapors as Driving Pump Fluids

If jet pumps operate on water vapor (steam), wastewater contaminated with product components develops from the motive vapor condensate even if surface condensers are used. In order to avoid even such small wastewater flows, certain processes in vacuum production call for product vapor (vapor of a substance used in the process) as motive medium. By this, in addition to preventing any possible wastewater, the danger of water or vapor backflow to the process is eliminated. This represents a strict requirement in many chemical processes.

In a system operating on product vapor, the motive vapor condensate, mixed with the condensate from the vapor pumped down from the process, is reevaporated in a cycle and again used as motive vapor. Excess condensate is fed back to the process. Therefore, no waste liquid occurs that would have to be disposed of. Viewed from the outside, such a system operates like a dry compressing vacuum pump. Systems of this type utilize surface condensers as well as direct contact condensers. In the direct contact condensers, motive vapor and coolant are of the same substance.

Organic vapors such as ethylene glycol, butane diol, butanol, monochlorobenzene, trichloroethylene, toluol, phenol, and others are appropriate motive media in product-vapor-driven jet pumps.

The applicability of a substance as motive medium is determined largely by its vapor-pressure curve. It is essential that the evaporation temperature remains below the decomposition temperature and that the condensation temperature is above the triple point. The temperature of the available coolant, which determines the necessary condensation pressure, may generally prohibit the use of certain motive media. In addition, modifying the vapor pressure by adding low boiling components from the suction stream should be considered.

These considerations show that accurate investigation and, possibly, testing prior to industrial scale application are suggested when utilizing product-vapordriven jet pumps.

Today, a number of processes have been investigated thoroughly, and flawless operation has been shown in numerous large-scale installations. Further, spread to new applications is surely desired in order to combine the advantages of vapor jet pumps in terms of operating safety and low investment costs with the advantage of a wastewater-free operation.

Typical applications are:

- vacuum generation for polycondensation in PET production;
- process vapor: ethylene glycol;
- process pressure: 0.1–0.5 mbar;
- synthetic fiber production;
- process vapor: butane diol;
- process pressure: <1.0 mbar;
- vacuum production for special products;
- process vapor: butanol, monochlorobenzene, phenol.

9.4 **Diffusion Pumps**

9.4.1

Design and Principle of Operation

The concept of diffusion pumps can be traced back to an invention by Gaede [2], who was also the first to use this term. Figure 9.10 shows a cross section of a diffusion pump. The cylindrical pump body PB terminates at the top in the highvacuum inlet flange Fin. A baffle BA is attached to the upper baffle flange. The impact plates of the baffle prevent vapor from entering into the vacuum chamber. At the bottom, the pump body is sealed with a ground plate PG. It forms the heated boiling chamber BC for the pump fluid. The fore-vacuum line FV is attached to the side and contains a small flange F_{FV} for connecting the fore pump. Above the fore-vacuum line, water flows through cooling tubes CT that cool the pump body. Cooling can also be provided by a cooling jacket or, in case of air cooling, by cooling fins. The pump body holds the internal part of the pump including the nozzle system. The image shows a four-stage pump with

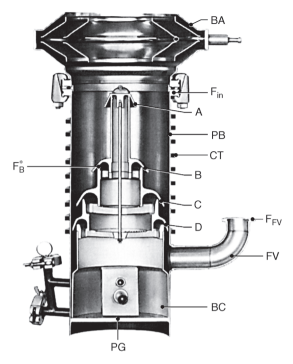


Figure 9.10 Section of a four-stage diffusion pump with attached baffle. A, B, C, D: concentric nozzles; BA: baffle; Fin: high-vacuum flange; F_{FV}: fore-vacuum flange; F_R^{*}: narrowest

cross section: CT: cooling tube (water cooling): PG: ground plate; PB: pump body; FV: forevacuum tube; BC: boiling chamber.

one high-vacuum stage (A), two medium-vacuum stages (B and C), and one forevacuum stage (D). Diffusion pumps with fewer or more stages are also available.

The operating principle is explained using Figure 9.11. A floor heater H or an immersion heater heats the pump fluid at the bottom of the pump body until a vapor pressure $p_0 = 0.1-1$ kPa develops in the boiling chamber BC. The vapor jet J moves upward inside the vapor tubes of the internal part, enters the annular nozzles A to D (see Figure 9.10) that are formed by the vapor tubes and the nozzle caps, and is deflected downward at this point. Behind the narrowest cross section (e.g., F_B^{*} in Figure 9.10), the vapor jet expands according to gas-dynamic laws and ultimately enters the chamber formed between the nozzle system and the cooled wall of the pump body. Here, expansion and velocity continue to increase.

In the volume below each nozzle cap, an umbrella-shaped vapor jet with annular cross section develops between the nozzle system and the cooled pump body wall. The jet moves downward at high supersonic velocity $(M \approx 3-8)$. The gas particles G, which are to be pumped, enter the pump from the top through the high-vacuum connection Fin and initially encounter the vapor jet of highvacuum nozzle A. They diffuse (Section 3.3.4) into the jet and accelerate downward due to impacting particles. The vapor condenses when it touches the

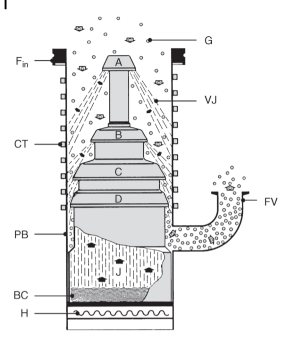


Figure 9.11 Operating principle of a diffusion pump. H: heater; BC: boiling chamber; PB: pump body; CT: cooling tubes; Fin: high-vacuum flange; G: pumped gas particles; VJ: vapor jet; FV: fore-vacuum port; A, B, C, D: nozzles; J: vapor jet.

cooled wall of the pump body PB. The noncondensable gas molecules enter the vapor jet of the intermediate stages B and C where they again accelerate and are transported to the chamber of the fore-vacuum stage D. The gas pressure increases from one stage to the next. The pressure ratio (compression ratio) for a stage can be expressed by the following equation: eject

$$\frac{p_{\text{before}}}{p_{\text{after}}} = \exp(\rho u d/D), \tag{9.8}$$

where ρ is the density of the pump fluid, u is its velocity, and d is the width of the jet. The diffusion coefficient D is approximated by using the molecular weights $M_{
m G}$ and $M_{
m M}$, and the molecular diameters $d_{
m G}$ and $d_{
m M}$ of the gas and the motive medium, respectively [3]:

$$D = \frac{3}{8\sqrt{2\pi}} \left(RT \frac{M_{\rm G} + M_{\rm M}}{M_{\rm G}M_{\rm M}} \right)^{0.5} \left(\frac{d_{\rm G} + d_{\rm M}}{2} \right)^{-2}.$$
 (9.9)

The compressed gas then enters the fore-vacuum tube FV and is pumped off by the fore-vacuum pump. The condensed motive medium drains down at the inside of the pump body until it reaches the evaporation chamber where it reevaporates in a cycle.

For constant mass flow, the volume flow rate of the pumped-down gas decreases as it travels from one stage to the next. Therefore, the inside of the pump is designed in such a way that the annular pump surface between the individual nozzle systems and the wall of the pump body decreases from one stage to the next. This has the advantage that the vapor expands less in the stages on the forevacuum side and, thus, a higher at-rest (or static) pressure ratio is obtained (see Section 4.2.7). This means a higher tolerable pressure on the fore-vacuum side.

Therefore, the first stage of a diffusion pump has the highest pumping speed and the lowest compression ratio. The opposite is the case for the last stage. Smaller diffusion pumps usually have three stages, while larger ones have up to five or six.

Proper cooling of a diffusion pump is crucial for its operation. The highest cooling demand occurs at points where the vapor jet hits the pump wall. If the cooling capacity is too low, the vapor condenses partially and may infiltrate the recipient to be evacuated (backflow). If cooling capacity is too high, the condensate cools too far and flows slower to the evaporation chamber. Thus, the maximum pumping speed of the pump drops and, additionally, unnecessarily high heating power is required for re-evaporation.

Figure 9.12 shows a plot of the pumping speed of a diffusion pump versus inlet pressure. Below a critical pressure, the pumping speed is constant because the probability for gas-molecule impact on the pump flange as well as their pumping probability in the vapor jet is pressure independent. On principle, this pumping speed remains unchanged even for arbitrarily low pressures. However, for very low pressures, the measured or apparent pumping speed drops because the compression ratio, that is, back diffusion from the fore pump, and outgassing of the pump determine the pressure in the pump. The main cause of outgassing in the pump is the backflow of pump fluid and its fugitive fractions.

Beyond the critical point (Figure 9.12) lies the area of constant particle flow: maximum throughput of the pump is reached. In the lower part of this overload

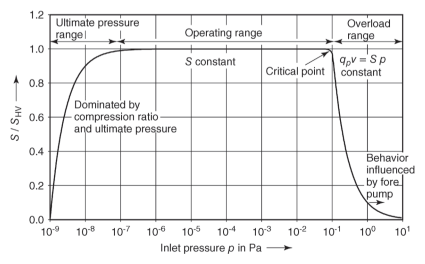


Figure 9.12 Diagram of relative pumping speed (S_{HV}: high-vacuum pumping speed) in a diffusion pump with supposed ultimate pressure of 10^{-9} Pa and the critical point at 0.1 Pa.

area, the size of the fore pump is already very important and can lead to an increase or decrease in S(p). In processes with known gas flow rates, the size of the diffusion pump is matched to obtain a pumping speed that is above this rate, and to deliver the desired process pressure.

Several factors may cause undesired pump fluid flow into the recipient (back diffusion):

- Vapor jet molecules from the top first stage accelerate toward the inlet flange due to interactions with gas particles or other motive medium molecules, or due to imperfect nozzle shape.
- Condensed vapor molecules re-evaporate and travel toward the inlet flange.
- Pump fluid oil creeps to the recipient along the walls.
- Just in front of the heater, oil droplets heat up high enough for them to accelerate toward the recipient as drops (similar to oil splattering in a frying pan).

Appropriate nozzle shapes as well as vapor traps and baffles (Section 9.4.3) minimize back diffusion. It should remain below 1×10^{-10} g cm⁻² min⁻¹. Corresponding values have in fact been measured [4]. The problem of back diffusion is also covered in [4-7].

Example 9.3

A diffusion pump with a pumping speed of $6000 \,\ell$ s⁻¹ (Figure 9.16) operates in a system at an inlet pressure p_{in} of 6×10^{-2} Pa (6×10^{-4} mbar). For this working pressure, how high is the throughput of the pump? Calculate the maximum throughput of the pump by using Figure 9.16. Which minimum pumping speed should the fore pump have if the intake pressure of the diffusion pump rises to 1 Pa and, at the same time, the fore-vacuum pressure is to remain below 30 Pa in order to remain safely below the critical backing pressure (see Section 9.4.6) of 50 Pa?

Throughput q_{pV} at 6×10^{-2} Pa is given by $q_{pV} = S p_{in} = 6000 \, \ell \, s^{-1} \times 10^{-2}$ $6 \times 10^{-2} \, \text{Pa} = 360 \, \text{Pa} \, \ell \, \text{s}^{-1}$. In the descending slope of the pumping speed curve, throughput is constant (compare Figures 9.12 and 9.16). Therefore, it is adequate to select a well-readable value: at 0.4 Pa (4 \times 10⁻³mbar), pumping speed is 2000 ℓ s⁻¹. Thus, the maximum throughput $q_{\rm max}=800\,{\rm Pa}\,\ell\,{\rm s}^{-1}$. The fore pump has to pump this flow rate. In order to reach a fore pressure p_F of 30 Pa, its pumping speed must amount to $S = q_{\text{max}}/p_F = 26.7 \, \text{f} \, \text{s}^{-1} = 96 \, \text{m}^3 \, \text{h}^{-1}$.

9.4.2

Pump Fluids

Until the early twentieth century, mercury was the only pump fluid used. However, the vapor pressure of mercury is relatively high (approximately 0.1 Pa) at cooling water temperature. Reaching high vacuum, thus, already required a low-

Table 9.3 Desired properties for pump fluids in diffusion pumps.

Property	Goal				
Low vapor pressure	Low ultimate pressure				
No volatile constituents	Low back diffusion, low ultimate pressure				
Low solubility for gases	Low ultimate pressure				
High molecular weight	Large momentum transfer, high pumping speed				
High surface tension	Reduced wall creeping (remigration)				
Appropriate viscosity at room temperature	Good flow characteristics, high pumping speed				
Low evaporation heat	Low heating power				
High flash point	Thermal stability, safety				
No decomposition in vacuum, due to electrical discharge	Long service life of oil				
No oxidation	Stability when exposed to air				
Stability against chemicals	Long service life of oil				
Nontoxic	Operational safety				

cooling trap between the pump and the vacuum container in order to reduce vapor pressure. Today, mercury diffusion pumps are used in special cases only. The most common diffusion pumps are oil diffusion pumps [8] that operate on pump fluids of high molecular weight based on refined crude oils, silicones, certain esters or fluorinated oils, for example, Fomblin (Table A.18). Figure 20.5 shows vapor pressure curves of selected pump fluids.

Desired characteristics of pump fluids are listed in Table 9.3; however, a single type of fluid cannot have all of these.

A detailed coverage of pump fluids has been provided in [9].

9.4.3 **Baffles and Vapor Traps**

Today, pump fluids of high molecular weight are available featuring extremely low vapor pressures at cooling water temperatures $(10^{-7}-10^{-9} Pa, see also$ Table A.18). Thus, low-cooling traps can be dispensable under certain conditions. However, a considerable number of oil molecules emerge upward, opposite to the pumping direction, from the vapor jet of the high-vacuum stage, particularly from the area of the upper nozzle cap. Thus, they reach the vacuum container (back diffusion) where they condense. Oil contamination, however, is undesired in most cases because oil vapor considerably disturbs nearly any vacuum process. A cooled nozzle cap baffle can trap more than 90% of the backflowing oil vapor (Figure 9.13).

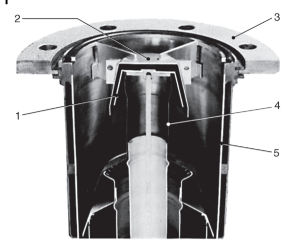


Figure 9.13 Nozzle cap baffle cooled via a heat-conducting connection to the pump body. 1: cap of top diffusion nozzle; 2: nozzle cap baffle with solid heat-conducting struts; 3: high-vacuum flange of diffusion pump; 4: ascending vapor pipe; 5: cooled pump body.

Reducing the oil vapor in the recipient down to the saturation pressure of the cooling water temperature requires a plate baffle, shell baffle, or chevron baffle refrigerated to this temperature and placed between the pump and the vacuum container (Figure 9.14). Such a vapor trap or baffle interrupts the line of sight between pump and container with a certain overlap so that each oil molecule traveling upward touches the cooled plates at least once. Deeper cooling of the baffle plates further reduces the oil vapor pressure in the vacuum chamber and is used occasionally for producing ultrahigh vacuum with diffusion pumps. If a low-cooled baffle is used at a temperature where the oil is so pasty that is does not flow back to the pump, a nozzle baffle should be inserted that condenses the larger portion of the ascending oil at cooling water temperature and feeds it back to the pump cycle. Vapor traps are usually equipped with a creep barrier that prevents the oil condensing at the walls from creeping into the recipient. A creep barrier is a thin piece of stainless sheet metal that connects the uncooled housing with the cooled shell of the baffle (Figure 9.14). The ascending, creeping oil

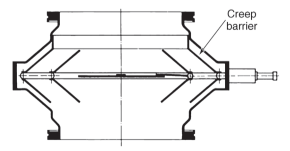


Figure 9.14 Section of a shell baffle.

film is guided across the cooled parts of the baffle and is, thus, prevented from entering the recipient.

Any vapor trap reduces the effective pumping speed S_{eff} at the vacuum chamber compared to the inherent pumping speed S of a diffusion pump without vapor trap. While $S_{\text{eff}} \approx 0.9 \, S$ in a pump with a nozzle cap vapor trap, optically tight vapor traps lead to $S_{\rm eff} \approx 0.5 \, S$.

9.4.4 Fractionating and Degassing

Pump fluids of high molecular weight are nonuniform substances and can, therefore, be fractionated. This is utilized for improving the ultimate pressure in fractionating diffusion pumps. After draining down at the wall of the pump body, the motive oil flows radially to the center of the heating chamber. Barriers divide this volume into several annular evaporation chambers (Figure 9.15).

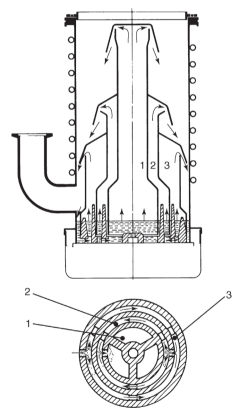


Figure 9.15 Three-stage diffusion pump with fractionating. 1: evaporation chamber of highvacuum stage; 2: evaporation chamber of intermediate stage; 3: evaporation chamber of forevacuum stage.

Initially, the oil enters the evaporation chamber of fore-vacuum stage 3. Here, mainly light components evaporate, while nonevaporating, heavier components reach the evaporation chamber of the intermediate stage 2. Here again, mostly lighter, volatile components evaporate so that only components that are most difficult to evaporate and have the lowest vapor pressure reach the evaporation chamber of the high-vacuum stage 1. Thus, the oil vapor pressure at the pump flange is lower than the oil vapor pressure in a pump without fractionating.

Due to thermal and chemical decomposition, pump fluids of high molecular weight continuously develop certain amounts of lighter fractions with considerably higher vapor pressure. These fractions do not condense in water-cooled vapor traps. Condensing these substances requires low-cooled traps. Occurrence of such light fractions can increase the ultimate vacuum of a diffusion pump by several powers of 10 compared to values which would correspond to the equilibrium vapor pressure of the actual pump fluid. Therefore, it is important to remove continuously light, volatile fractions from the oil. This is done by heating the oil film flowing down the inside walls of the pump body to 150 °C above cooling water temperature at the lower end, that is, below the fore-vacuum stage. For this, the coiled cooling pipe or other refrigerating components are designed to reach high enough, near the lower jet nozzle so that the vapor heats the downflowing condensate to the desired temperature. Volatile components then evaporate from the pump fluid flowing back to the boiling chamber and are discharged as gas through the fore-vacuum pipe together with the pumpeddown gas. Here, they cannot condense due to their high vapor pressure. Simultaneously, this degassing device removes volatile substances produced in the vacuum process from the pump oil as soon as they enter the pump. In addition, contaminants originally included in the pump fluid are eliminated.

9.4.5

Operating Suggestions

Diffusion pumps incorporating the described measures provide sufficient hydrocarbon purity for many applications when combined with matched vapor traps and high-grade pump fluids. However, cooling water interruptions, operating errors, and so on can lead to additional hydrocarbon introduction to the recipient. An automated pump system can reduce the risk of this happening. Here, unexpected events, such as cooling water interruption, power failure or recipient pressure increasing beyond certain limits, trigger appropriate actions, for example, closing a plate valve above the diffusion pump.

Switching on and off is automated as well. However, contact between a hot pump and air is restricted to pressures below several 10 Pa. This rule can be neglected if requirements concerning hydrocarbon purity are low and if the amount of air entering the hot pump is low (see Sections 18.5.3.1 and 18.5.4).

Particular emphasis in diffusion pumps is laid on shock-free, smooth evaporation and homogeneous heat transmission to the pump fluid so that excess temperatures that might cause decomposition are prevented.

Figure 9.16 shows the pumping speed S of selected diffusion pumps versus inlet pressure. A pumping speed constant across a wide pressure range is typical for diffusion pumps (compare also Figure 9.12). Pumping speed is given as nominal pumping

speed. Table 9.4 lists additional technical data of these pumps. The pumping speed depends on the gas species as well as on the heating power and the type of pump fluid.

Figure 9.17 illustrates the behavior of a diffusion pump for a desired intake pressure $p_{\rm in}$ with respect to rising pressure $p_{\rm F}$ on the fore-vacuum side. Up to a critical value p_{crit} of fore-vacuum pressure p_F , the pump is in normal operating condition, that is, the high-vacuum side is unaffected by the fore-vacuum side. At $p_{\rm F} = p_{\rm crit}$, however, the exceeding backpressure $p_{\rm F}$ destroys the supersonic vapor jet responsible for the pumping action (compare Section 9.4.7) because the jet no longer reaches the pump wall. Pumping action ceases and gas breaks through from the fore-vacuum side into the evacuated container. The critical pressure $p_{\rm F} = p_{\rm crit}$ is therefore referred to as the critical backing pressure or fore-vacuum tolerance (see Table 9.4).

The following section will explain that the shorter the distance between the jet nozzle and the pump wall the higher the fore-vacuum tolerance. Consequentially, a given value of critical backing pressure is more difficult to obtain in large diffusion pumps that generally afford more stages than smaller pumps.

Appropriate selection of fore pumps (see Example 9.3 and Section 18.5.4.1) must ensure that the critical backing pressure p_{crit} (typically several 10 Pa) is not exceeded during pump operation, particularly at high inlet pressure p_{in} . The pressure in the evaporation chamber of the pump amounts to approximately

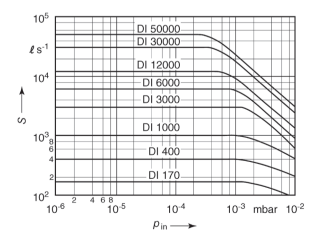


Figure 9.16 Pumping speed S versus inlet pressure p_{in} for selected oil diffusion pumps. Parameters on the curves give the nominal pumping speed S_n .

Table 9.4 Technical specifications of selected oil diffusion pumps.

Pump type	180	410	1010	3000	6000	12 000	30 000	50 000
High-vacuum connection DN	65 LF	100 LF	150 LF	250 LF	350 LF	500 LF	800 LF	1000 LF
Fore-vacuum connection DN	25 KF	25 KF	40 KF	50 KF	65 KF	100 LF	150 LF	150 LF
Pumping speed for air								
At 1 Pa (ℓ s ⁻¹)	100	200	400	600	950	1200	2400	3000
At 0.1 Pa (ℓs ⁻¹)	160	430	780	3000	6000	10 000	18 000	25 000
$<1 \text{ Pa } (\ell \text{ s}^{-1})$	180	410	1010	3000	6000	12 000	30 000	50 000
Operating range (Pa)	< 0.1	< 0.1	< 0.1	< 1	< 1	< 1	< 1	< 1
Fore-vacuum tolerance (Pa)	40	50	40	50	50	50	50	50
Motive-medium filling amount (min/max)	$30/70\mathrm{cm}^3$	$70/180\mathrm{cm}^3$	$0.1/0.5\ell$	$0.6/1.2\ell$	$1.2/2.4\ell$	$2.5/5\ell$	$8/16\ell$	$17/35\ell$
Heating power for pumping air (W)	450	800	1200	2200	3750	7500	19800	26 400
Approximate preheating time (min)	15	18	18	25	30	30	40	45
Minimum cooling-water flow (ℓh^{-1})	15	20	25	210	330	660	1200	2000
Weight (kg)	6	9	18	26	60	145	380	630
Suggested pumping speed for a roughing pun	np for inlet press	sures under conti	nuous operatior	ı				
>0.01 Pa (m ³ h ⁻¹)	8	16	30	100	200	250	500	1000
<0.01 Pa (m ³ h ⁻¹)	4	8	16	30	60	100	200	250

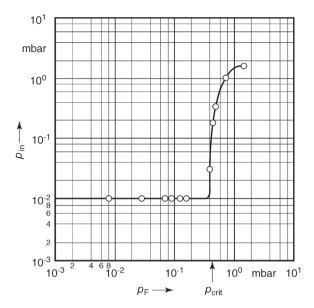


Figure 9.17 Inlet pressure p_{in} for rising fore-vacuum pressure p_{F} on the fore-vacuum side. For $p_{\rm F}=p_{\rm crit}$ (fore-vacuum tolerance), the gas breaks through from the fore vacuum into the inlet of the diffusion pump.

100–200 Pa, which means that the fore pump has to evacuate to approximately 50 Pa. As a general rule, diffusion pumps are designed so that $p_{crit} = 50 \, \text{Pa}$ as well, in order to utilize the maximum range.

Diffusion pumps are often combined with several vapor jet stages, termed ejector stages, to increase the fore-vacuum tolerance. These pumps are then referred to as hybrid pumps or diffusion ejector pumps. Compared to diffusion pumps, their fore-vacuum tolerance is higher. Hybrid pumps that use oil as pump fluid are preferred for pumping down large amounts of gas or water vapor. Here, pumps are employed with nominal pumping speeds of $10\,000\,\ell\,\mathrm{s}^{-1}$ and more. The critical backing pressure in such systems amounts to several 100 Pa.

9.4.7

Calculating Performance Characteristics of Diffusion and Vapor Jet Pumps by Using a Simple Pump Model

The pumping speed of a diffusion pump is calculated from the area of the free inlet cross section and the pumping probability or transmission probability P_{Ho} . Following a 1932 publication by Ho [10], P_{Ho} is also referred to as the Ho factor.

For a better understanding of the processes in diffusion and vapor jet pumps, a simple model is used for calculating pumping speed S (i.e., the pumped volume flow rate) as well as the transmission probability P_{Ho} and its dependence on the

gas species, pressure in the recipient, pump fluid data, as well as critical backing pressure and several other characteristic values.

The action in a diffusion pump relies on the momentum transfer during collisions of pumped-down gas molecules and vapor molecules traveling at jet velocity u. It can be attempted to calculate pumping speed S or pumping probability P_{Ho} directly from collision incidents. Approximations in such calculations lead to results moderately consistent with practical experience [11,12]. However, a relatively simple model based on an overall assessment of collision incidents already yields a viable correlation with experimental results. The model treats the interaction of pumped-down gas with the vapor jet as a diffusion process. This corresponds to Gaede's approach [13], which designated diffusion pumps, and later calculations by *Jaeckel* [3], who provided the earliest work that yielded realistic values of pumping speed. Using the term diffusion is reasonable because diffusion is in fact a macroscopically observable process relating to statistics of individual collision incidents (compare also Section 3.3.4).

Further discussion on the theory of diffusion pumps is found in [9,14], as well as [15–17].

The simplified model of a diffusion pump is derived from Figure 9.18, showing the upper stage of a diffusion pump. The vapor jet S-S originates at the annular jet nozzle JN (width δ , area A^*) formed by the upper end of the vapor pipe VP and the nozzle cap NC. It has the shape of a hollow cone, defined by VP on the inside and a blurred (diffuse) bell-shaped zone on the outside.20 For simplification, the outer boundary is assumed as a sharp jet edge JE formed by the envelope of a cone, which spreads, as an extension of the nozzle cap NC, from NC₀ to PB₂. A

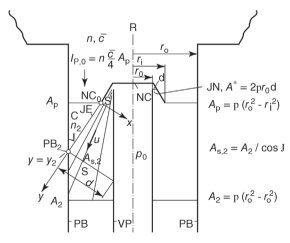


Figure 9.18 Quantities needed to calculate performance characteristics in the top stage of a diffusion pump.

²⁾ Ref. [18] discusses the expansion of the vapor jet under high vacuum as well as the jet boundary in detail.

surface line of this area is used as the y-axis (NC₀ being the origin of the coordinate and $y = y_2$ at the point of intersection with the pump body surface in PB₂).

The gas molecules (particle number density n_G , mean thermal velocity \overline{c}) pumped down from the large recipient R flow through the pump's annular aperture with an area A_P at the following rate (Section 4.4.1, Eq. 4.137):

$$I_{P,0} = n_G \frac{\overline{c}}{4} A_P. \tag{9.10}$$

They hit JE and diffuse into the motive jet S–S. The density n_v of the vapor jet molecules is much higher than the density n_G of the incoming, diffusing gas molecules. In addition to the thermal velocity \bar{c}_S , the vapor jet molecules also show a preferred velocity in the direction of the jet, termed the jet velocity u. They collide with the incoming gas molecules and impose a preferred velocity in jet direction on the gas molecules as well. The mean velocities of the molecules contained in the components of a gas mixture adjust rapidly after few collisions. Therefore, we can assume that the gas molecules also show a preferred velocity u in the direction of the jet. Because $n_v \gg n_G$ and $m_{\rm v} > m_{\rm G}$ (where m is the molecular weight), the jet is hardly decelerated; in fact, the gas molecules are entrained.

In this way, the gas flow I_2 is directed downward, through the jet cross section $A_{\rm J,2}$, and into the fore-vacuum chamber. Now, I_2 is not equal to $I_{\rm P,0}$ but smaller because not every molecule traveling downward from R through A_P diffuses into the jet and is then pumped. Some of the molecules return through A_P and back into R due to back diffusion, reflections at the wall, or other circumstances. If the probability of traveling back is $1 - P_{Ho}$, that is, the pumping probability is P_{Ho} , then, using Eq. (9.10), the pumping speed will be given as follows:

$$S = \frac{I_{P,0}}{n_G} P_{Ho} = \frac{\overline{c}}{4} A_P P_{Ho} = S_0 P_{Ho}. \tag{9.11}$$

So is the pumping speed of an ideal pump without any backflow. As stated above, pumping probability P_{Ho} is also referred to as the *Ho* factor [10,19].

An additional consequence of the backflow is that the particle number density n_2 of gas molecules in volume C (between A_P , PB, and JE) is lower than n_G . The total flow rate through A_P is equal to the flow rate from R to C minus the flow rate from C to R:

$$I_{\text{tot}} = n_{G} \frac{\bar{c}}{4} A_{P} - n_{2} \frac{\bar{c}}{4} A_{P}. \tag{9.12}$$

Using $I_{\text{tot}} = S n_{G}$,

$$S = \frac{\overline{c}}{4} A_{\rm P} \left(1 - \frac{n_2}{n_{\rm G}} \right),\tag{9.13}$$

and in combination with Eq. (9.11) we find the relationship

$$P_{\text{Ho}} = 1 - \frac{n_2}{n_G}.\tag{9.14}$$

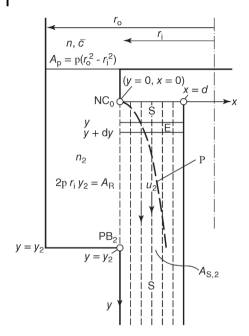


Figure 9.19 Simplified model of a diffusion pump for calculating pumping speed.

However, n_2 is still unknown. Determining its value—and thereby the value of P_{Ho} —requires investigating the diffusion process from C into S–S.

For calculating the diffusion process from C into S-S, the model is simplified further, which can be justified from diffusion theory. The deduction is not listed here for lack of space, but Figure 9.19 shows the simplification. The cone-shaped vapor jet is initially replaced by a tubular jet with the radii r_i and $r_i - d_i$, showing a parallel flow with the constant velocity u_2 (in the cone-shaped jet, the velocity at the nozzle differs from the velocity at the lower end). Furthermore, the particle density of the working vapor $n_{v,2}$ shall be constant. Finally, the tubular jet is unrolled to a band of length y_2 , width d, and rectangular area $A_{S,2}$. Now the situation may be treated as a plane diffusion problem.

We will consider an element E of the band jet between y and y + dy. At the left edge of this element, gas particle density is n_2 . If E was at rest and gas free (only consisting of working vapor molecules) at time t = 0, a distribution of incoming, diffusing gas molecules $n_G(x)$ according to Figure 9.20 would develop at time t. According to diffusion theory, the locus of the half-value is as follows:

$$x_{\text{diff}} = \sqrt{Dt} \tag{9.15}$$

if D is the diffusion coefficient of the gas in the vapor jet.

For further simplification, the curve for $n_G(x)$, illustrated in Figure 9.20, is replaced by a discontinuity at $x = x_{\text{diff}}$ so that the element is filled with $n_{\text{G}} = n_2$ between x = 0 and $x = x_{\text{diff}}$, and is empty for $x > x_{\text{diff}}$. Following the motion of

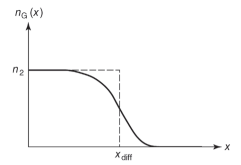


Figure 9.20 Particle number density in the motive jet and idealized square distribution.

jet element E in the y-direction from NC₀ to PB₂, it travels to the point $y = u_2 t$ within the time *t* and is then filled to the following coordinate:

$$x_{\text{diff}} = \sqrt{Dy/u_2},$$

which is represented by the parabola Π (Figure 9.19). Now, the flow of gas particles transported through $A_{S,2}$ can be calculated. However, we must consider that the flow of gas particles in our model is restricted to the fraction $x_{\text{diff},2}/d$ of the area $A_{S,2}$ (compare Figure 9.19). We find

$$I_2 = n_2 u_2 A_{5,2} \frac{x_{\text{diff},2}}{d} = n_2 u_2 \frac{A_2}{\cos \vartheta} \times \frac{x_{\text{diff},2}}{d}.$$
 (9.16)

 I_2 reaches a maximum when $x_{\text{diff},2} = d$. In a diffusion pump design, this condition is obtained by matching the width δ of the nozzles and the heating power. Then, u_2 and D can be controlled and

$$I_2 = n_2 u_2 \frac{A_2}{\cos \theta}.\tag{9.17}$$

Now we can calculate the pumping speed S and the pumping probability P_{Ho} by equating the particle number flow through A_P according to Eq. (9.12) and through A_2 according to Eq. (9.16). The obtained expression for n_2 , put in Eq. (9.14), yields the pumping probability

$$P_{\text{Ho}} = \frac{1}{1 + \frac{A_{\text{P}}\cos\theta}{A_2} \times \frac{\overline{c}}{4u_2} \times \frac{d}{x_{\text{diff},2}}}$$
(9.18)

and the pumping speed

$$S = A_{\rm P} \frac{\overline{c}}{4} \times \frac{1}{1 + a \frac{\overline{c}}{4u_2}} \tag{9.19}$$

with $a = A_P \cos \vartheta \times d/(A_2 x_{\text{diff},2})$.

In certain diffusion pumps, the pumping speed is increased by a belly-type expansion of the housing, near the first and second stages. This increases the contact length between the motive vapor jet and the pumped gas and, thus,

In vapor jet pumps, the vapor pressure is considerably higher than in diffusion pumps. Therefore, the diffusion coefficient D of the gas in the vapor is correspondingly low so that $x_{\text{diff},2} \ll d$ because of Eq. (9.15). Then, the constant 1 in the denominator of Eq. (9.18) may be neglected. Thus, pumping probability

$$P_{\text{Ho,vaporjetpump}} = b \frac{4u_2 \, x_{\text{diff,2}}}{A_P \overline{c}} \tag{9.20}$$

and pumping speed

$$S_{\text{vaporietpump}} = b \, u_2 x_{\text{diff},2} \approx 2\pi r r_0 x_{\text{diff},2} u_2,$$
 (9.21)

using the abbreviation

$$b = \frac{A_2}{d\cos\theta} \approx 2\pi r_0$$
 (compare Figure 9.18). (9.22)

Equation (9.19) expresses that S is independent of the gas pressure p_{in} or of the particle number density n at the inlet. Thus, it correctly describes the horizontal part of the curve in Figure 9.16. The drop of S at higher pressures will be discussed below. The influence of the gas species on the pumping speed is included in $\overline{c} \propto 1/\sqrt{M_{\text{molar}}}$ (compare Eq. 3.43), which explicitly appears twice in Eq. (9.19) and implicitly in the factor a where it is hidden in the diffusion length according to Eq. (9.15) and, here, in the diffusion coefficient D (Eq. 3.103). Therefore, with $a_{gas} \propto M^{1/4}$, the ratio of the pumping speed for any gas to the pumping speed for air is given as follows:

$$\frac{S_{\text{gas}}}{S_{\text{air}}} = \left(\frac{M_{\text{r,air}}}{M_{\text{r,gas}}}\right)^{1/2} \frac{1 + a_{\text{air}} \overline{c}_{\text{air}} / (4u_2)}{1 + (a_{\text{air}} \overline{c}_{\text{air}} / (4u_2)), (M_{\text{r,air}} / M_{\text{r,gas}})^{1/4}}.$$
 (9.23)

Figure 9.21 shows pumping-speed curves for selected gas species [20]. The model used here not only gives the value of the horizontal part of the pumpingspeed curve, but, as will be described qualitatively, it also explains the behavior of a diffusion pump under pressure variations on the fore-vacuum side, the drop of pumping speed at high inlet pressures, and the appearance of the maximum in pumping speed in fine-vacuum diffusion pumps and vapor jet vacuum pumps.

We will differentiate between the fore-vacuum tolerance p_{crit} , as a critical value for the pressure $p_{\rm F}$ on the fore-vacuum side, and the compression ratio $p_{\mathrm{F}}/p_{\mathrm{in}}.$ The fore-vacuum tolerance is the threshold value, which determines the maximum value of $p_{\rm F}$ up to which pumping action occurs (Figure 9.17). The compression ratio is usually high enough to not influence the ultimate vacuum and pumping speed. The only case where the compression ratio can have a relevant effect and might have to be considered is for light gases such as hydrogen.

Similar to vapor jet pumps, the rise in pressure from high vacuum to fore vacuum in diffusion pumps can occur in a compression shock (see Section 4.2.7). If the pressure on the fore-vacuum side rises due to increased gas flow through the pump or due to inflow to the fore-vacuum side of the pump, the shock moves

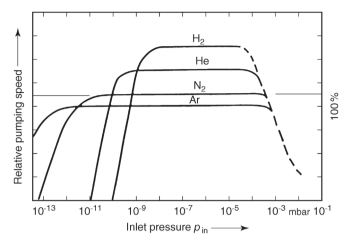


Figure 9.21 Typical pumping speeds of a diffusion pump for selected gas species versus inlet pressure p_{in} [20].

upstream until it reaches PB₂ (Figure 9.18). At this point, the area ratio of the motive vapor jet is A^*/A_2 (Figure 9.18). According to one curve in Figure 20.9, the static pressure ratio can be determined as a function of the area ratio $\iota = A^*/A_2$ (= current density ratio):

$$\frac{\hat{p}_0}{p_0} = f\left(\frac{A^*}{A_2}\right). \tag{9.24}$$

It follows the static pressure \hat{p}_0 behind the compression shock:

$$\hat{p}_0 = p_0 f\left(\frac{A^*}{A_2}\right),\tag{9.25}$$

which appears to be proportional to the static pressure p_0 of the driving jet in the evaporation chamber.

For any pressure value $p_{\rm F}$ on the fore-vacuum side that conforms to $p_{\rm F}<(\hat{p}_0)_2$ (\hat{p}_0) at the point PB₂), the vapor jet reaches the pump wall W and seals the highvacuum chamber from the fore-vacuum chamber. The pump is in normal operating condition. If though the pressure on the fore-vacuum side rises beyond the static pressure of the shock at the point PB₂, that is, $p_F > (\hat{p}_0)_2$, then the shock front shifts upward so that the supersonic jet no longer reaches the wall and the pumped gas flows back in between the jet and the pump wall. The pumping action then ceases. Thus, fore-vacuum tolerance

$$p_{\text{crit}} = (\hat{p}_0)_2 = p_0 f\left(\frac{A^*}{A_2}\right).$$
 (9.26)

The geometry of the pump determines the area ratio A^*/A_2 , and the static pressure p_0 depends linearly on the input heating power. Therefore, within certain boundaries, a linear relationship exists between the fore-vacuum tolerance and heating power.

At zero pumping flow (pumped volume), that is, when $I_N = \dot{N} = 0$ for any cross section A_y at a point $y > y_2$ (Figure 9.19), the compression ratio $(p_F/p_{in})_0$ is given by equating the downward convection current through A_{ν} with the diffusion current, which is directed upward:

$$A_{y}n(y)u_{2} = A_{y}D\frac{\mathrm{d}n}{\mathrm{d}y}.$$
(9.27)

Integration of this differential equation with the boundary conditions $n(y_2) =$ n_2 and $n(y_2 + L) = n_L$, and employment of the ideal gas law, Eq. (3.19), lead to

$$\frac{p_{\rm F}}{p_{\rm in}} = \frac{n_L}{n_2} = \exp\left(\frac{u_2 L}{D}\right). \tag{9.28}$$

For this integration, u_2 and $D \propto n_v^{-1}$ (because $n \ll n_v$) were assumed constant (compare Eq. 9.28 with Eq. 9.8 where D was defined slightly differently). Since vapor density $n_{\rm v}$ is approximately proportional to the heating power \dot{Q} , Eq. (9.28) can be rewritten as

$$\log p_{\rm in} = \log p_{\rm F} - {\rm constant} \times \dot{Q}. \tag{9.29}$$

Thus, for a given fore-vacuum pressure p_F , the decrease in high-vacuum-side pressure

$$\frac{\log p_{\text{in},2} - \log p_F}{\log p_{\text{in},1} - \log p_F} = \frac{\dot{Q}_2}{\dot{Q}_1}.$$
(9.30)

Equation (9.29) represents a straight line with gradient 1. Measured data shown in Figure 9.22 are described fairly well by this relationship. Equation (9.30) also

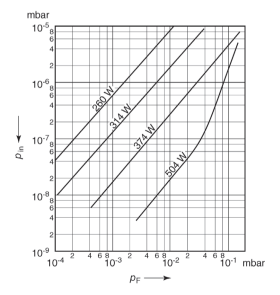


Figure 9.22 High-vacuum pressure p_{in} for a particle flow rate $\dot{N} = 0$ (ultimate pressure) versus fore-vacuum pressure $p_{\rm F}$, $p_{\rm F}$ is controlled by the air introduced to the fore vacuum. Parameter: heating power Q.

may be verified by the results: from Figure 9.22, for example, for $p_{\rm F} = 0.1 \, {\rm Pa}$ (10^{-3}mbar) , we obtain $p_{\text{in}} = 6 \times 10^{-5} \text{Pa} \ (6 \times 10^{-7} \text{mbar})$ for $\dot{Q} = 260 \text{ W}$, and $p_{\text{in}} = 6 \times 10^{-5} \text{ Pa}$ 1.5×10^{-6} Pa $(1.5 \times 10^{-8} \text{ mbar})$ for $\dot{Q} = 374$ W. With these values, the left side of Eq. (9.30) equates to 0.67, and the right side to 0.7, thus showing fair accordance. Figure 9.22 shows that the fore-vacuum tolerance, the maximum pumping speed, and the compression ratio all increase with rising heating power. Ultimate pressure, however, increases due to the increased density of the particles in the motive jet. The pumping speed also drops beyond a certain value of \dot{Q} . Locus and height of the maximum depend on the gas species.

9.5 **Diffusion Pumps Versus Vapor Jet Pumps**

The simple diffusion pump model (Section 9.4.7) is suitable for describing the processes in vapor jet pumps as well. The transition from a diffusion pump to a vapor jet pump is smooth. Figure 9.23 shows three pumping speed curves of a vapor jet pump for three different heating powers \dot{O} . For the low heating power of 50 W, the typical flat curve is produced in the low inlet-pressure range, and a steeper slope at $p_{in} = 0.1 \text{ Pa} (10^{-3} \text{mbar})$. Thus, the vapor jet pump operates at lower heating power than the diffusion pump. When the heating power is increased to 500 W, a transition condition is reached where the curve is flat, as in a diffusion pump, for inlet pressures $p_{\rm in}$ below $10^{-2}{\rm Pa}$ ($10^{-4}{\rm mbar}$), but then increases for higher pressures until a maximum is reached at $p_{\rm in}=1\,{\rm Pa}$ (10⁻²mbar), and finally drops steeply, as in a diffusion pump. For a standard heating power of a vapor jet pump ($\dot{Q} = 5 \text{ kW}$), the typical pumping-speed maximum of a vapor jet pump develops, as well as a drop in pumping speed to $S \approx 0$ at lower pressures.

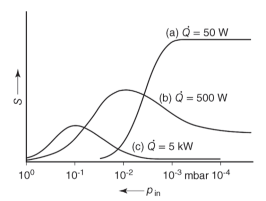


Figure 9.23 Pumping speed S of a singlestage jet pump versus inlet pressure p_{in} for selected heating powers Q. (a) Typical characteristic for the high-vacuum stage of a

diffusion pump. (b) Typical characteristic for the fore-vacuum stage of a diffusion pump. (c) Typical characteristic for a vapor jet pump.

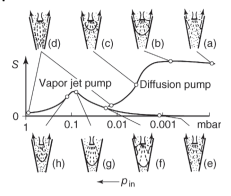


Figure 9.24 Pumping speeds S and diagrams of processes in pumps versus inlet pressure p_{in} . Point density resembles air density, dashed lines are flow patterns, and continuous lines are compression shocks.

In Section 9.4.7, the value for the pumping speed of a diffusion pump in the flat part of the curve was derived from the model. Now we will show that the model can be used to explain qualitatively the decline of the curve at higher pressures as well and, also, the maximum of a vapor jet pump's curve.

In diffusion pumps, inlet pressure is so low that the expansion of the jet is practically unconstrained by the surrounding gas. In contrast, it is the interference on the expansion of the vapor jet under higher inlet pressures p_{in} that is responsible for the development of the maximum in the pumping-speed curve in vapor jet pumps. Therefore, in vapor jet pumps, the influence of inlet pressure on the shape of the motive vapor jet must be considered. Here, alteration of the jet boundaries, represented by the γ -axis in Figure 9.18, is of particular concern. Figure 9.24 gives a schematic illustration.

The figure shows typical curves of pumping speeds S versus inlet pressure $p_{\rm in}$ for diffusion and vapor jet pumps. For the points of the curves denoted by (a)-(h), the diagram gives schematic illustrations of corresponding nozzle systems and shapes of the vapor jet. Obviously, in the low-intake pressure range, points (a), (b), (e), and (f), the shape of the vapor jet is practically independent of inlet pressure p_{in} . A compression shock on the pressure side in the lower part of the jet develops only under increased inlet pressures, points (b) and (f). Here, the pressure jumps up to the value on the fore-vacuum side, similar to the process described in Section 9.4.7. In a diffusion pump, points (a) and (b), the jet boundary represented by the y-axis hits the pump's wall at a grazing angle. Additionally, the gas penetrates deep, and thus, most of it is transported downward with the vapor jet. Pumping speed is high; only a negligible amount of gas emanates from the jet boundary and flows back upward.

For the vapor jet pump, however, the pumped-down gas penetrates only the outer edge of the jet because of the higher vapor pressure (compare Eqs. 9.20-9.22). At low inlet pressures, $p_{
m in} < 1$ Pa (10^{-2} mbar), points (e) and (f), the vapor jet expands more or less freely. Thus, the jet boundary hits the wall of the pump at a large angle ($\theta \approx 90^{\circ}$) and the transported gas, instead of flowing downward, flows back upward into the container. Pumping speed is negligible.

If inlet pressure rises to $p_{\rm in} \approx 10 \, \text{Pa}$ (0.1 mbar), point (g), the outer edge of the vapor jet cannot expand as far as for low inlet pressures, (e) and (f). Thus, the vapor jet is sharply focused. The intersection line of the jet surface (y-axis) cuts the pump wall at a considerably lower angle ϑ . This causes the gas to be transported downward in spite of the fact that it penetrates only the outer edge of the vapor jet and, thus, produces the maximum in the pumping-speed curve characteristic to vapor jet pumps. If inlet pressure $p_{\rm in}$ increases further, backflow from the fore-vacuum side is observed. This is because the compression shock, which develops just as in a diffusion pump, moves upstream far enough for a connection to be established between the inlet volume and the volume on the fore-vacuum side, near the pump wall, point (h). This is analog to the case in a diffusion pump, (c). If inlet pressure increases even further, (d), hardly any pumping action is producible. This is due to the fact that nearly all of the gas transported by the jet streams back in between the pump's wall and the vapor jet, which now is restrained heavily in terms of expansion. This applies equally to diffusion pumps and vapor jet pumps. These considerations lead to the conclusion that Eq. (9.22) for the pumping speed of a vapor jet pump applies to the value of S at the curve's maximum. A quantitative description of the complete curve would have to consider the transition of the vapor jet with inlet pressure and the pressure on the fore-vacuum side and, additionally, would have to acknowledge the amount of gas flowing upward from the boundary regions of the vapor jet, depending on the angle of incidence. The shape of the vapor jet has been investigated experimentally by visualizing mercury and oil vapor jets by means of a high-frequent gas discharge, as well as theoretically using gas-dynamic methods, under the conditions in diffusion and vapor jet pumps [21,22].

References

- 1 VDMA 24292 sheet 2 (1971) Dampfstrahl-Vakuumpumpen und Dampfstrahl-Verdichter, Messregeln.
- 2 Gaede, W. (1913) German Patent 286404.
- 3 Jaeckel, R. (1950) Kleinste Drücke, Springer, Berlin-Göttingen-Heidelberg, pp. 140 ff.
- 4 Rettinghaus, G. and Huber, W.K. (1974) Backstreaming in Diffusion Pumps. Trans. 6th International Vacuum Congress, Kyoto. Vacuum, 24, 249.
- 5 Holland, L. (1970) A review of some recent vacuum studies (related to pumps and ion beam sputtering systems). Vacuum, 20, 175.

- 6 Hablanian, M.H. and Maliakal, J.C. (1973) Advances in diffusion pump technology. J. Vac. Sci. and Technol., 10, 58.
- 7 Meyer, D.E. (1974) Residual gas analysis studies during sputtering of reactive metals. J. Vac. Sci. and Technol., 11, 168.
- 8 Burch, C.R. (1978) Oil, greases and high vacua. Nature, 122, 729.
- 9 O'Hanlon, John F. (1989) A User's Guide to Vacuum Technology, 2nd edn, John Wiley & Sons, New York, p. 215.
- 10 Ho, T.L. (1932) Physics, 2, 386.
- 11 Florescu, N.A. (1960) Vacuum, 10, 250; (1963) Vacuum, 13, 560.

- 12 Toth, G. (1960) Eine Übersicht über die Theorie der Diffusionspumpen. Vakuumtechnik, 16, 41-47.
- 13 Gaede, W. (1913) Die Molekularluftpumpe. Ann. Phys., 346, 337-380; (1915) Die Diffusion der Gase durch Quecksilberdampf bei niederen Drucken und die Diffusionsluftpumpe. Ann. Phys, 351, 357-392.
- 14 Dayton, B.B. (1998) Diffusion and diffusion-ejector pumps, in Foundations of Vacuum Science and Technology (ed. J.M. Lafferty), John Wiley & Sons, New York,
- 15 Dushman, S. and Lafferty, J.M. (1962) Scientific Foundations of Vacuum Technique, John Wiley, New York.
- 16 Power, B.D. (1966) High Vacuum Pumping Equipment, Chapman & Hall, London.

- 17 Beck, A.H. (1966) Handbook of Vacuum Physics: vol. 1. Gases and Vacua, Pergamon Press, Oxford.
- 18 Nöller, H.G. (1966) Approximate calculation of expansion of gas from nozzles into high vacuum. J. Vac. Sci. Technol., 3 (4), 202.
- 19 Fowler, P. and Brock, F.J. (1970) Accurate, wide range ultrahigh-vacuum calibration system. J. Vac. Sci. Technol., 7, 507.
- 20 Hablanian, M. (1986) Performance characteristics of displacement type vacuum pumps. J. Vac. Technol. A, 43, 286-292.
- 21 Nöller, H.G. (1966) Handbook of Vacuum Physics: vol 1, part 6: Theory of Vacuum Diffusion Pumps, Pergamon Press, Oxford.
- 22 Wutz, M. (1969) Molekular-kinetische, Vieweg & Sohn, Braunschweig.

10

Molecular and Turbomolecular Pumps

Dr. Frank Leiter

formerly: Pfeiffer Vacuum AG, Berliner Straße 43, 35614, Asslar, Germany

The molecular pumps having a strong significance in high-vacuum technology are introduced in this chapter. Due to the market penetration and extent of application, the turbomolecular design will be focused upon. The description of the technical construction follows a representation of the physical fundamentals. Indications regarding parameters, operation and applications of the pumps conclude the chapter.

10.1 Introduction

Molecular and turbomolecular pumps are connected with each other through their fundamental physical working principles [1]. In both pump types, gas is transported as momentum is transferred to the gas molecules thereby achieving a directed movement.

The momentum transfer takes place through a quickly moving wall or by blades of a quickly turning rotor. Indeed, this principle can only be in the molecular flow range because in this, the mean free path of the gas molecules to be pumped is greater or, better still, substantially greater than the distances within the pump. If one reduces the dimensions accordingly, additional regimes are attainable that usually would be ascribed to the transition range between molecular and viscous flow areas (several hPa). If the dimensions between the walls are decreased accordingly, additional ranges are attainable that usually would be ascribed to the transition range between molecular and viscous flow (several hPa).

Due to this circumstance, a molecular pump is usually not capable of compressing and ejecting against atmospheric pressure but requires a backing pump that compresses from the outlet pressure of the (turbo)molecular pump to ambient pressure. Positive displacement pumps, as described in Chapter 7, that compress to atmospheric pressure are used as backing pumps.



Rotor discs

Figure 10.1 Periphery of the rotor/stator stack in Becker's turbomolecular pump. The stator disk at the inlet of the pump increases the compression ratio but reduces the pumping speed.

Turbomolecular pumps are built up of fast rotating rotor disks with blades and mirror-symmetrical stator disks lying in between [2] (Figure 10.1). Gas particles are transported through the channels between the blades by an additional momentum transferred by the rotor blades.

Today, turbomolecular pumps are often combined with molecular pump stages, which are designed to exhaust gas at higher pressure. This allows the employment of cheaper, dry positive displacement pumps as backing pumps.

In contrast to sorption pumps (Chapter 11), which are limited in terms of their gas storage capacity and require regeneration phases, turbomolecular pumps are ready for operation quickly and transport gas through the pump continuously.

Due to the multistage axial pumping principle of the turbomolecular pump, low pressures can be generated at the inlet flange. Desorption of gases, which limits the ultimate pressure, can be reduced by baking. High compression ratios for heavy gases yield ultimate pressures in the area of 10^{-9} Pa. The same pumps can also be used to pump high gas throughputs in the inlet pressure range of 10^{-1} –1 Pa. Nowadays, applications of turbomolecular pumps in vacuum process technology with high gas loads (coating, semiconductor production) are economically far more important than pure vacuum production. The latter was the main field of application at the time such pumps were introduced on a greater scale.

In 1956, *W. Becker* [2] invented the turbomolecular pump (Figure 10.2). He explained the basic operating principle of his *new molecular pump* by the aid of *Gaede's* theory of the molecular pump. *Becker's* turbomolecular pump was the so-called double-flow pumps: Two multistage rotor–stator systems with turbine blades (Figure 10.1) pumped the gas from the inlet flange in the center to the fore-vacuum chambers, one of which also contained the drive unit. A common fore-vacuum line is connected to the backing pump. In this way, the ball bearings at both ends of the pump shaft could be housed in the fore-vacuum chambers. Advantages over the previously designed *Gaede* pump [3] are: high

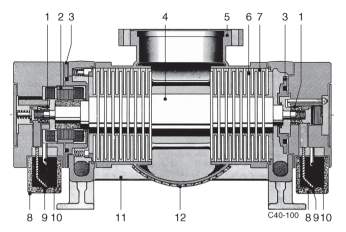


Figure 10.2 Section of Becker's double-flow turbomolecular pump TPU 200. 1: ball bearing; 2: motor; 3: labyrinth box; 4: rotor; 5: high-vacuum flange; 6: rotor disk; 7: stator disk; 8: oil reservoir; 9: oil supply wick to bearing; 10: oil-backflow line; 11: fore-vacuum channel; 12: heating jacket.

pumping speed, large distances between rotor and stator (≈ 1 mm), and a very high compression ratio due to the multistage design.

Further developments led to single-flow pumps with a smaller, lighter, and more economical design (Figure 10.3). Direct flange mounting to the recipient reduced conductance losses.

10.2 **Molecular Pumps**

Molecular pumps are also found in the product portfolios of the large manufacturers, but are primarily used in niche applications [4,5]. Generally, the



Figure 10.3 Single-flow turbomolecular pump HiPace 700 by Pfeiffer company with attached drive electronics for 48 V DC. 1: high-vacuum flange; 2: fore-vacuum flange; 3: venting valve; 4: purge-gas port; 5: electronic drive unit with remote control socket.

construction methods used follow Gaede, Siegbahn, and Holweck in forevacuum-sided pumping stages within turbomolecular pumps to reach higher output pressures. Such types of modified turbomolecular pumps are frequently referred to as *compound pumps*. Because these turbomolecular pumps are based on historically older molecular pumps and the operating principle of turbomolecular pumps can be well explained by considering this type of pump, we will first discuss the molecular pump.

The type of gas flow is the core difference between the molecular pump stage construction methods: In Gaede stages, it is pumped in a circumferential direction, following Siegbahn, radially along a disk surface, and with Holweck, in a thread-type groove in the surface of a cylinder.

In addition, regenerative pump designs also have been combined with molecular pump stages and are commercially available [6,7], see Section 10.3.

10.2.1

Gaede Pump Stage

The molecular (vacuum) pump was invented by Gaede (1913) [3]. He used the concept that molecules, which hit a wall, are not reflected directly but become adsorbed for a time interval, the dwell time, before they desorb (compare Chapter 6). On desorption, their velocity distribution is isotropic and corresponds to the wall temperature. The mean velocity \bar{c} is given by Eq. (3.43). If the wall moves with velocity u, then the velocity distribution will be superimposed by this drift velocity. A moving wall must therefore produce a flow and, thus, create a pumping effect.

Figure 10.4 shows the principle of Gaede's molecular pump. Molecules originating at inlet port A hit the rotor R with radius r, revolving at rotary frequency f. The molecules acquire a predominant velocity $u = 2\pi rf$ at which they move through the pump channel PC, with height h and width b, until they reach the fore-vacuum port FV. In order to prevent high backflow, FV must be separated from A by a locking slot LS of several 1/100 mm. This also applies to the gaps between the lid faces and the rotor.

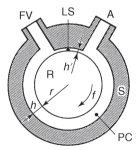


Figure 10.4 Principle of Gaede's molecular pump. The pumping effect is based on the tangential momentum transferred to the impinging gas molecules by the rotor. A: inlet port; FV: forevacuum port; R: rotor; S: stator; PC: pump channel; LS: locking slot.

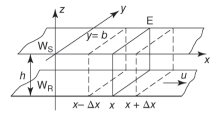


Figure 10.5 In reference to the principle of Gaede's molecular pump, Ws: stator wall; WB: rotor wall moving with velocity u; h: channel height; b: channel width; E: cutting plane.

The function and parameters of the pump can be described with a simple model that makes all the physically significant parameters of the molecular and turbomolecular pumps clear. For simplification, Figure 10.5 shows a plane section of the rotor wall W_R moving with velocity u, and the stationary stator wall W_S . The distance h between W_R and W_S shall be small compared to the mean free path \bar{l} of the gas with particle number density n. Therefore, the particles collide with the walls only, and not among one another. Thus, at any point in time, half of the particles in the x-direction carry the drift velocity u, whereas the other half show zero drift velocity. The particle flow moving in the channel PC (Figure 10.4) is

$$q_{(-1pt)N} = \frac{\mathrm{d}N}{\mathrm{d}t} = \frac{1}{2}nubh. \tag{10.1}$$

This particle flow density leads to the volume flow rate q_V , which is equal to the pumping speed *S* (compare Section 4.1.2):

$$S_0 = q_V = \frac{q_{(-1pt)N}}{n} = \frac{1}{2}ubh. \tag{10.2}$$

This equation for the pumping speed is valid for pumping without any pressure difference between FV and A. However, the influence of the front and back boundary walls of the channel remains unconsidered. For a basic approach, this may be neglected as long as $b \gg 4$. Thus, the pumping speed of an arrangement according to Figure 10.4 is proportional to the circumferential velocity u of the rotor and the cross section bh of the channel PC. For an examination of the basic principle, the backflow through LS, which can be described by Eq. (10.1) also, was neglected as well.

The gas flow from A to FV in Figure 10.4 according to Eq. (10.1) produces a pressure gradient dp/dx (Figure 10.5) or a density gradient dn/dx, which causes a backflow. Due to the drift velocity u, a particle flow $q_{(-1vt)N}$, according to Eq. (10.1), moves from left to right through the y-z-plane E at the point x where the particle number density shall be n(x). To the right of x, at the point $x + \Delta x$, the particle number density, therefore, is $n(x + \Delta x) > n(x) \leftarrow$ and analog $n(x - \Delta x) < n(x)$. Thus, according to Eq. (3.49), the random thermal particle flow from right to left through E is

$$q_{N, \text{th}, l} = bh \frac{\overline{c}}{4} n(x + \Delta x),$$

and from left to right accordingly is

$$\vec{q}_{(-1pt)N,\text{th},r} = bh\frac{\overline{c}}{4}n(x - \Delta x).$$

Thus, the excess backflow is

$$q_{N,\text{th}} = bh \frac{\overline{c}}{4} [n(x + \Delta x) - n(x - \Delta x)] = bh \frac{\overline{c}}{4} \times \frac{dn}{dx} 2\Delta x.$$

The quantity Δx is chosen on the basis of the same considerations as in Section 3.3.2: $n(x + \Delta x)$ and $n(x - \Delta x)$ have to be picked at the place where the particles had their last collision because here they got their mean isotropic distribution so that Eq. (3.48) is valid. In Section 3.3.2, this was at a distance \bar{l} , the mean free path in the gas, in front of the considered plane. However, here $\bar{l} \gg + \bar{l}$; the mean collision distance is, therefore, slightly higher than h. So we must set $\Delta x = \leftarrow$ gh (with g > 1) and obtain

$$q_{N,\text{th}} = gbh^2 \frac{\overline{c}}{2} \times \frac{dn}{dx},\tag{10.4}$$

and, thus, the output flow

$$q_{(-1pt)N} = \frac{1}{2}nubh - g\frac{\overline{c}}{2}bh^2\frac{\mathrm{d}n}{\mathrm{d}x}.$$
 (10.5)

If we set the output $q_{(-1pt)N}$ to zero, that is, drift flow and backflow are equal, we find

$$\frac{\mathrm{d}n}{n} = \frac{\mathrm{d}p}{p} = \frac{u\,\mathrm{d}x}{\bar{c}gh}\,. (10.6)$$

In this case, the pump delivers no net gas output, but produces maximum compression between inlet pressure p_A and exhaust pressure p_{FV} . Integration of this equation along channel length L yields the maximum compression ratio:

$$K_0 = \frac{p_{\text{FV}}}{p_{\text{A}}} = \exp\left(\frac{uL}{\overline{cgh}}\right). \tag{10.7}$$

The compression ratio without throughput K_0 , thus, increases exponentially with the velocity ratio u/\overline{c} and the ratio of channel length to channel height L/h. Because most molecules move through the channel at an angle, the factor g, describing the free path to the next wall collision, will be in the range 3 > g > 1. Equation (10.7) always produces too high values because it neglects the backflow through the narrow slot LS and also the backflow at the lid faces. For approximation, the exponent in Eq. (10.7) can be written as 34gSC. Here, S = ubh/2 is the pumping speed according to Eq. (10.2) and

$$C = \underbrace{\frac{4\overline{c}(bh)^2}{3 \times 2L(b+h)}}_{\leftarrow}$$

is the conductance of the pump channel under the condition $h \ll +b$. The main conclusion from Eq. (10.7) is that the ratio u/\bar{c} must be as high as possible in order to obtain high K_0 . Thus, molecular pumps, regardless of their type, call for high rotary frequencies.

Example 10.1

For r = 0.05 m and f = 1000 Hz, it follows that u = 314 m s⁻¹. For nitrogen at 300 K, $\overline{c} = 475 \text{ m s}^{-1}$ and the channel dimensions $L = 0.9 \times 2\pi r = 0.28 \text{ m}, h = 0.003 \text{ m}, b = 0.008 \text{ m}, g = 2$; it follows: $K_0 = 2.49 \times 10^{13} \text{ and } S_0 = 3.76 \text{ }\ell\text{ s}^{-1}$.

A molecular pump proposed by Gaede and built in 1913 by E. Leybolds Nachf had a pumping speed of $1.5 \,\ell \, {\rm s}^{-1}$ and a compression ratio of $K_0 = p_{\rm FV}/p_{\rm A} = 10^5$ at a rotational speed of 8200 rpm.

Equation (10.7) also shows that very high compression ratios will be generated for heavy gases due to $\overline{c} \sim M_r^{-1/2}$. Using fluorinated pump oil with an average molecular weight of 2100 on the fore-vacuum side and a vapor pressure of 10 Pa at 200 °C, extremely high compression ratios are obtainable. Comparing hydrogen and perfluorinated polyether (PFPE) oils, we find

$$\frac{(K_0)_{\text{PFPE}}}{(K_0)_{\text{H}_2}} = \exp\sqrt{\frac{2100}{2}} = 1.18 \times 10^{14}.$$

Thus, under standard operating conditions, a molecular pump will hold back heavy gases and vapors, as may be released by lubricating oil in the fore-vacuum chambers, from the high-vacuum side.

Under laminar flow conditions, the same equations apply for drift and backflow. However, since the mean free path $\bar{l} \ll H$ at higher pressure, the assumption $\Delta x = gh$ with g > 1 in Eq. (10.3) is wrong. Backflow in the pump channel as well as in the locking slot and at the lateral lids increases considerably due to the reduced free path. This causes a drastic decline in compression and, thus, limits operation of the highly effective Gaede stages to the molecular flow regime.

Today, Gaede pump stages are used by some manufacturers in turbomolecular pumps; however, the area of the gas seal with a narrow locking slot Sp'and the unsymmetrical gas flow represent hurdles.

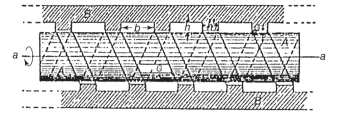


Figure 10.6 Gaede's thread pump or Holweck pump with a smooth rotor A. By the thread-like design of the pumping channel in the stator (B), the length of the channel is increased and also the compression ratio. Losses occur due to backflow underneath the barriers.

10.2.2

Holweck Pump Stage

In 1923, *Holweck* [8] developed a molecular pump with an operating principle analog to the thread pump suggested by *Gaede* (Figure 10.6). In a *Holweck* pump, the rotor R is made up of a cylindrical drum with a smooth surface. The coaxial, cylindrical stator S is equipped with a thread-type groove on the inside. With this arrangement, Holweck circumvented the problem of the gas seal; he nevertheless spoke of a Gaede pump himself. This pump had the inlet port in the center and the rotor transported the gas through two opposing grooves to the fore-vacuum chambers, which also contained the bearings. The pumping speed of this pump was $6 \,\ell$ s⁻¹ and the compression ratio for air reached 2×10^7 .

Gaede's equations for pumping speed, Eq. (10.2), and the compression ratio, Eq. (10.7), apply to Holweck stages. Instead of the losses due to backflow through the locking slot and lateral lids, however, losses here occur due to backflow through the gap between the rotor and the barriers of adjacent pump channels in the stator. The clearance below the barrier as well as the channel height h must be small compared to the mean free path in the desired pressure range. It is reasonable to reduce the channel height and clearance for higher pressures. The minimum clearance in real Holweck stages depend on rotor expansion caused by centrifugal forces and temperature changes. Radial rotor motion, which is possible due to soft suspension of the bearings and gaps around the safety bearings of magnetically suspended pumps, requires certain clearances as well. Using carbon fiber sleeves as *Holweck* rotors with very low coefficients of thermal expansion and low dilatation due to centrifugal forces allows gap widths of 0.3-0.5 mm. Corresponding channels should be approximately five times as high. If a mean free path of $\bar{l} = 0.25 \,\mathrm{mm}$ (Knudsen number = 0.5 for the gap) is assumed for such dimensions, the critical pressure $p_{\rm crit}$ for nitrogen with $\bar{l}p = 5.9 \times 10^{-3}$ Pa m is according to Eqs. (3.56) and (3.57): $p_{\rm crit} = 23.6$ Pa. If the pressure in the Holweck stage increases further, then the flow through the gaps and backflow in the pump channel rise until, by a pressure increase by a factor of approximately 100, pumping action in the Holweck stage ceases. The calculations of the Holweck stages are subject of the research. The pumping speed and

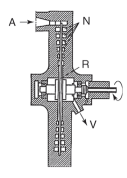


Figure 10.7 *Siegbahn's* molecular pump. The decreasing velocity toward the shaft, pumping against centrifugal forces, and difficulties arising when combined with turbomolecular pump stages are disadvantages compared to

the Holweck pump and have impeded their widespread technical use. A: inlet port: V: vacuum port; N: spiral pumping channel in the stator; R: rotor disk.

compression ratio are modeled in a work with moderate computational effort and in good correspondence with the measured values, cf. Section 5.7 and [9]. Good results for the viscous pressure range can be achieved with the CFD (computational fluid dynamic) methods [10].

10.2.3

Siegbahn Pump Stage

As an additional variant, Siegbahn [11] developed a molecular pump in 1943 (Figure 10.7), in which plane rotor disks with axially milled ring-type or spiraltype stator channels form the pumping system. Compared with the construction method by Gaede, the Siegbahn stages are more symmetrically constructed. Disadvantages due to the design of this system are low circumferential speed near the shaft, pumping against centrifugal forces in multistage systems, and the use of diametrically split stator disks, which are expensive to produce, as well as the relatively narrow gap between the axially adjacent rotor and stator components, which can become a problem due to the thermal expansion of the rotor. The Siegbahn stages have been developed over recent years with the goal of approaching the efficiency of the Holweck stages and in the best case, achieving it [12].

10.3 Molecular and Regenerative Drag Pump Combination¹⁾

In the 1990s, the molecular drag mechanism was combined with the mechanism of a regenerative or side channel pump in a high-speed single-shaft vacuum pump capable of delivering to atmospheric pressure [13].

1) Section 10.3 by courtesy of Nigel Schofield, Edwards company, UK.

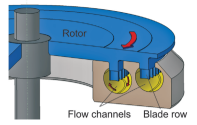


Figure 10.8 Two stages of a regenerative pump indicating the helical gas flow.

10.3.1

Regenerative Pump Mechanism

The regenerative pump mechanism uses fast moving blades to impart kinetic energy to the gas that is then converted into pressure rise and a pumping action. The name "regenerative" derives from the way gas circulates repeatedly through a blade row, each time "regenerating" the kinetic energy. Other names are also used: side channel, vortex, or peripheral flow pump.

A single stage of regenerative pump consists of a row of blades mounted on a rotor that protrude into a flow channel which is significantly larger than the blades [13] (see Figure 10.8). As the blades move forward through the gas, as a result of exchange of momentum and centrifugal forces, they generate a vortex, or helical, flow within the channel that spirals the gas through the blades. This allows the blades to repeatedly drive the gas forward, transferring kinetic energy to generate the pressure rise. The blades are shaped to catch the incident gas as smoothly as possible and deflect it forward into the channel. Blade speeds are typically between 100 and 250 m s^{-1} . Within a single stage the gas may circulate several times through a blade row, creating a compression ratio of typically 4:1.

Gas enters the circumferential flow channel at the inlet and moves around the channel until it is deflected away from the blade row into the exhaust or next stage of the pump (Figure 10.9). At this point the blades pass through a

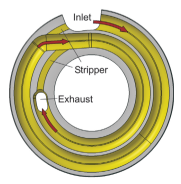


Figure 10.9 Two stages of a regenerative pump indicating flow deflection by the "stripper."



Figure 10.10 A typical pump cross section with inlet Holweck drag stages.

restrictive passage, generally known as the stripper, back to the inlet region. Here the channel cross section changes to a shape similar and just slightly larger than the blade cross section. Small pockets of gas between the blades, however, are carried through the stripper and create a carry-over volume, limiting the efficiency and compression of the machine.

Sixsmith [14] revolutionized the understanding and design of these machines and showed that practical compressors and vacuum pumps could be made. He developed some of the features described above and demonstrated a prototype machine with a maximum compression of 10 and a pumping speed of $250 \text{ m}^3 \text{ h}^{-1}$.

10.3.2

Pump Design and Applications

A regenerative vacuum pump typically uses a combination of regenerative stages and molecular drag stages to give good performance over a wide pressure range. Regenerative stages do not work in molecular flow and require the addition of drag stages to achieve pressures below about 1 mbar.

Pumps in which molecular drag stages are combined with regenerative stages are commercially available from several manufacturers. The example depicted in Figure 10.10 uses six concentric regenerative stages, exhausting to atmosphere and operating down to about 1 mbar. This is combined with five Holweck drag stages to take the inlet pressure down to 1×10^{-6} mbar. The resultant pump is very compact, and with no grease or oil in the vacuum region offers an ultraclean mechanism.

The pump speed curves of two different pumps of two different sizes are shown in Figure 10.11.

These pumps have found wide application in pumping semiconductor load locks and physics experiments such as accelerators and beam lines.

A further application of regenerative stages is in the exhaust of a turbomolecular pump. Here, one or two stages of regenerative mechanism may be used to increase the critical backing pressure and throughput of a turbomolecular pump.

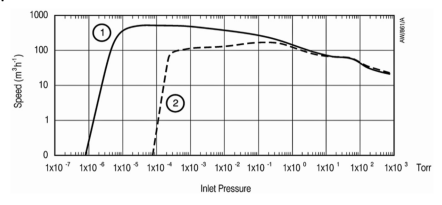


Figure 10.11 Pump speed curves of two types of regenerative pumps by Edwards, EPX500 (1) and EPX 180 (2). 1 Torr is equal to 133 Pa orl .33 mbar.

10.4 Physical Fundamentals of Turbomolecular Pump Stages

In the following, the pumping mechanism of turbomolecular pumps is more precisely examined and the important parameters such as the suction capacity and compression ratio are derived. In this instance, models are applied following *Becker* as well as *Kruger* and *Shapiro*. The thermal equilibrium of turbomolecular pumps is to be considered in their application and is, therefore, illustrated with the aid of the occurring effects.

10.4.1

Pumping Mechanism

We will first consider the pumping mechanism of a turbomolecular pump stage by investigating Figure 10.12.

A row of blades moving from left to right at a velocity u separates the spaces 1 and 2. Looking at the incoming particles as an observer moving with the blades, the blade velocity u will be added to the particle velocity \overline{c} vectorially. In the case of approximately equal velocities u and \overline{c} , many particles pass through the blade channel from space 1 without touching the blades. Particles that do touch the blades remain there for the dwell time and desorb according to the cosine law (Section 4.4.1 and [15]). For high velocities u, many particles touch the bottom surface of the blades and desorb mainly into space 2. If the same concept is applied to space 2, obviously only few particles will move in the direction of the channel and, therefore, only a small fraction enters space 1. These considerations lead to the conclusion that the transmission probability P_{12} for particles passing from 1 to 2 must be higher than P_{21} in the opposite direction. Thus, a pumping effect is produced.

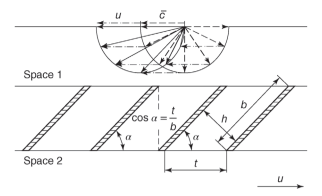


Figure 10.12 In reference to the principle of turbomolecular pumps. A row of blades with distances t between the blades, blade angle α , blade width b, and channel height h moves toward the right at velocity u. For an observer moving with the blades, blade velocity u is

added to the thermal velocity \overline{c} . For $u \approx \overline{c}$, nearly all molecules move in the direction of the channel or to the bottom side of the blades from where backflow into chamber 1 is improbable. $\cos \alpha = t/b$ is valid for an optically tight blade design.

10.4.2

Pumping Speed and Compression Ratio

In the molecular flow regime, P_{12} and P_{21} depend on the blade angle α , the ratio between the blade width and blade distance b/t, and the velocity ratio u/\bar{c} . If these values are known, the pumping speed S_0 and compression ratio K_0 of a turbomolecular pump stage can be calculated as shown next.

The transported particle number in the two spaces complies with

$$n_1 P = n_1 P_{12} - n_2 P_{21}. (10.8)$$

Considering $n_1 = n_2$, that is, equal pressure to both sides of the blade row, the pumping probability for zero pressure difference, also referred to as the Ho factor, is obtained:

$$P_{\rm H_0} = P_{12} - P_{21}. ag{10.9}$$

Using the inlet conductance $C = A\bar{c}/4$ (Eq. 4.143) with A = inlet cross section between two blades, the pumping speed of a blade channel is

$$S_0 = CP_{\text{Ho}} = A\frac{\overline{c}}{4}(P_{12} - P_{21}). \tag{10.10}$$

As in Eq. (10.7), the maximum compression ratio is derived for the case that the numbers of pumped and backflowing molecules are equal and, thus,

$$n_1 P_{12} = n_2 P_{21}, (10.11)$$

and because of p = nkT,

$$K_0 = \frac{p_2}{p_1} = \frac{P_{12}}{P_{21}}. (10.12)$$

If, in Eq. (10.8), n is expressed by pressure and the equation is multiplied by the inlet conductance C, the real pumping speed under pumping against a pressure difference is

$$S = CP = CP_{12} - \frac{p_2}{p_1} CP_{21}. \tag{10.13}$$

Setting $p_2/p_1 = K$, the real pressure ratio, it follows with Eq. (10.12):

$$S = CP_{12} \left(1 - \frac{K}{K_0} \right). \tag{10.14}$$

For K = 1, which means that $S = S_0$, we find

$$CP_{12} = \frac{S_0}{1 - 1/K_0}. (10.15)$$

Putting Eq. (10.15) in Eq. (10.14) yields

$$S = \frac{S_0(K_0 - K)}{K_0 - 1}.$$
 (10.16)

For pump stages in a series connection with equal gas throughputs as realized in a turbomolecular pump, $q = p_1 S = p_2 S_F$, and thus, $K = S/S_F$, Eq. (10.16) gives the real pumping speed of a turbomolecular pump stage with the pumping speed S_F of the preceding stage:

$$S = \frac{S_0}{1 - 1/K_0 + S_0/K_0 S_F}.$$
 (10.17)

This equation can be used as a recurrence formula for calculating the pumping speed of a multistage turbomolecular pump. For known pumping speed of the backing pump or a preceding pump stage S_F , as well as K_0 and S_0 of the succeeding turbomolecular pump stage, the pumping speed S of the latter one can be calculated.

10.4.3

Gaede and Statistical Theory of the Pumping Effect

10.4.3.1 **Consideration with Gaede**

For calculating pumping speed from the geometrical dimensions of pump stages (blade angle α , overlap ratio t/b, and velocity ratio u/\overline{c}), Becker has applied the Gaede theory to the suction capacity of the turbomolecular pump. A comprehensive description is given by Bernhardt [16]. Considering a turbomolecular pump disk (Figure 10.12) with z pump channels between blades with blade angle α and channel height h, rotating at velocity u, it follows:

$$S_0 = \frac{1}{2} z u_{\rm C} h l k_{\rm e},\tag{10.18}$$

with

 $z = \leftrightarrow blade number.$ $u_{\rm C} = \iota u \cos \alpha = \pi f (R_{\rm o} - R_{\rm i}) \cos \alpha = \text{velocity in channel direction},$ $h = \epsilon t \sin \alpha = \frac{\pi}{z} (R_i + R_o) \sin \alpha = \text{channel height},$ $l = \epsilon R_o - R_i = \text{blade length} (R_o \text{ and } R_i, \text{ outer and inner blade radii,})$ respectively).

The factor 1/2 in Eq. (10.18) accounts for the circumstance that only half of the gas molecules show a velocity component toward the pump. $0 < k_e \le 1$ is a factor describing the ratio of pumped and backflowing particles. For blade angles between 30° and 60°, velocity ratios $0.5 < u/\overline{c} \le 1$, and optically tight blade design $(b = t/\cos \alpha), k_e = 1$ is a good approximation. These assumptions lead to

$$S_0 = \frac{\pi}{2} (R_o^2 - R_i^2) \pi f(R_o - R_i) \cos \alpha \sin \alpha.$$
 (10.19)

$$S_0 = \frac{1}{2} A u \sin \alpha \cos \alpha \tag{10.20}$$

with A the area of the pumping disk covered by the blades.

Taking into account the conductance $C = \overline{c}/4 \times A$ of the area covered by the blades, the pumping speed S_R is reduced due to the inlet conductance:

$$S_{\rm R} = \frac{CS_0}{C + S_0} = \underbrace{\frac{A(\overline{c}/4) \leftarrow}{1 + \overline{c}/(2uk_e \sin\alpha \cos\alpha)}}.$$
 (10.21)

The pumping probability P_{Ho} , also referred to as the Ho factor, is obtained by dividing this quantity by the conductance:

$$P_{\text{Ho}} = \frac{1}{1 + \overline{c}/(2uk_{\text{e}}\sin\alpha\cos\alpha)}.$$
 (10.22)

Gaede's formula, Eq. (10.7), can be used to calculate the compression ratio K_0 if it is applied to the pump channel of a turbomolecular pump. By replacing u with $u \cos \alpha$, L with b, and h with t sin, we find

$$K_0 = \exp\left(\frac{ub}{\overline{c}gt\tan\alpha}\right). \tag{10.23}$$

The blades in modern turbomolecular pumps are more or less optically tight, which means that the blade channels comply with the condition $\cos \alpha = t/b$ (Figure 10.12). Larger blade distances decrease the compression ratio. Overlapping blades reduce pumping speed and are difficult to manufacture due to the narrow pump channels. If $t/b = \cos \alpha$, Eq. (10.23) yields

$$K_0 = exp\left(\frac{u}{\overline{cg}\sin\alpha}\right). \tag{10.24}$$

The factor g in this equation is unknown (see also Eq. 10.7 and Figure 10.14). Furthermore, losses due to backflow remain unconsidered so that accurate values for compression ratio cannot be expected from this equation.

10.4.3.2 Statistical Consideration

In 1960, Kruger and Shapiro [17] developed a statistical theory of turbomolecular pumps and calculated the transmission probabilities P_{12} and P_{21} for gas particles passing through a row of blades rotating at velocity u (Figure 10.12). They assumed:

- The mean free path is greater than the distance between blades (molecular flow).
- Particles in spaces 1 and 2 show *Maxwell* velocity distribution.
- Their average thermal velocity remains constant when they pass through the
- Desorption from the blade surface follows the cosine law.

Maulbetsch and Shapiro [18] tabulated transmission probabilities for velocity ratios $0.1 < u/c_{\text{prob}} < 5$ (c_{prob} is the most probable velocity), blade distance width ratios b/t between 0.25 and 2, as well as blade angles α in the range of $10^{\circ}-60^{\circ}$.

For both theoretical approaches, Eqs. (10.9) and (10.22) may be used to compare Ho factors, and Eqs. (10.12) and (10.24) to compare compression ratios. At a blade speed $u = 400 \,\mathrm{m \, s^{-1}}$, and using nitrogen with a mean thermal velocity $\overline{c} = 470 \,\mathrm{m \, s^{-1}}$, ratios of $u/\overline{c} = 0.85$ are obtained so that Figures 10.13 and 10.14 describe realistically the conditions in modern turbomolecular pumps.

Transmission probabilities P_{12} and P_{21} were determined according to [18] from tables by interpolation for the condition $b/t = \cos \alpha$. Figure 10.13 shows Ho factors for the velocity ratios $u/c_{\text{prob}} = 1$ and $u/c_{\text{prob}} = 0.5$ versus the blade angle. Curves 2 and 4 were calculated from Eq. (10.22) with $k_{\rm e}=1$, and curves 1 and 3 were obtained using Eq. (10.9). Experimental investigations verify curves 2 and 4 and, particularly, the maximum pumping speed has proven to exist at a blade angle of 45°, as shown in the curves 2 and 4.

In Figure 10.14, compression ratios according to Eq. (10.12) are plotted against the velocity ratio u/c_{prob} . For values $u/c_{\text{prob}} < 1$, K_0 rises exponentially. For large values, curves tend to be flatter. If we adopt the curves for $u/c_{\rm prob}=1$ and $\alpha=30^{\circ}$ to statistically calculate values using Eq. (10.24), we find g = 1.438. If this factor is used to calculate K_0 according to Eq. (10.24), curves 1-5 are obtained. For $u/c_{\text{prob}} = 1$ and typical blade angles of turbomolecular pumps between 20° and 60° , these curves correspond well with the statistically calculated curves 6-10.

Using the obtained values of K_0 and $S_0 = CP_{Ho}$ as well as Eq. (10.17), the pumping speed of a turbomolecular pump can be calculated for different gases.

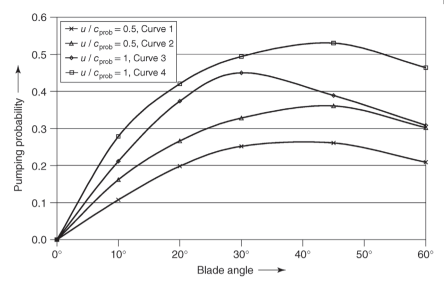


Figure 10.13 Pumping probabilities (*Ho* factors) $P_{\text{Ho}} = P_{12} - P_{21}$ versus blade angle α for optically tight blade designs. Curves 2 and 4 are calculated from geometrical data, Eq.

(10.22), for $k_{\rm e}=1$. Curves 1 and 3 are obtained from Eq. (10.9) using statistical data by $\it Maulbetsch$ [18]. Measured values fit better to curves 2 and 4.

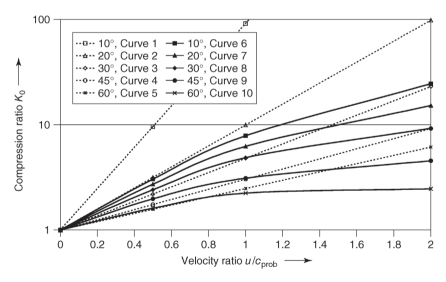


Figure 10.14 Compression ratio K_0 versus to *Gaede's* equation, velocity ratio u/c_{prob} for optically tight blade designs and selected blade angles α . Compression ratios are given by the group of straight lines (curves 1–5) calculated according for *Gaede's* equation.

to *Gaede's* equation, Eq. (10.24), and by curves 6–10 calculated according to *Maulbetsch* [18]. Matching the compression ratio for $\alpha = 30^{\circ}$ and $u/c_{\text{prob}} = 1.1$ to curve 8 yields g = 1.438 for *Gaede's* equation.

Blades with small angles α are used on the fore-vacuum side and blades with $\alpha = \leftarrow$ 45° are used on the suction side in order to maximize pumping speed.

Models used for the calculation and design of turbomolecular pumps today are based on the aforementioned molecular calculation methods. As a result of the increased meaning of the transitional range and the viscous pressure range for turbomolecular pumps, current developments are aimed at the integration of the direct simulation Monte Carlo (DSMC) methods. With these, attempts are made to include interactions between the molecules [19]. An additional advantage is the better graphic presentability of the calculation results that makes the understanding of the processes easier. However, a difficulty with the DSMC method lies in the necessary computing time [20]. With this method, for example, the influence of the rotor blade end faces could be simulated [21].

10.4.4

Thermal Balance

Due to the high rotary frequency and the accompanying high mechanical stresses, the rotors of molecular and turbomolecular pumps must be produced from a highly loadable material, for example, from aluminum alloys. In continuous operation, certain material-specific temperatures may not be exceeded because otherwise, stability losses and creep effects occur that, in conjunction with the mechanical stresses, may lead to an imbalance of the rotors and, in the worst case, to the total loss of the rotors [22].

The thermal equilibrium of the rotors in a turbomolecular pump is determined by the following effects: (1) first and foremost, gas friction heats the rotor [23] and (2) heat is produced by means of conduction, radiation, and heat flow. Thermal flow through the temperature increase of the pumped gas is negligible compared to the thermal conduction and radiation, even under the assumption that the gas takes on the temperature at each stage of the pump. This is caused by the low mass flows.

Exemplarily, the aforementioned effects should be calculated for a disk pair. In the turbomolecular pump, flows are nonvortical either in the molecular or in the viscous pressure range because the Reynolds numbers are very small. Axial gas mixing does not occur except through thermal movement of the molecules. Therefore, the formulae derived in Section 3.3.2 can be used for friction and thermal conduction.

For the calculation of the heat-generating gas friction, we use Eq. (3.77) that is a turbomolecurlar pump approximate for the rotor and stator disk pair. The frictional force FR between the two plates with A that move at a speed difference of $u = u_2 - u_1$ is valid following formula (3.77) after insertion of Eqs. (3.65) and (3.68) and resolving according to F_R :

$$F_R = \frac{4}{\pi} \times \frac{A \overline{l} p u}{\overline{c} (d + 2 \overline{l})}.$$

The frictional power P arises through the multiplication of the frictional force by the speed. Because rotary disks are considered, an integration of the radius r in R_i to R_a limits is to be performed. In addition, the relations $A = 2\pi r dr$ and $u(r) = 2\pi r f$ are to be considered, where f is the rotary frequency. Therefore, the starting point of the integration is

$$dP = dFu(r) = \frac{32\pi^2 f^2 p\bar{l}}{\bar{c}(d+2\bar{l})} r^3 dr.$$
 (10.25)

Here, the impulse accommodation factor $\sigma_t = 1$ t is set in Formula (3.65), cf. Section 3.3.2. Then, the integration yields

$$P_{\text{friction}} = \frac{2A\left(u_a^2 + u_i^2\right)p\bar{l}}{\bar{c}\pi(d+2\bar{l})} \propto \frac{1}{\frac{d}{p} + 2\text{const}},$$
(10.26)

at the blade speeds outside and inside u_a and u_i . In the last step, $\bar{l}p = \text{const}$ is used. Following Eq. (3.56), the constant is merely dependent on the temperature. The generated frictional power between the rotor and stator disks is evenly distributed between the rotor and stator. For low pressures with $2\bar{l} \gg d$, the frictional power is proportional to the pressure, for high pressures and $2\bar{l} \ll d$ in the laminar range, it is constant (Figure 10.15).

In the next step, the two mentioned heat transport mechanism should be considered for the delivery of heat in the stator. The first mechanism consists of thermal conduction through the gas between the disks. For the calculation, the relations are used in Eq. (3.94) after Eqs. (3.86) and (3.89).

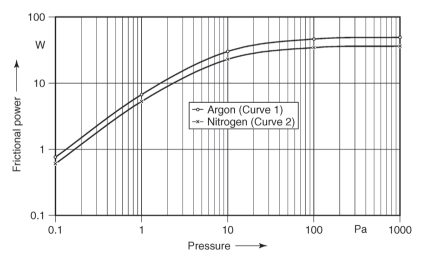


Figure 10.15 Frictional power between rotor and stator disks in a HiPace 1500 turbomolecular pump by Pfeiffer versus argon pressure (curve 1) and nitrogen pressure (curve 2), calculated from Eq. (10.27).

Resolving, thus, yields

$$P_{\text{conduction}} = \underbrace{\frac{p\overline{c}A\frac{\kappa+1}{\kappa-1}}{\left(\frac{4}{a_{\text{E}}} + 2\frac{d}{\overline{l}}\right)}}_{\text{T}_{2} + T_{1}} \times \underbrace{\frac{\overline{c}A \times \frac{\kappa+1}{\kappa-1} \times p\overline{l}}{\left(2(\overline{l}+d)(T_{2} + T_{1})\right)}}_{\text{(10.27)}},$$

where $2T = T_1 + T_2$ is used and, as explained in Section 3.3, the energy accommodation factor is set as $a_E = 1$. The isentropic coefficient κ is a measure of the degrees of freedom of the gas molecules. The pressure dependence of the power transported by means of thermal conduction takes place similar to the generated frictional power.

The second mechanism is the heat dissipation of the rotor in the stator disk through radiation. The heat loss between the two parallel disks with surface A, with distance d, and with temperatures T_1 and T_2 , according to the Stefan-Boltzmann law [24], results in

$$P_{\text{Strahlung}} = EA\sigma \left(T_2^4 - T_1^4\right) = \underbrace{A\sigma}_{1/\varepsilon_1 + 1/\varepsilon_2 - 1} \times \left(T_2^4 - T_1^4\right) \tag{3.90}$$

with $\sigma = 5.67 \times 10^{-8} \text{ W m}^{-2} \text{ K}^{-4}$.

With dull-finished aluminum alloys as they are used in turbomolecular pumps, the emission ratios $\varepsilon_1 = \varepsilon_2 = 0.3$ apply. For black-coated disks, an ε-value close to 1 can be obtained. However, these coatings generally show high gas desorption rates and are, therefore, only suitable for higher working pressures.

In the steady-state case without gas loads, the heat generated by gas friction is led away by means of thermal conduction through the gas and radiation. Then,

$$P_{\text{Reibung}} = P_{\text{Leitung}} + P_{\text{Strablung}} \tag{10.28}$$

applies under the use of Eqs. (10.26), 10.27 and (3.90) follows:

$$\frac{A(u_a^2 + u_i^2)p\bar{l}}{\bar{c}\pi(d+2\bar{l})} = \underbrace{\frac{p\bar{c}A\frac{\kappa+1}{\kappa-1}}{\kappa-1}}_{\leftarrow \frac{d}{\bar{l}}} \times \frac{T_2 - T_1}{T_2 + T_1} + \underbrace{\frac{A\sigma}{1/\varepsilon_1 + 1/\varepsilon_2 - 1}}_{(10.29)} \times \underbrace{(T_2^4 - T_1^4)}_{\sim 1}.$$

This equation cannot be solved analytically for $T_2 - T$. If for assessment, the radiation amount is neglected and $\bar{c}^2 = 8RT/(\pi M)$ and $T_1 + T_2 = 2T$ are used, then one sees the influence of the gas type on the difference in temperature:

$$T_2 - T_1 = \frac{(u_a^2 + u_i^2)M}{2R} \times \frac{\kappa - 1}{\kappa + 1}$$
 (10.30)

For gases of high molecular weight M and only three degrees of freedom, $f_g = 3$ (heavy noble gases), high temperature differences can be expected. The influence of molecular weight is significantly larger than the one from degrees of freedom.

Example 10.2

Argon:
$$M = 40$$
, $f_a = 3$, $u_o = 452$ m s⁻¹, $u_i = 151$ m s⁻¹ : $T_2 - T_1 = 182$ K!

Example 10.3

Nitrogen:
$$M = 28$$
, $f_q = 5$, $u_o = 452 \text{ m s}^{-1}$, $u_i = 151 \text{ m s}^{-1}$: $T_2 - T_1 = 76 \text{ K}$!

The temperature difference is pressure independent because both the produced frictional power and the heat transported by the gas are proportional to the pressure. However, the produced thermal power P drops at decreasing pressure (Figure 10.15) so that radiation can contribute considerably to the transport of produced thermal power at low pressures. Thus, in this case, temperatures remain below fair limits.

A single pair from the rotor and stator disks was considered above. As a general rule, turbomolecular pumps possess from 5 to 12 such pairs that work in different pressure ranges. At the same time, the largest heat quantity originates in the pair that is next toward the pump outlet. Hence, a complicated heat distribution and thermal conduction effects within the rotor occur [25]. Because turbomolecular pumps are used in processes and less in the end pressure operation and are operated under high gas loads, the temperature of the process gas can play a greater role and, in addition, efficiency losses can appear in the driving mechanism [26]. Gas type-dependent interactions between gas molecules and surface also play a role and are the subject of research as well as the deeper investigation of thermal conduction [27]. Moreover, attempts are being made to develop models for the Gaede and Holweck pumps [23].

10.5

Turbomolecular Pumps

After the physical fundamentals, the most important technical components and the practical application of the turbomolecular pumps are introduced in the following. Special weight has been given to the description of the rotor and its bearing.

10.5.1

Design and Function

In contrast to the model shown in Figure 10.2, today's turbomolecular pumps are designed as single-flow systems, that is, the first high-vacuum stage is arranged directly below the suction flange in order to reduce conductance losses.

A compound pump, that is, a turbomolecular pump with integrated molecular pumping stage, is shown in Figure 10.16. In the housing (1), the bearings (8) and (11) are supporting the rotor (3). The rotor is moved by a motor (13) in quick

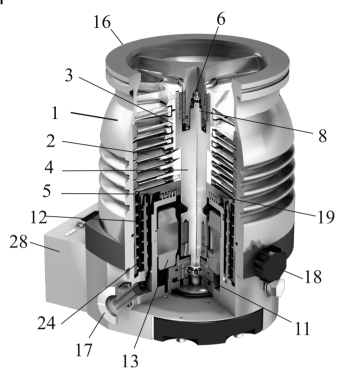


Figure 10.16 Turbomolecular pump with rotor disks shrunk onto the shaft. Bearings at both ends (UHV compatible permanent-magnet bearing on the inlet side and oil-lubricated bearing on the fore-vacuum side) provide favorable dynamic behavior of the rotor. 1: housing; 2: stator; 3: rotor; 4: rotor shaft; 5:

labyrinth seal; 6: safety bearing; 8: radial magnetic bearing; 11: lower ball bearing; 12: *Holweck* stators; 13: motor; 16: suction flange; 17: fore-vacuum port; 18: vent valve; 19: *Holweck* hub; 24: *Holweck* sleeves; 28: electronic drive unit.

rotation that is driven by flange drive electronics (28), see also 10.5.5. Between the rotor disks, which mounted on a shaft (4), diametrically split stator disks (2) are inserted. These are held at an axial distance by distance rings. The molecular stage is executed as a Holweck stage, in which a Holweck sleeve (24) attached in at least one Holweck hub (19) rotates within a Holweck stators (12). Particularly with a multistage concentric design, very high compression ratios K_0 of up to 10^8 are achieved at a low overall length whereby for low gap widths between rotor and stator, exhaust pressure can rise up to approximately 1500 Pa.

The gas enters the pump through the suction flange (16), is compressed by several turbomolecular stages, and is fed to the backing pump via the fore-vacuum port (17). Genuine turbomolecular pumps produce fore-vacuum pressures of approximately 50 Pa under high gas loads.

In compound pumps, the gas compressed by the turbomolecular stages is transported to the *Holweck* stages with lower pumping speed where it is

compressed to a fore-vacuum pressure of 100-1500 Pa. After pumping is completed, the pump can be vented via the vent valve (18).

10.5.2

Rotors of Turbomolecular Pumps

10.5.2.1 Rotor Design

Today two rotor designs are common. The first design is shown in Figure 10.16 and consists of a rotor with disks shrunk onto the shaft. Hence, these are frequently also referred to as "disk rotors." The shaft is mounted at its two ends: A permanent magnet bearing suited for ultrahigh vacuum (UHV) is found on the suction side and a ball bearing is provided on the fore-vacuum side. The second design is generally referred to as a "bell-shaped rotor" such as is shown in Figure 10.17. With this, the blades are milled from the external circumference of a cylindrical base metal. In the center of the base metal, a hollow cavity is created that contains a central shaft and houses the motor and bearing.

With both designs, a part of the fore-vacuum-side turbomolecular pump disks can be replaced by one or more concentric Holweck stages (12) in order to produce higher exhaust pressures, whereby diaphragm pumps can be used as backing pumps.

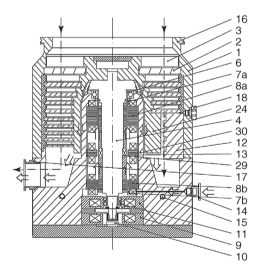


Figure 10.17 Magnetically levitated turbomolecular pump with bell-shaped rotor. The safety bearings support the rotor in case of a defect in the magnetic bearings. Pumps with ball bearings of the same design use greaselubricated bearings which require a thin shaft yielding unfavorable dynamic behavior of the rotor. 1: housing; 2: stator; 3: rotor; 4: rotor shaft; 6: safety bearing; 7: radial sensor; 8:

radial magnetic bearing; 9: axial magnetic bearing; 10: axial sensor; 11: pair of safety bearings: 12: Holweck stator: 13: motor: 14: purge-gas port; 15: cooling water channel; 16: suction flange; 17: fore-vacuum flange; 18: vent valve; 24: Holweck sleeve; 29: drive magnet; 30: motor support; \Rightarrow : purge gas flow; \rightarrow : process gas flow.

Disk rotors offer the advantage that they can be connected in parallel in order to obtain high gas throughput and fore-vacuum pressures of approximately 100 Pa. For this, openings are arranged in the Holweck hub (14) between the two Holweck cylinders, and the thread grooves in the stator are designed accordingly. A part of the Holweck hub can be used to dynamically seal the bearing range and motor range from the pumping space by means of a labyrinth seal (5).

10.5.2.2 Rotor Materials and Mechanical Requirements

Rotor disks and shafts (Figure 10.18) are made from high-strength aluminum alloys. They comply with special criteria in terms of purity and homogeneity of materials.

Carbon fiber sleeves are used for *Holweck* rotors. Their low expansion due to thermal load and centrifugal forces guarantees nearly constant inside and outside gap widths in *Holweck* stages under any permitted operating condition. These rotors reach circumferential speeds of up to 500 m s⁻¹ without exceeding tolerable material stresses.

In bell-shaped rotors, the blades are milled from solid bulk material. The large inside diameter with space for motor and bearings leads to high tangential stresses in the bell. Therefore, the rotational speed is limited. Optimum machining and shaping permits circumferential speeds of 400 m s⁻¹.

10.5.3

Safety Requirements

The stored rotation energy is substantial in the rotor of a molecular pump due to the high nominal speeds. Although the release of this energy is unlikely during normal operation, it cannot be entirely excluded. The pump manufacturers attempt to prevent instances of damage through constructive measures, for example, through the material choice and defining the operating conditions as well as through careful manufacturing and the establishment of maintenance intervals. Overloading through exceeding operational limits or the entry of objects into the pump interior can still lead to the destruction of the rotor.



Figure 10.18 Rotor and Holweck stators of a compound pump.

In the ISO Standard 27892 [28], the conceivable cases of damage, its production in the laboratory, and the subsequent documentation are established. A distinction is made between a "burst" in a bell-shaped rotor that is torn into several pieces or the shaft of a disk rotor breaks and a "crash" with which only the blades are separated from the rotor.

The torque of a rotor with the moment of inertia *J* and the nominal speed *f* is [28]

$$M = \frac{\mathrm{d}}{\mathrm{d}t}(J2\pi f) = 2\pi J \times \frac{\mathrm{d}f}{\mathrm{d}t}.$$
 (10.34)

The moment of inertia *J* can be derived from the geometry of the rotor or can be determined experimentally. In comparison to the bell-shaped rotor, the disk rotor possesses a lower moment of inertia in the same motor-force class because the masses are closer to the axis of rotation. Because the rotor is braked from nominal speed up to the shutdown, df = f can be set. The deceleration time dtcan only be determined experimentally by destroying a pump. It depends on the rotor type; extensive tests with bell-shaped rotor resulted in a few milliseconds and roughly 10-fold with disk rotors.

Example 10.4

Rotor fracture of a turbomolecular pump has a pumping speed of 1 500 ℓ s⁻¹, moment of inertia $J = 1.5 \times 10^{-2}$ kg m², and rotary frequency f = 600 Hz. With a deceleration time $dt = 10^{-2}$ s (disk rotor), the torque amounts to M = 5 655 N m, with a deceleration time dt = 3 ms (bell-shaped rotor), even $M = 18\,850$ N m. In this instance, the influence of the rotor construction on the moment of inertia was neglected.

The flanged joints are designed for the occurrence of torques in the case of damage. Compliance with all assembly instructions is mandatory. This is likewise valid for test runs.

The housings can withstand the rotor pieces due to the wall thicknesses and materials selected. Because toxic gases (e.g., PH₃ is pumped in the semiconductor production) are pumped in some applications, the housings are designed to remain sealed in the event of a crash.

Due to the moment of inertia and the slower expiry, pumps with disk rotors have advantages in comparison to pumps with bell-shaped rotors.

10.5.4

Bearing Arrangements for Rotors in Turbomolecular Pumps

Operating safety and service life of a turbomolecular pump depend highly on the quality of the bearing arrangements. This is due to a number of different demands such as high rotary frequency, high temperatures, continuous

operation, intermittent operation, arbitrary mounting position, low-vibration operation, and air-inrush protection. However, the fundamental condition for high service life is a well-balanced rotor, that is, under any operating conditions and temperature variation, balance quality G2 according to ISO 1940-1 [29] should not be exceeded.

Current state-of-the-art bearing arrangements include oil- or grease-lubricated ball bearings with ceramic balls, permanent magnet bearings made of SmCo or NeFeB, and active magnetic bearings. Often, combinations of these bearings are used. Unfortunately, none of the bearing types meets all of the desired requirements.

10.5.4.1 Shaft with Two Ball Bearings

Contamination on the high-vacuum side is not tolerable. Therefore, any lubricated bearings must be arranged at the fore-vacuum side. Such designs use bell-shaped rotors with ball bearings, (3) and (5) (Figure 10.19).

Bearings are usually filled with grease. Due to the particular operating conditions (vacuum, high rotor temperatures), special lubricating grease is required for such bearings. For sufficient service life of the bearings, maximum tolerable temperatures have to be reduced considerably compared to oil-lubricated ball bearings. Rotor dynamics require a certain minimum diameter of the shaft. The corresponding mean bearing diameter $d_{\rm m}$ and the rotary frequency f in rpm give the speed factor $d_{\rm m}f$. Due to the required continuous stability and the high rotational speeds, speed factors $> 5 \times 10^5$ mm min⁻¹ are necessary. Such values are in many cases not available with grease lubrication [30,31]. Furthermore, the limited service life of the grease has to be considered.

With oil lubrication, speed factors $>\!10^6\,\mathrm{mm\,min^{-1}}$ are obtainable. However, oil supply for the upper bearing is difficult to realize and certain mounting positions are not allowed.

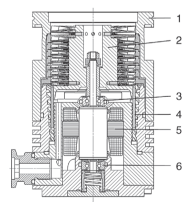


Figure 10.19 Turbomolecular pump TW 70 H with grease-lubricated ball bearings. 1: housing; 2: rotor; 3: ball bearing; 4: *Holweck* sleeve; 5: motor; 6: ball bearing.

10.5.4.2 Shaft With Permanent Magnet Bearing and Ball Bearing

Arranging the bearings of the rotor shaft at both ends eliminates the problems described in the previous section. Since the development of sintered SmCo magnets, a UHV-suitable material for permanent-magnet bearings on the high-vacuum side of pumps is available. In a turbomolecular pump according to Figure 10.16, the permanent-magnet bearing is made up of concentric, axially magnetized magnet rings, arranged so that poles of the same sign face each other [32]. The rotor rings (8a) are pressed into the rotor. Inside the rotor rings, the stator rings (8b) are stacked onto a pivot, and connected to the housing with three struts. A safety bearing (6) protects the rotor against radial movement during transport or in the event of shocks and strong vibrations. On the fore-vacuum side, a ball bearing, also absorbing axial forces exerted by the upper bearing, supports the rotor shaft. The ball bearing is oillubricated.

In small pumps ($S \le 500 \,\ell\,\mathrm{s}^{-1}$), an oil-soaked felt package is used, which gives off small amounts of oil to the bearing through a sliding contact. The felt package filters the backflowing oil. With this lubricating principle, where the oil is stored in the felt package, pumps can be arranged in any operating position.

Large pumps $S > 500 \,\ell\,\mathrm{s}^{-1}$ use oil pumps and perfluorpolyether PFPE oils for bearing lubrication. For the oil to flow back to the oil pump, operating positions are restricted to certain angles. In modern turbomolecular pumps, the oil guide is optimized with regard to a possibly very large angle position range. Special designs are available for overhead operation.

With oil lubrication, speed factors $d_{\rm m}f>10^6\,{\rm mm~min^{-1}}$ are possible under the described conditions in vacuum and with continuous operational stability. Thus, neither temperatures, nor service life of the lubricant or the existing speed factors present a problem for the service life of the bearings.

10.5.4.3 Magnetic Bearings

The requirements in terms of vacuum purity, desired maintenance-free operation, and requests for operation in any position promoted the development of rotors with noncontact magnetic-bearing arrangements. Due to the higher costs of active magnet bearings, magnetic-levitated pumps are more expensive than pumps with ball bearings. For a long time, magnet bearing systems were only attractive for large pumps. Thanks to the decreased costs for the control electronics, the distribution has strongly increased and, today, even midsize pump designs are available as magnetically mounted.

Bearings of a rotor have to stabilize motion in five degrees of freedom: two radial directions each in the planes of the bearings and, additionally, the axial direction [33]. Bearing arrangements based solely on permanent magnets are unfeasible; thus, electromagnets are used. Their electrical current is controlled using position sensors and amplifiers so that the distance between shaft and stator remains constant. In Figure 10.20, the circuit principle of a magnetically mounted pump according to Figure 10.17 is shown.

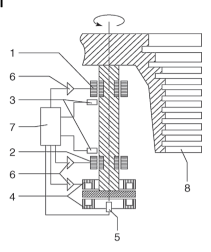


Figure 10.20 Model of an active five-axis magnetic rotor bearing system. Two additional radial sensors and lifting magnets are arranged perpendicularly to the plane of

projection. 1, 2: radial magnets; 3: radial sensors; 4: axial magnets; 5: axial sensor; 6: current amplifiers; 7: controller; 8: rotor blade.

The radial magnetic bearings (8a) and (8b) with sensors (7a) and (7b) stabilize the rotor shaft in the plane of projection as well as in the corresponding perpendicular direction by similar coils and sensors, not illustrated in the graphic. The two electromagnets (10), with a rotating disk in between, stabilize the shaft in the axial direction with sensor (7c). At the lower and upper end of the shaft, safety bearings (6, 11) prevent the rotor from touching the stator during transport, pump standstill, or in the event of shocks and strong external vibrations. A pump with noncontact magnetic bearings is also obtained by replacing the ball bearing in Figure 10.16 with the lower magnetic bearing (8b) and the axial bearing (9). Both designs are common. The regulation of rotor movement originating from the imbalance requires actuating forces that induce housing-sided vibrations. If the deviation control is abandoned, it is possible to achieve operation with low vibration [33].

Power supply and required current amplifiers as well as the motor drive are integrated into the magnetic bearing electronics, which are connected to the pump via an electric cable. In the event of a power loss during operation, the motor operates as a generator and supplies auxiliary power to the bearings down to approximately 20% of nominal speed. Subsequently, the rotor continues rotating in the safety bearings and gradually comes to a stop. In the cases of magnetic bearing electronics failure or heavy air inrush, the rotor drops into the safety bearings and will be braked by the motor. A particularly critical situation arises when the rotor coasts in the safety bearings under vacuum (in case cables are unplugged during operation) without any braking force by the motor because, then, the complete rotor energy has to be dissipated in the safety bearings. Due to high loads caused by the latter

described reasons, safety bearings show significant wear and must be replaced for reasons of precaution.

10.5.5

Drives and Handling

The high rotary frequencies in turbomolecular pumps are produced by drives using electronic frequency converters. Due to their higher efficiency, brushless direct current (DC) motors with permanent magnets mounted directly on the shaft replaced the formerly used three-phase asynchronous motors. In view of a vibration-optimized operation, brushless, sinusoidally commutated synchronous drives are also used. In brushless DC motors, no current warms up the motor. Therefore, small pumps with $S = 50-100 \,\ell\,\mathrm{s}^{-1}$ are capable of operating on convection cooling and obtain high bearing service life. Motors are available with Hall probes and, also, sensorless variants, where the rotor position is determined by the curve of the coil voltage. Sensorless systems can be more easily used when chemical effects exist or the pumps are exposed to high-energy radiation.

In applications with high-energy radiation (particle accelerators), which can destroy semiconductor components, electronic drive units have to be arranged at a safe location and long cables (up to 100 m) are used to connect them to the pump. Alternatively, asynchronous drives with mechanical frequency converters are used.

Nowadays, advances in miniaturization have promoted designs in which electronic drive units and sometimes the power supply are mounted directly to the pump in order to save an expensive cabling. A DC power supply or mains voltage is used for energy supply (Figure 10.3). Water-cooled pumps require waterprotected electronics for directly mounted systems because of water vapor condensation.

Several solutions are available for handling and controlling turbomolecular pumps (see Section 10.7): hand-operated systems, remote control via relays, and computer control through a serial interface (manufacturer-specific) or fieldbus system with standard interface, often combined with a programmable logic controller (PLC).

For operation within a system, inputs and outputs are equipped for controlling the most important functions. Adjustments are made using handheld equipment or a computer. Some manufacturers offer analog inputs with which parameters, particularly the rotary frequency, can be adjusted. Relay outputs signalize nominal speed and possible errors.

A computer over an interface, for example, a serial interface, offers the most comprehensive control possibilities such as with the RS 485. Direct communication is possible only with a manufacturer-specific transmission protocol. Universal control is supplied by a fieldbus system, which can also control other system components. This requires special fieldbus converters (Profibus, DeviceNet). Customer-specific interfaces are provided with the purchase of appropriate quantities.

10.5.6

Heating and Cooling

When very low pressures $< 10^{-6}$ Pa are produced, desorption of gases and water vapor limit the producible ultimate pressure (Chapter 6). Baking-out of the vacuum chamber and pump can reduce the desorption rate so that pressures $< 10^{-8}$ Pa can be obtained. For this purpose, a heating jacket (20) is arranged below the high-vacuum flange, which heats the housing to approximately 100 °C. Pumps with aluminum housing are not suitable for baking due to their thermal conductivity.

Large turbomolecular pumps with pumping speeds above 500 ℓ s⁻¹, for pumping process gases, are water cooled. Drive units, directly mounted to the pump, are connected to the cooling circuit as well. Pumps from 100 to 500 ℓ s⁻¹ can be either air or water cooled. In applications where vibrations present a problem, small turbomolecular pumps are often used without any forced cooling. This operation mode is possible when the pumps operate without gas load and idle power consumption is low.

10.5.7

Special Designs

Today, many turbomolecular pumps are used in process systems and analyzing equipment. Certain applications require special designs rather than standard pumps.

In semiconductor production, for example, for etching aluminum using chlorine, unused chlorine and aluminum chloride (Al₂Cl₆) appears; the latter condenses under standard operating conditions. Since aluminum components of the pumps are potentially threatened by corrosion, these parts are protected by coatings.

Bearings and motor chambers of corrosive-gas pumps are protected by feeding inert gas through port (14) (Figures 10.16 and 10.17). The flow resistance of the labyrinth seal (5) and the gap between the motor sleeve (19) and the rotor, through which the purge gas flows to the fore-vacuum port, increase the pressure in the motor chamber to several 100 Pa. Thus, a worth mentioning concentration of process gas due to diffusion against the direction of flow is not detectable in this area.

Condensation of process gases (Al₂Cl₆) can be prevented by heating (22) the fore-vacuum area. At the same time, rotor temperature increases due to gas load, and the bearings require cooling. Therefore, heat conducted to the motor and bearings is reduced by gaps (23) between the components. The temperature in the fore-vacuum chamber is controlled by means of a temperature sensor (21) and can be adjusted between 40 °C and 100 °C.

In helium counterflow leak detectors (Chapters 14 and 20), a mass spectrometer is mounted on the inlet flange of the turbomolecular pump (Figure 10.21). Depending on the pressure inside, the test piece is connected via valves to tappings or the fore-vacuum line. The turbomolecular pump has a high compression ratio for air, but not for helium. Thus, the pump avoids admission of air to the mass spectrometer and serves as a partial pressure amplifier for the mass spectrometer.



Figure 10.21 Special compound pump used in a helium leak detector. The test piece is connected via valves to ports 2 or 3 as required. The high-vacuum flange for the mass spectrometer for helium leak detection is

arranged at the back. 1: fore-vacuum connection; 2: port at Holweck stage; 3: port at turbo stage; 4: mass spectrometer flange; 5: electronic drive unit.

Certain analyzing equipment that uses mass spectrometers consists of several chambers with different pressures separated by orifices. In such applications, the so-called split-flow pumps (Figure 10.22) that possess multiple suction openings with adapted suction capacity have proved themselves. Each of these suction openings is connected with one of the chambers, so that with a fore-pump and a



Figure 10.22 Split-flow pump for mounting onto an LC-MS analyzing chamber.

split-flow pump, all chambers can be evacuated [34]. The skill of the vacuum pump manufacturer is evident in the layout of the pumping system of the split-flow pump [35].

10.6 Performance Characteristics of Turbomolecular Pumps

Users are particularly interested in pumping speed S, gas throughput q_{pV} , ultimate pressure or base pressure $p_{\rm b}$, compression ratio K_0 , as well as vacuum purity. Measurement of these data is described in international standard ISO 5302. Moreover, a new basic standard is available for measuring performance characteristics of vacuum pumps: ISO 21360.

10.6.1 **Pumping Speed**

Pumping speed S is the volume of gas flowing through the inlet of the pump per unit time. To measure the pumping speed, constant gas flows $q_{pV} = p S$ are admitted to a test dome and the pressures in the dome are recorded (Section 15.5).

For gases with a molecular weight of less than 50, pumping speeds do not vary significantly. With appropriate staging, the pumping speed for hydrogen can reach up to 80% of the value for nitrogen. For heavy gases (Ar), pumping speeds drop due to the higher inlet conductance of the pump. Figure 10.23

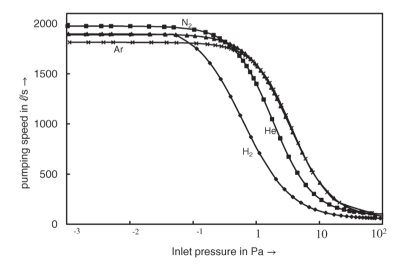


Figure 10.23 Pumping speeds of a turbomolecular pump TPH 2101 (Pfeiffer Vacuum). For lighter gases (H_2, He, N_2) the pumping speeds are nearly equal. Due to the lower impingement rate (mean thermal velocity) the pumping speed of argon is less.

shows the pumping speed of the genuine turbomolecular pump TPH 2101 for the gases hydrogen, helium, nitrogen, and argon. In the molecular flow range below 0.1 Pa, the pumping speed is constant and, between 0.1 and 10 Pa, it drops rapidly. This is caused by the compression ratio, which decreases in the transition to laminar flow. Using larger backing pumps shifts the drop in pumping speed toward even higher pressures. In the transition to laminar flow, the pumping speed of compound pumps is less dependent on the size of the backing pump because the preceding Holweck stages act as a large backing pump.

10.6.2

Compression Ratio, Ultimate Pressure, and Base Pressure

The maximum pressure against which a turbomolecular pump compresses is referred to as fore-vacuum tolerance p_c . In standard turbomolecular pumps, this pressure is in the range of 10-100 Pa, and up to 2000 Pa in compound pumps. If base pressures $p_{\rm b} < 10^{-8}$ Pa are desired, compression ratios exceeding the ratio p_c/p_b by approximately a factor of 10 are required. p_c is composed of the partial pressures of a number of gas species (N2, O2, H2O, H2). Therefore, the compression ratios for the individual partial pressures must comply with this condition as well. Base pressures listed in manufacturer's catalogues are often higher than the minimum producible ultimate pressures obtainable according to the above criteria. ISO 5302 requires the base pressure to be obtained after a maximum baking time of 48 h and an additional 48 h of waiting time.

Example 10.5

A pump unit including a compound pump and a diaphragm pump ($p_F = 500 \text{ Pa}$) shall produce an ultimate pressure of less than 10^{-8} Pa. Thus, for N_2 , a compression ratio of 5×10^{11} is required.

Desorbed hydrogen gas and water vapor from the recipient can accumulate on the fore-vacuum side and, thus, limit the ultimate pressure. Therefore, gas ballast is used in diaphragm pumps in order to reduce partial pressures of these gases. Due to the low hydrogen content of the atmosphere, hydrogen compression ratios of 10⁴ are sufficient for an ultimate pressure of 10⁻⁸ Pa in compound pumps. Higher compression ratios do not have any advantages over those calculated according to these criteria. In fact, increased technical expenditure would make such pumps more expensive.

Compression ratios are obtained by measuring the high-vacuum-sided partial pressure rise while gas is fed to the fore vacuum. Figure 10.24 shows typical curves for turbomolecular and compound pumps. While the former require a fore-vacuum pressure of 10-100 Pa, a pressure of 500-1000 Pa is sufficient for compound pumps.

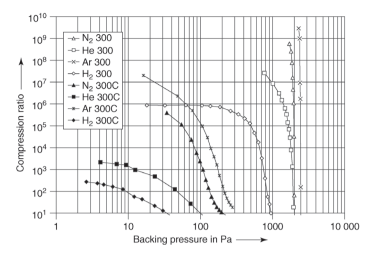


Figure 10.24 Compression ratios of turbomolecular and compound pumps. The left-hand group of 300C curves shows data for a corrosion-resistant turbomolecular pump (*HiPace* 300C) with turbo stages only. The right-hand 300 group shows values for a compound

pump (*HiPace* 300). Both have pumping speeds of $300 \ \ell \ s^{-1}$. Compression ratios in a compound pump are higher because fore-vacuum tolerance is higher than in a turbomolecular pump.

10.6.3 Pump-down Times for Vacuum Chambers

Turbomolecular pump systems without a high-vacuum valve can be used to evacuate vacuum chambers. Both pumps are started simultaneously at atmospheric pressure. Initially, the backing pump determines the speed of evacuation. Around 1000 Pa, the turbomolecular pump starts to accelerate and reaches nominal rotational speed within several minutes (run-up time). Figure 10.25 shows pressure—time curves for the evacuation of a $100-\ell$ vacuum chamber, using a turbomolecular pump with a pumping speed $S = 210 \ell$ s⁻¹ (flange size DN 100) with differently powerful drive units and different types of backing pumps. Apparently, the backing pump alone determines the pump-down speed down to a pressure of approximately 100 Pa. Different run-up times of the turbomolecular pump due to the selected drive unit (P = 120 W, 260 W) result in time differences of approximately two minutes till 10^{-2} Pa.

Pump-down time t between two pressures $p_{\rm A}$ and $p_{\rm E}$ in a chamber with volume V is calculated from:

$$t = \frac{V}{S} \ln \frac{p_{\rm A}}{p_{\rm E}}.\tag{10.38}$$

The pumping speed *S* has to be taken from the values of the pump in the corresponding pressure range.

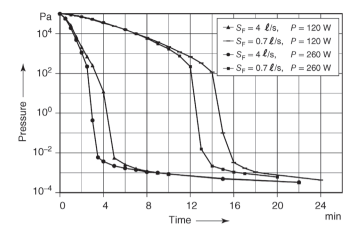


Figure 10.25 Pump-down curves of a 100-€ vessel for different backing pumps and a compound pump with high and low driving powers. The run-up time of the turbomolecular

pump strongly influences the pump-down time. In the 10⁻³ Pa range, desorption of water vapor in particular prevents a further significant drop in pressure.

Example 10.6

Pumping unit with a rotary vane pump ($S = 4 leq s^{-1}$ in the pressure range between 10^5 and 10^2 Pa) and a turbomolecular pump ($S = 200 \,\ell$ s⁻¹ in the pressure range between 10² and 10⁻² Pa) connected to a vacuum chamber with the volume $V = 100 \, \ell$. The times calculated from Eq. (10.38) for the rotary vane pump are

$$t_1 = \frac{100}{4} \ln \frac{10^5}{10^2} = 173 \,\mathrm{s},$$

and for the turbomolecular pump,

$$t_2 = \frac{100}{200} \ln \frac{10^2}{10^{-2}} = 4.6 \text{ s}.$$

Depending on the drive unit, run-up time varies between 40 and 160s. Thus, this time strongly affects the pump-down time. By using a bypass line with forevacuum and high-vacuum valves, the turbomolecular pump operating at nominal rotary frequency can be connected via the valves when 100 Pa are reached, and the pump-down time t_2 is reduced to the value calculated above.

10.6.4

Pumping of High Gas Throughputs

In vacuum process technology, considerable amounts of gas $q_{pV} = S p$ are often pumped continuously. Sp is also referred to as throughput or pV flow (Figure 10.26). In the range of constant pumping speed, throughput rises linearly with pressure. It continues to increase slowly as pumping speed drops, and then

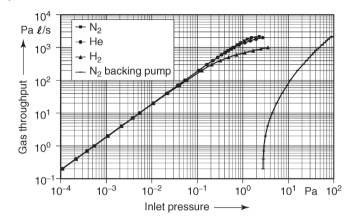


Figure 10.26 Throughputs or pV flows in a turbomolecular pump. For inlet pressures > 2 Pa, rotating speeds are instable due to the limited power for rotor protection. Thus, curves are not drawn beyond this point.

merges into the characteristic curve of the fore pump. The associated power consumption rises linearly with pressure in the molecular flow regime and approaches a limit in the laminar regime. In addition, the rotor temperature rises with power consumption and must be limited to approximately 120 °C for safe operation (Figure 10.27).

Unfortunately, inexpensive methods for measuring and monitoring rotor temperatures are not available. Certain manufacturers, therefore, monitor the temperature of the ball bearings to protect the rotor; others limit power

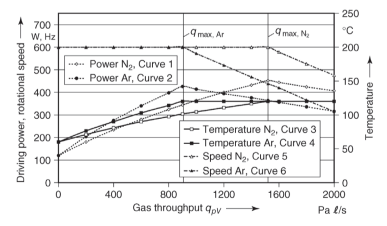


Figure 10.27 Rise in rotor temperature due to gas load. As gas load increases, driving powers (curves 1 and 2) as well as rotor temperatures (curves 3 and 4) increase. When the latter reaches the maximum value (120 °C) and gas load still increases, the driving power and also

the rotor speed are reduced far enough so that rotor temperature is kept roughly constant (curves 5 and 6). For heavy noble gases (argon), the maximum gas load $q_{\rm max}$ at nominal speed is lower than for polyatomic gases (nitrogen).

consumption as soon as the gas load exceeds maximum tolerable limits. For heavy noble gases (Ar), the maximum tolerable rotor temperature is reached at lower power consumption (see Section 10.3.4) than that for polyatomic gases (N_2) . Therefore, gas-specific limits for power consumption can be set.

The two vertical lines in Figure 10.27 represent the maximum throughputs at nominal rotational speed. Rotational speed fluctuates to the right of these lines. Therefore, a speed below the fluctuating frequencies is adjusted for higher gas throughputs. Thus, a speed-stable operation is obtained, that is, constant input pressure at constant throughput, but pumping speed is reduced.

10.7

Operation and Maintenance of Turbomolecular Pumps

10.7.1

Backing Pump Selection

The pumping speed of the backing pump shall lie between 1 and 10% of that of the turbopump. Larger backing pumps should be used for larger vacuum chambers and higher gas throughputs. Additionally, manufacturer's guidelines should be respected.

For genuine turbomolecular pumps, two-stage rotary vane pumps or dry forepumps should be used with an ultimate pressure < 10 Pa. Compound pumps require less powerful fore-pumps due to their higher fore-vacuum tolerance (critical backing pressure). If the recipient volume is less than $20 \,\ell$ and the pump-down time is irrelevant, a diaphragm pump with $S = 1 llet s^{-1}$ is sufficient for a compound pump with a pumping speed of $500 \,\ell\,\mathrm{s}^{-1}$.

10.7.2

General Notes

To avoid conductance losses, the inlet flange of the turbomolecular pump shall be directly mounted to the recipient. During installation, cleanness is important. UHV applications call for exclusive use of metal seals. Splinter shields can be mounted into the inlet flange to prevent parts from falling into the pump. However, such measures reduce pumping speed by 5–20%. For vibration-sensitive applications, anti-vibration bellows can be mounted between the pump and the recipient.

Manufacturer's installation instructions should be followed carefully due to the risks involved with possible unpredictable rotor fracture.

10.7.3

Startup

Generally, in case of an oil-sealed backing pump a turbomolecular pump should be started simultaneously with the backing pump to prevent oil-vapor backflow

to the recipient, but also in case of a dry backing pump this is an appropriate operation. Delayed starting of the turbomolecular pump is recommended only for pump-down times of the backing pump of more than 10min, as calculated using Eq. (10.38).

10.7.4

Obtaining Base Pressure

For desired pressures $< 10^{-6}$ Pa, the turbomolecular pump and the recipient require baking. For this, pumps are equipped with special heating jackets that produce tolerable pump temperatures. If the baking temperature for the recipient is higher than for the pump, the radiation power transmitted to the pump must be calculated according to Eq. (10.30). Radiation must be limited in compliance with manufacturer's guidelines. Depending on the temperatures, heating periods of 3-48 h are required to produce base pressures $< 10^{-8}$ Pa. Residual gas should then only contain masses 2 (hydrogen), 18/16 (water), 28 (CO), and 44 (CO₂). These constituents desorb continuously from the metal surface of a stainless steel recipient.

10.7.5

Operation in Magnetic Fields

Magnetic fields induce eddy currents in the rotating parts of a pump, which increase the temperature of the rotor. Because heat radiation is the only way of transporting the dissipated energy, magnetic induction must be limited to values between 5 and 10 mT by ferromagnetic shielding.

10.7.6

Venting

Turbomolecular pumps should be vented after shutdown in order to avoid back diffusion of hydrocarbons through the pump from the fore-vacuum side. For gas inlet, special venting ports are arranged between the lowest turbine disks or, in compound pumps, above the Holweck stage. For venting, automatic devices are available that vent for a predefined period only, after the speed drops below an adjustable speed-switch point. This also prevents accidental venting of the vacuum system in the case of short-time power loss. Venting with dry inert gas instead of ambient air avoids water vapor input to the pump. For restart, this measure reduces the time to reach the desired ultimate pressure. Additionally, it prevents chemical reactions between water vapor and possible deposits from the vacuum processes.

Mechanical bearings in turbomolecular pumps are designed to withstand accidental venting through large openings without showing any immediate defect. However, in such situations, the bearings for a short time are overloaded heavily, which reduces their service life. Therefore, for normal operation, certain gradients in pressure rise should not be exceeded during venting.

Axial forces on the bearings rise during venting and generate higher currents in the magnetic bearings (helicopter effect). If the current in the axial bearing of a pump reaches a limit, venting is interrupted, and is not continued before the current drops below a lower limit. Thus, the pump is rapidly vented but without any safety-bearing contact.

10.7.7

Maintenance

Turbomolecular pumps are usually used within very expensive installations (coating systems, semiconductor production, particle accelerators) where downtimes produce high costs. Users, therefore, often call for maintenance-free pumps. The minimum request though is that preventive maintenance guarantees reliable operation during a production period.

First, we will focus on pumps with ball bearings. When using hydrocarboncontaining oil, the oil should be replaced according to manufacturers' guidelines. Fluorinated oils do not require replacement under clean operating conditions because they are not subject to any aging. When running processes that produce contamination, special determination of oil-change intervals is advisable. Additionally, if necessary, bearings can also be replaced easily as long as this does not require rebalancing the rotor.

Grease-lubricated bearings have to be replaced after the service life of the lubricant because regreasing is not possible without risking a loss in service life. In pumps with bell-shaped rotors and two ball bearings, parts of the rotor shaft then must be disassembled which might require rebalancing of the rotor.

Safety bearings in pumps with magnetic levitation are subject to wear. They are designed to withstand approximately 20 rotor breakdowns at full speed and 200 gradual run-outs during long-lasting power failures. If such situations arise more often, safety bearings should be replaced preventively. Some manufacturers indicate the wear by monitoring the rotor-running period in the bearings.

In processes with heavy dust generation, the bearing chamber of a pump can be protected by a flow of purge gas in order to prevent contamination. Dust accumulating on rotor blades can cause increasing unbalance and large amounts of dust can clog pump channels. Then, dismounting and cleaning of the pump after certain process-dependent operating periods is necessary.

10.8 Applications

Today, the main areas of applications are vacuum process technology, analytics, and physical research. Fields of application and techniques are listed in Table 10.1. Requirements on turbomolecular pumps are particularly high in vacuum process technology. Overheating of rotors due to gas throughput at

Table 10.1 Applications of turbomolecular pumps.

Field of application	Operating pressure (Pa)	Process	Requirements
Analytics			
Materials and surface analytics	10^{-5}	AUGER, SIMS, LAMMA	High-purity vacuum, low vibration
Environmental analytics, medical science, process analytics	⟨10 ⁻² ⟩←	Mass spectrometry. LCMS, GCMS, ICPMS, TOF	Small light-weight pumps, low power losses, special pumps: split-flow with several ports
Electron microscopy	$\langle 10^{-2} \rangle \leftarrow$	REM, FEM	High-purity vacuum, low vibration
Leak detection	$\langle 10^{-2} \rangle \leftarrow$	Helium mass spectrometry	Small, light-weight, several ports
Industrial			
Tube and bulb manufacturing	10^{-4}	Automatic systems for evacuating, coating, and sealing	Frequent venting, vibration, and dust-particle tolerance
Vacuum locks	10^{-2}	Vacuum production	Frequent venting, quick pumping
Optical surface treatment, metallization	10 ⁻¹	Sputtering, physical vapor deposition (PVD)	High pumping speed, high gas loads, high-purity vacuum
Wear protection	10^{-1}	Hard coatings	High temperatures, dust. and vapor tolerance
Decorative coatings	10^{-1}	PVD	High gas loads
Solar cells	10^{-1}	PVD, CVD	High pumping speed, high gas loads
Semiconductor manufacturing	10 ⁻⁷ -10 ⁻¹	Ion implantation, dielectric layers, PVD, CVD etching, RIE	High-voltage operation, corrosion protection, high gas loads, heated fore-vacuum chamber
Research and Develop	ment		
Miscellaneous	10 ⁻⁸	UHV production	High compression ratio, metal seals, case heating
Accelerators	10^{-4}	Vacuum production	Radiation tolerance
Nuclear fusion	$10^{-4} - 10^{-1}$	Pumping of H_2 and He	Tritium tolerance, metal seals

required process pressure has to be avoided (see Section 10.6.4). Here, sufficiently large backing pumps should be selected. Manufacturers provide corrosion-protected pumps for pumping corrosive or toxic gases. In such applications, purge gas is used to protect the motor and the bearings. If condensing reaction products appear, the fore-vacuum chamber should be heated (see Section 10.5.7). Test runs are recommended for optimizing vacuum processes featuring cycles with different gas species and gas throughputs. Water cooling should be applied when high gas throughputs are expected.

Installations for semiconductor manufacturing as well as coating of foils, solar cells, and architectural glass include systems of 10 to 50 m length and up to 8 m width, operating 10 to 80 turbomolecular pumps (Figure 10.28). For the analytics, liquid chromatography mass spectrometry (LCMS) and gas chromatography mass spectrometry (GCMS) as well as leak detectors are produced in large scale. Such appliances are designed as desktop units for environmental analysis laboratories, food processing analytics, and medical examinations.

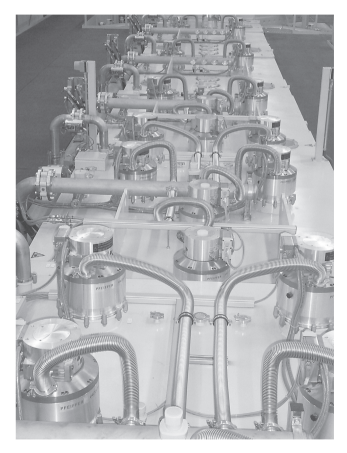


Figure 10.28 Aluminum foil coating system for high light reflection with TPH 2301 turbomolecular pumps. In this so-called air-to-air system, the aluminum foil is supplied into the continuous coating process system from air

and transported back out to air. Several layers of aluminum enhance the reflection of light to 95% which increases the effectiveness of lights by 10-20%.

Turbomolecular pumps used in physical research are often exposed to considerable doses of high-energy radiation. Such radiation can destroy semiconductor components as well as certain plastics. Electronic drive units should, therefore, be arranged at a safe distance and connected to the pumps via long cables. Otherwise, mechanical frequency converters are used.

Ready-for-use pumping units are available for evacuating vacuum chambers for numerous applications. These systems can be equipped with different types of turbomolecular and backing vacuum pumps.

References

- 1 DIN 28400, Teil 2 (1980) Vakuumtechnik Benennungen und Definitionen, Beuth Verlag, Berlin.
- 2 Becker, W. (1958) Eine neue Molekularpumpe (A new molecular pump). Vakuum-Tech., 7, 149-152.
- 3 Gaede, W. (1913) Die Molekularluftpumpe. Ann. Physik (Leipzig), 41, 337-380.
- 4 Bijker, M.D., Bosch, R.C.M., and Conrad, A. (2011) Aspects of fast load lock designs in vacuum manufacturing equipment. J. Vac. Sc. Technol. A, 29 (1), 011010.
- 5 Shirinov, A. (2010) Hochvakuumpumpe gegen Atmosphäre. Vakuum in Forschung und Praxis, 22, 10-14.
- 6 Chew, A.D. (2008) EPX die Ein-Pumpen-Lösung. Vakuum in Forschung und Praxis, 20 (6), 6-11.
- 7 Chew, A. et al. (2011) Improvements in the performance of turbomolecular pumps. Vakuum in Forschung und Praxis, 23 (3), 14-18.
- 8 Holweck, F. (1923) Helicoidal molecular pump. C. R. Acad. Science, Paris, 177, 43.
- 9 Sharipov, F., Fahrenbach, P., and Zipp, A. (2005) Numerical modelling of Holweck pump. J. Vac. Sci. Technol. A, 23, 1331-1339.
- 10 Boulon, O. and Mathes, R. (2001) Flow modelling of a Holweck pump stage in the viscous regime. Vacuum, 60, 73-83.
- 11 Siegbahn, M. (1943) A new design for a high vacuum pump. Arch. Math. Astr. Fys., 30B, p. 261.
- 12 Giors, S., Campagna, L., and Emelli, E. (2010) New spiral molecular drag stage design for high compression ratio,

- compact turbomolecular-drag pumps. J. Vac. Sc. Technol. A, 28 (4), 931-936.
- 13 Chew, A.D., Galtry, M., Livesey, R.G., and Stones, I. (2005) Towards the single pump solution: Recent development in high speed machines for dry vacuum pumping. J. Vac. Sci. Technol., A23, 1314.
- 14 Sixsmith, H. and Altmann, H. (1977) A Regenerative Compressor. J. Eng. Ind., 99, 637-647.
- 15 Knudsen, M. (1915) Das Cosinusgesetz in der kinetischen Gastheorie. Annalen der Physik, 48, 1113-1121.
- 16 Bernhardt, K.H. (1983) Calculation of the pumping speed of turbomolecular vacuum pumps by means of simple mechanical data. J. Vac. Sci. Technol., A1 (2), 136-139.
- 17 Kruger, C.H. (1960) The Axial-Flow Compressor in the Free-Molecular Range, Naval Research Report Contr. Nonr-1841 (55).
- 18 Maulbetsch, J.S. und Shapiro, A.H. (1963) Free-Molecule Flow in the Axial-Flow Turbo-Vacuum Pump, Final Report, Office of Naval Research Contract Nonr-1841(55).
- 19 Bird, G.A. (1994) Molecular Gas Dynamics and the Direct Simulation of Gas Flows, 2. Auflage, Oxford Science Publications.
- 20 Alexander, F.J. and Garcia, A.L. (1997). The direct simulation Monte Carlo method. Computer in Physics, 11 (6), 588-593.
- 21 Bird, G.A. (2011). Effect of inlet guide vanes and sharp blades on the performance of a turbomolecular pump. J. Vac. Sci. Technol. A, 29 (1), 011016-1-011016-6.

- 22 Kammer, C.A. (1998) Aluminium Taschenbuch Bd. 1, Grundlagen und Werkstoffe, Aluminium-Verlag Düsseldorf, ISBN 978-3870172749, p. 152 ff.
- 23 Cerruti, R., Spagnol, M., and Helmer, J.C. (1999) Power dissipation in turbmolecular pumps at high pressure. J. Vac. Sci. Technol. A, 17 (5), 3096-3102.
- 24 VDI-Wärmeatlas (1997) Berechnungsblätter für den Wärmeübergang, Verein Deutscher Ingenieure, Springer.
- 25 Blumenthal, R., Hillegas, S., Kalisch, D., Kuhn, M., and Latam, C. (2005) Vacuum requirements for advanced deposition processes. NanoS. The Nanotech Journal, 1, 28-32.
- 26 Froitzheim, M. and Heppekusen, J. (2008) Erweiterter einsatz von turbomolekular vakuumpumpen durch beschichtungen. Vakuum in Forschung und Praxis, 20 (3), 20 - 28.
- 27 Sharipov, F. and Bertoldo, G. (2006) Heat transfer trough a rarefied gas confined between two coaxial cylinders with high radius ratio. J. Vac. Sci. Technol. A, 24 (6), 2087-2093.

- 28 ISO, 27892 (2010) Measurement of rapid shutdown torque, International Organization for Standardization, Geneva, Switzerland.
- **29** Deutsche Norm DIN ISO 1940-1 (2003) Anforderungen an die Auswuchtgüte starrer Rotoren.
- 30 Produktinformation 5.429d, Firmenschrift: Klüber Lubrication München (Ausgabe 05.2000).
- 31 GfT-Arbeitsblatt 3, Wälzlagerschmierung, Gesellschaft für Tribologie e.V. Moers
- 32 Marinescu, M. (1982) Dauermagnetische Radiallager, Firmenschrift: Marinescu Ing.-Büro für Magnettechnik, Frankfurt/ Main.
- 33 Schweitzer, G., Bleuler und, H., and Traxler, A. (1994) Active Magnetic Bearings, Hochschulverlag AG an der ETH Zürich.
- 34 Conrad, A. and Ganschow, O. (1996) The impact of analytical instrument technology to vacuum pumping systems. Vacuum, 47, 575-672.
- 35 Conrad, A. and Ganschow, O. (2003) Vakuumpumpsystem, Deutsches Patent DE 43 31 589 C2.

11

Sorption Pumps

Dr. Karl Jousten

Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

In this chapter, the reader will learn about the working principles and characteristics of adsorption pumps, titanium sublimation pumps, getter pumps, particularly nonevaporable getter (NEG) pumps, and ion getter pumps. Physical fundamentals are covered in Chapter 6.

11.1 Introduction

Sorption pumps are arrangements in which impacting gas particles are bound to appropriate surfaces due to sorption (compare Chapter 6). This reduces the gas pressure in the container. Thus, sorption pumps act as gas traps without actually transporting the gas through the pump as in the true sense of the word. Sorption pumps are used throughout the entire vacuum pressure range, but chiefly in UHV technology, to produce hydrocarbon-free vacua.

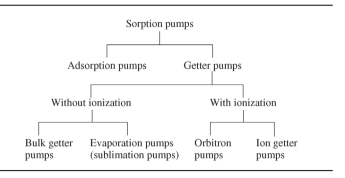
ISO 3529/2 [1] and DIN 28400, part 2 [2], differentiate between (Table 11.1) adsorption pumps (Section 11.2), getter pumps 11 *getter* derived from *to get* (Section 11.3), sublimation pumps (Section 11.3.3), and ion getter pumps, the latter divided into evaporation ion pumps (Section 11.5) and sputter ion pumps (Section 11.4), as well as the cryosorption pumps covered in Chapter 12.

For pumping action, *adsorption pumps* use the effect that certain solids, particularly at low temperatures, bind considerable amounts of gas. Solids with very large specific surface areas bind gas mostly due to physisorption. The pumps are used in the low and medium-high vacuum range.

A *getter pump* is a sorption pump in which gas binds to a getter material mainly due to chemisorption. This material is usually a metal or metal alloy, either bulk material or a freshly deposited thin layer.

Pumps that not only adsorb the gas at a surface but also use the effect of gas diffusing into a compact getter material are referred to as *bulk getter pumps*. For example, to improve or sustain vacua in small sealed systems such as CRTs, certain

Table 11.1 Classification of sorption pumps according to sorption principles.



solids are enclosed in the system as getter material that sorbs gases and vapors. Today, NEG pumps represent the predominant type of bulk getter pumps.

In contrast, so-called *evaporation getter pumps* (sublimation getter pumps) adsorb gas at the surface of continuously or intermittently fresh-deposited thin getter surfaces.

Bulk and evaporation getter pumps are not capable of pumping noble gases and other gases that are relatively passive in their chemical behavior such as methane. Ion getter pumps were developed in order to be able to pump such gases as well.

Ion getter pumps include an additional electrode arrangement that ionizes the gas particles and lets them bombard the getter surface by means of an acceleration voltage. These pumps include so-called *orbitron pumps* and *sputter ion pumps*. The latter are of considerable practical relevance and therefore covered separately in Section 11.4.

Orbitron pumps no longer have any commercial importance and are therefore covered at the end of this chapter. All *getter pumps* are used to produce high pumping speeds at relatively low pressures (low gas loads), for example, in UHV processing systems, particle accelerators, space simulation chambers, and surface analysis equipment. The latter particularly benefit from the vibration-free operation of this pump type.

Pumping action in *ion pumps* is produced solely by ionizing, accelerating, and implanting gas particles into solid surfaces by means of electrical fields. Due to low pumping speeds, they are not used in practice. However, the operating principle is used in *ion getter pumps* and *sputter ion pumps*, in addition to the getter effect, in order to also pump noble gases and other gases that are difficult to getter.

11.2 Adsorption Pumps

11.2.1

Working Principle

Certain porous substances, mainly activated carbon and zeolite, show very large specific surface areas (i.e., with respect to mass m of the porous solid) in the range of

 $A_{\rm m} = A/m \approx 10^6 \, {\rm m}^2 \, {\rm kg}^{-1}$. Thus, the adsorption capacity for gas at the inner surfaces of these substances (Sections 6.2 and 6.3) is substantial. According to Example 6.1, a monatomic layer of adparticles covering a surface corresponds to a particlenumber surface density $\overline{n}_{\text{mono}} = 10^{15} \text{ cm}^{-2}$, that is, approximately 10^{15} particles lie closely packed on each square centimeter. This corresponds to an adsorbed surfacerelated amount of substance $\nu_{\rm mono} = \overline{n}_{\rm mono}/N_{\rm A} \approx 10^{15} \, {\rm cm}^{-2}/(6 \times 10^{23} \, {\rm mol}^{-1}) =$ 1.7×10^{-9} mol cm⁻² (N_A is Avogadro's constant). The equation of state, Eq. (3.19), pV = nkT, provides the adsorbed surface-related pV-amount:

$$\tilde{b} = \frac{pV}{A} = \frac{N}{A}kT = \tilde{n}kT. \tag{11.1}$$

Multiplying with the specific surface A_m of the adsorbent yields the adsorbed mass-related pV-amount

$$\tilde{\mu} = \frac{pV}{A} \times \frac{A}{m} = \tilde{n}kT A_{\rm m}. \tag{11.2}$$

Equations (11.1) and (11.2) show that the temperature at which pV-amounts are measured must be stated together with values of these quantities. Therefore, it is useful and usually necessary to relate pV-amounts to standard temperature $T_{\text{extrmn}} = 273.15 \,\text{K}$ in order to obtain definite values. However, the reduction factor for calculating from room temperature to standard temperature 293/273 = 1.07 and its reciprocal are often insignificant because the quantities p (in particular) and V show a measuring uncertainty of more than 7%. Equations (11.1) and (11.2) thus read

$$\tilde{b}_{\rm n} = \frac{(pV)_{\rm n}}{A} = \tilde{n}kT_{\rm n} \tag{11.3}$$

and

$$\tilde{\mu}_{\rm n} = \tilde{n}kT_{\rm n}A_{\rm m}.\tag{11.4}$$

With

$$\begin{array}{ll} \overline{n}_{\rm mono} & = & 10^{15}~{\rm cm^{-2}}~{\rm and}~A_{\rm m} = 10^{6}~{\rm m^{2}~kg^{-1}},~{\rm we~find} \\ & \tilde{b}_{\rm n,mono} \approx 38~{\rm Pa}~\ell~{\rm m^{-2}}, \\ & \tilde{\mu}_{\rm n,mono} \approx 3.8~\times 10^{7}~{\rm Pa}~\ell~{\rm kg^{-1}}. \end{array}$$

At low temperatures, the probability for a gas atom that sticks to the surface to desorb back into the gas volume is lower than that at high temperatures. Therefore, the ability to capture gas is higher at low temperatures than at high temperatures. If the adsorption mass in contact with the recipient volume is cooled, it binds more gas than at normal temperature. Thus, the efficiency of the adsorption pump increases. After the adsorption process has completed, a valve is shut between the adsorption pump and the recipient. On warming of the adsorbent (usually warming to room temperature is sufficient), the gas adsorbed at lower temperature escapes through a venting valve (safety valve 3 in Figure 11.1).

Example 11.1

Calculate the pressure in an adsorption pump, which was sealed after pump-down, and reheated to room temperature. The pressure in the evacuated container with volume $V_1 = 30 \, \ell$ was reduced from atmospheric pressure p_1 to the pressure $p_2 = 1 \, \text{kPa}$ (10 mbar). The adsorption pump has an inherent volume $V_2 = 1 \, \ell$, which is filled about half way with adsorbent.

At room temperature, the *Boyle–Mariotte* law applies, Eq. (3.16), pV = constant:

$$\begin{split} &p_1(V_1+V_2/2) = p_2V_1 + pV_2/2, \\ &p = \frac{p_1(V_1+V_2/2) - p_2V_1}{V_2/2} = 6.04 \times 10^6 \text{ Pa (60.4 bar)} \end{split}$$

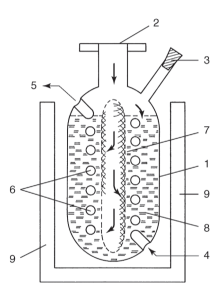


Figure 11.1 Diagram of an adsorption pump. 1 stainless-steel pump chamber, 2 inlet flange (suction port), 3 safety bung, 4 coolant inlet, 5 coolant discharge, 6 cooling coil, 7 tube sieve,

8 adsorbing medium (e.g., zeolite), 9 coolant vessel. Arrows indicate gas flow pumped by adsorption.

11.2.2

Design

An adsorption pump (Figure 11.1) is basically built up of a vacuum-tight container filled with adsorbing medium (adsorbent). A cutoff valve is placed

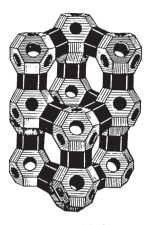


Figure 11.2 Model of an x-type molecular sieve structure [3].

in the tube connecting to the recipient. Generally, commercial adsorption pumps all use synthetic zeolite. Zeolites are M-aluminum silicates with M denoting sodium, calcium, or lithium. They are produced synthetically in large scale and serve as molecular sieves for separation of mixtures. These utilize the principle of characteristic adsorption for the individual components of the mixtures. Figure 11.2 shows the structure model of a zeolite crystal with cages and pores for gas adsorption. Using activated carbon is dangerous. If a sudden inrush of air occurs, the released heat of adsorption can heat the carbon rapidly so that it may explode in a reaction with atmospheric oxygen.

For cooling the adsorbent, the container is placed into a Dewar vessel filled with liquid nitrogen. If cooling of the adsorbent is interrupted after pumping and sealing, an automatic safety valve must provide a means for the released gas to escape. The simplest solution is a rubber bung in a tube (Figure 11.1).

A number of technical improvements are required to optimize the activity of the adsorbent. For rapid cooling and removal of the released adsorption heat during pumping, homogeneous cooling of the adsorbent is necessary [4]. Due to the low thermal conductivity of typical adsorption media, the pump body contains a system of cooling pipes or plates reducing the paths for heat transport. Additionally, an even distribution of gas load onto the complete surface of the adsorbent is mandatory [5]. For this, flow channels made of wire netting increase the surface area available for direct contact between gas and adsorbent. Furthermore, they reduce the flow paths through the closely packed adsorbent (Figure 11.1). Heating equipment regenerates the adsorbent, for example, an electrically heated jacket stretched around the outside of the pump body, which heats the adsorbent to 250-350°C.

11.2.3

Ultimate Vacuum and Pumping Speed

Ultimate Pressure with a Single Adsorption Pump

Figure 11.3 shows adsorption isotherms of a molecular sieve for nitrogen and neon. Adsorption isotherms of other gases such as oxygen, CO, or argon are similar to the isotherms of nitrogen. The light noble gases helium and neon are adsorbed much less; hydrogen is not adsorbed at all. A comparison of Figures 11.3 and 6.7 shows that the adsorption roughly follows a Langmuir adsorption isotherm (Section 6.2.4). For simplification, adsorption isotherms can be described by using the dashed curves in Figure 11.3. These simplified adsorption isotherms provide a rough approximation considerably streamlining the following calculations. However, calculated pressure values for $p > 0.1 \,\mathrm{Pa}$ (10⁻³ mbar) are generally slightly lower than the pressures producible in reality. For p > 0.1 Pa (10⁻³ mbar), the simplified adsorption isotherms often do not describe the course of adsorption correctly because even the slightest contamination disturbs in this pressure range. For high-purity adsorbents, overall, the simplified isotherms seem to characterize adsorption processes correctly even for low pressures.

The ascending branch of the simplified adsorption isotherms (compare Section 6.2.4), plotted in Figure 11.3 as dashed lines, can be specified by

$$\tilde{\mu}_{\rm p} = A^* p \tag{11.5}$$

and the descending branch by

$$\tilde{\mu}_{\rm n} = \tilde{\mu}_{\rm n,mono}.\tag{11.6}$$

Again, μ_n denotes the adsorbed mass-related pV-amount at standard temperature $T_n = 273.15$ K, Eq. (11.4).

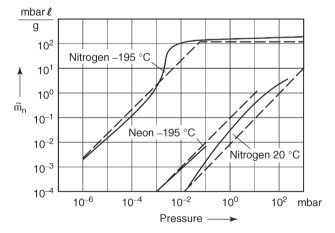


Figure 11.3 Adsorption isotherms of molecular sieve 5A for nitrogen and neon [6,7]. $\tilde{\mu}_n =$ $(pV)_n/m$ adsorbed mass-related pV-amount at standard temperature $T_n = 273.15$ K.

It is for nitrogen $A_{78~\rm K}^* \approx 2 \times 10^6~\ell~{\rm kg^{-1}}$, and for neon $A_{78~\rm K}^* \approx 100~\ell~{\rm kg^{-1}}$. By and 20 °C measurements measurements for nitrogen $A_{293 \text{ K}}^* \approx 10 \,\ell \text{ kg}^{-1}$.

For a mass m of adsorbent in the adsorption pump, volume V of recipient and adsorption pump, initial pressure p_1 of the gas (nitrogen), adsorbent and gas temperature before cooling T_1 , after cooling T_2 , as well as $V_0 = VT_n/T_1$ (standard temperature $T_n = 273.15 \text{ K}$), it follows for the ultimate pressure p_2 (without derivation) that

$$p_2 \approx p_1 \frac{\frac{V_0}{m} + A_{T_1}^*}{A_{T_2}^*}. (11.7)$$

Equation (11.7) is valid only as long as the adsorption isotherm is represented by Eq. (11.5). For nitrogen and an adsorbent temperature of T = 78 K, this is true if $p_2 < 7 \,\mathrm{Pa}$ (7 × 10⁻² mbar). Near saturation (monolayer coverage), Eq. (11.6) applies and instead of Eq. (11.7) we will use

$$p_2 \approx p_1 \frac{\frac{V_0}{m} + A_{T_1}^* - \frac{\mu_{\text{n,mono}}}{p_1}}{\frac{V_0}{m}}.$$
 (11.8)

If the adsorption pump is used *n* times without any intermediate heating, that is, without desorbing already adsorbed gas, the previously adsorbed amount of gas must be considered in calculations of succeeding pumping periods, and thus,

$$(p_2)_n \approx p_1 \frac{n_{\overline{m}}^{V_0} + A_{T_1}^*}{A_{T_2}^*}. (11.9)$$

Example 11.2

In many practical cases, 50 g of adsorbent is used per liter recipient volume, that is, when neglecting the volume of the pump, $V/m = 20 \ell \text{ kg}^{-1}$ $(V/m)(T_n/T_1) = 18.6 \ell \text{ kg}^1$ at 20 °C. For a starting pressure of $p_1 = 100 \text{ kPa}$ and nitrogen, according to Eq. (11.7), and using the values from Figure 11.3,

$$p_2 = 100 \text{ kPa} \frac{18.6 \,\ell \text{ kg}^{-1} + 10 \,\ell \text{ kg}^{-1}}{18.6 \,\ell \text{ kg}^{-1} + 2 \times 10^6 \,\ell \text{ kg}^{-1}} = 1.43 \,\text{Pa}.$$

This result is below 7 Pa, which means that using Eq. (11.7) has retrospectively turned out to be permitted.

