

Vacuum Technology for Particle Accelerators JUAS

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- 1. A quick view of vacuum technology
- 2. The basis of vacuum technology: pressure, conductance, pumping speed
- 3. Calculation of simple pressure profiles
- 4. Gas sources: thermal outgassing and beam-induced desorption
- 5. Gas pumping: momentum-transfer and capture pumps
- 6. Instruments
- 7. Seminar: Example of vacuum system design for particle accelerators (by Roberto Kersevan)



Selected books

Karl Jousten (Ed.), *Handbook of Vacuum Technology Book* (Verlag-VCH, Weinheim, 2008)

James M. Lafferty (Ed.), Foundations of Vacuum Science and Technology (John Wiley and Sons, 1998)

A.Berman, Vacuum Engineering calculation, Formulas, and Solved Exercises (Academic Press, 1992)

P. A. Redhead, J. P. Hobson, E. V. Kornelsen, *The Physical Basis of Ultrahigh Vacuum*, (Chapman and Hall, 1968)

John F. O'Hanlon, A User's Guide to Vacuum Technology

Handbook of Materials and Techniques for Vacuum Devices (AVS Classics in Vacuum Science and Technology)

A. Berman, *Total pressure measurement in vacuum technology*, Academic Press, 1985







Vacuum systems are made of **several components**. The connection between them is obtained by **flanges**.

Flange

Beam pipe

Pump (sputter ion pump)





Beam pipe & NEG coating

A vacuum sector is delimited by **gate valves**.

The pressure is monitored in each vacuum sector by **gauges**.

The beam pipes can be at different temperatures

Pressure measurement

Flange

Pump (sputter ion pump)





Beam pipe & NEG coating

Gauge

Pump (sputter ion pump)

Flange







Example of vacuum components:

- pumping ports
- bellows
- RF shield
- flanges



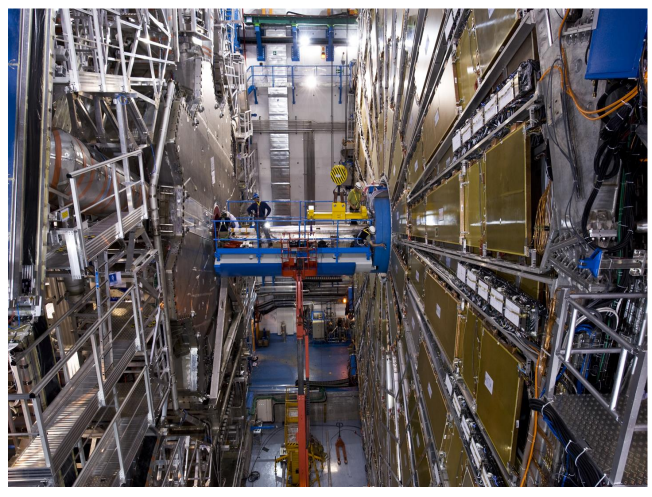




Example of vacuum components:

Copper beam pipes with stainless steel flanges

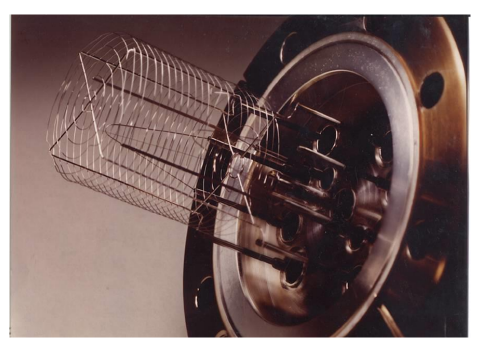




Example of vacuum components:

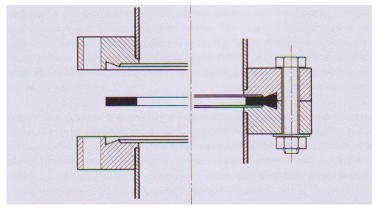
 Beryllium beam pipes in the collision area of the ATLAS experiment (CERN)







In general the leak-tight connection is made by copper gaskets that are squeezed between two stainless steel knives.











In general vacuum systems for particle accelerators are made of metals:

- Austenitic stainless steel
- Copper and its alloys
- Aluminum and its alloys
- Inconel (a Ni alloy)
- Beryllium

In several components the characteristics of the surface is modified by chemical treatment and coating.





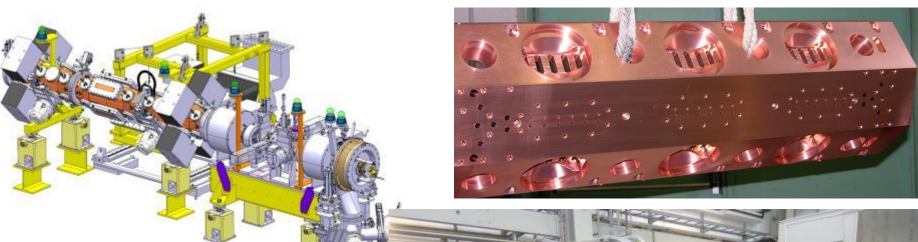
- Electropolishing
- Passivation
- Electroplating
- Thin-film coating by sputtering



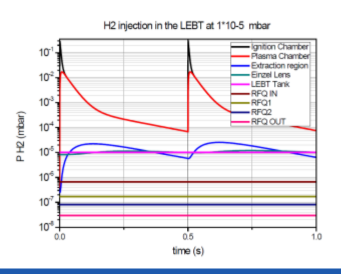
Steps in the production of vacuum systems:

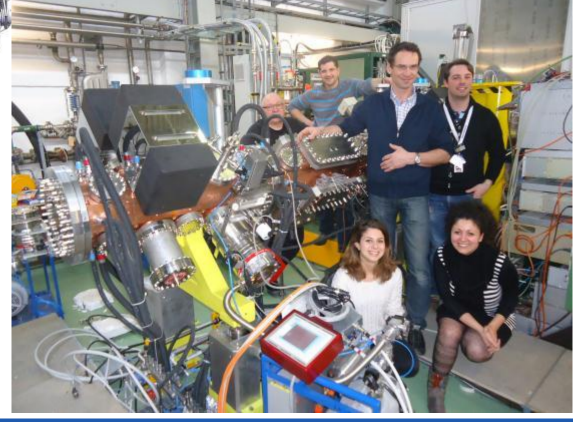
- Identification of requirements
- Choice of materials and treatments
- Design, mechanical evaluation and pressure profile calculation
- Machining & assembly (welding, brazing)
- Surface treatments
- Leak test and global assessment.
- Installation and in situ final check.





Example of vacuum system: The H⁻ source of the Linac4

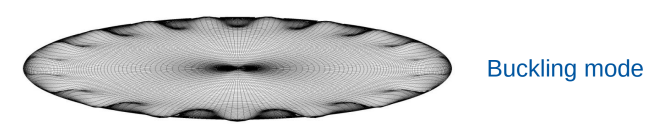


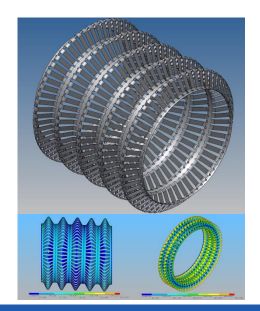


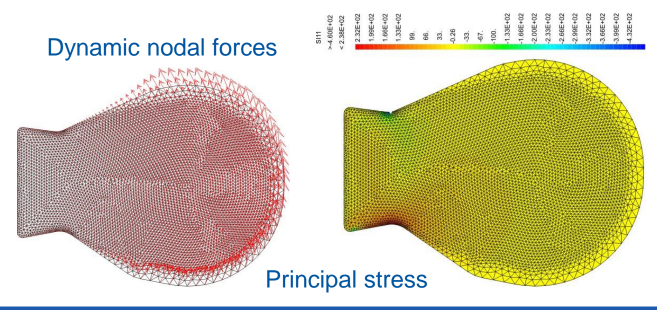


Mechanical engineering for vacuum technology











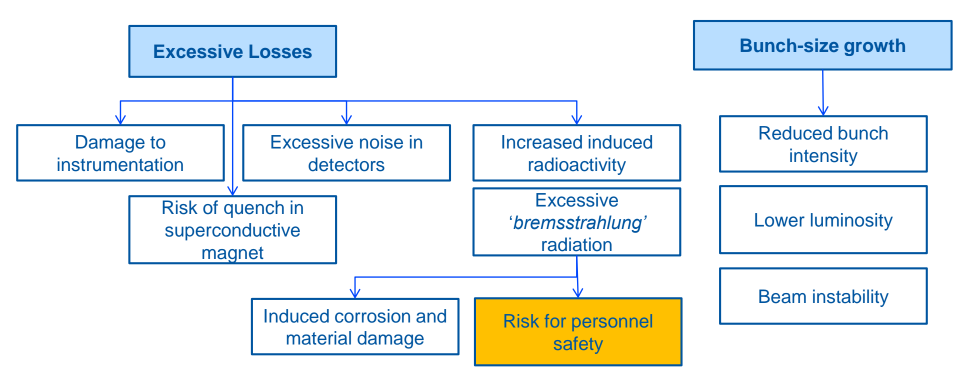
Part 2 The basis of vacuum technology



Why do we need vacuum in accelerators?

Collisions between gas molecules and particles have to be minimized, otherwise:

Particle energy is reduced and trajectories are modified, so that:



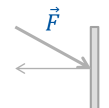
Vacuum is also necessary:

- to avoid electrical discharge in high-voltage (tens di MV/m)...
- To thermically isolate cryogenic devices



The basis of vacuum technology: pressure

Definition of pressure: |Force component in normal direction| Surface area



Unit of measurement: $\frac{[Force]}{[Surface]} \rightarrow \frac{N}{m^2} = Pa \rightarrow 10^5 Pa = 1 \ bar \rightarrow 1 \ atm = 1.013 \ bar$

In vacuum technology: mbar or Pa

Still used in vacuum technology: 1 *Torr* = pressure exerted by a column of 1 mm of Hg; 1 atm = 760 Torr

Conversion Table

	Pa	bar	atm	Torr
1 Pa	1	10 ⁻⁵	9.87 10-6	7.5 10 ⁻³
1 bar	10 ⁵	1	0.987	750.06
1 atm	1.013 10 ⁵	1.013	1	760
1 Torr	133.32	1.33 10 ⁻³	1.32 10 ⁻³	1



The basis of vacuum technology: pressure Degree of Vacuum

	Pressure boundaries [mbar]	Pressure boundaries [Pa]
Low Vacuum LV	1000-1	10 ⁵ -10 ²
Medium Vacuum MV	1-10 ⁻³	10 ² -10 ⁻¹
High Vacuum HV	10 ⁻³ -10 ⁻⁹	10 ⁻¹ -10 ⁻⁷
Ultra High vacuum UHV	10 ⁻⁹ -10 ⁻¹²	10 ⁻⁷ -10 ⁻¹⁰
Extreme Vacuum XHV	<10 ⁻¹²	<10 ⁻¹⁰

Pressures and gas quantities are correlated by the gas equation of state. In vacuum the ideal gas law is always fulfilled:

$$P~V = n~R~T$$
 (thermodynamic) $P~V = N~k_B \mathsf{T}$ (statistical mechanics)

P pressure, **V** volume, **T** temperature, quantity of gas in moles (\mathbf{n}) and number of molecules (\mathbf{N}), **R** gas constant, $\mathbf{k}_{\mathbf{B}}$ Boltzmann constant



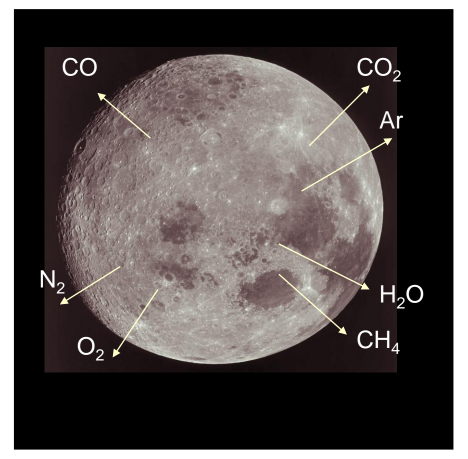
The basis of vacuum technology: gas density

 $\frac{molecules}{cm^3}$

	Pressure [Pa]	Gas Density 293 K	Gas Density 4.3K
Atmosphere	1.013 10 ⁵	2.5 10 ¹⁹	1.7 10 ²¹
Plasma chambers	1	2.5 10 ¹⁴	1.7 10 ¹⁶
LINAC pressure upper limit	10 ⁻⁵	2.5 10 ⁹	1.7 10 ¹¹
Lowest pressure ever measured at room T	10-12	250	1.7 10 ⁴

$$P V = Nk_B T \rightarrow P = \left(\frac{N}{V}\right) k_B T$$
 $k_B = 1.38 \ 10^{-23} \ \frac{Pa \ m^3}{K} = 1.04 \ 10^{-22} \frac{Torr \ l}{K}$





Gas density on the Moon: $10^5 \, \text{cm}^{-3} \, (10^{-10} \, \text{Pa})$ during night and $10^7 \, \text{cm}^{-3} \, (10^{-8} \, \text{Pa})$ during lunar day.



Intergallactical vacuum: 10⁻¹⁷ Pa Vacuum in Via Lattea: 10⁻¹⁵ Pa

Lowest pressure ever measured at room temperature: 10⁻¹² Pa

Lowest air pressure variation perceptible by human ears: 2 10⁻⁵ Pa

-> about 1/10¹⁰ of the atmospheric pressure



The basis of vacuum technology: gas quantity

Gas quantities can be expressed in:

- Number of molecules: N
- Moles: $\frac{N}{N_A} \left(N_A = 6.022 \ 10^{23} \left[\frac{molecules}{mole} \right] \right)$
- PV quantities if the temperature is known and constant : $PV = N(k_BT)$

Example: 1 Pa m³ at 293K contains
$$N = \frac{1 [Pa.m^3]}{1.38.10^{-23} [\frac{J}{K}] 293 [K]} = 2.47 \ 10^{20} \text{molecules}$$

1 Torr I = $3.3 \cdot 10^{19}$ molecules

1 mbar $I = 2.47 \cdot 10^{19}$ molecules

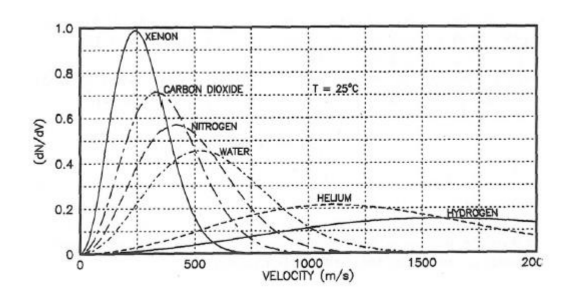


The basis of vacuum technology: gas kinetics

In the kinetic theory of gas the mean speed of a molecule is the mathematical average of the speed distribution:

$$\langle v \rangle = \sqrt{\frac{8 k_B T}{\pi m}} = \sqrt{\frac{8 R T}{\pi M}}$$

m is the molecular mass [Kg] M is the molar mass [Kg]





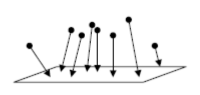
The basis of vacuum technology: gas kinetics

Gas	$\langle v \rangle$ at 293 $K \left[\frac{m}{s} \right]$	$\langle v \rangle$ at 4.3 $K\left[\frac{m}{s}\right]$
H ₂	1761	213
He	1244	151
CH ₄	622	75
N_2	470	57
Ar	394	48



The basis of vacuum technology: impingement rate

$$\varphi = \frac{1}{4} n \langle v \rangle = \frac{1}{4} n \sqrt{\frac{8 k_B T}{\pi m}} \qquad \varphi[cm^{-2}s^{-1}] = 2.635 \ 10^{22} \frac{P \ [mbar]}{\sqrt{M[g]T[K]}}$$



Gas	Pressure [mbar]	Impingement rate 293 K [cm ⁻² s ⁻¹]
	10 ⁻³	1.1 10 ¹⁸
H ₂	10 ⁻⁸	1.1 10 ¹³
	10 ⁻¹⁴	1.1 10 ⁸
N	10 ⁻³	2.9 10 ¹⁷
N_2	10 ⁻⁸	2.9 10 ¹²
A	10 ⁻³	2.4 10 ¹⁷
Ar	10 ⁻⁸	2.4 10 ¹²



The basis of vacuum technology: mean free path

The molecular collision rate ω in a gas is:

$$\omega = \sqrt{2} \, n \, \langle v \rangle \, \sigma_c$$

where σ_c is the collision cross section.

For a single gas, in case of elastic collision of solid spheres:

$$\sigma_c = \pi \delta^2 \rightarrow \omega = \sqrt{2} \pi n \langle v \rangle \delta^2$$

and δ is the molecular diameter.

The mean free path λ , i.e. the average distance travelled by

a molecule between collisions, is given by:

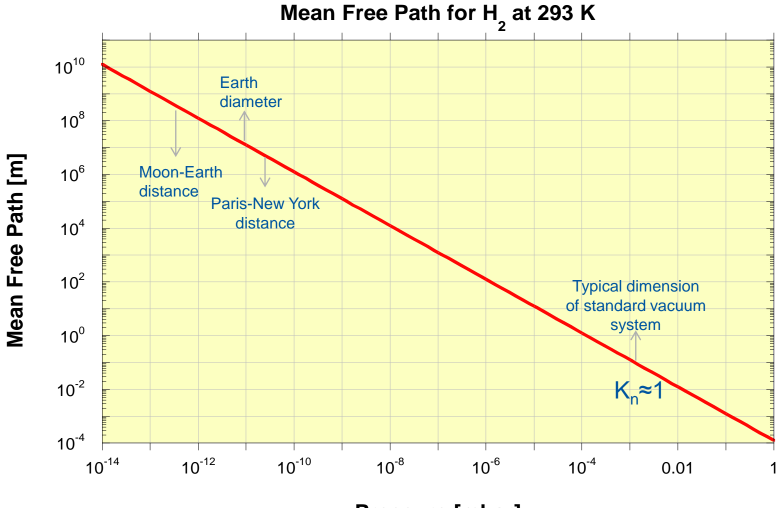
$$\bar{\lambda} = \frac{\langle v \rangle}{\omega} = \frac{1}{\sqrt{2} \pi n \delta^2} = \frac{k_B T}{\sqrt{2} \pi P \delta^2}$$

$$\overline{\lambda_{H_2}}[m] = 4.3 \ 10^{-5} \frac{T[K]}{P[Pa]}$$

Gas	$\sigma_c \left[nm^2 \right]$
H ₂	0.27
He	0.21
N_2	0.43
O ₂	0.40
CO ₂	0.52



The basis of vacuum technology: mean free path







The basis of vacuum technology: mean free path

When the mean free path is of the order of typical dimensions of the vacuum vessel, for example the diameter of cylindrical beam pipes, molecular collisions with the wall of the vacuum envelope become preponderant.

For even longer $\bar{\lambda}$, the gas dynamics is dominated by molecule-wall collisions; intermolecular interactions lose any effect on the gas displacement.



The basis of vacuum technology: Knudsen number

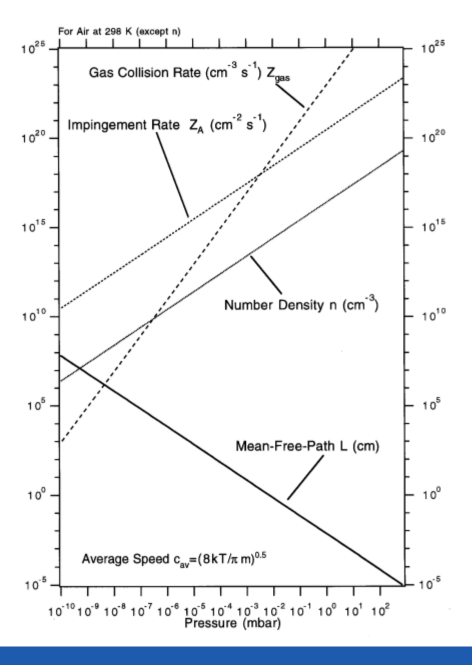
$$K_n = \frac{\bar{\lambda}}{D}$$

 $\bar{\lambda}$ is the mean free path and **D** is a characteristic dimension of a vacuum system (p.ex. the diameter of a beam pipe).

K _n range	Regime	Description
K _n >0.5	Free molecular flow	The gas dynamic is dominated by molecular collisions with the walls of the system
K _n < 0.01	Continuous (viscous) flow	The gas dynamic is dominated by intermolecular collisions
$0.5 < K_n < 0.01$	Transitional flow	Transition between molecular and viscous flow

Except for ion source's plasma chambers, vacuum systems of accelerators operate in free molecular regime; only this vacuum regime is considered in this lesson.







In molecular regime the net flux of molecules from one point (1) to another (2) is proportional to the pressure drop:

$$Q = C (P_1 - P_2)$$

C is called the gas conductance from the two points. It does not depend on pressure.

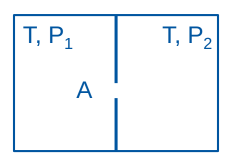
It depends only on the mean **molecular speed** and vacuum system **geometry**.

The unit of conductance is volume per unit time, i.e. **I s⁻¹** or m³ s⁻¹.



The simplest example is the gas flow through a small wall slot of area A and

infinitesimal thickness



$$\varphi = \frac{1}{4}n\langle v \rangle = \frac{1}{4}n\sqrt{\frac{8 k_B T}{\pi m}}$$

Gas flow 1
$$\Rightarrow$$
 2 : $\varphi_{1\rightarrow2}=\frac{1}{4}A\;n_1\langle v\rangle$

Gas flow 2
$$\Rightarrow$$
 1 : $\varphi_{2\rightarrow 1} = \frac{1}{4} A n_2 \langle v \rangle$

Net flow:
$$\frac{1}{4}A$$
 (n_1)

Net flow:
$$\frac{1}{4}A (n_1 - n_2)\langle v \rangle = \frac{1}{4}A \frac{\langle v \rangle}{k_BT} (P_1 - P_2)$$

In PV units
$$(PV = Nk_BT) \rightarrow Q = \frac{1}{4}A\langle v\rangle(P_1 - P_2) \rightarrow C = \frac{1}{4}A\langle v\rangle$$

$$C = \frac{1}{4} A \langle v \rangle \propto \sqrt{\frac{T}{m}}$$



The conductance of the wall slots is **inversely proportional to the square root of the molecular mass**.

Therefore, for equal pressure drop the gas flow of H_2 is the highest.

For gas molecules of different masses, the **conductance scales as the square root of the inverse mass ratio**:

$$\frac{C_1}{C_2} = \sqrt{\frac{m_2}{m_1}}$$



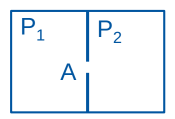
This table collects conductance values, for an orifice, **per** <u>unit surface area</u> (**C**') at room temperature for common gas species.

Conductance of a wall aperture in PV units, per unit area: $C' = \frac{1}{4} \langle v \rangle$

T = 293 K

Gas	$\langle v \rangle$ at 293 $K\left[\frac{m}{s}\right]$	$C'at 293 K \left[\frac{m^3}{s m^2} \right]$	$C' \text{ at 293 } K \left[\frac{l}{s cm^2} \right]$
H ₂	1761	440.25	44
He	1244	311	31.1
CH ₄	622	155.5	15.5
H ₂ O	587	146.7	14.7
N_2	470	11.75	11.75
Ar	394	98.5	9.85





Example: $H_2 P_1 = 5 \cdot 10^{-4} mbar$, $P_2 = 7 \cdot 10^{-5} mbar$, $A = 0.8 cm^2$

$$\rightarrow Q = 44 \times 0.8 \times (5 \ 10^{-4} - 7 \ 10^{-5}) = 1.5 \times 10^{-2} \frac{mbar \ l}{s}$$

$$\rightarrow Q = 1.5 \times 10^{-2} \frac{mbar \, l}{s} \times 2.47 \, 10^{19} \frac{molecules}{mbar \, l} = 3.74 \, 10^{17} \frac{molecules}{s}$$

If the gas was N_2 , the flow would have been $\sqrt{\frac{28}{2}} = 3.7$ times **smaller**



For more complex gas flow restrictions, the transmission probability τ is introduced.



