# **Cryogenic Engineering**

# Chapter 4.

# **Separation and Purification Systems**

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※ reversible mixing/unmixing – semipermeable membrane (free passage for one gas)



Reversible isothermal work

$$-\frac{W_i}{m} = T_m(s_1 - s_2) - (h_1 - h_2)$$

1 – Before separation2 – After separation



$$\begin{aligned} & -\frac{W_{i}}{m_{m}} = -\frac{W_{ia}}{m_{a}}\frac{m_{a}}{m_{m}} - \frac{W_{ib}}{m_{b}}\frac{m_{b}}{m_{m}} \\ & = \{T_{m}(s_{1a} - s_{2a}) - (h_{1a} - h_{2a})\}\left(\frac{m_{a}}{m_{m}}\right) + \{T_{m}(s_{1b} - s_{2b}) - (h_{1b} - h_{2b})\}\left(\frac{m_{b}}{m_{m}}\right) \end{aligned}$$

Ideal work of separation for an ideal gas mixture

For an ideal gas, 
$$pV = mRT$$
  
 $h = C_pT$   
 $s = C_plnT - Rlnp$ 

$$\therefore -\frac{W_i}{m_m} = T_m R_a \left( ln \frac{P_m}{P_{1a}} \right) \frac{m_a}{m_m} + T_m R_b \left( ln \frac{P_m}{P_{1b}} \right) \frac{m_b}{m_m}$$

$$\frac{P_{\rm m}}{P_{\rm 1a}} = \frac{m_{\rm m}R_{\rm m}T_{\rm m}/V_{\rm m}}{m_{\rm a}R_{\rm a}T_{\rm m}/V_{\rm m}} = \frac{m_{\rm m}R_{\rm m}}{m_{\rm a}R_{\rm a}}$$

Since 
$$mR = n\overline{R} = m\overline{R}/M$$

$$\frac{P_{m}}{P_{1a}} = \frac{n_{m}}{n_{a}} = \frac{1}{y_{a}}$$
  $y_{a}$ : mole fraction of gas A

Work requirement per unit mole of mixture

$$-\frac{W_{i}}{n_{m}} = \overline{R}T_{m}\left(y_{a}ln\frac{1}{y_{a}}+y_{b}ln\frac{1}{y_{b}}\right)$$
$$= \overline{R}T_{m}\sum_{j}y_{j}ln\frac{1}{y_{j}}$$

#### Ideal work of separation of gas mixtures

Table 4.1. Ideal work of separation of gas mixtures. The mixture is assumed to be separated into two parts: the desired gas, which has the given mole fraction in the original mixture, and the remainder of the gas mixture. The mixture is to be separated at 300 K (80°F)

		Work per Mix	r Mole of ture	Work per Unit Mass of Separated Gas	
Gas	Mole Fraction	J/mol	Btu/lbmole	J/g	Btu/lb <sub>m</sub>
Nitrogen	0.78084	1312	564	60.0	25.8
Oxygen	0.20946	1280	550	191.0	82.1
Argon	0.00934	132.1	56.8	353.9	152.2
Carbon dioxide	0.00033	7.42	3.19	511.0	219.7
Neon	$1.818 \times 10^{-5}$	0.540	0.232	1472.5	633.1
Krypton	$1.14 \times 10^{-6}$	0.0404	0.0174	438.1	188.4
Xenon	$8.60 \times 10^{-8}$	0.00370	0.001593	328.1	141.0
Hydrogen deuteride	$3.12 \times 10^{-4}$	7.06	3.035	7483	3217
Deuterium	$1.56 \times 10^{-4}$	3.80	1.634	6090	2618
Helium-3	$1.00 \times 10^{-7}$	0.00427	0.00184	14 222	6114

 For a single substance in a single phase, we need only 2 independent thermodynamic properties are required to Specify thermodynamic state

#### Gibbs phase rule

The phase rule is

Where

- f = degree of freedom
- c = number of components
- p = number of phases

At equilibrium for a system of any composition.

