Design and Fabrication of Prototype Internal 80 K helium gas purification system for helium plant

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ABSTRACT

The Helium Refrigerator/Liquefier (HRL) plant is normally operated with helium gas having purity better than 99.999 % by volume which is equivalent to having 10 PPM (parts per million) impure gas in the helium stream. In the process of gas transfer or due to some other processes before reaching to cold box of helium plant, impurity level sometimes can reach to as high as 500 PPM averaging to about 100 PPM. These impurities consist of mainly gases present in the air, hydrocarbons and H2 and traces of Ne. These gases condense at significantly higher temperature compared to the LHe temperature (4.5 K). If such high level of impurity enters the process equipment placed inside the cold box of the HRL, then it can condense and choke the pipe lines and valves leading to large pressure drop and inefficient liquefaction process. Some time, condensed and frozen impurity can destroy the blades of turbines of HRL. Hence, to be on safer side generally, internal purifiers are placed at two temperature levels (at 80 K and 20 K) inside the cold box to take care the operational problem due to impurities. Purifier at ~80 K removes Oxygen, Nitrogen and Argon from helium gas. This project is about the design and fabrication of the 80 K prototype purification system with its associated elements. Activated charcoals will be used to adsorb impure gases from cold helium gas at ~80 K and ~14 bar. The nominal flow rate of helium gas is ~120 g/s for the actual size purifier bed. Appropriate helium mass flow rate will be decided for the prototype adsorber bed. The pressure drop across the bed and filters will be measured. The break through curve or mass transfer zone (MTZ) data will be collected during this experiment and analyzed.

Keywords: Helium Purification, Adsorber bed

1. INTRODUCTION

Institute for Plasma Research (IPR) is the place where efforts are being made in one of the most challenging and necessary tasks of this century; “controlling nuclear fusion”. This institute in its experimental activity has embarked on an ambitious project of building the first Indian Steady State Superconducting (SST-1) Tokamak. The helium refrigerator/liquefier (HRL) of 1.3 kW capacity for the Steady State Superconducting Tokamak (SST-1) is successfully operating independently along with integrated flow control and distribution system.

The Helium Refrigerator/Liquefier (HRL) is normally operated with helium gas having purity better than 99.999 % by volume which is equivalent to having 10 PPM (parts per million) impure gas in the helium stream. Although sufficient precautions and impurity removal procedures are used, still, in the process of gas transfer or due to some other processes before reaching to liquefaction, impurity level sometimes can go as high as 100 PPM.

These impurities consist of mainly gases present in the air, like N2, O2, Ar, H2O, CO, CO2, H2 and traces of Ne. These gases condense at significantly higher
temperature compared to the LHe temperature (4.5 K). If such high level of impurity enters the process equipment placed inside the cold box of the HRL, then it can condense and choke the pipe lines and valves leading to large pressure drop and inefficient liquefaction process. Some time, condensed and frozen impurity can destroy the blades of turbines of HRL. Hence, to be on safer side generally, internal purifiers are placed at two temperature levels inside the cold box to take care the operational problem due to impurities. One is at ~80 K to remove nitrogen, oxygen and other gases having normal boiling points above 77 K. Another one is at ~20 K to remove hydrogen and neon gases.

II. SEPERATION AND PURIFICATION SYSTEM

To produce LHe, there must be used the pure helium. So, to produce that purity the helium separation & purification system can be used. As a rule, the partition of gases at cryogenic temperatures is the most efficient separation system for all; hence, industrially gas separation frameworks have been all around grown over the past half century. Practically, all the commercials produced by following separation and purification systems:

i. Rectification
ii. Adsorption
iii. Refrigeration purification

III. ADSORPTION & TYPES OF ADSORPTION [1]:

Whatever the nature of the forces holding a solid together, it can be regarded as producing a field of force around each ion, atom or molecule. At the surface of the solid, these forces cannot suddenly disappear and thus reach out in space beyond the surface of the solid. Due to these unsaturated and unbalanced forces, the solid has a tendency to attract and retain on its surface molecules and ions of other substances with which it comes in contact. Thus when a gas is brought in contact with solid, generally a part of the gas is taken up by the solid. If the gas molecules penetrate into the solid, the process is usually called absorption. If the gas molecules stick to the surface of the solid in one or more layers, the process is called adsorption.