Gas flow 1
$$\Rightarrow$$
 2 : $\varphi_{1\rightarrow 2} = \frac{1}{4}A_1n_1\langle v\rangle \tau_{1\rightarrow 2}$

Gas flow 2
$$\Rightarrow$$
 1: $\varphi_{2\rightarrow 1} = \frac{1}{4}A_2 n_2 \langle v \rangle \tau_{2\rightarrow 1}$

In absence of net flow: $\varphi_{1\to 2} = \varphi_{2\to 1}$ and $P_1 = P_2 \to A_1 \tau_{1\to 2} = A_2 \tau_{2\to 1}$

$$Q = \frac{1}{4} A_1 n_1 \langle v \rangle \ \tau_{1 \to 2} - \frac{1}{4} A_2 \ n_2 \langle v \rangle \ \tau_{2 \to 1} = \frac{1}{4} A_1 \langle v \rangle \ \tau_{1 \to 2} \ \frac{(P_1 - P_2)}{k_B T}$$

In PV units:

$$Q = \frac{1}{4} A_1 \langle v \rangle \, \tau_{1 \to 2} (P_1 - P_2) = C \, (P_1 - P_2)$$
$$C = C' A_1 \, \tau_{1 \to 2}$$

 $C = [conductance\ of\ the\ aperture] \times [molecular\ transmission\ probability]$



 $C = [conductance\ of\ the\ aperture] \times [molecular\ transmission\ probability]$

The molecular transmission probabilities:

- Depend only on the geometry of the vacuum system;
- They are accurately obtained by Monte Carlo methods.
- They can be calculated analytically only for simple geometry;



Example:

• **Tubes** of uniform circular cross section (*L* length, R radius); **Santeler formula** (max error 0.7%):

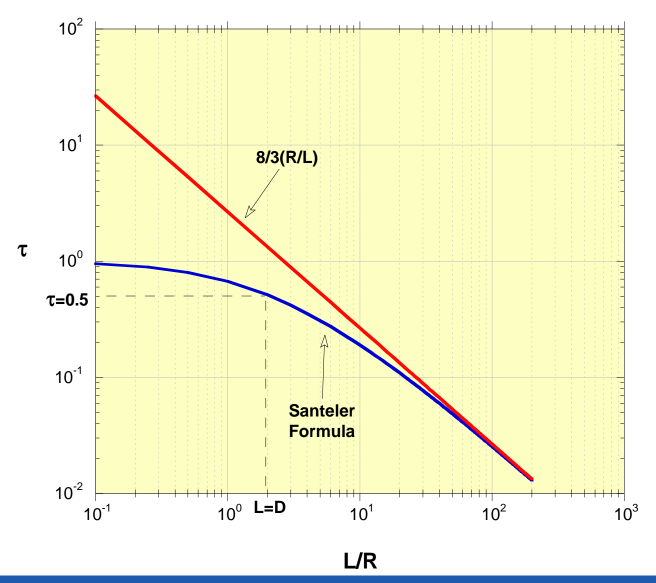
$$\tau = \tau_{1 \to 2} = \tau_{2 \to 1} = \frac{1}{1 + \frac{3L}{8R} \left(1 + \frac{1}{3\left(1 + \frac{L}{7R}\right)} \right)}$$

For long tubes
$$(\frac{L}{R} \gg 1)$$
: $\tau \approx \frac{1}{1 + \frac{3L}{8R}} \approx \frac{8R}{3L}$

For N_2 and $\frac{L}{R} \gg 1 \rightarrow$

$$C = C'A \tau \approx 11.75 \times \frac{\pi D^2}{4} \times \frac{4D}{3L} = 12.3 \frac{D^3}{L} \begin{bmatrix} l \\ s \end{bmatrix}$$
([D] and [l] in cm)





Monte Carlo

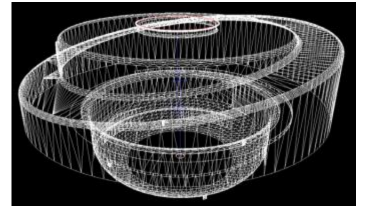
	110 00110
L/R	Cole mean value ^e
0.1	0.952 398 907
0.2	0.909 215 028
0.3	0.869 928 148
0.5	0.801 271 420
0.7	0.743 409 790
0.9	0.694 043 732
1.0	0.671 983 902
1.2	0.632 228 253
1.4	0.597 364 010
1.6	0.566 507 335
1.8	0.538 974 541
2.0	0.514 230 527
3.0	0.420 055 30
4.0	0.356 572 25
5.0	0.310 524 62
6.0	0.275 438 20
7.0	0.247 735 33
8.0	0.225 262 78
9.0	0.206 640 69
10.0	0.190 941 0
20.0	0.109 304
30.0	0.076 912
40.0	0.059 422
50.0	0.048 448
60.0	0.040 913
70.0	0.035 415
80.0	0.031 225
90.0	0.027 925
100.0	0.025 258
200.0	
500.0	
1000.0	0.002 6461

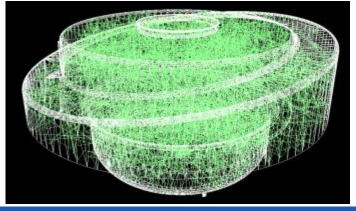


Other analytical expressions for the transmission probability can be found for ducts of circular, rectangular and elliptical cross section (see for example J. M. Lafferty, Foundation of Vacuum Science and Technology, Wiley Interscience).

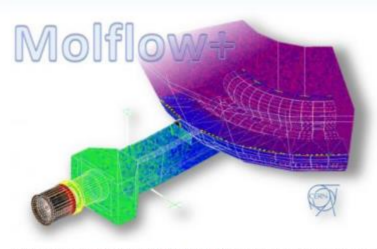
For more complicated geometry, **Test-Particle Monte Carlo methods** (TPMC) are

used.

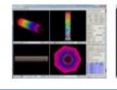


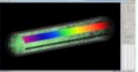
















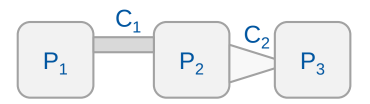
The TPMC:

- generates "random" molecules according to the cosine distribution at the entrance of the tube;
- then follows their traces until they reach the exit of the tube
- each collision with the walls is followed by a random emission
- many simulated molecules are needed in order to reduce the statistical scattering

The software used at CERN is Molflow+, written and maintained by **Roberto Kersevan** and Marton Szakacs (CERN, TE-VSC)



Conductance of components connected in series:



$$Q_{1} = C_{1}(P_{1} - P_{2})$$

$$Q_{2} = C_{2}(P_{2} - P_{3})$$

$$Q_{TOT} = C_{TOT}(P_{1} - P_{3})$$

In stable conditions, there is no gas accumulation in the whole system: $Q_1 = Q_2 = Q_{TOT}$

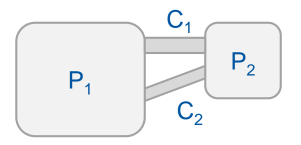
It can be easily verified that: $C_{TOT} = \frac{c_1 c_2}{c_{1+} c_2}$ and $\frac{1}{c_{TOT}} = \frac{1}{c_1} + \frac{1}{c_2}$:

In general for N vacuum components traversed by the same gas flux, i.e. placed **in** series :

$$\frac{1}{C_{TOT}} = \sum_{1}^{N} \frac{1}{C_i}$$



For components connected in **parallel** (same pressures at the extremities):



$$Q_{1} = C_{1}(P_{1} - P_{2})$$

$$Q_{2} = C_{2}(P_{1} - P_{2})$$

$$Q_{TOT} = C_{TOT}(P_{1} - P_{2})$$

$$Q_{TOT} = Q_1 + Q_2 \rightarrow C_{TOT} = C_1 + C_2$$

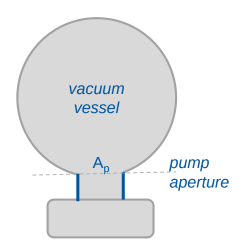
$$C_{TOT} = \sum_{1}^{N} C_{i}$$



In vacuum technology a pump is any 'object' that remove gas molecules from the gas phase.

The pumping speed S of a pump is defined as the **ratio** between the **pump** throughput Q_P (flow of gas definitively removed) and the **pressure P at the** entrance of the pump:

$$S = \frac{Q_P}{P}$$



$$[S] = \frac{[Volume]}{[Time]} = [conductance]$$



The gas removal rate can be written as:

$$Q_P = \frac{1}{4} A_P \, n \langle v \rangle \, \sigma = A_P \, C' n \, \sigma$$

 A_P : is the area of the pump aperture

C': is the conductance of the unit surface area

n: the gas density

 σ : the capture probability, i.e. the probability that a molecule entering the pump is definitively captured

As usual, in term of pressure and PV units:

$$Q_P = A_P C' n \sigma(k_B T) = A_P C' \sigma P$$

From the definition of pumping speed:

$$S = A_P C' \sigma$$

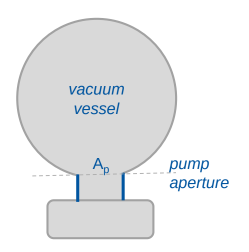


S depends on the conductance of the pump aperture A_PC' and the capture probability σ .

 σ is <u>in general</u> not a constant; it may depend on many parameters including pressure, kind of gas and quantity of gas already pumped.

$$S = A_P C' \sigma$$

The **maximum pumping speed** is obtained for $\sigma = 1$ and is equal to the conductance of the pump aperture.



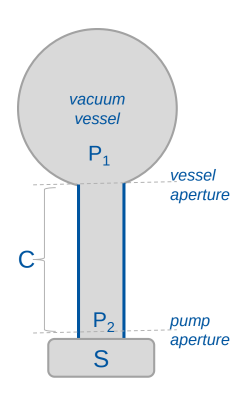
Maximum pumping speed [I s⁻¹]for different circular pump apertures

ID [mm]	H ₂	N ₂	Ar
36	448	120	100
63	1371	367	307
100	3456	924	773
150	7775	2079	1739



A gas flow restriction interposed between a pump and a vacuum vessel reduces the 'useful' pumping speed.

The effective pumping speed S_{eff} seen by the vacuum vessel is easily calculated:



$$Q = C_1(P_1 - P_2) = SP_2 = S_{eff}P_1$$

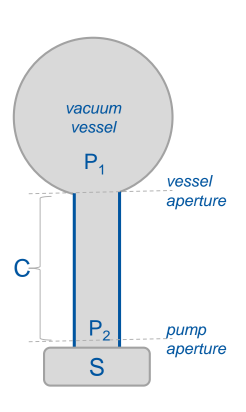
$$\frac{1}{S_{eff}} = \frac{1}{S} + \frac{1}{C}$$

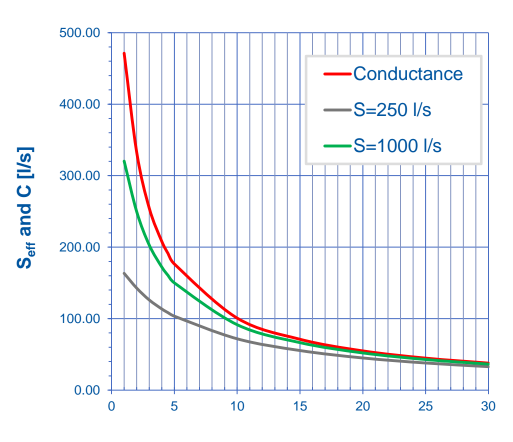
For C<<S: $S_{eff} \approx C$



Example:

Vessel and pump connected by a **10 cm diameter** tube; N_2 , S=250 l/s and 1000 l/s.





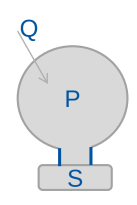




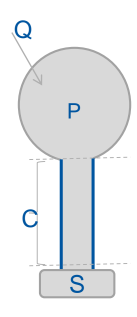


Part 3 Calculation of simple pressure profiles





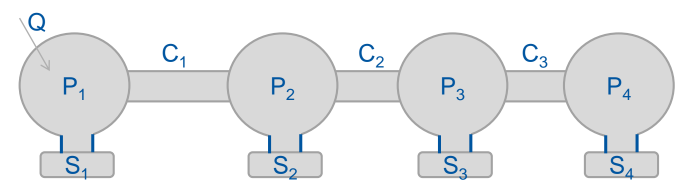
$$P = \frac{Q}{S} + P_0$$



$$P = \frac{Q}{S_{eff}} + P_0 = \frac{Q(C+S)}{C \times S} + P_0$$

P₀ is the pressure attained in the vacuum system without any gas load. It is an intrinsic limitation of the pump.





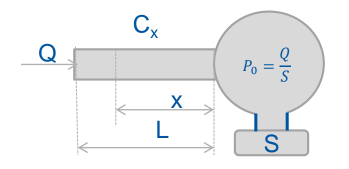
Flux balance at the connexions (node analysis):

$$Q = P_1 S_1 + C_1 (P_1 - P_2)$$

$$C_1 (P_1 - P_2) = C_2 (P_2 - P_3) + P_2 S_2$$

$$C_2 (P_2 - P_3) = C_3 (P_3 - P_4) + P_3 S_3$$

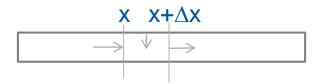
$$C_3 (P_3 - P_4) = P_4 S_4$$



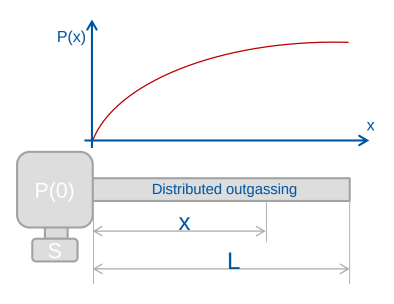
$$P_x = P_0 + \frac{Q}{C(x)}$$
$$C(x) = C(L)\frac{L}{x}$$



Pressure profiles with **distributed gas load** can be calculated analytically (for simple geometry), by electrical analogy or by Monte Carlo simulation.



Distributed gas load, one pump at one extremity only



$$Q(x + \Delta x) - Q(x) = 2\pi R \Delta x \cdot q \to \frac{dQ}{dx} = 2\pi R \cdot q$$

$$Q(x + \Delta x) = -C \frac{L}{\Delta x} \left(P(x + \Delta x) - P(x) \right) = -CL \frac{dP}{dx}$$

$$\to CL \frac{d^2 P}{dx^2} = -2\pi R \cdot q$$

$$P(0) = \frac{2\pi R L \cdot q}{S} = \frac{Q_{TOT}}{S}$$

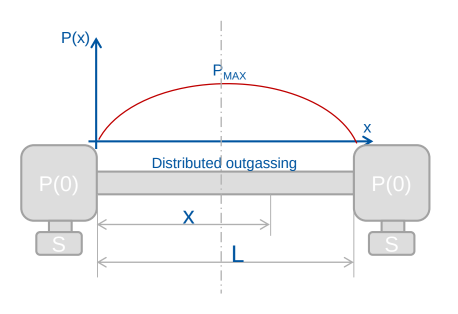
$$\left(\frac{dP}{dx}\right)_{x=L} = 0$$

$$P(x) - P(0) = -\frac{Q_{TOT}}{C} \left[\left(\frac{x}{L}\right) - \frac{1}{2} \left(\frac{x}{L}\right)^2 \right]$$

$$P(L) - P(0) = \frac{Q_{TOT}}{2C}$$



Distributed gas load, one pumps at each extremity



$$CL\frac{d^{2}P}{dx^{2}} = -2\pi R \cdot q$$

$$P(0) = \frac{2\pi RL \cdot q}{2S} = \frac{Q_{TOT}}{2S}$$

$$\left(\frac{dP}{dx}\right)_{x=L/2} = 0$$

$$P(x) - P(0) = -\frac{Q_{TOT}}{2C} \left[\left(\frac{x}{L}\right) - \left(\frac{x}{L}\right)^{2}\right]$$

$$P(\frac{L}{2}) - P(0) = \frac{Q_{TOT}}{8C}$$



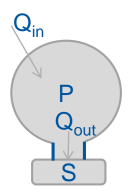
In time dependent condition:

$$Q_{in} \neq Q_{out}$$

Therefore, the variation of the gas density in the system has to be taken into account.

From the ideal gas equation:
$$PV = Nk_BT \rightarrow V\frac{dP}{dt} = k_BT\frac{dN}{dt}$$

A gas balance equation can be written as: $\frac{dN}{dt} = Q_{in} - Q_{out}$

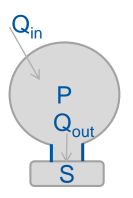


$$V\frac{dP}{dt} = k_B T(Q_{in} - Q_{out});$$

$$V \frac{dP}{dt} = k_B T (Q_{in} - Q_{out});$$
 in PV units: $V \frac{dP}{dt} = (Q_{in} - Q_{out})$

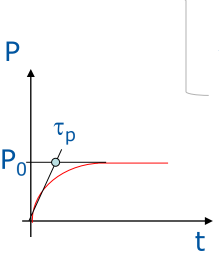
The pumped gas rate is:
$$Q_{out} = SP \rightarrow V \frac{dP}{dt} = Q_{in} - SP$$





$$V\frac{dP}{dt} = Q_{in} - SP$$

$$P(t) = Ae^{-\frac{t}{\tau_p}} + \frac{Q_{in}}{S}$$
 A = integration constant



$$P(0) = P_0 \to P(t) = \left(P_0 - \frac{Q_{in}}{S}\right)e^{-\frac{t}{\tau_p}} + \frac{Q_{in}}{S}$$

$$P(0) = 0 \to P(t) = \frac{Q_{in}}{S} \left(1 - e^{-\frac{t}{\tau_p}} \right)$$

 $\tau_p = \frac{V}{S}$ characteristic time of pumping

 $\tau_{\boldsymbol{p}}$ characterizes all pressure transients in a vacuum system



When Q_{in} is a function of time:

$$P(t) = \frac{\int e^{\frac{t}{\tau_p}} \frac{Q_{in}(t)}{V} dt + A}{e^{\frac{t}{\tau_p}}} = \frac{\int Q_{in}(t) dt}{V} - \frac{e^{-\frac{t}{\tau_p}}}{V\tau_p} \int e^{\frac{t}{\tau_p}} \left[\int Q_{in}(t) dt \right] dt + Ae^{-\frac{t}{\tau_p}}$$

$$A = \text{integration constant}$$

For a network of vacuum chambers, systems of coupled differential equations for each chamber have to be solved.

However, a simpler method exists. It is based on the **analogy between vacuum systems and electrical networks**. Very powerful software is available for the time dependent analysis of electrical networks.

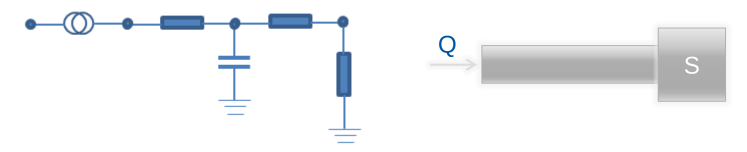


Vacuum element	Electrical elei	ments
Conductance C	Conductance 1/R	o
Gas Flow Q	Current I	
Pressure P	Voltage V	
Volume V	Capacitance C	
Pump	Conductance to ground	
Gas source	Current generator	•——
Constant pressure source	Voltage supply	o——o
Vacuum chamber with conductance and volume		〕

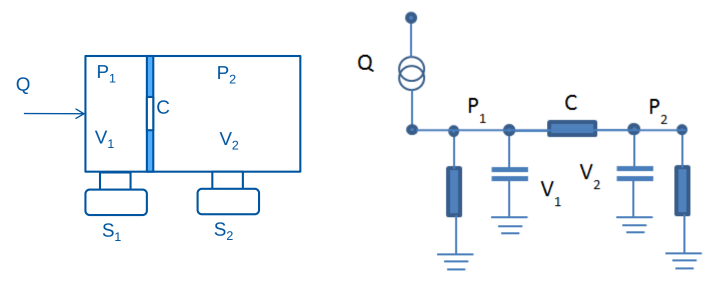
- The ground potential is equivalent to zero pressure.
- Long tubes are subdivided in smaller units and considered as single vacuum chambers (conductance + volume) in series.
 The conductance of a single small unit is equal to the conductance of the entire
 - small unit is equal to the conductance of the entire vacuum chamber times the number of units
- Non-linear electric characteristics can be used to simulate pressure and time dependent conductances and pumping speeds.



Electrical analogy of a vacuum chamber with gas injection and pumping

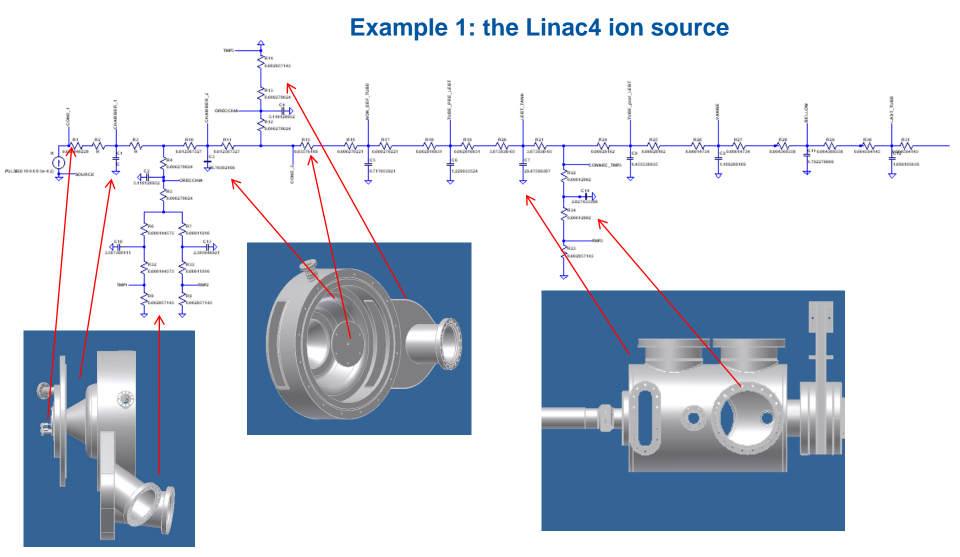


Electrical analogy of a vacuum system with differential pumping



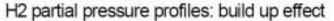
The electrical network is solved by dedicated software, for example LTSpice

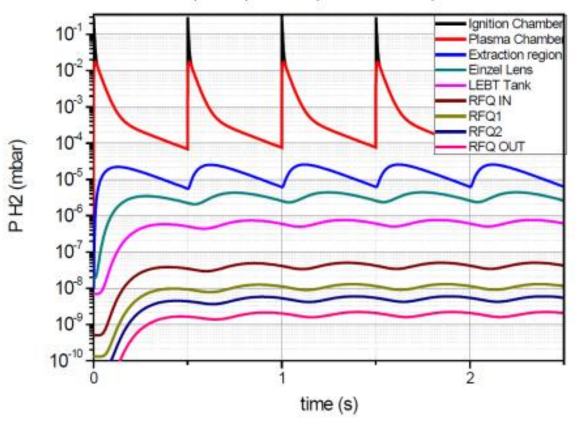






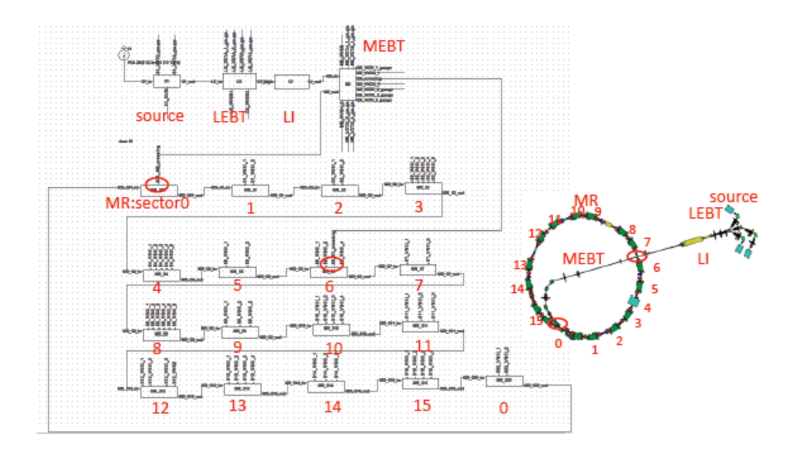
Example 1: the Linac4 ion source





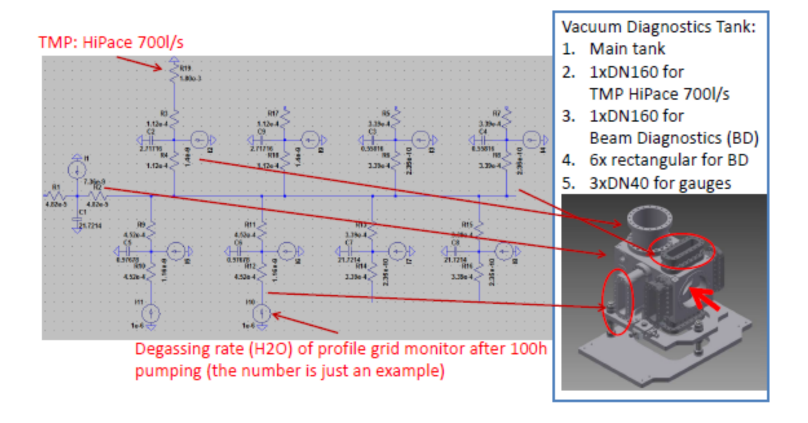


Example 2: the MedAustron facility



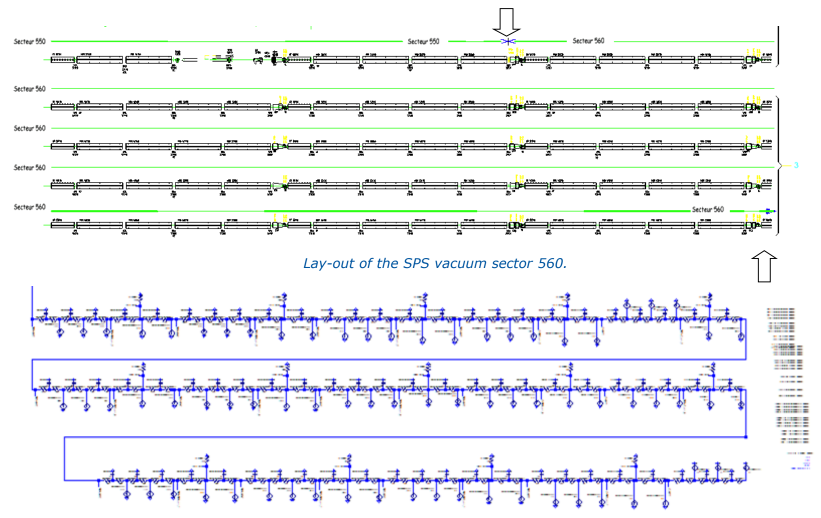


Example 2: the MedAustron facility





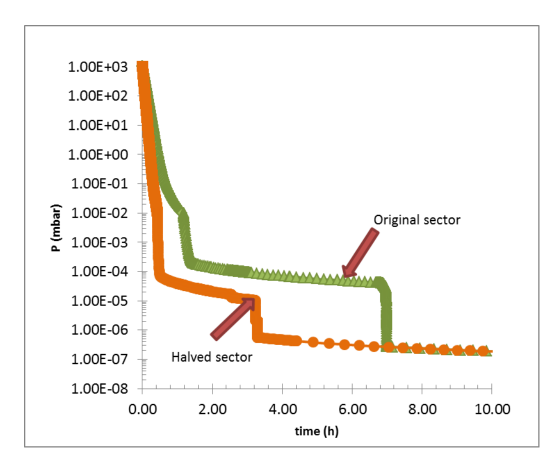
Example 3: pumpdown of one vacuum sector of the SPS







Example 3: pumpdown of one vacuum sector of the SPS



Current dependent resistances simulate conductance variations in the whole pressure range, even in viscous regime.