- Gibbs phase rule
  - Degree of freedom (f) = It is the number of external variables that can be changed independently without disturbing the number of phases in equilibrium. These are pressure, temperature and composition.
  - **Component** = A chemically independent constituent of a system.
  - Number of components (c) = The minimum number of independent species necessary to define the composition of all the phase present in the system.
  - Phase (p) = A state of matter that is uniform throughout in chemical composition and physical state.

Typical temperature-composition diagrams for binary mixtures



 (a) Typical curve for pressures less than the critical pressure for both components



(b)Typical curve for pressures less than the critical pressure for one component, but above the critical pressure for the other component

Azeotrope

not boil change

Azeotrope is a particular mixture (two or more miscible liquids) which does not change the composition during the distillation.



(c) Azeotropic mixture



Figure 6-16 Atkins Physical Chemistry, Eighth Edition © 2006 Peter Atkins and Julio de Paula

#### Water-alchol distillation

- An-dong soju(안동소주) : 23~45% alcohol
- Won-soju (원소주) : 22% alcohol





Distillation apparatus





The basics of distilling alcohol





#### How Alcohol distillation works

#### How a pot still works

- 1. Mixture with alcohol.
- Vapors from the boiling alcohol after heating to 173ºF.
- The captured vapors pass through coiled copper tubing which is submerged in cold water.
- The coiled tubing and cold water cool the vapors back to a liquid form.
- The resulting liquid is much higher in alcohol content than the previous mixture.



Multistage Distillation Principle



Stage	The beginning	The end	
1	11% (base alcohol, for example wine)	32%	
2	32%	55%	
3	55%	70%	
4	70%	78%	
5	78%	83%	

Two different shapes of temperature-composition diagrams

As shown below, in case of the narrow diagram, a substance cannot be purified quickly, so more steps are required to get purer substance in liquid composition



# **4.3 Enthalpy-Composition Diagrams**



 Another set of thermodynamic curves important in the study of mixtures

Fig. Enthalpy-composition for nitrogen-oxygen mixtures at 101.3 kPa

### **4.3 Enthalpy-Composition Diagrams**



Fig. Enthalpy-composition diagrams

### **4.3 Enthalpy-Composition Diagrams**

Mole Fraction N <sub>2</sub>	101.3 kPa (1 atm)			506.6 kPa(5atm)				
	Temp. (K)		h		Temp. (K)		L	
	Liquid	Vapor	J/mol	J/mol	Liquid	Vapor	/J/mol)	. (J/mol)
0.00	90.2	90.2	419	7252	108.9	108.9	1315	7536
0.10	87.7	89.5	461	7231	106.3	107.9	1403	7507
0.20	85.7	88.7	519	7210	104.2	106.7	1499	7478
0.30	84.1	87.7	599	7185	102.5	105.6	1591	7448
0.40	82.5	86.7	682	7151	100.9	104.3	1675	7415
0.50	81.3	85.6	779	7118	99.7	103.0	1758	7377
0.60	80.4	84.3	879	7084	98.5	101.5	1851	7339
0.70	79.6	83.1	984	7042	97.4	100.0	1947	7298
0.80	78.8	81.5	1084	6992	96.5	98.4	2052	7252
0.90	78.1	79.7	1181	6933	95.6	96.6	2152	7201
1.00	77.4	77.4	1273	6871	94.2	94.2	2248	7147

Table. Enthalpy of nitrogen-oxygen mixtures. h= enthalpy of the saturated liquid at the bubble point; H= enthalpy of the saturated vapor at the dew point

Rectification is synonymous with countercurrent distillation. This special distillation separation process enables the individual components of a mixture to be separated with a high purity combined with a good yield, even when their boiling points are relatively close to each other.



Distillation column in a cryogenic Air separation plant



- Consider two mixtures **AB** and **CD** at 1 atm whose temperature-composition diagrams are as shown.
  - Gas A + Gas B
  - Gas C + Gas D
- The respective boiling points of each gas are as shown.
- The boiling point difference of the mixture **AB** is less than that of the mixture **CD**.



- Let us consider a mixture with composition at state 1 as shown.
- If this mixture is rectified on **AB** diagram at 2, we have

 $y_{liq} = 2f_{AB}$ 

Again, if this mixture is rectified on
 CD diagram, we have

 $y_{liq} = 2f_{CD}$ 



- The separation is more effective when the difference in the boiling points is more.
- For such mixtures, almost pure product is obtained in one or two condensations.