If there is a very strong reaction between the solid and the gas, the adsorption process is called the chemical adsorption. If only weak intermolecular forces, called van der waals forces are brought into play, the process is called physical adsorption.

As physical adsorption has low heat of adsorption and it is rapid and reversible than chemical adsorption it will be a preferable choice for us. Physical Adsorption is also very significant at low temperature like 80K and 20K which will also beneficial to us.

IV. ADSORBENT

In physical adsorption the substance attached to the surface is called adsorbate, and the substance to which it is attached is known as adsorbent.

Desire properties of adsorbent,

- High internal volume – Which is accessible to the component removed from the fluid.
- Good mechanical properties - strength and resistance to attrition
- Good kinetic properties - capable of transferring adsorbing molecules rapidly to the adsorption sites.
- Regeneration can be carry out effectively.
- Raw material and production should be inexpensive.
- High internal surface area

Some of the adsorbent are,

- Activated carbon
- Carbon molecular sieve
- Silica gel
- Activated alumina
- Zeolite

Fig 2. General Structure of an Adsorbent Particle
V. ADSORPTION ISOTHERM

If a quantity \( q \) of a gas or vapour is adsorbed by a porous solid at constant temperature and the steady state equilibrium partial pressure is \( p \) then the function \( q(p) \) is the adsorption isotherm.

Adsorption isotherms indicate the manners by which the different adsorbate get adsorbed on the adsorbents. There are mainly six types of Adsorption Isotherms:

1. Microporous materials (e.g. Zeolite and Activated carbon) – monolayer formation
2. Non porous materials (e.g. Nonporous Alumina and Silica) – multilayer formation. It can be seen that after the point B formation of multilayer starts.
3. Non porous materials and materials which have the weak interaction between the adsorbate and adsorbent (e.g. Graphite/water) – adsorbent interactions are comparatively weak so weak bond form.
4. Mesoporous materials (e.g. Mesoporous Alumina and Silica) - The most distinguishing feature of the type IV isotherm is the hysteresis loop because of pure condensation.
5. Porous materials and materials that have the weak interaction between the adsorbate and adsorbent (e.g. Activated carbon/water) – weak interaction as in type III.
6. Homogeneous surface materials (e.g. Graphite/Kr and NaCl/Kr) – It is a special case, which represents stepwise multilayer adsorption on a uniform, non-porous surface, particularly by spherically symmetrical, non-polar adsorptive.

VI. ISOTHERM THEORY MODELS

1. **Langmuir Model**

   This model is portrayed by Irving Langmuir (1916) for gasses adsorbed into solids. It is utilized to evaluate the measure of adsorbate adsorbed into the adsorbent is a component of fractional weight at that temperature.

   The equation of the model is given by
   
   \[
   W = W_{\text{max}} \left( \frac{K_c}{1 + K_c} \right)
   \]

   Where
   - \( W \) = adsorbate filling / loading
   - \( K_c \) = Adsorption constant
   - \( C \) = concentration of fluid
   
2. **BET Model**

   This model is intended by Brunauer et al. (1938). This is in light of the multilayer adsorption model. It accept that there is meet vitality of adsorption for every layer aside from the first layer.
The equation of this model is given by

\[ \frac{V}{M_a} = \frac{V_m Z P}{P_{sat}} \left( 1 - \frac{P}{P_{sat}} \right) \left( 1 + (Z - 1) \frac{P}{P_{sat}} \right) \]

Where
- \( V \) = volume of gas adsorbed
- \( M_a \) = mass of adsorbent
- \( V_m \) = volume of gas per unit mass of adsorbent required to form a monomolecular layer over the entire adsorbent surface
- \( P \) = partial pressure of gas being adsorbed
- \( P_{sat} \) = saturation pressure of the gas being adsorbed at the temperature of the adsorbent
- \( Z \) = it is a function of energy of adsorption and temperature of adsorbent
- \( Z = \exp \left( \frac{\theta_a}{T} \right) \)

The values of \( \theta_a \) and \( V_m \) are different for various gases and adsorbents.

