Vacuum Technology for Particle Accelerators

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CERN Accelerator School: Introduction to Accelerator Physics

8th October 2016, Budapest, Hungary

Contents

• What is vacuum and why do we need it in particle accelerators?
• Basics of vacuum technology,
• Gas sources,
• Pumping technology,
• Pressure profile evaluation,
• Outlook of accelerator vacuum systems.
What is vacuum?

Vacuum is a space with no matter inside.

Vacuum in engineering and physics is a space in which the pressure is lower than atmospheric pressure and is measured by its absolute pressure.

Pressure is $\frac{\text{Force [N]}}{\text{Area [m}^2\text{]}}$ measured in Pascals, $1 \text{ [Pa]} = 1 \text{ N/m}^2$

The force exerted on the walls of an evacuated vessel surrounded by atmospheric pressure is: $1 \text{ kg/cm}^2$
What is vacuum?

Conversion table: units of pressure

<table>
<thead>
<tr>
<th></th>
<th>Pa</th>
<th>bar</th>
<th>atm</th>
<th>Torr</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Pa</td>
<td>1</td>
<td>$10^{-5}$</td>
<td>$9.87 \times 10^{-6}$</td>
<td>$7.5 \times 10^{-3}$</td>
</tr>
<tr>
<td>1 bar</td>
<td>$10^2$</td>
<td>1</td>
<td>$0.987$</td>
<td>750.06</td>
</tr>
<tr>
<td>1 atm</td>
<td>$1.013 \times 10^5$</td>
<td>1</td>
<td>1</td>
<td>760</td>
</tr>
<tr>
<td>1 Torr</td>
<td>133.32</td>
<td>$1.33 \times 10^{-3}$</td>
<td>$1.32 \times 10^{-3}$</td>
<td>1</td>
</tr>
</tbody>
</table>

In vacuum technology: mbar or Pa

Why do we need vacuum in particle accelerator?

Less beam-gas interaction:

- Increase beam lifetime,
- Prevents to increase beam size,
- Reduces radiation hazard,
- Lowers background in detectors.

The total beam lifetime in a particle accelerator is given by:

$$\frac{1}{\tau} = \frac{1}{\tau_{\text{elastic}}} + \frac{1}{\tau_{\text{inelastic}}} + \frac{1}{\tau_{\text{Touschek}}} + \frac{1}{\tau_{\text{quantum}}}$$

The interaction between beam particles and residual gas molecules consist of two main mechanisms: elastic and inelastic scattering which contribute to total beam lifetime.
Elastic scattering

Elastic scattering with residual gas molecules alter transverse motion of particles increasing their betatron oscillations (energy of particles is conserved).

- The particles are lost when the oscillation amplitude exceeds physical acceptance aperture.

\[
\frac{1}{\tau_{\text{elastic}}} \sim \frac{Z^2}{\gamma^2} \langle \beta_y \rangle \beta_y \frac{\Delta p}{p} n_g
\]

where:
- \( Z \) - atomic number of the residual gas (careful which gas is dominant)
- \( \langle \beta_y \rangle \) - average vertical beta function
- \( \beta_y \) - vertical beta function at the limiting vertical aperture
- \( \gamma \) - relativistic factor of the particles in the stored beam
- \( a_y \) - limiting vertical aperture
- \( n_g \) - residual gas density

Not only the absolute pressure is important but also what are the gas species in the system.

Inelastic scattering

Inelastic scattering with residual gas molecules decreases speed of particles causing photon emission called Bremsstrahlung (energy of particles is not conserved).

- The particle are lost if the energy loss exceeds the momentum acceptance of the accelerator.

\[
\frac{1}{\tau_{\text{inelastic}}} \sim \frac{Z^2}{\Delta p/p} n_g
\]

where:
- \( Z \) - atomic number of the residual gas (careful which gas is dominant)
- \( \Delta p/p \) - momentum acceptance
- \( n_g \) - residual gas density

Not only the absolute pressure is important but also what are the gas species in the system.
Vacuum Technology for Particle Accelerators, CAS Budapest, Hungary, October 2016
Marek Grabski