- As done earlier, rectification of this mixture at point 3.
  - Vapor is enriched in low boiling point component  $(N_2)$
  - Liquid is enriched in high boiling point component (O<sub>2</sub>)
- This process forms the fundamental step for the rectification column.



- As mentioned earlier, gas separation is a process of repeated rectification.
- The equipment which carries out these processes is called as a Rectification column.
- The figure shows the schematic of a Rectification column.



- It is vertical column which is closed by spherical domes, both at the top and at the bottom.
- Theses are spherical in shape in order to minimize surface area(less heat in-leak) and accommodate high pressures(1~5 atm)
- The column is well insulated because, it is usually operated at cryogenic temperatures.



- The top dome houses a **Condenser** and the bottom done houses a **Boiler**.
- The two phase mixture is first expanded isenthalpically. It can be liquid or liquid + vapor or vapor.
- This expanded product is introduced into the column as **Feed**.



- The rectification of the mixture occurs across each
  Plate and Downcomer assembly as shown in the figure.
- These **Plates** have **holes** for the vapor mixture to pass through and ultimately reach the condenser.
- Similarly, the **Downcomer** takes the mixture in liquid phase towards the boiler.



#### Rectification Column



• During this motion, vapor and liquid flow in opposite directions and exchange heat in a counter-flow manner.

• Hence, there is a temperature gradient across the length of the column.



- The low boiling component(LBC) is condensed in the **Condenser** and the high boiling component(HBC) is evaporated in the **Boiler**.
- The low and high boiling components are collected at the top and bottom respectively.



- For the ease of understanding, consider a mixture of N<sub>2</sub> and O<sub>2</sub> at 1 atm.
- Let the Feed in saturated vapor condition enter the column at 1, as shown in figure.
- Let us assume a steady state and an ideal operation of column.
- When the mixture condenses or evaporates, its composition remains constant.



- Hence for any plate, the vapor rising and the liquid on the plate have same molar composition, although the liquid is at lower temperature.
- Hence, the liquid on the plate above the feed inlet(point 2) has same composition as the vapor.
- Therefore, extending a constant composition line about point1, we have point 2.



- Also, the vapor leaving point 3 is in thermal equilibrium with liquid at 2. (same plate)
- Extending an isotherm about the point2, we have the point 3.



- Similarly, liquid at point 4 has same composition as point3.
- Extending a constant composition line about 3, we have point 4



- On the other hand, we have the points 1, 2' and 3' as shown in the figure.
- The point 1 and 2' are in thermal equilibrium(same plate).
- Also, liquid at point 2' has same composition as point 3' and liquid is at lower temperature.



- Therefore, the liquid moving down is enriched in high boiling point component( $O_2$ ).
- On the other hand, the vapor moving up is enriched in low boiling point component(N<sub>2</sub>).


- In order to keep the process running, some heat is supplied to the boiler, to continuously evaporate the high boiling component.
- Similarly, some heat is withdrawn from the condenser to condense the low boiling component.

Condensation and Evaporation in bubble.



- 1. When vapor bubbles through the liquid layer, the high temperature vapor transfers heat to the liquid.
- 2. Heat transfer from vapor bubble results in condensation of as little bit of high boiling point component(here,  $O_2$ ) from bubble.
- Also this heat exchange causes an evaporation of a little bit of low boiling point component(here, N<sub>2</sub>) from the bulk liquid.
- 4. Thus as the vapor bubble moves up, it becomes richer in low boiling point component, that is  $N_2$ .
- 5. And as liquid moves down, it gets richer and richer in high boiling point component, that is  $O_2$ .

- Real Rectification Column.
- In an ideal rectification column, the vapor and the liquid are in thermal equilibrium.
- But in actual rectification columns, the vapor does not leave the plate as the same temperature of the liquid.
- To ensure the required heat exchange, more number of plates are required than the theoretical prediction.
- Hence, there is a need to study efficiency of an actual system with respect to and ideal system.



- Let the mixture of N<sub>2</sub> and O<sub>2</sub> at 1 atm be subjected to rectification.
- Consider a j<sup>th</sup> plate of the rectification column as shown in the figure.
- Across this plate, the vapor mixture rises up and the liquid mixture flows down.