11.2.3.2 Ultimate Pressure with Two or More Adsorption Pumps

For lower ultimate pressures, the mass of the absorbent can be increased as indicated by Eqs. (11.7)-(11.9), for example, by using two adsorption pumps (Figure 11.4).

A reasonable procedure of pumping-down to produce low ultimate pressure is as follows: both valves V1 and V2 are open. Initially, cooling is only applied to adsorption pump P1. As soon as the pressure ceases to drop, valve V1 is shut

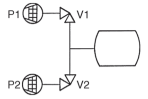


Figure 11.4 Evacuation using two adsorption pumps.

and absorption pump P2 is cooled. In this procedure, pump 2 is initially evacuated by pump 1, and thus, very low ultimate pressures develop. Analogous to Eq. (11.7), ultimate pressure

$$p_2 \approx p_1 \frac{\frac{V_0}{m_1 + m_2} + A_{T1}^*}{\frac{m_1}{m_1 + m_2} A_{T2}^*}.$$
 (11.10)

Example 11.3

To simplify, we will consider two adsorption pumps of the same type with $V/m_1=V/m_2=20\,\ell\,{\rm kg}^{-1}$ and $p_1=100\,{\rm kPa}$ at 20 °C. After cooling P1, ultimate pressure

$$p_2 = 100 \text{ kPa} \frac{10 \ell \text{ kg}^{-1} + \frac{1}{2} \times 18.6 \ell \text{ kg}^{-1}}{\frac{1}{2} \times 2 \times 10^6 \ell \text{ kg}^{-1}} = 1.93 \text{ Pa}$$

After adsorption, pump 1 is disconnected, and adsorption pump 2 is cooled, the resulting ultimate pressure, according to Eq. (11.10),

$$p_3 = 1.93 \, \text{Pa} \, \frac{10 \, \ell \, \text{kg}^{-1} + 18.6 \, \ell \, \text{kg}^{-1}}{2 \, \times \, 10^6 \, \ell \, \text{kg}^{-1}} = 2.8 \, \times \, 10^{-5} \, \text{Pa}.$$

Unfortunately, however, it is impossible to evacuate a recipient filled with atmospheric air this far using adsorption pumps. The main reason is neon and helium contents in atmospheric air, with partial pressures in the atmosphere of approximately 1.8 Pa and 0.52 Pa (Table A.6), respectively. Additionally, even minute contamination of the adsorbent has an effect at very low pressure.

Another workable method is as follows: both pumps P1 and P2 are cooled simultaneously while valves V1 and V2 are shut. Then, V1 is opened. When the pressure has almost reached the ultimate pressure of pump 1, valve V1 is closed and V2 opened. This procedure has the advantage that the airflow directed from the recipient to P1 carries along the neon and helium, and delivers it to pump P1. It is beneficial to shut V1 and open V2 at a pressure slightly higher than ultimate pressure in order to avoid any back diffusion of noble gases into the recipient.

11.2.4 **Operating Suggestions**

Besides using two adsorption pumps in order to produce very low pressures, it is advisable to start by using a dry fore pump to evacuate the recipient and the adsorption pump (or pumps) as far as possible. For example, if the total pressure is lowered to 100 Pa, neon partial pressure in the recipient drops to approximately 2×10^{-3} Pa. In practice, using two adsorption pumps with a pre-evacuated recipient yields ultimate pressures in the range of 10⁻³ Pa. By heating the adsorbent to approximately 450 °C during pre-evacuation for improved purity, pressures of 10^{-7} Pa are obtained with a two-stage adsorption pump [8].

A different approach to reduce noble-gas partial pressure in the recipient is to rinse or flood it with a well-adsorbable gas (e.g., nitrogen) prior to first evacuation. Again, thoroughly cleaned adsorbent is crucial for obtaining very low pressures [8].

The pumping speed of an adsorption pump depends strongly on the placing and type of the adsorbent [9]. Additional important factors are pressure and previous history of the pump, indicating that no constant value can be given for pumping speed. Figure 11.5 shows a typical pump-down curve for an adsorption pump.

Adsorbed gas has to be removed from the adsorbent, when the pump starts to saturate. This degassing is achieved by warming up the adsorbent [10]. With Zeolites, a 15- to 30-min warm-up period to room temperature is normally sufficient. However, at room temperature, adsorbed water is not removed (Table 11.2). Zeolites take up water very well. It reaches the surface of the adsorbent during evacuation of moist recipients or when recipients are re-

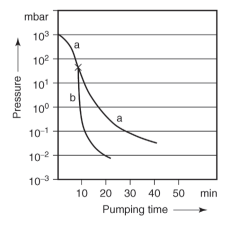


Figure 11.5 Pressure p versus pumping time for adsorption pumps arranged according to Figure 11.4. Recipient volume V=8 ℓ . Each adsorption pump is filled with m=400 g Linde molecular sieve. Curve a: one pump, curve b for switching from pump 1 to pump 2 after 8 min.

Table 11.2 Temperatures of desorption maxima during warming up of molecular sieve Linde 5A (aluminum–calcium–silicate) and 13X (aluminum–natrium–silicate).

Molecular sieve 5A 13X

Molecular sieve Adsorbed gas	5A Desorption maximum at	13X	
O_2	108 K	108 K	
Ar	110 K	103 K	
N_2	145 K	137 K	
CH_4	151 K	137 K	
CO	187 K	-	
CO_2	249 K	255–285 K	
H ₂ O	410–430 K	400–425 K	

From [10].

evacuated repeatedly (desorption gas flows from the walls of the recipient contain mostly water). Zeolite has to be heated to $300\,^{\circ}\text{C}$ for some time for water to be removed. For very low pressures, the best method is to pump down the developing vapors using a simple sliding vane rotary pump or a dry rough pump at a pressure of approximately $100\,\text{Pa}$ to $2\,\text{kPa}$.

11.3 Getter

11.3.1

Mode of Operation and Getter Types

The action of getter materials relies on gas accumulation due to

- adsorption, that is, accumulation of gas molecules at the surface;
- absorption, that is, solvation of gas molecules in the solid;
- · chemical binding.

In contrast to an adsorption pump using mainly physisorption, a getter pump for the most part depends on chemisorption (for definition see Chapter 6). Other than physisorption, chemisorption is irreversible at the temperatures that pumps are exposed to. At times, it is difficult to determine whether adsorption or absorption is predominant. Therefore, the processes are generally referred to as sorption (compare Chapter 6).

In many cases, the getter effect relies on adsorption as well as absorption. Here, impinging molecules and atoms initially bind to the surface of the getter material due to adsorption. However, they do not remain in place in a stationary

state but diffuse into the solid where the particles are trapped in grain boundaries, interstitial loci, and lattice defects.

In chemical binding, the adsorbed gas reacts with the surface atoms of the getter material (chemical adsorption) or with certain or all types of atoms in the getter material. The collection of gas in getter material is complicated, diverse, and to some extent not clarified.

Investigations are difficult due to the following conditions: any gas molecule or atom collected by the getter material initially adsorbs at the surface of the getter material. Adsorption strongly depends on individual surface characteristics. High specific surface areas of porous substances usually bind more gas through adsorption than smooth surfaces. Oxide layers, previously adsorbed layers, as well as lattice defects have differently strong effects on adsorption. Oxide layers can impede diffusion. The adsorption properties of a solid surface depend on its previous history. Adsorption characteristics are different for each gas species. Even for equal combinations of gas and getter material, adsorption characteristics can vary considerably depending on the condition of the surface. The chemically inactive noble gases are subject to the lowest collection rates. They are bound only due to physical adsorption.

Gas collection due to absorption and chemical binding is not only influenced by adsorption but also by temperature. High temperatures promote the ability to collect gas. However, in many cases, gas that becomes dissolved in a solid but not firmly bound at low temperature is released when the temperature rises beyond certain limits.

Generally, the temperature at which maximum gas collection occurs is characteristic to each gas/solid system. Thus, at a certain temperature, the situation may arise where a getter material shows its maximum gas receptivity for one gas species while it acts as a gas source for another. Combined getters where the getter material is assembled from several constituents often help to reduce this effect. A combined getter is used at a temperature where gas receptivity reaches a maximum for any gas species involved.

For getter efficiency, reversibly dissolved gas should (and often must) be released from the getter material prior to first use. Getter materials that have accumulated gas by physisorption and solvation for longer periods and at higher pressures can act as gas sources at lower pressures and thus limit the ultimate pressure of a vacuum system.

Due to different working principles, bulk getters and evaporation getters are differentiated. Bulk getters are made of solid, and in part, regenerative getter material, referred to as nonevaporable getter, or shortly, NEG (Section 11.3.2). As the name suggests, they are made of material that cannot be evaporated.

Evaporation getters (Section 11.3.3), in contrast, use getter material that is evaporated under vacuum and deposited as a thin and thus highly gas-receptive but not regenerative coating.

11.3.2 **NEG Pumps**

Fundamentals of Bulk Getters/NEG

Requirements on Bulk Getters/NEG

Getter materials in bulk getters bind gas by chemical reaction. Thus, they must be chemically reactive toward residual gases typically occurring in vacua (CO, CO₂, N₂, O₂, H₂O, H₂, etc.). Additionally, however, they should provide easy handling in contact with atmospheric air when being mounted. Metals, in particular, fulfill the former condition. The second property is ascribed to evaporable getter materials because the reactive surface of the getter is produced under vacuum by evaporation. For example, the common barium getters are used to generate elemental barium through a reaction of a mixture of a BaAl4 alloy and nickel at 800-1250 °C under high vacuum. The barium evaporates from the getter container and condenses as a reactive layer on the opposing surface (see Section 11.3.3.1).

Bulk getters or NEG are used primarily in applications where either evaporation of metal under vacuum is undesired or a surface for depositing a metal film is unavailable.

Bulk getters meet the second requirement by providing a thin protective, for example, passivating layer of oxides and nitrides. It forms spontaneously when the metal surface reacts with air, and protects the underlying metal from further reactions. Under vacuum and higher temperature, this protective layer dissolves due to diffusion into the bulk getter. Therefore, the materials used must also provide appropriate diffusion conditions for those gases to be bound.

Activating Bulk Getters/NEG

Getters bind gases at the surface. Thus, large surfaces are desired. After being placed in a vacuum, such surfaces, as any other surface, require decontamination from physically bound gases. This is done by baking.

The oxide/nitride layer on the passivating surface has to be dissolved in order to allow reactions between the getter material and gases. This step involves further heating under vacuum. In contrast to physically bound gases, the chemical bonds between NEG and oxygen as well as nitrogen atoms are too strong to be separated by heating. Therefore, even at temperatures around 1000 K, equilibrium pressures of, for example, oxygen and nitrogen, above their corresponding compounds are in the range of only 10⁻¹⁵ Pa [11].

However, high temperature increases the diffusion rates of oxide and nitride ions in the getter. Following the concentration gradient, the ions migrate into the bulk getter and the surface returns to its metal state, and thus, regains the ability to bind gases. The temperature required for the diffusion process to take place within a given time depends on the type of

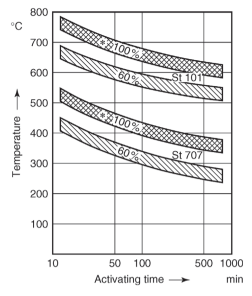


Figure 11.6 Activation diagram for St 101 and St 707 Zr alloys (see text). *Standard activating conditions.

getter material. The maximum time frame is usually predefined by the application. It spans from only a few seconds (lamp industry) to days (accelerator applications). In certain cases, partial activation of the getter surface is sufficient (Figure 11.6).

Binding Different Gas Species with Bulk Getters/NEG

Gas species, generally appearing in different proportions in vacuum applications, can be differentiated with respect to sorption by bulk getters.

a) Chemically active gases that react with the getter surface.

This group includes CO, CO₂, O₂, and N₂. These gases bind to the surface in several steps where they dissociate. This process takes place at room temperature and lower temperatures. The produced ions (carbides, oxides, and nitrides) diffuse from the surface and into the bulk of the getter material only at high temperature. Bonding is irreversible. The same applies to H₂O₂ which dissociates at the surface and is bound as O and H.

b) Chemically active gases that dissolve physically within the complete getter volume.

Hydrogen and its isotopes (deuterium and tritium) react in this way. Hydrogen binds to the activated metal surface of the getter and diffuses throughout the getter volume at or even below room temperature. The amount of hydrogen in the volume getter is in a temperature-dependent equilibrium with the outside partial pressure of hydrogen. Sievert's law

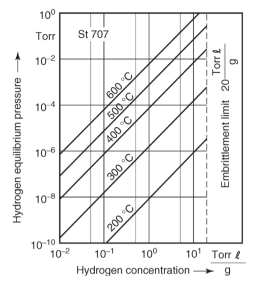


Figure 11.7 Behavior of Zr alloy (example: St 707) toward hydrogen at selected temperatures according to Sievert's law.

describes this equilibrium (Figure 11.7):

$$\log p_{\rm H} = A + 2\log q_{\rm H} - \frac{B}{T},\tag{11.11}$$

where $p_{\rm H}$ is the hydrogen partial pressure; A,B are constants; and $q_{\rm H}$ is the amount of hydrogen in the getter.

If pressure is given in Torr, amount in Torr $\ell/(g$ getter), and temperature in Kelvin, for, for example, the alloy St 707, A = 4.8 and B = 6100. Metal hydrides form only at high concentrations of hydrogen in the getter (approximately 25 hPa ℓ g⁻¹), and cause embrittlement of the getter material.

c) Chemically passive gases that do not react with the getter material or react only under high-temperature conditions.

Hydrocarbons are relatively inactive toward NEG materials. This is particularly true for methane, CH₄. They react at higher temperatures if the chemical bonds between carbon and hydrogen crack near hot surfaces. Binding then occurs and produces H2 or carbides. Long-chain hydrocarbons also physisorb to the getter surface. Getters do not sorb noble gases because they are chemically inactive. This behavior is utilized to purify noble gases with getters.

Composition of Bulk Getters/NEG

Following the list of required properties mentioned above, metals are suitable, particularly group 4 elements of the periodic table (titanium, zirconium,

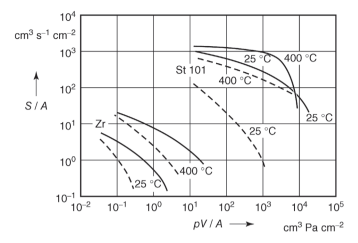


Figure 11.8 Surface-area-related pumping speeds S/A for St 101 zirconium alloy and unalloyed Zr at selected getter temperatures for CO (dashed line) and H₂ (solid line) versus previously sorbed surface-area-related gas amount pV/A. Courtesy of SAES Getters S.p.A.

hafnium), and also, tantalum, niobium, and thorium. In practice, apart from titanium, typical materials are specially developed alloys of zirconium.

Alloy compositions include Zr 84%, Al 16% (brand name St 101) and Zr 70%, V 24.6%, Fe 5.4% (brand name St 707). Compared to pure metals they have the advantage of showing higher sorption ability and lower activating temperatures θ_a (Figures 11.6 and 11.8): for tantalum, $\theta_a = 2000$ °C; for zirconium, $\theta_a = 1000$ °C; for St 101, $\theta_a = 750$ °C; and for St 707, $\theta_{xtrma} = 450$ °C.

Bulk Getter/NEG Stock

A pulverized getter alloy serves as raw material for getter production (the term getter refers to the ready-made product including getter material as well as holding devices). The particle size is in the range of 50-150 µm. Manufacturers provide getter products as getter strips, getter pellets, getter rings, and sintered bulk getters (known as porous getters).

- Getter strips (Figure 11.9).
- Here, the getter material is rolled on a carrier tape from one or both sides using high force. The carrier is made of a nickel-plated iron sheet or constantan (nonmagnetic).
- Getter pellets.
- If the getter material is not too brittle, it can be compressed to pellets.
- Getter rings.
- Getter material is pressed into metal rings with U-shaped cross section. The shape of the ring improves activating when heated with high-frequency excitation.
- Sintered bulk getters (porous getters).

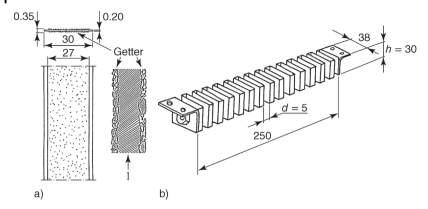


Figure 11.9 Strip getter: (a) standard strip getter 30D, dimensions given in millimeters; (b) strip getter folded to meander shape with a fixing device (getter module). Courtesy of *SAES Getters S.p.A*.

Mixing getter alloys with suitable metal powders yields sinterable mixtures that are sintered under vacuum to highly porous (up to 50% by volume), self-supporting sintered bodies. For mounting, either these getters are sintered onto metal foil, or carriers and heating elements are sintered into the material. Commercial sintered getters are made, for example, from zirconium (St 171), from zirconium and St 707 (St 172), as well as titanium and vanadium (St 185).

11.3.2.2 Design of NEG Pumps

Ready to be installed combinations of getters and resistance heaters for activation are referred to as NEG pumps. Generally, the heating element is mounted in the center of a CF flange equipped with lead throughs. Replaceable getter cartridges are fixed to the flange. The getter cartridges are built up either of folded getter strips or sintered getter disks (Figure 11.10). In both cases, the cartridges are designed to provide optimal access and contact between the getter surface and the gases to be bound. Getter strips, for example, are arranged in the cartridge as bellow-type folded rings similar to automobile air filters. The getter cartridge is a cylinder with a central hole for a heating element. The setup in NEG cartridges that contain sintered getter disks is similar. Sintered getter material does not require carriers. Therefore, they usually carry more getter material in a smaller volume.

Special electrical power supplies (NEG pump controllers) are used for activation and are wire-connected to the pump via socket connectors. Larger getter pumps are equipped with temperature probes that allow automatic control of the getter material's temperature. One manufacturer's new series (MK5, Figure 11.10, and Table 11.3) features a flange that is bakeable to 400 °C after the socket connector is removed.

NEG-alloys may also directly be applied to the inner vacuum chamber wall by coating [12]. By this method, which was developed and patented by CERN, the

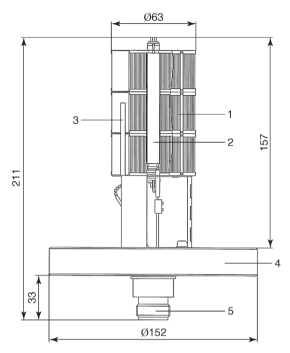


Figure 11.10 Side view with partial section of a bulk getter pump (GP200 MK5 type). Courtesy of SAES Getters S.p.A. 1 Getter cartridge, 2 heating element, 3 thermocouple, 4 base flange (CF), 5 electrical connectors for heater and thermocouple.

vacuum wall is turned from an outgassing source into a pump. It is helpful in particular where for geometrical reasons conductances to local pumps are small, for example, for insertion devices in high energy accelerators. The coating also reduces the outgassing rate and the secondary electron yield when the walls are exposed to photon (X-ray), electron, and ion bombardement. The getter is activated by a bake-out at 180 °C. The NEG alloy coating of the walls with a depth of about 1 µm is carried out by magnetron sputtering [12].

11.3.2.3 Pumping Speed and Getter Capacity

As described in Section 11.3.2.1, a bulk getter initially must undergo activation in vacuum and at high temperature before it shows any pumping action.

Table 11.3 Nominal pumping speed S_n for bulk getter pumps (SAES getters) of the design shown in Figure 11.10; mounted nude in a vacuum system. Getter material St 707.

Size	CF35	CF63	CF100	CF150
S_n for H_2 (ℓ s ⁻¹)	200	600	900	1900
$S_{\rm n}$ for CO $(\mathcal{E}{\rm s}^{-1})$	100	300	350	650

Pumping speed and getter capacity of volume getter pumps are usually measured at room temperature, a pressure of the gas to be bound of several 10⁻⁴ Pa (10⁻⁶ mbar), and with free placement of the pump in a vacuum chamber (according to ASTMF798-82). Characteristics are exemplified using two gases with different binding behaviors: hydrogen and carbon monoxide. Pumping speed is measured in liters per second, getter capacity is usually determined as Torrl or mbarl (pV-value at 23 °C, see Section 3.1.2). For hydrogen that diffuses into the bulk of the getter material, pumping speed is nearly independent of the amount of gas already bound in the getter. For carbon monoxide and other gases that bind to the surface of the getter, pumping speed drops as reaction products gradually occupy more and more surface area. Thus, a getter can be characterized by giving the pumping speed with respect to the amount of previously sorbed gas. The amount of gas accumulated at the time when pumping speed has dropped to 5% of its initial value is termed getter capacity (Figure 11.8).

Increasing temperature has an effect on all phases of gas binding. It promotes dissociation of gas molecules, and thus, a slight increase in pumping speed is generally observed at higher temperature. On getter capacity, however, temperature has a much greater effect. This is because diffusion rates of surface-bound gas atoms into the bulk of the getter material increase considerably at higher temperature. Whereas initial activation of an St 707 alloy requires 400 °C-450 °C, approximately 280 °C are sufficient to sustain diffusion when the getter operates under vacuum. This keeps the getter surface active continuously, and thus, getter capacity rises considerably for gases that bind only to the surface due to chemical reactions (Figure 11.8). Sorption continues for longer periods of time until saturation with gas particles spreads into deeper zones.

Getter capacity for hydrogen, however, does not increase with temperature because its diffusion rates in the getter material are already high at low temperature. Rather contrary, according to Sievert's law, Eq. (11.11), a rise in temperature in fact reduces the hydrogen content in the bulk of the getter material, which depends on the hydrogen partial pressure outside the getter.

As the gas content gradually approaches getter capacity, the pumping speed of the getter drops accordingly. Depending on gas species, several reasons for this can be identified:

- 1) The surface of the getter becomes covered with reaction products of getter material and absorbed gas, that is, the surface passivates. Additional chemically active gas cannot react with the surface. Access of hydrogen that could dissolve in the getter is aggravated as well.
- 2) The surface is clean but the concentration of hydrogen in the getter has grown too far (close to equilibrium concentration).

For both cases, reactivating is carried out by increasing the temperature:

1) In reactivation (e.g., at 450 °C for St 707, 750 °C for St 101), chemically bound gases diffuse into the bulk of the getter material. Up to 20 reactivation

- cycles are possible until a noteworthy increase in concentration of bound gas in the bulk of the getter material starts to reduce pumping efficiency.
- 2) Very high temperatures alter the equilibrium of hydrogen bound in the getter and surrounding hydrogen in the gas volume. Hydrogen partial pressure in the vacuum increases considerably and the hydrogen can be pumped down using mechanical pumps. After cooling, hydrogen can be absorbed anew. Theoretically, this procedure can be repeated indefinitely. However, small amounts of other gas species usually exist, which react at the surface.
- 3) The third approach is termed continuous reactivation, that is, after activation, the getter pump continues to operate permanently at higher temperature. This operating mode is suggested only for high gas loads (not H₂). Operating temperatures are well above activating temperatures (e.g., 280 °C for St 707, 400 °C for St 101).

11.3.2.4 Applications of NEG Pumps

NEG materials are used in many applications that require sealed vacua with long service life. These include any type of electron tubes [13], lamps, and stainless steel thermos flasks. Additionally, such materials are used for producing gas purifiers that purify process gases for semiconductor fabrication down to the ppt range [14]. Genuine NEG pumps are mainly used in UHV applications because of their high pumping efficiency for hydrogen that limits the ultimate pressure in such applications. Here, they operate together with other pump types, for example, turbomolecular pumps. In addition, ready-for-use combinations of ion getter pumps and NEG pumps are commercially available.

Applications cover surface analytics, very large physical experiments such as particle accelerators, experimental UHV setups, and so on. Furthermore, bulk getters have the ability of purifying noble gases. Such applications include industrial sputter systems (in situ purification) and equipment in geochronology laboratories.

Power consumption of getter pumps is low (after initial activation, further energy supply is often unnecessary). Thus, they are frequently used in mobile analysis equipment (GCM/MS) and space experiments. Bulk getter pumps can also be used to adjust a particular hydrogen partial pressure in a vacuum by setting getter temperature.

11.3.2.5 Safety and Operating Recommendations

Air Inrush During Activation

During activation or reactivation of getter material, sudden air inrush must be prevented. At temperatures above 450 °C (for St 101) and 200 °C (for St 707), the getter material would react completely with the atmospheric oxygen. This combustion progresses gradually; the material does not explode. Incomplete reaction is observed at lower temperature. However, formation of a thicker passivation layer may be observed. In such a case, reactivating requires higher than standard temperatures.

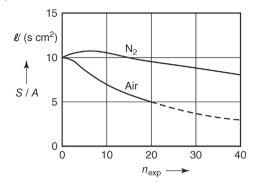


Figure 11.11 Influence of the number of exposures n_{exp} to air and nitrogen with subsequent reactivating on surface-related pumping speed S/A for hydrogen. Getter material St 101 [11].

Ventilation

It is safe to vent a volume getter pump with reactive gases (air) at temperatures below 50 °C. Afterwards, the getter simply requires reactivation. In each ventilation, however, the getter sorbs gas irreversibly, and thus, its capacity is lower after reactivating. Ten reactivating cycles do not yet severely reduce sorption characteristics of a getter. Use of nitrogen for ventilating is beneficial because pumping speed then remains high even after several reactivating cycles (after approximately 30 cycles of N_2 ventilation, S still reaches 80% of $S_{\rm initial}$; when using air for ventilation, it drops to 40% of $S_{\rm initial}$, Figure 11.11).

A further option would be ventilating with argon as protective gas for the getter to remain active. However, this procedure is rarely utilized in practice.

Overheating of Getter Cartridges

Getter cartridges using getter strips that are made of constantan should not be exposed to temperatures above 750 °C because copper and manganese could evaporate. Additionally, eutectic compositions of zirconium and copper developing above 850 °C could lead to melting of the getter cartridge.

Embrittlement of Getter Material

Beyond certain hydrogen concentrations in the getter, hydrides form in the bulk getter material. This changes the crystalline structure causing embrittlement of the material. However, such high hydrogen concentrations develop only if additional hydrogen sources are present in the system. Getter pump operation including frequent reactivation is safe for hydrogen concentrations up to $2700~{\rm Pa}~\ell~{\rm g}^{-1}~(20~{\rm Torr}~\ell~{\rm g}^{-1})$ getter material. If the concentration in the getter rises beyond this limit, the material embrittles, fractures, and in extreme cases, disintegrates into fine powder.

Evaporation/Sublimation Pumps

11.3.3.1 Evaporation Materials

Usually, evaporation getters are made from the metals barium, manganese, aluminum, thorium, or titanium. Evaporation getters just as bulk getters are initially degassed using an auxiliary pump, preferably under high vacuum. For this, the getter is heated as far as possible without noticeable evaporation occurring. Analog to subsequent heating for evaporation, heat here is produced either by a high-frequency sleeve coil arranged at the outside of the recipient or by a current-carrying suitable refractory base plate (e.g., tungsten) that holds the getter material. Degassing is finished when the pressure, after rising considerably during heat up, drops close to the initial pressure.

After degassing, the getter material is evaporated using thermal energy. Sufficient evaporation rates are obtained at a vapor pressure of approximately 1 Pa. Evaporated getter material deposits at the walls of the recipient. The produced fresh surface takes up large amounts of gas. Figure 11.12 exemplifies the procedure with a barium getter, the classic evaporation getter, in a television CRT. The Ba-Al alloy getter material BaAl₄, designed to produce approximately 200-300 mg of pure barium, is powder-mixed with nickel and pressed into a nickel-coated steel ring. This is mounted at a suitable point in the tube. After evacuating and baking out, the ring heats up to 800 °C due to the action of an outside high-frequency coil. At this temperature, a slightly exothermal reaction occurs in the powder (1.0-1.5 g), the BaAl₄ alloy dissociates, and barium is released and evaporates. The barium vapor condenses at the cold inner surface of the tube walls and produces a thin reflective coating that reaches a temperature of approximately 60 °C during tube operation.

The relatively large amount of barium yields a getter layer with a geometrical surface area of several to many 100 cm². This keeps the residual gas pressure in the tube below 10⁻⁵ Pa throughout its service life: ultrahigh vacuum in your living room!

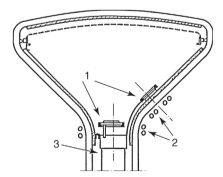


Figure 11.12 Evaporation getter in a television tube [14]. 1 Evaporation getter, arranged either in the neck of the tube or laterally at the so-called antenna position, 2 RF coil, 3 electron gun.

11.3.3.2 Pumping Speed

Volumetric flow of a gas against a container wall of area A, that is, against a getter surface as well, under high vacuum is given by $q_V = A\overline{c}/4$, as in Eq. (3.50). Gas molecules hitting a getter surface stick (adsorb) to the surface with a sticking probability s (Section 6.2.1), and reflected with a probability (1 - s). Thus, the pumping speed of a getter with surface A

$$S = sA\frac{\overline{c}}{4}. (11.12)$$

For air at 20 °C (Example 3.5), $j_V = 11 \ell (\text{s cm}^2)^{-1}$, and for hydrogen at 20 °C, $j_V = 43 \ell (\text{s cm}^2)^{-1}$. Sticking probabilities s on titanium are summarized in Table 11.4. These values represent only approximate values because they depend heavily on test conditions (compare also Table A.11). At low getter temperature, s is lower than at higher temperature. s drops as surface coverage θ increases. Figure 11.13 shows results of measurements on pumping speed S for hydrogen of a titanium evaporation getter at room temperature with surface area $A = 800 \,\mathrm{cm}^2$ and four different constant pressures p in the recipient [15]. Hydrogen volumetric flow onto the getter is constant for all four cases, $q_V = 35\,000\,\ell\,\mathrm{s}^{-1}$; pV flows $q_{pV} = pq_V$ are given for the individual sections of the image.

At first inspection, the individual images in Figure 11.13 show that the pumping speed of the getter at the beginning of gas intake (t = 10 s) is $S_0 = 700 \,\ell\text{ s}^{-1}$ and independent of p. According to Eq. (11.12), the sticking probability of the getter surface at the start of pumping is $s_0 = 0.02$. Additionally, Figure 11.13 reveals that the higher the pressure in the recipient, that is, the higher the incoming gas flow, the quicker pumping speed drops. This means that s drops as surface coverage increases. In the case of monolayer coverage, the amount of hydrogen adsorbed to the surface is $(pV)_{mono} = b_{mono} A = 3 \operatorname{Pa} \ell$, according to Eq. (11.3). If the sticking probability was constant and equal to s_0 , the getter surface would be covered within monolayer time $t_{\text{mono}} = (pV)_{\text{mono}}/(s_0q_{nV})$

Table 11.4 Sticking probabilities s for selected gas species on titanium [16] and for selected
values of surface coverage θ .

Surface coverage: Gas species	θ =	$\theta = 0.5$		$\theta = 1$		$\theta = 2$		$\theta = 5$	
	s (77)	s (300)	s (77)	s (300)	s (77)	s (300)	s (77)	s (300)	
$\overline{H_2}$	0.06	0.04	0.03	0.02			0.02	0.015	
H ₂ [17]	0.2	0.01							
N_2	1	0.3	0.005	0.001					
O_2	1	1	1	1	0.8	0.9	0.05	0.001	
CO	1	1	0.8	0.2	0.1				

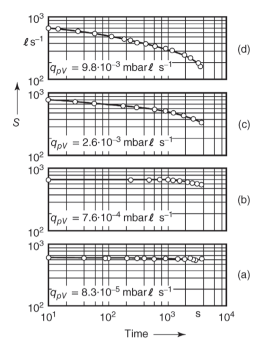


Figure 11.13 Time dependence of the pumpselected hydrogen pressures: (a) $p = 2.44 \times 10^{-7} \text{ Pa}$; (b) $p = 2.25 \times 10^{-6} \text{ Pa}$; ing speed S (getter speed) of a titanium (c) $p = 7.59 \times 10^{-6} \text{ Pa}$; (d) evaporation getter (getter surface area $A = 800 \text{ cm}^2$, getter temperature 20 °C) for $p = 2.87 \times 10^{-5} \text{ Pa } [15].$

(Section 6.2.6). For the cases featured in images (a)–(d) of Figure 11.13, this time would correspond to $t_a \approx 18\,000\,\text{s}$, $t_b \approx 1900\,\text{s}$, $t_c \approx 560\,\text{s}$, and $t_d \approx 150\,\text{s}$, respectively. For (a), test time is only approximately one fifth of monolayer time; s is constant and equal to s_0 . In case (d), pumping speed drops from $S_0 =$ 700 ℓ s⁻¹ to $S = 500 \ell$ s⁻¹ within $t_d \approx 150$ s, and thus, sticking probability drops from 0.02 to 0.014. Therefore, monolayer time can be expected to be slightly higher, approximately $t_d \approx 180$ s. Testing time in this case is about 20-fold of monolayer time. However, s is reduced to $s_0/4$.

It can be estimated that the getter has taken up five to ten monolayers within this time. Building up many layers is inconceivable because such a massive coating would represent solid hydrogen at room temperature. Thus, the getter process must be thought of differently: molecules hit the surface where they become adsorbed. Subsequently, they diffuse into the bulk material as H atoms. This diffusion proceeds rapidly so that surface coverage θ remains very low $\theta \ll 1$ for considerable time, up until a gradient of particle number density $\partial n/\partial x$ (x perpendicular to the surface) has developed that hampers diffusion current density (Section 3.3.4). The surface cannot take up more hydrogen than can diffuse into the bulk material.

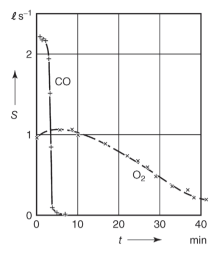


Figure 11.14 Time-dependent pumping speeds S for oxygen and carbon monoxide of a barium evaporation getter. Surface area $A=8~{\rm cm}^2$, getter temperature $\theta=50~{\rm ^{\circ}C}$, vessel pressure $p = 6.7 \times 10^{-2} \text{ Pa } [18].$

Pumping speed

$$S = s(\theta)q_V = AD\frac{\partial n}{\partial x}.$$
 (11.13)

As $\partial n/\partial x$ gradually decreases with time, pumping speed S drops proportionally. If the getter is saturated with particles, S approaches zero.

Figure 11.14 shows the time dependence of pumping speed S for a bariumevaporation getter with surface area $A = 8 \text{ cm}^2$ for oxygen and carbon monoxide at a getter temperature of $\theta = 50$ °C and a pressure of $p = 6.7 \times 10^{-2}$ Pa in the container. Pumping speed S_{O_2} for oxygen remains constant for approximately 10 min, but then drops gradually, and reaches 15 of the initial value after 40 min. S_{CO} , however, is practically zero after 5 min.

This very dissimilar behavior is because oxygen reacts with the complete barium layer to form barium oxide, whereas carbon monoxide produces a thin protective (passivating) layer that prevents further gas take-up. At temperatures θ < 40 °C, oxygen too forms a protective layer.

For temperatures $\theta > 80$ °C, the formation of a protective layer by carbon monoxide can also be prevented. The conditions in bulk getters are similar to evaporation getters, as shown in Figure 11.15.

11.3.3.3 Getter Capacity

The total amount of gas or vapor that a getter can take up is referred to as getter capacity. Often, it is so high that the number of bound atoms or molecules corresponds roughly to the number of atoms (molecules) in the getter material. For noble gases, which are absorbed at the surface only, getter capacity is lower by

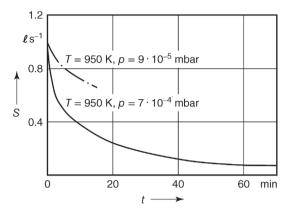


Figure 11.15 Time-dependent pumping speeds S for oxygen of a thorium bulk getter at two selected inlet pressures. Surface area $A = 4 \text{ cm}^2$, getter temperature T = 950 K [18].

many orders of magnitude. Table 11.5 gives an overview of specific getter capacities C_{getter} for high-vacuum deposited evaporation getters.

Example 11.4

According to Table 11.5, the specific getter capacity of titanium for hydrogen $C_{\text{getter}} = 27 \, \text{Pa} \, \ell \, \text{mg}^{-1}$. The molar mass of titanium $M_{\text{molar}} = 48 \, \text{g mol}^{-1}$. Thus, amass m = 1 mg of titanium contains $N = 1.25 \times 10^{19}$ titanium atoms. Because of pV = NkT, Eq. (3.19), the pV-amount $(pV) = 27 \text{ Pa } \ell$ contains $N = 6.7 \times 10^{18}$ particles, that is, 6.7×10^{18} hydrogen molecules dissociated into twice as many hydrogen atoms, 1.34×10^{19} , are distributed among 1.25×10^{19} titanium atoms. Thus, each titanium atom in the getter correlates to approximately one interstitial hydrogen atom (proton).

11.3.3.4 Design of Evaporation Getters

Evaporation getter pumps (sublimation getter pumps) use the adsorption of chemically active gases at the surface of a thin getter film for pumping. Pumping speed is particularly high when the getter film is freshly deposited and cooled to low temperatures. Since pumping speed drops as surface coverage θ increases, the getter layer is renewed after showing approximately half of monolayer coverage ($\theta = 0.5$) with gas particles.

According to Eq. (11.12), the area-related pumping speed

$$S_{\mathcal{A}} = \frac{S}{A} = S\frac{\overline{c}}{4},\tag{11.14}$$

Getter material	Pumped gas species	$C_{ m getter}$ (Pa ℓ mg $^{-1}$)	Reference
Titanium	H_2	27	[15]
	O_2	4.4	[19]
	N_2	0.85	[19]
Aluminum	O_2	1	[20]
Magnesium	O_2	2.7	[20]
Barium	H_2	11.5	[20]
	O_2	2	[20]
	N_2	1.26	[20]
	CO_2	0.69	[20]
Composition metal (cerium/ lanthanum)	H_2	6.13	[20]
	O_2	2.8	[20]
	N_2	0.43	[20]
	CO_2	0.29	[20]

Table 11.5 Mass-related or specific getter capacity $C_{\rm getter}$ for selected materials and various gas species at $\theta = 20^{\circ}$ C.

and if neglecting the surface-coverage dependence of sticking probability s, that is, $s = s_0(\theta = 0)$,

$$S_{\mathcal{A}} = s_0 \frac{\overline{c}}{4}.\tag{11.15}$$

This is independent of particle density and thus of the pressure also, but only as long as continuous or discontinuous deposition provides $\theta \ll 1$. If the maximum deposition rate determined by performance and arrangement of evaporation sources is obtained, higher pressures lead to decreased pumping speeds. Figure 11.16 describes this behavior of S(p). For pressures at which the mean free path \bar{l} of evaporated getter atoms in the pumped-down gas is in the range of the average distance between evaporator and getter screen L or lower $(\bar{l} < L)$, the deposition rate on the getter screen drops. This additionally decreases pumping speed and ultimately leads to ceasing pumping action.

The design of evaporation getters is remarkably simple (Figure 11.17). They include an evaporator for the getter material and a screen to capture the getter film, the sorption surface for pumped-down gas. Nearly all getters use titanium as getter material tebib19, which is produced cheaply in large scale. The simplest design features a getter screen, which is part of the vacuum container wall. Particularly in UHV equipment that operates at pressures $p < 10^{-6}$ Pa, the surface collision rate is low and coverage of the getter surface increases slowly so that only small amounts of titanium need to be deposited. Heat transfer to the getter

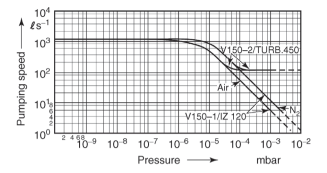


Figure 11.16 Pumping speed of a titanium evaporation pump versus pressure during pump-down of air and nitrogen. Evaporation pump V 150-1 (Figure 11.17) combined with

ion getter pump IZ 120 (see Section 11.4). Evaporation pump V 150-2 combined with turbomolecular pump TURBOVAC 450.

screen (heat of condensation) is low and heat removal through the container wall and to the surrounding is sufficient to maintain room temperature at the getter. If necessary (for higher pressures), the container can be cooled by ventilating or via an outside coiled cooling pipe. Higher pumping speeds, which require correspondingly higher deposition rates and thus produce higher thermal condensation output, are better produced by using a getter screen that is designed as a device separate of the container wall (Figure 11.18).

Due to its very high thermal conductivity, copper is an appropriate material. If it is cooled with liquid nitrogen, it has to be arranged separately from the vacuum container to keep cooling-medium consumption low.

The limited Ti reservoir of a Ti evaporator calls for economical use. This includes:

a) Equipment for maintaining constant evaporator temperature, for example, for stabilizing electron emissions of the hot Ti wire analogous to emission stabilizing in hot cathode ion gauges (see Section 13.7.3) [21].

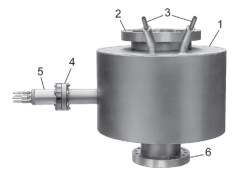


Figure 11.17 Titanium evaporation pump V 150-2 (see also Figure 11.18). 1 Pump housing, 2 connection flange for vacuum vessel, 3 coolant inlet and outlet (water or LN₂), 4 connecting flange for evaporator unit 5, 6 connecting flange for auxiliary pump.

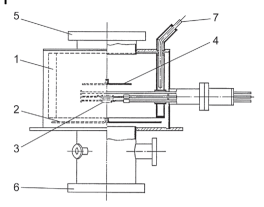


Figure 11.18 Section of the titanium evaporation pump shown in Figure 11.17 (diagram). 1 Cylindrical, cooled getter screen, 2 screening plate cooled by thermal conductance, 3 evaporator coil, 4 screening plate, 5 connector for vacuum vessel, 6 pump connection, 7 LN₂ feed line.

b) Electrical controls that operate the titanium evaporator(s) intermittently according to a predefined program. Here, the duration of operating interruptions is controlled automatically depending on the pressure in the container.

The pumping speed of the pump is determined not only by the getter surface area but also by the geometry of the screen. Strictly speaking, Eq. (11.15) applies only if the gas particles have access to every surface element of the getter surface from the entire half-volume, and show an undisturbed Maxwell distribution.

However, inside the pump's hollow body, this distribution is definitely disturbed. In practice, the hollow body shape is predetermined by the fact that the titanium emanating toward all directions from the evaporation source shall be utilized as much as possible.

Example 11.5

A getter screen consists of a cylinder of length l=0.3 m, diameter d=0.25 m, and its circular base plate. Thus, it has the surface area $A = \pi Id + \pi d^2/4 = 0.285 \text{ m}^2$. The circulartop surface $A' = 0.049 \,\mathrm{m}^2$ represents the inlet port of the getter pump. The getter screen is cooled with liquid nitrogen and coated completely with deposited titanium providing a constantly fresh surface. According to Table 11.4, the sticking probability for $\theta = 0.5$ and N₂ is s = 1; following Table A.10, the mean thermal velocity $\bar{c} = 470 \,\mathrm{m \, s^{-1}}$ for nitrogen at 20 °C. Equation (11.12) yields a pumping speed

$$S = s A \overline{c}/4 = 1 \times 0.285 \text{ m}^2 \times 470 \text{ m s}^{-1} \times 0.25 \approx 33500 \, \text{e}^{-1}$$

According to Eq. (4.143), the inlet's intrinsic conductance

$$C = 11.56A' \ell (s cm^2)^{-1} \approx 5700 \ell s^{-1}.$$

Thus, using Eq. (4.34), the effective pumping speed of the arrangement

$$\frac{1}{S_{eff}} = \frac{1}{S} + \frac{1}{C}; \quad S_{eff} = \frac{SC}{S+C} = \frac{33\,500\,\times\,5700}{39\,200} \; \ell' \; s^{-1} \approx 4870 \; \ell' \; s^{-1}.$$

Thus, the pumping speed is practically determined by the inlet port. Without nitrogen cooling, Table 11.4 suggests s = 0.3 and pumping speed amounts to $S = 0.3 \times 33500 \,\ell\,\text{s}^{-1} = 10050 \,\ell\,\text{s}^{-1}$, yielding an effective pumping speed $S_{\rm eff} = 3500 \, \ell \, {\rm s}^{-1}$, a reduction of only 25%.

Two methods are commonly used for evaporation and sublimation.

A resistance evaporator is particularly simple in terms of design and operation. Therefore, it conforms perfectly with the rest of the pump component's simplicity. It is made of a titanium wire, heated directly by means of an electrical current. The titanium wire is fixed at two contact points mounted to a separate flange (see Figure 11.18) providing easy replacement. One of the contacts is connected to the outside via a vacuum-tight electrical leadthrough; the other is connected to ground. An electrical current of approximately 40-50 A is required for a wire of 2 mm diameter for producing an evaporation rate per unit length of approximately 0.02 g h⁻¹ cm⁻¹. In order to reduce the spatial requirements for the evaporation chamber, the wire is coiled up to a spiral. Three or more such spirals are mounted on the flange. If the titanium reservoir of one wire is depleted, the succeeding wire is activated so that opening of the vacuum chamber is unnecessary. If titanium consumption is high, two coils can be operated simultaneously.

Temperatures required for sufficiently high evaporation rates (1200–1500 °C) are so high that negative effects on the mechanical strength of the titanium wire become noticeable. The wire softens, sags, and embrittles due to ongoing recrystallization. For increased mechanical strength, molybdenum or tungsten carrier wires spooled with titanium are common. However, the best solution prevailing so far uses a titanium wire alloyed with 15% molybdenum [22].

In narrow tubes (e.g., storage rings), titanium evaporators consist merely of a long, stretched wire (linear pump) sputtered in an argon gas discharge [23].

Electron beam evaporators are more complex but carry higher titanium reservoirs. Therefore, they allow long uninterrupted operation. They include a titanium anode (titanium block or rod) and one or more glow cathodes as electron sources. Electrons, here, are accelerated with a voltage of 2-4 kV. Special beam guidance or beam shaping is usually unnecessary. Initial heating of the titanium anode after starting the glow cathode takes about 1-3 min. At high evaporation rates, additional heating of the cathodes due to the bombardment with ionized titanium atoms must be taken into account. Thus, automatic emission-current stabilization is required. The shape of the titanium anode can call for a feeding mechanism that compensates for the growing distance between the titanium anode and the cathode due to titanium evaporation. As long as evaporation occurs from the solid phase only, both types of evaporators can usually be arranged in any desired position. However, this is not the case if liquid-phase evaporation is utilized in order to produce particularly high evaporation rates as in electron beam evaporators.

Apart from the actual getter screen, a getter pump includes several additional screen plates for preventing titanium from migrating from the evaporator, through the pump inlet, and into the recipient. These plates are designed to impair conductance of the inlet port as little as possible. Thus, they are usually placed close to the evaporator and are cooled similar to the getter screen (Figure 11.18).

A titanium evaporation pump operates in conjunction with a different type of pump that pumps down nongetterable gases such as CH₄ and noble gases. In this context, a problematic behavior of titanium evaporators arises. A titanium evaporator itself produces CH₄ [16]. Titanium sublimation pumps can be combined with any high-vacuum pump offering sufficient pumping speed for nongetterable gases (noble gases), that is, a diffusion pump, turbomolecular pump, or sputter ion pump.

Titanium evaporation pumps are often used for producing hydrocarbon-free vacua. Thus, combinations with diffusion pumps are avoided. Vacuum technology usually relies on combinations of sputter ion pumps or turbomolecular pumps and titanium evaporation pumps. The additional pump needs a noblegas pumping speed of only a few percent of the pumping speed of the titanium evaporation pumps. This is true as long as the content of gases, nongetterable for titanium, in the pumped-down gas mixture is low, which is the case, for example, for air.

A turbomolecular pump evacuates particularly quickly and, if dimensioned generously, operates up to working pressures of 0.1 Pa. Due to the high hydrogen pumping speed of the getter pump, producible ultimate pressure is no longer limited by the hydrogen compression ratio of the turbomolecular pump. Thus, combinations of titanium evaporation pumps and turbomolecular pumps produce pressures around 10^{-10} Pa.

Cooling the getter screen with liquid nitrogen has the advantage that the pumping speed for nitrogen rises twice as high compared to a system using water cooling. However, it has the disadvantage that venting of the vacuum system requires either both preheating and subsequent cooling of the getter screen or disconnecting the titanium evaporation pump with a valve, that is, an additional component.

It is also important to note that deposition of Ti on an LN2-cooled surface leads to much higher sticking probability and thus pumping speed compared to deposition onto a surface at room temperature [16,24–26].

11.4 Ion Getter Pumps

11.4.1

Working Principle

Sorption in ion getter pumps relies on (cathodic) sputtering of a getter material inside a gas discharge, and additionally, on bombardment (implantation) of ions from the gas discharge. Utilization of these effects for development and design of vacuum pumps [27] was encouraged by investigations aimed at preventing such processes (gas depletion and erroneous pressure measurement) in ionization vacuum gauges.

Gas discharge in an ion getter pump is of the *Penning* type [28] (Section 13.7.4.1). Figure 11.19 illustrates an electrode arrangement, two parallel cathode plates K₁ and K₂, and an anode cylinder A with the z-axis arranged perpendicularly to the cathode planes. A magnetic field of flux density $B \approx 0.1-0.2 \,\mathrm{T}$ is applied in the *z*-direction.

Operating voltage U between anode and cathodes is approximately 6 kV. Knauer [29] and Schuurman [30] thoroughly investigated the Penning discharge in such arrangements. Section 13.7.4.1 describes its mechanism in detail.

In detail, the pumping effects are as follows:

a) Ion implantation. The applied electrical potential (6 kV) accelerates the ions produced in the discharge to several kV, depending on their point of origin. Acceleration occurs nearly along a straight path toward the cathode because the ions, due to their large mass compared to electrons, are hardly influenced by the magnetic field. The ions penetrate the crystal structure of the cathode by approximately 10 atomic layers (ion implantation). This corresponds to a gas depletion that affects any species of gas ion including atomic and molecular ions of noble gases and other gas species. However, very large molecular

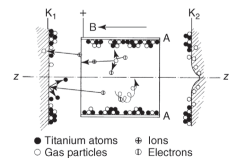


Figure 11.19 Diagram of the pumping action in a *Penning* cell (diode). K₁, K₂ cathode plates made of getter material (titanium), A anode cylinder with the z-axis, B magnetic field. The

getter film with buried gas particles is visible on A and toward the ends of K_1 and K_2 . Implanted gas particles in the center of K₂ (and K₁ as well, not drawn).

02	C ₂ H ₂	C ₂ H ₄	со	H ₂	CO ₂	N ₂
A	A	A	A	A	A	A
A	A	A	A	A	A	NA
A	A	A	A	A	NA	NA
A ^{a)}	A	A	A	NA	NA	NA
A	A	NA	NA	NA	NA	NA
A	NA	NA	NA	NA	NA	NA
NA	NA	NA	NA	NA	NA	NA
	A A A A A A A A	A A A A A A A A A A A A A A A A A A A	A A A A A A A A A A A A A A A A A A A	A A A A A A A A A A A A A A A A A A A	A A A A A A A A A A A A A A A A A A A	A A A A A A A A A A A A A A A A A A A

Table 11.6 Adsorption behavior of selected gas species on metals and metalloids.

A: adsorption, NA: no adsorption. Generally, adsorption energy drops from left to right.

From [31,32].

ions, such as hydrocarbons, do not penetrate the lattice structure. Of these ions, only the fraction disintegrating during surface impingement is pumped. The penetration depth of such fragments, however, is lower because their kinetic energy is low.

- b) Cathode sputtering. The ions hitting the cathode are implanted in part and sputter individual or larger numbers of lattice atoms. These atoms are released and deposit on surrounding surfaces where they form the getter film when the cathode is made from getter material (e.g., titanium). The mass of the sputtered material is roughly proportional to the pressure in the pump so that the pumping speed adjusts to this pressure. Pumping action, as any getter effect (Section 11.3), is strongly influenced by the gas species. Table 11.6 provides an overview of the adsorption of selected components in residual gas onto a number of metals. Depending on the field configuration caused by the electrons and the volume charge in the discharge, the ions accelerated toward the cathode can become focused to the z-coordinate. This produces a sputter crater in the center of the cathode (Figure 11.19, cathode K₂). In any case, getter action takes place mainly at the edges of the cathodes and at the anode; implantation occurs mostly in the center of the crater because the getter film here is resputtered.
- c) Neutral particle implantation. When ions, particularly noble-gas ions, impinge the surface, they can be reflected if they become neutralized in the metal. Indeed, this occurs often in an ion getter pump. Neutralized gas particles then become implanted at other spots because they still carry high kinetic energy.

The getter effect is the predominant effect in an ion getter pump. Nevertheless, the two implantation processes are very important because they represent the cause for noteworthy pumping speed of the ion getter pump for noble gases.

a) Not on Au.

For estimating the pumping speed of a Penning cell, we shall consider the following: the number of ionized gas molecules can be expected to be proportional to the pressure and to the number of electrons $Q_{\rm e}(p)$ in the volume charge cloud (see Section 13.7.4.1). The latter is slightly pressure dependent. Thus, the discharge current I amounts to

$$I = K_1 Q_e(p) p \tag{11.16}$$

with the proportionality constant K_1 . The quotient I/p is also referred to as the sensitivity of the Penning cell.

Furthermore, it can also be expected that the rate of pumped molecules q_{pV} is proportional to the discharge current:

$$q_{pV} = K_2 I = K_1 K_2 Q_e(p) p. (11.17)$$

Because pumping speed $S = q_{pV}/p$, it follows that

$$S = K_1 K_2 Q_{\rho}(p). \tag{11.18}$$

Section 13.7.4.1 also states that

$$I = Kp^m$$
 with $m = 1 - 1.4$. (11.19)

Equating Eqs. (11.16) and (11.19) yields

$$Q_{\rm e}(p) = \frac{K}{K_1} p^{m-1}$$
 with $m = 1 - 1.4$ (11.20)

so that

$$S = KK_2 p^{m-1}$$
 with $m = 1 - 1.4$. (11.21)

From this derivation, pumping speed can be expected to drop, even though slightly, with pressure p.

Equation (11.18) shows that maximizing volume charge Q_e is beneficial for obtaining high pumping speed. This circumstance has promoted many investigations focusing on the parameters that influence $Q_{\rm e}$ (Table 11.7).

Table 11.7 Setting values in commercial ion getter pumps that influence the pumping speed of a Penning cell.

Quantity	Symbol	Variation range	
Anode voltage	$U_{ m H}$	3.0-7.0 kV	
Magnetic field strength	В	$0.1-0.2\mathrm{T}$	
Cell diameter	d	1-3 cm	
Cell length	l	1-3.2 cm	
Distance between anode and cathode	а	0.6-1.0 cm	
Pressure	p^m	m = 1 - 1.4	

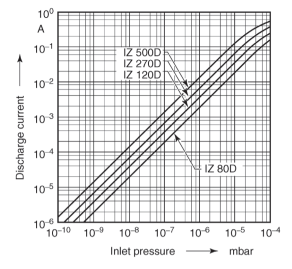


Figure 11.20 Discharge currents in ion getter pumps versus inlet pressure.

Under otherwise constant parameters, a minimum field strength B_{\min} (≈ 0.03 T) is required for sustaining the discharge. For $B > B_{\min}$, S initially rises fairly linearly up to a maximum and subsequently drops. In the region rising with B, S also increases linearly with the high-voltage $U_{\rm H}$. Investigations have shown that the distance a between the anode and cathode should not be too short so that passing of gas particles through the *Penning* cell is not hindered too much. Similar considerations apply to the length l of the cell. If l is too high, conductance for the gas is reduced thus counteracting the benefit of the higher volume charge. For constant l/d, larger values for d produce higher pumping speeds but only for pressures below 10^{-4} Pa [33].

Since the discharge current I in the Penning discharge is proportional to gas pressure within certain pressure ranges (Figure 11.20, compare also Figure 13.63, kink at approximately p=0.01 Pa $(10^{-4} \, \mathrm{mbar})$), measurement of I can be used to determine the pressure in the pump as in a Penning vacuum gauge (Section 13.7.4.1). However, it was observed often that, for hydrogen in particular [34], I (and thus S) varies considerably in spite of constant pressure, depending on the condition of the pump. Therefore, this pressure reading should be interpreted very carefully. The discharge continues to glow at pressures $p < 10^{-8}$ Pa. Therefore, baked-out ion getter pumps can be used to produce extremely low gas pressures.

Service life of an ion getter pump is determined mainly by the depletion of getter material. Depletion of the titanium cathode plates by sputtering in the electrode arrangement shown in Figure 11.21 is very inhomogeneous due to the crater formation described above. Thus, more than 90% of the total getter material is not exposed to the discharge and therefore remains unutilized. This *efficiency* can be increased by arranging the anode system (3) in Figure 11.21 so

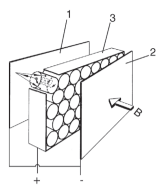


Figure 11.21 In reference to the design of a diode-type ion getter pump. 1, 2 Titanium cathode plates, 3 frame holding cylindrical anode cells. B Magnetic field produced by permanent magnets.

that it can slide parallel to the cathode plates [35]. Then, areas that have seen less or no exposure are sputtered as well. For working pressure p_w , approximate values for the service life t_s of commercial iongetter pumps are given by

$$\frac{t_{\rm s}}{\rm h} \cdot \frac{p_{\rm w}}{\rm Pa} \approx 4.$$
 (11.22)

11.4.2

Technical Design (Diode Type)

If a single *Penning* cell provides a small pumping speed $S_{P.c.}$, then n such cells can be connected in parallel to produce a pumping speed nS_{P.c.}. Reikhrudel et al. [36] described the first multicell ion getter pump of this kind. Today, anode cells are connected in parallel in a honeycomb design (Figure 11.21). The common cathode plates are arranged at a distance of a few millimeters. The complete electrode system represents a diode and lies in a vacuum-tight, nonmagnetic housing that is placed in the gap of a permanent magnet arranged outside of the vacuum. Pockets in the housing are either included in a pump housing with a flange or placed directly into the wall of the recipient. The common magnet system for all electrode systems has an annular yoke. This keeps stray field losses low and the magnetic flux density in the air gap, that is, in the electrode system, as high as possible. The electrode systems are commonly connected to an electrical current feed through in the wall of the pump housing, which is easily detached and replaced. Electrode systems in large pumps are replaceable as well. After the sputtered titanium has become depleted at the cathode, the electrode systems can be replaced by new systems.

A power supply provides high voltage, usually 3–7 kV, for the pump. A current limiter is required to protect the pump from overload as the discharge current

rises proportionally to the pressure (Figure 11.20). For this, a constant-current transformer is often utilized. Voltage and discharge current are measured by monitoring equipment built into the power supply. For electrical current, a logarithmic measuring scale is usually available with the reading calibrated in pressure units analogous to the design in a *Penning* ionization vacuum gauge.

11.4.3 Pumping Speed

As derived in Eq. (11.21), the pumping speed of sputter ion pumps is moderately pressure dependent and reaches a smooth maximum at a pressure of approximately 10^{-4} Pa (10^{-6} mbar) (Figure 11.22).

The different discharge shapes occurring in *Penning* discharges throughout the wide pressure range of many orders of magnitude chiefly determine the shape of the pumping speed curve [30]. Additionally, pumping speed, which depends on the amount of gas bound to the surface, changes during the service life of the pump. After longer periods of operation, pumping speed drops. Baking (regenerating) restores pumping speed. Obviously, saturation proceeds more slowly if the pump operates in the UHV range ($p < 10^{-5} \, \mathrm{Pa}$) only. However, when pressures are very low and approach the ultimate pressure of the pump, low saturation can already lead to relatively high desorption, that is, a drop in effective pumping speed.

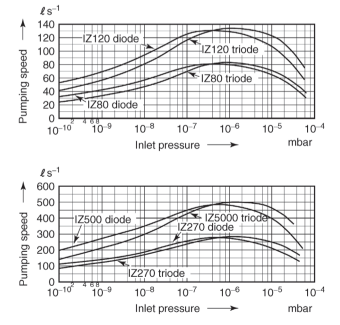


Figure 11.22 Pumping speeds for air versus inlet pressure of differently sized ion getter pumps.

Table 11.8 Pumping speeds of ion getter pumps for selected gas species with respect to the pumping speed for air (approximate values).

Gas species	Diode pump	Triode pump ^{a)}
Air	1	1
Nitrogen	1	1
Oxygen	1	1
Hydrogen	1.5–2	1.5-2
Carbon monoxide	0.9	0.9
Carbon dioxide	0.9	0.9
Water vapor	0.8	0.8
Light hydrocarbons	0.6–1	0.6-1.2
Argon	0.03	0.25
Helium	0.1	0.3

a) Section 11.4.5.

Pumping speed in an ion getter pump is gas-species dependent. Two main groups of gas species are differentiated with respect to their influence on pumping speed:

- a) Getterable gases that can be pumped by chemisorption, for example, nitrogen, oxygen, carbon oxides, light hydrocarbons, water vapor. According to Table 11.8, pumping speed for these gases is far higher than for the other species. The values in Table 11.8 are normalized, referring to the value for air or nitrogen. Differences are due to the variations of ionization probability of gas particles in the gas discharge and in sticking probability of gas particles on the getter layer. In addition, unequal sputter rates of ions at the cathode surface determine the values.
- b) Gases pumped only due to ion implantation, particularly noble gases. Due to this mechanism, the pumping speed of an ion getter pump is lower for noble gases than for other gas species. Furthermore, it drops after relatively short operating time because both processes, ion implantation and cathode sputtering, counteract in the *Penning* cell (Figure 11.19). Slight deviations in the volume charge inside one or more of the *Penning* cells can lead to a situation in which new parts of the cathode are bombarded that previously worked as getter only. Released gas particles lead to locally increased pressure again increasing the bombardment. Thus, positive feedback occurs. This causes socalled noble-gas instabilities, short pressure risings of up to two orders of ten with a periodicity of several to many minutes during early pumping phases. Later, after electrode temperature rises, this periodicity drops to fractions of a minute or several seconds. Figure 11.23 shows two such bursts. Obviously, a rise in hydrogen partial pressure follows each rise in argon partial pressure with a time lag of several seconds. During stationary pump operation at low

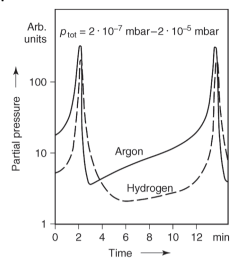


Figure 11.23 Noble-gas instabilities for pumping argon. Ordinate: partial pressure in arbitrary units, logarithmic scale. Abscissa: time.

pressure, the pumping speed for hydrogen $S_{\rm H_2}$ is approximately 1.5–2 times as high as the pumping speed for air. In contrast, $S_{\rm H_2}$ can drop considerably in the pressure range $p > 5 \times 10^{-5}$ Pa. Just as the hydrogen bursts, this effect can also be explained with heating of the electrodes under high discharge currents (heavy ion bombardment!). The diffusion coefficient and solubility of hydrogen in titanium is high (compare Section 11.3.3.3 and Example 11.4). Thus, as temperature increases, hydrogen diffuses to the surface and desorbs.

One more comment on hydrogen pumping: for H_2^+ to be taken up by getter material, it must initially dissociate into two H atoms because only about 2.5% of the hydrogen dissociates in the *Penning* discharge. For dissociation, appropriate surface loci (activation sites) have to be available that reduce the necessary activation energy of dissociation far enough for dissociation to actually take place. However, if the cathode surface becomes contaminated with TiN or Ti oxides, only few activation sites are available and dissociation of H_2^+ is suppressed. Activation sites are created by cathode sputtering. When pumping H_2 with ion getter pumps, producing a clean surface can take quite a while because H_2 itself has a low sputter rate. In these cases, a continuous rise in the pumping speed of the pump is observed for H_2 .

11.4.4 The Differential Ion Pump

The effect of noble-gas instability described in the previous section can be eliminated by a simple measure that replaces one of the titanium cathodes with a

tantalum cathode [37]. Tantalum is a very hard material with high atomic mass that reflects noble gas ions with much higher energy as neutral particles than titanium. The kinetic energy of reflected argon atoms is up to 50 times higher than for titanium [38,39], depending on the reflection angle, and reaches up to 50% of the energy of incidence. This leads to an implantation depth in the anode or opposing cathode high enough for the atoms to be trapped for the rest of the serviceable life of the ion getter pump. This type of an ion getter pump in diode design is referred to as a differential ion pump (DIP). However, the getter effect of tantalum is lower than that of titanium.

This slightly reduces pumping speed compared to a conventional diode pump. However, in addition to noble-gas stabilization, the pumping speed for noble gases is increased to approximately 25% of the pumping speed for nitrogen (compared to approximately 5% in conventional diode pumps). The success of this diode arrangement proved that neutral-particle implantation as described in Section 11.4.1 is in fact an important mechanism, at least when considering noble gases. The inventors of the pump, Tom and James [37], initially tried to explain the higher pumping speed and stability for noble gases with a difference in sputter rates for the two materials, which gave the pump its name. In this case, noble gas atoms would become trapped mainly in the tantalum cathode with the lower sputter rate because here more titanium atoms impinge from the opposing cathode compared to the rate of tantalum atoms being released. However, roughly equal sputter rates were observed soon [39] and the model of the inventors proved wrong. The name, however, prevailed.

Modifications of cathode arrangements were tested using individual pellets for each Penning cell [40], mixtures of titanium and tantalum, or designs with a 1mm-thick perforated tantalum plate on top of the titanium cathode [41]. Just recently, a commercial pump manufacturer introduced a system with a variable Ti/Ta ratio. However, it should be remembered that each individual *Penning* cell with Ti cathodes at both ends bares the continuous risk of argon instability.

11.4.5

Triode Pumps

Considerably increased pumping speeds for noble gases in ion getter pumps and better constancies are obtained by utilizing a so-called triode arrangement [42-44] as shown in Figure 11.24. Cathodes K here consist not of bulk plates but of mesh. A collector plate F on anode potential is arranged behind K. Often, the inner wall of the vacuum housing serves for this third electrode, and thus, A and F are at ground potential. The discharge in this setup is restricted to the volume of the anode cylinder and has the same shape as in a diode pump. Between K and F, the discharge is suppressed.

Ions accelerated from the discharge toward the cathode participate in mostly glancing angle collisions (large angles of incidence) with the cathode surface (compare Figure 11.24). Thus, they penetrate the cathode weakly but do cause a certain degree of sputtering of cathode material (titanium). Therefore, a getter

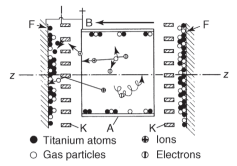


Figure 11.24 Diagram showing pumping action in a triode-assembly discharge cell. K Cathode mesh, A anode, F collector plates, Z anode axis, B magnetic field.

film deposits on F (and to a lower degree on A), and the associated gas consumption proceeds.

In this glancing reflection, the probability for ions to loose their charge is high [45] (they pick up electrons). However, their kinetic energy hardly drops; they hit F, and become implanted (important for noble-gas pumping [46]). Ions that have not lost their charge do not travel against the electric field between K and F; they return and again face the probability to impinge K and sputter. The arrangement provides a high sputter rate at K and thus high getter effect but a low sputter rate at F (neutral particles can sputter as well but not as intensively due to the loss of kinetic energy experienced during the collisions at K). Therefore, the probability of implanted particles to be released is much lower.

The combination of both effects leads to a pumping speed for noble gases, with respect to nitrogen, of 20-30% in a triode pump compared to 1-10% in a diode pump (compare Table 11.8).

Analogous to diode pumps, several triode cells are combined to compact electrode systems that form pockets, as in diode pumps, with the permanent magnets arranged in between (Figure 11.25). Ion getter pumps following the diode and triode principles look alike from the outside. Figure 11.26 shows a standard ion getter pump with a nominal pumping speed of $500 \, \ell \, \rm s^{-1}$, always referring to air or nitrogen (Figure 11.22). The pumping speed of this pump in diode design for argon and helium is 5 and $50 \, \ell \, \rm s^{-1}$, respectively, and for the triode design 125 and $150 \, \ell \, \rm s^{-1}$, respectively. In the triode design, starting pressure (see Section 11.4.8) is 1 Pa and thus 10-fold lower than in diode designs.

Production of triode pumps used to be very time consuming and thus expensive. Each individual Ti cathode strip had to be hand-fixed to the cathode frame. *Varian's* [47] StarCell triode pump (Figure 11.27) solved this cost problem. Furthermore, it also eliminated an operational problem of triode pumps that often malfunctioned or overheated when pumping hydrogen or water vapor at higher pressure due to short circuits between anode and cathode citebib44 because they are thermally and electrically isolated from the pump body. The StarCell pump has a considerably larger surface area so that these problems are minimized.

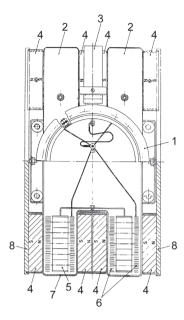


Figure 11.25 Diagram showing the design of an ion getter pump in triode configuration. Upper half of the image: top view. 1 High-vacuum flange, 2 electrode pockets, 3 power supply, 4 permanent magnets. Lower half of

image: Section 4. Permanent magnets, 5 anodes A in frame, 6 cathode mesh, 7 collector plate also part of the nonmagnetic pump case, 8 magnet yoke.



Figure 11.26 Ion getter pump IZ 500/IZ 500D (diode). Main dimensions in mm: width 408, depth 480, height (without sealing flange) 436. Baking temperatures: 350 °C with magnet, 450 °C without magnet. Eight-electrode system. Weight 135 kg.



Figure 11.27 Cathode of the StarCell triode pump by Varian. StarCell is patented and a copyrighted brand name of Varian.

11.4.6

Distributed Ion Pumps

Due to the conductance of tubes, efficiency of pumps in the long tubes of highenergy accelerators is limited. A linear pump, however, capable of pumping throughout the entire tube, pumps down the gas near its place of origin (photo desorption) [48–50]. Ion getter pumps can be used as such linear arrangements, referred to as distributed ion pumps or linear getter pumps, where the *Penning* cells are distributed along the tube. Furthermore, the magnetic field of the deflecting magnets can be utilized for operating the *Penning* cells. Figure 11.28 illustrates a cross section of the accelerator tube in the DESY electron storage ring.

11.4.7 Residual Gas Spectrum

Figure 11.29 shows a typical residual gas spectrum of an ion getter pump. Main species in the residual gas are hydrogen from the metal walls, carbon monoxide

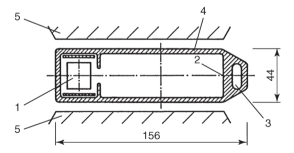


Figure 11.28 Section of a storage ring with a linear ion getter pump (IZ pump, diode) at DESY. 1 Ion getter pump (diode), 2 radiation absorber for synchrotron radiation, 3 coolant channel, 4 vacuum chamber, 5 beam-control magnet.

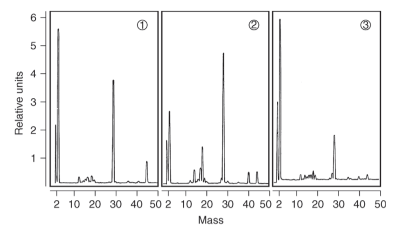


Figure 11.29 Residual gas spectrum of an ion getter pump in triode configuration. 1 Air inflow, pump unbaked, $p_{\text{tot}} = 8.6 \times 10^{-8} \, \text{Pa}$, 2 hours, $p_{\text{tot}} = 7.4 \times 10^{-8} \, \text{Pa}$. inflow of 40 kPa ℓ air and 370 Pa ℓ argon.

pump unbaked, $p_{\text{tot}} = 1 \times 10^{-7} \text{ Pa}$, 3 as 2 but after baking the pump at 300 °C for several

and carbon dioxide, as well as methane produced from hydrogen and carbon oxides in the electrical discharge.

However, Reich [51] and Lichtman [52] state that titanium also includes carbon as contaminant and thus an ion getter pump can produce methane itself. Increasing pump saturation affects the residual gas spectrum.

Here, a so-called memory effect occurs [46,53]. The spectrum contains species that were absorbed in the previous history of the pump even after attached to a completely different container in the meantime, which does not contain any of these gas species. Often, even baking does not decrease the partial pressures of such gases below detection limits. The memory effect is related to the discharge of buried gas particles during warming or sputtering of electrode material and is particularly high for noble gases. Triode pumps have a much lower tendency of showing a memory effect and noble gas instability.

11.4.8

Operation

Ion getter pumps are used to produce vacuum with low contents of hydrocarbons. Triode pumps indeed are capable of pumping down oil vapor, however, using diode or triode pumps in vacuum systems that contain oil vapors is not profitable. The reason is that the operation of diode pumps can become severely restrained by oil vapor contamination: they start more slowly and show lower pumping speeds.

Ion getter pumps comply with the requirement of very high operational reliability. These pumps form a thoroughly sealed unit together with the vacuum recipient, which prevents air from penetrating even if pump operation stops (e.g., due to power failure). High ambient temperature, radioactive radiation, and strong magnetic stray fields hardly disturb pump operation.

Ion getter pumps require only a single high-voltage wire and no additional supply lines. Via this wire, operation of the pump is monitored, started, or stopped from a remote control site (e.g., remote control in particle accelerators). Due to their vibration-free operation, ion getter pumps are applicable to vibration-sensitive measuring equipment. Compared to other types of UHV pumps (turbomolecular pumps, cryopumps), they also provide the best ratio of electrical energy consumption and pumping speed.

Possible disturbances include the magnetic stray field of the pump as well as the emission of ionized particles, neutral particles (titanium), and soft X-radiation by the gas discharge. The radiation can influence the readings on, for example, mass spectrometers and ionization vacuum gauges. Additional disadvantages are that ion getter pumps are heavy, and most prominently, are hardly applicable to pressures $> 10^{-2}$ Pa. Indeed, the pumps can cope with short periods of higher pressures, but degassing due to warming of the electrodes prevails after longer operating periods in the pressure range $> 10^{-2}$ Pa. Thus, the preferred pressure range for continuous operation is p < 1 mPa.

Ion getter pumps should not be started before the pressure drops below a certain starting pressure. For diode pumps, this threshold is $0.1 \,\mathrm{Pa}$ ($10^{-3} \,\mathrm{mbar}$), and for triode pumps, 1 Pa (10^{-2} mbar). Pumping down the vacuum system to starting pressure is carried out best by using pumps that produce an oil-free vacuum. For containers up to a volume of 300 ℓ , adsorption pumps (Section 11.2) are appropriate. Sliding vane rotary pumps with preconnected adsorption traps, however, are also suitable if the adsorption medium is regenerated by baking and care is taken that no oil vapor condenses in the connection line on the high-vacuum side. Usually, turbomolecular pumps are used for preevacuation, particularly, when large vacuum chambers are evacuated or if the conductance of pump connections is low for technical design reasons that would require the ion getter pump to operate in the pressure range of $0.5-5 \times 10^{-3}$ Pa for longer periods. In this case, the turbomolecular pump evacuates down to approximately 10⁻² Pa. After the ion getter pump is started, both pumps continue to operate simultaneously for some time.

Starting of an ion getter pump is different for diode and triode pumps. After the high voltage is connected at 1 Pa, a glow discharge is initiated spreading throughout the diode pump and even into the recipient because the complete housing is at ground potential. In the triode pump, however, the anode completely encapsulates the cathode restricting the discharge to the space between the electrodes right from the start. Therefore, from the beginning, the triode sputters titanium whereas sputtering in the diode is established fully only after the discharge has ceased in the vicinity of the electrons after the pressure drops.

Standard ion getter pumps are available for nominal pumping speeds up to $500 \,\ell\,\mathrm{s}^{-1}$. They are utilized as stand-alone pumps in relatively small vacuum systems (e.g., vacuum deposition systems, zone melting equipment). Large systems use such pumps in situations when vacuum containers, due to their geometry, require multiple small pumps instead of a single large pump (e.g., particle accelerators, storage rings).

11.5 **Orbitron Pumps**

An orbitron pump (Figure 11.30) is an ion getter pump utilizing the energy of the electrons both for ionizing molecules and for evaporating getter material. A radial electrical field is established between the cylindrical vacuum housing as cathode (ground potential) and a rod-shaped anode is placed along the center axis of the housing.

In this field, electrons from a glow cathode arranged between the cathodic cylinder and the anode follow multiple orbits along hypocycloid paths around the anode before they hit its surface. This increases the path length of the electrons dramatically, and thus, each electron collides with and ionizes gas particles several times. In the electrical field, the ions travel toward the housing where they become implanted in the getter layer.

An evaporation source is used to produce the getter layer. This source is usually heated with the ionizing electrons; in rare cases, an independent power supply is used. For the former case, the evaporator is combined with the anode of the ionizing equipment: on the rod-shaped anode, a titanium body is placed, heated, and evaporated by the impinging electrons (Figure 11.30).

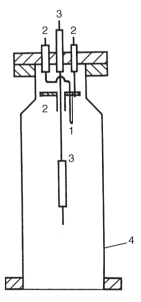


Figure 11.30 Diagram of an orbitron pump [54]. 1 Glow cathode with protective shield toward anode supply, 2 shield at cathode potential, 3 anode with titanium body, 4 housing at cathode potential.

However, the pumping effect in a system using this type of ionization is low. This becomes apparent in a relatively low pumping speed for noble gases, amounting to only a few percent of the pumping speed for nitrogen. In this sense, a combination of a simple getter pump and an additional ion getter pump or turbomolecular pump is superior. However, one important advantage is that orbitron pumps do not require a magnetic field as found in ion getter pumps, which makes them suitable for certain applications. Miniaturized orbitron pumps [55,56] have been discussed to pump micro-electrical-mechanical systems, which may be another field of application.

References

- 1 ISO 3529/2 (1981) Vacuum technology; Vocabulary; Part 2: Vacuum pumps and related terms. Trilingual edition.
- 2 DIN 28400 (1980) Part 2, Vacuum technology; terms and definitions, vacuum pumps, Beuth, Berlin.
- 3 Grubner, D.M. et al. (1968) Molekularsiebe, VEB Deutscher Verlag der Wissenschaften, Berlin.
- 4 Dobrozemsky, R. (1973) Vakuum-Technik, 22, 41-48.
- 5 Visser, J. and Scheer, J.J. (1973) Ned. Tijdschrift. Vac. Technol., 11, 17-25.
- 6 Turner, F.T. and Feinleib, M. (1961) Eighth Nat. Vac. Symp. and Second. Int. Nat. Vac. Congress, Pergamon Press, Oxford.
- 7 Stern, S.A. and Paolo, F.S. (1967) J. Vac. Sci. Technol., 4, 347-355.
- 8 Windsor, E.E. (1963) Sorption pumping using zeolithes, in Physik und technik von Sorptions- und desorptionsvorgängen bei niederen drücken (ed. G. Kienel). Rudolf A. Lang, Esch, Germany, pp. 279-283.
- 9 Creek, D.M. et al. (1968) J. Sci. Instr. (J. Phys. E), 2, 582-584.
- 10 Miller, H.C. (1973) J. Vac. Sci. Technol., **10**, 859–861.
- 11 Ferrario, B. (1998) Getters and Getter pumps, in Foundations of Vacuum Science and Technology (ed. J.M. Lafferty), John Wiley & Sons, New York, Chapter 5, pp. 261-315.
- 12 Benvenuti, C. (2001) Vacuum, 60, 57-65.
- 13 Boffito, C. and Sartorio, E. et al. (1986) An update of non-evaporable getters in electron tubes. Vakuum-Technik, 35, 212 - 217.
- 14 Juhr, W. (1987) Einsatz von Gettern zur Aufrechterhaltung von Vakua, in

- Vakuumtechnik in der industriellen Praxis (ed. Kerske et al.) Expert, Ehningen, pp. 145-169.
- 15 Kienel, G. and Lorenz, A. (1960) Vakuum-Technik, 9, 1-6.
- 16 Gupta, A.K. and Leck, J.H. (1975) An evaluation of the titanium sublimation pump. Vacuum, 25, 362-372.
- 17 Eisworth, L. et al. (1965) Vacuum, 15, 337 - 345.
- 18 Wagener, S. (1954) Z. Angew. Physik, 6, 433 - 442.
- 19 Lückert, J. (1961) Vakuum-Technik, 10,
- 20 Ehrke, L.F. and Slack, C.M. (1957) J. Appl. Phys., 28, 1027-1030.
- 21 Strubin, P. (1980) J. Vac. Sci. Technol., 17, 1216-1220.
- 22 McCracken, A.M. and Pashley, N.A. (1966) J. Vac. Sci. Technol., 3, 96 - 98
- 23 Blechschmidt, D. and Unterlechner, W. (1979) Vakuum-Technik, 28, 130 - 135.
- 24 Sweetman, D.R. (1961) The achievement of very high pumping speeds in the UHV region. Nucl. Instr. Meth., 13, 317.
- 25 Grigorov, G.I. and Tzatzov, K.K. (1983) Theory of getter pump evaluation. Sticking coefficients of common gases on continuously deposited getter films. Vacuum, 33, 139.
- 26 Grigorov, G.I. (1984) Apparent and real values of common gas sticking coefficients on titanium films and application to getter pump devices with periodic active films renovation. Vacuum, 34, 513.
- 27 Hall, L.D. (1958) Rev. Sci. Instr., 29, 367 - 370.

- 28 Penning, F.M. (1937) Physica IV, 2, 71-75, Philips Techn. Rundschau 2 (1937), 201 - 208.
- 29 Knauer, W. (1961) J. Appl. Phys., 33, 2093-2099.
- 30 Schuurman, W. (1966) Rijnhuizen-Report 66-28, University of Utrecht.
- 31 Bond, G.C. (1962) Catalysis by Metals, Academic Press, New York, p. 69.
- 32 Winters, H.F., Horne, D.E., and Donaldson, E.E. (1964) Absorption of gases by electron impact. J. Chem. Phys., 41, 2766.
- 33 Rutherford, S.L. (1964) Sputter ion pump for low pressure operation, Proc. 10th Nat. AVS Symposium 1963, Macmillan Company, New York p. 185.
- 34 Singleton, J.H. (1969) Hydrogen pumping speed of sputter ion pumps. J. Vac. Sci. Technol., 6, 316.
- 35 Henning, H. (1980) Proc. 8. Intern. Vac. Congress, Cannes Suppl. Rev., "Le Vide", No. 201 143-146.
- 36 Reikhrudel, E.M., Smirnitskaya, G.V., and Burnisenica, G.V. (1956) Ion pump with cold electrodes and its characteristics, Radiotekh. Electron, 2, 253.
- 37 Tom, T. and James, B.D. (1969) Inert gas ion pumping using differential sputter yield cathodes. J. Vac. Sci. Technol., 6, 304.
- 38 Jepsen, R.L. (1968) Proc. 4th Int. Vac. Congr. London 1968, Vol. I 317.
- 39 Welch, K.M. (1991) Capture Pumping Technology, Pergamon Press, Oxford, pp.
- 40 Baechler, W. and Henning, H. (1968) Proc. 4th Int. Vac. Congr. London 1968, Vol. I
- 41 Komiya, S. and Yagi, N. (1969) J. Vac. Sci. Technol., 6, 54.
- 42 Brubaker, W.M. (1959) Transact. of the 6th Nat. Vac. Symp, Pergamon Press, Oxford, pp. 302-306.
- 43 Vaumoran, J.A. and Biasio, M.P. (1970) Vacuum, 20, 109-111.
- 44 Singleton, J.H. (1971) Hydrogen pumping by sputter-ion pumps and getter pumps. J. Vac. Sci. Technol., 8, 275-282.
- 45 Oechsner, H. (1966) Z. Naturf., 21a, 859.
- 46 Bance, U.R. and Craig, R.D. (1966) Vacuum, 16, 647-652.
- 47 Pierini, M. and Dolieno, L. (1983) A new sputter-ion pump element. J. Vac. Sci. Technol. A, 1, 140.

- 48 Cummings, U. et al. (1971) Vacuum system for the stanford storage ring. SPEAR. J. Vac. Sci. Technol., 8, 348.
- 49 Pingel, H. and Schulz, L. (1980) Proc. 8. Intern. Vac. Congress, Cannes, Suppl. Rev., "Le Vide", Nr. 201, 147-150.
- 50 Blechschmidt, D. et al. (1980) Proc. 8. Intern. Vac. Congress, Cannes Suppl. Rev., "Le Vide", Nr. 201, 159-163.
- 51 Reich, G. (1963) Investigation of titanium sheets for sputter ion pumps. Supplemento Al Nuovo Cimento, 1, 487.
- 52 Lichtman, D. (1964) Hydrocarbon formation in ion pumps. J. Vac. Sci. Technol., 1, 23.
- 53 Henning, H. (1975) Vakuum-Technik, 24, 37 - 43.
- 54 Douglas, R.A. et al. (1965) Rev. Sci. Instr., **36**, 1–6.
- 55 European Patent EP000001403903A2, Inventor: Koops, H. (2012) Orbitronpumpe, published 2004-03-31.
- 56 Grzebyk, T. and Górecka-Drzazga, A. (2011) Field-emission electron source for vacuum micropump. Vacuum, 86, 39 - 43.

Further Reading

- Audi, M. and de Simon, M. (1987) Ion pumps. Vacuum, 37, 629-636.
- Audi, M. et al. (1987) A new ultrahigh vacuum combination pump. J. Vac. Sci. Technol., A, 5, 2587-2590.
- Benvenuti, C. and Francia, F. (1988) Roomtemperature pumping characteristics of a Zr-Al non-evaporable getter for individual gases. J. Vac. Sci. Technol. A, 64, 2528-2534.
- Boffito, C. et al. (1987) Gettering in cryogenic applications. J. Vac. Sci. Technol., A, 54, 3442-3445.
- della Porta, P. (1972) J. Vac. Sci. Technol., 9, 532 - 538.
- Ferrario, B. et al. (1985) A new generation of porous non-evaporable getters. Vacuum, **35**, 13.
- Jepsen, R.L. (1969) The physics of sputter ion pumps, Proc. of the Fourth Intern. Vac. Congress, Manchester, IOP Conference Series No. 5, London, 317-324.

12

Cryotechnology and Cryopumps

Dr. Christian Day

Karlsruhe Institute of Technology, Institute for Technical Physics, Hermann-von-Helmholtz-Platz 1, 76344, Eggenstein-Leopoldshafen, Germany

This chapter covers the physical and technical fundamentals of refrigeration and how they can be utilized for vacuum technology by means of cryopumps.

12.1 Introduction

"Cryo" is derived from the Greek word kryos meaning "cold." Thus, cryotechnology refers to refrigeration technology. Although "cold" is generally understood as temperatures below ambient temperature, it is agreed that only refrigeration in the temperature range $T < 120 \, \mathrm{K}$ is referred to as cryotechnology or cryogenics. Cryotechnology and vacuum technology are closely interrelated:

- Vacuum is indispensable for thermal insulation in the application of low temperatures in the cryo range: the lower the working temperature, the more important the quality of thermal insulation. In a refrigerated, vacuum insulated system, heat enters from outside through thermal radiation, from thermal conduction by the gas in the vacuum chamber, and from thermal conduction over solid connectors between parts with different temperatures. If the mean free path of the gas molecules is greater than the vessel dimensions, as is the case in the fine vacuum range, the thermal conductivity of the gas decreases linearly with the pressure (cf. Section 3.3.3) and is negligible with pressures in the high-vacuum range compared with the other heat transport processes. These vacuum pressures should be reached as a minimum for an effective insulation. Known design principles from high-vacuum and ultrahigh-vacuum technology are applied in order to ensure the necessary low leak rates in cryogenics.
- On the other hand, low temperatures as such can serve for the production of vacuum. At sufficiently low temperatures, all gases transform into a liquid or solid phase with an accordingly low vapor pressure or sublimation pressure. The particles leave the gas phase and the gas pressure decreases accordingly.

This thermal principle is the most direct method of producing vacuum without any moving parts.

Both fields of activity have always promoted each other's development. Only the application of vacuum to the thermal insulation made the liquefaction of hydrogen and helium possible. Later, the condensation trap cooled with liquid nitrogen became an indispensable accessory for the diffusion pump. Space research required large amounts of liquefied gases for space simulation and as rocket fuel. This led to substantial progress in cryotechnology and broadened the potential range of applications for cryopumps toward other vacuum technological purposes. After a short time, the expansion of the low-temperature research and the introduction of new communication systems that required even lower temperatures triggered the development of reliable refrigeration equipment of varying power levels. Today, they are available to vacuum engineers as refrigeration units for cryopumps.

The technical application of superconductivity for generators, wiring, power generation, energy storage, or in the high-field magnetic technology and nuclear fusion again represents a great challenge to vacuum technology and cryotechnology. The long service life demanded for the superconductivity systems and the high operational safety can only be realized when one also succeeds in keeping the leak rates from the cold seals extremely low and in utilizing the existing cold surfaces in a suitable manner as cryopumps. Thus, a deep understanding of the processes of the condensation and adsorption of gases at low temperatures is essential. The success with high-temperature superconductors is strongly dependent upon the availability of suitable refrigeration [1].

12.2 Methods of Refrigeration

Different refrigeration processes can be applied for the production of low temperatures in the range $T < 120 \,\mathrm{K}$. Two temperature levels are commonly worked with, namely, 5-20 K for the low-temperature stage and 50-80 K as a temperature of the thermal shield that is required to keep the incidence of thermal radiation on the low-temperature surface sufficiently low.

Today, commercial cryopumps are almost exclusively equipped with their own gas refrigeration machine that is operated with an internal working medium in the cycle, primarily helium. In addition, the pump must only be additionally equipped with a compressor and be supplied with electric power. As a result, the cryopumps are similarly simple to use as other high-vacuum pumps.

In addition, there are direct processes in which the surface to be cooled in special cryostats is brought into immediate contact with the boiling refrigerants. This technology is complicated to use, particularly when it concerns liquid helium. Moreover, a regular and controlled replenishment is necessary to compensate for vaporization losses.

Gas Refrigeration Processes

To cool without liquid cryogens, the cyclic processes are used in which the working gas utilized experiences a periodic change of state. In the cyclic process, work is used to produce cold or, more precisely, to withdraw a specific amount of heat from a cooled object in order that this can be held at a lower temperature T below the ambient temperature $T_{\rm m}$.

The ratio of the spent work W and the heat Q that is taken from the system at a temperature T indicates the efficiency η of the machine. The theoretical maximum efficiency can be calculated directly from both temperature levels for the case of the completely reversible ideal process that is known as the Carnot cycle:

Carnot efficiency
$$\eta_{\text{Carnot}} = \frac{W}{|Q_{\text{Carnot}}|} = \frac{T_{\text{u}} - T}{T}$$
. (12.1)

In the Carnot process, the largest possible portion of the supplied work is converted into cold. The formula illustrates the second law of thermodynamics, which states that the working effort strongly increases with the decrease in the desired cryogenic temperature T. Thus, at least 2.6 W mechanical drive power is required for 1 W of refrigeration power at 80 K, while 1 W at 4 K already requires at least 72 W of mechanical drive power.

Typical efficiencies of real gas refrigerating machines lie at 80 K in the order of magnitude of no more than 10–20% of the Carnot efficiency [2].

The coefficient of performance ε for a refrigerating machine refers to the ratio of generated cold to the mechanical work W utilized for it; this corresponds to the reciprocal value of the machine efficiency η :

$$\varepsilon = \frac{1}{\eta} = \frac{|Q|}{W}.\tag{12.2}$$

In contrast to the traditional refrigeration, for example, in the refrigerator where the compression refrigeration cycle with evaporating refrigerant is used for the heat absorption, gas refrigeration equipment is preferentially used for cryo-vacuum applications and is therefore characterized by the fact that the refrigerant continually exists above the two-phase range.

The cyclic processes used here continuously run between the two temperature levels, ambient temperature and the low temperature, and at two pressure levels. A temperature decrease is achieved when reducing the pressure. The work to be expended serves to compress the expanded gas again. In the simplest case, a throttling valve is used for the expansion of the working gas in which the pressure reduction is not used to extract any technical work and is idealized to happen without thermal exchange with the surroundings. Following the law of conservation of energy (the first law of thermodynamics), the energy content of the system therefore remains constant. Because for real gases the energy content is a function of pressure and temperature, the expansion is connected with a

temperature change (Joule-Thomson effect), whereas the temperatures remain constant with ideal gases.

Nevertheless, at a given pressure, there is a dedicated temperature for every gas at which the Joule-Thomson effect disappears. It is called inversion temperature T_{inv} . Above the inversion temperature, the Joule-Thomson effect is negative; that is, a warming of the gas occurs with the expansion. Below the inversion temperature, the Joule-Thomson effect is positive and the gas cools down. The inversion temperature is dependent upon pressure (inversion curve $p(T_{inv})$). To reach the desired temperature drop by Joule-Thomson expansion, the quantity pair (p, T) upstream the release valve must lie below the inversion curve of the gas concerned (Figure 12.1).

Nitrogen liquefaction from room temperature is possible through Joule-Thomson expansion alone because the inversion curve (see Figure 12.1c) includes the temperature of 293 K. Hydrogen and helium gases, on the other hand, must be precooled, for example, through heat transfer at lower temperatures. Helium, the traditional working medium in cryogenic gas refrigeration equipment with a closed cycle, must be precooled to values below the temperature of liquid nitrogen. This occurs within the cyclic process through heat transfer between the colder low-pressure gas and the inflowing warmer highpressure gas. Either counter flow heat exchangers (recuperators) or regenerators

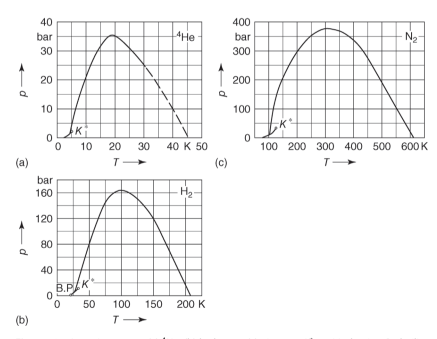


Figure 12.1 Inversion curves: (a) 4 He; (b) hydrogen; (c) nitrogen. K^{*} = critical point; S = boiling point. With value pairs (p, T) above the inversion curve, warming appears with Joule-Thomson expansion.

are used for this. While in recuperators the high-pressure gas and the lowpressure gas stand in continuous thermal contact with each other via a thermally well-conducting wall, in regenerators the two gas streams flow successively into the same system. The regenerator alternately absorbs heat and delivers it again. Regenerators also work very efficiently with small driving temperature differentials [3]. Through the temporal separation of the release and compression processes, one can manage with only one regenerator and needs no potentially vulnerable changeover devices.

Because no liquefaction of the working gas occurs, only a sliding temperature level can be generated on the refrigeration side of the gas process. If more demanding requirements for the temperature stability exist, of course one can utilize the cryostat technology with boiling refrigerant (Section 12.2.2).

With regard to design details of refrigerating machines, there exists extensive literature [4,5]. In the following, only three of the most frequently used processes for cooling cryopumps are described. All three concern regenerative gas compression refrigeration processes. This refrigeration equipment is also referred to as refrigerators.

12.2.1.1 Stirling Process

The Stirling process is an important process for the production of low temperatures that had been improved in the middle of the nineteenth century by the Philips Company and was used for air liquefaction for the first time [6].

As Figure 12.2 illustrates, a cylinder is employed containing a compression piston and a displacement piston that are driven by a common crankshaft. The working gas is in a closed system. It is compressed between the two pistons, then releases the heat of compression $|Q_{u}|$ while it flows through the cooler, and afterward is cooled in the regenerator to approximately the temperature at which the

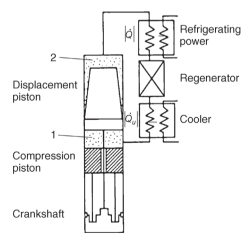


Figure 12.2 Stirling process. Dotted: working gas. (1) Compression volume; (2) expansion volume.

refrigeration power is to be released. Subsequently, the further cooling occurs through expansion between the displacement piston and the sealed cylinder end (volume 2). While flowing back, the expanded cold gas absorbs the heat Q from the object to be cooled in an open contact heat exchanger (cold head). While the gas continues to heat up still cooling the regenerator, it ultimately enters compression volume 1 between the two pistons in a warm state. The high-pressure side of the machine usually works between 10 and 30 bar. A given amount of working gas is only pushed between the compression volume 1 and the expansion volume 2 back and forth and has also in the expanded state a relatively high pressure. This means that the displacement compressor performs no compression work, but must merely overcome the flow resistance of the regenerator. Without the displacement piston that separates the gas movement spatially and timewise from the pressure and temperature oscillation, no heat transport would be possible. There is only a low pressure difference at the displacement piston, but a large temperature differential.

With this process, temperatures up to approximately 12 K can be reached in two-stage machines. The special advantages of the Philips version of a Stirling refrigerator are the high efficiency (at 80 K up to 25% of the Carnot efficiency [1]) and the simple mechanical construction. However, the necessary dry pistons require a high maintenance effort.

12.2.1.2 Gifford-McMahon Process

Also with this refrigeration process, as illustrated in Figure 12.3 [7], the working gas is pushed back and forth through a regenerator by means of a displacement piston. Here, however, the compression occurs outside, at ambient temperature, in a compressor separated from the refrigeration system. With this process, no work is done by the system, but an amount of heat $|Q_{ij}|$ is led to the surrounding,

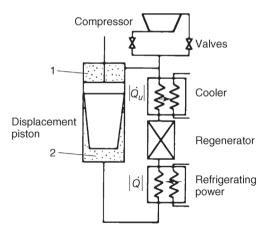


Figure 12.3 Gifford-McMahon process. Dotted: working gas. (1) Compression volume; (2) expansion volume.

corresponding to the amount of established cold |Q|. In the same manner, as the gas on the cold side 2 of the displacement piston is cooled due to expansion, heat originates on the ambient temperature side 1 of the displacement piston, namely, from the compression of the gas. The gas released from the system has a higher temperature than the gas supplied to the system.

The Gifford-McMahon process has the advantage that except for the slowmoving displacement piston, no other moving parts are necessary at low temperatures. In contrast to the Stirling process, the system has two control valves, but these are located in a place at ambient temperature. The control valves reduce the efficiency of the refrigerator, but this concept has the great advantage that oil-lubricated compressors can be used. In the same way as for the Stirling process, the refrigeration power in the Gifford-McMahon process also decreases with the decrease in temperature.

The Gifford-McMahon principle allows a spatial separation of the compressor and the cold side. The connection can be accomplished with pipes. Therefore, constructively, the displacement compressor piston, the cold heat exchanger, and the regenerator are united as a so-called cold head in an apparatus as reproduced in Figure 12.4. A two-stage Gifford-McMahon cooler is shown. It consists of a water-cooled compressor unit and a cold head with two temperature stages that is connected with the compressor unit with two flexible pressure pipes. In the process, the helium gas is pushed through the cold head at high pressure

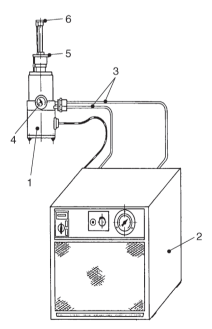


Figure 12.4 Two-stage Gifford-McMahon refrigerator: (1) cold head; (2) compressor unit; (3) flexible pressure pipes; (4) feedthrough for pressure and temperature measurement; (5) the first stage; (6) the second stage (low-temperature stage).

by the compressor. In this process step, the compressor outlet valve is opened and the intake valve is closed. When the displacement piston has reached the top dead end, the valve switch is reversed. As a result, the gas can flow back through the displacement piston from the compression volume via the expansion volume to the compressor; in the cold head, the gas is expanded successively in two stages and is thereby cooled down.

This setup is the traditional arrangement for a so-called refrigerator cryopump where the first stage is connected with a thermal shield and the second with the real pumping surface. This arrangement also allows for any installation position of the cold head. The typical refrigeration power in a two-stage machine amounts to 3-20 W at 20 K and up to 100 W at 80 K.

A disadvantage with Stirling machines is the transmission of the compressor vibration to the cold head and, with this, to the coupled structures. In the described Gifford-McMahon refrigerator, the spatial separation of compressor and cold head leads to a remedy since only the remaining vibrations of the displacement piston are transferred to the cold head. The refrigerator cryopumps addressed in Section 12.4.3.2 function according to the Gifford-McMahon process; the helium refrigerant flows in a closed cycle so that no refrigerant losses occur.

12.2.1.3 Pulse Tube Process

In the modern pulse tube cooler whose development in this form had only begun in the middle of the 1980s [8], the concept of substituting all moving parts in the refrigeration area is consistently applied. This process has conquered the market since about the year 2000, particularly as a miniaturized refrigeration machine. The displacement piston is entirely substituted with a phase-delayed movement of a gas column in a so-called pulse tube (Figure 12.5). This results in the pulse tube cooler being low in maintenance and low in vibration and, with this, makes it very attractive especially for demanding and sensitive applications, as, for example, in medical diagnostics, infrared spectroscopy (night vision device), and superconductivity.

The correct phase response between gas movement and pressure oscillation that is vital to the heat transfer is reached through the fact that the working fluid is collected cyclically in a reservoir volume that is connected via a nozzle. The nozzle takes over the role of the displacement piston and provides for a damping of the pressure oscillation during the gas movement and, with this, for a spatial separation of the temperature effects.

As in Stirling and Gifford-McMahon machines, the helium working gas is in a closed system. In the pulse tube cooler, however, the relevant heat transfer is based on a thermoacoustic oscillation of a small gas portion (reservoir) [9] that "undulates" back and forth. After the compression step, gas flows into the pulse tube and on the way delivers heat to the surroundings through a heat exchanger at the warm end of the pulse pipe. Afterward, the cooling occurs through expansion and the cold gas is pushed by the gas flow from the reservoir to the cold end of the pulse tube where it absorbs the heat of the object to be cooled at the cold

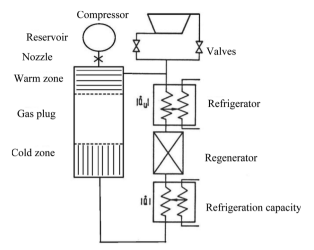


Figure 12.5 Pulse tube process.

surface of the cold head. As well as with the other gas refrigeration processes discussed here, the regenerator takes over the precooling of the inflowing highpressure gas.

For this, the pulse tube consists of three zones. The gas in the warm zone always moves back and forth between the warm end and the pulse tube middle, while analogously the gas of the cold zone always oscillates between the cold end and the middle range. The gas plug in the actual middle range builds up the temperature gradient and, as a result, functions exactly like a displacement compressor piston. The remaining gas filling works as a transformer of the pressure oscillation of the compressor. The advantage of the pulse pipe cooler exists in the clearly diminished vibration transmission to the surface to be cooled.

Concerning the size and weight of the refrigerator, the pulse tube cooler and the Gifford-McMahon cooler are very similar to each other. With a two-stage process and with ³He as working means, 1.27 K could be reached as the lowest temperature [10]. The refrigeration capacity lies in the order of magnitude of some 0.1 W at 2 K to 200 W at 80 K [10,11]. For a given refrigeration power, the prices of pulse tube coolers are still higher, but the pulse tube cooler absolutely has the potential to substitute for the Gifford-McMahon cooler in the medium term.

12.2.2

Cryostat with Liquid Cryogens

Cryostats are understood as low-temperature cooled, usually cylindrical vessels in which a specimen to be cooled or apparatus is introduced. Coolants for refrigeration are mostly liquid nitrogen (LN₂) and liquid helium (LHe). Following the refrigeration principle, a distinction is made between evaporator cryostats and bath cryostats. Through the high latent heat, the temperature can also be maintained in harsh environment until all cryogen has evaporated. Today, baths for cryopumps with liquid refrigerants for cooling the cold surfaces are only used in such exceptional cases that require high-temperature stability because their use is complicated and requires an access to the installation position to refill the cryogen.

Vacuum-insulated glass bath cryostat vessels named after Dewar are available in various versions. According to the experimental requirements, they are either completely silver-plated for the reduction of the heat input due to radiation or equipped with window strips or remain unplated without any silver coating. It is a special advantage of the glass cryostat that the experiment is visually accessible. Disadvantages are the brittleness of glass and the low but existent permeability of glass for helium.

Metal cryostats are usually made of stainless steel and copper. The thermal conductivities λ of these two materials show a considerable difference. Design takes advantage of this in the construction, on the one hand, in order to maintain a reduced heat input due to thermal conductivity as much as possible and, on the other, to permit optimum thermal equilibrium between the larger apparatus components (radiation shields, specimen carriers, etc.). The construction in Figure 12.6 illustrating a simple metal cryostat is analogous to that of a glass cryostat. Because metal containers are not permeable for helium, they can be built with a low leak rate ($<10^{-9}$ mbar ℓ s⁻¹) and follow-up evacuation is generally dispensable. For an increase in the service life, it is common to place an adsorbent (1) in the vacuum chamber.

The temperatures that are realized in cryostats typically correspond to the boiling temperature at normal pressure. However, if one designs the metal container as pressure vessel, every higher temperature can also be driven along the vaporization curve stably [12]. Pumping out in the gas chamber above the liquid phase is a proven process for realizing temperatures below the normal boiling temperature [13]. Another alternative is the use of the so-called evaporator cryostats [14]. For use as a cryopump, one often works with open cryostat vessels whose walls represent the pumping cold surface (cf. Section 12.4.3.2).

12.2.3

Measurement of Low Temperatures

Table 12.1 provides an overview regarding the temperature measuring methods in the cryo range.

12.3

Working Principles of Cryopumps

It has been known for a long time that gases and vapors bind to cooled surfaces [15]. While this effect has been used for the vacuum improvement

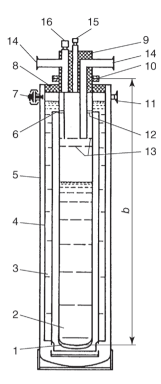


Figure 12.6 Bath cryostat from metal: (1) pocket with adsorbent; (2) helium Dewar with wide neck pipe; (3) nitrogen radiation shield; (4) tube with copper jacket; (5) vacuum jacket; (6) thermal bridge; (7) boiler safety valve; (8) covering ring; (9) lid; (10) Dewar

flange; (11) evacuation port with sealing valve; (12) LN₂-cooled baffle with spring contacts; (13) baffle cooled with helium exhaust gas; (14) exhaust gas pipe port; (15) sample entrance port; (16) helium inlet port.

practically since its discovery (cooling traps, baffles), the production of a highlevel high vacuum or ultrahigh vacuum with the aid of low-temperature cooled surfaces, so with cryopumps, has only found an increasing interest in the last 50 years.

According to DIN 28400, Part 2, "a cryopump is a gas-binding vacuum pump in which gases are condensed on low-temperature cooled surfaces and/or adsorb on low-temperature cooled sorbents (solids or condensate). The condensate and/or adsorbate is held at a temperature where the pressure of the gaseous phase is lower than the desired low pressure in the vacuum chamber."

Only vacuum pumps operating below 120 K are referred to as cryopumps. The selected temperature depends on the gas species to be pumped out and the desired vacuum level. The condensation pumps that work at higher temperatures are referred to as vapor condensers or simply condensers (see Chapter 8).

In contrast to all other pumps, a cryopump is able to achieve the theoretical pumping speed (e.g., for steam) and the pumping surface can be arranged

Table 12.1 Temperature measuring methods for temperatures from 1 to 300 $\rm{K}^{.1}$

Processes	Area (K)	V (cm ³)	Energy input (W)	<i>U(T)</i> (K)	Rep. (K)	Difference measurement	Measuring effort
Gas thermometer	2-800	>10	<10 ⁻⁴	10^{-2}	10^{-3}	Very poor	Very large ^{a)}
Vapor pressure thermometer	0.65–5 14–44 From 53	>≈1	<10 ⁻⁴	10^{-3} 10^{-2} 10^{-2}	10^{-4} 10^{-3} 10^{-2}	Very poor	Small
Industrial Pt resistor thermometer	20-500	<1	10^{-6} to 10^{-4}	10^{-1} b)	10^{-1}	Poor	Large
Industrial RhFe resistor thermometer	1-400	<1	10^{-5} to 10^{-4}	10^{-1} b)	10^{-1}	Poor	Large
Thermistor	70-400	10^{-2} to 10^{-1}	<10 ⁻⁵	10 ^{-1 b)}	10^{-1}	Poor	Small
Carbon resistor/thick film resistor (RuO_2)	0.1-300	10^{-2} to 10^{-1}	<10 ⁻⁵	10 ^{-2 b),c)}	10 ^{-2 c)}	Poor	Small
$Cernox^{^{TM}}\ resistance\ thermometer^{d)}$	1-10 10-400	10^{-1}	<10 ⁻⁵	10 ^{-2 b),c)} 10 ^{-3 b),c)}	10 ^{-2 c)} 10 ^{-3 c)}	Poor	Small
Carbon glass resistance thermometer	1 - 300	10^{-1}	<10 ⁻⁵	10 ^{-2 b),c)}	$10^{-2 \text{ c}}$	Poor	Small
Semiconductor diode (Si, GaAs)	1-30 30-400	10^{-2}	10 ⁻⁵	10 ^{-2 b)} 10 ^{-1 b)}	10^{-2} 10^{-1}	Poor	Small
Capacitive thermometer (SrTiO ₃)	1-290	10^{-1}	<10 ⁻⁵	12	1	Poor	Large
Thermo couples: Cu constantan AuFe-chromel	70-400 2-300	10 ⁻⁴ e)	<10 ⁻⁴	10 ^{-2 b), f}	10 ^{-2 f)}	Very well	Small

V: volumes of the sensor; U(T): measurement uncertainty with approximately 95% confidence interval; Rep.: repeatability of the measurement.

<sup>V: volumes of the sensor; U(1): measurement uncertainty with approximately 95% a)
a) For calibrating purposes.
b) Individual calibration.
c) Relative uncertainty U(T)/T/repeatability.
d) Cernox™ is a registered trademark of Lake Shore Cryotronics, Inc.
e) Only the measuring point, packaging, and thermal contact are not considered.</sup>

f) Relative uncertainty of differences in temperature $U(\Delta T)/\Delta T$.

¹⁾ Fellmuth, B. (2006) Low temperature metrology. PTB, Berlin, private communication.

directly inside the recipient, that is, without additional conductance losses through connection flanges. This is why the first cryopumps were used for space simulation chambers. Only here, conventional pumps were not able to achieve the required pumping speeds, so that the high expenditure for cryogenic technology seemed justified. The impressive progress in the technology of the regenerative gas refrigeration equipment (Section 12.2) has made refrigeration easier in the meantime to the extent that today there are an entire series of applications with which cryopumps with lower pumping speeds successfully compete with conventional pumps or frequently even represent the most advantageous technical solution.

Because the form of the cold surface can be adapted entirely to the available space for installation, the full pumping speed is also available for pumping processes in places that are otherwise barely accessible. As an additional advantage, no working fluid vapors enter the recipient. The capacity of a cryopump is limited because the pumped out gas remains bound to the cold surface. This is not a disadvantage in the high- and ultrahigh-vacuum regime, the primary range of application of the cryopumps, due to the low gas volumes involved. For applications at higher pressures or with processes that require a continuous gas flow being pumped, cryopumps must be regenerated regularly.

The underlying pumping mechanism in cryopumps is the gas binding to the cold surfaces via van der Waals forces [16]. The energetic interaction is strongly dependent upon the type of the binding surface. Accordingly, two cryopump mechanisms are primarily distinguished, the condensation and the (ad-)sorption. In practice, it is often not possible to clearly separate these mechanisms.

12.3.1

Gas Condensation

Condensation describes, in the strict thermodynamic sense, only the gaseous/liquid phase transition. Because it is frequently not relevant to the pumping effect whether the pumped gas is solid or liquid, the name "condensed phase" is used in the linguistic usage for both cases. In the best case, the ultimate pressures achievable at the given temperature of the cold surface through condensation result from the sublimation equilibrium pressure of the solid phase. Figure B.8 illustrates the equilibrium curves of gases and other substances relevant to vacuum technology; corresponding numerical values are summarized in Table 12.2. Aside from the components in air (Table A.6) including water, some hydrocarbons and mercury are listed.

Three groups of gases are to be distinguished. While the saturation pressure of water, carbon dioxide, and the higher hydrocarbons already reaches values p $< 10^{-7}$ Pa at the temperature of the liquid nitrogen (T = 77 K), the saturation pressures of methane, argon, oxygen, and nitrogen drop down to this region only at temperatures in the range $T \approx 20-30$ K. Finally, reducing the saturation pressure of neon and hydrogen down to values $p < 10^{-7}$ Pa requires temperatures in the range of liquid helium, so $T = 4.2 \,\mathrm{K}$ and lower are necessary.

Table 12.2 The vapor pressures of substances relevant to vacuum technology [17–19] extrapolated in the low-pressure range.

		Temperature in K for vapor pressure in Torr (133 Pa)																	
Symbol	Substance	Bp (K)	Fp (K)	10-12	10-11	10-10	10-9	10-8	10-7	10-6	10-5	10-4	10-3	10-2	10-1	10°	10 ¹	10 ²	10 ³
He	Helium	4.2	-	0.268	0.288	0.310	0.335	0.366	0.403	0.45	0.50	0.57	0.66	0.79	0.99	1.27	1.74	2.64	4.52
H_2	Hydrogen	20.4	13.4	2.88	3.01	3.21	3.45	3.71	4.03	4.40	4.84	5.38	6.05	6.90	8.03	9.55	11.7	15.1	21.4
Ne	Neon	27.1	24.6	5.79	6.11	6.47	6.88	7.34	7.87	8.45	9.19	10.05	11.05	12.30	13.85	15.8	18.45	22.1	27.5
N_2	Nitrogen	77.3	63.2	19.0	20.0	21.1	22.3	23.7	25.2	27.0	29.0	31.4	34.1	37.5	41.7	47.0	54.0	63.4	80.0
Ar	Argon	87.3	83.8	21.3	22.5	23.7	25.2	26.8	28.6	30.6	33.1	35.9	39.2	43.2	48.2	54.4	62.5	73.4	89.9
O_2	Oxygen	90.2	54.4	22.8	24.0	25.2	26.6	28.2	29.9	31.9	34.1	36.7	39.8	43.3	48.1	54.1	62.7	74.5	92.8
CH_4	Methane	111.7	90.1	25.3	26.7	28.2	30.0	32.0	34.2	36.9	39.9	43.5	47.7	52.9	59.2	67.3	77.7	91.7	115.0
Kr	Krypton	120.0	116.0	29.3	30.9	32.6	34.6	36.8	39.3	42.1	45.4	49.3	53.9	59.5	66.3	74.8	86.0	100.8	129.4
O_3	Ozone	161.3	22.2	40.0	42.1	44.3	46.9	49.8	53.0	56.7	60.8	65.8	71.6	78.6	87.2	101.1	116.0	136.2	165.4
Xe	Xenon	165.0	161.4	40.5	42.7	45.1	47.7	50.8	54.2	58.2	62.7	68.1	74.4	82.1	91.5	103.5	118.5	139.5	170.0
C_2H_6	Ethane	184.6	89.9	46.4	48.8	51.4	54.4	57.8	61.5	65.8	70.8	76.5	83.4	91.4	101.9	113.7	130.3	153.9	189.8
CO_2	Carbon dioxide	194.6	-	62.2	65.2	68.4	72.1	76.1	80.6	85.7	91.5	98.1	106.0	114.5	125.0	137.5	153.5	173.0	198.0
NH_3	Ammonia	239.8	195.4	74.1	77.6	81.5	85.8	90.6	95.9	102.0	108.5	116.5	125.5	136.0	148.0	163.0	181.0	206.0	245.0
C_3H_6O	Acetone	329.5	178.2											a)	191.0	213.8	242.1	280.9	337.0
C ₂ HCl ₃	Trichlorethylene	360.4	200.2											a)	211.5	234.5	264.4	305.8	369.2
H_2O	Water	373.2	273.2	118.5	124.0	130.0	137.0	144.5	153.0	162.0	173.0	185.0	198.5	215.0	233.0	256.0	284.0	325.0	381.0
H_2O_2	Hydrogen peroxide	431.0	271.5												a)	289.0	323.3	368.6	431.5
I_2	Iodine	456.2	386.8	147.5	154.0	161.5	169.5	178.5	188.5	199.5	212.0	226.0	243.0	262.0	285.0	312.0	345.0	389.0	471.0
Hg	Mercury	630	234.4	161.2	169.6	178.8	189.2	200.8	214.1	229.1	246.4	266.8	290.8	320.0	355.2	399.6	457.2	534.8	645.1

a) No vapor pressure equation is provided for the solid phase. Bp: boiling temperature; Fp: melting temperature.

In principle, any gas can be pumped when the selected temperature is sufficiently low. The temperature $T = 20 \,\mathrm{K}$ is sufficient to reach UHV conditions for all gases except neon, hydrogen, and helium. However, hydrogen is particularly interesting for vacuum technology because many materials are able to release it in vacuum. As a result, it is the most relevant residual gas component in the UHV range.

In the use of the vapor or sublimation pressure curves, it has to be considered that these represent the state of the phase equilibrium. In the equilibrium, a net mass flow of zero is reached at the cold surface, that is, no more usable pumping effect appears. Hence, the curves describe the theoretically possible ultimate pressure. In order to achieve a sufficiently high pumping speed in an application with gas flow constantly being pumped in the vacuum, a nonequilibrium must therefore be maintained while the gas to be pumped is thermodynamically oversaturated, usually two decades in pressure. Temperatures of $T < 3.5 \,\mathrm{K}$ are necessary for the binding of hydrogen to a cold surface in UHV if one does not involve the binding mechanisms of cryotrapping and cryosorption that are addressed in the following.

Example 12.1

Using the values from Table 12.2 yields the following: To reach a pressure of 10⁻⁶ Torr with the pumping from H₂ (according to an equilibrium temperature of 4.4 K), a temperature of 3.7 K must be made available at the cold surface (according to an equilibrium pressure of 10^{-8} Torr).

The properties of the formed ice layers depend on the pumping history in a complicated manner and change during growth [20,21].

The capacity of a condensation cryopump is almost infinite provided that the temperature of the cold surface can be maintained. Of course, the growth of a thick layer of ice increases the need for refrigeration; the finite thermal conductivity of the ice leads to an increase of the temperature on the ice surface that can lead to instabilities [22]. In practice, condensate layers of more than 10 mm in thickness can readily be formed. The efficiency of a condensation pump is close to the theoretical maximum [23].

12.3.2

Cryosorption

With cryotrapping and cryosorption, the efficiency of a cryopump can be noticeably raised and, hence, the operational temperature range can be moved to higher temperatures. Indeed, the attainable capacities are smaller than those with cryocondensation. The point of the vapor pressure or sublimation pressure curve that refers to the theoretically attainable pressure at a given temperature

on the surface of the condensate will be replaced by the sorption phase equilibrium curve. This is described with the aid of the sorption isotherms that describe the dependence of the pressure on the gas amount at the sorbing surface at a given temperature. The generally applied fundamental relationships are addressed in Chapter 6.

With cryosorption, the gas binds to the cold surface through physical adsorption forces. The interaction is complex and depends not only on the temperature, but also on the type of the surface. Specific sorbent materials are used for this, such as activated carbon, zeolite, or nanoporous materials that possess an especially high active surface through their porosity. Also a condensate layer condensed prior to the beginning of the pumping of a higher boiling gas (e.g., argon snow) has a sufficiently high porosity.

At a given temperature, the adsorption phase equilibrium always lies at pressures below the saturation vapor pressure. Hence, through adsorption, gas can also be pumped in the undersaturated range, that is, at considerably higher temperatures than would be necessary for condensation. This is of great significance to the economic pumping for the barely condensable gases such as hydrogen, helium, and neon. By means of cryosorption, at 5 K vacua in the 10^{-7} Pa range can also be easily achieved for these gases.

Figure 12.7a illustrates the cryosorption of hydrogen to solid ammonia. In Figure 12.8, the molar adsorption enthalpy ΔH of hydrogen to different gas condensates is plotted for various condensation temperatures $T_{\rm K}$ of the adsorbent. It is shown that carbon dioxide and ammonia are particularly suitable as an adsorbent for hydrogen. Furthermore, it is evident that for all adsorbents, a range of the condensation temperature T_K exists in which layers with particularly strong bindings are formed [24].

The use of gas condensates as a cryosorbent offers certain advantages compared with the use of solid adsorbents. First, the required good thermal contact with the cold surface is provided with condensates. Second, the cryopumps can be regenerated through the evaporation of the condensate and the production of a new layer at relatively low temperatures in a simple manner. Nevertheless, due to the clearly more complicated process and the need to accept a foreign gas into the system, cryosorption in gas condensates is only used in special cases [25,26]. All commercial refrigerator cryopumps use solid adsorbents.

12.3.2.1 Solid Adsorbents

The main challenge with the application of solid adsorbents (molecular sieves, activated carbon) for pressure reduction in high and ultrahigh vacuum through cryosorption is the heat transfer of the adsorbent to the cold surface. Cooling can only occur through the thermal conductivity in the adsorbent itself because the heat transfer through the gas to be pumped is negligible under low-pressure conditions. This is why it is essential to establish a high-quality thermally conducting contact, normally by adhesive bonding. The selection of a suitable adhesive must be ensured. Before cooling down the apparatus, the adsorbent must be degassed through bake-out as much as possible.

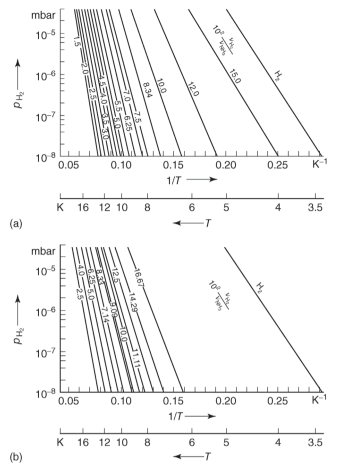


Figure 12.7 (a) Cryosorption of hydrogen to solid ammonia: adsorption equilibrium of H_2/NH_3 condensates for different molar ratios $v_{\rm H_2}/v_{\rm NH_3}$; production of the NH₃ condensates at T=7.99 K [17]. (b) Cryotrapping of hydrogen

with ammonia vapor: hydrogen partial vapor pressure over mixed NH₃/H₂ condensates with different molar ratios $v_{\rm H_2}/v_{\rm NH_3}$; production of the mixing condensates at $T=7.99\,\rm K$ [24].

The desorption during the regeneration as well as the total cryosorption pumping process strongly depends on the selected cryosorbents. Due to the great relevance that cryopumps had gained in large nuclear fusion experiments, large-scale qualification programs for a vast number of potential cryosorption materials (sinter metals, porous ceramics and metal meshs, silica gel, molecular sieves, activated carbons) were established in the 1990s [27–29]. Alongside this, not only the sorbing materials were examined, but an attempt was also made to find an optimum joining technology (resistance to the thermal cycling during regeneration and a good thermal conductivity) on the cooled surfaces (bonding, brazing, plasma coating). These programs were necessary since no experience in

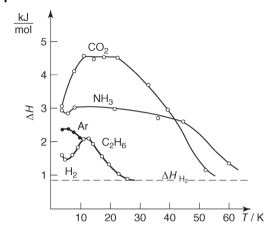


Figure 12.8 Molar sorption enthalpy ΔH of hydrogen to selected adsorbents (gas condensates) versus condensation temperature T_K of the adsorbent. Dashed line: condensation of $H_{2,aas}$ to H_{2,solid}.

usage within the cryogenic temperature range existed with the manufacturers of industrial adsorbents.

In practice (e.g., with all commercial pumps), activated carbon has become the most important sorbent, particularly the nonpolar, microporous variations [30]. The types that are gained from the carbonization of coconut shells are usually used. The natural source product results in a not too narrow pore size distribution, so that the interaction with the gas particles to be pumped is equally good for all particle types (Figure 12.9).

As a general rule in commercial pumps, activated carbon material pressed into cylindrical pellets is bonded to the cold surface with an epoxy resin, as Figure 12.10 indicates.

The transferability of the sorption equilibrium data from one carbon type to another is limited; therefore, literature data must be used that implies a high degree of uncertainty. Figure 12.11 clarifies this as an example for the pairing



Figure 12.9 Pore structures of granular coconut activated carbon. The scale is 100 µm (left), 10 μm (middle), and 1 μm (right).





Figure 12.10 Activated carbon-coated cryosurface of a refrigerator cryopump. The left image shows the complete arrangement with nine pumping surfaces (covered with glued

activated carbon pellets), and the right image shows a single surface (covered with granular material).

hydrogen on microporous coconut activated carbon at 77 K. For other activated carbon types, differences within an entire order of magnitude result. The cryosorption of helium below 10 K depends very strongly on the temperature (Figure 12.12).

Other aspects to be considered in the correct choice of cryosorption materials are the following:

- The cross-sensitivity of the cryosorption materials; that is, a reduction in the nominal pumping speed when the heavily volatile trace gases that require particularly high regeneration temperatures accumulate during the running time [32].
- The mobility of the sorbed particles; that is, the capacity of the pump can be increased when the particles that are initially pumped on the surface can be transported deeper into the pores through a defined energy input (heating) [33].

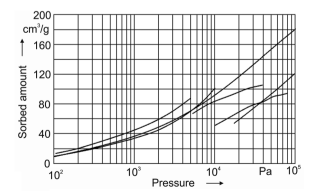


Figure 12.11 Sorption equilibrium for H₂ on activated carbon at 77 K. Different literature data are shown (summarized in Ref. [31]). The substance amount is indicated in cm³ sorbed hydrogen (at standard conditions) per gram of activated carbon.

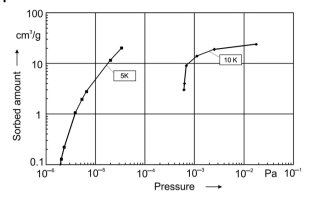


Figure 12.12 Sorption equilibrium for helium on microporous coconut activated carbon (Figure 12.9).

In the rough vacuum range, particularly as clean fore pumps for ion getter pumps, liquid nitrogen-cooled cryosorption pumps with zeolite filling have been demonstrated to work very effectively (see Section 11.2).

12.3.2.2 Cryotrapping

The cryotrapping pumping method can be utilized to reach even lower equilibrium pressures. This refers to the pumping of a lower boiling and thus at the same time difficult to condense gas with the aid of a higher boiling gas. In contrast to cryosorption where the gas to be pumped binds on a surface, with cryotrapping it is trapped in mixed condensate snow [34]. Argon, methane, carbon dioxide, ammonia, and higher hydrocarbons have been examined as a condensation partner for hydrogen thus far. At the given temperature of the cold surface, the resulting mixed condensates have a hydrogen partial pressure that is by several orders of magnitude smaller compared with the vapor pressure $p_{\rm H_2}$ above the surface of the pure condensate (cf. Table 12.2). Figure 12.7b demonstrates this exemplarily with mixed condensates from ammonia and hydrogen. The comparison with Figure 12.7a reveals that with the same molar ratio of ammonia to hydrogen, the drop in vapor pressure is by approximately 30% smaller with cryosorption than with the cryotrapping.

For practical application, cryosorption offers the advantage over cryotrapping that the gas condensate serving as adsorbent, with a negligible vapor pressure at the operating temperature of the cryopump, can be produced before starting the actual pumping process. In addition, cryosorption with solid adsorbents offers the advantage that no assist gas flow is required. Hence, cryotrapping is applied only on the laboratory scale.

Of course, with the condensing of gas mixtures on cold surfaces, cryosorption naturally also occurs together with cryotrapping. Therefore, an exact distinction between the processes according to the mentioned mechanisms is not possible.

12.4

Design of Cryopumps

Aspects of cryotechnology and vacuum technology must be combined in the design of cryopumps. Cryotechnology implies the generation of lower temperatures and the steady-state cooling of the cold surfaces used for pumping. Vacuum technology implies the creation of a high conductance value for the path that the particles have to take on their way to the pumping surface as well as an efficient interaction of the particles to be pumped with the cold surfaces.

12.4.1

Cryotechnological Constructive Parameters

The housing of a cryopump is usually at ambient temperature while the cold surfaces are actively cooled. As well as with other cryogenic components, the total thermal load of the cold surfaces $\dot{Q}_{\rm tot}$ has different contributions:

- solid-state thermal conduction through connected pipes, wiring, and so on, perhaps also through the originating ice layers $(\dot{Q}_{\rm E})$,
- gas thermal conduction through the residual gas between the cold surface and housing (\dot{Q}_G) ,
- radiant heat (\dot{Q}_{Rad}) , and
- enthalpy input of the gas to be pumped (cooling down to the cold temperature and phase transition enthalpy) ($\Delta \dot{H}$).

$$\dot{Q}_{\text{tot}} = \dot{Q}_{\text{F}} + \dot{Q}_{\text{G}} + \dot{Q}_{\text{Rad}} + \Delta \dot{H}. \tag{12.3}$$

The solid-state thermal conduction is commonly minimized through the selection of suitable materials with low thermal conductivity and/or small pipe diameters. The effect of the gas thermal conduction becomes negligible as soon as one works within the molecular flow range. Figure 12.13 qualitatively shows the gas heat transfer in relation to the pressure with different Knudsen numbers. In HV and UHV range, the condensation heat for the gases relevant here also lies in the $80-800\,\mathrm{J\,g^{-1}}$ range due to the negligibly small condensing volume flows.

With this, the radiant heat of 300 K on the cold surfaces is commonly the dominant contribution to the thermal load.

12.4.1.1 Thermal Radiation

In addition to other quantities, the heat transferred via thermal radiation also depends on the emissivity ε of the cold surface. With the construction of cryostats, it is suitable to lower the emissivity of the cold parts as far as possible by polishing or gold plating. Nevertheless, this only matters for cryopumps if low gas amounts are pumped so that the condensate layer deposited on the cold surface remains thin. With an increasing layer thickness of the gas condensate, the emissivity of the cold surface increases. While a polished copper surface has $\varepsilon = 0.03$, $\varepsilon = 0.9$ is to be calculated with a layer thickness of the gas condensate of 1 cm.

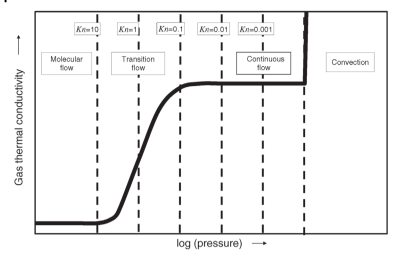


Figure 12.13 Influence of the pressure on the gas heat transfer.

According to the Stefan-Boltzmann law, the radiated power from the surface element dA of a black or gray body of the temperature T, that is, the total energy (heat) flux emitted by dA in all directions dI, if $\sigma = 5.67 \times 10^{-8} \,\mathrm{W m^{-2} K^{-4}}$ is the Stefan-Boltzmann constant, is

$$dI = \varepsilon \sigma T^4 dA. \tag{12.4}$$

The emissivity for a black body is $\varepsilon = 1$; for a gray body, $\varepsilon < 1$ regardless of the wavelength or frequency in the radiation spectrum. Equation (12.4) is only valid for $\varepsilon \leq 1$ = constant. For the evaluation of the radiation losses occurring in cryogenic apparatuses, this validity can be assumed with sufficient proximity. As Eq. (12.4) suggests, a body B2 at higher temperature T_2 will radiate more energy than a cooler body B1 with the lower temperature T_1 than vice versa so that a net energy flow $I_{\text{Radiation}}$ flows from B2 to B1. The value of $I_{\text{Radiation}}$ also depends on the geometry of the bodies 1 and 2; simple radiation formulas are obtained only for simple geometrical arrangements. Such simple arrangements include concentric cylinders, concentric spheres, and parallel planes, if it is assumed that edge disturbances due to the finite dimensions of the arrangements (edges) are negligible. For that, the linear dimensions of the radiant surfaces A_1 and A_2 of bodies B1 and B2 must be large in comparison with the (constant throughout the entire arrangement) distance between A_1 and A_2 .

For the case that body 2 (radiating surface A_2 , temperature T_2 , emissivity ε_2) is concentrically encasing body 1 (A_1 , $T_1 < T_2$, ε_1), we find the radiation flow [36]

$$I_{\text{radiation},2\to 1} = A_1 \frac{\sigma}{(1/\varepsilon_1) + (A_1/A_2)[1/(\varepsilon_2 - 1)]} (T_2^4 - T_1^4).$$
 (12.5)

If $T_1 > T_2$, then I is negative; that is, the energy is directed from 1 to 2. The radiation flow between parallel planes results from Eq. (12.5) for $A_1 = A_2$.

Introducing a "radiation parameter" C_S characterizing the geometrical arrangement

$$C_{S} = \frac{\sigma}{(1/\varepsilon_{1}) + (A_{1}/A_{2})[(1/\varepsilon_{2}) - 1]},$$
(12.6)

Eq. (12.5) can be written in the well-known form:

$$I_{\text{radiation},2\to 1} = A_1 \cdot C_S(T_2^4 - T_1^4).$$
 (12.7)

Example 12.2

The helium container (internal part of the Dewar vessel (2)) of the cryostat shown in Figure 12.6 has a radiating surface (cylinder plus plane bottom (or sphere zone)) $A_1 = 1.42 \,\mathrm{m}^2$, and its temperature is $T_1 = 4 \,\mathrm{K}$. It is concentrically surrounded by a similarly formed jacket of the surface $A_2 = 1.58 \,\mathrm{m}^2$ that is held by nitrogen bath (3) at the temperature $T_2 = 77$ K. The containers and jacket consist of copper that has been polished during production and thus, according to experience for long-term operation, is characterized by an emissivity $\varepsilon_1 = \varepsilon_2 = 0.1$. For $A_1/A_2 = 0.9$, one calculates $C_5 = 0.313 \times 10^{-8} \,\mathrm{W m^{-2} \, K^{-4}}$. Then Eq. (12.7) provides the radiation flow $I_{\text{radiation},2\rightarrow1}=0.16\,\text{W}$. If the nitrogen cooling of the external jacket is omitted, then $T_2 \cong 300 \text{ K}$ and this results in $I_{\text{radiation},2 \to 1} = 36.8 \text{ W}$.

In most cases in which two surfaces are in radiant heat exchange, one cannot assume as in the preceding section that the total radiation emitted by one surface hits the other surface. The calculation of the radiation exchange between these surfaces is then required to find out which energy contribution of the energy emitted by one surface hits the other and is then absorbed. This is a task to be purely geometrically solved. In Figure 12.14, the situation of two surfaces at an arbitrary position to each other is shown.

The angle β refers to the angle between the connecting line of the length s between both surface elements and their surface normals. For this general case, the radiation parameter C_{12} results in

$$C_{12} = \frac{\sigma \cdot \varepsilon_1 \cdot \varepsilon_2 \cdot \varphi_{12}}{1 - (1 - \varepsilon_1) \cdot (1 - \varepsilon_2) \cdot \varphi_{12} \cdot \varphi_{21}},$$
(12.8)

with the quantities φ_{ii} referred to as viewing factor:

$$\varphi_{12} = \frac{1}{\pi \cdot A_1} \int_{A_1, A_2} \frac{\cos \beta_1 \cdot \cos \beta_2}{s^2} dA_1 dA_2.$$
 (12.9)

Also, this formula is valid only when the reabsorption of the radiation from one surface reflected by the other can be neglected, which is well fulfilled if both

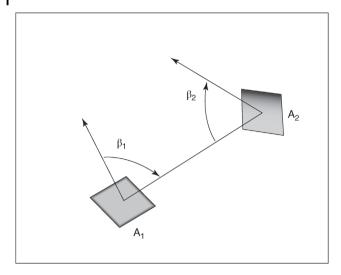


Figure 12.14 Derivation of the radiant heat exchange between two arbitrary surface elements.

surfaces have high emissivities or large spacing. Exact calculations are difficult and complicated [36]. In practice, one takes the viewing factors from lookup tables [37].

Diffuse radiation, that is, that with a constant emissivity independent of direction, follows the cosine distribution that was introduced in Section 4.4.1 for the particle flow. In principle, the photons of the radiation can be treated like particles in the range of the free molecular flow. For exotic geometries, one can thus calculate the viewing factors with Monte Carlo methods; the approach to this is quite similar to the calculation of particle trajectories (cf. details on its implementation in Section 5.2).

The radiative heat transfer is reduced when the radiant interior surface of the housing is given a small emissivity (e.g., by electropolishing with ε = 0.01). Nevertheless, the resultant radiant heat input must be even further reduced in most cases.

If the cold surface is enclosed with a nitrogen-cooled radiation shield (baffle) at a temperature $T\!=\!77\,\mathrm{K}$, the heat flow occurring through radiation is less than 1% of the value that would occur at a wall temperature of $T\!=\!300\,\mathrm{K}$ (see Example 12.2). Such a baffle primarily consists of blind covers or angled plates (chevrons). These are optically tight and often blackened (absorption a near 1) in order to allow as little reflected radiation to reach the cold surface as possible. Then, the resultant thermal load consists of the transmitted radiation of 300 K through the shield and the full radiation from the shield temperature on the pumping cold surfaces. The shield also leads to a minimization of the enthalpy portion in Eq. (12.3) since the shield design is chosen such that the pumped

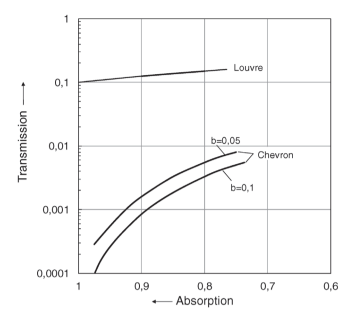


Figure 12.15 Transmission coefficient for heat radiation through a baffle (opening angle 60°. diameter/height ratio = 6.8) with a surface overlap b and a given absorption [38].

particles are completely thermally accommodated, that is, not impinging with ambient temperature, but rather with the temperature of the shield.

Figure 12.15 demonstrates an example of the dependence of the transmission on the absorption for two often used baffle geometries. Baffles have a flow resistance and therefore reduce the pumping speed of the cryopump.

12.4.1.2 Thermal Conduction

For the thermal conduction heat flow $I_{conduction}$ through a wire or rod of constant cross section A and the length L with a temperature difference $T_2 - T_1$ $(T_2 > T_1)$ between both ends and that shows no lateral heat loss (such as through radiation), we find the following equation:

$$I_{\text{conduction}} = \frac{A}{L} \int_{T_1}^{T_2} \lambda(T) dT.$$
 (12.10)

 $\lambda(T)$ is the temperature-dependent thermal conductivity of the rod material that is different along the rod as a result of the temperature gradient.

In Table 12.3, values for the "thermal conduction integrals" according to Eq. (12.10) of a number of materials that are frequently used in cryotechnology are listed. The lower limit of the integral was set to 4K, while the upper limit

Table 12.3 Thermal conduction integrals for selected materials frequently used in cryotechnology [35].

	$\int_{T_1=4}^{T_2} K \lambda(T) dT \text{ (W cm}^{-1})$											
	Сорре	Manganin	Brass	Aluminum	Stainless steel AISI 303, 304, 316, 317	Glass, Pyrex, quartz, borosilicate	Plastics					
<i>T</i> ₂ (K)	Electrolytic cold- drawn copper	Phosph. desoxid.						Teflon	Nylon			
6	8.0	0.176		0.053	1.38	0.0063	2.11	1.13	0.321			
10	33.2	0.785		0.229	6.07	0.0293	6.81	4.4	1.48			
20	140	3.95		1.12	27.6	0.163	20.0	16.4	8.23			
40	406	16.4	1.54	4.76	96.2	0.824	58.6	50.8	38.5			
60	587	35.5	3.74	10.4	170	1.98	115	93.6	85.9			
76	686	53.9	5.76	16.2	220	3.17	175	130	131			
80	707	58.9	6.28	17.7	232	3.49	194	139	142			
100	802	85.8	8.98	26.5	284	5.28	292	187	204			
120	891	115	11.8	36.5	330	7.26	408	237	269			
140	976	146	14.7	47.8	376	9.39	542	287	336			
160	1060	180	17.8	60.3	420	11.7	694	338	405			
180	1140	215	21.0	73.8	464	14.1	858	390	475			
200	1220	253	24.3	88.3	508	16.6	1030	442	545			
250	1420	353	33.4	128	618	23.4	1500	572	720			
300	1620	461	43.8	172	728	30.6	1990	792	895			

runs gradually from 6 to 300 K. The value of the integral of an interval T_3 – T_4 results from subtraction:

$$\int_{T_3}^{T_4} \lambda(T) dT = \int_{T=4 \text{ K}}^{T_4} \lambda(T) dT - \int_{T=4 \text{ K}}^{T_3} \lambda(T) dT.$$
 (12.11)

The thermal conduction heat load over parts of an apparatus can therefore be minimized by appropriate design measures.

Example 12.3

The stainless steel neck pipe of a cryostat has the following dimensions: external diameter D = 50 mm, wall thickness S = 0.4 mm, and length L = 320 mm. Thus, the cross section is determined by

$$A = 3.14(25^2 - 24.6^2) \text{ mm}^2 = 62.3 \text{ mm}^2 = 0.623 \text{ cm}^2.$$

The temperature at the bottom end of the neck pipe is helium temperature $T_1 = 4 \text{ K}$, while at its upper end a thermoconducting connection to a nitrogencooled radiation shield holds the temperature at $T_2 = 80 \, \text{K}$. The value of the thermal conduction integral of stainless steel for the interval between T_1 and T_2 can be found from Table 12.3:

$$\int_{T_{-}=4K}^{T_{2}=80 \text{ K}} \lambda(T) dT = 3.49 \text{ W cm}^{-1}.$$

With this, Eq. (12.10) yields the thermal conduction flow to be $I_{\text{conduction}} = 68 \text{ mW}$.

In Figure 12.16, the thermal conductivity λ of different gas condensates is shown in relation to the temperature T. Included for comparison are thermal conductivity curves of several other materials. Apparently, solid gases can show a comparably high thermal conductivity.

The curves shown here are valid for very pure specimens, partially present as a single crystal and partially as polycrystalline specimens. The thermal conductivity can be reduced by one to two orders of magnitude by deviations in the microstructure of the layers, particularly through formation of loose, snowy-like or amorphous condensates. With the buildup of thick layers, the thermal conductivity of the condensate becomes a limiting factor. The surface temperature of the condensate increases with the growing layer thickness, that is, increasing thermal impedance. This means that with increasing layer thickness, the pumping speed of the cryopumps drops continuously and can even reach the value zero. Within certain limits, a reduction of the cold surface's temperature can compensate for the effect. Nevertheless, this leads to an increase in the refrigeration power.

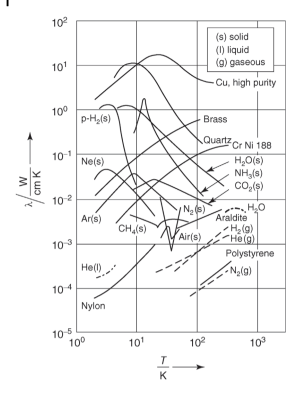


Figure 12.16 Thermal conductivity of solid gas condensates.

12.4.2

Vacuum Technology: Design Parameters

As explained above, cryopumps are characterized by large pumping probabilities. The maximum pumping speed $S_{\rm id}$ that can be provided with a cryopump in high vacuum over the inlet cross section A is identical to the maximum possible volume flow that is impinging on this surface. This was derived from the kinetic theory of gases for the free molecular flow in Section 3.2.4 (Eq. 3.50). In this ideal case, all particles are pumped, and there are no reflections or back flow. In the real case, the effective pumping speed is always smaller. This decrease is described by the capture coefficient P_c that indicates which proportion of the inflowing particles of the gas is pumped. Following Eq. (3.50) (cf. the limitation on the use of this equation in Section 12.5.3, if the pump is operated near the ultimate pressure), the following expression is obtained:

$$S = P_{c} \cdot S_{id} = P_{c} \cdot \frac{\overline{c}}{4} \cdot A = P_{c} \cdot \sqrt{\frac{R \cdot T}{2 \cdot \pi \cdot M}} \cdot A. \tag{12.12}$$

The pumping speed is a function of the pumping geometry (P_c, A) and the gas to be pumped (T, M). The capture coefficient is a complicated function of the gas

and particularly of the geometry that must be individually calculated for every pump with Monte Carlo simulations. The capture coefficient of one and the same pump can be by more than a factor of 5 smaller for He in comparison with N₂; nevertheless, this is partially compensated by the reciprocal square root dependence on the molar mass.

The ideal pumping speed for a real pump is in the first place decreased by the limited transmission probability of the path of the particles from the vacuum vessel to the pumping surface, and afterward through the not 100% sticking probability of the particle on the pumping surface. The sticking probability denotes the ratio of the particles that adhere to the pumping surface, in relation to the number of all particles that hit on the pumping surface.

While the transmission probability is a geometrical constant of cryopump design (as long as layers of ice do not significantly narrow the opening slit through the baffles), the sticking probability is only constant for condensation pumps. For cryosorption pumps, the sticking probability is dependent in a complex manner on the loading (gas to be pumped and possibly previously accumulated other gas) and therefore changes during the operation time pump. Thus, the capture coefficient P_c of a pump can be attributed to the geometric influence (expressed by the transmission probability w) and the physical interaction (expressed by the sticking probability α). Both influences can be optimized separately. For complicated geometries, the dependence can only be quantified by means of a proper Monte Carlo simulation [39].

In the simplistic case of a parallel geometry (i.e., the shield is arranged in parallel with the cold surface), the following equation can be used for an estimation:

$$\frac{1}{P_c} = \frac{1}{\alpha} + \frac{1}{w} - 1. \tag{12.13}$$

Figure 12.17 shows numerical values for the transmission probability of the most frequently used baffle construction elements [40,41]. In connection with Figure 12.15, it is clear that for a sound cryopump design, a compromise between the demand for low thermal loading and high pump efficiency must be found, which can result in very different results depending on the specific application.

For condensation (heavy gases), the sticking probabilities take a value of unity. For sorption (light gases), sticking probabilities that are smaller than unity are found, but they can normally be still considered as constant in the usual operating temperature range of the cryopump, as exemplarily shown in Figure 12.18 with the cryosorption in activated carbon. An exemption to this is the sorption of helium for which a strong dependence on the temperature must be considered in order to design the cycle times optimally.

The concept of the capture coefficient is very helpful in the design of vacuum systems because it allows us to uncouple the design of the pump itself from the rest of the system ("reverse engineering"). Normally, a set of requirements are given to the vacuum system (e.g., to maintain a specific pressure with a specific gas flow at a specific location in the recipient) and information on the size of the cryopumps is requested. Here, with suitable computational programs, one can now describe the

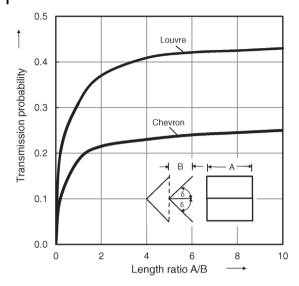


Figure 12.17 Transmission probability for a baffle (angle $\delta = 45^{\circ}$) depending on the ratio of diameter A and overlap B.

cryopump via a capture coefficient that is varied until the system requirements are fulfilled. Once the required capture coefficient is known, one can proceed to the next step and develop the design details of the cryopump [42].

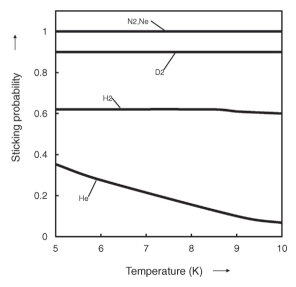


Figure 12.18 Sticking probability of different gases in unladen coconut activated carbon (granular, glued to the surface) [39].

Construction Principles

Every cryopump consists of three essential subsystems, the pumping surface, the thermal shield, and the refrigeration supply. For the user, the type of the refrigeration supply represents the primary classification parameter. For cryopumps, the same design and construction guidelines apply as described for cryostats in Section 12.2. There are cryopumps that contain a cryogen bath, there are those that are constructed following the evaporator principle, and, finally, those that are operated with a cryorefrigerator. Designs of the pump and the type of the cryogenic supply can be varied within broad limits and, with this, the principle of cryopumping can be matched to fit very different practical application cases. This is illustrated in the following sections with some selected proven designs.

Besides the traditional LHe bath cryopumps and the cryogen-free refrigerator cryopumps, there are also multitude of special constructions that are supplied with forced flows of cryogens (either liquid (LHe, LN) or with supercritical helium gas at the customary temperatures of 4 and 80 K). Because the design of the cryopump subsystems with the tools described above and some experience can be done sufficiently well, cryopumps can be developed in a very specific manner adapted to the available space. Moreover, there is the possibility to install the cryogen-driven cryopumps directly within the vacuum recipient. In this case, the conductance-limiting effects of a connection flange can be avoided. Section 12.6 presents selected special constructions more precisely.

12.4.3.1 Bath Cryopumps

This type of pump is no longer relevant for commercial applications, but is well suited to illustrating the typical aspects of the cryopump design. The pumping surface facing the vacuum is on its backside directly contacted with a bath of liquid helium. According to the specific application, a blank metal plate (copper, aluminum, stainless steel) or a plate coated with activated carbon can be used as a pumping surface. In order to increase the service life and control the temperature well, the pumping surface is shielded with passively cooled baffles that are connected to a bath container filled with liquid nitrogen.

The simplest forms of cryopumps as illustrated in Figure 12.19 correspond in principle to the well-known cold trap. The wall of the internal container (1) filled with liquid helium acts as the pumping cold surface (8). In order to reduce helium consumption, the helium container is surrounded by a nitrogen-cooled radiation shield (7). Because this is formed as an opaque baffle, the full pumping speed of the cold surface cannot be utilized. Nevertheless, when pumping gas mixtures that include higher boiling components, the radiation shield is also a selectively working pump.

The fact that bath cryopumps can likewise be adapted to cope with special requirements is illustrated in Figure 12.20 with a bath cryopump that has been optimized in any possible way, developed by Benvenuti in the 1970s at CERN. It

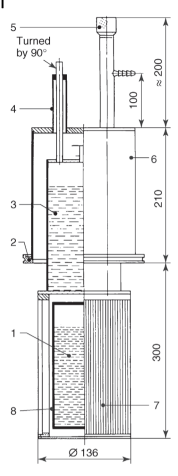


Figure 12.19 Bath cryopump: (1) liquid helium condensation surface. Pumping speed for N₂: in tank 8 (volume 1.25 ℓ); (2) connection flange DN 150; (3) liquid nitrogen (tank volume 1.5 ℓ); (4) LN₂ filling and exhaust gas ports; (5) LHe inlet with lateral exhaust ports; (6) outer housing; (7) baffle; (8) helium-cooled

2250 ℓ s⁻¹; pumping speed for H₂: 7000 ℓ s⁻¹; LHe consumption at 4.2 K and $p < 10^{-5}$ mbar: $0.035 \ell h^{-1}$; LHe lifetime at 4.2 K and $p < 10^{-5}$ mbar: 35 h; LN_2 consumption at 4.2 K and p $< 10^{-5}$ mbar: 0.75 ℓ h⁻¹.

can be operated at a temperature of the cold surface of 2.3 K and has, in this case for hydrogen, according to pump size, a pumping speed of 4500 and $1100\,\ell\,\mathrm{s}^{-1}$. The accessible ultimate pressure amounts to approximately 10⁻¹¹ Pa. Pumps with this design are equipped with an insulating vacuum; the consumption of liquid helium is, thus, very low, particularly since the radiation shield is also cooled with cold helium gas.

Figure 12.21 shows photographs of a manufactured bath cryopump.

For special applications, cryopumps can be "tailor-made" using standard helium condensers in the refrigerator operation. Such pumps were developed for applications in nuclear fusion (cf. Section 12.6).

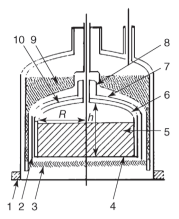


Figure 12.20 CERN - bath cryopump with long operational duration to generate extremely low gas pressures: (1) connection flange; (2) clearance; (3) baffle; (4) pumping surface (condensation surface); (5) liquid

helium; (6) Ne filled volume for generating insulating vacuum; (7) radiation shield (Ag plated on interior, exterior blackened); (8) neck, copper-plated; (9) liquid nitrogen; (10) protective housing (silver-plated).

12.4.3.2 Refrigerator Cryopumps

The modern alternative to the bath cryopump is the refrigerator cryopump whose refrigeration supply is produced by a gas refrigerator and generally follows the Gifford-McMahon principle (Section 12.2.1.2). The refrigerator works with helium gas in a closed cycle and consists of a force-cooled (water or air) compressor unit and a cold head with two temperature stages that is connected to the compressor unit by flexible pressure pipes (spatially separated). Hence, this pump model is also called cryogen-free.

The cold head of a two-stage Gifford-McMahon refrigerator allows for a refrigerator cryopump (Figure 12.22) to be mounted easily. The radiation shield and baffle of the cryopump are mechanically contacted with the first stage. The cold surfaces are contacted with the second stage. These are designed as plane

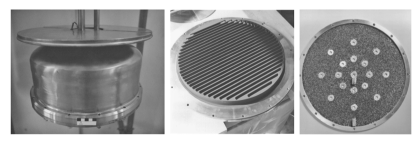


Figure 12.21 Construction elements of a LHe bath cryopump. Left: LHe container made of stainless steel. Middle: baffle made of copper, optically blackened. Below: pumping surface made of aluminum, with activated carbon glued on it.

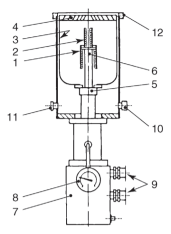


Figure 12.22 Refrigerator cryopump with integrated cold head: (1) cold surface of second stage (pumping surface); (2) adsorbent (interior coating of the cold surface); (3) radiation shield; (4) baffle; (5) cold head, first stage; (6) cold head, second stage; (7) cold head motor

with housing and electrical connections; (8) He gauge for hydrogen vapor pressure thermometer; (9) He gas connections to compressor; (10) boiler safety valve; (11) fore-vacuum connecting flange; (12) high-vacuum connecting flange.

cranked copper metal sheets that are placed in parallel at a short distance. The interior sides of these sheets that are turned toward each other are coated with activated carbon. At the temperatures given above for the second stage, the geometry of the arrangement and the surface structure of the low-temperature cooled plates (cold surfaces) ensure that any easily condensable gases will primarily condense on the exterior surfaces in the solid phase, while the gases that are difficult to condense (hydrogen, neon, helium) are sorbed on the interior surfaces that have been coated with activated carbon.