Vacuum ranges

1 Atm. = 1013 mbar =~ 1 bar

<table>
<thead>
<tr>
<th>Vacuum Range</th>
<th>Pressure range [mbar]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low Vacuum</td>
<td>$10^3$ - 1</td>
</tr>
<tr>
<td>Medium Vacuum</td>
<td>1 - $10^{-3}$</td>
</tr>
<tr>
<td>High Vacuum (HV)</td>
<td>$10^{-3}$ - $10^{-9}$</td>
</tr>
<tr>
<td>Ultra High Vacuum (UHV)</td>
<td>$10^{-9}$ - $10^{-12}$</td>
</tr>
<tr>
<td>Extreme High Vacuum XHV</td>
<td>&lt; $10^{-12}$</td>
</tr>
</tbody>
</table>

Ideal gas law

Ideal gas equation:

\[ P \cdot V = N_{\text{moles}} \cdot R \cdot T \]  \hspace{1cm} \text{macroscopic}

\[ P \cdot V = N_{\text{molecules}} \cdot k_b \cdot T \]  \hspace{1cm} \text{microscopic}

- \( P \) - the pressure of the gas,
- \( V \) - the volume of the gas,
- \( N_{\text{moles}} \) - the number of moles of the substance,
- \( N_{\text{molecules}} \) - the number of gas molecules,
- \( R \) - the universal gas constant,
- \( T \) - the absolute temperature of the gas,
- \( k_b \) - the Boltzmann constant

1 drop of water contains: $10^{20}$ molecules

Each cubic centimeter of gas at room temperature contains:

\(~10^{19} \text{ molecules/cm}^3\) - at 1 atm (~1 Bar),
\(~10^6 \text{ molecules/cm}^3\) - at $10^{-10}$ mbar (similar pressure as on the moon)
Maxwell-Boltzmann model

Mean speed of gas molecules:

\[
\langle v \rangle = \frac{\sqrt{8k_BT}}{\pi m_0}
\]

\(\varphi\) - impingement rate,
\(n\) - gas density (molecules/volume),
\(k_b\) - the Boltzmann constant \(1.38 \times 10^{-23} [\text{J/K}]\),
\(T\) - the absolute temperature of the gas [K],
\(m_0\) - molecular mass [kg],
\(\langle v \rangle\) - the average gas molecules speed [m/s],

Number of molecules striking unit surface in unit time (Impingement rate):

\[
\varphi = \frac{1}{4}n\langle v \rangle = \frac{1}{4}n \frac{8k_BT}{\pi m_0}
\]

Flow regimes

\[
K_n = \frac{l}{d}
\]

- Mean free path [m]
- Diameter of flow channel [m]
- Knudsen number dimensionless

<table>
<thead>
<tr>
<th>Flow Regime</th>
<th>(K_n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Viscous</td>
<td>(&lt; 0.01)</td>
</tr>
<tr>
<td>Transitional</td>
<td>(0.01 &lt; K_n &lt; 0.5)</td>
</tr>
<tr>
<td>Molecular</td>
<td>(K_n &gt; 0.5)</td>
</tr>
</tbody>
</table>

Mean free path:
- At atm. Pressure = 6.5x10^{-8} m
- At 10^{-9} mbar (storage ring) = 66 km

Increasing pressure:
- Typically: \(> 1 \text{ mbar}\)
- \(< 10^{-3} \text{ mbar}\)
Flow regimes

Particle accelerators operate in molecular flow regime.

Gas flow in molecular regime

In molecular flow regime the gas flow \( Q \) from one point to the other is proportional to the pressure drop:

\[
Q = C(P_1 - P_2)
\]

\[
[Q] = \frac{mbar \cdot l}{s}
\]

Conductance depends on the gas molecule velocity thus its molar mass and temperature (not on pressure).

\[C = \frac{1}{4} A \langle v \rangle = C' A \left[ \frac{l}{s} \right]\]

\[C' = \frac{1}{4} \langle v \rangle \left[ \frac{l}{s \ cm^2} \right]\]

\[\langle v \rangle\] - the average gas molecules speed [m/s]

Conductance \( C \) at 295 K for nitrogen (\( N_2 \) - molecular mass 28):

\[C = C' A = 11.8 A \left[ \frac{l}{s \ cm^2} \right]\]
Gas flow in molecular regime

Combination of conductances:

a). For components 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 steady state \( Q_1 = Q_2 \) \( \Rightarrow C_{TOT} = \frac{C_1 C_2}{C_1 + C_2} \) and \( \frac{1}{C_{TOT}} = \frac{1}{C_1} + \frac{1}{C_2} \) \( \Rightarrow \frac{1}{C_{TOT}} = \sum_1^N \frac{1}{C_i} \)

b). In parallel:

\[ Q_1 = C_1 (P_1 - P_2) \]
\[ Q_2 = C_2 (P_1 - P_2) \]
\[ Q_{TOT} = C_{TOT} (P_1 - P_2) \]

\[ Q_{TOT} = Q_1 + Q_2 \Rightarrow C_{TOT} = C_1 + C_2 \Rightarrow C_{TOT} = \sum_1^N C_i \]

Gas flow in molecular regime

In vacuum technology a pump is an object that permanently removes gas molecules from the gas phase.