Comparison between calculated pumpdown curves for original (green) and halved (orange) longest SPS vacuum sector.

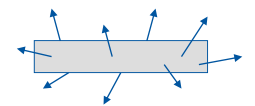


Part 4 Gas sources in particle accelerators

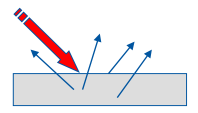


Gas sources

Outgassing is the spontaneous evolution of gas from solid or liquid.



Degassing is the deliberate removal of gas from a solid or a liquid.

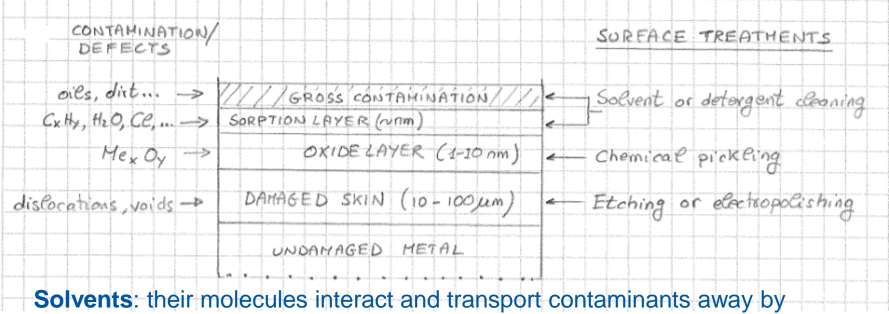


<u>Desorption</u> is the release of adsorbed chemical species from the <u>surface</u> of a solid or liquid.



Gas sources

After production, the **surface** of vacuum components is **always contaminated**. They must be **thoroughly cleaned** before installation.



diffusion (dilution) -> quite selective! (C₂Cl₄, wide spectrum; HFC, more restricted action)

Detergents in water: allows organics and water to combine by forming micelle (surfactant: **surf**ace **act**ing **agent**). Based on molecule with hydrophilic heads and lipophilic tail: less selective than solvents



Gas sources: outgassing

The **bulk of materials** is also a source of gas.

Gas molecules dissolved in the materials diffuse towards the surfaces and are then released

Polymers dissolve **huge quantity** of molecules in the voids between the molecular chains.

Metals can dissolve only **limited quantities** of small atoms that are immobile in the lattice at room temperature, except for hydrogen.

In one day, H atoms travel in average 4 μm in austenitic stainless steels, while O atoms travel the same distance in 1000 years.

Therefore, amongst the dissolved elements in the bulk of metals, **only H₂ is** released at room temperature.

For comparison, in one day, H_2O molecules move along about 20 μm in PEEK, a high performance polymer.



Gas sources: outgassing

Materials	Characteristics that determine outgassing rates	
Polymers	Dissolve entire molecules, for example H ₂ O, O ₂ , CO ₂ , solvents, etc.	
	The solubility in the bulk can be very high in particular for H ₂ O (up to a few weight percentage)	
	The dissolved molecules have a relatively high mobility	
Metals	Only single atoms are dissolved (H, O, C, S, etc.)	
	The atom solubility is in general very low at room temperature (typically up to 10 ³ at. ppm)	
	Only H has a significant mobility at room temperature	



Gas sources: outgassing

Metals

After state-of-art surface cleaning

 If not heated in situ: mainly H₂O for the first months in vacuum, then also H₂.

$$q_{H_2O} \approx \frac{3 \times 10^{-9}}{t[h]} \left[\frac{mbar \ l}{s \ cm^2} \right]$$

The source of H₂O is recharged after each venting to air.

• If heated in situ (baked-out): mainly

H₂. The outgassing rate can be
assumed as constant; it depends
on the accumulated effect of the
previous thermal treatments

Organics (Polymers)

- High solubility of gas in the bulk, in particular H₂O.
- In general, the outgassing process is dominated by H₂O release.
- In the initial phase of pumping:

$$q_{H_2O} \propto \frac{1}{\sqrt{t}}$$

- Heavier gas molecules can be outgassed (remnant of polymerisation, fraction of polymeric chains)
- The permeation of light molecules is not negligible, in particular He.



Gas sources: outgassing



Neoprene (10 h pumping):

 $q_{H2O} \approx 10^{-5} \text{ Torr I s}^{-1} \text{ cm}^{-2}$ $q_{H2O} = 3.3 \times 10^{14} \text{ molecules cm}^{-2}$

Unbaked stainless steel (10 h pumping):

 $q_{H2O} = 2x10^{-10} \text{ Torr } \text{I s}^{-1} \text{ cm}^{-2}$

 q_{H2O} =6.6x10⁹ molecules cm⁻²

Baked stainless steel (150° C x 24 h):

 $q_{H2}=2x10^{-12} \text{ Torr I s}^{-1} \text{ cm}^{-2}$

 q_{H2} =6.6x10⁷ molecules s⁻¹ cm⁻²

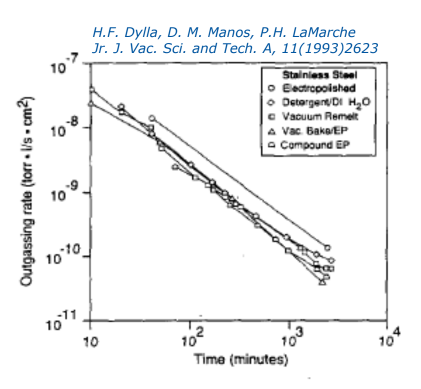
Baked OFS Copper (200° C x 24 h):

 $q_{H2}=2x10^{-14} \text{ Torr } \text{I s}^{-1} \text{ cm}^{-2}$

 q_{H2} =6.6x10⁵ molecules s⁻¹ cm⁻²



Outgassing rate of water vapour for unbaked metals



$$q_{H_2O} \approx \frac{3 \times 10^{-9}}{t[h]} \left[\frac{mbar \ l}{s \ cm^2} \right]$$

The most effective way to accelerate the water vapour outgassing rate is by in situ bakeout.

At least 120°C for 12 hours

Water vapour outgassing rate of stainless steel that underwent four different surface treatments



When water vapour outgassing is strongly reduced, by either long pumping or bakeout, the **outgassing process is led by H₂**.

This gas is dissolved in metals as single H atoms. Its **diffusion is relatively fast** and, after recombination on the surface, it can be released as molecular hydrogen

Most of the **H atoms are dissolved in metals at the liquid state**, during the production process.

H atoms mobility and solubility in the liquid are higher than in the solid.

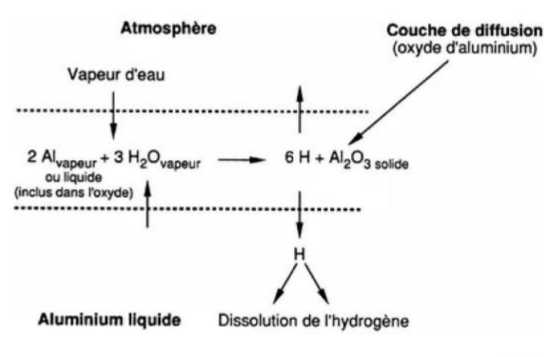
Typical sources of H are:

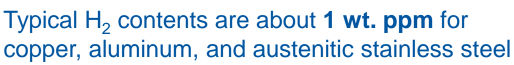
- metals ores;
- tools needed for fusion;
- refractory materials of furnaces;
- combustion and treatment gas;
- water vapour and fluids used for quenching (for example the hyperquench of austenitic stainless steels is carried out from 1100°C in water, air, or oil).

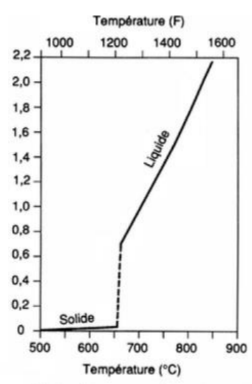


Example of a process of H₂ dissolution in liquid Al

If the solidification is fast, **the gas is trapped in the solid** far form the equilibrium







1 Ncm³ correspond à 1 cm³ de gaz mesuré dans les conditions normales de température (°C) et de pression (10⁵ Pa).



As for water vapour, hydrogen-outgassing rate is **reduced by heating**.

The high temperatures **increase the H atoms mobility** and, as a result, accelerate the depletion of the residual hydrogen content.

However, there is a crucial difference between water vapour and hydrogen.

Each time the vacuum system is exposed to air, water molecules re-adsorb on the surface, while **hydrogen is not recharged** in the bulk of the metal.

For most of the materials used for the manufacturing of vacuum chambers, the H solubility is very low in the solid state.

For example, to recharge **1 wt. ppm** of hydrogen at room temperature in stainless steel, the material has to be in equilibrium with the gas at **7 bar**. The hydrogen pressure in air is roughly 10⁻⁴ mbar, which gives a maximum recharging of about 2.10⁻⁴ wt. ppm.



For **copper and aluminium alloys**, a few bakeout at 150-200°C for 24 hours are sufficient to reduce the hydrogen-outgassing rate to less than 10⁻¹³ mbar I s⁻¹ cm⁻².

For **austenitic stainless steel**, higher temperatures are needed to have a similar effect for a few mm thick vacuum chambers. Repeated bakeout at temperature higher than 200°C may have a significant influence.

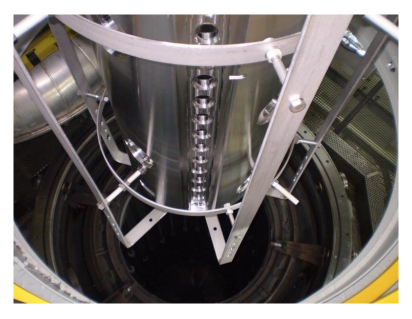
Materials	Bakeout T[°C] x 24 h	q [mbar I s ⁻¹ cm ⁻²]
Austenitic st. steel	150	3 10 ⁻¹²
Austenitic st. steel	200	2 10 ⁻¹²
Austenitic st. steel	300	5 10 ⁻¹³
Copper Silver added (OFS)	150	3 10 ⁻¹²
Copper Silver added (OFS)	200	≈ 10 ⁻¹⁴
Beryllium	150	< 10 ⁻¹⁴
Al alloys	150	< 10 ⁻¹³



For austenitic stainless steels, a radical effect is obtained by heating in a vacuum furnace to temperature up to about 1000°C.

Such a treatment is called 'vacuum firing'. At CERN it is carried out at 950°C for 2 h in a vacuum better than 10⁻⁵ mbar at the highest temperature.

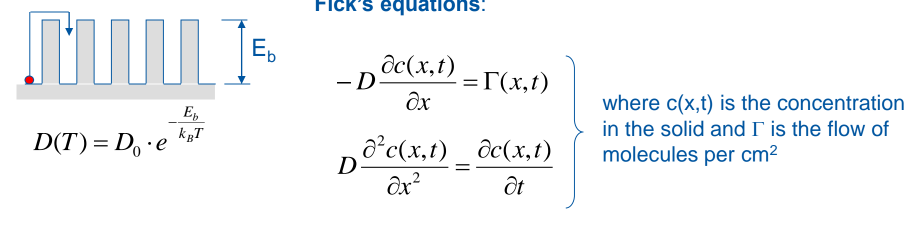




The **CERN's large furnace**: useful height and diameter: 6 m and 1 m, respectively. Maximum charge weight: 1000 Kg. Ultimate pressure: about 10⁻⁷ mbar; pressure at the end of the 950°Cx2h treatments: 10⁻⁵mbar.



Diffusion model of H₂ outgassing



Diffusion, in most of the cases of interest, is described by the Fick's equations:

$$-D\frac{\partial c(x,t)}{\partial x} = \Gamma(x,t)$$

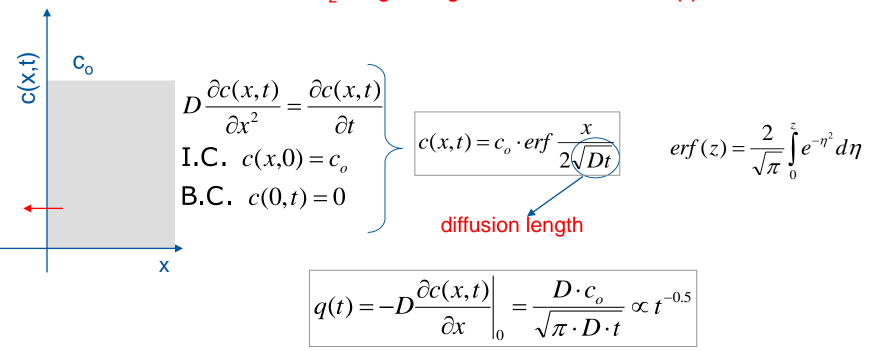
$$D\frac{\partial^2 c(x,t)}{\partial x^2} = \frac{\partial c(x,t)}{\partial t}$$

In the limit of this model, the outgassing rate is equal to a half of the flux of atoms arriving at the surface by diffusion (2 H atoms = $1 H_2$ molecule):

$$q(t) = -\frac{1}{2} D \frac{\partial c(x,t)}{\partial x} \bigg|_{x = SURF.}$$



Diffusion model of H₂ outgassing: semi-infinite solid approximation



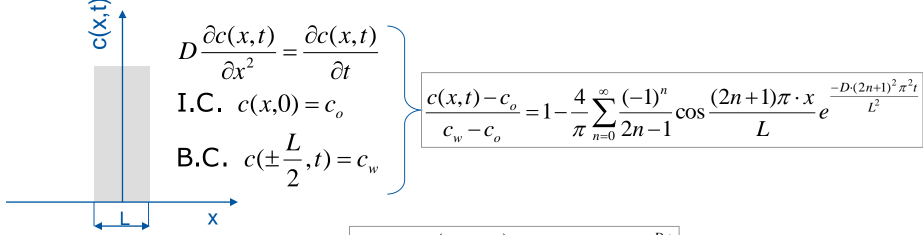
The $t^{-0.5}$ evolution holds also for solid of finite dimension L (slab) when L>>(Dt)^{0.5}

The total amount of substance M(t) which has left the solid is:

$$M(t) = \frac{2c_o}{\sqrt{\pi}} \cdot \sqrt{D \cdot t}$$



Diffusion model of H₂ outgassing: slab approximation



$$q(t) = \frac{4 \cdot (c_0 - c_w) \cdot D}{L} \sum_{n=0}^{\infty} e^{-(2n+1)^2 \pi^2 \cdot \frac{D \cdot t}{L^2}}$$

For Dt>0.05 L² only the first term of the series is relevant :

$$q(t) \approx \frac{4 \cdot (c_0 - c_w) \cdot D}{L} e^{-\pi^2 \cdot \frac{D \cdot t}{L^2}}$$



Diffusion model of H₂ outgassing: Sievert's law

If hydrogen in the gas phase and on the surface are in equilibrium, the surface concentration c_w may be calculated by the Sievert's law:

$$c_w = K_s(T) \sqrt{P_{H_2}} \qquad \begin{array}{l} \mathbf{P_{H2}} \text{ is the pressure of hydrogen in the gas phase} \\ \mathbf{K_s(T)} \text{ is the hydrogen solubility in the metal} \\ K_s(T) = K_0 e^{\frac{-E_s}{2k_BT}} \qquad \mathbf{K_s} \text{ is the H}_2 \text{ solubility energy} \\ \mathbf{K_0} \text{ a constant.} \end{array}$$

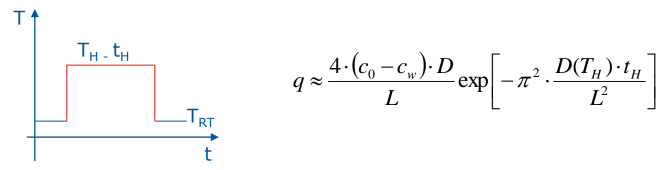
For stainless steel, E_s is positive; therefore, the solubility increases with temperature. At **room temperature** and for typical hydrogen pressure, $\mathbf{c_w}$ is **negligible** with respect to the residual hydrogen content. This is not true at higher temperature; for example during a vacuum firing treatment.

$$c_{H}[at.ppm] = 71.8 \cdot \sqrt{P_{H_{2}}[Torr]} \cdot e^{\frac{-0.114[eV]}{k_{B} \cdot T}}$$
 Sievert's law for austenitic stainless
$$c\left[\frac{Torr\ l\ (H_{2})}{cm^{3}}\right] = 8.21 \cdot 10^{-2} \sqrt{P_{H_{2}}[Torr]} \cdot e^{\frac{-2650}{1.99T[K]}}$$
 steel



Diffusion model of H₂ outgassing: Fourier number

If the slab is heated to a temperature T_H for a time t_H , the outgassing rate, when back at room temperature is:



$$q \approx \frac{4 \cdot (c_0 - c_w) \cdot D}{L} \exp \left[-\pi^2 \cdot \frac{D(T_H) \cdot t_H}{L^2} \right]$$

For an arbitrary thermal cycle T=T(t), it can be show that:

$$q \approx \frac{4 \cdot (c_0 - c_w) \cdot D}{L} \exp \left[-\pi^2 \cdot \frac{\int_0^{t_H} D(T) \cdot dt}{L^2} \right] = \frac{4 \cdot (c_0 - c_w) \cdot D}{L} \exp \left[-\pi^2 \cdot F_0 \right]$$

$$\int\limits_{t_H}^{t_H} D(T) \cdot dt$$
 The dimensionless number:
$$F_o = \frac{0}{I_c^2}$$
 is called the Fourier number



Diffusion model of H₂ outgassing: consecutive bakeouts

F_o 'records' the thermal history of the material and determines how much of the initial concentration is depleted.

When thermal treatment of $F_o \approx 3$ are applied, the solid is actually emptied or in equilibrium with the surrounding gas phase.

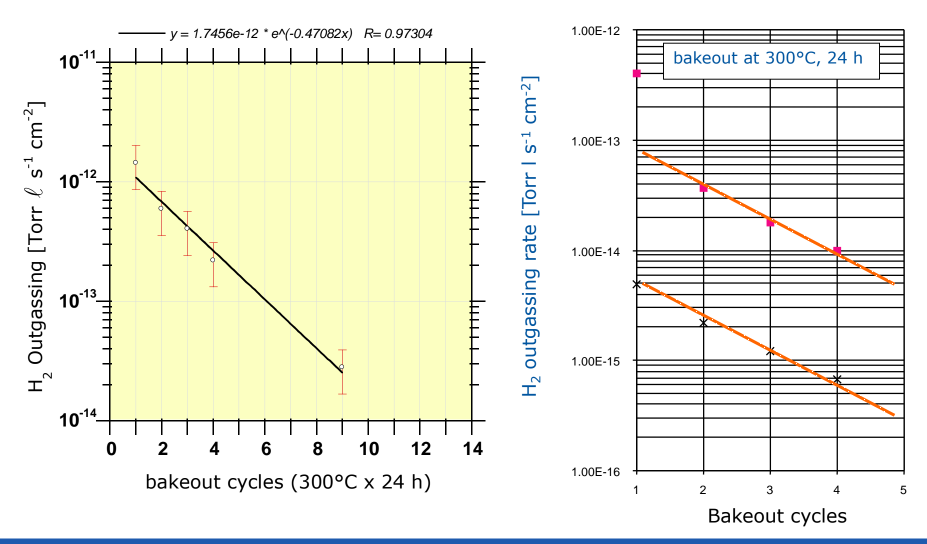
In case of **multiple identical thermal treatments** (for example bakeouts) with a given constant F_0 , the calculated ratio of the outgassing rate at room temperature after two successive thermal cycles is:

$$\frac{q_{n+1}}{q_n} = exp[-\pi^2 F_0]$$

That is to say that each bakeout reduces the outgassing rete of hydrogen by the same constant factor. This is in line with the experimental results.

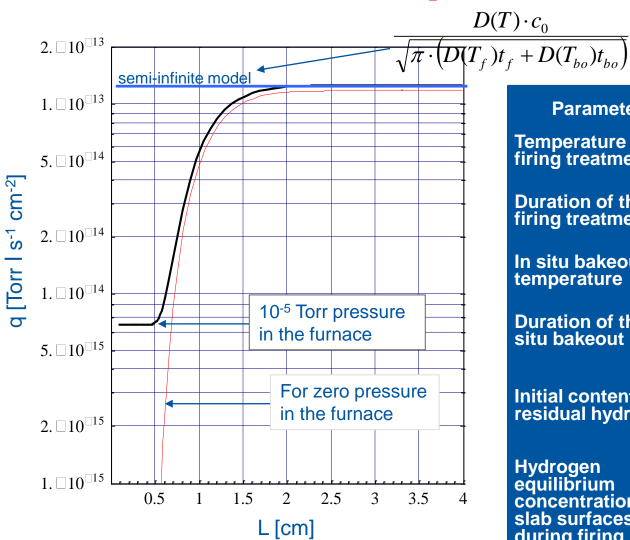


Diffusion model of H₂ outgassing: consecutive bakeouts





Diffusion model of H₂ outgassing: vacuum firing



Parameters	Symbol	Values
Temperature of the firing treatment	T _f	950°C
Duration of the firing treatment	t _f	2 hours
In situ bakeout temperature	T_bo	150°C
Duration of the in situ bakeout	t _{bo}	24 hours
Initial content of residual hydrogen	C _o	1 ppm wt (≈50 ppm at.)
Hydrogen equilibrium concentration on slab surfaces during firing	C _w	0.06 ppm wt. Equivalent to P _{H2} =1.3x10 ⁻⁵ mbar



Polymers, in particular their amorphous structures, can **dissolve huge quantities of gas**.

The water vapour solubility is very high; for example for common materials like Viton, PEEK, and Vespel the content of water in equilibrium with 50% humidity air at 20°C is 0.21, 0.2, and 1 wt. %, respectively.

The huge quantity of dissolved gas and the relatively high mobility through the polymeric chains result in much higher outgassing rates than the ones for metals. This is particularly marked for **water vapour**.

Polymer	Unbaked, 1 h pumping	Baked, ultimate
		A A A A A A A A A A A A A A A A A A A
Fluoroelastomer	$4 \times 10^{-7} \cdot 2 \times 10^{-5}$	$3 \times 10^{-11} - 2 \times 10^{-9}$
Buna-N	$2 \times 10^{-7} - 3 \times 10^{-6}$	
Neoprene	$5 \times 10^{-5} - 3 \times 10^{-4}$	
Butyl	$2 \times 10^{-6} - 1 \times 10^{-5}$	
Polyurethane	5×10^{-7}	PARTITION
Silicone	$3 \times 10^{-6} - 2 \times 10^{-5}$	_
Perfluoroelastomer	3×10^{-9}	$3 \times 10^{-11} - 3 \times 10^{-10}$
Γeflon	$2 \times 10^{-8} - 4 \times 10^{-6}$	NATIONAL PROPERTY OF THE PROPE
KEL-F	4×10^{-8}	3.5×10^{-10}
Polyimide	8×10^{-7}	3×10^{-11}

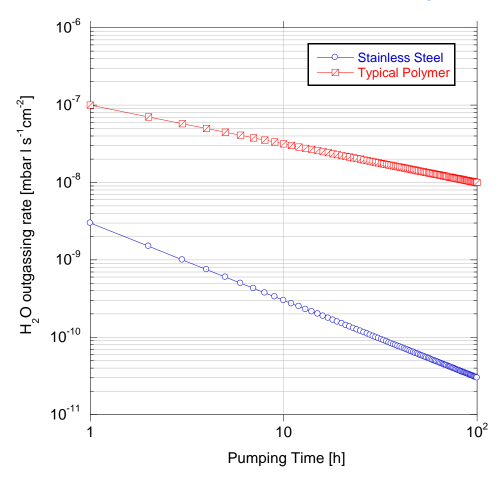
The maximum bakeout temperature depend on the type of polymer; it is limited to about 200 °C for Viton®.