The first stage is cooled with a refrigeration capacity between 10 and $80\,\mathrm{W}$ – according to refrigerator type and in relation to the load – to a temperature from 50 to maximum $80\,\mathrm{K}$. In the second stage, a refrigeration capacity of $2-5\,\mathrm{W}$ at a temperature between 8 and $20\,\mathrm{K}$ is available – according to load. These days, GM refrigeration equipment with sufficient refrigeration capacity (several watts) down to $4\,\mathrm{K}$ is also available. Together with the use of a sorbent material for the second stage, helium can also be pumped efficiently.

Example 12.4

For a refrigerator pump shielded with a louvred baffle and having the cryopanel at a temperature of 10 K (Figure 12.22), the following vacuum technology-specific values arise.

Heavy gases are pumped directly at the baffle with the following specific values: $\alpha = 1$, w = 1, and $P_c = 1$. With Eq. (12.12), the pumping speed related to the

surface, for example, for water (applied at 300 K), can be calculated as $14.9 \,\ell\,\mathrm{s}^{-1}\,\mathrm{cm}^{-2}$.

Nitrogen is pumped on the front side of the cryopanel by condensation. Assuming a transmission probability of the baffle of w=0.4 (Figure 12.17) and using the approximate formula (12.13), the pump-specific values result in $\alpha=1$, w=0.4, and $P_c=0.4$. The pumping speed related to the surface is calculated as $11.9 \, \ell \, \text{s}^{-1} \, \text{cm}^{-2}$.

Hydrogen and helium are pumped on the backside of the cryopanels (coated with activated carbon) by sorption.

The pump-specific values for hydrogen amount to α = 0.6, w = 0.4, and P_c = 0.32 (the additional conductance loss between the cryopanel front and back is neglected). The surface-related pumping speed is calculated as 13.4 ℓ s⁻¹ cm⁻².

An even smaller sticking probability yields for helium (Figure 12.18): $\alpha = 0.05$, w = 0.4, and $P_c = 0.05$. The pumping speed related to the surface in this case is calculated as $1.6 \, \ell \, \text{s}^{-1} \, \text{cm}^{-2}$.

The example clearly shows that the cryopumps pump the different gas species at different locations with different pumping speeds specific to the gas. With a sound cryopump design, the transmission probability and sticking probability should stand in a "suitable" ratio. With the clearly smaller sticking probability of helium in comparison with all other gases, pumps always show helium enrichment effects within the pump when exposed to helium-containing mixtures [43].

The interactions between the different processes can be very complex. Thus, the buildup of a frost layer at the thermal shield can, on the one hand, lead to a lower transmission probability, for example, in the case of carbon dioxide, and, on the other hand, lead to an additional cryosorption effect in the condensate. If there is an intensified condensation pumping at the sorbent layer, the pumping speed can be reduced for lighter gases. In view of this complexity, it is not easy to predict accurately the pumping of gas mixtures. On the other hand, these effects can also be utilized to separate gas mixtures into individual fractions (cf. Section 12.5.9).

Refrigerator cryopumps of varying sizes whose pumping speeds for air reach from 800 to $60\,000\,\ell\,s^{-1}$ are mass produced. Depending on where the connection flange is mounted, the refrigerator cryopump can be directly built into the vacuum vessel (built-in installation model, Figure 12.23a) or as the appendage pump connected to the vacuum vessel (built-on attachment model, Figure 12.23b).

Refrigerators with accordingly high refrigeration power are necessary for the operation of large cryopumps with high pumping speed. Nevertheless, there is also the possibility to use a small refrigerator for the cooling of the cold surfaces and to cool radiation shields and baffle with liquid nitrogen that is taken from a container integrated into the radiation shield (Figure 12.23a). Through the high refrigeration capacity of LN_2 , such pumps can also pump substantial amounts of

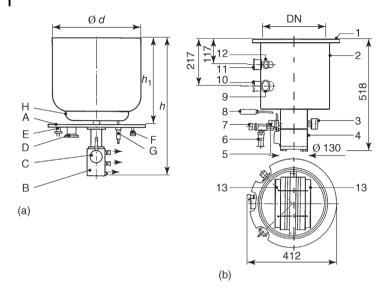


Figure 12.23 (a) Built-in type refrigerator cryopump: (A) connection flange for vacuum vessel; (B) refrigerator cold head; (C) thermometer; (D) fore-vacuum connection; (E) thermocouple; (F) boiler safety valve; (G) supply to ring tank (H) for LN_2 ; h = 540-900 mm, h_1 (radiation shield) = 240–530 mm, d (radiation shield) = 146–484 mm. (b) Built-on

type, DN = 292 mm; (1) connection flange; (2) housing; (3) vapor pressure thermometer; (4) cold head; (5, 7) He connection, (6) boiler safety valves; (8) electrical supply wiring; (9) fore-vacuum connection; (10) connection for regeneration heater; (11) connection for temperature sensor; (12) connection for vacuum gauge; (13) baffle (for radiation shielding).

water vapor at higher pressures. The losses of liquid nitrogen can be substituted for with an automatically functioning replenishment device that can be connected to the nitrogen container of the cryopumps through a vacuum jacketed pipe (near G in Figure 12.23a).

For the processes with which gas continuously flows in the cryopump or is flushed, for example, when sputtering with Ar, a maximum inlet pressure may not be exceeded. This is dependent upon the refrigeration capacity of the refrigerator and the dimensions of the cold surfaces. As approximate values for the maximum allowed tolerable vessel pressure $p_{\rm max}$, the following can be used for normal cryopumps:

$$p_{\text{max}} \approx 0.5 \, \text{Pa for N}_2, \text{Ar},$$

 $p_{\text{max}} \approx 0.15 \, \text{Pa for H}_2.$

For the user, refrigerator cryopumps can be used in the same way as other conventional pumps, that is, the only external interface is the electricity supply (for the helium compression cycle) and the connection flange. As a result, these devices are much easier to use than bath cryopumps and have almost entirely substituted them in the industrial practice. In the United States, cryopumps are

traditionally used for semiconductor applications [44], while in Europe, the turbomolecular pumps are given preference.

12.5 Characteristics of a Cryopump

As with any other vacuum pump, the cryopumps also have a set of characteristics: starting pressure p_{St} , ultimate pressure p_{end} , pumping speed S, service life $\overline{t}_{\rm B}$, gas capacity C, crossover value, maximum tolerable pV flow, and regeneration time. For these values, the manufacturers of refrigerator cryopumps have agreed on generally accepted guidelines for the measurement [45] that are the basis for the numerical values listed in the catalogues.

12.5.1

Starting Pressure

The explanations of adsorption pumps for the rough vacuum range in Section 11.1 demonstrate that, in principle, a cryopump can also be started at atmospheric pressure. Nevertheless, this is unsuitable for several reasons. While the mean free path of the gas molecules is smaller than the dimensions of the recipient (typically with p > 0.1 Pa), the thermal conductivity through the gas is so large that the thermal load onto the cold surface would be unacceptably high. Moreover, even if the temperature could be maintained, a relatively thick condensate layer would already form on the cold surface at startup. The capacity available for the actual operation phase of the cryopump would be noticeably reduced as a result.

The "start" pressure p_{St} must be selected in such a way that the heat load transferred to the cold surface through the condensation flow - up to a brief initial phase – is not greater than the cryogenic power Q_2 in the second stage of the pump. With a small volume V (e.g., with cooling of the closed pump), choosing a higher p_{St} is permitted since with the low pV amount in the pump housing, the initial phase is brief.

With a large volume, a very low p_{St} must be required; thus, the recipient must be evacuated with a fore pump to the required p_{St} before the start. The latter can be switched off after achievement of the start pressure. With cooled refrigerator cryopump, the $p_{St}V$ value lies significantly higher than that with a cooling pump due to the above considerations. This is of large relevance for high-vacuum systems in batch operation mode, where during the purging of the vacuum vessel the cryopump is locked by a valve and remains in operation all the time.

Experience has shown that the empirical relationship

$$\frac{p_{\text{St}} \cdot V}{\dot{Q}_2} \le 3 \,\text{kPa} \,\ell \,\mathbf{W}^{-1} \tag{12.14}$$

is sufficient to consider all factors for practical purposes. \dot{Q}_2 is the refrigeration capacity of the refrigerator in the second stage at T = 20 K.

Example 12.5

A cryorefrigerator pump with a refrigeration capacity $\dot{Q}_2 = 5 \,\mathrm{W}$ at $T_{\mathrm{K}} = 20 \,\mathrm{K}$ is used for the evacuation of a ventilated container with a volume of V=50 ℓ , while the already cold cryopump is separated from the container by means of a valve. The container must be pre-evacuated according to the above relationship at a pressure of 0.3 kPa before the valve to the cryopump may be opened.

12.5.2

Ultimate Pressure

The lowest pressure that can be reached in a given system with a given pump arrangement is called the ultimate pressure. In a sealed isotherm system where the surrounding vessel and the evaporating substance have the same temperature T, a "saturation vapor pressure" $p_s(T)$ or a saturation particle number density $n_s = p_s(T)/kT$ (steady-state equilibrium) is attained. A steady-state equilibrium also exists in the case of cryocondensation, and the condensation particle number flux density $j_{N,\text{cond}}$ on the cold surface is equal to the vaporization particle number flux density $j_{N,\text{evap}}$; both are provided by Eq. (3.129). While for the evaporating gas in Eq. (3.129), the temperature T_{cp} of the cold surface and the saturation particle number density $n_s = p_s(T_{cp})/kT_{cp}$ are to be used, for the condensing gas the (higher) temperature of the wall $T_{\rm w}$ and the particle number density $n_g = p_g/kT_w$ are to be used (approximately, because the gas in thermal contact with the wall is mixed with the evaporating gas), whereby p_g is the gas (vapor) pressure in the vessel. In the final state that corresponds to the ultimate pressure p_{ult} , the condensation particle number flux density

$$j_{N,\text{cond}} = \sigma_{c} \cdot n_{g,\text{ult}} \frac{\overline{c}_{w}}{4} = \frac{\sigma_{c}}{4} \cdot \frac{p_{g,\text{ult}}}{kT_{w}} \cdot \sqrt{\frac{8kT_{w}}{\pi m_{a}}}$$
(12.15)

and the evaporation particle number flux density

$$j_{N,\text{evap}} = \frac{\sigma_{\text{c}}}{4} \cdot \frac{p_{\text{s}}}{kT_{\text{cp}}} \cdot \sqrt{\frac{8kT_{\text{cp}}}{\pi m_{\text{a}}}}$$
(12.16)

must be equal to each other. The subscript "w" at the mean velocity of \overline{c} of the gas molecules indicates that the gas temperature is set to the wall temperature $T_{\rm w}$ of the recipient. From that, one finds the ultimate pressure

$$p_{\rm ult} = p_{\rm s} \sqrt{\frac{T_{\rm w}}{T_{\rm cp}}}. (12.17)$$

Here, p_s is the saturation vapor pressure of the gas to be pumped at the cold surface temperature T_{cp} of the cryopump and T_{w} is the wall temperature of the recipient. Because the latter is almost always higher than the cold surface temperature $T_{\rm K}$, the accessible ultimate pressure is almost always higher than the saturation vapor pressure of the condensate.

Example 12.6

For a given recipient temperature $T_{\rm w} = 80 \, \rm K$ (shield temperature), the cold surface temperature T_{cp} determines the ultimate pressure p_{ult} of a cryopump:

$$T_{\rm cp} = 2.5 \, {\rm K}, \quad p_{\rm ult} = p_{\rm s}(2.5 \, {\rm K}) \cdot \sqrt{rac{80 \, {\rm K}}{2.5 \, {\rm K}}} pprox 11 p_{\rm s}(2.5 \, {\rm K}), \ T_{\rm cp} = 20 \, {\rm K}, \quad p_{\rm ult} = 2 \cdot p_{\rm s}(2.5 \, {\rm K}).$$

In the case of nitrogen for which the saturation vapor pressure p_s at 20 K according to Table 12.3 has the value $p_s = 1.3 \times 10^{-11}$ mbar, this results in an ultimate pressure $p_{\rm end}$ of 2.6×10^{-11} mbar.

According to the guidelines, the ultimate pressure is measured 24 h after the start of the cool-down process.

12.5.3

Pumping Speed

The pumping speed of a cryopump is traditionally measured in a Fischer-Mommsen dome (cf. Section 15.5). A constant gas flow is established with this that, at most, should amount to 30% of the maximum allowed pV flow rate. This restriction ensures that the flow within the cryopumps is in the free molecular range. As soon as one enters into the transitional range, the pumping speed rises as long as the temperature of the cold surfaces can be maintained. This effect is frequently used to receive more pumping speed from an available cryopump (cf. Section 12.6.1).

If the final state is not reached yet, $j_{N,\text{cond}}$ according to Eq. (12.15) is greater than $j_{N,\text{evap}}$ according to Eq. (12.16), the system pumps and the difference of the two expressions represents the particle number pumping flux density at the pressure *p* in the recipient:

$$j_{N,\text{pump}} = j_{N,\text{cond}} \cdot \left(1 - \frac{j_{N,\text{evap}}}{j_{N,\text{cond}}} \right) = j_{N,\text{cond}} \cdot \left(1 - \frac{p_s}{p} \sqrt{\frac{T_w}{T_{cp}}} \right). \tag{12.18}$$

The surface-related pumping speed S_A is given by the pumping speed (Eqs. 4.19 and 4.6) of a pumping surface A divided by A. Because

$$S_{A} = \frac{S}{A} = \frac{q_{V}}{A} = \frac{q_{N}}{nA} = \frac{j_{N}}{n},$$

it follows that

$$S_{\rm A} = \frac{1}{4} \sigma_{\rm c} \overline{c}_{\rm w} \cdot \left(1 - \frac{p_{\rm s}}{p} \sqrt{\frac{T_{\rm w}}{T_{\rm cp}}} \right). \tag{12.19}$$

Again, the subscript "w" at the mean velocity of \overline{c} of the gas molecules indicates that the gas temperature is set to the wall temperature $T_{
m w}$ of the recipient. For the condensation coefficient σ_c (cf. also Table A.11), the ratio of the number of gas molecules condensing on the cold surface to the number of the gas molecules impinging on the surface, the value can be set to $\sigma_c = 1$ with sufficient accuracy for the low temperatures in question. Similarly high values for σ_c are valid for sorption with the exception of helium (Figure 12.18).

The condition for the validity of Eq. (12.19) is that the thermal equilibrium, that is, the probability distribution of the particle speeds, is not or is only slightly disturbed by the pumping process. This condition is fulfilled when the dimensions of the pumping cold surface are small in comparison with the vessel surface.

Following Eq. (12.19), a cold surface of the area $A_{\rm K}$ has the (theoretical) pumping speed

$$S = A_{\mathcal{K}} \cdot S_{\mathcal{A}}. \tag{12.20}$$

This is equally valid for condensation and sorption.

Equation (12.19) for S=0 yields the ultimate pressure following Eq. (12.17). With $p \gg p_s \sqrt{T_w/T_{cp}}$, S or S_A is practically equal to the maximum possible value determined by the volume flow density, $S_{A,\text{max}} = \overline{c}_{\text{w}}/4 = \sqrt{RT_{\text{s}}/2\pi M_{\text{molar}}}$. This term has already been used in Eq. (12.12) and in Example 12.4.

For the different gases, different values of the surface-related maximum pumping speed arise, some of which are listed in Table 12.4. An additional reason for them to be maximum values is that, in practice, the condition of the nearly undisturbed thermal equilibrium is usually not met because large cold surfaces are needed to provide short pumping times and a good ultimate vacuum. Deviations also arise if the cold surface is surrounded with a cooled radiation shield.

12.5.4 Service Life

The mean operational life of a cryopump with constant pressure p_R in the recipient up to the "saturation" is referred to as service life \overline{t}_{S} .

The cases with condensation are considered in the following. For the case of sorption, the service life is directly limited by the capacity.

Table 12.4 The maximum surface-related pumping speed for selected gases at different
temperatures.

<i>T</i> _w (K)	$S_{A,max}(\mathscr{C} s^{-1} cm^{-2})$							
	O ₂	N ₂	Ne	H ₂				
293	11.0	11.8	13.9	43.9				
77	5.7	6.1	7.2	22.5				

If the pumping effect is based on condensation, it fundamentally depends on the condensation coefficient σ_c and, with this, on the surface temperature and structure of the condensate layer. \bar{t}_{S} is therefore a function of the thermal conductivity λ (Section 12.4.1.2) and the layer thickness x or the growth speed G_c of the layer. An estimated value for \bar{t}_S is obtained when the layer thickness is set to $x_{\text{lim}} = 0.5 \text{ cm}$ as an empirical value for the "limiting" layer thickness; thus,

$$\overline{t}_{S} = x_{\lim}/G_{c}. \tag{12.21}$$

A layer thickness of x originates on the cold surface through condensation; the temporal increase in the layer thickness (growth speed) dx/dt can be derived from the increase in the mass distribution of the layer $m/A = Nm_A/A$:

$$\frac{\mathrm{d}x}{\mathrm{d}t} = \frac{m_{\mathrm{A}}}{\rho} \cdot j_{N,\mathrm{cond}},$$

where m_A is the particle mass, ρ is the density of the condensate, and $j_{N,\mathrm{cond}}$ is the condensation particle number flux density following Eq. (12.15).

The density of the originated layer depends on the structure of the layer and this again from the condensation conditions; the structure becomes looser with lower temperatures. Values of the density of solid gas condensates are shown in Table 12.5. They can only be used for the calculation of G_c with a certain degree of uncertainty. For example, nitrogen at the pressure $p = 10^{-4}$ Pa produces a condensate layer with the thickness of 1 mm in approximately 2000 h.

Table 12.5 Density of liquid and so	lid gas condensates: calculated values.
-------------------------------------	---

Gas	$T_{Siede}(K)$	$ ho_{ m f1}({ m g~cm}^{-1})$	⁻³)at <i>T</i> (K)	$T_{Schmelz}(K)$	$ ho_{fest}(g\;cn$	n ⁻³)at <i>T</i> (K)
Ar	87.27	1.59	86	83.77	1.59	83.77
					1.65	40
					1.81	0 (rep.)
Ne	27.17	1.204	27.17	24.54	$0.8^{a)}$	
H_2	20.39	0.071	20.39	13.95	0.087	13.95
		0.076	14.8		0.088	2.1
N_2	77.33	0.81	77.4	63.15	0.95	63.15
					1.03	20.6
					1.14	0 (rep.)
O_2	90.18	1.14	90.18	54.36	1.43	20.6
		1.22	74.8		1.57	0 (rep.)
CO_2	194.7	_	_		1.56	194.2
					1.63	84.2
⁴ He	4.22	0.125	4.22		0.188	0 (rep.)
		0.146	1.62			

a) Assessed: $\rho(Ne) = \rho(Ar) \cdot M_r(Ne) / M_r(Ar)$.

12.5.5

Capacity (Maximum Gas Intake)

The amount of gas that can be absorbed by condensation or sorption until the pumping speed drops to 50% of the initial value after a complete regeneration is referred to as the capacity of a cryopump.

In the case of the continuous operation of a cryopump, the capacity is a limiting value. Should large gas volumes be pumped out continually, two cryopumps connected to the recipient through valves or gates are utilized in alternating operation. While one pump is in operation, the other is being regenerated.

12.5.6

Crossover Value

The crossover value is a characteristic parameter of an already precooled refrigerator cryopump that is connected to the vacuum chamber via a valve. It is defined as the maximum pV gas amount of nitrogen or argon that the vacuum chamber may contain so that, when opening the valve and initiating a suddenly developing gas load, over 3 s the temperature of the pumping surfaces does not rise above $T \approx 20$ K. The crossover value is usually given in Pa ℓ or mbar ℓ .

12.5.7

Maximum Tolerable pV Flow

The maximum tolerable pV flow q_{pV} is the gas flow into the continuously operating cryopump that heats the pump surfaces to a maximum of $T \approx 20 \,\mathrm{K}$ and not higher. \dot{Q} is usually given in mbar ℓ s⁻¹ and depends on the gas species.

12.5.8

Resistance to Thermal Radiation

This characteristic parameter shows the maximum temperature at which black radiation can reach the cold surfaces through the inlet flange so that the pumping surfaces do not heat to more than $T \approx 20 \,\mathrm{K}$ and the thermal shield to no more than $T \approx 130$ K. For this measurement, a heated blackened plate (emissivity $\varepsilon > 0.9$) is attached in a simple dome arrangement (see Section 15.5) at the pump inlet. This parameter characterizes the capability of the pump to "see" the temperatures, which is particularly important to the use of cryopumps in warm recipients (bake out).

12.5.9

Regeneration

As with all gas-binding vacuum pumps, the limited capacity requires a regeneration of the (refrigerator) cryopump after a certain period of operation, the duration of which depends on the operating conditions [46]. At regeneration, the cryopumps are isolated from the vacuum system and the cold surfaces are heated (primarily through direct electric heating or purging with warm gas) in order to release the accumulated gases that can then be pumped out by the connected fore-pump system. Should the vacuum system be continuously pumped, at least two pumps that are alternately operated are required.

The regeneration cycle consists of the following individual steps:

- i) Shut off the refrigeration supply.
- ii) Heat the pump (cold surfaces) to desorption temperature.
- iii) Pump out the pumping volume with a fore pump.
- iv) Connect the refrigeration supply and cool down to ultimate temperature.

There are different processes for heating the cold surfaces. Electric heaters that have the best thermal contact to the cold surfaces are frequently used. With refrigerator cryopumps, the heating elements are attached to the first and second stages of the cold head. Alternatively, the cold surfaces can also be heated by direct resistance heating or by purging with an inert warm gas. Recooling of the cold surfaces occurs in the same manner as at the startup of the cryopumps.

With commercial refrigerator cryopumps, the time required for regeneration can amount to between 30 and 300 min according to size of the pump and heating process utilized [47].

In many applications, the cryopump is already regenerated before reaching the capacity limit. Thus, it is generally aimed that after regeneration the ultimate pressure does not lead to an unacceptable pressure increase. From safety considerations, the type of the gas pumped can also be the reason to start regeneration, for example, to avoid an overly large amount of toxic or corrosive gases being stored. This is particularly valid for gases that can form explosive mixtures when in contact with air, such as, for example, hydrogen. In these cases, the partial pressure after regeneration has to be selected such that the minimum ignition pressure or, even in the instance of an air ingress event, the explosion limit is not reached [48,49]. Otherwise, the entire vacuum system must be designed as explosion pressure-proof, which is usually avoided.

The regeneration temperatures depend on the type of the gases pumped. The gas release curves are different; condensed gas is instantaneously released while the sorbed gas follows release kinetics that is dependent upon the pore size distribution of the adsorbent used and the temperature control, and does often lie in the order of magnitude of some minutes. In addition, whether the desorption takes place against the increasing pressure in the closed cryopump or whether the fore pump is already switched on naturally also plays a role. The optimum design of the regeneration step can be absolutely complicated. With commercial refrigerator cryopumps, the regeneration is one, fully automatic operation mode preconfigured by the manufacturer so that the user need not be concerned with details.

It is also to be considered that the sorption capacity is still so large at temperatures above the boiling temperature that a readsorption can occur; that is, the

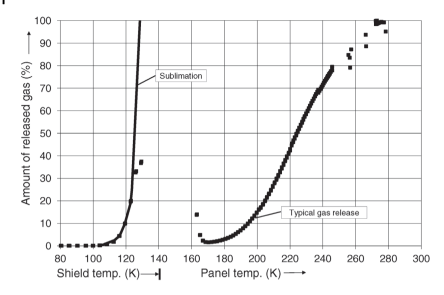


Figure 12.24 Example CO₂: illustration of the readsorption of a sublimated gas condensate on the activated carbon of the cryopanels during the cryopump regeneration.

gas produced by sublimation of condensate is immediately adsorbed again. The release characteristic of these gases also follows the desorption, not the sublimation (see Figure 12.24).

Figure 12.25 shows typical desorption release curves of selected gases from activated carbon. Regeneration above ambient temperature is only necessary for

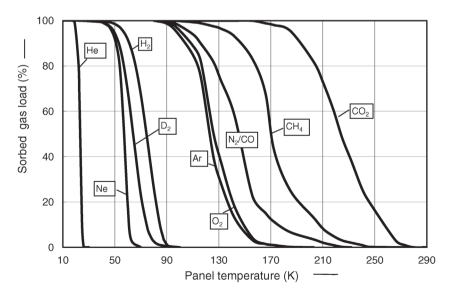


Figure 12.25 Gas release from activated carbon through desorption.

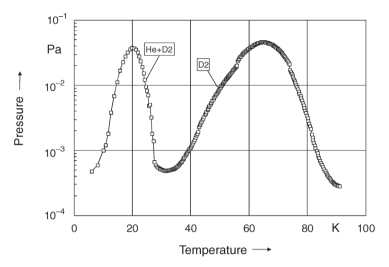


Figure 12.26 Temperature-controlled separation of helium (pumped gas mixture with 10% He and 90% deuterium) during regeneration. The heating rate amounts to approximately $0.05\,\mathrm{K\,s^{-1}}$.

water, higher hydrocarbons, solvent vapors, and so on [50]. For practical applications, 150 °C is sufficient. With refrigerator pumps, the sorption stage can also be completely exchanged after a certain time.

Figure 12.26 illustrates the possibility of also inducing a separation of the pumped gases [30] by suitable selection of the temperature control during the regeneration.

12.5.10

Performance Comparison

Table 12.6 compares the typical numerical values of a commercial refrigerator cryopump (DN 320). The numerical values reflect the range of the catalog data found.

12.6 Application Examples

Cryopumps are high- and ultrahigh-vacuum pumps. Some particular features are that they generate a hydrocarbon-free vacuum, show a high specific pumping speed (in relation to the size of the pump), have a high surface-related pumping speed for hydrogen, can be built in virtually any size, and, finally, can also be inserted *in situ*, that is, mounted directly in the recipient. Hence, cryopumps are also used in industrial facilities, particularly, however, in the vacuum facilities in large-scale research – nuclear fusion technology, space technology, particle

Table 12.6 Comparison of the parameters of DN 320 refrigerator cryopumps of different manufacturers.

Pumping speed (ℓ s ⁻¹)	9000-10 500
For water	3000-3200
For air	4500-5200
For hydrogen	2500-2700
For argon	1500-2300
For helium	
Maximum throughput (Pa m ³ s ⁻¹)	1.0-2.5
For argon	1.2
For hydrogen	
Capacity (Pa m ³)	1.5×10^5 to 3×10^5
For argon	1500-5000
For hydrogen	10-100
For helium	
Ultimate pressure (N ₂ equivalent) (Pa)	10^{-9} to 10^{-10}
Cool-down time (h)	1.5-2.5
Crossover (Pa m³)	35-50
Weight (kg)	30-50

accelerators, and beam guiding systems. Here, the cryopumps are specially custom-made and adapted for the specific application. The choice of the pump (size and type), also in view of the alternative use of other high-vacuum pumps [51], is determined completely by the particular operational requirements and conditions.

12.6.1 Cryopumps with Forced Cooling Using Supercritical Helium

Fusion technology has been a great driver in the development of very large cryopumps. The special requirements (exclusion of all organic materials, longest possible maintenance intervals due to the need for remote handling, and compatibility with high magnetic fields) are only to be fulfilled by cryopumps specifically developed for this. The pumping gas consists primarily of hydrogen (and its isotopes) and helium.

In the ITER fusion experiment, the cryopumps are force cooled with cold helium [52]. The advantage of the use of gaseous cryogenic helium compared with the liquid phase exists in the lower pressure losses and lower sensitivity compared with oscillating thermal loads. For the cryopanels, supercritical helium is used at 0.4 MPa and 4.5 K, and for the thermal shields, cold helium is used at 1.8 MPa and 80 K. Another specific feature in the operation of this cryopump is the particularly high gas flow that leads to the fact that there is no free molecular flow in the pumping volume, but rather it is a transitional flow that dominates. With this, it is possible to realize higher pumping speeds because the number of particle density increases in the pumped volume accordingly (Figure 12.27). The

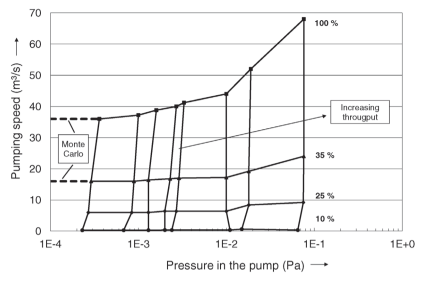


Figure 12.27 Pumping speed curves of the ITER cryopumps (for hydrogen) that are operated in the transitional range. The percentage numbers indicate the opening degree of the built-in valve.

price to be paid for this is the likewise increased thermal load on the cryosurface. Figure 12.28 shows the design of such a cryopump.

ITER has a total of six such pumps in a plasma chamber (1300 m³ volume). For explosion safety reasons, the maximum hydrogen amount to be pumped is limited so that, for example, the pumps must be regenerated every 10 min, which is clearly before reaching the capacity limit. Hence, of the six connected pumps, only four pumps are in continual pumping operation while the remaining two are being regenerated. An integrated valve is used for closing the pumps during the regeneration as well as for the control of the inflowing gas flow.

The heating of the fusion plasma occurs with extremely powerful particle beams. The world's largest cryopumps are required for the operation of these beam systems. They are used directly within the beam vacuum vessel and are not connected through a flange. Figure 12.29 shows the cryopump used for this at the JET nuclear fusion facility in England. Similarly, large systems are also currently being developed for ITER [53,54]. For these cryopumps, the reverse engineering method (cf. Section 12.4.2) yielded the required capture coefficient of 32%. In addition, high thermal loads from hot components (surface temperatures up to 150 °C) have an effect on the pump so that a simple louvred shield geometry is no longer sufficient. Nevertheless, the better shielded chevron geometry does not have a sufficient transmission probability (cf. Example 12.4, Figure 12.17). This is why an entirely new arrangement of the shield and pumping surfaces had to be developed [55].

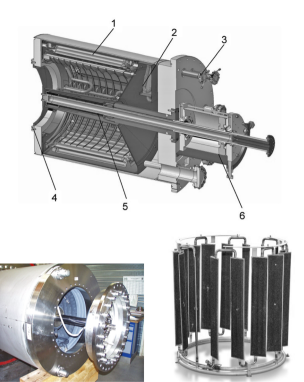


Figure 12.28 Design of the ITER cryopump. (1) Cryopanels (4.5 K); (2) thermal shields (80 K); (3) connections to the refrigeration supply (Johnston coupling); (4) inlet valve

(DN 800); (5) valve stem; (6) pneumatic valve actuators. The photos show the valve (left) and the cryopanel arrangement (right) from a reduced-size prototype.

12.6.2 Combined Refrigerator/Liquid Cryopumps

Refrigerator cryopumps are also used to evacuate large-space simulation chambers because they can generate an extremely clean high vacuum and be built as an individual unit for a high (nitrogen) pumping speed. The pumps for these applications are three-stage pumps [56]. The radiation shield and baffle are cooled in a separate cycle with liquid nitrogen at 77 K (Figure 12.30). Two twostage GM coolers are installed inside the pump and are connected with a condensation panel in the first stage at 50 K and with an adsorption panel (activated carbon) at 20 K in the low-temperature stage. The regeneration can be done with attached electric heaters or through forced ventilation with warm gas. With an opening diameter of 1.25 m, the nominal pumping speed amounts to 50 m³ s⁻¹ for nitrogen (capture coefficient $P_c = 0.35$) and hydrogen (capture coefficient $P_{\rm c} = 0.09$).

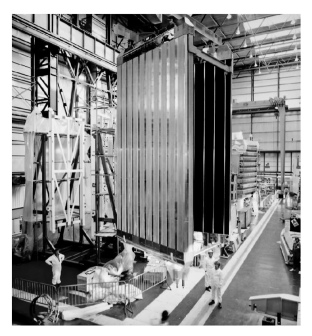


Figure 12.29 The cryopumps of the neutral beam heating system in the JET nuclear fusion facility (pumping speed is 6000 m³ s⁻¹ for hydrogen).

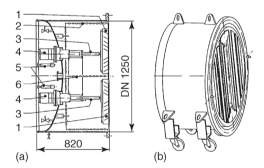


Figure 12.30 Cryopumps for evacuating a space simulation chamber. $S(N_2) = 5.5 \times 10^4 \, \ell \, s^{-1}$. (1) Baffle; (2, 6) radiation shields; (3) cold surfaces; (4) cold heads; (5) LN₂ connections.

12.6.3 Cryopumps with Forced Cooling Using Liquid Cryogen

Figure 12.31 shows a cryopump with forced liquid cooling developed for installation within the vacuum chamber [57]. Centrifugal compressors are used for both cryogenic cycles (LHe, LN2). The geometry was given and predefined since titanium sublimation pumps were to be substituted. With the change to the cryopumps, the capacity could be increased by a factor of 1000 and the pumping speed by a factor of 10.



Figure 12.31 Cryopumps with forced liquid cooling (height 4.50 m, width 1.50 m, depth 0.25 m). Pumping speed for hydrogen is $500 \, \text{m}^3 \, \text{s}^{-1}$.

The central part of the pump is a system of hydroformed activated carbon-coated panels that are supplied with liquid helium. The panels on the gas side are shielded with chevron baffles and on the rear with a closed shield. The two shield systems are supplied with liquid nitrogen. The nominal pumping speed amounts to approximately $500\,\mathrm{m}^3\,\mathrm{s}^{-1}$ for hydrogen. The cryopanels are coated with electric resistance heaters for defined regeneration.

In total, 2×37 vertical chevron sections per pump were manufactured. All cryopanels are connected in a series. While cooling the cryopanel circuit, a pressure loss of up to 0.5 bar is to be overcome.

12.6.4

Cryopumps in Large Research Applications

Cryopumps with gas condensates as adsorbents often find application in cases where the regeneration should take place at lower temperatures than the desorption of the solid adsorbent requires. Argon, in particular, has been successfully utilized to pump helium and hydrogen (as a precondensed snow for the cryosorption or as an assist gas spray for the cryotrapping). In the JET fusion facility, it has been found that a particle surplus in argon by a factor of 20 is necessary for the quantitative cryosorption of helium [58]. In the KATRIN neutrino experiment that is currently being constructed, argon snow is also used for the pumping of tritium (ultraheavy hydrogen). Partial pressures of less than 10^{-11} Pa [59] can be expected here.

The tubular vacuum chamber of particle accelerators (length of the chamber » dimensions of their cross section) is evacuated by a vast number of small high-vacuum pumps (turbomolecular pumps, sputter ion and NEG pumps) so that the use of cryopumps is limited on a (few) special cases. An example of this is the use of cryosorption pumps with activated carbon for the pumping of energetic particles (ions, electrons, synchrotron radiation) in the Large Hadron Collider at CERN [60].

Very large cryopumps are also used in test chambers for the development of ion thrusters for propulsion. Here, a pumping speed in the order of magnitude of 500 m³ s⁻¹ is required for xenon.

12.6.5

Cryopumps in Industrial Facilities

Small- and mid-sized (refrigerator) cryopumps are also coming into use in increasing numbers in production plants. Vacuum coating facilities [61] count among these, as well as cover sputtering and thin film deposition applications. Refrigerator cryopumps with nominal pumping speeds $S_n(N_2) = 5-30 \times 10^3 \, \ell \, s^{-1}$ have been recently used, particularly because of their high pumping speed for H_2O vapor $(S(H_2O) \approx 3S_n(N_2))$, for generation of a hydrocarbon-free vacuum for the evacuation of high-vacuum ovens. However, the connection of the cryopumps must be chosen with care to ensure that the thermal load from the oven is manageable.

12.6.6

Development Trends for Cryopumps

Concerning the design and the operating methods, cryopumps demonstrate a far larger variety than any other pump type. This is why the area of cryopumps is continuously evolving. Thus, for example, cryocondensation pumps with possibilities for continuous regeneration [62] were developed. For this, slowly rotating rotors were incorporated that mechanically scrape off the forming condensate and collect it or thermally evaporate and evacuate it locally. Furthermore, a cryopump with a thermal diffusion stage was developed for applications with an especially high throughput that leads to a viscous flow area within the cryopump [63].

Cryopumps are unmatched in their flexibility and versatility to be adapted to given conditions and specific requirements. One can return to commercial solutions that are available in the form of versatile standard models or develop individually tailored designs. Moreover, the cryopumps are characterized by a good scalability that allows for the construction of very large units. The demand for high-vacuum pumps with very high pumping speeds is continually rising with the result that the significance of the cryopumps will still increase in the future.

References

- 1 Rühlich, I. (2002) Gaskältemaschinen für HTSL-Anwendungen. KI Luft- und Kältetechnik, 8, 361-365.
- 2 Organ, A.J. (2005) Stirling and Pulse-Tube Cryocoolers, John Wiley & Sons, Inc., New York.
- 3 Ackermann, R.A. (1997) Cryogenic Regenerative Heat Exchangers, Plenum Press, New York.
- 4 Frey, H., Haefer, R., and Eder, F.X. (1992) Tieftemperaturtechnologie, VDI-Verlag, Düsseldorf.
- 5 Flynn, Th. (2004) Cryogenic Engineering, 2. Auflage, CRC Press, Boca Raton, FL.
- 6 Köhler, J.W.L. and Jonkers, C.O. (1954) Fundamentals of the gas refrigerating machine. Philips Tech. Rev., 16, 69-105.
- 7 McMahon, H.O. and Gifford, W.E. (1960) A new low temperature gas expansion cycle parts 1 and 2. Adv. Cryogen. Eng., 5, 354-368.
- 8 Radebaugh, R. et al. (1986) Comparison of three types of pulse tube refrigerators: new methods for reaching 60 K. Adv. Cryogen. Eng., 31, 779-789.
- 9 Xiao, J.H. (1995) Thermoacoustic heat transportation and energy transformation. Parts I-III. Cryogenics, 35, 15-35.
- 10 Jiang, N. et al. (2004) A ³He pulse tube cooler operating down to 1.27 K. Cryogenics, 44, 809-816.
- 11 Zia, J.H. (2004) Design and operation of a 4 kW liner motor driven pulse tube cryocooler. Adv. Cryogen. Eng., 49, 1309-1317.
- 12 Haas, H., Day, C., and Herzog, F. (2012) TIMO-2 - a cryogenic test bed for the ITER cryosorption pumps. Adv. Cryogen. Eng., 57, 1699-1705.
- 13 Streibl, B. et al. (2011) Operational behaviour of the ASDEX upgrade in-vessel cryopump. Fusion Eng. Des., 56-57, 867 - 872.
- 14 Klipping, G. (1973) Relations between cryogenics and Vacuum Technology up to now and in the future. Cryogenics, 13, 197.
- 15 Tait, P.G. and Dewar, J. (1874) Farther researches in very perfect vacua. Proc. R. Soc. (Edinburgh), 8, 348-628.

- 16 Maitland, G.C., Rigby, M., Smith, E.B., and Wakeham, W. (1981) Intermolecular Forces, Clarendon Press, Oxford, UK.
- 17 Honig, R.E. and Hock, H.O. (1960) Vapor pressure data for some common gases. RCA Rev., 21, 360.
- 18 Pollack, G.L. (1964) The solid state of rare gases. Rev. Mod. Phys., 36, 748.
- 19 Landolt-Börnstein, 6. Auflage, 1960, II/2a.
- 20 Haefer, R.A. (1989) Cryopumping, Clarendon Press, Oxford, UK.
- 21 Anashin, V.V. et al. (1997) Photon induced molecular desorption from condensed gases. Vacuum, 48, 785-788.
- 22 Syssoev, S.E., Eacobacci, M.E., and Bartlett, A.J. (2010) Controlled formation of condensed frost layers in cryogenic high vacuum pumps. J. Vac. Sci. Technol. A, 28, 925-930.
- 23 Hirth, J.P. and Pound, G.M. (1963) Condensation and Evaporation, Pergamon Press, Oxford, UK.
- 24 Becker, K. et al. (1972) Cryopumping of hydrogen by adsorption on condensed gases. Proceedings of the 4th International Cryogenic Engineering Conference, Eindhoven, 319 and S. 323.
- 25 Kachalin, G.V., Kryukov, A.P., and Nesterov, S.B. (1998) Adsorption of gaseous helium near Tλ at low pressures. Low Temp. Phys., 24, 97.
- 26 Yuferov, V.B. (1993) Sorption properties of vacuum cryocondensates. Low Temp. Phys., 19, 413.
- 27 Perinić, D., Haas, H., and Mack, A. (1994) Development of cryosorption panels for cryopumps. Adv. Cryogen. Eng., 39, 1553.
- 28 Sedgley, D.W., Tobin, G., Batzer, T.H., and Call, W.R. (1987) Characterization of charcoals for helium cryopumping in fusion devices. J. Vac. Sci. Technol. A, 5, 2572.
- 29 Anashin, V.V. et al. (2004) Molecular cryosorption properties of porous copper, anodized aluminium and charcoal at temperatures between 10 and 20 K. Vacuum, 76, 23-29.
- **30** Day, C. (2001) The use of active carbons as cryosorbent. Colloids Surf. A, 187-188, 187.

- 31 Day, C. and Hauer, V. (2002) Pore characterisation of cryosorbent carbon materials at cryogenic temperatures, in Fundamentals of Adsorption (eds K. Kaneko et al.), vol. 7, pp. 1093-1100.
- 32 Day, C. et al. (2001) New operational aspects of the ITER-FEAT primary vacuum pumping system. Fusion Eng. Des., **58-59**, 301.
- 33 Day, C., Kammerer, B., and Mack, A. (1998) Pumping performance of cryopanels coated with activated carbon. Adv. Cryogen. Eng., 43, 1327.
- 34 Boissin, J.C., Thibault, J.J., and Richardt, H. (1972) Method of an apparatus for cryopumping. Le Vide, 157, 103.
- 35 Johnson, R.W., Collins, S.C., and Smith, J.I., Jr. (1971) Hydraulically operated twophase helium expansion machine. Adv. Cryogen. Eng., 16.
- 36 Siegel, R. and Howell, J.R. (1981) Thermal Radiation Heat Transfer, 2nd edn, Hemisphere Publishing, New York.
- 37 Howell, J.R. (1982) A Catalog of Radiation Configuration Factors, McGraw-Hill, New York.
- 38 Benvenuti, C. et al. (1974) Characteristics, Advantages, and Possible Applications of Condensation Cryopumping. Le Vide, 169,
- 39 Day, C. et al. (2007) R&D and design for the cryogenic and mechanical vacuum pumping systems of ITER. Vacuum, 81, 738.
- 40 Obert, W. and Perinić, G. (1992) Pumping speed and thermal load analysis of the JET appendage cryopump by Monte Carlo calculation. Vacuum, 43, 163.
- 41 Saksaganskii, G.L. (1988) Molecular Flow in Complex Vacuum Systems, Gordon and Breach, New York.
- 42 Luo, X. and Day, C. (2008) Test particle Monte Carlo study of the cryogenic pumping system of the Karlsruhe tritium neutrino experiment. J. Vac. Sci. Technol. A, 26, 1319.
- 43 Day, C. (1998) The use of a highresolution quadrupole gas mass spectrometer system for selective detection of helium and deuterium. Vacuum, 51, 21-30.
- 44 Engers, B. and Bauer, H.U. (1999) Cost-Effective PVD Coatings in batch system. Surf. Coat. Technol., 116, 705.

- 45 Welch, K.M. et al. (1999) Recommended practices for measuring the performance and characteristics of closed-loop gaseous helium cryopumps. J. Vac. Sci. Technol. A, 17, 3081-3095.
- 46 Longworth, R.C. and Bonney, G.E. (1982) Cryopump regeneration studies. J. Vac. Sci. Technol., 21, 1022-1027.
- 47 Mundinger, H.J. et al. (1992) A new cryopump with a very fast regeneration system. Vacuum, 43, 545-549.
- 48 Edesky, F.J. and Stewart, W.F. (1996) Safety in the Handling of Cryogenic Fluids, Plenum Press, New York.
- 49 Graham, W.G. and Ruby, L. (1979) Cryopump measurements relating to safety, pumping speed, and radiation outgassing. J. Vac. Sci. Technol. A, 16, 927.
- 50 Day, C. and Haas, H. (2009) Experimental assessment of the ITER cryosorption pump high temperature regeneration. Fusion Eng. Des., 84, 665.
- 51 Bentley, P.D. (1980) The modern cryopump. Vacuum, 30, 145-158.
- 52 Day, C., Murdoch, D., and Pearce, R. (2009) The vacuum systems of ITER. Vacuum, 83, 773-778.
- 53 Day, C. et al. (2011) Design progress for the ITER torus and neutral beam cryopumps. Fusion Eng. Des., 86, 2188-2191.
- 54 Luo, X. and Day, C. (2010) 3D Monte Carlo vacuum modeling of the neutral beam injection system of ITER. Fusion Eng. Des., 85, 1446.
- 55 Dremel, M. et al. (2009) Cryopump design for the ITER heating neutral beam injector. Nucl. Fusion, 49, S075035.
- 56 Nuss, H.E. (1988) Vacuum system for a space simulation facility. Vacuum, 38, 617-620.
- 57 Dremel, M. et al. (2005) Design of cryosorption pumps for testbeds of ITER relevant neutral beam injectors. Fusion Eng. Des., 74, 205-209.
- 58 Obert, W. et al. (1995) Performance of the supercritical helium cooling loop for the JET divertor cryopump. Adv. Cryogen. Eng., 45.
- 59 Kazachenko, O. et al. (2008) TRAP a cryopump for pumping tritium on precondensed argon. Nucl. Instrum. Methods Phys. Res. A, 587, 136-144.

- 60 Anashin, V.V. et al. (2004) Vacuum performance of a beam screen with charcoal for the LHC long straight sections. Vacuum, 72, 379-283.
- 61 Klein, H.-H. (1986) Der Einsatz von Kryopumpen in Produktionsanlagen. Vak.-Techn., 35, 203-211.
- 62 Foster, C.A. (1987) High throughput continuous cryopump. J. Vac. Sci. Technol. *A*, **5**, 2558–2562.
- 63 Hemmerich, J.L. and Küssel, E. (1990) The cryogenic diffusion pump and its implementation in a complete fusion reactor forevacuum system. J. Vac. Sci. Technol. A, 8, 141-144.

13

Total Pressure Vacuum Gauges

Dr. Karl Jousten

Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

This chapter introduces common vacuum gauges and explains their operating principle. Gas flowmeters are covered as well.

13.1 Introduction

Vacuum technology measures pressure p either directly according to its defining equation, Eq. (3.1), by measuring the force F = pA exerted to an area A or indirectly by measuring a physical quantity proportional to the pressure, for example, particle number density n, particle impingement rate $n\bar{c}$, and thermal conductivity, among others.

Measuring pressure directly by assessing the effect of a force is limited to pressures greater than approximately 1 mPa. At this pressure, the force exerted to $1~\rm cm^2$ amounts to only about $10^{-7}~\rm N$. Measuring such low forces requires an electrically amplified signal.

The pressure range measured in vacuum technology spans across 15 powers of 10. No single gauge type covers the whole range. Figure 13.1 classifies common vacuum gauges according to their physical measuring principles. Table A.19 provides an overview of the gauges common to individual pressure ranges.

Calibration of vacuum gauges is usually necessary in order to guarantee correct measurements (Chapter 15). Calibration may not be required only in cases where the measuring signal can be traced back to pressure's defining equation. In addition, any physical processes and determining quantities leading to the measuring signal must be sufficiently well known. Particularly, precise knowledge of these data is essential if the measuring signal is used as a primary standard (Chapter 15). U-tube manometers and piston manometers are considered suitable for primary standards. However, operating them is elaborate and impractical for everyday measuring tasks, particularly in an industrial environment. Because they are mostly used as primary standards, they are covered in Chapter 15.

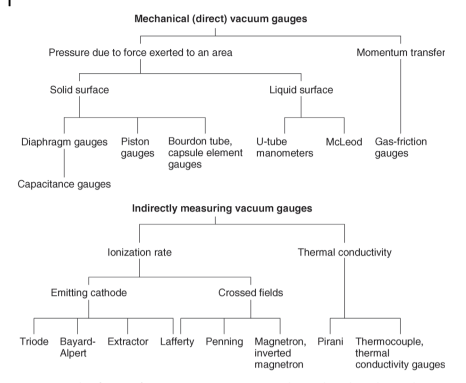


Figure 13.1 Classification of common vacuum gauges according to their physical principles. Crossed fields mean crossed electric and magnetic fields.

For gas-friction vacuum gauges, physical processes involved are thoroughly investigated and the measuring signal can be traced back directly to particle density and particle impingement rate. However, one parameter, the accommodation coefficient, cannot be given a priori, and thus, calibration is mandatory. Therefore, gas-friction vacuum gauges are covered in this chapter.

13.2 Mechanical Vacuum Gauges

13.2.1

Principle and Classification

Figure 13.2 schematically illustrates a possible principle of direct mechanical pressure measurement. The diaphragm D with surface area A separates two volumes, 1 and 2, with pressures p_1 and p_2 , respectively. The force exerted to the diaphragm toward the right side is given according to Eq. (3.1):

$$F = (p_1 - p_2)A \tag{13.1}$$

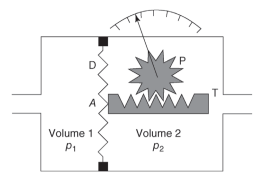


Figure 13.2 Diagram of mechanical pressure measurement using the deflection of a diaphragm.

and deflects the diaphragm. If the displacement stretch x is translated into an angular movement ϕ , the gauge directly displays the pressure difference (p_1-p_2) . For the principle in Figure 13.2, this is done using a toothed rack T and pinion P. When volume 2 is evacuated down to a reference pressure $p_2 \ll p_1$, the vacuum gauge directly registers the absolute pressure p_1 in volume 1.

Mechanical vacuum gauges operating on this principle as well as any liquid manometers (Section 15.2.1.1) generally measure a difference between two pressures. The reference pressure can be negligibly small compared with the measured pressure. The reading on a directly measuring mechanical vacuum gauge is independent of the gas species as long as the temperatures of the measuring instrument and the measured container are equal. If they are not, the reading is gas-species dependent if the measured pressure p is below the so-called viscous regime ($p < 50 \, \text{Pa}$).

Common types of mechanical, differentially measuring vacuum gauges are categorized into three groups according to the type of reference pressure and the location of the sensor.

- a) Reference pressure is atmospheric pressure. The sensor is placed on the reference-pressure side.
- b) Reference pressure is zero (i.e., below the resolution of the instrument). The sensor is placed on the measuring side, that is, at a place connected to the volume in which the pressure is measured.
- Reference pressure is zero. The sensor is placed on the reference-pressure side.

13.2.2

Corrugated-Diaphragm Vacuum Gauges

The diaphragms used here are circular, corrugated membranes. They are mounted in between two flanges, either clamped or welded at their edges. One side of the membrane is exposed to the volume whose pressure is measured. The diaphragm deflection caused by the pressure difference is used as a measure of

pressure, and is displayed by a motion work (see principle in Figure 13.2). Diaphragm pressure gauges are of the types (a) and (b) listed in Section 13.2.1.

The deflection force for diaphragms is relatively high (compared with tube springs/ Bourdon tubes; see Section 13.2.4) and their annular fixing makes them comparably insensitive to vibration. A diaphragm can be protected against overload by allowing it to lean against a safety plate or the flange on the low-pressure side. Coating provides a means of protecting the measuring instrument against corrosive gases.

13.2.3 Capsule Element Vacuum Gauges (Measuring Range 1-100 kPa)

A capsule element is made up of either two circular, corrugated diaphragms or a circular solid wall combined with a circular, corrugated diaphragm, arranged as a cell (Figure 13.3). The cell is evacuated and sealed vacuum-tight. Capsule element vacuum gauges are type (b) mechanical vacuum gauges according to Section 13.2.1.

Thus, the reading of a capsule element vacuum gauge is independent of the ambient pressure. If the test pressure drops, the distance between both walls increases according to their elastic forces or due to an integrated compression spring. This deflection represents the measured quantity, which is translated into a reading by an appropriate sensor. In the instrument shown in Figure 13.3, the measured value is transmitted by a level system. Sensor and display are arranged in the volume where the pressure is monitored. For higher actuating

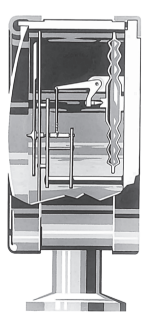


Figure 13.3 Capsule element vacuum gauge.

forces, several capsule elements can be mechanically connected in series. The main advantage of the measuring principle is that the deflection is, to a large extent, proportional to the pressure. A disadvantage is that they are destroyed by gases condensing in the measuring unit or by corrosion in the unit. End-scale deflections of 2 kPa (20 mbar), 10 kPa (100 mbar), and 100 kPa (1000 mbar) are typical in the vacuum range.

Generally, the measuring accuracy of such an instrument can be improved for a certain pressure range by filling the cell with a gas of predefined pressure. This allows measurement accuracies of, for example, 1% in the pressure range between 10 and 11 kPa. However, the disadvantage of this type of operation is that the pressure in the diaphragm cell is temperature dependent. From Eq. (13.1), it follows that a rise in temperature $\Delta T = 3$ K produces a relative change in the pressure reading of 1%.

Slightly modified, the design of capsule element vacuum gauges is used in diaphragm pressure switches as well (see Section 13.2.6).

13.2.4 Bourdon Tube Vacuum Gauges (Measuring Range 1-100 kPa)

A Bourdon tube vacuum gauge, shown in Figure 13.4, is a typical type of group (a) mechanical vacuum gauge according to Section 13.2.1. The inside of a tube (T) curved to a 270° circular arc (usually with oval cross section) is connected to the volume where the pressure is measured. If the pressure drops inside tube T, the bending radius of the arc changes because the force exerted to the (larger) outside surface is higher than the force on the inside of the arc. The ambient pressure reduces the bending radius [1,2]. A level system (L) transfers the resulting deflection directly to a pointer that indicates the reading on a scale mounted to the gauge. If the surrounding of the tube is at ambient pressure, the reading on instruments of this type depends on the surrounding atmospheric pressure (meteorological conditions, height above sea level). This error can be corrected by rotating the scale around the pointer axis. Bourdon tube vacuum gauges are robust and quite corrosion resistant.

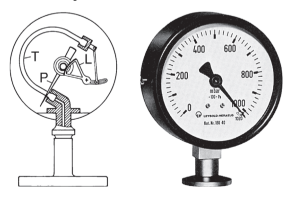


Figure 13.4 Bourdon tube vacuum gauges. Section and front view.

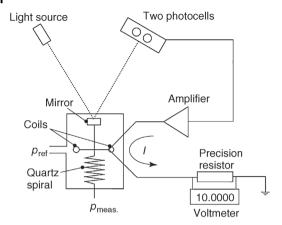


Figure 13.5 Diagram of a quartz Bourdon tube difference is measured with a beam of light. vacuum gauge by Ruska Corporation, Houston, Electrical coils hold the guartz spiral in the Texas (now GE Sensing). The deflecting force exerted on a quartz spiral due to pressure

zero position. The coil current is proportional to the pressure.

13.2.4.1 Quartz Bourdon Tube Vacuum Gauges

From a measuring technology perspective, quartz Bourdon tube vacuum gauges are a particularly ingenious variant of Bourdon tube vacuum gauges (Figures 13.5 and 13.6). They are type (a) or (c) mechanical vacuum gauges, according to Section 13.2.1.

A helically bent quartz tube is deflected similar to the Bourdon tube vacuum gauge by a pressure difference between the outside and inside. It is repositioned

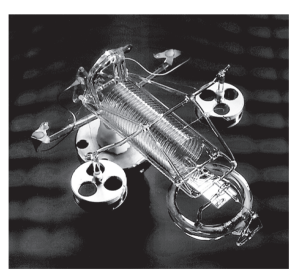


Figure 13.6 Photograph of quartz spiral and balancing coils. (Courtesy of Ruska Corporation, Houston, Texas (now GE Sensing).)

by an electromagnetic coil. The necessary coil current is a measure of the pressure difference. A beam of light optically determines the initial position. A mirror arranged at one end of the spiral reflects the beam and differential balancing of two photodiodes defines the mirror's angle. The reading is simple to linearize and the long-term stability of the instrument is very high because the spiral always returns to its initial position. Typically, readings show relative deviations $< 2 \times 10^{-4}$ within 1 year of operation and throughout a large measuring range (3–100% of end-scale deflection).

The resolution of the instrument is 10^{-5} of end-scale deflection, and reproducibility is approximately 2×10^{-5} . Instruments are available with end-scale deflections between 7 and 100 kPa.

13.2.5

Diaphragm (Membrane) Vacuum Gauges

Elastic diaphragms deflected reversibly by a pressure difference between both sides of the diaphragm have been used for pressure measurements in vacuum since 1929 [3]. Approaches of measuring diaphragm deflection are manifold and covered in the following sections.

13.2.5.1 Diaphragm (Membrane) Vacuum Gauges with Mechanical Displays (Measuring Range 0.1-100 kPa)

If we evacuate volume 2 of the system shown in Figure 13.2 down to the pressure p = 0 and subsequently seal it, we obtain a diaphragm vacuum gauge with a reading that is temperature independent (except for temperature-dependent mechanical properties) and independent of atmospheric pressure. Sensor and display are not exposed to the measured gas. Thus, the instrument is a type (c) mechanical vacuum gauge (Section 13.2.1). The system is relatively corrosion resistant because sensitive parts are not exposed to the measured gas. Only the corrugated membrane made from a copper-beryllium alloy might require corrosion protection, for example, by gold plating, in particularly corrosive environments.

As indicated in Section 13.2.3, deflection of a corrugated diaphragm is largely proportional to the pressure difference. Thus, the scale in such an instrument is linear in the case of a proportional conversion of deflection into reading. However, in many cases it is convenient to use a trick and dilate the reading in the low-pressure range (see Figure 13.8): if the pressures are equal on both sides of the diaphragm, that is, at low pressures, the entire diaphragm surface area is ready to accept the pressure. As the pressure rises into the range of 1–1.5 kPa, the first fold in the membrane comes to rest at the base plate that is manufactured with a corresponding contour (Figure 13.7). The surface area of the membrane thus decreases, the membrane's stiffness increases, and sensitivity drops. In the range 5-6 kPa, the next fold touches the base plate and the membrane's surface area decreases further. This process action repeats a third time in the

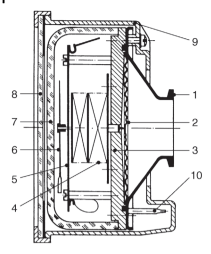


Figure 13.7 Section of a diaphragm vacuum gauge with mechanical display (diagram). (1) Connecting flange and gas inlet, (2) diaphragm, (3) base plate, (4) transfer system for

diaphragm deflection, (5) indicating disk, (6) index hand, (7) vacuum-tight glass cap, (8) front plate, part of housing, and (9, 10) evacuating ports.

range 15–20 kPa. The described trick yields a scale dilated across a wide pressure range.

13.2.5.2 Diaphragm (Membrane) Vacuum Gauges with Electrical Converters

Diaphragm vacuum gauges of the described type are suitable for incorporating electrical sensors as well. Figure 13.9 shows three examples of electrical signal generation. Sensors using wire resistance strain gauges are outdated; however, they are included in the figure to illustrate the measuring concept.



Figure 13.8 Front plate of a standard diaphragm vacuum gauge with expanded scale.

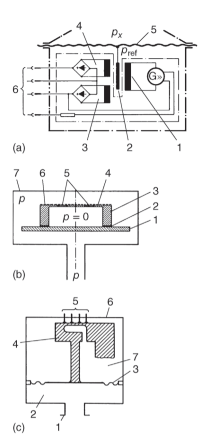


Figure 13.9 (a) Diaphragm vacuum gauge for remote display with inductive displacement transducer. p_{ref} : reference pressure; p_x : measured pressure. (1, 3, 4) Differential transformer, (2) ferromagnetic immersion rod, (5) diaphragm, and (6) electrical connectors. (b) Diaphragm vacuum for remote display with piezoresistive sensor. (1) Silicon base plate, (2) vacuum-tight Cu–Si joint, (3) n-silicon cap, (4)

diaphragm, (5) resistance bridge of p-silicon integrated into diaphragm by diffusion, with wire connection, (6) flexible protective layer, and (7) housing. (c) Diaphragm vacuum gauge with wire resistance strain gauges. (1) Connecting flange, (2) measuring volume, (3) diaphragm, (4) deflection rod, (5) wire resistance strain gauges, (6) housing, and (7) volume with reference pressure p = 0.

A second principle uses a *capacitive* displacement sensor. It is widespread and thus covered separately in Section 13.2.5.5. It provides an electrical signal, which can be electronically linearized and teletransmitted.

The *inductive* displacement sensor in Figure 13.9a uses a ferromagnetic pin extending more or less far into a differential transformer coil. The produced signal is proportional to the deflection of the diaphragm. Such sensors are available down to an end-scale deflection of 25 Pa but measurement accuracy due to temperature variations is limited to 1 Pa. Inductive sensors show low hysteresis and high reproducibility, and are relatively insensitive against mechanical disturbance

(vibration). They are available with electrical current (4-20 mA) as well as voltage outputs (up to 10 V).

The design in Figure 13.9c uses a deflecting rod (4) to pick up the diaphragm deflection. Wire resistance strain gauges in a bridge circuit are applied to the rod providing the electrical output signals. The arrangement allows accurate pressure measurements up to 200 kPa.

Diaphragm (Membrane) Vacuum Gauges Using the Piezoresistive Principle

The piezoresistive effect (Figure 13.9b), due to which a pressure change causes a change in electrical resistance, is being used increasingly for pressure measurements, even under vacuum. In metals, the change in resistance is determined by the geometrical change of the conductor (cross section, length). Semiconductors additionally show a change in the specific resistance of the material, thereby increasing the piezoresistive effect. Thus, semiconductor materials are preferred for pressure measurements relying on this principle.

Crystalline silicon has extraordinary elastic properties. It can be loaded nearly to the fracture limit, shows hardly any hysteresis, and is very stable. Silicon is thoroughly investigated due to its applications in microelectronics, and processing including doping, thin-film coating, and etching is well under control. Thus, it represents the material of choice.

For measuring resistance changes, doped low-impedance conductor patterns in radial and tangential directions are applied to one side of the circular diaphragm (Figure 13.10). The thin silicon disk is etched down to a thin diaphragm from the other side. The resistors are connected in series and are part of a resistance bridge adjusted to p < 0.1 kPa. Changes in gas pressure deform the silicon diaphragm and the following resistance change detunes the bridge. The electronically linearized signal is proportional to the absolute pressure and

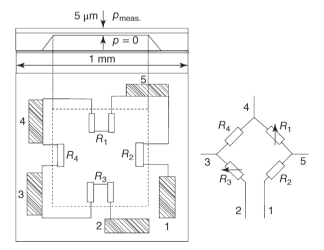


Figure 13.10 Piezoresistive vacuum gauge.

independent of the ambient atmospheric pressure as well as of the gas species. The measuring head has a very small measuring volume of only 1 cm³. The measuring range is 0.1–100 kPa. Integrated circuit technology allows one to integrate amplifiers directly into the sensor element. As in any diaphragm vacuum gauge, absolute pressure is measured if the reference side is evacuated down to below the resolution limit of the measuring instrument, and differential pressure measurement is obtained if the reference side is exposed to any desired pressure value.

However, compensation of the strong temperature dependence of semiconductor resistors is required (see Ref. [4]).

13.2.5.4 Piezoelectric Vacuum Gauges

If a force is applied to a quartz crystal, the piezoelectric effect produces charges on the crystal surface that are conducted by electrodes and measured with suitable instruments (Figure 13.11). A pressure exerting forces from all sides does not create any charges. Therefore, a piezoelectric pressure sensor uses a diaphragm that transforms the pressure into a force related to the elastic surface of the diaphragm. The force is measured by employing a quartz crystal rod. An electrode conducts the produced charges to a connector plug from where highly isolated connecting wires transfer the charges to the input of a charge amplifier that transforms the charge into a voltage.

13.2.5.5 Resonant Diaphragm Vacuum Gauges

The measuring principle of a resonant diaphragm vacuum gauge [5] is based on the frequency change of a resonator due to changes in a solid. The strain is changed by pressure changes across a diaphragm. Figure 13.12 shows the basic setup of such an instrument, manufactured by micromachining. Two H-shaped resonators are placed onto a thin-etched diaphragm: one in the middle and the

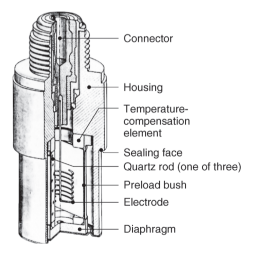


Figure 13.11 Piezoelectric pressure sensor.

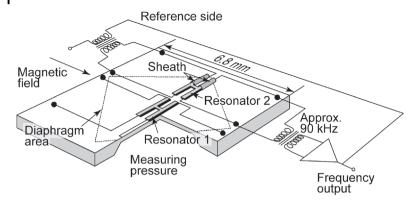


Figure 13.12 Diagram of the sensor chip in a resonant diaphragm vacuum gauge [5]. Image includes electrical coupling for resonator 2.

other at the edge of the diaphragm. Diaphragm and resonators are made of monocrystalline silicon with excellent elastic properties. The resonators are 30 μm wide, 500 μm long, 5 μm high, and placed in a specially manufactured vacuum housing with walls made of highly p-doped silicon, just as the resonators.

The two resonators are not exactly equal in size. Thus, their natural frequencies differ slightly and they are excited by self-oscillation. When pressure is applied, resonator 2 at the edge of the diaphragm reduces its resonance frequency f_2 (approximately 90 kHz) and f_1 of resonator 1 in the middle rises. The difference (f_1-f_2) is a measure of the differential pressure. The sum (f_1+f_2) is a measure of the total line pressure to both sides of the diaphragm. Isolation of the resonators and protection against the surrounding pressure by means of the vacuum housing are extremely important for the accuracy of the sensor because they prevent an impact of ambient pressure on resonator quality (Q factor) and the dependent resonance frequency. The deflection of the diaphragm alone determines the frequency change. If the volume above the diaphragm is evacuated down to pressures below the resolution limit, the sensor can be used as an absolute pressure gauge.

Additional temperature sensors placed on the chip compensate for null drifts and sensitivity changes caused by temperature variations. The frequency change is approximately 20% for a differential pressure of $100\,\mathrm{kPa}$. In this example, the relative strain change in the silicon crystal is 1×10^{-4} .

For the highest accuracy (approximately 1×10^{-4}), the measuring range of such sensors is 100 Pa to 100 kPa. Long-term and transport stability of these instruments has proven to be very high as well (3×10^{-5} at 1 kPa) [6]. Unfortunately, these gauges are presently (2015) out of the market.

13.2.5.6 Capacitance Diaphragm Vacuum Gauges

In capacitive pressure measurement, the measuring diaphragm, deflecting due to pressure, forms one electrode of a capacitor. The change in capacitance of this capacitor caused by the pressure difference from one diaphragm side to the other is measured. The systems are referred to as capacitance diaphragm gauges (CDGs) because the measuring signal relies on a capacitance measurement. The high sensitivity of this pressure measuring technique allows measurements of very small pressure differences, and thus absolute pressures, if the pressure on the reference side is below the resolution limit of the instrument. The resolution limit of these instruments is approximately 1 mPa.

Starting in 1949, development of capacitance pressure sensors went through several stages [7] leading to today's designs that show high measuring accuracy, long-term stability, and overload safety. The pressure sensors attached to the instruments are robust and compact. Capacitance vacuum gauges are used in industrial vacuum systems, particularly semiconductor industry, as well as for reference standards (Chapter 15) for calibration services.

A capacitance vacuum gauge is made up of a transducer and the electronics for processing and displaying the pressure-dependent signal. The latter can be either fully integrated into the transducer or placed inside a separate unit. For separately arranged electronics, the transducer includes a pre-amplifier.

A large variety of transducer designs are commercially available for different applications and measuring ranges. Figure 13.13 shows a schematic illustration of a capacitance vacuum gauge.

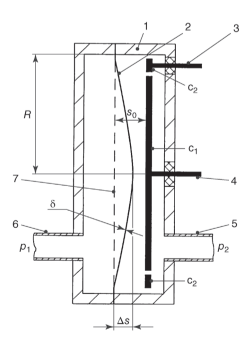


Figure 13.13 Diagram of a capacitance diaphragm vacuum gauge with Invar diaphragm. (1) Housing, (2) diaphragm, (3) leadthrough to capacitor ring c2, (4) leadthrough to capacitor ring c_1 , (5) gas inlet (reference pressure p_2), (6)

connecting port to vacuum vessel with measured pressure p_1 , and (7) diaphragm in zero position ($p_1 = p_2$). δ : diaphragm thickness; Δs : deflection of diaphragm; so: distance between diaphragm and capacitor plate c₁.

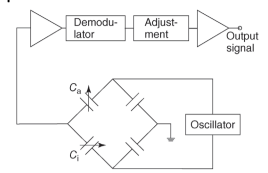


Figure 13.14 Measuring setup of a capacitance diaphragm vacuum gauge by a capacitor bridge as done in gauges of the MKS Company (Figure 13.13). C_i is the capacitance

between the diaphragm and the inner electrode, and C_a the capacitance between the diaphragm and the outer annular electrode. (Courtesy of MKS Instruments GmbH, Munich.)

The diaphragm (2: deflected; 7: zero position for equal pressure on both sides of the diaphragm) exposed to the measured pressure is a circular disk made of a material with a low coefficient of thermal expansion, for example, Invar or ceramic. Thickness is determined by the desired end-scale deflection and can be as low as $25 \, \mu m$. It is fixed by welding, cementing, or brazing. For improved zero-point stability, the membrane forms two capacitors with the circular electrode c_1 and the annular electrode c_2 . The difference in capacitance of the two capacitors is used as a measuring signal.

Figure 13.14 shows a possible measuring chain. Both capacitors are part of a measuring bridge made up of four capacitors. The oscillator sends a stationary signal (commonly used frequency 10 kHz, sometimes 85 kHz) to the pressure sensor (transducer) that is made up of the two capacitors with the capacitances C_i and C_a . The measuring bridge changes the amplitude and phase of the oscillator signal. The amplitude is linear with the pressure. For negative pressures ($p_2 > p_1$ in Figure 13.13), the phase position is shifted by 180° compared with positive pressure ($p_2 < p_1$). Demodulation with the oscillator signal produces a DC voltage signal, which is linearized and amplified.

Another method of measuring the capacity is the use of a capacitance to digital converter (Figure 13.15). A square wave voltage excitation source (voltage V) is connected to the capacitance C_x to be measured and injects a charge Q according to $Q = C_x V$. This charge is fed into a charge balancing modulator. As the integrator tries to offset the introduced charge, the following comparator compares at high frequency if the voltage is higher or lower than ground and sends a binary signal that is filtered for noise. Gain coefficients are applied.

In common pressure sensors of this type, the counter electrodes are deposited as metallic layers onto a common ceramic disk (forsterite). The housing (1), Figure 13.13, is equipped with two small gas inlet tubes (5 and 6) that receive appropriate vacuum-tight ports (screw joints, small flanges) for connecting to the apparatus. Such pressure sensors have typical inner volumes of

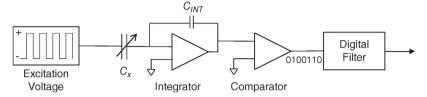


Figure 13.15 Measuring setup of a capacitance diaphragm vacuum gauge by a capacitance to digital converter as done in gauges of the *INFICON AG* (Figure 13.16). (Courtesy of *INFICON AG*, Liechtenstein.)

approximately 2.5 cm³ (without the inlet tubes). The bulged volume of the diaphragm amounts to be typically 0.18 cm³ [8].

The pressure sensors are capable of monitoring differential pressures $\Delta p = p_1 - p_2$, but also absolute pressures if the pressure on the reference side (right-hand side in Figure 13.13) is below the resolution limit of the instrument. Most commercially available pressure sensors feature today a pre-evacuated and sealed reference side. Getter material placed in the volume on the reference side guarantees long-term low pressure. Figure 13.16 shows such a capacitance vacuum gauge with a ceramic diaphragm.

The measuring range of the pressure sensor is varied by using differently thick diaphragms, keeping the dimensions of the sensor constant. The maximum possible pressure, the pressure at end-scale deflection, characterizes the range. Commercially available sensors have end-scale deflections of 13.3 Pa, 133 Pa, 1.3 kPa, 13 kPa, and 133 kPa; they display the measured values using 312 or 512 digits. If the measured pressure exceeds the maximum pressure (by approximately 20%), the deflected diaphragm touches the opposite capacitor electrode, thereby

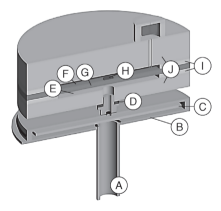


Figure 13.16 Absolute-pressure capacitance diaphragm vacuum gauge with ceramic diaphragm. (A) Connection to vacuum system, (B) protecting chamber with (C) plasma shield, (D) metal/ceramic joint (*Vacon*), (E) measuring

chamber, (F) reference chamber, (G) Al₂O₃ ceramic diaphragm, (H) gold electrode, (I) glass/ceramic joint, and (J) Al₂O₃ ceramic housing. (Courtesy of *INFICON AG*, Liechtenstein.)

preventing damage to the diaphragm. In certain types of instruments, however, such an overload may invalidate previous calibration.

Very sensitive capacitance pressure sensors carrying diaphragms with a thickness of only about 0.025 mm are capable of measuring differential and absolute pressures of approximately 1 mPa. Here, deflection of the diaphragm is just about 0.4 nm (slightly more than an atomic diameter!). The capacitance change caused by this deflection is approximately 10^{-4} pF. End-scale deflection of the diaphragm is several um.

When a capacitance vacuum gauge is calibrated to differential pressure, that is, reference pressure $p_{\rm ref} \neq 0$, and the same calibration is to be used at a different reference pressure, for example, $p_{ref} = 0$, then the changing pressure difference exerting force from the outside to the inside may change the state of stresses in the diaphragm. Thus, calibration done at the initial reference pressure may be nontransferable. Therefore, certain designs include a guard volume around the diaphragm fixing, where the pressure is always equal to the pressure on the reference side so that external stresses are not introduced into the diaphragm. However, it should be noted for $p_{ref} \neq 0$ that the dielectric constant of the considered gas does influence the capacitance of the sensor on the reference side [9].

13.2.5.7 Thermal Transpiration

Thermal transpiration occurs when parts of a vacuum system are at different temperature than others and when these regions are separated by small conductances with *Knudsen* numbers Kn > 0.5 (Figure 13.17). For example, thermal transpiration is observed in capacitance vacuum gauges that are held at a constant temperature T_2 for improved zero-point stability, but with T_2 being above the temperature of the remaining vacuum system $T_1 \approx T_{amb}$. Because of their high sensitivity, the effect is particularly important in capacitance vacuum gauges. The physics of thermal transpiration are covered in Section 4.6.

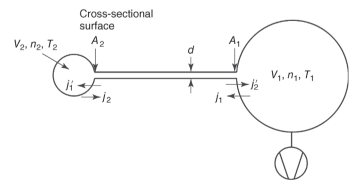


Figure 13.17 Diagram of thermal transpiration. Two vessels at unequal temperatures are connected via a tube with low conductance where the *Knudsen* number Kn > 0.5.

From Eq. (4.193), it follows that

$$\frac{n_2}{n_1} = \sqrt{\frac{T_1}{T_2}}, \quad Kn \gg 0.5,$$
 (13.2)

and because p = nkT,

$$\frac{p_2}{p_1} = \sqrt{\frac{T_2}{T_1}}. (13.3)$$

See also Eq. (4.194). For $Kn \gg 0.5$, that is, under molecular flow conditions, Eq. (13.3) indicates that the part of a vacuum system with the higher temperature is in fact at higher pressure. For a heated capacitance vacuum gauge and molecular flow, this means that the pressure reading is higher than the actual pressure in the recipient (Figures 4.52 and 15.21). For viscous flow $(Kn \ll 0.5,$ in practice p > 100 Pa), the pressure in the instrument is equal to the pressure in the recipient $(p_2/p_1 = 1)$.

A number of empirical formulas characterize the transition between these two ranges. Most commonly used is an equation by Takaishi and Sensui [10], derived from an equation first published by Liang [11]. Takaishi and Sensui's equation reads

$$\frac{p_2}{p_1} = \frac{AX^2 + BX + C\sqrt{X} + \sqrt{T_2/T_1}}{AX^2 + BX + C\sqrt{X} + 1},$$

$$X = 0.133p_2d, \quad [p_2] = \text{Pa}, \quad [d] = \text{m},$$

$$A = A^*T_{\text{m}}^{-2},$$

$$B = B^*T_{\text{m}}^{-1},$$

$$C = C^*T_{\text{m}}^{-0.5},$$

$$T_{\text{m}} = (T_1 + T_2)/2,$$
(13.4)

d is the diameter of the tube connecting T_2 and T_1 . For selected gases, Table 13.1 lists the constants A^* , B^* , and C^* found by *Takaishi* and *Sensui*.

Setina [13] provided another description of the transition range, which uses the Knudsen number and thus is applicable to any gas species (see Eq. (4.195) in Section 4.6).

Jitschin and Röhl [14] discovered that precise measurements reveal small systematical deviations between the measured curves and the Takaishi-Sensui equation. They concluded a material-dependent molecule-surface interaction, described by the constants A^* , B^* , and C^* . However, instead of recalculating these parameters, they suggested using adopted values for d_{eff} and $T_{2,\text{eff}}$ to yield a more accurate description of the transition range with Eq. (13.4). However, these effective values deviate considerably from the real physical quantities. Thus, the values obtained by calibrating with a certain gas species should not be used for calculating the transition range of any other gas species. For this, it is more advisable to use the method suggested by Poulter [12] or Eq. (4.195) by Setina [13].

Gas species	A*(Pa² m²)	<i>B</i> *(Pa m)	$C^*(Pa^{1/2} m^{1/2})$
H_2	1.24×10^{5}	8×10^{2}	10.6
Не	1.5×10^{5}	1.15×10^{2}	19
CH ₄	1.45×10^{5}	1.5×10^{3}	13
Ne	2.65×10^{5}	1.88×10^{2}	30
N_2	1.2×10^{6}	1×10^3	14
O_2	8×10^{5}	1.75×10^{3}	_
Ar	1.08×10^{6}	8.08×10^{2}	15.6
Kr	1.45×10^{6}	1.5×10^{3}	13.7
Xe	3.5×10^{6}	4.14×10^{3}	10
SF_6	1.53×10^{7}	2.42×10^4	04.4

Table 13.1 Values for the constants A^* , B^* , and C^* in Eq. (13.4).

Taken from Refs. [10,12].

Figure 15.21 shows a calibration curve for helium and nitrogen of a capacitance vacuum gauge.

13.2.6 Pressure Switches and Pressure Controllers

Often, when operating a vacuum system, certain processes such as opening or shutting valves, starting or stopping pumps, signal triggering, and starting or stopping heaters need to be activated at certain pressure levels. This is where socalled pressure switches are utilized, actuating the switching operation as soon as the pressure reaches the corresponding threshold value (switching pressure). Pressure controllers limit the process pressure in a vacuum process so that it remains below a certain predefined pressure value.

The designs used in vacuum technology mostly follow the designs of mechanical vacuum gauges. Thus, this section will focus only on the most common diaphragm pressure switches and diaphragm pressure controllers.

A diaphragm pressure switch (Figure 13.18) is made up of a thin-walled diaphragm (2) separating the switch volume into a measuring volume (8) and a reference volume (3). The latter holds a contacting pin (4) with an isolated leadthrough (5) to the environment. The diaphragm connected to earth provides the counter contact. A small, integrated valve (1) is used to adjust the desired switching point. The open valve allows intake of gas with the desired (reference) pressure p_0 via 7. Then, pressures in 3 and 8 are equal and the diaphragm relaxes. Now, the diaphragm touches the contacting pin (4). In order to guarantee the contact and to compensate for maximum manufacturing tolerances of several hundredths of a millimeter, the diaphragm is pressed slightly and elastically against the contact (4) using an adjusting screw (9). The principle of the relaxed diaphragm yields long service life and high switching accuracy of ± 10 Pa.

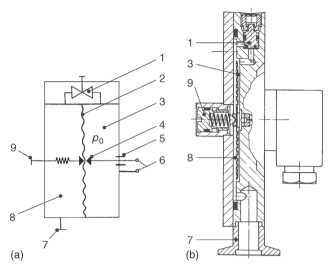


Figure 13.18 Diaphragm pressure switch: (a) principle; (b) section. (1) Bypass valve, (2) diaphragm, (3) reference volume, (4) contacting pin, (5) electrical leadthrough, (6) electrical connectors, (7) connecting flange, (8) measuring chamber, and (9) adjusting screw.

When the pressure in the measuring chamber falls short of the set reference value p_0 by more than 10 Pa, the contact opens and actuates the relay in the connected switching amplifier. The amplifier circuit is usually connected to a preceding time-delay circuit causing a time lag of approximately 0.5 s. This prevents relay flutter in the output. For safety reasons, the cutoff cycle is usually not delayed.

The diaphragm pressure controller in Figure 13.19 is a further development of the previously described diaphragm pressure switch. It includes a throttle port in the controlled chamber. Adjusting reference pressure p_0 is analogous to the procedure described for the diaphragm pressure switch. If pressures are equal in both chambers 3 and 7, the throttle port touches the diaphragm (2). If the pressure in the recipient rises due to gas load and valve 8 is open, the diaphragm lifts off from the throttle port and the pumping line is opened via the open valve 9. Now, the pump is connected to the recipient and reduces the pressure here. As soon as the predefined process pressure p_0 is obtained, the diaphragm again touches the throttle port. The pumping line is shut and thus prevents further pressure drop in the recipient.

Analogous to the diaphragm vacuum gauge, the control range (range of adjustable reference pressures p_0) for the diaphragm pressure switch is 1-100 kPa. The reaction time is a few milliseconds and control accuracy is a few percent of the adjusted reference pressure. Using the valve sizes given in Figure 13.19 for valves 8 and 9 (DN 16), the maximum gas flow is 16 m³ h⁻¹ under standard conditions ($T_{\rm m}$, $p_{\rm m}$).

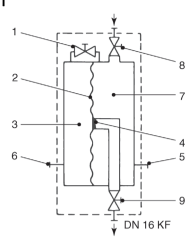


Figure 13.19 Principle of diaphragm pressure controllers. (1) Bypass valve, (2) diaphragm, (3) reference volume (reference pressure p_0), (4) throttling port, (5) process pressure measuring

port, (6) reference pressure measuring port, (7) control chamber, (8) recipient valve, and (9) valve to vacuum pump.

13.3 Spinning Rotor Gauges (Gas-Friction Vacuum Gauges)

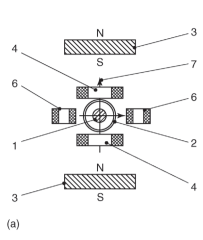
Section 3.3.2 explains the proportionality between the frictional force of a gas in between a moving and a stationary wall, and gas density n or pressure p, for low pressures. More precisely, the frictional force is proportional to the impingement rate $n\bar{c}/4$ of gas particles. Here, low pressure means that the mean free path \bar{l} of gas particles is greater than the distance d between the walls. If this effect is to be used in a vacuum gauge where wall distances of 1 cm are realistic, Table A.9 shows that the pressure must be below 1 Pa.

Meyer [15], Maxwell [16], Kundt and Warburg [17], Sutherland [18], Hogg [19], and Knudsen tried to measure gas friction with disks suspended on thin glass torsion fibers. A disk of the same geometry rotating below accelerates the gas and turns the disk on the fiber until the moment of torsion balances the moment of friction. The torsion angle then is a measure of the pressure. However, in these early experiments, the friction in the disk suspension was too high. Only after 1937, when magnetic suspension allowed frictionless mounting [20], the effect of gas friction was successfully utilized for gas pressure measurements. The first instruments by Beams [21,22] were hard to handle. Later, Fremerey [23–26] developed a convincing vacuum gauge.

13.3.1

Measuring Setup and Measuring Principle

Figure 13.20 shows a schematic illustration of the most important components in a spinning rotor gauge and a section through the measuring head. In a tube



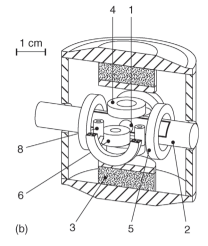


Figure 13.20 (a) Diagram and (b) section of the measuring head in a spinning rotor gauge. (1) Rotor, (2) vacuum tube, (3) permanent magnets, (4) two coils for stabilizing vertically,

(5) four driving coils, (6) two signal sensing coils, (7) spin axis, and (8) four coils for stabilizing horizontally.

(2, length 60 mm, inside diameter 7.5 mm) flange-mounted to the vacuum vessel, a steel (preferably stainless steel) sphere (1, diameter 4–4.76 mm) is freely suspended magnetically. The magnetic field of the permanent magnets (3) compensates for the main part of the sphere's weight. The superimposed magnetic field of the coils (4) holds the sphere in place.

Two coils (4) hold the sphere vertically; four coils (8) stabilize it horizontally [26]. Four drive coils (5) create a rotating field in the horizontal plane, cycling at a frequency ν_0 of approximately 415 Hz, that accelerates the sphere around the vertical axis (7) to its initial rotary frequency ν_0 . After this frequency is obtained, the rotary field is shut off. The rotary frequency of the sphere thus drops according to the retarding moments caused, among other effects, by impinging gas molecules. The sphere's relative frequency gradient $\Delta\nu/\nu$ is the measured signal (DCR signal, deceleration rate). It is picked up by two sensing coils (6) in the horizontal plane. The azimuthally inhomogeneous magnetic field of the sphere, characterized by a magnetic moment, creates an AC voltage with the rotary frequency of the sphere in the sensing coils (6). Retarding effects on the sphere, independent of the number of impinging gas particles, include the current induced in the sensing coils, eddy currents produced by the magnetic moment of the sphere (particularly in vacuum tube 2), and eddy currents produced in the sphere itself by nonhomogeneities of the magnetic field. This pressure-independent offset (residual drag) must be subtracted from the measured signal.

13.3.2

Retarding Effect due to Gas Friction

According to Eq. (3.48), a gas of pressure p = nkT (n: particle number density) in tube 2 (Figure 13.20) produces a particle flow onto each surface area element dA of the sphere of

$$\mathrm{d}q_N = \frac{n\overline{c}}{4} \; \mathrm{d}A.$$

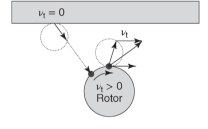
During interaction with the surface, each particle subject to short-term adsorption receives a linear momentum (Figure 13.21)

$$P = \sigma_t m \nu_t, \tag{13.5}$$

with $\nu_{\rm t}$ denoting the tangential velocity of the considered sphere-surface element dA, and $\sigma_{\rm t}$ denoting the tangential-momentum accommodation coefficient (see Section 3.3.2).

According to the law of conservation of momentum, an equally large, oppositely directed, and thus retarding momentum is transferred to the sphere. Therefore, the surface element dA of the sphere is subject to the retarding force $\mathrm{d}F = P\,\mathrm{d}N/\mathrm{d}t = P\,\mathrm{d}q_N,$

$$dF = \sigma_t m \nu_t \frac{n\overline{c}}{4} dA. \tag{13.6}$$



Wall

Figure 13.21 Principle of a spinning rotor gauge. Observed in the static wall system, gas particles leaving the wall show a random distribution following the $\cos \theta$ law. After leaving the rotor and due to accommodation, they

feature a privileged velocity direction parallel to the tangential velocity v_t of the rotor. Thus, when the particles impinge on the rotor, they transfer a linear momentum in the direction opposite to the rotor.

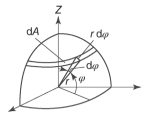


Figure 13.22 Geometry of a sphere for calculating the retarding force.

The higher the surface velocity ν_t or rotary frequency of the sphere, the higher the retarding force. The sphere zone between φ and $\phi + d\phi$ (φ : azimuthal angle, latitude of the sphere, that is, $\phi = \pm 90^{\circ}$ pole) has the surface area (Figure 13.22)

$$dA = 2\pi r \cos \phi \, r \, d\phi \tag{13.7}$$

and the velocity

$$\nu_{t} = \omega r \cos \phi \tag{13.8}$$

if r is the sphere's radius and $\omega = 2\pi\nu$ is the angular frequency.

Putting this into Eq. (13.6) yields the retarding force for the azimuthal wedge between φ and $\phi + d\phi$:

$$dF = \sigma_t \frac{n\overline{c}}{4} m\omega r \cos \phi r d\phi$$
 (13.9)

and the retarding angular momentum

$$dM = -r\cos\phi \,dF. \tag{13.10}$$

With $n\overline{c}m = 8p/\pi \cdot \overline{c}$ and the sphere's volume $V = 4\pi r^3/3$, integration of dM along the sphere's surface yields the retarding angular momentum induced by the gas:

$$M_{\rm gas} = -\frac{4\sigma_{\rm t}}{\pi \bar{c}} r V \omega p. \tag{13.11}$$

Equation (13.11) presumes that the particles hitting the sphere show an undisturbed velocity distribution (required for Eq. (3.48) to be valid). If the pressure in tube 2 is sufficiently low, particles having received an additional momentum in the rotating direction of the sphere when impinging the sphere hit the tube wall, accommodate here (compare Section 4.4.1), and are reemitted with a distribution corresponding to wall temperature $T_{\rm w}$. Thus, the mean velocity in Eq. (13.11) is

$$\overline{c} = \sqrt{8kT_{\rm w}/\pi \cdot m}.$$

This process repeats in the described way as long as the mean free path λ of the particles is higher than the dimensions of the container, that is, approximately $\lambda > 7.5$ mm. Table A.9 gives values for $\lambda p \approx 7.5$ mmPa as reference. Thus, Eq. (13.11) applies to pressures of up to approximately 1 Pa.

Now, the deceleration of the sphere due to gas friction is calculated from

$$\frac{\mathrm{d}}{\mathrm{d}t}(\Theta\omega) = \Theta\frac{\mathrm{d}\omega}{\mathrm{d}t} + \omega\frac{\mathrm{d}\Theta}{\mathrm{d}t} = M_{\mathrm{gas}} + M_{\mathrm{rest}},\tag{13.12}$$

with Θ being the moment of inertia of the sphere. The second term $\omega d\Theta/dt$ describes the change in moment of inertia of the sphere with respect to time and can be relevant when the temperature changes. However, generally, this term is small and will be neglected for the time being. When we neglect the moment of the pressure-independent residual retarding effect M_{rest} as well and use $\Theta = \frac{2}{5} m_{\text{rotor}} r^2$, Eqs. (13.11) and (13.12) yield the relative frequency deceleration with time:

$$-\frac{\dot{\omega}}{\omega} = \frac{10}{\pi} \cdot \frac{\sigma_{\rm t}}{r\rho} \cdot p\overline{c},\tag{13.13}$$

with $\rho = m_{\rm rotor}/V$ denoting the density of the sphere. This means that the frequency gradient with respect to time $-\dot{\omega}$ of the sphere is higher when the sphere rotates faster. However, the relative frequency gradient with respect to time (relative retarding rate) $-\dot{\omega}/\omega$ is constant if the pressure is constant. Thus, pressure p can be calculated from the measured relative retarding rate $-\dot{\omega}/\omega$ when Eq. (13.13) is rewritten:

$$p = \frac{\pi \bar{c} r \rho}{10 \sigma_{\rm t}} \cdot \left(-\frac{\dot{\omega}}{\omega} \right). \tag{13.14}$$

As experiments show, the retarding moment M_{rest} caused by the energy consumption in eddy currents is largely proportional to ω as well,

$$M_{\rm rest} \approx {\rm constant} \cdot \omega.$$
 (13.15)

Thus, if M_{rest} is not neglected in Eq. (13.12), we find

$$p = \frac{\pi \bar{c} r \rho}{10 \sigma_{\rm t}} \left(-\frac{\dot{\omega}}{\omega} - \text{RD} \right), \tag{13.16}$$

with RD denoting the residual drag, RD = constant/ Θ . However, it should be emphasized that Eq. (13.15) is an approximation and that quite often RD depends on the rotor frequency: $RD = RD(\omega)$. Usually, RD drops with the frequency $\omega/(2\pi)$; very rarely, an increase of RD with $\omega/(2\pi)$ is observed.

If the temperature rises, the rotor expands, and its moment of inertia increases, that is, it decelerates additionally, whereas it accelerates when the temperature drops (pirouette effect). Weller investigated the quantitative effect of a continuous temperature change dT_{rotor}/dt of the rotor on the measured signal [27]. As an approximation, it follows for constant pressure and constant RD that

$$-\frac{\dot{\omega}}{\omega} = 2\alpha \frac{\mathrm{d}T_{\mathrm{rotor}}}{\mathrm{d}t} \tag{13.17}$$

where α denotes the coefficient of thermal expansion of the rotor material. A rise in temperature of the rotor is observed mainly during initial acceleration of the sphere.

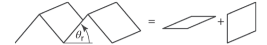


Figure 13.23 Simulation of rotor surface roughness according to Fremerey [28]. Surfaces at an angle θ_r are separated into surface elements oriented tangentially and normally to the direction of motion.

Subsequently, temperature drops. This is why accurate pressure measurements should be performed several hours after the initial acceleration of the sphere. Thus, without the approximations leading from Eq. (13.12) to Eq. (13.13), we find

$$p = \frac{\pi \bar{c} r \rho}{10\sigma_{\rm t}} \left(-\frac{\dot{\omega}}{\omega} - \text{RD}(\omega) - 2\alpha \frac{\text{d}T_{\rm rotor}}{\text{d}t} \right), \tag{13.18}$$

replacing Eqs. (13.14) and (13.16).

The tangential-momentum accommodation coefficient σ_t at the surface of technically smooth spheres from ball bearings is approximately 1 and slightly gas-type dependent. However, experiments have revealed that the roughness of the spheres is of considerable impact.

Fremerey [26] introduced a simple model, which takes into account the roughness of the sphere's surface by using plane surface-area elements that are skewed by a single angle θ_r against the geometrical surface (Figure 13.23). Each of these surface elements is allocated a fraction f_n , interacting with the gas particles in the direction of motion of the surface area element, that is, normally, and a fraction $f_t = 1 - f_n$ interacting with the gas particles perpendicular to the direction of motion of the surface element, that is, tangentially. For the tangential interaction, $\sigma_t(0-1)$ characterizes the degree of transfer of the tangential momentum. For the normal interaction, $\beta(0-1)$ describes the degree of transfer of kinetic energy. Figure 13.24 shows the dependence of the effective accommodation coefficient σ_{eff} on the roughness parameter θ_{r} for extreme values of σ_{t} and β . $\sigma_{\rm eff} > 1$ for higher roughness; however, $\sigma_{\rm eff}$ remains $\leq 4/\pi = 1.273$. In fact, experiments could not reveal any higher values. Normal technical surfaces are in the range $\sigma_t = 1.0$, $0 \le q\beta \le 1$. Very clean, polished surfaces yield $\sigma_t \le 1$. Experiments rarely find values of $\sigma_{\rm eff}$ < 1 (Figure 13.26).

Thus, σ_t in Eq. (13.18) must be replaced by σ_{eff} in order to take into account all influences. We find

$$p = \frac{\pi \bar{c} r \rho}{10 \sigma_{\text{eff}}} \left(-\frac{\dot{\omega}}{\omega} - \text{RD}(\omega) - 2\alpha \frac{\text{d} T_{\text{rotor}}}{\text{d} t} \right). \tag{13.19}$$

The mean velocity \bar{c} of the gas particles in a pure gas depends on $\sqrt{m_a}$. In a gas mixture of *n* constituents with known volume percentages $\chi_i = V_i/V$ and

$$\sum_{i=1}^{n} \chi_i = 1,$$

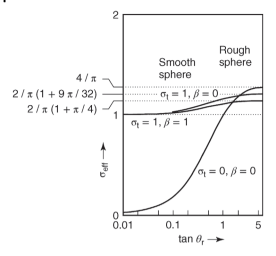


Figure 13.24 Effective accommodation coefficients $\sigma_{\rm eff}$ versus tan $\theta_{\rm r}$ (Figure 13.23) for selected values of the tangential momentum transfer coefficient $\sigma_{\rm t}$ and the energy transfer coefficient β according to *Fremerey* [28].

we can calculate an effective mass

$$m_{\text{eff}} = \left(\sum_{i=1}^{n} \chi_i(\sigma_{\text{eff}})_i \sqrt{m_i}\right)^2, \tag{13.20}$$

which allows total pressure measurements of gas mixtures when put into \overline{c} in Eq. (13.19).

13.3.3

Measuring Procedure

For measuring the DCR signal, an obvious method would be to measure the number of revolutions of the sphere in constant subsequent time intervals. However, because the zero crossings (either from plus to minus or vice versa) of the sinusoidal signal are measured in the sensing coils, measurement is simplified when a fixed number of revolutions N is predefined and the subsequent time intervals τ_i elapsing during N revolutions are measured.

The deceleration of rotary frequency with time is extremely low. Thus, differences can be tolerated in the equations instead of infinitesimal quantities. It follows [26]

$$-\frac{\dot{\omega}}{\omega} = \frac{\tau_j - \tau_i}{\tau_i \tau_i}.\tag{13.21}$$

As τ_i , τ_j also denotes a time interval including N revolutions, but for a later moment in time than τ_i . For a minimum standard deviation in the measured signal within a short period of time, the so-called accumulative multiperiod

average [26] has proven successful. Here, the average is taken over n time intervals τ_i in order to reduce spreading of measured values by \sqrt{n} .

The number of revolutions N should be predefined so that enough n-intervals can be measured at higher pressure, and that, for low pressures, $\Delta \tau_i$ -values (scattering of τ_i) with respect to τ_i are sufficiently small in order to reduce measurement uncertainty.

Time intervals are measured with a quartz clock (frequency approximately 10 MHz). A computer in the instrument calculates DCR values, standard deviations of individual and mean values, and the pressure of the gas, presuming adequate input values. Input data required for pressure calculation include the temperature, the molecular weight, and $\sigma_{\rm eff}$ of the considered gas species, as well as the previously determined residual drag (offset) RD.

The residual drag must be determined at a pressure below the lower measuring limit of the spinning rotor gauge, that is, $< 10^{-6}$ Pa. For precise pressure measurements $< 10^{-3}$ Pa, it is important that the residual drag is ω -dependent (see Section 13.3.2), which means that the dependence RD(ω) must be determined prior to measurement as well. Pressure measurement must then also take into account the frequency of this previous measurement. Figure 13.25 shows a plot of RD(ω). Between 415 and 405 s⁻¹, the offset varies by approximately 10%. In the 33 h measurement, the sphere was accelerated and then freely rotated without reacceleration until it reached 405 s⁻¹. The first measured value at $f = 415 \,\mathrm{s}^{-1}$ is significantly lower than the next: the system requires several minutes after reacceleration until a reliable, precise measuring signal is obtained. Scattering is caused by standard deviations of measured values around 10^{-10} s⁻¹ and by small temperature variations.

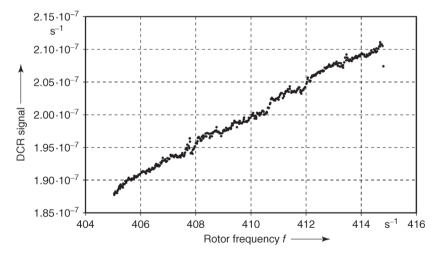


Figure 13.25 Measured dependence of the DCR signal in a spinning rotor gauge versus rotary frequency for the pressure range $p < 10^{-6}$ Pa. Residual drag RD(ω). During measurement, the speed decreased from 415 to 405 s⁻¹ within 33 h under UHV conditions.

13.3.4

Extending the Measuring Range toward Higher Pressures

At pressures p > 1 Pa, prerequisites for Eq. (13.6) are no longer met as explained in Section 13.3.1. However, precise measurements reveal that the influence of mutual collisions between gas particles is already detectable at pressures above approximately 0.1 Pa.

At pressures p > 1 Pa, the gas increasingly develops layer flow around the sphere. The final collisions of particles prior to impinging the surface of the sphere occur in a rotating gas layer and the retarding momentum transfer as well as the retarding moment are smaller than calculated with Eqs. (13.6) and (13.11).

For even higher pressures, where $\lambda \ll D = 7.5$ mm, the frictional force in laminar flow even becomes independent of pressure, according to Eq. (3.69). However, flow around the sphere is nonlaminar, alone due to the absence of rotational symmetry (the distance between the sphere and the wall is greater in the longitudinal direction than perpendicular to this direction). This leads to a slight pressure dependence of $-\dot{\omega}/\omega$ at high pressures as well, which can be utilized for measurements [29] if additional constants determined by the special geometry of the system (diameters of sphere and tube) are taken into account [30]. However, at higher pressures, continuous actuation of the sphere is required, creating considerable heat in the sphere, finger, gas, and measuring head. Thus, sufficiently accurate measuring calls for temperature monitoring and temperature correction of the measured values [29,30]. Following these problems, spinning rotor gauges are usually used for pressure measurements at pressures below 1 Pa because other simpler or more precise measuring instruments are available for higher pressures.

13.3.5

Measuring Uncertainty

Spinning rotor gauges are used mainly as reference standards and for precise pressure measurements (see Section 15.2.4). Therefore, contributions to measuring uncertainty are listed separately in this section. Measuring uncertainty is caused by the following:

- Sphere radius r and sphere density ρ : They can be determined accurately enough so that they become negligible compared with any other uncertainty. However, generally, measurement is omitted and manufacturers' nominal and average values are used. The actual values spread around these averages by approximately 0.12%. Nevertheless, if the effective accommodation coefficient $\sigma_{\rm eff}$ of the sphere is determined by calibration, then the r and ρ values of this calibration must be used. Deviations from nominal values can thus be introduced into calibration of σ_{eff} .
- Effective accommodation coefficient $\sigma_{\rm eff}$: Uncalibrated spheres from ball bearings that appear smooth show considerable scattering. For $\sigma_{\rm eff}$, Dittmann et al. found values between 0.97 and 1.06 for 67 steel balls of unknown

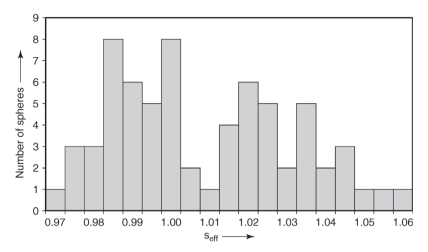


Figure 13.26 Histogram-like distribution of the effective accommodation coefficients for 67 smooth rotor spheres (steel) to different values in ranges of 0.005 width each. (Taken from Ref. [31].)

alloy composition [31] (Figure 13.26). In addition, the gas-type dependence of $\sigma_{\rm eff}$ must be taken into account when using uncalibrated spheres. Relative deviations between gas species (e.g., hydrogen and argon) can amount to 3%. The uncertainty of the calibration value $\sigma_{\rm eff}$ is only 0.3%. For the calibrated value to prevail during transport, back and forth movements as well as corrosion on the sphere should be prevented. The best way is to fix the sphere under vacuum for the period of transport [32]. Long-term stability during 1 year is generally between 0.3% and 0.5%, presuming optimum storage and transport. However, jumps in the range of 1% can occur.

- DCR value: In spite of the approximations made for deriving Eq. (13.21), uncertainties caused by neglecting the higher terms can be neglected. The scattering of measured values is by far greater and amounts to approximately 0.1-1% in offset measurements and 0.01% at 10^{-2} Pa.
- Residual drag RD: Temperature changes in the rotor can lead to false results when measuring the residual drag or the offset (see Eq. (13.17)). Therefore, temperature stability is very important, which also includes a run-in spinning rotor gauge (about 6 h). If the frequency dependence of the residual drag is not measured, its value can change by 10% compared with the measurement (Figure 12.25). If it is taken into account, then the relative uncertainty of RD reaches 0.1–1%, depending on the case considered.

13.4 **Direct Electric Pressure Measuring Transducers**

At this point, we will list a number of effects that, in principle, can be used for pressure measuring but that have not yet been implemented in widespread applications. These effects include the pressure-dependent natural frequency of oscillator quartz [33,34], pressure-sensitive transistors, pressure-sensitive tunnel diodes, pressure-sensitive photoresistors, and the pressure dependence of the resistance of thin, soft magnetic coatings [4].

13.5 Thermal Conductivity Vacuum Gauges

13.5.1

Principle

Thermal conductivity vacuum gauges are pressure measuring instruments for medium and low vacuum that measure the pressure-dependent thermal loss (loss of energy) of a heated element, usually a wire, through the gas.

The measuring principle is based on a 1906 publication by Pirani [35]: a heated wire is part of a Wheatstone bridge that supplies the necessary energy to the wire and measures its resistance or the dissipated power. English literature, in particular, often refers to thermal conductivity vacuum gauges based on a Wheatstone bridge simply as Pirani gauges.

Thermal conductivity vacuum gauges (Pirani gauges) offer several different modes of operation: thermal conductivity vacuum gauges that maintain constant wire temperature and measure the required heating power, which depends on the pressure, represent the most precise setup and show the largest linear measuring range (0.1 Pa to 10 kPa). However, measuring technology here is the most complex and thus expensive. Alternatively, the heating power or current is kept constant and the compensation current in the bridge can be used as a measure of pressure.

Also, circuits that keep heating power constant and measure the temperature, which depends on the gas pressure, are possible. Usually, a thermocouple is used for this temperature measurement. Here, however, the measuring range is only approximately 0.1 Pa to 1 kPa. Less sophisticated and cheaper versions do without a Wheatstone bridge and use a simple thermocouple for measuring the current that follows the contact voltage in the thermoelement if the heated wire receives constant current. Their measuring range is 0.1–100 Pa.

The principal design of a thermal conductivity vacuum gauge (compare Figure 13.27) usually includes a wire W with a diameter of 5-20 μm and length of 50-100 mm suspended axially in a cylindrical tube of 10-30 mm diameter. When the wire is heated electrically, it approaches an equilibrium temperature T_1 at which the supplied electrical power $\dot{Q} = IU$ equals the dissipated thermal power. The latter is made up of four components (Figure 13.28):

1) Thermal conduction $\dot{Q}_{\rm gas}$ via the gas between the warm wire ($T_1 \approx 400~{\rm K}$) and the wall at room temperature ($T_2 \approx 300 \,\mathrm{K}$). According to Eq. (3.98), the energy flux rate or dissipated thermal power is proportional to the pressure

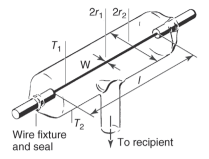


Figure 13.27 Basic setup of a thermal conductivity vacuum gauge.

p. In the viscous flow regime, $\dot{Q}_{\rm gas}$ is independent of pressure (see Eqs. (3.99) and (3.90)). In the transition range, a function of the type p/(1+gp) (g: constant) describes the pressure dependence of Q_{gas} well, and thus,

$$\dot{Q}_{\rm gas} = \varepsilon \frac{p}{1 + gp}.\tag{13.22}$$

The constant ε (sensitivity) includes gas characteristics such as C_{molar} and \overline{c} (compare Eq. (3.98)), and g is a factor determined by the geometry (as well as M_r) (Eq. (3.99)).

- 2) Thermal conduction \dot{Q}_{end} at the wire ends via the wire fixing.
- 3) Thermal radiation \dot{Q}_{rad} emitted by the hot wire (hot with respect to its surrounding).
- 4) Thermal conduction \dot{Q}_{conv} due to convection at pressures > 1 kPa.

 $Q_{
m end}$ and $Q_{
m rad}$ are disturbing effects that create the false impression of a gas pressure p_0 in the measuring cell even if the cell does not contain any gas, that is, if pressure is zero. We will set

$$\dot{Q}_{\rm end} + \dot{Q}_{\rm rad} = \varepsilon p_0, \tag{13.23}$$

which defines zero pressure. Parameters in the functional dependences of the thermal loss due to radiation include emissivity and geometry of the surface as

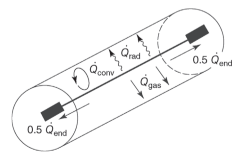


Figure 13.28 Heat fluxes from a heating wire in a thermal conductivity vacuum gauge: Q_{gas} via the gas, $\dot{Q}_{\rm rad}$ due to radiation, $\dot{Q}_{\rm conv}$ due to convection, and $\dot{Q}_{\rm end}$ due to thermal conduction.

well as the temperature distribution on the wire. Thus, on the one hand, calculation is difficult and, on the other hand, εp_0 might change from one day to the other or even during the course of a single measurement. For simplification, however, constant Q_{end} , Q_{rad} , and thus p_0 are assumed as long as the temperatures of wire and surrounding, T_1 and T_2 , respectively, are kept constant. This is one reason for measurement uncertainty. At pressures low enough for convection to be negligible, the power balance at the wire W reads

$$\dot{Q}_{\rm el} = \dot{Q}_{\rm end} + \dot{Q}_{\rm rad} + \dot{Q}_{\rm gas},\tag{13.24}$$

or using Eqs. (13.22) and (13.23),

$$\dot{Q}_{\rm el} = UI = \frac{U^2}{R} = \varepsilon \left(p_0 + \frac{p}{1 + gp} \right). \tag{13.25}$$

Thus, for pressures at which $\dot{Q}_{\rm gas} \propto p$, measuring the electrical heating power \dot{Q}_{el} supplied to the wire in a measuring cell (resistance R) according to Figure 13.27, or a related quantity, with respect to pressure p, yields a linear proportional signal (Figure 13.29). At higher pressures where thermal conductance Q_{gas} is pressure independent, heat dissipation rises only due to convection. At low pressures, thermal radiation, which is also pressure independent, dominates so that the measured signal merges into a constant offset (Figure 13.29).

Figure 13.30 shows a measuring example. For selected gases, heating powers $Q_{\rm el}$ were measured versus pressure p using a tungsten wire with a length of l=160 mm and a diameter of 7.7 μ m, at a temperature difference $\Delta T = T_1 - T_2 =$ 100 K. The plots show that the sensitivity of the vacuum gauge, given by the position of the curves (dashed lines), depends on the gas species. Furthermore, zero pressures p_0 (at the intersection of the dashed lines and the asymptotic

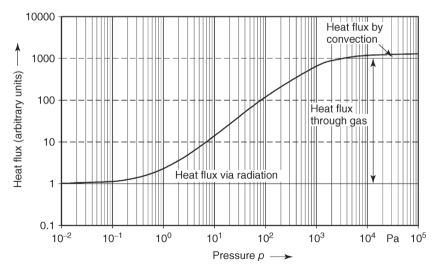


Figure 13.29 Diagram of heat flux in a thermal conductivity vacuum gauge versus pressure.

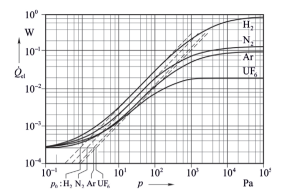


Figure 13.30 Electrical heating powers \dot{Q}_{el} for maintaining constant heating-wire temperature T_1 ($\Delta T = T_1 - T_2$) versus gas pressure p for selected gas species. Straight dashed lines indicate the linear range of thermal conduction ($qp \ll 1$).

horizontal line) vary between 0.9 and 2.2 Pa, depending on the gas species. For $\Delta T = 100 \,\mathrm{K}$, the value of p_0 is weakly dependent on the temperature T_1 but it does change with the total emissivity of the wire surface. In the measurements yielding Figure 13.30, the radiation proportion was 10% of the zero power.

Zero pressure determines the lower limit for the measuring range of a thermal conductivity vacuum gauge. For a gas with high relative molecular mass M_r , for example, UF₆, the upper limit p_{max} is approximately 1 kPa. For hydrogen with $M_{\rm r}=2$, in contrast, it is approximately 50 kPa. It should be noted that for UF₆ (high M_r), measurement never reaches the linear range $\dot{Q}_{el} \propto p$. Table 13.2 lists data on the sensitivity ε as well as lower and upper measuring limits of commercially available thermal conductivity vacuum gauges. Due to the low thermal dissipation via convection, the measuring range can be extended to 100 kPa for heavy gases as well.

13.5.2 Thermal Conductivity Vacuum Gauges with Constant Wire Temperature

Here, a thin wire made of tungsten or nickel with a diameter of $7-10 \,\mu m$ is used for measuring. As Figure 13.31 shows, the wire represents one arm Rw of a

Table 13.2 Sensitivities ϵ , lower limits (zero pressures p_0), and upper limits p_{max} for the measu	r-
ing range of standard thermal conductivity vacuum gauges.	

Gas species	ε (W Pa $^{-1}$)	p ₀ (Pa)	p _{max} (kPa)
H_2	2.9×10^{-4}	0.90	50
N_2	1.9×10^{-4}	1.37	20
Ar	1.3×10^{-4}	2.16	15
UF ₆	1.1×10^{-4}	2.20	1

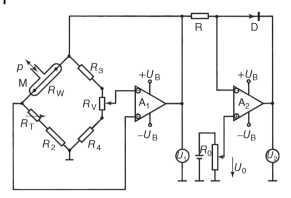


Figure 13.31 Electrical diagram for a thermal conductivity vacuum gauge with constant wire temperature. M: measuring cell; R_w : resistance of measuring wire.

Wheatstone bridge including the resistances $R_2 \approx R_3 \approx R_4 \approx R_w$ and a resistor for temperature compensation R_T . Amplifier A_1 controls the voltage U_1 at the bridge so that the resistance of the measuring wire, and thus its temperature, is heat flux independent, that is, independent of the pressure. In this case, the bridge is balanced at all times.

Electronics required for displaying the dissipated thermal power $\dot{Q}_{\rm el}$, or the square of heating voltage, are complex. Therefore, the bridge is often not fully balanced; instead, the voltage U_1 at the bridge is displayed using an appropriately calibrated voltmeter. In this case, the relation between the shown voltage U_1 and pressure p is approximately given by (Eq. (13.25))

$$U_1 = 2\sqrt{R_{\rm w}\varepsilon\left(p_0 + \frac{p}{1+gp}\right)}. (13.26)$$

Thus, a nonlinear scale is obtained, with zero pressure at approximately 1 Pa so that the pressure range between 0.1 and 1 Pa defies measurement. A more complex circuit (right-hand side in Figure 13.31) allows compensating for zero pressure using a voltage U_0 at potentiometer R_0 . Taking the logarithm of the voltage $U_1 - U_0$ by means of a diode D yields a scale as shown in Figure 13.32. Today,

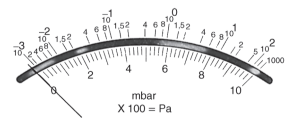


Figure 13.32 Analog scale in an older thermal conductivity vacuum gauge with constant wire temperature and zero-point compensation, indicating pressure dependence of signal sensitivity. Today, digital scales have replaced the former analog scales.

such analog scales have been replaced by digital scales. However, at a glance, Figure 13.32 clearly demonstrates the signal sensitivity to pressure in a thermal conductivity vacuum gauge.

Digital readings often depend on a characteristic curve where a change of 1 V corresponds to a change of one decade in pressure (e.g., the range 1-9 V corresponds to pressure range of 10^{-3} to 10^{5} Pa). This, on the decades equally distributed curve, however, should not mislead the user that due to the reduction of sensitivity at the limits of the measuring range measurement uncertainties greatly increase. Voltmeters can be used to indicate output voltage of "active" thermal conductivity vacuum gauges, which is then converted to a pressure reading by a formula or table.

At gas temperatures around 300 K and wire temperatures > 400 K, sensitivity ε is no longer temperature dependent. A means of calibration is often provided because sensitivity ε and zero pressure p_0 vary from one measuring setup to the next. Calibration involves the following steps.

At atmospheric pressure, the temperature set point of the heating wire, and thus sensitivity ε_i , is adjusted using potentiometer R_V (Figure 13.31) so that the display shows 100 kPa. Next, the measuring tube is evacuated to a pressure that is small compared with the lowest measurable pressure, in this case, approximately 0.01 Pa. Then, compensation voltage U_0 is adjusted with R_0 until the display indicates that U_2 equals zero. In some cases, the calibration potentiometers $R_{\rm V}$ and $R_{\rm 0}$ are integrated into the head of the measuring tube, together with the remaining resistances of the Wheatstone bridge and the resistor for temperature compensation $R_{\rm T}$, as shown in Figure 13.33. Then, the measuring tube can be calibrated prior to delivery so that it can be hooked up to any measuring instrument without requiring any individual adjustment.

This calibration generally prevails as long as the heating wire does not suffer any irreversible changes. There is a possibility of increased blackening on the heating wire due to deposition of contaminants. Such an increase causes a rise

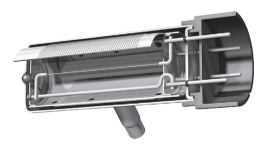


Figure 13.33 Section of a measuring tube in a thermal conductivity vacuum gauge (constant wire temperature) with gold-plated tungsten heating wire and a measuring range from 0.1 Pa to 100 kPa. The surrounding cylindrical housing (see heating wires) is temperature

controlled to approximately 40 °C for increased measurement accuracy. At higher pressures, heat transport due to convection (Figure 13.30) is used as a measuring signal. (Courtesy of Granville-Philips/VACOM GmbH.) in zero pressure, which leads to too high values displayed by the instrument in the low-pressure range, that is, the zero point is never reached. Subsequent calibration using voltage U_0 can compensate this deviation throughout a wide range without having any considerable impact on measuring accuracy.

The tungsten wire in the thermal conductivity vacuum gauge shown in Figure 13.33 is gold plated in order to avoid changes in the heating wire.

An additional method of zero-point compensation is given by designing resistor R_3 (Figure 13.31) as a pressure/vacuum-tight encapsulated heating wire element (compensation element), analogous to the measuring-wire cell. If these two cells are thermally short-circuited, temperature changes in both arms of the bridge will have analogous effects: the bridge remains balanced, but only for a certain pressure, namely, the pressure in the compensation element's capsule, because the temperature variation in the heating wire depends on the surrounding pressure. Thus, it is advisable to realize compensation by incorporating variable resistors.

Compared with simple designs (see following sections), sophisticated thermal conductivity vacuum gauges with constant wire temperatures provide wider measuring ranges and have the advantage that they react quicker to pressure changes. This is because their components are not subject to any temperature changes, and thus, no thermal relaxation periods arise, which would delay measuring signal acquisition.

Instead of a hot wire, modern thermal conductivity vacuum gauges manufactured by microtechnology use a 1 mm² hot silicon plate and measure its dissipated heat (Figure 13.34). Two meander-shaped, vapor-deposited resistor "wires" are used for heating the plate as well as for temperature measurement. Here, thermal conduction occurs at the complete edge of the plate. Again, the sensor plate is wired (Figure 13.34b) so that a constant temperature difference is established between the plate and the surrounding shell. Two temperature sensors arranged in the surrounding substrate provide temperature compensation in the event of variations in ambient temperature.

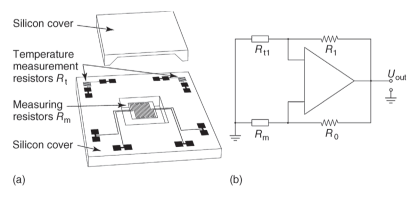


Figure 13.34 (a) Diagram of the sensor in a MicroPirani, manufactured by means of microtechnology. The 1 mm × 1 mm plate is heated to approximately 60 °C. (Courtesy of MKS Instruments, Denmark.) (b) Electrical circuit diagram for the MicroPirani shown in (a). $R_0 \approx R_{\rm m}$.

The small size of microtechnologically fabricated thermal conductivity vacuum gauges has the advantage that convection does not develop so that the measured values are independent of the mounting position. Due to its stable geometry, such an instrument can operate in accelerated systems as well. The lower limit p_0 (Eq. (13.23)) of the measuring range is reduced, compared with a thermal conductivity vacuum gauge using a macroscopic wire, because the lowered operating temperature of approximately 60 °C of the plate reduces thermal radiation as well as thermal conduction.

13.5.3

Thermal Conductivity Vacuum Gauges with Constant Heating

In addition to the operating mode of a thermal conductivity vacuum gauge using constant wire temperature as described in Section 13.5.2, the heating wire can also be supplied with constant heating voltage (or constant electrical heating current) (Figure 13.35). Then, the resistance of the wire is acquired as a measure of its pressure-dependent temperature.

Pirani's [35] antiquated principle uses a heating wire that forms one arm of a Wheatstone bridge operating at constant voltage (U_B) . If the temperature in the heating wire changes due to pressure variations, resistance R_w in the wire also changes. This change in resistance detunes the bridge, thus creating a voltage signal U on the indicating instrument in the bridge's diagonal (Figure 13.35). Using appropriate calibration, the voltage can be translated and displayed as pressure p. The temperature coefficient of the electrical resistance in the measuring wire's material should be as high as possible. Pure metals or semiconductor resistors (thermistors with positive or negative temperature coefficient) usually meet this condition.

The display's pressure dependence is derived similarly to the method described in Section 13.5.2. Neglecting second-order quantities, we find the equation

$$U = AU_{\rm B} \frac{p}{1 + Bd/l} \approx \frac{AU_{\rm B}}{Bd/l} p = \varepsilon p, \tag{13.27}$$

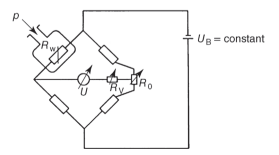


Figure 13.35 Electrical circuit in a thermal conductivity vacuum gauge using constant supply voltage and variable wire temperatures. Top left: measuring cell; R_w : resistance of measuring wire.

with the constants A and B. The primary parameter for the measuring range of these thermal conductivity vacuum gauges is the ratio of wire diameter d and wire length l.

The upper limit of the measuring range of such instruments is reached when wire temperature T_1 is just above ambient temperature T_2 . Since the balancing current in the bridge's diagonal is proportional to the temperature difference $T_1 - T_2$, which drops as the pressure rises, sensitivity ε also drops with rising pressure and the scale becomes nonlinear. The linear range for a *Pirani* gauge with constant heating power is smaller than that in a thermal conductivity vacuum gauge operating at constant temperature. The lower measuring limit is generally one thousandth of the maximum measurable pressure.

If a thin long wire is used, the factor d/l in the denominator in Eq. (13.27) is small, and thus, sensitivity ε is high. However, the highest measurable pressure is fairly low. In contrast, a short wire with larger diameter shifts the measuring range toward higher pressures. Advances in electronics have led to low-priced solutions for measuring small currents. Thus, the reduced sensitivity for lower values of d/l is no longer significant and the general approach is to use the wider linear measuring range [36].

Possible temperature changes in the gas or in the surrounding have considerable impact on the measuring inaccuracy of a Pirani gauge. For Pirani gauges with constant heating power, in particular, this effect increases as pressure rises because the temperature difference between the wire and the environment drops. At wire temperatures of 400 K (p = 0) and a measured pressure of 100 Pa, a temperature change of 1 K produces a deviation in the measured value of approximately 10 Pa. At 600 K wire temperature, the deviation is still between about 1 and 5 Pa [36]. As the ambient temperature changes by 1 K, the zeropoint signal changes by approximately 0.1 Pa. The same change is observed when the voltage on the bridge changes by 0.1%.

13.5.4

Thermocouple Vacuum Gauges

The temperature of the heating wire can also be measured directly using a thermocouple as shown in Figure 13.36. The thermocouple is attached to the heating wire and its thermoelectric voltage is directly monitored. Spot welding or brazing with silver filler metal is used to join the thermocouple, for example, chromel/alumel, and the heating wire. The typical current fed to the heating wire is 150 mA. Here, the wire is considerably shorter than that in a sophisticated *Pirani* gauge, and thus, sensitivity at low pressures is reduced. Typical measuring ranges lie between 0.1 and several 100 Pa. The system is calibrated at low pressure by adjusting the temperature of the heating wire using a controllable dropping resistor so that the display shows end-scale deflection (highest voltage level). Additional line-up at atmospheric pressure is often dispensable because then the temperature of the heating wire is just slightly above ambient temperature. Here also, it can be necessary to compensate for changes in the surface condition of

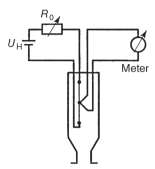


Figure 13.36 Diagram of a thermal conductivity vacuum gauge with thermocouple.

the heating wire by readjusting the compensation potentiometer. Above considerations apply likewise to the measuring range and its dependence on wire diameter and wire length.

13.5.5

Thermistors

Thermistors are thermal conductivity vacuum gauges in which the heating wire is replaced by oxidic semiconductors usually showing a negative resistance coefficient $d\rho/dT$. Its positive value is considerably above that of tungsten and platinum. Thus, in a bridge circuit, the compensation currents under unbalanced operation are considerably above the currents appearing in glow wires. For improving zero-point stability disturbed by changes in ambient temperature, the same semiconductor material is encapsulated under high vacuum and integrated into the bridge.

13.5.6

Guidelines for Operating Thermal Conductivity Vacuum Gauges

The main application of thermal conductivity vacuum gauges is in the mediumvacuum pressure range, and here, in particular, for monitoring the fore-vacuum of multistage pump systems and roughing pump operation. Thermal conductivity vacuum gauges, particularly those operating in constant-temperature mode, are well suitable for control purposes because their response time is very low, 20-50 ms, and their output signal is wide, 0-10 V. These properties as well as the fact that thermal conductivity vacuum gauges are the cheapest electrically displaying vacuum gauges are the reasons for their predominant position among other vacuum gauges.

For control processes, however, it should be noted that the acquired value in thermal conductivity vacuum gauges might be pressure independent at pressures exceeding 10 kPa. This applies particularly to gas species with a molecular weight higher than that of nitrogen. Thus, in venting processes,

Gas species	Range (Pa)	Mean CF	Variation (±)	
H_2	0.1–13	0.62	0.07	
He	0.1-13	1.04	0.12	
Ne	0.5-30	1.39	0.07	
CO	0.1-200	0.98	0.02	
N_2	0.1-10 000	1.00		
Ar	0.1-90	1.62	0.12	
CO_2	0.1-30	0.95	0.03	
Kr	0.5-30	2.22	0.16	
Xe	0.5–13	2.71	0.20	

Table 13.3 Correction factors CF and their spread for the pressure indication of four different thermal conductivity gauges with constant-temperature mode of the heated element for several gas species, when the reading for nitrogen is correct (pure gases).

The second column indicates the range where CF can be applied. CF_{H2} = 0.62 means that the reading of the vacuum gauge, calibrated for nitrogen, is too high for hydrogen. Therefore, the reading in the range 0.1-13 Pa has to be multiplied by (0.62 ± 0.07) to obtain the true hydrogen pressure. From Ref. [38].

there is a danger of overpressure load, and therefore, using a pressure relief valve is advisable.

Thermal conductivity vacuum gauges are made of W, Pt, Ni, and alloys of Cr and Ni as well as Cr and Al. Pt and Ni are catalytic and might initiate ignition of explosive gas mixtures. Hot spots and fractured wires as well as short circuits can additionally cause ignition so that thermal conductivity vacuum gauges should not be used when exposed to explosive gas mixtures.

Changes on the heated surface such as oxidation and contamination influence heat transport and thus cause indication errors. If these exceed tolerable limits, the vacuum gauge must be cleaned and recalibrated or replaced. Also, heated elements coated with ceramic substances are offered for use with chemically aggressive gases.

Scales in the displays of commercial thermal conductivity vacuum gauges are valid for air and nitrogen. For other gas species, manufacturers provide calibration curves standardized to air or nitrogen. Figure A.14 shows an example, and Table 13.3 gives correction factors for several gas species.

It has to be mentioned that, because of the energy accommodation of the gas molecules at the heated surface, the correction factors may depend on the surface state of the heated element and dirt may affect the factors.

13.6 Thermal Mass Flowmeters

Using the effect of heat transport in a gas is not restricted to pressure measurement but can be used for assessing gas flow as well. Whereas pressure

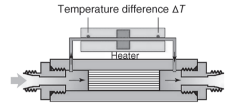


Figure 13.37 Thermal mass flowmeter with a heating element and two temperature sensors measuring the difference between upstream and downstream temperatures, $\Delta T = T_h - T_{lo}$. (Design courtesy of *MKS Instruments*.)

measurement utilizes the heat transport from a hot element via thermal self-movement of gas particles, flowmeters utilize the heat transport in a collective flow field of gas particles directed toward low-pressure regions. A thermal mass flowmeter measures the heat transferred from a small, heated tube to a gas flow on its inside.

Gas flowmeters with measuring ranges 7×10^{-8} to 4×10^{-2} mol s⁻¹ $(0.1-5\times 10^4 \text{ sccm}; \text{ for conversion see Table 4.1})$ are used mainly in semi-conductor industry for measuring and controlling gas flows in process reactors. Here, thermal flowmeters based on the introduced principle have prevailed.

A thermal gas flowmeter is made up of a small capillary (usually stainless steel) with an inside diameter of 0.25–1 mm, covered with a coiled heating wire that shows high resistance and a high temperature coefficient of electrical resistance. Such a wire can be used as a heating element and as a temperature sensor. A number of designs are available:

- Two temperature sensors that measure the downstream and upstream capillary temperatures (Figure 13.37) frame a single heating element.
- Two heating elements are integrated into a *Wheatstone* bridge and react to changes in the temperature profile caused by the gas flow.
- Three heating elements at temperatures T_1 , T_2 , and T_3 are held at constant temperatures with $T_2 > T_1 > T_3$, for a direction of gas flow from 1 to 3. The voltage for the second heating element is the measuring signal.

For the first approximation, the mass flow rate \dot{m} of the gas can be calculated from

$$\dot{m} = \frac{\dot{Q}}{c_p(T_{lo} - T_{h})}. (13.28)$$

 \dot{Q} is the heat flow from the capillary or the heating element into the gas, c_p is the molar heat capacity of the gas, $T_{\rm h}$ is the upstream temperature of the gas (high-pressure side), and $T_{\rm lo}$ is the downstream temperature. Either the temperature difference $(T_{\rm lo}-T_{\rm h})$ is held constant and the required heating power \dot{Q} is measured, or \dot{Q} is kept constant and $(T_{\rm lo}-T_{\rm h})$ is measured.

Nonlinearities (deviations from Eq. (13.28)) can have different causes:

- c_p not only depends on the gas species but is also a function of temperature.
- The gas temperature can be different from the measured wall temperature of the capillary.
- The heat transfer coefficient $\dot{Q}_{gas}/\dot{Q}_{wall}$ can depend on the gas flow.
- As in thermal conductivity vacuum gauges, disturbing effects can be caused by heat transport in the capillary itself, by radiation losses, and by thermal losses due to convection outside of the capillary.

The flow is separated in order to increase the measuring range of a thermal mass flowmeter (Figure 13.37): a bypass allows free gas flow, and a branch feeds the gas to the actual sensor. An appropriate flow barrier is arranged in the bypass in order to guarantee laminar flow throughout the entire measuring range.

It is absolutely crucial to encapsulate the flow sensor with a temperature-stabilized housing in order to, on the one hand, precisely define T_h and, on the other hand, minimize some of the disturbing effects described above as well as provide zero-point stability. Usually, the temperature of the housing is set to 60 °C.

Figure 13.38 shows the basic temperature distribution in a flow sensor with heating element [37]. Curve 1 schematically shows the temperature distribution along the capillary for zero gas flow. Curve 2 shows the temperature distribution at the capillary wall for a flow of \dot{m} through the capillary. Curves 3 and 4 illustrate the temperature of helium and nitrogen, respectively, flowing through the capillary with m. Helium behaves differently from nitrogen because the so-called diffusivity of helium, the ratio of thermal conductivity and specific heat, is higher than that for nitrogen. The higher this quantity, the sooner the temperature of the gas will adopt to the temperature of the surrounding capillary.

If gas flow changes, the temperature profile of the gas will change rapidly, within milliseconds, whereas the temperature profile of the capillary, due to its

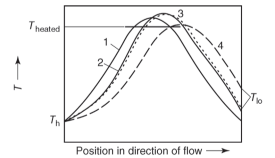


Figure 13.38 Basic temperature sequence in a thermal mass flowmeter according to Figure 13.37 versus position in the sensor. (1) Temperature of capillary wall without any gas flow, (2) temperature of capillary wall with gas flow, (3) (dashed) temperature of the flowing

helium, and (4) temperature of flowing nitrogen. The position of the tube element heated to T_{heated} is marked with a solid line. T_h: temperature on the high-pressure side (upstream); T_{lo} : temperature on the low-pressure side (downstream).

much higher heat capacity, takes several seconds to obtain a new thermal equilibrium. Thus, changes in flow produce time constants of several seconds that must be taken into account when used in control circuits.

Thermal gas flowmeters are usually calibrated for nitrogen. Manufacturers provide correction factors $K_{\rm gas}$ for other gas species. However, these factors have to be replaced by correction functions $K_{gas}(\dot{m})$ if measuring accuracies of less than 20% are required [38,39] because the given correction factors do not consider thermal conduction through the gas.

The bypass can create additional dependences of gas correction factors: for different gas species, viscosity is differently temperature dependent. Thus, for different gas species, gas flows are not distributed among the measuring capillary and the bypass in the same way.

Metrological investigations showed [39] that the orientation of flowmeters with respect to the gravitational field has an influence on the zero point (the temperature distribution of the capillary changes due to variations in convection). However, measured values remain correct if the zero point is corrected. Also, the dependence of sensitivity toward inlet pressure is negligible (0.75% deviation per MPa pressure change). Long-term stability is quite high and the relative measurement deviations remain below ±1% during an operational period of 1 year [39]. Yet, uncalibrated instruments showed measuring deviations from correct gas flows of up to 17% [39].

13.7 **Ionization Gauges**

13.7.1

Principle and Classification

Ionization gauges measure pressures indirectly by determining an electrical quantity proportional to the particle number density n. In order to make this quantity available, the gas whose pressure is to be measured is partially ionized in the measuring head of the ionization gauge. Depending on the method of ionization, the measured electrical quantity is either a pure ion flow (hot- or emitting-cathode ionization gauge, Sections 13.7.3 and 13.7.3.2) or a gas discharge flow (crossed-field ionization gauge, Section 13.7.4).

In the hot-cathode gauge, the electrons used for ionizing the gas are emitted from an emissive cathode, usually a glow cathode (C, Figure 13.41), and accelerated toward a surrounding anode screen. The electrons emitted by cathode C with a current I^- collide with gas particles that subsequently become ionized with a certain probability. The resulting positive ions reach the ion collector (IC) and are measured as an ion current I^+ .

In ionization gauges with crossed electromagnetic fields, a gas discharge is created by establishing a sufficiently high DC voltage (several kilovolts) between two metal electrodes (cathode, anode). In order to sustain the gas discharge at low pressures also, electron paths are extended considerably by superimposing a magnetic field crosswise over the electrical field. The gas discharge current is pressure dependent and is used as a measure of gas density, that is, gas pressure.

The term cold-cathode ionization gauge should be avoided because it may cause some misunderstanding. Cold cathodes are cathodes that emit electrons not via a glowing wire but by cold field emission. Such cold cathodes may be once capable of replacing the hot cathodes in hot-cathode ionization gauges.

13.7.2

History of Ionization Gauges

The birth of ionization gauges dates back to 1909, when von Baeyer [40] showed that a triodic vacuum tube can be used as a vacuum gauge. However, Buckley [41] is usually named as the inventor of the ionization gauge. Later, he further developed the triode as an ionization gauge down to a measuring limit of 10^{-6} Pa.

Three electrodes are required for an ionization gauge: the cathode as an electron source, the anode, and the collector gathering the emitted positive ions. Classic glass-encapsulated triode tubes featured three electrodes (Figure 13.39). The grid could be used as ion collector. More commonly, however, the anode plate was used as collector because here ion flux was higher, and thus, the triode was more sensitive.

Some of the basic ideas used for the electrical triode circuit in Figure 13.39 are still used today. With respect to the cathode, the ion collector has to be at negative potential so that it attracts ions only, and not electrons. The acceleration voltage for electrons, that is, the voltage between the cathode and anode, has to be around 100 V.

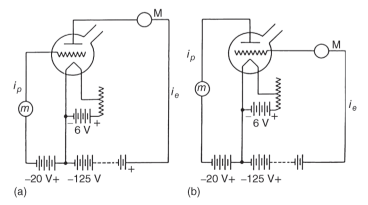


Figure 13.39 Using a triode as an ionization gauge [42]. (a) Controlling electrode (grid) acts as ion collector; (b) controlling electrode acts as anode for accelerating electrons. The former anode works as collector.

The reason for this is that the probability of ionizing neutral gas molecules depends on the electron energy and, for most gases, reaches a maximum at 100 V (Figure 13.43).

Because high electron energy along the entire path is desired, the voltage is usually adjusted to slightly above 100 V. This has the additional advantage of reducing the difference in ionization probability for different gas species.

The main setup of the triode remained unchanged for 30 years although physicists wondered why no vacuum vessel could be evacuated to a pressure below 10⁻⁶ Pa. Vacuum pumps improved constantly and measurements of change rates such as work function or thermionic emission indicated during the 1930s and 1940s that indeed much lower pressures had been produced than those measured by the triodes.

During the first International Vacuum Congress (IVC) in 1947, Nottingham suggested that the lower pressure limit might not be due to the pumps but caused by an X-ray effect in the triode tubes: soft X-rays developing when the electrons hit the anode would release photoelectrons when they reach the collector. This current of photoelectrons emanating from the collector was indistinguishable from an ion current directed toward the collector.

Bayard and Alpert soon confirmed this hypothesis. They reduced the size of the collector by constructing it as a thin wire on the axis of a cylindrical mesh anode instead of using a cylinder that encloses the remaining electrodes. This elegant solution reduced the lower measuring limit by two orders of 10 and is still used in today's so-called Bayard-Alpert tubes.

Penning [43] is considered the inventor of the crossed-field ionization gauge. It was based on his 1936 patent on sputter coating [44]. He used a high voltage of 2 kV to ignite a discharge between a cathode and an anode. An additional magnetic field, perpendicular to the electrical field, considerably increased the path length of the electrons by forcing them onto helical paths, and thus, the discharge was maintained at low pressures (Figure 13.40).

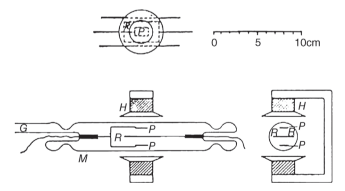


Figure 13.40 Historical Penning tube dating back to 1937 [43]. M: glass housing of manometer; H: magnets; R: ring anode; P: cathode; G: glass tube to vacuum system. Later, Penning extended the anode ring used

in the first design to an open cylinder [45]. This geometry is used in today's ion getter pumps, but only in simple, robust measuring devices.

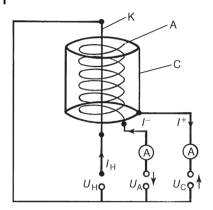


Figure 13.41 Basic design of an ionization gauge with emitting cathode. U_H: heating voltage; $I_{\rm H}$: electrical heating current in the emitting cathode; $U_{\rm A}$: anode voltage; I^- : electron current to the anode; I⁺: ion current to the collector; K: cathode; A: anode; C: collector.

13.7.3

Emitting-Cathode Ionization Gauges (Hot-Cathode Ionization Gauges)

13.7.3.1 Measurement Principle

On their path Δl through a gas of particle number density n (compare Figure 13.42 and Section 3.2.5), N^- electrons participate in

$$\Delta N^{-} = N^{-} n \sigma \Delta l \tag{13.29}$$

collisions and thereby produce the same amount of ions $\Delta N^+ = \Delta N^-$. Here, σ is the ionization cross section, which depends on the energy of the colliding electrons and the gas species. The quantity $n\sigma$, termed differential ionization P_{ion} , describes the number of ions produced by each electron per unit length of its path in a gas with particle density n. P_{ion} is often given for a pressure of p_n = 1 Torr = 133 Pa and the temperature T_n = 273 K. According to the equation of

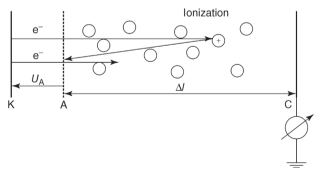


Figure 13.42 Diagram of an ionization gauge for calculating ion current I^+ . U_A : accelerating voltage for electrons; eUA: electron energy; K: cathode; A: anode; C: collector.

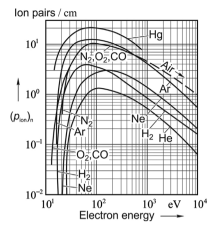


Figure 13.43 Differential ionizations $(P_{ion})_n$ of electrons versus energy levels in selected gas species at a pressure of 133 Pa and a temperature of 0 °C.

state of an ideal gas (Eq. (3.19)), $n_n = 3.54 \times 10^{22} \text{ m}^{-3}$. Figure 13.43 shows the differential ionization $(P_{ion})_n$ of selected gas species in this state.

Dividing Eq. (13.29) by time yields the electrical current. Thus, the electron current I_e produces the ion current

$$I^{+} = I_{e} n \sigma \Delta l = I_{e} P_{ion} \Delta l, \tag{13.30}$$

with $P_{\text{ion}} = (P_{\text{ion}})_n n / n_n$. Putting the value of n, given by Eq. (3.19), into Eq. (13.30) yields the relationship between the ion current I^+ to collector C and pressure:

$$I^{+} = I_{e} \frac{\sigma \Delta l}{kT} p = I_{e} \frac{P_{\text{ion}}}{p_{p}} \cdot \frac{T_{n}}{T} \Delta l p = I_{e} S p.$$
(13.31)

This introduces the gauge sensitivity (ISO 3529:2013), also referred to as sensitivity coefficient, S:

$$S = \frac{I^{+}}{I_{e}p} = \frac{P_{\text{ion}}}{p_{\text{n}}} \cdot \frac{T_{\text{n}}}{T} \Delta l, \tag{13.32}$$

which depends on the gas species, the mean free path Δl of the electrons in the measuring head, that is, on system geometry, and so on. Δl defies sufficiently accurate determination (except in the simple system shown in Figure 13.42, where $\Delta l \approx l$). Therefore, *S* has to be determined by calibration.

It should be noted that the ion current I^+ has a pressure-independent component. Furthermore, S is well defined only for a particular gas species. In contrast, the residual gas of pressure p_0 is a mixture of gases with usually unknown composition. Thus, S is more accurately defined by

$$S = \frac{I^{+} - I_{0}}{I_{e}(p - p_{0})},$$
(13.33)

with I_0 denoting the ion current at residual pressure p_0 .

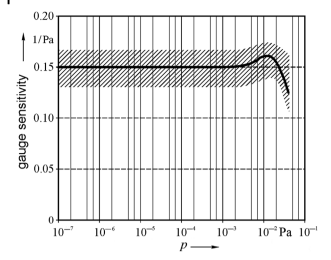


Figure 13.44 Gauge sensitivity S in Pa⁻¹ for an ionization gauge versus gas pressure p (schematic).

Equation (13.32) also shows that S depends on temperature T, a fact worth considering for accurate measurements, particularly in cases where operating temperature differs considerably from calibration temperature.

The gauge sensitivity S is given, for example, in Pa^{-1} . Figure 13.44 schematically shows S as a function of p between 10^{-7} and several 10^{-2} Pa. S drops at higher pressures because the ion current, together with the produced secondary electron current, reaches the same magnitude as the ionizing electron current (emission current). The emission current is held stable, and thus, the ionization rate drops. In addition, the free path of the electrons and ions decreases. The increasing number of collisions leads to a decline in ionization probability for each electron and to a higher probability for each ion not to reach the collector. All these effects reduce the gauge sensitivity factor at higher pressures and cause nonlinear readings on an ionization gauge. This nonlinearity starts between 10^{-2} and 1 Pa, depending on the type and geometry of the ionization gauge.

As Figure 13.44 shows schematically, there is a possibility that S rises slightly before it drops at high pressures. This is because the ion capturing probability of the collector increases due to a higher collision rate restricting axial particle motion. Then, fewer ions leave the tube without reaching the collector. This is explained in more detail in Section 13.7.3.5.

Another important gauge parameter is the ionization sensitivity *K*:

$$K = \frac{\Delta I^+}{\Delta p}. ag{13.34}$$

Here, ΔI^+ is the current change following the pressure change Δp .

According to Eq. (13.31), in emitting-cathode ionization vacuum gauges [46],

$$K = SI_{e}. (13.35)$$

Following Eq. (13.31), ion current I^+ is the pressure-dependent measured quantity. The electron current I_e emitted by the cathode can be subject to a pressure-dependent change at constant heating power due to surface coverage and additional surface effects (work function). Electronic control of the cathode heater keeps the electron current constant at an adjustable level within the range of $10 \,\mu\text{A}$ to $10 \,\text{mA}$. For a typical value $I_{\rm e} = 1 \,\text{mA}$ and with a gauge constant $S = 0.1 \text{ Pa}^{-1}$, sensitivity $K = 10^{-4} \text{ A Pa}^{-1}$, that is, an ion current of $I^+ = 10^{-8} \text{ A}$ corresponds to a pressure $p = 10^{-4}$ Pa.

Gas depletion can occur in an ionization gauge system causing a pressure drop in the measuring head. An estimate for gas depletion is obtained when assuming that each captured ion is pumped down. For this case, the pumped particle number with respect to time is calculated from Eqs. (13.32) and (13.35):

$$\dot{N} = \dot{N}^{+} = \frac{I^{+}}{e} = \frac{I_{e}}{e} Sp = \frac{K}{e} p.$$
 (13.36)

And with the equation of state, Eq. (3.19),

$$V = N \frac{kT}{p}$$
, and accordingly $\dot{V} = \dot{N} \frac{kT}{p}$, (13.37)

pumping speed is obtained:

$$S_{\text{pump}} =: \dot{V} = K \frac{kT}{e}; \tag{13.38}$$

or for $T = 296 \, \text{K}$,

$$\frac{S_{\text{pump}}}{\ell \, \text{s}^{-1}} = 26 \frac{K}{\text{A Pa}^{-1}}.$$
 (13.39)

Thus, gas depletion of the gauge is proportional to its sensitivity *K*.

13.7.3.2 Design of Emitting-Cathode Ionization Gauges (Hot-Cathode Ionization Gauges)

Based on the principle of an emitting-cathode ionization gauge shown in Figure 13.41, a variety of types have been developed, in particular, when considering measuring range [47] and manufacturing effort. In the following, the four most important designs of measuring systems are introduced, differing mainly in terms of measuring range. Figure 13.45 schematically shows the measuring ranges of the considered types. The lower measuring limit is reached as soon as the indicated pressure p_{ind} remains constant in spite of a change in the surrounding pressure $p_{\rm true}$. At the upper measuring limit, ion current, that is, indicated pressure, drops as the pressure increases.

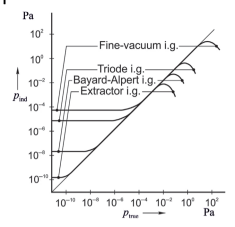


Figure 13.45 Pressure readings p_{ind} versus true pressure p_{true} for selected types of ionization

13.7.3.3 Concentric Triode

This oldest type of ionization gauge system emerged from an amplifier triode (Figure 13.39). The central cathode is surrounded by a cylindrical mesh (anode) for electron collection, enclosed by the cylindrical ion collector. The electrons emitted from the cathode accelerate toward the anode and can oscillate around the mesh wires several times before they reach the anode.

Ions produced in the process between the anode and the ion collector by electron collisions approach the ion collector, and thus produce a DC signal that is proportional to the pressure. Figure 13.46 (left) shows such an ionization gauge tube. The lower measuring limit of about 10⁻⁵ Pa is due to the so-called X-ray effect (compare Sections 13.7.2 and 13.7.3.5): electrons impinging on the anode release X-ray photons. If these photons hit the collector cylinder, they release secondary electrons that cannot be differentiated from the ion current.

13.7.3.4 Fine-Vacuum Ionization Gauges

Development of burn-out-proof oxide cathodes allowed an increase in service life of ionization gauge systems and resolved the 1 Pa maximum working pressure limit. However, in the early development stages, the measuring range of triodes could not yet be extended toward higher pressures because gauge

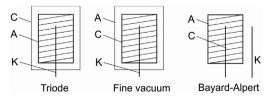


Figure 13.46 Diagram of electrode arrangement of three types of cylindrical ionization gauges with emissive cathode. A: anode; C: collector; K: emissive cathode.



Figure 13.47 Fine-vacuum ionization gauge HPG 400 by INFICON with a measuring range of 1 mPa to 100 Pa, combined with a Pirani gauge. A: anode (+180 V); C: collector (0 V, ground potential); K: cathode (+37 V); P: Pirani gauge. (Courtesy of INFICON AG, Liechtenstein.)

coefficients dropped at higher pressures. Calibration curves deviate considerably from linear functions, as shown in Figure 13.45 (triode) [48,49].

This problem was resolved by switching the functions of the anode and the ion collector and by reducing their distance. The center cathode is surrounded by an ion collecting mesh, at negative potential with respect to the cathode. An electron collector, at positive potential with respect to the cathode, encloses the ion collector. This operating mode extends the measuring range from 1 to 100 Pa. Figure 13.47 shows such a measuring system as a nude gauge. The cathode is made of an iridium band plated with thorium oxide. Such a cathode is safe from burning out, even at atmospheric pressure. However, it is much more expensive than a cathode manufactured from a thin tungsten wire. For materials used in heating wires, see Section 14.2.2.

The slope of the characteristic curve at higher pressures leads to ambiguous pressure readings. In measuring practice, it is often difficult to determine the point at which the system is on the characteristic curve. Experimental investigations help to resolve this issue: in the right part of the characteristic curve, increasing pressure reduces the indicated value.

Schulz and Phleps [50] introduced a historically famous and remarkably simple fine-vacuum ionization gauge, shown in Figure 13.48. Depending on gas species, it provided linear readings up to 130 Pa.

13.7.3.5 Bayard-Alpert Ionization Gauges

Electrons impinging on matter release photons (X-rays). Their number is proportional to the electron current. When these photons hit metal surfaces, they

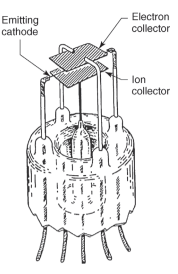


Figure 13.48 Fine-vacuum ionization gauge according to *Schulz* and *Phleps* [50]. The emitting cathode is at 60 V, the anode (electron collector) at 120 V, and the ion collector at 0 V.

release photoelectrons that leave the surface if an appropriate electrical field is present. In an ionization gauge, such a field exists at the ion collector. The pressure-dependent photoelectron current leaving the collector is proportional to the electron current and adds a constant value to the positive ion current toward the ion collector. Thus, pressure readings are too high. This yields a lower pressure limit between 10^{-4} and 10^{-5} Pa, for the measuring range in the systems introduced in Sections 13.7.3.3 and 13.7.3.4.

For reducing the X-ray effect, *Bayard* and *Alpert* [51] suggested building an ionization gauge in which the surface of the ion collector was particularly small (Figure 13.49). The collector is made of a very thin wire (diameter several 10 µm)

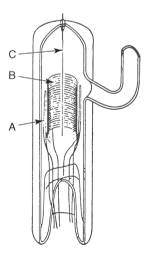


Figure 13.49 Original Bayard-Alpert ionization gauge [51]. A: cathode; B: anode grid; C: collector.

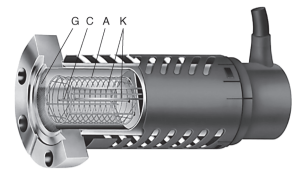


Figure 13.50 Commercially available Bayard-Alpert ionization gauge. K: cathodes (two band cathodes, each fixed at both ends); A: anode; C: central ion collector; G: mesh at ground

potential protecting against changes in electrical potential. (Courtesy of Granville-Philips/ VACOM GmbH.)

and is arranged on the axis of a cylindrical anode mesh, B. The cathode, A, is outside of the anode. Bayard-Alpert ionization gauges show an X-ray limit of 10^{-8} to 10^{-9} Pa.

Figure 13.50 shows a commercial measuring system of the kind. This sophisticated instrument with its highly stable gauge sensitivity is designed specifically for precise measurements [52,53].

Figure 13.51 shows the block diagram of an ionization gauge. The block denoted with H controls the emission current of cathode C (electron current I_e can be stepwise adjustable). Block A provides the anode potential and block C amplifies the ion current so it can be displayed on standard current meters. The emission-current control circuit adjusts the heating current of the directly heated cathode in the measuring system such that the electron current is kept precisely constant with a tolerance of 2%.

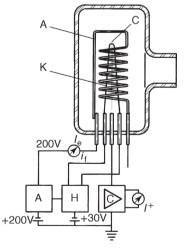


Figure 13.51 Block diagram for operating an emitting-cathode ionization gauge. Electrode system (K, A, C) according to Bayard and Alpert.

A disadvantage of the open cylinder is that ions can escape along the center axis of the cylinder. The electrical field exerts a collector-directed force onto ions produced inside the mesh. In addition, however, the angular momentum must remain constant, and thus, neutral particles with sufficiently high kinetic energy in tangential direction, which turned into ions, can in fact circulate around the collector without impinging on it. If they carry an additional axial velocity component, they thus might exit the anode mesh in the axial direction. Furthermore, escaping is promoted by the lines of the electrical field in the open mesh, which are directed axially, away from the collector. Collisions preventing this efflux at higher pressures lead to the transitional rise in the gauge sensitivity described in Section 13.7.3.1 (Figure 13.44).

Thus, the anode mesh is closed in order to increase the gauge sensitivity S. But then, however, a closed mesh produces nonlinear ion current (with respect to pressure) above approximately 1 mPa, probably due to space-charge effects that are not observed in open cylindrical meshes until pressures reach approximately 10 mPa [54].

Reducing the diameter of the collector wire leads to even smaller pressure values for the lower measuring limit. Van Oostrom [55], for example, used a wire with a diameter of 4 µm. However, a disadvantage of smaller diameters is that gauge sensitivity drops, again due to the conservation of angular momentum described above.

When operating Bayard-Alpert measuring systems, and ionization gauges in general, parasitic errors can cause pressure readings that are too high, particularly under low-pressure conditions. The pressure-independent background current [56-58] usually includes not only the photoelectron current produced by the X-ray effect (Figure 13.52) but also additional inputs from other physical

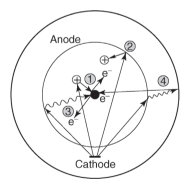


Figure 13.52 Physical effects in a Bayard-Alpert ionization gauge (collector wire in the center). (1) Electron ionizing a gas particle, subsequent release of a secondary electron when the ion impinges on the collector; (2) electron-stimulated ion desorption; (3) X-ray

effect, the X-ray photon produced during electron bombardment at the anode releases an electron at the collector; (4) inverse X-ray effect, X-rays release electrons at the wall, which hit the collector (at equal potential as the wall).

effects and parasitic errors [59-61]. The latter mainly include the so-called ESD (electron-stimulated desorption) [62], that is, desorption of positive ions caused by electrons impinging on a solid surface. In ionization gauges, the electrons hitting the anode release positive ions that reach the ion collector and thus contribute to background current I_0 . Also, negative currents can occur at the collector if the so-called inverse X-ray effect is predominant (Figure 13.52). Furthermore, dissociation of gas particles at hot surfaces (Section 13.7.3.8) and gas released by heated parts of measuring equipment are important issues.