3. Polanyi Potential Theory \(^5\):

The main disadvantage in Langmuir model is the guess of monomolecular adsorption. And BET theory multilayer adsorption is based on the assumption of higher pressure ratio. Polanyi Potential Theory is founded on the multi molecular pore filling which is appropriate theory for adsorption on highly porous resources like activated carbon and zeolites.

This volume adsorbed (\( W \)) is a function of adsorption potential (\( e \)), \( W = f(e) \).

The adsorption potential is equal to work essential to remove one molecule from its location in adsorbed phase to gas phase and can be associated to pressure. The adsorption potential characterises the molar free energy change with the change in vapour pressure from that over pure liquid phase \( P_0 \) to equilibrium pressure \( P \) at a given coverage of adsorbent surface.

The mathematical formula can be written as

\[ e = RT \ln \left( \frac{P_0}{P} \right) \]

![Fig.4 Monolayer & Multilayer Adsorption Phenomenon \(^4\)]

4. Dubinin-Radushkevitch (D-R) Equation \(^6\):

The Dubinin-Radushkevitch (1947) calculation is the adaptation of the Polanyi potential model. The D-R equation highpoints on the micro pore volume filling of the adsorbent material rather than layer formation on the wall of the adsorbent material. According to this model, adsorbent materials like activated carbon, silica gel, zeolites contain varieties of pore sizes but it is the micropore volume that controls the amount of adsorption.

Another parameter \( \theta \) is presented such a way that

\[ \theta = \frac{W}{W_0} \]

Where,
- \( W_0 \) = total volume of the micropore
- \( W \) = volume that has been filled when the relative pressure is \( P/P_0 \)

Here in this equation \( \theta \) is a function of adsorption potential \( e \) in such a manner that

\[ \theta = f \left( \frac{e}{E_0} \right) \]

Where, \( E = E_0 \beta \)
- \( E \) = characteristic adsorption energy
- \( E_0 \) = Adsorption energy of the reference vapor
- \( \beta \) = affinity coefficients

Assuming the pore size distribution is Gaussian, Dubinin and Radushkevitch found out the expression

\[ \frac{W}{W_0} = \exp \left[ -\frac{E}{E_0} \right]^2 \]
Where $\varepsilon = Rln \left( \frac{E}{E_0} \right)$

$E = E_0 \beta$,

Dubinin-Astakhov (D-A) equation is widespread form of D-R equation, is articulated as [7]

$$\frac{W}{W_0} = \exp \left[ -\frac{E}{E_n} \right]$$

Here, $n$ = Heterogeneity factor depends upon the surface.

- $n = 2$ for Carbonaceous materials.
- $n = 4$ to 6 for Zeolites.

VII. BREAKTHROUGH CURVE & MASS TRANSFER ZONE (MTZ) [4]:

In very few beds the outline of the beds can be assumed internally, though these profiles can be forecast and used in calculating the curve of concentration v/s time for fluid leaving the bed. When the breakthrough of the adsorbate begins to happen it is essential to take the bed offline so that the adsorbent can be regenerated.

Breakthrough time, $t_b$ is the time when the $C/Co$ value becomes (proposed outlet impurity/Inlet impurity) & saturated time, $t_b$ is the time when $C/Co$ value becomes 1.

At any prompt in time in the adsorption, it is clear from fig.5 that the adsorbent particles upstream and downstream of the mass transfer zone (MTZ) don’t contribute in mass transfer process. Upstream of the MTZ, the adsorbent will be in equilibrium with the feed and unable to adsorb further adsorbate molecules. Downstream of the MTZ, the adsorbent will not have been in contact with any adsorbate molecules and therefore, despite having the capability of doing so.

**Fig 5. Concentration profile, mass transfer zone and break through curve**

VIII. BREAKTHROUGH CURVE PREDICTION [3]:

Mathematical model for prediction of mass transfer is based in following assumption,

- Gas behaves like an ideal gas.
- Isothermal adsorption.
- Mass and velocity gradients are negligible in radial direction of the bed.
- Bed is tube-like and the axial dispersion is considered in the bulk phase.
- Particles of the charcoal are spherical and they are packed regularly into the fixed bed.

1. Axial Dispersion Model:

The axial dispersion model is based on the concept that the concentration front is dispersed by both hydrodynamic and kinetic (finite transport rates) factors.