The outgassing rates of thick (L>Dt) slab of polymers, as expected by the diffusion model, decrease with the inverse of the square root of the pumping time t:

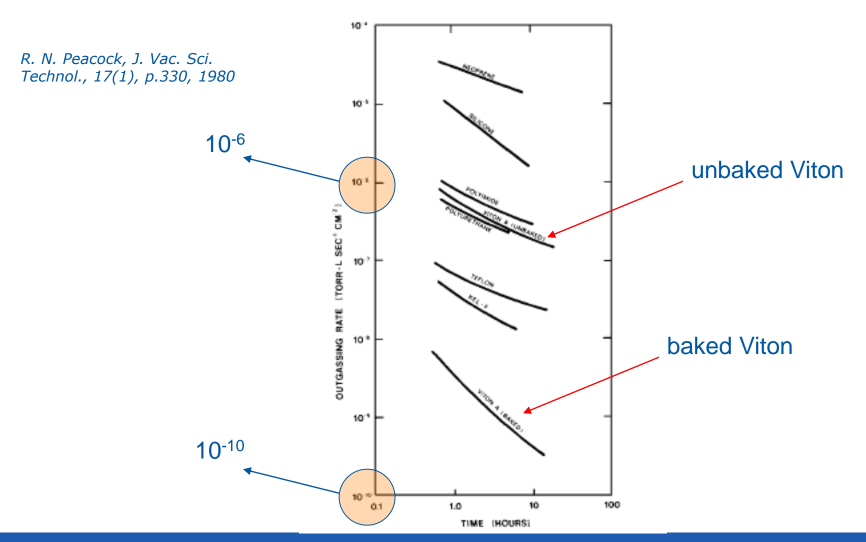
$$q_{H_2O} \propto \frac{1}{\sqrt{t}}$$

Compared to unbaked metals, thick polymers experience a much slower reduction of water outgassing rate.





Outgassing rate of water vapour for polymers





- Another important limitation of polymers used as seals is the high gas permeability.
- Gas penetrates into the material and diffuses towards the vacuum system. The
 permeation flow may limit the ultimate water vapour pressure in vacuum
 systems and affect the sensitivity of helium leak-detection.
- As an example, the permeation flow of atmospheric water through a Viton O-ring, 5 mm cross section diameter, 6 cm torus diameter, is about 10⁻⁷ mbar I s⁻¹
 This value is equivalent to the outgassing of 1 m² of stainless steel after 100 h of pumping.



Application of polymers in UHV technology is hindered by the **huge outgassing** and **permeability**.

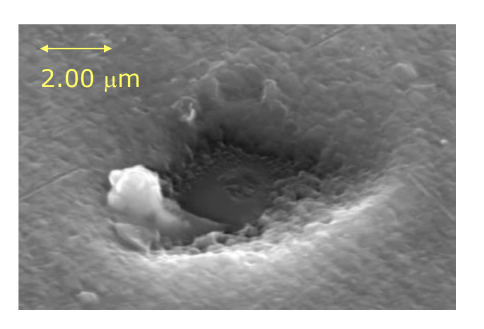
This drawback has been partially overcome in other technological domains (i.e. packaging) by **coating polymers with metals** or metal oxides (more recently also with a:C-H).

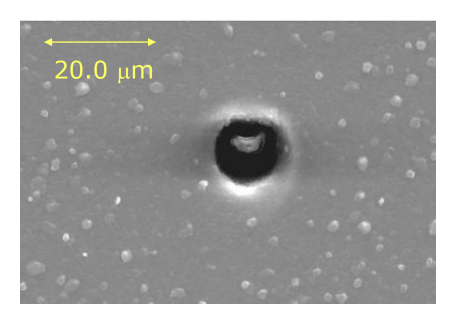
How effective are coatings in reducing outgassing and permeation rate?

- Since the permeability of metals is negligible for all gases, metallic coatings should entirely block the polymer outgassing and permeation.
- ➤ However, experimental results show that only a partial reduction of the flux is attained.
- This is attributed to defects on the coating (pinholes or scratches) that cause discontinuity on the surface coverage.
- ➤ Pinholes are produced during the deposition process and they are presumably due to atmospheric dust particles.



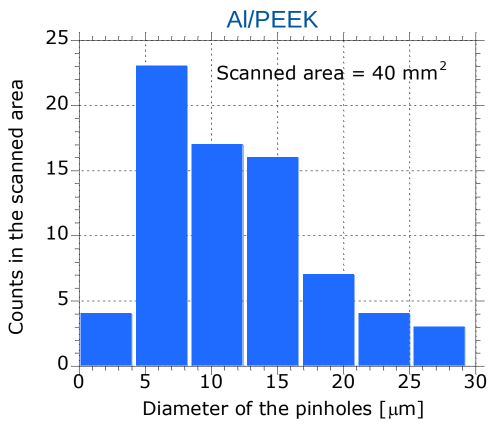
Examples of pinholes on Al coating deposited on PEEK







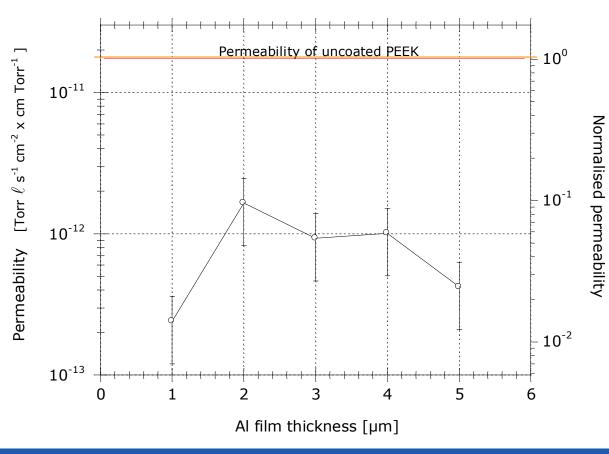
Transmitted light optical microscopy has been used to detect uncoated surfaces on metal films, showing that they mainly consist of pinholes of different diameters.



Normalised uncoated surface areas of the order of 10⁻⁴ are measured



However, experiments have shown that the permeability is **reduced much less than expected** from the uncoated fraction

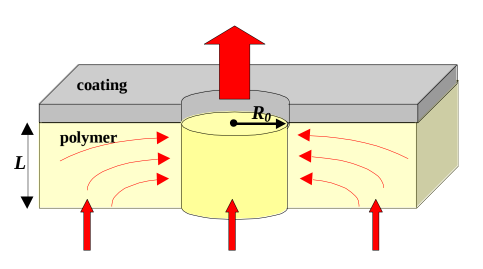


Example: Al/PEEK(0.125 mm thick) The uncoated fraction is **10**-4, but the permeability of He is **10**-2 of that of the uncoated polymer.



This apparent inconsistency can be justified considering that the pinhole gas throughput is enhanced by lateral diffusion.

- ρ =Normalised permeability \rightarrow order of 10⁻²
- Θ =Normalised uncoated area \rightarrow order of 10⁻⁴



For $L/R_0 > 0.3$:

$$\rho = \Theta \times \left(1 + 1.18 \frac{L}{R_0}\right)^*$$
Amplification factor



After W.Prins and J.J.Hermans, J.Phys.Chem., 63 (1959) 716.

In the literature the barrier efficiency is called "barrier improvement factor" or BIF. It is the inverse of normalized permeability.

Normalized permeability:

$$\rho = \Theta \times \left(1 + 1.18 \frac{L}{R_0}\right)$$

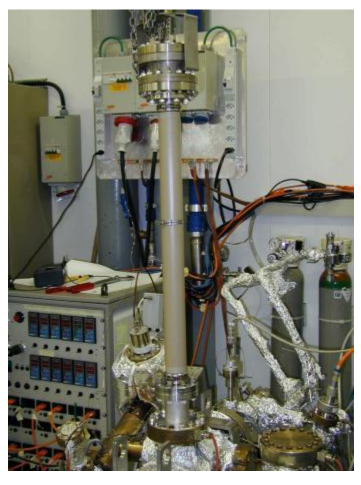
Barrier improvement factor:

$$BIF = \frac{1}{\rho} = \frac{1}{\Theta \times \left(1 + 1.18 \frac{L}{R_0}\right)}$$

The fact that ρ depends on L means that for L>10R the flux value does not depend on the polymer thickness. In other terms, all the concentration gradient is localized near the pin-hole.

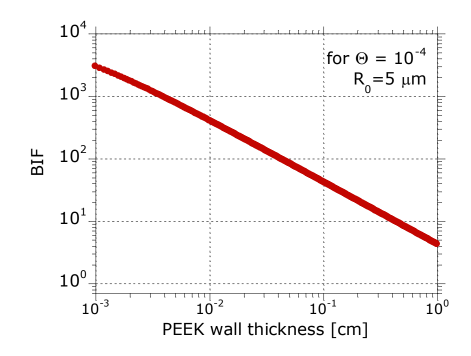
It follows also that, for similar coating, the improvement due to the coating is less significant for thicker polymer substrates.





The BIF for AI coated PEEK.

For thickness larger than 5 mm, the coating is useless.

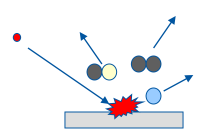


In conclusion, metallic coatings are very efficient in the reduction of the permeation and outgassing rate in thin polymers. For polymers thicker than 5 mm, the benefit of the coating is negligible.



Gas sources: beam induced desorption

The impingement of electrons and photons, of energy higher than some eV, on solid surfaces can result in ions and neutral molecules desorption.



The first observations of induced desorption are very old and they go back to the origin of modern vacuum technology.

 In 1918, Dempster observed ion desorption from electron bombarded salts (Phys. Rev. 11, 323)

ESD: electron stimulated desorption

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POSITIVE RAY ANALYSIS.

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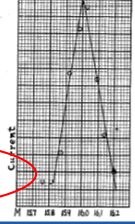
Positive Ions from Electron Bombardment.

It was thought that the bombardment of salts by electrons might break up the chemical compounds and give rise to many positive ions. At first a Wehnelt cathode was used; the ions formed passed beside the cathode (Fig. 1) and were then accelerated by a large potential difference. Aluminium phosphate on a piece of platinum foil was first bombarded. The intensity of the rays increased very rapidly with a slight increase in the amount or energy of the bombarding electrons, indicating that the salt needs to be heated to a certain degree before the ions are separated. Although the aluminium phosphate was chemically pure, the rays ob-

tained under the bombardment of 128 volt electrons were very complex; the following ions were observed besides a couple of unresolved groups; H_1 , H_2 , Li (weak), O_1 (strong), Na (strong), O_3 (?) (weak), M = 62 (weak, possibly Na_2O), M = 67 (strong, possibly $H_3PO_2 = 66$), M = 76 (strong), M = 86 (weak, possibly Rb = 85.5), M = 112 (strong, possibly $P_2O_3 = 110$).

The experiments indicated the convenience of the method of obtaining positive rays and opened up an interesting field for investigation.

The experiments were however first directed





Gas sources: beam induced desorption

Millikan reported the first evidence of **photon induced desorption** in 1909 during the measurement of the photoelectric current of metals exposed to ultraviolet radiation. The first interpretation is given by Winch in 1930 (Phys. Rev. **36**, 601).

Winch was the first to see the implication of photoelectrons on photon induced desorption.

PSD: photon stimulated desorption



While attempting to study the photoelectric emission from the two sides of an unbacked film of gold, 2×10^{-6} cm thick, it was found that the emission did not hold constant but increased with exposure to ultraviolet. With 360 hours of exposure of the film to ultraviolet the photo-current from the side where the light was incident increased 136 fold and at the same time the long wave limit shifted from 2000A to 2537A.

The emission from the unexposed face increased, in the first 50 hours, about one tenth as much as from the exposed face. It was found that the film was transmitting down to a wave-length of 2259A but only about 10 percent of the energy in the lines below 2537A was being transmitted. On exposure of the second face to the direct radiation of the arc its emission increased as in the case of the first face.

It was found that with the film charged positively, so that the electrons returned to the film after being ejected, the increase in emission was more rapid and the long wave limit shift was greater than when the film was charged negatively. In either case, when the film was exposed through a filter which absorbed all wave-lengths below 2800A but transmitted 55 percent of the total energy of the arc, there was no increase in emission.

A plate of solid gold was treated in like fashion and showed a similar increase in photo-current, which took place much more slowly, accompanied by a shift in long wave limit from about 2000A to 2482A.

At intervals during the exposure of the

specimen to ultraviolet fatigue curves, taken by leaving the specimen in a vacuum of 10⁻⁷ mm of Hg unexposed, showed during the first stages a rapid decrease in photocurrent with time of standing, but, after 360 hours of exposure for the film and 160 hours for the solid gold, the photo-current from the former held constant for 3 hours, and from the latter 1½ hours. This seemed to indicate that a fairly stable equilibrium had been reached, and the subsequent fatigue was consistent with the idea that it was due to return of gas to the surface.

The experiment was repeated, using a silver filament approximately 0.025 mm thick, and an increase in emission comparable to that for the gold film was obtained.

The probable explanation is that photoclectrons, both when ejected and returned to the surface by a reverse field, remove adsorbed gas from the surface.

Millikan¹ noted an increase in photoelectric emission on exposure of certain metals to ultraviolet, but did not note the corresponding change in long wave-length limit or that the photoelectrons themselves apparently play an important part in the outgassing.

Work is being carried forward testing this explanation and obtaining more data on photoelectric properties of thin films.

RALPH P. WINCH

Laboratory of Physics, University of Wisconsin, Madison, Wisconsin, July 15, 1930.

Millikan, Phys. Rev. 29,185 (1909).



Gas sources: beam induced desorption

The **desorption yield** η , i.e. the number of molecules desorbed per impinging particle, is needed to design the vacuum system of particle accelerators.

$$\eta = \frac{\text{number of molecules desorbed}}{\text{number of particules impinging on the surface}}$$

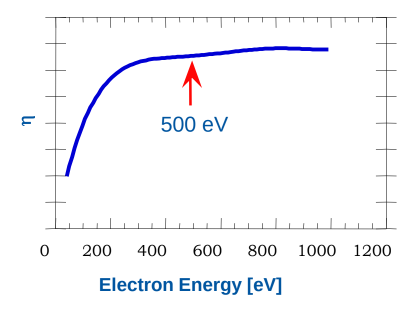
η depends on many parameters, in particular:

- on the nature and energy of the impinging particle;
- the material of the vacuum chamber;
- the nature of the desorbed gas;
- quantity of particles that have already impinged on the surface, namely the dose D [particles/cm²].

The cleanliness of the surfaces has also a crucial influence.



Typical values of η for baked structural vacuum materials vary between 10⁻¹ and 10⁻³ molecules per electron at 500 eV. Neutral desorption is much more likely than ion desorption, about 100 times higher.



For typical metals used in vacuum technology, η of hydrogen is the highest followed by those for CO, CO₂ and CH₄, in decreasing order.

$$\eta_{H_2} > \eta_{CO} > \eta_{CO_2} > \eta_{CH_4}$$



Electron bombardment cleans metallic surfaces by gas desorption.

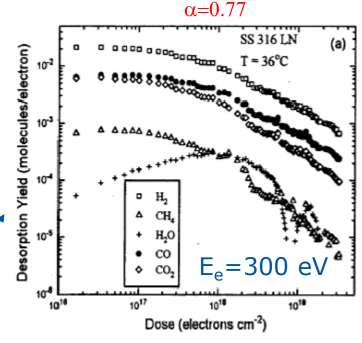
As a consequence, the desorption yields decrease when the electron dose increases.

 η_e values have a power law dependence on the dose of electrons D_e (for doses higher than 10^{18} e⁻ cm⁻²):

$$\eta = \eta_o \cdot D^{-\alpha}$$

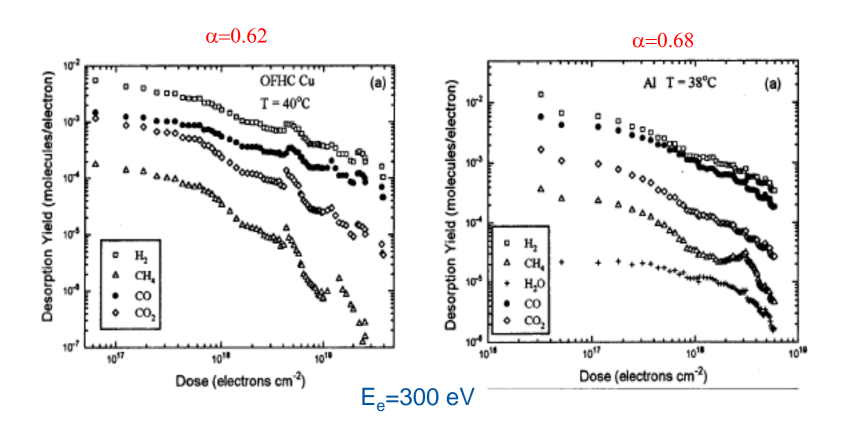
where α varies between 0.6 and 1.

- J. Gomez-Goñi and A. G. Mathewson
- J. Vac. Sci. Technol. A 15, 3093 (1997)



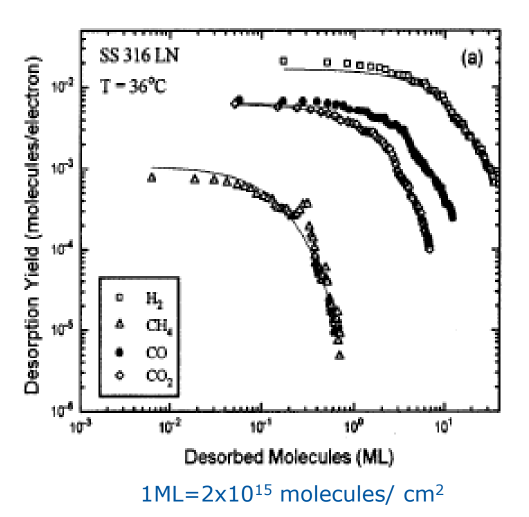
The samples were baked at 150 C for 24 h and at 300 C for 2 h





J. Gomez-Goñi and A. G. Mathewson J. Vac. Sci. Technol. A 15, 3093 (1997)





The total quantity of gas desorbed exceed by far one monolayer.

In order to explain the huge quantity of gas desorbed, some authors considered stimulated gas diffusion from the oxide layer

The accumulated dose effect is lost when the surface is vented to air for long time. For short exposure to air (for example a few hours) a residual effect is measured; it is called 'memory effect': η values similar to those attained before venting are achieved after a smaller dose.

J. Gomez-Goñi and A. G. Mathewson J. Vac. Sci. Technol. A 15, 3093 (1997)



The ESD process can be considered as an **isolated interaction between an electron and an adsorbate**.

The maximum energy ΔE transferred during the collision between a low energy electron of mass m_e and a molecule of mass M is calculated by classical kinematics. For hard-sphere scattering and m_e <<M, one can estimate that:

$$\frac{\Delta E}{E_e} \cong \frac{4m_e}{M}$$

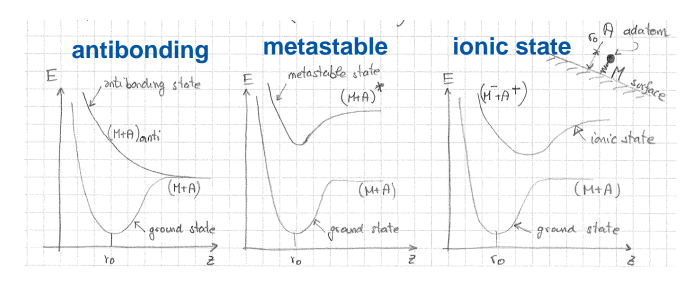
For H_2 the fraction of energy transferred to the molecules is about 2/1840. For typical electron energy of about 300 eV, ΔE is of the order of 0.1 eV.

These values are much lower than the measured energies of desorbed molecules, typically in the **2-10 eV range**. Therefore, **the direct momentum transfer is not the dominant process in ESD**. Electronic energy transfers are then considered.



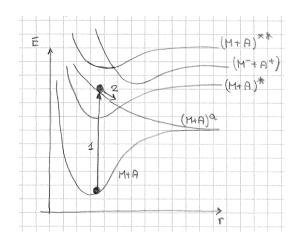
One of the earliest and general models to explain ESD from surfaces was introduced independently by Menzel, Gomer, and Redhead (MGR) in 1964.

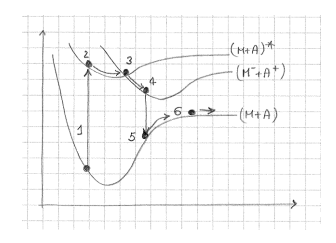
The interaction with the incident electron provokes an adiabatic transition (Franck-Condon principle) to exited states of different nature:



The adiabatic transition implies a change of electronic energy in about **10**⁻¹⁶ **s** with a frozen distance z between nuclei. The transition is vertical in potential energy vs. distance diagram.







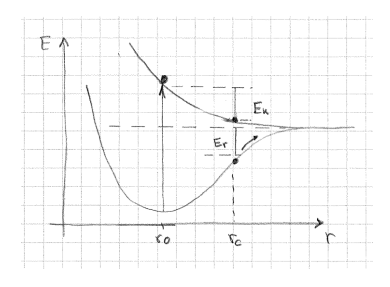
After the excitation, **nuclear motion may occur** over a time scale of about 10⁻¹³-10⁻¹⁴ s: part of the **potential energy is converted into kinetic energy**.

The crossing of potential curves is possible resulting in different de-excitation pathways.



Whenever the original state is reached during the downward transition (rebonding), the molecule can escape if it has acquired enough kinetic energy to overpass the potential binding energy at the position of transition.

The **desorption as a stable neutral** is possible when the downward transition occurs at a distance from the surface which is larger than **a critical distance** \mathbf{r}_{c} .





Particle beams emit photons whenever they are accelerated, namely whenever their velocity vectors are changed.

Synchrotron radiation (SR) is emitted when charged particles moving with relativistic speeds are forced to follow **curved trajectories in magnetic fields**.

The emitted photons impinge on the wall of the vacuum system and may **desorb** gas molecules.

There is experimental evidence that the photon stimulated desorption is generated by a two-step process:

- 1. The impinging photons extract photoelectrons.
- 2. The extraction and recapture of the photoelectrons stimulate the desorption of gas molecules by the ESD mechanism.



A particle of charge 'e', energy E and rest mass m_0 , moving on a circular orbit (radius ρ) radiates electromagnetic radiation with the following power P_{rad} :

$$P_{rad} = \frac{e^2 c}{6\pi\varepsilon_0 (m_0 c^2)^4} \frac{E^4}{\rho^2}$$

where ε_0 and c are the vacuum permittivity and the speed of light, respectively.

The emitted power depends strongly on the **beam energy**, the **radius of the bent trajectory**, and **the mass of the charged particle**.

Consequently, electrons emit much more synchrotron radiation power than protons for the same bending radius and energy:

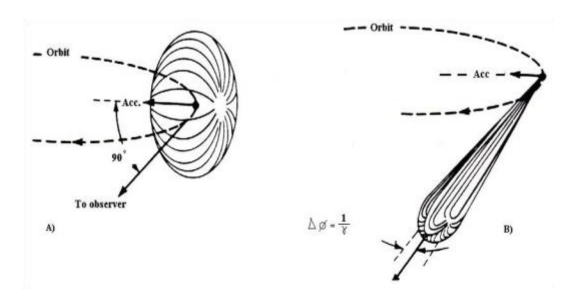
$$\frac{\left(P_{rad}\right)_{electrons}}{\left(P_{rad}\right)_{protons}} = \left(\frac{m_p c^2}{m_e c^2}\right)^4 = 1.13 \times 10^{13}$$



Synchrotron emission is **strongly beamed** along the direction of motion, which is perpendicular to the acceleration.