The anode mesh has a surface area of approximately 10 cm². Thus, the number of adsorbed molecules can be considerable (1016 molecules in a single monolayer). This is why a clean anode is important: after venting or exposure to high pressures, a special degassing procedure activated by the control equipment cleans the anode mesh by means of electron bombardment. In addition, the electron current into the anode should not be too low so that the vacuum gauge is permanently self-cleaning.

Redhead [63] suggested another approach for reducing the lower measuring limit. The so-called modulator method measures background current I₀ and subtracts it from the measured signal. This requires an additional electrode in the grid space, referred to as modulator, usually a wire parallel to the collector and close to the cylindrical mesh. A modulator at anode potential has little or no effect on a Bayard-Alpert tube. The measured collector current is

$$I_1 = I_0 + I_G, (13.40)$$

with I_0 denoting the background current and I_G the ion current.

Applying a strong negative potential, approximately -100 V, most ions travel to the modulator so that the collector current

$$I_2 = I_0 + mI_G (13.41)$$

drops (m < 1). Thus,

$$I_{\rm G} = \frac{I_1 - I_2}{1 - m} \tag{13.42}$$

and

$$I_0 = \frac{I_2 - mI_1}{1 - m} \,. \tag{13.43}$$

The modulation factor *m* is determined at pressures where $I_G \gg I_0$.

However, it appeared that I_0 also changes considerably with modulator voltage because the latter influences electron trajectories. Hobson [64] estimated that the corresponding error of measurement would amount to 3×10^{-10} Pa. Thus, the idea had no significant commercial success because the additional effort could not improve the lower measuring limit substantially.

The measuring limit was finally lowered by changing the geometry of the Bayard-Alpert tube. This led to the development of the extractor ionization gauge.

13.7.3.6 Extractor Ionization Gauges

In the extractor principle, the collector is placed in a chamber separate from the volume in which the ions and X-ray photons are released. The aim is to further reduce the X-ray effect. The ions are extracted (hence, extractor ionization gauge) to this separate chamber. The spatial separation considerably reduces the exposure of the collector to X-rays.

Figure 13.53 shows the design of an extractor. An annular cathode envelopes the lower part of the cylindrical anode. At the bottom end, the anode is closely opposed to an aperture of a negative extractor cathode at ground potential. The space below the extractor electrode holds a small wire-shaped ion collector shrouded by a reflector electrode at anode potential. In this electrode arrangement, combined with appropriate design, the extraction electrode focuses the ions produced inside the anode grid onto the ion collector. The solid angle under which photons produced at the anode reach the ion collector is far lower than that in Bayard-Alpert systems. The X-ray effect corresponds to a pressure reading of approximately 1×10^{-10} Pa. The upper measuring limit is approximately 10^{-2} Pa.

Benefits of extractor systems are not limited to pressures below 10⁻⁸ Pa. For the pressure range of less than 10⁻⁶ Pa, the described parasitic errors for such vacuum gauge systems are extremely low.

Helmer [65,66] further developed the extractor principle by introducing a 90°-deflection unit for ions behind the extraction electrode (Figure 13.54). In this design, there is no line of sight between the anode and the collector, and thus, the rate of X-rays reaching the collector again drops dramatically.

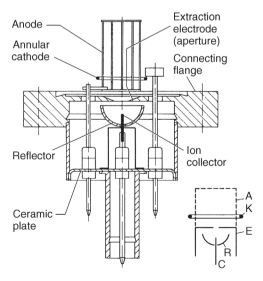


Figure 13.53 Extractor measuring system. Side image: A: anode (300 V); K: cathode (200 V); E: extraction electrode (0 V); R: reflector (290 V); C: ion collector (0 V).

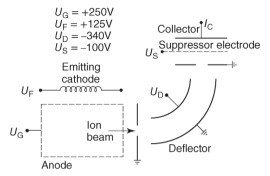


Figure 13.54 Ionization vacuum gauge according to Helmer [65].

Watanabe [67] refined Helmer's deflection principle by using the deflection unit as an energy filter. Ions produced at the anode mesh by electron-stimulated desorption carry somewhat higher energy than ions produced in the free space of the anode mesh. This is caused by the space charge of the electrons in the mesh.

In order to increase this effect of space charge, Watanabe used a spherical anode grid (Figure 13.55), which on average provides a higher electron density in the center of the sphere than at the edges. In Watanabe's ion spectroscopy gauge, the deflection unit is a hemispherical deflector with an inner electrode at ground potential and an outer, variable-potential positive electrode.

Figure 13.56 shows that this vacuum gauge is capable of differentiating between ions from the gas phase and ions produced by electron-stimulated desorption. Those parts of the ion spectroscopy gauge adjacent to the hot cathode are available for degassing by means of direct electrical heating or electron bombardment. The remaining parts of an ion spectroscopy gauge are

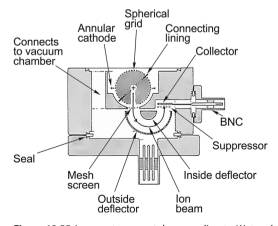


Figure 13.55 Ion spectroscopy tube according to Watanabe [67].

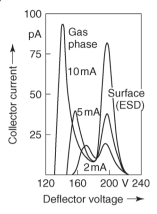


Figure 13.56 Energy spectrum of deflected ions in the ion spectroscopy tube shown in Figure 13.55.

manufactured from well-thermoconducting materials such as copper or aluminum so that heating and thus outgassing (of hydrogen, in particular) is prevented as far as possible. *Watanabe* specified a lower measuring limit of 2×10^{-12} Pa.

A simple and commercially available procedure for separating ESD ions from ions produced in the free space uses an energy analyzer referred to as *Bessel box* [68]. It has the advantage of using a straight, cylindrically symmetric arrangement allowing compact design of ionization gauges (Figure 13.57). The designers of this tube called their system AxTRAN gauge (axial symmetric transmission gauge) [69]. Depending on the voltage $U_{\rm BE}$, only ions with a particular energy level pass the *Bessel* analyzer and are detected at the secondary electron multiplier (SEM). ESD ions are thus suppressed by optimizing this voltage. The disk charged to cylinder potential and arranged in the center of the *Bessel* analyzer prevents direct line-of-sight contact for photons from the anode grid to the

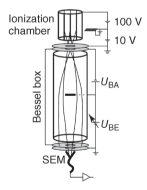


Figure 13.57 Diagram of the AXTRAN gauge, which suppresses ESD ions and X-rays by using a cylindrically symmetric energy analyzer (Bessel box). The image shows the

situation at a voltage $U_{\rm BE}$ that suppresses ESD ions from the anode mesh (see trajectories in the image).

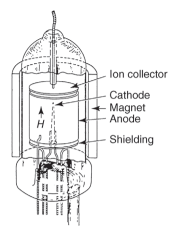


Figure 13.58 Lafferty gauge [71].

SEM, and thus suppresses the X-ray effect. The estimate for the lower measuring limit is 3×10^{-12} Pa (nitrogen equivalent) [74]. Manufacturer's data for the commercially available instrument specify 5×10^{-11} Pa [70].

13.7.3.7 Additional Types of Emitting-Cathode Ionization Gauges

Many other measuring systems in addition to those described in the previous sections have been developed, though, without having found noticeably broad applications.

Lafferty [71] built an ionization gauge with a hot cathode arranged in the axis of an anodic cylinder grid (Figure 13.58). A magnetic field, axial as well, forces the electrons to travel on a circular path, which increases their mean free path compared with a Bayard-Alpert tube by several orders of magnitude. Emission current was limited to 10 µA for stable operation. For this ionization gauge, the calculated X-ray limit was 2×10^{-12} Pa.

Further ionization gauges are only listed here in brief: Schuemann suppressor gauge [72], orbitron system (very long electron paths) [73], and Lafferty magnetron gauge with hot cathode [74].

These and other measuring instruments were mainly used for measuring extremely low pressures ($< 10^{-8} \text{ Pa}$) [75–82].

13.7.3.8 Operating Suggestions for Emitting-Cathode Ionization Gauges

From a vacuum-technological view, hot glow cathodes are chemical factories. Figure 13.59 shows important reaction channels on a hot, 2000 °C tungsten cathode. Water vapor and hydrocarbons in a vacuum system develop hydrogen, carbon, and oxygen. Additional carbon impurities are released via diffusion by the tungsten cathode. In this way, an ionization gauge can itself produce residual gas and contaminate the vacuum.

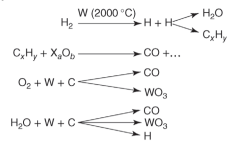


Figure 13.59 Reaction channels on a hot, 2000 °C tungsten cathode under high vacuum.

Emissive tungsten cathodes require protection against too high operating pressure (> 1 Pa). Otherwise, tungsten wires might burn out in oxygen-containing gases. Many control units include corresponding protective circuits, which shut off the cathode heater as soon as the adjusted measuring range or the maximum tolerable pressure is exceeded.

Degassing of measuring systems reduces the parasitic errors described here and in the previous sections (ESD, reactions occurring at the hot cathode, secondary electrons, etc.). Usually, engaging an appropriately adjusted press-button switch starts the degassing procedure. Then, electron bombardment cleans the anode and thus removes impurities (adsorbed layers). In some instruments, degassing uses a direct current through the anode for heating.

Features of commercial instruments include automatic measuring-range switches, digital displays for measured values and measuring ranges, and pressure-switch units with adjustable switching points. Apart from linear measuring ranges, many instruments can be equipped with a logarithmic measuring range. Certain applications require emission current to be adjusted stepwise, for example, from 0.1 to 1 and 10 mA.

Today, compact transmitter tubes deliver voltage amplitudes of 0-10 V without any additional operating equipment required. A variation of one decade in pressure produces a change of 1 V.

Often, ionization gauges are combined with measuring instruments for higher pressure ranges such as thermal conductivity gauges.

Caution: If an electrode with a potential > 100 V with respect to ground potential is used in an *ungrounded* metal vacuum vessel, and if the pressure in the container is greater than approximately 0.1 Pa, then grounding of the container can lead to a discharge inside the container. Pre-ionization in between the hot cathode and corresponding anode (e.g., in an ionization gauge) promotes ignition. If grounding is caused by contact with the human body (touching), then the touching person can suffer considerable injuries due to the electrical current passing through the body. Manufacturers attempt to reduce and prevent such damage as far as possible by designing their equipment appropriately. However, 100% safety requires the experimenter or user to take into account the mentioned facts and to appropriately build and operate the equipment [83].

13.7.4

Crossed-Field Ionization Gauges

13.7.4.1 Penning Gauges

The operation principle of these gauges for low pressures uses a gas discharge ignited between two metal electrodes (anode, cathode) by applying sufficiently high DC voltage (in the kilovolt range). The gas discharge current is pressure dependent and thus used as measured quantity. However, if no additional measures are taken, the lower measuring limit is only about 1 Pa. At lower pressures, the number of carriers produced is insufficient for sustaining the gas discharge. The so-called *Penning* discharge maintains gas discharge even down to very low pressures. For this, it uses a sufficiently powerful magnetic field arranged so that the electron paths from the cathode to the anode are stretched considerably by forcing the electrons onto spiral paths. This leads to higher ion yield. Due to their higher mass, ions are hardly distracted by the magnetic field and travel directly to the cathode. Secondary electrons, produced when the ions impinge on the cathode, nourish the discharge. Within a broad range, discharge current I is a measure of pressure p:

$$p = KI^m. (13.44)$$

The exponent m depends on the precise design of the gauge and is in the range m = 1 - 1.4.

For a detailed discussion of the operating principle in a *Penning* gauge, please consider Figures 13.60 and 13.61: a DC voltage of, for example, 3 kV produced in high-voltage generator HV is applied to the annular anode AA of the Penning instrument P via the protective resistor R. The Penning tube has a grounded metal housing H. The two plane walls parallel to the annular anode AA form the cathodes C. The magnetic field $\rightarrow B$ is applied such that its field lines from one cathode to the other pass through the anode. Ammeter A measures the discharge current I. Measures must be taken in order to prevent isolation currents between the anode and the cathode because the gauge would indicate these too.

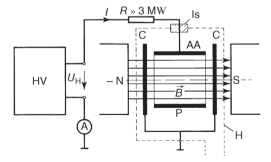


Figure 13.60 Diagram of a Penning gauge. AA: annular anode; C: cathode plates; H: isolating wall of housing; N, S: magnet pole pieces; HV: high-voltage supply; $U_{\rm H} \approx 3000 \, \rm V$; $B = 0.1 - 0.2 \, \rm T$.

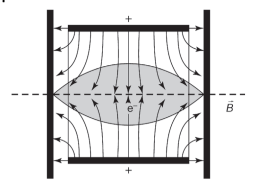


Figure 13.61 Diagram showing the orientations (not the strengths) of the electrical field in a *Penning* gauge according to Figure 13.60. Shaded area: electron space charge.

Among other effects, the protective resistor R (several $M\Omega$) also serves to limit discharge current particularly at high pressures.

Figure 13.62 shows a typical calibration curve for a *Penning* gauge in the range of 10^{-8} to 1 mbar (10^{-6} to 100 Pa). As indicated in Figure 13.62, two different discharge mechanisms occur with a transition at about 10^{-4} mbar (10^{-2} Pa). The discharge is characterized by a negative ring current at lower pressure, and by plasma at higher pressure. In both cases, the magnetic field ($B \approx 0.1-0.2$ T) predominantly influences discharge because it strongly impedes motion of the electrons perpendicular to the field lines, that is, to the anode (compare Figure 13.60). According to *Knauer* [84,85] (compare Figure 13.63 giving typical values), at low pressures, a rotating electron volume charge concentric to the axis of the anode cylinder develops, that is, a ring current of approximately 1 A (shaded area in Figure 13.61). A strong electrical field develops between the anode and this space

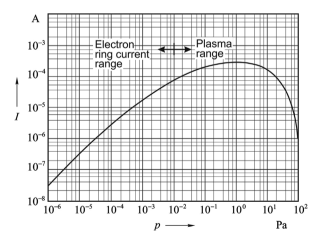


Figure 13.62 Typical calibration curve for a *Penning* gauge. *I*: discharge current; *p*: pressure. In the linear range at low pressures, sensitivity $K \approx 3$ A mbar⁻¹. Transition between annular electron current and plasma at $p = 10^{-2}$ Pa (10^{-4} mbar).

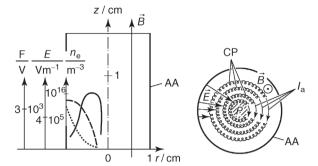


Figure 13.63 Mechanisms in a Penning discharge at low pressures ($p < 10^{-4}$ mbar) according to *Knauer* [84,85]. AA: anode; \rightarrow B: magnetic field; \rightarrow E: electrical field; I_a : annular electron current (approximately 1 A); CP: cycloid paths of electrons; dashed: E electrical

field strength; dotted: Φ potential; n_e : electron density versus r(cylindrical coordinates, r = 1radius anode cylinder, z along cylinder axis in cm); typical values. P: plasma. Pitch circle radius of electron cycloid paths is about 0.1 mm.

charge, in which almost the entire voltage applied to the electrodes in the range of 3 kV falls off. The space charge considerably enhances the radial component of the electrical field (Figure 13.61).

Within the space charge ring, a plasma P develops with homogeneously distributed, small number densities of ions and electrons. Its potential is only a few hundred volts above cathode potential; it is of minor importance for the discharge mechanism. Electrons in the ring current, influenced by the crossed fields, follow cycloid paths and approach the anode stepwise only due to collisions with gas atoms. Thus, the magnetic field heavily impedes electron diffusion to the outside. The ring current would persist constantly if the volume were gasfree. Collisions with gas atoms in the volume, apart from causing diffusion as described above, can also cause ionizing. An ionizing collision produces another electron and an ion. The electron merges into the volume charge cloud, that is, the ring current, whereas field E accelerates the ion toward the inside. In contrast to the electrons with trajectory radii in the area of 0.1 mm for combined electrical and magnetic fields, the radius of ion trajectories is higher by a factor determined by the ratio of ion mass to electron mass, and thus in the range of meters. Therefore, ions reach the cathode guickly. Both the number of ions produced per unit time (ionizing rate) and the diffusion coefficient perpendicular to the magnetic field are proportional to gas density n. Thus, an equilibrium between carrier production (by ionizing) and carrier depletion (by diffusion) can develop in the ring current such that the electron density in the ring current I_R , and thus I_R itself, remains constant and nearly independent of n. Then, the outside current is proportional to *n* throughout the entire pressure range $(p < 10^{-2})$ Pa), and thus proportional to pressure p. The fact that the ring current, that is, the axial space charge, indeed slightly follows n leads to m > 1 in Eq. (13.44).

The ring current in a *Penning* system has the same function as the electron current in an emitting-cathode ionization gauge. However, sensitivity is much higher because the electron current is higher. Typical Penning system sensitivities are in the range 0.02-0.05 A Pa⁻¹, for pressures below 0.01 Pa. *Penning* cells for ion getter pumps (Section 11.4.1) can be optimized to 0.1 eA Pa⁻¹.

As the pressure drops, sensitivity is maintained as long as the ring current continues. Obviously, the ring current depends on an additional supply of electrons released from the cathodes. Otherwise, electrons are depleted from the ring current, which ultimately ceases as the pressure continues to drop. Depending on geometry as well as magnetic and electrical field strengths, such discharges are maintained down to pressures far below 10⁻⁹ Pa.

Ion getter pumps operate on the same principle. At low pressures, their pumping speed also requires maintaining a continuous discharge.

At pressures $p > 10^{-2}$ to 10^{-1} Pa, the density of positive carriers in the ring current rises to levels at which the discharge type described above is no longer stable. The entire anode cylinder is filled with more or less equipotential plasma, which now determines the discharge mechanism. Its potential lies between those of cathode and anode. It is separated from both electrodes by space charge layers. In the cathode layer, ions from the plasma accelerate toward the cathode where they release electrons. Thus, the ions resupply electrons, that is, ionizing carriers, to the system, replenishing such electrons that became depleted from the plasma and trapped by the anode due to diffusion perpendicular to the magnetic field. In this mechanism, fluctuations (plasma oscillations) promote such diffusion. Insight into these diffusion-promoting processes is necessary for understanding the high currents observed in such discharges [86]. In the considered pressure range, discharge current and pressure are no longer proportional (Figure 13.62). At higher pressures, readings on such a Penning gauge depend on the condition of the cathode surfaces because electron release during ion collisions is influenced by the state of the cathode surface. Thus, frequent cleaning of the cathodes can be advisable in certain cases. For easier cleaning, cathodes are often made of two thin, replaceable stainless-steel sheets.

The main advantage of Penning gauges is that the discharge current in a Penning system is high enough for current measurements, that is, pressure measurements, down to 10^{-4} Pa without using an amplifier if a sensitive ammeter is employed (see also Figure 13.62). Thus, Penning gauges are cheaper and more trouble-free than emitting-cathode ionization gauges. Utilizing Penning gauges is advisable for simple control tasks if the pressure in a vacuum system is to be checked rather than precisely measured. Figure 13.64 shows a measuring tube. The anode and the cathode plates can be pulled out for cleaning purposes. In order to counteract the contamination issue described above, a baffle as shown in Figure 13.65 should protect the *Penning* gauge.

The measuring range of a *Penning* system as shown in Figure 13.62 can be enhanced for measuring lower pressures. For this, an additional amplifier, preferably having logarithmic characteristic, is added to the supply unit.

As a result, the measuring scale is approximately logarithmic throughout a pressure range of 10^{-7} to 10^{-2} Pa. Penning gauges are particularly useful when



Figure 13.64 Penning gauge. (Courtesy of INFICON Inc.)

combined with a thermal conductivity vacuum gauge. The benefit is that the thermal conductivity gauge controls the *Penning* metering point such that the Penning gauge is activated if the pressure drops below 0.5 Pa, and shut off if the pressure exceeds this threshold value. This yields a continuous pressure scale from 10⁻⁴ to 100 kPa. Furthermore, the arrangement prevents operator's errors with the Penning gauge. Such errors occur when the Penning system operates at pressures above 1 Pa, as shown in the calibration curve in Figure 13.62, because discharge current drops in this pressure range, and thus, readings are ambiguous.

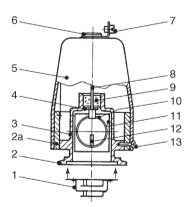


Figure 13.65 Section of a Penning vacuum gauge. (1) Pulled out vapor trap with centering, (2) small flange, vacuum connection, (2a) permanent magnet, (3) housing, (4) protective plate for isolator (9), (5) sealing lid, (6)

connector for high-voltage operating power, (7) ground connector, (8) anode connector, (9) compression glass seal, (10) annular anode, (11) cathode plate, (12) ignition pin, and (13) connecting bolt for sealing (5).

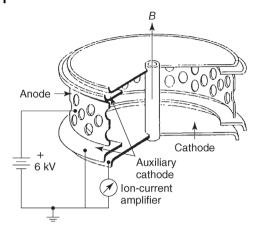


Figure 13.66 Diagram of Redhead's magnetron [87].

13.7.4.2 Magnetron and Inverted Magnetron

In order to stabilize the discharge and improve starting behavior, Redhead developed the magnetron [87] and Hobson and Redhead designed the inverted magnetron [88].

In a magnetron, the anode is an open cylinder. The cathode forms axis and both end plates of the cylinder (Figure 13.66). In the inverted magnetron (Figure 13.67), the anode is a rod on the axis of a nearly closed cylinder serving as cathode. In the magnetron, two annular rings at cathode potential shield the end disks of the cathode from the high electrical fields. The ion

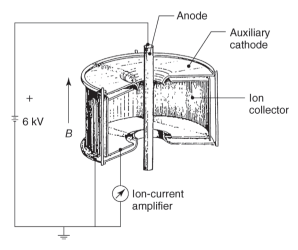


Figure 13.67 Inverted magnetron according to Hobson and Redhead [88].

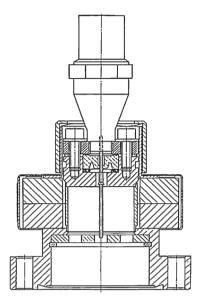


Figure 13.68 IKR020 inverted magnetron by INFICON. (Courtesy of INFICON Inc.)

current amplifier does not detect any possible field emission currents produced by the rings.

In inverted magnetrons, guard rings prevent field emission currents between the cathode and the anode. The magnetic field is parallel to the anode. This ionization gauge can be operated at up to 6 kV and 0.2 T.

Both types of gauges with crossed electromagnetic fields trap electrons more efficiently than the *Penning* type. This improves starting conditions, interrelationships of p, B, and V follow theoretical predictions more accurately, and the discharge is stable down to much lower pressures. Redhead and Hobson specified a usable pressure range of 10^{-11} to 10^{-2} Pa for their ionization gauges.

Equation (13.44) applies to these ionization gauges, too. However, m is pressure dependent, which makes measurements complicated. Generally, m is higher at high pressures than at low pressures. Therefore, at low pressures, an ionization gauge produces too low pressure readings if the meter uses extrapolation of p(I) or m from higher to lower pressures. Thus, at readings of 10^{-10} Pa, an error of one order of magnitude can be expected.

Figure 13.68 shows the design of a modern inverted magnetron.

Comparison of Both Types of Ionization Gauges

On the one hand, crossed-field ionization gauges have the general advantage that they show no metering-relevant X-ray limit and electron-stimulated desorption

Туре	Gas species	Pumping speed (ℓ s ⁻¹)	Reference
Penning	Air	0.25	[89]
Magnetron	Helium	0.17	[90]
	Nitrogen	0.25	
Magnetron	Helium	0.2	[91]
	Nitrogen	0.14	
	Oxygen	0.15	
Inverted magnetron	Helium	0.03	[92,93]

Table 13.4 Published pumping speeds for selected ionization gauges with crossed electromagnetic fields.

is negligible. Due to their strong, intense magnetic field, they react less sensitively to outside magnetic fields (e.g., fields from bending magnets in accelerators) compared with emitting-cathode ionization gauges. Furthermore, outside magnetic fields can even be used for crossed-field ionization gauges.

On the other hand, the internal magnetic field of crossed-field ionization gauges may be disturbing, for example, in electron optics in electron microscopes, and thus might require appropriate shielding.

Crossed-field ionization gauges show four general disadvantages:

- The measuring signal (discharge current) does not show strictly linear pressure dependence.
- High negative electron volume charge easily leads to dynamic discharge instabilities (high-frequency oscillations, mode jumping).
- They are sensitive toward contamination in industrial processes.
- They pump. Pumping speed here is usually one to two orders of magnitude higher than that in emitting-cathode ionization gauges (Table 13.4).

The latter means that the measuring instrument itself influences the measured quantity.

The problem appearing at low pressures that crossed-field ionization gauges might not start or only start with high time delay is solved in most of today's magnetrons or inverted magnetrons, for example, by including a weak radioactive source.

Emitting-cathode ionization gauges are preferred when accurate measurements (measuring uncertainty less than 10%) are desired. They can be used as reference standards because their measuring-signal linearity with pressure is better. The gauge sensitivity is obtained by comparing it with a spinning rotor gauge at pressures between 10^{-3} and 10^{-2} Pa.

However, under clean vacuum conditions, magnetrons and inverted magnetrons have proven successful as reference standards as well because their characteristic I(p) is reproducible and long-time stable [94]. For

Table 13.5 Comparison between ionization vacuum gauges with emitting cathode and with
crossed electromagnetic fields.

Criterion	lonization vacuum gauges with crossed electromagnetic fields	lonization gauges with emitting cathode
Errors due to pumping speed	High	Low
Accuracy, stability	Moderate	Good
Size, mechanical stability	Good	Moderate
Sensitivity toward external magnetic fields	Low	High
Produces outside mag- netic field	Yes	No
Susceptibility to contamination	High	Moderate
Startup problems	Occur	None
X-ray limit	None	10^{-10} to 10^{-6} Pa
Electron-stimulated desorption	Negligible	Yes, gas species dependent
Price	Low	High

hydrogen as calibration gas, they even seem more appropriate than emittingcathode ionization gauges.

Lower susceptibility to contamination is an additional benefit of emittingcathode ionization gauges. This is particularly important in semiconductor manufacturing. Electron bombardment and direct current passage provide effective cleaning procedures for the basic parts of ionization gauges. High emission currents during operation enhance the self-cleaning effect.

Table 13.5 summarizes advantages and disadvantages of both types of ionization gauges.

13.7.6

General Suggestions

Readings of all ionization gauges depend on the gas species. If the gas species and gas composition are known, gas-type dependence can be taken into account by multiplying the pressure reading with the appropriate correction factor listed in Table 13.6. These factors were first gathered for emitting-cathode ionization gauges. However, within measuring accuracy, the factors are valid for Penning gauges as well.

Gas species	CF _i (N ₂)	Gas species	$CF_i(N_2)$
N_2	1.00	СО	0.97
Не	7.24	CO_2	0.70
Ne	4.55	I	0.17
Ar	0.85	CH_4	0.71
Kr	0.59	C_2H_6	0.37
Xe	0.41	C_3H_8	0.22
H_2	2.49	CF_2Cl_2	0.36
O_2	1.07	Oil vapors	0.1
Air	1.02		

Table 13.6 Correction factors CF for taking into account the influence of gas species on the indicated pressure p_{ind} in ionization vacuum gauges calibrated to nitrogen, $p = p_{ind}$ CF.

For mixed gases with relative amount-of-substance fractions χ_i , $1/\text{CF} = \sum_i \chi_i (1/\text{CF}_i)$. Deviations of up to 10% from listed CF values may occur.

Example 13.1

The pressure of a 70% argon and 30% helium gas mixture at 23 °C is 1×10^{-2} Pa. How high is the pressure reading in a Bayard-Alpert system if the scale is calibrated to nitrogen? The gas consists of 7×10^{-3} Pa argon and 3×10^{-3} Pa helium. For argon, the ionization gauge shows $7 \times 10^{-3} / 0.85 \, \text{Pa} = 8.4 \times 10^{-3} \, \text{Pa}$, and for helium, $3 \text{ to } 10^{-3} / 7.24 \text{ Pa} = 4.1 \times 10^{-4} \text{ Pa}$, and thus, a total of $8.8 \times 10^{-3} \text{ Pa.}$

The accuracy of the values in Table 13.6 should be considered carefully. The differential ionization probability leading to these values depends on the anode voltage. Additional dependences specific to the instrument type include ioncapturing probability, dissociation effects, and secondary-electron production. True values may differ from those in Table 13.6 by up to 10%. For higher accuracy, it is advisable to calibrate the gauge to the considered gas species.

Figure 13.69 shows one example of occurring effects. The plot contains the ratio of the gauge sensitivities for hydrogen and deuterium [95]. For the electron energy in an ionization gauge, the electron structure is the same for hydrogen and deuterium. Thus, the ratio of gauge sensitivities should be expected to be equal to 1. Obviously, this only applies approximately to certain cases. Furthermore, the ratio even changes for a single tube depending on its previous history. This is in fact quite surprising because neither geometry nor voltages are changed.

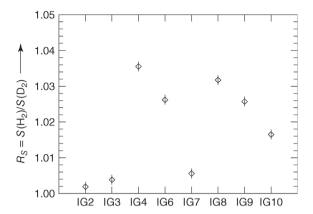


Figure 13.69 Ratios of gauge sensitivities for hydrogen/deuterium for selected emittingcathode ionization gauges [95].

The reason for the deviations in sensitivity for hydrogen and deuterium is that hydrogen, due to its lower mass and higher velocity, produces more secondary electrons than deuterium in the same electrical field. Electron yield at the cathode depends strongly on its surface condition, which explains why the ratio $S(H_2)/S(D_2)$ can be subject to change. If secondary electrons are suppressed entirely, the ratio is equal to 1 [95].

Gas-type dependence is a problem in combinations of thermal conductivity gauges and ionization gauges. In the presence of hydrogen, as Table 13.6, Section 13.5, and Figure 13.30 show, gas-type dependences in both measuring principles act in opposite directions producing too high readings on thermal conductivity gauges and too low readings on the ionization gauge. In contrast, with a gas showing higher relative molecular mass M_r than nitrogen, ionization gauges usually indicate too high values compared with the true pressure, whereas readings of thermal conductivity gauges are too low. This circumstance should be taken into account in the transition range in such combination pressure gauges.

An additional problem of ionization gauges is that they measure gas density n instead of pressure p. We will consider a sealed chamber at room temperature in high vacuum, filled with hydrogen of pressure p_1 . The chamber includes an ionization gauge for pressure measuring. If the entire chamber is submersed in liquid nitrogen, pressure p_2 will drop according to

$$\frac{p_2}{p_1} = \frac{T_2}{T_1} = \frac{77}{300} = 0.26,\tag{13.45}$$

and thus to about 1/4 of p_1 . However, the reading on the ionization gauge will remain unchanged because density $n_1 = n_2$ (Eq. (3.19)). This example shows how important it is to take into account the temperature when measuring pressure. Precision measurements in fact require respecting temperature changes within the room-temperature range (> 1 K) [96].

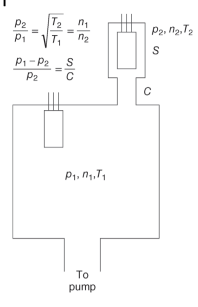


Figure 13.70 Influence of pumping speed S in (pressure p_1 , temperature T_1) and ionization an ionization gauge (see Eg. (4.33)) and thermal transpiration on the indicated pressure. C: the ionization gauge measures gas density n2. conductance between vacuum vessel

gauge. The reading drops if $T_2 > T_1$ because

When a chamber is evacuated continuously, molecular flow develops in an emitting-cathode ionization gauge's measuring head with internal temperature T₂, and thus, the equation of continuity applies. This means that thermal transpiration (Section 4.6) occurs if chamber and gauge are connected via a thin tube (Figure 13.70). If p_1 denotes the gas pressure in the chamber at temperature T_1 ,

$$\frac{p_1}{p_2} = \sqrt{\frac{T_1}{T_2}}. (13.46)$$

Pressure p_2 in the emitting-cathode ionization gauge thus rises (Figure 13.70), whereas the indicated pressure follows n_2 and drops.

$$\frac{n_2}{n_1} = \sqrt{\frac{T_1}{T_2}}. (13.47)$$

Usually, the effect of thermal transpiration is taken into account during calibration of the gauge sensitivity in emitting-cathode ionization gauges.

Figure 13.70 also explains the influence of pumping speed in an ionization gauge. If the instrument is not immersed like the lower gauge in Figure 13.70, and a finite conductance C is present between the chamber and the gauge, the indicated pressure p_2 is reduced as well.

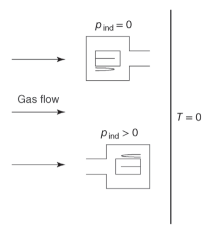


Figure 13.71 Influence of the orientation of an upper ionization gauge indicates pressure ionization gauge on the indicated pressure. The wall to the right shall be assumed to pump ideally at absolute zero (T = 0). The

p = 0, and the lower ionization gauge indicates a pressure unequal to zero.

Ionization gauges experience an additional problem appearing when the pressure distribution in a container is inhomogeneous, that is, when net flow occurs in the chamber. Figure 13.71 shows that, in certain cases, the orientation of the ionization gauge to the gas flow can be influential. Obviously, this applies to any type of gauge. Any open vacuum system (path to the pump is not sealed) shows net gas flow.

In Figure 13.71, gas flows from left to right toward a wall with a temperature at which all gas particles condense. The upper ideal gauge (without internal gas sources) will indicate zero pressure, whereas the same gauge in the bottom position will indicate a finite pressure. This pressure is the equilibrium pressure established when the rates of gas particles entering and leaving the measuring head are equal. None of the two positions actually indicate the true pressure.

Outgassing and reemission of particles previously pumped by the ionization gauge can present a severe problem at low pressures. An ionization gauge exposed to higher pressure for long time will show considerable relaxation times of hours or even days before low-pressure readings stabilize at constant values. Outgassing rates of emitting-cathode ionization gauges typically vary between 10^{-9} and 10^{-7} Pa ℓ s⁻¹ [67].

13.8 **Combined Vacuum Gauges**

Operators of vacuum systems like to have a compact vacuum gauge that can cover the whole vacuum range from high vacuum to atmospheric pressure.



Figure 13.72 Combined vacuum gauge BCG 450 by *INFICON* with a range from 5×10^{-8} Pa to 150 kPa. It is composed of a capacitive diaphragm sensor (left), a Pirani gauge (right),

and an ionization gauge of the Bayard–Alpert type (center part). (Courtesy of *INFICON AG*, Liechtenstein.)

For this need, manufacturers developed vacuum gauges that combine different measurement principles in one gauge. Figure 13.47 already gave us an example where a fine vacuum ionization gauge was combined with a Pirani gauge. Common is also the combination of Bayard–Alpert ionization gauge with Pirani gauge. Also quartz oscillators have been combined with ionization gauges.

A very compact vacuum gauge, where three different measurement principles were combined, is depicted in Figure 13.72. In the range from 150 to $1\,\mathrm{kPa}$, the pressure is measured to be gas species independent by a capacitive diaphragm gauge, from $1\,\mathrm{kPa}$ to $2\,\mathrm{Pa}$ the pressure is measured by a Pirani gauge, the reading, however, being dependent on the gas species, as this is the case for the third principle, a Bayard–Alpert ionization gauge with a range down to $10^{-8}\,\mathrm{Pa}$. In the overlapping ranges of the measurement principles, from $100\,\mathrm{Pa}$ to $1\,\mathrm{kPa}$ and from 0.5 to $2\,\mathrm{Pa}$, electronic circuits smooth out eventual jumps between the indications obtained from the different measurement principles. These jumps are caused partly by the different gas species dependences of the measurement principles, partly by insufficient

calibration, and partly by long-term instabilities of the different gauges. These jumps are a significant problem for combined vacuum gauges, so that their accuracy is rather modest.

References

- 1 Wolf, A. (1946) An elementary theory of the Bourdon gage. J. Appl. Mech., 68, 207 - 210.
- 2 Jennings, F.B. (1956) Theories on Bourdon tubes. Trans. ASME, 78, 55-64.
- 3 Olson, A.R. and Hurst, I.L. (1929) A new differential pressure gage. J. Chem. Phys., **51** (8), 2378.
- 4 Bonfig, K.W. (1988) Technische Druckund Kraftmessung.
- 5 Harada, K. (1999) Various applications of resonant pressure sensor chip based on 3-D micromachining. Sens. Actuators A, 73, 261–266. See also Refs. [6,7] herein.
- 6 Miiller, A. et al. (2002) Final report on key comparison CCM.P-K4 of absolute pressure standards from 1 Pa to 1,000 Pa. Metrologia, 39 (Tech. Suppl.), 07001.
- 7 Redhead, P.A. (1984) The measurement of vacuum pressures. J. Vac. Sci. Technol. A, 2, 132-138.
- 8 Arai, K., Yoshida, H., and Kobata, T. (2014) Measurement of volume change induced by a bulging of a diaphragm inside a differential type capacitance diaphragm gauge. Measurement, 48, 149-154.
- 9 Sharma, J.K.N., Sharma, D.R., and Gupta, A.C. (1991) Zero shift in the differential capacitance diaphragm gauges due to the change in line pressure. J. Vac. Sci. Technol. A, 9, 2389-2393.
- 10 Takaishi, T. and Sensui, Y. (1963) Thermal transpiration effect of hydrogen, rare gases and methane. Trans. Faraday Soc., 59, 2503-2514.
- 11 Liang, S.C. (1951) Some measurements of thermal transpiration. J. Appl. Phys., 22, 148 - 153.
- 12 Poulter, K.F. et al. (1983) Thermal transpiration correction in capacitance manometers. Vacuum, 33, 311-316.
- 13 Setina, J. (1999) New approach to corrections for thermal transpirationeffects

- in capacitance diaphragm gauges. Metrologia, 36, 623-626.
- 14 Jitschin, W. and Röhl, P. (1987) Quantitative study of the thermal transpiration effect in vacuum gauges. J. Vac. Sci. Technol. A, 5, 372-375.
- 15 Meyer, O.E. (1865) Über die innere Reibung der Gase. Pogg. Ann., 125, 177.
- 16 Maxwell, J.C. (1866) The Bakerian Lecture: on the viscosity or internal friction of air and other gases. Philos. Trans. R. Soc., 156, 249.
- 17 Kundt, A. and Warburg, E. (1875) Über Reibung und Wärmeleitung verdünnter Gase. Pogg. Ann., 155, 337.
- 18 Sutherland, W. (1896) On thermal transpiration and radiometer motion. Philos. Mag. Ser. 5, 42, 373.
- 19 Hogg, J.L. (1906) Friction and force due to transpiration as dependent on pressure in gases. Proc. Am. Acad. Arts Sci., 42, 113.
- 20 Holmes, F.T. (1937) Axial magnetic suspensions. Rev. Sci. Instrum., 8, 444.
- 21 Beams, J.W., Young, J.L., and Moore, J.W. (1946) The production of high centrifugal fields. J. Appl. Phys., 17, 886.
- 22 Beams, J.W., Spitzer, D.M., Jr., and Wade, J.P., Jr. (1962) Spinning rotor pressure gauge. Rev. Sci. Instrum., 33, 151.
- 23 Fremerey, J.K. (1973) Cavity type permanent magnet suspension. Rev. Sci. Instrum., 44, 1396-1397.
- 24 Fremerey, J.K. (1983) Gas friction vacuum meter and method of making measuring body US Patent 4395914.
- 25 Fremerey, J.K. (1982) Spinning rotor vacuum gauges. Vacuum, 32, 685-690.
- 26 Fremerey, J.K. (1985) The spinning rotor gauge. J. Vac. Sci. Technol. A, 3, 1715-1720.
- 27 Weller, A. (1983) Temperature determination of freely rotating bodies. Rev. Sci. Instrum., 54, 952-957.

- 28 Fremerey, J.K. (1980) Theoretical model for calculation of molecular drag on tangentially moving rough surfaces. Proceedings of the 8th International Vacuum Congress, 4th ICSS, and 3rd ECOSS, Cannes, vol. II, pp. 869-872.
- 29 Lindenau, B.E. and Fremerey, J.K. (1991) Linearization and temperature compensation up to one atmosphere for the SRG. J. Vac. Sci. Technol. A, 9, 2737-2743.
- 30 Setina, J. and Looney, J.P. (1993) Behaviour of commercial SRGs in the transition regime. Vacuum, 44, 577-580.
- 31 Dittmann, S., Lindenau, B., and Tilford, C.R. (1989) The molecular drag gauge as a calibration standard. J. Vac. Sci. Technol. A. 7, 3356-3360.
- 32 Röhl, P. and Jitschin, W. (1988) Performance of the spinning rotor gauge with a novel transport device as a transfer standard for high vacuum. Vacuum, 38, 507.
- 33 Ono, M. et al. (1985) Design and performance of a quartz oscillator vacuum gauge with a controller. J. Vac. Sci. Technol. A, 3, 1746.
- 34 Hirata, M., Kokobun, K., Ono, M., and Nakayama, K. (1985) Size effect of a quartz oscillator on its characteristics as a friction vacuum gauge. J. Vac. Sci. Technol. A, 3, 1742-1745.
- 35 Pirani, M. (1906) Selbstzeigendes Vakuum-Meßinstrument. Dtsch. Phys. Ges. Verh., 8, 686.
- 36 Leck, J.H. (1989) Total and Partial Pressure Measurement in Vacuum Systems, Blackie, pp. 49 ff.
- 37 Jousten, K. (2008) On the gas species dependence of Pirani vacuum gauges. J. Vac. Sci. Technol. A, 26, 352 - 359.
- 38 Hinkle, L.D. and Mariano, C.F. (1991) Toward understanding the fundamental mechanism and properties of the thermal mass flow controller. J. Vac. Sci. Technol. A, 9, 2043–2047.
- 39 Tison, S.A. (1996) A critical evaluation of thermal mass flow meters. J. Vac. Sci. Technol. A, 14, 2582-2591.
- 40 von Baeyer, O. (1909) Phys. Z., 10, 168.
- 41 Buckley, O.E. (1916) An ionization manometer. Proc. Natl. Acad. Sci. USA, 2, 683.

- 42 Dushman, S. and Lafferty, J.M. (1962) Scientific Foundations of Vacuum Technique, 2nd edn, John Wiley & Sons,
- 43 Penning, F.M. (1937) Physica, 4, 71. Penning, F.M. (1937) Ein neues manometer für niedrige Gasdrücke, insbesondere zwischen 10-3 und 10-5 mm. Philips Tech. Rev., 2, 201.
- 44 Penning, F.M. (1936) Die Glimmentladung bei niedrigem Druck zwischen koaxialen Zylindern in einem axialen Magnetfeld. Physica, 3, 873. Penning, F.M. (1939) US Patent US2197079 A.
- 45 Penning, F.M. and Nienhuis, K. (1949) Construction and application of a new design of the Philips vacuum gauge. Philips Tech. Rev., 11, 116.
- 46 Tilford, C.R. (1985) Sensitivity of hot cathode ionization gauges. J. Vac. Sci. Technol. A, 3, 546-549.
- 47 Edelmann, C. and Engelmann, P. (1982) Möglichkeiten der Messbereichserweiterung bei Glühkatoden-Ionisationsvakuummetern. Vak.-Tech., 31, 2-10.
- 48 Kno, Z.H. (1981) An approach to the nonlinearity of an ionization vacuum gauge at the upper limit of the measured pressure. Vacuum, 31 (7), 303-308.
- 49 Wang, Y.-Z. (1984) A fundamental theory of high pressure hot cathode ionization gauges. Vacuum, 34, 775-778.
- 50 Schulz, G.J. and Phleps, A.V. (1957) Ionization gauges for measuring pressures up to the millimeter range. Rev. Sci. Instrum., 28, 1051.
- **51** Bayard, R.T. and Alpert, D. (1950) Extension of the low pressure range of the ionization gauge. Rev. Sci. Instrum., 21,
- 52 Arnold, P.C., Bills, D.G., Borenstein, M.D., and Borichevsky, S.C. (1994) Stable and reproducible Bayard-Alpert ionization gauge. J. Vac. Sci. Technol. A, 12,
- 53 Schmidt, K. and Bergner, U. (1996) Stabilität von Hochvakuum-Meßröhren. Vak. Forsch. Prax., 8, 177-182.
- 54 Peacock, R.N. and Peacock, N.T. (1990) Sensitivity variation of Bayard-Alpert gauges with and without closed grids from

- 10-4 to 1 Pa. J. Vac. Sci. Technol. A, 8,
- 55 Van Oostrom, A. (1961) Vacuum Symposium Transactions, vol. 1, Pergamon Press, New York, 443 pp.
- 56 Repa, P. (1986) The residual current of the modulated BA-gauge. Vacuum, 36, 559-560.
- 57 Chou, T.S. and Tang, Z.Q. (1986) Investigation on the low pressure limit of the Bayard-Alpert gauge. J. Vac. Sci. Technol. A, 4, 2280-2283.
- 58 Filipelli, A.R. (1987) Residual currents in several commercial ultra high Bayard-Alpert gauges. J. Vac. Sci. Technol. A, 5, 3234-3241.
- 59 Berman, A. (1985) Total Pressure Measurements in Vacuum Technology, Academic Press.
- 60 Grosse, G. et al. (1987) Secondary electrons in ion gauges. J. Vac. Sci. Technol. A, 5, 3242.
- 61 Harten, U. et al. (1988) Surface effects on the stability of hot cathode ionization gauges. Vacuum, 38, 167-169.
- 62 Redhead, P.A. (1962) Electron stimulated desorption. Vacuum, 12, 267.
- 63 Redhead, P.A. (1960) Modulated Bayard-Alpert gauge. Rev. Sci. Instrum., 31, 343.
- 64 Hobson, J.P. (1964) Measurements with a modulated Bayard-Alpert gauge in aluminosilicate glass at pressures below 10⁻¹² Torr. J. Vac. Sci. Technol. A, 1, 1.
- 65 Helmer, J.C. and Hayward, W.D. (1966) Ion gauge for vacuum pressure measurements below 1×10^{-10} torr. Rev. Sci. Instrum., 37, 1652.
- 66 Han, S.-W. et al. (1988) Performance of the bent beam ionization gauge in ultrahigh vacuum measurements. Vacuum, 38, 1079-1082.
- 67 Watanabe, F. (1992) Ion spectroscopy gauge: total pressure measurements down to 10⁻¹² Pa with discrimination against electron-stimulated-desorption ions. J. Vac. Sci. Technol. A, 10, 3333-3339.
- 68 Craig, J.H. and Hock, J.H. (1980) Construction and performance characteristics of a low cost energy prefilter. J. Vac. Sci. Technol., 17, 1360-1363.

- 69 Akimichi, H. et al. (1995) Development of a new ionization gauge with Bessel box type energy analyser. Vacuum, 46, 749-752.
- 70 ULVAC Cooperation catalog, Japan, January 2006. Available at http://www. ulvac.co.jp/eng/.
- 71 Lafferty, J.M. (1961) Hot-cathode magnetron ionization gauge for the measurement of ultrahigh vacua. J. Appl. Phys., 32, 424.
- 72 Schuemann, W.C. (1963) Ionization vacuum gauge with photocurrent suppression. Rev. Sci. Instrum., 34,
- 73 Messer, G. (1980) Long term stability of various reference gauges over a three years period. Proceedings of the 8th International Vacuum Congress, Cannes, Vide Suppl. 201, pp. 191–194.
- 74 Lafferty, J.M. (1960) Transactions of the American Vacuum Society Symposium, vol. 7, p. 97.
- 75 Chen, J.Z. et al. (1982) An axial-emission ultra-high vacuum gauge. J. Vac. Sci. Technol., 20, 88-91.
- 76 Chen, J.Z. et al. (1983) Proceedings of the 9th International Vacuum Congress, Madrid, p. 99.
- 77 Ohsako, N. (1982) A new wide-range B-A gauge from UHV to 10-1 Torr. J. Vac. Sci. Technol., 20, 1153-1155.
- 78 Watanabe, F. (1987) Point collector ionization gauge with spherical grid for measuring pressures below 10⁻¹¹ Pa. J. Vac. Sci. Technol. A, 5, 242-248.
- 79 Gentsch, H. (1987) Inertes Ionisationsvakuummeter mit extrahierendem Kollektor (EXKOLL). Vak.-Tech., 36 (3), 67-74.
- 80 Redhead, P.A. (1987) Ultrahigh vacuum pressure measurements: limiting processes. J. Vac. Sci. Technol. A, 5, 3215-3223.
- 81 Madey, T.E. (1987) Surface phenomena and their influence on ultrahigh vacuum gauges. J. Vac. Sci. Technol. A, 5, 3249 (summary abstract).
- 82 Oshima, C. and Otuko, A. (1994) Performance of an ionization gauge with a large-angle ion deflector. I. Total pressure measurement in extreme high vacuum. J. Vac. Sci. Technol. A, 12, 3233.

- 83 Morrison, D. (1986) Lethal voltages from ion/gas discharge interactions. Le Vide, 41, 297-304.
- 84 Knauer, W. (1962) Mechanism of the penning discharge at low pressures. J. Appl. Phys., 33, 2093.
- 85 Knauer, W. et al. (1963) Instability of plasma sheath rotation and associated microwave generation in a Penning discharge. Appl. Phys. Lett., 3, 111.
- 86 Bohm, D. et al. (1949) Theoretical considerations regarding minimum pressure for stable arc operations. Natl. Nucl. Energy Ser. 1, 5, 77 ff. and 173 ff.
- 87 Redhead, P.A. (1958) The townsend discharge in a coaxial diode with axial magnetic field. Can. J. Phys., 36 (3), 255 - 270.
- 88 Hobson, J.P. and Redhead, P.A. (1958) Operation of an inverted-magnetron gauge in the pressure range 10⁻³ to 10⁻¹² mm Hg. Can. J. Phys., **36** (3), 271 - 288.
- 89 Leck, J.H. (1953) Sorption and desorption of gas in the cold-cathode ionization gauge. J. Sci. Instrum., 30, 71.

- 90 Barnes, G., Gaines, J., and Kees, J. (1962) Relative sensitivity and pumping rate of the redhead magnetron gauge. Vacuum, **12**, 141.
- 91 Rhodin, T.N. and Rovner, L.H. (1960) Gasmetal reactions in oxygen at low pressures. Transactions of the 7th National Symposium on Vacuum Technology, p. 228.
- 92 Kornelsen, E.V. (1960) A small ionic pump employing metal evaporation. Transactions of the 7th National Symposium on Vacuum Technology, p. 29.
- 93 Li, D. and Jousten, K. (2003) Comparison of some metrological characteristics of hot and cold cathode ionization gauges. Vacuum, 70, 531-541.
- 94 Li, D. and Jousten, K. (2003) Comparison of the stability of hot and cold cathode ionization gauges. J. Vac. Sci. Technol. A, 21, 937-946.
- 95 Jousten, K. (1995) Comparison of the sensitivities of ionization gauges to hydrogen and deuterium. Vacuum, 46, 9-12.
- 96 Jousten, K. (1998) Temperature corrections for the calibration of vacuum gauges. Vacuum, 49, 81-87.

14

Partial Pressure Vacuum Gauges and Leak Detectors

Dr. Karl Jousten, 1 Dr. Robert Ellefson, 2 and Werner Große Bley3

In this chapter, the reader learns about detecting gas species in vacuum gas mixtures. Leak detectors are also covered here because they are capable of detecting special gases as well.

14.1 Introduction

Besides being aware of total pressure, it is important to know the partial pressure constituents in a vacuum system. Knowing the components and their partial pressures is useful for assessing the pumping status of a vacuum system or for controlling a particular vacuum process. Measurement of partial pressures of the residual gases in a vacuum is principally done using small mass spectrometers, more commonly called residual gas analyzers (RGAs, covered in Section 14.2). Recently, optical absorption techniques to measure partial pressures of specific gas species have proven successful (Section 14.3). This chapter addresses both methods of measurement.

Leak detectors identify gas used as tracer gas. The tracer gas is the gas the detector is searching for. Tracer gas detectors thus present an important issue. Often, mass spectrometers are used as well. Leak detectors are covered in the final section (Section 14.4).

14.2 Partial Pressure Analysis by Mass Spectrometry

A mass spectrometer for measuring the residual gases in a vacuum system has five main components in its design: an ion source, a mass analyzer, an ion

¹Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany ²Consultant, formely INFICON, 6898 Vienna Woods Trail, 45459, OH Dayton, USA

³Inficon GmbH, Bonner Str. 498, 50968, Köln, Germany

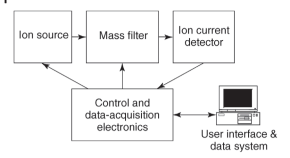


Figure 14.1 Elements of a mass spectrometer for partial pressure analysis.

detector to measure the ion currents of the mass-separated ions, control and data-acquisition electronics, as well as a computer and software providing a user interface (see Figure 14.1).

The ion source uses electron impact ionization similar to a total pressure gauge. Ions from the ionization region are extracted from the source by voltages on plates that have slits or orifices to direct and focus the ion beam into a mass analyzer.

The mass analyzer is tuned to a mass-to-charge ratio, m/e, of interest and a detector measures the transmitted ion current associated with ions showing the selected ratio m/e. Scanning through a range of m/e values with the mass analyzer and recording the ion currents as a function of mass produces a mass spectrum as shown in Figure 14.2. The ions in the spectrum are produced during ionization of gas species by electron impact. They represent the distribution of the total ion current in the source among its constituents. Ionization produces a characteristic pattern of m/e values for any specific gas species. Three possible categories of ions are formed.

- The parent ion formed by removal of a single electron from the molecule or atom.
- Fragment ions developing when chemical bonds in molecules break during ionization.
- Multiply charged ions formed by the loss of more than one electron from the atom or molecule.

The composite of all these ions appears in the mass spectrum. The relative abundances of ions in the mass spectrum are related to the partial pressures of the parent molecules or atoms present in the residual gas of the vacuum system. Thus, the mass spectrum can be used to infer the gas composition.

The partial pressure p_i of gas species i is related to a corresponding ion current, I_i , by

$$p_i = I_i/K_i, (14.1)$$

where K_i , determined by calibration, is the partial pressure sensitivity for the considered ion species.

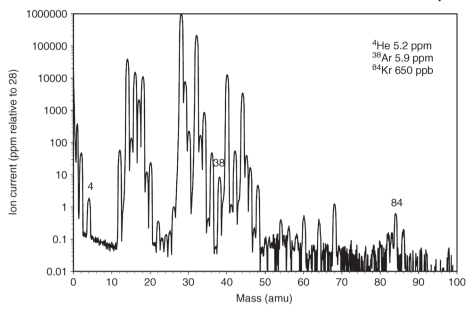


Figure 14.2 QMS spectrum of air. (CIS using 40 eV electron energy, electron multiplier with 1 s signal integration). Courtesy of *Inficon Inc*.

Patterns of ions are recorded in reference libraries, and with some experience, they become recognizable. This enables the user to qualitatively evaluate the species present. For example, Figure 14.2 shows air components from an air sample including minorcomponents of air such as helium and krypton. Methods to analyze mass spectra and to monitor selected ion species as a function of time are presented later in Section 14.2.6.2.

Partial pressure gauges are characterized by a number of fundamental terms [1,2]:

- Ion peak line width Δm , or resolving power $A = m/\Delta m$ (also called mass resolution).
- Sensitivity K_i for a reference gas (e.g., Ar or N_2).
- Maximum operating pressure, p_{max} .
- \bullet Minimum detectable partial pressure, $p_{\min} = \text{ion current noise/sensitivity}.$
- Abundance sensitivity or contribution of neighbor peak, AS, at a given mass.
- Mass range of operation for the mass spectrometer (e.g., 2–80 amu,2–200 amu).

The shape of an ion peak in a mass spectrum is a result of apertures or ion transmission features of the mass analyzer and is characterized by the width of the peak in mass units (amu). Peak width, Δm , is commonly defined as the width at 10% of the peak amplitude although sometimes the width at 50% height is used. The width Δm_{10} should be < 1 amu to avoid ion current contribution to neighboring ions and to allow measuring neighboring ion currents without any bias. Abundance sensitivity, AS, is a measure of this

contribution of a peak tail to the nearest neighbor mass:

$$AS = I_{m+1}/I_m,$$
 (14.2)

where I_{m+1} is the ion current measured at mass $m \pm 1$ due to a beam of ions of mass m. As an example, Figure 14.2 shows the tail of the mass-40 peak contributing \sim 2 ppm to the mass-41 peak when the peak width has a Δm of 0.90 amu and abundance sensitivity I_{41}/I_{40} of 2 ppm. Similarly at mass 39, the abundance sensitivity I_{39}/I_{40} is also ~2 ppm. Argon is a good choice for this measurement because most other gas species produce directly adjacent peaks.

Sensitivity K_i for specific gas species is determined by measuring the ratio of current for the reference ion and a known partial pressure for the reference gas species:

$$K_i = I_i/p_i. (14.3)$$

Defined pressure of a gas species is produced in a system by adding the gas to the vacuum system and noting the change in total pressure. If a gas-speciessensitive total pressure gauge is used, a correction factor for the species is needed (Section 13.7.6). The ionization in the source of the RGA is similarly species-sensitive, but correction factors differ from those in total pressure gauges because RGAs typically use 70 eV electron energy and ion gauges use 150 eV. It is common to determine the sensitivity for one particular gas (e.g., Ar or N_2) to characterize the basic sensitivity of the RGA and use a table of relative ionization probabilities K_{relative} to estimate species sensitivity:

$$K_i = K_{N_2} K_{\text{relative}}.$$
 (14.4)

Table 14.1 lists common relative ionization probabilities. Given values are calculated from the product of frequency of the substance fragment [3] and the relative sensitivity for ionization gauges [4], see Section 13.7.6.

Maximum operating pressure, p_{max} , is the pressure at which the sensitivity drops to 20% below low-pressure sensitivity. This loss of linear response (constant sensitivity) is typically due to space charge effects reducing ion transmission from the ion source through the mass analyzer.

The noise level of the current-measuring system and the sensitivity of the mass spectrometer in A Pa-1 ultimately limit the partial pressure detection limit for any ion species at very low pressures. A measure of this low-pressure detection capability is minimum detectable partial pressure (MDPP), which is defined as

$$MDPP = noise \ level (1\sigma)/(KG), \tag{14.5}$$

where G is the gain of an electron multiplier detector (if present) or unity for Faraday cup detection. The noise level or noise (1σ) measured depends on the integration time for current measurement, and thus, it is important to define the measurement interval or dwell time spent during current measurement. A typical dwell time is 1s for the longest practical time spent to measure an ion current. More often, shorter dwell times are used and the resulting noise level and thus MDPP rise.

Table 14.1 Compilation of common ion species and their relative ionization probabilities at 70 eV. Sensitivity for a considered species is calculated from the product of fragment frequency for the main ion and the relative ionization probability of the substance, $K_i = K_{N_2} K_{relative}$.

Substance	Chemicalnotation	Main ion	K_{relative}	Substance	Chemicalnotation	Main ion	K_{relative}
Acetone	(CH ₃) ₂ CO	43	2.09	Krypton	Kr	84	0.97
Air		28	0.71	Methane	CH ₄	15	0.64
Ammonia	NH_2	17	0.69	Methanol	CH ₂ OH	31	0.77
Argon	Ar	40	1.06	Neon	Ne	20	0.21
Benzole	C_2H_2	78	3.13	Nitrogen	N_2	28	1.00
Bromine	Br_2	81 or 79	2.53	Nitrogen oxide	NO	30	1.09
Carbon dioxide	CO_2	44	1.19	Dinitrogen monoxide	N_2O	44	1.10
Carbon disulfide	CS_2	76	2.67	Oxygen	O_2	32	0.91
Carbon monoxide	CO	28	0.96	Phosphine	PH_3	35	1.44
Carbon tetrafluoride	CF_4	69	1.19	Propane	C_3H_8	29	1.16
Dichlorodifluoromethane	CCl_2F_2	85	1.68	Hydrosilicon	SiH_4	28	1.13
Ethan	C_2H_6	27	1.16	Silicon tetrachloride	SiF ₄	85	1.1
Ethanol	C ₂ H ₅ OH	31	1.76	silicon oxyfluoride	SiOF	63	1.1
Helium	He	4	0.14	Sulfur dioxide	SO_2	64	1.26
Hydrogen chloride	HCl	36	1.07	Sulfur hexafluoride	SF ₆		1.64
Hydrogen fluoride	HF	20	1	Water	H_2O	18	0.75
Nitrogen trifluoride	NF_3	52	0.83	Xenon	Xe	134	0.78
Hydrogen sulfide	H_2S	34	1.47				

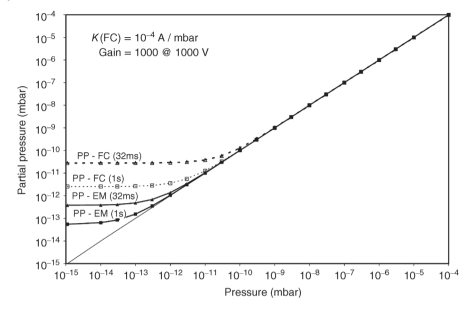


Figure 14.3 Minimum detectable partial pressures (MDPP) for selected ion-current integration times (32 ms and 1 s) for the ion-current signals in a *Faraday* detector (FC) and in an SEM (EM). MDPP is determined mainly by the noise level in the amplifier.

Figure 14.3 shows a modeling of measured noise levels for *Faraday* cup (FC) and electron multiplier measurements for two different dwell times: 32 ms and 1 s. It is clear from this set of curves that the partial pressure detection limit is raised by shorter dwell times associated with higher sampling rates. Thus, the user must trade off detection limit with sampling rate to see small changes in composition. For electron multipliers, signal gains range from 10^2 to 10^6 , which in principle dramatically lowers the minimum detectable partial pressure. However, the noise level (1σ) in MDPP calculation also scales with gain such that a typical improvement in signal-to-noise by using an SEM (secondary electron multiplier) is a factor of about 100 over an FC measurement. The other factor that can reduce MDPP is increased FC sensitivity. This is often achieved by increasing electron emission current, but at the expense of high-pressure linearity due to space charge effects in the ion source. If best low-pressure detection limit is the goal, such increases in sensitivity with emission current coupled with modest SEM gain (\sim 1000) are recommended.

14.2.1 Ion Source Design

Electron impact ion sources have evolved from early models by *Dempster* [5], *Bleakney* [6], *Nier* [7,8], as well as *Ingraham* and *Hayden* [9] for magnetic sector mass spectrometers. Descriptions of these early ion sources are given in

Duckworth et al. [10]. Austin, Holmes, and Leck [11] published variants of the ion sources for combinations with quadrupole mass spectrometers (QMS) based on work done by Brubaker [12]. The dominant partial pressure analyzer is the QMS, so the focus here is on ion sources for this instrument. The nature of the quadrupole mass filter is to accept a cylindrical ion beam from the source. Thus, ion sources for the QMS have elements with cylindrical symmetry as a characteristic of their design. For RGAs, three categories of ion sources are available: open ion sources (OIS), closed ion sources (CIS), and molecular beam ion sources (MBIS).

The OIS ionizes the residual gas of the vacuum chamber into which the RGA is inserted. The chamber pressure of the process being monitored is lower than the operating pressure of the RGA (typically $< 10^{-2} \, \mathrm{Pa}$) so that the RGA with OIS can be inserted directly into the vacuum system.

An RGA with a CIS samples the process gas that is at a pressure higher than the operating pressure of the analyzer. Flow-controlling orifices or channels are used to introduce a small gas stream from some higher pressure source through the CIS region (where ionization takes place) and into the lower pressure region of the mass analyzer and detector. An independent pumping system connected to the analyzer is used to establish the flow of gas through the CIS RGA.

A molecular beam ion source is characterized by a collimated beam of process gas focused through the ionization region without colliding with the walls of the ion chamber. This minimizes accumulations on the ion chamber walls that can contribute to background (memory) from earlier sample compositions or can cause reactions on the surfaces leading to insulating layers (e.g., SiO₂) that affect ion optics. Figure 14.4 shows examples of different types of ion sources.

14.2.1.1 Open Ion Sources (OIS)

The term open ion source refers to a high gas conductance between the ionization region and the region surrounding it. This allows the gas molecule density in the ion source region to be the same as the density in the vacuum chamber (no pressure differential). This is the basis for residual gas analysis of the species in a chamber. An open ion source is characteristic to all RGAs. Figure 14.5 shows two structures of open ion sources.











Figure 14.4 Commercial RGA ion sources. Left to right: open ion source, UHV ion source, molecular beam ion source, and two types of CIS.

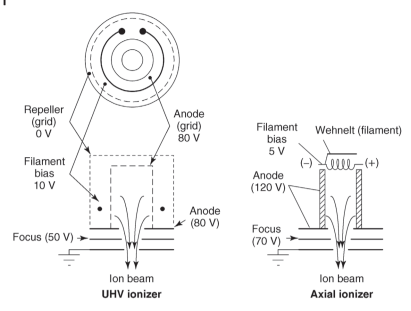


Figure 14.5 Open ion source examples and typical operating potentials.

The UHV ion source shown is similar to an ionization gauge. The open grid structure assures high conductance to the vacuum region surrounding the ion source. Anode voltage is 80 V or higher. This assures that the filament is positive with respect to ground while biased 70 V below anode potential. Electrons from a filament outside of the anode grid structure are then accelerated toward and through the grid while producing ions within the enclosed anode region. These ions, instead of being collected on a center wire as is the case in an ion gauge, are extracted by the ion focus lens(es) to form an ion beam directed toward the mass filter.

One advantage of the UHV ion source is that it has less surface area for outgassing from the grid structure than an axial ionizer [11,13]. This reduces the residual gas background in the ion source and allows direct measurement of lower partial pressures down to the 10^{-13} Pa range (see Figure 14.8). Appropriate selection of materials and pretreatment (e.g., vacuum baking) further reduce outgassing.

The axial ionizer [14] is open on both ends with a relatively high conductance between the ionization region and the vacuum outside. Again, the anode is charged with the highest potential in the ion source (120 V in Figure 14.5). The repeller (*Wehnelt*) tied to the negative side of the DC filament supply repels electrons from the filament. The filament is biased by 70 V with respect to the anode so electrons are attracted to the anode. The electrons drift through the anode region and collide with gas molecules to form ions, which are drawn into the mass filter by the lower potentials of the focus lens and ion exit plates. A focus plate, more positive than anode potential, repels electrons that drift through the

anode region, while the same plate attracts and focuses ions to form the ion beam. The resulting axial ion source shows high efficiency for ion formation with sensitivity as high as 10^{-5} A Pa⁻¹ (10^{-3} A mbar⁻¹).

14.2.1.2 Closed Ion Sources (CIS)

Imagine the anode of the UHV ion source in Figure 14.5 as a full-metal cylinder instead of the shown grid structure. Sealing the spaces between focusing elements with ceramic insulators and adding a hole on the side of the anode allowing electrons to enter the enclosed region yields a CIS [15-17]. Figure 14.6 shows such a CIS where the gas to be analyzed enters through a supply tube sealed to the closed anode. The gas exits through the electron beam hole and the ion exit hole.

The basic advantage of the CIS is that pressure in the ionization region is higher than in the analyzer region. The pressure factor is equal to the ratio obtained by dividing the analyzer pumping speed by the gas-exit pumping speed of the CIS. Typically, this ratio ranges from 10 to 100, producing CIS pressures up to 10^{-1} Pa (10^{-3} Torr) while the analyzer operates at 10^{-3} Pa (10^{-5} Torr). High ion source pressure allows measuring low-level impurities whereas analyzer pressure is maintained at a pressure low enough to minimize ion losses due to gas collisions in the analyzer section. For a throughput of gas through the CIS and analyzer of, for example, $0.02 \, \mathrm{Pa} \, \ell \, \mathrm{s}^{-1}$, the CIS exit conductance of $1 \ell \, \mathrm{s}^{-1}$ produces 0.02 Pa in the CIS ionization region. Background components in the CIS due to outgassing or sample-induced desorption are released mainly from the small surface area inside the ionization region where the electron beam terminates. Thus, their contribution to contamination of the target gas sample is low. These smaller backgrounds allow measuring low-level impurities in gas samples. The filament is located externally to the ionization region thus

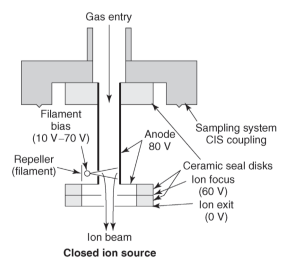


Figure 14.6 Principle of a CIS. Note ceramic gas seals between focus plates and sampling system for maintaining high pressure inside the anode region where the electron beam forms ions.

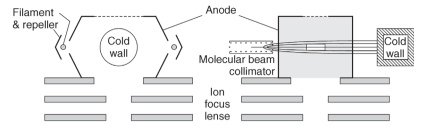


Figure 14.7 Principle of a crossed-beam ion source. The left-hand side shows the glow filament, the repeller, the anode, as well as the corresponding extraction and focus lenses. The molecular beam enters perpendicularly to the plane of projection through the center of

the anode. The right-hand side illustrates the molecular beam as it travels through the ionization chamber and hits a cold surface. Molecules travel perpendicularly to the electrons as well as to the ion beam.

minimizing the diffusion of filament-surface reaction products back into the ionization region and their subsequent contribution to the composition of the analyzed gas sample.

14.2.1.3 Molecular Beam Ion Sources (MBIS)

Molecular beam ion sources (MBIS) represent the third variant of ion sources. In an MBIS, the gas enters the ionization chamber with an electron beam oriented perpendicularly to the gas flow [18]. Produced ions are extracted in a direction perpendicular to the gas flow as well as to the electron beam as shown in Figure 14.7. The aim in this type of design is to ionize the molecules in the gas jet before they impinge on the surface of the ion source. This is particularly advantageous when analyzing condensable gases, gases with high sticking probability, and gases that tend to produce insulating layers. A molecular emitter collimates the gas flow so that it passes through the ionization chamber and onto a cooling surface or into a pump. For gases such as UF₆, this condensation on a cooling surface is important because it prevents contamination of the walls in the ionization chamber, and thus, minimizes memory effects due to outgassing of previously analyzed gas mixtures.

14.2.2

Filament Materials

The choice of filament materials to use in an RGA or CIS should be directed by the analysis application. Table 14.2 shows a list of filament materials sorted according to the frequency of current usage. The most popular filament materials for general RGA use are yttria-coated iridium (Y_2O_3/Ir) and thoria-coated iridium (ThO_2/Ir) . This is due to two main reasons:

1) The oxide coating provides high tolerance to oxidizing atmospheres often present with residual air components and water vapor, which can shorten the lifetime of tungsten or rhenium filaments.

Table 14.2 Filament materials and their gas compatibility.

Material	Operating temperature [19]	Gas tolerance	Comments
Y ₂ O ₃ /Ir	1200°C	Inert gases, air/ O_2 , NO_x , SO_x	Short life in halogens
			Generates some CO/CO ₂
			O_2 with high H_2O background
ThO ₂ /Ir	1200°C	Inert gases, air/ O_2 , NO_x , SO_x	Short life in halogens
			Generates some CO/CO ₂
			O_2 with high H_2O background
			[Low level radioactivity (α)]
W, W/ 3% Re	1800°C	Inert gases, H ₂ , halogens freons	Short life in oxidizing gas
			Generates copious CO/CO ₂ from O ₂ or H ₂ O background
			WO_x is volatile and may evaporate
Re	1800 °C	Inert gases, hydrocarbons H ₂ , halogens, freons	Months of lifetime due to evaporation
			Used for hydrocarbon but consumes O_2
			Generates moderate CO/CO_2 from O_2 or H_2O background

2) The operating temperature for electron emission is lower [19] than for tungsten so thermally induced outgassing is lower as well.

However, both oxide-coated filaments are not tolerant to reducing atmospheres of halogens or hydrogen. Thus, these filament materials should be avoided for semiconductor etching processes or freon measurements. Fortunately, tungsten or tungsten with 3% rhenium [20] (the rhenium reduces embrittlement and improves oxygen tolerance) can be used for hydrogen, halogens, and freons. The main problem with tungsten is short service life in air/oxygen. However, for partial pressures of oxygen remaining below 10^{-4} Pa (10^{-6} Torr), service lives of greater than a year can be expected for tungsten filaments. Pure rhenium has been used extensively for hydrocarbon analysis applications but it suffers from high evaporation rates (10 times higher than for tungsten filaments) at standard operating temperatures for electron emission [20]. Users of RGAs in clean vacuum systems regard the relatively short service lives of rhenium filaments (2 to 4 months) and the associated frequent filament replacement as disadvantages. In contrast, users of hydrocarbon analysis do not seem to mind; they have to clean their ion source often so replacing a filament, for them, is part of the process.

14.2.3

Artifacts in the Mass Spectrum due to the Ion Source

A hot filament generating electrons for ionization can also produce undesired effects. The hot filament can induce many surface reactions. A typical reaction involves surface carbon on the filament reacting with oxygen from O2 or H2O thus producing CO and CO2. High levels of water vapor in contact with Y2O3 (and to a lesser extent ThO2) can lead to low levels of O2 from surface reactions on the filament.

Presence of water vapor can additionally lead to formation of hydronium, H₃O⁺. The time of passage of water vapor ions in the ion source is long enough for H_2O^+ to react to H_3O^+ in further ion–molecule reactions [21].

Miniature mass spectrometers commonly operate in the 10^{-2} –1 Pa (10^{-4} – 10⁻² mbar) pressure range and exhibit ion-molecule reaction species (with abundances up to 1000ths of a ppm with respect to the parent ion) [22]. Examples are N^{3+} when N_2 is dominant, ArH^+ and Ar_2^+ when Ar is the dominant gas present at high pressure. Reaction products of these chemistries can appear in the mass spectrum of a high-pressure RGA directly and to a lesser extent in a CIS as artifacts of the mass spectrometer. Figure 14.2 shows an example of ionmolecule reactions in a CIS with $m/e = 42, N_3^+$ mass present in addition to the true species present in the vacuum.

The heat from the filament can also desorb species (H₂O, CO₂, H₂) from nearby surfaces, or by bulk heating of the surrounding metal, creating a high concentration of off-gassed species not representing the composition in the rest of the vacuum chamber.

Finally, the electrons emitted can strike vacuum surfaces with enough energy to desorb species such as O⁺, Cl⁺, and F⁺ from the ion source grids or walls, which then appear in the mass spectrum as artifacts of the measuring process [23]. These ions released by electron-stimulated desorption (ESD) can be identified by reducing the ion energy and monitoring the drop in ion currents. Peaks that do not drop as rapidly as those of typical ions in the gas phase relate to ESD ions. The ESD ions observed in the spectrum leave the surface and carry a certain initial energy promoting their passage through the mass filter even when ions from the gas phase are already repelled due to their lower energy. Detection of F and Cl among ESD ions indicates that parts of the vacuum system, in their previous history, were exposed to freons or other chemicals containing F or Cl.

These thermal and ESD artifacts are particularly noticeable in UHV and XHV systems. Efforts to minimize these effects include a copper heat sink to transport the filament heat [24,25] as well as Pt-coated or Pt–Ir wire, fine mesh anode grids for minimizing ESD [26,27], see Figure 14.8.

Other reactions induced by the filament include the formation of HD and/or HT when D2 and/or T2 are present due to isotopic exchange on the hot filament or wall surfaces [28].

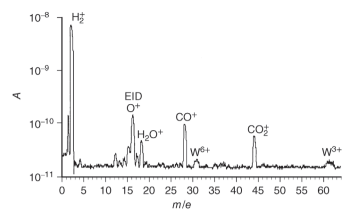


Figure 14.8 Mass spectrum recorded in an XHV system at 2×10^{-10} Pa. Apart from the H_2^+ peak dominant in this pressure range, the spectrum shows further residual gas constituents as well as ions produced by ESD with masses 16 (O⁺), 19 (F⁺), and 35 (CI⁺) [26].

14.2.4 Mass Analyzers

The most common mass spectrometer for partial pressure measurement in vacuum systems is the QMS. It is available in many mass ranges and quad sizes relating to performance and operating pressure. Small magnetic sector mass spectrometers have been extensively used for (helium) leak detection (see Section 14.4) and some for residual gas analysis. A new mass analyzer design is the autoresonant trap mass spectrometer (ARTMS). Other small mass analyzer types include $E \times B$ -cycloid mass analyzers [29,30] and the omegatron [31,32]. Both of these mass spectrometers types are currently not commercially available. Time-of-flight (TOF) mass spectrometers are used extensively for rapid analysis of high-mass hydrocarbons for pharmaceuticals and biological molecules. Small TOF mass spectrometers have been built for portable instruments but to date not for residual gas analysis [33]. In this section, design and operation of a quadrupole mass analyzer, a magnetic sector mass analyzer, and the ARTMS are presented. All types are employed in RGAs for vacuum analysis.

14.2.4.1 Quadrupole Mass Analyzers

Ideally, a QMS or mass filter is based on four hyperbolically shaped cylindrical surfaces arranged 90° apart (as shown in Figure 14.9) to generate a pure quadrupole electric field on the axis of the filter [34]. A hyperbolic pole mass analyzer has been built for a miniature mass spectrometer [35] but most mass filters are made from round rods. The rods are connected to a voltage source as shown in Figure 14.9 which is a sum of a DC rod potential, an RF voltage of frequency $f = \omega/2\pi$, plus a constant pole zero axis voltage (field axis FA). The ion energy of ions that are injected into and drift along the axis of the mass analyzer is the

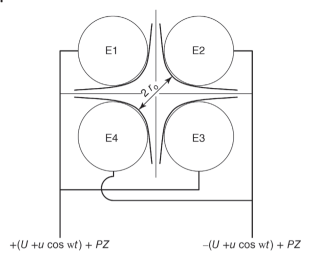


Figure 14.9 Diagram of electrodes E1, E2,E3, and E4 in a quadrupole mass filter made up of The opposite poles E1 and E3 are connected round rods or hyperbolic pole rods. Round rods are a reasonable approximation and

produce electric fields similar to quadrupoles. to one side of the RF/DC voltage input, and E2 and E4 to the other side.

difference between the anode potential and the pole zero, PZ, applied to the quadrupole and is typically in the range of $5-15 \, \text{eV}$.

The rod RF and DC voltages acting on the drifting ions cause oscillation and deflection of ions perpendicular to the drift axis as shown in Figure 14.10 [36]. For ions to drift successfully through the rod structure, their oscillatory amplitudes must be $r < r_0$ so they do not hit a rod and receive an electron that neutralizes their charge and causes the ion to be lost back to the gas phase.

Note that in Figure 14.10 most ions are lost near the entrance. A small fraction (a few per cent) of the ions formed in the ion source are mass analyzed and measured in the detector. This large rejection of ions explains why the sensitivity of a typical RGA with $\Delta m = 1$ is about 1×10^{-6} A Pa⁻¹ while a total pressure ion gauge with similar electron current has a sensitivity of 1×10^{-4} A Pa⁻¹.

Conditions for stable trajectories through the quadrupole structure are given by solutions to *Mathieu's* equations [37] describing the motion. The stability region is characterized by two parameters:

$$a = \frac{8\xi eU}{Mr_0^2 \omega^2}$$
 or $a = 0.194 \frac{\xi U}{mr_0^2 f^2}$, (14.6a)

$$q = \frac{4\xi eu}{Mr_0^2 \omega^2} \text{ or } q = 0.097 \frac{\xi u}{mr_0^2 f^2},$$
 (14.6b)

where ξ is the charge state of the ion, $(\pm)U$ is the DC potential applied to adjacent rods, u is the peak voltage of the RF applied to the rods, M is the actual

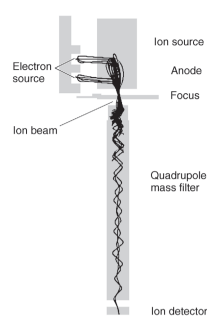


Figure 14.10 Ion-optics-software plot (Simion 7.0) of electron trajectories and the trajectories of the ions formed in the ion source as they are focused into the quadrupole mass analyzer. Note that most of the ions are rejected

near the entrance to the filter and only ions of correct mass, position, and velocity pass successfully (oscillatory) through the quadrupole structure to the detector.

mass in kg, m is the nominal mass in amu, and $\omega = 2\pi f$, where f is the frequency of the RF.

Figure 14.11 shows the stable region described by Eqs. (14.6a) and (14.6b). For constant Δm ion transmission (the most common mode of QMS operation), the voltage applied to the quadrupole is

$$V = U + u\cos\omega t,\tag{14.7}$$

where the ratio a/q = 2U/u is constant and in the range $0 \le 2U/u \le 0.336$.

Figure 14.11 shows a scan line with slope a/q = 0.283 as an example of scanning the RF amplitude u and keeping the DC value U = 0.283u/2 = 0.1415u. This allows transmission of a small portion of all ions producing an ion transmission peak centered around mass m with a peak width Δm . The area between the scan line and the apex (a = 0.237, b = 0.706) is proportional to the sensitivity of the QMS. As the scan line ratio a/q approaches 0.336, peaks become very narrow but sensitivity also drops. In general, for a QMS, the product of the resolving power of the mass filter, $m/\Delta m$, and sensitivity, K, is a constant. Additionally, for $\Delta m = 1$, this relation predicts K = constant/m. Figure 14.12 shows the result of this prediction plotting the sensitivities of common gases in relation to $K(N_2)$. A transmission factor T(m) = 28/m is shown to represent both the

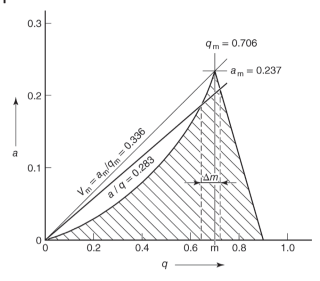


Figure 14.11 Plot of *a* versus *q*, showing the stability region of a quadrupole mass filter (i.e., ions with parameter combinations in the shaded region pass through the filter).

effect of the mass filter and the relative ionization probability $K_{\rm relative}$ of gas species, the latter accounting for deviations from the transmission curve. Ion current transmission in a quadrupole depends on the inside radius r_0 , the rod length, RF frequency, ion energy, and the injection conditions under which the ions enter the quadrupole from the ion source.

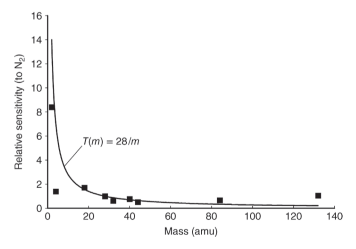


Figure 14.12 Ion transmission and relative sensitivity for common gas species: 2-H₂, 4-He, 18-H₂O, 28-N₂, 32-O₂, 40-Ar, 44-CO₂, Kr, and Xe. Note the remarkably high sensitivity for H₂. The sensitivity for He is about the same as for

 N_2 due to the effect of high ion transmission (28/4) which compensates for the low ionization probability (~17 of ionization probability for N_2).

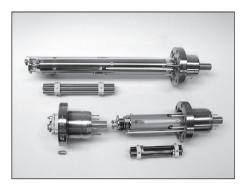




Figure 14.13 (a) Examples of small QMS sensors and their associated quad rod assemblies and electron multipliers. Clockwise from the top right: a 300 amu sensor with open ion source, $6 \text{ mm} \times 130 \text{ mm}$ quad, and an optional CIS (note ceramic seal disk). A compact 100 amu sensor with $6 \text{ mm} \times 90 \text{ mm}$ quad with a small microchannel plate EM. A high-

pressure QMS with 12 mm long quad with hyperbolic poles. (b) Rod assemblies with the following dimensions: $6 \text{ mm} \times 100 \text{ mm}$, $8 \text{ mm} \times 200 \text{ mm}$, and $16 \text{ mm} \times 300 \text{ mm}$. Compared to smaller sensors, large sensors allow higher mass operation (> 2000 amu) and their larger r_0 leads to higher transmission and sensitivity. Courtesy of *Inficon Inc*.

Examples of commercial QMS are shown in Figure 14.13. The sizes vary primarily due to quadrupole lengths and electron multiplier sizes. The ones shown are designed to fit into CF $40\,\mathrm{mm}$ flanges and UHV tubing.

14.2.4.2 Miniaturized Quadrupole Mass Analyzers

Another factor in the operation of a OMS is the operating pressure of the RGA. Ions are accelerated, drift through gas and can encounter collisions and be lost if the mean free path of an ion is shorter than the flight path between ion source and detector. Flight paths in common QMS for residual gas analysis with 100 mm long rods have an ion flight path of \sim 250 mm. As a result of this mean free path, argon collisional losses are approximately 63% at $3 \times 10^{-2} \, \text{Pa}$ $(3 \times 10^{-4} \text{ mbar})$. To operate an OIS QMS at higher pressure for direct monitoring of physical vapor deposition (PVD) processes requires reducing the ion path length to the detector while maintaining the mass separation function. An example of a high-pressure RGA is the 12 mm long quad and RGA shown in Figure 14.13 [35]. Evidence of the extended pressure range is given in Figure 14.14 showing FC sensitivity as a function of pressure for a standard RGA and a miniature (high-pressure) RGA. Both separate masses from 0 to 100 amu with peak widths $\Delta m = 1$. The pressure associated with the mean free ion path is shown for the standard and miniature RGA. Note that the loss of sensitivity occurs at lower pressures than predicted by scattering alone. An additional mechanism for loss of sensitivity is the shielding of draw out potentials by the buildup of ion space charge plus some Coulomb repulsion of the ion beam before entering the quadrupole [38-41]. Note that the tradeoff for achieving an extended pressure range in RGA operation is lower sensitivity. This stems from

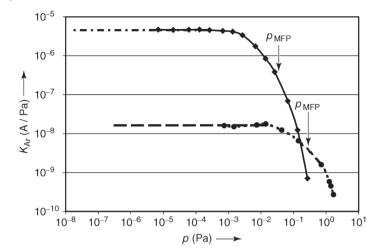


Figure 14.14 Sensitivity versus pressure for a standard RGA (♦) and a high-pressure RGA (♦). each RGA. Additional ion current fall-off at The pressure corresponding to one mean free path (p_{MFP}) for complete ion motion

between source and detector is noted for lower pressures is due to space-charge shielding in the ion source and to ion repulsion.

the scaling of r_0 and L by a factor 1/10 which reduces the area of the quad entrance for ion transmission by a factor of 1/100.

Thus, the nominal 2×10^{-6} A Pa⁻¹ $(2 \times 10^{-4} \text{ A mbar}^{-1})$ sensitivity for standard RGA changes to 2×10^{-8} A Pa⁻¹ (2×10^{-6} A mbar⁻¹) for the high-pressure RGA. This lower sensitivity is made up for by the extended pressure range. Operation of the high-pressure RGA at 0.5 Pa $(5 \times 10^{-3} \text{ mbar})$ with $1.3 \times$ 10^{-8} A Pa⁻¹ (1.3 × 10^{-6} A mbar⁻¹) sensitivity still produces 1 × 10^{-8} A, the same current as obtained in a standard RGA operating a 2×10^{-3} Pa $(2 \times 10^{-5} \text{ mbar})$ with 5×10^{-8} A Pa⁻¹ (5×10^{-6} A mbar⁻¹) sensitivity.

A number of approaches are available to making a miniature quadrupole or quadrupole array for producing RGAs that operate at pressures of 10⁻³ mbar and above. They are summarized in a publication by Badman and Cooks [42]. The quadrupole for the high-pressure RGA in Figures 14.13 and 14.14 is a composite of Inconel and ceramic with hyperbolic pole faces machined in place by electrode discharge machining [43]. Another commercially available highpressure RGA uses the traditional mechanical alignment of four miniature rods. A third approach is an array of 16 pins sealed in a glass feedthrough to form nine quadrupoles operating in parallel [44]. A new approach is to micromachine (etch) precise features into a silicon wafer, bond metallized (fiber optic) glass rods to the silicon, provide electrical contacts with indium, and then to assemble the quadrupole structure by facing the two plates together with a spacer [45,46]. The result is a micro-electro-mechanical structure (MEMS) as a basis for a QMS. Another approach aims at producing a MEMS Wien mass filter (crossedfield) [47]. Some of these miniature mass spectrometers are commercially available while others will develop further and will probably be available in the future.

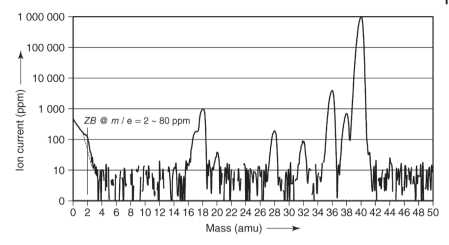


Figure 14.15 Analog scan of Ar at 0.7 Pa 8 eV of ion er $(7 \times 10^{-3} \text{ mbar})$ with a high-pressure RGA an 18 mm hy operating with FC ion detection. The spectrum $r_0 = 0.5 \text{ mm}$. is taken with 40 eV/200 μA electron current,

8 eV of ion energy with ions focused through an 18 mm hyperbolic quadrupole with

A practical goal for a high-pressure RGA is to produce a mass spectrum with 1 amu peak-width mass resolution and enough sensitivity to measure ion currents of low-level species in the residual gases of plasma processes operating at pressures around 1 Pa. Figure 14.15 shows a mass spectrum of 0.7 Pa (7×10^{-3} mbar) argon recorded by a high-pressure RGA with the sensitivity shown in Figure 14.14. *Faraday* cup ion detection shows a noise level of about 10 ppm for the baseline.

14.2.4.3 Magnetic Sector Analyzers

A second important type of mass analyzer for small mass spectrometers is the magnetic sector. Large-radius magnetic sector mass spectrometers have been used for high (mass) resolution mass spectrometry for decades. Interest in magnetic sectors for small mass spectrometers has waned with the success of the QMS but is renewed with the availability of rare-earth permanent magnets capable of producing very high (e.g., 1 T) magnetic fields with small but useful pole gaps.

A magnetic sector mass spectrometer separates ions by magnetic deflection. Ions formed in the ion source are accelerated with a voltage U into the magnetic sector. Each ion thus has an energy

$$\xi eU = \frac{1}{2}mv^2,\tag{14.8}$$

where ξ is the charge state of the ion of mass m, and ν is the speed of the ion. The ion moving through the magnetic sector in the plane perpendicular to the magnetic field experiences a magnetic force directed toward the center of the radius of the sector. This force is equal to

$$\xi evB = \frac{mv^2}{R},\tag{14.9}$$

where *B* is the magnetic field strength and *R* is the radius of the path traced by the ion's motion in the uniform magnetic field. By eliminating ν from the two equations above, the mass-to-charge ratio m/e is related to the radius R, magnetic field strength B, and ion accelerating voltage U:

$$\frac{m}{\xi e} = \frac{R^2 B^2}{2U}. (14.10)$$

This is the traditional result of a single-focusing magnetic sector mass spectrometer. A recent development by *Diaz et al.* [48] added an electrostatic sector, as shown in Figure 14.16, that exerts an outward radial force to the ions in the sector. Thus, the net force on an ion is

$$Mv^2/R = \xi evB - \xi eE, \tag{14.11}$$

with E denoting the radial electrical field in the sector. The condition for energy focusing is met when the inward magnetic force is twice the outward electrostatic force on the ion, which means that

$$vB = 2E. \tag{14.12}$$

Solving for m/e by again eliminating ν gives the double-focusing magnetic sector relation

$$\frac{m}{\xi e} = \frac{R^2 B^2}{8U}.\tag{14.13}$$

A practical interpretation of this result is that with a double-focusing sector of radius R and field B, the ion accelerating voltage is 1/4 of that in a single-focusing magnetic sector. The theoretical resolving power is also enhanced by a factor of

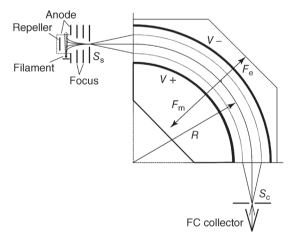


Figure 14.16 Double-focusing magnetic sector mass spectrometer according to Diaz, Giese, and Gentry. The magnetic field emerges perpendicularly from the plane of projection. (J. Am. Soc. Mass Spectrom. 2001, 12, 619–632). Courtesy of Mass Sensors Inc.

2 over single-focusing and is given by

$$A = \frac{m}{\Delta m} = \frac{2R}{S_s + S_c},\tag{14.14}$$

where S_s and S_c are the source and collector slit widths, respectively, that define transmission through the magnetic sector.

This design, together with NdFeB permanent magnets producing a 1T magnetic field, allows a small 8 mm radius magnetic sector to be built that produces a mass spectrum as shown in Figure 14.17. This mass spectrum is taken by scanning the ion accelerating voltage, U, defined in Eq. (14.12).

The resolving power A of a magnetic sector is constant which means that the step size of U to scan over a peak must vary with mass. Starting with Eq. (14.13), the change in mass transmitted with a voltage increment dU is

$$dm = \frac{-R^2 B^2 dU}{8U^2},$$
 (14.15)

where the negative sign implies that to move in positive dm on the mass scale, the voltage change dU must be negative. If X individual steps over a peak width Δm are desired, then

$$dm = \frac{\Delta m}{X} = \frac{m}{AX},\tag{14.16}$$

where A is the resolving power. Combining Eqs. (14.15) and (14.16), and solving for d*U* gives

$$dU = -\frac{1}{AX}U. (14.17)$$

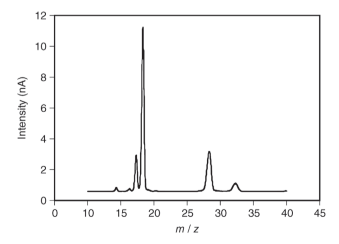


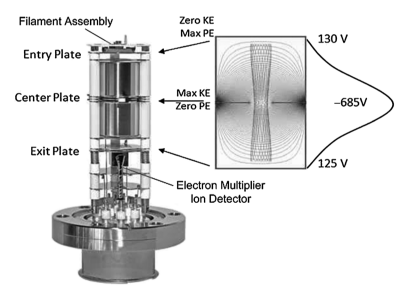
Figure 14.17 Double-focusing magnetic sector mass spectrometer with radius r = 8 mm, B = 1 T, slot widths $S_s = 0.1 \text{ mm}$ and $S_c = 0.2 \text{ mm}$. Peaks shown are background peaks at 10⁻⁶ mbar measured with an EM featuring a gain of 1000.

Thus, by choosing the desired number of steps X over a peak, and knowing the resolving power A, Eq. (14.14), the increments dU can be calculated to produce the scan in Figure 14.17.

14.2.4.4 Autoresonant Ion Trap Mass Analyzers

A new design for mass analysis is the autoresonant ion trap mass spectrometer (ARTMS) [49-51] where the ions are created by electron impact ionization within an electrostatic field that traps the ions into a confined region as indicated in Figure 14.18. The ion trap boundaries are defined by the entry plate and exit plate potentials, and the depth of the potential well is defined by the voltage on the center plate.

The resonant frequency that adds energy to the ion for ejection to the electron multiplier detector is proportional to $(M/Z)^{-1/2}$, where M is the mass and Z is the charge on the ion. Electrostatic scans over a frequency range can be very fast such that the whole mass range can be scanned within 100 ms. The ion trap as configured holds about 20 000 ions which limit the dynamic range of measurement for minor species to two orders of magnitude for a single scan. With averaging of multiple scans the dynamic range extends to three orders of magnitude of species abundance or more. Species abundances can be measured at all



trometer, where ions formed within the potential of the entry plate are trapped in the potential well defined by the center plate. The electron emitting filament assembly is on the upper end. The trapped ions have maximum potential energy (PE) at the entrance and exit plates and maximum kinetic energy (KE) going

Figure 14.18 An autoresonant trap mass specthrough the center plate aperture. The ions are ejected through the lower exit plate to an electron multiplier by applying a fixed RF voltage on the center plate and varying the frequency over the resonant frequencies of the masses of interest. Courtesy of Granville-Phillips Division of MKS Instruments, Inc.

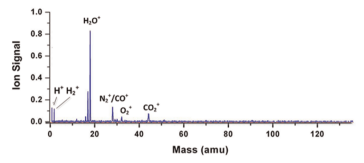


Figure 14.19 A single 80 ms ARTMS scan of residual gas at 10^{-7} mbar pressure. The signal-tonoise ratio is 400. *Courtesy of Granville-Phillips Division of MKS Instruments, Inc.*

operating pressures because the ion trap fills with ions at all pressures. The operation of the ART mass spectrometer (ARTMS) is best at 10^{-7} mbar pressure but reaches into the UHV and XHV pressure regimes producing mass spectra like that shown in Figure 14.19 [52]. The upper pressure range limit is about 10^{-5} mbar. At higher pressures, the ion production rate exceeds the resonant ion extraction rate and the excess ions appear as an elevated baseline. Ion extraction efficiency is constant over the mass range in contrast to the 28/M mass dependence for QMS ion transmission shown in Figure 14.12. As a result, the relative ionization probabilities for the ARTMS are different from those given in Table 14.1. To obtain partial pressures, the relative abundances measured are multiplied with a total pressure measurement from a suitable gauge.

14.2.5 **Ion Detectors**

A key element in a mass spectrometer is the measurement of the ion current of mass-separated ions. Two basic types of detection methods are used in RGAs: *Faraday* cup detector (FC) and (secondary) electron multiplier, (S)EM. The category of electron multipliers includes several types, described next, that are used for residual gas analysis.

14.2.5.1 Faraday Cups

The Faraday cup (FC) is the simplest detection method for ion current measurements because it is simply a detection plate and an electrometer. It is generically termed Faraday cup detector in honor of Michael Faraday, a nineteenth century physicist who in his experiments collected electrical charges in a metal cup. Ion detection occurs when electron current flows to the detection plate to neutralize the charge of the ions that arrive; an appropriate electrometer circuit measures the current. The input resistance and the distributed capacitance of the current-measurement circuit dictate the time constant for a Faraday type of ion current

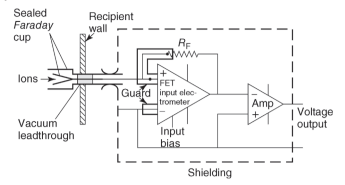


Figure 14.20 Electrometer circuit that connects to a shielded and immersed *Faraday* cup. Both the vacuum feed-through and the complete electrometer circuit are shielded to prevent measuring errors caused by interference from the surrounding.

measurement. A time constant $\tau = RC \approx 0.1\,\mathrm{s}$ is typical for Faraday cup/electrometer systems. The detection limit of a *Faraday* detector is typically $3\times 10^{-16}\,\mathrm{A}$ $(2\times 10^3\,\mathrm{ions\,s^{-1}})$, the noise limit in modern field-effect transistor electrometers for 1 s integration time (see Figure 14.20 for a basic circuit diagram). A *Faraday* detector is the simplest and least expensive ion detection device and is often used in low-cost mass spectrometers.

14.2.5.2 SEM Detection

Small partial pressures produce small ion currents possibly below the detection limit of a Faraday detection system. In (secondary) electron multipliers, (S)EM, the electrons released from a single ion incident on the detector surface are converted into a larger electron current using multi-stage amplifying of the electron current. A performance feature of the EM is the gain defined as the ratio of the output electron current to the ion current incident on the entrance to the EM and the noise associated with that gain. The gain can range from 10^2 to 10^7 depending on EM design and operating high voltage. Three types of electron multipliers are used in RGA: discrete dynode, continuous dynode, and microchannel plate electron multiplier. For each of the multiplier types, variants are available that operate in either analog output mode or pulse-counting mode. In the analog mode, a fixed gain ranging from 10² to 10⁶ is set by adjusting the applied high voltage. The output electron current is linearly proportional to the ion current incident on the EM. The source of electrons for the output is the resistive layer applied either to the string of dynodes or the inside surface of the EM through which a current flows. The dynode current is the applied high voltage (1–3.5 kV depending on EM) divided by the dynode resistance (\sim 100 M Ω). For very high ion and electron currents, the EM output can be less than predicted by a linear response. This signal droop is due to depletion of electrons in the dynode current layer when EM output current is greater than a few per cent of the dynode current. A practical limit for analog EM output is $10^{-6}-10^{-5}$ A (with some signal droop) while the lower detection limit is due to the electrical noise of the electrometer measurement which is 10⁻¹³ A, giving an ion current measurement dynamic range of 107.

In pulse counting mode, the EM design has a very high gain ($\sim 10^7$) such that each arriving ion produces a large output pulse of electrons. The pulses can be amplified and those pulses exceeding a threshold value are counted. The ion current measured then is related linearly to the count rate of ions. Deviation from linearity occurs when the count rate grows too high and a second ion arrives during the dead time τ (~10 ns) while the first is being counted. Therefore, the observed count rate of ions is

$$I_{\text{measured}} = I(1 - I\tau) \text{ in s}^{-1},$$
 (14.18)

or for low count rates.

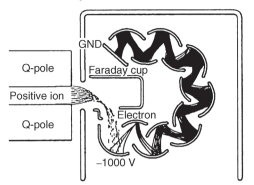
$$I \approx \frac{I_{\text{measured}}}{1 - I_{\text{measured}} \tau} \quad \text{in s}^{-1}.$$

For $\tau = 10$ ns, the measured count rate of 10^7 s⁻¹ is 90% of the true count rate which corresponds to an ion current of $10^7 \text{ s}^{-1} \times 1.6 \times 10^{-19} \text{C} = 1.6 \times 10^{-12} \text{A}$ as a practical limit for ion counting. For an RGA sensitivity of 5×10^{-6} A Pa⁻¹, this corresponds to a pressure limit of 3×10^{-7} Pa $(3 \times 10^{-9} \text{ mbar})$. The real advantage of the ion counting technique is the extremely low noise when no ions arrive. Typical dark-current count rates of $< 1 \,\mathrm{s}^{-1}$ are limited by stray photons and cosmic rays. This corresponds to 3×10^{-14} Pa $(3 \times 10^{-16}$ mbar) detection limits applicable in XHV monitoring applications in accelerators and electron synchrotron storage rings.

14.2.5.3 Discrete Dynode Electron Multipliers

Figure 14.21 shows a schematic representation of a discrete dynode electron multiplier (DDEM). Numbers of dynodes vary according to application from 12 in miniature DDEM [53] to 17 dynodes in low-noise high-gain units for pulse counting, Figure 14.22. Discrete dynodes are typically made of Cu-Be (2%) or Ag-Mg (2%-4%) activated in air to form a stable beryllium-oxide (or manganese-oxide) film on the surface which controls the secondary electron yield. This layer can become chemically altered or contaminated during use in a mass spectrometer, giving rise to changes in multiplier gain. The accelerating potential between each stage is 50-100 V for increased electron current. Two factors determine the electron gain *G* of the multiplier:

- 1) Conversion rate of the incident ions to electrons at the first dynode or point of incidence on a continuous dynode device. The number of electrons released in this primary event, P, is typically between 1 and 5 depending on ion energy, mass, molecular structure and even ionization potential of the gas species.
- 2) Electron gain per stage, q, as the secondary electrons are accelerated between dynodes or through the continuous dynode structure.



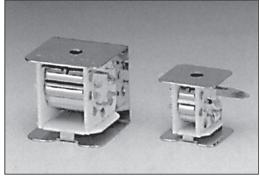


Figure 14.21 Miniature discrete dynode electron multipliers with *Faraday* cup for use with small mass spectrometers. Length and width of the smaller system are only about 10 mm each. Courtesy of *Hamamatsu Corp*.

This is summarized in the relation

$$G = pq^n, (14.19)$$

where n is the number of discrete dynodes following behind the initial conversion dynode.

14.2.5.4 Continuous Dynode Electron Multipliers (CDEM)

In Figure 14.23, the continuous dynode film is formed on the inside of a glass horn-like structure with the resistive film (approximately $10^8\Omega$) extending to the outside on both ends to provide connection points for high voltage and ground. The semiconductive film on a CDEM is chemically stable and adsorbed gases are desorbed under continuous use thus producing a more stable gain over time than many discrete dynode EMs. The nonconductive zone indicated is on the



Figure 14.22 SEM with 17 discrete CuBe dynodes. The resistor chain containing individual resistors of 1 M Ω is distinguishable in the right image. Without connectors, the height of the SEM is 90 mm. Courtesy of *Inficon Inc.*

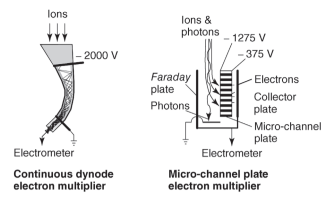


Figure 14.23 Diagrams of EM operation with a are caused by photons from the ion source CDEM (left) and an MCP (right). The MCP features off-axis FC operation by applying the anode voltage of the RGA to the MCP. Off-axis current [56]. ion detection avoids baseline offsets, which

plasma striking the FC collector, ejecting photoelectrons, and thus, creating a false ion

outside of the glass structure and isolates the ends electrically. The applied voltage on the horn end is between -1000 and -2000 V with 10-20 μA of resulting current through the film [54]. This establishes the potential gradient inside the channel. This gradient transports and amplifies secondary electrons from the first ion-to-electron conversion event to the capturing of the electrons by a collector plate connected to the electrometer/preamplifier. The output electron current of the CDEM is $10^2 - 10^6$ times the ion current at the first dynode. This is easily measured with an electrometer with lower input resistance than that required by the Faraday detector, resulting in shorter time constants and larger signal-to-noise ratios. High ion currents on the EM and electron currents through the EM are reported to degrade gain. EMgain loss by a factor of 10 can occur for a total transported electron charge of 1000 µAh [55]. Additional gain loss can occur from deposits, especially hydrocarbons, from an ion beam or from contamination. Thus, it is preferable to keep the EM operating in a clean vacuum to minimize contamination and prolong its lifetime.

14.2.5.5 Microchannel Plate Detectors

The gain of a continuous dynode EM depends on the length-to-diameter ratio, which allows miniaturizing this type of EM. Combining a large number of small channels (5-25 µm in diameter) in a planar array results in a microchannel plate (MCP) where miniature continuous dynode EMs in parallel form a compact multiplier detector for ions. The short length of the microchannels limits the applied voltage to about 1200 V producing a gain of up to 25 000. Figure 14.23 shows a schematic of a microchannel plate EM in combination with an off-axis FC detection plate [56]. The front of the MCP is biased to -1200 V while the collector side of the MCP is biased from -600 to -50 V in order to vary the gain of the EM from 100 to 25 000. The electrons exit and are collected on the electron collector assembly, which is connected to a bipolar electrometer. This same EM assembly can collect ions in FC mode if the high voltage to the EM is turned off and the entrance side of the MCP is biased with the anode voltage to deflect ions onto the FC plate. The signal from the bipolar electrometer when the MCP produces electrons is inverted to produce a positive response to ion current. The EM gain factor G can be measured by measuring the same (small) ion peak in FC mode and EM mode. For correct results, the net peak height obtained by subtracting possible zero-point drifts is taken:

$$G = \frac{I_{\rm EM} - BS_{\rm EM}}{I_{\rm FC} - BS_{\rm FC}},\tag{14.20}$$

where the net peak height for EM mode is I_{EM} , the baseline with EM on is BS_{EM} , while the net peak height for the FC mode is I_{FC} , and the baseline with EM off is BS_{FC} . One of the positive features of the MCP is tolerance to operating pressures as high as 1 Pa for MCPs with very small pore sizes (5 µm). This allows operating an EM with high-pressure RGAs over the full range of RGA operation [57]. At such high pressures, the ion source plasma containing ions and electrons also emits photons that can shine through the quadrupole onto the Faraday cup. The MCP in Figure 14.23 has an off-axis Faraday plate and a grounded photonbeam stop to terminate the photons without recording false ion current. In FC operating mode, the EM high voltage is turned off and the MCP is biased to anode potential, which is sufficient for deflecting the ions to the Faraday plate and electrometer. Results for ion current are accurate without requiring (difficult) correction of the photo-electron current.

14.2.6

Software for Mass Spectrometer Control

An essential element of a modern RGA is the command set and software that controls operation of the instrument. The software is literally the front panel for controlling operation and displaying results of operations such as scanning a mass spectrum, monitoring selected peaks over a period of time, leak detecting, and acquisition of process data with ion current versus time for a specific wafer.

Analog Scan, Ion Current versus Mass

The fundamental output of mass spectrometry data is mass spectra such as the ones shown in Figures 14.8 and 14.15. Any software that controls the analog scan must provide means to define the mass range to be scanned, the number of measurements per amu, and the integration time for ion current measurement. For the RGA electronics, these commands provide the starting mass, mass scanning rate, ion-measurement times, and ending mass necessary to acquire the

data and display the spectrum as the scan progresses. The minimum number of points per amu required to identify the peak shape for a QMS is 5. Some RGAs and their software provide 10, 25, or even 100 points per amu thus yielding more detailed information on ion transmission capabilities, effects of contamination, and information on abundance sensitivity. The tradeoff is time to obtain a spectrum scale with the higher number of points per amu. With a magnetic sector mass spectrometer, the peak width is proportional to the mass such that lowmass peaks are very narrow. The step size for the voltage varies according to Eq. (14.17).

Mass scans with multiple points per amu take more time than usually available for monitoring vacuum processes. Then, one point per amu yields the best results. This measuring mode is often referred to as scan bar graph. Here, the goal has to be that each single point represents the maximum of the ion current. Thus, a stable mass scale is required calibrated to locate the maxima of nominal mass. With this calibration performed with an appropriate known gas, users can reliably survey the entire mass range at each nominal mass. This is particularly valuable when looking for unknown gases and can be done in a matter of seconds. With repetitive scans, any desired mass can be selected for display as a trend plot of intensity versus scan (time).

14.2.6.2 Selected Peaks, Ion Current versus Time

After the key masses that characterize the process are identified, selected masses can be scanned much faster to show specific information. Thus, monitoring the course of complex processes is possible, Figure 14.25.

14.2.6.3 Leak Detection Mode

Leak detection done with an RGA is principally done with helium as the selected ion mass. However, flexibility and broad measuring range of modern RGAs allow other test gases such as H2 or SF6. Much software available provides user features that indicate the magnitude of a leak with the pitch of a sound as well as a trend plot to show the history of helium-signal intensity.

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Further Applications of Mass Spectrometers

Apart from traditional residual-gas analysis and process gas analysis, mass spectrometry can provide additional important information on vacuum processes. MBE (molecular beam epitaxy) systems are capable of directly analyzing molecular beams from selected evaporation sources using appropriate apertures and a crossbeam ion source. Results are used to control the sources for producing stoichiometric coating compositions [27]. The high sensitivity of a mass spectrometer even allows controlling processes with extremely low deposition rates of less than 0.01 nm s⁻¹ [58,59]. Combinations of QMS and electrostatic energy filters are also used for plasma diagnosis (Figure 14.24) [60,61], and for controlling etch processes (finish point determination), Figure 14.25.

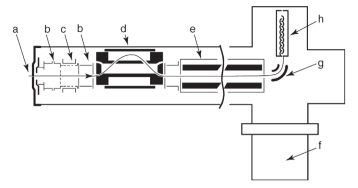


Figure 14.24 Differentially pumped QMS for combined ion and neutral-particle detection in plasma processes. (a) Electrically isolated extraction aperture adjustable to any desired potential, (b) ion lenses, (c) electron-collision ion source with controllable electron energy for process-gas analysis and chemical radical

detection, (d) electrostatic energy filter CMA (cylindrical mirror analyzer), (e) quadrupole analyzer with a mass range up to 2048 amu, (f) turbomolecular pump, (g) ion deflection unit for reduced photon-induced background effects, (h) liftable SEM for detecting positive and negative ions. Courtesy of *Inficon Inc.*

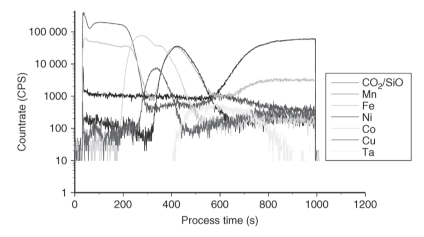


Figure 14.25 Count rates for selected constituents in argon etching of a wafer with complex layer architecture. Analysis was carried out using a QMS with preconnected ion

optics. Such analyses are vital for finding the correct termination times in etching processes. Courtesy of *Inficon Inc.*

14.3 Partial Pressure Measurement Using Optical Methods

Since the 1990s, optical methods for partial pressure analysis have found growing interest because the swift progress in optical storage and communication technology provides laser sources, mirrors, detectors, and so on that could

promote future development of affordable optical partial pressure gauges. Certain leak detectors already utilize absorption methods (Section 14.4.6.3).

Common molecules in high-vacuum systems include H₂O, CO, CO₂, CH₄, and complex hydrocarbons. The diatomic molecules show rotational-vibration transitions in the near- (wavelength $\lambda > 0.7 \, \mu m$) and mid-infrared (IR) range (> 2 μm), and thus, are available to IR absorption measurements. Absorption of an isolated IR beam in a gas follows Lambert-Beer's law:

$$I(\lambda^{-1}, L) = I_0 \lambda^{-1} \exp(-S\Phi(\lambda^{-1} - \lambda_0^{-1})nL), \tag{14.21}$$

where λ^{-1} is the wave number, $I(\lambda^{-1}, L)$ is the intensity transmitted through the gas after the absorbing path $L, I_0(\lambda^{-1})$ is the initial intensity, n is the number density of the absorbing species, S is the wave-number-independent line strength per unit molecule density, and Φ is the so-called line shape function, normalized to

$$\int_{-\infty}^{\infty} \Phi(\lambda^{-1} - \lambda_0^{-1}) d\lambda^{-1} = 1.$$
 (14.22)

The product $S\Phi n$ is usually referred to as the linear absorption coefficient.

Thus, measuring the transmitted intensity with respect to the wave length (wave number), and taking into account the known line strength S, the partial pressure of the absorbing gas is obtained from

$$p = \frac{kT}{S(T)\Phi(\lambda^{-1} - \lambda_0^{-1})L} \ln \frac{I_0(\lambda^{-1})}{I(\lambda^{-1})}.$$
 (14.23)

The advantage of this optical measuring approach is that the absolute partial pressure is obtained directly from measurement without requiring any previous calibration, as is necessary when using, for example, a QMS.

Investigations using standard lock-in technique and a multiple-reflection cell showed [62,63] (Figure 14.26) that measuring of CO partial pressures of 10^{-5} Pa (λ approximately 4.6 µm) is possible with a measuring uncertainty of only 3% at 10⁻² Pa. Here, total pressure can be as high as 1 kPa, in a pressure range not available for QMS (Section 14.2). For CO2 near 30 Pa a measurement uncertainty of only 0.5% was achieved [64].

McAndrew and Inman used IR absorption for humidity measurements of high-purity gases and reactive gases for semiconductor industry in the ppb range [65,66], and to test the cleaning methods used for etch chambers [67].

Humidity and other partial pressures can also be measured precisely with a method referred to as cavity ring-down spectroscopy (CRDS) [68-70] (Figure 14.27). Here, the absorption spectrum, and thus, n and p, are derived accurately from the course of a laser light signal fading with time from a resonator after introducing a pulse.

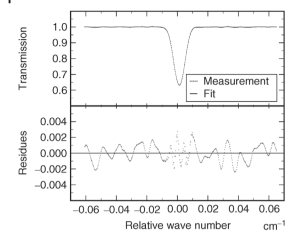


Figure 14.26 Normalized absorption signal for the R(10) line ($\lambda^{-1} = 2183.22 \, \text{cm}^{-1}$) of CO in the fundamental oscillating-rotating band at $p = 1.07 \times 10^{-2}$ Pa. L = 27.236 m. The lower image shows the residues with respect to the theoretical adopted curve [62].

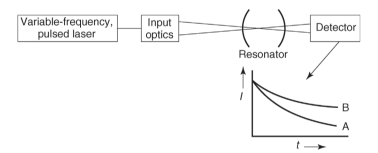


Figure 14.27 Cavity ring-down spectroscopy for quantitative absorption and partial pressure measurements. The course of a fading laser light signal in a resonator with time is

measured. The stronger the slope (case A), the more absorption occurs (absorption is higher for A than for B).

Furthermore, resonance-enhanced multiphoton ionization of gas particles, combined with a TOF (time-of-flight) mass spectrometer, has been used for partial pressure analysis, and this, down to 10^{-10} Pa [71].

The disadvantage of the presented methods is that they only measure a single gas species, which, in addition, must be known ahead of time. However, using Fourier transformation spectrometers, several gases have been simultaneously detected in a CVD process [72]. Currently, narrow-band sources are being developed (OPOs, optical parametric oscillators) that provide means for spanning the entire spectrum in IR. Thus, there is a chance that in the future it might be possible to perform a wavelength scan instead of a mass scan to characterize a vacuum gas mixture.

14.4 **Leak Detectors**

14.4.1

Basic Principles and Historical Overview

Leak detectors are instruments capable of detecting a particular tracer gas. Detection can take place in the atmosphere or in the vacuum. For the former, tracer gas concentration is measured. The latter involves measuring a flow of tracer gas.

Helium is the most commonly used tracer gas (see Section 19.1.5). Detecting a particular tracer gas has the tremendous advantage that leaks can be detected so small that their contribution to a change in total pressure of a system is below any detection limit. In such cases, leaks do not produce sudden increase or drop in pressure but rather lead to long-term damage to the system or process. In vacuum technology, small leaks represent a very important issue because, at low pressures, even very small gas flows from leaks can have tremendous impact on vacuum processes (e.g., in semiconductor manufacturing).

Using helium and its detection with a mass spectrometer as a means of detecting leaks was already suggested in the 1940s. The first detection systems were not yet leak detectors as known today. They were relatively small mass spectrometers attached to the tested vacuum system via a throttling port to reduce the introduced gas flow so that the mass spectrometer could operate at pressures below 10⁻² Pa. Nier built one of the first mass-spectrometer helium leak detectors. He used a sector field spectrometer, which, for the first time, was not made of glass but of metal, and thus, was comparably robust [73]. A publication by Nerken [74] covers the history of helium leak detectors.

1442 **Helium Leak Detectors**

14.4.2.1 Requirements and General Functions of Vacuum Leak Detection

Generally, a leak detector has to be capable of quantitatively measuring a flow of tracer gas. For this, helium leak detectors utilize a mass spectrometer requiring a pressure less than 10⁻² Pa for smooth operation. Therefore, leak detectors are equipped with a high-vacuum pump system for producing and maintaining this pressure (often, this pumping system is already sufficient for evacuating smaller testing volumes down to the required inlet pressure of the leak detector). Test samples are either restricted to components that allow evacuation down to a certain vacuum pressure (inlet pressure of the leak detector), or the leak detector is attached directly to a vacuum chamber enclosing the specimens filled with tracer gas. In both cases, the leak detector measures the helium gas flow into the vacuum and displays the results as leak rates.

An electrical analogy may be used for better understanding the measuring principle of such leak detectors (Figure 14.28). Consider measuring an electrical current with a voltmeter. The voltmeter is analogous to the partial

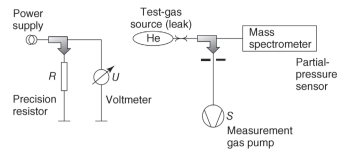


Figure 14.28 Electrical analogy to the measurement principle of a helium leak detector.

pressure gauge (mass spectrometer) in the leak detector, and the electrical current corresponds to the flow of tracer gas. Such an electrical measurement requires a precision resistor at which a voltage falls off that is proportional to the electrical current (*Ohm's* law). This voltage is measured with the voltmeter. Analog to this measuring resistor, the leak detector uses a conductance (aperture) producing a partial pressure proportional to the helium pressure. The mass spectrometer monitors the partial pressure. Electrical ground is the analogy to the pump in a vacuum system to where every gas atom ultimately travels.

14.4.2.2 Helium Sector Field Mass Spectrometers

Helium leak detectors use mass spectrometers as partial pressure sensors. Most common are magnetic sector mass spectrometers; only few manufacturers use QMS. The general operating principles of both spectrometers are described in the first section of this chapter. The main reason for the predominant use of sector field spectrometers in helium leak detectors is the relatively high insensitivity to contamination and total pressure. This characteristic is mainly due to the comparably high ion energy of several 100 eV, as opposed to approximately 10 eV in a QMS. This high energy reduces the influence of interference on the ion paths, caused by insulating contamination layers on the walls, and thus, on the sensitivity and resolving power. These properties are particularly important in a leak detector operating at high pressures and with high requirements in terms of signal stability.

A sector field spectrometer for a helium leak detector has certain characteristic features due to its special operating tasks. While robust and simple design is desired, the predominant characteristic should be high partial pressure sensitivity for helium, and this, at high total pressures also, that is, the instrument should be able to measure very small concentrations or partial pressure ratios. Furthermore, temperature and long-time stability are important characteristics of measuring instruments such as leak detectors.

These requirements are usually met by a double-focusing system with permanent magnets (see Figure 14.29). Double-focus means that the magnetic sector

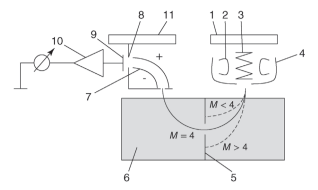


Figure 14.29 Diagram of a double-focusing mass spectrometer for helium leak detectors. 1 Flange of ion source; 2 filaments (operating alternately); 3 anode (heated if necessary); 4 shielding extractor cap; 5 intermediate aperture; 6 magnetic field (magnetic sector field,

momentum dispersion); 7 cylindrical capacitor (electrical sector field, energy dispersion); 8 exit aperture: 9 ion collector plate: (10) preamplifier system (current/voltage converter); (11) flange of ion collector system.

field, apart from focusing ions with the same momentum (i.e., the same mass for equal velocities), also sorts out ions of equal energy (i.e., the same velocity for equal masses) into an electrical sector field (cylinder condenser). Standard mass spectrometers perform the energy selection before the ions enter the magnetic field in order to enhance resolving power. In contrast, helium mass spectrometers utilize it after magnetic separation. This is intended to suppress wrong ions at high total pressures, approaching the ion collector due to scattering at gas molecules although they do not meet separating prerequisites.

Today, so-called suppressors are predominantly used instead of expensive cylinder condensers. They include an aperture charged to the anode potential of the ion source. This prevents any ion having lost energy in collisions from reaching the ion collector. In addition to this electrical separation, certain spectrometers include intermediate apertures that mask any non-helium ions and thus improve resolving power. The described measures for helium mass spectrometer design guarantee full sensitivity at pressures up to 10^{-2} Pa (as opposed to analytical sector field spectrometers requiring total pressures of less than 10⁻⁴ Pa). Typical helium sensitivity is several 10^{-6} A Pa⁻¹ (10^{-4} A mbar⁻¹), the lowest measurable partial pressure is approximately 10^{-10} Pa, and the lowest measurable concentration is approximately 0.1 ppm.

14.4.2.3 Inlet Pressure of Helium Leak Detectors

In contrast to measurements of electrical currents, measuring tracer gas flows additionally faces disturbance caused by the total pressure. Thus, a leak detector has a maximum tolerable inlet pressure up to which operation is possible. During measurement, pumps in a leak detector fulfill several tasks: apart from acting

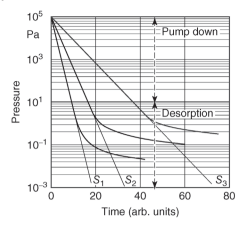


Figure 14.30 Pressure trend during pump-down of a test object with three pumps of different pumping speeds $S_1 > S_2 > S_3$.

as a measuring-gas pump (analogous to electrical ground) they assure that the mass spectrometer continuously operates at its working pressures below 10^{-2} Pa.

When measurement starts, the leak detector and in some cases an additional auxiliary pump evacuate the specimen down to the tolerable inlet pressure. The time required for this initial step should be as short as possible, thus ideally, approaching zero. For this, high pumping speeds are beneficial. However, it should be noted that water vapor desorption from the walls of the specimen becomes noticeable at pressures below approximately 10 Pa. This means that the pressure now follows a 1/t-characteristic and does no longer drop exponentially as in the higher pressure range, see Figure 14.30. Therefore, low specimen pressures should be avoided if short measuring cycles are desired. The following sections on direct flow and counterflow leak detectors describe how different measuring principles cope with this requirement.

14.4.2.4 Time Response of Helium Leak Detectors

The response time of a leak detector, that is, the time until the indicated leak rate rises to a certain fraction of full scale deflection, depends on the time constant τ of the test volume V_0 . When $S_{0,\text{He}}$ denotes the pumping speed for helium at the inlet port of the leak detector, then τ is given by

$$\tau = \frac{V_0}{S_{0,\text{He}}}. (14.24)$$

Within this time, the leak rate signal rises to 63% of its final value. The time required for obtaining 95% of the final value is three times as long (the helium tracer gas must be present during this time, i.e., constant spraying or integral testing is required). A common mistake in local leak detection is premature termination of spraying, producing readings of only a fraction of the true leak rates or even zero readings.

Response time is determined by the pumping speed for helium at the leak detector inlet port. This pumping speed has to be differentiated from the pumping speed of the auxiliary or roughing pump effective during pre-evacuation. The pumping speed for helium at the inlet port is always less than the net pumping speed of the high-vacuum pump because connecting valves reduce pumping speed. It is also smaller than the net pumping speed for air of the auxiliary pump because the pumping speed of a roughing pump is lower in the molecular range (low pressures) than in the viscous range.

14.4.2.5 Operating Principles of Helium Leak Detectors

After helium leak detectors operated solely according to the electrical analogy shown in Figure 14.28 for many years (direct flow principle), the idea of the counterflow principle emerged in the 1970s. This development did not require any liquid helium in the cold trap of leak detectors and thus paved the way for an industrial application of helium leak detectors.

Direct Flow Leak Detectors

The term direct flow leak detector indicates that here the entire gas flow O from the specimen is fed through the high-vacuum pumping system, and thus, the tracer gas helium directly enters the mass spectrometer. Corresponding to the electrical analogy, the helium flow $q_{\rm He}$ is converted into a helium partial pressure $p_{
m He}^{
m MS}$, which is recorded by the mass spectrometer. In this case, the indicated leak rate q_i for a given high-vacuum pumping speed S_{He}^{HV} ,

$$q_{\rm i} \propto p_{\rm He}^{\rm MS} = \frac{q_{\rm He}}{S_{\rm He}^{\rm HV}},$$
 (14.25)

where the sensitivity is determined by the pumping speed of the high-vacuum pump $S_{\text{He}}^{\text{HV}}$. As opposed to the electrical analogy, a large amount of other gases, water vapor in particular, emanates from the specimen. After pumping starts, these other species prevent rapid establishment of a pressure below 10⁻² Pa, and thus, delay measuring readiness of the mass spectrometer. In order to reduce this time delay, direct flow leak detectors usually require a cold trap with liquid nitrogen, in which water vapor condenses thus providing a pressure of less than 0.01 Pa in the mass spectrometer. As an accompanying effect, the cold trap additionally prevents contamination of the mass spectrometer with oil from the diffusion pump or other condensable gases possibly released by insufficiently cleaned specimens.

Figure 14.31 only shows the detection system. In practice, an auxiliary pump is required for pre-evacuating the specimen because the high-vacuum pump features sufficient pumping speed to maintain vacuum in the specimen only below a pressure of approximately 1 Pa. In certain cases, bypass lines and additional valves are used to run the roughing pump as auxiliary pump and to temporarily disconnect the high-vacuum pump from the fore vacuum. Then, however, specimen volumes are limited because the high-vacuum pump cannot operate indefinitely without a roughing pump.

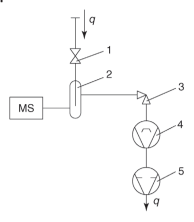


Figure 14.31 Detection system of a direct flow helium leak detector: the test gas flows through the cold trap and entirely passes through the high-vacuum system. Total pressure in the mass spectrometer is the sum of the test-gas partial pressure and the partial

pressures of permanent gases, water vapor, and pump oil condensed in the cold trap.

1 Inlet valve; 2 cold trap; 3 high-vacuum valve; 4 high-vacuum pump (diffusion pump, or possibly, turbomolecular pump); 5 fore-vacuum pump.

Advantages of direct flow leak detectors include high pumping speed for helium at the inlet port and thus quicker response times even for larger specimen volumes as well as very low detection limits down to the $10^{-10}\,\mathrm{Pa}\,\ell\mathrm{s}^{-1}$ range. On the other hand, handling liquid nitrogen is complicated, and managing the transition from pre-evacuation to measuring operation is difficult to automate (this is because, for varying water vapor contents, the pressure at which the inlet valve to the high-vacuum pump can be opened is not defined and has to be determined by trial and error). Both disadvantages prevented widespread use of direct flow leak detectors in industrial applications.

Counterflow Leak Detectors

In counterflow leak detectors, the total gas flow q from the specimen is fed only through the roughing pump from where it flows backward, in counterflow, via the high-vacuum pump and into the mass spectrometer (Figure 14.32). Here also, the helium flow q_{He} is transformed into a helium partial pressure p_{He}^{F} , in analogy with the electrical example, but over the fore-vacuum pump with the pumping speed S_{F} . This helium partial pressure produces a helium partial pressure in the mass spectrometer reduced by a factor equal to the compression factor K_{He} in the high-vacuum pump. Thus, the indicated leak rate q_{i} follows

$$q_{\rm i} \propto p_{\rm He}^{\rm MS} = \frac{q_{\rm He}}{K_{\rm He} S_{\rm He}^{\rm F}}.$$
 (14.26)

Such a counterflow leak detector has the same sensitivity as a direct flow leak detector if the denominator in Eq. (14.26) is equal to the denominator in Eq. (14.25), that is, if the product of fore-vacuum pumping speed and

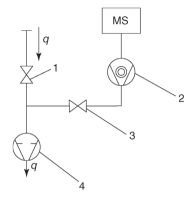


Figure 14.32 Detection system of a counterflow helium leak detector: the test gas flows only through the fore-vacuum pump. The test gas partial pressure in the mass spectrometer (MS) is the sum of the partial pressure of all gas species and helium partial pressure. The molecular-weight-dependent compression of

the turbopump generates strong enrichment of helium and air; most of the water vapor and pump oil remain in the fore-vacuum region. 1 Inlet valve; 2 high-vacuum pump (preferably a turbomolecular pump); 3 fore-vacuum valve; 4 fore-vacuum pump.

compression in the turbopump is equal to the pumping speed of the highvacuum pump in a direct flow leak detector.

A cold trap with liquid nitrogen is dispensable in such an arrangement because any gas with higher molecular weight than helium is kept away efficiently from the high vacuum due to the exponentially increasing compression (compare Chapter 10). In practice, the speed of turbomolecular pumps is adjusted so that, for example, $K_{\text{He}} = 50$. Then, typical values for water and nitrogen are $K_{\rm H_2O} = 4000$ and $K_{\rm N_2} = 30\,000$, respectively. This means that the partial pressure ratio of helium to nitrogen in the mass spectrometer is 600 times the ratio in the fore vacuum, and still 80 times as high in relation to water vapor. Such conditions yield water vapor suppression comparable to the suppression obtained when using a cold trap.

The main advantage of counterflow leak detectors is that they obtain measuring readiness very quickly without requiring liquid nitrogen because, during pumpdown, the specimen is only brought down to the fore-vacuum pressure required by the turbomolecular pump (10 Pa). Thus, when measuring starts, the pressure in the specimen is far from the desorption range (Figure 14.30). In a counterflow leak detector, the transition from pre-evacuation to measuring is independent of the water vapor content and thus simple to control according to the pressure. For the first time, this principle allowed fully automating leak detection processes.

Requirements on the fore pump in terms of helium tightness and stable pumping speed are higher compared to direct flow leak detectors because the fore pump now replaces the high-vacuum pump as measuring pump. Recently, compound turbomolecular pumps were able to reduce the requirements on the fore-vacuum pump because the *Holweck* stages in the turbomolecular pump with their high compressions serve as stable fore pumps when combined in series with the fore-vacuum pump.

When testing large specimen volumes, the helium pumping speed of a fore pump (directly active at the inlet port to the leak detector) is not sufficient for producing low response times of several seconds. This disadvantage is eliminated by connecting an additional high-vacuum pump, referred to as booster pump, to the inlet. To date, use of a booster pump was economical only when it was part of the counterflow pump. A number of different solutions were developed. All of them divide the turbopump rotor into several sections with appropriate additional inlet ports [75]. Figure 14.33 shows an example of an innovative approach: here, a relatively small $(50 \,\ell\,\text{s}^{-1})$ separate turbomolecular pump was connected to the inlet as booster pump. The uncommon approach is to avoid a (large and expensive) inlet valve above this pump and thus utilize the full pumping speed at the inlet. The turbopump operates intermittently. It is vented (and throttled) together with the specimen and not restarted until the following preevacuation cycle starts.

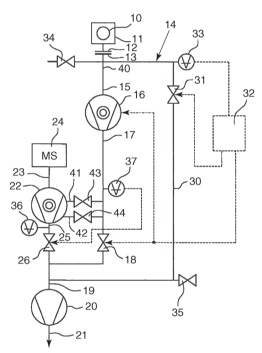


Figure 14.33 Vacuum diagram of a modern helium leak detector with a booster turbomolecular pump at its inlet and a counterflow turbopump with molecular drag stage and several intermediate inlets (UL5000, Inficon Inc.). The figure is an excerpt from the patent application DE10319633 describing the use of a booster turbopump without inlet valve.

Important elements are - 13 inlet flange; 31 roughing valve; 33 inlet pressure gauge; 34 vent valve; 16 booster turbopump; 37 intermediate pressure gauge; 43 and 44 inlet valves; 22 counterflow turbo pump; 24 mass spectrometer; 36 fore-vacuum pressure gauge; 35 purge gas inlet; 20 scroll fore-vacuum pump.

Detection limits of modern counterflow leak detectors typically lie in the low $10^{-8} - 10^{-9} \text{ Pa} \ell \text{s}^{-1} \ (10^{-10} - 10^{-11} \text{ mbar} \ell \text{s}^{-1})$ range, and thus, nowadays are comparable to early direct flow leak detectors. For practical use, it was essential to extend the measuring range toward high leak rates, and additionally, inlet pressures of up to several hPa have become common, indispensable for industrial applications.

Wet versus Dry Leak Detectors

Helium leak detectors considered so far use a standard high-vacuum pump system with a diffusion or turbomolecular pump and an oil-sealed sliding vane rotary pump. Ca. since 1990, semiconductor industry has massively promoted using so-called dry pumping systems in chip manufacturing in order to avoid contamination. This need also called for the development of dry leak detectors. These are helium leak detectors operating according to the counterflow principle but with dry pumps, for example, diaphragm pumps, as fore pumps [76]. Using such pumps became feasible after so-called compound turbomolecular pumps with an additional Holweck stage were utilized that tolerate the relatively high ultimate pressures (typically 1 hPa) of such dry fore pumps.

However, for helium leak detection, considerable problems arise from the need to detect very low helium partial pressures in such high base pressures. Difficulties are mainly due to the fading effective pumping speeds for helium under ultimate-pressure operation in the diaphragm pump. Dry leak detectors thus generally include an air inlet for flooding gas in the fore vacuum to ensure gas transport, and thus, provide acceptable response times. Figure 14.34 shows atypical vacuum block diagram of such a leak detector designed as a portable instrument.

The dry leak detector shown in Figure 14.34 compensates for another problem of dry pumping systems namely the instability of the diaphragm pump's pumping speed by modulating the incoming gas flow. A phase-sensitive lock-in method is used for detecting the ion current in the pre-amplifier of the mass spectrometer. Here, only such currents are regarded that conform to the modulated gas flow in terms of frequency and phase. Any dissimilar variations of the ion current are effectively suppressed. This extends the detection limit down to $10^{-9} \text{ Pa } \ell \text{ s}^{-1}$.

Utilizing more sophisticated dry roughing pumps such as scroll pumps or piston pumps [77] with ultimate pressures around 1 Pa yields detection limits as provided by wet leak detectors. The leak detector in Figure 14.33 has a detection limit in the range of $10^{-11} \, \text{Pa} \, \ell \, \text{s}^{-1}$. However, to date, such pumps are used only for testing semiconductor-manufacturing equipment because they are expensive and are susceptible to considerable wear.

14.4.2.6 Sniffing Devices for Helium Leak Detectors

Helium leak detectors are also used for overpressure leak detection. For this, they are equipped with a sniffing device. Section 19.1.2 explains the principle of overpressure leak detection.

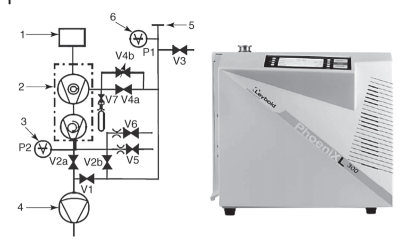


Figure 14.34 Vacuum diagram and view of a portable dry counterflow leak detector with compound turbomolecular pump and inlet flow modulation (*Leybold* PhoeniXL300): 1 mass spectrometer; 2 compound turbopump; 3 fore-vacuum gauge; 4 diaphragm fore-

vacuum pump; 5 inlet flange; 6 inlet pressure gauge; V1 roughing valve; V2a and V2b counterflow valves; V3 vent valve; V4a inlet valve; V4b inlet control valve; V5 HV-vent valve; V6 purge gas valve; V7 valve for internal test leak.

The sniffing device for a helium leak detector contains a hose with a fine tip including a throttling port for admitting a stream of air into the detection system of the leak detector. If this stream of air takes up atmospheric helium, this helium is detected in the leak detector's mass spectrometer. The detection limit is determined mainly by the constancy of the atmospheric helium.

The first sniffing devices included a small throttling valve at the sniffing tip. Later, membranes were used that produced a slight helium-enrichment effect. They were highly sensitive to the helium cloud in front of the leak. Thus, leaks could be measured accurately. However, locating a leak was difficult because readings varied considerably with the distance from the leak. The system was improved by increasing the gas flow through a hose providing laminar flow. Yet, this gas flow was too high for directly entering the detection system of the leak detector so only a small fraction of the sniffing flow could be allowed into the high vacuum of the mass spectrometer. This equipment is known under the brand name *Quicktest (Leybold Vacuum)*. In modern counterflow leak detectors, allowing relatively high gas flows at the inlet, throttling is simply adjusted by selecting a sniffing hose with appropriate length and diameter so that, on one hand, enough gas is taken up at the tip, and on the other hand, direct inflow to the detection system is allowed. This principle paved the way for very simple, robust, but at the same time sensitive sniffing devices.

14.4.2.7 Applications of Mass Spectrometer Helium Leak Detectors

Mass spectrometer helium leak detectors (MSLD) are used in applications that call for either high sensitivity (research) or ISO 9000 certified measuring





Figure 14.35 Mobile dry leak detector for service on semiconductor systems in clean rooms. Left: display and operating unit. Right: overall view.

procedures (industry). Quantitative leak rate measurements usually utilize the vacuum method whereas the sniffing method with search-gas overpressure is common for leak locating. The latter method is convenient for testing any industrial components exposed to overpressure during normal operation as well.

The semiconductor industry uses mobile MSLD as standard preventive maintenance instruments for vacuum coating equipment in clean rooms (Figure 14.35). During fabrication of such systems, stationary MSLD with large pumps rapidly evacuate and inspect the relatively large chamber volumes, often with considerable internal surface area. Today, the semiconductor industry accepts dry pumping systems only, thus leak detectors also face this challenge.

14.4.3 Refrigerant Leak Detectors

14.4.3.1 Design and Operating Principle

Refrigerant leak detectors operate solely on the principle of sniffing. This is because discharge of refrigerant tracer gas into the atmosphere is no longer tolerable due to its ozone-harming characteristics and greenhouse-effect potential. Refrigerant leak detectors use a number of different methods for sensitive detection of refrigerants leaking from filled refrigerant cycles. The most sophisticated instruments, such as helium leak detectors, are equipped with a mass spectrometer. The alkali ion sensor is a simpler instrument that uses an electrochemical principle. A newly introduced method is based on infrared absorption. This section covers refrigerant leak detectors using mass spectrometers. The remaining principles are discussed briefly further below.

Refrigerant leak detectors use QMS only, with a measuring range of 0-200 amu as a partial pressure gauge, because investigated refrigerants produce characteristic lines in this atomic mass range. Counterflow operation is not an option because here refrigerants with masses in the range of 58 (butane) to 127 (SF₆) cannot be separated from other, often lighter gases (particularly water

Table 14.3 Mass spectra of refrigerants: R134a, a completely fluorinated refrigerant (FHC) mostly used in automobile air conditioners, R600/600a = butane and isobutane, pure hydrocarbons (HC) without chlorine and fluorine used in refrigerators, and R22, a refrigerant containing both chlorine and fluorine (FCHC) being phased out soon (in these fractional patterns, the proportionate intensities of mass peaks are normalized to the highest peak of the respective gas species).

Code Designation	Chemical name	Chemical notation	Molar mass	М	%	М	%	М	%	М	%	М	%
R134a	1,1,1,2-Tetrafluoroethane	CH ₂ F-CF ₃	102.03		100		72		63		14		07
R600	Butane	CH ₃ -CH ₂ -CH ₂ -CH ₃	58.12		100		44		32		13		12
R600a	Isobutane	CH(CH ₃) ₂ -CH ₃	58.12		100		37		35		04		17
R22	Chlorodifluoromethane	CHClF ₂	86.47		100		15		12				

vapor). Refrigerants produce distinguished, characteristic individual lines in the spectrum. Thus, detection often relies on a selected individual line. Table 14.3 shows the spectra of typical refrigerants. Individual peak analysis does not require any sophisticated analysis algorithms and provides very short response times, which are important for leak locating.

Obviously, QMS are also capable of detecting helium. Here, the counterflow principle with gas intake in the fore vacuum is applicable, and thus, a refrigerant leak detector for helium is nearly as sensitive as a standard leak detector with a sector field spectrometer. Therefore, a single instrument can be used for helium pretesting as well as for finally testing the filled refrigerant cycle; however, both require sniffing mode.

Refrigerant leak detectors are generally limited to leak locating because sniffing inspection is strongly distance- and velocity-dependent (see Figure 19.2). Practical industrial applications usually practice a semiquantitative test procedure with a fixed threshold value. Apart from visual indication, instruments are required to provide a distinct acoustic alarm. Whereas helium leak detectors usually produce a frequency-variable tone depending on the size of the leak, refrigerant leak detectors require only a simple alarm that starts as soon as the threshold value (rejection limit) is exceeded.

14.4.3.2 Applications

Refrigerant leak detectors with mass spectrometers are used for final testing in air-conditioner manufacturing for automobiles, domestic refrigerators, and deep freezers. They are universal and capable of detecting nearly any gas species. Thus, any newly introduced refrigerant or mixture can be measured easily and, above all, accurately. Hence, in spite of their high cost compared to simple service equipment, such instruments have prevailed as an important factor for quality assurance for modern refrigerating and air-conditioning component manufacturing.

14.4.4

Reference Leaks

Reference leaks are used for checking leak detectors. They are sources of tracer gas, analogous to sources of electrical current (see also Figure 14.28). Similar to the electrical analogy, they consist of a pressurized gas reservoir (analogous to the electrical voltage supply) and a downstream constriction (analogous to the high-value resistor in a source of electrical current). Reference leaks are distinguished mainly by the geometry of their constriction. The latter is designed either as a hole (conductance leak) or as a diaphragm (permeation leak).

14.4.4.1 Permeation Leaks

For helium, permeation leaks are built from quartz-glass membranes (Figure 14.36). Such a membrane is usually designed as a closed glass tube, open at

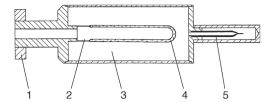


Figure 14.36 Cross section of a helium permeation leak with gas reservoir. 1 Connecting flange, 2 glass to metal seal, 3 helium gas volume, 4 quartz glass tube determining flow conductance, 5 crimped copper tube for helium supply.

one end, and surrounded by a helium reservoir pressurized to several bar. After a run-in period of several hours, such a leak is extremely long time stable and immune to clogging. However, due to the permeation current, the temperature coefficient is exponential, but susceptible to an approximation of about $3.5\%~\rm K^{-1}$ [78]. Helium permeation leaks are mostly produced for around $10^{-6}~\rm Pa\ell s^{-1}$ ($10^{-8}~\rm mbar\,\ell s^{-1}$).

Using appropriate synthetic membranes, permeation leaks can be fabricated for refrigerants. These can be filled with liquid refrigerants, and thus, service life of such equipment is high.

14.4.4.2 Conductance Leaks

Most conductance leaks are designed as capillaries. Glass tubes are drawn out to capillaries, metal tubes are crimped, that is, squeezed with a pair of pliers until the desired throttling effect is obtained. However, smooth glass capillaries with constant diameters provide the most precise leaks because flow conditions are reliably laminar and changes in leakage flow due to turbulent flow transitions are unlikely [79]. Conductance leaks can also be manufactured with compressed metal powders or sintered ceramic. However, such structures are usually not as long-time stable and reproducible as capillary leaks.

The temperature coefficient in capillary leaks is low, approximately 0.3% K⁻¹ [80] so that correction is usually unnecessary. Any conductance leak is susceptible to clogging, and obviously, the smaller the conductance, the higher the danger of clogging. Therefore, particularly when heavy climatic changes are to be expected, such conductance leaks must be kept in sealed containers and not opened until shortly before use. Leak rates in conductance leaks should be greater than 10^{-5} Pa ℓ s⁻¹ (10^{-7} mbar ℓ s⁻¹) in order to prevent clogging.

14.4.4.3 Practical Realization of Reference Leaks

In practice, designs of reference leaks differ considerably due to the different principles and according to application. Figure 14.37 shows selected examples.

Generally, leaks with and without gas reservoir are differentiated. Permeation leaks always require a gas reservoir because after filling they require several hours of run-in before the leak rate is stable. Conductance leaks can also be



Figure 14.37 Selected test leak designs. From left to right: capillary-type leak (range $> 10^{-4} \text{Pa} \, \ell \text{s}^{-1}$) without reservoir; capillarytype leak with manual valve, test gas reservoir, and adjustable upstream pressure for leakage

rates from 10^{-4} to 10^{-2} Pa ℓ s⁻¹; built-in test leak with solenoid valve (permeation or capillary type); permeation-type test leak (range $< 10^{-6} \text{Pa} \, \ell \text{s}^{-1}$) with manual valve and helium reservoir.

produced without gas reservoir, which makes them particularly suitable for simulating leaks in equipment.

14.4.4.4 Calibrating Reference Leaks

Reference leaks are the basis for traceable calibration of helium leak detectors. Thus, calibration is inevitable. General procedures for reference leak calibration are standardized in EN 13192. Three methods are differentiated. Method A: comparison with reference leak, method B: comparison with two reference leaks. and method C: flow measurement by means of capillary measurement.

Method B is used in accredited calibration laboratories within the German accreditation system DAkkS (Deutsche Akkreditierungsstelle). Leaks calibrated by accredited laboratories guarantee that their value can be traced back to a national standard (see Section 15.4). Other countries have similar organizations that guarantee traceable calibration services for reference leaks. For details on the accuracy of helium reference leaks, see [81].

14.4.5

Measuring Characteristics and Calibration of Leak Detectors

14.4.5.1 Leak Detectors as Test Equipment According to ISO 9001

Leak detectors are applicable as test equipment in accordance with ISO 9001. This standard specifies examinations of test equipment on a regular basis in which function and calibration are verified. In helium leak detectors, as opposed to most other test equipment, calibration is not immanent to the instrument itself but to one or more reference leaks used for adjusting a helium leak detector. The reference leaks can be interpreted as sources of helium *current*, in analogy with a source of electrical current. They set free a well-defined flow of helium, to which the reading of the leak detector is adjusted during calibration (see section on reference leaks). It is important for calibration that initially the zero point of the scale is correctly adjusted with the reference leak sealed. Then, the leak is opened and the calibration factor, that is, the multiplier to calculate the correct reference leakage from the reading, is determined after the reading has reached a constant value. Modern leak detectors usually control this procedure on a fully automated basis and often use integrated reference leaks.

14.4.5.2 Calibration Uncertainty

Calibration yields the position and slope of a calibration line. The slope of this line can be interpreted as sensitivity; the position is determined by the zero signal, that is, the reading with the reference leak sealed. Three individual effects influence measuring uncertainty: deviations from the straight line (nonlinearity), deviation from the zero point (zero-point drift), and deviations in the slope (sensitivity drift). The extent of the deviations, caused mainly by temperature variations, determines how often a leak detector requires calibration with a reference leak. Calibration should generally be verified after a change of position and in the event of strong temperature changes at the place of installation. Modern leak detectors are ready for measuring only just a few minutes after they are switched on. However, accurate measurements should not be performed until one or two hours later. Calibration, as well, should not be started any earlier because otherwise the produced result may even be worse than the initial condition.

Note that the standard procedure for adjusting a leak detector is a single-point calibration, with the zero point as the second defining point for the calibration line. This means the detection system is assumed to be linear. For checking the linearity of a leak detector, readings would have to be plotted versus the reference leak rates for a number of reference leaks across the desired range. However, considering the given measurement uncertainties of reference leaks, typically in the range of $\pm 5\%$ to $\pm 15\%$, it seems obvious that no accurate method for linearity checking of a leak detector is available today because the deviations from linear behavior are of the same order of magnitude: nonlinearities in helium leak detectors can be as high as 20%. Thus, for most accurate measurements, a reference leak from the same decade as the measuring results should be selected in order to keep deviations from linear behavior negligible.

The second uncertainty of calibration is the zero point of the calibration line. The thermal shift is the predominant factor for deviations observed in measurements of small leak rates. Instruments usually automatically correct shifting leading to negative values. The only method to prevent shifting to positive values is to check the zero point regularly by sealing the reference leak valve. In practice, however, deviations from the zero signal due to background helium concentrations are usually higher than the thermal zero-point drift of the instrument itself.

The sensitivity of a leak detector is also subject to temperature drift. This drift is determined mainly by the temperature coefficient of the magnet in the mass spectrometer and the properties of the pre-amplifier. In modern instruments, high-grade magnet materials and well-temperature-compensated amplifiers keep these influences low. After stronger changes in ambient temperature, readjustment of a leak detector with a temperature-corrected reference leak is advisable.

14.4.6

Leak Detectors Based on Other Sensor Principles

14.4.6.1 Helium Sniffers with Ouartz Glass Membrane

In the 1960s, the first attempts were made to build helium leak detectors without mass spectrometers. A 1966 Karlsruhe Nuclear Research Center diploma thesis introduced a sniffer for detecting helium by using the increase in total pressure in a sealed high vacuum, measured in a gas discharge or an ion getter pump. For separating, that is, selective helium admission, the system featured a quartz (SiO₂) membrane heated with a tungsten coil [82]. The membrane was fabricated by simply drawing out a thin (approximately 100 µm) glass capillary on the face of a glass tube.

In the 1990s, Varian Inc. introduced the battery-powered HeliTest based on this principle [83]. It uses a thin glass capillary heated with a wire coil as inlet membrane for helium, and a miniature ion getter pump for detection. The system is capable of detecting leak rates down to the range of 10^{-4} Pa ℓ s⁻¹.

In 2005, Inficon introduced the first quartz-membrane leak detector for industrial testing of refrigerator components (Protec P3000), quantitatively detecting helium leak rates down to $2 \times 10^{-5} \, \text{Pa} \, \ell \, \text{s}^{-1}$ and with very short response times.

Here, the SiO₂ membrane is formed by a 7 µm oxide coating on a silicon chip. By means of micromachining, the chip is equipped with numerous etched boreholes that are covered by the SiO₂ layer. The silicon chip is anodically bonded to a glass housing in which the total pressure without any helium is approximately 10⁻¹⁰ Pa, and heated to several hundred degrees centigrade by sputtered platinum heating meanders. The ion current of a Penning discharge in the glass housing is a measure of pressure increase due to introduced helium, and thus, of the incoming leak rate.

This detection principle is only just in the early development stages and an increase in sensitivity can be expected so that leak detectors for helium vacuum detection according to this principle should be available in the future.

14.4.6.2 Halogen Leak Detectors with Alkali Ion Sensors

The halogen leak detector with alkali ion sensor is the classic refrigerant sniffer (the detection principle was also utilized in fine vacuum for leak-sensing probes until use of refrigerants as tracer gas was prohibited). Its sensor detects gaseous halogen compounds with an alkali-ion-emitting electrode heated to 800-900 °C. The emitted current rises in the presence of halogens. Detection sensitivity is in the 1 ppm range. However, due to the strong nonlinearity of the measuring principle, the instrument does not yield absolute values of leak rates but rather requires threshold calibration with a reference leak precisely giving the reject leak rate. This value is taken as trigger value and the instrument only indicates excess values (usually with an acoustic siren). The sensor principle is still quite



Figure 14.38 Sniffer leak detector for refrigerants operating on infrared absorption (the complete detection system is integrated into the sniffer handle).

common; however, it does not differentiate between different refrigerant species and reacts sensitively to disturbing gases (vapors of water, benzene, alcohol, etc.). Due to today's requirements in terms of sensitivity and selectivity, phase-out of these instruments can be foreseen.

14.4.6.3 Halogen Leak Detectors with Infrared Sensors

A new principle for sniffing detection of refrigerants utilizes light absorption in the mid-infrared range. With this principle, a number of modern refrigerants can be detected sensitively and with high selectivity.

Here, the gas is fed through a cell where it is exposed to infrared radiation. A detector determines the absorbed light intensity. By alternately pumping down gas from the vicinity of the leak and from the actual leak, background influences are reduced effectively, making this principle particularly suitable for industrial environments (Figure 14.38).

In contrast to the alkali ion measuring cell, this principle yields considerably higher service lives and provides linear readings. Thus, it is applicable to traceable calibration in industrial testing according to ISO 9000.