Pumping speed \( S \) of a pump is defined as the ratio between the pump throughput \( Q_p \) and the pressure \( P \) at the entrance to the pump.

\[ S = \frac{Q_p}{P} \quad \text{[l/s]} \]

Gas removal rate can be written as:

\[ Q_p = \varphi A_p \alpha = \frac{1}{4} A_p n \langle v \rangle \alpha = A_p C' P \alpha \]

From the definition of pumping speed:

\[ S = A_p C' \alpha \]

\( \varphi \) - impingement rate
\( A_p \) – is the area of the pump aperture [cm²]
\( C' \) – is the conductance of the unit surface area for given gas
\( n \) – gas density
\( \alpha \) – is the capture probability
Gas flow in molecular regime

Introduced limitation between pump and pumped vacuum volume limits the nominal pumping speed of chosen pump.

\[
\frac{1}{S_{\text{eff}}} = \frac{1}{S} + \frac{1}{C}
\]

Example:

300 l/s ion pump connected to a vacuum chamber through a 90 deg elbow of given dimensions can result in ~30 \% pumping speed reduction.

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Generic vacuum system

\[
P = \frac{Q}{S_{\text{eff}}}
\]

- \( P \) – gas pressure,
- \( Q \) – gas load (outgassing),
- \( S_{\text{eff}} \) – Effective pumping speed.
Sources of gases

Sources of static gas loads in vacuum system:

Vacuum chambers are source of gas

Processes \( \rightarrow Q \)

What process defines pressure

What process defines the pressure over time?

Outgassing:
- Material
  - Binding energy
- Surface condition
  - As delivered
  - Machined
  - Polished
  - Cleaning
  - Heat treatment…

Diffusion:
- Material
- Heat treatment
  (Vacuum firing)
- Inner surface barrier
  (Air baking, Film deposition)

Thermal outgassing

Thermal outgassing (static outgassing)

For metals:

- If not baked (heated to 200°C) in-situ water is the dominant gas specie.

\[ q_{H_2O} \approx \frac{3 \times 10^{-9}}{l} \left( \text{mbar l} \right) \left( \text{s cm}^2 \right) \]

- If baked (heated to ~200°C) in-situ hydrogen H\(_2\) is the dominant gas.

Outgassing rates \( q \left( \text{torr l} \right) \left( \text{s cm}^2 \right) \) at 20°C:

<table>
<thead>
<tr>
<th>Material</th>
<th>Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Austenitic stainless steel not baked,</td>
<td>3x10^{-10} (H(_2)O)</td>
</tr>
<tr>
<td>after 10 h pumping</td>
<td></td>
</tr>
<tr>
<td>Austenitic stainless steel baked in-situ</td>
<td>2x10^{-12} (H(_2))</td>
</tr>
<tr>
<td>for 24 h at 150°C</td>
<td></td>
</tr>
<tr>
<td>OFS copper baked in-situ for 24 h at</td>
<td>~10^{-14} (H(_2))</td>
</tr>
<tr>
<td>200°C</td>
<td></td>
</tr>
</tbody>
</table>

Dynamic outgassing

In particle accelerators energized particles impinging on vacuum surfaces induce desorption of molecules. Usually such dynamic gas load dominate over thermal outgassing.

Beam stimulated desorption is characterised by \( \eta \) - the desorption yield:

\[ \eta = \frac{\text{number of desorbed molecules}}{\text{number of particle impinging the surface}} \]

\( \eta \) – depends on many parameters:

- incident particle: type and energy,
- material,
- surface roughness,
- cleanliness of the surface,
- history of the material (dose),
- Particle flux.

The desorption may be stimulated by:

- electrons,
- ions,
- synchrotron radiation (photons).
When charged particles (moving at relativistic speeds) are accelerated they emit synchrotron radiation in a narrow cone. This photon flux impinging on vacuum surfaces produces strong outgassing thus a large dynamic pressure increase.