- Let  $y_{j-1}$  be the mole fraction of LBC in vapor phase rising to the j<sup>th</sup> plate.
- The composition of liquid at this plate is given by  $X_j$
- In thermal equilibrium, the mole fraction of LBC in vapor phase leaving the j<sup>th</sup> plate is  $\mathcal{Y}_{0,j}$



- But due to the non-ideality, the mole fraction of LBC in vapor phase leaving the  $j^{th}$  plate is  $y_j$ .
- Hence, the maximum possible and the actual change in mole fractions are
  - $y_{0,j} y_{j-1}$
  - $y_j y_{j-1}$

Murphree efficiency



- Murphree efficiency of a plate is defined as the ratio of actual change in mole fraction to the maximum possible change that can occur.
- Mathematically,

$$\eta_M = \frac{y_j - y_{j-1}}{y_{0,j} - y_{j-1}}$$

Murphree efficiency

- Heat and mass transfer analysis between the vapor and the bulk fluid is important to understand the underlying physics.
- In order to achieve high Murphree efficiency, the following conditions are required.
  - Small bubbles
  - Long time of contact
  - Large values of the overall mass and heat transfer coefficients.

## 4.6 Flash Calculations



Fig. rectification column

 When the feed stream is introduced into a rectification column, it is important to determine the condition of the feed stream if the stream enters the column as a twophase fluid. Flash calculation is used to solve this type of problem. Applying a mass balance for component j, we obtain

$$x_{Fj}F = y_jV + x_jL = y_j(F - L) + x_jL$$

The mole fraction of the jth component in the liquid phase  $x_j = y_j/K_j$ , so We may solve the mole fraction  $y_j$ 

$$y_j = \frac{x_{Fj}}{1 + \frac{L}{F} \left(\frac{1}{K_j} - 1\right)}$$

#### **4.6 Flash Calculations**

If we apply an energy balance to the system, we obtain

$$h_FF = hL + HV = hL + H(F - L)$$

$$\frac{L}{F} = \frac{H - h_F}{H - h}$$

The sum of the mole fractions for the vapor phase is equal to unity.

$$\sum_j y_j = 1 = \sum_j \frac{x_{Fj}}{1 + \frac{L}{F} \left( \frac{1}{K_j} - 1 \right)}$$





Rectification column

Perforated plate

Two basic methods used to determine the number of theoretical plates

- Ponchon (1921) and Savarit (1922)
- McCabe and Thiele (1925)

#### Ponchon (1921) and Savarit (1922) method

- Rigorous and capable of handling all distillations, but requires detailed enthalpy data for its application
- Simultaneous material and energy balance calculations
- Detailed calculations of equilibrium conditions
- More accurate, but difficult to use
- Largely superseded by rigorous computer simulation calculations
- Requires detailed enthalpy data in its application

#### McCabe and Thiele (1925) method

- Less rigorous, enthalpy data not required.
- Adequate for many applications, more commonly use because of its simplicity
- Uses graphical solution for binary mixture on equilibrium diagram (x-y plot).
- Provides the number of theoretical (ideal) trays required for a given separation.
- Pressure is assumed constant throughout the entire column.
- Less general, but it requires only equilibrium concentration data in its application



Upper enriching section (suppose a two-component mixture for simplicity)



Conservation of Mass Principle

$$V_n = L_{n+1} + D$$
 (4.38)

$$y_n V_n = x_{n+1} L_{n+1} + x_D D$$
 (4.39)

#### Upper enriching section (suppose a two-component mixture for simplicity)

• First Law of Thermodynamics

Assuming

- 1. no heat in-leaks from ambient
- 2. no work transfer
- 3. steady-state operation
- 4. negligible changes in kinetic and potential energies

Upper enriching section (suppose a two-component mixture for simplicity)

$$\overset{(3)}{\not E} = \dot{Q} - \overset{(2)}{\not W} + \sum \dot{m}_{i}(h_{i} + \frac{1}{2}v_{i}^{2} + gz_{i}) - \sum \dot{m}_{e}(h_{e} + \frac{1}{2}v_{e}^{2} + gz_{e})$$

$$= \dot{Q}_{n} - \dot{Q}_{out} + \dot{m}_{i}(h_{i} + \frac{1}{2}v_{i}^{2} + gz_{i}) - \dot{m}_{e}(h_{e} + \frac{1}{2}v_{e}^{2} + gz_{e})$$