References

- DIN 28410 (1968) Vacuum Technology; Mass Spectrometer Partial Pressure Gauges; Definitions, Characteristics, Operating Conditions.
- 2 Basford, J.A., Boeckmann, M.D., Ellefson, R.E., Filippelli, A.R., Holkeboer, D.H., Lieszkovszky, L., and Stupak, C.M. (1993) Recommended practice for the calibration of mass spectrometers for partial pressure analysis. J. Vac. Sci. Technol. A, 11, A22.
- 3 NIST, EPA (1995) NIH Mass Spectral Data Base, Version 1, 1995 (Software File) or Internet search http://webbook.nist. gov/chemistry/ormser.html.
- 4 Summers, R.L. (1969) NASA Technical Note, NASA TN D5285.
- 5 Dempster, A.J. (1918) A new method of positive ray analysis. *Phys. Rev.*, 11, 316.
- **6** Bleakney, W. (1929) A new method of positive ray analysis and its application to

- the measurement of ionization potentials in mercury vapor. Phys. Rev., 34, 157.
- 7 Nier, A.O.C. (1940) A mass spectrometer for routine isotope-abundance measurements. Rev. Sci. Instrum., 11, 212.
- 8 Nier, A.O.C. (1947) A mass spectrometer for isotope and gas analyses. Rev. Sci. Instrum., 18, 398-411.
- 9 Inghram, M.G. and Hayden, R.J. (1954) A Handbook on Mass Spectrometry, Nuclear Science Series, Report No. 14, National Academy of Science, Washington.
- 10 Duckworth, H.E., Barber, R.C., and Venkatasubramanian, V.S. (1990) Mass Spectrometry, vol. 45, Cambridge University Press, Cambridge, UK.
- 11 Austin, W.E., Holme, A.E., and Leck, J.H. (1995) Quadrupole Mass Spectrometry and Its Applications (ed. J.H. Dawson), American Institute of Physics, Woodbury, NY, pp. 121-152.
- 12 Brubaker, W.M. (1970) NASA Report, NASW 1298.
- 13 Hofmann, G.A. (1974) Quadrupole mass spectrometers of various performance groups. Vacuum, 24, 65.
- 14 INFICON, Quadrupolanalysatoren und zugehörige Ionenquellen, Company brochure vina07d1.
- 15 Blessing, J. (1987) Research & Development September.
- 16 INFICON (1992) Closed Ion Source, Bulletin BR31D38K.
- 17 Koprio, J., Muralt, P., Rettinghaus, R., and Strasser, G. (1990) Mass spectrometric detection of low-ppm contaminants in sputter process systems at 10⁻² mbar, using a directly exposed ion source. Vacuum, 41, 2106-2108.
- 18 INFICON, Quadrupolanalysatoren und zugehörige Ionenquellen, Company brochure vina07d1.
- 19 Kohl, W.H. (1995) Handbook of Materials and Techniques for Vacuum Devices, American Institute of Physics, Woodbury, NY, pp. 487-502.
- 20 Ibid. pp. 262-264; 240.
- 21 Cole, C.R. et al. (2004) Contribution and origin of H3O+ in the mass spectral peak at 19 amu. J. Vac. Sci. Technol. A, 22, 2056-2060.
- 22 INFICON (2002) Ion-Molecule Reactions, Technical Note.

- 23 Redhead, P.A., Hobson, J.P., and Kornelsen, E.V. (1993) The Physical Basis of Ultrahigh Vacuum, American Institute of Physics, Woodbury, NY, pp. 174-175.
- 24 Watanabe, F. (1990) Very-low outgassing rate, separable, aluminum-flanged, hotcathode ion source for a residual gas analyzer. J. Vac. Sci. Technol. A, 8, 3890.
- 25 Watanabe, F. and Kasai, A. (1995) Low outgassing residual gas analyzer with a beryllium-copper-alloy-flanged ion source. J. Vac. Sci. Technol. A, 13, 497.
- 26 Huber, W.K., Müller, N., and Rettinghaus, G. (1990) Total and partial pressure measurement in the low 10-12 mbar range. Vacuum, 41, 2103-2105.
- 27 Müller, N. (1993) Use of quadrupole mass spectrometers in ultrahigh vacuum systems. Vacuum, 44, 623-626.
- 28 Ellefson, R.E., Moddeman, W.E., and Dylla, H.F. (1981) Hydrogen isotope analysis by quadrupole mass spectrometry. J. Vac. Sci. Technol., 18, 1062.
- 29 Bleakney, W. and Hipple, J.A. (1938) A new mass spectrometer with improved focusing properties. Phys. Rev., 53, 521.
- 30 Robinson, C.E. and Hall, L.G. (1956) Small general purpose cycloidal-focusing mass spectrometer. Rev. Sci. Instrum., 27, 504.
- 31 Sommer, H., Thomas, H.A., and Hipple, J.A. (1951) The measurement of e/M by cyclotron resonance. Phys. Rev., 82, 697.
- 32 Alpert, D. and Buritz, R.S. (1954) Ultrahigh vacuum. II. Limiting factors on the attainment of very low pressures. J. Appl. Phys., 25, 202.
- 33 Brydon, W.A., Benson, R.C., Ecelberger, S.A., Phillips, T.E., Cotter, R.J., and Fenselau, C. (1995) The tiny-TOF mass spectrometer for chemical and biological sensing. Johns Hopkins APL Tech Dig., 16, 296.
- 34 Gibson, J.R. and Taylor, S. (2000) Prediction of quadrupolar mass filter performance for hyperbolic and circular cross-section electrodes. Rapid Commun. Mass Spectrom., 14, 1669-1673.
- 35 Holkeboer, D.H., Karandy, T.L., Currier, F.C., Frees, L.C., and Ellefson, R.E. (1998) Miniature quadrupole residual gas analyzer for process monitoring at milliTorr pressures. J. Vac. Sci. Technol. A, 16, 1157.

- 36 Dahl, D.A. (2004) SIMION 3D Version 7.0 Ion Optics Software.
- 37 Dawson, P.H. (1995) Quadrupole Mass Spectrometry and Its Applications, AVS Classics Series, American Institute of Physics, Woodbury, NY, pp. 13-36.
- 38 Austin, W.E., Leck, J.H., and Batey, J.H. (1992) Study of the performance of a group of quadrupole mass spectrometers. J. Vac. Sci. Technol. A, 10, 3563.
- 39 Lieszkovszky, L., Filippelli, A.R., and Tilford, C.R. (1990) Metrological characteristics of a group of quadrupole partial pressure analyzers. I. Vac. Sci. Technol. A, 8, 3838.
- 40 Cowen, M.C., Allison, W., and Batey, J.H. (1993) Electron space charge effects in ion sources for residual gas analysis. Meas. Sci. Technol., 4, 72.
- 41 Cowen, M.C., Allison, W., and Batey, J.H. (1994) Nonlinearities in sensitivity of quadrupole partial pressure analyzers operating at higher gas pressures. J. Vac. Sci. Technol. A, 12, 228.
- 42 Badman, E.R. and Cooks, R.G. (2000) Miniature mass analyzers. J. Mass Spectrom., 35, 659-671.
- 43 Holkeboer, D.H. (Dec. 22, 1998) Method of Manufacturing a Miniature Quadrupole Using Electrode-Discharge Machining, US Patent 5852270.
- 44 Ferran, R.J. and Boumsellek, S. (1996) High-pressure effects in miniature arrays of quadrupole analyzers for residual gas analysis from 10^{-9} to 10^{-2} Torr. J. Vac. Sci. Technol. A, 14, 1258.
- 45 Syms, R.R.A., Tate, T.J., Ahmad, M.M., and Taylor, S. (1998) Design of a microengineered electrostatic quadrupole lens. IEEE Trans. Electron Devices, 45, 2304.
- 46 Taylor, S., Tindall, R.F., and Syms, R.R.A. (2001) Silicon based quadrupole mass spectrometry using microelectromechanical systems. J. Vac. Sci. Technol. B, 19, 557.
- 47 Freidhoff, C.B., Young, R.M., Sriram, S., Braggins, T.T., O'Keefe, T.W., Adam, J.D., Nathanson, H.C., Syms, R.R.A., Tate, T.J., Ahmad, M.M., Taylor, S., and Tunstall, J. (1996) Chemical sensing using nonoptical microelectromechanical systems. J. Vac. Sci. Technol. A, 17, 2300.

- 48 Diaz, J.A., Giese, C.F., and Gentry, W.R. (2001) Sub-miniature ExB sector-field mass spectrometer. J. Am. Soc. Mass Spectrom., 12, 619-632.
- 49 Ermakov, A.V. and Hinch, B.J. (2010) An electrostatic autoresonant ion trap mass spectrometer. Rev. Sci. Instrum., 81, 013107.
- 50 Brucker, G.A., Van Antwerp, K.D., Rathbone, G.J., Heinbuch, S.C., Schott, M.N., Hinch, B.J., and Ermakov, A.V. (Nov. 19, 2013) Electrostatic Ion Trap, US Patent 8586918.
- 51 Brucker, G.A. and Rathbone, G.J. (2010) Autoresonant Trap Mass Spectrometry (ART MS) for remote sensing applications. Int. J. Mass Spectrom., 295, 133.
- 52 Brucker, G.A. and Rathbone, G.J. (2011) Autoresonant Trap Mass Spectrometry, 8th Harsh Environment Mass Spectrometry Workshop.
- 53 Hamamatsu, Inc., Model R5150 series Compact Ion Detector Brochure.
- 54 Burle Technologies, Inc. (2001) Channeltron⁽ Electron Multiplier Handbook for Mass Spectrometer Applications.
- 55 Detector Technology, Inc. Tech. Note: Theoretical Life Equation for Channel Electron Multipliers.
- 56 Parfitt, W.E., Karandy, T.L., Frees, L.C., and Ellefson, R.E. (July 18. 2000) Ion Collector Assembly, US Patent 6,091,068.
- 57 Laprade, B. and Cochran, R. (1997) Operation of Microchannel Plate Based Detectors at Elevated Pressure, American Society for Mass Spectrometry Conference.
- 58 Peter, G., Koller, A., and Vazques, S. (1991) A new electron beam evaporation source for Si molecular beam epitaxy controlled by a quadrupole mass spectrometer. J. Vac. Sci. Technol. A, 9, 3061.
- 59 Koprio, J.A., Peter, G., and Fischer, H. (1988) Quadrupole mass spectrometer monitored and controlled thin film deposition processes under ultrahigh vacuum conditions. Vacuum, 38, 784.
- 60 Wieers, E. (June 2002) Thesis: Limburgs Universitair Centrum, Dipenbeek, Belgium.

- 61 Kersten, H., Deutsch, H., Steffen, H., Kroessen, G.M.W., and Hippler, R. (2001) The energy balance at substrate surfaces during plasma processing. Vacuum, 63, 385-431.
- 62 Lanzinger, E., Jousten, K., and Kühne, M. (1998) Partial pressure measurement by means of infrared laser absorption spectroscopy. Vacuum, 51, 47-51.
- 63 Jousten, K., Lanzinger, E., and Kühne, M. (2002) Genaue Linienstärkebestimmung von CO-Übergängen im mittleren Infrarot zur Teilchendichtemessung, VDI Bulletin No. 1667, Optische Analysenmesstechnik, pp. 99-104.
- 64 Jousten, K., Padilla, G., and Bock, Th. (2008) Investigation of tunable diode laser absorption spectroscopy for its application as primary standard for partial pressure measurements. J. Phys.: Conf. Ser., 100, 092005
- 65 McAndrew, J.F., Inman, R.S., and Jurcik, B. (1995) Gaseous contaminant measurement for semiconductor processing by diode laser spectroscopy. J. Inst. Environ. Sci., 38, 22-29.
- 66 Inman, R.S. and McAndrew, J.F. (1994) Application of tunable diode laser absorption spectroscopy to trace moisture measurements in gases. Anal. Chem., 66, 2471-2479.
- 67 McAndrew, J.F. and Inman, R.S. (1996) Using diode laser spectroscopy to evaluate techniques for acceleration of etch chamber evacuation. J. Vac. Sci. Technol. A, 14, 1266-1272.
- 68 Zalicki, P. and Zare, R.N. (1995) Cavity ring-down spectroscopy for quantitative absorption measurements. J. Chem. Phys., **102**, 2706-2717.
- 69 Hodges, J.T., Looney, J.P., and van Zee, R.D. (1996) Laser bandwidth effects in quantitative cavity ring-down spectroscopy. Appl. Opt., 35, 4112-4116.
- 70 Hodges, J.T., Looney, J.P., and van Zee, R.D. (1996) Response of a ring-down

- cavity to an arbitrary excitation. J. Chem. Phys., 105, 10278-10288.
- 71 Looney, J.P. et al. (1993) Measurement of CO pressures in the UHV regime using resonance-enhanced mulitphoton-ionization time-of-flight mass spectroscopy. J. Vac. Sci. Technol. A, 11, 3111-3120.
- 72 Neill, J.A., Passow, M.L., and Cotler, T.J. (1994) Infrared absorption spectroscopy for monitoring condensible gases in chemical vapor deposition applications. J. Vac. Sci. Technol. A, 12, 839-845.
- 73 Nier, A.O. and Stevens, C.M. (1947) Mass spectrometer for leak detection. J. Appl. Phys., 18, 30.
- 74 Nerken, A. (1991) History of helium leak detection. J. Vac. Sci. Technol. A, 9, 2036.
- 75 Reich, G. (1987) The principle of Helium enrichment in a counter flow leak detector with a turbo molecular pump with two inlets. J. Vac. Sci. Technol. A, 5, 2641.
- 76 Hablanian, M. (1991) Use of oil-free mechanical pumps with leak detectors. J. Vac. Sci. Technol. A, 9, 2039.
- 77 Liepert, A. and Lessard, P. (2001) Design and operation of scroll-type dry primary vacuum pumps. J. Vac. Sci. Technol. A, 19,
- 78 Jitschin, W. and Wandrey, D. (1988) Temperature dependence of the leak rate of helium diffusion leak artefacts. Vacuum,
- 79 Chamberlin, J.L. (1989) The modelling of standard gas leaks. J. Vac. Sci. Technol. A, 7, 2408.
- 80 Bley, W.Große. (1990) Temperature dependence and long-term stability of helium reference leaks. Vacuum, 41, 1863.
- 81 Große, G. and Messer, G. (1987) Summary abstract: calibration and long-term stability of helium reference leaks. J. Vac. Sci. Technol. A, 5, 2661.
- 82 Spiess, J. (1966) Die Lecksuchröhre, Diploma thesis, KFK Karlsruhe.
- 83 Audi, M. (1990) An ion pump based leak detector. Vacuum, 41, 1856.

15

Calibrations and Standards

Dr. Karl Jousten

Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

This chapter deals with the calibration of vacuum gauges, with primary and secondary standards for vacuum pressure, and with the question how to measure pumping speeds.

15.1 Introduction

Calibration determines the relation of the reading of a measuring instrument (e.g., a vacuum gauge) and the corresponding value of the measurand (e.g., physical vacuum pressure), as established by a standard under well-defined conditions. Calibrations are necessary when the output quantity Y of an instrument cannot be described by known input quantities X_i via a physical formula, or when the X_i are not well known or defy measurement. In most cases, the output quantity Y of an instrument is not measured directly but determined by a set of input quantities,

$$Y = F(X_i)$$
.

Some of the X_i may be physical constants. If the X_i are unknown or defy determination, a new relation is set up using a calibration constant.

As an example, let us examine an ionization gauge. The reading of such a gauge shall be given by Eq. (13.31):

$$p = \frac{I^+}{I_0} \cdot \frac{kT}{\sigma \Delta I},$$

where I^+ is the measured ion current, I_e the current of electrons entering the ionization region, T the temperature of the gas, σ the ionization cross section depending on the energy of the bombarding electrons and the gas species, Δl the mean path length of electrons through the ionization space, and k Boltzmann's constant. Only k, as a physical constant, and I^+ as the measured quantity

are well known, while the other quantities are not. For example, it is impossible to calculate Δl with sufficient accuracy by means of computer simulations. Even if this would be possible, experiments [1] show that the spatial emission distribution of the electrons on the cathode changes with time so that the information may be lost.

A calibration constant serves to define a new relation between an input quantity or measured quantity (in this case, ion current I^+) and the output quantity (here pressure). For the case of the ionization gauge, this was accomplished by employing Eq. (13.33) via the sensitivity constant, which is determined by calibration.

The purpose of a calibration is to assure correct readings on a gauge by making the indicated quantity directly or indirectly traceable to SI units. This is important not only for physical but also economical reasons. If for example a manufacturer of a high-vacuum pump tests the system using an ionization gauge indicating too high values above the suction flange, the manufacturer will quote too low values of pumping speed and may be at disadvantage compared to a competitor who measured correctly (see Figure 15.1).

For this reason, the International Organization for Standardization (ISO) in the 1990s and during the last decade adopted standards to ensure that calibration laboratories work according to quality management rules and make their measuring instruments traceable to the SI units. In order to overcome trade barriers, many of the states of the meter convention signed an arrangement by which they mutually recognize their calibration certificates of their national metrological institutes, provided these comply with the quality management rules of ISO.

Two principal methods are available for calibrating vacuum gauges (Table 15.1):

1) The reading of the gauge under calibration is compared to a known pressure of a primary standard. This is the more accurate of the two methods. A pri-

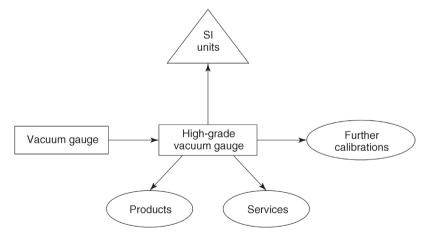


Figure 15.1 Purpose of the calibration of a vacuum gauge.

Pressure Primary standards		Secondary standards		
10 ⁵ Pa	Mercury column manometer	Quartz-Bourdon manometer		
	Rotary piston gauge	Resonance silicon gauge		
	Pressure balance	Capacitance diaphragm gauge		
	Static expansion system	Spinning rotor gauge		
10^{-10} Pa	Continuous expansion system	Ionization vacuum gauge		
		Extractor gauge		
		Mass spectrometer		

Table 15.1 Primary and secondary standards in use for pressures in the vacuum regime.

There is no strict one-to-one relation between the primary and secondary standards.

mary standard is defined [2] as a standard designated or widely acknowledged as having the highest metrological quality and whose value is accepted without reference to other standards of the same quantity. By using primary standards, pressures can be generated most accurately or with the lowest uncertainty, since they derive the unit of pressure directly from the SI units mass, length, and time according to Eq. (3.1).

A single type of primary standard is not capable of covering the entire technically relevant regime of vacuum pressures spanning about 15 decades (10⁻¹⁰-10⁵ Pa). The various methods available for the different ranges are explained in Section 15.2.1. Some national metrological institutes operate primary standards for vacuum (Figure 15.2), for example, the National

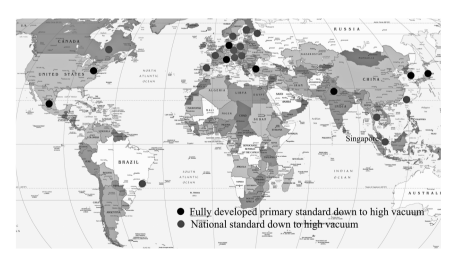


Figure 15.2 National or designated metrological institutes, which operate primary (black circles) or secondary (gray circles) standards for vacuum down into the high-vacuum

regime and which are listed with an approved entry in the international data base of the BIPM (www.bipm.org).

Institute of Standards and Technology (NIST) in the United States, the Physikalisch-Technische Bundesanstalt in Germany, but primary standards are also kept in large companies or calibration service laboratories.

2) The reading of the gauge under calibration is compared to another calibrated vacuum gauge. Such a device is called secondary standard when calibrated to a primary standard, or generally, reference standard. This method is referred to as comparison method and is described in Section 15.2.2.

An important requirement for using a vacuum gauge as reference standard is good long-term and transport stability to ensure that the information gained during calibration is not lost too early. In addition, the measurement uncertainty must be low compared to other vacuum gauges used in the considered pressure range. It follows that for each vacuum pressure range there is one particularly suitable type of vacuum gauge. These types are covered in Sections 15.2.3–15.2.5 in terms of their use as secondary standards.

Despite the fact that partial pressure gauges cannot be calibrated for their full measurement range, calibrations are possible and sensible under well-defined conditions. Section 15.3 covers this subject. Readings of leak detectors are calibrated by employing test leaks. Calibration of the latter is described in Section 15.4.

The values of pumping speed and other important parameters of vacuum pumps depend strongly on the measuring method. In order to make the specifications of different manufacturers comparable, written standards are available that are addressed in the final section of this chapter.

15.2 Calibration of Vacuum Gauges

15.2.1

Primary Standards

Pressures around atmospheric pressure down to about 10 Pa can be measured directly as force per unit area (Eq. (3.1)), for example, in a mercury manometer.

Below 10 Pa, the idea of all vacuum primary standards used today is the following: the gas is measured precisely at a pressure as high as possible, and then the gas is expanded to the desired pressure in a manner well treatable by calculation. When the gas is expanded from one enclosed volume into another, the method is referred to as static expansion. During expansion, all pumps are disconnected from the vessels. When the gas is expanded into a vacuum pump in a stationary flow through orifices of different conductances, the method is called continuous expansion method. In this method, the vacuum pumps are an inherent part of the method. The static expansion method is applied in the fine and high vacuum, and the continuous expansion method in the high and ultrahigh vacuum.

Scaling the pressure in the vacuum regime by means of primary standards is comparable to climbing down a ladder. Starting from atmospheric pressures, which can be measured with high accuracy, each further step represents an expansion to a lower value, and each new step again provides the basis for the next pressure reduction step.

By definition, primary standards are independent tools and there is no direct way to decide whether their value is correct or not. Each value of the produced measurand, however, is attributed an uncertainty assumed valid across the entire range covered by the physical value of the measurand. However, whether this is actually the case or whether an error occurred can only be revealed by a comparing between primary standards. For this reason, the International Committee of Weights and Measures (CIPM) carries out key comparisons at regular intervals by which the degree of equivalence is determined. Such comparisons are also conducted for vacuum pressures (Table 15.2).

15.2.1.1 Liquid Manometers

Liquid manometers are both the oldest and the most accurate vacuum gauges: in 1644, Torricelli was first to prove the existence of vacuum with his mercury tube (Figure 1.2). Until 1873, the *Torricelli* tube (Figure 1.9) was the only vacuum gauge available. This is why vacuum pressures were given as mm mercury column.

The accuracy of liquid manometers is due to the employment of the very dense sealing liquid mercury and to the accuracies of today's methods of length measurement. Mercury provides best capabilities of accurate measurement because of the stability of its specific density and the accuracy by which the latter is known. With modern methods of laser interferometry or distance measurement with ultrasound, heights of mercury can be measured with a resolution of 10 nm, so that pressures of 100 kPa (100 Pa) can be measured with a relative uncertainty of 3×10^{-6} (3×10^{-5}).

The Torricelli tube is one of the so-called open liquid manometers (Figure 15.3). Here, one side of the U-shaped tube is open to the atmosphere. This has the disadvantage that the height of the column in the closed shank is subject to atmospheric pressure and its changes.

In the closed U-tube manometer, one side of the U is evacuated. Depending on the vapor pressure of the liquid used, a certain pressure will develop that has to be measured and taken as correction.

Figure 15.3 shows a schematic diagram of a liquid manometer. The different pressures in volumes 1 and 2 above the liquid have to be compensated by different heights of the liquid in the two shanks. Equilibrium exists when the pressure on any level N (Figure 15.3) below the two liquids is equal:

$$p_1 + \frac{m_1 g}{A_1} = p_2 + \frac{m_2 g}{A_2},\tag{15.1}$$

where m_1 and m_2 are the masses of the liquid columns above N, and $A_{1,2}$ the cross-sectional areas of the shank at level N. If ρ denotes the density of the liquid

Table 15.2 Published international comparisons of national standards between 1989 and 2014.

Year of publication	Pressure range (Pa)		
1989	10 ⁻⁴ to 1	CMU, ETL, IMGC, LNE, NIM, NIST, NPL, NPL/I, PTB	[3]
1989	10^{-3} to 1	NPL/I, PTB	[4]
1992	2×10^{-2} to 0.2	LIP, PTB	[5]
1997	3×10^{-7} to 9×10^{-4}	NIST, NPL, PTB	[6]
2002	1 to 10 ³	CSIRO-NML, IMGC, KRISS, NIST, NPL, NPL/I, PTB	[7]
2005	3×10^{-4} to 9×10^{-2}	CENAM, PTB	[8]
2005	3×10^{-4} to 9×10^{-2}	CEM, IMGC, IMT, LNE, NPL, PTB, UME	[9]
2005	1 to 10 ³	CEM, IMGC, LNE, MIKES, NMi, NPL, OMH, PTB, SP, UME	[10]
2007	3×10^{-1} to 7×10^{3}	CMI, PTB	[11]
2010	$3 \times 10^{-6} \text{Pa to}$ $9 \times 10^{-4} \text{Pa}$	KRISS, NIST, NPL, NPL/I, PTB	[12]
2011	3×10^{-6} Pa to 9×10^{-4} Pa	KRISS, NMIJ	[13]
2014 2014	1 Pa to 15 kPa 3×10^{-6} Pa to 9×10^{-4} Pa	CMI, INRIM, LNE, MIKES, PTB NIST, PTB	[14] [15]

Laboratories represent the following countries: CENAM - Mexico; CEM - Spain; CMU - former Czechoslovakia; CMI - Czech Republic; ETL - Japan; IMGC and INRIM - Italy; IMT - Slovenia; KRISS - Korea; LIP - China; LNE - France; MIKES - Finland; NIST - USA; NMi - Netherlands; NPL - United Kingdom; NPL/I - India; OMH - Hungary; PTB - Germany; SOGEV - France; SP -Sweden; UME - Turkey.

and g the local acceleration due to gravity, it follows:

$$p_1 + \frac{\rho A_1 h_1 g}{A_1} = p_2 + \frac{\rho A_2 h_2 g}{A_2}.$$
 (15.2)

Thus, the pressure difference $p_1 - p_2$ between both volumes,

$$p_1 - p_2 = \rho g(h_2 - h_1) = \rho g \Delta h. \tag{15.3}$$

Using this equation makes sense for measuring a pressure difference $p_1 - p_2$ either against a fixed and well-defined pressure p_2 (differential mode) or against atmospheric pressure $p_2 = p_{\text{atm}}$ (gauge mode).

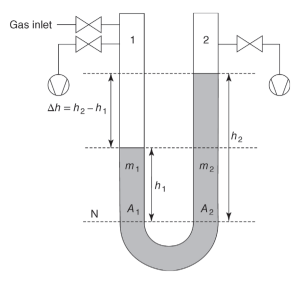


Figure 15.3 Diagram of a U-tube manometer: if volume 2 above the mercury surface is closed and evacuated as shown in the figure, the manometer is of the closed type, if it is open to atmosphere, it is referred to as open type.

For measuring absolute pressures with a U-tube manometer (absolute mode), we rewrite Eq. (15.3) as

$$p_1 = \rho g \Delta h + p_2. \tag{15.4}$$

Using mercury with a density $\rho = 13545.84 \, \mathrm{kg \ m^{-3}}$ (Table 15.3) as sealing liquid, we find the standard pressure $p_0 = p_1 - p_2 = 101\,325\,\mathrm{Pa}$ with $g = 100\,\mathrm{Pa}$ 9.81253 ms⁻² (Braunschweig, Germany) at $\Delta h = 762.307$ mm.

An important prerequisite for accurate pressure measurement is avoiding capillary depression: adhesion forces on the walls would influence the height of the mercury column. This effect is inversely proportional to the diameter of the liquid column. When using U-tube mercury manometers as primary standards, this diameter must exceed 15 mm.

Modern measurements of the height difference Δh can be performed so accurately that the uncertainty of the value of the mercury density due to the uncertainty of temperature measurement is the limiting factor. The mercury

Table 15.3 Density of mercury $\rho(t,p_0)$ at standard pressure $p_0=$ 101.325 kPa [16,17] versus temperature t.

t (°C)	17	20	23	26
$\rho(t,p_0)~(\mathrm{kg}\mathrm{m}^{-3})$	13 553.207	13 545.840	13 538.480	13 531.125

The standard uncertainty at 20 °C is 0.0068 kg m⁻³.

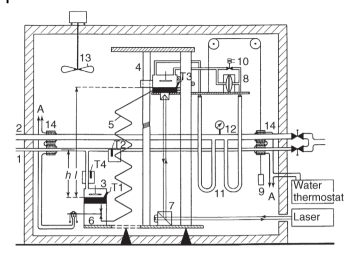


Figure 15.4 Diagram of the high-accuracy mercury U-tube of the *Physikalisch-Technische Bundesanstalt* in Braunschweig, Germany. 1 Gas inlet. 2: vacuum tube. 3: fixed cistern. 4: movable cistern with spindle and support column. 5: flexible tube containing the mercury and connecting the two columns. T1 to T4:

temperature test points. 6: mercury valve. 7: beam splitter for interferometer. 8: capacitance diaphragm gauge. 9: counter weight. 10: bypass valve. 11: flexible metal tube. 12: Penning vacuum gauge. 13: Fan. 14: thermostatically controlled body for temperature stabilization.

surface itself cannot serve as reflecting surface, since vibrations on the surface would limit the resolution [18]. Figure 15.4 shows a setup for height measurement. The height of the liquid is fixed by a capacitor, one electrode being a metal plate above the mercury, the other being the mercury surface itself. Subsequently, the height difference Δh to the movable capacitor electrode is determined by optical interferometry [19,20].

Floats on the mercury surface can be used as an alternative to capacitive coupling. They reflect the laser beam coming from the top the tube. The disadvantage of using floats is their slight friction with the tube walls, which limits the resolution.

A third possibility of measuring the height difference is by using ultrasonic pulses (Figure 15.5). The sound wave impulse is coupled in from the bottom of the shank via a piezoelectric crystal and a phase-sensitive measurement of the impulse's transit time is performed [21]. Instead of two, three aligned shanks can be used in a mercury U-tube. The middle shank is evacuated and the two outer shanks are loaded with equal test pressures. Any small common tilt of the shanks against the vertical direction is eliminated if the average of the height differences between the outer shanks and the middle shank is considered.

15.2.1.2 Compression Manometer after McLeod

For extending the measuring range of liquid manometers to lower pressures, *McLeod* invented the so-called compression gauge in 1873. Here, the gas is

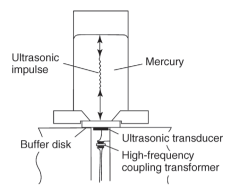


Figure 15.5 Injection of an ultrasound impulse disk avoids propagation of the ultrasound into the mercury column of a U-tube manometer. The pulse is used to measure the height via its running time in the column. The buffer

pulse into the supporting structure. Data taken from Ref. [22].

compressed before measuring its pressure. For known compression ratios, the pressure prior to compressing can be calculated from the value after the compression.

Until the 1960s, the McLeod gauge was the only primary standard for the fine and high-vacuum regimes. Its measuring range extended down to 10⁻⁴ Pa. In fact, until the 1930s, it was the only vacuum gauge available for this regime. Today though, it is hardly used. However, because of its importance in the past and the interesting measuring principle, we will briefly explain it in this section.

Figure 15.6 shows a diagram of a McLeod vacuum meter. The connection to the recipient whose pressure p is to be measured is arranged at the top. Mercury can be injected from a reservoir at the bottom, for example, using ambient atmospheric pressure. Initially, the mercury is significantly below the A-A level. It has to be assured that the pressure p to be measured is present throughout the entire volume of the McLeod gauge, especially in the compression volume V and in the capillaries K1 and K2. The mercury is admitted, and as soon as it reaches level A-A, the compression volume Vcomprising S_1 , V_1 , and K_1 is disconnected from the recipient. As the mercury level rises, the pressure of the test gas in the compression volume increases. The mercury is allowed to rise further until it reaches the zero line 0-0 in reference capillary K2. The measuring capillary K1 and the reference capillary K2 must have the same cross-sectional area in order to feature the same capillary depression (Section 15.2.1.1). Because of the higher pressure due to compression, the mercury level in K_1 is lower, by the height h. The higher the initial pressure p, the lower the level and the higher the value of h measured from 0-0. The amount of gas enclosed in capillary K1 fills the volume $V' = A_K h$ at pressure $p' = \rho g h + p$. The Boyle-Mariotte law (3.16) states that, just before compression (when the mercury level is at A-A), pV

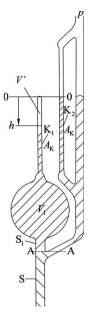


Figure 15.6 Compression vacuum meter after McLeod. V Compression volume comprising the volumes of the sphere V_1 , capillary K_1 , and the neck S_1 . Capillaries K_1 and K_2 have the same cross-sectional area A_K . S Riser tube. A-A level of disconnection. 0-0 zero level.

equals p'V':

$$pV = p'V' = (\rho gh + p)hA_{K}. \tag{15.5}$$

This yields the pressure p to be measured:

$$p = \frac{\rho g A_{\rm K}}{V - h A_{\rm K}} h^2. \tag{15.6}$$

If $hA_K \ll V, p$ follows a square law in h:

$$p = \frac{\rho g A_{\rm K}}{V} h^2. \tag{15.7}$$

Linear scales are also possible with the McLeod. For this, the mercury is raised to a fixed level in capillary K_1 thus yielding a fixed compression ratio. The height difference of the mercury levels in K_1 and K_2 is then proportional to p. The disadvantage of this method is that it covers only two pressure decades whereas the quadratic scale is capable of covering four decades.

Equation (15.7) is not applicable if the gas or part of it (e.g., water vapor) condenses under compression. Many effects, particularly the *Gaede–Ishii* effect [23–25], hamper operation of a McLeod: to avoid recipient pollution with mercury vapor, an LN^2 cold trap is placed between recipient and McLeod. This,

however, causes a steady flow of mercury into the cold trap, with the flow acting like a diffusion pump. Depending on the species, this falsifies pressures by up to 50%.

As a measuring device, the McLeod almost disappeared since it measures discontinuously and requires a multistep procedure to obtain the pressure value. The static expansion method replaced the McLeod as a primary standard due to the hazards involved with its operation (glass apparatus, large amount of mercury).

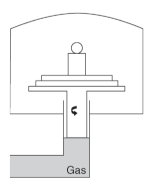
15.2.1.3 Piston Gauges and Pressure Balances

Since the 1990s, pressure measurements with piston gauges and so-called pressure balances provide uncertainties of only a few 10⁻⁶. Figure 15.7 shows the principle of a gas-operated piston gauge. A cylindrical piston rotates in a closely fitted circular cylinder. The pressure at the base of the piston is defined as the ratio of the total downward force on the piston to the effective area of the piston when floating at its operating level.

As with many vacuum primary standards, a piston gauge is a pressure generator, not a vacuum meter. In addition, to be more specific, it generates a pressure difference between the piston's bottom and the volume above the piston-cylinder assembly. If this volume is enclosed by a bell jar and evacuated, the piston gauge can be used as an absolute pressure generator. Typical pressure ranges of piston gauges used for vacuum gauge calibrations range from 2 kPa to 300 kPa.

The gap between piston and cylinder typically spans a few tenths of a micrometer for piston cross sections of 10 cm². Thus, obviously, manufacturing such a piston-cylinder assembly requires sophisticated precision techniques and only few materials such as tungsten carbide or ceramics can cope with the requirements.

The downward force shall be determined solely by the weight of the piston and the weight of additional masses placed on top of it. The rotary movement minimizes friction effects. Magnetic and electrostatic forces between piston and cylinder must be avoided as well. The effective area can be determined by



piston gauge. The force exerted by the gas pressure onto the piston's cross-sectional area is equal to the weight force of the piston and

Figure 15.7 Operational principle of a rotating the additional masses. The gap between piston and cylinder is strongly exaggerated in the drawing and amounts only to a few µm in practice.

dimensional measurements, or for known weight, from pressure comparisons with liquid manometers. Comparisons with liquid manometers are more convenient and more precise in determining effective piston areas because disturbing effects such as misalignments of the piston axis against the cylinder axis, eccentricity of parts, frictions effects, and so on, are calibrated into the effective area. In addition, possible influences of gas species, operating height, and pressure [26,27] on the effective area are easily determined by comparisons with liquid manometers. Since the mid-1990s, however, it is possible to consider these aspects in calculations and to manufacture dimensions of piston-cylinder systems to an accuracy providing piston gauges with a metrological quality equal to that of mercury manometers.

Neglecting the buoyancy of the piston in vacuum, the pressure generated by the piston gauge is given by

$$p = \frac{\sum_{m_i g}}{A_{\text{eff}} \left[1 + (\alpha_{\text{cyl}} + \alpha_{\text{pist}})(T - T_0) \right]} + p_{\text{res}},$$
(15.8)

where m_i are the pieces of mass contributing to the weight force, g is the local acceleration due to gravity, Aeff is the effective cross-sectional area of the piston, $\alpha_{
m cvl}$ and $\alpha_{
m bist}$ are the thermal expansion coefficients of cylinder and piston, respectively, T is the prevailing temperature, and T_0 is the temperature at which A_0 was determined. $p_{\rm res}$ denotes the residual pressure in the bell jar above the piston.

Example 15.1

What is the effective cross-sectional area of an ideal piston-cylinder system in a first approximation calculation (piston radius r_p , cylinder radius r_c)?

The upward force acting on the piston has two components: the force caused by the pressure difference $\pi r_p^2(p-p_{res})$ and the frictional force F exerted to the cylindrical surface of the piston by the gas molecules flowing upward under the influence of the pressure difference. Both components together equalize the weight force G:

$$G = \pi r_{\rm p}^2 (p - p_{\rm res}) + F.$$

For viscous flow, the maximum of the bulk velocity develops approximately in the center between the piston radius and the cylinder radius (not exactly in the middle due to the cylinder geometry). At this point, friction between two neighboring layers is zero. This cylindrical surface is characterized by its radius $r_{\rm eff}$. For the cylindrical ring between r_p and r_{eff} , the inherent weight force of the gas and the friction F of the gas on the piston must be equal to the upward force of the gas due to the pressure difference:

$$G_{\rm gas} + F = \pi (r_{\rm eff}^2 - r_{\rm p}^2)(p - p_{\rm res}).$$

Adding both equations yields

$$G + G_{\text{gas}} = \pi r_{\text{eff}}^2 (p - p_{\text{res}}).$$

This indicates that the effective area of the piston can be described by a virtual piston with radius $r_{\rm eff}$, lying approximately in the center between $r_{\rm c}$ and $r_{\rm p}$. For gas-operated piston gauges, G_{gas} is negligible compared to G (error $\ll 10^{-6}$).

Typical uncertainties of p (confidence interval 95%) amount to (smaller uncertainties are possible)

$$\Delta p = 1 \,\text{Pa} + 3 \times 10^{-5} \,p. \tag{15.9}$$

Piston gauges can serve for calibrating vacuum gauges in the range of 2–100 kPa, or for generating precise initial pressures in a static expansion scheme.

In the late 1990s, so-called pressure balances became commercially available, resolving pressures down to 1 mPa, see Figure 15.8. Here also, a piston-cylinder

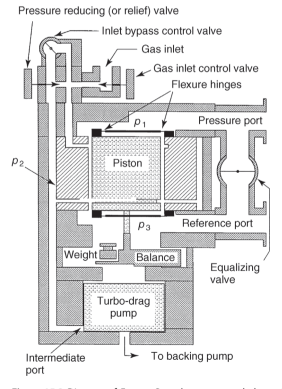


Figure 15.8 Diagram of Furness-Rosenberg pressure balance. The pressure at the top of the piston exerts a force onto the piston that is measured by means of a balance located in the vacuum space underneath the piston. Courtesy of Furness Controls.

assembly defines the effective area A_{eff} . In contrast to the principle of a rotating piston gauge, however, the force exerted by the gas (here from the top of the piston) is not measured by pieces of mass but by a balance (force meter). A rod transmits the force to the balance. Since this design prevents rotation of the piston, the clearance between piston and cylinder must be larger than for the rotating piston in order to avoid static frictional forces. The measurement range lies between 1 Pa and 7 kPa [28], later extended to 11 kPa.

In a different type of pressure balance (force balanced pressure gauge [FPG] by DHI) the force exerted by the pressure is also measured by a balance; the piston, however, centers itself by the forces of a "lubricating" gas flow. Ooiwa [29,30] developed the operating principle. The disadvantage of this type compared to the first is the relatively high gas flow needed limiting somewhat the applications of this type as an absolute pressure-measuring device.

15.2.1.4 Static Expansion Method

The static expansion method employs the model used in the Boyle-Mariotte law (3.16): the product of pressure and volume of a fixed amount of gas is constant at constant temperature. If gas contained in a small volume at relatively high pressure is expanded into a much larger evacuated volume, the pressure drops according to the volume ratio (Figure 15.9). The initial pressure prior to the expansion should not exceed values significantly violating ideal gas behavior. For the rare gases and nitrogen, this is the case at a pressure of 300 kPa.

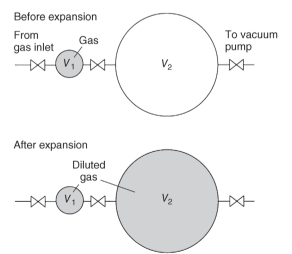


Figure 15.9 The static expansion method: by expanding a fixed amount of gas from a small volume into a large evacuated volume, the initial pressure drops according to the volume ratio.

Generally, it is impossible to provide precisely equal temperatures for both volumes including their connection pipes. This is why calculations are based on the ideal gas law (3.17) instead of the Boyle-Mariotte law (3.16).

If p_1 , V_1 , and T_1 denote pressure, volume, and temperature, respectively, prior to expansion, p_2 the pressure after and V_2 the volume including the connecting tubes into which the gas expands, it follows that

$$\frac{p_1 V_1}{T_1} = \frac{p_2 (V_1 + V_2)}{T_2},\tag{15.10}$$

if we approximate that the gas, after expansion, takes on the temperature T_2 of the much larger volume. The pressure after expansion therefore amounts to

$$p_2 = p_1 \frac{V_1}{V_1 + V_2} \cdot \frac{T_2}{T_1}. (15.11)$$

The so-called expansion ratio $V_1/(V_1+V_2)$ is the crucial parameter of an expansion system and has to be determined with the highest possible accuracy. Knudsen was first to apply the method of static expansion in 1910 [31].

Example 15.2

What is the equation of expansion for a real gas? The approximating equation for the real gas, Eq. (3.113), is

$$pV_{\text{mol}} = RT(1 + B''p),$$

where B" denotes the second virial coefficient. Since the pressure after expansion is much smaller than before, the term B''p is small, and thus, it is sufficient to correct solely for the gas prior to expansion. Therefore,

$$p_2 = p_1 \frac{V_1}{V_1 + V_2} \cdot \frac{T_2}{T_1} \cdot \frac{1}{1 + B'' p_1}.$$

The initial pressure, which ranges typically from 1 to 300 kPa, is generated with a piston gauge for highest accuracy or a suitable secondary standard such as a quartz Bourdon spiral manometer (Section 13.2.4.1). A fraction of the gas expanded to pressure p_2 can be used for a further subsequent expansion. Such repeated expansions provide pressures down to about 10⁻⁶ Pa. Figure 15.10 shows a five-stage expansion system.

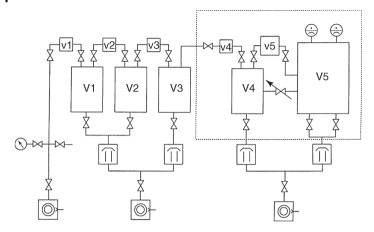


Figure 15.10 Example of a five-stage expansion system as developed by the *National Physical Laboratory (NPL)* in England. Pressures down to 10^{-6} Pa can be generated in volume V5.

It is also possible to repeat the same expansion by closing the valve to the large volume so it can be evacuated after the expansion.

This approach is required for obtaining pressures below 10^{-2} Pa with the two-stage expansion system of the *Physikalisch-Technische Bundesanstalt (PTB*) shown in Figure 15.11.

It is advisable to design the expansion ratios of a static expansion system such that there is an overlap of generated pressures via different expansion paths: a certain pressure range should be producible both with a given expansion ratio and low initial pressure, as well as with a high initial pressure using the next lower expansion ratio.

This overlap provides a means to check whether pressures generated by different expansion paths are consistent. There should also be an overlap of the pressure range after the first expansion with the pressures measured directly with the initial-pressure gauge.

Calibration equipment attached to the vessel filled with calibration pressure changes the volume and therefore the corresponding expansion ratio. This additional volume has to be estimated with sufficient accuracy. In the case of the PTB expansion system shown in Figure 15.11, a volume V_7 was introduced for this estimation. The valve to V_7 is shut in an evacuated state, and subsequently, an arbitrary but simple to measure pressure p_x is adjusted and measured under calibration. The valve to V_6 is closed. Opening the valve to V_7 initiates an expansion. Denoting the corresponding pressure after expansion with p_x , we obtain the unknown value of volume V_x by using the Boyle-Mariotte law:

$$V_x = \frac{p_y}{p_x - p_y} V_7. {(15.12)}$$

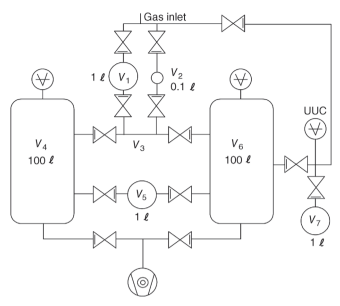


Figure 15.11 Example of a two-stage expansion system used at the Physikalisch-Technische Bundesanstalt in Germany. (UUC: unit under calibration.) The pressure routinely generated in V_6 lies between 10^{-2} and 10^{3} Pa.

The smallest pressures obtainable with sufficient accuracy with the static expansion method depend on the following required conditions.

- The pressures shall be approximately two orders of magnitude above the level of residual pressure in the system.
- The outgassing rate from inner surfaces of the calibration vessels must not lead to any significant pressure increase during measurement time.
- Significant loss of calibration gas due to adsorption on the chamber walls must be ruled out.

Due to the last-mentioned aspect, not all gases are suitable as calibration gas in the static expansion method. In practice, rare gases and nitrogen do not exhibit significant adsorption down to pressures of 10⁻⁶ Pa whereas, for example, hydrogen can only be used down to about 10^{-2} Pa. Oxygen or oxygen-containing gas species are hardly applicable in the static expansion method.

The most challenging but essential aspect when employing the static expansion method is an accurate determination of the expansion ratios. These may range from 1/100 to 1/100 000. The methods published to date can be categorized into three techniques:

1) Gravimetric technique. In this method, the individual volumes are determined by weighing the liquid they contain, which is of known density. For volumes $> 0.1 \,\ell$, this is the most accurate method. Highly distilled water or other liquids such as alcohol or mercury [32] may be used. To remove any

- air from the water as well as air bubbles sticking to the walls, the volume is evacuated. Fiber optics provide a means to check any inaccessible spots for remaining bubbles. When water temperatures including gradual temperature gradients in larger vessels are measured accurately and buoyancy corrections are applied, volumes can be determined with relative uncertainties in the low 10⁻⁴ range. The disadvantage of this method is that complex connection tubes with valves can only be measured with great difficulties and poor accuracy.
- 2) Constant pressure technique. Here, a variable volume of known size is used to determine the unknown volume. After expansion into the small, evacuated volume to be measured, the variable volume is adjusted such that the gauge indicates the same reading as prior to the expansion. This procedure is well suited for volumes $< 0.1 \,\ell$ [33]. A piston driven by a micrometer screw may serve as variable volume. The screw is driven into a volume connected with the volume to be measured via a valve. A stable vacuum gauge monitors the pressure inside the volume, and subsequently, the gas is expanded into the unknown volume. To compensate for the effect of the valve volume, the valve to the small volume then has to be sealed carefully (slowly). The change in the known variable volume required to reproduce the previous pressure is just equal to the volume to be measured.
- 3) Expansion technique. In this technique, the pressures before and after the expansion are used for determining the expansion ratio. This procedure is applicable only if the pressure after expansion can be measured with about the same accuracy as the pressure before expansion. For this reason, expansion is repeated as often as necessary. If the expansion ratio is about 1/100 and the initial pressure is 100 kPa, a pressure that can be measured with a relative uncertainty of a few 10⁻⁴ is reached after about 20 expansions. To determine the expansion ratio corrections to the ideal gas law, temperature gradients between the vessels and the timely drift of the temperatures need to be considered [34,35]. An expansion ratio of 1/100 can be determined with an uncertainty of less than 1×10^{-3} . This method was introduced by Elliott and Clapham [36]. Instead of using two calibrated gauges, a single uncalibrated gauge with a strictly linear pressure response can be utilized as well. The procedure was first reported by Berman and Fremerey [37]. They used a spinning rotor gauge for measuring the pressure before and after expansion.

Figure 15.12 shows the basic experimental setup. A first expansion provides a sufficiently low initial pressure. After closing valve V2, the large volume is evacuated through the open valve V3, and after disconnecting the vacuum pump, the remaining gas amount in the small volume is expanded into the large volume. Above 10⁻² Pa, the deceleration rate of the spinning rotor gauge is not strictly linear proportional to pressure, and thus, requires linearization. This method is capable of determining expansion ratios down to 1/250 with a relative uncertainty of 1×10^{-3} [35].

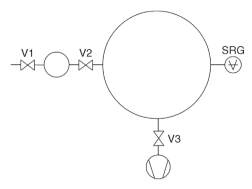


Figure 15.12 Measuring a volume or expansion ratio by means of a spinning rotor gauge.

The main advantage of the expansion method is that it is carried out in situ. However, the volumes are not determined to their absolute values, which is inconvenient when adding of volumes is desired. Therefore, the method is often combined with the gravimetrical method.

The uncertainties of the generated pressures in static expansion systems are mainly determined by the uncertainties of the expansion ratios and by the uncertainties in temperature measurements. Figure 15.13 shows the relative uncertainties exhibited in pressure generation using the PTB system in Figure 15.11.

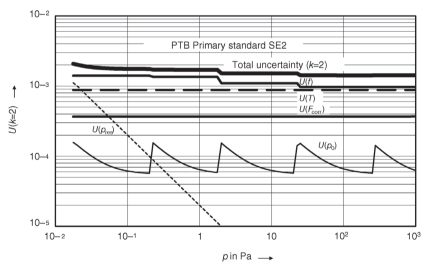


Figure 15.13 Uncertainty of pressure generation in the static expansion system of Figure 15.11. The indicated uncertainty is twice the relative standard uncertainty, typically corresponding to a confidence interval of 95%. The individual components of uncertainty are U(f) uncertainty of realized pressure

due to expansion ratio. U(T) due to temperature, $U(F_{corr})$ due to correction factors as virial coefficient, valve wear, adsorption, and so on, $U(p_0)$ due to initial pressure, $U(p_{res})$ due to residual pressure and the outgassing (unbaked system).

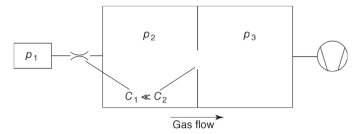


Figure 15.14 Basic principle of the continuous expansion method. Gas originating from a reservoir is pumped continuously through two flow elements of largely different conductances into a vacuum pump. *p* pressures in the volumes, *C* conductances.

15.2.1.5 Continuous Expansion Method

While the static expansion method uses two very differently sized volumes, the continuous expansion method employs two largely different conductances for pressure reduction (Figure 15.14).

The calibration gas is expanded continuously from a volume at high pressure into the vacuum pump via two conductances. If there are no sinks or sources of gas between the two conductances, the equation of continuity is valid and the net flow through the two orifices must be equal (isothermal conditions, for meaning of variables see Figure 15.14):

$$(p_1 - p_2)C_1 = (p_2 - p_3)C_2. (15.13)$$

With the approximation that p_2 and p_3 are negligible compared to p_1 and p_2 , respectively, we obtain

$$p_2 = p_1 \frac{C_1}{C_2}. (15.14)$$

According to the ratio of the two conductances, the initial pressure p_1 is reduced to p_2 , representing just the desired calibration pressure.

For a primary standard according to the continuous expansion method, conductance C_1 is very small $(10^{-6}-10^{-5}\,\ell\,\mathrm{s}^{-1})$ compared to C_2 $(10-100\,\ell\,\mathrm{s}^{-1})$, resulting in considerable pressure reduction.

Even though the principal procedure seems similarly simple as the static expansion method, its realization requires considerably more effort. For example, the approximation that p_3 is negligible compared to p_2 is usually unacceptable, considering the desired accuracies. Another difficulty is that the very small leak rates produced by C_1 depend on pressure, that is, the type of flow through the conductance, and that they lack stability in time. For highest accuracies, it is thus necessary to measure C_1 a new prior to each pressure generation. This is done in a so-called gas flowmeter. Such a flowmeter comprises a gas inlet system, a vacuum system in which pressure p_1 is adjusted and measured, as well as measuring equipment for determining conductance C_1 .

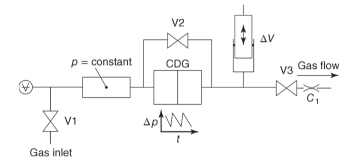


Figure 15.15 Basic principle of a gas flowmeter: from the working volume enclosed by V2, CDG, and V3, the gas flows into the calibration system through C_1 . The pressure in the working volume can be changed by adjusting

the variable volume ΔV (see the text). The pressure on the reference side of the CDG (capacitance diaphragm gauge) remains constant

Figure 15.15 shows the diagram of a flowmeter of constant pressure as listed in the classification provided by Peggs [38]. The pressure is measured by secondary standards calibrated using a piston gauge, pressure balance, or static expansion system. The measuring equipment for determining the conductance consists of a working volume (right-hand side) and a reference volume (left-hand side, p_0 = constant), separated by the differential pressure gauge CDG and bypass valve V2. The working volume containing displacement bellows as variable volume is additionally connected to valve V3 and leak C_1 through which the gas is transported to the calibration system.

Conductance measurements make use of a procedure described by *Bennewitz* and *Dohmann* [39]: initially, both volumes are pressurized equally with valves V1 to V3 open. After closing valves V1 and V2, pressure p_1 is measured using a secondary standard suitable for this pressure range. After closing V2, the pressure in the reference volume remains constant. Gas leaks out of the working volume through the open valve V3 producing a drop in pressure measured with the CDG. The obtained signal is used to drive the bellows such that pressure inside the working volume remains constant. For this, two methods are available. The variable volume provided by a piston or bellows is reduced continuously at constant speed so that the pressure remains constant. Alternatively, the volume can be adjusted by a fixed amount ΔV at intervals so that the pressure will vary slightly $(\pm 5 \times 10^{-4})$, following a saw-tooth characteristic. The interval Δt between equal values of pressure is measured. In both cases, the value of conductance C_1 is given by the volume speed of the variable element,

$$C_1 = \frac{\Delta V}{\Delta t}.\tag{15.15}$$

Since variations in the temperature differences between working and reference volumes would also change the signal on the CDG, the temperature in the

flowmeter should be constant. A temperature drift of only 1 mK min⁻¹ already affects measuring results [40]. The pressure differences caused by such drifts can be measured with valves V2 and V3 shut.

Example 15.3

Calculate the effect on the conductance measurement of $C_1 = 10^{-6} \, \ell \text{s}^{-1}$ for a temperature change in the reference volume of +1 mK min⁻¹ at 23 °C with respect to the temperature of the working volume ($V_{\text{workingvolume}} = 0.1 \ell$).

The relative change of pressure in the reference volume

$$\frac{\frac{\Delta p}{\Delta t}}{p} = \frac{\frac{\Delta T}{\Delta t}}{T}$$

produces a corresponding signal on the CDG. A pressure increase on the reference side is indistinguishable from a gas loss $\Delta V/\Delta t$ (drop in pressure) in the working volume V,

$$\frac{\frac{\Delta p}{\Delta t}}{p} = \frac{\frac{\Delta V}{\Delta t}}{V}, \quad \frac{\Delta V}{\Delta t} = V \frac{\frac{\Delta p}{\Delta t}}{p} = V \frac{\frac{\Delta T}{\Delta t}}{T} = 0.1 \ \ell \frac{\frac{1 \text{mK}}{60 \text{ s}}}{296.15 \text{ K}} = 7 \times 10^{-9} \ \ell \text{ s}^{-1}.$$

Comparison with $10^{-6} \ell s^{-1}$ yields an error of 0.7%.

The throughput q_{pV} produced in the flowmeter at a given temperature is given by

$$q_{nV} = p_1 C_1. (15.16)$$

This gas flow is fed to the calibrating system.

Figure 15.16 shows the diagram of a fully automated flowmeter as used by the PTB. In this setup, dosing valve V4, modified for constant conductance independent of the adjusting screw, provides the small conductance C_1 .

An important factor for the accuracy of the determined calibration pressure p_2 is the accuracy by which the larger conductance C_2 can be determined. As shown in Chapter 4, highest accuracy is achievable only in the molecular regime where the conductance of an ideally thin orifice is simply determined by its cross-sectional area A. Therefore, the continuous expansion method is mostly limited to pressures where flow is molecular.

In the calibration system, the calculated pressure and the pressure at the equipment to be calibrated must be equal. This is nontrivial since the system is an open system where gas is injected and pumped off.

In the calibration system, four components can be differentiated: the inlet system from the flowmeter, the calibration chamber, the pump orifice with conductance C_2 , and the pumping system.

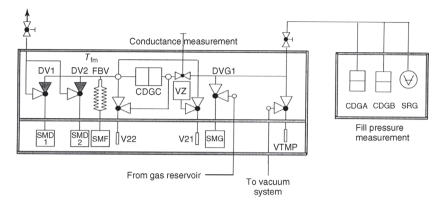


Figure 15.16 A flowmeter built by the *Physikalisch-Technische Bundesanstalt* according to the basic diagram in Figure 15.15. During operation, the system runs fully automated. Valves V21, V22, and VTMP operate pneumatically, the modified dosing valves DV1 and 2 are driven by stepping motors SMD1 and 2.

Dosing valve DVG1 for controlled gas admission is operated by stepping motor SMG, the variable volume is produced by the formed bellows FBV whose length is adjusted with a micrometer screw driven by stepping motor SMF. CDG: capacitance diaphragm gauge, SRG: spinning rotor gauge.

Inlet system. The inlet system has to be designed such that any beam effect of the molecular flow through the tubing from the flowmeter does not spread into the calibration chamber. This can be provided by building a separate prechamber with a small orifice to the calibration chamber. In the prechamber, the directions of the molecules are again uniformly distributed so that the flow through the orifice follows a Maxwellian distribution. Another solution is to install a baffle plate in front the inlet tube. This baffle plate also inhibits the direct path of molecules from the gas inlet to the outlet (pump orifice), a correction that has to be taken into account in the first case. It is also possible to form the inlet tubing such that escaping particles hit a portion of the wall of the calibration chamber distant from the pump orifice.

Calibration chamber. Both the gas inlet and the pump orifice disturb the *Maxwellian* distribution inside the calibration chamber. To minimize these disturbances, the following precautions have to be taken:

- The ideal shape for the chamber is a sphere; however, a more practical approach uses a cylinder with equal length and diameter. These shapes minimize the ratio of inner surface to volume, as well as pressure gradients within the vessel.
- The surface area of the largest inscribed sphere in the chamber should be at least a factor of 1000 larger than the orifice area. This is to ensure that pressure in homogeneities within the chamber is limited to approximately 10⁻³. However, Monte Carlo simulations of molecular scattering in the chamber that can be performed in a reasonable amount of time with the computer power available yield an estimation of the particle density distribution in the chamber and allow determining corrections for the pressures at the flanges

used for the gauges under calibration [41]. Furthermore, it is also possible to measure density distributions [42].

- The volume of the chamber should be considerably larger (by a factor of 50) than the added volumes of the attached gauges to be calibrated.
- The flanges carrying the gauges to be calibrated (test flanges) have to be oriented such that there is no direct interaction between attached gauges (no line of sight) and that their axes face neither the gas inlet nor the pump orifice.

Pump orifice. The conductance of the pump orifice can be described by the following equation:

$$C_2 = A \frac{\overline{c}}{4} K_1 K_2 \tag{15.17}$$

where \bar{c} is the mean thermal speed of molecules, K_1 a correction factor considering the reflection of molecules at the orifice edge of thickness l_1 and diameter d_2 see Eq. (4.151),

$$K_1 = 1 - \frac{l}{d}. ag{15.18}$$

Correction factor K₂ considers intermolecular collisions and may be approximated by

$$K_2 = 1 + a\frac{d}{\bar{l}},\tag{15.19}$$

where a is a constant and \overline{l} is the mean free path. According to older publications [43–45], a ranges between 0.01 and 0.125, a reported experimental value is a = 0.08 (Looney, J.P. NIST, private communication.). The smaller the two correction factors can be made, the smaller are the resulting uncertainties of the conductance. Therefore, the orifice is manufactured as thin as possible, for example, with a tapered edge, and pressures are kept low enough for the mean free paths to be large compared to the orifice diameter.

Pumping system. Pressure p_3 is negligible compared to calibration pressure p_2 only if the conductance C_2 of the pump orifice is small compared to the effective pumping speed of the high-vacuum pump. This is true, for example, for cryocondensation pumps (see Section 12.4.3.2) featuring cold surfaces at a temperature of 2.7 K behind the pump orifice [42]. When using such pumps, a separate chamber behind the pump orifice is dispensable. The pumping speed of turbomolecular or diffusion pumps, however, is not high enough to meet the assumption mentioned above. In this case, conductance C_2 of the orifice must be replaced by

$$C_2' = C_s \left(1 - \frac{p_3}{p_2} \right). \tag{15.20}$$

The ratio p_3/p_2 , often referred to as back streaming factor, depends on pressure and gas species and has typical values of 10⁻³ for heavier gases and 0.2 for lighter gases. This ratio can be measured if a chamber of about the same size as the calibration chamber is arranged behind the pump orifice so that the pressure distribution in this pump chamber is sufficiently homogeneous for measuring p_2 .

To correct for non-isothermal conditions, it is possible to refer both the temperatures of the flowmeter and of the calibration chamber to a common temperature of 23 °C. This leads to the difficulty that different correction terms are necessary for gauges, depending on whether they respond to gas density (e.g., ion gauges), impingement rates (spinning rotor gauges), or directly to pressure. This problem can be avoided by calculating the pressure actually present at the temperature inside the chamber. Ref. [46] covers the corresponding temperature correction in more detail.

A major benefit of the continuous expansion system method is the presence of a stationary equilibrium: the number of particles leaving the calibration chamber just balances the number entering the chamber. Therefore, ad- and desorption effects are relevant only until this equilibrium is obtained, and a wider selection of gas species are applicable with this method than with static expansion. Even water vapor has been used successfully [47].

As mentioned above, the upper limit of calibration pressure in continuous expansion is determined by the transition from molecular to viscous flow. In practice, this means that the upper pressure limit amounts to a few 10^{-2} Pa. With some effort, this limit can be shifted to higher values, as first demonstrated at NIST, by using small capillaries of a few microns in diameter for realizing C_2 in a channel plate structure.

The lower limit of calibration pressure in continuous expansion systems is determined by the residual pressure in the calibration chamber and by the lowest flow rate generable with sufficient accuracy in the flowmeter. This is typically 10^{-5} Pa ℓ s⁻¹, so that according to Eqs. (15.15) and (15.16), about 10^{-7} Pa can be generated in the calibration chamber.

For lower pressures, the flow divider principle may be applied [48-50], as shown in Figure 15.17.

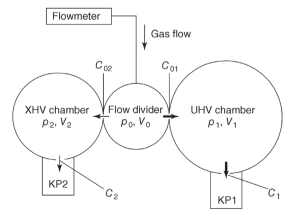


Figure 15.17 Calibration system CE3 at the Physikalisch-Technische Bundesanstalt for the continuous expansion method with flow divider. KP: Cryo-condensation pump. Pressures between 10^{-10} and 3×10^{-2} Pa can be generated in the two chambers.

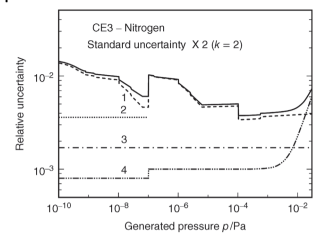


Figure 15.18 Uncertainties of the pressures generated by the calibration system CE3 (Figure 15.17). The individual uncertainties are due to inaccurate knowledge of 1: the flow

rate, 2: the conductance ratio C_{01}/C_{02} in the flow divider, 3: temperature, 4: conductances of the pump orifices C_1 and C_2 .

The flow from the flowmeter is fed into divider chamber V_0 containing two orifices of largely different conductances (nominal values $C_{01}=5\ell {\rm s}^{-1}$, $C_{02}=0.05\ell {\rm s}^{-1}$). When both chambers are pumped with the cryo-condensation pumps KP1 and KP2, about 99% of the gas flows into calibration chamber V_1 , but only 1% into chamber V_2 . This means that the pressure in V_2 is lower than in V_1 by about a factor of 100. For equal total flow from the flowmeter, this method is capable of generating calibration pressures 100 times smaller. The only trade-off is that the uncertainty of measurement of the conductance ratio C_{02}/C_{01} has to be added.

The uncertainties of the generated pressures in such a system are shown in Figure 15.18. In most of the range, the uncertainty of the gas flow rate dominates; only above 10^{-2} Pa does the uncertainty of the conductance due to the transition to viscous flow have a significant effect.

15.2.1.6 Other Primary Standards

In addition to the described primary standards, there are others that operate with molecular beams or pressure versus time methods. Optical methods might also play some role in the future.

Grosse and Messer [51] developed a primary standard with a molecular beam. Its operation, however, is elaborate and has the inherent handicap that relatively small conductances to the pump have to be installed in the calibration chamber so that the residual pressure is limited by the outgassing rate of the ion gauge under calibration.

Pressure versus time methods were reviewed by *Kuz'min* [52]. In a defined volume, the pressure increase or pressure decrease produced using a small conductance from a vessel at constant pressure or to a vacuum pump, respectively, is precisely predictable when ad- and desorption effects are negligible. The

methods' pressure ranges, however, are rather small and calibration is timeconsuming.

Optical methods were investigated to be suitable as primary standards [53–58], but have not yet been utilized. Laser absorption spectroscopy [56] or cavityring-down spectroscopy [57] (see also Section 14.3) is capable of determining absolute molecular densities if the line strengths of the corresponding absorption transitions of the gas species are known. Recently, there have been promising research projects that used refractive index measurements to measure pressure for a pure gas in a fundamental manner in the range 100 Pa to 100 kPa [59]. This method may have the capability to replace mercury manometers in the future.

15.2.2

Calibration by Comparison

In the comparison method, the device under calibration is compared to a calibrated reference standard. It is the easiest and fastest calibration method and therefore the one most often used in spite of the larger uncertainties involved. However, these uncertainties are acceptable in most customers' applications.

The readings of the two vacuum meters can be compared only if both are exposed to the same pressure. Thus, the vacuum chambers and the position of the test flanges must be designed to fulfill this condition. ISO 3567 [60] provides the corresponding guidelines. When the pressure is established using a stationary gas flow, neither the reference standard nor the gauge under calibration and the corresponding flange openings must be hit by the incoming gas flow. Similarly, a line-of-sight connection to the pump outlet is inappropriate. All measuring devices must be positioned symmetrically to both gas inlet and pump outlet.

If the pressure is not established by means of a stationary flow, but statically, that is, by disconnecting any vacuum pump from the chamber, the geometrical requirements mentioned above can be disregarded, since the pressure throughout an enclosed system inconstant (however, any thermal effusion effects must be ruled out, see Section 4.6). This static method is usually applied for pressures higher than 100 Pa. The stationary flow method calls for sufficient pressure stability during the measuring interval for the reference standard and the calibrated unit.

To minimize any disturbances caused by adsorption, desorption, and pump effects, the tubing to the measurement devices should be as short as possible and have a diameter equal to or greater than the entrance flange of the measurement unit. For the same reason, the chamber's surface-to-volume ratio should be small, best provided by using a sphere. ISO 3567 also permits a ratio as in a straight cylinder with a length of twice its diameter. The residual pressure should not exceed 10% of the lowest pressure.

Preferably, calibrations shall be carried out at 23 °C. Figure 15.19 shows an example of a calibration system.

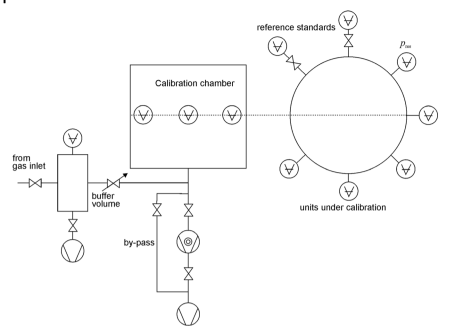


Figure 15.19 Diagram of a comparison calibration system that realizes the ISO 3567. The calibration chamber has to have cylindrical symmetry (including a sphere as possibility), gas in- and outlet must be located on the cylindrical axis, the flanges to adopt the

reference standards and units to calibrate must lie on the same equatorial plane. The buffer volume helps to stabilize the gas flow in, a bypass of the turbomolecular pump to evacuate the calibration chamber after venting.

The quality of the calibration result is determined mainly by the quality of the reference standard. The latter is characterized by its measurement accuracy, repeatability, long-term stability, as well as by the way it was calibrated. The following sections cover these aspects in more detail.

When a reference standard has been calibrated using a primary standard, it is called secondary standard. Most of the accredited calibration service laboratories use secondary standards. Both the German Calibration Service [61] and the American Vacuum Society [62,63] have issued guidelines for calibrating vacuum gauges.

15.2.3

Capacitance Diaphragm Gauges

Section 13.2.5.6 describes the operating principle of capacitance diaphragm gauges. Now, as well as in the following sections for other equipment, we will discuss the aspects of successful and accurate calibration of these vacuum gauges and procedures for optimizing their long-term stability.

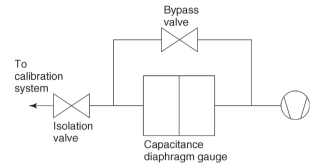


Figure 15.20 Vacuum setup for calibrating differential-type capacitance diaphragm gauges.

Section 13.2.5.6 introduces the two types of capacitance diaphragm gauges: absolute and differential capacitance diaphragm gauges. For the first type, the reference side of the deflecting diaphragm containing the measuring condenser (see Figures 13.13 and 13.15) is an enclosed and evacuated volume kept at a pressure below the resolution limit of the capacitance diaphragm gauge by a getter material.

In the differential type, the reference side is open. This allows measuring differential as well as absolute pressures. For the latter, the reference side must be evacuated below the resolution limit of the device via an outside pump.

Figure 15.20 shows the setup for calibrating differential type capacitance diaphragm gauges as reference standards. Transducers of the absolute type with full-scale deflections of 1 kPa or less should be protected by a valve in order to avoid exceeding the full scale, especially by atmospheric pressure. In some types of gauges, such overloads may invalidate previous calibrations, particularly for gauges containing metal diaphragms.

This is not necessary for a differential type gauge but it has to be taken into account that when the gauge is disconnected from the vacuum system both sides of the diaphragm are pressurized equally so that it does not suffer any overload. The measuring (test) side should be disconnected first.

The calibration setup in Figure 15.20 allows a correct zero (offset) measurement. The isolation valve to the system is closed and the bypass valve opened so that both sides are pumped using an additional pump.

Before adjusting the first calibration pressure, the bypass valve is shut and the isolation valve opened. If the offset value changes, then depending on the direction of the change, either the pressure on the reference side or on the test side is higher than the lower resolution limit of the gauge, for example, due to a leak.

Since the modulus of elasticity and the inner geometry of the measuring head are temperature dependent (note that the capacitance diaphragm gauge is capable of detecting deflections as low as 0.5 nm), high-accuracy gauge heads are kept at a temperature of about 45 °C, with a control variation of only 0.02 °C.

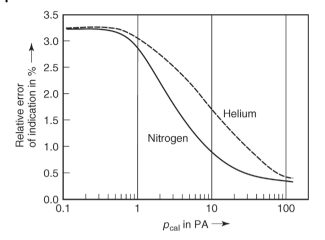


Figure 15.21 Example of a calibration curve of a capacitance diaphragm gauge at the *Physikalisch-Technische Bundesanstalt (PTB)*.

This elevated temperature T_2 of the transducer leads to thermal transpiration, covered in more detail in Sections 13.2.5.7 and 4.6.

Figure 15.21 shows a typical calibration curve of a capacitance diaphragm gauge with a pressure-independent error of indication in the molecular and viscous flow regimes as well as the transition regime. With helium, the molecular regime reaches higher pressures compared to nitrogen because the mean free path for helium is larger, considering comparison at equal temperatures and pressures.

The American Vacuum Society (AVS) published useful guidelines for calibration of capacitance diaphragm gauges [62]. Table 15.4 summarizes important aspects.

The gauge head should be calibrated in the same orientation as used in later operation, since gravitational forces change the deflection of the membrane. For differential gauge heads, as a rule, the test port should be exposed to calibration pressure because the instruments are only linearized in this direction, and the reference port should be kept as clean as possible. Vibrations and strong local airflows affect zero-point stability.

Due to the thermostatic control of the gauge head, a run-in time between 12 and 24 h is advisable. Afterward, the zero point should be adjusted. With most high-quality gauges of this type, no hysteresis effects were reported. If they are known to occur, they may be reduced by applying a preload to the membrane prior to calibration.

The result of a calibration of a capacitance diaphragm gauge is either the determination of the error of reading e or the correction factor CF, defined as

$$e = \frac{p_{\text{ind}} - p}{p} \tag{15.21}$$

Table 15.4 Selected important aspects to consider for calibrating capacitance diaphragm gauges.

Calibration of capacitance diaphragm gauges

Installation:

Low-vibration area

No strong airflow around device

Orientation of the membrane (diaphragm)

Valve for absolute gauge heads with full-scale deflection < 100 kPa

Leak test

Before calibration:

Run-in time 12 h for thermostatically controlled devices

Zero adjustment or measurement

Membrane exposed shortly to full-scale deflection

During calibration:

Ascending pressure points

Test of hysteresis, if necessary

Zero check

Dismounting:

Close isolation valve before venting

Switch heater off 1 h before venting

Always pressurize from the test side when using differential gauge heads

and

$$CF = \frac{p}{p_{ind}},\tag{15.22}$$

where p_{ind} is the indicated pressure of the device and p the generated pressure. When calibrating the voltage output, an analogous procedure is used, sometimes including a scaling factor.

Calibration is suggested to begin with the lowest pressure and to be continued to higher pressures. At each target point, a waiting time is necessary for thermal equilibrium of the gas to establish between the head and room temperature (approximately 30-60 s). The zero reading should be checked whenever possible, ideally, after each target point. In the transition regime between molecular and viscous flow as well as in the molecular regime, the temperature of the gas has to be recorded. If it deviates considerably from 23 °C, the error of indication should be corrected to the error occurring at 23 °C [46],

$$e(T_0) = e_{\text{vis}} + (e(T_1) - e_{\text{vis}}) \frac{\sqrt{\frac{T_H}{T_0}} - 1}{\sqrt{\frac{T_H}{T_1}} - 1}.$$
 (15.23)

Here, e_{vis} is the error of reading in the viscous flow regime, T_{H} the temperature of the gauge head (usually, it is sufficient to use a nominal value), and T_1 the temperature, at which calibration was performed. The same equation allows calculating e for any other measurement temperature T by replacing T_0 with T.

The correction factor CF can be replaced by

$$CF(T_0) = CF(T_1)\sqrt{\frac{T_1}{T_0}}$$
 (15.24)

in the molecular regime. In the transition regime, the ratio $CF(T_0)/CF(T_1)$ is pressure dependent and can be approximated analogously to $e(T_0)$ in Eq. (15.23) by replacing e with CF.

In the higher pressure range (> 5 kPa), relative uncertainties for readings of capacitance diaphragm gauges are dominated by their reproducibility, in the medium-high range (0.1 Pa-5 kPa), by the uncertainties of the generated or measured pressures of the primary standard, and at lower pressures, by their zero-point stability. Typically, they are in the range from 0.3% at 0.1 Pa to 0.01% at 100 kPa (95% confidence interval).

Long-term stabilities of capacitance diaphragm gauges, as for any type of vacuum gauge, are rather individual and have to be estimated by recalibrations (annually) over longer periods of time. Changes are usually characterized by jumps, but also by a continuous drift. They depend on the full-scale deflection of the considered capacitance diaphragm gauge. In the viscous flow regime, long-term stability is generally better. For a device with a full-scale deflection of 133 kPa to 13 kPa and in the viscous flow regime, long-term instability of about 0.1% per year should be expected, for a full-scale deflection of 1 kPa and 100 Pa, 0.3%-0.4% per year, for newer type capacitance diaphragm gauges with ceramic membranes, 0.1%-0.2%.

15.2.4

Spinning Rotor Gauges

The measuring principle of the spinning rotor gauge is outlined in Section 13.3.2 in detail. The measurand that has to be determined by a calibration is the effective accommodation coefficient of tangential momentum $\sigma_{\rm eff}$ (Section 13.3.2, Eq. (13.19)). It is determined by the considered surface and its roughness. The rougher the surface, the higher σ_{eff} .

Note that the spinning rotor gauge is the only vacuum meter below 10 Pa providing relatively accurate values of absolute pressure even without any calibration: if a smooth stainless-steel rotor is used as fabricated, $\sigma_{\rm eff}$ varies by about 1.5% (68% frequency interval) [64] (see Figure 13.25 in Section 13.3.5). This means that pressures can be measured with this uncertainty as far as other characteristic parameters such as diameter and density of the rotor are known. Other benefits of the spinning rotor gauge are its inertness (no molecules pumped or cracked, no outgassing), its insensitivity against thermal transpiration, and its long-term stability. These advantages make the spinning rotor gauge very useful as a secondary and reference standard. As a trade-off, relatively long measurement times must be taken into account (at very low pressures, up to 5 min per pressure point).

Spinning rotor gauges are applicable as secondary standards in the pressure range between 10⁻⁴ and 1 Pa. At lower pressures, fluctuations of the residual drag grow too large; at pressures > 1 Pa, gas friction causes unacceptably large temperature changes in the measuring system. Generally, two methods are available for determining the effective tangential accommodation coefficient $\sigma_{\rm eff}$:

1) The spinning rotor gauge is exposed to a pressure of about 0.01 Pa and its reading p_{ind} is compared to the actual pressure p. After all parameters such as diameter and density of the rotor, temperature, offset, and $\sigma_{\rm eff}=1$ have been entered into the controller, σ_{eff} is given by (see Eq. (13.19))

$$\sigma_{\text{eff}} = \frac{p_{\text{ind}}}{p}.\tag{15.25}$$

2) The second method uses the fact that $\sigma_{\rm eff}$ drops linearly with pressure up to about 2 Pa [65] if no viscosity correction is made in the controller (viscosity = 0 entered). If $\sigma_{\rm eff}$ is determined at several points between 0.1 and 1 Pa, $\sigma_{\rm eff}(p)$ may be extrapolated for $p \to 0$. An example of such a calibration is shown in Figure 15.22. Since higher pressures can be generated with lower uncertainty, $\sigma_{\rm eff}$ is determined more accurately, which is an advantage of this calibration method.

Table 15.5 lists a few points to consider when calibrating spinning rotor gauges. Vibrations on the measurement system have to be avoided. These would increase measurement uncertainty considerably. Because of the temperature effects mentioned in Section 13.3.2 during the first acceleration of the rotor, an initial run-in time of at least 6 h has to be allowed after starting before calibration can be initiated. Temperature drifts around the measurement head must be avoided under all circumstances.

As mentioned earlier, the reading of the spinning rotor gauge is indifferent against the thermal transpiration effect. This is understandable since the condition of equilibrium for calculating the effect is that the same numbers of molecules enter and exit the volume with the higher temperature. Since the spinning rotor gauge measures the impingement rate of molecules, its indication is independent of a local rise in temperature. Thus, the mean temperature of the large calibration vessel has to be entered in the controller and not the local temperature (e.g., the temperature at the thimble of the gauge).

For applying spinning rotor gauges as reference standards at low pressures, it must be considered that the magnitude of the residual drag may depend on the frequency of the rotor (Section 13.3.3, Figure 13.24). Either this dependence

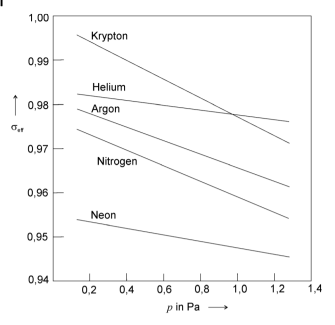


Figure 15.22 The effective accommodation factor $\sigma_{\rm eff}$ (Eq. 15.25) of a spinning rotor gauge for several gas species according to a calibration at PTB. $\sigma_{\rm eff}$ depends linearly on pressure up to about 2 Pa.

should be determined before measurement or the residual drag should be determined immediately before and after a measurement.

To obtain high long-term stability, any circumstance changing the surface should be avoided. Mechanical friction may change surface roughness so that the rotor should be fixed during transport. For rotors made of standard steel, corrosion must be avoided. Keeping the rotor under vacuum at all times is advisable. Both requirements are fulfilled by employing a special transport device as developed by $R\ddot{o}hl$ and Jitschin [66]. Also baking may change the value of σ_{eff} by up to 2%.

Table 15.5 Important aspects to consider for calibrating spinning rotor gauges.

Calibration of spinning rotor gauges (SRG)

- Absence of strong vibrations
- Stable temperatures around gauge head
- Baking may change accommodation factor
- Run-in time at least 6 h
- Frequency dependence of residual drag (significant for lower pressures)
- During transport: rotor stands still and is not subject to any corrosion

The relative uncertainty of $\sigma_{\rm eff}$ during calibration is typically 0.3% (95% confidence interval). The long-term instability of well-treated rotors lies between 0.3% and 0.5% over a one-year period. However, occasional step changes of 1% may occur. For a rotor calibrated for the first time, the latter value should be applied.

15.2.5

Ionization Gauges

Section 13.7 explains the measuring principle and designs of ionization gauges. Even though ionization gauges exhibit several metrological disadvantages, among them mediocre reproducibility and long-term stability, there is no alternative to using them as secondary standards below 10⁻⁴ Pa. Hitherto, only ionization gauges with a hot emissive cathode have been employed as reference standards.

In principle, the ion gauge sensitivity S (Eq. (13.33)) can be calibrated. For this, however, ion currents and emission currents have to be measured with highquality calibrated electrometers, and the distribution of emission currents along the cathodes during calibration must be equal to that prevailing in future applications.

The last point is particularly difficult to realize. Thus, calibrating ion gauge sensitivity became less and less important. Instead, the measuring head, wire, as well as control and monitoring equipment are treated as a single calibration unit. In this case, calibration of the electrical devices is dispensable thus yielding cost advantages. The calibration unit is then calibrated for the error of reading e or the correction factor CF according to Eqs. (15.21) and (15.22).

Unfortunately, the electrometers used in the measuring units are usually not of the highest quality, and thus, e or CF changes from one decade to the next, since the measuring resistors are insufficiently accurate. If an external top-grade electrometer is used, these changes disappear.

Physically, an ionization gauge measures gas density; however, it indicates pressure. For high accuracies, it is therefore necessary to consider the temperature. The hot cathode leads to a considerable rise in temperature in the gauge head compared to the temperature in the calibration chamber so that the thermal transpiration effect also plays a role. If the heat transferred from the measuring head to the vacuum system is independent of the latter, this effect is calibrated into the calibration value. Temperature corrections can be made as follows [46,67]:

$$e(T_0) = \frac{p_{\text{ind}} \frac{T_{\text{ch}}}{T_0} - p_{\text{ch}}}{p_{\text{ch}}},$$
(15.26)

where $p_{\rm ch}$ is the pressure generated in the calibration chamber at temperature $T_{\rm ch}, p_{\rm ind}$ is the indicated pressure at $T_{\rm ch}$, and T_0 the desired reference

Table 15.6 Important aspects to consider for calibrating ionization gauges.

Calibration of ionization gauges

Tubulate nude gauges

Controller plus gauge head = measuring unit

Orientation of gauge head

Warm-up period 12 h

Degassing by electron bombardment

Cleaning of ion collector by operation at 'high' pressure

Residual current measurement

Ascending pressure sequence

temperature, usually 23 °C. The modified correction factor

$$CF(T_0) = CF(T_{ch}) \frac{T_0}{T_{ch}}.$$
 (15.27)

For vacuum sensitivity S,

$$S(T_0) = S(T_{\rm ch}) \frac{T_{\rm ch}}{T_0}$$
 (15.28)

Table 15.6 summarizes important aspects for calibrating ionization gauges as secondary standards.

The distribution of potential inside the gauge head depends on the distance to the surrounding grounded walls. For a nude gauge calibrated as secondary standard, it is therefore advisable to enclose the gauge head with a non-removable tube throughout its full length. In order to transport the gauge head under vacuum, this tube may be sealed using a UHV-compatible valve. The orientation of the gauge head should be identical during calibration and use, since geometrical deformations due to different orientations may affect the distribution of potential as well as electron trajectories in the gauge head.

Since the ion current depends significantly on the surface state of the electrodes, for example, due to secondary electron yield [68-71] the electrode surfaces should be kept as clean as possible at all times in order to obtain reproducible results. For this reason, the ion gauge is baked out and cleaned in 'degassing mode' during the cooling phase: compared to operation, the cathode is cleaned using much higher temperatures, and the anode by employing simultaneous electron bombardment.

The ion collector is cleaned with ion bombardment by operating the ionization gauge at high argon pressure $(10^{-3}-10^{-2} \text{ Pa})$ for about 1 h [72]. After these conditioning procedures, the gauge has to be operated in normal mode (regular emission current) for 12h before calibration. The residual pressure in the calibration system should be at least one order of magnitude below the lowest calibration pressure. This requirement cannot be met for pressures of 10⁻⁹ Pa or less. Generally, a relevant signal at residual pressure should be subtracted from the signal obtained at calibration pressure.

Although, in principle, it should be possible to calibrate a gauge for one gas species and use this calibration for some other gas species using the ionization probability ratio for the two gas species as scaling factor, investigations have shown [73,74] that this is not feasible if high accuracy (uncertainty < 10%) is required. Even for isotopes of the same gas species, for example, hydrogen and deuterium, significant differences in sensitivities were found and explained [70] (see Section 13.9).

Ionization gauges should be calibrated with three points per decade in a rising sequence throughout their operating range. The sensitivity is at least slightly pressure dependent (with about a few percent over several decades), even below 10⁻³ Pa [75]. The relative uncertainties of calibration for the reading and for sensitivity depend on the pressure and amount to 40% at 10⁻¹⁰ Pa, 2%-3% at 10^{-6} Pa, and 0.5%-1% at 10^{-2} Pa.

The long-term instability of any IG is characteristic for each gauge and depends on its use [76]. For a high-quality ionization gauge used as reference standard under clean conditions, a value of 3%-6% over a one-year period may serve as reasonable estimate for its instability during this period [77]; however, according to measurements at PTB 1% can also be obtained.

15.3 Calibrations of Residual Gas Analyzers

Residual gas analyzers (Section 14.2) are used extensively for qualitative and quantitative analyses of gas mixtures in many applications in industry and research. Especially in the microelectronics industry, their correct and accurate readings are of significant economic importance. The purity of gases has been improved continuously for large-scale integration of devices, and monitoring of impurities in the process gases is required on a routine basis.

Today, most residual gas analyzers for general purposes are of the quadrupole type, dominating about 95% of the market [78]. Therefore, the following sections focus on quadrupole mass spectrometers. The requirement to achieve reproducible results in gas purity analyses is opposed by the relatively mediocre metrological quality.

It is impossible to calibrate quadrupole mass spectrometers such that quantitative interpretations of any mass spectrum would be possible. The sensitivity S_i , Eq. (14.3), for one gas species may significantly depend on the presence of other gas species of higher pressures [79]. This is due to space charge effects in the ion source that change the probability for ion extraction. Collisions between ions and neutral particles may alter the ratios of different ion species.

Investigations on the stability of quadrupole mass spectrometers have shown that their sensitivities may change by a factor of 2 over a period of 220 days [80]. In this case, ion collectors of the Faraday principle were used. The use of electron multipliers aggravates the problem due to the alteration of electron multiplication by aging and bake-out cycles [81]. Another problem is outgassing of the residual gas analyzers themselves. Many of the constituents of the residual gas spectrum are generated in the device itself. Several investigations showed that gases such as methane, water vapor, carbon monoxide, and carbon dioxide develop when hydrogen is introduced [81–83].

For these reasons, mass spectrometers are normally not calibrated against primary standards of the continuous expansion type, although this would be possible by using two or more flowmeters. With each flowmeter, a known gas flow of a different gas species may be introduced, and the corresponding partial pressure calculated with Eq. (15.14). However, the effort involved with using several flowmeters is rather large and rarely justified.

The simplest calibration method for a mass spectrometer is to compare it to an ionization gauge. The reading of the latter is more stable and features better reproducibility and linearity [79]. The sensitivity S_i for gas species i is determined by

$$S_i = \frac{I_A - I_0}{p_i - p_0},\tag{15.29}$$

where I_A and p_i denote the ion collector current and the pressure as measured with the ionization gauge after gas has been introduced, I_0 and p_0 are the ion collector current and the pressure prior to gas introduction, respectively, (reading in equivalents of the introduced gas species i). It also possible to calibrate gas mixtures in ratios of 1:1 to 1:10 using this method if the gases are introduced consecutively, since below 10⁻³ Pa, ionization gauges react linearly to adding of a further gas constituent. For implementing such a comparison system, the guidelines of Section 15.2.2 apply as well.

Another possibility is calibrating by comparison with a spinning rotor gauge. The overlap of the operating ranges of the two devices from about 10⁻⁴ to 10⁻² Pa, however, is in a range where the reading of the quadrupole mass spectrometer is affected by space charge effects and non-linearities, so that extrapolation to lower pressures is usually impossible.

In addition to total pressure gauges, test leaks [84] and capillary leaks may also be applied for calibration [85]. For known gas flow rate and effective pumping speed of the calibration chamber, partial pressure can be determined [84]. This method is primarily used for calibrating a tracer gas in a matrix gas. An ionization gauge measures the pressure of the latter. Quite often, capacitance diaphragm gauges measuring relatively high pressures are used to establish gas mixtures in a reservoir in front of the calibrated

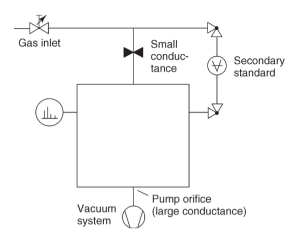


Figure 15.23 Calibration of a mass spectrometer with a known small conductance and a secondary standard measuring the pressure difference across the conductance.

capillary. When there is no calibrated leak or capillary, a secondary standard may be used to measure the differential pressure across the leak or a small conductance (Figure 15.23).

For this procedure, the pressure ratio must be independent of pressure and gas species, that is, the flow through the leak must be molecular. When dosing valves are used, it has to be assured that their conductance is stable. Run-in times of several hours may be necessary.

A committee of the American Vacuum Society (AVS) published recommendations on the calibration of mass spectrometers [85,86]. Table 15.7 summarizes important points to be considered for calibration. Note, in particular, that voltage settings, resolution, and so on influence sensitivity values significantly and have to be recorded during calibration.

The relative uncertainties for determining the ε_i for defined gas mixtures are hardly below 10%. Lower uncertainties can be achieved only with repeated in situ calibration of mass spectrometers.

15.4 Calibration of Test Leaks

Section 14.4.4 discusses designs and applications of test leaks. At constant temperature, test leaks provide a constant gas flow, usually of helium because leak detectors are calibrated with helium. These helium test leaks are based mainly on the principle of permeation (Section 14.4.4.1). Capillary leaks (14.4.4.2) may be used with any gas species that does not clog the capillary or react with its surface.

Table 15.7 Important aspects to consider for calibrating quadrupole mass spectrometers.

Calibration of quadrupole mass spectrometers

- For UHV:
- Cleanliness of gauge head and calibration system
- Thorough bake-out
- Residual partial pressure < 1/10 of calibration pressure
- Before calibration:
- · Connect to ground
- Warm-up time 6 h
- Optimize settings (resolution, scan speed, ion energy, etc.)
- Shape and repeatability of mass peaks
- During calibration:
- · Record all settings
- · Measure background offset
- · Measurement of scatter and drift of background offset
- · Outgassing of mass spectrometer
- · Fragmentation of molecules
- Linearity
- Dependence of sensitivity on total pressure and gas mixture

Both types of leaks are temperature dependent. The permeation-type leak changes its flow rate by about 3% K⁻¹ so that it should be temperaturecontrolled to a variation of 0.1 °C during calibration.

For users of test leaks, it is often convenient to obtain the value of the flow rate at a given temperature of the permeation leak or the capillary as throughput q_{nV} in the unit Pa ℓ s⁻¹. This quantity, however, is incomplete if the temperature T at which the throughput was measured is not given. The value of this temperature is redundant if the flow rate is given as molar flow rate q_{ν} , that is, as the number of moles passing through the leak per unit time.

The two quantities can be converted:

$$q_{\nu} = \frac{d\nu}{dt} = \frac{q_{pV}}{RT} = \frac{d(pV)}{dt}(RT).$$
 (15.30)

Thus, it is advisable to record q_{ν} for a test leak [87–89].

Rewriting this equation already indicates the two main calibration methods:

$$q_{\nu} = \frac{p_{\frac{dV}{dt}}^{dV} + V_{\frac{dp}{dt}}^{dp}}{RT}.$$
 (15.31)

- 1) The pressure is maintained constant to make the second term equal to zero.
- 2) The volume is kept constant to set term one to zero.

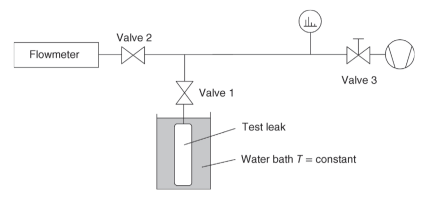


Figure 15.24 Experimental setup for calibrating a test leak by comparison with a flowmeter.

Since flowmeters operate on the same principle, it is possible to calibrate test leaks by comparing their gas flow rates with that of a flowmeter. A mass spectrometer adjusted to the corresponding gas species serves as indicator. It does not require any calibration. Figure 15.24 shows the measurement diagram.

Similarly as in the calibration chamber of the continuous expansion or comparison method, the mass spectrometer must not be exposed directly or preferentially to one of the gas flows. Instead, it should either be positioned symmetrically to both gas sources or it should be assured, for example, using elbows or long tubing, that equal flow rates from the two sources yield equal signals on the indicator.

By operating one of the flowmeters at PTB with the method of constant conductance [40], test leaks can be calibrated down to $10^{-15} \, \mathrm{mol s^{-1}} \, (2 \times 10^{-9} \, \mathrm{Pa} \ell \, \mathrm{s}^{-1}$ at 23 °C). If there is no flowmeter available, either the method of constant pressure or of constant volume must be applied directly.

Method of Constant Pressure:

Necessary equipment is shown in Figure 15.25.

The gas from the test leak is introduced to volume V_1 until a certain pressure p_1 measured by a secondary standard is reached. After opening a valve to the previously evacuated volume V_2 , the pressure will drop. The time period t_1 until p_1 reestablishes is measured. The flow rate is then given by

$$q_{\nu} = \frac{p_1 V_2}{t_1} (RT). \tag{15.32}$$

Instead of a fixed volume V_2 , a piston–cylinder system may also be used [90]. Similar to the procedure using a flowmeter, the volume speed required for maintaining constant pressure is measured.

Method of Constant Volume:

By measuring the pressure rise caused by the test leak in a not too small volume ($\geq 50 \, \mathrm{cm}^3$), q_{ν} can be calculated with the second term in Eq. (15.31). For long measurement periods such as several weeks, flow rates down to

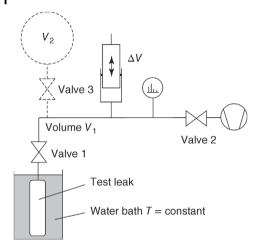


Figure 15.25 Calibration of a test leak via the method of constant pressure, using either a vessel of known volume (dashed line) or a volume displacer (piston–cylinder system AV).

 10^{-18} mol s⁻¹ are measurable. In this case, and generally for test leak measurements, the following possible disturbing effects should be considered:

- adsorption on and desorption from the walls;
- · outgassing from the walls;
- · leaks in the vacuum system;
- temperature drift;
- instability and drift of background noise;
- pumping and outgassing effects of the vacuum device.

Mainly due to the last point, spinning rotor gauges are used for these kinds of measurements.

If permeation leaks are isolated by valves for longer periods of time, there is a steady pressure rise on the downstream side of the leak that reduces the concentration gradient within the permeating material. When the valve is opened, some relaxation time is needed until a stationary equilibrium is obtained.

15.5 Standards for Determining Characteristics of Vacuum Pumps

Several sections of Chapters 6 to 11 cover important characteristics of vacuum pumps. Table 15.8 summarizes these characteristics.

Pumping speed S, Eq. (4.19), is determined by measuring the pressure p above the pump inlet, when a known throughput rate q_{pV} is injected into

Parameter	er Symbol Applies to		Corresponding standards (DIN)	Corresponding standards (ISO)
Pumping speed	S	All pumps	28426 (part 1), 28427, 28428, 28429, 28431, 28432	1607 (part 1), 1608 (part 1), 5302, 21360 (2 parts)
Ultimate pressure	$\mathcal{P}_{ ext{ult}}$	All pumps	28426 (part 1), 28427, 28428, 28429, 28431, 28432	1607 (part 2), 1608 (part 2), 5302
Ultimate partial pressure		Oil-sealed rotary vane pumps	28426 (part 1)	1607
Starting pressure	$p_{ m start}$	All high-vacuum pumps	28428, 28429	
Foreline pressure (maximum admissible)	$(p_{ m fore})_{ m max}$	Diffusion pumps Turbomolecular pumps	28427	1608 (part 2), 5302
Compression ratio	K	Roots pumps Turbomolecular pumps	28426 (part 2) 28428	5302
Water vapor tolerance pressure	$p_{ m w}$	Rotary vane pumps	28426 (part 1)	1607 and 21360 (part 2)

Table 15.8 Important vacuum-technological characteristics of vacuum pumps.

the pump,

$$S = \frac{q_{pV}}{p - p_0}. (15.33)$$

Here, p_0 is the pressure when $q_{pV} = 0$. p should be at least twice as high as

While q_{pV} is comparably simple to determine using a flowmeter or other methods (see below), measuring p just in front of the pump inlet is relatively problematic due to the directed gas flow. In a thermodynamic sense, pressure pis well defined only in an enclosed system under equilibrium conditions. In the ideal physical model, an infinitely large volume above the pump inlet is needed where the opening of the inlet would have a negligible effect on the isotropic Maxwellian distribution in this volume [91,92]. Since the inlet flanges of highvacuum pumps are of considerable size, practical chambers above the inlet cannot be designed large enough to meet this ideal condition. Instead, a real chamber always shows a significant pressure gradient and the motion of the gas molecules is not isotropic.

For this reason, it is necessary to agree on international standards defining at which place above the inlet and in which orientation pressure p should be measured. Two basic ideas guided the development of these standards: first, in a standard chamber (test dome) a similar value should be measured as in the ideal case (infinite volume). Second, a standard chamber should be similar to a practical chamber, so that the pumping speed is a meaningful quantity for the development of vacuum systems.

Tables 15.8 and A.22 (in the Appendix) give an overview of national and international standards. In 1987 [93] and 1989 [94], the American Vacuum Society (AVS) changed their recommendations for the determination of pumping speeds in order to meet the corresponding standards of ISO, PNEUROP, and DIN. Since 2004, ISO are revising their standards; the basic content, however, remains unchanged.

In all standards or recommendations, the test dome must be of the same inner diameter as the pump's inlet flange. ISO, however, requires this for flange diameters down to 100 mm, while the AVS recommendation extends down to 50 mm. ISO specifies both the shape and the size of the adapter from the test dome to the pump's inlet flange to be < 100 mm. The upper end of the test dome must be rounded, conical, or inclined. This shape specification was mainly meant to provide that any oil from diffusion pumps possibly condensing on the cap drains to the side of the dome and does not drip back into the pump where it could cause pressure spikes. For other oil-free or nearly oil-free pumps, this aspect is irrelevant.

Two methods to measure q_{nV} are recommended by the ISO/PNEUROP/DIN standards. Where no suitable gas flowmeters with sufficient accuracy are available, a two-dome configuration can be used (Figure 15.27) to determine q_{nV} by the pressure difference across the orifice of known conductance.

The pumping speed can then be calculated from

$$S = \frac{q_{pV}}{p_2} = C\left(\frac{p_1}{p_2} - 1\right),\tag{15.34}$$

where C is the conductance between the two domes, and p_1 and p_2 are the pressures above and below. The conductance is calculated from Eq. (15.17) (thin orifice).

Where suitable flowmeters are available (typically at inlet pressures above $10^{-4}\,\mathrm{Pa}$), a single dome (Figure 15.26) is used and the measurement of q_{nV} is carried out by the flowmeter. The results obtained by the two methods do not completely agree in an overlapping range [95], which is not too surprising since different molecular flows exist in the two kinds of test domes.

As an alternative to using the single dome according to the throughput method at constant pressure, ISO 21360 and DIN 28432 state a further method, mainly for determining the pumping speeds of diaphragm pumps. To curb the relatively elaborate measurement of $q_{\nu V}$, a large volume is pumped out intermittently via a quick acting valve and the pressure is measured before and after each

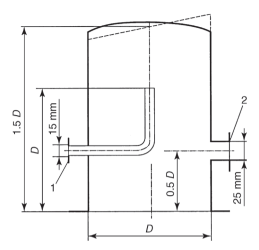


Figure 15.26 Single dome for measuring pumping speeds of high-vacuum pumps according to DIN 28428 and 28429.

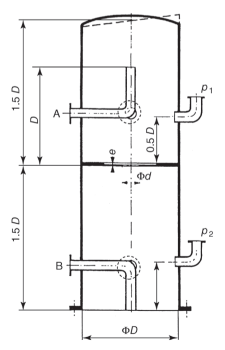


Figure 15.27 Double dome after Fischer-Mommsen for measuring the pumping speed of highvacuum pumps (DIN 28429).

pump-down cycle. Due to the temperature changes during a cycle, a sufficiently long period of time has to elapse before the measurement of pressure can take place. This method is simple to automate.

Other methods of measuring pumping speeds are available, for example, the method of conductance modulation [96,97]. As mentioned, however, results need to be comparable, so that sole use of standardized methods is strongly recommended.

References

- 1 Jousten, K. and Röhl, P. (1995) Instability of the spatial electron current distribution in hot cathode ionization gauges as a source of sensitivity changes. J. Vac. Sci. Technol. A, 13, 2266-2270.
- 2 JCGM200: (2012) International vocabulary of metrology - Basic and general concepts and associated terms, 3rd edition with minor corrections from 2008, available from www.bipm.org.
- 3 Messer, G., Jitschin, W., Rubet, L., Calcatelli, A., Redgrave, F.J., Keprt, A., Weinan, F., Sharma, J.K.N., Dittmann, S., and Ono, M. (1989) Intercomparison of nine national high-vacuum standards under the auspices of the bureau international des poids et mesure. Metrologia, 26, 183.
- 4 Sharma, J.K.N., Mohan, P., Jitschin, W., and Röhl, P. (1989) Intercomparison of vacuum standards between the PTB and the NPL (India) using two spinning rotor gauges. J. Vac. Sci. Technol. A, 7, 2788.
- 5 Wangkui, L., Quiang, Liu., Zhenhai, Li., Messer, G., and Grosse, G. (1992) Intercomparison of vacuum standards between LIP and PTB. Vacuum, 43, 1091.
- 6 Jousten, K., Filipelli, A.R., Tilford, C.R., and Redgrave, F.J. (1997) Comparison of the standards for high and ultrahigh vacuum at three national standards laboratories. J. Vac. Sci. Technol. A, 15, 2395-2406.
- 7 Miiller, A. et al. (2002) Final report on key comparison CCM.P-K4 of absolute pressure standards from 1 Pa to 1,000 Pa. Metrologia, 39, Tech. Suppl., 07001.
- 8 Jousten, K. et al. Results of the regional key comparison Euromet.M.P-K1.b in the pressure range from 3· 10⁻⁴ Pa to 0.9 Pa,

- Technical Supplement to Metrologia, 2004, 42 Tech. Suppl. 07001.
- 9 Jousten, K., Romero, L.A.S., and Guzman, J.C.T. (2005) Results of the key Comparison SIM-Euromet.M.P-BK3 (bilateral comparison) in the pressure range from 3×10^{-4} Pa to 0.9 Pa. Metrologia, 42, Tech. Suppl., 07002.
- 10 Calcatelli, A. et al. (2005) Final report on key comparison EUROMET.M.P-K1.a, Euromet project 442 A, pressure range: 0.1 Pa-1000 Pa. Metrologia, 42, Tech. Suppl., 07001.
- 11 Jousten, K., Bock, Th., Dominik, Pražák., and Zdeněk, Krajíček. (2007) Final report on the supplementary comparison EUROMET.M.P-S2 (bilateral comparison) in the pressure range 30 Pa to 7000 Pa. Metrologia, 44, Tech. Suppl., 07007.
- 12 Olson, D.A., Abbott, P.J., Jousten, K., Redgrave, F.J., Mohan, P., and Hong, S.S. (2010) Final report of key comparison CCM.P-K3: absolute pressure measurements in gas from 3×10^{-6} Pa to 9×10^{-4} Pa. Metrologia, 47, Tech. Suppl., 07004.
- 13 Yoshida, H., Arai, K., Akimichi, H., Hong, S.S., and Song, H.W. (2011) Final Report on key comparison APMP.M.P-K3: Absolute pressure measurements in gas from 3×10^{-6} Pa to 9×10^{-4} Pa. Metrologia, 48, Tech. Suppl., 07013.
- 14 Krajíček, Z., Bergoglio, M., Jousten, K., Otal, P., Sabuga, W., Saxholm, S., Pražák, D., and Vičar, M. (2014) Final report on EURAMET.M.P-K4.2010 - key and supplementary comparison of national pressure standards in the range 1 Pa to 15 kPa of absolute and gauge pressure. Metrologia, 51, Tech. Suppl., 07002.

- 15 Fedchak, J.A., Bock, Th., and Jousten, K. (2014) Bilateral key comparison CCM.P-K3.1 for absolute pressure measurements from 3×10^{-6} Pa to 9×10^{-4} Pa. Metrologia, 51, Tech. Suppl., 07005.
- 16 Adametz, H. and Wloka, M. (1991) Measurements of the absolute density of mercury in the ASMW. Metrologia, 28, 333-337.
- 17 Sommer, K.-D. and Poziemski, J. (1993/94) Density, thermal expansion and compressibility of mercury. Metrologia, 30, 665-668.
- 18 Bennet, S.J., Clapham, P.B., Dadson, J.E., and Simpson, D.I. (1975) Laser interferometry applied to mercury surfaces. J. Phys. E: Sci. Instrum., 8, 5-7.
- 19 Jäger, J. (1993/94) Use of a precision mercury manometer with capacitance sensing of menisci. Metrologia, 30, 553-558.
- 20 Bauer, H. (1979) Die Darstellung der Druckskala im Bereich von 0,01 bar bis 2 bar durch das quecksilbermanometer der PTB. PTB-Mitteilungen, 89, 248-255.
- 21 Heydemann, P.L. (1971) A fringe-counting pulsed ultrasonic interferometer. Rev. Sci. Instrum., 42, 983-986.
- 22 Tilford, C.R. (1987) The speed of sound in a mercury ultrasonic interferometer manometer. Metrologia, 24, 121–131.
- 23 Gaede, W. (1915) Die diffusion der gase durch quecksilberdampf bei niederen drucken und die diffusionsluftpumpe. Ann. Phys, 46, 357-392.
- 24 Ishii, H. and Nakayama, K. (1962) 8th AVS Nat. Vac. Symp, 519.
- 25 Meinke, C. and Reich, G. (1963) Influence of diffusion on the measurement of low pressure with the McLeod vacuum gauge. Vacuum, 13, 579-581.
- 26 Klingenberg, G. and Lüdicke, F. (1991) PTB-Mitteilungen, 101, 7-18.
- 27 Tilford, C.R., Hyland, R.W., and Sheng, Y.T. (1989) Yi-Tang, BIPM. BIPM Monogr., 89/1, 105-113.
- 28 Rendle, C.G. and Rosenberg:, H. (1999) New absolute pressure standard in the range 1 Pa to 7 kPa. Metrologia, 36, 613-615.
- 29 Ooiwa, A. and Ueki, M. (1993) Development of novel air piston gauge for medium vacuum and fine differential

- pressure measurement. Vacuum, 44, 603-605.
- 30 Ooiwa, A. (1993) Novel nonrotational piston gauge with weight balance mechanism for the measurement of small differential pressures. Metrologia, 30, 607-610.
- 31 Knudsen, M. (1910) Thermischer molekulardruck der Gase in Röhren und porösen Körpern. Ann. Phys. (Leipzig), 31,
- 32 Schuman, S. (1962) Trans. Natl. Vac. Symp., 9, 463.
- 33 Bergoglio, M., Calcatelli, A., Marzola, L., and Rumanio, G. (1988) Primary measurements down to 10⁻⁶ Pa. Vacuum, **38**, 887.
- 34 Jitschin, W., Migwi, J.K., and Grosse, G. (1990) Pressures in the high and medium vacuum range generated by a series expansion standard. Vacuum, 40, 293-304.
- 35 Jousten, K., Aranda Contreras, V., and Röhl, P. (1999) Volume ratio determination in static expansion systems by means of a spinning rotor gauge. Vacuum, 52, 491-499.
- 36 Elliott, K.W.T. and Clapham, P.B. (1978) The accurate measurement of the volume ratios of vacuum vessels, NPL Report MOM.
- 37 Berman, A. and Fremerey, J.K. (1987) Precision calibration of a static pressure divider by means of a spinning rotor gauge. J. Vac. Sci. Technol. A, 5, 2436-2439.
- 38 Peggs, G.N. (1976) The measurement of gas throughput in range 10^{-4} to 10^{-10} Pa m³/s. Vacuum, **26**, 321–328.
- 39 Bennewitz, H.G. and Dohmann, H.D. (1965) Dynamische druckbestimmung. Vak. Tech., 14, 8.
- 40 Jousten, K., Messer, G., and Wandrey, D. (1993) A precision gas flowmeter for vacuum metrology. Vacuum, 44, 135-151.
- 41 Szwemin, P., Szymansky, K., and Jousten, K. (1999) Monte Carlo study of a new PTB primary standard for very low pressures. Metrologia, 36 (6), 561-564.
- 42 Grosse, G. and Messer, G. (1970) Experiences with cryopumps in calibration devices. Vacuum, 20, 373-376.

- 43 Poulter, K.F. (1978) Vacuum gauge calibration by the orifice flow method in the pressure range 10^{-4} to 10 Pa. Vacuum, 28, 135.
- 44 Liepmann, W. (1961) Gas kinetics and gas dynamics of orifice flow. J. Fluid Mech., 10, 65.
- 45 Sharma, J.K.N., Mohan, P., and Sharma, D.R. (1990) Comparison of two primary pressure standards using spinning rotor gauges. J. Vac. Sci. Technol. A, 8,
- **46** Jousten, K. (1998) Temperature corrections for the calibration of vacuum gauges. Vacuum, 49, 81-87.
- **47** Tison, A. and Tilford, C.R. (1993) RL/NIST Workshop on Moisture Measurement and Control for Microelectronics (eds B.A. Moore and J.A. Carpenter, Jr.), NISTIR, NIST, Washington DC, pp. 19-29.
- 48 Roehrig, J.R. and Simons, J.C. (1962) Trans. Natl. Vac. Symp., 9, 511.
- 49 Feakes, F. and Torney, F.L. (1963) Trans. Natl. Vac. Symp., 10, 257.
- 50 Jousten, K., Menzer, H., and Niepraschk, R. (1999) New fully automated, primary standard for generating pressures between 10^{-10} Pa and 3×10^{-2} Pa with respect to residual pressure. Metrologia, 36, 493-497.
- **51** Grosse, G. and Messer, G. (1981) Calibration of vacuum gauges at pressures below 10⁻⁹ mbar with a molecular beam method. Vak. Tech., 30, 226.
- 52 Kuz'min, V.V. (1995) Development of pressure-time dependence methods for the calibration of vacuum gauges: a review. Vacuum, 46, 251.
- 53 Looney, J.P. (1993) Measurement of CO pressures in UHV regime using resonance-enhanced mulitphotononization time-of flight mass spectrosocopy. J. Vac. Sci. Technol. A, 11, 3111-3120.
- 54 Shimizu, H., Hashizume, H., Ichimura, S., and Kokubun, K. (1988) Detection of sputtered neutral atoms by nonresonant multiphoton ionization. Japan. J. Appl. Phys., 27, 502-505.
- 55 Ichimura, S., Kokubun, K., Shimizu, H., and Sekine, S. (1996) Possibility and current status of absolute XHV

- measurement by laser ionization. Vacuum, 47, 545-552.
- 56 Lanzinger, E., Jousten, K., and Kühne, M. (1998) Partial pressure measurement by means of infrared laser absorption spectroscopy. Vacuum, 51, 47 - 51.
- 57 Hodges, J.T., Looney, J.P., and van Zee, R.D. (1996) Response of a ringdown-cavity to an arbitrary excitation. J. Chem Phys., 105, 10278-10288.
- 58 Jousten, K., Padilla, G., and Bock, T. (2008) Investigation on tunable diode laser absorption spectroscopy for its application as primary standard for partial pressure measurements. J. Phys.: Conf. Ser., 100, 092005.
- 59 Egan, P.F. et al. (2015) Performance of a dual Fabry - Perot cavity refractometer. Opt. Lett., 40 (17) 3945-3948.
- 60 ISO 3567 (2011) Vacuum Gauges -Calibration by direct comparison with a reference gauge.
- 61 Guideline DKD-R 6-2, Calibration of Measuring Devices for Vacuum, 03/2002, Part 1 Fundamentals, Part 2 Measurement Uncertainties, Part 3 Electrical Diaphragm Gauges, Part 4 Ionization Gauges, Part 5 Pirani Gauges, available from www.dkd.eu or Deutscher Kalibrierdienst, Accreditation Body, Bundesallee 100, 38116 Braunschweig, Germany.
- 62 Hyland, R.W. and Shaffer, R.L. (1991) Recommended practices for the calibration and use of capacitance diaphragm gages as transfer standards. J. Vac. Sci. Technol. A, 9, 2843.
- 63 Ellefson, R.E. and Miiller, A.P. (2000) Recommended practice for calibrating vacuum gauges of the thermal conductivity type. J. Vac. Sci. Technol. A, **18**, 2568–2577.
- 64 Dittmann, S., Lindenau, B., and Tilford, C.R. (1989) The molecular drag gauge as a calibration standard. J. Vac. Sci. Technol. A, 7, 3356-3360.
- 65 Messer, G. and Röhl, P. (1984) Druckabhängigkeit der Koeffizienten fürden tangentialen Impulsaustausch an Gasreibungsmanometer-Kugelrotoroberflächen, PTB Annual report 226.

- 66 Röhl, P. and Jitschin, W. (1988) Performance of the spinning rotor gauge with a novel transport device as a transfer standard for high vacuum. Vacuum, 38,
- 67 Abbott, P.J., Looney, J.P., and Mohan, P. (2005) The effect of ambient temperature on the sensitivity of hot-cathode ionization gauges. Vacuum, 77, 217-222.
- 68 Becker, H.U. and Messer, G. (1980) Sensitivity dependence on collector surface properties in ion gauges. Vide Suppl., 201, 234-237.
- 69 Grosse, G., Harten, U., Jitschin, W., and Gentsch, H. (1987) Secondary electrons in ion gauges. J. Vac. Sci. Technol. A, 5, 3242-3243.
- 70 Jousten, K. and Röhl, P. (1995) Comparison of the sensitivities of ionization gauges to hydrogen and deuterium, Vacuum, 46, 9-12.
- 71 Ave, H., Becker, H.U., and Messer, G. (1985) Stabilisierung der Vakuummeterkonstanten von lonisationsvakuummetern. PTB Mitteilungen, 951, 20.
- 72 Becker, H.U. and Messer, G. (1983) Proc. 5th Int. Vac. Congress/9th Int. Conf. on Surface Science, Madrid, 84 (unpublished).
- 73 Holanda, R. (1973) Investigation of the sensitivity of ionization-type vacuum gauges. J. Vac. Sci. Technol., 10, 1133.
- 74 Filipelli, A. (1988) A brief review of some recent sensitivity and residual current measurements for several commerical ionization gauges. AIP Conf. Proc., 171, 236-243. http://dx.doi.org/10.1063/ 1.37305.
- 75 Tilford, C.R., Mc Culloh, K.E., and Seung Woong, H. (1982) Performance characteristics of a broad range ionization gauge tube. J. Vac. Sci. Technol., 20, 1150.
- 76 Wood, S.D. and Tilford, C.R. (1985) Longterm stability of two types of hot cathode ionization gauges. J. Vac. Sci. Technol. A, 3, 542.
- 77 Filipelli, A.R. and Abbott, P.J. (1995) Longterm stability of bayard-alpert gauge performance: results obtained from repeated calibrations against the national institute of standards and technology primary vacuum standard. J. Vac. Sci. Technol. A, 13, 2582.

- 78 Lichtman, D. (1990) Residual gas analysis: Past, present, and future. J. Vac. Sci. Technol. A, 8, 2810.
- 79 Lieszkovsky, L., Filipelli, A.R., and Tilford, C.R. (1990) Metrological characteristics of a group of quadrupole partial pressure analyzers. J. Vac. Sci. Technol. A, 8, 3838.
- 80 Blanchard, W.R., Mccarthy, P.J., Dylla, H.F., La Marche, H., and Simpkins, J.E. (1986) Long-term changes in the sensitivity of quadrupole mass spectrometers. J. Vac. Sci. Technol. A, 4,
- 81 Bennet, J.R. and Elsey, R.J. (1993) Anomalies in the measurement of the residual gases in a large UHV system using a quadrupole mass analyser. Vacuum, 44, 647.
- 82 Fremerey, J.K. (1994) Qualifying an XHV system by the residual gas accumulation method. J. Vac. Soc. Jpn., 37, 718.
- 83 Nakashima, Y., Tsuchiya, K., Ohtoshi, K., Shoji, M., Yatsu, K., and Tamano, T. (1995) Evaluation of the background signal for mass spectrometry under high ambient hydrogen pressures in plasma discharges. J. Vac. Sci. Technol. A, 13, 2470.
- 84 Santeler, D.J. (1987) Gas dynamics in residual gas analyzer calibration. J. Vac. Sci. Technol. A, 5, 129.
- 85 Basford, J.A., Boeckmann, N.D., Ellefson, R.E., Filipelli, A.R., Holkeboer, D.H., Lieszkovsky, L., and Stupak, C.M. (1993) Recommended practice for the calibration of mass spectrometers. J. Vac. Sci. Technol. A. 11, A22.
- 86 Ellefson, R.E., Cain, D., and Lindsay, C.N. (1987) Calibration of mass spectrometers for quantitative gas mixture analysis. J. Vac. Sci. Technol. A, 5, 129.
- 87 Solomon, G.M. (1986) Standardization and temperature correction of calibrated leaks. J. Vac. Sci. Technol. A, 4, 327 - 333.
- 88 Ehrlich, C.D. (1986) A note on flow rate and leak rate units. J. Vac. Sci. Technol. A, 4, 2384.
- 89 Ehrlich, C.D. and Basford, J.A. (1992) Recommended practices for the calibration and use of leaks. J. Vac. Sci. Technol. A, 10, 1-17.

- 90 Thornberg, S.M. (1988) Stepped linear piston displacement fundamental leak calibration system. J. Vac. Sci. Technol. A, 6, 2522.
- 91 Fischer, E. and Mommsen, H. (1967) Monte Carlo computations on molecular flow in pumping speed test domes. Vacuum, 17, 309.
- 92 Feng, Yu-guo. and Xu, TingWie. (1980) The appropriate test domes for pumping speed measurement. Vacuum, **30**, 377.
- 93 Hablanian, M.H. (1987) Recommended procedure for measuring pumping speeds. J. Vac. Sci. Technol. A, 5, 2552.
- 94 Kendall, B.R.F. (1989) Recommended procedures for measuring the performance

- of positive-displacement mechanical vacuum pumps. J. Vac. Sci. Technol. A, 7, 2403-2407.
- 95 Grosse, G., Jitschin, W., and Wandrey, D. (1990) Procedures for measuring pumping speeds. Vacuum, 41, 2120.
- 96 Terado, K., Okano, T., and Tuzi, Y. (1989) Conductance modulation method for the measurement of the pumping speed and outgassing rate of pumps in ultrahigh vacuum. J. Vac. Sci. Technol. A, 7, 2397.
- 97 Tuzi, Y., Okano, T., and Terano, K. (1990) Accuracy of the conductance modulation method for the measurement of pumping speed and outgassing rate. Vacuum, 41, 2004.

16

Materials

Dr. Karl Jousten

Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

This chapter explains the requirements for materials of vacuum technology, the materials for particular applications, as well as processing and manufacturing techniques.

16.1 Requirements and Overview of Materials

Depending on the particular type of usage, requirements for materials of vacuum technology are manifold. For example, materials used to build vacuum vessels must be absolutely gas tight, that is, ambient air must be held back from the vacuum. Vessel material has to be strong enough not to deform when evacuating produces a pressure differential of 100 kPa (1 bar), corresponding to the force exerted by a weight of 10 t per 1 m². At the same time, gas emissions from the material into the vacuum are an issue. The desired vacuum pressure determines the maximum tolerable gas emission rates. For built-in components surrounded by vacuum, in contrast, tightness is of less or no concern at all. Rotor blades of turbomolecular pumps are subject to extreme accelerations. Significant warming of such blades due to gas friction is not tolerable. Materials for helium bath cryostats have to withstand temperature variations down to only a few K; materials for high-bakeable ultrahigh-vacuum equipment are exposed to up to 450 °C.

Thus, the main requirements specific to materials for vacuum technology are as follows:

- sufficient mechanical strength;
- corrosion resistance;
- high gas tightness (leak rates, e.g., $< 10^{-9}$ mbar ℓ s⁻¹ (10^{-7} Pa ℓ s⁻¹));
- low intrinsic vapor pressure;

Table 16.1 Overview of commonly used materials for vessels and seals as well as corresponding vacuum ranges.

Pressure and vacuum ranges		Application examples	Materials	
10 ² mbar (10 ⁴ Pa)	Rough or low vacuum	Drying	Structural steel	
$1\text{mbar}(10^2\text{Pa})$		Degassing	Stainless steel	
		Distillation	Ceramics	
			Aluminum	
			Elastomer seals	
$1\mathrm{mbar}(10^2\mathrm{Pa})$	Medium or fine vacuum	Vacuum process technology	Aluminum	
10 ⁻³ mbar (10 ⁻¹ Pa)			Stainless steel	
			Ceramics	
			Elastomer seals	
$10^{-3} \text{ mbar } (10^{-1} \text{ Pa})$	High vacuum	Coating technology	Stainless steel	
10 ⁻⁷ mbar (10 ⁻⁵ Pa)		Molecular beam epitaxy	Aluminum	
			Al ₂ O ₃ ceramic	
			Elastomer seals	
			Tantalum, molybdenum	
10 ⁻⁸ mbar (10 ⁻⁶ Pa)	Ultrahigh vacuum	Deposition of high- purity coatings	Stainless steel	
		Materials analysis	Aluminum	
$10^{-12} \text{ mbar } (10^{-10} \text{ Pa})$		Accelerator technology	Al ₂ O ₃ ceramic	
			Copper seals	
			Special seals	
			Gold, silver, tantalum, molybdenum	

- low foreign-gas content;
- favorable degassing properties;
- high melting and boiling points;
- clean surfaces;
- adopted thermal expansion behavior;
- high thermal fatigue resistance.

Additionally, materials must provide sufficiently high chemical resistance against gases and vapors developing in the processes.

Requirements in terms of gas emissions and tightness increase when approaching the ultrahigh-vacuum range. Table 16.1 provides an overview of materials used in vacuum technology.

16.2 Materials for Vacuum Technology [1]

16.2.1

Metals

Metals and metal alloys in the solid state are made up of atoms arranged regularly in characteristic lattice structures. Only in special cases does this lattice structure fill the complete metal part. Components are rather composed of numerous small crystal grains, so-called crystallites with sizes ranging from several cubic micrometers to several cubic millimeters. Such materials are thus termed polycrystalline materials.

The grains are separated by grain boundaries with at times considerably different chemical and mechanical properties than the crystallites. The shape and size of the crystallites determine mechanical properties. They are influenced by the processing techniques chosen for manufacturing, also including heat treatments. Fine, stretched crystallites make the material hard and brittle. The strength of the material is high. Such materials are cold-rolled, work-hardened, and cold-worked.

When material is heated beyond its recrystallization temperature T_R for longer periods, large crystallites form, making the material soft and ductile (annealing). Usually, this recrystallization temperature is given by $T_R \approx 0.4 T_E$, with $T_{\rm E}$ denoting the melting point, both temperatures given in K.

Table 16.2 lists processing techniques as well as their corresponding advantages and disadvantages for applications in vacuum technology.

16.2.1.1 The Most Important Metals and Metal Alloys

Mild Steel/General Structural Steel

General structural steels are known as S23511 and S355.21 Usually, they are contaminated with carbon, phosphorous, and sulfur. They are used for high-vacuum applications down to 10^{-4} Pa (10^{-6} mbar) if corrosion resistance is not required. Due to processing technology, such steels emit CO gas continuously.

Grades must be picked carefully. Generally, an operating experiment is conducted in order to investigate the behavior of selected sample components in terms of weld ability and helium tightness. Transfer of working techniques from boiler making and tank construction to vacuum-tank construction is limited. For example, multiple welding beads and machining down into segregation zones (agglomerations of undesired chemical elements in the metal) should be avoided. Tools should be held separate from tools used for stainless-steel processing in order to prevent microscopic residue of mild steel from contaminating stainlesssteel parts. Selection has to take into account material characteristics, for example, in terms of weld ability.

¹⁾ DIN EN 10028-2.

²⁾ DIN EN 10025-2.

Table 16.2 Processing technologies for metals and corresponding characteristics.

Processing technologies	Characteristics		
Casting of molten metal into casting molds	Series production possible		
	Volume shrinkage during solidification		
	Possible inclusions and shrinkage cavities (exhalation sources, leakage)		
	Applicable to components in fore- and low-vacuum range or for low requirements		
Drawing and rolling	Compression of materials		
	Applicable to any components down to UHV		
Special melting (electroslag remelting, ESR) and additional forging or rolling	Production of high-purity and homogeneous materials		
	Additional materials compression during forging		
	Expensive		
	Required for special applications		
Vacuum melting	Production of materials with low contents of foreign gases		
	Expensive		
	Required for special materials (e.g., titanium)		
Extruding	Special geometries producible, not feasible with any other technique		
	Restricted to appropriate materials (aluminum)		

Starting materials include sheets, rods, and tubes. Cast parts (e.g., pumps and valve housings) are restricted to low- and fine-vacuum applications.

Mild steel is also of interest for certain special applications, for example, for providing a particular magnetic shielding effect. If such applications call for corrosion resistance, the steel surface has to be coated.

Stainless Steel

Stainless steel is common to vacuum technology. The reason is that stainlesssteel surfaces, due to their surface microstructure, show sufficient passivation, and thus, are protected adequately against corrosion during baking and vacuum processing. At the same time, the group of stainless-steel materials also includes grades providing sufficient strength for flange joints exposed to baking procedures (typically, 200-300 °C in the UHV range).

A wide range of stainless steels is available (see also Table A.24): nonstabilized (high-carbon) grades such as 1.4301 (DIN, AISI: 304), low-carbon grades such as 1.4306 (304 L), as well as stabilized grades such as 1.4541 and 1.4571 (316 Ti). The latter include alloying elements that react with carbon during welding and thus prevent a drop in corrosion resistance usually caused by this element. For welding, stainless steels with low carbon contents or stabilized grades are recommended. Analyzer and accelerator technologies mostly use low-magnetic permeability stainless steels such as 1.4429 (316LN), 1.4404 (316 L), and 1.4435 (316 L).

Steel - Special Alloys

Special alloys are used for glass-metal joints or for joints to premetallized ceramics. This is because the differences in coefficients of thermal expansion between commercial stainless steel and glass or ceramics are so high that manufactured joints, if produced at all, suffer cracks or offsets.

In terms of thermal expansion, employed special alloys are adopted to glass and ceramics. Such alloys mainly comprise binary iron-nickel alloys or ternary iron-nickel-cobalt alloys known as Kovar, Fernico, Nilo-K, or Vacon. Note that these materials are highly magnetic, which might cause problems in certain physical applications.

Titanium

The material is highly gas binding. Therefore, titanium is produced in vacuummelting processes. Commercial grades show low carbon contents and high ductility. Titanium's density lies between those of iron and aluminum. Above approximately 150 °C, titanium reacts easily with atmospheric gases such as oxygen and nitrogen. This behavior is disadvantageous in certain situations (e.g., when welding without shielding). On the other hand, it is just this reactivity that paves a way for applications of titanium in vacuum technology: titanium hydride is used for premetallization of ceramics providing reliable bonds between the ceramic and subsequent metal coatings (e.g., nickel). Applying pure titanium coatings onto appropriate ceramics allows active brazing without separate metallization procedures. Titanium is also used as getter material in pump technology (Sections 11.3.3 and 11.4). Titanium evaporates very easily even below its melting point of 1670 °C; thus, it is evaporated from solid phase in titanium evaporation pumps (evaporation at approximately 1350 °C), heated by an electrical current passing through. Typical coatings reach thicknesses of 5 µm after one hour of evaporation. Ion getter pumps feature titanium plates as cathodes sputtered by ion bombardment.

Aluminum

Aluminum is used mainly in fore and high vacua. Small-flange components are often made from aluminum bulk material. Furthermore, aluminum is used for seals operating at temperatures up to 100 °C. Alloys used should be free of elements with high vapor pressures such as lead or zinc. Mostly, alloys are composed of aluminum and silicon. Vapor pressure is low and reaches approximately 10⁻⁶ Pa (10⁻⁸ mbar) at the melting point of 660 °C. Disadvantages include high thermal expansion, a porous aluminum-oxide surface layer, and high thermal conduction. This causes particular susceptibility to porosity and cracking in welding processes, and high distortion due to preheating required for welding. New applications utilize aluminum-UHV components [2]. Such applications call for special processing techniques in order to overcome some of the negative properties of aluminum, for example, surface porosity or low hardness.

Copper

Predominant features of copper include high thermal and electrical conductivity. These properties are required in cryotechnology and for heat sinks (e.g., radiation absorbers in particle accelerators) as well as for high-performance electrical conductors. Such components are used frequently in many fields of vacuum technology.

A negative property of copper is associated with hydrogen embrittlement. Hydrogen appearing in heat treatments reacts chemically with the oxygen contained in copper. Developing water fissures the copper's microstructure. Thus, UHV applications use so-called OFHC copper (oxygen-free, high conductivity) or copper with reduced oxygen content.

Materials selection should pay particular attention to avoid alloy constituents with high vapor pressure. Alloys such as tombac and brass contain tin or zinc as alloying elements that easily evaporate at higher temperatures under vacuum and thus degrade the vacuum.

On a large scale, UHV technology uses metal seals made of copper, bakeable to approximately 450 °C. Grades used here provide controllable hardness (CF-flange seals) and good cold weld ability (for COF-flange seals). Filler metals for high-temperature brazing include Cu/Au and Cu/Au/Pd alloys.

Gold and Silver

Vapor pressures of gold and silver are sufficiently low for the metals to be used in ultrahigh vacuum.

Gold is used for seals in appropriately manufactured flange connections, for sealing valve faces in all-metal valves, or as coatings for high-performance electrical conductors and connectors. Similar to copper, gold is used in hightemperature brazing filler metals made of Cu/Au and Cu/Au/Pd alloys. Silverplated stainless-steel screws yield prolonged service lives and show reduced tendency to cold welding. Also the copper CF-sealings are silver plated for some applications.

Indium

Indium is used for metal seals in high-vacuum applications, as solid sealing material between flanges, and to provide high heat transmission between different materials.

Its vapor pressure is low even though the melting point is quite low (156 °C). Special applications (e.g., cryotechnology) call for the material in cases where other sealing materials are inappropriate. Indium is soft, and thus, required sealing forces are low.

Materials for Special Vacuum Processes

Many vacuum processes require materials that differ from standard materials. Examples of such processes include measuring processes, heating, and cooling, all of which depend on special materials and components.

Special materials such as tungsten, tantalum, or molybdenum are employed for measuring systems, in high-temperature applications (shielding, evaporator crucibles), in accelerator technology (radiation absorbers), as well as in lowtemperature applications.

Heating Equipment, Radiation Shields, Insulators

Heaters are made of refractory materials such as tungsten or tantalum. Additionally, graphite is used. Thermal shields are made of, for example, molybdenum sheet metal.

Brittleness is typical for refractory elements. Thus, they are not welded or brazed but mostly riveted. Heating elements, once heated to high temperatures, should not be exposed to vibrations, let alone moved.

Insulators are made of appropriate high-melting oxides (e.g., alumina). For lower temperature applications, Macor is also used.

Materials for Heat Sinks

Copper, preferably OFHC-grade, is usually the material of choice for heat sinks. High-temperature applications, for example, synchrotron-radiation technology, employ compound materials made of copper with traces of aluminum oxide. Due to the high melting point of the alumina, the material withstands temperatures of up to approximately 900 °C.

Materials for Radiation and Magnetic Shielding

Lead bulk material is usually used for radiation shielding. However, lead is less eligible for vacuum, and thus, lead shielding is hermetically sealed in appropriate stainless-steel coverings. Synchrotron-radiation equipment uses compound materials of refractories for shielding purposes.

Magnetic shielding utilizes alloys of certain stoichiometric nickel and iron compounds in conjunction with other elements (Mu-metal).

They are used for shielding plates (arranged inside vacuum vessels as single or multilayers) and for vacuum vessels where the outer housing (main and side tubes, upper and lower lids) is made of the shielding material. Flanges are made of stainless steel.

Such components, after finishing, require a special heat treatment under vacuum for adjusting the desired shielding properties. For shielding, magnetism in the components must be high. After heat treatment, deformation must be kept low because it would impair the adjusted properties.

16.2.2

Technical Glass

16.2.2.1 Basics

Until well into the 1960s, vacuum equipment, for high and ultrahigh vacuum in particular, was manufactured almost exclusively from glass. Small mercury and oil diffusion pumps, valves, cold traps, and vacuum gauges, for example, McLeod vacuum gauges and U-tube manometers, were made of glass too. Glass was also used for electrical insulations in high-vacuum leadthroughs. Today, glass is found only in certain special applications of vacuum technology, for example, for optical reasons (windows, beam input to vacuum), for vacuum deposition equipment (glass bells), permeation elements in test leaks, or certain process-technological applications. Chemical stability of glass is excellent because only very few organic or inorganic compounds act corrosively to glass.

In contrast to metals, the structure of glass is amorphous, producing completely different physical behavior. Constituents form an irregular, nonsymmetric network. Glass does not show a fixed melting point, and thus, is regarded more as an undercooled liquid. Glass is brittle and easily cracks when exposed to shocks. However, in contrast to common belief, it is highly elastic. The resistance to pressure of glass is similar to that of metals whereas tensile strength is more than one decade lower (approximately $10^7 \, \text{N m}^{-2}$). *Young's* modulus of elasticity is in the same range as for metals, $E = 5 \times 10^{10} - 10^{11} \, \text{N m}^{-2}$.

Even at room temperature, the viscosity of glass is low enough to be perceived. A glass rod, leaning against the corner of a room, becomes bent after a few weeks. If glass is heated, viscosity initially drops slowly, then quicker, and finally again slower. This behavior is illustrated in Figures 16.1 and 16.2.

With rising temperature, glass undergoes a transition from brittle to tough. The corresponding temperature range is termed transition temperature range. Start and end points of the transition range are characterized by the lower $T_{\rm lo}$ and upper $T_{\rm hi}$ cooling temperatures, respectively. Glass can be used continuously

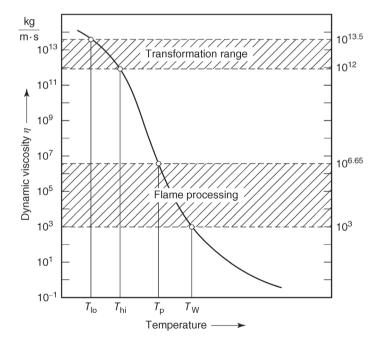


Figure 16.1 Dynamic viscosity η of glass versus temperature (schematic). Lower cooling temperature T_{lo} , upper cooling temperature T_{hi} , softening temperature T_{W} , and processing temperature T_{p} (1 kg m⁻¹ s⁻¹ = 1 Pas = 10 Poise (P)).

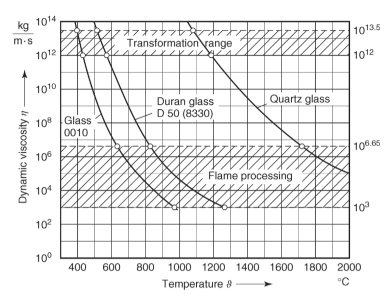


Figure 16.2 Dynamic viscosities η of soft glass 0010 (*Corning Glass Works*, Corning, NY, USA), hard glass DURAN 50 (*Jenaer Glaswerke Schott*), and quartz glass (*Heraeus Quarzglas*) versus temperature.

at temperatures up to approximately 30 K below the lower cooling temperature. If heated beyond the so-called devitrification temperature for longer periods or repeatedly, glass becomes nontransparent. It crystallizes and devitrifies, suffering volume changes, cracks, or offsets.

The thin water film on glass surfaces has to be removed by heating (flame-scarfing or baking in a furnace) for high-vacuum applications. Here, it can be necessary to heat close to the softening temperature of the glass.

Knowing the specific properties of glass is very important because a wide selection of glasses is available and their individual behaviors (e.g., maximum tolerable temperatures) differ significantly.

16.2.2.2 Properties of Important Glasses

Glasses are categorized according to their mean coefficients of linear thermal expansion α as follows:

- soft glass $(60 \times 10^{-7} 120 \times 10^{-7} \text{ K}^{-1})$;
- hard glass ($< 50 \times 10^{-7} \text{ K}^{-1}$);
- quartz glass ($\approx 5 \times 10^{-7} \text{ K}^{-1}$).

Additionally, we differentiate

- · sintered glass;
- · crystallized glass.

They differ significantly in terms of physical and chemical properties, thus qualifying for appropriate applications:

Soft Glass

SiO₂ content in soft glass is comparably low, whereas the content of alkali elements is relatively high. Typical amounts of substance are, for example, 65-70% SiO₂, 2.5–15% Na₂O, and 5–15% CaO. Typical soft glasses include Ruhrglas AG AR Glass in Germany and Dow Corning Glass Works 0010 in the United States, with coefficients of thermal expansion of 90×10^{-7} K⁻¹. Usually, the lower cooling temperature is in the range 370-450 °C.

Soft glasses are used mainly because of their good electrical insulation properties and low permeabilities for H₂ and He. The permeability of lead glass, a particular type of soft glass, is only 1/10 000 of the permeability of common hard glasses.

Soft glasses show certain characteristics limiting utilization and processing. Their resistance to thermal shocks is poor, that is, cooling and heating have to be performed very slowly. Thus, after processing, intense stress relief in the transformation range is required, followed by slow cooling. Figure 16.2 shows that processing temperatures for soft glasses are lower than for any other type of glass and that soft glasses show viscosities at low temperatures for which hard glasses would require heating to higher temperatures.

Thus, their use in high-vacuum technology is limited because softening occurs even at low temperatures. Therefore, they cannot be baked out completely.

Hard Glasses

SiO₂ contents in hard glasses usually exceed 70%. Hardness generally increases with boron content; thus, so-called borosilicate glass is hard glass. Other brand names are Pyrex and Duran. Their coefficients of expansion are approximately $30 \times 10^{-7} \text{ K}^{-1}$. One glass of this group, Corning 7056, is particularly popular in vacuum technology.

Their resistance to thermal shocks is good because lower cooling temperatures are above 500 °C. Likewise, the transition range is at relatively high temperature (Figure 16.2). Thus, hard glasses are better suitable than soft glasses for the highand ultrahigh-vacuum range.

Quartz Glass

Quartz glass is pure SiO₂ glass. Its resistance to thermal shocks is very high because the coefficient of expansion is only $5 \times 10^{-7} \text{ K}^{-1}$.

Temperatures for processing and transition ranges are the highest of all known glasses (see Figure 16.2). It devitrifies at temperatures above 1100 °C and can be used continuously up to approximately 1050 °C. Thus, quartz glass is used in vacuum furnaces (muffle furnaces) for high temperatures.

Comparably low amounts of alkali vapors lower the temperature for devitrification considerably. Therefore, quartz glass has to be processed carefully (separate from other types of glass).

Due to its optical properties, quartz glass is also used for sophisticated windows (see Section 17.4.5).

Sintered Glass and Crystallized Glass

Sintered glass is finely ground glass, sintered after being pressed, and thus, is easily formed into any desired shape. Appropriate powder selection allows adjusting the coefficient of thermal expansion to any value obtainable with glass.

Crystallized glasses are produced by adding crystallizing agents. After shaping at relatively low temperatures as with standard glasses, further heating initiates crystallization. Materials produced show beneficial thermal (and usually also electrical) behavior compared to the starting material.

16.2.3

Ceramic Materials

16.2.3.1 Basics

Ceramics are nonmetal inorganic materials (more than 30% crystalline) used in vacuum technology mainly for insulation purposes. This section provides an overview of their most prominent properties.

Burning of ceramics for vacuum applications must provide high tightness. Appropriate burning temperatures and durations have to be chosen. Often, the exhalation material is dried, preferably under vacuum. Ceramics are frequently glazed to provide proper tightness. Hard glazing (enamel) is applied to dried or preburnt pieces, and soft glazing to ready-burnt parts. Shrinking should always be expected when baking ceramics. Usually, ready-baked ceramic parts are cut with diamond saws and finished by grinding. Utilizing ceramics requires employment of appropriate custom molds.

16.2.3.2 Properties of Important Ceramics

Ceramics are categorized according to their chemical compositions:

- silicate ceramics;
- pure-oxide ceramics;
- special ceramics, for example, glass ceramics.

Their properties differ, thus determining individual fields of applications in vacuum technology.

Silicate ceramics are mixtures of inorganic crystalline substances with glassy fluxing or binding agents. Typical are porcelain (mass fractions approximately 6-8% undissolved quartz fragments, approximately 26% mullite, and approximately 66% feldspar silicic acid), steatite, forrestite, and ceramics with high Al_2O_3 contents.

They do not show fixed melting points and most of their properties are similar to those of hard glasses. However, their thermal resistance is higher and mechanical as well as electrical properties are better. Baking temperatures are between 1300 °C and 1500 °C. Nearly any porcelain, steatite, and forrestite can be used up to approximately 1000 °C, and high-Al₂O₃ ceramics up to approximately 1350 °C.

Pure-oxide ceramics are made of crystalline aluminum-, beryllium-, zirconium-, or magnesium oxides. Their melting point is well defined. Baking temperatures are in the range of 1800-2000 °C. Vacuum technology mostly uses alumina (Al₂O₃). In contrast to silicate ceramics, they are more temperature resistant (maximum working temperature 1800 °C) and show better resistance to thermal shocks. Additionally, their mechanical and electrical properties are better. Zirconium oxide is used in cases where required thermal resistance is even higher than provided by Al₂O₃. Due to its toxicity, beryllium oxide is used far less, and magnesium oxide is limited to lower temperature applications.

Glass ceramics are crystalline ceramics modified from the standard amorphous condition using crystallizing agents and heat treatment. The baked material can be machined with standard tools [3]. It is available under the brand name Macor and Corning no. 9658. The mean coefficient of linear thermal expansion is equal to expansion coefficients of standard technical soft glasses (see Table 16.3). Maximum operating temperature is 1000 °C. Macor can be metallized and is used for electrical leadthroughs, for electrical insulators, and vacuum-tight ceramic molds. Applications should be picked carefully because Macor softens at high temperatures.

16.2.3.3 Ceramics in Vacuum Technology

Vacuum technology predominantly employs ceramics based mainly on Al₂O₃ (amount of substance > 92%). This is due to the above-mentioned advantages of Al₂O₃ ceramics and their low costs. Al₂O₃ ceramics are used for heavy-duty thermal and electrical components (transmitting tubes), for vacuum vessels in accelerators, and electrical leadthroughs. Al2O3 ceramics are particularly widespread as electrical insulators for high-bakeable electrical leadthroughs (more than 300 °C).

Sapphire is a special kind of monocrystalline Al₂O₃. Its radiation transparency (for infrared and UV) is high. Sapphire disks are used for bakeable windows.

16.2.3.4 Ceramic/Metal Joining Technologies

Special techniques and processes are necessary for joining ceramics and metals because ceramics usually are not weldable or brazeable. Ceramic surfaces at the joining interface require pretreatment by metallization or activation. Otherwise, reliable joints, generally created by brazing, could not be produced.

Techniques for manufacturing vacuum-tight, permanent ceramic/metal joints are known as molybdenum manganese and titanium hydride processes, covered in Chapter 17.

16.2.3.5 Zeolite

Zeolites are highly porous alumina silicates with alkali metals. Industrial applications use them as drying agents. Artificially produced zeolites show homogeneous porosity. In vacuum technology, they are utilized for adsorbing water, oil vapors, and a number of other gases. Adsorption is increased by liquid-nitrogen

Table 16.3 Glasses common to vacuum technology.

Glass type	Density gcm ⁻³	$\alpha^{a)}$ (20°C – 300°C) 10 ⁻⁷ K ⁻¹	°C	θ _g c) °C	θ _a d) °C	θ _e e)	θ _{σ100} f) °C	Comments
Soft glass								
Schott AR 8418	2.52	90	520	320	708	198		
Hard glass								
Schott 8250	2.28	50	185	495	380	715	384	VACON 70, KOVAR and Mo-sealing glass
Schott 8482	2.34	49		493	393	738	416	
Schott 8487	2.25	39		523	420	760	275	for fusing with W wires
Schott 8330 DURAN	2.23	32.5	250	530	430	815	248	typical W sealing glass

a) mean coefficient of thermal expansion in indicated range

b) thermal shock endurance

c) transformation temperature

d) maximum heating temperature e) softening point f) temperature at which the glass' electrical conductivity is σ = 10-6 Ω $^{-1}$ m $^{-1}$

cooling (see Section 11.2). Vice versa, zeolites can be regenerated nearly completely by heating.

16.2.4

Plastics

16.2.4.1 **Basics**

Vacuum technology employs plastics mainly for sealing purposes. Plastics are made of more or less cross-linked organic macromolecules. Specific properties are created by additives, for example, softening agents for reducing chemical linkage forces between molecules, or fillers.

16.2.4.2 Properties of Major Plastics

The main groups of plastics used in vacuum technology are elastomers, thermoplastics, and duroplastics. They differ in terms of structures and properties (Table 16.4).

The filamentous macromolecules in *elastomers* are cross-linked with more or less strong chemical bonds. Showing rubbery behavior, the materials are easily deformed by tension as well as compression and return to their initial shape after load relief thereby maintaining their total volume. This behavior makes the materials particularly suitable for seals. Typical examples are Perbunan (NBR), Viton, Kalrez, Chemraz, and silicone. Natural rubber is rarely used due to its sensitivity toward oil and wear [4,5].

Thermoplastics feature filamentous macromolecules as well. At room temperature, the materials show more or less moderate hardness. They can be soft gluey when dominated by short macromolecules, or when containing long macromolecules, tough-hard, elastic, or rubbery, depending on type and amount of softeners. At higher temperatures (approximately 100 °C), the mobility of macromolecules increases, making the materials ductile.

Duroplastics feature a network of spatially cross-linked macromolecules. They retain their hardness even at higher temperatures. Vacuum technology uses epoxy resins (e.g., araldite), thermally irreversible fillers made from resin solution and hardening agent. The constituents are stored separately and mixed just before joining. Cold-hardening grades operate at temperatures up to 100 °C, and warm-hardening types at up to approximately 180 °C. Their adhesion to metals, glass, and ceramics is high. Carefully processed parts (fully hardened, dried) show gas emissions comparable to those of Teflon or Hostaflon.

Sections 16.3.3 and 16.3.2.3 cover gas emissions and gas permeability of rubbery materials (elastomers).

16.2.5

Vacuum Greases

Often, vacuum technology calls for sealing or lubricating greases (Table 16.5). Requirements on such greases include low vapor pressures and high viscosities.

Table 16.4 Important elastomers (*) and thermoplastics (**) for vacuum technology.

Symbol (DIN ISO 1629)	Explanation	Brand name	Characteristics	Applications in vacuum technology
1. Rubber NR	Natural rubber			Flexible connections, thick-wall tubing
2. PVC soft**	High-polymer polyvinyl chloride		Chemically resistant, inexpensive	Flexible connections, vacuum tubing for low and fine vacuum
3. NBR	Butadiene acrylonitrile rubber	Perbunan N	Resistant towards oil, favorable mechanical properties, inexpensive	Sealing material for general high- vacuum use, seal- ing rings, tempera- ture range -25 to +80 °C
4. CR*	Polychlorobutadiene	Neoprene	See 3	See 3
5. MVQ	Silicon rubber	Siloprene	Constantly temperature stable up to 150 °C	Used rarely today for temperatures up to 150 °C
6. FPM*	Vinylidene fluoride-hexa- fluoropropene copolymers (fluorocaoutchouc)	Viton	Temperature stable between -10 °C and +200 °C	Used in high-vacuum technology mainly for seals between bakeable flanges $(\theta < 200^{\circ}\text{C})$
7. PTFE*	Polytetrafluoroethylene	Teflon	Temperature stable up to 300 °C, very low gassing (unloaded)	Sheathing material for high and ultra- high vacuum
		Hostaflon		
		Halon		
8. CFM (PCTFE)*	Polychlorotrifluoroethylene	Kel F	Similar to 7, but employ- able down to very low temperatures	Valve seals in valves for cryogenics
9. —	Copolymer of tetrafluoro- ethylene and perfluoro- methyl vinyl ether	Kalrez [4]	High temperature stability (300 °C) and chemically resistant	O-ring seals, tubing, plates

Table 16.5 Vacuum-suitable greases and oils as well as their characteristics.

Туре	Applications	Vapor pressure at 20°C (Pa)	Dripping tem- perature (°C)	Maximum operating temperature (°C)
Ramsay grease, thick Ramsay grease, soft	Lubrication of ground- in connections and taps for > 1 Pa	10-2	> 56	30
Gleitlen	Lubrication of stirring shafts	10^{-2}	> 50	30
Lithelen	Lubrication of ground- in connections and taps below 1 Pa	10^{-8}	> 210	150
Silicone high- vacuum grease	Lubrication of ground- in connections and taps below 1 Pa	n.s.	> 200 polymerization	180
Dynafat	Lubrication of sealing rings	n.s.	148	110
Diffelen	Rotary transmission leadthroughs	2×10^{-7}	n.s.	120
Ultratherm 2000	Lubrication	< 10 ⁻⁷	n.s.	At least 150

See also Section 16.6.4.

However, the requirements, in part, contradict each other, and thus, gas emissions of greases can be substantial. Grease should therefore always be applied as thin as possible. Applications in vacuum technology are limited, particularly in high and ultrahigh vacua.

16.2.6 Oils

Oil is used as a lubricating and sealing agent in roughing pumps, sliding vane rotary pumps, in particular, and as working fluid in oil diffusion pumps (see Tables A.17 and A.18). In roughing pumps, oil vapor pressure at room temperature is of less concern than viscosity, which should be of appropriate values (see Section 7.5.1.2). Typical oils for such applications are hydrocarbon-based mineral oils. Special applications (oxygen handling) call for special oils. Oils for diffusion pumps must be picked carefully (see Section 9.4). Here, silicone oils are used almost exclusively. Main requirements on such oils are low vapor pressures at room temperature, high thermal stability, high resistance to atmospheric oxygen when hot, high molecular weight, no combustibility, and no toxicity. Otherwise, pumping speeds, ultimate vacua, and environmental impact would be impaired.

Table 16.6 Important vacuum coolants and associated applications.

Coolants	Applications			
Water	Large mechanical pumps			
	Diffusion pumps			
	Certain types of oil catchers, cooling temperature approximately 10 $^{\circ}\mathrm{C}$			
	Single- or two-stage refrigerators, cooling temperature approximately -20 to $-40^{\circ}\mathrm{C}$			
Liquid	Sorption pumps			
nitrogen	Cold shields			
Liquid helium	Bath cryostats			
	Evaporator cryostats			
	Cryopumps			

16.2.7

Coolants

Coolants used are water (in diffusion pumps, turbopumps, oil separators), liquid nitrogen (sorption pumps, traps), and liquid helium (cryopumps, cryostats, superconductors) (Table 16.6). Using liquid air is extremely dangerous and should be avoided. Nitrogen has a boiling point of $-196\,^{\circ}\text{C} = 77\,\text{K}$, whereas the boiling point of oxygen is -183 °C = 90 K. Thus, from a reservoir of fresh liquid air, nitrogen would evaporate rapidly. The mixture would subsequently become enriched with oxygen and present a serious threat.

16.3 Gas Permeability and Gas Emissions of Materials

16.3.1

Fundamentals

The obtainable ultimate pressure determines performance data of a vacuum vessel. This pressure is generally controlled by the gas tightness of the housing, by gas emissions from the inside of the chamber, and gas permeability through possible pores, capillaries, or other leaks. Knowing these mechanisms and the specific behavior of materials is very important for designing and operating vacuum systems.

16.3.2

Gas Permeability

Gas permeation includes the individual processes of adsorption at wall surfaces on the atmospheric side, diffusion through walls, and desorption from the wall

surfaces on the vacuum side. Think of gas permeability of a wall as a conductance of permeation. It can be derived in a model based on the assumption of an ohmic dependence from wall surface area A and wall thickness d.

The considered wall shall be of thickness d, surface area A, and temperature T. Pressures on both sides of the wall are p_1 and p_2 . The gas permeability of such a wall is given by the permeation flux q, which is proportional to the wall surface area A, and as a fair approximation, inversely proportional to the wall thickness d. Furthermore, q follows a complicated dependence on wall temperature T as well as on the pressures p_1 and p_2 to both sides of the wall. Thus,

$$q = f(T)f(p_1, p_2)\frac{A}{d}. (16.1)$$

Permeation flux q can be given as a pV flow q_{pV} in Pa ℓ s⁻¹. Often, it refers to a surface area $A = 1 \text{ m}^2$, thickness d = 1 mm, and a pressure gradient of 101.3 kPa. This specific gas permeation flux (conductance of permeation) is given by

$$\overline{q}_{\text{perm}} = q_{pV} \frac{d}{A} \cdot \frac{1}{p_1 - p_2}.$$
(16.2)

The following sections describe the conductance of permeation for metals, glasses, ceramics, and plastics in more detail.

16.3.2.1 Gas Permeability of Metals

Figure 16.3 shows permeabilities $\overline{q}_{\mathrm{perm}}$ for selected gas—metal combinations versus temperature (see also Figure A.12).

Obviously, gas permeability is very low at room temperature. Hydrogen permeates through metals more rapidly than any other gas species, and palladium is the metal with the highest gas permeability for hydrogen. Due to this, heated palladium tubes can serve to feed pure hydrogen to a vacuum vessel. Fusion technology intensely investigates the permeation of deuterium and tritium.