The emission is concentrated into an angle of the order of $2/\gamma$ rad along the direction of motion:

$$\gamma = 1/\sqrt{1 - (v/c)^2} = E/mc^2$$





Synchrotron radiation from bending magnets has a **very broad energy spectrum**, which is characterised by the **critical energy** ε_{c} :

$$\varepsilon_c = \frac{3}{2} \frac{\hbar c}{\rho} \gamma^3 = \frac{3\hbar c}{2(mc^2)^3} \frac{E^3}{\rho}$$

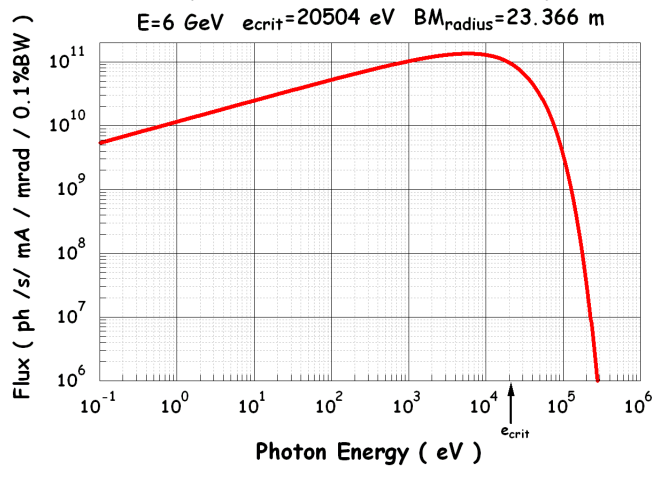
The critical energy subdivide the photon spectrum in two parts of equal emitted power.

For electrons
$$\mathcal{E}_c[KeV] = 0.665 \times E^2[GeV^2] \times B[T]$$

For protons
$$\varepsilon_c[KeV] = 1.1 \times 10^{-10} \times E^2[GeV^2] \times B[T]$$

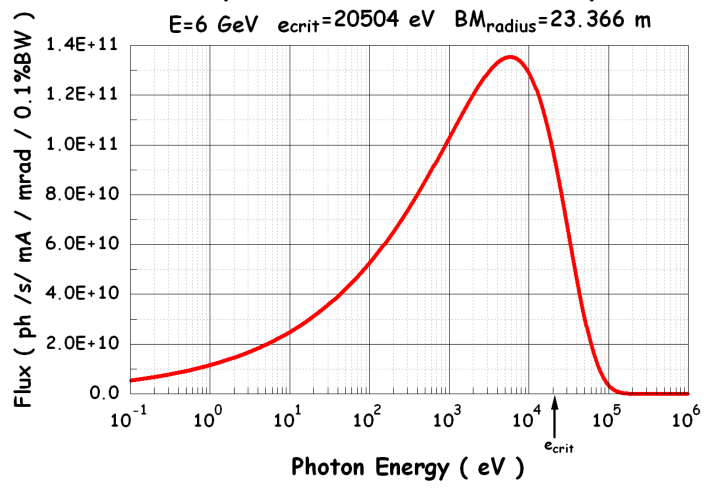


ESRF Synchrotron Radiation Flux Spectrum





ESRF Synchrotron Radiation Flux Spectrum





It can be shown that the number of emitted photon per unit time:

$$\dot{N} = \frac{15\sqrt{3}}{8} \frac{P_{rad}}{\varepsilon_c}$$

$$\dot{N} = \frac{P_{rad}}{\langle \varepsilon \rangle} \rightarrow \langle \varepsilon \rangle = \frac{8}{15\sqrt{3}} \varepsilon_c$$
 Mean photon energy: max of the distribution

For electrons

$$\dot{N} = 8.08 \times 10^{17} I [mA] E [GeV]$$

The linear flux, i.e. the emitted photons per second and meter, is given by

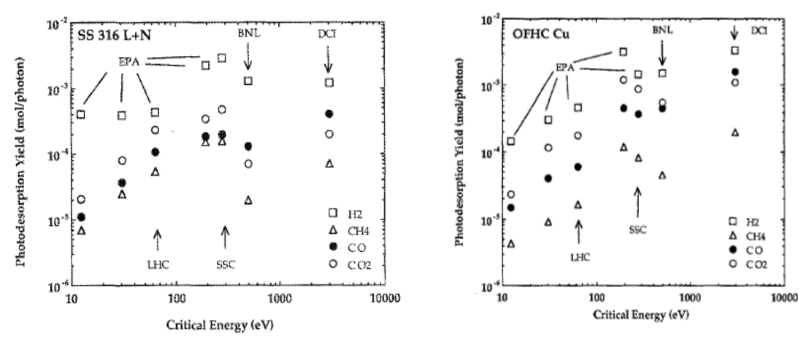
For electrons
$$\frac{d\dot{N}}{ds} = 1.28 \times 10^{17} \frac{I[mA]E[GeV]}{\rho[m]}$$



Photons with energy lower than the photoelectron extraction threshold (workfunction, about 5 eV for metals used in UHV) do not stimulate gas desorption.

 $\eta_{\rm ph}$ depends linearly on the critical energy of the photon spectrum when $\varepsilon_{\rm c}$ < 280 eV:

$$\eta_{ph} \propto arepsilon_c^eta$$

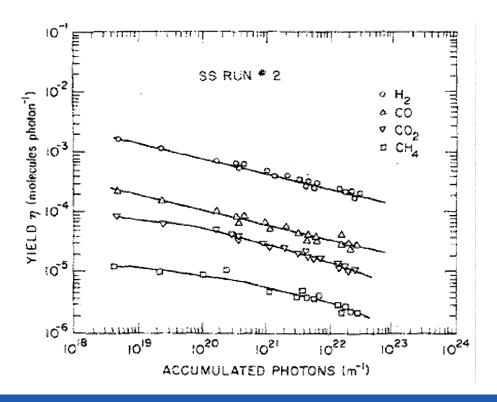


J. Gomez-Goñi, O. Groebner and A. G. Mathewson J. Vac. Sci. Technol. A 12, 1714 (1994)



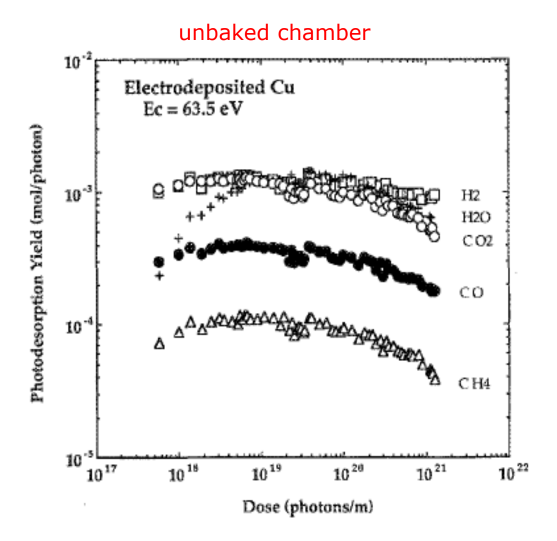
 η_{ph} does not depend significantly on the photon dose D up to about 10²⁰ photons/m. For higher doses, η_{ph} varies as a **power law function of the dose**:

$$\eta_{ph} \propto D^{-lpha}$$



Photon desorption yields for baked stainless steel as a function of the accumulated dose of photons.





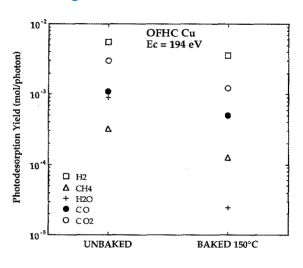


Methods for the reduction of electron and photon induced desorption

The gas desorbed by electrons or photons is extracted from the surface of vacuum chambers or from the first tens of atomic layers underneath.

Therefore, a state of art **surface cleaning** is essential to avoid excessive desorption.

A beneficial effect is obtained by **in situ bakeout** at temperature higher than 120°C; heating the material results in a cleaner surface; however, the effect on the η values is limited and, in general, less than a factor of 10.



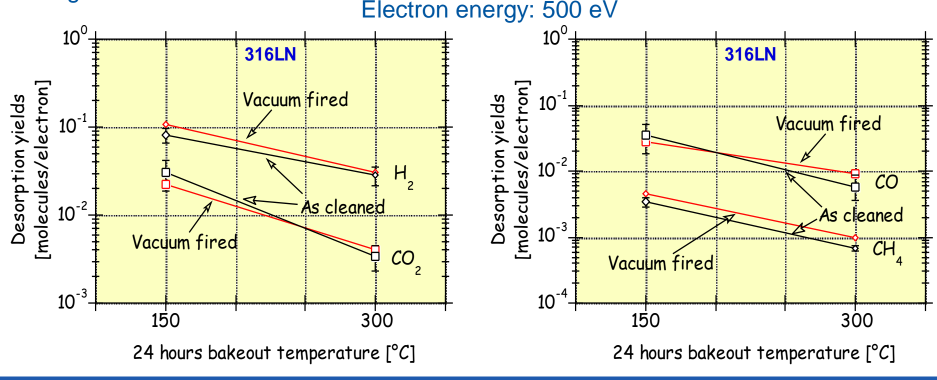


Methods for the reduction of electron and photon induced desorption

Electropolishing and **vacuum firing** have a very limited effect on the η values.

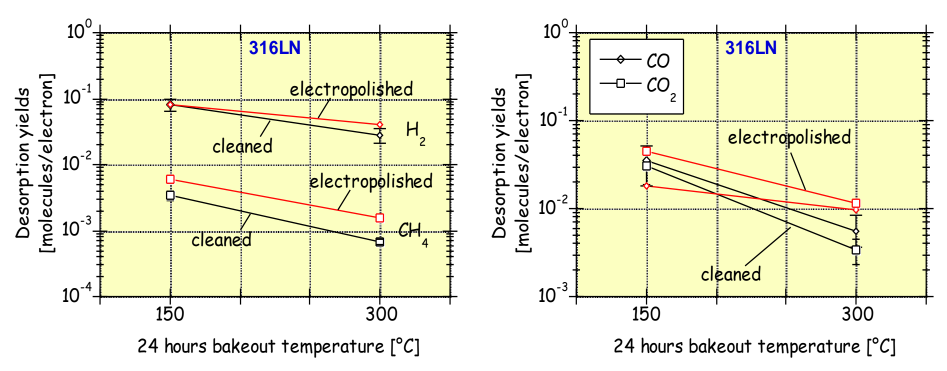
Air exposure after these treatments oxidizes again the surface and regenerates

the gas source.







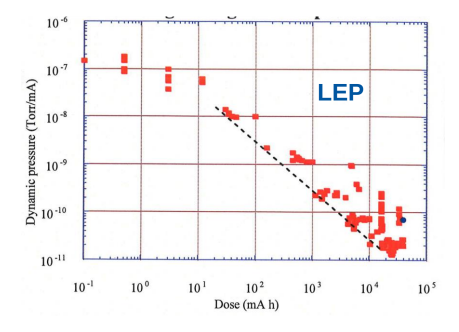


Electropolishing does not influence the desorption yield of H₂. Very small variations are recorded for the other gasses.



The accumulated dose effect lead to a significant reduction of the desorption yields. Power law reductions are measured for both accumulated electron and photon

impingements.



Particle accelerators, whose performance is limited by beam induced desorption, are run at progressively increasing current to accumulate photon or electron dose.

This operation is called 'beam conditioning' of the vacuum chambers. It can last several hours to several days.



Another radical method to reduce stimulated desorption is **coating** the inner wall of the beam pipes with a **thin non-evaporable getter film**.

After in situ bakeout at temperature higher than 180°C, the oxide layer covering the film is dissolved leaving an extremely clean surface.

This treatment is considered in the chapter dedicated to pumps.



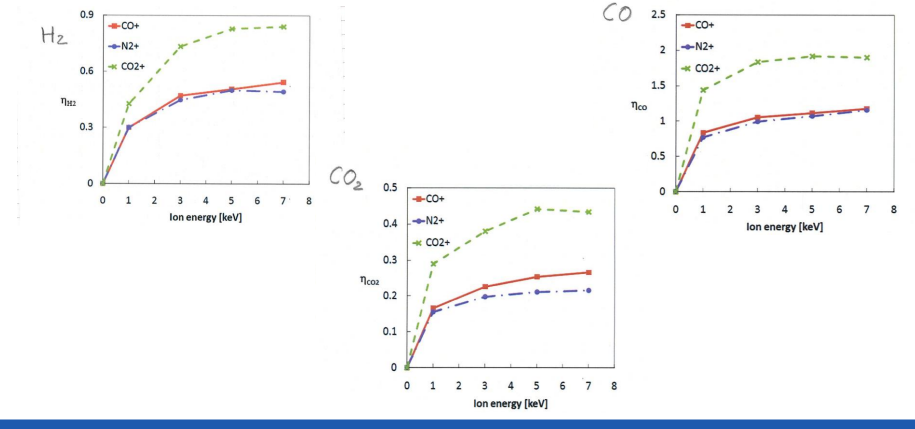
There are two sources of energetic ions:

- 1. Beam particles may collide with the **residual gas** and create ions (H₂+, CH₄+, CO+, CO₂+, etc.). If the beam is positively charged, the ions are accelerated by the beam potential and collide on the nearby walls with energy between **1 eV to several KeV**. The collision results in gas desorption.
- 2. When **heavy ions are accelerated**, beam losses may lead to collision of high-energy heavy ions with the wall of the vacuum system. Typical ions are Pb⁵³⁺, U⁷³⁺, and Ar¹⁰⁺. Experimental studies have been carried out with beam energy in the range from **1 MeV/nucleon to 100 GeV/nucleon**. In this case, the desorption yields may be orders of magnitude higher than those for residual-gas ions.



Residual-gas ion

lons are more effective in desorbing gas than electrons and photons. Typical η_i values for baked copper and ion energy of about 1 KeV are about 1 for H₂ and CO; 5 and 10 times lower for CO₂ and CH₄, respectively.

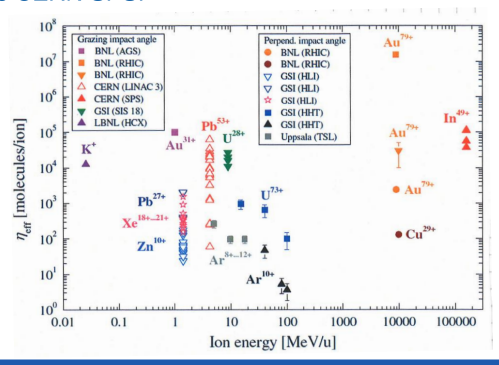




High-energy heavy-ions

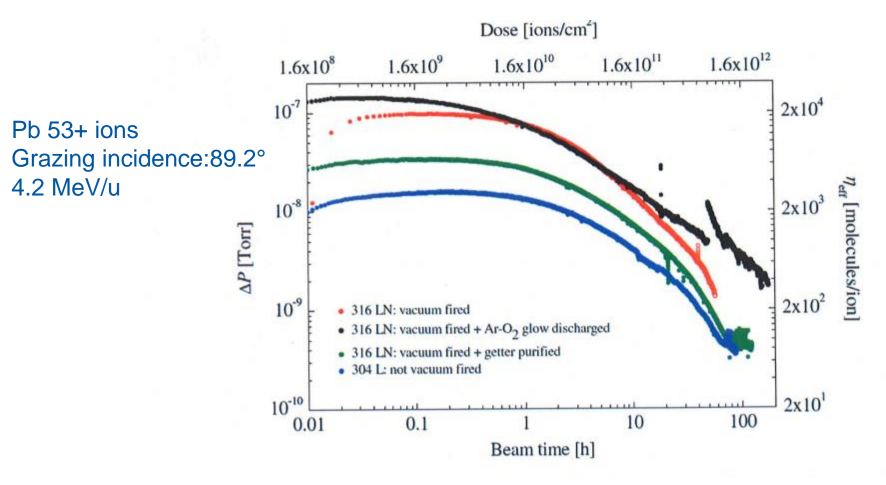
In the last 15 years, several experiments have shown that high energy (**E** > **1 MeV/nucleon**) ions can induce the desorption of a **huge quantity of gas molecules**.

Desorption yield up to **10**⁵ molecules/ion were measured for In⁴⁹⁺ at 158 GeV/nucleon at the CERN SPS.





High-energy heavy-ions

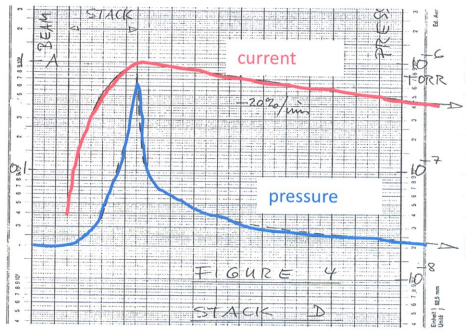


The accumulated dose has a beneficial effect also on the ion induced desorption



Ion induces desorption can trigger a rapid pressure rise in accelerators where positive beams circulate.

This phenomenon was shown first at the CERN ISR when increasing the proton current to about 1 A

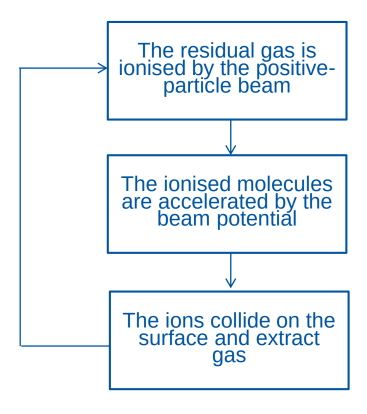


First documented pressure bump in the ISR

E. Fischer/O. Gröbner/E. Jones 18/11/1970



The pressure rise is generated by a positive feedback process that can be depicted with the following block diagram.





The critical current at which the runaway process starts can be obtained in the following way:

1. The ionisation probability for one proton is $\sigma_i n = \sigma_i^P/k_B T$ and the flow of circulating protons is I/e. Therefore, the total gas flow of desorbed molecules per unit length of vacuum chamber is:

$$Q = \eta_i \cdot \sigma_i \frac{PI}{e} + Q_{th}$$

2. The pressure in the beam pipe is:

$$P = \frac{Q}{S_{eff}} = \eta_i \sigma_i \frac{PI}{S_{eff}} + \frac{Q_{th}}{S_{eff}} \rightarrow P = \frac{Q_{th}}{S_{eff}} - \eta_i \sigma_i \frac{I}{e}$$

3. The pressure diverges when:

$$S_{eff} - \eta_i \sigma_i \frac{I}{e} \rightarrow 0$$



Critical current:

$$I_c = rac{S_{\it eff} e}{\eta_i \sigma_i}$$

Gas	$\sigma_{\rm i}[10^{\text{-}18}~{ m cm}^2]$	$\sigma_{\rm i} [10^{-18}~{\rm cm}^2]$
	26 GeV	7 TeV
H_2	0.22	0.37
He	0.23	0.38
CH_4	1.2	2.1
CO	1.0	1.8
CO_2	1.1	2.0
Ar	1.6	2.8

Example:

For an effective pumping speed of 10 l s⁻¹ and an ionisation cross section of 10⁻¹⁸ cm² (CO ionised by protons at 26 GeV), the critical current is about 16 A if the desorption yield is 1.



Part 5 Gas pumping in particle accelerators



Gas pumping

In molecular regime:

- gas molecules cannot be removed by suction: the molecules do not transfer energy and momentum amongst them; pumps act on each molecule singularly;
- pumps are classified in two families:
 - 1. momentum transfer pumps;
 - 2. capture pumps.
- In the first, molecules receive a momentum component pointing towards the pump outlet (foreline) where the gas is compressed and evacuated by pumps working in the viscous regime (for example: rotary vane, diaphragm, and scroll pumps).
- Capture pumps remove molecules from the gas phase by fixing them onto an internal wall.



In molecular pumps, molecules are headed for the outlet by moving surfaces.

Molecules impinge and adsorb on the moving surface; on desorption, the velocity distribution is superimposed to the **drift velocity** of the wall →

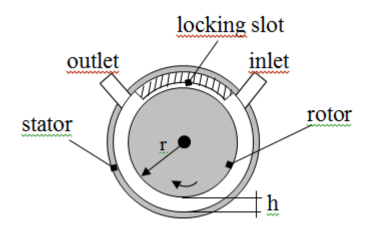
As a result, molecules are preferentially redirected towards the direction of the wall movement, i.e. the **density of molecules increases** in the same direction.

A moving wall produces a **gas flow**.





The first molecular pump was invented by W. Gaede. In the original design, the moving surface is a rotor revolving at high frequency



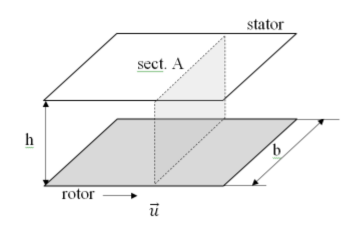
To prevent backstreaming, the inlet and the outlet were separated by a **very thin slot** (locking slot). The slot height was of the order of **10**⁻² **mm**.

The most important characteristics of molecular pumps are:

- 1. Pumping speed S
- 2. Maximum compression ratio $K_0 = \left(\frac{P_{OUT}}{P_{IN}}\right)_{MAX}$

The parameters affecting S and K_0 can be identified by a simple model.





At any time, **half of the molecules** have just collided with the moving surface (rotor) and drift in the direction of the velocity \vec{u} . The **other half** hits the stator where the drift component is lost.

As a result, the drifted molecular flow Q_p towards an imaginary section A is:

$$Q_p = \frac{1}{2} n \cdot u \cdot b \cdot h = \frac{1}{2} \frac{P}{k_B T} \cdot u \cdot b \cdot h$$

Where 'bh' is the area of the cross section A and 'ubh' is the gas volume drifting through section A in one second.



Converting in pressure-volume units and dividing by P, the pumping speed is obtained:

$$S = \frac{Q_p}{P} = \frac{1}{2}u \cdot bh$$

The pumping speed of molecular pumps:

- Depends linearly on the speed of the moving wall
- Does not depend on the nature of the gas (in the frame of this model)→molecular pumps are not selective!



The moving wall produces a higher gas density near the pump outlet. In turn, the pressure gradient generates a gas backflow Q_{bf} that can be written as:

$$Q_{bf} = -\bar{c} \, \nabla P$$

where \bar{c} is the conductance of the unit-length duct of cross section area bh.

For h<
b:

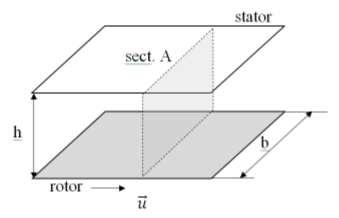
$$\bar{c} = \frac{2}{3} \frac{(bh)^2}{b+h} \langle v \rangle$$

The net flow is therefore given by:
$$Q = Q_p - Q_{bf} = \frac{1}{2} P \ u \ bh - \bar{c} \ \nabla P$$

When the net flux is zero, the maximum compression ratio is reached; in this case:

$$\frac{dP}{P} = \frac{1}{2} \frac{u \ bh}{\bar{c}} dx$$





Integrating between inlet and outlet (distance L) the maximum compression ratio is obtained:

$$K_{0} = \left(\frac{P_{OUT}}{P_{IN}}\right)_{MAX} \propto \exp\left[\frac{u}{\langle v \rangle} \times \frac{L}{h}\right] \propto \exp\left[\left(u\sqrt{m_{i}} \frac{L}{h}\right)\right]$$

L and h length and width of the pump duct

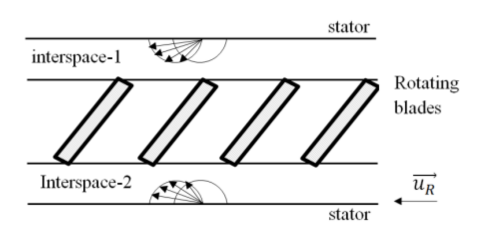
High compression ratio for molecular pumps can be obtained for:

- Fast moving surfaces: at least of the same order of the average molecular velocity (>100 ms⁻¹).
- Narrow and long pumping ducts (lower backstreaming).
- Heavy masses → the maximum compression ratio depends strongly on the molecular mass, the lowest being for H₂ → The ultimate pressure of molecular pumps is dominated by H₂.



For many years, the industrialisation of molecular pumps was hindered by the machining of the locking slot. The narrow gap between rotor and stator created mechanical issues due to tolerances and thermal expansion.

The problem was overcome in 1957 when W. Becker invented the turbomolecular pump (TMP). In the TMP, the momentum transfer is produced by rapidly rotating blades rather than parallel surfaces.



The molecules **seen from the blades** have a velocity oriented toward the blades' channels when they come from space 1.

From space 2, most of the molecules hit the blades and are backscattered.

A significant gas flow is set only if at least $\langle v \rangle \approx u$.



In the real turbomolecular pump, the gas is compressed by **several series of rotating blades**.

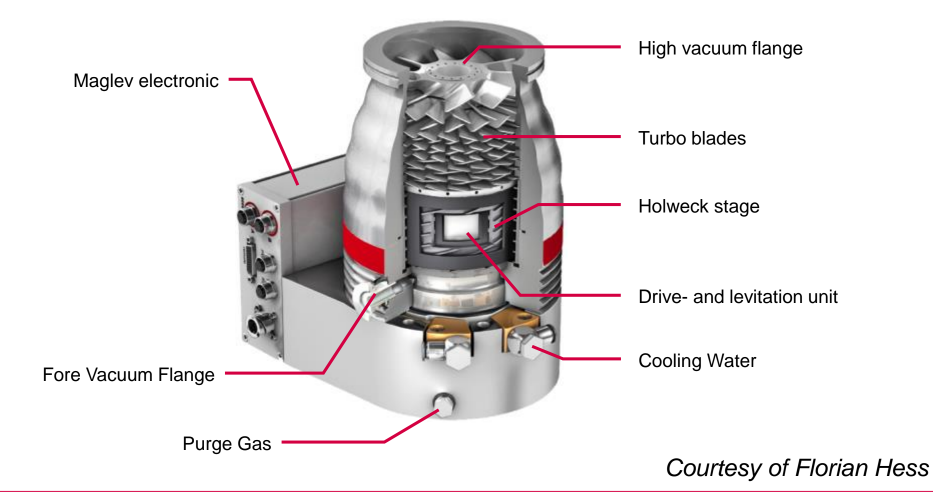
Every series of rotating blades is followed by a series of static blades. Molecules transmitted through the rotating blades' channels hit the static blades; as a result, the angular distribution of velocity is randomized and the molecules are ready for the **next compression stage**.