The equation for the axial dispersion model is given by
\[ \frac{C}{C_0} = \frac{1}{2} \left[ 1 - \text{erf} \left( \frac{1 - \frac{t}{t'}}{\sqrt{2(D_z t/ue)}} \right) \right] \]

\[ t' = \frac{L}{u} \left[ 1 + \frac{K_H}{\varepsilon} \right] \]

\[ D_z = D_e \left( \frac{\gamma_1 + \gamma_2}{\varepsilon} \right) \]

\[ \gamma_1 = 0.45 + 0.55(\varepsilon) \]

\[ \gamma_2 = 0.5 \left( 1 + \frac{13\gamma_1 \varepsilon}{R_e S_e} \right) \]

Where, \( t \) = time

\( C \) = Outlet concentration

\( C_0 \) = Inlet concentration

\( D_z \) = Axial dispersion coefficient

\( u \) = interstitial velocity

\( L \) = length of total bed

\( K_H \) = dimensionless Henry constant

\( \varepsilon \) = void porosity

\( D_e \) = effective diffusivity

\( D_m \) = molecular diffusivity

\( D_k \) = Knudsen’s Diffusivity

\( S_c = \nu / D_e \)

2. Rosen Model:

The Rosen Model, developed by Rosen assistant in finding analytical results to adsorption difficulties.

The basic assumptions for this model are as below [5]

1. No Axial dispersion
2. Isothermal conditions and linear isotherm
3. Constant flow velocity & Constant effective diffusivity

\[ \frac{C}{C_0} = \frac{1}{2} \left[ 1 + \text{erf} \left( \frac{3U/2V - 1}{2\left(1 + 5\nu\right)^\frac{1}{2}} \right) \right] \]

\[ V = \frac{D_e K}{R_p k} \]

\[ V = \frac{3D_e K L}{u R_p^2 \left( \frac{\varepsilon}{1 - \varepsilon} \right)} \]

\[ U = \frac{2D_e \left( t - \frac{L}{u} \right)}{R_p^2} \]

Where, \( C \) = Outlet concentration

\( C_0 \) = Inlet concentration

\( U \) = dimensionless contact parameter

\( V \) = dimensionless bed length parameter

\( v \) = dimensionless film resistance

\( R_p \) = pellet radius

\( K_H \) = Henry constant

IX. DESIGN OF ADSORBER BED

Design of adsorber bed can carry out in following way-

1) Selection of adsorbent
2) Find mass of Adsorbent required
3) Calculate the length and diameter of adsorber bed
4) Calculate breakthrough time and mass transfer zone (MTZ)
5) Optimize the length and diameter of adsorber bed on different parameter like MTZ, breakthrough time, pressure drop, velocity, mass flow rate etc.

<table>
<thead>
<tr>
<th>Adsorbent (Carbon)</th>
<th>Shape of particles</th>
<th>Internal porosity %</th>
<th>Bulk density, Kg/L</th>
<th>Surface area, Km²/kg</th>
<th>Sorptive capacity, kg/kg (dry)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shell based</td>
<td>G</td>
<td>60</td>
<td>0.45-0.55</td>
<td>0.8-1.6</td>
<td>0.4</td>
</tr>
<tr>
<td>Wood-based</td>
<td>G</td>
<td>~80</td>
<td>0.25-0.30</td>
<td>0.8-1.8</td>
<td>~0.7</td>
</tr>
<tr>
<td>Petroleum-based</td>
<td>G,C</td>
<td>~80</td>
<td>0.45-0.55</td>
<td>0.9-1.3</td>
<td>0.3-0.4</td>
</tr>
<tr>
<td>Peat-based</td>
<td>G,C,P</td>
<td>~55</td>
<td>0.3-0.5</td>
<td>0.8-1.6</td>
<td>0.5</td>
</tr>
<tr>
<td>Lignite based</td>
<td>G,P</td>
<td>70-85</td>
<td>0.4-0.7</td>
<td>0.4-0.7</td>
<td>0.3</td>
</tr>
<tr>
<td>Bituminous coal-based</td>
<td>G,P</td>
<td>60-80</td>
<td>0.4-0.6</td>
<td>0.9-1.2</td>
<td>0.4</td>
</tr>
<tr>
<td>Carbon molecular sieve</td>
<td>C</td>
<td>35-50</td>
<td>0.5-0.7</td>
<td></td>
<td>0.05-0.20</td>
</tr>
</tbody>
</table>

6) Design the filter

1. Selection of adsorbent

Table shows Physical Property of Adsorbents (Courtesy: Perry’s Chemical Handbook)

From the studies of different research paper and books we can say that the coconut shell based charcoal is most optimised adsorber as per our operating condition.