 $= \dot{Q}_{out} + \dot{m_i}h_i - \dot{m_e}h_e = -\dot{Q}_D + VnHn - L_{n+1}h_{n+1} - h_DD$ 

 $\mathbf{V}_{\mathbf{n}}\mathbf{H}_{\mathbf{n}} = \mathbf{h}_{\mathbf{n+1}}\mathbf{L}_{\mathbf{n+1}} + \mathbf{h}_{\mathbf{D}}\mathbf{D} + \dot{\mathbf{Q}}_{\mathbf{D}}$ 

Upper enriching section (suppose a two-component mixture for simplicity)

$$V_{n}H_{n} = h_{n+1}L_{n+1} + h_{D}D + \dot{Q}_{D}$$

$$V_{n}H_{n}/D = h_{n+1}L_{n+1}/D + h_{D} + \dot{Q}_{D}/D$$

$$\frac{V_{n}H_{n}}{D} = \frac{h_{n+1}(V_{n} - D)}{D} + h_{D} + \frac{\dot{Q}_{D}}{D}$$

$$\frac{V_{n}}{D}(H_{n} - h_{n+1}) = \frac{\dot{Q}_{D}}{D} + h_{D} - h_{n+1}$$

$$\frac{D}{V_{n}} = \frac{H_{n} - h_{n+1}}{(\dot{Q}_{D}/D) + h_{D} - h_{n+1}}$$

Upper enriching section (suppose a two-component mixture for simplicity)

Using eqn. (4.38) 
$$V_n = L_{n+1} + D \longrightarrow \frac{L_{n+1}}{V_n} = 1 - (D/V_n)$$
 (4.43)

Using eqn. (4.39) 
$$y_n V_n = x_{n+1}L_{n+1} + x_D D \longrightarrow y_n = \frac{L_{n+1}}{V_n} x_{n+1} + \frac{D}{V_n} x_D$$
 (4.44)

#### Operating line (upper enriching section)



Enthalpy-composition for nitrogen-oxygen mixtures at 1atm

The bubble line and the dew line



- 1. The molar heat of vaporization for the two pure components were identical
- 2. The enthalpy of mixing were negligible  $\downarrow$ Then, Exactly horizontal lines  $\downarrow$   $h_{n+1}$  and H are constans for a particular column

Operating line (upper enriching section)

$$\frac{D}{V_{n}} = \frac{H_{n} - h_{n+1}}{(\dot{Q}_{D}/D) + h_{D} - h_{n+1}} = \text{constant}$$
(4.42)

$$\frac{L_{n+1}}{V_n} = 1 - (D/V_n) = constant$$
 (4.43)

Operating line (upper enriching section)

$$\frac{D}{V_{n}} = \frac{H_{n} - h_{n+1}}{(\dot{Q}_{D}/D) + h_{D} - h_{n+1}} = \text{constant}$$
(4.42)

$$\frac{L_{n+1}}{V_n} = 1 - (D/V_n) = \text{constant}$$
(4.43)

The operating line



Operating line (upper enriching section)



The y-intercept value (at x=0,  $y=x_D D/V_n$ ) may be used to construct the operating line for the enriching section of the column



Similar analysis applied to the stripping section

Conservation of mass principle



<Control volume for the stripping section>

Similar analysis applied to the stripping section

Applying lower boiling-point components yields,

$$\mathbf{x}_{m+1}\mathbf{L}_{m+1} = \mathbf{y}_m\mathbf{V}_m + \mathbf{x}_B\mathbf{B}$$

 $(x_B = mole \ fraction \ of \ the \ lower - boiling - point \ component \ in \ the \ bottom \ product)$ 

Applying First law of thermodynamics,

$$\mathbf{h}_{m+1}\mathbf{L}_{m+1} + \dot{\mathbf{Q}}_{\mathrm{B}} = \mathbf{H}_{\mathrm{m}}\mathbf{V}_{\mathrm{m}} + \mathbf{h}_{\mathrm{B}}\mathbf{B}$$