**Photon Stimulated Desorption**

Total photon flux $\dot{\Gamma}$ [photons/s] around electron storage ring:

$$\dot{\Gamma} = 8.08 \times 10^{17} IE \quad \rightarrow \quad \frac{\text{photons}}{s}$$

$I$ – machine current [mA]

$E$ – energy [GeV]

Gas flow $Q_{PSD}$ due to photon induced desorption:

$$q_{PSD} = \eta \dot{\Gamma} \quad \rightarrow \quad \frac{\text{molecules}}{s}$$

$\eta$ – desorption yield

$$Q_{PSD} = K\eta \dot{\Gamma} \quad \rightarrow \quad \frac{\text{mbar} \times \text{l}}{s}$$

$K$ – converts number of molecules to pressure units $4.04 \times 10^{-20}$ [mbar×l/molecule]

Knowing the gas load (outgassing) due to PSD ($Q_{PSD}$), thermal outgassing ($Q_{thermal}$) and the target pressure ($P$) the effective pumping speed $S_{eff}$ can be calculated.

$$S_{eff} = \frac{Q_{PSD} + Q_{thermal}}{P}$$
Photon Stimulated Desorption

Evaluating Photon Stimulated Desorption (PSD):

- **PSD yield effect of the dose:**
  \[ \eta = \eta_0 D^{-\alpha} \]
- \( \alpha \) is between 0.6 and ~1

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Vacuum scrubbing

Vacuum conditioning:

Average pressure normalized to machine current vs accumulated beam dose (or photon dose)

- Dynamic pressure is proportional to current:
  \[ P \propto I \]
- Dynamic pressure rise:
  \[ \frac{\Delta P}{I} \]
Pump classification

- **Vacuum Pumps (molecular regime)**
  - **Momentum transfer pumps**
    - Example: Turbomolecular Pump
    - Principle: Molecules impinge on fast moving surfaces which direct them towards the pump outlet where they are evacuated by pumps operating in viscous flow. The molecules do not transfer energy to each other.
  - **Capture pumps**
    - Example: Sputter Ion Pump, Getter pump, Cryo pump
    - Principle: gas molecules are fixed to a surface inside vacuum (pump has no moving parts).

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**Turbomolecular Pump**

(Momentum transfer pump)

- **Pressure range:** $10^{-1}$ till $10^{-10}$ mbar, (with backing pump connected in series).
- **Usual operational pressure:** $< 10^{-5}$ mbar.
- **Outlet to primary pump**
- **Inlet**

- **Outlet**

**Blade rotational speed:** 1000 – 1500 Hz

**$S$ (pumping speed)** does not depend significantly on the mass of the molecule.

$$K_o = \frac{P_{\text{outlet}}}{P_{\text{inlet}}} \text{(compression ratio)}$$

$K_o$ depends exponentially on the wall speed and square root of the gas molecule mass.
Turbomolecular Pump

Turbo molecular and roughing pump connected in series:
from 1 bar (atmospheric pressure) until ~10^{-10} mbar

Turbo Molecular Pump (range: 10^{-1} to 10^{-10} mbar)

Turbomolecular pumping speeds:
10 l/s - 25,000 l/s.

Turbomolecular pumps are widely used in particle accelerators for:
• evacuating vacuum systems from atmospheric to ultra high vacuum,
• Testing (leak tests),
• Conditioning (bakeouts),
• High gas loads,
• For accelerator operation with beam capture pumps take over,

Usually they are not permanent part of the vacuum system (attached when needed).

Capture pumps: getters

Getter materials adsorb gas molecules on their surface which is contamination and native oxide layer free. Such surface can be produced in two ways:

• sublimating the reactive meta in situ: evaporable getters or sublimation pumps,
• dissolving the surface contamination into the bulk of the getter material by heating: non-evaporable getters (NEG); the dissolution process is called activation.

Getter surface is characterized by the sticking probability ‘α’:

\[
\alpha = \frac{\text{number of molecules captured}}{\text{number of molecules impinging}}
\]

Getter pumping speed (S):

\[
S = \alpha A_\text{getter} C'
\]

Where:
- \(A_\text{getter}\) surface area of active getter surface,
- \(C'\) conductance for given gas of unit surface area.