2. Find the mass of adsorber

We can select the appropriate calculation model from the relative pressure. As relative pressure of impurity is fall in range of 10⁻⁴, we will go with D-R model for the calculation of mass of adsorber.

\[ n_i = \frac{W_o}{\sum X_i V_{mi} \exp \left[ \frac{-BT^2}{(\sum X_i \beta_i)^2} \left( \sum X_i \ln \left( \frac{P_{adi}}{P_i} \right) \right)^2 \right]} \]

For mass flow rate of 1 g/s, operation period of 3 hour and impurity of 100 ppm the required mass of adsorber is around 250 gram.

3. Calculate the length and diameter of adsorber bed

There is a pressure drop in the helium flow across the bed because of the adsorbent material, carbon SC11 in this case. First thing we keep in mind while designing the bed is that the pressure drop across the bed should not be more than 30 mbar.

Fig 7. Pressure drop vs diameter of bed
The graph for pressure drop across the bed with various bed diameter is shown. So the diameter of bed should be more than 1 inch for pressure drop less than 30 mbar. Initially taking diameter of bed 1 inch (0.0254 m). Length can be calculated as follow,

Now, \( v = \frac{L}{D^2} \) and taking \( D = 0.0508 \) m 
\( L = 0.90 \) m 
\( L/D = 35.43 \)

Which we will be optimized further.

4. Calculate MTZ length and breakthrough time

From the axial dispersion or from the Rosen model as we discussed earlier we can find the breakthrough curve and so the mass transfer zone. We can also find the breakthrough time and saturation time for our operation.

![Rosen model](image1)

![Axial dispersion model](image2)

Rosen and Axial dispersion model are shown in the graph. Mass transfer zone for Rosen and Axial dispersion model are 0.65 and 0.50 respectively. Breakthrough time for Rosen and Axial dispersion model are 3.6 and 5.3 hour respectively.

We will consider Rosen model for our calculation because breakthrough time for Rosen is less which we can take as a safer side and diameter of our pipe is less so no axial dispersion will take place, which is basic assumption of Rosen model.

5. Optimization

![Effect of different mass flow rate on breakthrough curve](image3)

![Effect of particle diameter on breakthrough curve](image4)

![Pressure Drop variation with Superficial Velocity](image5)

Fig 8. Rosen model

Fig 9. Axial dispersion model

Fig 10. Effect of different mass flow rate on breakthrough curve

Fig 11. Effect of particle diameter on breakthrough curve
Effect of different parameters like mass flow rate, particle diameter,

As in fig 10 as we move from left to right mass flow rate is decrease. As mass flow rate decrease the impurities to be adsorbed are decrease and breakthrough time increase.

As in fig 11, as diameter of particle increases the mass transfer zone increases, because as the particle diameter increase the surface area available per unit volume is decrease. So the adsorption area decrease and the length of adsorption will increase. So mass transfer zone will increase.

As in fig 12, as we move from left to right void fraction decrease so the adsorbent become more dense so more impurity can adsorb. So breakthrough time increases. But it also leads to more pressure drop.

As from fig 12, 13 and 14 we can see the effect of different parameter like mass flow rate, flow velocity and L/D ratio on Pressure drop. These are used to optimize the parameters.

6. Dimension of vessel and vacuum jacket:

Since the pressurized helium gas has to be passed through the adsorber vessel, it can be designed considering a pressure vessel. According to the ASME Section VIII, the minimum thickness of the cylindrical shell can be given by:

\[ t = \frac{pD}{2S_a e_w - 1.2p} \]

Where

- \( p \) = design internal pressure,
- \( D \) = inside diameter of shell,
- \( S_a \) = allowable stress,
- \( e_w \) = weld efficiency.