The momentum transfer is effective only if the molecules **do not experience intermolecular collisions** after hitting the blades; this is equivalent to say that the mean free path has to be larger than the blade distances. As a result, this type of pumps works at full pumping speed **only in molecular regimes** (P < 10⁻³ mbar).





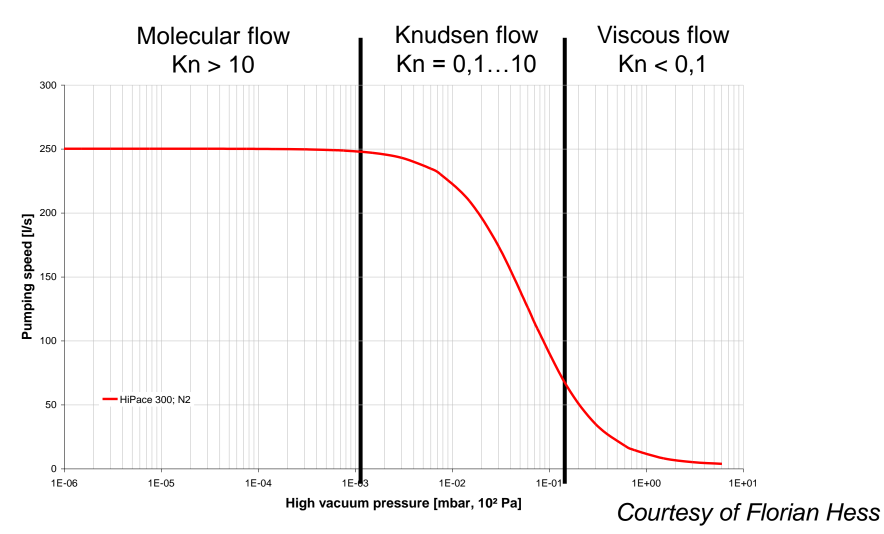
Maglev Turbopump Design







Different Flow Areas

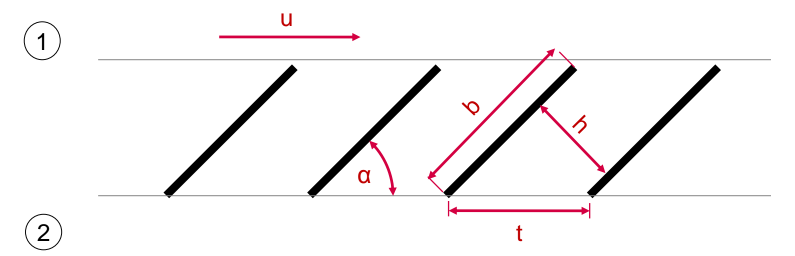






Theory of Turbo Pumping Effect

Courtesy of Florian Hess



Compression ratio

$$\mathbf{K}_0 = \exp\left(\frac{u}{\langle \mathbf{v} \rangle \cdot \mathbf{g} \cdot \mathbf{t} \cdot \sin \alpha}\right)$$

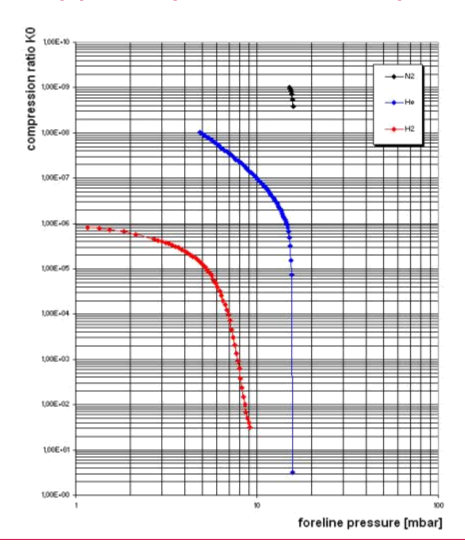
$$\langle \mathbf{v} \rangle = \sqrt{\frac{8 \cdot \mathbf{R} \cdot \mathbf{T}}{\pi \cdot \mathbf{M}}}$$

g = geometric factor

Source: Karl Jousten, Wutz Handbuch Vakuumtechnik, edition 9, Vieweg, p. 329



Gas Type dependant Compression Ratio

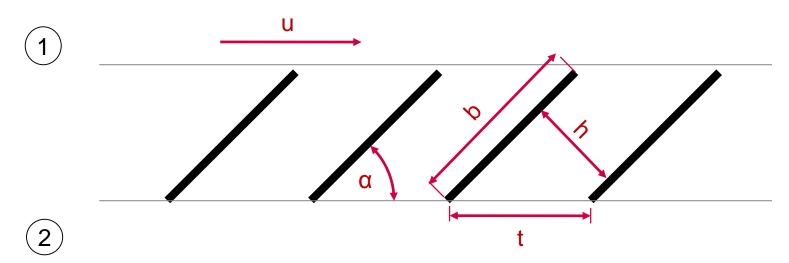


$$K_0 = \frac{p_{FV}}{p_{HV}}$$





Theory of Turbo Pumping Effect



Pumping Speed

$$\mathbf{S}_0 = \frac{1}{2} \cdot A \cdot u \cdot \sin \alpha \cdot \cos \alpha$$

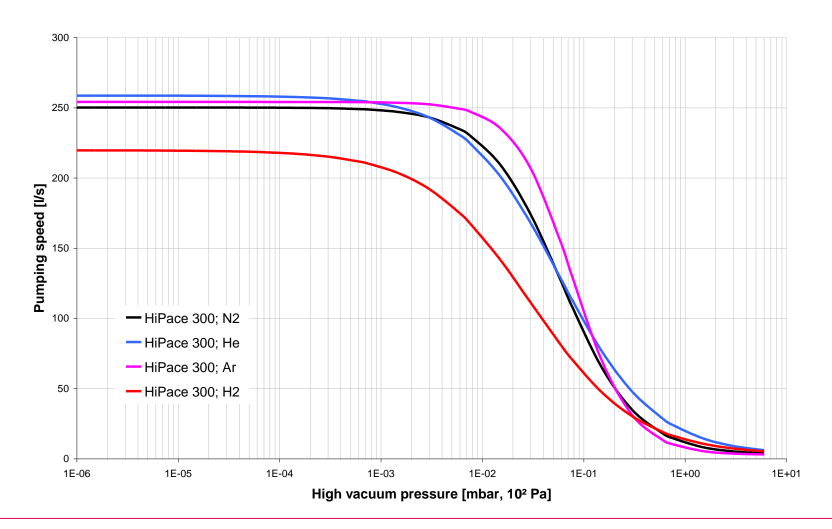
$$S_{\text{eff,i}} = \frac{S_{0,i}}{1 - \frac{1}{K_{0,i}} + \frac{S_{0,i}}{K_{0,i} \cdot S_{\text{eff,i-1}}}}$$

Source: Karl Jousten, Wutz Handbuch Vakuumtechnik, edition 9, Vieweg, p. 328 f





Gas Type dependant Pumping Speed





History of the Turbopump

- In 1958 Dr. W. Becker invents the worlds first Turbopump
- Invention was done by chance

TVF 400

Pumping speed 140 l/s for N₂

Weight: approx. 60 Kg





HiPace 300

Pumping speed 260 l/s for N₂

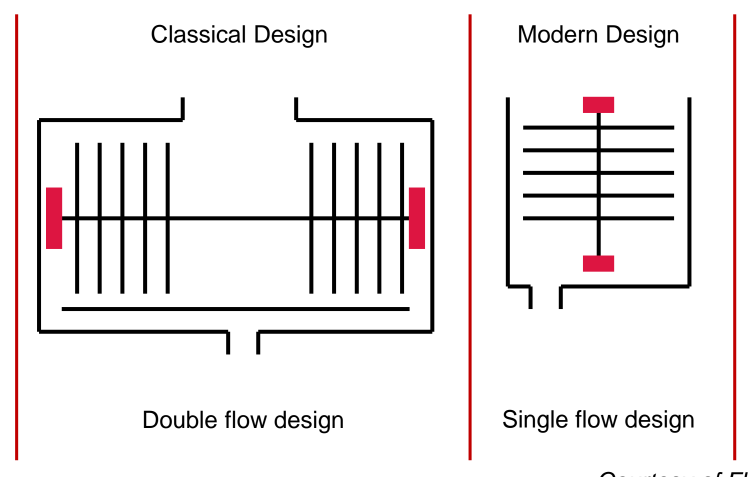
Weight: 6.2 Kg







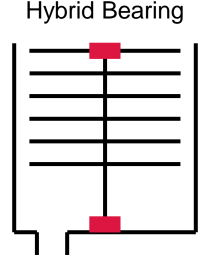
General Turbopump Design





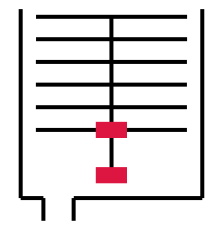






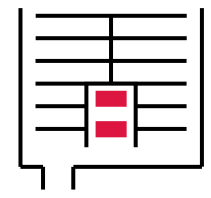
- Robust design
- Less vibrations
- Easy maintenance
- Oil lubrication

Cantilever Bearing



- No field service
- Grease lubrication





- Compact size
- Optimized crash design



Different Rotor Concepts

Pure Turbo



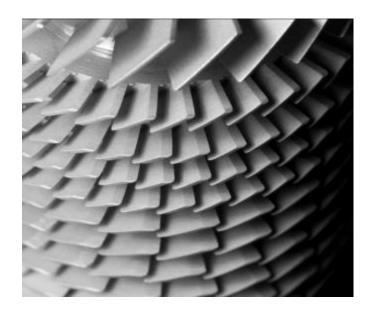
- Gas flow optimized
- Suitable for process applications

With Holweck Stage



- Compression optimized
- Ultra High Vacuum





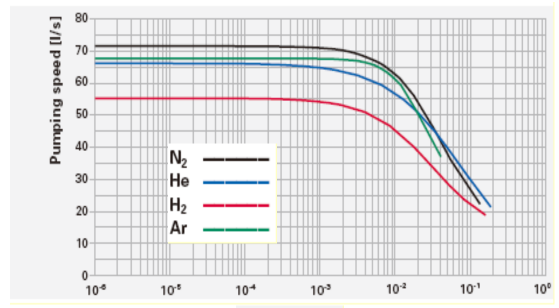
The indications given by the simplified model for molecular pumps are valid also for the TMP:

S: no strong influence of gas molecular mass + linear dependence on revolution speed

K₀: **exponentially** dependent on: revolution speed and square root of gas molecular mass



Gas pumping: momentum transfer

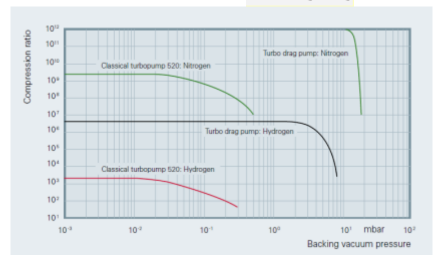


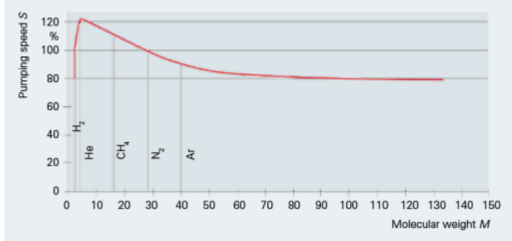
TMP pumping speeds are in the range from 10 l/s to 25,000 l/s.

Their ultimate pressure (H₂) is of the order of 10⁻¹⁰, 10⁻¹¹ mbar

Courtesy of Pfeiffer Vacuum
http://www.pfeiffer-vacuum.com
Vacuum Technology KnowHow









At the pump outlet, the compressed gas is evacuated by a **mechanical pump** (often called backing pump) operating in viscous regime. The turbomolecular and its backing pump are in general **assembled in a single unit** that includes power supplies, controls and instrumentation.





The main drawback of TMP is related to possible **mechanical failures** leading to definitive damage of the high-speed rotor.





Advantages of TMP:

- 1. constant pumping speed in a large range of pressure
- 2. no memory effect (the gas is definitively evacuated) nor gas selectivity
- 3. start working at relatively high pressure (as soon as molecular regime is attained)

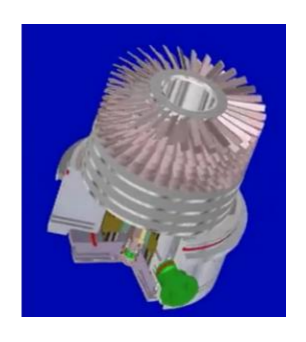
Disadvantages of TMP:

- 1. mechanical fragility
- 2. risk of contamination from the backing pump
- 3. need of venting anytime the pump is stopped to block backstreaming of contaminations→need of valve between TMP and vacuum vessel
- 4. intrinsic limitation in ultimate pressure of H₂

Present trend:

- Use of dry pumps as backing pumps (but lower compression ratio than oil pumps)
- 2. Increase compression ratio by adding molecular drag stages below the set of TMP blades (very compact design)
- 3. Remove all lubricated mechanical bearing by magnetic rotor suspension (higher cost).





http://www.youtube.com/watch?v=1xZe1H2XHhM&feature=youtube_gdata_player

http://www.youtube.com/watch?v=XOca57gPsq4



Gas pumping: capture pumps

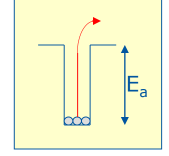
Capture pumps remove gas molecules by fixing them on an internal wall.

To be efficient, capture pumps must block the gas molecules for an average time (sojourn time t_s) much longer than the typical running time of the accelerator.

To do so the sojourn time on the wall has to be much longer than the typical time of the accelerator run. An estimation of sojourn time is given by the Frenkel law *J. Frenkel, Z. Physik, 26, 117 (1924*):

$$t_s = t_0 e^{\frac{E_a}{k_B T}}$$

where E_a is the adsorption energy and $t_0 \approx \frac{h}{k_B T} \approx 10^{-13} s$.



E_a >> k_BT → Chemical pumps (**getter pumps**)

$$T \ll \frac{E_a}{k_B} \rightarrow \mathbf{cryopumps}$$



Gas pumping: capture pumps

In **chemical pumps**, the binding force involves electron exchange between gas molecules and surfaces (**chemisorption**). T

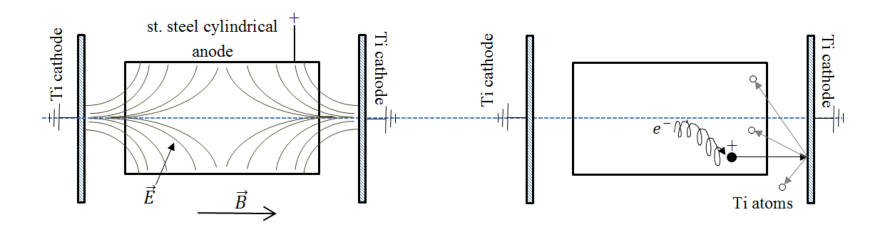
Typical binding energies for getter pumps are higher than 1 eV/molecule.

In cryogenic pumps, Van der Waals interactions might be sufficiently strong to fix molecules to cold surfaces (**physisorption**).

In this case, the binding energy can be **much lower than 0.5 eV/molecule**.

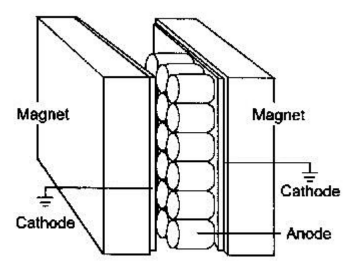


- In SIP the residual gas is ionized in a Penning cell.
- The ions are accelerated towards a cathode made of reactive metal.
- The collisions provoke sputtering of reactive metal atoms that are deposited on the nearby surfaces.





In the diode SIP, the anode is composed of **several open cylinders** made of stainless steel at a positive electrical potential (from 3 to 7 kV). The **cathodes are plates of Ti** -at ground potential- placed a few millimetres from both anode extremities.

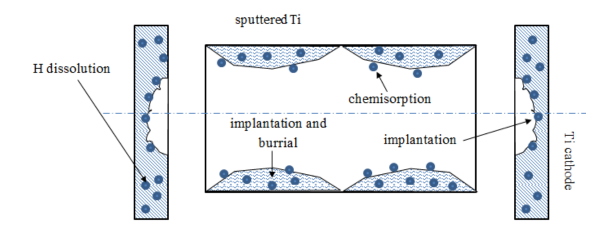




- The pumping action is given by:
 - chemical adsorption onto the reactive metal layer and subsequent burial by additional metallic atoms of gas molecules: all gases except rare gases
 - 2. implantation of gas ions in the cathode and of energetic neutrals bounced back from the cathode in the deposited film: only mechanism of pumping for rare gases
 - 3. diffusion into the cathode and the deposited film: only H₂

http://www.youtube.com/watch?v=_snzYepQTjl





Molecular **implantation in the cathodes is not a permanent pumping**; in fact, the progressive erosion due to gas ion sputtering soon or later frees the implanted gas.

Once released from the cathodes, reactive gas species are chemisorbed while rare gases remain in the gas phase until the next implantation.



Pressure instabilities have been reported after excessive pumping of rare gas:

the continuous **erosion liberates gas** that increases the pressure, which in turn increases the erosion.

Consequently, a pressure rise is generated; this is stopped when most of the rare gas is implanted in a deeper zone of the cathode.

This type of instability is typically observed when large quantities of air are pumped (Ar 1% in air) and are known as 'argon disease'.

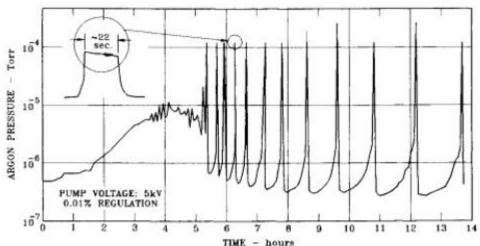


Figure 2.4.12. Example of instabilities in a system with one sputter-ion pump with titanium cathodes, after pumping argon at 5 x 10⁻⁷ Torr for a few hours



To increase the pumping efficiency of noble gas, the rate of ions implantation in the cathode has to be reduced while increasing the rate of energetic neutrals impingement on the anode and their burial probability.

Two different approaches:

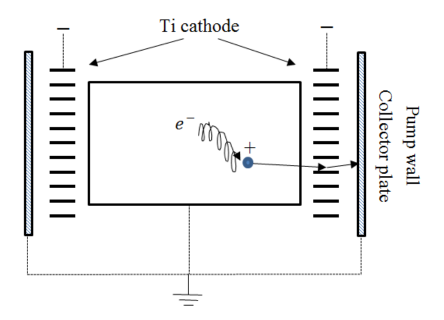
1. Heavier atoms for the cathode

Ta (181 amu) is used instead of Ti (48 amu). The ions, once neutralized, bounce back at higher energy and rate → these pumps are called 'noble diode'



2. Different geometry of the Penning cell.

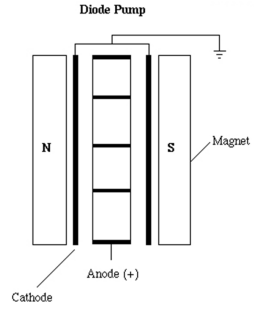
- a) Three electrodes are used: **triode pumps**. The cathodes consists of a series of small platelets aligned along the cell axis.
- b) The collisions ion-cathode are at glancing angle → higher sputtering rate of Ti atoms + higher probability of neutralization + higher energy of bouncing + lower probability of implantation in the cathode.



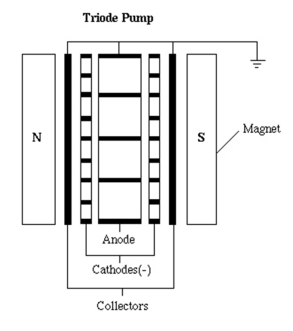


Gas pumping

DIODE

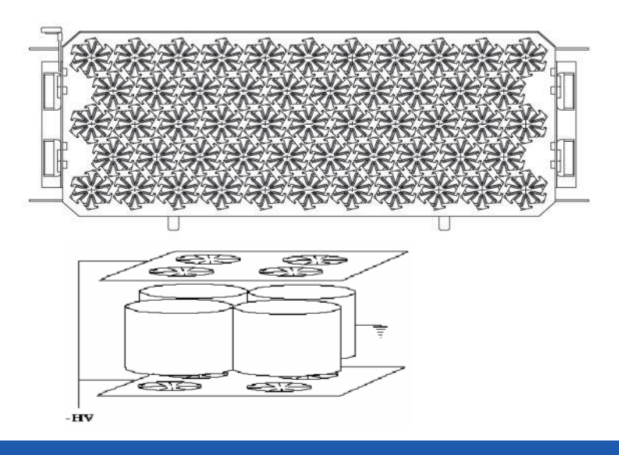


TRIODE





An improved triode pump is the **StarCell® produced by Agilent**. Two sheets of Ti that are cut and bent to form radial blades replace the series of small plates. This increases the rigidity of the cathode and reduces the risks of short circuits





Pumping speed for SIP depends on the pressure at the pump inlet and the nature of the gas.

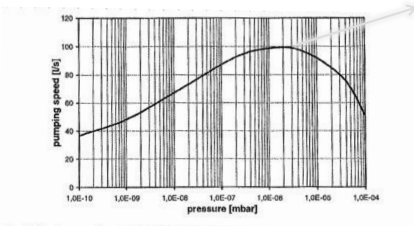


Fig. 5 Pumping speed vs pressure for a standard diode with S_n= 100 l/s (for air after saturation).

Nominal pumping speed for N₂: Agilent StarCell®

DN	S [I s ⁻¹]
63	50
100	70/125
150	240/500

Nominal pumping speed

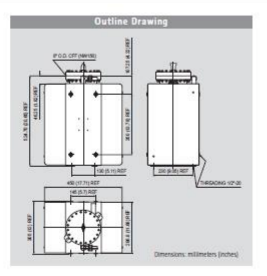
GAS	DIODE PUMPS	TRIODE PUMPS	
AIR	1	1	
N_2	1	1	
O ₂	1	1	
H ₂	1.5-2	1.5-2	
CO	0.9	0.9	
CO ₂	0.9	0.9	
H ₂ O	0.8	0.8	
CH ₄	0.6-1	0.6-1	
Ar	0.03	0.25	
He	0.1	0.3	

Pumping speed normalized to air



VacIon Plus 500





	StarCell®	Noble Diode	Diode
Nominal pumping speed for Nitrogen (*) (Vs)	410	440	500
Operating life at 1x10-6 mbar (hours)	80,000	50,000	50,000
Maximum starting pressure (mbar)	≤ 1x10 ⁻³	≤ 1x10 ⁻³	
Ultimate pressure	Below 10- ^M		
Inlet flange	8" CFF (NW 150) AISI 304 ESR		
Maximum baking temperature (°C)	350		
Weight, kg (fbs)	120 (264)		







Hydrogen pumping by SIP

- H₂ is mainly pumped by diffusion into the cathode.
- To be adsorbed, H₂ must be dissociated. Only 2.5% of the ions created in a low-pressure H₂ Penning discharge are H⁺ ions.
- The dissociation is possible only on atomically clean Ti.
- H₂⁺ ions have poor sputtering yield: 0.01 at 7 KeV on Ti.
- When H₂ is the main gas, it takes a long time to clean the cathode surface by sputtering.
- As a consequence, at the beginning of the operation the pumping speed for H₂ is lower than the nominal and increases gradually with time.



Hydrogen pumping by SIP

- The simultaneous pumping of another gas has strong effects on H₂ pumping speed.
 - Higher sputtering yield→faster cleaning→ higher pumping speed
 - Contaminating of the Ti surface → lower pumping speed
 - Desorption of implanted H ions→ lower pumping speed
- When the concentration of H₂ is higher than the solubility limit in Ti, hydride precipitates are formed → Ti expansion and hydrogen embrittlement → short circuits and cathode brittleness (for 500 l/s pumps: typical value are 13000 mbar.l of H₂)



High Pressure Operation

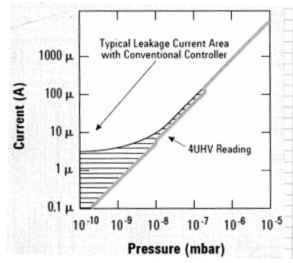
- High pressure (>10⁻⁵ mbar) operation can generate thermal run-away. It is frequently noticeable during the pumping of H₂ or after the absorption of high quantity of H₂ (for example due to pumping of H₂O).
- The Penning discharge heats the cathode and provokes gas desorption, which enhance the discharge. This positive feedback mechanism can melt locally the cathode.
- The total electrical power given to the pump has to be limited at high pressure.