Getter materials do not pump noble gases and methane (CH\(_4\)) at room temperature. Therefore, they need auxiliary pumping to keep a stable pressure.
Evaporable Getters

Evaporable getters: TSP – Titanium Sublimation Pump

Ti is the **sublimated** metal. Ti filaments are heated up to 1500°C reaching Ti vapour pressure which is deposited on the surrounding surfaces creating a chemically active surface where gas molecules are captured.

When the deposited film is saturated, a sublimation is needed to recover the initial pumping speed. A single filament with hundreds of sublimation cycles

\[
\text{Sticking probabilities:} \\
H_2: 0.01 \leq \alpha \leq 0.1 \\
CO: 0.5 \leq \alpha \leq 1
\]

Film capacity:
- For CO, one monolayer adsorbed,
- For O₂ several monolayers,
- For N₂ fraction of monolayer

Hydrogen diffuses in the Ti film $\rightarrow$ much higher capacity

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Non-Evaporable Getters (NEG)

On activation the oxide layer at the surface of NEG is diffused to the bulk of the material creating clean, chemically active surface where gas molecules are captured.
Non-Evaporable Getters (NEG)

NEG materials are produced industrially by powder technology. The powder is sintered to form discs or strips.

A typical alloy produced by SAES Getter is **St707** made of Zr (70%), V (25%), Fe (5%).

The high porosity of NEG materials allows pumping of relatively high quantities of gas without reactivation. After 40 venting cycles (with nitrogen) and reactivation 80% pumping speed is conserved.

Full pumping speed is obtained after heating at 450°C for 45’ or 350°C for 24h.

Distributed pumping

- Pressure profile with lumped pumps
- Pressure profile with distributed pumps
Distributed pumping can be obtained by installing NEG strips (antechamber needed)

**Distributed pumping**

*LEP dipole vacuum chamber (first application of NEG strip for distributed pumping)*

**NEG coatings**

NEG-coating transforms a vacuum chamber from a gas source to a vacuum pump.

NEG coating transforms a vacuum chamber from a gas source to a vacuum pump. The technology of coating vacuum chambers by magnetron sputtering was developed at CERN for the warm sections of LHC. Nowadays it is also widely applied in synchrotron radiation sources.

NEG film characteristics:
- Film composition: Ti (30%), Zr (40%), V (30%).
- Thickness ~1 µm,
- Activation temperature 200°C for 24 h,
- Low PSD (Photon stimulated desorption),
- Sticking probability similar to TSP.

Disadvantage of NEG: has limited capacity and activation cycles.
NEG coatings

Photon stimulated desorption (PSD) measurements at ESRF (beamline D31).

*‘Synchrotron Radiation-Induced Desorption from a NEG-Coated Vacuum Chamber’*, P. Chiggiato, R. Kersevan

\[ \eta = 4.2 \times 10^{-5} \ D^{0.38} \]

\[ \eta = 0.38 \ D^{-1} \]

CERN chamber, 316LN, 2 m, Ø60mm

\[ \text{Linear Photons Dose (ph/m)} \]

\[ \text{Accumulated Dose (mA h)} \]

Sputter ion pumps

- Voltage 3 to 7 kV,
- Cathodes are plates of Ti at ground potential,
- Magnetic field generated by external permanent magnets ~0.1 T.

*Agilent ion pump catalogue*
Sputter ion pumps

Pumping mechanisms:

- Chemical adsorption onto the reactive metal layer (Ti) deposited on anode and subsequent burial by additional metallic atoms of gas molecules: all gases except rare gases,
- Implantation of gas ions in the cathode (not permanent), and of energetic neutrals bounced back from the cathode in the deposited film: only mechanism of pumping for rare (noble) gases,
- Diffusion into the cathode and the deposited film: only H₂.

Diode configuration (cell cross-section)

Effective pumping speed:

Pumping speed vs pressure for a diode ion pump with nominal pumping speed 100 l/s (for air after saturation)

Nominal (catalogue) pumping speed 100 l/s at ~1e-6 mbar

L. Schulz, Sputter ion pumps, CAS 1999 proceedings
**Pressure measurement**: Ion pumps can be used for pressure measurement up to $10^{-10}$ mbar if the voltage is reduced to 3 kV. The current is proportional to pressure.