Stainless steel shows lower gas permeability than standard steel. Nevertheless, even for iron, hydrogen permeation at room temperature is usually insignificant due to the low hydrogen content of atmospheric air. As for any other diffusion process, hydrogen diffusion and thus hydrogen-permeation rates rise with temperature [6].

Among metals, silver represents a special case with high permeability for oxygen.

16.3.2.2 Gas Permeability of Glasses and Ceramics

Figure 16.4 shows permeabilities \overline{q}_{perm} of selected glass–glass systems. Unfortunately, no comparable experimental data are available for ceramics.

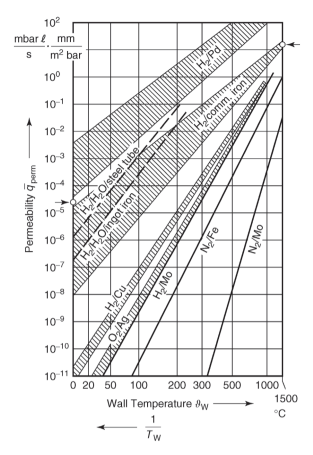


Figure 16.3 Permeabilities for selected gas species through metals versus wall temperature.

The gas permeability of quartz glass is approximated according to Example 16.1:

Example 16.1

Estimation of helium permeation of quartz glass under atmospheric ambient conditions. For nonmetals, the gas permeation flux q_{pV} is proportional to the pressure gradient $p_1 - p_2$. Ohm's law applies: $q_{pV} = \overline{q}_{perm}(T)(p_1 - p_2)A/d$ (Eq. (16.1)).

Thus, the gas permeation flux can be calculated using \dot{q}_{perm} values from Figure 16.4. Helium partial pressure in air is 0.53 Pa. At 20 °C, according to Figure 16.4, \dot{q}_{perm} amounts to

$$\overline{q}_{\rm perm} = 5.3 \times 10^{-5} \, {\rm mbar} \, \ell \, {\rm s}^{-1} \, {\rm mm \, m}^{-2} \, {\rm bar}^{-1} \ (5.3 \times 10^{-8} \, {\rm Pa} \, \ell \, {\rm s}^{-1} \, {\rm mm \, m}^{-2} \, {\rm Pa}^{-1}).$$

For a thickness of the glass of 2 mm and a surface area of 0.5 m², leakage flow through the wall is

$$\begin{split} q_{pV} &= 5 \times 10^{-5} \; \mathrm{mbar} \, \ell \; \mathrm{s}^{-1} \; \mathrm{mm} \; \mathrm{m}^{-2} \; \mathrm{bar}^{-1} \cdot 5.3 \times 10^{-3} \\ &\quad \mathrm{mbar} \cdot 0.5 \; \mathrm{m}^2 / 2 \; \mathrm{mm} \\ &= 6.6 \times 10^{-11} \; \mathrm{mbar} \, \ell \; \mathrm{s}^{-1} (6.6 \times 10^{-9} \; \mathrm{Pa} \, \ell \; \mathrm{s}^{-1}). \end{split}$$

Even at room temperature, quartz glass shows comparably high permeability for helium and a smaller permeability for hydrogen. However, permeability for helium drops when SiO2 contents are reduced. Thus, soft glasses feature lower helium permeability than hard glasses.

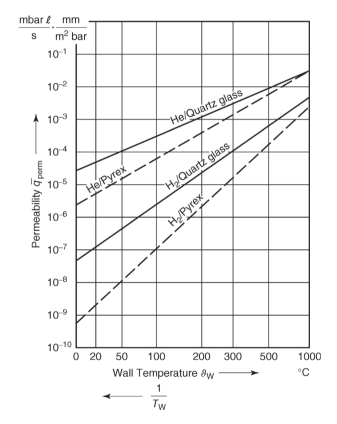


Figure 16.4 Permeabilities $\overline{q}_{\text{perm}}$ of selected glasses and gas species versus wall temperature. Scaling of the abscissa corresponds to outlining $1/T_{\rm W}$ from right to left.

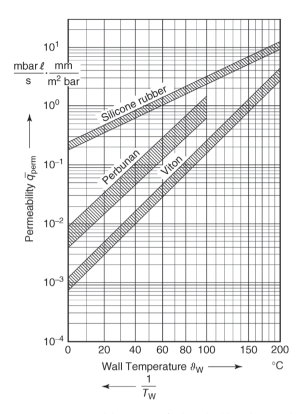


Figure 16.5 Permeabilities $\overline{q}_{\text{perm}}$ of selected rubber-elastic sealing materials versus temperature. Scaling of the abscissa corresponds to outlining $1/T_{\rm W}$ from right to left. Notice by W. Beckmann, Carl Freudenberg, Weinheim, Bergstrasse, Germany.

16.3.2.3 Gas Permeability of Plastics [7]

Gas permeabilities $\overline{q}_{\text{perm}}$ of plastics are shown in Figure 16.5. Permeability for atmospheric air increases rapidly with air humidity. Figure 16.5 gives values for 60% humidity. Listed data should be treated as approximate values because gas permeabilities of plastics depend greatly on the type of plastic and its additives.

Gas permeabilities of toroidal sealing rings are estimated using Eq. (16.2):

Example 16.2

Estimating permeability of toroidal sealing rings. The length of the considered toroidal sealing is 1 m, with a diameter of 5 mm. When pressed, the height of the seal is 2.5 mm with a width of 8 mm. Thus, the permeation surface is

$$A = 1 \text{ m} \cdot 2.5 \times 10^{-3} \text{ m} = 2.5 \times 10^{-3} \text{ m}^2.$$

According to Figure 16.5, permeability at 20 °C amounts to

$$\begin{split} \overline{q}_{\rm perm} &= 2 \times 10^{-2} \; {\rm mbar} \, \ell \; {\rm s}^{-1} \; {\rm mm} \; {\rm m}^{-2} \; {\rm bar}^{-1} \\ &\qquad \qquad (2 \times 10^{-5} \; {\rm Pa} \, \ell \; {\rm s}^{-1} \; {\rm mm} \; {\rm m}^{-2} \; {\rm Pa}^{-1}) \, {\rm for} \; {\rm Perbunan} \\ \overline{q}_{\rm perm} &= 3 \times 10^{-3} \; {\rm mbar} \, \ell \; {\rm s}^{-1} \; {\rm mm} \; {\rm m}^{-2} \; {\rm bar}^{-1} \\ &\qquad \qquad (3 \times 10^{-6} \; {\rm Pa} \, \ell \; {\rm s}^{-1} \; {\rm mm} \; {\rm m}^{-2} \; {\rm Pa}^{-1}) \, {\rm for} \; {\rm Viton}, \\ \overline{q}_{\rm perm} &= 3 \times 10^{-1} \; {\rm mbar} \, \ell \; {\rm s}^{-1} \; {\rm mm} \; {\rm m}^{-2} \; {\rm bar}^{-1} \\ &\qquad \qquad (3 \times 10^{-4} \; {\rm Pa} \, \ell \; {\rm s}^{-1} \; {\rm mm} \; {\rm m}^{-2} \; {\rm Pa}^{-1}) \, {\rm for} \; {\rm silicone} \; {\rm rubber}. \end{split}$$

Equation (16.2) yields the following permeation rates for the selected sealing materials:

$$\begin{split} q_{pV} &= 2\times 10^{-2}~\text{mbar}\,\ell~\text{s}^{-1}~\text{mm}~\text{m}^{-2}~\text{bar}^{-1}\cdot 1~\text{bar}\cdot 2.5\times 10^{-3}~\text{m}^2/8~\text{mm} \\ &= 6.25\times 10^{-6}~\text{mbar}\,\ell~\text{s}^{-1}~(6.25\times 10^{-4}~\text{Pa}\,\ell~\text{s}^{-1})~\text{for Perbunan}, \\ q_{pV} &= 9.4\times 10^{-7}~\text{mbar}\,\ell~\text{s}^{-1}~(9.4\times 10^{-5}~\text{Pa}\,\ell~\text{s}^{-1})~\text{for Viton}, \\ q_{pV} &= 9.4\times 10^{-5}~\text{mbar}\,\ell~\text{s}^{-1}(9.4\times 10^{-3}~\text{Pa}\,\ell~\text{s}^{-1})~\text{for silicone rubber}. \end{split}$$

Helium permeates plastics such as Viton, Perbunan, and silicone rubber, preferably used as O-ring seals. This should be taken into account during helium leak testing. Corresponding regions have to be shielded from too high helium loads. Here, Viton provides the lowest and silicon rubber shows the highest permeability. Perbunan is in a middle position. Epoxy resins feature water-vapor permeabilities at room temperature of approximately $q_{\rm perm} = 10^{-1} \, {\rm mbar} \, \ell \, {\rm s}^{-1} \, {\rm mm} \, {\rm m}^{-2} \, {\rm bar}^{-1} \, (10^{-4} \, {\rm Pa} \, \ell \, {\rm s}^{-1} \, {\rm mm} \, {\rm m}^{-2} \, {\rm Pa}^{-1})$. Thus, such materials always represent a source of (undesired) water vapor in vacuum processes.

16.3.3

Gas Emissions

16.3.3.1 **Basics**

Under vacuum, any solid releases gas [8,9]. Gas emissions are made up of three sources: intrinsic vapor pressure (saturation vapor pressure) of the solid, desorption of surface-adsorbed gas, and diffusion of dissolved or absorbed gas from the bulk to the surface of the solid. These physical processes are covered in Chapter 6. Here, we will focus on numeric-value equations and actual values for practical use.

16.3.3.2 Saturation Vapor Pressure (See Also Section 3.5.1)

Any substance in a sealed system is in thermodynamic equilibrium with its vapor phase. The resulting vapor pressure p_s is termed intrinsic vapor pressure or saturation vapor pressure. It rises with temperature. A numeric-value equation for the saturation pressure can be derived from the following concepts:

The surface area-related particle-evaporation rate is given by Eq. (3.129); the surface area-related mass-evaporation rate is given by Eq. (3.130). The

corresponding numeric-value equation reads

$$(j_m)_{\text{max}} = 0.438 \,\sigma_{\text{c}} \sqrt{\frac{M_{\text{t}}}{T}} p \, \text{s}(T).$$
 (16.3)

Here, j_m is given in kg m⁻² s⁻¹, T in K, and p_s in mbar (10² Pa). M_r is the molecular weight, and σ_c is the condensation probability (compare Section 3.5.2 and Table A.11).

Thus, Eq. (16.3) yields the saturation vapor pressure

$$p_{\rm s}(T) = 2.28 \frac{1}{\sigma} \sqrt{\frac{T}{M} (j_m)_{\rm max}}.$$
 (16.4)

A saturation vapor-pressure curve, particularly important for vacuum technology, describes the relationship between saturation vapor pressure and the temperature. We will now discuss the saturation vapor pressures of metals and nonmetals in more detail.

Saturation Vapor Pressure of Metals

Figure 20.3 plots vapor-pressure curves of selected metals. Table 16.7 lists melting points and the temperatures where the corresponding chemical elements show a vapor pressure of 1.33×10^{-3} Pa. Species marked with * are molten when they reach 1.33×10^{-3} Pa.

Table 16.7 Melting points θ_E and temperatures $\theta(10^{-5})$ at vapor pressure $p_v = 10^{-5}$ Torr = 1.33×10^{-5} mbar for selected materials relevant to vacuum technology.

Material	∂ _E (°C)	ϑ(10 ⁻⁵) (°C)
Mercury ^{a)}	-39	-15
Cadmium	321	145
Zinc	419	210
Magnesium	650	280
Lead ^{a)}	327	500
Indium ^{a)}	156	650
Silver	961	785
Tin ^{a)}	232	850
Aluminum ^{a)}	659	895
Copper	1083	955
Steel (iron)	1535	1045
Nickel	1453	1050
Gold ^{a)}	1063	1090
Titanium	1690	1327
Molybdenum	1622	1930
Tantalum	2696	2400
Tungsten	3382	2570

a) For these materials, $\vartheta(10^{-5}) > \vartheta_{\rm E}$.

Apparently, there is no clear relationship between the melting point and the saturation vapor pressure of a substance. Considering saturation vapor pressures and melting points of metals, we find that cadmium, zinc, and magnesium feature relatively high vapor pressures, impairing the vacuum, whereas tin has a very low melting point. Thus, these metals are inappropriate for vacuum technology. The same is true for alloys of these metal species.

Saturation Vapor Pressures of Nonmetals

Figures 20.4–20.7 show vapor-pressure curves of selected nonmetals. Nonmetals such as glasses, ceramics, greases, and oils are chemically complex substances made up of molecular groups with various molar masses. Thus, vapor pressures can only be defined for certain molecular groups. These define the vapor pressure depending on the current temperature. It appears that rubber-elastic materials, greases, resins, and oils show vapor pressures that have to be considered when calculating the ultimate pressure of a vacuum system. In contrast, vapor pressures of technical glasses, ceramics, and quartz are so low that they are usually irrelevant to vacuum technology.

16.3.3.3 Surface Desorption, Gas Diffusion from Bulk Material, Reference Values for Gas Emissions [8,9,11]

An adsorbed layer (see Chapter 6) covers any surface that has been exposed to ambient air. The layer mainly consists of molecules of water, nitrogen, and oxygen. In order to produce deep vacua, this layer has to be removed as thoroughly as possible. A monolayer of adsorbed gas corresponds to a gas amount of approximately 40 Pa ℓ m⁻². For an inner surface area of 1 m² and volume of 200 ℓ , this gas amount of only a single monolayer would create a pressure of 0.2 Pa when released from the surface. Usually, several monolayers are produced during venting.

Solid surfaces are far from homogeneous. Even the most highly polished surfaces feature a true surface area always larger than calculated geometrically. Even low levels of contaminants increase gas emissions to the vacuum. Surfaces exposed to humid air for longer periods release particularly high amounts of gas. Often, up to 90% of the desorbed gas is water vapor (see Section 6.2.3).

Raising the temperature considerably accelerates desorption of particles bound to the surfaces of metals, glasses, and ceramics. As a rule of thumb, gas emissions after baking drop by an order of 10 per 100 degrees of temperature increase during baking. Gas emissions from metal, glass, or ceramic surfaces can be estimated as approximately 10^{-8} – 10^{-10} mbar ℓ s $^{-1}$ m $^{-2}$ $(10^{-6}-10^{-8} \text{ Pa } \ell \text{ s}^{-1} \text{ m}^{-2})$ after baking at 450 °C (for several hours).

16.3.4

Gas Diffusion from Bulk Material

From a technical point of view, foreign-gas contents are more difficult to remove than adsorption layers. Complete removal is possible only under vacuum and at

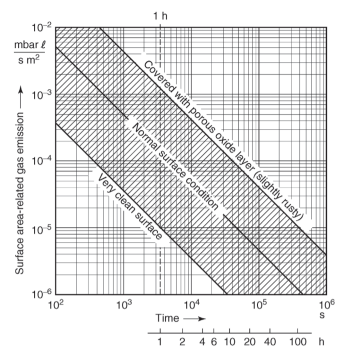


Figure 16.6 Approximate reference values of surface area-related gas emissions of metal surfaces versus time at room temperature.

very high temperatures, a problem when handling elastomers that prohibit exposition to high temperatures. The diffusion coefficient D determines the gas emissions originating in the deep bulk of the material. The gradient with time of the outgassing flux (see also Sections 6.2.3 and 6.3) follows several principles, not investigated in detail. As an empirically confirmed result for metals, we can assume an outgassing behavior with time that is proportional to t^{-1} (Figure 16.6).

In plastics, the drop is approximately proportional to $t^{-1/2}$, but also proportional to e^{-t/t_0} , where $t_0 \propto d^2 D$, if d is the thickness of the material in the direction of diffusion. Figure 16.7 shows both behaviors. Perbunan and Viton follow the $t^{-1/2}$ law for longer periods, whereas silicone rubber shows exponential behavior.

Figure 20.11 comprises a larger number of outgassing curves. Apparently, outgassing is a complex process, depending on many prerequisites and parameters. Thus, the uncertainties expressed in Figure 20.11 should be taken into account when quantitative assessments of outgassing processes and periods are made.

The principles described in Section 6.3 seem most suitable to characterize the behavior of plastics. Figure 16.8 shows a calculated outgassing curve (I) and a calculated permeation curve (II).

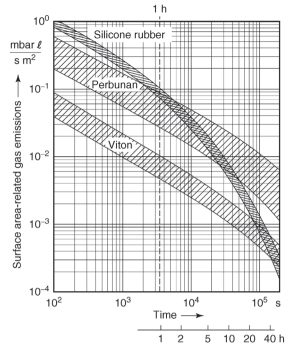


Figure 16.7 Surface area-related gas emissions of common rubber-elastic materials versus time [10].

Calculation I is based on the model of a plastic material (data for Viton) with thickness d, saturated with air and enclosed by vacuum on both sides. Then, until approximately

$$t \,\mathrm{W} \approx 0.05 \frac{d^2}{D}$$
 (SI units), (16.5)

the outgassing rate drops with the square root of time t. Gas emission rates increase for $t > t_{\rm W}$ (i.e., now the exponential behavior dominates), see Figure 16.8, curve I.

Calculation II (Figure 16.8) assumes a completely degassed plastic material with one side in physical contact with atmospheric air, and the other exposed to vacuum. At the time

$$t > t_{\rm K} \approx 0.3 \frac{d^2}{D}$$
 (SI units), (16.6)

a constant permeation flux develops through the plastic material according to Eq. (16.2).

For a plastic material, after having absorbed or dissolved gas, exposed to the atmosphere on one side and to vacuum on the other (sealing), gas emissions

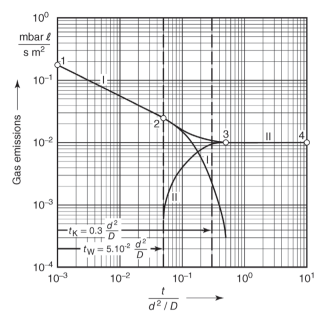


Figure 16.8 Calculated gas emissions of plastic (d in m and D in m² s⁻¹). Curve I: outgassing, curve II: permeation.

drop according to curve 1-2 (Figure 16.8). Along curve 2-3, permeation becomes noticeable. On curve 3-4, the permeation flux determines the gas flux into the vacuum.

Some plastics containing substances with high vapor pressures (e.g., softeners) are inapplicable to vacuum systems. Outgassing would continue until the entire amount of constituents with high vapor pressures is released. Depletion of softeners would thus change the properties of the plastic materials.

Figure 16.7 shows outgassing of common rubber grades versus time. Specimens used for these investigations were rods with 13 mm diameter and 6 mm height, previously conditioned under atmospheric air for a week at 20°C and 60% humidity.

At 20 °C, values for the diffusion coefficients D in the most important plastics are in the following ranges:

$$\begin{array}{lll} \text{Perbunan N} & D & = (1.7-2.5)\times 10^{-11} \text{ m}^2 \text{ s}^{-1}, \\ \text{Viton} & D & = (3.8-4.2)\times 10^{-12} \text{ m}^2 \text{ s}^{-1}, \\ \text{silicone rubber} & D = (5.9-8.1)\times 10^{-10} \text{ m}^2 \text{ s}^{-1}. \end{array}$$

For polycarbonate, $D = (1.7 \dots 2.4) \times 10^{-12} \text{ m}^2 \text{ s}^{-1}$ was published [12]. Thus, according to Eq. (16.5) and for a rubber thickness of d = 5 mm, times $t_{\rm W}$ up until outgassing follows $t^{-1/2}$,

```
Perbunan N t_W = (14 - 20)h,
Viton t_{W} = (83 - 91)h,
silicone rubber t_{\rm W} = (0.4 - 0.6)h.
```

In the case of sealing, the time at which constant permeation gas flux develops under otherwise equal conditions is given by Eq. (16.6):

```
Perbunan N t_{K} = (80 - 120)h,
Viton t_{\rm K}
                 = (500 - 550)h
silicone rubber t_K = (2.6 - 3.5)h.
```

Even though the obtained diffusion coefficients (representing average values for a number of specimens) should be treated as approximations only, it does seem apparent that, for example, the high diffusion coefficient for nitrogen in silicone rubber allows relatively short outgassing periods. Thus, gas permeability of silicone rubber is high. Outgassing of plastic materials can be accelerated considerably by thermal treatment because diffusion coefficients increase rapidly with temperature. Of course, baking under vacuum is beneficial but heat treatment in a hot-air chamber already provides a drop of initial outgassing rates by an order of 10, due to water-vapor desorption. Obviously, baking temperatures must be limited to the maximum tolerable operating temperature of the considered plastic material.

16.3.5

Reference Values for Total Gas Emission Rates [8,11]

Under vacuum, adsorbed and occluded gases are initially released rapidly, and subsequently, emission rates drop with time. For comparison, the total gas emission rate is given after a 10-hour exposition to vacuum at room temperature. The following reference values apply (see also Table A.5 and Figure A.11):

Metals
$$10^{-7} \text{ Pa } \ell \text{ s}^{-1} \text{ cm}^{-2} (10^{-9} \text{ mbar } \ell \text{ s}^{-1} \text{ cm}^{-2}),$$

elastomers $10^{-5} \text{ Pa } \ell \text{ s}^{-1} \text{ cm}^{-2} (10^{-7} \text{ mbar } \ell \text{ s}^{-1} \text{ cm}^{-2}).$

Teflon and Hostaflon are exceptions, with outgassing rates approximately between the two ranges given.

Due to the relatively high gas emissions of elastomers, they are used as little as possible in vacuum systems, and never as construction materials (insulations). In cases where there is no alternative material available, for example, in sealing of flange joints or valve seats, special plastics are used. Even such special plastics are then arranged so that only the least possible surface area of such substances is exposed to the vacuum.

References

1 Espe, W. (1961) Werkstoffkunde der Hochvakuumtechnik, VEB Deutscher Verlag der Wissenschaften, Vol. 1 (Metalle und metallisch leitende Werkstoffe) 1959;

- Vol. 2 (Silikatwerkstoffe) 1960; Vol. 3 (Hilfswerkstoffe).
- 2 Ishimaru, H. (1990) Developments and applications for all-aluminum ally vacuum systems. MRS Bull., XV (7), 23-31.
- 3 Mog, D. (1976) Machinable glass-ceramic: a new material for vacuum equipment. Vacuum, 26, 25 ff.
- 4 Chernatony, L. (1977) Recent advances in elastomer technology for UHV applications. Vacuum, 27, 605-609.
- 5 Peacock, R.M. (1980) Practical selection of elastomer materials. J. Vac. Sci. Technol., 17, 330-336.
- 6 Esser, H.-G. (1984) DEUPERM eine anlage zur messung von festkörper-diffusion und permeation für wasserstoffisotope. Vak.-Techn., 33 (8), 226-237.
- 7 Beckmann, W. and Seider, J.H. (1967) Gasdurchlässigkeit von gummielastischen Werkstoffen für Stickstoff. Kolloid Zsch. und Zsch. f. Polymere, 220, 97-107.
- 8 Elsey, R.J. (1975) Outgassing of vacuum materials. Vacuum, 25, 299 ff. and 347 ff. (detailed Tables).
- 9 Edelmann, C. (1989) Gasabgabe von Festkörpern im vakuum. Vak.-Techn., 38, 223 - 243.
- 10 Beckmann, W. (1963) Gasabgabe von gummielastischen werkstoffen im vakuum. Vacuum, 13, 349-357 (in German and English).

- 11 Erikson, E.D. et al. (1984) Vacuum outgassing of various Materials. J. Vac. Sci. Technol., A2, 206-210.
- 12 Kwon, J.-S., Jung, H., Yeo, I.S., and Song, T.H. (2011) Outgassing characteristics of a polycarbonate core material for vacuum insulation panel. Vacuum, 85, 839-846.

Further Reading

- Caporicio, A. and Steenrod, R.A. Jr. (1978) Properties and use of perfluorethers for vacuum applications. J. Vac. Sci. Technol., 152, 775 ff.
- Cherepnin (1978) Treatment of Materials for Use in High Vacuum, 3rd edn, Holon/ Israel, Ordentlich, 192 p.
- Dauphin, J. (1982) Materials in space: working in vacuum, 32, 669-673.
- Diels, K. and Jaeckel, R. (1962) Leybold Vakuum Taschenbuch, 2nd edn, Springer,
- Halliday, B.S. (1987) An introduction to materials for use in vacuum. Vacuum, 37, 583-585.
- Perkins, W.A. (1973) Permeation and outgassing of vacuum materials. J. Vac. Sci. Technol., 10, 543-556.
- Waldschmidt, E. (1954) Gasabgabe und Gasdurchlässigkeit metallischer Vakuumbaustoffe. Metall, 8, 749-758.

17

Vacuum Components, Seals, and Joints

Dr. Ute Bergner

VACOM GmbH, In den Brückenäckern 3, 07751, Großlöbichau, Germany

In this chapter, the reader learns about vacuum components, joining and sealing technologies, dimensioning of vacuum vessels, and cleanliness needed for vacuum components.

17.1 Introduction

This chapter provides an overview of the construction elements and components used in vacuum technology and their connections that are required to monitor, control, and measure processes within the vacuum. Because the pressure range includes low atmospheric pressure of more than 15 decades, varied requirements arise for the components and joining technologies as well as for their cleanliness. With this, the total volume of a vacuum system, the outgassing behavior of the materials exposed to vacuum, the occurring leak rates, and the choice of the pumps are decisive for the achievable ultimate pressure. The users of vacuum technology who would like to construct and operate, for example, a coating plant, an apparatus for surface analysis, or an accelerator line must consider a large number of aspects in the planning phase. In addition to the desired pressure range as well as the operation and maximum/minimum temperatures (e.g., through baking out/cooling) of the equipment, the materials used in the vacuum and media (solids, liquids, gases) must be sufficient for the cleanliness requirements of the process and must not release disruptive elements.

In Section 17.2, the basic rules regarding clean handling with vacuum components are presented. Contaminations are classed and explained for the purposes of vacuum technology and the methods and processes with which the cleanliness of the components can be made measurable and comparable are likewise indicated.

Special requirements are made for connecting and joining technologies in the vacuum technology. Alongside the mechanical stability of the connection, a leak tightness according to the requirements of the final application must also be

ensured. In Section 17.3, the established joining technologies (welding, brazing, and bonding) and detachable connection types (KF, ISO, CF, QCF, and others) are discussed and their advantages, limits, and applicability are illuminated.

In addition to pumps and measuring systems, a vacuum apparatus primarily consists of a chamber as well as selected connecting components and transfer components that are addressed in Section 17.4. The chamber represents the core element for processes in vacuum. When designing and constructing vacuum chambers, it is always to be guestioned which tolerances are truly necessary and make sense in order to achieve an optimal cost performance ratio. When the vacuum is reached in the chamber, it is necessary to communicate from the vacuum to the outside world, so that processes can be run and monitored. In addition to this, the transition between atmosphere and vacuum should take place in a defined manner, unaltered through external influences. If it is necessary to initiate and control mechanical movements in the vacuum from outside, mechanical feedthroughs come into use. If gases, electrical current, and light are to be brought into the vacuum and to be led out again, then valves as well as electrical and optical feedthroughs, among other things, are used. These component types as well as the subject of heat supply and dissipation in vacuum chambers are also discussed in detail in Section 17.4.

17.2 Vacuum Hygiene

Vacuum is often applied to run processes under undisturbed or well-defined ambient conditions. To this end, the evacuated system must be protected against penetrating gases (through atmospheric gases or from leaky liquid lines) and be free from or low in contaminations. Penetrating gases and contaminations of the parts exposed to vacuum can disturb the processes running in the vacuum and limit the achievable ultimate pressure. However, the ultimate cleanliness necessary in the vacuum, the vacuum hygiene, always depends on the specific requirements of the respective process.

Figure 17.1 provides an overview of possible disturbing factors from the process surroundings affecting the vacuum.

The penetration of atmospheric gases can occur through three possible ways: leakage, permeation, and virtual leakage. Atmospheric leakages primarily originate in junctions while joining or mounting. Other leaks can originate from leakages in liquid cooling lines that are laid in the vacuum. Due to their signature in the residual gas spectrum, the latter can be clearly distinguished from the atmospheric leaks. The permeation (see Section 6.3) of ambient gases through the wall of vacuum recipients can be minimized through the use of suitable materials and the observance of minimum wall thicknesses (to a negligible level compared with desorption effects). The so-called virtual leaks refer to mistakenly trapped gas volumes in the vacuum. They originate, for example, while mounting a screw in a blind tapped hole under atmospheric pressure. Then, in the vacuum, the trapped gas flows along the thread guide very slowly and continuously into the chamber. Over a (too) long period, the trapped

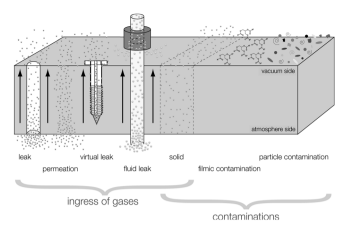


Figure 17.1 Sources of disturbances in vacuum.

volume generates a continuous gas source with the properties of an atmospheric leak. The gas can be easily evacuated with a ventilation bore or through slots in the screw thread and thus eliminate this apparent leak.

In addition to the penetrating gases, the other disturbance sources can be divided into three classes: particles, molecular contaminations (also referred to as filmic contaminations), and contaminations from the solid-state material. Table 17.1 provides an overview of the type of these contaminations, possible sources, and established detection methods.

Particles prevent the sealing effect of the surfaces to be fitted or mounted and can disturb processes in the vacuum, while, for example, they are deposited on the surface that is to be examined or situated in the process. A particularly great challenge exists in avoiding particle generation during movements in the vacuum or its border areas due to the limited application of lubricants. Thus, under many conditions, grease is suitable as a lubricant for the avoidance of galling ("seizing") of the screw connections and friction losses. In contrast, low outgassing lubricants are to be used in vacuum or modern materials are to be worked with that show neither galling nor abrasion [1]. Also, traditional materials, such as stainless steels, can be prepared through surface finishing for application without lubricant in vacuum. An example of this is the Kolsterising[®] of stainless steel surfaces [2].

In addition to particles, we also consider substances as contaminants that are located in or on the surface of solids exposed to vacuum and are found in a solid or liquid phase under atmospheric conditions at room temperature. Then, in vacuum, these are able to desorb into the gas phase with or without a catalytic effect (process-induced outgassing), particularly when the vapor pressure of the substance falls short of the chamber pressure. A process-induced outgassing is present, for example, in particle accelerators where highly energetic particles hit the chamber wall and cause desorption of the atoms or molecules of the surface and/or from the solid state (bulk) into the gas phase.

In contrast to the contaminations that are present on solid-state surfaces or in the solids themselves, *molecular contaminations* or *filmic contaminations* refer

Table 17.1 Overview of the sources and detection methods for contaminations.

	Particles	Filmic contaminations	Solid-state contaminations
Examples	Turnings Dust Micro- and nanodust	Water/lime Oils Cooling lubricants Proteins Skin oils	Gases bound in solid states (H ₂) Alloying elements (Si, Pb, Zn, SnN, etc.) Plasticizers Oxide and corrosion layers
Sources	Machining Milling, turning, sawing Polishing, grinding, sandblasting Exposure in atmosphere Floating particles in the cleaning liquid Lint from cloths Particle abrasion during handling Packaging material	Lubricants from machining Milling, turning, sawing Handling of the components without gloves, skin oils, sneezing Minerals and hydrocarbons in the cleaning liquid Residuals from cleaning agents	Welding (annealing colors) Natural affinity of elements of the PSE Contaminations in production process Alloy admixtures
Detection methods Qualitatively	Examination under black light Examination under white light (oblique light and side light)	Residual gas analysis in the vacuum with QMS TDS	Residual gas anal- ysis in the vac- uum with QMS
Quantitatively	Particle counter (optical) SEM AFM	Residual gas analysis in the vacuum with QMS Fluorescence measurement Outgassing accumula- tion method	Residual gas analysis in the vac- uum with QMS XPS, SIMS, AES TML

to such contaminations that additionally arise on the surface, for example, through manufacturing, storage, or handling steps. According to the required vacuum regime or planned process, very different requirements for the material composition and the cleanliness of the components as well as for the use of the components exist. Thus, for example, grease can have a sealing effect in rough vacuum (above the vapor pressure of the substance), but can be disruptive in high vacuum and at lower pressures (below the vapor pressure of the substance) as a result of the then volatile hydrocarbons. Therefore, in HV and UHV and especially XHV, it is essential to work with gloves in order to avoid contamination of the components with skin oils. The semiconductor industry, particularly EUV lithography, has extremely high cleanliness requirements in high vacuum. For these high cleanliness requirements in moderate pressure ranges, the term ultraclean vacuum (UCV) has been established.

The different processes that can be applied for the detection of contaminations are shown in Table 17.1. A series of different qualitative and quantitative valuation and measuring methods can be applied according to component material and component geometry and depending on the contaminations to be examined and the required detection boundaries. From the basic principle, the contaminations are first solved from the surface or from the volume (e.g., upon transition into the gas phase with TML, RGA, and TDS, or by bombardment with highly energetic (charged) particles or radiation with SIMS, XPS, and AES) in order to be analyzed afterward (e.g., through a quadrupole mass spectrometer (OMS) with RGA and SIMS or with an electron detector with XPS and AES). However, the comparability of the results of different processes is only restrictedly provided due to the type of the extraction and the specific analysis method. For detection of molecular or filmic contaminations (water, hydrocarbons, cooling lubricants, etc.), RGA has been established as a gas chromatographic method [3]. In doing so, the substances outgassing from components and chambers in vacuum are qualitatively and quantitatively examined with QMS. This process is very well suited to the requirements of different vacuum applications because during the examination the components are exposed to conditions similar to those in the later operation of the vacuum system.

In addition to the detection of contaminations or the examination of the component cleanliness, the cleaning of vacuum components and the preservation of the state of cleanliness through suitable packaging have become an important part of vacuum technology [3]. Table 17.2 provides an overview of the material and surface treatment methods established in vacuum technology. Detailed information regarding the mechanisms of action and the attainable cleaning success of the individual processes can be taken from the literature shown within the table. While plain surfaces of stainless steel are relatively easy to clean, there is an increase in the degree of difficulty and effort of cleaning with the complexity of the geometry (e.g., blind tapped holes, diaphragm bellows, thin bent pipes, spark-eroded slits) and the variety of materials (aluminum, copper, elastomers). Building upon the information presented, appropriate cleanliness requirement profiles and cleaning processes can be determined for specific vacuum applications. Detailed information regarding the established cleaning processes for UHV/XHV components is found, for example, in the publications of the particle accelerators developed by DESY [4], FAIR [5], and CERN [6].

17.3 Joining Technologies in Vacuum Technology

Vacuum technology applications need an evacuable and hermetically sealed system that consists of vessels (Section 17.4.1), pipelines (Section 17.4.1), and other

Wet-chemical cleaning	Electrochemical cleaning	Mechanical cleaning	Dry cleaning
Aqueous solutions Water Deionized water Surfactant admixtures (basic, acidic, neutral) Solvents Isopropanol Acetone Supportable by Ultrasound [7] High and low	Electrolytic polishing [8] Plasma polishing [9,10] Pickling	Wiping with cloth [11] ■ Dry ■ Moist ■ Solvent-saturated Polishing ^{a)} [12] Grinding Glass bead blasting ^{a)} [13] CO ₂ -snow blasting [14] Blowing off, for exam-	Baking out under atmospheric pressure Baking out under vacuum [15] Low H ₂ annealing ("vacuum firing") [16] Plasma cleaning [17] Ozone cleaning [18]
pressure		ple, with dry nitrogen	

Table 17.2 Overview of cleaning process established in vacuum technology.

components (Sections 17.4.2–17.4.6). In addition to the selection of suitable materials (Chapter 16) and the correct material and surface treatment (Section 17.2), the joining technologies applied are crucial for the quality of the generated vacuum.

In mechanical and plant engineering, connections are classified according to different criteria. Static and dynamic connections are classed according to the function, that is, fixed or moveable in different degrees of freedom. If the connections are classed according to manufacturing aspects, a distinction is made between detachable and permanent connections (see Figure 17.2).

In addition to the mechanical stability, a vacuum system requires leak tightness and low outgassing in all junctions that correspond to the targeted ultimate pressure. Restrictions for the possible joining processes arise from this.

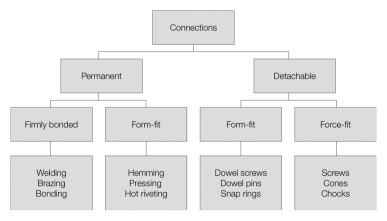


Figure 17.2 System of the most commonly used connections in mechanical and plant engineering.

a) Not applicable in clean room.

17.3.1

Permanent Connections

The German standard DIN 8593-0 defines permanent connection as a "connection made by joining that can only be removed again under acceptance of a damage or destruction of the joined parts" [19].

In vacuum technology, the material-bonded connections that are realized through welding, brazing, or bonding primarily count among the relevant permanent connections.

Application and ultimate pressure determine which joining processes are used for vacuum components. Determining criteria for the selection are

- materials to be joined (Table 17.3);
- outgassing behavior of the joining agents (e.g., adhesive/welding filler/solder);
- gas impermeability of the joining agents (permeation, pore formation, packing density, particle generation);
- temperature resistance of the joining agents (melting ranges, aging);
- mechanical stability of the component (mechanical stress through joining, mechanical loads during the operation);
- accessibility of the joint in the joining process;
- particle generation potential of the joining materials.

The joining technologies such as brazing, welding, and bonding have application preferences depending on the materials of the joining partners that should be considered in their selection (see Table 17.3).

Table 17.3	Applicable	ioinina te	chnoloaies f	for different	material pairings.

Material pairing	Brazing	Welding	Bonding	Examples of use
Fe metals/Fe metals	0	+	0	Components made of corrosion- resistant stainless steel
NF metals/NF metals	+	0	0	Aluminum components, copper tubes in stainless steel flanges
NF metals/Fe metals	+	0	+	Aluminum components with flanges made of corrosion-resistant stainless steel
Metal/glass	+	_	+	Viewports
Metal/ceramic	+	_	+	Insulators, electrical feedthroughs
Glass/ceramic	_	_	+	Optical fiber feedthroughs
Plastic/metal	-	-	+	Insulating constructions, attached elastomer seals

⁽⁺⁾ Frequently applied joining process; (0) possible/uncommon joining process; (-) unsuitable/ impossible/possible joining method with high technological effort.

17.3.1.1 Welding

Welding is a "joining process with which two or more parts are connected and a continuity of the materials of the connecting parts is produced under application of heat or force or both and with or without welding fillers" [20]. In vacuum technology, common welding techniques are tungsten inert gas welding, microplasma welding, electron beam welding, laser beam welding, friction welding, explosion welding, and diffusion bonding whose functionality and areas of application in vacuum technology are briefly explained in the following [21,22]:

- Tungsten inert gas welding (briefly, TIG welding or argon arc welding). With this fusion welding technique, an electric arc is used as heat source between the material and the nonmelting tungsten electrode. TIG welding can be used very flexibly and with low technical expenditure by use of a manual torch guidance.
- Microplasma welding. Microplasma welding is used similar to TIG welding; nevertheless, it uses an electric arc of higher energy density with a plasma gas nozzle constricting the electric arc. This also burns stably with low current and allows the welding of low wall thicknesses up to the foil range (0.01–1 mm).
- Electron beam welding. With this process, electrons within an evacuated chamber are accelerated through high voltage on the material surface where they convert their kinetic energy with the impact into heat. If the electron beam from the vacuum chamber is led out through pressure stages and directed upon the workpiece under free atmosphere, we talk of nonvacuum electron beam welding. The quick, complicated pendulum movements that allow the parallel welding of several connections can be generated by deflecting coils.
- Laser beam welding. Laser beam welding uses monochromatic, coherent, and strongly focused light of high energy density as a heat source that enables the welding of electrically conductive and nonconductive materials. Very narrow and deep welded joints (deep weld effect) can be generated without contact. Due to the very local, punctual (concentrated) energy entry, laser beam welding is particularly suitable for components that only admit a low heat input as well as for connections demanding a very low distortion of the component geometry.
- Friction welding. The workpiece surfaces to be joined are moved relative to each other under application of force and are connected through the originating frictional heat and distortion heat without additives. With a fully automated process sequence, short weld times and a high positional accuracy of the joining partners are possible. Friction stir welding is a special method with which the joining partners are plasticized by a rotating tool.
- Explosion welding. The workpieces come into contact with each other and are joined (cladded) under use of an explosion-induced shock wave collision. With explosion welding, no structural changes appear beyond the direct joining zone.
- Diffusion bonding. Diffusion bonding is a special joining technique that is based on the principle of solid diffusion and grain boundary migration. The materials are physically joined with each other under heat and application of pressure without a fusion process. Diffusion bonding can be a recommendable connecting process with new construction materials, intermetallic connections, complex ceramic-based materials, and refractory superalloys.

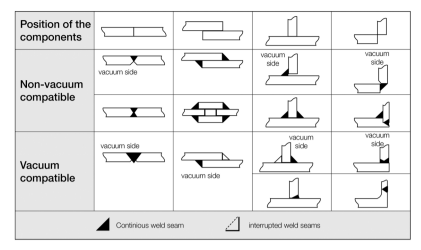


Figure 17.3 Examples of welded joints compatible with vacuum. (According to Ref. [22].)

Certain construction rules are to be followed in the vacuum-compatible welding of components (Figure 17.3) [22]. If the component geometry permits, the seams are always to be welded vacuum-sided to avoid outgassing sources (virtual leaks) and collection points for contaminations. One-ply welded seams are to be preferred over multilayer seams in order to avoid inter-run fusion and decrease the risk of inclusions. Different wall thicknesses and component geometries of the joining partners can complicate welding due to the divergent fusion behavior.

The heat can be stopped through the insertion of a groove in the joint, and therefore, the heat flux can be controlled (heat insulating groove, Figure 17.4).

The generated bridge can additionally lead to the distortion and thus diminish the stresses of the welding joint. To increase the mechanical loading capacity of the connection, supporting seams on the atmosphere side are used in addition to the vacuum-sided sealing seams. These must be welded with staggering to ensure the accessibility of the seal seam for the test gas (Section 19.4.4).

Also in vacuum technology, the quality assessment of welded joints is to be questioned after the welding. Special attention is placed on the following criteria.

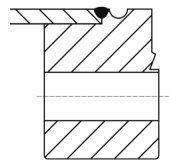


Figure 17.4 Sketch of a flange-to-tube weld with a heat insulating groove.

The *tightness* of the connection is reached through the absence of fissures, pores, bonding defects, and similar irregularities. Through their corrosive effects, oxidations such as scaling and annealing colors can directly or indirectly negatively influence the *outgassing behavior* of the connection. The use of welding fillers in vacuum technology is to be critically examined because extremely high cleanliness requirements are made for the weld seams. The application of welding fillers with high cleanliness and without fluxes can counteract against the melting loss of alloy elements and therefore diminish the negative influence on the mechanical stability of the joint. The mechanical stability of a joint is based on its cohesion and is closely linked - alongside the materials used and the seam geometry - with the dimensions of the connection. In addition to this, the originating distortions and stresses must be considered with larger cross sections. Shape and position tolerances are of increased importance in many vacuum applications. Examples of this are the requirements in accelerator technology, the closely tolerated positioning of focus points in the analysis measurement technology, or the need for flat sealing surfaces. Narrow form and position tolerances can require cost-intensive adjustment units, additional construction expenditure (e.g., greater wall thicknesses, heat insulating grooves), or intermediate machining steps (e.g., refacing of flanges and pipe ports).

In some applications of vacuum technology, particularly in accelerator technology, the change of the magnetic *permeability* during the welding process can have a negative influence on the application. In such cases, the microstructural state of the material as well as the heat influence by the weld process must be considered.

According to the material, welding techniques, and areas of application, different standards are used for the conventional quality assessment of welded joints. For fusion welded joints (except beam welding of steel), DIN EN ISO 5817:2014-06 [23] is frequently applied. Because this standard places the mechanical stability of the connection in the center, adaptations and enlargements must be made that embrace the previously named specific requirements for the respective vacuum application.

Welding techniques are qualified and standardized in order to achieve connections with high quality and reproducibility. In DIN EN ISO 15607:2004-03, instruction can be found for the development of a qualified provisional welding procedure specification (qWPS) up to the qualification for the finished welding procedure specification (WPS) [24].

17.3.1.2 **Brazing**

The thermal process for a material-bonded joining is referred to as brazing in which the materials are connected with the aid of the liquefaction of an additional component that is called solder. The melt temperature of the base material, the so-called solidus temperature $T_{\rm S}$, is not reached with brazing. The permanent connection originates from wetting the solid base material with the liquid solder and from diffusion of the molten solder into the base material. Solder components form no intermetallic phases with base materials.

Because a fusion of the base materials is not necessary with brazing, materials with different melting ranges can be connected with each other (e.g., steel with copper, ceramics/glass with stainless steel). Brazing is preferentially used in case of formed components with low wall thicknesses or narrow tolerances or hardly accessible joints (Figure 17.6). In vacuum technology, brazing is frequently used to realize metal to ceramic connections (e.g., in electrical feedthroughs), steelcopper connections (e.g., with coolant feeds), or for components made of nonferrous metals (e.g., titanium and aluminum).

A requirement for a successful brazing is the choice of a solder material suited to the basic materials and operating conditions. The following criteria have to be considered:

- · Melting range of the solder material.
- Wettability of the base metals by the liquid solder.
- Solubility of the solder components in the base material.
- Corrosion resistance of the brazing.
- Producibility as solder rod, tape, wire, foil, paste, or powder.

For a successful joining, suitable brazing constructions (narrow gaps, force transmission) and precisely defined brazing processes (defined surfaces, brazing atmosphere, temperatures, hold times) are also necessary with vacuum applications. Figure 17.5 shows an overview of brazing constructions compatible with vacuum.

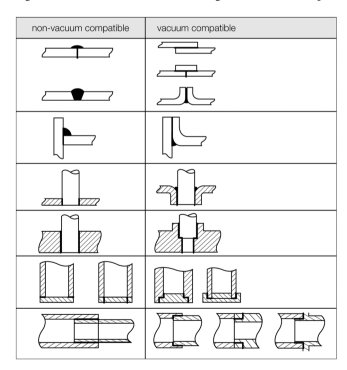


Figure 17.5 Examples of suitable brazing constructions. (Following Ref. [22].)

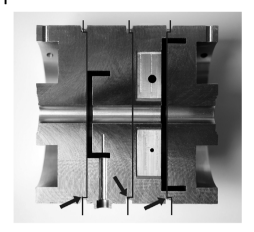


Figure 17.6 Cavity beam position monitor (E-XFEL prototype) in cutaway view, brazed with copper foil 2.0091 DIN EN 1044 CU 104 as an example of a vacuum component with hardly accessible joints. (Courtesy of the Deutsches Elektronen Synchrotron Hamburg.)

Metallic solders are divided according to the temperature of their complete liquefaction, the so-called liquidus temperature $T_{\rm L}$, in soft solders ($T_{\rm L} < 450\,^{\circ}{\rm C}$), hard solders ($450\,^{\circ}{\rm C} < T_{\rm L} < 900\,^{\circ}{\rm C}$), and high-temperature solders ($T_{\rm L} > 900\,^{\circ}{\rm C}$). High-temperature solders are free of fluxes. The brazing process takes place under inert gas or vacuum atmosphere. The so-called *active solders* that are used for the brazing of ceramic materials form a special group of metallic solders. Ceramics are generally not sufficiently well wetted by metallic fusions. With the addition of active, reactive (oxygen-affine) elements such as titanium or zircon, the wetting and the brazeability of ceramics are improved through the change in the interfacial chemistry.

Solder materials must comply with additional requirements for the application in vacuum technology:

- Formation of a gas-tight connection.
- Low outgassing of the solder components.
- Low contamination with vacuum-affecting elements.
- Mechanical stability of the connection (stability with pressure differences ≥1 bar).
- Heat stability (stability with baking out processes).

Soft solders (primarily based on zinc, lead, or cadmium) are not used in vacuum technology due to the high vapor pressures of the individual solder components and the resulting high outgassing rates as well as the low heat stability.

Hard solders that are suitable for vacuum applications are alloys based on *copper*, *silver*, *gold*, or *nickel*. For special vacuum requirements, solder materials with very low contaminations in carbon, cadmium, phosphorus, lead, zinc, manganese, and indium are primarily used. The contamination limits are listed in DIN EN ISO 17672:2010-11 "Brazing – Solders" [25]. In this standard, the exact compositions and characteristics of hard solders are also explained. Table 17.4 shows an overview of selected hard solders that are used in vacuum technology.

Table 17.4 Selected hard solders for application in vacuum technology.

Brazing group	Abbreviation according to DIN EN ISO 17672:2010-11	Composition	Operating temperature (approximately) (°C)	Remarks	Ranges of application
Aluminum hard solders	Al 110	Al 90 Si 10	600	Unsuitable for anodized components	Rolled/cast aluminum alloys, connection aluminum with Cr–Ni steels
	Al 112	Al 88 Si 12	590	Solidus temperature of the base materials must be >630 °C Unsuitable for anodized components	Rolled/cast aluminum alloys with Mg concentration ≤3%, connection aluminum with Cr–Ni steels
	CuP 279	P 6.3 Ag 2 Residual Cu	740	Potential contaminations max. 25% Unsuitable for steel and NiFe alloys, unsuitable for vacuum	For induction furnaces or inert gas furnaces electrical industry/air-conditioning (from -50 to $+150$ °C)
	CuP 284	P 7.05 Ag 18 Residual Cu	650		
Silver hard solder	Ag 272	Ag 72 Cu 28	780	Extremely versatile in application; also for alloyed and unalloyed steels, nickel	Vacuum technology connections, copper/ copper, copper/stainless steel (nickel-plated), stainless steel/stainless steel (nickel-plated), copper/ceramic connections
Copper hard solders	Cu 104	Cu 99.9 P <0.025	1100	Also suitable for gap brazings	Alloyed and unalloyed steels and for vacuum application, brazing of stainless steels with- out nickel plating
	Cu 105	Cu 98 Ni 2	1110	High tensile strength of the connection	Alloyed and unalloyed steels, hard metals, W, Mo, Ta
	Cu 922	Cu 94 Sn 6	950–1000		For copper, iron, and nickel materials

(continued)

Table 17.4 (Continued)

Brazing group	Abbreviation according to DIN EN ISO 17672:2010-11	Composition	Operating temperature (approximately) (°C)	Remarks	Ranges of application
	Cu 925	P 0.2 Cu 88 Sn 12 P 0.2	950–1030		
Gold hard solders	Au 295	Au 30 Cu 70	1000-1020	Good high-temperature oxide resistance	Vacuum tubes, electrical tubes, vacuum applications
	Au 375	Au 37.5 Residual copper	990		
	Au 752	Au 75 Residual Ni	960-985	High stability of connection and good high-temperature corrosion resistance	Engine construction, vacuum applications similar to 827
	Au 827	Au 82 Residual Ni	950	High stability of connection, good high- temperature corrosion resistance, alloyed steels, Ni alloys, stainless steels	Engine construction, vacuum applications; material pairing: Fe/Cr, Mo/W, Ni, Ni/Cu, Ni/Fe, Fe/Co, Kovar, and Vacon®
Palladium- containing solders	Pd 288	Ag 95 Pd 5	1015	Vacuum melting solder, oxide-free	Metallized ceramics, zircon, stainless steels, copper alloys, and nickel alloys
	Pd 647	Pd 60 Ni 40	1237	Poor resistance to HF (hydrogen fluo- ride), very good flow properties	Metallized ceramics, stainless steels, copper alloys, and nickel alloys
Active solders		Ag 70.5 Cu 26.5 Ti 3	803–1477	Brazing under vacuum or argon, silver is easily able to diffuse under vacuum (solder temperature max. 900 °C depending on the pressure)	Ceramic metal connection without activation coatings such as nickel, graphites, glass, diamond

Courtesy of the Deutsches Elektronen Synchrotron Hamburg.

Different brazing processes are applied according to solder temperature, shape, and material of the components to be joined. In addition to furnace brazing, flame brazing, resistance brazing, induction brazing, and dip brazing are to be named. These primarily take place with the aid of fluxes for the removal of metal oxide on the surface. Nevertheless, flux residuals can be integrated in the surface and, with this, show a vacuum-affecting or corrosive effect.

Vacuum components are preferentially soldered with high-temperature brazing, that is, in the furnace under vacuum or inert gas atmosphere and without flux. Here, the entire component is fixed with fitted, applied, or inlaid solder and heated up to solder temperature. Several joints of complex, large components can be soldered at the same time and thus show minimal distortion. Small brazing components can be produced in large quantities in a single batch. The furnace allows constant and defined heating up and cooling down, which is of advantage for the fusion and solidification processes of the brazing alloys. Through an adjustable inert gas atmosphere (CAB - controlled atmosphere brazing) or under vacuum, reactions of the base material and solder with the environment can be eliminated or selectively promoted. The desorption rates of the workpiece surfaces are already lowered during brazing as a result of the high process temperatures. If necessary, a heat treatment can be directly connected to the brazing process in the furnace.

Independent of the brazing process, attention is absolutely to be paid to the fact that while heating up the components the solder temperature is reached not only at the heat source or at the temperature measuring thermocouple, but also truly on the surfaces to be soldered. At too low temperatures, insufficient wetting or flowing of the solder will entail incomplete gap filling. The results are leaky brazing joints or considerable strength losses. A too quick cooling down leads to segregations (local decompositions) of the molten solder that favor the formation of undesirable microstructures with brittle phases, microcracks, and significant concentration fluctuations. Poor or unpredictable mechanical properties of the soldered joint are the result.

17.3.1.3 Adhesive Bonding

Adhesive bonding is a nonthermal, material-bonded joining process in which an additional component (adhesive) wets the surfaces of the base material. In particular, adhesive bonding finds application:

- if the materials and material combinations to be joined are difficult or not weldable (e.g., titanium, tantalum with stainless steel),
- with nonbrazable materials with low temperature resistance (e.g., plastic, aluminum).
- if a low-distortion and low-stress joining is necessary,
- when joining materials are sensitive to temperature.

The base material is not modified with bonding – unlike with brazing and welding. Because no diffusion processes take place between the materials used, a vast number of material combinations can be joined.

A distinction is made between physically and chemically bonding adhesives. For vacuum applications, bonding is only used with chemically curing adhesives because physically bonding adhesives reach adhesion through the evaporation of solvents or dispersion media. Chemically reacting adhesives produce their adhesive properties through chemical reactions, for example, through crosslinking [26]. Therefore, a distinction is made between two- (or more) component systems and one-component systems.

With two-component adhesives (2C-adhesives), two spatially separated preparations are used. One of the preparations contains resinous monomers (binders), while the other contains hardeners. The chemical reaction of the adhesive polymer begins with the contact between the binder and the hardener. Therefore, 2C-adhesives can only be processed within a defined period of time. After insertion of the adhesive in the joint, the curing time in which the final strength of the bonding is built follows. This curing time is influenced by temperature and electromagnetic waves (e.g., UV light).

With one-component adhesives (1C-adhesives), the ready-for-use adhesive is inserted directly into the adhesive joint. The adhesive hardens with the change in the ambient conditions. This can happen, for example, through a temperature rise, admission of air humidity, exclusion of air oxygen, or contact with the substrate surface. Also with the chemically hardening 1C-adhesives, chemical reactions between the binder and hardener are responsible for the formation of the polymer. As opposed to the 2Cadhesives, the binder and hardener of 1C-adhesives cannot or only extremely slowly react under the storage conditions recommended by the manufacturer. In the databases of the NASA [27] and the European Space Agency ESA [28], approximately 2500 adhesives are listed and characterized with regard to the mass loss in vacuum. A low TML value indicates a good vacuum suitability by a low outgassing rate.

Adherence has extremely high requirements for the cleanliness and freedom from grease of the surfaces to be joined. Contact with the hands alone is sufficient to diminish the adhesive effect and render the bonding useless. The bonding strength can be increased by machining, such as, for example, by grinding, brushing, or blasting. Through roughening and the enlargement of the surface associated with it, the adhesive is additionally form-fitted in the surface. Chemical (e.g., pickling) or physical pretreatments (e.g., corona and plasma processes, flame treatments) can also increase the bonding strength. For the after-treatment of the adhesive seam, for example, the use of primers (preservation of the adhesive bond) is available. The use of primers improves the adhesion quality and protects the bonding against humidity and corrosion. In this process, the primer must always be adapted to the adhesive.

As with all other joining processes, a suitable bonding design for the components that are to be joined must be found. This means that sufficiently large bonding surfaces must be provided in the component design. In the application, the adhesive seam itself may only be stressed on shear (a), and tension (b), peel (c) and bending (d) stresses must be avoided (see Figure 17.7).

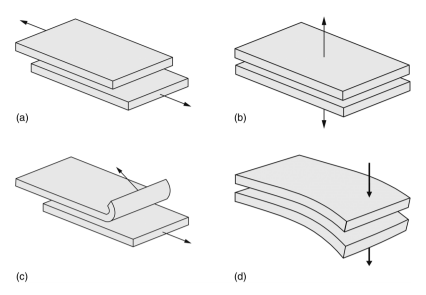


Figure 17.7 Adhesive seam stresses.

Bondings are always subject to aging and their durability is determined by mechanical (static and dynamic forces), chemical (humidity, solvents), and physical (heat, UV, and other radiation) influences. In the selection of the adhesive, the possible temperature load must also be considered alongside the ambient conditions. The adhesive will become brittle and crack at too low temperatures. At too high temperatures, the adhesive will soften and can melt/degrade.

17.3.2

Detachable Connections

For reasons of flexibility and a simple maintenance, the individual components of a vacuum facility should be detachably connected with each other. Form-fit and force-fit connections must be suitable for the many application-specific aspects in vacuum technology. The connections must comply with the different process-related requirements:

- pressure range,
- outgassing behavior,
- · leak tightness,
- temperature resistance,
- chemical resistance,
- · magnetizability,
- particle generation.

All detachable connecting systems in vacuum technology need a sealing medium, generally referred to as seal, to ensure the necessary leak tightness.

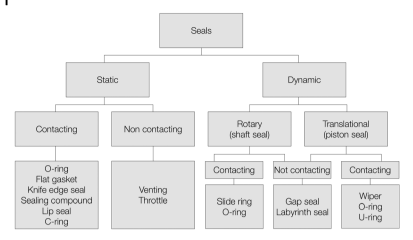


Figure 17.8 Classification of seals in mechanical and plant engineering.

Seals count among the design elements in mechanical and plant engineering and are divided into static and dynamic elements (see Figure 17.8). In both cases, contacting seals are common in vacuum technology; dynamic, noncontact gap seals are known for special applications [29]. Static seals for vacuum applications are O-ring seals, flat seals, knife edge seals, and lip seals; sealing compounds are used in special cases. O-ring seals and lip seals are only activated through a pressure difference between the spaces to be sealed, whereas flat and knife edge seals require an external pressing force for the sealing effect. Differently from that common in mechanical engineering, in vacuum technology, such seals that are loosened or connected during the operation of a vacuum system are considered as dynamic seals, for example, valve plate seals. Although this classification is systematically not quite exact, a valve plate seal also has its fundamental function in the static sealing and thus must never seal and allow a relative movement of the seal partners at the same time. Nevertheless, this widespread classification should be used here (Figure 17.8).

The detachable connections that are widespread in vacuum technology can be most suitably classed according to their permeation (leak) rates and therefore according to the pressure range in which they can be used. Table 17.5 provides an overview of the most common connecting systems and elements.

The division of the different connecting systems according to the pressure range reveals that varying values of leak rates of the connections limit the application.

17.3.2.1 Elastomer-Sealed, Static, Detachable Connections (KF, ISO, and Others)

Elastomer-sealed connections are based on force-activated seal materials and are applicable from the overpressure up to the vacuum range. The material composition of the elastomer seals and the geometry determine the lower pressure operation limit in the vacuum. With suitable selection and

 Table 17.5 Detachable connections of vacuum technology and their areas of application.

Area of application	Connecting system	Seal partner	Seal	Typical applications	Typical He permeation/ leak rate (Pa ℓ s ⁻¹)
Fine vacuum	Conical ground glass joint	Glass cone surfaces	Grease	Glass appara- tuses for chem- istry/biology	Not specified
High vacuum		Flat sealing surfaces	Metal C-ring coated	Special applica- tions, large nominal diameters	10 ⁻⁵
	Lip seal	Flat sealing surfaces	Elastomer lip seal	Vacuum vessel doors	10^{-6}
	ISO/KF flange system	Flat sealing surfaces	Elastomer O-ring	Vacuum sys- tems in high vacuum	$10^{-6 \text{ a}} (10^{-7})$
	Swagelok [®] Ultra-Torr	Metal cone surfaces	Elastomer O-ring	Pipe screw connections	4×10^{-7}
	Swagelok [®] VCR [®]	Metal seal lips	Metal flat seal	All-metal fittings	4×10^{-9}
Ultrahigh vacuum		Flat sealing surfaces	Metal O-ring concave, gas- filled, coated	Special applica- tions, large nominal diameters	10 ⁻⁷
	KF flange system	Flat sealing surfaces	Metal edge seal		$<4 \times 10^{-7}$
	Bostec® H-Seal	Knife edge	Metal profile seal	Special applications	Not specified
	Helicoflex [®]	Flat sealing surfaces	Covered metal spiral ring	Special applica- tions, large nominal diameters	<10 ⁻⁸
	VATSEAL	Flat sealing surface	Metal profile seal		<10 ⁻⁸
	COF flange system	Conical sealing surfaces	Metal round wire	All-metal seals, large nominal diameters	<10 ⁻⁸
Extreme ultrahigh vacuum	CF/QCF flange system	Knife edge	Metal flat seal	Vacuum sys- tems in ultra- high vacuum and XHV	<10 ⁻⁹

a) Leak rate with ISO and KF flange systems depending on the flange material.

arrangement, it is quite possible to use elastomer seals for an ultimate pressure of up to 10^{-8} Pa.

The basic principle of elastomer-sealed connections consists in pressing an elastic material between two sealing surfaces. By the deformation of the elastic material, a flat seal develops so that the direct through-flow of gas is prevented or minimized. The geometry of the seals and the sealing surfaces can be shaped in a variety of forms. In addition to the most widespread toroidal sealing ring (O-ring), the X-form, H-form, or different lip forms find application in the sealing technology (Figure 17.12). Because with vacuum applications the pressure difference is 1 bar at maximum, the determining challenges lie in the leak tightness. In mechanical engineering, a distinction is made between main load connections (Figure 17.9 a and b) and off-load connections (Figure 17.9 c, d, e, and f). In the main load connections, the full mounting force presses on the seal, which leads to a distinctive setting behavior (stress relaxation). Consequently, a time-dependent decrease in the sealing effect and an increase in the

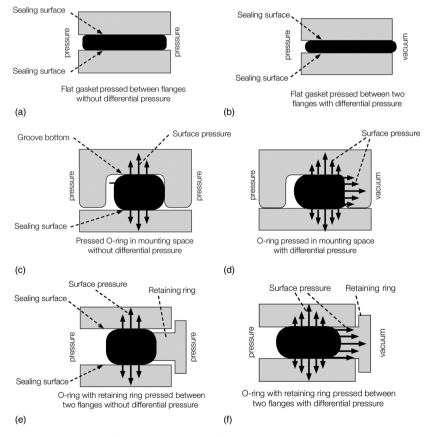


Figure 17.9 Charting of the sealing principles for elastomer-sealed connections.

leakage rate occur. Therefore, the off-force sealing principle primarily finds application in vacuum technology. The plastic deformation is reduced through suitably designed grooves or retaining rings (Figure 17.9e and f) and, with this, the preservation of the sealing effect is extended. Attention is to be paid to the fact that the O-ring is in contact with the groove wall in which it is later pressed through the pressure difference.

The hardness of the elastomer material used has a considerable influence on the technical tightness of an elastomer seal. Softer materials can better seal gaps in sealing surfaces, whereas harder materials generate a higher pressure on the surfaces. Elastomer materials with a hardness from Shore A 60 to Shore A 85 are commonly used in vacuum technology [30].

The sealing effect of elastomers is limited to a material-dependent temperature range. Below the glass transition temperature $T_{\rm G}$, the material is in a hard, glassy state. The elastic properties and, with this, the sealing effect get lost entirely. This process is reversible. In contrast, polymer chains above the decomposition temperature $T_{\rm Z}$ are destroyed and the sealing effect is permanently eliminated as a result. As a rule, the operating temperatures of elastomers are specified for a period of application of 1000 h. Should a seal hold longer, it may not be operated on the upper limit of the operating temperature for the material.

NBR, FKM (formerly FPM), or FFKM elastomers are predominantly used in vacuum technology whereby FKM and FFKM elastomers are characterized by a higher chemical resistance and temperature resistance (Figure 17.10). They primarily consist of vulcanized rubber and a filler (e.g., soot). In its vulcanized form, the rubber provides for the elasticity, and the hardness of the elastomer is adjusted by the proportion of the filler. Other components such as plasticizers, sulfur as a vulcanizing agent, and zinc oxide as a vulcanizing activator as well as various processing agents and antioxidants are additionally included. There are no standards for elastomer materials in which the exact composition is specified. Hence, the composition is strongly dependent upon the manufacturer. Products with similar names can have very different properties (particularly with regard to the outgassing rate). In addition to the operating temperature, the surrounding atmosphere considerably influences the aging of elastomers. Ozone, in particular, causes rapid aging of elastomers. The strong oxidation agent splits the polymer

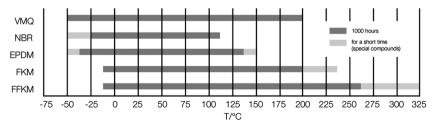


Figure 17.10 Elastomer materials in vacuum technology and their operating temperatures/durations (manufacturers' data): FFKM (perfluoro rubber), FKM (fluoro rubber), EPDM

(ethylene propylene diene monomer rubber), NBR (acrylonitrile butadiene rubber), and VMQ (methyl vinyl silicone rubber).

Polymer type	Temperature (°C)	$Q_{\rm P}(\text{helium}) (\times 10^{-9} \text{cm}^2 \text{s}^{-1} \text{bar}^{-1})$
NBR	25	80
NBR	80	660
FKM	25	130
FKM	80	1310
FKM	150	4900
FFKM	25	40

Table 17.6 Gas permeability coefficient Q_P (in 10^{-9} cm² s⁻¹ bar⁻¹) of various elastomer materials depending on the temperature (manufacturers' data).

chains and can cause fissures also at low mechanical loads. This must be considered for the storage of elastomer seals; that is, circulating air and direct sunlight are to be avoided. In general, it is recommended to control elastomer seals regularly for damage such as surface fissures. By no means cracked seals may be used.

The gas permeation is determined by the material properties (base material and filler material) and is measured with standardized test procedures [31]. The gas permeation is dependent upon gas type and temperature. In Tables 17.6 and 17.7, selected gas permeability coefficients are listed depending on different base materials, material compositions, temperatures, and gas types.

The effect of the permeation can be reduced if differentially pumped double seals are used because the pressure difference influences the permeation. Thus, a circumferential, separately evacuable space is created between two elastomer seals.

The outgassing behavior of O-rings can be reduced through suitable processes. With the example of FKM and FFKM O-rings, Figure 17.11 demonstrates how the outgassing behavior can be significantly reduced through suitable tempering processes.

Table 17.7 Gas permeability coefficient Q_P (in 10^{-9} cm² s⁻¹ bar⁻¹) from NBR elastomers depending on its acrylonitrile concentration (ACN) for different gases and temperatures (manufacturers' data).

Gas		Air		Nitrogen		Carbon dioxide	
Temperature (°C)	60	80	60	80	60	80	
ACN concentration NBR (%)	Gas permeability coefficient Q _P						
18	140	0	60	80	60	80	
28	75	120	90	_	750	_	
24	35	70	40	70	580	970	
39	25	55	20	55	560	630	
			10	25	300	480	

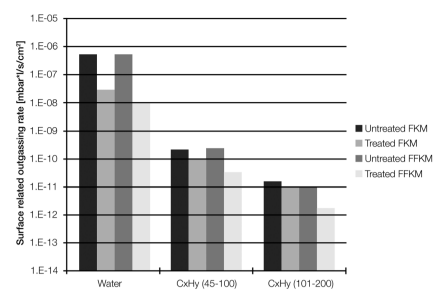


Figure 17.11 Outgassing rates in units of the generated partial pressure for untreated, low outgassing FKM and FFKM O-rings.

In many applications, elastomer O-rings are provided with a grease before the assembly. The grease serves as a protective film, supports deviating movements in excessive force effects, and seals roughness valleys as well as smaller scratches on the sealing surfaces. In the HV range, attention is to be paid to the fact that only suitable greases are used, such as, for example, the PTFE-filled lubricants that are declared as suitable for vacuum. No greases may be used in the UHV and XHV ranges and with ultraclean vacuum.

Specific directives must be considered in the design of elastomer-sealed connections. With flat seals, merely the intact surface is vital and the surface roughness should be Ra 1.2 µm or better. In contrast, profile seals must be fixed, which usually occurs with a groove. Lip seals primarily have another bead that can be pressed into the groove. For technological reasons, O-rings are fixed in rectangular grooves and the groove must always be wider than the cord thickness of the O-ring. Elastomers are approximately incompressible materials, which is why the cross section of the groove must be able to take up the complete seal cross section. Additional effects such as swelling and thermal expansion, which can cause considerable cross-sectional changes in elastomers, are to be considered. FKM O-rings, for example, expand by up to 30% between 200 and 300°C. At the same time, the O-ring may be stretched or compressed at most by up to 5% depending on the subsequent pressure direction. With the assembly, attention must be paid to the fact that the stretching or compression is evenly distributed throughout the entire length of the seal. This is achieved while the seal is first pressed into the groove in some places over the full length and only then completely inserted. In order to avoid damages in the seals, the outside edges of the

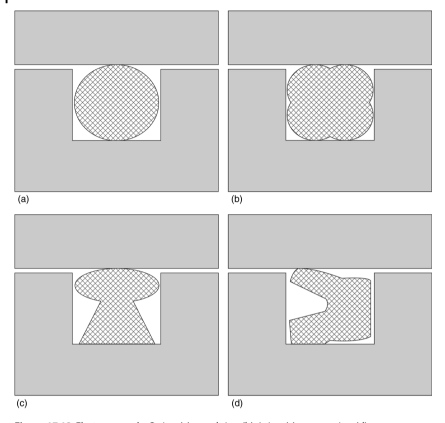


Figure 17.12 Elastomer seals: O-ring (a), quad ring (b), I-ring (c), groove ring (d).

groove should be rounded or chamfered. Fundamentally, the groove depth is to be selected in such a way that with static sealings a compression (cross-section change) of the O-ring between 15 and 30% is possible.

In addition to rectangle grooves, trapezoid-shaped grooves or V-grooves also find application. The named directives are to be considered/observed in the dimensioning of these grooves. The roughness of the sealing surfaces must be Ra $0.8\,\mu m$ or better here (Figure 17.12).

Elastomer O-rings are also used in screw connections with which the sealing force is introduced through a thread. Crimped O-rings that are additionally exposed to torsion forces can be spared by lightly greasing, which, however, limits their application at low pressures as well as the low outgassing.

In vacuum technology, the existing standardized clamp flanged joints (Figure 17.13, right) and Klein flange quick release couplings (Figure 17.13, left) are described in DIN 28404:1986-10 "Vacuum technology; flanges; dimensions" [32] or ISO 1609:1986-03 "Vacuum technology – Flange dimensions" [33]. With the ISO flanged joints, a distinction is made between clamp flanges (ISO-K) with clamps or support brackets and ISO-F flanges with screw holes. Both variations

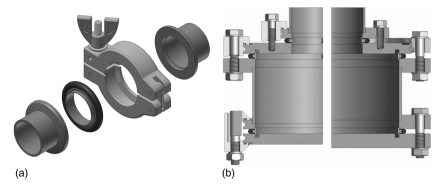


Figure 17.13 Schematic drawings of Klein flange quick release couplings (KF, a) and clamp flanged joints (ISO, b).

are easily combinable with each other, while an ISO-K flange with a collar flange becomes an ISO-F flange. The sealing forces for the different nominal diameters are specified in the appendix of the standard. From the number of the required screws and the nominal diameter of the used O-ring seal, a sealing force per mm of seal length is obtained. Klein flange couplings (KF) are described in DIN 28403:1986-09 "Vacuum technology; quick release couplings; clamped type couplings" [34] or ISO 2861:2013-05 "Vacuum technology – Dimensions of clamped-type quick-release couplings" [35] for the nominal sizes DN 10, DN 16 (DN 20), DN 25 (DN 32), DN 40, and DN 50.

In addition to the components explained in the standards, a large number of other designs of the clamping element, clamping chains made of plastic, aluminum, or stainless steel, collar half shells, and claws for the direct assembly on chambers exist. Clamping chains or half shells of stainless steel are necessary for sealing KF flanges with metal seals (e.g., aluminum, copper). With KF and ISO flange systems, the elastomer O-ring seal is usually held by an internally lying centering ring and its dimension is compressed by maximal 22–25% by the axial force, which is generated between the two sealing planes. The elastic quality of the seal generates a reset force that constitutes a hermetic seal between the sealing surfaces and the seal. By evacuation of the inner volume, the originating pressure difference generates an additional force component that increases the sealing effect.

In addition to these standardized flange systems, there are other established connections for pipe connections using an elastomer seal with diameters of up to approximately 50 mm. With the Ultra-Torr connections by Swagelok®, a tube can be vacuum-tight clamped into a weld fitting, for example. They are designed in such a way that the crimped O-ring has as little as possible contact with the vacuum.

The so-called vulcanized seals represent a specific feature of the elastomer seals. This seal is nondetachably connected with one of the two seal partners. This variation finds use especially with vacuum valves.

17.3.2.2 Metal-Sealed, Static, Detachable Connections

In the UHV and XHV ranges, seals made of a metallic base material are used whereby the metallic seals must be plastically deformed for the achievement of the necessary sealing effect. The plastic deformation provides for the fact that the production-related unevenness and waviness of the sealing surfaces are compensated and roughness valleys are closed. As seal materials for CF and QCF connections, oxygen-free, high-purity copper (99.99%, OFHC quality, CU-OFE, material number CW009A) is primarily used; pure aluminum (99.5%, EN AW 1050) is more rarely used. In special applications, other metal materials are also used. If the connections are baked out at more than 200 °C, silver-plated copper seals should be used to protect the seal against oxidation.

The most widespread metal seal in vacuum technology is the combination of a flat seal and sealing surfaces with a knife edge (CF flange system, Figure 17.14a). In the ISO 3669 standard "Vacuum technology - Bakeable flanges - Dimensions," the external flange dimensions and the number and position of the screws are defined depending on the flange diameter [36]. The sealing forces to be

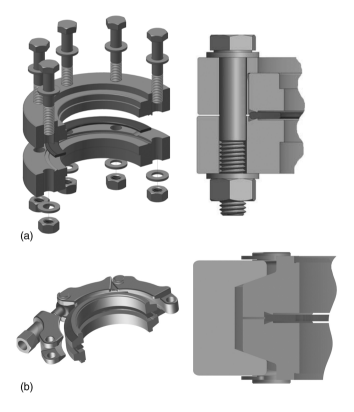


Figure 17.14 Schematic drawings of the CF flange system (a) and the QCF flange system for the simple, quick, and space-saving implementation of CF flanged joints (b).

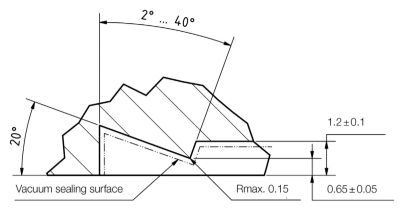


Figure 17.15 CF flange system: cross-section drawing of the knife edge.

applied in reality and the resulting contact pressure are dependent upon the hardness of the flange and sealing materials as well as upon the knife edge geometry. From 2007 and in supplement to the previously named standard, there is ISO/TS 3669-2 "Vacuum technology - Bakeable flanges - Part 2: Dimension of knife-edge flanges" that is used worldwide as technical specification (TS) [37]. In addition to the external dimensions of the flanges, it also defines the geometry of the knife edge that is shown in Figure 17.15. For the generation of the sealing force, the use of stainless steel screws grade A2 strength class 70 with a tensile yield strength of 640 MPa is common. If flanges as well as bolts and nuts are made of stainless steel, suitable measures are to be taken to avoid galling (seizure) of the threads. Galling describes the formation of a connection between components on the basis of adhesion. Due to their adhesive properties and the lower hardness of Cr-Ni steels, they are especially vulnerable to galling in comparison with most other steels. The application of lubricants that are suitable to vacuum technology or Kolsterising® [38] of the connecting elements reduces the danger of the galling [39].

The QCF connection demonstrates the further advancement of the existing CF flange system (Figure 17.14b). This flange system unites the simple assembly principle of the KF connections with the sealing principle of the CF connections. With this flange system, the sealing force is applied through a VaCFix[®] clamp chain on conically shaped flanges. The seal and sealing surface or knife edge of the flanges correspond to ISO/TS 3669-2. The use of the VaCFix[®] clamp chain, which ensures the same tightness and operational safety, enables a substantially easier, quicker, and more space-saving assembly of the flanged joint than the conventional CF flanges to be connected with screwing.

In addition to flat seals, profiles and concave rings with an open (C-ring) or round cross section as well as diamond-shaped (rhombic) cross sections (e.g., VATSEAL) are used. A great advantage of this seal form is the applicability for any flange geometry.

To achieve a better sealing effect with hollow ring seals, they are often provided with soft metallic coatings on the exterior side, for example, silver or aluminum. A variation of the metal C-rings are referred to as Helicoflex® seals in which an additional spiral spring is brought into the ring cross section. The spring causes an enlargement and a steadier distribution of the reset force.

At flange nominal sizes of 400-800 mm, wires with circular cross section, known as COF connections, are mainly used (Figure 17.16). A metal wire that is realized as a round seal is pressed between two conically formed sealing surfaces.

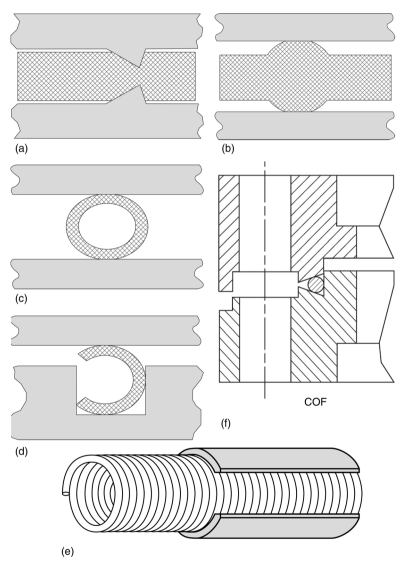


Figure 17.16 Metal seals: Flat seal and sealing surfaces with knife edge (a), profile seal (b), metal O-ring seal (c), C-ring (d), spiral Helicoflex® (e), COF flange system (f).

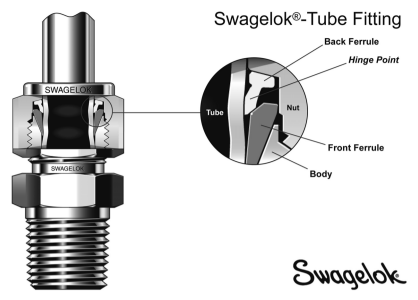


Figure 17.17 Sealing principle of Swagelok® (advanced) ferrule connection. (Courtesy of Swagelok® Company.)

The advantage of the connection lies in the lower assembly forces and the lower susceptibility of the connection to deformations of the seal.

KF and ISO flanges can also be metal sealed. The flanges must be able to merely transfer the required forces. Edge seals as well as flat seals are available, but are not standardized.

There are different metal-sealed solutions for pipe connections. The most widespread systems are known under the brand names Swagelok[®] (Figure 17.17) and VCR® (Figure 17.18). In this instance, two cone surfaces that are pressed together swage a knife edge into the exterior surface of the pipe.

Swagelok®-VCR® Assembly

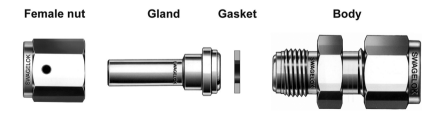




Figure 17.18 Typical VCR® connection with body gasket, gland, and female nut. (Courtesy of Swagelok® Company.)

17.4 Components

17.4.1

Standard Components and Chambers

The design of a vacuum chamber decisively influences the success and the repeatability of processes or experiments and at the same time is an essential expense factor. With the materials and manufacturing methods available today, a large part of the requirements in chamber building can be realized with standard components. Various components are required for the construction of vacuum systems (Figure 17.19). Leak-tight connections are primarily achieved through standardized flanges. The functional components are joined through connecting and manifold pieces, such as, for example, crosspieces or elbows.

In addition to the design of vacuum chambers applicable to vacuum, the principle of important functional components such as valves and electrical, mechanical, and optic feedthroughs is described. Furthermore, the possibilities of the heat supply and dissipation to ensure the process stability are explained.

Vacuum-Applicable Designing and Joining

Before and also during the planning and design of a vacuum recipient, the user should also consider the potentially appearing mechanical loads, process-related mechanical stresses, the material choice, the necessary wall thicknesses, the stability of the joints, and necessary tolerances of the interfaces (flange outlets). The clarification of these questions enables the optimal design of the vacuum chamber for the subsequent application.

Undesirable outgassing can be minimized through the use of materials with low inherent vapor pressure and nonporous surfaces. In addition to the required mechanical stability against atmospheric pressure as well as corrosion resistance, the surfaces should be easy to clean. Depending on the application, stainless steel, aluminum, copper, glass, special elastomers (FKM, e.g., Viton®; PTFE, e.g.,



Figure 17.19 Vacuum components.

Teflon®), and ceramics are commonly used in vacuum technology. From a manufacturing point of view, the vacuum chambers can be divided into three groups:

• Cylindrical chambers. The simplest design is the cylindrical chamber (Figure 17.20). It is characterized by a very stable geometry. The resulting advantage with cylindrical chambers is that relatively low wall thicknesses can be realized. Cylindrical chambers can be closed through weld flanges, specially produced "lids," "dished heads," or domed disks. For the construction of cylindrical chambers, the industry offers standard elements in terms of tube diameters and wall thicknesses as well as for the closure of the chamber.



Figure 17.20 Cylindrical chamber.



Figure 17.21 Spherical chamber.

• Spherical chambers. The spherical chamber is a commonly used variation (Figure 17.21). Spheres dispose of an especially stable geometry against external and internal loads (pressure). In spite of the manufacturing-related larger tolerance of the spheres or hemispheres, exact focus points are realizable with modern technologies.

With the cylindrical as well as the spherical chambers, the manufacturing expenditure disproportionately increases with the reduction of the tolerances. In the concept phase, it should always be checked whether the necessary high accuracy can be achieved more cost effectively with adjustment possibilities for the components to be assembled.

Rectangular chambers. Rectangular chambers (Figure 17.22) are the third variation. With rectangular chambers, significantly more attention must be placed on the mechanical stability depending on the application. Due to their geometry, rectangular chambers are vulnerable to buckling during the pumping out. For the mechanical layout of rectangular chambers, the application of FEM software for the calculation of the deflection of the side walls is recommended.

Figure 17.23 shows how wall thicknesses and deflections can be minimized and costs can be saved through constructive measures and by means of examination with FEM calculations.

Concerning their accuracy (focus points), rectangular chambers are less complex to manufacture than cylindrical or spherical chambers. With their flat surfaces, the rectangular chambers offer optimum possibilities for components to be mounted.

• Double-walled chambers. For various processes, it is necessary to heat chambers or also to dissipate heat. This can be realized through the welding of

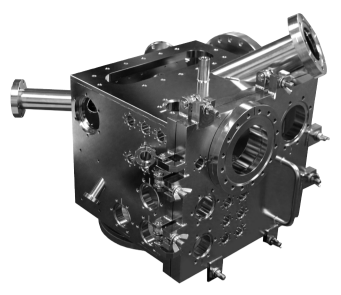


Figure 17.22 Rectangular chamber.

cooling channels (Figure 17.24) or with double-walled chambers. In order to prevent turbulences and to achieve an efficient temperature exchange, suitable guide plates that provide a forced direction to the medium are to be installed between the walls (Figure 17.25).

• Design of vacuum vessels. In view of the occurring stresses and pressure differences, vacuum vessels must be configured safely and reliably. In contrast to overpressure vessels, there are no mandatory regulations for vacuum vessels. Independent of this, the material choice, calculation, configuration, production, checking, and commissioning occur according to approved rules of engineering. For the orientation, for example, large-scale research institutes such as CERN, DESY, BESSY, or ESRF define their requirements for materials, production, and testing very specifically.

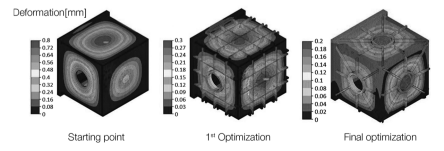


Figure 17.23 FEM simulation of the buckling of a rectangular chamber.



Figure 17.24 Cross with welded cooling channels.

For the design of the chambers, the leaflets of the "Arbeitsgemeinschaft Druckbehälter" (Pressure Vessels Working Committee) in the "AD 2000 Code for Pressure Vessels" [40] that specify all essential safety standards according to the European Pressure Equipment Directive can be referred to. The calculation fundamentals are also clearly summarized therein:

- B0 Design of pressure vessels (general part).
- B3 Domed ends subjected to internal and external overpressure.
- B4 Dished heads.
- B6 Cylindrical shells subjected to external overpressure.

The pages B3, B4, and B6 also contain equations for the calculation and examination of wall thicknesses with external overpressure that can be used for the configuration of vacuum chambers.

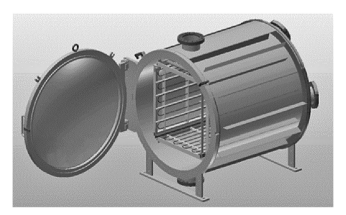


Figure 17.25 Double-walled vacuum furnace with guide plates.

Example 17.1

Dimensioning of Vacuum Vessels with a Calculation Example

For the dimensioning, the pressure in the vessel is to be set at zero. The external pressure amounts to 100 kPa (1 bar). The vessel has the form of a cylinder with domed ends (dished heads) (Figure 17.26). The design provides for the fact that the permissible stress in the wall is not exceeded. At the same time, the safety against plastic buckling must be provided. The tension σ is calculated from the equation

$$\sigma = \frac{pR}{S_{w}},$$

with p denoting the outside pressure, R the cylinder radius, and $S_{\rm w}$ its wall thickness.

Example calculation for a cylindrical chamber:

Diameter: 750 mm Length: 1000 mm Material: 1.4301 Temperature: 20°C

 $195\,000\,\mathrm{N}\,\mathrm{mm}^{-2}$ E-module: $210 \, N \, mm^{-2}$ Tensile strength value at 20 °C:

Safety factor S: 2 (safety factor at calculation temperature) Safety factor S_k : 3 (as a rule 3, with an unroundness of 1.5%)

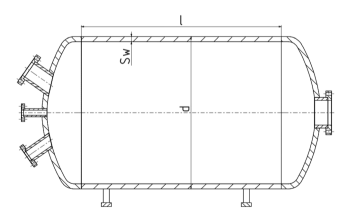


Figure 17.26 Drawing of a cylindrical chamber: *p* indicates the external pressure (100 kPa) and $S_{\rm w}$ the wall thickness. The wall thickness S_w of cylindrical bodies can be determined from Figure 17.27 (safety

against elastic buckling) and Figure 17.28 (safety against plastic deformation). The larger value from both calculations determines the necessary wall thickness.

Calculation of the Resistance against Elastic Buckling

From the diagram (necessary wall thickness s for the calculation against elastic buckling) follows:

"x" - value
$$\frac{p \cdot S_k}{E} \times 10^5 = \frac{0.1 \text{ N mm}^{-2} \times 3}{195\,000 \text{ N mm}^{-2}} \times 10^5 = 0.15,$$

$$\frac{D_{\rm a}}{I} = \frac{750 \text{ mm}}{1000 \text{ mm}} = 0.75 = 0.8.$$

The diagram shows approximately 7.25 (Y-axis) for 0.15 (X-axis) and 0.8 (curve). The formula $D_a/[(S_w-c_1-c_2)\times 10^2]$ after conversion results in (with S_w – $c_1 - c_2$ as the wall thickness including all safety values) 750 mm/(7.25 × 10²) = 1.03 mm (where c_1 is the allowance considering the deviation of the wall thickness and c_2 is the wear allowance). The necessary wall thickness for elastic buckling resistance amounts to 1.03 mm.

This is valid for austenitic sheets (with DIN EN 10029 as measuring standard) only for negative tolerances up to values of the lower dimension according to class A. In case of metal sheets with larger negative tolerances, the remaining difference is only to be considered. To simplify the matter, the example is calculated with $c_1 = 0$.

For austenitic plates and nonferrous metals, the wear allowance is $c_2 = 0$.

Calculation of the Resistance against Plastic Buckling

Now the necessary wall density for resistance to plastic buckling is yet to be calculated.

From the diagram (Figure 17.27) follows:

$$10 \times \frac{p \cdot S}{K} = 10 \times \frac{0.1 \text{ N mm}^{-2} \times 2}{210 \text{ N mm}^{-2}} = 0.01.$$

In the diagram, the value 0.01 is not readable. If one calculates with the value 0.03 and places for

$$\frac{D_a}{I} = \frac{750 \text{ mm}}{1000 \text{ mm}} = 1,$$

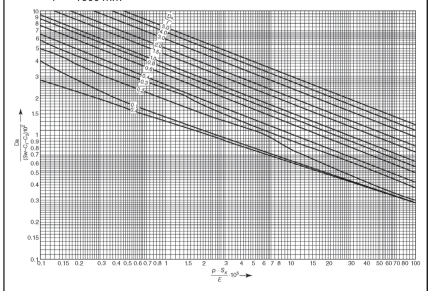


Figure 17.27 Necessary wall thickness S with calculated resistance to elastic buckling.

the value 5.8 arises. From the equation

$$\frac{D_a}{(S_w - c_1 - c_2) \times 10^2} = 5.8,$$

a wall thickness of 1.3 mm follows.

The minimum wall thicknesses designated in the AD 2000 Code with steel of 3 mm and with aluminum of 5 mm refer to external pressures over normal pressure (Figure 17.28).

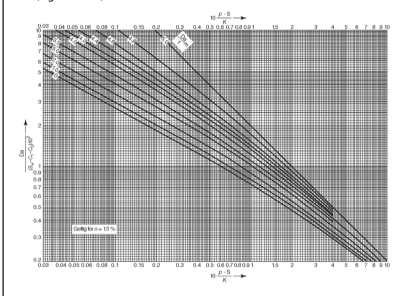


Figure 17.28 Necessary wall thickness S for the calculation of the resistance against plastic buckling.

With the design of vacuum chambers, in addition to the calculated wall thicknesses, the available wall thicknesses of tubes and hemispheres are also to be considered. The application of standard semifinished products is recommended. They must lie, in any case, above the calculated wall thicknesses.

17.4.2

Mechanical Feedthroughs

Mechanical feedthroughs are required to transfer the movements that are generated outside a vacuum chamber into the vacuum. The air-sided drive can occur manually, electrically, or also pneumatically. Already ensuring that the movement in the vacuum is low in loss and low in wear - without or with low outgassing lubricants - is quite a special challenge. At the same time, there is the additional requirement for a leak rate of the feedthrough that has to be as low as possible.

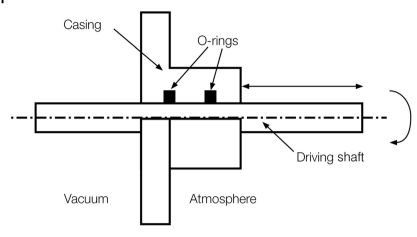


Figure 17.29 O-ring sealed rotary-linear feedthrough.

A mechanical feedthrough basically consists of a force-transferring element (shaft), a seal, and a flange. Mechanical feedthroughs are divided into three groups according to their operating principles:

- feedthroughs with dynamic seals,
- · feedthroughs with flexible elements,
- feedthroughs with magnetic force coupling.

The best known examples of dynamic seals are O-ring sealed (Figure 17.29) and magnetofluid sealed (Figure 17.30) feedthroughs whereby the latter are only

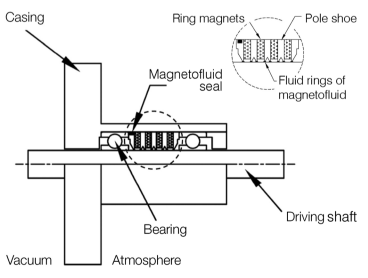


Figure 17.30 Magnetofluid sealed rotary feedthrough.

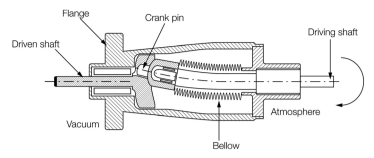


Figure 17.31 Bellow-sealed rotary feedthrough (so-called cat tail).

used for rotary motions. It is typical of these feedthroughs that the seal zone always shows a dynamic leak rate between the fixed element and the moved element that, as a rule, is greater than the static leak rate. O-ring sealed feedthroughs are subjected to a continuous wearing of the seal.

In feedthroughs with flexible elements, edge welded bellows or hydroformed bellows are primarily used. Many and also precise movement types can be realized with these feedthroughs. They are characterized by the fact that they show no dynamic leak rate. The service life is determined by the dimensioning and stress of the bellow (Figure 17.31).

Feedthroughs with magnetic force coupling likewise show no dynamic leak rate, but are not suitable for all applications due to the magnetic stray fields and the limited transferable forces or torques (Figure 17.32).

Table 17.8 provides an overview of the properties and area of application of mechanical feedthroughs.

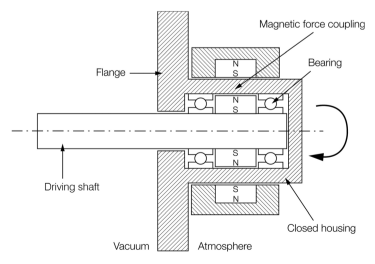


Figure 17.32 Magnetic force coupling.

Table 17.8 Qualities and areas of application of mechanical feedthroughs.

Sealing principle	Movement		Typical	Pressure	Materials	Outgassing	Particle
	Rotary	Translatory	leak rate (Pa ℓ s ⁻¹)	range	exposed to vacuum	behavior	generation
O-ring	X	Х	10 ⁻⁷ static 10 ⁻³ dynamic ally	LV Upper HV	Aluminum stainless steel elastomers	Permeation Leak	Very high
Magnetofluid seal	X		5×10^{-7}	Lower HV	Stainless steel Magnetofluid (on the basis of synthetic oil or PFPE)	Volumes Steam pressure synthetic oils and PFPE: 1×10^{-8} Pa	Slight
Diaphragm bellow (spring bellow)	X	X	<10 ⁻⁸	HV UHV XHV	Stainless steel	Very low	Very low
Magnetically coupled force transmission	X	X	<10 ⁻⁸	HV UHV XHV	Stainless steel MoS ₂ PTFE Sinter bronze PEEK Ceramics	Very low	Low

For complex movements, different mechanical feedthroughs are often combined (Figures 17.33 and 17.34).

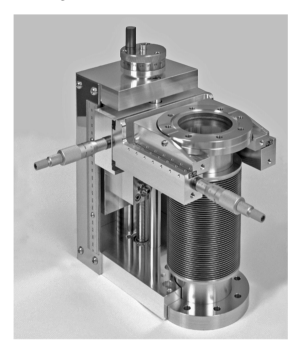


Figure 17.33 Diaphragm bellow sealed XYZ adjustment unit.

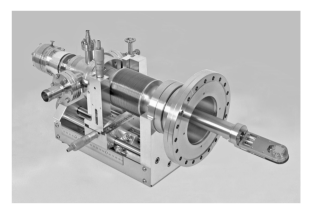


Figure 17.34 Five-axis sample manipulator with integrated sample heating and sample cooling.

17.4.3 Valves

Valves are feedthroughs for gases that, according to demand, can admit, regulate, throttle, or interrupt the gas flow between two volumes. They are placed between two or more systems with different pressures. Conductance, leak rate, number of the guaranteed load cycles, and media-exposed materials characterize the valves in vacuum technology. A valve consists of a valve case, valve seal, mechanical movement feedthrough, and an external drive (mechanically, electrically, or pneumatically). The basic construction of a valve is shown in Figure 17.35.

Many valve types are used in vacuum technology. In Table 17.9, the most current valves are shown with their essential pressure ranges, sealing principles, and materials that determine their application characteristics.

Valves can be used either for opening and closing or for regulating. The actual conductance of valves has a decisive influence on processes and on process times, which is why knowledge of them is of great importance. With the

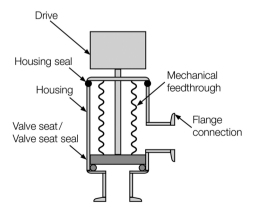


Figure 17.35 Schematic presentation of a right-angle valve.

Table 17.9 Valve models with pressure ranges and typical application possibilities.

Valve model/flange offsetting	Housing material	Seals		Pressure range	Max. tempera- ture (°C)	Qualities and Applications	
		Case	Valve seating	Movable	92	ture (c)	
Angle valve 90°	SS/aluminum	Elaston	er	- Hydroformed bellow- Edge welded bellow- O-ring sealed/slide feedthrough	HV	150	Open, closed intermediate positionsSeparation/connection of two systems
	SS	Metal	Elastomer	· ·	UHV	200	,
		Metal			UHV	200	
In-line valve 180°	SS/aluminum	Elaston	ner	- Hydroformed bellow - Edge welded bellow - O-ring sealed/slide feedthrough	HV	150	 Open, closed intermediate positions Separation/connection of two systems No free optical path
	SS	Metal	Elastomer		UHV	200	1 1
		Metal			UHV	200	
Butterfly valve 180°	SS/aluminum	Elaston	ner	O-ring sealed/slide feedthrough	HV	150	 Open, closed intermediate positions Throttling of pumps Downstream regulation Low space requirements
Gate valve 180°	SS/aluminum	Elaston	ner	- Hydroformed bellow - Edge welded bellow - O-ring sealed/linear feedthrough	HV	150	- Open, closed intermediate positions - Throttling of pumps - Downstream regulation - Low space requirements - Transfer valve

		Metal Elastomer		UHV	200	
		Metal		UHV	200	
Ball cock 180°/90°	SS Brass	Elastomer/PTFE	Sphere seal	GV	150	- Open, closed - Separation/connection of two systems - Two-way, three-way
Needle valve 180°/ 90°	SS/aluminum	Elastomer Metal Elastomer Metal	Conical Needle/round opening	HV	150	- Intermediate positions, open - Introduction of process gases - Ventilation - Only small nominal sizes
Diaphragm valves 180°/90°	SS/aluminum	Elastomer	Diaphragm seal	HV	120	- Open, closed intermediate positions - Separation/connection of two systems - Gas metering - Compact
Mass flow controller 180°	SS	Elastomer Metal	- Diaphragm seal - Needle valve	HV	50	- Open, (closed), intermediate positions - Upstream regulation - Introduction of process gases
Daving rulus 00°	SS		Diamburana anal	HV	200	On an aloned intonnedi
Dosing valve 90°	22	Metal	Diaphragm seal	пv	200	- Open, closed intermediate positions - Introduction of process gases - Fine actuation
				UHV		
				UHV		

application of an angle valve, the effective pumping speed of a pump can be reduced by up to 50% in comparison with the application of an in-line valve of comparable geometrical dimensions.

Example 17.2

Comparison of a DN63CF angle valve and a DN63CF gate valve. A turbomolecular pump with a nominal pumping speed of 300 ℓ s⁻¹ is connected with a chamber through a valve. An angle valve or a gate valve has to be selected. Both have a nominal size of DN 63CF. The effective pumping speed arises from

$$\frac{1}{\mathsf{S}_{\mathsf{eff}}} = \frac{1}{\mathsf{S}_{\mathsf{TMP}}} + \frac{1}{L_{\mathsf{Valve}}} \,.$$

The following differences arise from the calculation of the conductance for inline valves (long pipe in molecular flow) and for angle valves (90° bent pipe) (Figures 17.36 and 17.37):

	Straight pipe/in-line valve	Pipe elbow/angle valve
Formula:	$L \sim \frac{D^3}{I}$	$L \sim \frac{D^3}{2SL + 0.7D}$
Conductance:	524 ℓs^{-1}	$160 \ \ell \ s^{-1}$
S _{eff} :	190 ℓ s ⁻¹	$104 \ \ell \ s^{-1}$

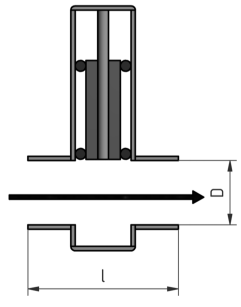


Figure 17.36 In-line valve.

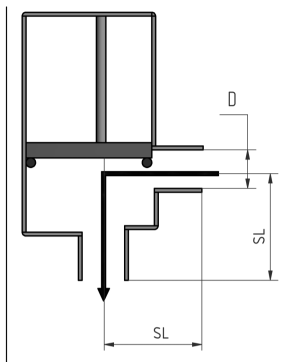


Figure 17.37 Angle valve.

To prevent turbulences, the conductance should be slowly increased. This applies to venting as well as to pumping of sensitive systems. Prevention of turbulence reduces stirring up of particles and dust or the destruction or misalignment of sensitive components. An undesirable condensation can also be counteracted through slow venting and/or evacuating.

Valves can be controlled mechanically (handwheel, lever, hexagon head), electrically (electromagnetically NC/NO, step motor-controlled), or electropneumatically (single-acting NC/NO, double-acting). The abbreviation "NC" stands for normally closed (in rest position) and "NO" for normally open (in rest position).

The driving mode is decided by the maximum operating temperature, the demanded automation, the controller used, and the controllability of the pressure (Table 17.10).

Manually operated valves are frequently used in laboratory equipment. Through the manual drive, it is possible to apply intermediate positions and to adjust a pressure or gas flow manually.

Valves with pneumatic drive are optimal for a partial automation or full automation. It is possible to control the vacuum valves through a valve island or through a separate control valve. In this instance, attention is to be paid to the fact that there are only two states (open/closed). In single-acting pneumatic andelectromagnetic drives, the valves are brought into the neutral position

Driving mode	Possible positions	Temperature (°C)	Automation	Pressure regulation
Manual	Any	<200	_	++
Pneumatic	Open/closed	50-200	+	_
Electropneumatic	Open/closed	50	++	_
Three-position pneumatic	Open/closed inter- mediate position	50-200	+	+
Electromagnetic	Open/closed	50	++	_
Step motor	Intermediate positions limited by step number	50	++	++

Table 17.10 Driving mode possibilities and operating conditions for valves.

through a return spring in the drive. According to construction, the valve is therefore automatically closed (NC) or open (NO) in the pressureless/deenergized state. In contrast, double-acting actuators consist of two pressure chambers. For both movements (open or closed), the actuator must be supplied with air pressure. As a result, an automatic reset is not provided with this drive.

Electromagnetic valves can be used in partially or fully automated controls and at places relevant to safety. The valve plate is connected with an armature that can be moved by a magnet coil. In the de-energized state, the valve plate moves back to its initial position through a return spring in the armature. With this, for example, these drives allow for a safety ventilation or the separation from fore-vacuum systems in case of a power failure.

Step motor controllers are used in dosing or gate valves. The valve can be operated by this electrical drive in small steps and adjusted to a desired conductance or flow.

Pressure Regulation

The pressure in a vacuum system results from the equilibrium between the inflowing gas and the pumping speed of the pump. Vacuum valves can influence the amount of the inflowing gas as well as the outflowing gas and, with this, the equilibrium pressure. This occurs in combination with pumps and sensors.

When the gas inflow is regulated in the chamber, we speak of an upstream regulation, while in contrast when the pumping speed of the pump is changed through a valve a downstream regulation exists. Different systems find application depending on the response time and sensitivity. With these regulations, the process gas composition and decomposition phenomena as well as the gas type-dependent pumping speed of pumps are to be considered.

A change of the effective pumping speed through a conductance change to the pump serves the rather rough regulation. Gate valves are ideal for the throttling of pumps because in open position the pumping speed is barely influenced by the high conductance.

According to the size of the system, pressure range, and pumping speed, needle valves, that is, valves with small variable apertures or conductances as well as with capillaries, come into use for upstream regulation.

Dosing valves are suitable for very sensitive applications due to their smooth actuation and low conductance. The regulation through valves is no longer sufficient for an extremely high accuracy and a stable process gas composition. Mass flow controllers (MFCs) are used for this. The mass flow rate of the gas is measured directly in the device through a thermal measuring system. It is to be noted that the MFCs mostly cannot close completely, and therefore, an additional closing valve is necessary.

17.4.4

Electrical Feedthroughs

Electrical feedthroughs provide the transfer of electric energy into the vacuum or out of it.

They consist of one or several conductors, a dielectric, and a metal housing. These components must be joined with each other through a hermetic seal so that a leak rate that is sufficiently low for the respective application is ensured.

An electrical feedthrough is hermetically connected with its metal housing to a vacuum flange or the vacuum chamber wall through a welded joint, but sometimes also by means of a bulkhead fitting or a screw-in thread. The dielectric serves as an electric insulation of the contacts of the ideally grounded vacuum chamber (Figure 17.38).

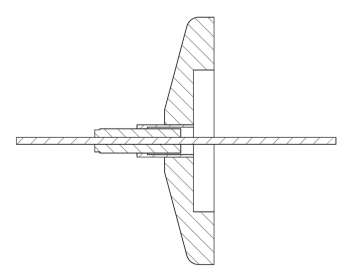


Figure 17.38 Cross section of a BNC coaxial feedthrough with Al₂O₃ insulation and KF25 flange.

	Al ₂ O ₃	Glass	Glass ceramics	Resin casting
Specific electrical resistance @ $20 ^{\circ}\text{C} (\Omega \text{cm})$	>10 ¹⁴	>10 ¹⁴	>10 ¹⁴	>10 ¹⁴
Specific electrical resistance @ 250° C (Ω cm)	>10 ¹²	$10^9 - 10^{11}$	$10^9 - 10^{11}$	-
Max. baking out temperature (°C)	350-450	200-250	350-450	80–150
Typical leak rate (Pa ℓ s ⁻¹)	$<1 \times 10^{-7}$	$<1 \times 10^{-7}$	$<1 \times 10^{-7}$	10^{-5} to 10^{-7}
Outgassing behavior	Very good	Very good	Very good	Poor to moderate

Table 17.11 Vacuum suitable insulation materials

Paschen's law is to be followed in the selection of a suitable electrical feedthrough (Figure B.15). It describes the dependence of the breakdown voltage in a gas atmosphere in relation to the gas type, the pressure, and the electrode gap. For a typical conducting feedthrough distance in the order of magnitude of 1 mm, a minimal breakdown is shown for air in the pressure range of 100-1000 Pa, so that a voltage breakdown can already occur with a few hundred volts (Figure B.15). With higher pressures, the breakdown voltage rises again linearly, and with lower pressures, even exponentially. The maximum voltage load specified by manufacturers for an electrical feedthrough is primarily only valid for a residual gas pressure of $p < 10^{-2}$ Pa.

Insulating Materials

Four groups of insulation materials are commonly used in vacuum technology. Table 17.11 provides an overview of these materials and their suitability for different areas of application.

Electrical Feedthroughs with Aluminum Oxide Ceramics

In case of very high requirements for insulation resistance, outgassing behavior, and hermeticity as well as bakeabilty, aluminum oxide ceramics (>94% Al₂O₃) have primarily proved themselves as a suitable dielectric.

Many metals that are commonly used for housings and conductors - for example, stainless steel and copper - show a very divergent thermal expansion behavior compared with aluminum oxide ceramics. To minimize the mechanical stresses thereby appearing at high or low temperatures and to avoid any damage, additional metal sleeves are required as a "tension buffer" whose thermal coefficient of expansion is adapted to that of the ceramics. The individual components are material bonded in a complex multistage joining process (high-temperature brazing).

Electrical feedthroughs with ceramic insulation are bakeable up to a minimum of 350 °C, some even to 450 °C, and, with an adequate cleaning, can reach a total



Figure 17.39 HV feedthrough with wave-shaped ceramics.

outgassing rate of $<1 \times 10^{-8}$ Pa ℓ s⁻¹ cm⁻² by which its range of application goes up to the deep UHV.

Up to 300 °C, the specific electrical resistance of aluminum oxide ceramics only decreases by two decades so that electrical feedthroughs with ceramic insulation are suitable under temperature load as well as for the transfer of a high power (> $10^{14}\Omega$ cm at 25 °C, > $10^{12}\Omega$ cm at 300 °C, > $10^{8}\Omega$ cm at 600 °C) [41]. For cryogenic applications, aluminum oxide ceramics are also well suited as an insulation material if the maximum cooling rate is not exceeded.

Because ceramics can be produced in many different shapes, the design of an electrical feedthrough with ceramic insulation can be accordingly adapted to nearly any application and is thereby better suited for high voltage than other insulation materials (Figure 17.39–17.41).

Electrical Feedthroughs with Glass or Glass Ceramics

In the use of glass or glass ceramics as a dielectric, the seal is normally not material bonded, but is based merely on mechanical tension (compression). This requires a careful choice of materials with adapted thermal expansion coefficients, which limits the eligible housing materials and conducting materials. With glass or glass ceramics as a dielectric, very good values are achieved concerning insulation resistance (<250 °C), hermeticity, and outgassing.



Figure 17.40 Multiple feedthrough with glass ceramic insulation.



Figure 17.41 Resonant beam position monitor with electrical feedthrough.

Electrical feedthroughs with glass and glass ceramic insulation are bakeable up to at least 200 °C, with some special materials even up to 450 °C. Total outgassing rates of $<1\times10^{-8}\,\mathrm{Pa}~\ell~\mathrm{s}^{-1}~\mathrm{cm}^{-2}$ can also be achieved with glass/glass ceramic insulated electrical feedthroughs so that they are also applicable up to the low UHV. Nevertheless, unlike with aluminum oxide ceramics, the specific resistance quickly decreases with the rising temperature by several orders of magnitude (> $10^{14}\,\Omega$ cm at 25 °C, between 10^9 and $10^{11}\,\Omega$ cm at 250 °C, between 10^7 and $10^9\,\Omega$ cm at 350 °C) [42]. Thus, electrical feedthroughs with glass/glass ceramics can be sufficiently well baked out, but they should not be heated up to more than $150-200\,^{\circ}\mathrm{C}$ during the operation and only moderate power should be transferred. For cryogenic applications, glass or glass ceramics are also well suited as an insulation material if the maximum cooling rate is not exceeded.

Glass and glass ceramics offer advantages as a result of their less costly joining processes with metals, particularly in case of electrical feedthroughs with a high conducting path density such as the sub-D feedthrough or similar multiple feedthroughs.

Electrical Feedthroughs with Resin Casting

A favorable alternative for lower requirements, for example, for rough or fine vacuum applications, can be electrical feedthroughs that use a hardening casting compound, often an epoxy resin, as dielectric.

Current casting compounds compared with ceramics, glass, and glass ceramic insulating bodies show disadvantages in the outgassing behavior, with the hermeticity and the maximum operating and baking out temperature. Degeneration

processes over periods of months or years can also have negative effects on the functionality [43].

Electrical Parameters and Their Influence on the Design of Feedthroughs

According to application, other electrical parameters in the transfer of electrical energy are central. Different requirements for the feedthrough with effects on the design can arise thereby.

Current

Fundamentally, the current-carrying capacity depends on the conductor material and conductor diameter. The most common conductor materials are copper, nickel, stainless steel, molybdenum, and some special nickel alloys. The specified upper limit for the current load of an electrical feedthrough refers to a specific, maximum permissible heating defined by the manufacturer. With very high current, waveguides for liquid cooling are also sometimes used.

In contrast, if very low electrical currents are to be detected, the use of coaxial feedthroughs (10⁻⁹ to 10⁻¹² A) and triaxial feedthroughs (avoidance of internal leakage currents up to the subpicoampere range) is recommended because these offer a shielding against electromagnetic interference due to their design (Figure 17.42-17.44).

Voltage

The dielectric strength of a feedthrough is determined by the distance of the potential-carrying parts, the potential difference, the specific electrical resistance of the dielectric, the "creepage currents" along the surface of the dielectric, and the vacuum-sided gas pressure and gas composition.

The disruptive voltage – related to a breakthrough through the dielectric – is determined by the material and the temperature of the dielectric. In case of disruptive voltage discharges as a result of creepage currents along the surface, the disruptive voltage primarily depends on the creepage path between the electrodes. It can be extended by the design of the dielectric, for example, in wave form or with grooves (Figure 17.39). In addition to this, the availability of ionizable material and, with this, the surface cleanliness of the dielectric play a large role. The surface should be as flat as possible and therefore well cleanable and as



Figure 17.42 High-current feedthrough for the transfer of current.



Figure 17.43 Triaxial feedthrough for the measurement of very low currents.



Figure 17.44 Sub-D feedthroughs in CF flange. (Source: VACOM.)

clean as possible. The transfer of the so-called thermotension demonstrates a specific feature by which the local temperature in the vacuum is determined based on the Seebeck effect. Thermocouple feedthroughs (Figure 17.50), which consist of two different conducting materials permitting a direct connectivity to standardized measuring lines for temperature measurement on the air side as well as at the vacuum side, are widely used in industry and research. Thermocouple feedthroughs ensure an undistorted transfer of measuring signals through the chamber wall.

Frequency

If alternating voltage signals ("AC") are to be worked with, the use of coaxial feedthroughs in the MHz to GHz frequency range is recommended, in particular with a $50\,\Omega$ impedance adjustment. With the outer conductor or "shield," the coaxial design contains a shielding of the live internal conductor against external electromagnetic interference. The $50\,\Omega$ impedance adjustment ensures a low damping and low reflection connection to the feedthrough to the 50Ω wiring standard so that sensitive AC signals can be transferred largely undistorted through a chamber wall. Most coaxial feedthroughs use glass/glass ceramics as dielectric due to the low dielectric constant.

17.4.5

Optical Feedthroughs [44]

Electromagnetic radiation, from the infrared (IR) through the visible (VIS) and up to the ultraviolet and extreme ultraviolet (UV/EUV) range, is taking on greater importance in vacuum. With optical feedthroughs (viewports, Figure 17.45, and optical fiber feedthroughs, Figure 17.46), the transfer of electromagnetic radiation from the atmosphere into the vacuum and vice versa is being realized in different optical qualities.

In practice, it is about processes that are induced by electromagnetic radiation as well as about the processes in which electromagnetic radiation is generated. In addition, the detection of electromagnetic radiation is an important means for the process observation and process control.

An optical feedthrough basically consists of the optical material (window or fiber) and a frame that is usually metallic. These materials with very different material properties (e.g., thermal expansion coefficient) must be hermetically joined with suitable connecting materials. The optical materials have the properties that are necessary to lead the electromagnetic radiation into a vacuum chamber or out of it as undisturbed as possible and in the optically required quality. Therefore, values for attenuation, absorption, dispersion, and plane parallelism, among others, are defined. The defined optical properties can be changed in their quality during the joining process (Section 17.3).

The technical characterization of a viewport usually occurs according to the intended purpose. Viewports that only serve the visual observation are



Figure 17.45 Viewport in CF flange.



Figure 17.46 Optical fiber feedthroughs in CF flange.

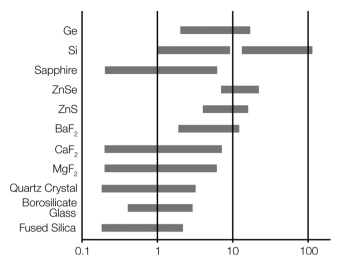


Figure 17.47 Transmission ranges of common optical materials.

specified with the type of the optical material (ordinarily borosilicate glass), the view diameter, the helium leak rate, and the operating temperature/heating rate.

The properties/quality of the optical material for viewports that meet higher optical requirements are further specified. Common parameters are

- transmission range (see Figure 17.47),
- refractive index,
- surface flatness.
- surface defects,
- parallelism.

These parameters describe, among others, effects such as reflection, absorption, dispersion, and image defects.

Viewports for optical applications are often provided with a coating in order to optimize the reflection or the transmission according to the use. With coated viewports, the coated diameter also matters alongside the view diameter. The transmission range of important optical materials is shown in Figure 17.47.

A viewport normally consists of three components: the flange (installation of the viewport into the chamber), the optical material (transfer of the radiation), and the connecting element. The connection between the optical material and the flange has to be vacuum-tight, mechanically stable (pressure difference >1 bar), and low in outgassing. In addition, it must provide for the equilibrium of the mechanical stresses, for example, through thermal expansion (Table 17.12). It can be

- elastomer sealed,
- bonded,

Flange material	Thermal expansion coefficient (µm m ⁻¹ K ⁻¹)	Optical material	Thermal expansion coefficient (μm m ⁻¹ K ⁻¹)	Weld ring material	Thermal expansion coefficient (µm m ⁻¹ K ⁻¹)
Stainless steel 1.4301	16–18 [45]	Borosilicate	7.1 [46]	Kovar	4.9-6.2 [47]
Stainless steel 1.4305	16–18 [48]	Fused silica	0.55 [46]	Indium	32.1 [49]
Stainless steel 1.4404	16–18 [50]	Sapphire	5.0-5.6-7 [46]	Titanium	8.6 [51]
		Calcium fluoride	18.85 [46]		

Table 17.12 Thermal expansion coefficients of metals and optical materials.

- soldered, or
- fused.

As to not overburden this junction, the use of soft seals (e.g., of annealed copper seals) is recommended for the flange mounting of the viewports. When designing a viewport, attention must be paid to the fact that the optical material withstands reliably the mechanical load of 1 bar. An evenly loaded circular plate with clamped edge follows the equation

$$\sigma = 0.48 \ \Delta p \ r^2/d^2,$$

where σ is the mechanical tension that must not exceed the modulus of rupture, Δp is the pressure difference (as a rule, 1 bar), r is the radius of the clamped plate (approximately the view diameter), and d is the viewport thickness [52].

Optical Fiber Feedthroughs

Optical signals can also be transported through fiber-optic cables (also called optical fiber) into a vacuum chamber. Monochromatic as well as polychromatic light can be guided precisely and flexibly.

Alongside the standard demands of vacuum technology (see Sections 17.2 and 17.3), an optical fiber feedthrough (OFF) for vacuum applications must meet the optical demands, which is why the feedthroughs are additionally characterized by insertion loss, reflection loss, and intrinsic loss per fiber length. The type of the fiber can be characterized by the core or expanded beam diameter or the mode field diameter. The fiber type must correspond with the feedthrough and with the connecting cable so that greater losses are prevented.

A fiber can be joined into a flange in complete length, which, however, is problematic for a failure-free operation due to the high rupture susceptibility of unprotected fibers. On the other hand, feedthroughs with plug connectors stand



Figure 17.48 Complete OFF.



Figure 17.49 Flange with OFF and wiring.

out due to a high flexibility and operational reliability. In addition to the optical fiber feedthrough, protected fibers are used as a cable on both the air side and the vacuum side. According to application, high requirements are made on the vacuum side for the low outgassing of the protective sleeve of the fiber. The overall balance of the attenuation is determined by the plug type as well as the quality of the plugs (Figures 17.48 and 17.49).

Common plug standards are

- FSMA plug (preferred for multimode fibers);
- FC plug (preferred for single-mode fibers)

These plug systems distinguish themselves through a high robustness and are also available in an entirely metal version. In addition, FC plugs are exceptionally precise and low in attenuation. Plugs according to the ST standard are less widespread, but are also available for vacuum technology applications.

17.4.6

Heat Supply and Dissipation

The transmission of heat occurs through thermal conduction, convection, and heat radiation. Because heat radiation is independent of substance, it is the only heat transport possible in pressure ranges <1 mPa. Thermal conduction and convection are dependent upon a substance and gas type, respectively. In the pressure range >1 mPa, the pressure dependence on thermal conduction and heat convection follows complex interrelationships (Sections 3.3 and 3.4). Because specific heat supply and dissipation are often relevant to the process in the vacuum, this dependence must be considered and also be controlled, which can be realized through separate operation devices or the entire process control system. In addition, baking out processes play a crucial role for the cleanliness in the vacuum, the avoidance of contaminations through decomposition, and the achievement of the HV, UHV, and XHV ranges. Heat supply and dissipation can only be specifically controlled if a measurement of temperature is possible at the points relevant to the process.

Temperature Measurement in the Vacuum

Temperatures in the vacuum are primarily measured by a thermocouple or pyrometer. The use of thermocouples is based on the Seebeck effect [53] with which an electric voltage originates in a closed circuit with different conductors at the contact points with different temperatures.

When choosing conducting materials for the suitable temperature range, attention is to be paid to the fact that the thermotension changes nearly linearly. Electrical as well as thermal insulation has an influence on the measuring accuracy and reproducibility.

In order to outwardly transport the thermovoltage of the measuring point in the vacuum, a suitable thermocouple feedthrough (Figure 17.50) is required for the type used. Here, the suitable conductors are externally connected to the fitting legs, whereas on the external side a regular cable is sufficient to determine the voltage in the measuring device. As described in Section 17.4.4, here, analog conditions are valid for the vacuum suitability of the feedthrough.

Pyrometers work on the fact that each body that has a temperature >0 K emits heat radiation. The intensity, the spectral distribution, and the position of the emission maximum are dependent upon the temperature of the material to be measured and the material itself. Based on the Stefan-Boltzmann law (Eq. (12.4)), the total radiation power is proportional to the fourth power of the



Figure 17.50 Thermocouple feedthrough.

absolute temperature. A pyrometer can directly measure the external temperatures of vacuum chambers and vacuum systems. In order to be able to carry out pyrometric measurements in the vacuum, optical feedthroughs such as viewports or optical fiber feedthroughs (Section 17.4.5) for the respected wavelength range are required. While with viewports a direct view contact must exist between the measured object and the pyrometer, with fiber optical feedthroughs the emitted heat radiation is coupled into the fiber by a collimator lens and guided to the pyrometer in a flexible path.

Heat Dissipation from the Vacuum

Process heat can be dissipated either integrally through the chamber wall or directly at the place of the origin.

The integral heat dissipation through the external chamber wall can be achieved either by cooling lines with a good thermal connection to the wall or by the use of double-walled chambers (Section 17.4.1). The temperature range and the dissipated heat quantity determine the construction design, the cooling liquid, and the flow velocity.

For the local heat dissipation, cooling liquids are commonly used and, if necessary, supported by Peltier elements [54]. A liquid feedthrough is required for cooling with cooling liquids. While with refrigerant temperatures at >0 °C, a simple pipe in the flange is sufficient (Figure 17.51), refrigerant temperatures at <0 °C with icing effects near the flange seal are to be avoided through the use of an insulating vacuum (Figure 17.52).

Baking Out Processes and Heat Supply

The heat supply into a vacuum system can occur through heat generation inside and outside the vacuum.

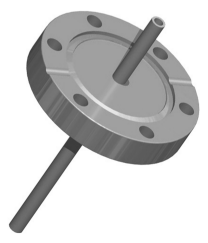


Figure 17.51 CF liquid feedthrough.



Figure 17.52 CF liquid feedthrough with insulating vacuum for liquid nitrogen.

With heat generation outside the vacuum, three proven possibilities are distinguished (Tab. 17.13):

- heating tapes (Figure 17.53),
- heating tents or heating boxes (Figure 17.54),
- heating collars (Figure 17.55).



Figure 17.53 Heating tapes.

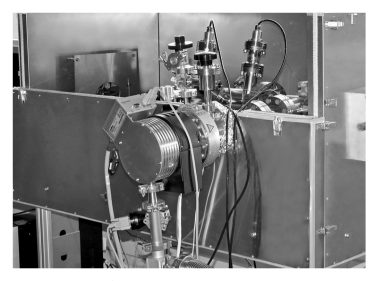


Figure 17.54 Heating box.

Heating tapes are flexibly applicable for various geometries. The rated powers usually lie between 20 and 250 W m⁻¹ and, according to implementation, they are suitable for temperatures of up to 900 °C. However, it is to be noted that heating tapes overheat themselves above decomposition temperature if they are not regulated. For the secure and failure-free operation of heating tapes, the minimum bending radius may not be undercut while, at the same time, the correct laying distance and the thermal insulation toward the air side are to be ensured. With the application of various heating tapes, different temperature ranges can be generated at several zones and temperature-sensitive components can be omitted.



Figure 17.55 T-piece with heating collar.

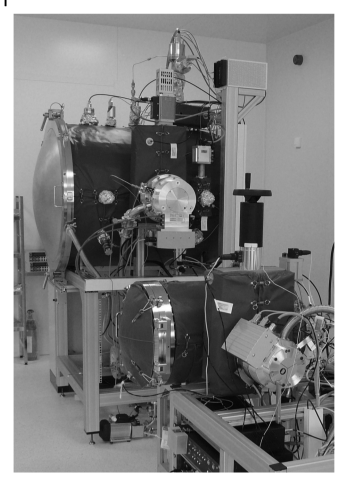


Figure 17.56 RGA with clean room applicable heating collars.

Heating tents and heating boxes dispose of inside heating surfaces with good external insulation and generate a constant temperature of the air around the chambers and equipment whose homogeneous distribution can be supported by internal ventilating fans. They allow no specific differences in temperature in various areas and require the temperature resistance of all included components.

Heating collars for chambers and components are tailor-made heating devices that heat directly on the surface and are thermally well insulated on the outside (Figure 17.56). They allow for different heating zones as well as the omission of temperature-sensitive components. The typically used materials are, for example, glass silk, silicone, PTFE, and aluminum-coated or silicone-coated materials. For the application in the clean room, there are silicone-free and low in particle solutions for temperatures up to $200\,^{\circ}\text{C}$.

Characteristic	Heating tape	Collar	Heating tent
Price	Reasonable	Cost-intensive	Medium costs
Flexibility	Flexible	Precise	Limited
Deployability	Elaborate	Simple	Continual construction and dismantling
Time involved	High	Slight (<50%)	Medium
Insulation	No	Integrated	Integrated
Combustion protection	No	Yes, through insulation	Yes, if insulated sufficiently
Temperature sensor	No	Optionally better adjustable	Only in the heating element
Clean room suitability	No	Possible	No
Several heating circles	Possible (complicated)	Yes	No
Power savings	_	Up to 75% in comparison with heating tapes	Not specified

Table 17.13 Overview of heating possibilities in vacuum technology.

The evaluation of the necessary heating capacity for the achievement of a certain temperature in a defined time follows the equation

$$P = K \frac{m \cdot c \cdot \Delta T}{\Delta t},$$

where P is the total power in watts, c is the specific heat capacity, m is the mass of the component to be heated, ΔT is the temperature difference (starting temperature – end temperature), Δt is the desired heating duration in seconds, and *K* is the correction factor for heat losses and thermal bridges.

Typical heat capacities c for aluminum amount to $897 \,\mathrm{J\,kg^{-1}\,K^{-1}}$, copper, 385 J kg⁻¹ K⁻¹, and stainless steel, 500 J kg⁻¹ K⁻¹. Due to additional heat losses and thermal bridges difficult to quantify, to be on the safe side, the ascertained value should be increased by 50%.

The heat generation in vacuum occurs, for example, through electrically operated ceramic, graphite, stainless steel, tantalum, or titanium heaters. The heatgenerating materials are selected according to the necessary temperatures and, in all instances, must be low in outgassing.

Abbreviations

ACN	Acrylonitrile
AES	Atomic emission spectroscopy
AFM	Atomic force microscopy

Ethylene propylene diene monomer rubber **EPDM**

EUV Extreme ultraviolet light

FC Fiber connector Perfluoro rubber **FFKM** FKM Fluoro rubber

FSMA Fiber subminiature version A MVQ/VMQ Methyl vinyl silicone rubber **NBR** Acrylonitrile butadiene rubber Optical fiber feedthrough OFF PEEK Polyether ether ketone PFPE Perfluoropolyether

PSE Periodic system of elements PTFE Polytetrafluoroethylene

OMS Quadrupole mass spectrometer

RGA Residual gas analysis

Scanning electron microscopy SEM SIMS Secondary ion mass spectrometry TDS Thermal desorption spectroscopy

Total mass loss TML UCV Ultraclean vacuum

XPS X-ray photoelectron spectroscopy

References

- 1 CEROBEAR GmbH, CERABEAR ceramic bearing technology, Herzogenrath 2014, http://www.cerobear.de/fileadmin/images 3/ Prospekte/ CEROBEAR_company_brochure. pdf
- 2 BODYCOTE WÄRMEBEHANDKLUNG GMBG, Kolsterising Corrosion resistant surface hardening of austenitic stainless steel, Nürnberg 2005, internet.bodycote. org/kolsterising/brochures/147-302 body kolst_rd_gb_finr.pdf
- 3 Posth, O., Wunderlich, H., Flämmich, M., and Bergner, U. (2012) Reinigung von Vakuumbauteilen für UHV- und UCV-Anwendungen. Vak. Forsch. Prax., 24 (6),
- 4 U. Hahn and K. Zapfe: Richtlinien für UHV-Komponenten bei DESY, Technische Spezifikationen Nr.: Vakuum 005/2008 Version 1.6/ 22.09.2010 https:// mhf-e-wiki.desy.de/images/c/c6/Vakuum_ 005_DESY_UHV_Richtlinien_1-5_final_ stamp eng.pdf

- 5 GSI (2011) Testing the Cleanliness of Cryostat Insulation Vacuum Components. Technical Guideline. Available at http:// indico.gsi.de/getFile.py/access? resId=20&materialId=0&confId=1420.
- 6 Taborelli, M. (2007) Cleaning and Surface Properties. CERN. Available at http://cds. cern.ch/record/1047073/files/p321.pdf.
- 7 Henglein, A. and Gutierrez, M.J. (1993) Sonochemistry and sonoluminescence: effects of external pressure. J. Phys. Chem., 97, 158.
- 8 Kao, P.S. and Hocheng, H. (2003) Optimization of electrochemical polishing of stainless steel by grey relational analysis. J. Mater. Process. Technol., 140, 255-259.
- 9 Anders, H. (2010) Prozessabhängige Oberflächenqualität plasmapolierter Werkstücke aus Chrom-Nickel-Stahl. Diplomarbeit BA Glauchau.
- 10 W. Rähse, Industrial Product Design: Materials for the Machinery, Wiley Online

- Library, ChemBioEng Rev 2014, 1, No. 3, p. 6-10
- 11 Middleman, K.J., Herbert, J.D., and Reid, R.J. (2007) Cleaning stainless steel for use in accelerators - phase 1. Vacuum, 81 (6), 793-798.
- 12 Bornmyr, A. and Holmberg, B. (1995) Handbook for the Pickling and Cleaning of Stainless Steel, AvestaPolarit Welding.
- 13 van Os, T. (2005) Ceramic Bead Blasting vs. Glass Bead Blasting. Vecom Group Technical Bulletin Number 2005/07.
- 14 Sherman, R., Grob, J., and Whitlock, W. (1991) Dry surface cleaning using CO₂ snow. J. Vac. Sci. Technol. B, 9 (4), 1970-1977.
- 15 Tuzi, Y., Kurokawa, Y., and Takeuchi, K. (1993) Effect of bake-out on the adsorption kinetics of gases in a vacuum chamber. Vacuum, 44 (5-7), 447-449.
- 16 Westerberg, L., Hjörvarsson, B., Wallén, E., and Mathewson, A. (1997) Hydrogen content and outgassing of air-baked and vacuum-fired stainless steel. Vacuum, 48 (7-9), 771-773.
- 17 Jones, A.W., Jones, E., and Williams, E.M. (1973) Investigation by techniques of electron stimulated desorption of the merits of glow discharge cleaning of the surfaces of vacuum chambers at the CERN intersecting storage rings. Vacuum, 23 (7), 227 - 230.
- 18 Vigh, J.R. (1985) UV/ozone cleaning of surfaces. J. Vac. Sci. Technol. A, 3 (3), 1027-1034.
- 19 DIN 8593-0:2003-09 (2003) Fertigungsverfahren Fügen - Teil 0: Allgemeines; Einordnung, Unterteilung, Begriffe, Beuth Verlag, Berlin.
- 20 DIN 1910-100:2008-02 (2008) Schweißen und verwandte Prozesse - Begriffe - Teil 100: Metallschweißprozesse mit Ergänzungen zu DIN EN 14610:2005, Beuth Verlag, Berlin.
- 21 Dorn, L. (2007) Hartlöten und Hochtemperaturlöten, Expert Verlag, Renningen.
- 22 Roth, A. (1994) Vacuum Sealing Techniques, American Vacuum Society Classics, AIP Press, New York.
- 23 DIN EN ISO 5817:2014-06 (2014) Schweißen - Schmelzschweißverbindungen an Stahl, Nickel, Titan und deren

- Legierungen (ohne Strahlschweißen) Bewertungsgruppen von Unregelmäßigkeiten, Beuth Verlag, Berlin.
- 24 DIN EN ISO 15607:2004-03 (2004) Anforderung und Qualifizierung von Schweißverfahren für metallische Werkstoffe – Allgemeine Regeln (ISO 15607:2003), Beuth Verlag, Berlin.
- 25 DIN EN ISO 17672:2010-11 (2010) Hartlöten - Lote, Beuth Verlag, Berlin.
- **26** DIN 8593-8:2003-9 (2003) Fertigungsverfahren Fügen - Teil 8: Kleben; Einordnung, Unterteilung, Begriffe, Beuth Verlag, Berlin.
- 27 NASA: MAPTIS Materials and Processes Technical Information System NASA's Authorized Guide to Materials. http:// maptis.nasa.gov/
- 28 ESA/ESTEC/TOS-QM, Materials and Processes Division: SME INITIATIVE COURSE MATERIALS, SME1-_Intro.ppt. http://esmat.esa.int/SME/sme.html Class 10 Materials Group Adhesives, Coatings, Varnishes: http://esmat.esa.int/Services/ Preferred_Lists/Materials_Lists/ materials lists 9.html
- 29 Patentschrift DE 10 2008 019 681 B4. Schenk, C.; Risse, S.; Harnisch, G.; Peschel, T.; Bauer, R. Mittels aerostatischer Lagerelemente geführter Tisch für Vakuumanwendungen
- **30** DIN ISO 7619-1:2012-02 (2012) Elastomere und thermoplastische Elastomere – Bestimmung der Härte – Teil 1: Durometer-Verfahren (Shore-Härte) (ISO 7619-1: 2010), Beuth Verlag, Berlin.
- 31 DIN 53536:1992-10 (1992) Prüfung von Kautschuk und Elastomeren; Bestimmung der Gasdurchlässigkeit, Beuth Verlag,
- 32 DIN 28404:1986-10 (1986) Vakuumtechnik; Flansche; Maße, Beuth Verlag, Berlin.
- 33 ISO 1609:1986-03 (1986) Vacuum technology - Flange dimensions, Beuth Verlag, Berlin.
- 34 DIN 28403:1986-09 (1986) Vakuumtechnik; Schnellverbindungen; Kleinflansch-Verbindungen, Beuth Verlag, Berlin.
- 35 ISO 2861:2013-05 (2013) Vacuum technology - Dimensions of clamped-type

- quick-release couplings, Beuth Verlag,
- **36** ISO 3669:1986 (1986) Vacuum technology - Bakable flanges - Dimensions. Withdrawn. Available at http://www.iso. org/iso/catalogue_detail.htm? csnumber=9124.
- 37 ISO/TS 3669-2:2007-09 (2007) Vacuum technology - Bakable flanges - Part 2: Dimensions of knife-edge flanges, Beuth Verlag, Berlin.
- 38 Bach, F.-W., Laarmann, A., Möhlwald, K., and Wenz, T. (2004) Moderne Beschichtungsverfahren, 2. Auflage, Wilev-VCH Verlag GmbH.
- 39 Wiegand, H., Kloos, K.-H., and Thomala, W. (2007) Schraubenverbindungen (Untertitel: Grundlagen, Berechnung, Eigenschaften, Handhabung), 4. Auflage, Springer.
- **40** TÜV e.V. (2011) AD 2000-Regelwerk, TÜV e.V., Berlin.
- 41 Barat Ceramics GmbH: Ceramics meet Precision, Materials - Material Datas -Surfaces - Certificates, Auma http://www .barat-ceramics.com/files/12CBA97633D/ Werkstoffe.pdf Accessed 01.02.2016
- 42 Technical Glasses. Physical and Technical Properties, Schott AG, Mainz, 2014; Dow Corning Insulation Glass Manual, Dow Corning Corporation, Midland, 2011.
- **43** Loctite (2014) The Adhesive Sourcebook, vol. 18, Henkel Corp., Rocky Hill.
- 44 Rietmann, T., Gottschall, S., Sahib, Ch., Flämmich, M., and Bergner, U. (2013)

- Optik braucht Vakuum. Vak. Forsch. Prax., 25 (1), 18-26.
- 45 Deutsche Edelstahlwerke (2008) Datenblatt 1.4404.
- 46 Matthews, K. (ed.) (2008) The Crystran Handbook of Infra-Red and Ultra-Violet Optical Materials, Crystran Ltd, Poole.
- 47 Parmaco (2009) Datenblatt Kovar.
- 48 Deutsche Edelstahlwerke (2008) Datenblatt 1.4305.
- 49 WebElements: the periodic table on the WWW, http://www.webelements.com/ indium. Mark Winter, The University of Sheffield and WebElements Ltd, UK, 1993-2016.
- 50 Deutsche Edelstahlwerke (2008) Datenblatt 1.4301.
- 51 WebElements: the periodic table on the WWW, http://www.webelements.com/ titanium. Mark Winter, The University of Sheffield and WebElements Ltd, UK, 1993-2016.
- 52 Grote, K-.H. and Feldhusen, J. (eds) (2004) Dubbel: Taschenbuch für den Maschinenbau, Springer, Berlin.
- 53 Bergmann, L. and Schäfer, C. (1971) Lehrbuch der Experimentalphysik, Bd. 2, Elektrizitätslehre und Elektromagnetismus, 6th edn, Gruyter, Berlin, Chapter III 29.
- 54 Apertet, Y., Ourdane, H., Glavatskaya, O., Goupil, C., and Lecoeur, P. (2012) Optimal working conditions for thermoelectric generators with realistic thermal coupling. Europhys. Lett., 97, 28001.

18

Operating Vacuum Systems

Dr. Karl Jousten¹ and Uwe Meissner²

18.1

Electronic Integration of Vacuum Systems

18.1.1

Control by Means of Process Sensors and Automated Data Processing

18.1.1.1 Requirements and Applications

Vacuum process systems and their components such as gas flow regulators, vacuum pumps, vacuum monitoring equipment, plasma generators, and evaporator units provide large amounts of data for monitoring and controlling processes in vacuum technology. A number of special monitoring solutions are also available for reliable supervision of processes.

For electronic monitoring, the main components of a vacuum process chamber are integrated into a bus system. It is necessary to monitor every process-critical parameter. Integrated sensors should transmit the required information not only to the system's control but also to higher level systems (Figure 18.1).

There is a continuous demand for more productive, more economical, and more energy efficient electronic devices. This requires a continuous reduction of the pattern size on wafers of larger size and challenges the vacuum systems for continuous improvement.

For economic reasons, chip manufacturers must reduce times to mature yield (85% in processors, 90% in memory components). This requires continuous process monitoring and automated control. Considering approximately 40 mask steps per wafer, the complete production process can take several weeks. A leading processor manufacturer in Dresden, Germany, has therefore developed automated processes for production and material handling in his production site, coming close to an autonomous factory. Apart from appropriate software, this approach requires sophisticated procedures for process monitoring as well as sensors.

¹Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

²MKS Instruments, Zur Wetterwarte 50 Haus 337/F, 01109, Dresden, Germany

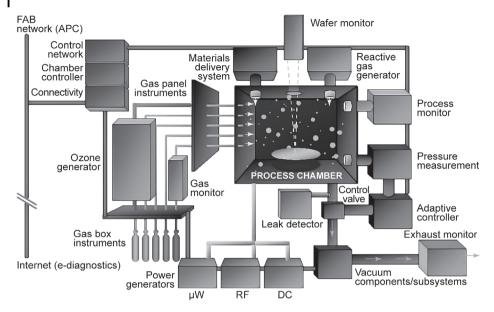


Figure 18.1 Possible components of a vacuum process chamber. Processes operated determine installed components.

In other areas as well, sophisticated process monitoring is a prerequisite, for example, for constant coating thicknesses across the entire surface in large-area glass coating and solar panel manufacturing.

Sensor and measuring principles that have prevailed for the various tasks in practical applications are listed in Table 18.1.

Table 18.1 Process sensors/measurement principles and application	Table 18.
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Process sensors/ measurement principles	Processes	Applications
Mass spectrometer/RGA	PVD/sputtering, ion implantation	Monitoring of vacuum conditions Residual and process gas composition End-point determination Outgassing behavior of substrates
Plasma emission spectrometer	PECVD	End-point determination
	Plasma etching	Plasma analysis
Infrared (FTIR/NDIR) spectrometer	Plasma etching	End-point determination Process residual-gas monitoring
VI probe/RF probe	Plasma processes	Plasma characterization via electrical parameters
Particle detectors	Vacuum processes	Monitoring of process vessel contamination

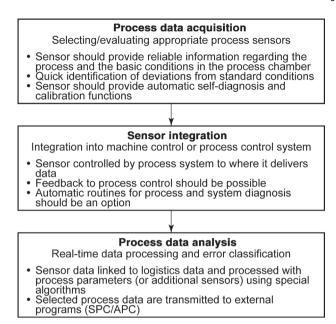


Figure 18.2 Criteria for efficiently implementing and optimizing process control.

Very complex processes require high levels of sensor integration into the automated system. Stand-alone sensor units with additional operator attention are inappropriate for such production environments. Effective process monitoring should therefore respect and continuously optimize the criteria shown in Figure 18.2 in order to improve error analysis and product quality.

18.1.2 Integrated Solutions

As mentioned, sensor integration is important due to two main reasons:

- Process monitoring and systems control must be capable of being automated.
- Data transfer to production controls or external systems such as APC (advanced process control), SPC (statistical process control), or MES (manufacturing executing system).

This strategy allows reliable monitoring and control of processes and systems as well as further data handling for thorough data analysis and quality control in production.

In the past, sensors often provided simple analog or digital signals only (e.g., for identifying end-points or deviations from set point values). Today, digital interfaces and protocols provide bidirectional data transfer between sensors and system controllers. Many sensor manufacturers provide convenient software solutions for data handling, however, often involving manufacturer-specific protocols. In spite of ongoing standardization, particularly in semiconductor industry, protocols for process sensors have not yet been standardized.

The following sections describe available concepts of integration in brief.

18.1.2.1 ASCII Protocols

Some of today's process-monitoring systems such as mass spectrometers (residual gas analyzers—RGA) are equipped with an embedded processor including system software. The process sensor can thus operate without any additional computer or software required. This applies to both the acquisition of measurement data such as partial pressures as well as to measurement configuration data including mass range, ionization, and further parameters. Comprehensive electronic control of an RGA also features an integrated web server, easily controllable from remote locations via a standard web browser.

The ASCII protocol contains sensor-specific functions and is thus published by the manufacturer. Optional tools provide integration into programming environments such as Java or C + +. Accompanying TCP/IP Ethernet protocol implementation integrates the system into the local network, thus resolving common limitations of serial interfaces (Figure 18.3).

18.1.2.2 Standardized Bus Systems

Initially, sensor networks were developed for transmitting digital information such as "on" or "off" to machines or facilities. The most common bus type, especially in Europe, is the ASI system [1], providing solutions for simple and lowcost system controls.

In addition to such simple signals, the next higher grade fieldbus is capable of transmitting more complex information, for example, set points or actual values, and it allows regulation loops. Typical examples of fieldbus systems [2] developed for industrial automation about 20 years ago include DeviceNet [3], Interbus, and Profibus [4]. New developments focus on TCP/IP and real-time processing. Due to its standardized protocols, Ethernet TCP/IP provides

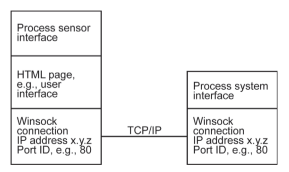


Figure 18.3 Two RGAs (residual gas analyzers) connected to an Ethernet network hub. Courtesy of MKS Instruments Deutschland GmbH.

universal networking between office computers, higher level enterprise networks, and down to the individual sensor.

The successor to *Profibus* (data transfer rate 16 megabits per second) is called Profinet (data transfer rate 100 megabits per second) and is based on the Industrial Fast Ethernet with TCP/IP. Utilizing a special chip (switch), Profinet guarantees cycle times of one millisecond at a jitter of one microsecond even when many sensors and actuators are connected [5]. SEMI (Semiconductor Equipment and Materials Institute) already accepted Profibus/Profinet in its SEMI E54.8/ E54.14 standard for applications in semiconductor industry in April 2005.

A disadvantage is the limited length of a message. Integrating a larger number of complex process sensors is limited. Thus, these bus systems are restricted to integration of simple sensors with just few measurement values and simple components such as gas flow regulators.

18.1.2.3 Sensor Integration According to SECS and HSMS Standards

In semiconductor industry, processes predominantly define necessary protocols. Thus, for historical reasons, protocols based on the serial interface RS232 [6] such as GEM (generic equipment model) and SECS (semiconductor equipment communication standard) have established. Many suppliers of semiconductor manufacturing equipment have adapted these standards and provide corresponding software for process integration.

For example, the SECS standard enables a host computer to start measurement cycles and allows transfer of measured values. SECS-I is based on RS232 and has a limited message length of 8 megabyte. SECS-II messages pass through the network as structured binary data without wasting bandwidth. If the HSMS standard (high-speed message services) is used on a TCP/IP network, the maximum message size is upgraded to 16 megabytes. The GEM standard defines a sequence of SECS-II messages for certain scenarios predefinable by the manufacturer of the equipment.

A disadvantage of serial-interface communication protocols is that the number of simultaneously communicating peers is limited to two. Furthermore, integration of process sensors is constrained because usually only two serial-interface ports are present. One of them maintains communication to the external manufacturing execution system (MES) whereas the other handles the process. Primarily, MES is responsible for the entire process control. It transfers logistics data (product IDs, recipes), and collects and prepares systems' process data.

The HSMS protocol, introduced several years ago, was not able to resolve these limitations [7]. Some commercial products overcome the restrictions by utilizing software multiplexers that allow multiple accesses to the protocol. Figure 18.4 shows an example of how the interface limitations can be resolved using the MKS® Blue Box®.

Here, sensors and/or complex process-monitoring equipment are connected via an independent (process-specific) network around the process control. Sensors using different protocols and communication technologies can be integrated, independent of whether the interfaces are analog, serial-digital, or

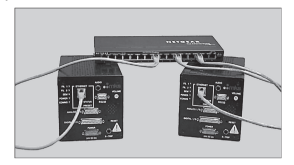


Figure 18.4 Sensor integration using SECS multiplexer in semiconductor manufacturing (MKS® Blue Box®).

TCP/IP. Gateways provide entire and manufacturer-independent sensor integration for the data acquisition system.

Independent networks are necessary in order to synchronize sensor and process data. Any data are equipped with a timestamp and assigned to the manufactured product and process step. These requirements are met by the MKS ToolSide/protocol based on industry standards TCP/IP and XML. Additionally, it sends data such as information regarding conditions in the process vessel and logistics data from the production facility to other applications. Sensor data can be retrieved from integrated web servers.

For extensive process-data analysis, according to Section 18.1.2.4, such data have to reach the production network. Figure 18.5 shows the principle of a production environment featuring a network of several processing units.

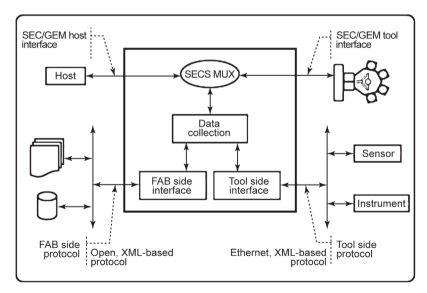


Figure 18.5 Process equipment and sensor network in semiconductor industry.

18.1.2.4 Process-Data Analysis

Performing process-data analyses is suggested if many process parameters are monitored.

Data volumes and information complexity from process sensors (e.g., mass spectra, adsorption spectra) often call for real-time data processing so that these data are utilizable for process monitoring and control. For this, sensor- and process-specific algorithms consider raw data of sensors and other measured quantities or information (Figure 18.6).

Higher level systems for process monitoring and optimization make use of a set of procedures, specialized on the complexity of semiconductor manufacturing, and providing control throughout several process steps. Some of these techniques are listed below:

SPC (Statistical Process Control). Error frequencies are correlated with machine and sensor data to localize and resolve long-term influences on final product quality. Process-wide statistical analyses check whether process tolerances are met and which impact on quality is to be expected from deviations in individual process steps.

AEC (Advanced Equipment Control). Monitoring of process chambers or processing systems using active sensors that are capable of taking corrective actions. If variations from set point values occur, running processes are terminated or warnings or alerts are raised.

APC (Advanced Process Control). Networked systems process data in a higher level system and dynamically adjust preceding and succeeding

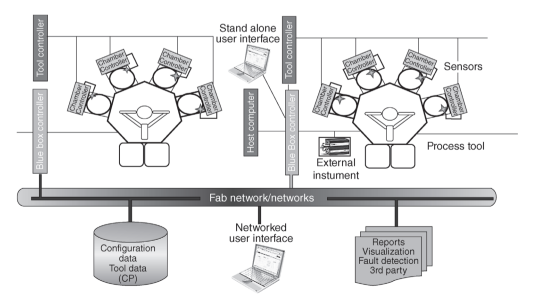


Figure 18.6 RGA-based algorithm for process control.

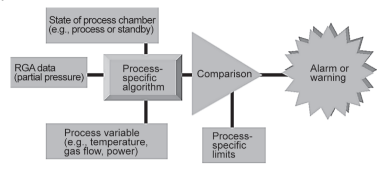


Figure 18.7 Projection of measured data in PCA.

processes. A combination of all involved sensors identifies trends and drifts, and correlates them to machine, system, and production data of the entire cvcle.

FDC (Fault Detection and Classification). Sensors control product quality during production in real time, and by using process-specific models, identify negative trends that could lead to quality loss even in succeeding processes. Machine and sensor data help to predict characteristics such as coating properties and to determine whether a product is still within a tolerable process window.

In APC applications, data volumes are often too high for univariant (with respect to a single parameter) sensor-data analysis. Thus, recently, multivariant methods for process-data analysis [8] have become available, using statistical interpretations of correlated data. Figure 18.8 shows the simplified principle. Each data point represents a measured process value. For processes with several specific steps, separate analyses are recommended.

For an analysis of the main components (PCA—principle component analysis), measured values are projected to the first two components, yielding the coordinates characterizing the current data cloud. This considerably compresses data volumes and provides a basis for identifying critical process parameters and their mutual correlations (Figure 18.7). In complex processes, several components are investigated by using the described method, thus yielding the best projection for the process and its determining quantities.

A considerable advantage is that the method not only detects the amplitude of the sensor signal but also the change in time and correlation to other process parameters. In fact, most practical process phenomena are multivariant, that is, determined by more than one parameter.

A straight example for a multivariant dependence is a virtual leak in a vacuum vessel. Although detected by an RGA due to variations in partial pressures, identification as a definite cause of failure is possible only with a correlation to machine parameters.

Some FDC systems available today also facilitate this analysis method for realtime process monitoring.

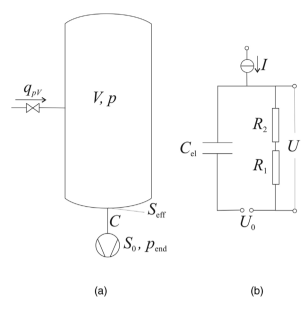


Figure 18.8 Simple vacuum system (a) with gas inlet flow q_{pV} , chamber volume V and pressure p, with a conductance C to the vacuum pump having an intrinsic pumping speed of So and its electrical analog on (b), see Example 18.2.

18.2 Calculation of Vacuum Systems

Any construction of a complex vacuum system should be accompanied by its calculation. Subject of the calculation mainly are the determination of ultimate pressure, the achievement of the required working pressure (possibly atmospheric pressure, e.g., when venting), and the required time periods to obtain these pressures. To be successful, the conductances in the system have to be determined; the pumps with their pumping speeds, ultimate pressure, and compression ratios have to be correctly selected; and the gas inlet has to be designed in an appropriate manner. Positioning the vacuum and partial pressure gauges and the valves is another important design feature so that the vacuum system can be safely and correctly controlled.

The foundation for such calculations has been laid down in Chapters 3-6. As we did in previous chapters we will explain in this one how to apply the knowledge to praxis.

18.2.1

Analytical and Numerical Calculation

For vacuum applications the ideal gas law is an appropriate assumption. The general differential equation of pressure p in a local volume cell V is

$$V\frac{dp}{dt} = q_{p\nu,\text{in}} - q_{pV,\text{out}} \tag{18.1}$$

with $q_{pV,\text{in}}$ the gas throughput in and $q_{pV,\text{out}}$ the gas throughput out of the volume. Under the additional assumption of a constant $q_{pV,in}$ and a constant pumping speed, this equation is solved by

$$p(t) = \left[p_0 - p_{\text{ult}} - \frac{q_{pV,\text{in}}}{S_N} \right] \exp\left(-\frac{S_N}{V}t\right) + p_{\text{ult}} + \frac{q_{pV,\text{in}}}{S_N},$$
(18.2)

where p_0 is the initial pressure in the volume at time zero and the pumping speed S is approximated by

$$S(p) = S_{\rm N} \left(1 - \frac{p_{\rm ult}}{p} \right). \tag{18.3}$$

Here S_N is the nominal pumping speed and p_{ult} the ultimate pressure of a pump (see Section 18.2.3).

In real cases, the above equations may be too simplified. S should be replaced by the effective pumping speed of the volume (see Section 4.1.4), but still, S may have a much more complex relationship with pressure than expressed by Eq. (18.3). q_{pV} in is not constant but due to desorption and outgassing a rather complex function of time, pressure, and temperature (Section 6.2). In a complex system of pumps and volumes, $q_{pV,in}$ may also depend on the pressure in the neighboring chamber.

To make the problem easier, it is possible to apply step functions for S and $q_{pV,\text{in}}$, that is, a constant value in a period of time and/or pressure range. With numerical methods, the periods and ranges can be partitioned rather short/ small. When the accuracy level for the calculation of complex systems is not high, the model can be simplified [9].

When pump-down times have to be calculated, it is helpful to perform this in two steps. In the first step, the evacuation time of the gas in the volume is calculated (higher pressures down to about 100 Pa); in the second step, the time to evacuate the desorbing gas (Eqs. (6.13) and (6.14)) is calculated. Finally, the two times are added. This method normally overestimates the evacuation time for a given pressure, but gives a sufficient estimate.