Pressure measurement by ion pumps

- The discharge current of the penning cells can be used for pressure measurement.
- In the low pressure range, the current measurement is **limited by field emission** (leakage current): pressure reading limitation in the 10⁻⁹ mbar range.
- By reducing the applied voltage in the lower pressure range, the pressure measurement is possible down to 10⁻¹⁰ mbar.

Courtesy of Agilent Vacuum





Gas pumping: capture pumps / getter

The surface of getter materials reacts with gas molecules by forming **stable chemical compounds**.

This is **possible only if** the surface is clean, **free of contamination and native oxide**.

The clean metallic surface is obtained by:

- Sublimating the reactive metal in situ → Evaporable Getters, Sublimation Pumps
- 2. **Dissolving** the surface contamination into the bulk of the getter material by heating in situ (activation): Non-Evaporable Getters **NEG**.



Gas pumping: capture pumps / getters Sticking probability

Getter surfaces are characterized by the sticking probability α :

$$\alpha = \frac{number\ of\ molecules\ captured}{number\ of\ molecules\ impinging}$$

$$0 \le \alpha \le 1$$

$$S = \alpha A_{getter} C'$$

For α =1, the pumping speed of the surface is equal to its **maximum theoretical** pumping speed.

Getter materials do not pump rare gases and methane at room temperature.

In that respect, they always need auxiliary pumping to keep a stable pressure



Gas pumping: capture pumps / getters Sticking probability

The sticking probabilities of homonuclear diatomic molecules are in general lower than that of molecules composed of dissimilar atoms.

The values of α for common gas species in vacuum systems may be ordered in a growing series as follows:

$$N_2 < H_2 < O_2 < H_2 O < CO < CO_2$$

For N_2 typical α are in the high 10^{-3} to low 10^{-2} range; for CO and CO₂, α close to 1 may be measured (for negligible coverage).



Gas pumping: capture pumps / getters Sticking probability

The sticking probability is **strongly dependent on the gas surface coverage**, i.e. the quantity of gas previously pumped on the getter material.

As the gas surface coverage increases, the sticking probability decreases. The pumping becomes **negligible when the surface is fully covered** by gas molecules. The pumping is restored by regenerating the surface.

As an example, complete saturation of a smooth NEG surface is obtained when about 10^{15} CO molecules cm⁻² are adsorbed. Quantities about 5-10 times higher are reported for H_2O and O_2 .

H₂ dissociates on the getter surface and diffuses into the bulk of the NEG material. As a result, its pumping does not block the adsorption of other molecules.

The pumping capacity of hydrogen is orders of magnitude higher than that for other gas species.



Gas pumping: capture pumps / getters Sievert's law

Due to the high mobility of H atoms and the relatively low binding energy, the **pumping of this gas is reversible**: H₂ may be released by heating.

The opposite effects of pumping and desorption set up an **equilibrium pressure** P_{H2} that depends on heating temperature T and hydrogen concentration c_H .

The three variables are correlated by the **Sievert's law**. In the logarithmic form:

$$\log P_{H2} = A + 2\log c_H - \frac{B}{T}$$

A and B are typical values for a given getter material.



Gas pumping: capture pumps / sublimation pumps

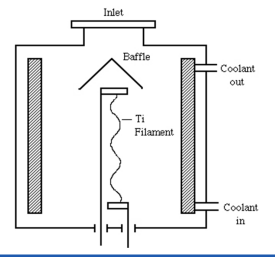
For particle accelerators Ti is the sublimated metal.

Ti alloy rods are heated up to 1500°C attaining a Ti vapour pressure of about 10⁻³ mbar.

The Ti thin film is deposited onto the pump vessel.

In some cases, the wall of the sublimation pumps may be cooled to liquid nitrogen temperature. In this case, the sublimated film is rougher and consequently the sticking probabilities are higher.





Courtesy of Kurt J. Lesker Company http://www.lesker.com/newweb/Vacuum_Pumps



Gas pumping: capture pumps / sublimation pumps

The sticking probabilities depend on the nature of the gas and the quantity of gas already pumped.

$$\max \begin{cases} H_2: \ 10^{-2} \le \alpha \le 10^{-1} \\ CO: 5 \times 10^{-1} \le \alpha \le 1 \end{cases}$$

The sticking probability is negligible:

- For CO, one monolayer adsorbed
- For of O₂ several monolayer
- For N₂ fraction of monolayer

A new sublimation is needed.

Hydrogen diffuses in the Ti film→much higher capacity

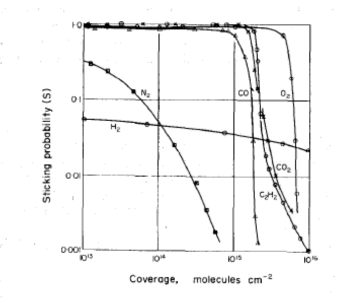
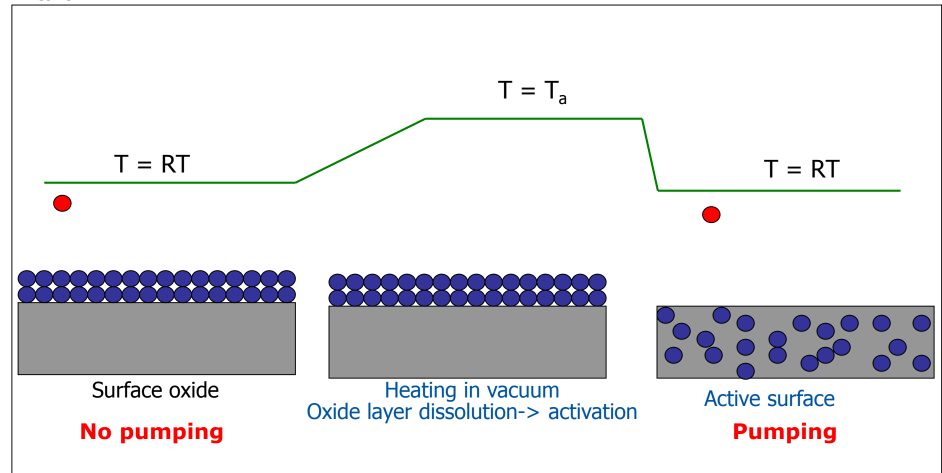


Figure 4. Room temperature sorption characteristics for pure gases on batch evaporated clean titanium films.

A K Gupta and J H Leek, Vacuum, 25(1975)362

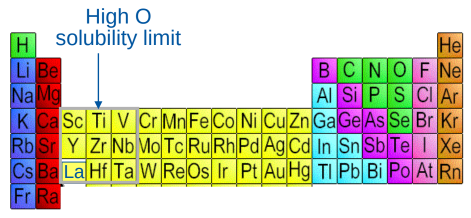


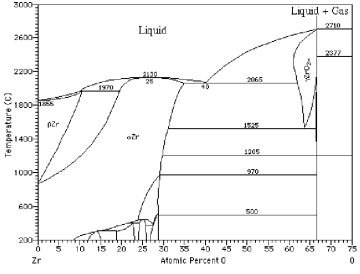
The dissolution of the native oxide layer (**ACTIVATION**) is possible only in metals having very high oxygen solubility limit, namely the elements of the 4th group: Ti, Zr and Hf.





Gas pumping: capture pumps / sublimation pumps







The activation temperature of the 4th group elements can be decreased by adding selected elements that increase oxygen diffusivity.

NEG materials are produced industrially by powder technology. Small grains are sintered to form pellets, discs or plates. The grains can also be pressed at room temperature on metallic ribbon.

A typical alloy produced by SAES Getter is **St707**:

Element	Concentration [wt. %]	Main role in the alloy
Zr	70	High O solubility limit.Chemical reactivity
V	24.6	Increases O diffusivity,Chemical reactivity
Fe	5.4	- Reduces pyrophoricity

Full pumping speed is obtained after heating at 400°C for 45' or 300°C for 24h



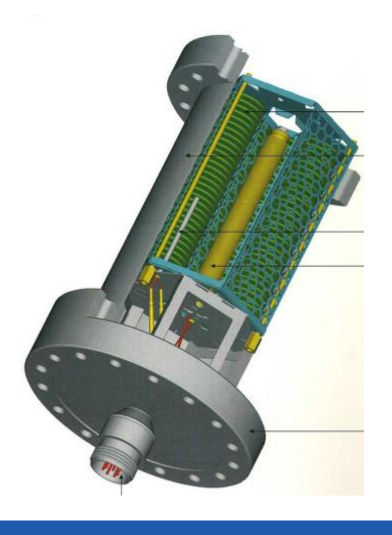
Courtesy of SAES Getters, www.saesgetters.com

Gas pumping: capture pumps / NEG





Different shapes of NEG pumps



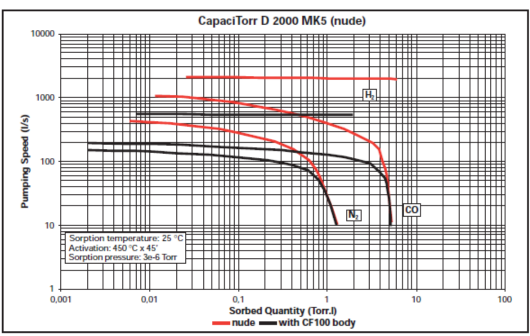


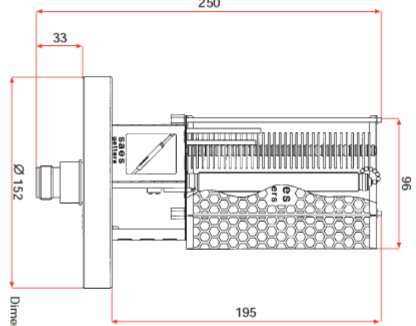
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33

The **high porosity of NEG materials** allows pumping of relatively high quantities of gas without reactivation: for CO about 100 times higher than those for sublimation pumps per unit of geometrical surface of active metal.

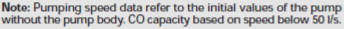
Courtesy of SAES Getters, www.saesgetters.com





Typical Pump Characteristics

Typical Fullip Characteristics				
Alloy Type	St 172®			
Alloy Compo	ZrVFe			
Getter Mass	225			
Getter Surface (cm²)		1900		
Pumping Speed (I/s)	H ₂	2000		
Speed (I/s)	со	1000		
Sorption	H ₂	4500		
Capacity (Torr I)	CO Room Temperature	5		
(10111)	CO Total	2000		





The maximum H₂ sorption capacity is limited by H₂ embrittlement of the NEG elements.

In general a safe limit is 20 Torr I/g is given by the supplier.

The stored H₂ can be desorbed by heating and pumping out with an auxiliary pump (for example a turbomolecular pump).



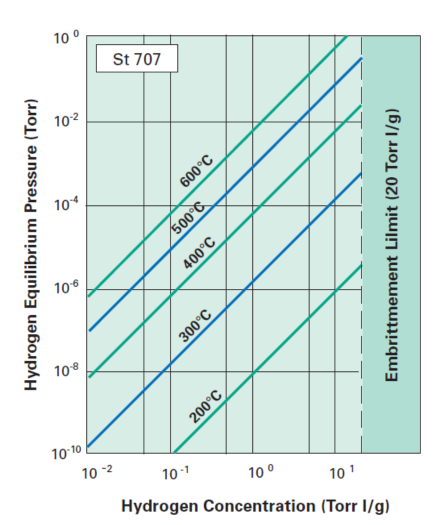


Figure 3 - Hydrogen equilibrium pressure for the St 707 alloy

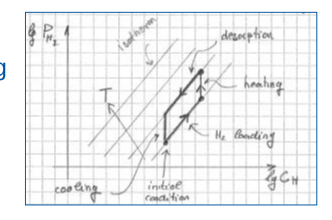


The time t_R needed to regenerate a NEG pump is given by:

$$t_R = \frac{M_{NEG}}{S_{aux}} \left(\frac{1}{q_f} - \frac{1}{q_i} \right) \cdot 10^{-\left(A + \frac{B}{T}\right)}$$

Where:

 q_f and q_i are the final and initial concentrations in Torr I/g S_{aux} is the pumping speed of the TMP M_{NEG} the mass of the NEG in g A and B typical values of the NEG material For St707: A=4.8, B= -6116

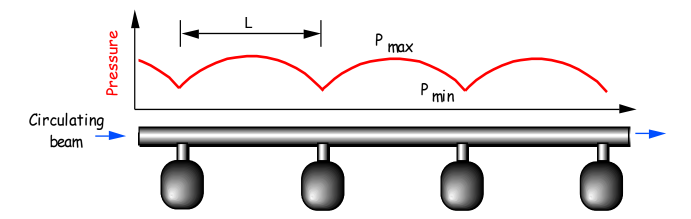


The equilibrium pressure during the regeneration heating is given by:

$$Log(P) = A + 2Log(q) + \frac{B}{T}$$

q is the actual hydrogen concentration





In general in particle accelerators the **gas source is uniformly distributed** along the beam pipe

Pumps located at regular distances from each other give rise to a **parabolic pressure profile** with minima at the pumps location.

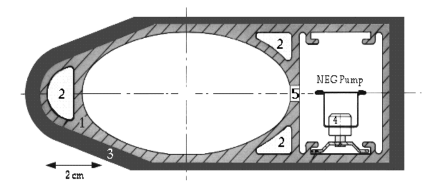
Even for large aperture, the average pressure is quickly limited by the chamber conductance when increasing the spacing between the pumps: the benefit of increasing the lumped pumping speed is minimal.

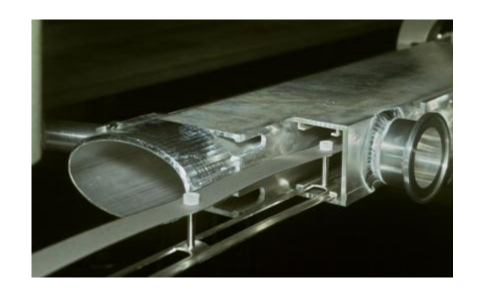
It is more efficient to increase the number of pumps. But this can be very expensive or not possible due to space limitation. Linear pumping is the solution.



Linear pumping may be obtained by NEG ribbons.

LEP dipole vacuum chamber





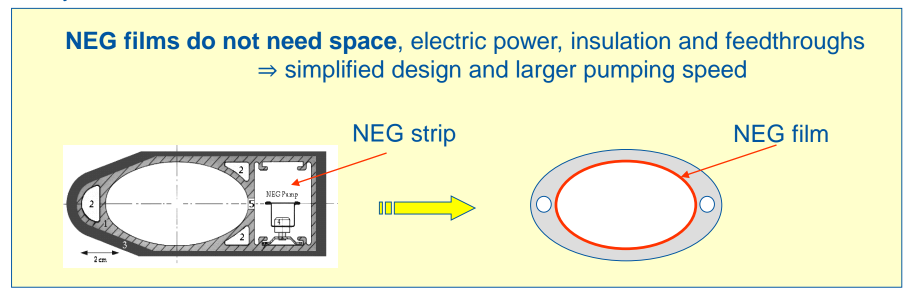
The first application was in the LEP.



The trend in vacuum technology consists in moving the pump progressively closer to the vacuum chamber wall.

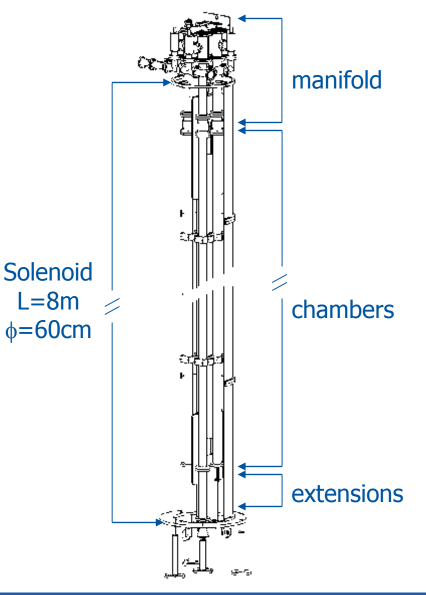
The ultimate step of this process consists of transforming the vacuum chamber from a gas source into a pump.

One way to do this is by "ex-situ" coating the vacuum chamber with a NEG thin film that will be activated during the "in situ" bakeout of the vacuum system.



The coating techniques is **sputtering**.

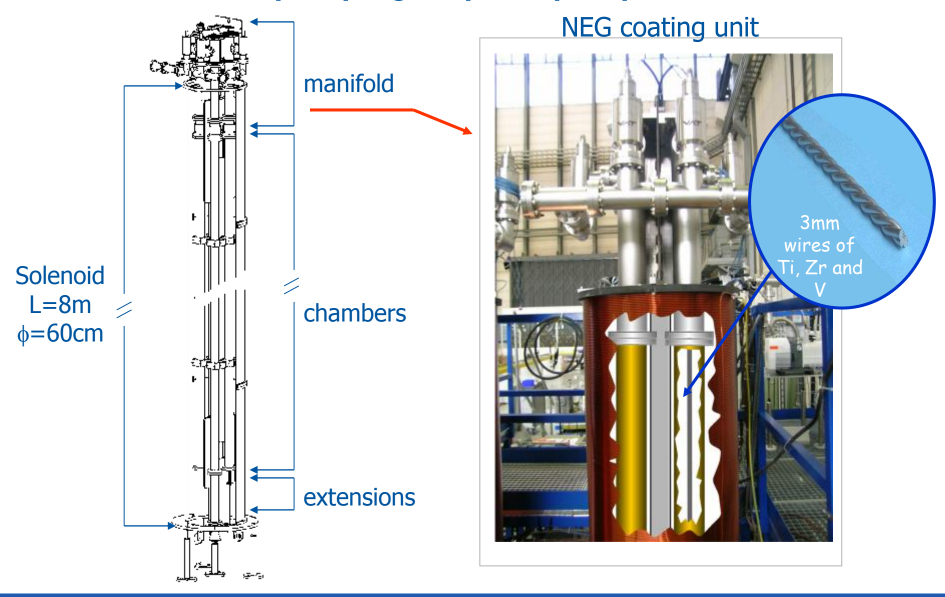




NEG coating unit at CERN











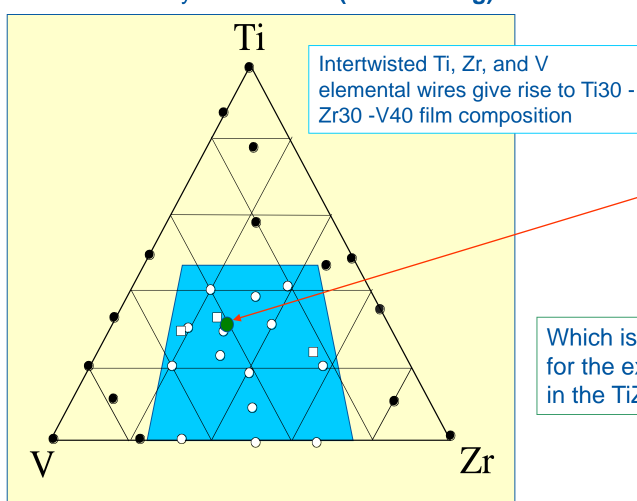


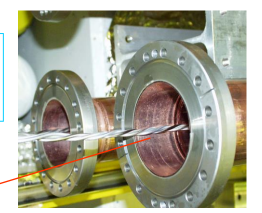






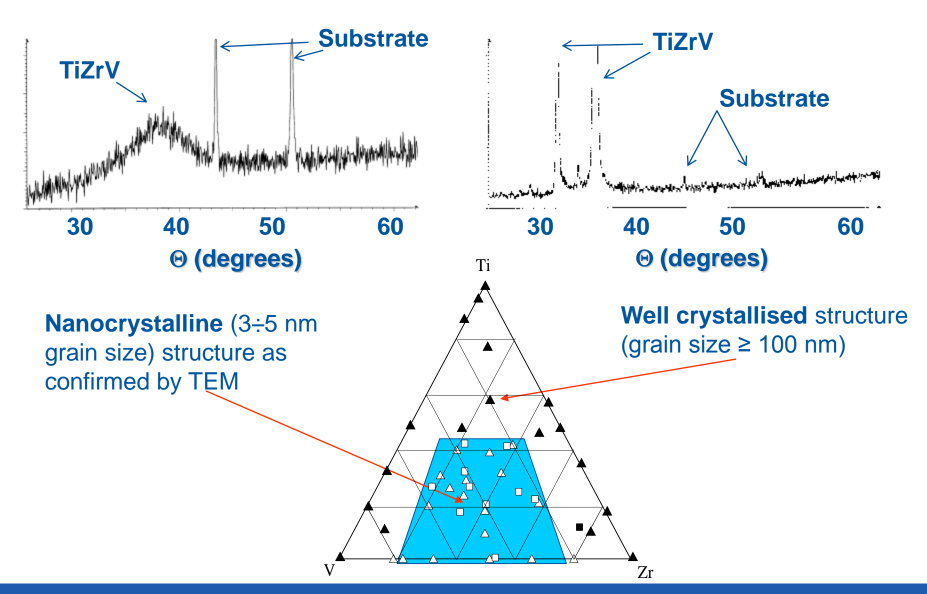
The lowest activation temperature has been found in a wide range of composition in the **Ti-Zr-V** system: **180** °C (**24 h heating**).





Which is the necessary condition for the exceptional fast activation in the TiZrV system?



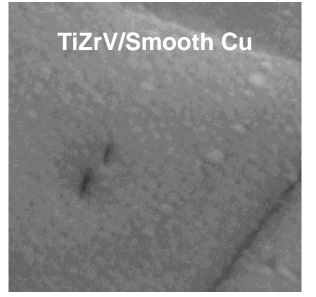


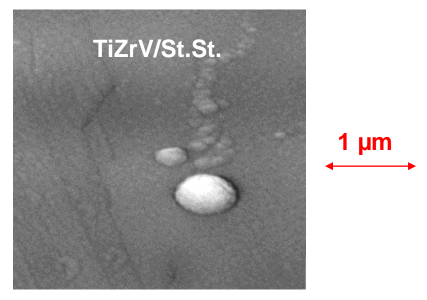


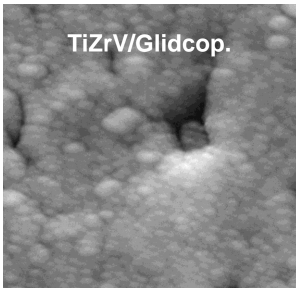
About 1400 chambers for the warm zones of the LHC were coated with a Ti-Zr-V film.