L. Schulz, *Sputter ion pumps*, CAS 1999 proceedings

**Lifetime**: cathode is sputtered away (eroded) by impacting ions. If operating at high pressures ($10^{-4}$ mbar) the pump lifetime is 400 h whereas at $10^{-6}$ it is 40000 h (4.5 year).
Sputter ion pumps

Wide variety of ion pumps to choose:
- Electrode material and configuration: Diode, noble diode, triode,
- Pumping speeds: from 0.2 l/s (weight 0.6 kg) till 500 l/s and more (260 kg)

Nominal pumping speed normalized to that of air for diode and triode ion pump:

<table>
<thead>
<tr>
<th>Gas</th>
<th>Air</th>
<th>N₂</th>
<th>O₂</th>
<th>H₂</th>
<th>CO</th>
<th>CO₂</th>
<th>H₂O</th>
<th>CH₄</th>
<th>Ar</th>
<th>He</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diode</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1.5-2</td>
<td>0.9</td>
<td>0.9</td>
<td>0.8</td>
<td>0.6-1</td>
<td>0.03</td>
<td>0.1</td>
</tr>
<tr>
<td>Triode</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1.5-2</td>
<td>0.9</td>
<td>0.9</td>
<td>0.8</td>
<td>0.6-1</td>
<td>0.25</td>
<td>0.3</td>
</tr>
</tbody>
</table>

Advantages of ion pumps: clean, bakeable, vibration free (no moving parts), continuous operation, wide operating range $10^{-4}$ to $10^{-11}$ mbar, low power consumption, long lifetime at low pressure.

Pressure profile simulation

To ensure that the pressure distribution in an accelerator is satisfactory, the pressure profile must be evaluated. For this purpose simulations programs based on test particle Monte-Carlo are widely used (eg.: Molflow+).

Component
- CAD model of surfaces exposed to vacuum
  - Imported to Molflow+
- Boundary conditions defined
  - Set: desorption, pumping surfaces (sticking probability), opacity.
  - Particle trajectories in green.

https://test-molflow.web.cern.ch/content/about-molflow
Pressure profile simulation

CAD model with highlighted absorber (Abs) locations:

Arbitrary cross-sections can be modelled:

Vacuum model in Molflow+
with highlighted pumps:

• Photon flux [photons/s] for each surface intercepting the beam (absorbers) is calculated,
• Photon Stimulated Desorption (PSD) yield [molecules/photon] for assumed accumulated
dose in [Ah] or [photons/m] is estimated from published data:
• Having the PSD yield and photon flux impinging on each surface, local outgassing [mbar*l/s]
can be calculated for each irradiated area that is the input to the simulation.

To evaluated static pressure due to thermal outgassing the total surface area is multiplied by the
outgassing rate for chosen material.

To calculate dynamic pressure profile due to Photon Stimulated Desorption (PSD):
• Photon flux [photons/s] for each surface intercepting the beam (absorbers) is calculated,
Partial pressure measurement

Residual gas analyzer assembly

Sensor head exposed to vacuum

Residual gas analyzer – (mass spectrometers) used to monitor the quality of vacuum i.e. which gas species are present in the system.

Quadrupole mass filter:
- Ions entering the quadrupole field experience potential differences deflecting them from their original trajectory.
- The extent of deflection of ions is related to its mass to charge (m/e or m/z) ratio.
- At each instance only one m/e ratio resonates with the field allowing the ion to pass along its axis.
- All other species are deflected and neutralised by impact on the quadrupole rods.

G. J. Peter et al., 'Partial pressure gauges', Vacuum in Accelerators, CAS 2006 proceedings

Partial pressure measurement

Residual gas spectrums of an UHV system:

at total pressure 4x10^-9 mbar

![Graph showing residual gas spectrum at 4x10^-9 mbar]

at total pressure 4x10^-11 mbar

![Graph showing residual gas spectrum at 4x10^-11 mbar]

Paolo Chiggiato, Vacuum Technology for particle accelerators, 2013
Vacuum systems in accelerators

Vacuum systems are constructed of many components joined together with flanges.

Vacuum systems are divided in sectors by gate valves.

The pressure is monitored by vacuum gauges.

Bayard-Alpert hot cathode ionization gauge
Connection between UHV components is made by copper gaskets that are squeezed between two flanges.

RF shielded bellows are essential to ensure low impedance and allow vacuum chamber compression and expansion. Beam diagnostic devices are often integral part of vacuum system (example: beam position monitor.)
Thank you for your attention

Acknowledgments:
Paolo Chiggiato (CERN), Esra Al-Dmour (MAX IV),

Some references:

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The physical basis of ultra-high vacuum, P.A. Redhead, J.P. Hobson, E.V. Kornelsen. AVS.