Both analytical and numerical methods need reliable input data. Pumping speed curves S(p) are often available for turbo molecular pumps and diffusion pumps with well-defined rough pumps, but rarely available for combined systems (e.g., with boosters and Roots pumps), because these ad hoc created pump systems have not been characterized before.

18.2.2

Calculation by Analog Electrical Network

There exists an analogy between the quantities in an electrical networks and a vacuum system. Table 18.2 lists the most important corresponding quantities. This correspondence allows one to apply software designed to develop electrical circuits to calculate a vacuum network.

Vacuum technology	Electrical engineering
Pressure difference Δp	Voltage <i>U</i>
Gas flow q_{pV}	Current I
Flow resistance Z	Electrical resistance R
Conductance C	Electrical conductance G
Volume V	Capacity Cel

Table 18.2 Analogy between vacuum technological and electrical quantities.

Example 18.1

Suppose the simple vacuum system of Figure 18.8a. (a) Draw the electrical analogon. (b) Calculate the corresponding quantity of S_{eff} . (c) Calculate the analog on to the characteristic pump-down time τ with q_{pV} (i.e., l) = 0.

- a. The chamber volume V is simulated by an electrical capacity according to Table 18.2. The chamber is evacuated via two flow conductances C and S_0 (the pumping speed of the pump alone). The electrical capacity is drained via two resistors R_2 (corresponds to C) and R_1 (corresponds to S_0). The pressure difference p(chamber) - p(pump inlet) corresponds to the voltage U. The ultimate pressure p_{ult} of the vacuum pump is best simulated by a fixed voltage U_0 against ground. The electrical current source I simulates the gas inlet flow q_{pV} .
- b. See Eq. (4.34). The total effective electrical conductance is calculated by the sum of the inverted individual conductances:

$$\frac{1}{G_{\text{tot}}} = \frac{1}{G_1} + \frac{1}{G_2}; \quad G_{\text{tot}} = \frac{G_1 G_2}{G_1 + G_2} = R^{-1} = \frac{1}{R_1 + R_2}$$

c. The vacuum technological differential equation (see also Section 18.3.6)

$$V\frac{d(p-p_{\text{ult}})}{dt} = -S_{\text{eff}}(p-p_{\text{ult}})$$

is solved by
$$p(t) = p_{\text{ult}} + p(0) \exp\left(-\frac{t}{\tau}\right)$$

with $\tau = V/S_{eff}$. The analog equation for the electrical network is $C_{el} \frac{d(U-U_0)}{dt} = -G_{tot}(U-U_0)$ with the solution

$$U(t) = U_0 + U(0) \exp\left(-\frac{t}{\tau}\right)$$

with
$$\tau = C_{\rm el}/C_{\rm tot}$$
.

The analogy, however, does have some important limits:

- The flow resistance (conductance) may depend on pressure. The electrical resistance, on the other hand, does not depend on voltage as long as Ohm's law is valid.
- The flow resistance (conductance) depends on gas species while there is only one electrical current.
- The series connection of flow resistances (conductances) has to consider the beam effect which is not necessary for the electrical analog on.
- In vacuum technology there is no analog on to imaginary impedances.

If one considers these limits, it is possible to quickly obtain important design criteria for a vacuum system by applying tools for an electrical network.

18.2.3

Ultimate and Working Pressure in Vacuum Systems

The ultimate pressure of a vacuum system is the lowest pressure that the vacuum system approaches asymptotically after a long time when no gas enters the system. It cannot be smaller than the ultimate pressure of the vacuum pumps present in the system.

The working pressure is the pressure necessary to run a process in vacuum and usually is several orders of magnitudes above ultimate pressure.

18.2.3.1 Ultimate Pressure

The ultimate pressure of a vacuum pump is the lowest pressure that the vacuum pump approaches asymptotically when no gas enters on the suction side. Ultimate pressures are measured using internationally standardized setups specific to each pump category (compare Table 15.8).

Since the ultimate pressure p_{ult} , per definition, is an asymptotic quantity which would require infinitely long waiting time for measuring, practical applications use the so-called ultimate working pressure $p_{W,ult}$. It refers to the ultimate pressure obtained after an individually specified (by standards) finite time. Times (pump-down times) required for measuring $p_{W,ult}$ increase as pressures drop, and amount to approximately 24 hours for high-vacuum (HV) pumps, where $p_{W \text{ ult}}$ is already in the ultrahigh-vacuum (UHV) range. For rotary vane pumps, it is distinguished between ultimate total pressure (with oil vapor pressure) and ultimate partial pressure (without oil vapor pressure). For pumps equipped with gas ballast (see Section 7.6.1), both pressures are commonly measured with and without gas ballast.

The ultimate pressure of a vacuum apparatus or system is the lowest pressure establishing asymptotically in a clean and dry vacuum apparatus or system. In practice, the ultimate working pressure $p_{W.ult.S}$ is measured analogously to the ultimate working pressure of a vacuum pump. This pressure is determined by

a) the type of vacuum pump or vacuum pump combination, including additional components (e.g., cold traps, sorption traps, etc.);

- b) the gas flow desorbing from the walls of the vacuum vessel and from built-in components as well as outgassing from the inside (Chapter 6 and Section 16.3);
- c) the tightness of the apparatus or system, given mainly by the type and number of flanges and their seals.

The ultimate working pressure $p_{W.ult,S}$ is generally higher, but at least equal to the measured ultimate pressure produced by the employed vacuum pump under standard conditions.

18.2.3.2 Operating Pressure

During a vacuum process, the time- and location-dependent pressure in a vacuum vessel is referred to as working pressure $p_{\rm w}$. In practice, the locationdependent pressure gradient is out of consideration because vacuum vessels usually provide a number of fixed pressure-measurement points. For pressure critical processes, this is a matter of consideration.

This time dependence of working pressure follows the equation

$$q_{pV,\text{out}}(t) + q_{pV,\text{evap}}(t) + q_{pV,\text{leak}}(t)$$

$$p_{\text{w}}(t) = \frac{+q_{pV,\text{perm}}(t) + q_{pV,\text{process}}(t)}{S_{\text{eff}}(t)} + p_{\text{W, ult, S}}, \tag{18.4}$$

with the outgassing flux $q_{pV,\text{out}}$, that is, the gas flow desorbing from the walls of the vacuum vessel and from built-in components as well as outgassing from the inside (Chapter 6 and Section 16.3). $q_{pV,evap}$ is the gas flow produced by evaporating volatile substances and is pumped off by the pump. $q_{pV,\mathrm{leak}}$ (Section 19.2.2) and $q_{pV,\mathrm{perm}}$ (Section 16.3) are the (usually time-independent) leakage and permeation gas flows, respectively. $q_{pV,\mathrm{process}}$ is the gas flow developing in the vacuum process (e.g., annealing, melting, evaporating). S_{eff} is the effective pumping speed at the vessel (Section 4.1.4). $p_{\mathrm{W,ult,S}}$ is the ultimate working pressure developing in the vessel when all $q_{pV} = 0$.

When operating a vacuum process, the main pressure of concern is the base pressure established in the vessel prior to process initiation. This base pressure or starting pressure $p_{\rm w}(0)$ is obtained from Eq. (18.1) if $q_{pV,{\rm process}}$ is set to zero:

$$p_{\rm w}(0) = \frac{q_{pV,\rm out}(0) + q_{pV,\rm evap}(0) + q_{pV,\rm leak} + q_{pV,\rm perm}}{S_{\rm eff}} + p_{\rm W,ult,S}.$$
 (18.5)

Depending on the type of the operated vacuum process, tolerable values for this starting value differ significantly. For drying, values in the fine-vacuum range are sufficient. Vapor deposition requires high vacuum. The higher the tolerable starting pressure $p_{w}(0)$ and the lower the process gas flow, the more economic the vacuum process. However, at times, process gas flow can reach considerable values thus representing an important selection criterion (maximum process gas flow) for vacuum-pump or pump-combination selection. For assigning the effective pumping speed mostly to the vacuum process, the added system-related gas

flows $q_{pV,\mathrm{out}}$ + $q_{pV,\mathrm{evap}}$ + $q_{pV,\mathrm{leak}}$ + $q_{pV,\mathrm{perm}}$ should remain below 0.1 $q_{pV,\mathrm{process}}$ if

If the working pressure is determined mainly by the gas flow resulting from the vacuum process, Eq. (18.1) yields

$$p_{\rm w} = \frac{q_{pV, \rm process}}{S_{\rm eff}}.$$
 (18.6)

Universally valid amounts for gas flows developing in individual vacuum processes cannot be given because they differ considerably depending on the conducted process.

Example 18.2

Steel with a mass fraction w = 50 ppm hydrogen is molten at a rate s =0.02 kg min⁻¹ in a vacuum furnace. If the total amount of hydrogen is released during melting, the mass flow of pumped-down hydrogen is

$$q_m = \dot{m} = sw = 0.02 \text{ kg min}^{-1} \times 50 \times 10^{-6} = \frac{10^{-6} \text{ kg}}{60 \text{ s}} = 1.67 \times 10^{-8} \text{ kg s}^{-1}.$$

According to the equation of state (3.19) or Eq. (4.14), this corresponds to a pV

$$q_{pV} = \frac{\dot{m}}{M_{\text{molar}}} RT.$$

With $M_{\text{molar}} = 2 \text{ kg kmol}^{-1}$ for H_2 , $R = 83.14 \text{ mbar } \ell \text{ mol}^{-1} \text{ K}^{-1}$, and T = 293 K $(\vartheta = 20 \, ^{\circ}\text{C})$, we find

$$=q_{pV,\text{process}} = \frac{1.67 \times 10^{-8} \text{ kg s}^{-1} \times 83.14 \text{ mbar } \ell \text{ mol}^{-1} \text{ K}^{-1} \times 293 \text{ K}}{2 \text{ kg kmol}^{-1}}$$
$$= 2.03 \times 10^{-1} \text{ mbar } \ell \text{ s}^{-1}.$$

For working pressure to remain constant ($p_w = 1 \times 10^{-4}$ mbar) throughout the entire process, the effective pumping speed required according to Eq. (18.3) is

$$S_{\text{eff}} = \frac{q_{pV,\text{process}}}{p_{\text{w}}} = \frac{2.03 \times 10^{-1} \text{ mbar } \ell \text{ s}^{-1}}{1 \times 10^{-4} \text{ mbar}} = 2030 \ \ell \text{ s}^{-1}.$$

However, if a working pressure $p_{\rm w}=1\times10^{-6}$ mbar is desired for the process, the effective pumping speed amounts to

$$S_{\text{eff}} = \frac{2.03 \times 10^{-1} \text{ mbar } \ell \text{ s}^{-1}}{1 \times 10^{-6} \text{ mbar}} \approx 2 \times 10^{5} \ell \text{ s}^{-1}.$$

As long as evaporable substances with vapor pressures above the desired ultimate pressure are present in a vacuum vessel, the pressure in the apparatus is determined, on the one hand, by the evaporation rate q_V of the substances, Eq. (3.133), and on the other, by the effective pumping speed $S_{\rm eff}$ of the pump arrangement. If $q_V \gg S_{\text{eff}}$, the pressure in the apparatus remains just barely below the saturation vapor pressure p_s of the substance until evaporation has completed. In any other case, a pressure $p < p_s$ establishes which is calculated from an equation equating the rate of change of the vapor amount in the recipient and the difference between evaporation rate q_V and pumped gas S_{meff} (compare Section 18.2.6, Eq. (18.4)). According to Table A.13, the saturation vapor pressure of water at room temperature $\theta = 20$ °C is $p_s = 23.35$ mbar. Thus, for a vessel containing water at room temperature, regardless of whether it is distributed or rather a film of water covering the walls, the pressure during evacuation initially does not drop significantly below 23.4 mbar. Only after the vessel dries due to continuous pump-down of the water vapor, the pressure approaches the ultimate value determined by other limiting effects.

The evaporation draws heat from the water which leads to a cooling of the water and to a drop in vapor pressure with correspondingly prolonged evaporation times.

Example 18.3

When cooling from $\theta = 20$ °C to $\theta = 0$ °C, that is, by $\Delta T = \Delta \theta = 20$ K, a mass m =1 g of water releases the quantity of heat $\Delta Q = mc_p \Delta T$, with the specific heat capacity of water $c_p \cong 4.2 \text{ J g}^{-1} \text{ K}^{-1}$. However, supplying $\Delta Q = 1 \text{ g} \times 4.2 \text{ J g}^{-1} \text{ K}^{-1} \times 10^{-1} \text{ g}$ 20 K = 84 J only evaporates $m' = \Delta Q/\Lambda_v = 84 \text{ J}/(2500 \text{ Jg}^{-1}) = 3.4 \times 10^{-2} \text{ g}$, that is, 3.4% of the water. Thereby, the saturation vapor pressure drops from $p_s =$ 23.3 mbar to $p_s = 6.1$ mbar. Now the water freezes and releases $\Delta Q' = (m - m')\Lambda_s$, with Λ_s being the solidification heat, causing $m'' = \Delta Q'/\Lambda_v \approx$ $1 \text{ g} \times 336 \text{ J} \text{ g}^{-1}/(2500 \text{ J} \text{ g}^{-1}) \approx 1.4 \text{ g}$ of ice to evaporate. Ongoing evaporation of the ice (evaporation heat of ice $\Lambda_{v,ice}\approx 2900\,J\,g^{-1})$ then cools the ice, and thus, further reduces the vapor pressure.

Evaporating $m''' = m - m'' = 0.86 \,\mathrm{g}$ of ice requires the heat amount $\Delta Q^{\prime\prime\prime} = 0.86 \,\mathrm{g} \times 2900 \,\mathrm{J}\,\mathrm{g}^{-1} = 2500 \,\mathrm{J}$. However, now the specific heat capacity of ice is $c_p \approx 2 \, \text{Jg}^{-1} \, \text{K}^{-1}$ (at $\vartheta \approx 0 \, ^{\circ}\text{C}$, it drops with temperature!). If 0.01 g of ice evaporates, then the temperature of the remaining 0.85 g would drop by $\Delta T =$ 20 K to $\theta = -20$ °C, and the vapor pressure would be reduced from $p_{\text{s,ice}}(0$ °C) = 6.1 mbar to $p_{\text{sice}}(-20\,^{\circ}\text{C}) = 1$ mbar. Without any further calculations, it seems obvious that removing the water from the vessel without providing any heat from the surrounding via convection, radiation, or controlled heating would require an infinite amount of time.

In the low-vacuum regime, desorption (adsorbed gas) and outgassing (occluded gas) usually do not disturb working pressures, and in the medium vacuum, generally, their influence is still low. Under high vacuum, outgassing has considerable negative impact on working pressures and on pump-down times. In the UHV regime, for producing extremely low pressures, it is necessary to remove adsorbates and, in particular occluded gas, as far and rapidly as possible to push their residual gas flows below disturbing values. Outgassing rates of clean metal, glass, and ceramic surfaces at room temperature after an hour of pumping are in the range (Figure 16.7)

$$j_{pV, {
m out}} \approx 10^{-4} \, {
m mbar} \, \ell \, {
m s}^{-1} \, {
m m}^{-2} (10^{-2} \, {
m Pa} \, \ell \, {
m s}^{-1} \, {
m m}^{-2}).$$

For an outgassing surface area of $A = 1 \text{ m}^2$, this outgassing rate and Eq. (18.1) yield the pumping speeds required for desired working pressures:

$p_{\text{W}}(0) = 10^{-7} \text{mbar} (10^{-5} \text{Pa})$	$S_{\rm eff} = 103 \ell \rm s^{-1}$
$p_{\rm W}(0) = 10^{-9} \mathrm{mbar} (10^{-7} \mathrm{Pa})$	$S_{\rm eff} = 105 \ell \rm s^{-1}$
$p_{\text{W}}(0) = 10^{-11} \text{mbar} (10^{-9} \text{Pa})$	$S_{\rm eff} = 107 \ell {\rm s}^{-1}$

This explains why baking is necessary. Without such a procedure, uneconomically large pumps would be required.

The permeation gas flow entering a vessel through metal, glass, or ceramic walls at room temperature does usually not impede the producible base or ultimate pressure.

Elastomers are the most common sealing materials for low, medium, and high vacuum. Their high gas emissions (Figures 16.7 and B.12) and permeabilities, particularly for helium, have to be taken into account (Figure 16.5). This leads to the basic rule that the use of elastomers in vacuum apparatuses and systems should be reduced to the absolutely necessary minimum. Design of an elastomer seal should aim at keeping both the surface areas of the rubber-elastic material exposed to high pressure as well as the surface areas exposed to low pressures as small as possible. In addition, it is advisable to dry elastomers at approximately 60 °C prior to installation.

Reusing elastomer O-ring seals is limited. Producing tight seals with old Oring seals requires high contact pressures because they harden with time. Hardened O-ring seals should not be reused. If absolutely necessary, external greasing of a hardened O-ring seal with appropriate HV grease may provide sufficient sealing if mechanical stresses are low.

Elastomers are inappropriate for use in UHV applications. Here, not only permeation gas flows are disturbing, also, the comparably low thermal stability of elastomers is a problem. UHV apparatuses have to be bakeable at up to $\theta = 400$ °C, making employment of special metal seals a necessity.

For economic reasons, and with a particular focus on the necessary pumping speed to be installed in a system, the total of all leakage gas flows should not exceed 5%-10% of the gas flow pumpable by the vacuum pump at the lowest working pressure p_{extrmw} .

18.3 **Pressure Control**

Pressure in a vacuum system can be controlled in two ways [10]:

- by the gas injected into the system,
- by the effective pump speed.

Control by the gas inlet has the advantage that the time constant $\tau = V/S_{\text{eff}}$ (see Example 18.1) of the system remains unchanged. Such an additional change places high demands on a controller. A constant effective pumping speed also has the advantage that composition and total pressure of the residual gas remain constant. This is particularly important, if the residual pressure may disturb the process. For surface processes (e.g., evaporation), however, where the gas flow cannot be changed, pressure control via the effective pump speed is the only choice. Also in coating processes, where the total gas flow as well as its composition (carrier gas, active gas) must remain constant, the effective pump speed is controlled.

To control the gas flow in, dosing valves and mass flow controllers are used. The latter are composed of a flowmeter (Section 13.6) and a control valve. In Section 13.6, we mentioned that the linearity of a flowmeter is rather limited. With digital technique and look-up tables nonlinearities cannot be completely compensated. For the purpose of calibration, repair, or replacement mass, flow controllers can be easily replaced.

Control of effective pump speed may be performed by the pump itself or a variable conductance between process chamber and vacuum pump, for example, via a dosing valve.

The pump speed of only a few pumps and only in a few pressure ranges can be controlled. The pump speed of membrane pumps, turbomolecular pumps, and in particular Roots pumps can be changed by controlling their frequency. The times for adjustment are normally relatively slow so that this type of pressure control is limited to processes where time constants are not critical.

When control or dosing valves are being used, one needs to consider that the effective pump speed consists of three contributions: the conductance of the control valve C_{CV} , the conductance of the connection tubes to the pumps C_{tube} and the intrinsic pump speed S_0

$$\frac{1}{S_{\text{eff}}} = \frac{1}{S_0} + \frac{1}{C_{\text{tube}}} + \frac{1}{C_{CV}}.$$
 (18.7)

For given $q_{pV,process}$, the lower limit of the controllable pressure range is given by $p_{\min} = q_{pV,\text{process}}/S_{\text{eff}}$ with C_{CV} in maximum value. The upper limit is determined by the smallest conductance C_{CV} of the control valve close to its vacuum tight position (if applicable). Suitable control valves are iris diaphragms (these are not vacuum tight), butterfly valves, gate valves, and Chevron baffles with adjustable blade angles. The linearization of the conductance is difficult and depends on the valve type. It is achieved by an appropriate gear between driving motor and valve mechanics.

18.4 **Techniques for Operating Low-Vacuum Systems**

18.4.1

Overview

Many, particularly industrial vacuum processes are carried out under low vacuum. Typical examples are clamping, holding, handling, and sorting of small and larger flat workpieces, pouring of liquids, deep-drawing of plastic components, drying, impregnating, evaporating, and condensing, packaging of foodstuff and luxury foods such as meat, fruits, coffee, and so on. Even LCL (Less than Container Load) packaging increasingly relies on low-vacuum equipment.

Many of these processes utilize the pressure of atmospheric air for the process by establishing a pressure difference Δp . Other processes require reduced amounts of oxygen and/or humidity in the air without calling for extreme vacua. However, in any case, mechanical stresses on the vacuum vessels (chambers) must be considered (Section 17.4). Regardless of how intense a vessel is evacuated, the load on containers never exceeds the ambient atmospheric pressure of approximately 1 bar.

Example 18.4

A shop-window glass with the mass $G = 2000 \,\mathrm{kg}$, for example, a square, 6-mlong pane with a thickness of 2 cm, is to be lifted off a stack and transported to machining equipment. The suction cap (Figure 18.9) is evacuated with a vacuum pump offering a nominal pumping speed of $S_n = 30 \,\mathrm{m}^3 \,\mathrm{h}^{-1} = 8.3 \,\ell\,\mathrm{s}^{-1}$ within $\Delta t = 5$ s (suction stroke) down to a pressure difference $p_{\text{mout}} - p_{\text{in}} = 900$ mbar.

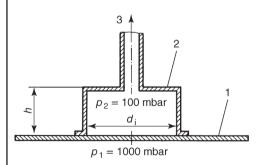


Figure 18.9 Vacuum lifting device: 1 lifted item (window glass), 2 suction cap, 3 connection to vacuum pump.

For the cap to lift the pane, the minimum internal diameter of the cap can be calculated from the equilibrium condition (F_G : weight):

$$(p_{\text{out}} - p_{\text{in}}) \frac{d_{\text{in}}^2 \pi}{4} = F_{\text{G}} = 2000 \text{ kg} \times 9.81 \text{ m s}^{-2} = 9 \times 10^4 \text{ Pa} \frac{\pi}{4} d_{\text{in}}^2,$$

which yields $d_{\rm in}=0.53$ m. Thus, in practice, a minimum of $d_{\rm in}=0.6$ m would be

Equation (18.15) gives the evacuating time for a vessel of volume V, pumped by a pump with the pumping speed S_{eff} :

$$t_{\rm p} = \frac{V}{S_{\rm eff}} \ln \frac{p_1}{p_2}.$$

Assuming $S_{\text{eff}} = 0.5 S_{\text{n}}$, the volume of the suction cap

$$V = S_{\text{eff}} t_p / \ln \frac{p_1}{p_2} = 0.5 \times 8.3 \, \ell \, \text{s}^{-1} \times 5 \, \text{s} / \ln \frac{1000 \, \text{mbar}}{100 \, \text{mbar}}$$
$$= 9 \, \ell = 9 \times 10^{-3} \, \text{m}^{-3}.$$

This determines the height of the suction cap, whose inner cross-sectional area $A = d_{\rm in}^2 \pi/4 = 0.283 \,\mathrm{m}^2$, and thus, $h = V/A = 9 \times 10^{-3} \,\mathrm{m}^3/(0.283 \,\mathrm{m}^2) = 31.8 \times 10^{-3} \,\mathrm{m}^3$ $10^{-3} \text{ m} = 32 \text{ mm}.$

Tightly sealed evacuating equipment is a prerequisite for the calculations in Example 18.4. In practice, however, unavoidable surface roughness and contaminations of handled parts cause a certain leakage gas flow that should be considered in dimensioning. In addition, note that the resistance of the suction tubing (Section 4.1.4) reduces pumping speed at the suction cap. (In the example, this was taken into account by introducing the factor 0.5 but special cases may require different correction factors.) Both effects prolong cycle times. Also, the process relies on underpressure in the suction cap to prevail throughout the handling time. This is guaranteed easily by integrating a safety valve in the suction line. Generally, any equipment utilizing atmospheric pressure for establishing pressure gradients is subject to a drop in air pressure with absolute altitude (Figure B.1).

18.4.2

Assembly of Low-Vacuum Systems

From a vacuum-technological view, requirements concerning tightness and gas emissions in a low-vacuum system are comparably low. Therefore, components meeting just these low requirements are utilizable. Joining and sealing such components often involves screwed joints, sealed with Teflon tape. Shut-off elements are standard valves as used for gas lines. However, to avoid unpleasant surprises and the delays they produce, it is advisable to use vacuum flanges and elastomer seals for any joints in the low vacuum as well, and to seal rotary leadthroughs with rotary shaft seals or cup leather shaft seals.

18.4.3

Pumps: Types and Pumping Speeds

Numerous pump types as listed in Table 18.3 are available for producing low vacua. The specific operating functionalities, conditions, and economic considerations determine designs and utilization of pumps (Table 18.3).

As a rough, but not compulsory guideline for pump selection, we can differentiate between working pressures $p_{\rm w} < 50$ mbar and $p_{\rm w} > 50$ mbar. Liquid (water) ring vacuum pumps and dry-running multiple vane pumps are used mostly for $p_{\rm w} \ge 50$ mbar, whereas oil-sealed sliding vane rotary pumps and rotary plunger pumps are common to applications with $p_{\rm w}$ < 50 mbar. Screw-type pumps and vapor jet pumps are used in both pressure ranges. Roots pumps (usually combined with oil-sealed rotary vacuum pumps) are rarely employed in low-vacuum technology (only, for example, if considerable amounts of process gases develop at low working pressures of approximately 10 mbar).

The pumping speed applied to a vacuum apparatus or system is determined by the desired working pressure, pump-down times, amounts of gases and vapors released, and the leakage gas flow. If the amount of gas or vapor produced in a process is unknown, it is often appropriate to select pumping speed from Figure 18.10. Note, however, that Figure 18.10 is based on empirical values and can only provide a rough guideline. Particularly for vessel volumes $V < 100 \,\ell$ and in the low-vacuum range, lower pumping speeds than obtained from Figure 18.10 are often sufficient. In pump assembly, appropriate dimensioning of tubing diameters is mandatory (see Chapter 5).

18.4.4 **Low-Vacuum Pump Stands**

Due to the variety of vacuum-technological applications in industry and research, demands in terms of pump systems and vacuum apparatuses vary considerably. For example, even for purely economic reasons, pumping speeds and working pressures should be matched carefully to the planned process.

Table 18.3 Pumping-speed ranges of commercial vacuum pumps
for producing rough vacuum.

Pump types	Pumping-speed ranges	
Diaphragm pump	$1-20 \text{ m}^3 \text{ h}^{-1}$	
Liquid ring vacuum pump	$10-20000~\text{m}^3~\text{h}^{-1}$	
Multiple vane pump	$1-5000~\text{m}^3~\text{h}^{-1}$	
Sliding vane rotary pump	$1-600 \mathrm{m^3 h^{-1}}$	
Rotary plunger pump	$100-500 \text{ m}^3 \text{ h}^{-1}$	
Screw-type pump	$70-2500 \text{ m}^3 \text{ h}^{-1}$	
Vapor jet pump	$20{-}100000m^3\;h^{-1}$	

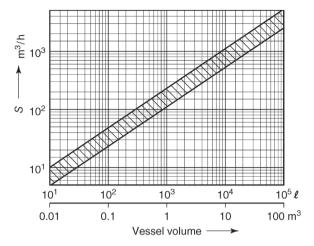


Figure 18.10 Suggested effective pumping speeds $S_{\rm eff}$ (guidance values) at vacuum vessels for rough-vacuum (and fine-vacuum) operation.

Vacuum industry has developed pump systems, listed under the collective name pump stands in manufacturer catalogs, in order to simplify users' pump selection and pump combination. By adding standard accessories, such pump stands are easily adapted to meet any desired special requirements. For example, Figure 18.11 shows a low-vacuum pump stand used in a small,

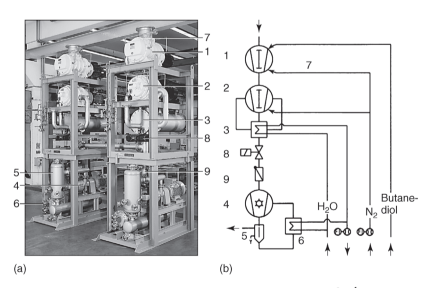


Figure 18.11 (a) Rough-vacuum pump stand for polyester processing (overall dimensions width \times depth \times height: 4350 mm \times 2500 mm × 1200 mm. 1 Roots pump no. 1 $(S_n = 4000 \text{ m}^3 \text{ h}^{-1})$; 2 Roots pump no. 2,

gas-cooled ($S_n = 3000 \text{ m}^3 \text{ h}^{-1}$); 3 gas cooler to no. 2; 4 liquid ring pump ($S_n = 800 \text{ m}^3 \text{ h}^{-1}$); 5 separator to 4; 6 tube-bundle cooler to 4; 7 seal gas tubing; 8 nonreturn valve; 9 check valve. (b) Diagram of pump stand in Figure 18.11.

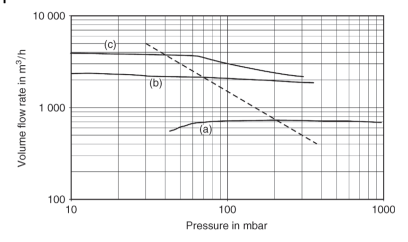


Figure 18.12 Pumping speed curves of rough-vacuum pump stand in Figure 18.11. (a) Liquid ring pump, (b) Roots pump 2 plus liquid ring pump, (c) Roots pump 2 plus Roots pump 1 plus liquid ring pump. Dashed line: pump stand's design condition.

discontinuously operating system (batch operation) for polyester processing. It comprises a series connection of two Roots pumps as well as a liquid ring pump (Section 7.3.1).

Figure 18.12 plots the characteristic values of pumping speed S versus inlet pressure p_{in} for this pump stand. The lowest suction pressure is 40 mbar. Continuously operating, larger systems for polyester production usually employ vapor jet pumps for producing vacuum.

18.4.5

Low-Vacuum Pressure Measurement

Most measuring instruments available for low vacuum are different types of mechanical gauges, for example, diaphragm vacuum gauges (Section 13.2.5). In the low-vacuum range, pressure readings of mechanical vacuum gauges are independent of the gas or vapor species.

18.4.6

Pressure and Pump-Down Times in Low Vacuum

A number of factors determine the pump-down time $t_{\rm p}$ required by a vacuum pump of pumping speed S_{i} , or effective pumping speed S_{eff} if the resistance of the tubing (Section 4.1.4) between pump and vessel is relevant, to evacuate a vessel with volume V from starting pressure $p_0(t=0)$ down to a desired pressure p(t). The pump-down time is affected by the influx (compare

Section 18.2.2) due to leakage, permeation, outgassing, evaporation, and gas- or vapor production in the process:

$$q_{nV,\text{in}} = q_{nV,\text{leak}} + q_{nV,\text{perm}} + q_{nV,\text{out}} + q_{nV,\text{evap}} + q_{nV,\text{process}}.$$
 (18.8a)

Then, at any point in time, the change with time of the pV amount d(pV)/dt in the vessel is equal to the influx $q_{pV,in}$ minus the outflow $q_{pV,out} = p S_{eff}$. Thus, since V = constant,

$$\frac{\mathrm{d}(pV)}{\mathrm{d}t} = V \frac{\mathrm{d}p}{\mathrm{d}t} = q_{pV,\mathrm{in}} - pS_{\mathrm{eff}}.$$
(18.8b)

S and S_{eff} are usually pressure dependent. In the low-pressure range, below the pressure corresponding to the (often constant) maximum pumping speed S_{max} , the pumped gas amount pS_{max} is reduced due to leakage and backflow from the compression chamber so that we can approximate

$$pS = pS_{\text{max}} - q_{pV,\text{back}+L}. \tag{18.9}$$

If $pS_{\text{max}} = q_{nV \text{ back+L}}$, then S = 0, and the system reaches its ultimate pressure:

$$p_{\text{ult.P}}S_{\text{max}} = q_{nV.\text{back+L}}$$
.

Putting this into Eq. (18.12) yields an approximation for the pressure dependence of the pumping speed at low pressures:

$$S = S_{\text{max}} \left(1 - \frac{p_{\text{ult,P}}}{p} \right)$$
 and $S_{\text{eff}} = S_{\text{eff,max}} \left(1 - \frac{p_{\text{ult,P}}}{p} \right)$. (18.10a,b)

Figure 18.13 shows the goodness of the approximation. If it is used for calculating $S_{\rm eff}$, Eq. (18.10b), in cases where the pump's tubing conductance is comparable to the pumping speed of the pump, the error remains within the same order of magnitude. Putting the term for $S_{\text{eff}}(p)$ from Eq. (18.10b) into the differential equation (17.4b), and assuming $q_{pV,in}$ to be constant in time, we find the solution for the initial condition $p = p_0$ for t = 0:

$$p = p_0 \exp\left(-\frac{S_{\text{eff,max}}}{V}t\right) + \left(\frac{q_{pV,\text{in}}}{S_{\text{eff,max}}} + p_{\text{ult,P}}\right) \left(1 - \exp\left(-\frac{S_{\text{eff,max}}}{V}t\right)\right). \quad (18.11)$$

Thus, the pressure in the vessel drops exponentially. After an infinite pumping time, it approaches the ultimate value (ultimate working pressure)

$$p_{\text{w,ult,A}} = p_{\text{ult,P}} + \frac{q_{pV,\text{in}}}{S_{\text{eff max}}}.$$
(18.12)

For a clean and vacuum-tight apparatus, in which the process does not develop any gas $(q_{pV,in} = 0)$, the producible ultimate pressure is equal to the ultimate pressure of the vacuum pump $(p_{w,ult,A} = p_{ult,P})$.

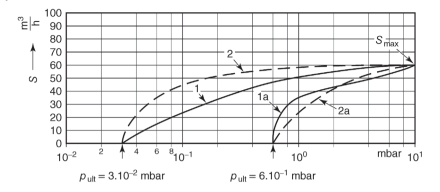


Figure 18.13 Pumping speed curves of twostage positive displacement pump with nominal pumping speed $S_n = 60 \text{ m}^3 \text{ h}^{-1}$. Measured data: curve 1 without gas ballast, curve 1a with gas ballast. Calculated according to

Eq. (18.6): curve 2 without gas ballast, curve 2a with gas ballast. Calculations assume ultimate pressures $p_{\rm ult}=3\times10^{-2}$ mbar (without gas ballast) and $p_{\rm ult}=6\times10^{-1}$ mbar (with gas ballast).

As Eq. (18.11) shows, the quantity determining the drop with time, the socalled e-value time or time constant.

$$\tau = \frac{V}{S_{\text{eff max}}}.$$
 (18.13)

For $t = \tau$, $\exp(-S_{\rm eff,max}t/V) = \exp(-t/\tau) = e^{-1} = 0.368$. Correspondingly, for $t = 3\tau$, $\exp(-3) = 0.050 = 5\%$, for $t = 4\tau$, $\exp(-4) = 0.018 = 1.8\%$, and for $t = 5 \tau$, $\exp(-5) = 0.007 = 0.7\%$.

This allows a quick estimation as to when ultimate pressure, Eq. (18.12), is obtained but a few percent. With Eq. (18.13), rearranging Eq. (18.11) for t yields the pumping time for a drop in pressure from p_0 to p:

$$t_{\rm p} = \tau \ln \frac{p_0 - p_{\rm ult, P} - \frac{q_{pV,\rm in}}{S_{\rm eff,max}}}{p - p_{\rm ult, P} - \frac{q_{pV,\rm in}}{S_{\rm eff,max}}}.$$
(18.14)

For low vacuum, it is usually sufficient to approximate pumping time using

$$t_{\rm p} = \tau \ln \frac{p_0}{p} = \frac{V}{S_{\rm eff, max}} \ln \frac{p_0}{p}. \tag{18.15}$$

Prerequisites for Eqs. (18.14) and (18.15) to be valid are a pump operating at working temperature (70 °C - 80 °C) and total pressure measurement. Additionally, Eq. (18.15) requires a dry and clean vessel, ventilated with atmospheric air. In practice, a safety factor of 20% is added to the calculated values of $t_{\rm p}$.

Example 18.5

A clean and dry vacuum system with volume $V = 1000 \,\ell$ is evacuated from atmospheric pressure $p_0 = 1013 \, \text{mbar}$ down to working pressure $p_w = p_1 = 1 \, \text{mbar}$ using a directly attached ($S_{xtrmeff} = S$) pump combination comprising an oil-sealed rotary pump R (without gas ballast) and a Roots pump W. Starting pressure for the Roots pump is $p_{W \text{ on}} = 20 \text{ mbar}$. The leakage rate of the system shall be $q_1 = 10^{-1} \text{ mbar } \ell \text{ s}^{-1}$. The following data of the rotary pump R and the pump combination W + R are needed for calculating pumping time t_p (see also Figure 18.14):

Rotary pump R

$$S_{\text{max}} = 150 \text{ m}^3 \text{h}^{-1} = 41.7 \, \text{\ell s}^{-1}, \qquad p_{\text{ult.P}} = 3 \times 10^{-2} \text{ mbar},$$

Combination W + R

$$S_{\text{max}} = 540 \text{ m}^3 \text{ h}^{-1} = 150 \, \ell \text{ s}^{-1}, \qquad p_{\text{ult.P}} = 4 \times 10^{-3} \text{ mbar.}$$

Calculations are based on the measured pumping speed curves shown in Figure 18.14. Evacuating, and thus calculation of pumping time t_p , both feature two succeeding steps:

- **a.** Pumping time t_{p1} for producing pressure $p_1 = 20$ mbar, using only the rotary
- **b.** Pumping time $t_{\rm p2}$ for obtaining pressure $p_2=1$ mbar, using the pump combination W + R in the pressure range between 20 and 1 mbar.

According to Figure 18.14, during step (a) (evacuation from $p_0 = 1013$ mbar to $p_1 = 20$ mbar), the pumping speed of the rotary pump R is pressure-independent and constant, $S = S_{\text{max}} = 150 \text{ m}^3 \text{ h}^{-1}$. As Eq. (18.6) shows, this circumstance can be taken into account by setting $p_{\text{ult,P}} = 0$. In the second step (evacuation from $p_1 = 20$ mbar to $p_2 = 1$ mbar), according to Figure 18.14, the pumping speed of the combination rises from $S(20 \text{ mbar}) = 375 \text{ m}^3 \text{ h}^{-1}$ to $S(1 \text{ mbar}) = 500 \text{ m}^3 \text{ h}^{-1}$, following a logarithmic slope with p. This fact is considered by using a mean constant pumping speed $S = 450 \text{ m}^3 \text{ h}^{-1} = 125 \text{ }\ell \text{ s}^{-1}$ (curve B in Figure 18.14) for calculation, and again, by setting $p_{\text{ult,P}} = 0$. Then, Eq. (18.13) yields

$$au_1 = \frac{1000 \, \ell}{41.7 \, \ell \, s^{-1}} = 24 \, s, \quad au_2 = \frac{1000 \, \ell}{125 \, \ell \, s^{-1}} = 8 \, s,$$

and with Eq. (18.14),

$$t_{p1} = 24 \text{ s} \times \ln \frac{1013 \text{ mbar} - \frac{10^{-1} \text{ mbar} \ell \text{ s}^{-1}}{41.7\ell \text{ s}^{-1}}}{20 \text{ mbar} - \frac{10^{-1} \text{ mbar} \ell \text{ s}^{-1}}{41.7\ell \text{ s}^{-1}}} = 24 \text{ s} \times 3.93 = 94.2 \text{ s},$$

$$t_{p2} = 8 \text{ s} \times \ln \frac{20 \text{ mbar} - 7 \times 10^{-4} \text{ mbar}}{1 \text{ mbar} - 7 \times 10^{-4} \text{ mbar}} = 24 \text{ s}.$$

Thus, the total pumping time

$$t_p = t_{p1} + t_{p2} = 118 \text{ s (multiplying by 1.2 yields 145 s)}.$$

Note that a certain amount of time is necessary for switching. If pumped only with the rotary pump R, evacuating from 1013 mbar down to 1 mbar would take $t_{\rm p} = 166 \, \rm s.$

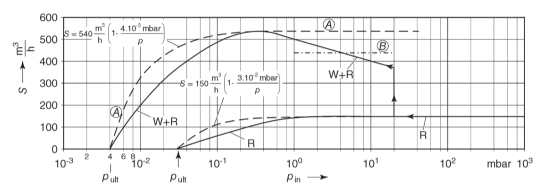


Figure 18.14 Pumping speed curves of a rotary pump (R) and a pump combination comprising a Roots pump (W) and a rotary pump. Solid lines: measured data. Dashed lines: Calculated from Eq. (18.6). p_{in} pressure at inlet. B practical average. The theoretical value A is too high.

This example also shows that the assumed leakage rate $q_1 = 10^{-1}$ mbar ℓ s⁻¹ does not influence pumping time as long as considering low-vacuum regimes. Here, leakage rates could be far higher before they impede the vacuum. If the pump combination is not attached to the vacuum vessel directly but via a connecting tube, then it is necessary to calculate the effective pumping speed $S_{\rm eff}$ according to Section 4.1.4. The nomogram in Figure B.13 is convenient for straightforward calculation of pumping times in the low vacuum.

18.4.7

Venting

Venting time is the time it takes to refill a vacuum system evacuated to the pressure p_2 with atmospheric air.

When the pump is shut off, that is, $S_{\text{eff}} = 0$, Eq. (18.8b) yields the rise in vessel pressure:

$$V\frac{\mathrm{d}p}{\mathrm{d}t} = q_{pV,\mathrm{in}}.\tag{18.8c}$$

Thus, if the temperature T of the gas in the vessel is kept constant, the venting time for changing from p_2 to p_1 is

$$t = \int_0^t dt = \int_{p_2}^{p_1} \frac{V dp}{q_{pV,\text{in}}}.$$

Until reaching critical pressure p_2^* , given by Eq. (4.90), that is, from $p_2 < p_2^*$ to p_2^* , the influx of gas remains constant (Sections 4.2.3 and 4.2.4), yielding a simple solution to the partial integral. As soon as the pressure in the vessel rises above the critical pressure p_2^* , the rate of incoming gas drops while the pressure rises. In this pressure range, the integral is solved graphically or numerically.

For the entire range $p_2 < p_2^*$ to $p_1 > p_2^*$, the result for air at $\theta = 20$ °C reads

$$t = 6.42 \times 10^{-2} \frac{V}{d^2} \left\{ \chi \left(\frac{p_1}{p_0} \right) - \chi \left(\frac{p_2}{p_0} \right) \right\} s, \tag{18.16}$$

with t = venting time in s, $V = \text{volume of vented vessel in } \lambda$, $d = \text{diameter of } \lambda$ venting aperture in cm, p_0 = pressure in front of vent valve, usually atmospheric pressure, p_2 = pressure in the vessel prior to opening the vent valve, and p_1 = final venting pressure. Figure 18.15 provides values for the numerically found function $\chi(p_1/p_0)$.

If pressurized air is used for venting instead of atmospheric air, p_0 must be set to the correct pressure existing in front of the vent valve.

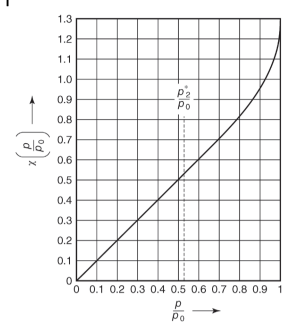


Figure 18.15 Factor $\chi(p/p_0)$ for calculating venting time t (for air at $\theta = 20$ °C). See Eq. (18.16).

18.5 **Techniques for Operating Fine-Vacuum Systems**

18.5.1

Overview

Most industrial vacuum processes operate in the fine (medium) vacuum range, that is, at pressures between 1 and 10^{-3} mbar. As long as working pressures in the vacuum systems remain above $p_{\rm w} \approx 1 \times 10^{-2}$ mbar, outgassing flows are mostly irrelevant. However, they can be disturbing in the deeper fine-vacuum range, that is, at working pressures between 1×10^{-2} and 1×10^{-3} mbar. Manufacturers protect their brand-new metal components against corrosion either with an airtight packaging or with a grease film. Before greased components are mounted in a vacuum system, they should be degreased and subsequently dried thoroughly.

18.5.2

Assembly of Fine-Vacuum Systems

Due to higher tightness demands of fine vacuum, every component must remain below the maximum tolerable leakage rate $q_{\rm L}=10^{-3}\,{\rm mbar}\,\ell\,{\rm s}^{-1}.$ If this threshold value is not met, lower working pressures are producible only with

disproportionately high pumping speeds, that is, with uneconomic pumps. In addition, systems must be kept dry (free of water vapor) and clean. Surfaces exposed to vacuum have to be free of corrosion and must be prepared appropriately, for example, by sandblasting. Each detachable joint is sealed with elastomer seals (Sections 16.2.4.2 and 17.3), and parts moved mechanically must be free of grease when they are inserted into the evacuated volume.

18.5.3

Pumps: Types and Pumping Speeds

Usually, two-stage rotary pumps, dry-running screw-type pumps, Roots pumps, and vapor jet pumps are used for producing medium vacuum. While those twostage, oil-sealed rotary pumps and screw-type pumps available feature relatively low pumping speeds (max. approximately $S_n = 2 \times 10^3 \text{ m}^3 \text{ h}^{-1}$), combinations of Roots pumps and multistage vapor jet pumps offer pumping speeds up to $S \approx 10^5 \,\mathrm{m}^3 (T_\mathrm{n}, p_\mathrm{n}) \,\mathrm{h}^{-1}$, or $S \approx 0.044 \,M_\mathrm{r} \,\mathrm{kg} \,\mathrm{h}^{-1}$ gas (or vapor). Technical and economic considerations determine whether an individual application should employ a Roots pump combination or rather a multistage unit of vapor jet pumps. No general suggestions can be given here. However, the following list of factors should simplify decision making:

- 1) medium or media to be pumped (gases, vapors, or gas-vapor mixtures: composition, and particularly, water vapor contents);
- 2) amounts of these pumped media, available time;
- 3) type and amount of possibly appearing corrosive media;
- 4) temperature of pumped medium;
- 5) starting pressure (usually ambient atmospheric pressure);
- 6) desired working pressure of the pump unit;
- 7) ambient temperature of the pump unit;
- 8) dimensions and weight of the pump unit (including motor), distance between the pump unit and the vacuum system;
- 9) inner volume of the vacuum system;
- 10) leakage rate:
- 11) type and size of connecting flanges on the system;
- 12) power supply voltage and kind of current (three-phase current or DC), tolerable peak (switch-on) current;
- 13) electrical power consumption of the pump unit;
- 14) available motive media (water vapor or oil), vapor costs.

18.5.4

Pressure Measurement

Vacuum gauges for technical fine-vacuum systems mainly include capacitance vacuum gauges, thermal conductivity vacuum gauges, and (fine-vacuum) ionization vacuum gauges (Chapter 13). Section 18.6.2 contains guidelines for mounting measurement tubes as well as for cleaning and contamination prevention.

18.5.5

Pump-Down Time and Ultimate Pressure

In the deeper fine-vacuum range, Eqs. (18.11) and (18.14) are no longer valid for calculating pump-down times because the pumped gas flow from the system $q_{nV \text{ in}}$ now contains high proportions of gases and vapors released by the walls of the apparatus (outgassing). This outgassing flux $q_{pV,out}$ drops with time, and thus, the prerequisite for Eq. (18.11) that $q_{pV,\mathrm{out}} = \mathrm{constant}$ is no longer true. However, compared to the change in gas flow pumped down from the volume of a system, the drop in outgassing flux of technical systems is low. Thus, as a sufficient approximation for the fine-vacuum range, replacing Eq. (18.11), the following simplified equation can be formed for the pressure drop in a recipient or system:

$$p = p_0 \exp\left(-\frac{S_{\text{eff}}}{V}t\right) + p_{\text{ult,P}} + \frac{q'_{pV,\text{in}}}{S_{\text{eff}}} + \frac{q_{pV,\text{out}}(t)}{S_{\text{eff}}}.$$
 (18.17)

The first term in Eq. (18.17) describes the pressure drop with time from starting pressure p_0 (usually atmospheric pressure) down to the pressure p, if only the vessel would have to be evacuated. The ultimate pressure of the pump is $p_{\text{ult.P}}$, and $q'_{pV,\text{in}}$ is the constant flow of incoming gas. $q_{pV,\text{out}}(t)$ is the time-dependent outgassing flux.

For a fair approximation in practical applications, the drop with time of the outgassing flux of metals and glasses as well as ceramics is given by

$$j_{\rm M} = \frac{K_{\rm M}}{t},\tag{18.18}$$

with the surface area-related outgassing flux j_M (outgassing flux density) and the constant $K_{\rm M}$ (subscript "M": metal). The range for the outgassing constant $K_{\rm M}$ is obtained from Figures 16.6 and B.11:

$$K_{\rm M} \approx (4-30) \times 10^{-5} \; {\rm mbar} \; \ell \; {\rm s}^{-1} \; {\rm m}^{-2} \times 3600 \; {\rm s}$$

 $\approx (0.15-1.1) \; {\rm mbar} \; \ell \; {\rm m}^{-2}.$

Thus roughly, the mean value reads

$$K_{\rm M} \approx 0.5 \,\mathrm{mbar} \,\ell\,\mathrm{m}^{-2}.$$
 (18.19)

For $t \to 0$, Eq. (18.18) yields the unrealistic $j_{\rm M} \to \infty$. Surely, the value of $j_{\rm M}$ is finite $(j_{0,M})$ at t=0. A reasonable approximation is obtained when $j_M=j_{0,M}=$ constant is assumed for $t \le t_0$, and Eq. (18.18) is used for $t \ge t_0$.

In plastics (subscript "K"), according to Figures 16.7 and B.11, the outgassing flux drops inversely proportional to the square root of time. Thus, as an approximation,

$$j_{\rm K} = \frac{K_{\rm K}}{\sqrt{t}}.\tag{18.20}$$

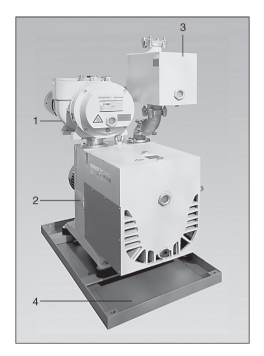


Figure 18.16 Fine-vacuum pump stand for coating technology. 1 Roots pump with canned motor ($S_n = 500 \text{ m}^3 \text{ h}^{-1}$); 2 two-stage sliding vane rotary pump ($S_n = 250 \text{ m}^3 \text{ h}^{-1}$); 3 oil-mist filter; 4 frame with oil tray.

Figure 16.7 provides mean values for the outgassing constant:

$$K_{\rm K} \approx 6 \times 10^{-3} \text{ mbar } \ell \text{ s}^{-1} \text{ m}^{-2} \sqrt{3600 \text{ s}}$$

 $\approx 0.4 \text{ mbar } \ell \text{ s}^{-1/2} \text{ m}^{-2}.$ (18.21a)

For Perbunan:

$$K_K \approx 6 \times 10^{-2} \text{ mbar } \ell \text{ s}^{-1} \text{ m}^{-2} \sqrt{3600 \text{ s}}$$
 (18.21b)
 $\approx 4 \text{ mbar } \ell \text{ s}^{-1/2} \text{ m}^{-2}.$

The starting value $j_{0,\mathrm{K}}$ is treated analogously to that of metals, see comments on Eq. (18.18).

No specific value can be given for t_0 . The correct value is determined by the duration it takes the pump to produce a pressure in the vessel that provides undisturbed outgassing, and by the assessment of outgassing in the previous pressure range.

Thus, the time-dependent outgassing fluxes are described by the following equations:

$$t \le t_0: \quad q_{pV,\text{out,M}}(t) = A_M \frac{K_M}{t_0},$$
 (18.18a)

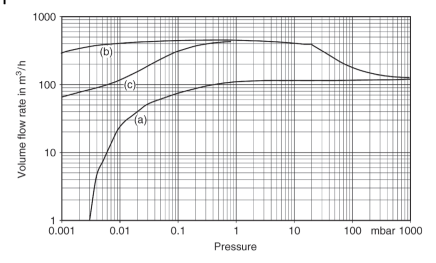


Figure 18.17 Pumping speed curves for the fine-vacuum pump stand in Figure 18.16. (a) Twostage sliding vane rotary pump. (b) Roots pump plus two-stage sliding vane rotary pump. (c) Influence of tube DN 100, $I = 5000 \, \text{mm}$ on configuration (b).

$$t \ge t_0: \quad q_{pV, \text{out,M}}(t) = A_M \frac{K_M}{t},$$
 (18.18b)

and

$$t \le t_0: \quad q_{pV,\text{out,K}}(t) = A_K \frac{K_K}{\sqrt{t_0}},$$
 (18.20a)

$$t \ge t_0: \quad q_{pV, \text{out, K}}(t) = A_K \frac{K_K}{\sqrt{t}},$$
 (18.20b)

with the surface areas of metal and plastics exposed to the vacuum, A_{M} and $A_{
m K}$, respectively. $K_{
m K}$ and $j_{0,
m K}$ are usually far greater than $K_{
m M}$ and $j_{0,
m M}$, respectively, so that, if plastics are present in the vessel, they determine pumping times.

We will now calculate an approximate value for pumping time t_p , applicable in practice. Consider the three processes of (1) pumping down the free gas from the volume, (2) outgassing of gases bound to metals, and (3) outgassing of gases bound to plastics. Each of these processes is initially treated independently to calculate a pumping time assuming that the considered process was singular and stationary. Subsequently, the three values for pumping times are added according to

$$t_{\rm p} = t_{\rm p,1} + t_{\rm p,2} + t_{\rm p,3}. ag{18.22}$$

For this approach, the slowest of the three processes is predominant. For process 1, the first three terms of Eq. (18.17) have to be considered, and thus, the time required for reducing the pressure from p_0 down to pis

$$t_{p,1} = \frac{V}{S_{\text{eff,max}}} \ln \frac{p_0}{p - p_{\text{ult}}},$$
 (18.23)

which uses the ultimate pressure obtained in all three cases:

$$p_{\text{ult}} = p_{\text{ult,P}} + \frac{q'_{pV,\text{in}}}{S_{\text{eff,max}}}.$$
(18.24)

For the second process, terms two, three, and four, the latter in the form of Eq. (18.18b), are relevant, and thus,

$$t_{\rm p,2} = \frac{A_{\rm M}K_{\rm M}}{S_{\rm eff,max}(p - p_{\rm ult})}.$$
 (18.25)

The third process is treated analogously (terms two, three, and four of Eq. (18.17), the latter in the form of Eq. (18.20b)); thus

$$t_{\rm p,3} = \frac{A_{\rm K}^2 K_{\rm K}^2}{S_{\rm eff.\,max}^2 (p - p_{\rm ult})^2}.$$
 (18.26)

 $t_{\rm p,2}$ and $t_{\rm p,3}$ are the times for lowering the pressure to p, if outgassing starts at t = 0 and the outgassing flux is pumped down by the pump continuously (stationary process). Once more, remember that Eqs. (18.23)–(18.26) present rough but practically applicable approximations.

Example 18.6

A steel vessel with volume $V = 2.5 \text{ m}^3$ and the inner surface area $A_{\rm M} = 15 \text{ m}^2$ is evacuated from the pressure $p_0 = 1013$ mbar down to $p_2 = 5 \times 10^{-3}$ mbar using a combination of a trochoidal pump TR (without gas ballast) and a Roots pump RO. Starting pressure of the Roots pump is $p_1 = 20$ mbar. The constant leakage flow shall be $q_L = 0.5 \, \text{mbar} \, \ell \, \text{s}^{-1}$. Evacuation is carried out in two subsequent

- **a.** Evacuation from $p_0 = 1013$ mbar to $p_1 = 20$ mbar using TR only. In this pressure range, the pumping speed of the TR is constant, $S(TR) = 380 \text{ m}^3 \text{ h}^{-1} =$ 105.6 ℓ s⁻¹. Ultimate pressure of the pump $p_{\rm ult,P} = 6.5 \times 10^{-2}$ mbar.
- **b.** Evacuation from $p_1 = 20$ mbar to $p_2 = 5 \times 10^{-3}$ mbar using the combination TR plus RO. In this pressure range, the pumping speed of the combination is practically constant, $S(TR + RO) = 1750 \text{ m}^3 \text{ h}^{-1} = 486 \text{ } \ell \text{ s}^{-1}$. Ultimate pressure of the combination $p_{\text{ult,P}} = 2.5 \times 10^{-3}$ mbar.

The pump unit shall be flange-mounted directly to the vessel, so that $S_{eff,max} =$ S is a close approximation.

a. Step 1. The factor in front of the logarithm in Eq. (18.23), that is, the e-value time (Eq. (18.13)), $\tau = V/S = 2500 \,\ell/(100 \,\ell\,\text{s}^{-1}) = 25 \,\text{s}$. Here, we round down pumping speed to $S = 100 \, \ell \, s^{-1}$ because our equations are rough approximations, that is, calculating with precise values would be senseless. According to Eq. (18.24), ultimate pressure $p_{\rm ult} = 6.5 \times 10^{-2} \, \rm mbar +$ 0.5 mbar ℓ s⁻¹/(100 ℓ s⁻¹) = 7 × 10⁻² mbar. Then, Eq. (18.23) yields

$$t_{\rm p,1} = 25 \, \rm s \, ln \frac{1000 \, mbar}{(20 - 7 \times 10^{-2}) \, mbar} \approx 100 \, \rm s,$$

and with the K_{M} -value from Eqs. (18.19) and (18.25),

$$t_{\rm p,2} = \frac{15 \,\mathrm{m}^2 \times 0.5 \,\mathrm{mbar} \,\ell \,\mathrm{m}^{-2}}{100 \,\ell \,\mathrm{s}^{-1} \times 20 \,\mathrm{mbar}} = 3.8 \times 10^{-3} \,\mathrm{s}.$$

Thus.

$$t_{p,a} \cong 100 \text{ s}.$$

b. Step 2. $S \approx 500 \,\ell \, s^{-1}$, and thus, $\tau = 2500 \,\ell / (500 \,\ell \, s^{-1}) = 5 \,s$. Ultimate pressure $p_{\text{ult}} = 2.5 \times 10^{-3} \text{ mbar} + 0.5 \text{ mbar} \, \ell \, \text{s}^{-1} / 500 \, \ell \, \text{s}^{-1} = 3.5 \times 10^{-3} \, \text{mbar}$, so

$$t_{\rm p,1} = 5 \, \rm s \, ln \frac{20 \, \rm mbar}{(5 - 3.5) \times 10^{-3} \, \rm mbar} \approx 50 \, \rm s$$

and

$$t_{\rm p,2} = \frac{15 \ \text{m}^2 \times 0.5 \ \text{mbar} \ \ell \ \text{m}^{-2}}{500 \ \ell \ \text{s}^{-1} \times 2.5 \times 10^{-3} \ \text{mbar}} \approx 6 \ \text{s}.$$

This yields $t_{\rm p,b} \cong 56$ s.

In both steps, removal of the free gas from the vessel is the predominant process determining pump-down time. Therefore, according to Eq. (18.22), a pumping time

$$t_p = 156 \,\mathrm{s} \approx 2.5 \,\mathrm{min}$$

should be anticipated.

Example 18.7

The same vacuum system as in Example 18.7 is now used for degassing plastic chips with a total surface area $A = 10 \text{ m}^2$. A pretest revealed surface area-related gas emissions of $j_K = 2 \times 10^{-2} \text{ mbar } \ell \text{ s}^{-1} \text{ m}^{-2}$ for the batch after $t = 3 \times 10^3 \text{ s}$. This yields the outgassing constant for the plastic material:

$$K_K = 2 \times 10^{-2} \; \text{mbar} \, \ell \; \text{s}^{-1} \; \text{m}^{-2} \sqrt{3 \times 10^3 \; \text{s}} = 1.1 \; \text{mbar} \, \ell \; \text{s}^{-1/2} \; \text{m}^{-2}.$$

For step 1, the pumping time calculated with Eq. (18.26) amounts to $t_{\rm p,3}=3\times10^{-5}$ s, which is negligible. If again $p_2=5\times10^{-3}$ mbar, then Eq. (18.26) yields a pumping time for step 2 of

$$t_{\rm p,3} = \frac{100~{\rm m}^4 \times 1.2~{\rm mbar}^2~\ell^2~{\rm m}^{-4}~{\rm s}^{-1}}{2.5 \times 10^5~\ell^2~{\rm s}^{-2} \times (2.5 \times 10^{-3}~{\rm mbar})^2} = 77~{\rm s} \approx 1.3~{\rm min}.$$

Thus, charging the vessel with plastic chips prolongs the 2.5 min pumping time calculated in Example 18.7 to approximately 4 min.

In the previous examples, the total leakage rate of the vessel was estimated to $q_{\rm L}=0.5\,{\rm mbar}\,\ell\,{\rm s}^{-1}$. However, when processed, certain plastic materials are extremely sensitive towards oxygen and a leakage rate of this magnitude could already cause oxidation due to the oxygen content of the atmospheric air. It is thus advisable to reduce the leakage rate of the vessel to $q_{\rm L}=10^{-3}\,{\rm mbar}\,\ell\,{\rm s}^{-1}$ in such cases.

18.5.6

Venting

For venting a fine-vacuum apparatus, the same general considerations apply as for low-vacuum systems. Refer to Eq. (18.16) for calculating venting time t.

18.5.7

Fine-vacuum Pump Stands

Ready-for-work pump stands for producing fine vacuum at nominal pumping speeds of several thousand $m^3 h^{-1}$ are designed as multistage pump combinations. Common combinations are: Roots pump(s) with an oil-sealed rotary pump (Figure 18.16), Roots pump(s) with a liquid ring pump (Figure 18.18), screw pumps, multistage water vapor jet pumps compressing up to atmospheric pressure, and multistage water vapor jet pumps with rotary pumps.

Figure 18.16 shows an example of a fine-vacuum pump stand comprising a Roots pump and a two-stage sliding vane rotary pump (Section 7.3.2). The pump stand is used in coating processes where it is exposed to corrosive gases and associated reaction products. Therefore, both pumps are corrosive-gas-protected types and the Roots pump is equipped with a canned motor. Figure 18.17 contains plots of characteristic pumping speeds S versus suction pressure $p_{\rm in}$ for this pump stand. As the example of a 5-m-long DN-100 tubing shows, tubing already has a significant impact on the effective pumping speed at the vessel in the fine-vacuum range.

Figure 18.18 shows a fine-vacuum pump stand for the chemical industry. All of the Roots pumps are protected against pressure shocks and are PTFE-sealed. Construction material is GGG (nodular graphite iron) 40.3 and components are nickel/PTFE-plated. The liquid ring pump is made of ceramic materials, and the separator of graphite. Lowest inlet pressure is 3 mbar. The pumping speed can be adjusted between 1100 and 2300 $\rm m^3~h^{-1}$ via pump frequency control (Figure 18.19).

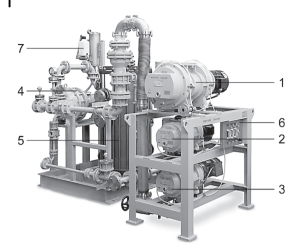


Figure 18.18 Fine-vacuum pump stand for chemical industry. 1 Roots pump $(S_n = 2000 \text{ m}^3 \text{ h}^{-1}); 2 \text{ Roots pump}$ $(S_n = 500 \text{ m}^3 \text{ h}^{-1}); 3 \text{ Roots pump}$ $(S_n = 250 \text{ m}^3 \text{ h}^{-1}); 4 \text{ liquid ring pump}$

 $(S_n = 130 \text{ m}^3 \text{ h}^{-1})$; 5 separator/heat exchanger; 6 seal-gas panel for Roots pumps; 7 sealing medium for floating ring seals. Not shown: flange DN 150 for flushing pumps.

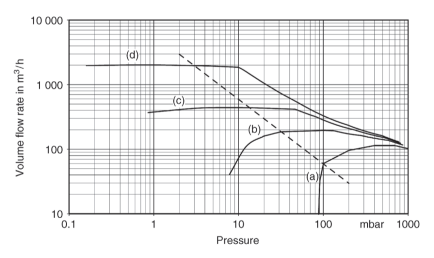


Figure 18.19 Pumping speed curves for the fine-vacuum pump stand in Figure 18.18. (a) Liquid ring pump. (b) Roots pump $(S_n = 250 \text{ m}^3 \text{ h}^{-1})$ plus liquid ring pump. (c) Roots pump $(S_n = 500 \text{ m}^3 \text{ h}^{-1})$ plus Roots

pump ($S_n = 250 \text{ m}^3 \text{ h}^{-1}$) plus liquid ring pump. (d) Roots pump ($S_n = 2000 \text{ m}^3 \text{ h}^{-1}$) plus Roots pump ($S_n = 500 \text{ m}^3 \text{ h}^{-1}$) plus Roots pump $(S_n = 250 \text{ m}^3 \text{ h}^{-1})$ plus liquid ring pump. Dashed line: design condition of pump stand.

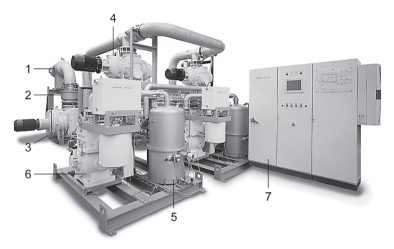


Figure 18.20 Fine-vacuum pump stand for space-simulation chamber. 1 Gas cooler (water-cooled). 2 Cold trap (cooled with liquid nitrogen). 3 Roots pump ($S_n = 25\,000\,\text{m}^3\,\text{h}^{-1}$). 4 Roots pump ($S_n = 6000 \,\mathrm{m}^3 \,\mathrm{h}^{-1}$). Not shown:

Roots pump ($S_n = 2000 \text{ m}^3 \text{ h}^{-1}$). 5 Condenser with automatic condensate drain. 6 Singlestage sliding vane rotary pump $(S_n = 500 \text{ m}^3 \text{ h}^{-1})$ with oil-mist separator DN 160. 7 Control cabinet with touch panel.

Figure 18.20 shows a fine-vacuum pump stand for evacuating a space-simulation chamber for testing satellite propulsion units. The pump stand includes two identical parallel branches that can be interconnected as desired using an SPC. Frequency transformers provide control of pumping speeds for the first stage. The lowest suction pressure is 1 mbar and the maximum pumping speed is 45 000 m³ h⁻¹ (Figure 18.21).

18.6 **Techniques for Operating High-Vacuum Systems**

When operating HV systems, outgassing of the inner surfaces of the apparatus or system is of much more concern than the free gas and vapor in the volume.

To achieve the necessary evacuation time and base pressure, it is necessary to consider the criteria for selection of materials as well as the manufacturing methods and cleaning procedures described in Chapters 16 and 17.

18.6.1

Pumps: Types and Pumping Speeds

The HV range is the domain of turbomolecular and still diffusion pumps. Since outgassing fluxes in high vacuum are made up of hydrogen mainly, pumping times are reduced by adding cryopumps, cooled with liquid nitrogen only.

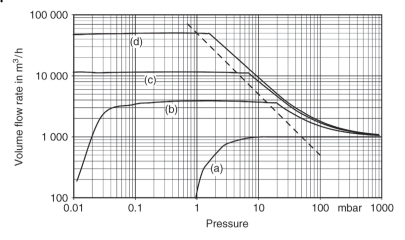


Figure 18.21 Pumping speed curves for the fine-vacuum pump stand in Figure 18.20. (a) Sliding vane rotary pump. (b) Roots pump $(S_n = 2000 \text{ m}^3 \text{ h}^{-1})$ plus sliding vane rotary pump. (c) Roots pump ($S_n = 6000 \text{ m}^3 \text{ h}^{-1}$) plus Roots pump ($S_n = 2000 \,\mathrm{m}^3 \,\mathrm{h}^{-1}$) plus sliding

vane rotary pump. (d) Roots pump $(S_n = 25\,000\,\mathrm{m}^3\,\mathrm{h}^{-1})$ plus Roots pump $(S_n = 6000 \text{ m}^3 \text{ h}^{-1}) \text{ plus Roots pump}$ $(S_n = 2000 \,\mathrm{m}^3 \,\mathrm{h}^{-1})$ plus sliding vane rotary pump. Dashed line: design condition of pump

Setups of pump units in terms of pump types and sizes are determined by

- working pressure;
- pump-down time until working pressure is obtained;
- system volume;
- amount of process gas flow;
- · amount of total leakage rate in the system;
- amount of outgassing fluxes.

If the values of these quantities are mostly unknown, the pumping speed to be installed for a given vessel volume can be approximated using Figure 18.22. The diagram is based on experience values that can only provide a rough guideline. It is applicable to the high and ultrahigh vacuum.

With time, nearly all surfaces in HV applications are subject to increased contamination. This leads to higher outgassing rates, and thus, prolonged pump-down times. Thus, when in doubt, HV pumps with higher pumping speeds are selected. As experience shows, such an approach ultimately saves on time and costs.

18.6.2

Cleaning of Vacuum Gauges

Contaminants and layer of contaminants affect the accuracy, the correctness of measurement, of all vacuum gauges. The sensitivity is changed, the zero stability and the residual reading (offset) are worsened. When in ionization gauges

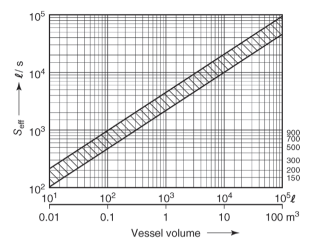


Figure 18.22 Suggested effective pumping speeds (guidance values!) at vacuum vessels for high- and ultrahigh-vacuum operation versus vessel volume.

contaminants cover the electrical insulators of feedthrough, leakage current develop and sparkovers may occur.

Depending on the instrument's value and the level and kind of contamination, it has to be decided if a cleaning or replacement makes sense. In the following, we will give some hints, if a cleaning procedure is preferred.

At all cleaning procedures, it is important to wear fuzz-free gloves and use fuzz-free fabrics and Q-tips.

The priority attempt for the cleaning of mechanical gauges and capacitive diaphragm gauges is to pour in an organic cleaning solvent (e.g., isopropanol). Oil and oily particles can be removed in this way. For best results, the solvent shall flow along the inner wall of the gauge. Then, the solvent shall be distributed within the gauge by gently shaking it (take care of sensitive lever mechanics), after which the solvent is poured out. This procedure has to be repeated until the solvent looks clean and does no more contain any contaminants. A Q-tip may be additionally used with caution to clean the connection tube and filter of a membrane gauge. You can expect success of the cleaning procedure only, if the contaminants can be easily removed. Finally, the vacuum gauge is dried in a dry box at moderate temperature (consider manufacturer's manual) or in a simple vacuum system.

The same described procedure may be applied for a thermal conductivity gauge (Pirani gauge), except that one should abstain from shaking the solvent due to the sensitive, thin heating element (wire).

Ionization gauges with crossed electric and magnetic fields ("cold cathode gauges") are relatively robust and easy to dismantle, so that the parts are well accessible for cleaning. This can be done mechanically by a nonwoven fabric (carded web) with successive application of isopropanol. In some cases (see gauge manual), dilute acids may be used for cleaning.

Ionization gauge with emissive cathodes cannot be efficiently cleaned, except by running the "degas" option or exchange of the cathode.

18.6.3

High-Vacuum Pump Stands

HV pump stands are fully assembled and vacuum-tested units that are provided ready-for-work. Such a stand contains a fore (roughing) pump, a HV pump, forevacuum valves, measurement ports, and sometimes oil-vapor traps, which makes them directly utilizable as HV units. Their compact, space-saving, easy-maintenance design, serial interfaces (RS232/485), and standardized software (e.g., Profibus) guarantee simple operation and integration into existent HV facilities.

HV pump stands are equipped with turbomolecular or diffusion pumps, and are available in different designs and sizes.

As in any other series connection of vacuum pumps, HV pump and fore pump must match so that critical values (e.g., fore-vacuum tolerance) for each pump are not exceeded during operation. This should be checked carefully, particularly after either systems or processes were altered.

18.6.3.1 High-Vacuum Pump Stands with Diffusion Pumps (See Also Section 9.4.6) During operation, the fore-vacuum pressure must remain below the at-rest

pressure of the decelerated vapor jet behind the compression shock (see Sections 9.4.6 and 9.4.7).

The pumping speed of the fore pump must be high enough so that the pressure p_{crit} , given by fore-vacuum tolerance, is not exceeded in any stage of operation. The pumping speed S_F of the fore pump is sufficient if

$$S_{\rm F} \ge \frac{Sp}{p_{\rm crit}} = \frac{\dot{Q}}{p_{\rm crit}},\tag{18.27}$$

with pumping speed S, pressure p, and the throughput \dot{Q} at the vessel. Usually, oil diffusion pumps are not used when inlet pressures are above $p_{in} \approx 10^{-3}$ mbar. Since pumping speed drops at higher pressures, the pumping speed of the fore pump is usually satisfactory if

$$S_{\rm F} \ge \frac{p_{\rm in}}{p_{\rm crit}} S = \frac{10^{-3} \text{ mbar}}{p_{\rm crit}} S = \frac{10^{-3} \text{ mbar}}{2 \times 10^{-1} \text{ mbar}} S = 5 \times 10^{-3} S.$$
 (18.28)

According to Eq. (18.28), a fore pump with low pumping speed S_F is sufficient if inlet pressure p_{in} is low. However, it is generally not advisable to select a fore pump with a pumping speed lower than calculated from Eq. (18.28). Evacuating a vessel always involves pumping-down pressure ranges producing higher inlet pressures. If they rise beyond the limit determined by the maximum tolerable fore-vacuum backing pressure, pumping action of the diffusion pump practically ceases, possibly prolonging pumping time considerably. In addition, sudden gas eruptions occur in most vacuum processes. If the fore-vacuum pump is too

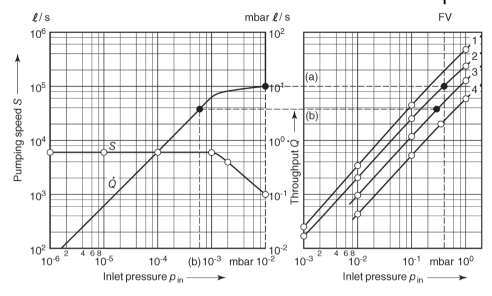


Figure 18.23 Diagram for graphic determination of appropriate fore pumps (see text).

weak, the critical backing pressure $p_{\rm crit}$ is exceeded, pumping action stops, and the erupted gas is only pumped down slowly. Furthermore, when fore-vacuum tolerance is passed, a subsonic vapor jet exits the nozzles thereby increasing oil backflow considerably (Section 9.5).

Diagrams provide a simple means for matching diffusion pumps with roughing pumps. Figure 18.23 exemplifies such matching for a pump combination comprising a $6000\,\ell\,\mathrm{s}^{-1}$ diffusion pump with a two-stage oil-sealed rotary vacuum pump as a fore pump.

The left-hand diagram contains a plot of pumping speed S of the diffusion pump and the derived throughput characteristic \dot{Q} .

The right-hand side shows throughputs 1–4 of commercial two-stage rotary vacuum pumps with $S_n = 135$, 70, 36, and $16 \text{ m}^3 \text{ h}^{-1}$. The dashed vertical line FV indicates the fore-vacuum tolerance of the diffusion pump in the example.

Matching shall be demonstrated for two cases (a) and (b):

- a) (maximum gas flow) $\dot{Q} = 10 \text{ mbar } \ell \text{ s}^{-1}$;
- b) (minimum gas flow) $\dot{Q} = 4 \, \text{mbar} \, \ell \, \text{s}^{-1}$.

In case (a), fore pump 2 should be used in order to comply with FV (see the right-hand diagram).

Sliding vane rotary pumps and rotary plunger pumps are best suitable as fore pumps. They are supplemented with Roots pumps if high pumping speeds are required. In single-stage oil-sealed pumps, pumping speed drops significantly below nominal pumping speed if inlet pressure reaches several 10^{-1} mbar. Thus, it is advisable to use two-stage fore pumps and to run them with gas ballast

during evacuation. Two-stage fore pumps have an additional advantage. For low inlet pressures at the diffusion pump $(p_{\rm in} < 10^{-6}\,{\rm mbar})$, pumping down light gases, helium and nitrogen in particular, requires very low fore-vacuum pressure, considerably lower than fore-vacuum tolerance p_{crit} . For very low inlet pressures and high fore-vacuum pressures, back diffusion of the light gas species opposed to the direction of the vapor jet becomes noticeable and disturbing. At low inlet pressures, two-stage fore pumps produce such low fore-vacuum pressures that back diffusion, even for low species, is harmless.

The smaller fore pump 3 is sufficient for case (b). Analogous considerations apply for matching the turbomolecular pump and the fore pump.

Assembly for valveless operation is the simplest kind of vacuum installation. Here, the diffusion pump and the corresponding fore pump are connected to the vacuum vessel without any valves in between. During evacuation, the vessel is initially pumped down to $p < p_{
m crit}$ with the cold jet pump. Then, the jet pump is heated. After heating, the jet pump continues with the evacuation. Just after heating, the pressure in the vessel rises slightly because gases or vapors are released due to warming. If the time for pump-down from atmospheric pressure to $p_{\rm crit}$ is considerably shorter than the heating time for the jet pump, and if the vessel is tight, then the jet pump is started together with the fore pump. After pumping is finished, the jet pump is shut off and the fore pump continues operating until the jet pump has cooled to a temperature at which the motive medium is not harmed by venting.

If a cooled or LN₂-cooled oil trap is present, the baffle surfaces should reach room temperature before venting because the humidity inherent to atmospheric air would otherwise tend to condense on the baffle surfaces. In this case, the water vapor flow released during subsequent pumping would reduce the effective pumping speed remaining for the vacuum vessel considerably, and might even prevent producing very low pressures within short time.

Cooling water control switches protect against water interruption. In the event of a power failure, the fore pump stops and vents the hot jet pump with the attached recipient. The best protection is achieved by integrating a self-shutting valve in between the fore pump and the jet pump. Combined safety and venting valves simultaneously vent the fore pump.

The fairly long cooling and pump-down times associated with valveless operation are often unacceptable. Therefore, pump stands with jet pumps often feature a bypass line (Figure 18.24).

When operated for the first time, HV valve 9 (large diameter), bypass valve 2, and venting valve 10 are shut. The fore pump evacuates the jet pump via valve 5, and after critical backing pressure p_{crit} is reached, heating of the jet pump is initiated. As soon as heating is completed, the pump stand is ready for operation.

Evacuation is performed according to the following sequence: valves 9 and 10 remain shut. Valve 5 is sealed (now the hot diffusion pump pumps only from volume V of the fore-vacuum line), and subsequently, valve 2 is opened. Using the fore pump, the recipient is now evacuated down to the pressure at which the jet pump starts operating. Now, valve 2 is shut, valves 5 and subsequently 9 are

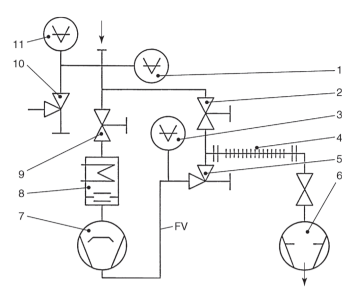


Figure 18.24 HV pump stand with bypass line. Diffusion pump. 8 Vapor trap. 9 HV valve. 1 Fine-vacuum gauge. 2 Valve in bypass line. 3 Fore-vacuum gauge. 4 Metal bellows. 5 Forevacuum valve. 6 Rotary vacuum pump. 7

10 Vent valve. 11 HV gauge. FV Fore-vacuum line (volume V). Note: gauges 1 and 11 can be integrated into a combination gauge.

opened to connect the recipient to the hot, ready-for-work jet pump. In order to decrease pumping time, switching to the jet pump should be performed at the maximum tolerable pressure. This is due to the circumstance that a diffusion pump with pumping speed $S=12\,000\,\ell\,\mathrm{s}^{-1}$ still features a pumping speed $S\gg$ $1000 \,\ell\,\mathrm{s}^{-1}$ at 10^{-2} mbar, whereas pumps usually utilized as fore pumps only yield $S = 180 \text{ m}^3 \text{ h}^{-1} = 50 \, \text{\ell s}^{-1}.$

For the vessel to be vented, valve 9 is shut first, and subsequently, vent valve 10 is opened. While the recipient is vented, the jet pump remains heated and thus operable.

For final shutdown, the first step is to switch off heating of the jet pump, then all valves are shut, and the fore pump is turned off. This pump is then vented via the included safety valve. If the system is not equipped with a safety valve, venting requires an additional valve. In larger jet pumps (above approximately $S \approx 3000 \,\ell\,\mathrm{s}^{-1}$), cooling water supply is maintained for an hour because otherwise the heat capacity of the heating element and boiling chamber would cause excessive heating of the entire pump. Manufacturers' operating instructions should be followed at all times.

In pump stands with bypass line, opening vent valve 10 results in additional venting of the hot diffusion pump if HV valve 9 was not shut previously. This is the most common operator's mistake in manually operated pump stands with bypass lines. Such operating errors are avoided by using remote-controlled, interlocking valves (Figure 18.25). In most cases, these valves are locked by limit

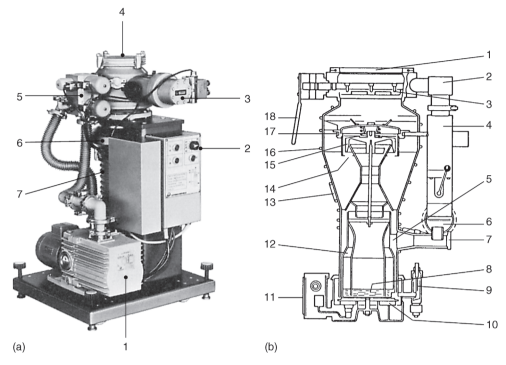


Figure 18.25 (a) HV pump stand on wheels with manually operated valve sequence switch (electropneumatic valves). 1 Two-stage oil-sealed rotary vacuum pump $(S_n = 10 \text{ m}^3 \text{ h}^{-1})$. 2 Sequence switch. 3 HV valve. 4 HV flange. 5 Valve block. 6 Vapor trap. 7 Water-cooled oil diffusion pump $(S_n = 180 \text{ e/s}^{-1})$. (b) Diffstak® compact HV pump stand [13]. 1 Inlet flange. 2 Bypass line.

3 HV valve. 4 Valve in bypass line. 5 Ejector stage. 6 Fore-vacuum connection with fore-vacuum baffle. 7 Thermal switch. 8 Motive medium reservoir. 9 Motive medium drain and refill port. 10 Heating plate. 11 Terminal box. 12 Oil fractionation. 13 Thermal safety switch. 14 Top nozzle. 15 Top nozzle baffle. 16 Pump body with cooling coil. 17 Water-cooled vapor trap. 18 Manual control for HV valve.

switches so that, for example, valve 2 does not open before valves 10, 5, and 9 are shut. Remote-controlled valves shut automatically in the event of power failure, and thus, a pump stand of the type shown in Figure 18.25 is protected sufficiently against mains failure.

All of the important components of the HV system shown in Figure 18.24 can be mounted directly to the diffusion pump, yielding a compact HV pump stand (Figure 18.25a). For operation, it only requires attachment of the fore pump. One of the main advantages of such a compact system is that it reduces the number of seals required. Commercial HV compact pump stands are equipped with diffusion pumps whose nominal pumping speeds S_n lie between 135 and $2000 \,\ell\, \rm s^{-1}$, depending on the type of pump used. The Diffstak® compact HV pump stand (18 in Figure 18.25a) can be designed for manual or pneumatic operation. Connection pieces for vacuum gauges are integrated.

If rough-pumping times, during which the fore-vacuum valve is shut, in a pump stand with bypass line according to Figure 18.24 or 18.25 are very long, volume V of the fore-vacuum line (see Figure 18.24) may not be sufficient to hold the pressure in the fore-vacuum line below the critical pressure p_{crit} of the jet pump. In order to prolong the time until pressure in the fore-vacuum line reaches critical pressure p_{crit} , a so-called fore-vacuum chamber is integrated to increase the volume of the fore-vacuum line (Figure 18.26).

At times, it is inconvenient to run (sometimes large) fore pumps while vacuum vessels are vented. Here also, use of a fore-vacuum chamber is an option, just as using a small so-called holding pump that only keeps pressures below forevacuum tolerance of the jet pump. Occasionally, vibrations produced by the fore pump are disturbing. In this case, the fore pump is shut off and the jet pump pumps the fore-vacuum chamber for some time.

The gas flow exiting the fore-vacuum port of the jet pump determines the size of the fore-vacuum chamber or the holding pump. Even when HV valve 11 is shut (assuming absolute tightness), this gas flow is not constant but is affected by degradation and deterioration of the motive medium due to pumped-down gases and vapors.

For example, shortly after a sudden inrush of air, the gas flow exiting the forevacuum port of the jet pump is greater than normal. However, experience shows that even in large jet pumps, gas flows from fore-vacuum ports are limited to $q_{nV \text{ FV}} = 2 \times 10^{-3} \text{ mbar } \ell \text{ s}^{-1} \text{ if inlets are sealed. Additional load develops from}$ gas released by the walls of the entire fore-vacuum line including the forevacuum chamber as well as leakage gas flow. The gas flow released from the walls of fore-vacuum lines is low since they are usually covered with oil. Likewise, leakage rates are usually far below $q_L = 10^{-3}$ mbar ℓ s⁻¹, and thus, are negligible.

The rise in pressure Δp in a fore-vacuum chamber of volume V within a time Δt

$$\Delta p = \frac{q_{pV,\text{FV}}}{V} \Delta t. \tag{18.29}$$

Using $q_{pV,\mathrm{FV}} = 2 \times 10^{-3} \; \mathrm{mbar} \, \ell \; \mathrm{s}^{-1}$, we calculate a time of approximately $2 \min \ell^{-1}$ volume until the pressure reaches the critical backing pressure $p_{\rm crit} \approx 2 \times 10^{-1}$ mbar. Thus, the fore-vacuum container should provide a volume of approximately $0.5 \ell \text{min}^{-1}$ downtime. If the HV valve is open, the volume of the fore-vacuum chamber must be increased accordingly.

The pumping speed of a holding pump is given by

$$S_{\rm H} = \frac{q_{pV}}{p_{\rm crit}},\tag{18.30}$$

with the gas flow from the fore-vacuum port q_{pV} , and the critical backing pressure $p_{\rm crit}$.

The holding pump is connected via valve 8 (Figure 18.26), which is opened either automatically (using remote control) or by hand after valve 6 is shut.

In most cases, a bimetallic switch is connected to the fore-vacuum chamber that starts the fore pump and opens valve 6 as soon as fore-vacuum tolerance

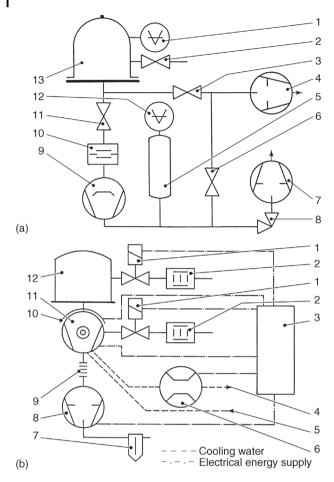


Figure 18.26 (a) HV pump stand with fore-vacuum vessel and holding pump. 1 HV vacuum gauge. 2 Vent valve. 3 Bypass line valve. 4 Fore pump. 5 Fore-vacuum vessel. 6 Fore-vacuum valve. 7 Holding pump. 8 Valve. 9 Diffusion pump. 10 Vapor trap. 11 HV valve. 12 Fore-vacuum gauge. 13 Vacuum vessel. (b) HV pump stand with turbomolecular pump. 1 Vent valve. 2 Sorption trap. 3 Pump-stand control. 4 Cooling water outlet. 5 Cooling water inlet. 6 Cooling water controller. 7 Oil separator. 8 Backing pump. 9 Spring baffles. 10 Heating jacket. 11 Turbomolecular pump. 12 Vacuum gauge. (c) HV pump stand on wheels with turbomolecular pump. Overall dimensions in mm, width \times depth \times height: $392 \times 400 \times 609$. 1 HV connection. 2 Turbomolecular pump $(S_n = 500 \,\mathrm{I \, s^{-1}})$. 3 Pump-stand control unit.

4 Two-stage oil-sealed sliding vane rotary pump $(S_n = 5 \text{ m}^3 \text{ h}^{-1})$. (d) Connections of the pump stand in Figure 18.26b. 1 Switch on/off. 2 Mains connector. 3 Measuring cable feedthrough. 4 Exhaust. 5 Connectors for seal gas and vent gas. 6 Connectors for water cooling. 7 Transport handle. (e) Configurations of a modular-design HV pump stand with turbomolecular pump. (a) Two-stage diaphragm pump ($S_n = 0.9 \,\mathrm{m}^3 \,\mathrm{h}^{-1}$). (b) Twostage diaphragm pump ($S_n = 2.1 \text{ m}^3 \text{ h}^{-1}$). (c) Four-stage diaphragm pump $(S_n = 3.3 \text{ m}^3 \text{ h}^{-1})$. (d) Two-stage sliding vane rotary pump ($S_n = 2.5 \text{ m}^3 \text{ h}^{-1}$). (e) Two-stage sliding vane rotary pump ($S_n = 5 \text{ m}^3 \text{ h}^{-1}$ or 10 m³ h⁻¹). (f) Turbomolecular pump $(S_n = 210 \,\ell\,s^{-1})$. (g) Turbomolecular pump $(S_n = 300 \,\ell \,s^{-1} \text{ or } 500 \,\ell \,s^{-1}).$

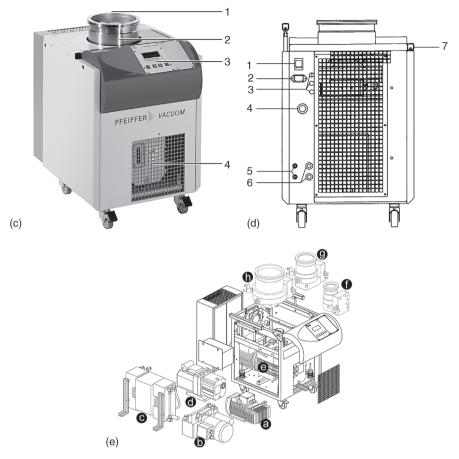


Figure 18.26 (Continued)

 $p_{\rm crit}$ is reached. The same switch closes valve 6 and shuts off the fore pump after a pressure of approximately 10^{-2} mbar is obtained.

18.6.3.2 High-Vacuum Pump Stands with Turbomolecular Pumps

The setup of such a pump stand is simpler than that of a typical HV pump stand with a diffusion pump, also because a bypass line is dispensable. The (usually two-stage) fore pump and the turbomolecular pump can be started simultaneously. Evacuation times are reduced because the turbomolecular pumping action is already effective while the rotor accelerates. Often, an oil-adsorption trap is mounted between fore pump 8 (Figure 18.26a) and spring bellows 9. A diagram analogous to Figure 18.23 helps to select the correct fore pump. Here, considering fore-vacuum tolerance (pressure p_{crit}), the fore-vacuum inlet pressure picked should correspond to the state at which the compression ratio of the turbomolecular pump drops steeply (see, e.g., Figure 10.20).

For producing extremely low pressures (down to the UHV range), not only the vacuum vessel but also the upper part of the turbomolecular pump require baking. For this, heating jacket 10 is integrated.

At one of the positions indicated in Figure 18.26a, vent valve 1 (with preconnected sorption trap 2) is arranged for venting.

Depending on ambient temperatures, HV pump stands with small turbomolecular pumps $(S_n < 500 \,\ell \,\mathrm{s}^{-1})$ either operate on air or water cooling (4, 5, 6). Corresponding threshold temperatures are listed in manufacturers' operating manuals. HV pump stands utilizing large turbomolecular pumps $(S_n > 500 \, \ell \, s^{-1})$ require water cooling.

Figure 18.26b shows an HV pump stand on wheels. The caption lists important technical data and dimensions. Figure 18.26c shows the connections on this pump stand.

Modern HV pump stands feature modular designs (Figure 18.26d) adaptable to individual applications. Turbomolecular pump, controls, and fore pump are selected for the specific purpose. Mounted in a closed, portable case, they provide a pump unit that is ready for connection and fully automatic. Display and keyboard are arranged at the front. All electrical connections are collected at the back (Figure 18.26c).

Serial interfaces provide information regarding operating states and error messages, as well as a means of adjusting operating parameters. An optional remote control can be connected as well.

Sidepieces are detachable providing easy access to individual components, for example, for oil changes on a sliding vane rotary pump or for removing a turbomolecular pump from the case to connect it to a vessel.

18.6.3.3 Fully Automatic High-Vacuum Pump Stands

Pump stands equipped with electromagnetic or electropneumatic valves are controlled fully automatically with pump-stand control units. The unit shown in Figure 18.27 is a microprocessor-controlled pump-stand control unit for HV pump stands. It is universally utilizable for controlling diffusion, turbomolecular, and cryopump stands. Together with control and monitoring equipment integrated into the pump stand, the control unit provides fully automatic execution of predefined programs. The desired program is selected either via buttons on the front plate or via remote control inputs, and executed with respect to necessary safety precautions. The control unit features an 80-digit alphanumeric display providing plaintext dialogues with the instrument. Up to five pressuredependent switching values can be free-programmed with the buttons on the front plate, and the microprocessor control handles the correct corresponding procedure. Possible error messages are indicated directly on the display in plaintext.

The control unit features inputs and outputs, connected in parallel, for remote-controlling procedures and data processing (e.g., monitoring of components' switching statuses, pressure values, etc.). The status of each input or output can be checked at any time, using the display.

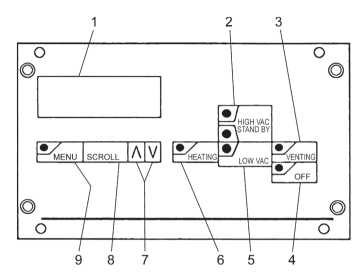


Figure 18.27 HV pump-stand control unit. 1 LCD display, 2 Pushbutton for a HV program and standby. 3 Pushbutton for venting the pump stand. 4 Pushbutton for shutting down pump stand. 5 Pushbutton for fore-vacuum

program. 6 Pushbutton for baking using connected external baking unit. 7 Pushbuttons for cursor control. 8 Pushbutton for scrolling through menus. 9 Pushbutton for menu selection.

Pump stands can be supplemented with an additional control unit (Figure 18.27), for example, for controlling HV valves and processing of pressure gauge tubes' values (0–10 V). A mains supply unit is a further option, able to supply the control unit as well as the corresponding pump-stand components with electrical energy.

Automated HV pump stands, equipped with a diffusion pump, require additional temperature-control equipment. Figure 18.28 shows the main components of a medium-sized, fully automatic HV pump system with a diffusion pump (vacuum vessel not shown).

18.6.4

Pump-Down Time and Venting

With generally sufficient accuracy, pumping time t_p until pressure p is obtained in a HV system is calculated using Eqs. (18.22)-(18.26). However, in Eq. (18.23), S_{eff} must be replaced by the pumping speed S_{extrmF} of the fore pump, and p by the pressure $p_{\rm F}$ at which the HV pump (diffusion or turbomolecular pump) is engaged. Equation (18.23) yields the so-called rough-pumping time.

Outgassing rates heavily influence pumping times in the HV range. Warming of the gas emitting surfaces increases outgassing, and thus, reduces pumping times considerably. Even just warming to $\theta \approx 70$ °C usually increases outgassing rates by approximately an order of 10, and therefore, reduces outgassing times and the corresponding pumping times accordingly. For smaller systems, an industrial blow drier is sufficient for warming to such temperature levels. Larger recipients are heated either with hot water tubing or with electrical heating strips.

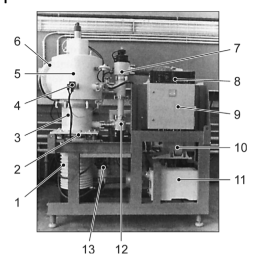


Figure 18.28 Fully automatic HV pump stand with oil diffusion pump. 1 Oil diffusion pump $(S_n = 6000 \, \ell \, \text{s}^{-1})$. 2 Water-cooled oil-vapor trap. 3 Intermediate piece. 4 Vent valve DN 10. 5 Corner valve DN 400. 6 Connecting flange DN 400 to vacuum vessel. 7 Bypass valve

DN 100. 8 Electronic control unit. 9 Power supply unit. 10 Roots pump ($S_n = 500 \, \text{m}^3 \, \text{h}^{-1}$). 11 Single-stage sliding vane rotary pump ($S_n = 160 \, \text{m}^3 \, \text{h}^{-1}$). 12 Corner valve (throttle valve) DN 65. 13 Corner valve (fore-vacuum) DN 65.

As research and manufacturing processes using high-vacuum or ultrahigh-vacuum environments have shown, not only low total pressures but also residual gas constituents of the vacuum play an important role. In many cases, contaminations of the working chambers with hydrocarbons, oxygen, carbon dioxide, carbon monoxide, and methane have negative effects on quality and service lives of vacuum-treated products. Often such contaminations also hamper progress in investigations or make it impossible to acquire reproducible results. Clean vacua with none of the mentioned contaminants are required mainly in semiconductor industry, vapor deposition, and sputter technology, in X-ray tube manufacturing, as well as for fabricating image converter tubes.

18.7 Techniques for Operating Ultrahigh-Vacuum Systems [11]

18.7.1

Overview

The very low gas pressures (i.e., gas densities) in UHV are obtained and maintained only if

- total leakage rates are extremely low;
- outgassing rates are very low;

- negative pump feedback, for example, motive medium backflow (diffusion pumps), is practically zero;
- re-emissions of previously pumped gas (ion getter pumps, cryopumps) are practically zero.

The low outgassing rates are met only if the UHV part of the system, including connected and accompanying components, are bakeable at high temperature $(\vartheta > 100 \,^{\circ}\text{C})$. Thus, UHV technology relies predominantly on stainless steel components (vacuum vessels, valves, tube seals), specially designed gauges, Viton- or metal (Cu, Al, In)-sealed flange connections, ceramic feedthroughs for electrical energy, and special windows. The following components for producing and maintaining UHV are fully or partially bakeable:

- turbomolecular pumps;
- ion getter pumps;
- titanium evaporation pumps (sublimation pumps);
- bulk getter pumps (Non evaporable getter pumps);
- adsorption pumps and dry-running positive displacement pumps (as fore pumps).

As well as combinations of these pumps.

However, utilizing such components is not the only prerequisite for successfully operating at extremely low gas pressures. In fact, many rules and procedures, as covered in this section, should be followed during assembly of a UHV system.

For the pump types listed above, leakage rates and pump feedback can be kept sufficiently low when employing carefully completed permanent and detachable joints.

After one or two hours of baking at approximately 450 °C, and subsequent cooling, outgassing rates of metals drop to approximately 10^{-8} – 10^{-9} mbar ℓ s⁻¹ m⁻² (see also Section 16.3.3). However, baking at $\theta \approx 300$ °C, but for longer periods of time, generally also yields outgassing rates in the range of 10^{-8} mbar ℓ s⁻¹ m⁻². Glasses even provide outgassing rates of only approximately 10^{-10} mbar 10^{-8} s⁻¹ m⁻² after baking for longer periods at 450 °C. As a general guideline, we may summarize:

Outgassing rates drop by an order of 10 per 100 K increase in baking temperature.

18.7.2

Operating Guidelines for UHV Pumps

Rapid production and maintaining of extremely low pressures are not the only aims of UHV technology. In many cases, residual gases must also be free of hydrocarbons. For this reason, UHV production relies nearly completely on oilfree pumping systems.

18.7.2.1 Adsorption Pumps

For large vessels, several adsorption pumps (ASP) should be utilized so that at least one ASP DN 20 per 30 ℓ vessel volume initially reduces vessel pressure from atmospheric pressure down to several mbar. After these saturated ASPs have been shut off from the vessel, a formerly closed valve is opened, connecting to an additional ASP carrying clean adsorbent. With this procedure, pressures below 10^{-2} mbar are produced easily.

Pumping times can be reduced considerably when a dry-running rotary pump is added to the adsorption pumps. This setup is shown in Figure 18.29.

The ultimate pressure producible with ASPs is determined chiefly by those gas species that are present in the vessel when pumping starts (usually atmospheric air) and are difficult or impossible to adsorb such as He and Ne (see also Figure 11.3).

18.7.2.2 Ion Getter Pumps

Ion getter pumps (Section 11.4) are used frequently in UHV systems. In these applications, they are equipped with metal-seal flanges and are bakeable at higher temperatures for reduced self-emissions of gases. Baking temperatures with attached magnet range up to $\theta = 350$ °C, with detached magnet up to $\theta = 450$ °C. Ion getter pumps are often combined with an (integrated) titanium evaporation pump featuring high pumping speed for hydrogen.

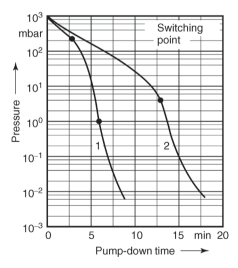


Figure 18.29 Pump-down times for a 180 ℓ vessel using adsorption pumps (ASP) and dry runner (DR, pump with oil-free pumping volume). Pumping intervals for curve 1: 1000 mbar - 200 mbar DR, 200 mbar - 1 mbar

ASP 1, 1 mbar - 10⁻² mbar ASP 2, precooling time of ASP approximately 10 min. Pumping intervals for curve 2: 1000 mbar - 4 mbar ASP 1, 4 mbar -10^{-2} mbar ASP 2, precooling time of ASP approximately 25 min.

Depending on operating conditions, ion getter pumps require cleaning from time to time, and regeneration when getter capability (pumping speed) changes.

Hydrocarbons disturb smooth operation of ion getter pumps. Crack products develop in the gas discharge and on the titanium surface, thus contaminating cathode surfaces and impeding titanium sputtering. Therefore, vacuum systems including ion getter pumps should at least be cleaned with grease-free organic solvents, or preferably, be steam-degreased. Grease-sealed joints (ground-in connections) are disadvantageous as well.

Ion getter pumps contaminated with hydrocarbons (oil vapors, vacuum grease) can be cleaned by baking at $\theta = 300$ °C. Then, released hydrocarbons have to be pumped off with an additional pump while the ion getter pump is not in operation. For subsequent, thorough cleaning of the electrodes, oxygen or air is fed to the system at $p = 1 \times 10^{-6}$ mbar, followed by argon.

If baking does not reestablish the initial pumping speed and ultimate pressure in an ion getter pump, the pump housing, and if cathode mesh is not too worn, the anodes of the electrode system require cleaning. Electrode systems with depleted cathode mesh should be replaced.

When pump housings and electrodes are cleaned, the high-voltage electrical feedthrough should also be checked in terms of insulation. This electrical feedthrough is easily demountable and thus replaceable if necessary.

18.7.2.3 **Titanium Evaporation Pumps**

Titanium evaporation pumps (Section 11.3.3) are used mainly in combinations with ion getter pumps and turbomolecular pumps. The pumps are usually operated intermittently due to their limited titanium reservoir. Evaporation and idle times are adjusted with the supply unit. Evaporation time (typically between 5 s and 5 min) is adjusted prior to conducting the experimental procedure. Idle times depend on pressure and gas load. Too short interruptions of operation may prevent sufficient cooling of the getter screen after vapor deposition. The screen gradually heats, leading to increased gas emissions. Too long interruptions can lead to saturation of the getter layer, and thus, to a loss in pumping action.

For low gas loads, the following reference values are suggested:

Pressure in mbar	Operating interruption		
1×10^{-5}	Several minutes		
1×10^{-7}	Several minutes		
1×10^{-9}	10-30 min		
1×10^{-10}	Several hours		

The heating current of the evaporator determines the rate of evaporation. It can be adjusted in several ways. New evaporator coils release large amounts of gas during first use. Therefore, the electrical current should be raised very gradually during initial operation of an evaporator coil. If the pressure rises too high, the fore pump should be engaged. If several new evaporator coils are employed, they should be degassed back-to-back.

If the producible ultimate pressure rises after longer operating periods, deposited titanium layers have to be removed, involving disassembly of the screen plates. The getter screen and the plates are best cleaned with a wire brush (preferably stainless steel) or via sandblasting.

18.7.2.4 Turbomolecular Pumps

For UHV production, turbomolecular pumps (Chapter 10) are utilized with metal-seal connecting flanges and detachable baking equipment.

Figure 10.21 shows typical pump-down curves for vessel pressures in the UHV. For UHV applications, in particular, note that heavy oil outgassing develops and potentially lasts for several hours during initial pump-down with a fresh-oil filled turbomolecular pump. Therefore, a turbomolecular pump should not be started before the necessary fore-vacuum pressure is obtained. After this, gas bubbles escaping from the oil reservoir no longer influence the UHV.

18.7.2.5 **Cryopumps**

All three types of cryopumps (Section 12.4), bath cryopumps, evaporator cryopumps, and refrigerator cryopumps, are used for producing UHV. Ultimate pressures $p < 10^{-4}$ mbar are obtainable even when pumping hydrogen, and even if the temperature of the cold surface $T > 4.2 \,\mathrm{K}$, as long as the cold surface is covered with a carbon layer working as absorbent. When utilizing bath and evaporator cryopumps, attention must be paid to helium consumption, liquid nitrogen consumption (if used), and particularly to the gas-species-dependent capacity of the pump. UHV cryopumps are either equipped with ConFlat flanges or welded directly onto the recipient.

18.7.2.6 Bulk Getter (NEG) Pumps

Bulk getter (NEG) pumps (Section 11.3.2) are used as the so-called flat getters in research facilities (Section 18.6.7) and industrial applications, for example, electron microscopes. The following example [12] illustrates application of NE flat getters.

Example 18.8

Base pressure (residual gas pressure) of a given research facility shall be 10⁻¹⁰ mbar. This very low pressure can be maintained with a standard vacuum pump stand featuring an effective pumping speed $S_{\rm eff} = 1000 \, \ell \, \rm s^{-1}$, corresponding to 1×10^{-7} mbar ℓ s⁻¹ gas evolution rate of the system. The sputter process requires an argon atmosphere with a pressure of 3×10^{-3} mbar. To keep the pressure at this level when a flow of argon of 40 cm³ (p_n , T_n) per minute is fed to the system, the effective pumping speed of the vacuum pump unit has to be reduced from 1000 to 170 ℓ s⁻¹. Gas emissions of the system are practically unaffected by the relatively thin argon gas flow; thus, throttling the pumping speed leads to an increase of residual gas pressure to 2.4×10^{-9} mbar, that is, residual gas pressure increases by a factor of 24. Adding to this are the contaminants in the argon gas, producing a total contamination of the sputter gas 1.5 orders of 10 higher than tolerable. Utilizing an (additional) NE getter in the immediate vicinity of the wafer (in situ) provides an ideal solution for reducing contamination so that the full pumping speed of the NE getter (not pumping down argon) is available.

18.7.3

Pressure Measurement

Ionization vacuum gauges (Section 13.7) are applicable to pressure measurements in the UHV range [13,14]. Bayard-Alpert gauges are used frequently. Note guidelines given in Sections ecr13.7.3.8 and 13.9 (dissociation at hot cathodes, pumping effect of gauges, ESD, etc.).

18.7.4

Pump-Down Times, Ultimate Pressure, and Evacuating Procedures

Pump-down times of unbaked UHV systems can be approximated using Eqs. (18.22)–(18.26). Pumping times for baked UHV systems depend strongly on the obtainable heating and cooling rates of the considered system, so general suggestions cannot be given here. High heating and cooling rates are producible if involved masses are low. The ultimate pressure producible with an UHV system can be approximated with Eq. (18.24).

Unbaked UHV systems, generally equipped with an additional cryopump, are initially evacuated with a pump stand carrying a turbomolecular pump. The cryopump is engaged subsequently at approximately 10^{-4} mbar.

Bakeable UHV systems featuring getter pumps are baked with the ion getter pump running.

Bakeable UHV systems operating with evaporator pumps are baked with the auxiliary pump running. Afterwards, this pump is turned off. This procedure is followed because it would be difficult, if not impossible, to cope with the outgassing rates during baking using the ion getter pump with its comparably low getter capacity.

An ion getter pump should be engaged at relatively high starting pressure, that is, 1×10^{-2} mbar, if lowest pressures in the range of 10^{-11} mbar are required quickly. Due to the high discharge current produced, temperatures of parts inside the pump rise, and this baking considerably reduces outgassing rates and thus promotes production of lowest pressures.

UHV systems including diffusion pumps are baked together with the oil trap. The latter has to return to normal operating temperature before the vacuum vessel cools because otherwise it would allow considerable amounts of motive medium to pass through. If two oil traps connected in series are reutilized after baking, first the water-operated oil trap closer to the pump is cooled down to the temperature of the cooling water, and subsequently, the second, for example, nitrogen-operated oil trap, while the system is still at baking temperature. The complete system is cooled as soon as the second oil trap has reached full efficiency.

18.7.5

Venting

If an apparatus contains cryopumps or other cooled surfaces, before venting, they have to be heated at least to room temperature. In order to reduce water vapor uptake, UHV systems should be vented with dried air or pure nitrogen.

18.7.6

Ultrahigh-Vacuum Systems

Such systems include low-cost, simple UHV systems used mainly in industrial applications. High-cost UHV systems are employed in accelerators for major elementaryparticle research, featuring one thousand or even more vacuum pumps of various types and the corresponding components used for monitoring and control.

Most of the pumps, measuring equipment, and components used here are covered in the previous sections. Thus, the following section provides a brief roundup with additional comments. Additionally, references listed in Section 18.7 contain special information providing in-depth coverage of the continuously developing state of the art in UHV technology.

18.7.7

Ultrahigh-Vacuum (UHV) Components

UHV components include

- CF flanges (Section 17.3.4.4, Figure 17.10);
- tube pieces, bends, T-, and crosspieces equipped with flanges (Section 17.3.4.7);

- flexible elements (Section 17.5);
- valves (Section 17.7);
- windows (Section 17.6.3.1):
- feedthroughs (Section 17.6).

See corresponding sections for more details.

18.7.8

Ultrahigh-Vacuum (UHV) Pump Stands

Preferred pump combinations for evacuating UHV vessels are listed in Table 18.4. Here, the most important steps in a pump-down procedure are covered considering the UHV pump stand shown in Figure 18.30 whose diagram is shown in Figure 18.31.

The individual stages of evacuation are summarized in Table 18.5. Figure 18.32 shows the corresponding course of falling pressure in the vessel. Pumping times (first column in Table 18.5) apply to evacuation of a clean and empty vessel with open intermediate valve and should be treated as guideline values. For repeated evacuation (charge changes), the intermediate valve is shut first, thereby reducing the volume and yielding shorter rough-pumping times. Table 18.5 comprises the main steps of evacuation omitting detailed descriptions of operations with valves, adsorption traps, and further measures. These should be acquired from manufacturers' manuals.

For baking the apparatus, heating jackets are arranged at appropriate positions. It should be taken care that tolerable temperatures are not exceeded at certain points such as elastomer seals of the intermediate valve and the permanent magnet of the ion getter pump, in the example. In continuous operation, temperatures of elastomer seals and permanent magnets should be limited to

Table 18.4 Common pump combinations for produci	ng ultrahigh vacuum.
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UHV pumps	Fore pumps
Ion getter pump plus (integrated) titanium evaporation pump	Adsorption pump(s) and oil-free rotary vacuum pumps
Turbomolecular pump (and titanium evaporation pump)	Two-stage sliding vane rotary pump and adsorption trap
Turbo-drag pump (turbomolecular pump with <i>Holweck</i> stage) plus Ti evaporation pump	Diaphragm pump
Bath, evaporator, or refrigerator cryopump	Two-stage sliding vane rotary pump or adsorption pump(s) or dry positive displacement pump

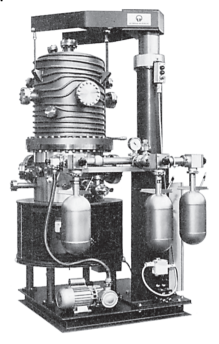


Figure 18.30 UHV pump stand with oil-free pumps. For configuration, see the diagram in Figure 18.31. Pumping speed (depending on type of cooling) 5000 ℓ s⁻¹ or 10 000 ℓ s⁻¹.

150 °C and 380 °C, respectively. Further critical points in UHV apparatuses are vacuum measuring tubes made of glass.

The vessel shown in Figure 18.30 is equipped with a water cooling system for preventing excessive heating of the bell, particularly the usually integrated window, and for reduced gas emissions. High or very high temperatures develop, for example, in annealing furnaces.

For venting a UHV pump stand and UHV systems in general, UHV measuring equipment and ion getter pumps have to be cut off from the recipient prior to venting using an intermediate valve. If a system does not include an intermediate valve, measuring equipment and ion getter pumps must be turned off prior to venting. Any surfaces that are at low temperature when operating (getter screens, cryosurfaces) should reach room temperature before the system is vented in order to prevent water vapor from condensing (formation of ice!).

A common approach for warming the cold surfaces is to the replace the coolants (water, liquid nitrogen) with pressurized air at ambient temperature. Actual flooding should employ dry nitrogen, if available, fed to the system via a gradually opened vent valve. Any vacuum-side surfaces should be exposed to atmospheric air only as shortly as possible.

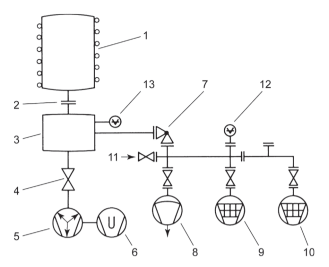


Figure 18.31 Configuration diagram of the UHV pump stand in Figure 18.30. 1 Stainlesssteel vacuum vessel DN 450 with cooling coil, height 450 mm. 2 UHV connecting flange DN 450, 3 Base part, 4 Valve integrated into 3, Vitilan®-sealed. 5 Ion getter pump with seven triode systems. 6 Titanium evaporation pump.

7 Bakeable all-metal corner valve DN 40. 8 Oilfree diaphragm pump with spring-bellows valve. 9 and 10 Adsorption pumps with one spring-bellows valve each. 11 Vent valve. 12 Vacuum gauge for rough and fine vacuum. 13 UHV gauge (built-in design).

18.7.8.1 Large Ultrahigh-Vacuum (UHV) Facilities

Typical large UHV installations are particle accelerators and storage rings. Across the globe, they are operated for elementary particle research. The systems differ according to the application in terms of size, design, and equipment. However, several generally characteristic parameters that are of interest for vacuum technology can be identified [15]. These are discussed in the following section.

The mean free paths of elementary particles need to be high for such particles to travel collision-free across large distances (Section 3.2.5). Only high- and ultrahigh-vacuum environments with low residual-gas pressures provide these conditions. In some systems, this lower threshold value for the pressure is just several 10⁻⁹ mbar.

In principle, the vacuum vessels considered are made up of tubes connected to form rings that can span several kilometers. For operating reasons, these tubes are fairly low in internal diameter (e.g., 100 mm), so that flow conductance (molecular flow) is low. In order to obtain sufficiently low pressures across the vessel, vacuum pump stands are attached all way around the vessel at short intervals. For maintenance and defect localizing, sliding valves are integrated that separate the rings into sectors. Dimensions of such sectors can be up to several hundred meters (see Figure 18.33). Gas emissions from the walls are a key issue because the inner surface areas of the vacuum (steel) tubes, with

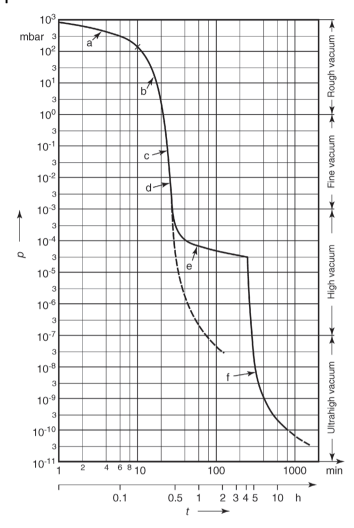


Figure 18.32 Pump-down curve for UHV pump stand according to Figures 18.30 and 18.31.

respect to their volume, are an order of 10 larger than in standard vacuum vessels.

CERN's SPS ring has a circumference of 7 km and comprises 73 sectors of different lengths. Figure 18.33 shows the vacuum-technology equipment of a typical sector [16].

In Figure 18.33 we find:

VR: stainless-steel (1.4429) vacuum tube. VPRS: vacuum pump assembly comprising a turbomolecular pump ($S_{\rm n}=450\,{\rm \ell\ s^{-1}}$) and a two-stage sliding vane rotary pump $(S_n = 35 \text{ m}^3 \text{ h}^{-1})$ for rough-evacuation of vacuum tube VR

Table 18.5 Evacuation procedure for operating the UHV pump stand in Figure 18.30, see also the pump-down curve in Figure 18.32.

Pumping times (min)	Branches of pump-down curve (Figure 18.32)	Pressure changes (mbar)	Pumps in operation	Remarks
000-10	a	1013 → 130	Diaphragm pump or oil- free sliding vane rotary pump	Adsorption pumps cooled previously
010–20	b	0130 → 1	Adsorption pump 1	Valve to dia- phragm pump shut
020-25	С	$0001 \rightarrow 2 \times 10^{-2}$	Adsorption pump 2	Valve to adsorp- tion pump 1 shut
020-30	d	$0002 \times 10^{-2} \rightarrow \approx 10^{-3}$	Ion getter pump	UHV corner valve (fore-vacuum valve) shut
030-250	e	$0010^{-3} \to \text{several } 10^{-5}$	Ion getter pump	Baking time ≈ 3.5 h, baking temperature 350 °C
250-800	f	$0010^{-5} \to \text{several } 10^{-10}$	Ion getter pump plus titanium evaporation pump	Titanium evaporation pump in intermittent operation

down to 10^{-7} mbar. The effective pumping speed at the vacuum tube is only $20 \,\ell \,\mathrm{s}^{-1}$. VPSA: ion getter pump (IG pump) with $S_{\rm n} = 30 \,\ell \,\mathrm{s}^{-1}$ per pump for deeper evacuation of the steel tube. For reduced pumping times, the IG pumps are operated simultaneously with additional titanium evaporation pumps (sublimation pumps), the latter running intermittently. The average operating pressure in the SPS is 3×10^{-9} mbar nitrogen equivalent pressure. This pressure is obtained after a pumping time of approximately two weeks. VGR: thermal-conductivity and glow-cathode gauges (as combined gauges) for measuring pressures in the range $10^3 - 10^{-8}$ mbar. VGC: vacuum gauge in the center of the sector, thermal-conductivity and cold-cathode vacuum gauges with a lower measuring limit of 10⁻¹¹ mbar. In addition, pressures are measured at individual positions throughout the steel tube via electrical current measurements of the IG pumps, providing a lower measuring limit in

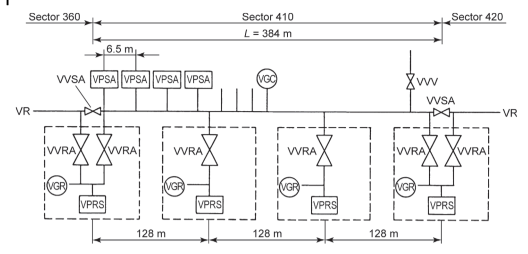


Figure 18.33 Vacuum-technology equipment for a typical sector in CERN's former Super Proton Synchrotron (SPS) [16].

the SPS of 7×10^{-9} mbar. VVRA: fore-vacuum valve DN 70. VVSA: sector cut-off valve DN 100 with indium valve-head seal. The latter is sufficient since the SPS ring is not designed for baking. VVV: vent valve for admittance of dry nitrogen. See [16] for further details.

References

- 1 AS-International Association, www.asinterface.net.
- 2 Wellenreuther, G. and Zastrow, D. (2002) Automatisieren mit SPS, Theorie und Praxis, Vieweg, Braunschweig/Wiesbaden, 2002, ISBN 978-3-528-13910-0.
- 3 Moyne, J.R., Najafi, N., Judd, D., and Stock, A. Analysis of Sensor/Actuator Bus Interoperability Standard Alternatives for Semiconductor Manufacturing, Sensors expo conference proceedings, Celveland, OH, September 1994.
- 4 Profibus International (2002) Profibus Technology and Application, System Description. www.profibus.com.
- 5 Profibus International (2002) ProfiNet Technology and Application, System Description. www.profibus.com/libraries
- 6 SEMI Standards SEMI E30 Generic Model for Communications and Control of

- Manufacturing Equipment (GEM), 2007, http://ams.semi.org/ebusiness/standards
- 7 SEMI Standards SEMI E37 High-Speed Message Services (HSMS) Generic Services, 2008, http://ams.semi.org/ ebusiness/standards.
- 8 L., Erikson, Multi- and megavariate Data Analysis: principles and appliactions, 2001, Umetrics Academy, ISBN 91-973730-1-X.
- 9 Hauer, V. and Day, C. (2009) Conductance modelling of ITER vacuum systems. Fusion Eng. Des., 84, 903-907.
- 10 Pöcheim, N. (1995) Druckregelung in vakuumsystemen. Vakuum in Forschung und Praxis, 1, 39-46.
- 11 Bergandt, E. and Henning, H. (1976) Methoden zur Erzeugung von Ultrahochvakuum. Vak.-Techn., 25 (5), 131-140.
- 12 Briesacher, J. et al. (1990) Non-evaporable getter pumps for semiconductor

- equipment. Ultra Clean Technol., 1, 49-57.
- 13 Tilford, C.R. (1983) Reliability of high vacuum measurements. J. Vac. Sci. Technol., A1 (2), S. 152-162.
- 14 Weston, G.F. (1979) Measurement of ultra-high vacuum. Vacuum, 29 (8/9), 277-292 and 30 (2) (1980), pp. 49-69.
- 15 Pupp, Hartmann (1991) Vakuumtechnik (Chapter 57) Hanser, Munich, Vienna, 558 p.
- 16 Wahl, H. (1989) Das Vakuumsystem des CERN am 450GeV super-protonensynchron und speicherring (SPS). Vakuum in der Praxis, 1, 43-51.

19

Methods of Leak Detection

Werner Große Bley

Inficon GmbH, Bonner Str. 498, 50968, Köln, Germany

This chapter describes how leaks in vacuum systems are identified and localized. Leak detectors are covered in Section 14.4.

19.1 Overview

Not only in research but also increasingly in the industry, tightness of vessels, tubing, components, and packaging are a key issue. Driving forces include environmental protection and the associated legislation as well as competition for ambitious customers calling for highly sophisticated, reliable products. Companies seek to certify their quality management systems, and thus, demand objective measuring methods with testing equipment calibrated traceable (to national standards).

Therefore, rather than developing more and more sensitive leak detectors for smaller and smaller leaks, progress seeks to provide quicker and certifiable procedures for economic survival in the competitive industrial environment. This also explains why tracer-gas leak detection using leak detectors is gaining in importance while historical procedures such as bubble emission techniques or pressure rise/drop are becoming less common. Employing leak detectors correctly and selecting tailored, application-oriented leak testing methods requires thorough knowledge of the fundamental physical principles involved. These shall be covered in the following sections, with a focus on practical applications.

19.1.1

Vacuum Leak Detection

If a vacuum system has a leak and is evacuated via a pumping system with the pumping speed S down to pressure p, small compared to ambient pressure p_0 ,

then a constant leakage flow penetrates through the leak channel. Given as pVflow q_{nV} , it is referred to as $leak\ rate$. For stationary pumping action, q_{pV}, p , and S are related according to Eq. (4.21):

$$p = \frac{q_{pV}}{S}. ag{19.1}$$

For leak detection and leak measurements, a vacuum system can be immersed in tracer gas (or test gas, subscript "T," tracer gas acts as "search gas"), or the tracer gas is sprayed onto the system. Then, the partial pressure

$$p_{\mathrm{T}} = \frac{q_{pV,\mathrm{T}}}{S_{\mathrm{T}}} \tag{19.2}$$

is detected or measured with a tracer-gas-specific detector namely a leak detector.

During operation of a vacuum system, air continuously passes through the leaks. Therefore, the leak rate $q_{pV,T}$ measured with the tracer gas has to be converted to the leak rate of air $q_{vV,
m air}$. In the first step, this calculation requires an assumption regarding the type of flow, based on the pressure conditions and the order of the leak rate. Then, the leak rate can be calculated from specific properties (relative molecular mass, viscosity) of the tracer gas and air (Chapter 4).

Since every vacuum system includes a pump with defined pumping speed, leak detection can be carried out for a system equipped with a partial-pressure gauge (Section 14.2) for measuring tracer-gas pressures. The latter is true in many cases, particularly for ultrahigh vacuum (UHV) systems. Other cases however, especially when testing individual components, utilize a separate high-vacuum pumping system with an integrated mass-spectrometer partial-pressure gauge.

The difference between such leak detectors and the partial-pressure gauges presented in Section 14.2 is that the former measure tracer-gas flows (and not just partial pressures) and that they are designed for a specific tracer gas, preferably helium.

19.1.2

Overpressure Leak Detection

Industrial components are often charged with an overpressure of tracer gas so that tracer gas escaping from possible leaks can be detected. If such a filled specimen is placed inside a vacuum vessel that is connected to a vacuum leak detector, the tracer gas flows into the vacuum and the above considerations for leak rates apply.

However, tracer gas released directly to the environment usually defies measurement of the total emitted gas flow, that is, the leak rate. Here, the developing (distance-dependent) concentration in front of the leak is measured with a sniffing device (Figure 19.1) as a measure for the leak rate. This also allows localizing leaks because the escaping tracer gas forms a stationary *cloud* in front of the leak opening, thus characterizing the location of the leak and to a certain extent allows determination of the leak rate.

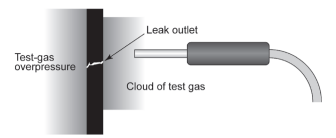


Figure 19.1 Sniffer tip in the test gas cloud in front of a leak opening.

19.1.3

Tracer-Gas Distribution in the Atmosphere in Front of a Leak

According to Fick's law, Eq. (3.102), the tracer-gas flow density j_{leak} for a tracergas partial pressure p^* in the vicinity of a leak opening is

$$j_{\text{leak}} = -D \operatorname{grad}(p^*), \tag{19.3}$$

with the diffusion coefficient D for helium in air. For a leak rate q_{nV} , and since the helium cloud in front of the leak has the shape of a hemisphere with radius r,

$$\frac{q_{pV}}{2\pi r^2} = -D\frac{\mathrm{d}}{\mathrm{d}r}p^*(r). \tag{19.4}$$

Integration of this differential equation yields the following expression for the tracer-gas concentration c(r) at a distance r from the leak at atmospheric pressure p_{atm} :

$$c(r) = \frac{p^*(r)}{p_{\text{atm}}} = \frac{q_{pV}}{2\pi D(r_0 + r)p_{\text{atm}}}.$$
 (19.5)

The radius r_0 is a minimum distance in practice. Setting $c(r_0) = 1$, that is, helium concentration is equal to 100% at distance r_0 , we find for c(r)

$$c(r) = \frac{q_{pV}}{q_{pV} + 2\pi Dr \, p_{\text{atm}}}.$$
 (19.6)

Example 19.1

Helium escapes from a specimen filled with helium at overpressure through a leak and into the ambient atmosphere. The helium leak rate is 1×10^{-6} mbar ℓ s⁻¹, the assumed atmospheric pressure is 100 kPa. For estimating the diffusion coefficient of helium in air, we will use the mean thermal velocity of helium $\overline{c}_{He} = 1245 \, \text{m s}^{-1}$ (at 20 °C) and the mean free path in air

$$\bar{l}_{\text{air}} = \frac{\bar{l}_{\text{N}_2} p}{p_{\text{atm}}} = \frac{5.9 \times 10^{-3} \text{ mPa}}{100 \text{ kPa}} = 5.9 \times 10^{-8} \text{ m}.$$

A practical estimate for the diffusion coefficient of helium in air is then given by the equation for self-diffusion of a gas while putting in the free path of air and the mean thermal velocity of helium:

$$D_{\text{He,air}} = \frac{\overline{c}_{\text{He}}\overline{l}_{\text{air}}}{3} = 2.4 \times 10^{-5} \text{ m}^2 \text{s}^{-1},$$

which yields a helium concentration at distance r = 1 mm from the leak of approximately 2.2 ppm.

This is the concentration measured with a sniffing device calibrated for helium and placed at a distance of 1 mm in front of the leak. At a distance of 10 mm, the concentration amounts to only 0.2 ppm and may be undetectable in industrial environments as soon as background is only slightly unstable.

19.1.4 Measurement Results with the Sniffing Method

Ideally, the sniffer measures the concentration of tracer gas at a particular spot in front of the leak and uses this value as a measure of leak rate. However, in practice this is feasible only under static conditions because a moving sniffing probe would disturb the cloud considerably. In addition, for a given leak rate, the signal indicated by the probe drops rapidly and soon defies reliable measurement as the crosswise velocity at which the probe moves through the cloud increases. The maximum signal is then produced at some considerable distance from the actual leak, as shown in Figure 19.2.

Far more convenient for practical applications are sniffers that aspirate noticeable gas flows. Compared to former types of sniffers utilizing small openings or

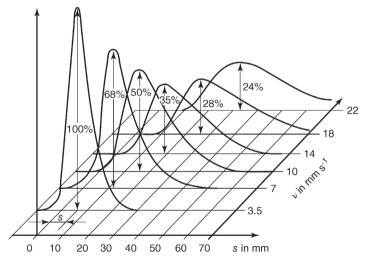


Figure 19.2 Dependence of signal height and apparent leak position s with sniffer tip moved at velocity v (position 0 on the x-axis designates true position of leak).

membranes at the tip for measuring the concentration of the tracer-gas cloud under static conditions very accurately, nearly all sniffers used today feature gas throughputs of $50-200 \,\mathrm{Pa} \,\ell\,\mathrm{s}^{-1}$ (0.5-2 mbar $\ell\,\mathrm{s}^{-1}$). Leak detectors employing the principle of countercurrent flow fostered such developments and tolerate such high gas loads, see Section 14.4.2.5.

Even though the resulting vacuum cleaner effect heavily influences the tracergas cloud, leak localization and quantitative assessment are improved considerably because practically the complete tracer gas escaping from the leak is sucked in, even from locations at some distance, and used for measurement. Quantitative leak rate assessments require calibration with those velocities and distances used in subsequent measurements.

19.1.5

Tracer-Gas Species

The most important requirements for a tracer gas are:

- 1) environmental compatibility;
- 2) chemical and physical inertness (neither reactions nor adsorption);
- 3) low content in standard environments (atmospheric air);
- 4) sensitive and selective detection in vacuum and atmospheric air.

19.1.5.1 Helium

The requirements listed are met largely by noble gases, particularly helium. Helium is absolutely nontoxic; it produces no oxidation, and hardly becomes adsorbed. Since 1998, the European Union has even permitted it as food additive (E939). Its content in atmospheric air is only 5.24 ppm and constant across the globe.

19.1.5.2 Noble Gases Other Than Helium

Of course, noble gases other than helium can also be used for leak detection. However, they either are more prominent constituents of atmospheric air (argon 1%) or are detected in parts of the mass spectrum including lines of the residual gas, particularly hydrocarbons; the latter applies to neon, argon, and krypton. Xenon finally is very expensive and therefore unsuitable as tracer gas, except for very special applications (xenon lamps).

19.1.5.3 **Hydrogen H₂**

Due to its highly explosive nature, hydrogen can be used as tracer gas only when working with forming gas (5% H₂ in N₂); according to ISO 10156, the lower explosion limit is 5.7% H₂ in N₂. In the special case of leak detection on power plant generators, hydrogen used for cooling purposes can be detected directly. Detectors used include sector field mass spectrometers and instruments with semiconductor sensors [1].

To date, hydrogen has not become widely accepted as tracer gas because the low H₂ content in forming gas impairs the detection limit and even slight H₂ concentrations in industrial environments, for example, from electroplating or battery-charging units, cause additional interference.

19.1.5.4 Methane CH₄

The petrochemical industry uses methane as tracer gas for components of chemical processing equipment because methane has similar properties as the substances employed during actual operation of the parts. It is detected with sniffers that use a flame ionization detector (FID) as sensor. This sensor principle is also utilized for leak detection with the organic operating media in field testing. The procedure is standardized in Method 21, Pt. 60, App. A guideline by the US Environmental Protection Agency (EPA).

19.1.5.5 Carbon Dioxide CO₂

Carbon dioxide is unsuitable as tracer gas due to its high and variable content in atmospheric air (exhaled air!). However, its possible applications as coolant for automobile air conditioning spark interests in detection principles that are capable of compensating the variable atmospheric background (e.g., via reference measurements of ambient air).

19.1.5.6 Sulfur Hexafluoride SF₆

Sulfur hexafluoride is used as quenching gas in high-voltage switches. For leak testing, it would therefore be interesting to be able to detect it when it escapes into the environment.

Just recently, the industry has started to use sulfur hexafluoride as tracer gas since photoacoustic infrared absorption allows very sensitive detection (in the ppb range) and because its high molecular weight provides precise leak localization (stable persistence in front of the leak due to low diffusion rate). Leak localizing utilizes either the standard sniffing procedure with a photoacoustic detector, or laser scanning of the specimen with local photoacoustic resonance.

The detection limit for SF₆ is generally good; however, its low diffusion rate in air may possibly prevent reliable localization of hidden leaks. In addition, SF₆ should not be used as tracer gas escaping freely into the atmosphere (or only in low dosages) because, although not ozone harming, it is a highly active greenhouse gas. This is why testing is restricted to overpressure testing (pressurized tracer gas inside the specimen) with highly diluted SF₆ tracer gas. Due to the sensitivity of measurement, even small leaks are detectable.

19.2 Properties of Leaks

19.2.1

Leak Rate, Units

Leak-detection techniques [2] generally use the concept of throughput q_{nV} in pVunits (mbar ℓ s⁻¹) for specifying leak rates q_L (DIN 28402, EN ISO 20484). However, the time-related pV product alone actually only describes the power loss from a volume or pressure change in a gas. Therefore, the gas species and

Unit	mbar ℓ s ⁻¹ $(T_n)^{a)}$	$cm^3(T_n, \boldsymbol{p}_n)s^{-1}$	sccm	Pa ℓ s ⁻¹ (T_n)	Torr ℓ s ⁻¹ (T_n)	kg h ⁻¹ air (20°C)
1 mbar ℓ s ⁻¹ (T_n)	001	000.99	60	100	000.75	4.3×10^{-3}
$1 \mathrm{cm}^3(T_{\mathrm{n}}, p_{\mathrm{n}}) \mathrm{s}^{-1}$	001.01	001	60	101	000.76	4.3×10^{-3}
Sccm	001.68×10^{-2}	001.67×10^{-2}	001	001.68	001.36×10^{-2}	7.77×10^{-5}
1 Pa ℓ s ⁻¹ ($T_{\rm n}$)	010^{-2}	010^{-2}	0.60	1	007.5×10^{-3}	4.3×10^{-5}
1 Torr ℓ s ⁻¹ (T_n)	001.33	001.32	73.5	133	001	5.7×10^{-3}
$1~\text{kg}~\text{h}^{-1}~\text{air}~(20^{\circ}\text{C})$	230	230	12800	23 000	175	1

Table 19.1 Conversion of selected leak rate units for pV flows $q_{\rm L}$ and mass flows $q_{\rm m}$ (see also

temperature T must be considered for the equation of state for an ideal gas (divided by time to give gas flow) to yield the corresponding molar flow rate ν/t (mol s^{-1}) [3]:

$$q_{\rm L} = q_{pV} = \frac{pV}{t} = \frac{\nu}{t}RT = \frac{m}{t} \cdot \frac{RT}{M}.$$
 (19.7)

The number of moles are replaced by $\nu = m/M$ (m = mass, M = molar mass of considered gas species) for calculating mass flow m/t (e.g., g a⁻¹ or kg h⁻¹).

Table 19.1 shows conversion factors for selected pV and mass flow units. The conversion factors refer to a gas at standard conditions, indicated by standard temperature T_n , Eq. (3.6), and standard pressure p_n , Eq. (3.3). Due to the measuring uncertainty involved in practical applications, differences between room temperature and standard temperature are usually negligible when specifying leak rates $q_{\rm L}$.

19.2.2

Types of Leaks

Depending on the predominant flow regime, different types of leaks are differentiated [4]: leaks with viscous gas flow (pores with leak rates $> 10^{-2} \, \mathrm{Pa} \, \ell \, \mathrm{s}^{-1}$ $(10^{-4} \text{ mbar } \ell \text{ s}^{-1})$, with molecular gas flow (pores with leak rates $< 10^{-5} \text{ Pa } \ell \text{ s}^{-1}$ $(10^{-7} \, \mathrm{mbar} \, \ell \, \mathrm{s}^{-1})$, and leaks with gas permeation (through interatomic spaces in elastomers).

The predominant types of flow developing in pore-like leaks depend on the pressure conditions and dimensions of the leak channel. The higher the pressure and the wider the diameter, the more viscous the flow, that is, similar to a thick liquid. The lower the pressure and the narrower the leak channel, the more molecular the flow, that is, gas particles travel independently through the leak. In practice, unevenly shaped leak channels do not show uniform flow conditions

a) (T_n) corresponds to 0.9869 cm³ (T_n, p_n) s⁻¹ of an ideal gas under standard conditions. In practice, it is sufficient to assume 1 mbar ℓ s⁻¹ $(T_n, p_n) = 1$ cm³ (T_n, p_n) . Furthermore, 1 mbar ℓ s⁻¹ $(T_n) =$ $4.41 \times 10^{-5} \text{ mol s}^{-1}$.

but rather viscous flow on the outer atmospheric side and more of a molecular flow on the inner vacuum side.

Permeation leaks do not feature a singular leak channel but are found in joints filled with elastomers (particularly adhesives, seals, etc.) and on the surfaces of plastic parts. Depending on predominant flow regimes, the different types of leaks show specific properties that determine design and procedures of leak testing methods.

Properties of Pore-like Leaks

When investigating pore-like leaks, it should be noted that pressure conditions change dramatically along the leak channels and range from atmospheric pressure down to fractions of a pascal, thus leading to nonuniform flow conditions. If we neglect the problem of turbulent flow for the moment (depending on the microroughness of the leak channel, such flow develops only in very large leaks exceeding approximately 1 Pa ℓ s⁻¹ (10⁻² mbar ℓ s⁻¹), the conditions can always be treated as combinations of initially viscous-laminar flow followed by molecular flow. The length featuring laminar flow depends on the diameter of the leak channel. In very thin leak channels, molecular flow develops nearly throughout the entire channel, as demonstrated in the two following examples.

Example 19.2

We will investigate a cylindrical leak channel with a diameter of $d = 3 \mu m$ and a length of I = 2 mm. The outside is at atmospheric pressure $p_0 = 100$ kPa, and on the inside there is vacuum of $p_i = 0$ Pa. Assuming laminar flow, Eq. (4.94) yields the pV flow through the leak channel $q_{pV} = 2.73 \times 10^{-4} \,\mathrm{Pa}\,\ell\,\mathrm{s}^{-1}$, and Eq. (4.96) gives the critical pressure for choked flow $p_2^* = 100 \,\mathrm{Pa}$. At this pressure, the mean free path of air $\bar{l} = 6.3 \times 10^{-5}$ m = 21 d, so that molecular flow develops for a certain stretch l_2 of the leak channel (starting at pressure p_2).

For air, this molecular flow is given by

$$q_{pV,\text{molec}} = 12.1 \frac{d^3}{l_2} (p_2 - p_i).$$

Along the first stretch of the leak channel (with length l_1), the developing laminar flow

$$q_{pV, \text{ lam}} = 135 \frac{d^4}{I_1} \cdot \frac{p_0^2 - p_2^2}{2}.$$

Setting $q_{pV,lam} = q_{pV,molec}$ and asking the free path at the transition point with pressure p_2 to be equal to the diameter of the leak channel d, we find $p_2 = 2.1 \text{ kPa}, \ l_2 = 0.012 \ l$, and $q_{pV} = 2.8 \times 10^{-4} \ \text{Pa} \ \ell \ \text{s}^{-1}$. Thus, flow is laminar throughout most of the leak channel and changes to molecular only at the very end.

Example 19.3

If we increase the diameter of the leak channel to $d = 10 \,\mu\text{m}$ and reduce its length to l = 1 mm while considering the same pressure conditions as in Example 19.2, then the laminar pV flow amounts to $q_{pV,lam} = 6.75 \times 10^{-2} \, \text{Pa} \, \ell \, \text{s}^{-1}$, and the critical pressure for choked flow $p_2^* = 2.3$ kPa. At this pressure, the mean free path is 2.7 μ m or approximately d/4, so that laminar flow conditions are predominant throughout practically the entire leak channel.

In practice, the leak rate itself can be used for estimating the type of flow because fore pressure is always atmospheric pressure and the vacuum pressure is always below critical pressure. As a rule of thumb, larger leak rates $> 10^{-2} \, \mathrm{Pa} \, \ell \, \mathrm{s}^{-1} \, (10^{-4} \, \mathrm{mbar} \, \ell \, \mathrm{s}^{-1})$ indicate practically complete laminar flow whereas small leak rates $< 10^{-5} \text{ Pa} \, \ell \, \text{s}^{-1} \, (10^{-7} \, \text{mbar} \, \ell \, \text{s}^{-1})$ suggest entirely molecular flow. Conditions are more complicated in the transition range (see Chapter 4); thus, the flow regime cannot be described this simply. As long as either laminar or molecular flow conditions can be assumed, the following equations apply for converting leak rates under different pressure conditions and for different gas species.

In pore-like leaks with higher leak rates and thus laminar flow, the escaping gas flow rises with the square of the pressure gradient. This means that the leak rate depends not only on the pressure difference but also on the actual mean pressure level, that is, the leak rates of two gas species A and B with the input and output pressures p_1, p_2 , and p_3, p_4 , respectively, behave like the differences of the squared pressures:

$$\frac{q_{\rm A}}{q_{\rm B}} = \frac{p_1^2 - p_2^2}{p_3^2 - p_4^2}.\tag{19.8}$$

Thus, raising tracer-gas pressure provides a means of increasing the gas flow overproportionally, with a possible considerable impact on the detection limit of sniffing methods.

The leak rates of two different gas species in a laminar leak behave according to their dynamic viscosities, that is, when the pressures at the input and output are constant, the leak rates of two gas species A and B behave like the viscosities:

$$\frac{q_{\rm A}}{q_{\rm B}} = \frac{\eta_{\rm B}}{\eta_{\rm A}}.\tag{19.9}$$

Since the viscosities of helium and nitrogen differ only by a few percent, the helium leak rate of laminar pore-like leaks can be assumed equal to that of air, for practical considerations! The time of passage of tracer gas through the leak channel is practically negligible because the flow velocity is high. Thus, the response of a leak detector will be immediate if helium is sprayed at such a leak.

In pore-like leaks with smaller leak rates and thus molecular flow, the flow only follows the pressure gradient linearly, that is, the leak rates of two gas species A and B with the input and output pressures p_1, p_2 , and p_3, p_4 , respectively, behave like the differences of the pressures:

$$\frac{q_{\rm A}}{q_{\rm B}} = \frac{p_1 - p_2}{p_3 - p_4}.\tag{19.10}$$

Thus, smaller leak rates at higher pressures also rise only linearly with pressure.

In molecular leaks, the leak rates of two different gas species behave like the square roots of their molar masses, that is, the leak rates of two gas species A and B with the molecular masses M_A and M_B comply with

$$\frac{q_{\rm A}}{q_{\rm B}} = \sqrt{\frac{M_{\rm B}}{M_{\rm A}}}.\tag{19.11}$$

For example, the leak rate for helium exceeds the leak rate of nitrogen or air by a factor of $\sqrt{28/4} = \sqrt{7} \approx 2.6$. This means that for smaller leaks, helium leak detection automatically includes a safety factor of approximately 3 compared to the actual leak rate of air.

As for viscous leaks, the time of passage of tracer gas through a leak channel is also practically negligible under molecular flow conditions because the individual gas particles travel at their thermal velocity (e.g., helium: \overline{c} = 1245 m s⁻¹ at 20 °C). Therefore, even for such small leaks, an attached helium leak detector will respond instantly after spraying, thus simplifying leak localization.

19.2.2.2 Permeation Leaks

At points not representing actual leaks but rather areas permeable for gases, the leak rate q rises linearly with the permeation surface area A and the pressure difference. It drops proportionally to the permeation distance L, Eq. (6.42):

$$q = K_{\text{perm}} \frac{A}{L} (p_1 - p_2). \tag{19.12}$$

The proportionality factor K_{perm} is the permeability and $K_{perm} A/L$ is the permeation conductance (similar to the specific resistance and the resistance of a geometrical body in Ohm's law). Permeation leaks behave similar to molecular leaks in terms of pressure differences. However, their time behavior differs significantly: for tracer gas penetrating through an elastomer, starting time, that is, the time until the permeating gas flow reaches nearly constant values, always spans several seconds in practice. This time increases with the square (!) of the permeation distance L and drops linearly with the material-specific diffusion coefficient D. Introducing the so-called induction period t_i ,

$$t_{\rm i} = \frac{L^2}{2D},\tag{19.13}$$

within which the permeating gas flow reaches 62% of its final value [5]. After $3t_i$, 99% are reached (see also the analogous description of diffusion in gases, Eq. (6.33)). After spraying with tracer gas, therefore, the immediate response is always caused by the real leaks, whereas it takes seconds or minutes until tracergas permeable areas become detectable, for example, cemented joints or seals. This allows clear identification of pore-like leaks even in permeable plastic components by interpreting only the immediate response after tracer-gas application without waiting for the gradually increasing permeation.

Leak rates of different gas species behave like their permeation coefficients, and thus, are highly material specific, which is particularly important for seals. Helium's permeation gas flows through silicone and Teflon are exceptionally high, a reason why these seal materials should be used for vacuum systems only in special cases (e.g., for high temperatures).

19.2.2.3 Virtual Leaks in Vacuum Vessels

A vacuum vessel has virtual leaks if its pressure rises after pumps are shut off but no leaks are detectable. This occurs either when unanticipated out gassing from the walls occurs (water vapor desorption) or a small, gas-filled cavity is present, which releases gas to the vacuum through a pore facing to the inside. Such cavities require long evacuation periods, resulting in long-term gas emissions that appear as leaks. Typical cavities of the kind include blowholes in cast iron or nonevacuated screws in blind holes. Virtual leaks are very difficult to localize once they are there and should therefore be avoided by designing vacuum vessels carefully.

19.2.2.4 Liquid Leaks

After having undergone gas leak detection, industrial applications often call for systems to provide liquid tightness during operation. It can be estimated whether a particular liquid can penetrate a gas leak, or whether the leak is absolutely tight in terms of escaping liquid due to surface tension. (However, evaporating vapor of the liquid may still emanate from the leak. If the leak opens into a vacuum vessel, e.g., a leak in a cold trap cooled with liquid nitrogen, the resulting gas leak is still considerable.)

A leak exposed to a pressure difference Δp is tight toward escaping liquid with a surface tension σ if the diameter d of the leak opening is smaller than d_{max} :

$$d_{\text{max}} = \frac{4\sigma\cos\Phi}{\Delta p},\tag{19.14}$$

where Φ is the so-called wetting angle between the channel wall and the surface of the liquid. For approximations, we can set $\cos \Phi = 1$, representing the worst possible case of the largest possible leak.

Assuming a cylindrical leak channel with diameter d, we can introduce d_{max} to the Hagen-Poiseuille equations (4.81) and (4.82) to calculate the leak rate limit that a liquid-tight leak must provide during gas leak detection when the leak channel is of length l (= wall thickness!) and the tracer gas is of viscosity η :

$$q_{lim} = \frac{\pi}{8} \left(\frac{d_{max}}{2} \right)^4 \frac{1}{l\eta} \Delta p \overline{p}, \tag{19.15}$$

where $\overline{p} = (p_1 + p_2)/2$ is the mean value of the pressures at both ends of the leak channel.

19.3

Overview of Leak-Detection Methods (See also EN 1779)

19.3.1

General Guidelines for Tightness Testing

Sufficiently high pumping speeds of attached pumps would balance the effects of leaks in a vacuum vessel and provide desired total pressures in spite of leaks. However, this approach is not followed in practice because air entering into the vacuum system generally disturbs the running vacuum processes (usually, it is oxygen that produces undesirable reactions and causes damage). Furthermore, it is inefficient to employ extremely large pumps just for pumping down leakage gas.

During leak detection on vacuum vessels (and other specimens), it is important that possible leaks are not clogged with liquids or contaminants. This is why specimens have to be dry and clean, for example, pressurized water testing should not have been performed prior to leak detection. Ideally, leak detection should be performed under the same conditions that are encountered when the specimen is employed in later operation. More precisely, this means that both the direction of the pressure gradient and the temperature should correspond to operating conditions. Leaks undetectable at room temperature can become particularly apparent when material shrinks or expands thermally, as observed for very low (cryostats!) and high temperatures (baking!).

Defining desired tightness values by specifying tolerable leak rate limits is a particularly challenging task for inspection engineers [6]. Basic rules are as follows:

- There is no such thing as absolute tightness, but rather leak rate limits.
- The leak rate limit is not determined by the detection limit of the utilized leak detector but rather by the application and its demands (e.g., service life of an apparatus or product loss).
- Testable leak rate limits must be at least five times as high as the detection limit of the leak detector used.

All of the testing procedures described here use gases for testing, either air or special tracer gas. Liquid tightness and water tightness in particular, are special issues since they additionally involve wettability, that is, surface tension. For many liquids, particularly water with its high surface tension, this means that liquid emissions can be ruled out with additional safety after gas leak detection is passed.

European standard EN 1779 provides a general guideline for selecting appropriate testing methods. Common methods are described in brief and boundary

Standard **English title** EN ISO Nondestructive testing - Terminology - Part 8: Terms used in leak tightness 20484 EN 1518 Nondestructive testing - Leak testing - Characterization of mass spectrometer leak detectors EN 1779 Nondestructive testing - Leak testing - Criteria for method and technique selection EN 1593 Nondestructive testing - Leak testing - Bubble emission techniques EN 13184 Nondestructive testing - Leak testing - Pressure change method EN 13185 Nondestructive testing - Leak testing - Tracer-gas method Nondestructive testing - Leak testing - Calibration of reference leaks for gases EN 13192 EN 13625 Nondestructive testing – Leak test – Guide to the selection of instrumentation for the measurement of gas leakage

Table 19.2 European standards for tightness testing.

conditions and limitations of practical procedures are pointed out. Detailed descriptions are found in the individual standards given in Table 19.2. Particular attention should be paid to standard EN 1330-8, listing all important terms of tightness testing in an English/French/German compilation.

When a component is tested for leaks, the first step should always be to perform an integral tightness test before possible leaks are localized. This approach avoids unnecessary work in terms of leak detection on components that are in fact tight. The procedure is best explained in a flow chart as shown in Figure 19.3. This procedure should not only be followed in industrial mass production but also in laboratory arrangements.

19.3.2 Methods without Tracer Gas (Pressure Testing)

19.3.2.1 Introduction

A number of methods are available for testing the tightness of specimens. The simplest procedures simply analyze the total pressure behavior inside the test object to determine tightness. Here, the specimen is filled with compressed air, that is, overpressure with respect to the ambient air pressure, or pumped down to a certain vacuum pressure. Leak detection improves considerably when tracer gas is utilized instead of air because such a gas is detectable by partial-pressure gauges or special leak detectors. After filling the specimen with tracer gas, submerging the specimen in tracer-gas atmosphere, or local spraying with tracer gas, the total leak rate and locations of possible leaks in a specimen become detectable, independent of any virtual leaks that may affect total pressure.

In vacuum technology, overpressure methods are less widespread due to the request for testing under near to operational conditions. General industrial components, however, are mostly tested by means of overpressure leak detection.

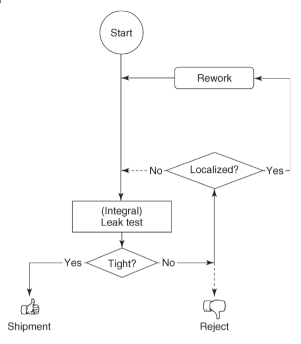


Figure 19.3 Flowchart showing the procedure for leak testing and localization of leaks.

19.3.2.2 Pressure Loss Measurement

Measuring the decline in pressure is the standard industrial mass-testing method. It is not covered here but details are given in standard EN 13184. The method basically measures the pressure loss produced by a leak after a specimen is filled with a certain test pressure. The leak rate q_{nV} is calculated from the known inner volume V and the rate of pressure loss $\Delta p/\Delta t$.

Pressure loss measurements can only identify comparably large leaks, particularly on parts that need to be liquid-tight during later operation (e.g., gear boxes and engines in the automotive industry). In smaller specimens, tolerable leak rates can be as low as 1 Pa ℓ s⁻¹, and reach a maximum of 100 Pa ℓ s⁻¹ in large containers of several m3.

Alternatively, the leak rate can be specified by stating the flowing gas amount required to sustain a certain pressure in the specimen. In this case, the specimen's volume and pressure can be unknown, but must both be constant; this approach is useful, for example, for testing pipelines.

19.3.2.3 Pressure Rise Measurement

Vacuum vessels, particularly very large chambers (chemical reactors), often call for pressure rise measurements for initial integral tightness testing. Generally, an influx of gas q_{pV} raises the pressure (dp/dt) in a sealed volume V:

$$q_{pV}(t) = V \frac{\mathrm{d}p}{\mathrm{d}t}.$$
 (19.16)

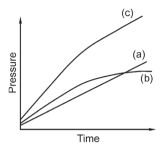


Figure 19.4 Pressure rise due to a leak in a sealed test object: (a) leak rate only; (b) water vapor desorption only; (c) both leakage and desorption.

If the increase in pressure is caused by a leak, the leakage gas flow is constant in time, and thus,

$$q_{pV}(t) = V \frac{\Delta p}{\Delta t} = \text{constant},$$
 (19.17)

yielding a means to calculate the leak rate q_{nV} from a change in pressure Δp during a time interval Δt .

Water vapor desorption from the walls, very noticeable below 10 Pa, is a problem when assessing pressure increase in vacuum vessels. For pure desorption, the pressure in a sealed volume initially rises and subsequently approaches an equilibrium pressure determined by the vapor pressure of water at the prevailing temperature and by the degree of surface coverage on the walls. An additional leakage gas flow affects the pressure conditions so that the pressure now rises nonlinearly, and after a longer period undergoes a transition to linear behavior as shown in Figure 19.4.

When assessing the leak rate of a vacuum system by means of pressure rise measurement, the measurement must prolong until the nonlinear part of the desorption curve is passed, and until the rise in pressure $\Delta p/\Delta t$ is obtained in the linear branch. The main disturbance is caused by temperature changes in the system, producing pressure variations of the same order as a leak. Therefore, it is often beneficial to extend measurement throughout longer periods (hours) and to monitor the courses of pressure and temperature.