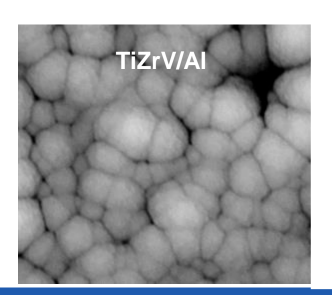




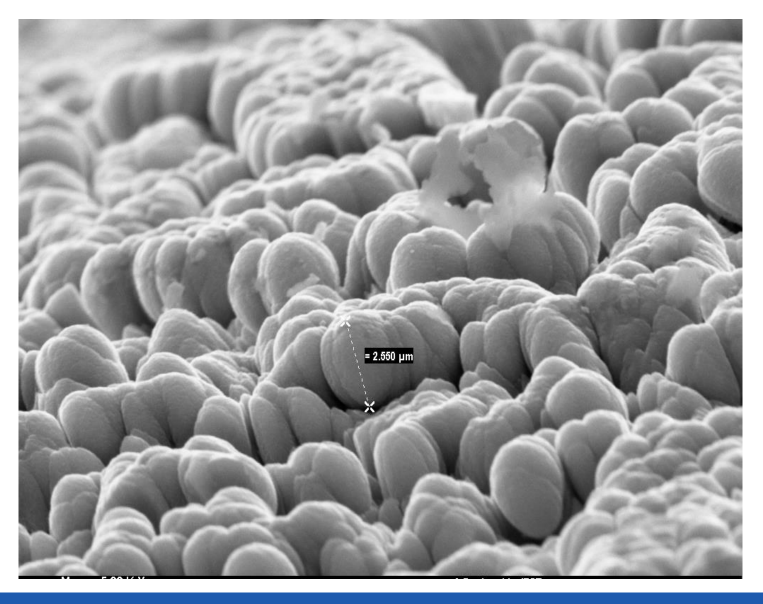




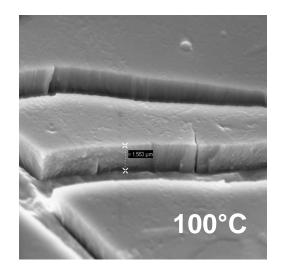


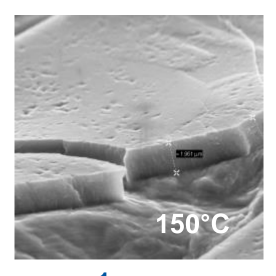


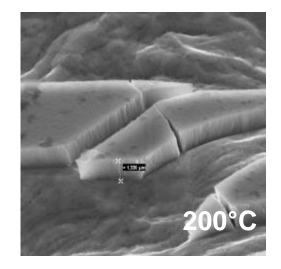


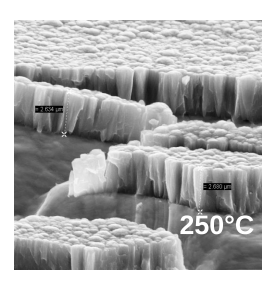


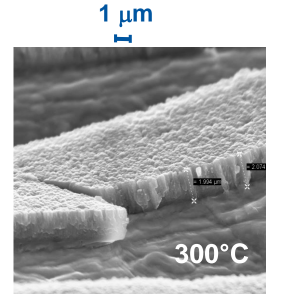


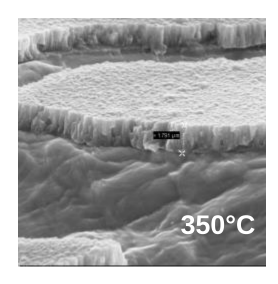












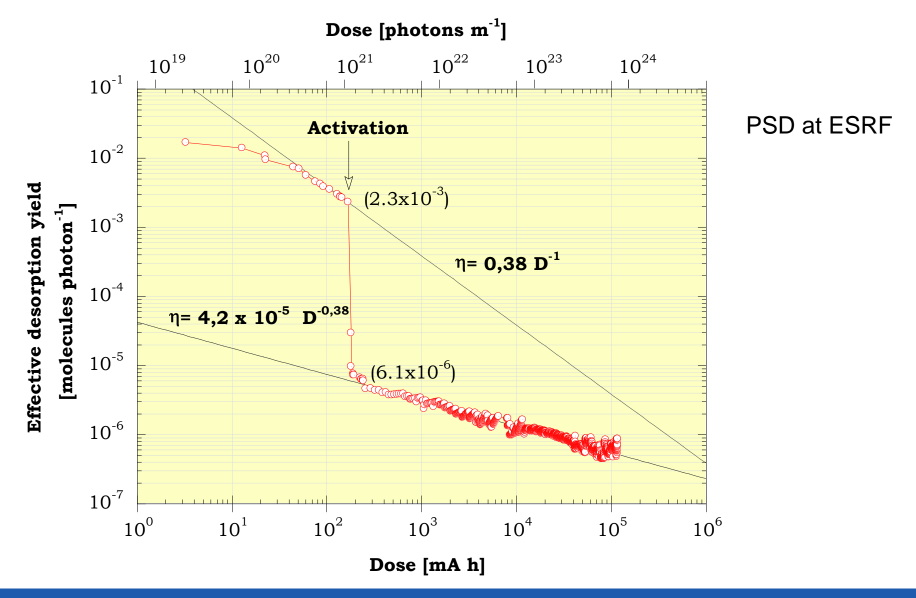


After activation a NEG film surface is very clean resulting in both a large pumping speed and **reduced degassing** (both thermal and ion/radiation/electron induced).

Very important bonus: low secondary electron yield.

NEG films trap the gas coming from the substrate material



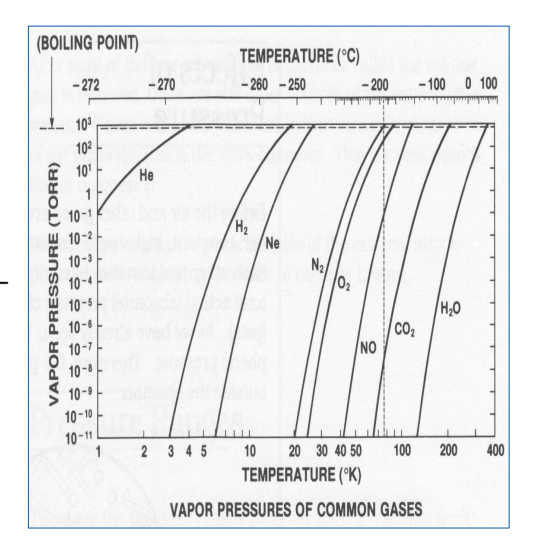


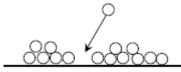


Cryopumps rely on three different pumping mechanisms:

- **1. Cryocondensation**: is based on the mutual attraction of **similar** molecules at low temperature:
 - a. the key property is the **saturated vapour pressure**, i.e. the pressure of the gas phase in equilibrium with the condensate at a given temperature. The attainable pressure is limited by the saturated vapour pressure.
 - b. Only **Ne**, **H**₂ and **He** have saturated vapour pressures higher than 10⁻¹¹ Torr at **20 K**.
 - c. The vapour pressure of H_2 at 4.3 K is in the 10⁻⁷ Torr range, at 1.9 lower than 10^{-12} Torr.
 - d. Large quantity of gas can be cryocondensated (limited only by the thermal properties of the condensate phase)



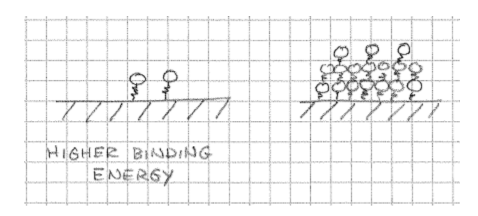




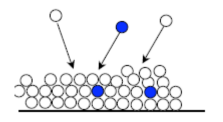


- 2. Cryosorption: is based on the attraction between molecules and substrate. The van der Waals forces are much stronger than those between similar molecules:
 - a) Gas molecules are pumped at pressure much lower than the saturated vapour pressure providing the adsorbed quantity is **lower than one monolayer**.
 - a) Porous materials are used to increase the specific surface area; for charcoal about 1000 m² per gram are normally achieved.
 - b) The important consequence is that significant quantities of H₂ can be pumped at 20 K and He at 4.3 K.
 - **c)** Submonolayer quantities of all gases may be effectively cryosorbed at their own boiling temperature; for example at 77 K all gases except He, H₂ and Ne.





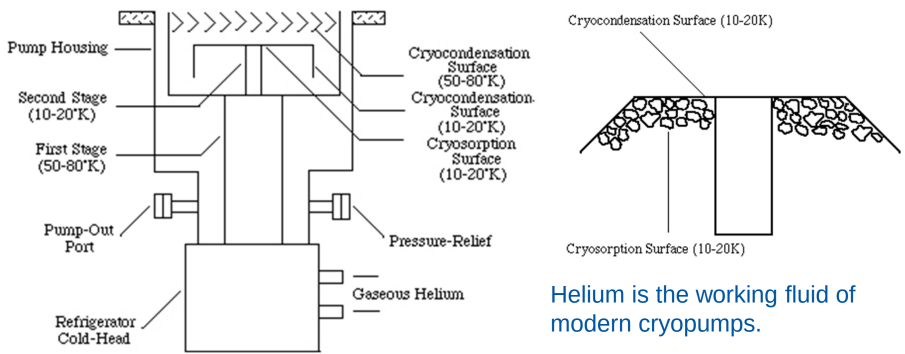
3. Cryotrapping: low boiling point gas molecules are trapped in the layer of an easily condensable gas. The trapped gas has a saturation vapor pressure by several orders of magnitude lower than in the pure condensate. Examples: Ar trapped in CO₂ at 77 K; H₂ trapped in N₂ at 20 K.





Modern cryopumps take advantages of the first two mechanisms.

- 1. The cryocondensation takes place on a cold surfaces, in general at 80 K for H₂O and 10 or 20 K for the other gases.
- 2. The cryosorption of H₂, Ne and He is localised on a hidden surface where a porous material is fixed. This surface is kept away from the reach of the other molecules.

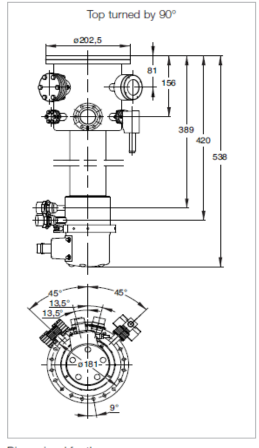




Technical Data

800 BL UHV (CF)

High vacuum flange	DN	160 CF
Fore vacuum flange	40 CF	
Flange for other purpos	16 CF (1x), 40 CF (1x)	
Safety valve with DN 40 connection for gas exh	burst disk mounted on DN 16 CF	
Pumping speed		
H ₂ O	I x s ⁻¹	2600
Ar/N ₂	l x s ⁻¹	640 / 800
H ₂ / He	I x s ⁻¹	1000 / 300
Capacity		
Ar / N ₂	bar x I (Torr x I)	300 (225 000) / 300 (225 000)
H ₂ at 10 ⁻⁶ mbar	bar x I (Torr x I)	4.3 (3225)
He	bar x I (Torr x I)	0.5 (375)
Built-in cold head	7/25	
Max. throughput		
Ar / N ₂ mbar	x l x s ⁻¹ (Torr x l x s ⁻¹)	4 (3) / 4 (3)
H ₂ mbar	x I x s ⁻¹ (Torr x I x s ⁻¹)	2 (1.5)
Crossover value	mbar x I (Torr x I)	150 (112)
Cool down time to 20 k	50	
Overall height	mm (in.)	538 (21.18)
Weight	kg (lbs)	12 (26.5)
Silicon diode for temper at second stage of the o	built-in to a DN 16 CF with 4 way with UHV feedthrough	



Dimensional for the COOLVAC 800 BL UHV (160 CF)

Courtesy of Oerlikon Leybold Vacuum www.oerlikon.com/leyboldvacuum



Characteristics of Cryopumps

1. Starting Pressure

- Cryopumps should be started when the mean free path of molecules is higher than the pump vessel diameter: P<10⁻³ mbar. Otherwise the thermal load is too high.
- In addition a thick condensate layer must be avoided.
- They always need auxiliary pumps.

2. Pumping speed

- **High effective pumping speed** for all gases. Pumping speed from 800 l/s up to 60000 l/s are commercially available .
- Pumping speed for water vapour close to the theoretical maximum.



3. Maximum Gas Intake (Capacity)

- At the maximum gas intake, the initial pumping speed of the gas is reduced by a factor of 2.
- Condensed gases: the limitation is given by the thermal conductivity of the gas layer.
- Adsorbed gases: the capacity depends on the quantity and properties of the sorption agent; it is pressure dependent and generally several orders of magnitude lower compared to that of condensable gases.



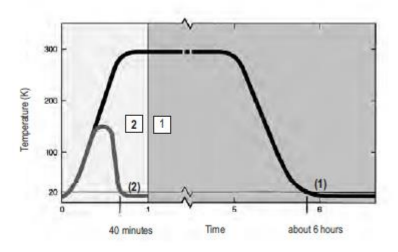
Cryopumps require **periodic regeneration** to evacuate the gas adsorbed or condensed.

To remove all captured gas, **the pump is warmed at room temperature**. The desorbed gas is removed by mechanical pumps (in general, for accelerators, mobile TMP).

During regeneration, the rest of the system must be separated by a valve.

In the majority of application, the performance deterioration is given by the gas adsorbed on the second stage (10-20 K). **A partial regeneration** may be carried out for a shorter time while water vapour is kept on the first stage at temperatures lower than 140 K.





Courtesy of Oerlikon Leybold Vacuum www.oerlikon.com/leyboldvacuum



Gas pumping

	Advantages	Disadvantages
ТМР	 No memory effects Constant pumping speed for pressures lower than 10⁻³ mbar Pumping speed independent of total gas load Starts working at high pressures (molecular regime) 	 Mechanical fragility Risk of contamination from the backing pump Need of venting anytime the pump is stopped Need of valve on the main flange Intrinsic limitation in ultimate pressure of H₂ Possible vibrations Maintenance
SIP	 Clean pumping No maintenance No vibrations Installation in any orientation Relatively long lifetime Relatively low cost Limited but high H₂ capacity The pump current gives a pressure reading 	 Low capture probability Gas Selectivity and limited capacity Memory effects (in particular for rare gases) Ignition in 10⁻⁵ mbar range Bulky Difficult starting for old pumps Production of charged particles in particular at start-up Field emission problems for old pumps Fringing magnetic field Safety issue: high voltage



Gas pumping

	Advantages	Disadvantages
NEG pumps	 Clean vacuum High pumping speed for reactive gases With SIP, extremely low vacuum can be achieve High gas capacity for porous NEG Low cost Electrical power needed only for activation; it works in case of power cut No maintenance No vibration 	 Selective pumping (no pumping of rare gases and methane) H₂ enbrittlement if regeneration is not applied Formation of dust particles is not excluded Safety issue: pyrophoric, it burns when heated in air at high temperature



Gas pumping

	Advantages	Disadvantages
Sublimation Pumps	 Clean vacuum High pumping speed for reactive gases With SIP, extremely low vacuum can be achieve Low cost Electrical power only for sublimation; it works in case of power cut Limited maintenance (filament change) No vibration 	 Very limited capacity Need frequent sublimations at high pressure Ti film peel-off for high sublimation rates Selective pumping (no pumping of rare gases and methane) Risk of leakage current in high voltage insulators Ideal for low pressure applications



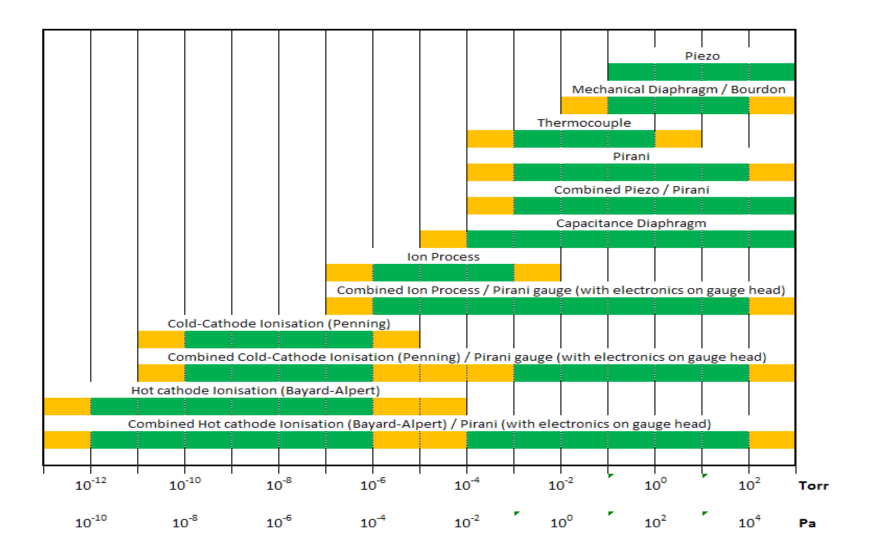
Gas pumping

	Advantages	Disadvantages
Cryopumps	 Very large pumping speed for all gases Clean vacuum High pumping capacity Limited selectivity 	 Cost and maintenance Relatively large volume needed (including refrigerator) Gas release in case of power cut Reduced pumping efficiency for H₂ for high quantity of gas adsorbed: regeneration needed Need of valve on the main flange



Part 6 Instrumentation







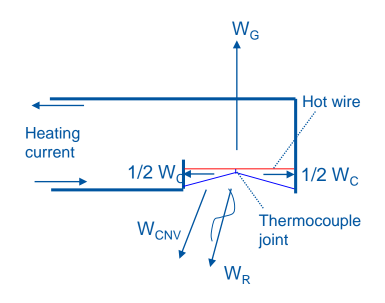
Thermal Conductivity Gauges

Principle of operation:

Pirani gauge: temperature dependent resistance of a hot wire

Thermocouple gauge: temperature dependent voltage from a thermocouple joint

- Four possible mechanisms to take heat away from sensing element:
 - Radiative heat transfer to surroundings
 - Conduction along the sensing element
 - Conduction through the residual gas
 - Convective heat flow to the residual gas





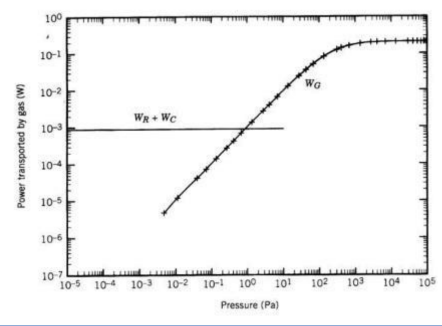
Thermal Conductivity Gauges

Radiative heat transfer: does not depend on pressure

Conduction along the wire: does not depend on pressure

Convection for pressure higher than 100 mbar

Conduction through the residual gas: linear with pressure. Gas properties determine variation in sensor response to pressure





Thermal Conductivity Gauges

Constant current mode:

As pressure decreases, less heat is removed by conduction through the gas from the sensing element, and temperature increases.

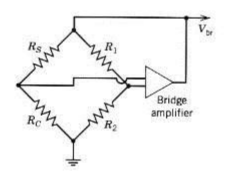
→ Measure temperature dependent change of resistance (or TC voltage).

Constant temperature mode:

Adjust current to heater to maintain constant sensor property (resistance or voltage)

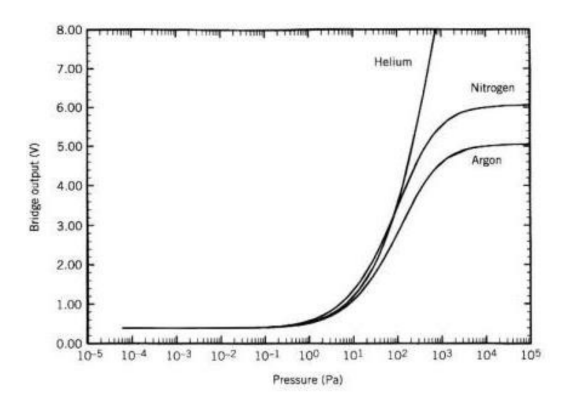
Use Wheatstone bridge arrangement to optimize sensitivity

→Correlate current flow against pressure





Thermal Conductivity Gauges





Ionization Gauges

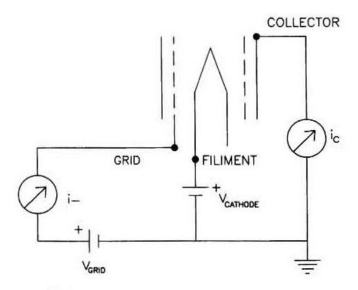
Hot cathode gauge

Thermionic source (electrons) cause inelastic collisions with gas, producing ions lons collected at (-) biased surface Electrons travel to a grid

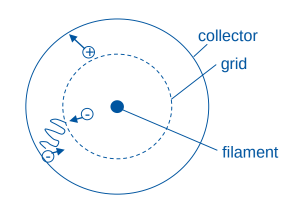
Pre-1950 lower limit of 10⁻⁸ mbar

Soft x-rays produced at grid result in photoelectron flow away from collector (same as ion flow to collector)

Bayard-Alpert modification: fine wire collector surrounded by grid \Rightarrow range extended to 10^{-12} mbar



Basic circuit for operating a hot-cathode ionization gauge



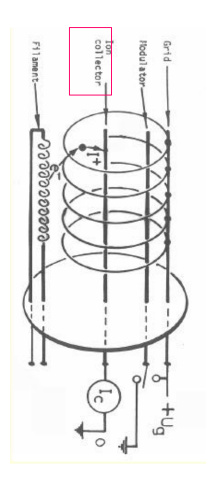


- Bayard-Alpert gauges are used for vacuum measurement purposes in the range 10⁻⁵ -10⁻¹² mbar.
- It is a hot filament ionization gauge. Electrons emitted by the filament perform oscillations inside the grid and ionize the molecules of the residual gas. Ions are then collected by an electrode.

 $I^+ = I^- \sigma n L$

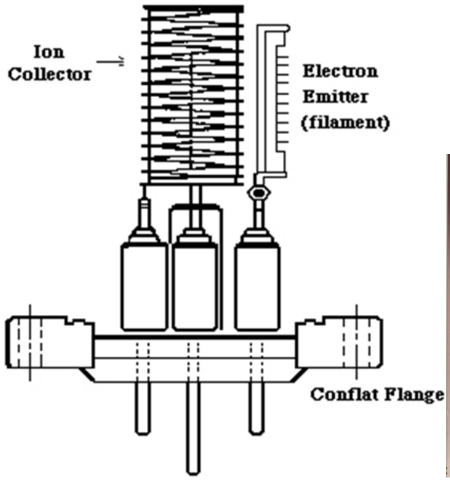
Where:

- I⁺ is the ion current
- I is the filament current
- σ is the ionisation cross section
- n the gas density
- L the electron path lenght
- The gauge needs to be calibrated
- X-ray limit of a few 10⁻¹² mbar

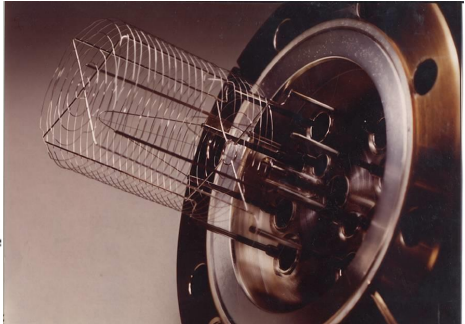




Ionization Gauges



Collisions between electrons, with sufficient energy, and gas molecules results in the ionization of the gas species. The resultant ion current can be measured and correlated with gas pressure.



SVT Bayard Alpert modulated gauge 10⁻¹² Torr



Ionization Gauges

Cold Cathode Gauge: Invented by Penning in 1937 - electrical configuration like diode SIP

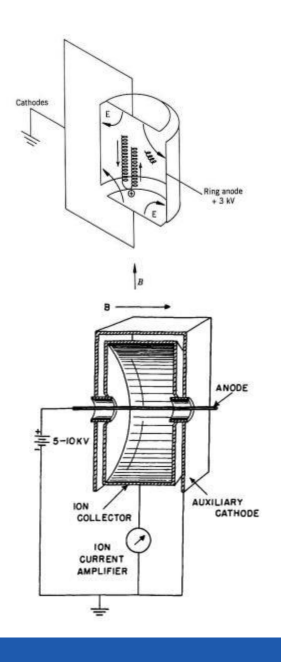
1952 design of wire-in-cylinder with reverse polarity (+ wire) provides key performance improvement

"Inverted Magnetron" - Magnetic field produces long helical path for electrons & ions ⇒ many collisions

Ionizing electrons part of self-sustaining gas discharge

No background current to mask the ion current

Power into CCG ~ 0.1 W





Ionization Gauges

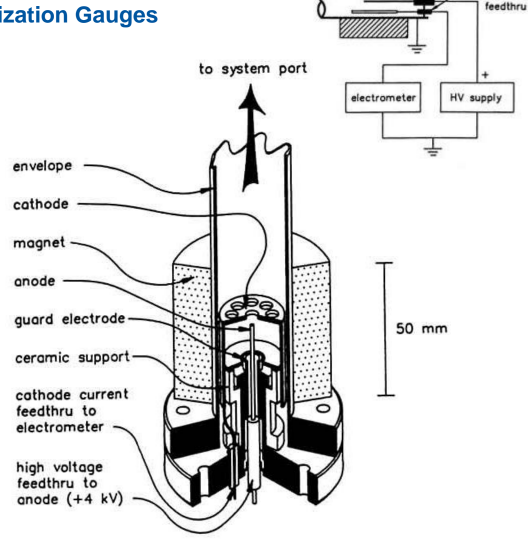
"Striking time" is delay before discharge starts.

At 10^{-4} Pa, $t_s \sim$ seconds; at 10^{-8} Pa, t_s ~ hours or days

Nearly linear dependence of ion current with pressure:

$$i_g = KP^n$$
; $n = 1.05 - 1.2$

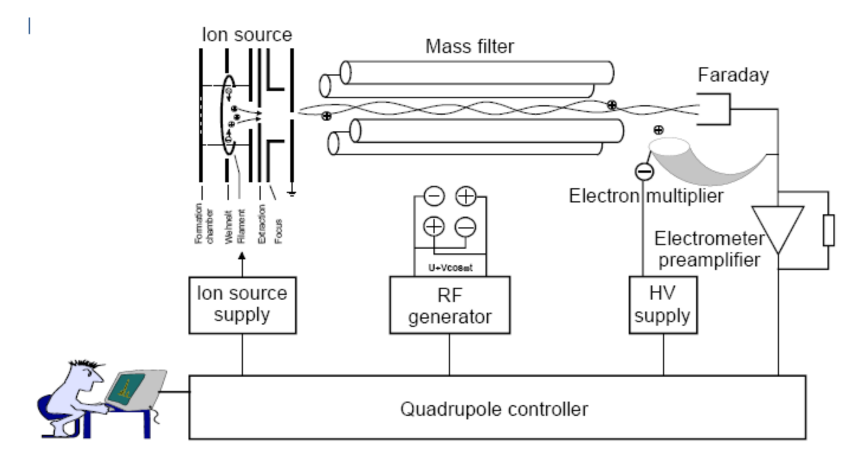
Typical anode voltage +3 kV





cathode

Residual gas analyzers









RGA