The internal design pressure is 20 bar = 2.0 MPa.

Considering the material to be SS304L, so, \( S_a = 120.6 \) MPa,

Taking weld efficiency \( e_w = 1 \)

\( t = 0.25 \) mm. For the safe side and market availability we take pipe of NSP-1 sch. 10 with outer diameter 33.40 mm, inner diameter 27.86 mm and,

Thickness of vessel is 2.77 mm

The minimum thickness for spherical shells, hemispherical heads is given by:

\[ t_h = \frac{pDK}{2S_a e_w - 0.2p} \]

Where, \( K = \text{constant} = 1/6 \left[ 2 + \left( \frac{D}{D_1} \right)^2 \right] \)

We have, hemispherical head thickness, \( t_h = 0.124 \) mm

Head thickness \( t_h = 0.280 \) inch

We will use 3 inch pipe for vacuum jacket which will vacuum by \( 10^{-4} \) mbar by rotary pump.

Total heat transfer in vacuum jacket is 14.53W. We can use multilayer insulation for better performance.

7. Design of filter:

Filter is required to separate fine charcoal particle of adsorber from the main helium stream. So the design of filter is also essential.
Consider that $D_p$ is the particle diameter and $w$ is the aperture of the mesh. Now from Pythagoras theorem we have,

$$w = \frac{D_p}{\sqrt{2}}$$

For the present work, the charcoal dust that is carried with helium has particle size ranging from 30 microns. So by using equation,

$$w = \frac{30}{\sqrt{2}} = 21.21 \text{ micron}$$

Fig 16. Mesh opening

In 1” we have 25400 microns, so

Mesh = \frac{25400}{51} = 498\text{~}500.

Mesh size = 500

<table>
<thead>
<tr>
<th>Filter</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Operating Pressure</td>
<td>14</td>
<td>bar</td>
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<tr>
<td>Operating Temperature</td>
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<td>K</td>
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<tr>
<td>Density of He</td>
<td>8.2196 Kg/m³</td>
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<tr>
<td>Mass flow rate, m</td>
<td>1</td>
<td>g/s</td>
</tr>
<tr>
<td>Vol.Flow rate, Q</td>
<td>1.22E-4 m³/s</td>
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</tr>
<tr>
<td>Operation time</td>
<td>3</td>
<td>Hr</td>
</tr>
<tr>
<td>Charcoal particle Dia,$D_p$</td>
<td>30</td>
<td>microns</td>
</tr>
<tr>
<td>Aperture Size, Width, $w$</td>
<td>21</td>
<td>microns</td>
</tr>
<tr>
<td>Diameter of wire, $d$</td>
<td>30</td>
<td>microns</td>
</tr>
<tr>
<td>Pitch of the Mesh, $P$</td>
<td>51</td>
<td>microns</td>
</tr>
<tr>
<td>~Mesh</td>
<td>550</td>
<td></td>
</tr>
<tr>
<td>ΔP Across Wire Screen</td>
<td></td>
<td></td>
</tr>
<tr>
<td>He flow rate, Q</td>
<td>1.22E-4 m³/s</td>
<td></td>
</tr>
<tr>
<td>Length, $L$</td>
<td>0.04</td>
<td>m</td>
</tr>
<tr>
<td>Diameter of Screen, $D_i$</td>
<td>0.03</td>
<td>m</td>
</tr>
</tbody>
</table>

Pressure Drop, ΔP

| Across Inlet Pipe                         |           |           |
| D                                        | 0.0254    | m         |
| Pre. Drop mbar                           | 0.0023    | mbar      |

Across Outlet Pipe

| D                                        | 0.0254    | m         |
| A                                        | 5.06E-4 m² |
| Pre. Drop mbar                           | 0.0023    | mbar      |

Across Annular Space Area

| Do                                       | 0.0580    | m         |
| Aannular                                 | 0.00131   | m²        |
| Uannular                                 | 0.00924   | m/s       |
| Pre.Drop                                 | 0.035     | Pa        |
| Pre.Drop mbar                            | 0.0003    | mbar      |
| Total P Drop                             | 3.375     | Mbar      |

8. Modeling :
9. Component of test facility

X. Acknowledgment

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