Example 19.4

A vessel of volume $V = 2 \ell$ is designated to accommodate a rough-vacuum process. Thus, a diaphragm gauge is available providing reliable readings for $\Delta p = 100$ Pa. The maximum tolerable leak rate is $q_L = 10^{-2}$ Pa ℓ s⁻¹. According to Eq. (19.17), measuring time for determining q_1 is at least $\Delta t = 2 \times 10^4$ s = 5.6 h.

As the example shows, pressure rise measurements under rough vacuum require considerable measuring times when leak rates q_L are low. The process is speeded up by performing the measurement at lower pressures. For example,

keeping all quantities constant but lowering the pressure to 10^{-2} Pa yields a waiting time of only 2 s, assuming the gauge provides a resolution of 1×10^{-2} Pa. However, at such low pressures, gas emissions from the walls disturb measurements considerably. In these cases, several measurements can be performed at selected pressure levels to determine whether out gassing flows occur or whether real leaks exist.

Even though this method is time-consuming and inaccurate, it often represents the only method available for very large systems that defy enclosed integral tests using tracer gas.

19.3.2.4 Additional Methods

The bubble emission method is another type of measuring technique that does not employ any special tracer gas. Here also, we will refer to EN 1593, containing a comprehensive overview of available variants. Bubble emission testing locates leaks using overpressure producing gas bubble formation at the leaks. Leaks are detected either by observing the chains of gas bubbles escaping from the leaks of a submerged specimen, or by locating froth on a component wetted with foaming agent. This method is attractive since there is no doubt concerning the location of a particular leak.

However, the bubble emission method is often overrated in terms of detection limit. Detection of leaks smaller than $10^{-2} \,\mathrm{Pa}\,\ell\,\mathrm{s}^{-1}$, if feasible at all, requires very experienced operators or special liquids. Tightness in a bubble emission test, therefore, at best means water tightness, which however is often sufficient. Note that bubble emission testing is a visual inspection method. Appropriate lighting is essential and personnel's eyesight should be checked regularly. Obviously, the method does not meet the requirements of ISO 9001 quality standards in terms of traceable inspection equipment. Any method fulfilling such standards yields objective leak rates with specifiable measurement uncertainty. Tracer-gas procedures employing leak detectors as described below meet these requirements.

19.3.3

Tracer-Gas Methods without Helium

19.3.3.1 Basics

As indicated previously, the chief benefit of using tracer gas is that real leaks can be differentiated clearly from virtual leaks and that even small leaks are identified without requiring extremely low total pressures. Particularly in industrial testing for small leaks (below simple water tightness), leak testing without tracer gas would lead to unjustifiable testing times since series productions feature typical cycle times per part of only a few seconds.

Apart from dramatically reduced testing times, tracer-gas methods allow precise leak localization. For this, either specimens can be inspected partially or the positions of individual leaks are determined directly. Both approaches are workable in vacuum, overpressure, or sniffing leak detection.

Not all tracer-gas procedures require helium leak detectors; however, detection sensitivity is limited without such detectors. We will now discuss simple tracer-gas methods (without leak detectors) and then go on to tracer-gas procedures with helium leak detectors, which are capable of detecting leak rates down to 10^{-10} Pa ℓ s⁻¹.

19.3.3.2 Vacuum Leak Detection with Non-helium Tracer Gas

In a vacuum system, the simplest detector for a tracer gas other than air is a gasspecies-dependent vacuum gauge such as a thermal conductivity vacuum gauge or an ionization vacuum gauge, which are usually available for controlling the pressure in a system. In both cases, the total pressure reading is determined by the equilibrium pressure produced by the system's pumps depending on desorption gas flows and air influx through leaks. If the air in front of a leak is replaced with tracer gas, the reading on the gauge changes slightly, while the change depends on the difference in sensitivity for tracer gas and the proportion of leakage air.

Example 19.5

A vacuum system shall have a total pressure of 0.1 Pa with an installed pumping speed $S = 10 \,\ell\,\mathrm{s}^{-1}$. For a leak of $q_1 = 0.1\,\mathrm{Pa}\,\ell\,\mathrm{s}^{-1}$, the resulting proportion of the total pressure reading

$$p = \frac{q_L}{S} = \frac{10^{-1} \text{ Pa } \ell \text{ s}^{-1}}{10 \ell \text{ s}^{-1}} = 10^{-2} \text{ Pa}$$

or 10% of the total reading. If the leak is sprayed with helium, then the reading on an ionization vacuum gauge for the leakage gas proportion is reduced by a factor of 7, that is, indicating approximately 0.14×10^{-2} Pa. Thus, the total pressure reading changes from 0.1 Pa to approximately 0.09 Pa. For gas species other than helium, for example, argon or methane (natural gas!), changes are correspondingly smaller.

As this example shows, the change in the reading of the gauge, though clearly visible at careful inspection, defies quantitative assessment. This method only allows detection and localization of large leaks, but no high-quality testing. It is limited to cases where other methods of leak detection are unfeasible.

Employing a real partial-pressure gauge, for example, a quadrupole mass spectrometer, provides a far more sophisticated means of tightness testing on a vacuum system. Such a measuring instrument, also referred to as residual gas analyzer (RGA), shows the gas composition in a so-called residual gas spectrum (see Chapter 14) with the density of ionized gas fragments indicated as ion currents versus the individual mass numbers of particles.

Figure 19.5, for example, shows the mass spectrum of a leaking vacuum system at a total pressure of 2.1×10^{-3} Pa. As the diagram shows, the total pressure is determined mainly by water with a typical group oflines at 16, 17, and

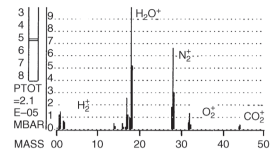


Figure 19.5 Mass spectrum of a leaky vacuum system at a total pressure of 2.1×10^{-3} Pa.

18 amu. Hydrogen ions are located at masses 1 and 2 where mass 1 (H⁺) is mainly an indicator of water vapor and thus features a higher peak than mass 2 (H₂⁺). CO⁺ and CO₂⁺ are located at masses 28 and 44, respectively, but are produced solely by the hot cathode of the mass spectrometer ion source (carbon oxidation). An unmistakable sign of a leak is the presence of oxygen (O₂⁺) on mass 32 in conjunction with a peak on mass 28 approximately four times the intensity of the O₂⁺ peak. Mass 28 indicates the presence of nitrogen (N₂⁺). Since the ratio of the ion currents i^+ on masses 28 and 32 corresponds approximately (except for a 10% deviation because O2 is better ionizable) to the ratio of nitrogen and oxygen in natural air (80%:20% or 4:1), these two mass peaks indicate an air leak. The size of the leak is simple to calculate if the mass spectrometer's sensitivity E_{N_2} for nitrogen (usually provided in technical specifications) and the effective pumping speed S at the vacuum system are known. Then, the leak rate

$$q_{pV} = \frac{1}{0.8} pS = \frac{i^+}{0.8 E_{N_2}} S. \tag{19.18}$$

Of course, the same result would be obtained when interpreting mass 32 using a factor of 1/0.2; however, usually the sensitivity is given only for nitrogen.

Investigating the spectrum, we instantly notice that the total pressure proportion of water vapor is higher than the proportion caused by the leak. Thus, using nothing more than a total pressure gauge, it would be impossible to decide whether a leak or too high desorption gas fluxes, which could possibly be eliminated by baking, cause the problem. Therefore, a residual gas analyzer provides an integral testing method utilizing air as tracer gas, and provides reliable distinction between on the one hand the most prominent disturbing gas water vapor, and on the other, leaks. However, this approach, that is, refraining from utilizing any special tracer gas, does not allow any leak localization.

19.3.3.3 Overpressure Leak Detection with Tracer Gases Other Than Helium

If not air but a different kind of gas is used in pressure testing, leaks can be identified by detecting this particular gas species. Most applications utilize such gas species that also occur as process gases during system operation. Typical gas species include, for example, sulfur hexafluoride (SF₆) in high-voltage switches, methane (CH₄) in chemical valves, refrigerants (Freons, ammonia) in refrigerators and air conditioners. These gases are usually detected with sniffers designed for a particular gas species. Chapter 14 covers the characteristics of sniffing leak detection in more detail. A so-called electron capture detector is usually used for detecting SF₆, a flame-ionizing detector for CH₄, and an alkali ion sensor for refrigerants (see halogen leak detectors in Section 14.4.6.2). Infrared absorption methods are also becoming increasingly popular for detecting these infraredactive gas species.

Utilizing chemical detection is common for ammonia, and at the same time provides a means of leak localization. For this, the specimen covered with ozalid paper (blueprint paper) or ozalid-impregnated swathe is filled with an overpressure of ammonia. This chemical substance blackens at leaks. Waiting times of 12 h might be necessary with a detection limit of up to 10^{-5} Pa ℓ s⁻¹. The procedure calls for safe handling and disposal of the highly corrosive ammonia, which is why this substance has not become widespread as tracer gas.

19.4 **Leak Detection Using Helium Leak Detectors**

19.4.1

Properties of Helium Leak Detectors

In the following, we will refer to as helium leak detectors such instruments that detect and quantitatively measure helium gas flows with a mass spectrometer. Operating principles are covered in detail in Section 14.4. Although a helium leak detector can be used for sniffing methods, its detection principle makes it a vacuum leak detector, that is, the specimen or testing chamber is evacuated. The helium penetrating the vacuum from the environment through leaks is detected. This detection is quantitative and yields a pV flow given in volume flow units, for example, Pa ℓ s⁻¹ or mbar ℓ s⁻¹. Calibrations use reference leaks emitting a precisely defined helium leakage flow. Due to this calibration, which is traceable to national standards, the helium leak detector is the only tightness testing instrument complying fully with the requirements of quality standard ISO 9001. Thus, companies certified to this standard increasingly use such instruments and thereby increasingly replace subjective or qualitative methods, for example, bubble emission techniques.

The most important specifications in terms of practical applications of a helium leak detector (according to standard EN 1518) comprise:

- The leak rate detection limit including the corresponding time constant.
- The helium pumping speed at the inlet, determining the response time for a connected specimen volume and the sensitivity during partial flow operation.
- The maximum tolerable inlet pressure, determining the connection point on the tested system.

The significance of these characteristics is explained in more detail in the following descriptions of procedures.

19.4.2

Testing of Components

Here, we will speak of components as parts that do not include their own vacuum pumping systems. This means that they are stable enough to be evacuated by a leak detector down to a pressure appropriate for leak detection within practicable time. The so-called pump-down time, until the necessary inlet pressure of the leak detector is produced, depends heavily on the condition of the specimen. Any contamination, particularly humidity, oils, and greases, prolongs pumping times due to the emanating vapors that have to be pumped. More than the specimen's volume, it is its entire inner surface area, which is determining. Complicated designs with built-in parts, large surface areas, and difficult to evacuate gaps and dead volumes thus impede leak detection considerably. On a microscopic scale, rough surfaces resemble far greater surface areas than smooth surfaces, therefore prolonging pumping times significantly.

19.4.2.1 Testing Procedure, Integral Testing

Following the flow chart in Figure 19.3, tightness testing generally starts with an integral test in order to avoid the effort of local testing in cases where it is dispensable. Depending on later applications, specimens are either filled with helium overpressure and placed in a vacuum vessel, or they are evacuated and submerged in helium. In both cases, helium will follow the pressure gradient and flow toward the vacuum where it becomes detected by a leak detector.

A simple method of integrally testing a vacuum component attaches the component to the inlet flange of a leak detector, evacuates, and envelopes the component with a soft jacket (the simplest approach uses a plastic bag) filled with helium. In order to prevent measuring permeation flow through the inlet flange and other elastomer seals, it is advisable not to pull the bag over the connecting flange to the leak detector, and to allow only a few seconds for the helium to penetrate before the bag is removed. Any pore-like leaks produce an immediate reaction in the leak detector whereas permeation leaks on cemented joints and seals require seconds or minutes before the ultimate reading is obtained, thus allowing clear differentiation (see the next section). Quantitative readings of leak rates are correct only if two requirements are met:

- 1) The leak detector has been previously calibrated with an (internal or external) reference leak.
- 2) The jacket was filled with 100% helium (air replaced completely).

The indicated leak rate then corresponds to the so-called *standardized leak* rate for a flow from an area at 1 bar into vacuum (see also EN ISO 20484) and requires no further correction. In a similarly simple procedure, components filled with helium overpressure can be tested in a vacuum vessel (e.g., built up of flange components) that is connected to the inlet of a leak detector. After placing the specimen in the vessel, the chamber is evacuated and the indicated leak rate corresponds to the (still uncorrected!) total leak rate of the specimen. It is advisable to measure the empty vessel previously in order to rule out leaks of the vessel itself. However, such leaks are of noticeable impact only in extreme cases since the helium concentration in the vessel's environment is only 5 ppm. In industrial testing, this testing chamber method is usually the method of choice because specimens are usually designed for overpressure.

For converting a result to the standardized leak rate, pressure and concentration corrections of leak rates are required for specimens filled with overpressure. Two corrections are needed:

- 1) Concentration correction: if the tracer gas contains only a percentage x of helium, then the standardized leak rate is obtained by multiplying with 100/x.
- 2) Pressure correction: this initially requires a decision whether flow is laminar viscous or rather molecular. As a rule of thumb, we can assume
 - Laminar viscous flow for measured leak rates exceeding 10^{-2} Pa ℓ s⁻¹.
 - Molecular flow for measured leak rates below 10^{-5} Pa ℓ s⁻¹.

Corresponding to the assessed flow regime, the leak rate is now converted by either dividing by the squared filling pressure (laminar case), or by the filling pressure (molecular case). For intermediate leak rates, the molecular value is taken as a conservative approximation (largest possible standard leak rate).

19.4.2.2 Procedure for Leak Localization

Apart from integral tightness testing, employing tracer gas also allows leak localization. This is feasible if tracer gas is sprayed locally onto the specimen from the outside. The tracer gas spreads rapidly within the vacuum in the specimen thus providing a clear correlation between the spraying action and the reaction of the leak detector. Of course, an important prerequisite for precisely locating a leak is a locally focused tracer-gas load with the highest concentration available. This is applied with an appropriate fine spraying tip that releases only a low amount of helium so that adjacent leaks do not add to the reading of the detector. Since the latter effect cannot be avoided fully in all cases, large leaks in particular should either be sealed beforehand or at least be closed provisionally (e.g., with alcohol that evaporates after some time). Locally restrained testing is also possible by enveloping the area of interest, for example, by sealing with foil.

If a leak on an evacuated specimen volume is sprayed with helium, a leak detector will always take a finite time to indicate the corresponding leak rate or its ultimate value. The course of this indication is determined by the physical mechanism during helium transport in the specimen. Particularly, the total pressure present in the specimen is of considerable impact. In terms of time response, three mechanisms are differentiated: dead time, accommodating time, and starting time.

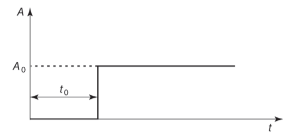


Figure 19.6 Dead time (delay time) with an ultimate leak-rate signal of A_0 .

By its nature, a dead time t_0 is a period occurring practically only during viscous flow. It is given by the ratio of distance l traveled and the velocity of flow ν of a tracer gas in viscous flow. As an approximation,

$$t_0 = \frac{l}{\nu},\tag{19.19}$$

where t_0 is the time passed until a reading is observed, as shown in Figure 19.6.

By its nature, an accommodating time is the period required until a certain volume becomes filled to equilibrium pressure with tracer gas. It is observed in both molecular and viscous flow regimes. The accommodating time τ is the ratio of the specimen volume V and the helium pumping speed S_{He} at the specimen (analogous to an electrical time constant resulting from a capacitor and discharge resistor):

$$\tau = \frac{V}{S_{\text{He}}}.\tag{19.20}$$

The accommodation time (or time constant) according to the above definition is the time required until the reading reaches 63% of its ultimate value (Figure 19.7).

The value 63% (= 1 - 1/e) is a result of the exponentially rising tracer-gas partial pressure in the evacuated specimen. Theoretically, the ultimate reading is approached asymptotically. However, 95% of the ultimate value is obtained after a duration of three time constants.

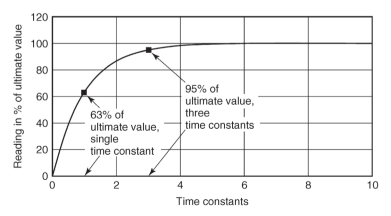


Figure 19.7 Signal rise during leak testing of a vessel with finite volume.

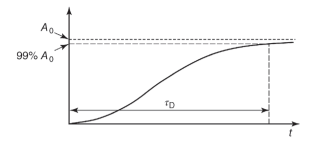


Figure 19.8 Starting time of diffusion for an ultimate leak-rate signal of A_0 .

By its nature, a starting time is the period during which tracer gas diffuses across a certain distance without any flow of the total gas occurring in the system. Diffusion takes place in both molecular and viscous flow regimes. However, noticeable deviations due to diffusion times are observed only at higher pressures (i.e., in the viscous regime). Diffusion of tracer gas develops particularly if the leak detector is connected via a comparably long tube to a region at higher pressure (>1 hPa) and a throttle is integrated just shortly in front of the leak detector.

Although such arrangements should not be used, they cannot be avoided in all cases. If utilized, the gas in the tube is practically stationary and the tracer gas reaches the throttle merely by diffusion, that is, transport occurs only due to partial-pressure gradients.

In this situation, the characteristic starting time is obtained from Eq. (19.13). According to the above definition, the starting time is the time required until the reading reaches 99% of its ultimate value (Figure 19.8). Note that the reading initially rises slower (with a horizontal tangent) than in the case of an accommodation time (with linear initial gradient). In addition, note that the time is determined by the square of the diffusion length L, that is, that short tube lengths are more important than lower pressures!

19.4.3

Testing of Vacuum Systems

Other than components, vacuum systems include their own pumping systems. Although it is possible to use the leak detector instead of the system's pumping unit for maintaining the vacuum in the system (this provides full helium sensitivity since all of the helium passes through the detector), the inlet pumping speed of a leak detector is usually far below the system's pumping speed, generally resulting in unacceptably long response times. However, if the pumping unit of the system operates together with the leak detector, then only part of the helium penetrating into the system volume through leaks actually reaches the detector. This is referred to as partial flow operation.

Partial flow operation is also applied if large containers (not necessarily vacuum vessels) are tested with the vacuum method. Here, additional auxiliary

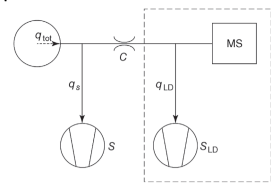


Figure 19.9 Partial flow method with throttled conductance C in the line to the leak detector.

pumps supplement the leak detector to provide the necessary pre-evacuation, and also to yield short response times. For dirty containers, an auxiliary pump is mandatory alone due to the high desorption gas flows that would exceed the capabilities of a leak detector.

19.4.3.1 General Considerations for Partial Flow Operation

Figure 19.9 shows the arrangement used for partial flow operation. The total leakage gas flow $q_{\rm leak} = q_{\rm tot}$ from the system separates into the partial flow $q_{\rm S}$ into the system's pump with pumping speed S and the partial flow $q_{\rm LD}$ into the leak detector with pumping speed $S_{\rm LD}$. The leak detector's partial flow $q_{\rm LD}$ is indicated and therefore has to be calculated from the pumping speed ratios. The following two equations apply to the gas flows for a given system pressure $p_{\rm Sys}$:

$$q_{\text{tot}} = (C + S)p_{\text{Sys}}$$
 (a) and $q_{\text{LD}} = Cp_{\text{Sys}}$ (b). (19.21)

Eliminating system pressure p_{Sys} yields

$$q_{\rm LD} = \frac{C}{C + S} q_{\rm tot} = \gamma q_{\rm tot}, \tag{19.22}$$

with the so-called partial flow factor γ characterizing the loss in sensitivity of this operating mode compared to direct connection of a leak detector. Usually, the conductance C of the throttle is low compared to the system's pumping speed S so that $\gamma = C/S$ can be assumed in most cases.

Throttle C is required only if system pressure $p_{\rm Sys}$ is higher than the tolerable inlet pressure of the leak detector because the effective inlet pressure for the leak detector behind the throttle is reduced by a factor $C/S_{\rm LD}$ compared to system pressure.

The above equation can only provide an approximation for the partial flow factor γ during the design phase of planned leak detection. For a quantitative interpretation of measured leak rates in practice, γ is obtained by connecting a defined reference leak to the system. The partial flow factor γ is then acquired by simply dividing the measured and the correct leak rates of the reference leak.

In modern leak detectors, the partial flow factor is determined automatically and stored during calibration with an external system leak so that subsequent measurements yield correct leak rates.

19.4.3.2 Points on Systems for Connecting Leak Detectors

The pumping speed of a vacuum system's pumps is usually high enough to produce practicable response times in partial flow operation. However, for leak-detection sensitivity to be high when response times are low, it is important to select an appropriate connection point for the leak detector to be attached to the system. Generally, three variants for connecting a leak detector to a vacuum system are available, each featuring a characteristic partial flow factor:

- In the high or working vacuum of the process vessel.
- In the fore vacuum of the pumping system.
- In the exhaust line of the pumping system.

Figure 19.10 shows these positions. While the pressure in the high or working vacuum (position a) is always low enough for the leak detector, the other two positions may require throttles in order to limit the inlet pressure to the leak detector. However, modern leak detectors usually provide sufficiently high inlet pressures for them to be connected to the fore vacuum (position b). Therefore, this generally is the connection type of choice and is often already provided by the manufacturer who integrates appropriate testing flanges.

If system pressure is above the maximum tolerable inlet pressure of the leak detector, the leak detector is not attached directly but via a throttle valve. By

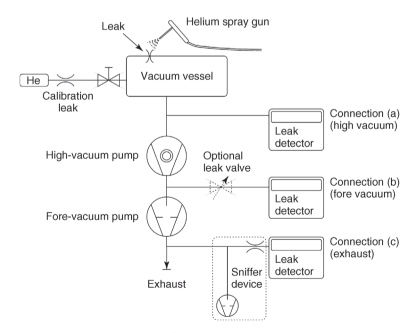


Figure 19.10 Connection points of leak detector when checking a system for leaks.

adjusting the valve opening, the pressure in the leak detector can be selected so that maximum sensitivity for a given pressure is obtained.

If connecting positions are available neither in the intermediate vacuum nor in the fore vacuum, or if pressures are even too high for throttle valves, then there is still the possibility of employing a sniffer at the system's exhaust (position c). Instruments best suitable feature their own small pump delivering gas from the exhaust to the leak detector. Although this produces an additional partial flow, the method allows leak detection in cases prohibiting any other type of connection to the system (e.g., power plant condensers).

If system pressure is low ($< 10^{-2} \text{ Pa}$) a leak detector could be connected without any negative impact on the detector; however, the system could then become contaminated due to oil backstreaming from the leak detector pumps.

Here it is advisable to connect the leak detector to the fore vacuum line, thereby using a connection point usually providing a beneficial partial flow factor because the pumping speed of the fore-vacuum pump is comparable to that of the leak detector. If this approach is not feasible, for example, in UHV systems, because there is no appropriate connection available or because a cryo or getter pump is used to evacuate the system, then the following measures represent alternative approaches:

- Place a cold trap or adsorption trap in front of the leak detector.
- Use counterflow leak detector with turbomolecular pump at inlet.
- Employ quadrupole mass spectrometer (at UHV with electron multiplier).

Working with throttles in partial flow operation is accompanied by corresponding sensitivity losses since only a small portion of the leakage gas flow actually passes through the leak detector. In order to assess detectable leak rates and the associated response times, it is essential to connect a reference leak featuring the desired order of leak rate at a position most remote to the leak detector. This procedure yields a reading and the response time from the moment the leak is closed until a certain residual reading is obtained, which are the chief parameters used for assessing a leak-detection method.

The following measures can be taken if sensitivities and/or response time are unsatisfactory:

Insufficient sensitivity:

- Reduce partial flow ratio in order to feed more gas to the leak detector.
- Connect leak detector at a position with a lower pumping speed of the system (fore vacuum, exhaust).
- Reduce pumping speed of system if the pressure is not too high (possibly down to zero; then, the entire gas flows through the leak detector).
- Increase pumping speed of the leak detector (open throttle valve further or remove throttle valve).
- Response time too long.
- · Reduce dead time by reducing the distance between the leak detector and a supposed leak.

- Reduce accommodating time by mounting the leak detector at a position where V/S is lower (e.g., fore vacuum!).
- Reduce starting time by arranging the inlet throttle in the middle of the gas flow (probe) and not at the walls of a tube or vessel.

Dirty System

Systems tested by means of vacuum technique but not designed as vacuum systems show the problem of leak detector contamination caused by the system. The tolerance of the leak detector's pumps toward dust, vapors, and corrosive gases is limited. Appropriate countermeasures have to be selected specifically for individual applications; however, helpful measures include:

- Utilize counterflow leak detector.
- Arrange a cold trap in front of the leak detector.
- Employ condensate separator and dust filter in front of leak detector.

19.4.3.3 Detection Limit and Response Time

Quick response times represent a main concern in local leak detection because only short times provide a clear correlation between spraying action and the signal response, and thus, provide reliable localization. Therefore, leak detectors should feature a high inlet pumping speed for helium (S_{He}) because, as we know, the response time constant $\tau = V/S_{He}$, with V denoting system volume. If response times reach several seconds, spraying durations are usually far below the time constant. Figure 19.11 shows the consequences: it appears as if the

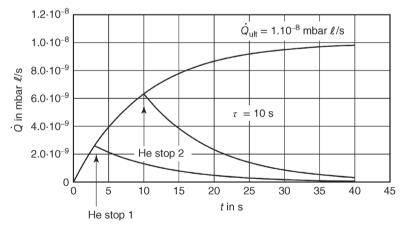


Figure 19.11 Signal rise until ultimate signal is produced after spraying helium. The rise comes to an end at a reduced level if the spraying is terminated too early: termination of spraying after 3 s (with a time constant of $\tau = 10 \text{ s}$) results in a reading of only 2.7 \times 10^{-9} mbar ℓ s⁻¹ (2.7 × 10^{-7} Pa ℓ s⁻¹) for a leak

of 1×10^{-8} mbar ℓ s⁻¹ (1×10^{-6} Pa ℓ s⁻¹). Spraying for the duration of one time constant yields 63% of the ultimate value or $6.3 \times$ 10^{-9} mbar ℓ s⁻¹ (6.3 × 10^{-7} Pa ℓ s⁻¹). Only after a period of three time constants or 30 s of spraying, the deviation from the true final value drops to 5%, which is acceptable.

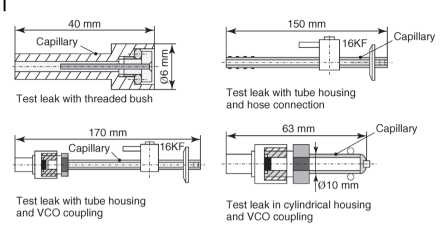


Figure 19.12 Industrial test leak designs suitable to be sprayed with helium directly. Such leaks are customized to the required threshold leak rate (INFICON GmbH, Cologne, Germany).

The element determining the leak rate is a glass capillary mounted in a small metal ferrule and cemented into the housing as chosen by the customer.

sensitivity was reduced because helium is not available long enough in order to provide full signal rise up to the ultimate signal. By definition, 63% of the signal is produced if spraying extends for only one time constant (He stop 2), and even far lower readings if spraying time is reduced further (e.g., He stop 1). If the time constant of the leak-detection arrangement is known, it is advisable to spray for approximately three time constants in order to produce approximately 95% of the ultimate reading.

This behavior can also be checked by equipping the system with a reference leak. However, note that the leak should be designed to be sprayed with helium as the types shown in Figure 19.12 (pure calibration leaks often include a large dead volume in front of the leak inlet that prolongs response times).

For determining the time constant of a vacuum system with a reference leak (in this case, a leak with a gas reservoir for constant leak rate), a novice would probably simply open the valve in front of the reference leak and interpret the exponential rise of the leak rate signal. In practice, however, the reverse procedure is recommended because this avoids signal errors due to helium accumulating in front of the closed reference leak. The reference leak rate of the system is observed for some time (at best with a chart recorder) until it stabilizes, and subsequently, the leak is shut as quickly as possible. The declining signal follows the same exponential law as the increase and can be interpreted in a similar way: after the time τ , the signal drops from equilibrium by 63% (i.e., down to 37%).

19.4.4

Overpressure (Sniffing) Leak Detection with a Helium Leak Detector

Helium as tracer gas also allows sniffer testing of components that are not filled with gas or that are filled with an undetectable gas. This occurs, for example, in

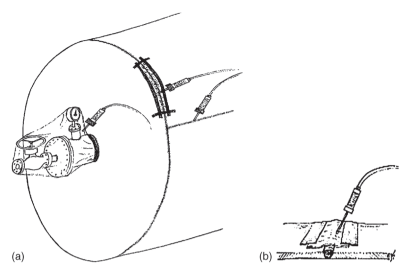


Figure 19.13 (a) Enveloping in a plastic bag or with film and adhesive tape and subsequent local sniffing of areas on a helium-filled heat exchanger. (b) Testing of a weld seam by inserting the sniffer tip into the volume separated by film and adhesive tape.

pretesting or in reworking of leaking components. In both cases, specimens are filled with helium overpressure, and a helium sniffer is used to detect escaping helium.

19.4.4.1 Integral Procedure (Total or Partial)

In integral sniffing procedures, either the specimen is enclosed in a soft envelope (total testing) or parts are enclosed or sealed with foil as shown in Figure 19.13.

This method detects the rising concentration in the envelope or the sealed part of the component a reasonable time after the test object was filled with helium. Initially, the method provides only a pure tightness test and does not allow quantitative assessment of leak rates. However, under atmospheric pressure $p_{\rm atm}$, for known envelope volume $V_{\rm H}$, and after a recorded waiting time Δt , the leak rate $q_{\rm He}$ leading to the measured increase in partial pressure Δp_{He} or in concentration Δc_{He} can be calculated. The leak rate is as follows:

$$q_{\rm He} = V_{\rm H} \frac{\Delta p_{\rm He}}{\Delta t} = V_{\rm H} p_{\rm atm} \frac{\Delta c_{\rm He}}{\Delta t}.$$
 (19.23)

The increase in helium concentration $\Delta c_{\rm He}$ is found simply by comparison with the natural helium concentration in air:

$$\frac{\Delta c_{\text{He}}}{5 \times 10^{-6}} = \frac{I_{\text{H}} - I_{\text{5 ppm}}}{I_{\text{5 ppm}}}.$$
 (19.24)

For calculating Eq. (19.24), the reading is acquired for atmospheric helium $I_{\rm 5\;ppm}$ as well as for the concentration in the envelope $I_{\rm H}$, after waiting time Δt .

19.4.4.2 Leak Localization with a Sniffer

For localizing a leak, one can utilize the circumstance that helium escaping into the atmosphere forms a stationary diffusion cloud in front of the leak (see section 19.1.3), which can be detected with a sniffing probe (Section 19.1.2). Leak localization by means of sniffing is the most widespread method for reworking faulty components in industrial leak detection because most components contain overpressure during operation, and thus, are tested under the same conditions.

In leak localization with a sniffing probe, sniffing velocity is particularly important because the inevitable response time of the sniffing equipment causes a virtual offset in the leak's position (Figure 19.2). The higher the velocity of the sniffing probe, the higher this offset. Thus, a low sniffing velocity is beneficial for precise leak localization, but requires a considerable amount of extra time. Therefore, note that here also an integral tightness test should always precede leak localization procedures (see Figure 19.3).

19.5 Leak Detection with Other Tracer Gases

19.5.1

Basics

If not using helium as tracer gas, the vacuum method is not practicable for tightness testing. Although a mass spectrometer in principle is capable of detecting any type of tracer gas, vacuum tightness testing requires separation of tracer gas and residual gas, particularly water vapor, in order to produce low evacuating times.

As known, this is done in a cold trap with liquid nitrogen when using a direct flow leak detector and by utilizing the differing compression of the turbo pump when employing a counterflow leak detector (Section 14.4.2.5). Both methods are inapplicable to condensable gas species and gases with higher molecular weights. Therefore, any tracer gas other than helium is always detected by means of sniffing.

Sniffing Leak Detection on Refrigerators and Air Conditioners

Tightness testing of refrigerators and air conditioners is a particularly important aspect of industry and craft. Industry, in particular, has to guarantee very low refrigerant losses in the order of several grams per year (g a⁻¹) since filling amounts down to 20 g or 30 g in refrigerators must still provide service lives of 20 to 30 years. Generally, a fully assembled refrigerating system is subject to an integral pretest, then loaded with refrigerant, and finally only tested for refrigerant loss at the sealed filling port.

Testing for refrigerant leaks is done by local sniffing leak detection using special equipment that is sensitive to refrigerants (see Section 14.4.6). As opposed to standard local sniffing leak detection, such applications do not detect the position of a leak (only one spot is tested), but only test for leak rates exceeding the tolerable leak rate limit.

The desired threshold for the leak rate limit is given by a test or reference leak and stored in the instrument during calibration. Simple good/bad test results are sufficient, and are usually indicated both optically as well as acoustically.

Particular problems in this application arise from indication errors produced by foreign gases. Such foreign gases include, on the one hand, the actual coolant as background concentration in the environment, and on the other hand, other gases or vapors that also produce a signal in the leak detector.

A large variety of coolants are used today and many of these are also utilized as foaming agents in insulating boards (in refrigerators) and padding materials (in cars). Many other substances occur such as solvents, cleaning solutions, brake fluids, and so on. All these substances produce signals in simple alkali ion leak detectors, which could be interpreted as leaks. Therefore, testing areas should be carefully shielded and ventilated. Just recently, modern instrument designs using infrared detectors and reference-gas methods have provided solutions to these problems (see also Section 14.4.6).

19.6 **Industrial Tightness Testing of Mass-Production Components**

19.6.1

Basics

Industrial tightness testing includes all methods for semi or fully automated testing of high-volume series components. Nearly all methods employ integral chamber testing using helium tracer gas. Depending on later operating conditions, specimens are either filled with helium overpressure, or evacuated and exposed to outside helium. High-priced components are reworked and the leaks are located and repaired by means of sniffing or helium spraying.

19.6.2

Industrial Testing of Series Components

Industrial series components are parts produced in high volumes that meet high requirements in terms of tightness. The large variety of parts ranges from simple exhaust manifolds to all types of valves and airbag sensors. Demands regarding shortest possible testing times are increasing steadily, and at the same time, testing of the objects is becoming more and more difficult in terms of prerequisites such as cleanliness, temperatures, adaptability, sealing, and so on.

19.6.2.1 Envelope Testing Method for Vacuum Components (Method A1 in EN 1779)

When testing vacuum components, these parts are connected to the massspectrometric detection system and evacuated. Subsequently, the testing chamber is used as a sheath filled with tracer gas. The rest of the test procedure is the same as for overpressure components except for the fact that the specimens have vacuum connectors and the chamber has pressure connectors. A detailed description of the procedure is given in the following section on overpressure components.

19.6.2.2 Vacuum Chamber Method for Overpressure Components (Method B6 in EN 1779)

Parts that are designed to operate at overpressure are also tested with helium overpressure. Depending on tightness requirements and testing times, tracer-gas mixtures containing only a certain percentage of helium in nitrogen are often used for economic reasons. The fully automatic procedure comprises the following steps:

- 1) Specimen is placed on test desk, adjusted, and connected to tracer-gas
- 2) Hood is lowered onto specimen to form a chamber around the specimen.
- 3) Chamber and specimen are evacuated.
- 4) Specimen is loaded with tracer gas (possibly mixture).
- 5) Connection to mass spectrometer detector is established; helium flow = leak rate from specimen into vacuum chamber is measured after accommodation time.
- 6) Acoustic or optical good/bad signal, depending on predefined leak rate threshold.
- 7) Pump down of tracer gas from the specimen (helium recovery, if applicable) and venting of the chamber.
- 8) Hood is lifted and specimen is removed.

As an example of such a fully automated tightness testing system, Figure 19.14 shows a photograph of a wheel testing system for tightness testing of light-metal automotive wheels. Such systems have already been available for 20 to 30 years, relying more or less on the same operating principles. A considerable improvement was the introduction of the counterflow principle since it no longer required a cold trap with liquid nitrogen or a refrigerating system.

Furthermore, service lives, particularly of system valves, were prolonged considerably by an order of 10 (from approximately 300 000-3 000 000 cycles) and speeds of handling systems were enhanced significantly. All these advances led to modern cycle times of just a few seconds and still continuously improving detection limits.

Figure 19.15 shows the vacuum diagram of this system. As the image shows, helium recovery is an integral part of the system. The measuring principle of the mass spectrometer (MS) leak detector is not shown in detail.



Figure 19.14 Fully automated test system for aluminum wheels (Von der Heyde, Stade, Germany). Apart from actual measurement, the system also performs the operations "load/unload," "sort," "label." Main technical specifications: leak rate: 10^{-4} mbar ℓ s⁻¹ $(10^{-2} \text{ Pa } \ell \text{ s}^{-1})$ with automatic calibration

system; testing pressure: controllable up to 4 bar; test gas: helium/air mixture with recovery of test gas; production cycle time: 200 parts h⁻¹ (400 parts h⁻¹ in twin system); wheel sizes: 13-20" (diameter); 3-12" height, testing of mixed types.

Table 19.3 compares technical specifications of the 1982 and 2002 wheel testing apparatuses. The counterflow principle yielded considerable improvements and made the costly utilization of a cold trap and refrigerator dispensable.

Other improvements involve engineering advances in the design of mechanical and control components. All developments are aimed at providing systems for three-shift operation with lowest possible downtime to the customers.

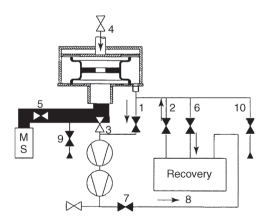


Figure 19.15 Vacuum schematic of wheel tester in Figure 19.14. 1 Pump-down valve, test chamber. 2 Test gas filling valve. 3 Pump-down valve, test gas volume. 4 Sealing system.

5 Measurement valve. 6 Test gas deflating valve. 7 Test gas return valve. 8 Test gas recovery system. 9 Vent valve, measurement chamber. 10 Vent valve, test gas volume.

Property	1982	2002
Method	Direct flow with cold trap/ refrigerator	Counterflow with turbopump
Service life of valves	Several 100 000 cycles	Several million cycles
Cycle time	Approximately 30 s	Approximately 18 s
Wheel sealing	Uncontrolled sealing pres- sure, standard elastomer profiles	Sealing pressure controlled according to wheel size, special elastomer profiles
Control system	Programmable logic controller (PLC) without graphics	Operator guidance with visualized procedures
Test gas recovery	Option	Integrated into system

Table 19.3 Comparison of important technical properties of industrial wheel testing systems between 1982 and 2002.

19.6.2.3 Testing of Hermetically Sealed Components by Means of Bombing (Method B5 in EN 1779)

Hermetically sealed parts such as semiconductor components or optoelectronic components are not testable with simple test chamber methods because they do not include connecting flanges for tracer gas or leak detectors. Therefore, testing of such components comprises the following two steps:

- 1) Exposure of the components to pressurized helium (approximately 8 bar) for several hours (bombing: helium penetrates into the inner hollow through possible leaks).
- 2) Specimens are placed inside a vacuum vessel and tested for escaping helium using a mass spectrometer (as in method B6, EN 1779).

For the parameters time t and pressure p during pressurized exposure of a test object with volume V, the rejection limit q_R , that is, the maximum tolerable leak rate indicated for a tolerable leak rate $q_{\rm L}$ of the test object is

$$q_{\rm R} = (q_{\rm L}p)/p_0 \left(1 - e^{-q_{\rm L} \frac{T}{p_0 V}}\right) e^{-q_{\rm L} \frac{T}{p_0 V}},\tag{19.25}$$

with atmospheric pressure p_0 . Time T is the waiting time between pressurized exposure and testing. This time is required for desorbing helium possibly adsorbed to the outside of the specimen (particularly for glass or plastic housings) since its signal could be misinterpreted as leaks [7].

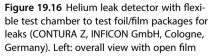
The bombing method yields similarly small signals for very large leaks as for fine leaks because the helium is released rapidly via large leaks after the pressurized exposure. Therefore, rough-leak pretesting is mandatory in order to reject heavily leaking parts. Often, such testing employs bubble emission methods (according to EN 1593), for which the specimens are submerged in liquids with low surface tension (water with surface-active agent, solvents, or oils) after the pressurized exposure. Hot liquids enhance bubble formation due to the temperature-dependent rise of inner pressure, thus yielding higher detection sensitivity. Details concerning the individual test methods are given in MIL standard 883 "Test Method Standard Microcircuits," method 1014.9.

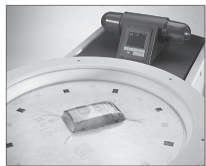
19.6.2.4 Testing of Food Packaging in Flexible Test Chambers

Food packaging made of flexible film or foil requires a special testing approach. Although such usually protective-gas filled packaging could be filled partially or completely with helium, it cannot be tested for tightness simply by placing it inside a vacuum chamber because the film or foil does not withstand any noteworthy pressure differences. For this application, a testing method was developed in which the testing chamber itself is made up of two flexible films that support the packaging film or foil safely during evacuation. A laid-in fleece allows free flow of escaping helium to the detection system. The procedure has by now been standardized for Germany in DIN 55533:2004-06 "Packaging testing – Integral test method for packaging made of films or foils for leakage using a flexible test chamber with tracer gas."

The remarkable sensitivity of the helium testing procedure allows detection of leaks down to 1×10^{-5} Pa ℓ s⁻¹ (1×10^{-7} mbar ℓ s⁻¹), thus providing (e.g., by sampling of packaging while the production is running and offline testing) a means of monitoring packaging machines (e.g., sealing processes). This enables corrective adjustments before possible rejects are produced. Considerable savings in terms of time and cost were achieved since the necessary waiting periods of several days of packages in quarantine have become unnecessary. Figure 19.16 shows a commercial instrument for testing flexible packages. The photographs show two circular







chamber; right: detailed view of evacuated film chamber containing a coffee package. The tightly fitting film supports the package thoroughly.

frames forming the food chamber into which the packaging is placed. An automated testing cycle starts as soon as the chamber is closed, and indicates the result as a good/bad signal. Recently, this method has been extended to a pure pressure rise method (int. patent application WO002013072173). Without any specific tracer gas in the packages this method allows leaks down to 10⁻³ mbarl/s to be found within a few seconds cycle time.

References

- 1 Nylander, C. and Lohm, U. (1995) Dichtheitsprüfung mit Wasserstoff. QZ, 40,
- 2 Moore, P.O. (ed.) (1998) Nondestructive Testing Handbook, vol. 1, 3rd edn, (Technical eds C.N. Jr. Jackson, C.N. Sherlock), Leak Testing, American Society for Nondestructive Testing, ISBN 1-57117-
- 3 Ehrlich, C.D. (1986) A note on flow rate and leak rate units. J. Vac. Sci. Technol. A,
- 4 Nerken, A. (1958) Versuche über die Strömung von Gasen durch Lecke. Vak. Techn., 7, 111.
- 5 Beckmann, W. and Seider, M.H. (1967) Gasdurchlässigkeit von gummielastischen

- Werkstoffen für Stickstoff. Kolloid Zeitschr., 220, 97.
- 6 Requirements on personnel for nondestructive testing, specified in European standard EN 473.
- 7 Howl, D.A. and Mann, C.A. (1965) The back-pressurising technique of leak testing. Vacuum, 15, 347.

Further reading

1 Calcatelli, A., Bergoglio, M., and Mari, D. (2007) Leak detection, calibrations and reference flows: practical example. Vacuum, 81, 1538-1544.

Appendix

Dr. Karl Jousten

Physikalisch-Technische Bundesanstalt, Vacuum Metrology, Abbestr. 2-12, 10587, Berlin, Germany

Table A.1 Pressure ranges in vacuum technology.

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Table A.20 Vacuum-technological applications and applicable pump systems.

Table A.21 Selected vacuum processes and techniques.

Table A.22 National and international standards and guidelines.

Table A.23 Common stainless steel types in vacuum technology.

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Table A.1 Pressure ranges in vacuum technology and their characteristics (numeric values apply to gas species with collision radii and molecular masses similar to those of air at 300 K, rounded to powers of 10).

Quantity	Symbol	Unit	Section	Equation	Low vacuum	Medium vacuum	High vacuum	Ultrahigh vacuum
Pressure	p	Pa	3.1.1	3.1	$100 - 10^5$	0.1 - 100	$10^{-5} - 0.1$	$< 10^{-5}$
Particle number density	п	cm^{-3}	3.1.1	3.10	$10^{16} - 10^{19}$	$10^{13} - 10^{16}$	$10^9 - 10^{13}$	$< 10^9$
Mean free path	Ī	m	3.2.5	3.55	$10^{-4} - 10^{-7}$	$00.1 - 10^{-4}$	$10^3 - 0.1$	$> 10^{3}$
Collision rate, molecular flow rate	j_N	$\mathrm{cm^{-2}\ s^{-1}}$	3.2.4	3.48	$10^{20} - 10^{23}$	$10^{17} - 10^{20}$	$10^{13} - 10^{17}$	$< 10^{13}$
Volume collision rate	χ	${\rm cm}^{-3}~{\rm s}^{-1}$	3.2.5	3.60	$10^{23} - 10^{29}$	$10^{17} - 10^{23}$	$10^9 - 10^{17}$	$< 10^9$
Monolayer time	$t_{ m mono}$	S	6.2.6	6.25	$10^{-8} - 10^{-5}$	$10^{-2} - 10^{-5}$	$100 - 10^{-2}$	>100
Flow regimes					Viscous flow	Knudsen flow (transition flow)	Molecular flow	Molecular flow

Table A.2 Conversion table for pressure units (units prohibited in SI: at, atm, Torr, psi) ("at" was technical atmosphere, "atm" standard atmosphere; example: 1 Torr = 1.333 22 mbar, 1 psi = 68.95 mbar; multiply with the corresponding number in the cell).

1 Pa	1 bar	1 mbar	1 at	1 atm	1 Torr	1 psi
1	10^{-5}	10-2	1.0197×10^{-5}	9.8692×10^{-6}	7.5006×10^{-3}	1.4504×10^{-4}
10^{5}	1	1000	1.0197	0.986 92	750.06	14.5032
10^{2}	10^{-3}	1	1.0197×10^{-3}	0.98692×10^{-3}	0.750 06	14.5032×10^{-3}
98 066.5	≈0.981	980.68	1	0.967 84	735.56	14.2247
101 325	1.013	1013.25	1.03323	1	760	14.6972
133.322	≈ 0.00133	1.33322	0.00136	1.3158×10^{-3}	1	0.01934
6894.8	0.06895	68.95	0.0703	0.06804	51.715	1
	1 10 ⁵ 10 ² 98 066.5 101 325 133.322	$\begin{array}{cccc} 1 & & 10^{-5} \\ 10^{5} & & 1 \\ 10^{2} & & 10^{-3} \\ 98\ 066.5 & \approx 0.981 \\ 101\ 325 & & 1.013 \\ 133.322 & \approx 0.00133 \end{array}$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

a) $1 \text{ at} = 1 \text{ kp cm}^{-2}$ (old definition).

b) $1 \text{ psi} = 1 \text{ pound inch}^{-2}$.

Table A.3 Conversion table for pumping speeds (cfm: cubic feet per minute); example: $1 \text{ m}^3 \text{ h}^{-1} = 0.278$ $\ell \, \text{s}^{-1}$; 1 $\ell \, \text{s}^{-1} = 3.6 \, \text{m}^3 \, \text{h}^{-1}$; 1 cfm = 0.471 $\ell \, \text{s}^{-1}$.

↓corresponds to→	m ³ h ⁻¹	ℓ s ⁻¹	m ³ s ⁻¹	${\mathscr C} {\mathsf {min}}^{-1}$	cfm
$m^3 h^{-1}$	1	0.278	2.78×10^{-4}	16.68	0.589
ℓs^{-1}	3.6	1	1000	60	2.12
m^3s^{-1}	3600	1000	1	60 000	2120
$\ell \mathrm{min}^{-1}$	0.06	0.0167	1.67×10^{-5}	1	0.0353
cfm	1.7	0.471	4.71×10^{-4}	28.3	1

Table A.4 Important equations for physics of ideal gases.

Quantity	Eq.	Quantity equation	Numeric-value equation	NVE for air ($M_{\rm r}=28.96$) and $\theta=20^{\circ}{\rm C}$
Most probable particle velocity c_{prob}	3.42	$c_{ m prob} = \sqrt{\frac{2RT}{M_{ m molar}}}$	$c_{\text{prob}} = 129 \sqrt{\frac{T}{M_{\text{r}}}} \text{m s}^{-1}$	$c_{\rm prob}=410~{\rm m~s^{-1}}$
Mean particle velocity \bar{c}	3.43	$\overline{c} = \sqrt{\frac{8RT}{\pi M_{ m molar}}}$	$\overline{c} = 146\sqrt{\frac{T}{M_{\rm r}}}\mathrm{m}\;\mathrm{s}^{-1}$	$\overline{c} = 463 \mathrm{m}\mathrm{s}^{-1}$
Mean square velocity \overline{c}^2	3.44	$\overline{c}^2 = \frac{3RT}{M_{\text{molar}}}$	$\overline{c}^2 = 24000 \frac{T}{M_r} \text{ m}^2 \text{ s}^{-2}$	$\bar{c}^2 = 25.2 \times 10^4 \text{ m}^2 \text{ s}^{-2}$
Effective velocity $c_{\rm eff}$	3.44	$c_{ m eff} = \sqrt{\overline{c}^2}$ $= \sqrt{\frac{3RT}{M_{ m molar}}}$	$c_{\rm eff} = 158 \sqrt{\frac{T}{M_t}} \mathrm{m} \cdot \mathrm{s}^{-1}$	$c_{\rm eff}=502~\rm ms^{-1}$
Equation of state for ideal gases	3.20	$pV = \nu RT$	$pV = 83.14 \nu T \text{ mbar } \ell$	$pV = 2.44 \times 10^4 \nu$ mbar $\ell^{\rm a)}$
	3.19	p = nkT	$p = 1.38 \times 10^{-19} \ n \ T \ \text{mbar}$	$p = 4.04 \times 10^{-17} n \text{mbar}^{a)}$
Gas pressure p	3.34	$p = \frac{1}{3}nm_{\rm a}\overline{c^2} = \frac{1}{3}\rho\overline{c^2}$	_	_
Particle number density <i>n</i>	3.19	n = p/(kT)	$n = 7.25 \times 10^{18} \frac{p}{T} \text{ cm}^{-3}$	$n = 2.5 \times 10^{16} p \mathrm{cm}^{-3} ^{\mathrm{a}}$
Collision rate j_N	3.48	$j_N = \frac{n\overline{c}}{4}$	$j_N = 2.63 \times 10^{22} \frac{p}{\sqrt{M_r T}} \text{ cm}^{-2} \text{ s}^{-1}$	$j_N = 2.85 \times 10^{20} p \text{ cm}^{-2} \text{ s}^{-1}$
		$=\sqrt{rac{N_{ m A}^2}{2\pi RTM_{ m molar}}}p$		
Surface area-related	_	$q_{m,A} = j_m = j_N m_a$	$q_{m,A}\equiv j_m=4.38\times 10^{-2}$	$q_{m,A} \equiv j_m = 1.38 \times$
		$=\sqrt{\frac{M_{\rm molar}}{2\pi RT}}$	$\sqrt{\frac{M_{\rm r}}{T}}ptrmg~{ m cm}^{-2}~{ m s}^{-1}$	$10^{-2} p \mathrm{g \ cm^{-2} \ s^{-1}}$
Volume collision rate χ	3.60	$\chi = \frac{1}{2} \cdot \frac{n\overline{c}}{\overline{l}}$ $= \frac{1}{\overline{l}p} \sqrt{\frac{8N_A^2}{\pi M_{\text{modar}} RT}} p^2$	$\chi = 5.27 \times 10^{22} \frac{1}{lp} \cdot \frac{p^2}{\sqrt{M_t T}} \text{ cm}^{-3} \text{ s}^{-1}$	$\chi = 8.6 \times 10^{22} p^2 \text{ cm}^{-3} \text{ s}^{-1}$

R: universal (molar) gas constant; T: thermodynamic temperature; $M_{
m molar}$: molar mass; $M_{
m r}$: relative atomic or molecular mass; p: pressure; V: volume; ν : amount of substance, number of moles; n particle number density; ρ : density; k: Boltzmann's constant; m_a : particle mass; N_A : Avogadro's constant; \bar{l} : mean free path. Units for numeric-value equations (NVE): T in K, M_r = dimensionless number, p in hPa, V in ℓ , ν in mol, n in cm⁻³, $\bar{l}p$ in cm hPa (see also Table A.9, there in m hPa).

a) For any ideal gas.

Table A.5 Internationally recommended values of physical constants (CODATA available at http://physics.nist.gov/cuu/Constants/index.html). Unless standard uncertainties are given in parentheses, values are exact, that is, as agreed.

Symbol	Quantity	Value
c	Speed of light	$2.997992459 \times 10^8 \text{ m s}^{-1}$
e	Elementary charge	$1.60217653(14) \times 10^{-19} \text{ C}$
$g_{\rm n}$	Standard acceleration of gravity	9.80665m s^{-2}
h	Planck constant	$6.6260693(11) \times 10^{-34} \mathrm{J s}$
$k^{a)}$	Boltzmann constant	$1.3806505(23)\times10^{-23}\mathrm{JK^{-1}}$
$m_{\rm e}$	Electron mass	$9.1093826(16) \times 10^{-31} \text{ kg}$
$m_{\rm p}$	Proton mass	$1.67262171(29) \times 10^{-27} \text{ kg}$
$m_{\rm n}$	Neutron mass	$1.67492728(29) \times 10^{-27} \text{ kg}$
$m_{ m p}/m_{ m e}$	Ratio of proton and electron masses	1.83615267261(85)
m_u	Atomic mass unit (1 u)	$1.66053886(28) \times 10^{-27} \text{ kg}$
N_{A}	Avogadro's constant	$6.0221415(10) \times 10^{23} \text{ mol}^{-1}$
$p_{\rm n}^{\rm b,c)}$	Standard pressure	$101325~{ m N}~{ m m}^{-2}$
R	Universal (molar) gas constant	$8.314472(15)\mathrm{J}\mathrm{mol}^{-1}\mathrm{K}^{-1}$
		$= 83.145 \text{ mbar } \ell \text{ mol}^{-1} \text{ K}^{-1}$
		$= 8.3145 \times 10^4 \mathrm{mbar} \ell \mathrm{kmol}^{-1} \mathrm{K}^{-1}$
		$= 8.3145 \times 10^{3} \text{ Pa m}^{3} \text{ kmol}^{-1} \text{ K}^{-1}$
$T_n^{(b)}$	Standard temperature	273.15 K
$V_{ m molar,n}$	Standard molar volume of ideal gas	$22.413996(39) \ell \text{ mol}^{-1}$
σ	Stefan-Boltzmann constant	$5.670400(40) \times 10^{-8} \text{ W m}^{-2} \text{ K}^{-4}$
$e/m_{\rm e}$	Electron charge to mass quotient	$-1.75882012(15) \times 10^{11} \mathrm{Ckg^{-1}}$

a) $k = R/N_A$.

b) DIN 1343 asks for using subscript "std" (standard) instead of "n" for standard conditions.

c) $p_{\rm n}$ is also termed standard atmosphere (symbol atm), according to IUPAP 25, SUNAMCO 87-1.

Table A.6 Volume percentages of air, partial pressures with respect to atmospheric pressure p = 1000 hPa, and percentages by mass of constituents. Values given for N₂, O₂, Ar, and CO₂ are taken from a recent publicationfn1, replacing the older recommended values^{b)}. The volume percentage of CO₂ varies according to regions and time; however, O₂ and CO₂ are locally anti-correlated; their added volume percentages yield a constant 20.982%. The density of dry air at 1013.2 hPa and 0 °C (standard conditions) is 1.2924 kg m⁻³. The mean molar mass of dry air is $28.968 \text{ g mol}^{-1}$.

Constituent		Volume percentage ^{c)} %	Partial pressure in hPa at $p_{\text{tot}} = 1000 \text{ hPa}$	Percentage by mass %
Nitrogen	N_2	78.083	780.83	75.5167
Oxygen	O_2	20.945	209.45	23.1385
Argon	Ar	0.9331	9.33	1.2869
Carbon dioxide	CO_2	0.0369	0.37	0.0561
Neon	Ne	1.8×10^{-3}	1.8×10^{-2}	1.3×10^{-3}
Helium	He	5.2×10^{-4}	5.2×10^{-3}	7.2×10^{-5}
Methane	CH_4	1.5×10^{-4}	1.5×10^{-3}	8.3×10^{-5}
Krypton	Kr	1.1×10^{-4}	1.1×10^{-3}	3.2×10^{-4}
Hydrogen	H_2	5×10^{-5}	5×10^{-4}	3.5×10^{-6}
Dinitrogen	N_2O	3×10^{-5}	3×10^{-4}	4.6×10^{-5}
monoxide				
Xenon	Xe	1×10^{-5}	1×10^{-4}	4.5×10^{-5}
Ammonia	NH_3	2.6×10^{-6}	2.6×10^{-5}	2.8×10^{-6}
Ozone	O_3	2×10^{-6}	2×10^{-5}	3.3×10^{-6}
Hydrogen peroxide	H_2O_2	4×10^{-8}	4×10^{-7}	4.7×10^{-8}
Iodine	I_2	3.5×10^{-9}	3.5×10^{-8}	1.5×10^{-8}
Radon	Rn	7×10^{-18}	7×10^{-17}	5.4×10^{-17}

Apart from these constituents, ambient air contains varying amounts of water vapor and carbon monoxide. The values below relate to the saturation state for water vapor at 293 K. The values for carbon monoxide apply to a typical large city.

Water vapor	H_2O	≤ 2.3	≤ 23.3
Carbon	CO	$\leq 2\times 10^{-5}$	$\leq 2\times 10^{-4}$
monoxide			

a) S. Y. Parket al., A redetermination of the argon content of air for buoyancy corrections in mass standard comparisons, Metrologia (2004) 387-395.

b) R. S. Davis, Equation for the determination of the density of moist air (1981/91), Metrologia (1992) 67-70.

c) Volume percentage= amount of substance.

Table A.7 Selected properties of gases.

No.	No. Gas, vapor, or mixture	Chemical notation	$A_{ m r}^{ m a)}$ or $M_{ m r}^{ m a)}$	m _a ^{a)} (10 ⁻²⁷ kg)	$ ho_{ m n}$ (kg m $^{-3}$)	c _p	c _V	$10^{3}\lambda$ (Wm ⁻¹ K ⁻¹)	$10^6 \eta^{9)}$ (kg m $^{-1}$ s $^{-1}$)	Sta	ndard boilin	g point	T _{melt} (K)
				-	-			(kJ kg ⁻¹ K, ⁻¹)		T _{B.P.} (K)	$\rho_{ m liq.}$ (kg m ⁻³)	$\Lambda_{v, \text{ spec.}}$ (kJ kg ⁻¹)	
							at θ :	= 20 °C, p = 100	kPa				
1	Hydrogen	H ₂	2.016	3.348	0.0899	14.32	10.14	182.6	8.8	20.38	71	454	13.95
2	Helium	He	4.003	6.647	0.1785	5.23	3.21	148	19.6	4.22	130	20.6	_
3	Methane	CH_4	16.043	26.64	0.7168	2.22	1.70	33.1	10.8	111.71	425	510	90.63
4	Ammonia	NH_3	17.031	28.28	0.7714	2.16	1.66	22	9.8	239.75	682	1370	195.45
5	Water vapor	H_2O	18.015	29.97	0.8042	1.94 ^{b)}	_	_	≈9	373.15	958.35	2255.5	273.15
6	Carbon monoxide	CO	28.011	46.51	1.250	1.04	0.74	24.5	17.6	81.68	792	216	68.08
7	Nitrogen	N_2	28.013	46.52	1.2505	1.04	0.74	25.5	17.5	77.35	808	198	63.15
8	Air	0.78	28.96	48.09	1.2929	1.01	0.72	25.6	18.19	81.75	-	_	_
		$N_2 + 0.21$											
		$O_2 + 0.01 Ar$											
9	Oxygen	O_2	31.999	53.14	1.4290	0.92	0.66	26.1	20.2	90.18	1134	213	54.36
10	Hydrogen chloride	HCl	36.461	60.55	1.6392	0.80	0.56	13.6	14.2	188.15	1194	443	158.95
11	Argon	Ar	39.948	66.34	1.784	0.52	0.32	17.3	22.11	87.29	1390	163	83.77
12	Carbon dioxide	CO_2	44.010	73.08	1.977	0.84	0.65	15.8	14.6	194.65 ^{c)}	1560 ^{e)}	136.8 ^{f)}	216.58 ^{d)}
13	Chlorine	Cl_2	70.906	117.7	3.214	0.75	0.35	8.6	13.2	239.05	1564	290	172.15
14	Difluorodichloro- meth- ane (Freon 12)	CCl ₂ F ₂	120.914	200.8	5.510				13.2	248.25	1484	162	114.95

 $[\]rho_{\rm n}$: standard density (at $T_{\rm n}=273.15\,{\rm K}$ and $p_{\rm n}=1013.25\,{\rm mbar}$); c_p and c_V : specific heat capacities at constant pressure and constant volume, respectively; λ : thermal conductivity; η : dynamic viscosity; $\Lambda_{v,spec}$: specific heat of evaporation (evaporation enthalpy).

a) Mean relative mass (column 4) and mean mass (column 5) of atoms or molecules in natural isotope composition. Molar mass $M_{\text{molar}} = A_{\text{r}}$ in kg kmol⁻¹ and M_{r} in kg kmol⁻¹. b) At $\theta = 20$ °C and p = 100 kPa.

of For $T_{B.P.} = 194.65$ K, the vapor pressure of solid carbon dioxide is equal to standard pressure $p_n = 101.325$ kPa. The liquid phase occurs only above the triple point $T_t = 216.58$ K, $p_t = 0.5$ MPa.

e) Density of solid CO₂ at $T_{B.P.}$ and p_n .

f) Sublimation heat. g) $1 \text{ kg m}^{-1} \text{ s}^{-1} = 10^{-2} \text{ mbars.}$

Table A.8 Second virial coefficient B" according to Eq. (3.113) for selected gas species at 23 °C, its uncertainty u(B''), and temperature coefficient $\Delta B''/\Delta T$ in the room-temperature range (18 °C to 30 °C). Values are based on measured and assessed data (from Dymond and Smith "The virial coefficients of pure gases and mixtures," Clarendon Press, 2nd Ed., Oxford, 1980).

Gas species		B"(23 °C) (cm³ mol ⁻¹)	$u(B^{\prime\prime})$ (cm ³ mol ⁻¹)	$\Delta B'' / \Delta T$ (18 °C to 30°C) (cm ³ (mol K) ⁻¹)
Hydrogen	H_2	14.7	0.5	0.035
Deuterium	D_2	13.4	0.5	0.02
Helium	He	11.7	0.5	-0.0045
Methane	CH_4	-43.7	1.0	0.44
Water vapor	H_2O	-1200	150	18
Neon	Ne	11.2	1.0	0.037
Ethylene	C_2H_2	-176.1	25	2.1
Nitrogen monoxide	NO	-22.2	2.0	0.22
Nitrogen	N_2	-5.1	0.5	0.25
Carbon monoxide	СО	-8.8	0.5	0.18
Oxygen	O_2	-16.9	1.0	0.24
Argon	Ar	-16.5	0.5	0.25
Carbon dioxide	CO_2	-126.5	2.0	0.98
Krypton	Kr	-52.7	1.0	0.5
Xenon	Xe	-136.5	3.0	0.92

Table A.9 Collision radii R=2r, Sutherland's constant $T_{\rm D}$, see Eq. (3.58), and mean free path \bar{I} of selected important gas species ($\bar{I}p$ values). R calculated according to Eq. (3.76) from values of dynamic viscosity η measured at T=273.15 K. Sutherland's correction using constant $T_{\rm D}$ is questionable, compare Ar and N₂ as well as H₂ and He. The temperature intervals for which $T_{\rm D}$ values are calculated from measured data are given in brackets. R_{∞} extrapolated from $R_{\rm T}$ values which are calculated from $\eta_{\rm T}$ values. Due to their inherent uncertainty, $\bar{I}p$ values are given with one decimal place only; within their accuracy, they are also valid for T=293.15 K ($\theta=20^{\circ}{\rm C}$).

Gas Species	$R_{\infty} \times 10^{10}$ (m)	$R_T \times 10^{10}$ (m) ($T = 273.15$ K from η_{273})	<i>T</i> _D (K)	<i>l̄p</i> (m Torr)	<i>lp</i> (m mbar)
				for <i>T</i> = 273.	15 K ^{a)}
H ₂	2.2 - 2.4	2.72	75 – 235 (90 – 1 000)	8.6×10^{-5}	11.5×10^{-5}
N_2	3.2	3.78	98 - 107 $(90 - 1000)$	4.4×10^{-5}	5.9×10^{-5}
O_2		3.62		4.9×10^{-5}	6.5×10^{-5}
Не	1.82 – 1.94	2.18	22 - 175 $(20 - 1000)$	13.1×10^{-5}	17.5×10^{-5}
Ne		2.56		9.5×10^{-5}	12.7×10^{-5}
Ar	2.86 - 2.99	3.66	132 - 144 (90 – 1 000)	4.8×10^{-5}	6.4×10^{-5}
Air ^{b)}			≈102	5×10^{-5}	6.65×10^{-5}
Kr		4.14		3.7×10^{-5}	4.9×10^{-5}
Xe		4.88		2.7×10^{-5}	3.6×10^{-5}
Hg				2.3×10^{-5}	3.1×10^{-5}
H_2O		4.14		5.1×10^{-5}	6.8×10^{-5}
CO		3.77		4.5×10^{-5}	6.0×10^{-5}
CO_2		4.62		3.0×10^{-5}	4.0×10^{-5}
HCl		4.51		3.3×10^{-5}	4.4×10^{-5}
NH_3		4.47		3.2×10^{-5}	4.3×10^{-5}
Cl ₂		5.52		2.1×10^{-5}	2.8×10^{-5}

a) $(\bar{l}p)_T/(\bar{l}p)_{T=273 \text{ K}} = \left(\frac{T}{273 \text{ K}}\right)^2 \frac{273 \text{ K} + T_D}{T + T_D} =: \Theta.$

b) Measured at 20 °C.

Table A.10 Most probable (c_{prob}), Eq. (3.42), mean (\overline{c}), Eq. (3.43), and effective (c_{eff}) velocities, Eq. (3.44), of important gas molecules and atoms, and for a hypothetical air particle according to the mean molecular mass of dry air.

Gas Species	Chemical Molar mass c_{prob} Notation (q mol ⁻¹)			<u>c</u>			
		(3			(m s ⁻¹)		
			23 °C	0°C	23 °C	100 °C	23 °C
Hydrogen	H_2	2.016	1563	1694	1764	1980	1914
Helium	He	4.003	1109	1202	1252	1405	1358
Methane	CH_4	16.043	554	600	625	702	679
Water vapor	H_2O	18.015	523	567	590	662	640
Neon	Ne	20.18	494	535	557	626	605
Nitrogen	N_2	28.0134	419	454	473	531	514
Air (dry)	$0.78 N_2 + 0.21$	28.965	412	447	465	522	505
	$O_2 + 0.01 Ar$						
Oxygen	O_2	31.9988	392	425	443	497	480
Argon	Ar	39.948	351	380	396	445	430
Carbon dioxide	CO_2	44.01	335	363	377	424	410
Krypton	Kr	83.8	242	263	274	307	297
Xenon	Xe	131.29	194	210	219	245	237

Table A.11 Condensation probabilities (condensation coefficients) $\sigma_{\rm c}$ of selected substances under various conditions. (Measured bibliographical data vary considerably.)

Substance		Temperature range (°C)	Saturation vapor pressure p_s (mbar)	Saturation ratio during measurement $\beta = p_v/p_s$	Condensation coefficient σ_{c}
Mercury Hg	Solid	−64 to −41	$5 \times 10^{-8} - 3 \times 10^{-6}$	0	0.8 - 1.0
Water H ₂ O	Solid	-13 to -2	2.0 - 5.2	0.5 - 0.9	0.011 - 0.022
	Liquid	-0.8 to $+4.1$	5.6 - 8.1	0.5 - 0.9	0.032 - 0.055
		40 to −100	74 - 1013	0.9 - 1	0.02 - 0.03
Acetyl alcohol C_2H_5OH	Liquid	-2 to +16	13 – 48	0.5 - 0.8	0.024
Benzole C_6H_6	Liquid	6	50	0.99	0.9

Table A.12 Specific volumes $V_{\text{spec.}}$, specific heat capacities c_p , and specific evaporation enthalpies $\Lambda'_{\text{spec.}}$ of saturated water vapor above water and ice. See Table A.13 for saturation vapor pressures and densities.

ϑ (°C)	<i>T</i> (K)	$V_{\text{spec.}} (\text{m}^2 \text{ kg}^{-1})$ (above ice)	V _{spec.} (m ² kg ⁻¹) (above water)	$\Lambda_{\rm spec.}^{\prime}(\rm kJkg^{-1})$	$c_p (\text{kJ kg}^{-1} \text{K}^{-1})$
-30	243.15	2860	1750		
-20	253.15	1111	926		
-10	263.15	465.1	424		
0	273.15	206.300	206.300	2500.5	1.858
20	293.15		57.840	2453.4	1.862
40	313.15		19.560	2406.2	1.871
60	333.15		7.682	2357.9	1.881
80	353.15		3.410	2307.8	1.901
100	373.15		1.673	2255.5	1.94

 $\Lambda_{\rm spec.}'(T) = \Lambda_{\rm spec.}(T) + p_{\rm s}(T) V_{\rm spec.}(T), \text{ and analogously, } \Lambda_{\rm molar}'(T) = \Lambda_{\rm molar}(T) + p_{\rm s}(T) V_{\rm molar}(T), \text{ with the heat of evaporation } \Lambda.$

Table A.13 Pressure p_s and density ρ_s of the saturated vapor above pure liquid water (undercooled liquid for $\theta < 0$ °C, above ice in brackets) in the temperature range $\theta = -30$ °C to 140 °C. p_s is calculated after D. Sonntag, Z. Meteorol. 70 (1990), 340–344.

ϑ (°C)	p _s (hPa)	$ ho_{ m s}~({ m g~m^{-3}})$
-100	1.40E-05	_
-80	5.47E-04	
-60	1.08E-02	
-40	0.1284	
-30	0.5103 (0.3800)	0.57 (0.35)
-20	1.2559 (1.0324)	1.08 (0.9)
-14	2.0776 (1.8120)	1.739 (1.5)
-12	2.4432 (2.1731)	2.029 (1.8)
-10	2.8652 (2.5989)	2.359 (2.15)
-8	3.3511 (3.0998)	2.739 (2.55)
-6	3.9092 (3.6873)	3.170 (3.01)
-4	4.5485 (4.3748)	3.660 (3.54)
-2	5.2791 (5.1772)	4.214 (4.15)
0	6.1121 (6.1115)	4.847
1	6.5708	5.192
2	7.0597	5.559
3	7.5806	5.947
4	8.1352	6.360
5	8.7254	6.797
6	9.3531	7.260
7	10.020	7.750

Table A.13 (Continued)

θ (°C)	p _s (hPa)	$ ho_{ m s}~({ m g~m^{-3}})$
8	10.729	8.270
9	11.482	8.819
10	12.281	9.399
11	13.129	10.01
12	14.028	10.06
13	14.980	11.35
14	15.989	12.07
15	17.057	12.83
16	18.187	13.63
17	19.383	14.48
18	20.647	15.37
19	21.982	16.31
20	23.392	17.30
21	24.882	18.34
22	26.453	19.43
23	28.110	20.58
24	29.858	21.78
25	31.699	23.05
26	33.639	24.38
27	35.681	25.78
28	37.831	27.24
29	40.092	28.78
30	42.470	30.38
31	44.970	32.07
32	47.597	33.83
33	50.356	35.68
34	53.252	37.61
35	56.292	39.63
36	59.481	41.75
37	62.825	43.96
38	66.331	46.26
39	70.005	48.67
40	73.853	51.19
41	77.882	53.82
42	82.100	56.56
43	86.513	59.41
44	91.129	62.39
45	95.956	65.50
46	101.00	68.73
47	106.27	72.10
48	111.78	75.61
49	111.78	79.26
17	117.33	(continued

Table A.13 (Continued)

θ (°C)	p _s (hPa)	$ ho_{ m s}~({ m g~m^{-3}})$
50	123.53	83.06
51	129.79	87.01
52	136.32	91.12
53	143.13	95.39
54	150.23	99.83
55	157.63	104.4
56	165.34	109.2
57	173.37	114.2
58	181.73	119.4
59	190.43	124.7
60	199.48	130.2
61	208.89	135.9
62	218.68	141.9
63	228.86	148.1
64	239.44	154.5
65	250.43	161.2
66	261.84	168.1
67	273.70	175.2
68	286.00	182.6
69	298.77	190.2
70	312.02	198.1
71	325.77	206.3
72	340.02	214.7
73	354.79	223.5
74	370.10	232.5
75	385.97	241.8
76	402.40	251.5
77	419.42	261.4
78	437.04	271.7
79	455.28	282.3
80	474.16	293.3
81	493.68	304.6
82	513.88	316.3
83	534.77	328.3
84	556.36	340.7
85	578.68	353.5
86	601.74	366.6
87	625.57	380.2
88	650.17	394.2
89	675.59	408.6
90	701.82	423.5
91	728.90	438.8

Table A.13 (Continued)

<u>θ (°C)</u>	p _s (hPa)	$ ho_{ m s}~({ m g~m^{-3}})$
92	756.85	454.5
93	785.68	470.7
94	815.42	487.4
95	846.09	504.5
96	877.71	522.1
97	910.31	540.3
98	943.91	558.9
99	978.53	578.1
100	1014.19	597.8
101	1050.93	618.0
102	1088.75	638.8
103	1127.70	660.2
104	1167.80	682.2
105	1209.06	704.7
106	1251.52	727.8
107	1295.20	751.6
108	1340.14	776.0
109	1386.35	801.0
110	1433.86	826.7
111	1482.71	853.0
112	1532.91	880.0
113	1584.51	907.7
114	1637.52	936.1
115	1691.98	965.2
116	1747.92	995.0
117	1805.36	1026
118	1864.34	1057
119	1924.89	1089
120	1987.04	1122
121	2050.82	1156
122	2116.27	1190
123	2183.41	1225
124	2252.28	1262
125	2322.91	1299
126	2395.34	1337
127	2469.59	1375
128	2545.71	1415
129	2623.73	1456
130	2703.69	1497
131	2785.61	1540
132	2869.54	1583
133	2955.51	1627
100	2/33.31	
		(continued

Table A.13 (Continued)

θ (°C)	p _s (hPa)	$ ho_{ m s}~({ m g~m^{-3}})$
134	3043.56	1673
135	3133.72	1719
136	3226.04	1767
137	3320.55	1815
138	3417.29	1865
139	3516.30	1915
140	3617.61	1967

Table A.14 Surface area-related gas emissions (outgassing flux) j_1 after a pumping time of t=1 h, according to R. J. Elsey. *Outgassing of vacuum materials*. Vacuum 25 (1975), 299–306 and 347–361.

(a) Metals, j_1 (10 ⁻⁹ mbar ℓ s ⁻¹ cm ⁻²)				
Aluminum ^{a)}	6.3			
Aluminum, differently treated	4.1 - 6.6			
Duraluminum	170			
Gold wire ^{a)}	15.8			
Copper ^{a)}	40			
Copper ^{b)}	3.5			
Copper OFHCCu	18.8			
Copper OFHCCu ^{b)}	1.9			
Brass	400			
Molybdenum	5.2			
Titanium	4 - 11.3			
Zinc	220			
Selected steel grades	$j_1 \ (10^{-9} \ \mathrm{mbar} \ \ell \ \mathrm{s}^{-1} \ \mathrm{cm}^{-2})$			
Ingot iron	540			
Ingot iron, slightly corroded	600			
Steel, de-scaled	307			
Steel, chromium plated ^{a)}	7.1			
Steel, chromium plated ^{b)}	9.1			
Steel, nickel plated ^{a)}	4.2			
Steel, nickel plated	2.8			
Steel, nickel plated ^{a)}	8.3			
Stainless steel ^{a)}	13.5			
Stainless steel, Sandblasted	8.3			
Stainless steel ^{b)}	1.7			
Stainless steel, Electropolished	4.3			

Table A.14 (Continued)

(b) Other materials, j_1 (10 ⁻⁸ mbar \mathscr{E} s ⁻¹ cm ⁻²)						
Sealing materials	Data from Figure 20.11					
	Elsey	Curves	j_1 from curves			
Kel-F	4	_	_			
Neoprene	3000 - 300	5, 13	480, 210			
Perbunan	350	8, 9, 10	440, 300, 270			
Silicone	1800	20	430			
Silicone rubber	_	6. 14	650, 330			
Vespel	90	_	_			
Viton ^{a)}	114	23, 27, 28	620, 380, 350			
Viton, degassed	0.4					
Other materials						
Araldite, cast	120	_	_			
Araldite, various	150 - 800	12, 25, 19	150, 120, 40			
Plexiglas	70 - 300	15	110			
Polyethylene	23	34	12			
Polystyrene	56	29, 30, 31	30, 27, 20			
PTFE	30	_	_			
Pyrex glass ^{a)}	0.74	d	0.62			
Pyrex glass, exposed to air	0.12	_	_			
for 1 month						
Pyrophyllite	20	a	21			
Steatite	9	b	8.8			

a) As produced.

b) Mechanically polished.

Table A.15 Overview of vacuum pumps (DIN 28400-2. ISO 3529-2) and their names.

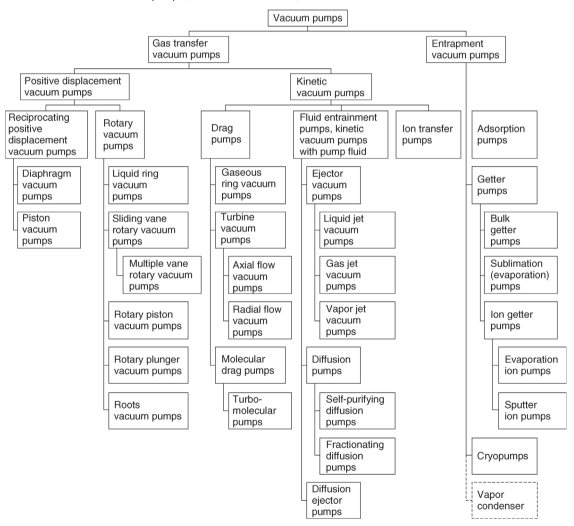
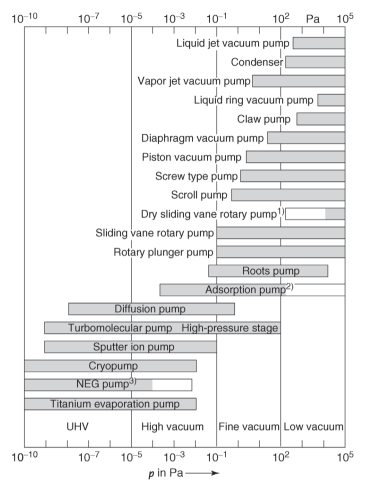


Table A.16 Overview of suction pressure ranges during continuous operation for typical pump types. Common series connections of several pump stages are taken into account; however, combinations of different pump types are not included.



- a) (Common) single-stage dry sliding vane rotary pumps produce approximately 10 kPa, (rare) multistage pumps can provide down to several 100 Pa.
- b) Adsorption pumps are not suitable for pumping continuously at pressures $> 100 \, \mathrm{Pa}$.
- c) NEG pumps are not suitable for pumping continuously at pressures $> 10^{-4}$ Pa.

Table A.17 Recommended oil types for selected applications of oil-filled positive displacement pumps.^{a)}.

Oil type	Universal oil	Special oil Protelen	Special oil NC 2	Special oil NC 1 Fomblin (PFPE)
Oil classification	Unalloyed machine oil (viscosity grade ISO VG 100)	Alkaline precharged machine oil	White oil. that is, unalloyed paraffin oil without olefins. aromatic compounds. and hetero compounds	Synthetic oil. perfluorinated polyether ^{b)}
Temperature range	up to 120 °C	up to 120 °C	up to 120 °C	up to 140 °C
Kinematic viscosity at 40°C	$83 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$	$131 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$	$70 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$	$87 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$
Density at 20 °C	$0.87~{\rm g~cm^{-3}}$	$0.90 \mathrm{g cm^{-3}}$	$0.887 \mathrm{g cm^{-3}}$	$1.90\mathrm{gcm^{-3}}$
Flashing point	260 °C	230 °C	225 °C	Non-flammable
Applications (examples)	Laboratory pumps operated with cold traps	Pumping of acidic gases or acid vapors	 Pumping of gases and vapors aggressive towards olefinic double bonds or aromatic compounds, for example, halogens, halogen hydra- cids, Lewis acids (BCl₃, AlCl₃, TiCl₄, etc.), halogenated hydrocarbons 	 Approved by BAM^{c)} for pumping pure oxygen
	 Pumping of air and chemically inert per- manent gases (e.g., noble gases) 	 Pumping of sub- stances producing acids during hydrol- ysis (e.g., organic acid chlorides) 		 Highly suitable for pumping strong oxidants (e.g., fluorine, other halogens, nitrogen oxides, etc.)
	 Pumping of water vapor Pumping of gases and vapors, weakly reactive towards olefins, and aromatic compounds 			

Comments

Alkyl sulphonic acid

• Extruder degassing

· Pumping of styrene,

other easily polym-

erizing substances

butadiene, and

ester (softener)

 $38 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$

up to 120°C

 $1.03 \, \mathrm{g \, cm^{-3}}$

224 °C

- · Oil according to DIN 51 506^{d)}
- Do not leave pumps out of operation for longer periods (standstill corrosion)
- · Chemical filters prolong service life
- Corrosion protection is consumed (well timed oil change)
- · Do not use chemical oil filters because they make additives ineffective

Diester-based synthetic up to 140 °C $95\times 10^{-6}\ m^2\ s^{-1}$ $0.95 \, \mathrm{g \ cm^{-3}}$

 Approved by BAM^{c)} for pumping pure oxygen

polychlortrifluoroethylene

Synthetic oil.

up to 120°C

 $1.9 \, \mathrm{g} \, \mathrm{cm}^{-3}$

 $52 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$

Non-flammable

• Highly suitable for pumping strong oxidants (e.g., fluorine, other halogens, nitrogen oxides, etc.)

- 266 °C • Operation at high ambient temperatures
- Pumping of solvent vapors at elevated operating temperature of vacuum pump

- · Can also be used in the presence of radically polymerizing substances if inhibitor is used (inquiry necessary)
- · Chemical oil filter advisable except for inhibited NC 2 because inhibitor is made ineffective

Polyglycol-(polyether-)based synthetic up to 140 $^{\circ}\mathrm{C}$ $85 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$ $1.01 \mathrm{\, g \, cm^{-3}}$ 270°C

- Operation at high ambient temperatures
- · Pumping of solvent vapors at elevated operating temperature of vacuum pump

- NC 1 produces an emulsion with mineral oil. Thus, when switching to NC 1, the pump has to be cleaned thoroughly from mineral oil residue. Demounting necessary.
- · Employment of chemical oil filter strongly recommended

Dioctylphthalate (softener)

up to 120°C $28 \times 10^{-6} \text{ m}^2 \text{ s}^{-1}$ 0.99 g cm^{-3} 210 °C

- Extruder degassing
- Pumping of styrene, butadiene, and other easily polymerizing substances

(continued)

Table A.17 (Continued)

Oil type	Universal oil	Special oil Protelen	Special oil NC 2	Special oil NC 1 Fomblin (PFPE)
Formation of crystalline, highly viscous organic decomposition products inside vacuum pumps		Formation of crystalline, highly viscous organic decomposition products inside vacuum pumps	Formation of crystalline, highly viscous organic decomposition products inside vacuum pumps	Formation of crystalline, highly viscous organic decomposition products inside vacuum pumps
Formation of tarry or resinous organic decomposition products inside vac- uum pumps		Formation of tarry or resinous organic decomposition products inside vac- uum pumps	Formation of tarry or resinous organic decomposition products inside vacuum pumps	Formation of tarry or resinous organic decomposition products inside vacuum pumps
			Degassing of and charging with polyether-based brake fluid	
Do not use chemical oil filters	 Impairs ultimate pressure 	 Do not use chemical oil filters 	 Glygoyle 11 is hygroscopic; this impairs ultimate pressure 	Do not use chemical oil filters
	Switching see NC 1		Mineral oil is hardly soluble in Glygoyle 11. Therefore, pumps should be cleaned thoroughly before switching to Glygoyle 11	Low viscosity!
	 Employment of chemi- cal oil filter strongly recommended 		Do not use chemical oil filters	

a) See also Laurenson L. Technology and applications of pump fluids. J. Vac. Sci. Techn. Vol. 20 (1982) No. 4. pp. 989–995.

a) Should no longer be used.
c) Translator's note: German Federal Institute for Materials Research and Testing (Bundesanstalt für Materialforschung und -prüfung).

d) Translator's note: corresponds to ISO 6521.

Table A.18 Specifications of common motive media for diffusion pumps.

1	2	3	4	5	6	7	8	9	10	11	12
Motive medium	Relative molecular mass	Dynamic viscosity (Pa s)	Refractive index	Density (g cm ⁻³)	A ^{g)}	В ⁹⁾ (К)	ϑ _{−5} ⁱ⁾ (°C)	ϑ _{−2} ⁱ⁾ (°C)	p _s (25 °C) (hPa)	Flashing point (°C)	Dripping point (°C)
butyl phthalate	278	2.1 × 10 ⁻² (20 °C)	1.4903 (20 °C)	1.035 (20°C)	13.96	5204	18	81	4.4×10^{-5}	190	_
Narcoi 40(A) ^{a)}	419	_	1.4828 (20°C)	0.973 (20°C)	11.54	5690	73	146	8×10^{-8}	_	=
Octoil S ^{b)}	426	_	_	_	11.39	5514	50	142	2.7×10^{-8}	_	_
Apiezon AP 201 ^{c)}	-	3 × 10 ⁻² (20 °C)	=	0.876 (12°C)	_	_	_	_	5×10^{-6} k)	196	=
DIFFELEN L ^{d)}	440	1.69 × 10 ⁻² (20 °C)	1.4807 (20°C)	0.8849 (20°C)	12.82	6098	71	142	2.4×0^{-8}	232	-27
DIFFELEN N ^{d)}	470	1.92 × 10 ⁻² (20 °C)	1.4802 (20°C)	0.8815 (20°C)	13.27	6329	76	145	1.1×10^{-8}	242	-27
DIFFELEN U ^{d)}	530	2.18 × 10 ⁻² (20 °C)	1.4832 (20°C)	0.8771 (20°C)	13.04	6410	85	156	3.5×10^{-9}	257	-29
DC 704 ^{e)}	484	4.17 × 10 ⁻² (25 °C)	1.5565 (25 °C)	1.07 (25 °C)	11.15	5570	74	155	2.8×10^{-8}	221	-38
DC 705 ^{e)}	546	1.91 × 10 ⁻² (25 °C)	1.579								
(25°C)	1.094 (20°C)	_	_	_	_	$4\cdot 10^{-10}$	243	_			
Polyphenylene	454	1.20×10^{-2} (25 °C)	=	1.2 (25 °C)	_	-	-	_	1.8×10^{-9}	350	=

(continued)

Table A.18 (Continued)

1	2	3	4	5	6	7	8	9	10	11	12
Motive medium	Relative molecular mass	Dynamic viscosity (Pa s)	Refractive index	Density (g cm ⁻³)	A ^{g)}	В ^{g)} (K)	9 _{−5} ⁱ⁾ (°C)	ϑ _{−2} i) (°C)	p _s (25 °C) (hPa)	Flashing point (°C)	Dripping point (°C)
ether ^{f)}											
Mercury	200.6	_	_	13.55 (20°C)	10.67 ^{h)}	3333 ^{h)}	-28	45	2.6×10^{-3}	-	-
Fomblin ^{k)}	_	0.36									
		(20 °C)	-	1.9 (20°C)	_	_	_	-	4×10^{-8}		-42

- Notes:
 a) Bis(3.5.5-trimethylhexyl) phthalate.
 b) Di(2-ethylhexyl) sebacate.
 c) Mixture of hydrocarbons.
 d) Mixture of saturated hydrocarbons.
- e) Organic Si compounds. f) Trade names Convalex 10, Santovac 5, Ultralen.

- 1) Trade finites Convace 10, Santowac 3, Ordinates 1.

 g) A,B: constants in vapor pressure equation $\log \frac{F}{L^2} = A \frac{B}{T}$.

 h) $\log \frac{F}{B^2} = 10.67 \frac{3338}{T} 0.848 \log \frac{T}{K}$ for Hg.

 i) Temperatures in 'C' where the vapor pressures are just 1.3×10^{-5} hPa and 1.3×10^{-2} hPa, respectively.
 j) At 160 °C.

 k) Y-HVAC 18/8; see also Table A.17.

Table A.19 Overview of measuring ranges of commercial vacuum gauges. In the ranges indicated, the specified measuring principles provide acceptable accuracy. IVG: ionization vacuum gauge.

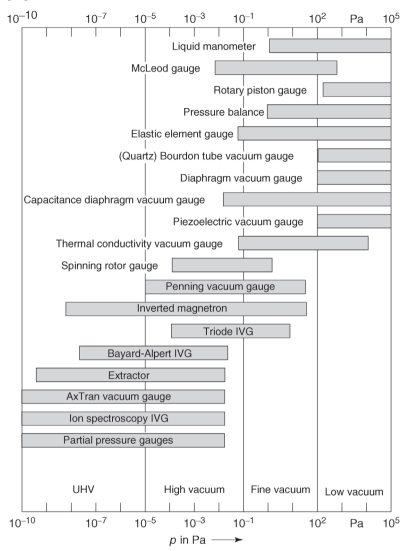


Table A.20 Selected applications of vacuum technology and corresponding pump systems, classified according to working pressure ranges (p_w) . ^{a)}

Typical Applications **Typical Pump Systems** (A) Process medium: air $(p_{w>10 \text{ hPa}})$ Pneumatic transport and handling systems Pumping systems usually comprise only a sin-(e.g., vacuum chucks, vacuum holding, and gle pump. Available pumps include liquid ring transport equipment for large glass panels), vacuum pumps (usually with water as working fluid), oil-free multiple vane vacuum sorting equipment, vacuum packaging, vacuum forming, and altitude cabins for rockets. pumps, and single-stage sliding vane rotary pumps (oil-sealed). See Table A.16 for pressure ranges. (B) Process media: mostly vapors $(p_{\rm w} > {\rm approximately} \ 10 \ {\rm hPa})$ Vacuum distillation, evaporation, crystallization, degassing of mixers, filtration, degassing of oil, and concentrating of aqueous solutions. Vacuum refining (VOB process). Pump systems similar to (A) but often with pre-connected condenser (see Chapter 10). Also applicable are two-stage water ring vacuum pumps with additional gas jet pump. Water jet pump (for small pumping speeds). (C) Process media: vapor air mixtures $(p_{w}$ in the transition range fine/low vacuum) Impregnating, vacuum treatment of trans-Pump systems comprise process-specific former oil, flash distillation, steel degassing, combinations of condensers, main pumps and vacuum drying. (Roots pumps, and vapor jet pumps), and fore pumps (water ring pumps, slidingvane rotary pumps, or rotary plunger pumps). (D) Process media: gases (p_w mainly in the fine-vacuum range) Vacuum-metallurgical processes with high gas Water vapor jet pump; Roots pump (possibly and dust formation (vacuum induction furnamultistage) with suitable forepumps (rotary cesand vacuum arc furnaces), cathode sputpumps and water ring vacuum pumps). tering, air tunnels (at low pressure), and vacuum brazing. (E) Process media: gas vapor mixtures $(p_{\rm w}$ in the fine-vacuum range) Freeze drying Molecular distillation Decorative metal plating Two-stage, oil-sealed rotary vacuum pumps with condensers; Roots pumps with fore pumps. Also vapor jet pumps with fore pumps. (F) Processes under high vacuum $(p_w < 0.1 \, \text{Pa s})$ Thin film technology: metal coatings for plas-Oil diffusion pumps (up to total pumping

speeds of 90 000 ℓ s⁻¹) with additional cold

surfaces in the vacuum vessel; combined oil

tic moldings, paper and plastic foils, hard

material coatings, decorative coatings, optical

Typical Applications	Typical Pump Systems
coatings. Coating of large glass panels (solar glass and thermal barrier coatings); conductive coatings, transparent coatings; data storage equipment, compact discs. Pump systems with turbomolecular pumps, at times combined with cryopumps, and with Roots pumps or rotary pumps as fore pumps	vapor jet pumps and oil-sealed rotary vacuum pumps.
depending on the type of system. Space simulation chambers	Bath cryopumps and refrigerator cryopumps
Space simulation chambers	with overall very high pumping speeds combined with turbomolecular pumps and Roots pumps.
Metallurgical HV processes; electron beam melting and casting; HV degassing; electron beam welding.	HV pump stands usually equipped with turbomolecular pumps, rarely with diffusion pumps. Cryopumps used in large systems.
Electron microscopy (instruments with hot cathodes)	
Electron tubes: receiver tubes, high-power transmitting tubes; klystrons; television picture tubes; image intensifier tubes.	High-vacuum pump stands with oil- or Hg-diffusion pumps (rare) or with turbomolecular pumps. The sealed tubes are often equipped with evaporation getters or small sputter ion pumps (referred to as appendix pumps) for maintaining the very low working vacua ($< 10^{-7}$ mbar).
X-ray tubes	Common HV pump stands. Tubes equipped with Ba evaporation getters.
(G) Processes under ultrahigh vacuum $(p_{\rm w} < 10^{-5} { m Pa})$	
Surface and thin film analysis	
Electron microscopy with cold emission cathodes and analyzing equipment	UHV systems with turbomolecular, sputter ion, and Ti sublimation pumps. Occasionally: oil diffusion pumps with LN ₂ -cooled baffles.
Particle accelerators and storage rings	UHV systems with (at times integrated) sputter ion pumps, Ti sublimation pumps, turbomolecular pumps, <i>in situ</i> NEG strips.
Thermonuclear fusion	Bath cryopumps with very high pumping speeds; turbomolecular pumps.
Semiconductor technology (in certain process steps)	Dry and corrosion-resistant pump systems, with hybrid turbomolecular pumps, claw pumps, diaphragm pumps, and NEG strips.

a) After N. T. M Dennis and T. A. Happell, Vacuum System Design. Chapman and Hall, 1968; N. S. Harris, School of Vacuum Technology, Edwards High Vacuum. Crawly, England, 1981; N. S. Harris, Modern Vacuum Practice. McGraw Hill, 1989, 315 p; W. Pupp and H. K. Hartmann, Vakuumtechnik. Grundlagen und Anwendungen. Carl Hanser, Munich, 1991, 558 p.

Table A.21 Overview of important vacuum processes (first part) and surface analysis techniques (beginning with AES).

Abbreviation	Name	Technique
PVD	Physical vapor deposition	Deposition of thin layers from the vapor phase by sputtering or ion plating
CVD	Chemical vapor deposition	Deposition of coatings involving chemical reactions on the substrate surface or in the process gas. Activation by plasma (PE-CVD), laser, ions, microwaves
RIE	Reactive ion etching	Etching of coatings by means of ions produced in high-frequency plasma
EBW	Electron beam welding	Welding of metals under vacuum, heating the weld seam by means of high-energy electrons
_	Ion implantation	Implanting of high-energy ions (> 1 keV) into a solid (usually semiconductor material).
_	Leak detection	Specimens sprayed with search gas from the outside. The gas penetrating through leaks (usually helium) is detected by a mass spectrometer
AES	Auger electron spectrometry	Chemical element detection using electron bombardment. Electrons from lower elec- tron shells are removed and replaced. The released electrons are analyzed
EELS	Electron energy loss spectroscopy	Energetic analysis of backscattered electrons from a surface, previously impinging the sur- face with low energy
ESCA	Electron spectroscopy for chemical analysis	Outdated designation. See XPS
GCMS/ LCMS	Gas and liquid chromatography. mass spectrometry	Chromatography for timed separation of substances topped with a mass spectrometer
ISS	Ion scattering spectroscopy	Energetic analysis of backscattered ions from a surface
LEED/ HEED/		
RHEED	Low/high/reflection high energy electron diffraction	Detection of diffraction reflections of electron beams
MS	Mass spectrometry	Molecules of chemical compounds or ele- ments are ionized and sorted according to their ratio of charge and mass in magnetic or high-frequency fields
PIXE	Proton induced X-ray emission	Measurement of the characteristic X-ray radiation produced by protons
RBS	Rutherford backscattering	Energy analysis of light ions backscattered from a solid

Table A.21 (Continued)

Abbreviation	Name	Technique
SEM/TEM	Scanning electron microscopy. transmission electron microscopy	Scanning of an object using a focused electron beam. Variations in absorption and scattering are analyzed and translated into a reflection or transmission picture (TEM) on a screen
STM	Scanning tunneling microscopy	Scanning of a surface using an atomic tip held at constant height above the surface by measuring the tunnel current
SIMS	Secondary ion mass spectrometry	MS detection of secondary ions produced by primary ion (noble gases) bombardment
SNMS	Secondary neutral mass spectrometry	MS detection of neutral particles produced by primary ion bombardment
UPS	UV photoelectron spectroscopy	Detection of photoelectrons emitted due to UV radiation
XPS	X-ray photoelectron spectroscopy	Detection of photoelectrons emitted due to X-radiation

Table A.22 National and international standards and guidelines for vacuum technology.

(a) German national standards. European standards denominated with EN.					
DIN	Title	Edition			
EN 1012-2	Compressors and vacuum pumps, safety requirements – Part 2: Vacuum pumps Units	07.1996			
1301	Part 1: Unit names, unit symbols	10.2002			
	Part 2: Submultiples and multiples for general use	02.1978			
	Part 3: Conversions of units no longer to be used	10.1979			
1304	Letter symbols for physical quantities	03.1994			
1306	Density; concepts, presentation of values	06.1984			
1313	Quantities	12.1998			
1314 1319	Pressure; basic definitions, units Fundamentals of metrology	02.1977			
	Part 2: Terminology related to measuring equipment Part 3: Evaluation of measurements of a single measurand	10.2005			
	Measurement uncertainty	05.1996			
	Part 4: Evaluation of measurements; uncertainty of measurement	02.1999			
EN 1330-8	Nondestructive testing – Terminology – Part 8: Terms used in leak tightness testing	07.1998			
1343	Reference conditions; normal condi- tions; normal volume; concepts and values	01.1990			
EN 1518	Nondestructive testing – Leak testing – Characterization of mass spectrome- ter leak detectors	06.1998			
EN 1779	Nondestructive testing – Leak testing – Criteria for the method and technique selection	10.1999			
EN 1593	Nondestructive testing – Leak testing – Bubble emission techniques	11.1999			
ISO 3567	Vacuum gauges – calibration by direct comparison with a reference gauge (in German)	05.2015			
EN 13184	Nondestructive testing – Leak test – Pressure change method	07.2001			

Table A.22 (Continued)

DIN	Title	Edition
EN 13185	Nondestructive testing – Leak testing – Tracer gas method	07.2001
EN 13192	Nondestructive testing – Leak test – Calibration of gaseous reference leaks	03.2002
EN 13463	Nonelectrical equipment for use in potentially explosive atmospheres – Part 5: Protection by constructional safety "c"	03.2004
EN 13625	Nondestructive testing – Leak test – Guide to the selection of instrumentation for the measurement of gas leakage	03.2002
24290	Jet pumps (ejectors); terms, classification	08.1981
24291	Jet pumps (ejectors); denomination of parts	04.1974
28400	Vacuum technology; terms and definitions	
	Part 1: General terms	05.1990
	Part 2: Vacuum pumps	10.1980
	Part 3: Vacuum gauges	06.1992
	Part 4: Vacuum coating technology	03.1976
	Part 5: Vacuum drying and lyophilization	03.1981
	Part 6: Surface analysis techniques	10.1980
	Part 7: Vacuum metallurgy	07.1978
	Part 8: Vacuum systems. components and accessories	11.1997
28401	Vacuum technology – Graphical symbols – Summary	04.2007
28402	Vacuum technology; quantities; symbols; units, summary	12.1976
28403	Vacuum technology; quick release couplings; clamped-type couplings	09.1986
28404	Vacuum technology; flanges; dimensions	10.1986
28410	Vacuum technology; mass spectrome- ter partial pressure gauges; definitions; characteristics; operating conditions	11.1968
28411	Vacuum technology; acceptance speci- fications for mass spectrometer-type leak detector; terms	03.1976
		(continued)

Table A.22 (Continued)

(a) German national standards. European standards denominated with EN.			
DIN	Title	Edition	
28416	Vacuum technology; calibration of vacuum gauges within the range of 10^{-3} to 10^{-7} mbar. general method: pressure reduction by continuous flow		
28417	Vacuum technology; measurement of throughput by the volumetric method at constant pressure	03.1976	
28418	Vacuum technology; standard method for vacuum gauge calibration by direct comparison with a reference vacuum gauge		
	Part 1: Basics	05.1976	
	Part 2: Ionization vacuum gauge	09.1978	
	Part 3: Thermal conductivity gauges	08.1980	
28426	Vacuum technology		
	Part 1: Acceptance specifications for rotary plunger vacuum pumps, sliding vane rotary vacuum pumps, and rotary piston vacuum pumps for the ranges of rough and medium vacuum	08.1983	
	Part 2: Acceptance specifications for roots vacuum pumps for the range of medium vacuum	03.1976	
28427	Vacuum technology; acceptance speci- fications for diffusion pumps and vapor jet vacuum pumps for vapor pressures of the pump fluid < 1 mbar	02.1983	
28428	Vacuum technology; acceptance speci- fications for turbomolecular pumps	11.1978	
28429	Vacuum technology; acceptance speci- fications for ion getter pumps	05.2014	
28430	Vacuum technology; rules for the mea- surement of steam jet vacuum pumps and steam jet compressors using steam as the working fluid	11.1984	
28431	Vacuum technology; acceptance speci- fications for liquid ring vacuum pumps	01.1987	
28432	Vacuum technology; acceptance speci- fications for diaphragm vacuum pumps	09.1996	
45635	Measurement of airborne noise emitted by machines; enveloping surface method Part 13: Compressors, vacuum pumps included (displacement, turbo- and jet compressors)	02.1977	
66038	Torr – millibar, millibar – torr; conversion tables	04.1971	

Table A.22 (Continued)

(b) Ge	erman national guidelines and agreements	
Name	Title	Edition
DKD-R 6-2	Calibration of measuring devices for vacuum	03.2002
	Part 1: Fundamentals	
	Part 2: Measurement uncertainties	
	Part 3: Electrical diaphragm gauges	
	Part 4: Ionization gauges	
	Part 5: Pirani gauges	
(c) Internationa	standards, technical guidelines, and draft standards	
ISO	Title	Edition
554	Standard atmospheres for conditioning and/or testing; specifications	07.1976
1000	SI units and recommendations for the use of their multiples and of certain other units	11.1992
1043	Plastics – Symbols and abbreviated terms – Part 1: Basic polymers and their special characteristics	12.2001
1607	Positive-displacement vacuum pumps; measurement of performance characteristics	
	Part 1: Measurement of volume rate of flow (pumping speed)	12.1993
	Part 2: Measurement of ultimate pressure	11.1989
1608	Vapor vacuum pumps	
	Part 1: Measurement of volume rate of flow (pumping speed)	12.1993
	Part 2: Measurement of critical backing pressure	12.1989
1609	Vacuum technology; flange dimensions	03.1986
2533	Standard atmosphere	05.1975
2861	Quick release couplings; dimensions	
	Part 1: Clamped type	08.1974
	Part 2: Screwed type	08.1980
3529	Vacuum technology – Vocabulary	
	Part 1: General terms	12.1981
	Part 2: Vacuum pumps and related terms	12.1981
	Part 3: Vacuum gauges	12.1981

Table A.22 (Continued)

(c) International standards, technical guidelines, and draft standards			
ISO	Title	Edition	
3530	Vacuum technology; mass spectrometer- type leak-detector calibration	09.1979	
3567	Vacuum gauges – calibration by direct comparison with a reference gauge	12.2011	
3669	Vacuum technology; bakeable flanges; dimensions	02.1986	
TS 3669-2	Vacuum technology – Bakeable flanges – Part 2: Dimensions of knife-edge flanges	08.2007	
5167	Measurement of fluid flow by means of pressure differential devices inserted in circular cross-section conduits running full. Part 1–4.	03.2003	
5298 (CD)	Vacuum technology; partial pressure analyzers of the mass spectrometer type – interpretation of results	02.1997	
5302	Vacuum technology – Turbomolecular pumps – Measurements of performance characteristics	07.2003	
9300	Measurement of gas flow by means of critical flow Venturi nozzles	08.2005	
9803	Vacuum technology – Mounting dimensions of pipeline fittings	05.2007	
	Part 1: Non-knife-edge flange type Part 2: Knife-edge flange type		
14291	Vacuum gauges – Definitions and speci- fications for quadrupole mass spectrometers	07.2012	
14617	Graphical symbols for diagrams Part 3: Connections and related devices	09.2002	
	Part 5: Measurement and control devices		
	Part 8: Valves and dampers		
21250	Part 9: Pumps, compressors and fans	04.000=	
21358	Vacuum technology – Right-angle valve – Dimensions and interfaces for pneu- matic actuator	04.2007	
21360-1	Vacuum technology – Standard methods for measuring vacuum-pump perform- ance – Part 1: General description	06.2007	

(c) International standards, technical guidelines, and draft standards

ISO	Title	Edition
21360-2	Vacuum technology – Standard methods for measuring vacuum-pump perform- ance – Part 2: Positive displacement pumps	01.2012
27892	Vacuum technology — Turbomolecular pumps — Measurement of rapid shutdown torque	01.2010
27893	Vacuum technology – Vacuum gauges – Evaluation of the uncertainties of results of calibrations by direct comparison with a reference gauge	08.2012
27894	Vacuum technology – Vacuum gauges – Specifications for hot cathode ionization gauges	11.2009
27895	Vacuum technology – Valves – Leak test	11.2009

(d) International guidelines and agreementsc

Name	Title	Edition
PNEUROP 5607	Vacuum Pumps. Rules of Acceptance: Part II – Vapor Pumps. Identical to DIN 28427	1972
PNEUROP 5608	Vacuum Pumps. Rules of Acceptance: Part III – Turbomolecular Pumps. Identical to DIN 28428	1973
PNEUROP 5615	Vacuum Pumps. Rules of Acceptance: Part IV – Sputter Ion Pumps. Identical to DIN 28429	1976
PNEUROP 6601	Application of National Standards for Acceptance and Capacity Measurement of Steam Jet Vacuum Pumps and Steam Jet Compressors. Identical to DIN 28430	1978
PNEUROP 6602	Vacuum Pumps. Rules of Acceptance: Part I – Positive Displacement Pumps – Roots Pumps. Identical to DIN 28426	1979
PNEUROP 6606	Vacuum Flanges and Connections — Dimensions. Identical to DIN 28403 and 28404	1985

(continued)

Table A.22 (Continued)

	I guidelines and agreementsc	
Name	Title	Edition
PNEUROP		
PN5ASRCC/5	Pneurop Acceptance Specification for Refrigerator Cooled Cryopumps	07.1989
OIML R 17	Indicating pressure gauges; vacuum gauges and pressure-vacuum gauges; ordinary instruments (in French – English translation available)	1981
ISO	Title	Edition
OIML R 101	Indicating and recording pressure gauges, vacuum gauges, and pressurevacuum gauges with elastic sensing elements (ordinary instruments) (in French – English translation available)	1991
94/9/EC	Directive 94/9/EC of the European Par- liament and the Council of 23 March 1994 on the approximation of the laws of the Member States concerning equip- ment and protective systems intended for use in potentially explosive atmospheres	03.1994
1999/92/EC	Directive 1999/92/EC of the European Parliament and of the Council of 16 December 1999 on minimum require- ments for improving the safety and health protection of workers potentially at risk from explosive atmospheres	12.1999
(e) Recommendations k	by the American Vacuum Society AVS	
Title of recommendation	Published in	
Recommended procedure for measuring pumping speeds	J. Vac. Sci. Technol. A 5 (1987), 2552	
American Vacuum Society recommended practices for pumping hazardous gases	J. Vac. Sci. Technol. A 6 (1988). 1226	
A survey of vacuum material cleaning procedures: A subcommittee report of the American Vacuum Society Recom- mended Practices Committee	J. Vac. Sci. Technol. A 9 (1991), 2025	
Recommended practices for the cali- bration and use of capacitance dia- phragm gauges as transfer standards	J. Vac. Sci. Technol. A 9 (1991), 2843 – 28	363
Recommended practices for the calibration and use of leaks	J. Vac. Sci. Technol. A 10 (1992), 1 – 17	

(e) Recommendations by the American Vacuum Society AVS Recommended practice for the calibra-J. Vac. Sci. Technol. A 11 (1993), A22-A40 tion of mass spectrometers for partial pressure analysis Recommended practices for measuring J. Vac. Sci. Technol. A 17 (1999), 3081 the performance and characteristics of closed-loop gaseous helium cryopumps Recommended practice for calibrating J. Vac. Sci. Technol. A 18 (2000), 2568 - 2577 vacuum gauges of the thermal conductivity type Recommended practice for measuring J. Vac. Sci. Technol. A 20 (2002), 1667 and reporting outgassing data

Short versions of many of the older standards listed are given in A. J. Schubert. Normen und Empfehlungen der Vakuumtechnik. In: Vakuum in der Praxis (1991). 211 - 217.

PNEUROP is the European committee of manufacturers of compressors; vacuum pumps; pneumatic tools and allied equipment.

DIN; DIN EN; and ISO standards are available from Beuth Verlag, Berlin; PNEUROP standards from www.pneurop.org.

 Table A.23 Common stainless steel (chromium/nickel steel) types in vacuum technology.

Materia	l number		Note		Relative comp	position (%)	
EN	AISI	DIN		С	Cr	Ni	Мо
1.4301	304 (V2A)	X5CrNi18-10	Stable up to 600 °C, weakly magnetic, easy to weld, very good to polish, not expensive	≤0.07	17.5 19.5	8.0 10.5	
1.4303	305	X4CrNi19-11	Easy to weld, cannot be hardened, polishable, electropolishable	≤0.06	17 19	11 13	
1.4305	303	X8CrNiS18-9	Alloy with sulfur, very easy to machine, cannot be shaped cold, cannot be welded or electropolished	≤0.1	17 19	8 10	
1.4306	304 L	X2CrNi19-11	Increased resistance to intercrystalline corrosion, can be shaped cold, easy to weld, well polishable, stable up to 500°C	≤0.03	18 20	10 12	
1.4307	304 L	X2CrNi18-9	As 1.4306, widespread, about same corrosion resistance but less expensive compared to 1.4301	≤0.03	17.5 19.7	8 10	
1.4310	301	X10CrNi18-8	Ferromagnetic	0.05 0.15	16 19	6 9.5	≤0.08
1.4401	316	X5CrNiMo17- 12-2	Can be shaped cold and hot, easy to weld, stable up to 500°C	≤0.07	16.5 18.5	10 13	2 2.5
1.4404	316 L (V4A)	X2CrNiMo17- 12-2	Stable up to 700 °C, easy to weld, well polishable, low magnetic permeability	≤0.03	16.5 18.5	10 13	2 2.5
1.4435	316 L	X2CrNiMo18- 14-3	As above, ferrit content is defined	≤0.03	17 19	12.5 15	2.5 3
1.4429	316 LN	X2CrNiMo17- 13-3	Very hard, very low magnetic permeability, easy to weld, well polishable, electropolishable, stable up to $900^{\circ}\mathrm{C}$	≤0.03	16.5 18.5	11 14	2.5 3
1.4541	321	X6CrNiTi18-10	Weakly magnetic, easy to weld, cannot be electropolished, stable up to 850 °C	≤0.08	17 19	9 12	
1.4568	631	X7CrNiAl17-7	Magnetic, suitable for foils and wires	≤0.09	16 18		6.5 7.8
1.4571	316 Ti	X6CrNiMoTi17- 12-2	Can be polished, but may show scratches	≤0.08	16.5 18.5	10.5 13.5	2 2.5

В Diagrams

Figure B.1a	Standard atmosphere: atmospheric pressure and temperature versus height above sea level.
Figure B.1b	Standard atmosphere: gas composition versus height above sea level.
Figure B.2	Gas-kinetic quantities versus pressure for air at 20 $^{\circ}\text{C}.$
Figure B.3a,b	Saturation vapor pressures of important substances in vacuum technology versus temperature.
Figure B.4	Saturation vapor pressures of solvents and detergents.
Figure B.5	Saturation vapor pressures of motive media.
Figure B.6	Saturation vapor pressures of vacuum greases.
Figure B.7	Vapor pressures of selected elastomers.
Figure B.8a	Saturation vapor pressures of selected substances in the temperature range $T\!=\!1\mathrm{K}\!-\!400\mathrm{K}.$
Figure B.8b	Saturation vapor pressures of selected substances for cryotechnology.
Figure B.9a–e	Fundamental quantities of gas flow.
Figure B.10	Flow conductances of circular apertures for molecular flow.
Figure B.11	Surface area-related gas emissions of selected materials versus time.
Figure B.12	Conductance of permeation for selected combinations of solids and gas species.
Figure B.13	Nomogram for determining pump-down times of a vessel in the rough-vacuum range.
Figure B.14	Calibration curves for thermal conductivity vacuum gauges.
Figure B.15	Break-through voltage (sparking voltage) for air (Paschen curve).

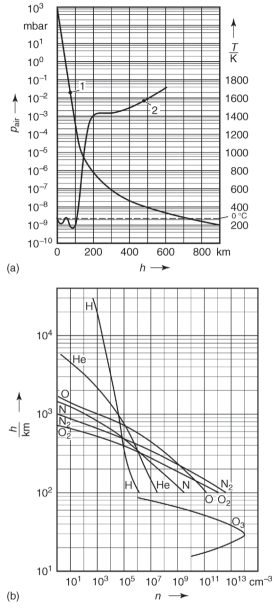


Figure B.1 (a) Standard atmosphere: atmospheric pressure $p_{\rm air}$ (1) and temperature T (2) versus height h above sea level. Note that the temperature scale is linear. (Guideline values.

For internationally standardizes values refer to ISO/DIN 2533). (b) Standard atmosphere: gas composition versus height *h* above sea level. *n*: particle number density.

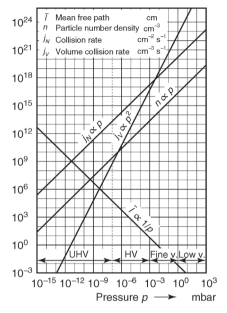


Figure B.2 Mean free path \bar{l} , Eq. (3.55), particle number density n, Eq. (3.19), collision rate j_N , Eq. (3.48), and volume collision rate j_V , Eq. (3.60) for air at $\vartheta = 20$ °C versus pressure.

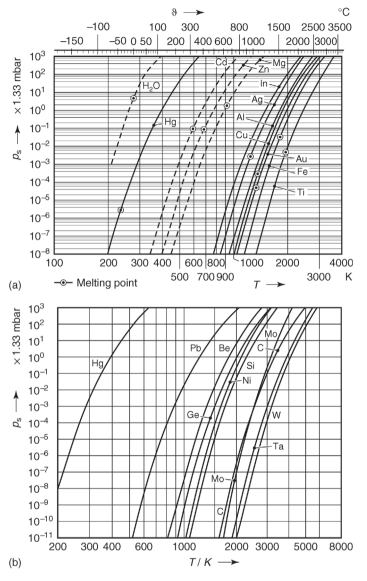


Figure B.3 (a) Saturation vapor pressures p_s of important substances in vacuum technology and of water versus temperatures T and ϑ . For H₂O see also Table A.13. Dashed lines: curves for disturbing substances in vacuum technology. After R. E. Honig, RCA Review XXIII (1962)

567 ff. (b) Saturation vapor pressures $p_{\rm s}$ of important metals in vacuum technology, of carbon, of the semiconductors silicon and germanium, and for comparison, of mercury versus temperature T. From Honig l.c.; — • — melting points.

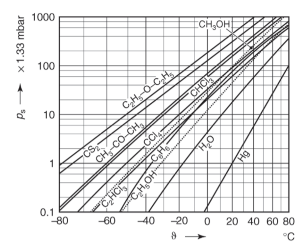


Figure B.4 Saturation vapor pressures p_s of common solvents and detergents versus temperature ϑ . Note that the p_s values obtained from the figure for Hg have to be divided by 1000. C₂H₅-O-C₂H₅ Diethyl ether. CS₂ Carbon

disulphide. CH₃-CO-CH₃ Acetone. CHCl₃ Chloroform. CCl₄ Carbon tetrachloride. CH₃-OH Methanol. C₂HCl₃ Trichloroethylene, Trilene. C₆H₆ Benzole. C₂H₅OH Ethanol.

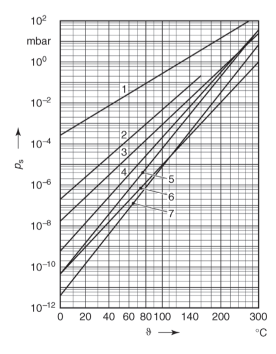


Figure B.5 Saturation vapor pressures p_s of motive media for fluid entrainment pumps versus temperature ϑ . See also Table A.10. 1 Mercury. 2 Fomblin Y-LVAC 06/6. 3 High-

vacuum oil 'light'. 4 High-vacuum oil 'standard'; Fomblin Y-HVAC 18/8. 5 Ultrahighvacuum oil. 6 Ultralen, Convalex 10, Santovac 5. Silicone oil DC 705.

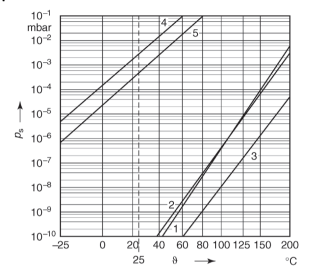


Figure B.6 Saturation vapor pressures p_s of vacuum greases and picein versus temperature ϑ . 1 Grease P. 2 Grease R. 3 Silicone grease. 4 Picein. 5 Ramsay grease.

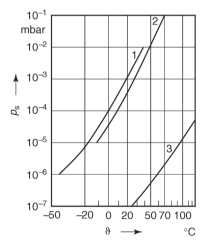


Figure B.7 Vapor pressures $p_{\rm s}$ of elastomers versus temperature ϑ . 1 Perbunan. 2 Silicone rubber. 3 Teflon.

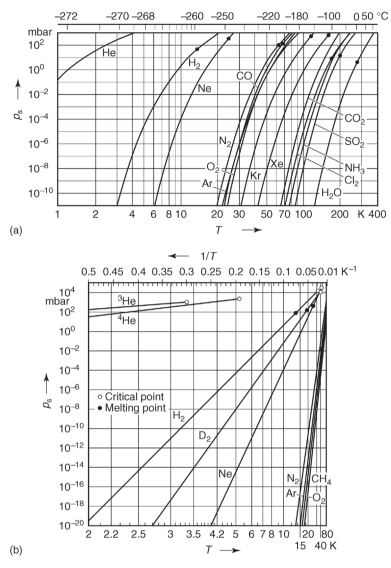


Figure B.8 (a) Saturation vapor pressures p_s of selected substances in the temperature range T = 1 K-400 K. $-\bullet$ - Melting points. (b) Saturation vapor pressures p_s of selected substances in cryotechnology in the temperature range T = 2 K-80 K.



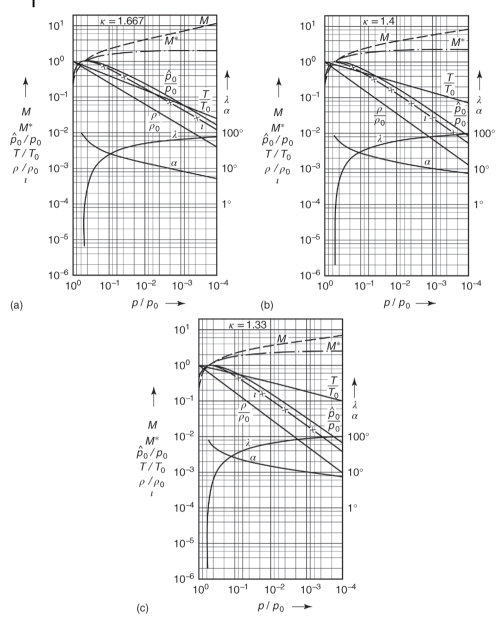


Figure B.9 (a)–(e) Fundamental quantities of gas flow in a compression shock (see Section 4.2) versus the expansion ratio p/p_0 for selected values of the isentropic exponent κ . Instead of the usual Ma, the Mach number here is denominated by M. \hat{p}_0 is the static pressure behind the shock (gas at rest), p_0 is the static pressure in front of the shock, p the static pressure in the tube, thus, $p/p_0 = 0-1$ is the expansion ratio.

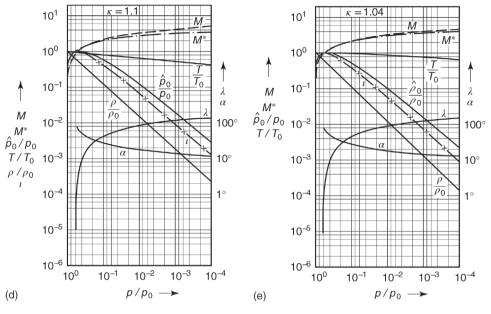


Figure B.9 (continued)

Details:

M = Ma (Mach number): see Eq. (4.57). $M^* = Ma^*$ (critical *Mach* number):

$$Ma^* = \frac{\nu}{\nu^*} = \left(\frac{\kappa + 1}{\kappa - 1} \left[1 - \left(\frac{p}{p_0}\right)^{\frac{\kappa - 1}{\kappa}}\right] \right)^{1/2}.$$

 \hat{p}_0/p_0 static pressure ratio:

$$\frac{\hat{p}_0}{p_0} = \frac{\kappa + 1}{\kappa - 1} \left(\frac{p}{p_0} \right)^{1/\kappa} \frac{\left(1 - \left(\frac{p}{p_0} \right)^{(\kappa - 1)/\kappa} \right)^{\kappa/(\kappa - 1)}}{\left(\frac{4\kappa}{(\kappa + 1)^2} - \left(\frac{p}{p_0} \right)^{(\kappa - 1)/\kappa} \right)^{1/(\kappa - 1)}}.$$

 T/T_0 temperature ratio: see Eq. (4.42). *ι* density of flow ratio:

$$i = \frac{\rho \nu}{\rho^* \nu^*} = \left(\frac{\left(\frac{\kappa+1}{2}\right)^{\frac{\kappa+1}{\kappa-1}}}{\frac{\kappa-1}{2}} \left[\left(\frac{p}{p_0}\right)^{2/\kappa} - \left(\frac{p}{p_0}\right)^{\frac{1+\kappa}{\kappa}} \right] \right)^{1/2}.$$

 $ho/
ho_0$ density ration: see Eq. (4.42). λ flow parameter (in degrees of angle): see Eq. (4.71). α Mach angle (in degrees of angle): $\sin \alpha = \frac{1}{Ma}$.

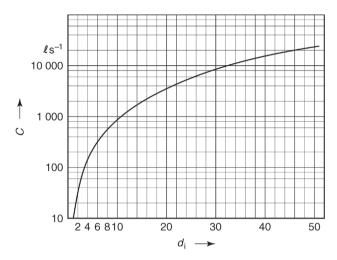


Figure B.10 Conductances *C* of a circular aperture with the diameter d_i for air at $\vartheta = 20\,^{\circ}$ C (see also Table 4.4) under molecular flow conditions.

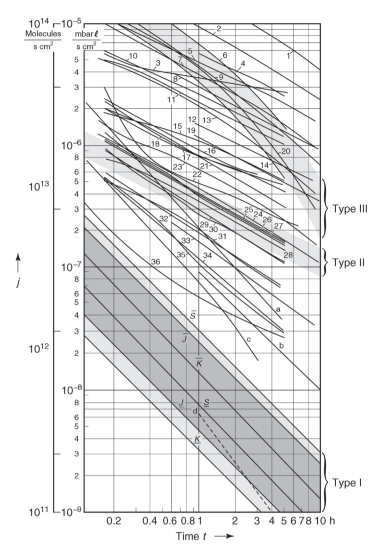


Figure B.11 Surface area-related gas emissions (outgassing flow rates) j of selected materials at $\theta = 20$ °C versus time t. Type I: metals $j \propto t_0/t$. Type II: plastics $j \propto \sqrt{t_0/t}$. Type III: plastics $j \propto \exp(-t_0/t)$. \overline{J} , J, \overline{K} , K, \overline{S} , S: upper and lower limits according to different authors. From K. Diels and R. Jaeckel, Vakuum-Taschenbuch, 2nd Ed., Berlin, 1962. 1 Vulkollan. 2 Perbunan and Buna. 3 Mowital. 4 Mowilith. 5 Neoprene (45/Ne 747). 6 Silicone rubber. 7 Natural rubber. 8 Perbunan. 9 Perbunan. 10 Perbunan. 11 Polyamide. 12 Araldite.

13 Neoprene (35/Ne 746). 14 Silicone rubber (O-ring seal). 15 Plexiglas. 16 Polyvinylcarbazole. 17 Polyvinylcarbazole. 18 Polycarbonate. 19 Araldite. 20 Silicone (37/Si 502). 21 Ultramid. 22 PVC. 23 Viton (25/Vi 575). 24 Teflon (3/Tf 528). 25 Araldite. 26 Polymethane. 27 Viton. 28 Viton. 29 Polystyrene. 30 Polystyrene. 31 Polystyrene. 32 Teflon. 33 Teflon. 34 Polyethylene. 35 Polyethylene. 36 Hostaflon. a Pyrophyllite. b Steatite (Al₂O₃). c Degussit (Al₂O₃). d Pyrex glass.

Notes on Figure B.11 and Table B.14:

Outgassing flow rates are (at times heavily) influenced by the previous history (fabrication process, thermal pretreatments), surface conditions, surface modification (mechanical, chemical, physical), occluded gas species, and of course, temperature. Deviations in measuring results are also due to measuring methods and the material's condition at the time when measurements are engaged.

Note that some publications use m^2/s as unit for \overline{q}_{perm} (ordinate in Figure B.12); conversion:

$$1\frac{\text{mbar }\ell}{\text{s}} \cdot \frac{\text{mm}}{\text{m}^2 \text{ bar}} = 1 \cdot 10^{-9} \frac{\text{m}^2}{\text{s}}$$

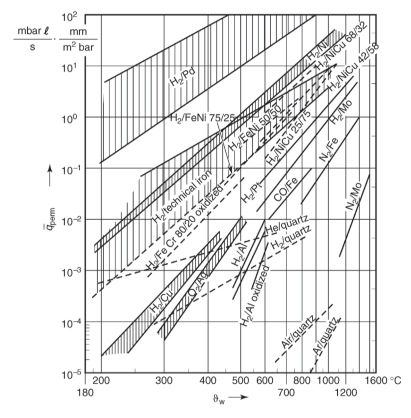


Figure B.12 Conductances of permeation (specific gas permeation fluxes, see Section 16.3 and Eq. (16.2)) \overline{q}_{perm} for selected metals and metal alloys for gas species H_2 , N_2 , O_2 , and O_1 , measured at O_2 and O_3 measured at O_4 measured at

For comparison, \overline{q}_{perm} of quartz glass for He, H₂, air, and Ar. ϑ_w : temperature of solid (wall temperature). According to E. Waldschmidt, [3] in Chapter 16.

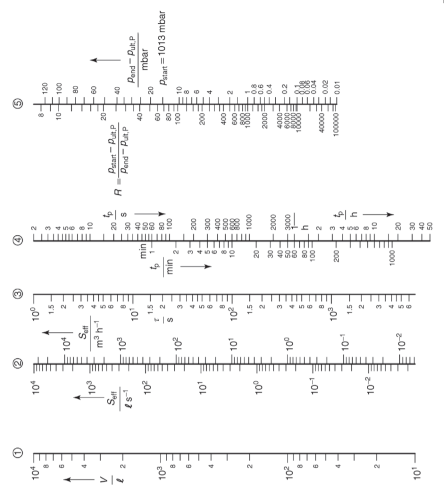


Figure B.13 Nomogram for determining pump-down times t_p of a vessel in the roughvacuum range. Scale (1): vessel volume V in liters. Scale (2): maximum of effective pumping speed Seff, max at the vessel in (left) liters per second and (right) cubic meters per hour. Scale ③: time constant, e-value time τ in seconds according to Eq. (18.9), $\tau = V/S_{eff,max}$. Scale (4): Pump-down time t_p in (top right) seconds and (left center) minutes and (bottom right) hours according to Eqs. (18.10) and (18.11). Scale (5): pressure p_{end} in millibar at the end of pumping, for atmospheric starting pressure at the beginning of pumping action $p_{\text{start}} \approx p_{\text{n}} = 1013 \text{ mbar}$. The desired final pressure p_{end} has to be reduced by the ultimate

pressure of the pump $p_{ult,P}$; the difference is the value used for the scale. In the case of in flow $q_{pV(i)}$, the value used on the scale is $p_{\text{end}} - P_{\text{ult, }P} - q_{pV,\text{in}}/S_{\text{eff, max}}$. Left: pressure reduction ratio $R = (p_{\text{start}} - p_{\text{ult,P}} - q_{pV,\text{in}}/S_{\text{eff, max}})/S_{\text{eff, max}}$ $(p_{\rm end} - p_{\rm ult,P} - q_{pV, \rm in}/S_{\rm eff, max})$,for a starting pressure of p_{start} and a final pressure of p_{end} . The pressure-dependency of the pumping speed is taken in to account in the nomogram according to Eq. (18.6), and is expressed in scale (5). If the pressure of the pump $p_{ult,P}$ is small compared to the desired final pressure p_{end} at the end of pumping action, then this corresponds to a constant pumping speed S or Seff throughout the entire pumping process; see Eq. (18.6).

Example 1 to nomogram 20.13

A vessel with volume $V=2000~\ell$ is evacuated from $p_{\rm start}=1000~{\rm mbar}$ (atmospheric pressure) down to $p_{\rm end}=10^{-1}~{\rm mbar}$ using a rotary plunger pump. The effective pumping speed at the vessel $S_{\rm eff,\ max}=60~{\rm m}^3~{\rm h}^{-1}=16.7~\ell~{\rm s}^{-1}$. Two steps are necessary to obtain the pump-down time from the nomogram:

- 1) Determination of τ : draw a straight line through $V=2000~\ell$ (scale ①) and $S_{\rm eff}=60~{\rm m}^3~{\rm h}^{-1}=16.7~\ell~{\rm s}^{-1}$ (scale ②). The value at the point of intersection of the line and scale ③ is $\tau=120~{\rm s}=2~{\rm min}$ (note that this procedure bares an uncertainty of approximately $\Delta \tau=\pm10~{\rm s}$, i.e., the relative uncertainty amounts to approximately 10 per cent).
- 2) Determination of t_p : according to manufacturer's data, the ultimate pressure of the rotary plunger pump $p_{\rm ult,P} = 3 \cdot 10^{-2}$ mbar. The system is clean and leaks are negligible $(q_{pV,\rm in}=0)$. Thus, $p_{\rm end}-p_{\rm ult,P}=10^{-1}$ mbar $-3 \cdot 10^{-2}$ mbar = $7 \cdot 10^{-2}$ mbar. Now draw a straight line through the point of intersection found in 1), $\tau=120$ s (scale ③), and $p_{\rm end}-p_{\rm ult,P}=7 \cdot 10^{-2}$ mbar (scale ④). The new point of the intersection with scale ⑤ reads $t_{\rm P}=1100$ s ≈ 18.5 min. (Again, the relative uncertainty of the procedure is in the order of 10 per cent, leading to an uncertainty for $t_{\rm P}$ of approximately 15 per cent.) With a safety factor of 20 per cent (see Section 18.3.6), the expected pumping time $t_{\rm P}=18.5$ min $\cdot (1+15\%+20\%)=18.5$ min 1.35=25 min.

Example 2 to nomogram 20.13

The same clean and dry vacuum system $(q_{pVin} = 0)$ as in Example 1 with $V = 2000 \ \ell$ is pumped down to $p_{\rm end} = 10^{-2} \, \rm mbar$. Since this pressure is below the ultimate pressure of the rotary plunger pump ($S_{\rm eff,max} = 60 \,\mathrm{m}^3 \,\mathrm{h}^{-1} = 16.7 \,\ell$ s⁻¹, $p_{\text{ult},P} = 3 \cdot 10^{-2}$ mbar) we will employ a series connection of a rotary plunger pump and a Roots pump. The latter has a starting pressure $p_1 = 20$ mbar, the pumping speed $S_{\rm eff,max}$ = 200 m³ h⁻¹ \approx 55 ℓ s⁻¹, and $p_{\rm ult,P}$ = 4 \cdot 10⁻³ mbar. Thus, the rotary plunger pump will be used for evacuating from $p_{\text{start}} = 1000 \,\text{mbar}$ down to p = 20 mbar and the Roots pump will be added for the pressure range from $p_1 = 20$ mbar down to $p_{\text{end}} = 10^{-2}$ mbar, while the rotary plunger pump (now the fore pump) continues operating. For the first pumping phase, the nomogram (straight line through $V = 2000 \ \ell$ and $S_{\rm eff} = 16.7 \ \ell \ {\rm s}^{-1}$) yields the time constant $\tau = 120 \text{ s} = 2 \text{ min}$, just as in Example 1. If this point on scale 3 is $p_{\text{ult,P}}$ is neglected, i.e., the pumping speed of the rotary plunger pump is constant throughout the entire pressure range 1000 mbar-20 mbar, see Eq. (18.6)), we find $t_{\rm P,1} = 7.7$ min. The Roots pump pumps down from $p_1 = 20$ mbar to $p_{\rm end} =$ 10^{-2} mbar, i.e., the pressure reduction ratio $R = (20 \text{ mbar} - 4 \cdot 10^{-3} \text{ mbar})/$ $(10^{-2} \,\mathrm{mbar} - 4 \cdot 10^{-3} \,\mathrm{mbar}) = 20/(6 \cdot 10^{-3}) = 3300.$

The time constant (straight line between $V=2000\ \ell$ on ① and $S_{\rm eff}=55\ \ell$ s⁻¹ on ② amounts to $\tau=37$ s (on ③). Connecting this point on ③ with R=3300 on ③ yields $t_{\rm p,2}=290$ s = 4.8 min on ④. Adding $t_{\rm sw}=1$ min for switching leads to a total pump-down time of $t_{\rm p}=t_{\rm p,1}+t_{\rm sw}+t_{\rm p,2}=7.7$ min + 1 min + 4.8 min = 13.5 min.

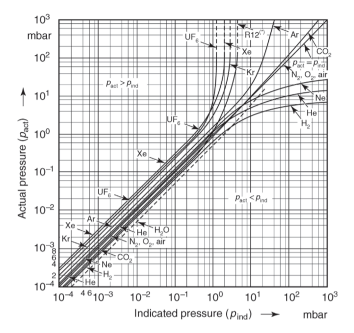


Figure B.14 Calibration curves for a thermal conductivity vacuum gauge with respect to air = $N_2 = O_2$.* R12 = Freon (see Section 13.5). For other thermal conductivity gauges, these

calibration curves represent guideline values only. Also, gas species dependencies (correction factors) normalized to nitrogen can differ well around 20 per cent between gauges.

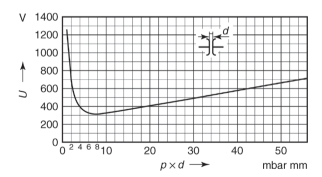


Figure B.15 Break-through voltage (sparking voltage) U between two parallel plates in a homogeneous electric field versus the product

 $p \times d$ (gas pressure \times plate distance) for air (Paschen curve).

Common Abbreviations

AISI	American Iron and Steel Institute
B.P.	Boiling point
CF	Conflat (flange)
DIN	Deutsches Institut fur Normung (German Institute for Standardization)
DN	Diameter nominal
F.P.	Fusing point
GB	Gas ballast
HV	High vacuum
ISO	International Standardization Organization
ITS-90	1990 International Temperature Scale
IUPAP	International Union for Pure and Applied Physics
IVC	International Vacuum Congress
KF	Adopted by ISO from the German Kleinflansch (small flange)
LF	Light flange = clamped flange
LHe	Liquid Helium
LN_2	Liquid Nitrogen
NEG	Non evaporable getter
NTP	Normal Temperature Pressure = at standard temperature and standard pressure, usually not used here; see (T_n, p_n)
OFHC	Oxygen-free high conductivity copper
PF	Pneurop flange
PNEUROP	European committee of manufacturers of compressors, vacuum pumps, pneumatic tools and allied equipment
PTB	Physikalisch-Technische Bundesanstalt (German National Metrology Institute)
SI	Système international d'unités = International system of units
(T_n, p_n)	Indicating that a gas amount refers to standard conditions, i.e., standard temperature T_n and standard pressure p_n (e.g., m^3 (T_n , p_n) used here instead of m^3 (NTP))
TP	Triple p oint (temperature)
UHV	Ultrahigh vacuum

D

Ouantities and Units

This book describes quantitative physical relationships with quantities and quantity equations according to DIN 1313 and ISO 31.

The following sections provide a selection of helpful advice and notes to the reader.

Physical Quantities

Measuring means quantitatively comparing analogous characteristics (properties such as length) of two objects, processes, or events. Using the property of interest (length) of an arbitrary object (gauge block) or instance (light wave) as unit, the result of the comparison yields the numeric value of the ratio of the property and the property unit. Vice versa, we obtain the definition:

Physical quantity = numeric value × unit

or, using symbols:

$$Q = \{Q\} \times [Q] \tag{1}$$

i.e., the physical quantity Q is in fact a product of the numeric value $\{Q\}$ and the selected unit [Q].

For example, in the special case of a *length* with the *symbol* l and the unit meter (unit symbol m),

$$l = 7 \times m = 7 \cdot m = 7 m \tag{2}$$

Comparing this with Eq. (1) yields:

Numeric value of the considered length is
$$\{l\} = 7$$

Selected unit for the length is $[l] = m$

According to Eqs. (1) and (2), appropriate notations for table headings or coordinate axes are:

$$\frac{l}{m}$$
, or generally, $\frac{Q}{[Q]}$, i.e., $\frac{p}{Pa}$ etc.

In addition, we may also write l in m, p in Pa.

2

Quantity Equations, Coherent units

Quantity equations describe interrelationships of physical quantities. They are valid independent of the units used. When using arbitrary units, each quantity is put into the quantity equation as a product of numeric value and unit. The result then is a product of a numeric value and a mix of units. Example: Eq. (4.138) with P = 1:

$$\begin{split} q_N &= A \frac{\overline{c}}{4} (n_1 - n_2) \end{split}$$
 If $A = 8 \text{ cm}^2$, $\overline{c} = 500 \text{ m} \cdot \text{s}^{-1}$, $n_1 = 10^{18} \text{ m}^{-3}$, $n_2 = 0$, then
$$q_N = 8 \text{ cm}^2 \cdot 500 \text{ m} \cdot \text{s}^{-1} \frac{1}{4} 10^{18} \text{ m}^{-3} = 10^{21} \text{ cm}^2 \text{ m}^{-2} \text{ s}^{-1} \end{split}$$

Mixed units have to be converted to a common unit.

Thus, it is appropriate to only use the units of a coherent system of units. In such a system, units are connected only by unit equations containing no numeric factors different from 1. The SI with its base units meter (m), kilogram (kg), second (s), ampere (A), kelvin (K), mole (mol), and candela (cd) as well as the derived units obtained from the defining equations of the other quantities is such a system. For the example using Eq. (4.138), we should then use $[A] = m^2$, $[\overline{c}] = \text{m} \cdot \text{s}^{-1}$, $[n] = \text{m}^{-3}$, and $[q_N] = [A] \cdot [\overline{c}] \cdot [n] = \text{m}^2 \,\text{m} \cdot \text{s}^{-1} \,\text{m}^{-3} = \text{s}^{-1}$.

It is advisable to always put in quantity values as numeric value × unit when calculating equations because this simultaneously provides a unit check, see, e.g., Example 4.9 etc.

When putting in units in expressions with long fraction bars, it is often safer to write units in the denominator (e.g., in m/s, read as meters per second, a fraction) by using negative exponents, i.e., $m \cdot s^{-1}$. Inorder to avoid mistaking this for milliseconds (ms), use either a blank between m and s or a multiplication point, i.e., $m \cdot s^{-1}$.

Fitted Quantity Equations

In fitted quantity equations, each quantity is divided by the unit specific to the particular case, e.g.,

$$\frac{q_N}{s^{-1}} = \frac{10^{-4}}{4} \cdot \frac{A}{cm^2} \cdot \frac{\overline{c}}{m \cdot s^{-1}} \cdot \frac{n}{m^{-3}}$$
(3)

Here, each quantity is put in using the unit specified in the quantity's denominator. A numeric factor is necessary when mixed units are used. In Eq. (3), this is the factor 10^{-4} .

Numeric-value Equations

Numeric-value equations explain relationships between numeric values of quantities. They include additional units applicable to the numeric values. An example is Eq. (4.105).

Ε Glossary, Symbols of Physical Quantities and their SI Units used in this Book

Symbol	Quantity	Unit	Defining equation	Section
a	Distance, length	m		
а	Thermal diffusivity, thermal conductivity, heat conductivity	m^2/s	3.96	3.3.3
a	Speed of sound	m/s	4.55	4.2.3
a_E	Energy-accommodation coefficient	1		3.3.3
A	Surface area	m^2		
$A_{\rm C}$	Surface area of capillary	m^2		
$A_{\rm c.s.}$	Surface area of cold surface	m^2		
A_m	Specific surface area	m ² /kg		11.2.1
$A_{\rm S}$	Surface area ofsphere	m^2		
$A_{\rm p}$	Effective pumping (suction) area ofa vacuum vessel, inlet area ofa pump	m^2		6.2.3
$A_{\rm r}$	Relative atomic mass	1	3.15	3.1.2
$A_{\rm S}$	Inner surface area of avacuum vessel	m^2		
$ ilde{b}$	Amount of adsorbed gas per unit area	Pa ℓ/m^2	11.1	11.2.1
$ ilde{b}_{ m n}$	Amount of adsorbed gas per unit area at standard temperature	Pa ℓ/m²	11.3	11.2.1
B	Magnetic field strength	T		
$B^{\prime\prime}$	Second virial coefficient	Pa^{-1}	3.114	3.4.1
c	Velocity, speed	m/s		
\overline{c}	Mean thermal speed	m/s	3.43	3.2.3
c_p	Specific heat capacity at constant pressure	$\rm J~K~kg^{-1}$	3.82	3.3.3
c_{mp}	Most probable speed of a <i>Maxwell-Boltzmann</i> ensemble	m/s	3.42	3.2.3
$c_{ m rms}$	Root-mean-square speed	m/s	3.44	3.2.3
c_V	Specific heat capacity at constant volume	$\rm J~K~kg^{-1}$	3.81	3.3.3
C	Conductance	ℓ/s	4.27	4.1.3
C	Heat capacity	J/K		3.3.3
C_m	Mass capacity of a pump	kg		
			(0	continued)

Symbol	Quantity	Unit	Defining equation	Section
$C_{ m molar}$	Molar heat capacity	J K ⁻¹ mol ⁻¹	,	
C_p	Heat capacity at constant pressure	$\rm J~K~kg^{-1}$	3.82	3.3.3
C_V	Heat capacity at constant volume	J/K		3.3.3
C_{pV}	pV capacity of a pump	Pa €		
CF	Correction factor of reading	1	15.22	15.2.3
d	Geometrical quantity such as diameter, thickness, etc.	m		
D	Diffusion coefficient	cm ² /s		3.3.4
D_0	Temperature-independent diffusion coefficient	cm ² /s	6.28	6.3
e	Error of reading	1	15.21	15.2.3
$E_{ m act}$	Activation energy	kJ/mol, eV		6.1
E_{ad}	Heat or energy of adsorption	kJ/mol, eV		6.1
$E_{ m des}$	Desorption energy	kJ/mol, eV		6.1
$E_{ m dif}$	Activation energy of diffusion	kJ/mol, eV	6.28	6.3
$E_{\rm kin}$	Kinetic energy	kJ/mol, eV		
$E_{\rm S}$	Enthalpy of solution	kJ/mol, eV	6.45	6.4
f	Degree of freedom	1		
f	Standardized fraction of physical unit	1		
F	Force	N		
$F_{ m R}$	Frictional force	N		
G	Gain factor	1	14.20	14.2.5.5
G_{c}	Growth rate of condensate layer	m/s	12.29	12.4.2.8
g	Acceleration due to gravity	m/s ²		3.1.1, 15.2.1.1
H	Enthalpy	J	12.5	12.2.1
h	Height	m		
h	Specific enthalpy of evaporation	J, eV	3.126	3.5.1
i	Counter for countable set	1		
I	Electrical current	A		
I^+	Ion current	A		
I_e	Electron current	A		
J	Surface area-related flow rate	$m^{-2} s^{-1}$	3.48	3.2.4
$j_{ m ad}$	Flow rate of adsorption	$m^{-2} s^{-1}$		
$J_{ m des}$	Flow rate of desorption	$m^{-2} s^{-1}$		
$j_{ m dif}$	Flow rate of diffusion	$m^{-2} s^{-1}$		
j_N	Molecular flow rate	$m^{-2} s^{-1}$		
$j_{ m rec}$	Rate of recombination given as molecules per unit time	s^{-1}	6.41	6.3

Symbol	Quantity	Unit	Defining equation	
$j_{ m perm}$	Permeation rate	$m^{-2} s^{-1}$		
j _V	Volume flow rate	m/s	4.179	4.5.2
k	Boltzmann's constant, black body constant	J/K	3.21	3.1.3
K	Constant of proportionality			
K	Compression ratio	1	7.13, 10.12	7.4.3.5, 10.3.2
$K_{ m perm}$	Permeation constant, permeation conductivity	$\ell \text{ s}^{-1} \text{ cm}^{-1}$	6.44	6.4
$K_{\rm S}$	Solubility constant	1	6.45	6.4
Kn	Knudsen number	1	4.1	4.1.1
l	Length	m		
Ī	Mean free path	m	3.55	3.2.5
т	Mass	kg		
m	Degree of modulation	1		
m	Exponent, e.g., for reading-pressure dependency	1		
$m_{ m P}$	Particle mass	kg		
M	Torque, angular moment	Nm		
M,	Molar mass	kg/mol		
$M_{ m molar}$				
$M_{ m r}$	Relative atomic mass	1		
n	Number of countable amount	1		
n	Number density of atoms or molecules	m^{-3}	3.10	3.1.2
ñ	Number density per unit area	m^{-2}		6.1
$n_{\rm G}$	Volume number density	m^{-3}	3.10	3.1.2
$n_{ m dis}$	Number density in bulk material	m^{-3}		
$\tilde{n}_{ ext{mono}}$	Surface area-related number density of monolayer	1	6.1	6.1
$n_{\rm n}$	Number density under standard conditions	m^{-3}		
$n_{\rm S}$	Surface area-related number density of particles on a surface	m^{-2}		
N	Number of particles	1		
N	Number of revolutions	1		
$N_{\rm A}$	Avogadro's constant	1	3.7	
$N_{\rm ad}$	Number of adsorbed particles	1		
p	Pressure	Pa		
$p_{ m amb}$	Ambient pressure	Pa		
$p_{\rm c}$	Chamber pressure, vessel pressure	Pa		
$p_{ m crit}$		Pa		9.4.6
			(continued)

Symbol	Quantity	Unit	Defining equation	Section
	Critical (backing) pressure, fore-vacuum tolerance			
$p_{ m F}$	Fore-line pressure, fore-vacuum pressure	Pa		
$p_{ m in}$	Inlet pressure	Pa		4.1.4
$p_{ m ind}$	Indicated pressure, reading	Pa		
$p_{ m m}$	Motive pressure, pressure of working medium	Pa		
$p_{\rm n}$	Pressure under standard conditions	Pa	3.3	3.1.1
$p_{\rm s}$	Saturation vapor pressure	Pa	3.127	3.5.1
$p_{ m tot}$	Total pressure	Pa		
$p_{ m ult}$	Ultimate pressure	Pa		
$p_{ m v}$	Vapor pressure	Pa		
$p_{ m w}$	Working pressure	Pa		
P	Momentum	kg m/s		
P	Power	W		
P	Transmission probability	1	4.135	4.4.1
P_{Ho}	Ho factor, pumping probability	1		9.4.7
P_{ion}	Differential ionization	cm^{-1}		13.7.3.1
q_N	Particle flow (rate)	s^{-1}	4.9	4.1.2
q_{pV}	Energy flow rate, pV flow, throughput d $(pV)/dt$	Pa ℓ/s	4.10	4.1.2
q_{ν}	Molar flow rate	mol/s	4.15	4.1.2
Q	Charge	C		
Q	Amount ofheat	J		12.2.1
Q_e	Charge ofelectron cloud	С		
Q	Heating power	W		
r	Radius	m		
R	Universal gas constant	$\rm J~mol^{-1}~K^{-1}$		
R	Electrical resistance	Ω		
$R_{\rm s}$	Specific gas constant	$\rm J \ kg^{-1} \ K^{-1}$	3.23	3.1.3
$R_{\rm w}$	Wire resistance	Ω		13.5.2
Re	Reynolds number	1	4.4	4.1.1
RD	Residual drag	s^{-1}	13.16	13.3.2
S	Sticking probability	1	6.3	6.2.1
S_0	Sticking probability on empty surface	1	6.3	6.2.1
$S_{ m W}$	Wall thickness	m		17.4.1.1
S, S _{pump}	Pumping speed	ℓ/s	4.19	4.1.2
S	Sensitivity, gauge constant	Pa ⁻¹	13.33	13.7.3.1
S	Entropy	J/K		

Symbol	Quantity	Unit	Defining equation	
S_A	Surface area-related pumping speed	$\ell \text{ s}^{-1} \text{ cm}^{-2}$	•	,
S_{back}	Loss due to backflow	ℓ/s		7.4.3.4
$S_{\rm c}$	Pumping speed at a chamber or vessel	ℓ/s		
$S_{ m eff}$	Effective pumping speed	ℓ/s		4.1.4
$S_{\rm n}$	Nominal pumping speed	ℓ/s		
$S_{\mathrm{P.c.}}$	Pumping speed of a Penning cell	ℓ/s		11.4.2
t	Time	s		
$t_{ m out}$	Outgassing time	s	6.33	6.3
$t_{ m mono}$	Monolayer time	s	6.25	6.2.6
$t_{ m s}$	Service life	s	11.	11.4.1
$t_{ m sw}$	Switching time	s		20.B
T	Thermodynamic (absolute) temperature	K	3.5	3.1.1
$T_{ m amb}$	Ambient temperature	K		
$T_{ m B.P.}$	Boiling point	K		
$T_{\rm c.s.}$	Temperature of cold surface	K		
$T_{ m D}$	Sutherland's constant	K	3.58	3.2.5
$T_{\rm n}$	Standard temperature	K	3.6	3.1.1
$T_{\rm p}$	Processing temperature	K, °C		16.2.2.1
$T_{\rm s}$	Temperature of dew point	K		8.1
$T_{ m W}$	Wall temperature	K		
и	Relative velocity	m/s		
и	Atomic mass unit, a.m.u.	kg	3.13	
ν	Velocity	m/s		
U	Electrical voltage, potential	V		
$U_{\rm H}$	High voltage	V		
V	Volume	m^3		
\dot{V}	Volume flow rate	ℓ/s		4.1.4
$V_{ m D.S.}$	Dead space, dead volume	m^3		7.2
$V_{ m mol}$	Molar volume	m ³ /mol		
$V_{ m molar,n}$	Molar volume under standard conditions	m ³ /mol	3.24	3.1.3
$V_{ m s,v}$	Specific volume of vapor	m^3/kg		18.2.5
W	Work	N m		12.2.1
x	Distance, length	m		
$x_{ m diff}$	Diffusion length	m	9.15	9.4.7
у	Distance, length	m		
y	Ratio of inherent gas particle volume to total volume	1	3.109	3.4.1
α	Coefficient of linear thermal expansion	K ⁻¹		13.3.2
β	Energy transfer coefficient	1		13.3.2
			(continued)

Symbol	Quantity	Unit	Defining equation	Section
ε	Sensitivity	,		
ϵ	Minimum of Lennard-Jones potential	J		3.4.2
η	Viscosity	Pa s	3.71	3.3.2
κ	Isentropic exponent, ratio ofheat capacities at constant pressure and constant volume	1	4.42	4.2.2
λ	Thermal conductivity (coefficient)	$W m^{-1} K^{-1}$	3.92	3.3.3
$ ilde{\mu}$	Adsorbed amount of gas per unit mass of adsorbent	Pa ℓ/kg	11.2	11.2.1
$\tilde{\mu}_{\mathrm{n}}$	Adsorbed amount of gas per unit mass of adsorbent under standard conditions	Pa ℓ/kg	11.4	11.2.1
ν	Frequency	s^{-1}		
ν	Amount of substance, number of moles	mol	3.8	3.1.2
ν_0	Vibrational frequency ofadsorbed particle	s^{-1}		6.2.2
θ	Surface coverage	1	6.2	6.1
ϑ	Celsius temperature	°C	3.5	3.1.1
ϑ_{C}	Coolant temperature	$^{\circ}$ C		
ρ	Density	kg/m ³		
σ	Ionization cross-section	m^2	13.29	13.7.3.1
σ	Stefan-Boltzmann's constant	$W\ m^{-2}\ K^{-4}$		12.3.4
σ	Surface tension	N/m = Pa m		19.2.2.4
$\sigma_{ m c}$	Probability of condensation	1		3.5.2
$\sigma_{ m eff}$	Effective accommodation coefficient	1	13.19	13.3.2
σ_{t}	Tangential-momentum accommodation coefficient	1	3.63, 13.5	3.3.2, 13.3.2
τ	Period, time interval	s		
τ	Mean time between collisions	s	3.59	3.2.5
$ au_0$	Period of vibrational motion of adsorbed particle	s		6.2.2
χ	Volume percent	1		
χ	Volume collision rate	$m^{-3} s^{-1}$	3.60	3.2.5
ω	Angular frequency	s^{-1}		
$\Lambda_{\mathbf{v}}$	Heat of evaporation	kJ/mol, eV		
Θ	Moment ofinertia	kg m ²		
Ω	Collision integral	1		3.4.2

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