 $(h_B = the enthalpy of the bottom product stream)$ 

Similar analysis applied to the stripping section

Operating line for the lower or stripping section of the column,

$$\mathbf{y}_{\mathrm{m}} = \left(\frac{\mathbf{L}_{\mathrm{m+1}}}{\mathbf{V}_{\mathrm{m}}}\right) \mathbf{x}_{\mathrm{m+1}} - \left(\frac{\mathbf{B}}{\mathbf{V}_{\mathrm{m}}}\right) \mathbf{x}_{\mathrm{B}}$$

$$\frac{B}{V_{m}} = \frac{H_{m} - h_{m+1}}{\left(\frac{\dot{Q}_{B}}{B}\right) - h_{B} + h_{m+1}}$$

If  $x_{m+1} = x_B = y_m$  on the 45° diagonal

$$\frac{L_{m+1}}{V_m} = 1 + \left(\frac{B}{V_m}\right)$$

Finding intersection of the operating lines for the upper and lower sections,



Define parameter q = the ratio of the difference in liquid flow between the upper and lower column sections to the feed flow

$$q = (L_{m+1} - L_{n+1})/F$$

Combine equations,

$$F = V_n - V_m + L_{m+1} - L_{n+1}, \qquad q = (L_{m+1} - L_{n+1})/F$$

The difference in vapor flows,

$$V_{\rm n} - V_{\rm m} = (1 - q)F$$

Using upper vapor flow and lower vapor flow equations, we can obtain the intersection of the two operating lines

$$y_n = \left(\frac{L_{n+1}}{V_n}\right) x_{n+1} + \left(\frac{D}{V_n}\right) x_D$$
 (upper section)

$$y_m = \left(\frac{L_{m+1}}{V_m}\right) x_{m+1} - \left(\frac{B}{V_m}\right) x_B$$
 (lower section)

$$V_n - V_m = (L_{n+1} - L_{m+1})(x/y) + (x_D D + x_B B)/y = (1 - q)F$$

For the column as a whole,

$$x_F F = x_D D + x_B B$$

Therefore,

$$-qF\left(\frac{x}{y}\right) + \frac{x_FF}{y} = (1-q)F$$

The locus of the intersection of the two operating lines is,

$$\mathbf{y} = \frac{\mathbf{x}_{\mathrm{F}}}{1-q} - \frac{q}{1-q}\mathbf{x}$$

Determine q value,

$$h_F F = V_n H_n - V_m H_m + L_{m+1} h_{m+1} - L_{n+1} h_{n+1}$$

Vapor and liquid enthalpies are assumed to be independent of composition,

$$H_n = H_m = H$$
 and  $h_{m+1} = h_{n+1} = h$ 

And,

$$h_FF = (V_n - V_m)H + (L_{m+1} - L_{n+1})h$$

And, we also can obtain,

$$\mathbf{h}_{\mathrm{F}} = (1 - q)\mathbf{H} + q\mathbf{h}$$

The value of q may be determined from the feed stream enthalpy  $h_F$  and the saturated vapor enthalpy H and saturated liquid enthalpy h

$$q = \frac{H - h_F}{H - h}$$

The value of the parameter q is as follows, depending upon the condition of the feed stream,



i ) Sub-cooled liquid feed : q > 1
ii ) Saturated liquid feed : q = 1
iii ) Two-phase mixture feed : 0 < q < 1</li>
iv ) Saturated vapor feed : q = 0
v ) Superheated vapor feed : q < 0</li>

#### Example 4.7 – Determine the number of theoretical plates required!

- ① Top product : 98% nitrogen
- ② Bottom product : 5% nitrogen, 95% oxygen
- ③ Feed stream : 79% nitrogen, 21% oxygen
- ④ Feed stream liquid Molar fraction : 0.831 mol liquid/mol mixture
- (5) The bottom, top product streams leave as sat.liquid
- 6 Bottom product flow rate : 25 mol/s
- $\bigcirc$  Heat removed at the top : 1071 kW
- (8) Mean pressure : 101.3 kPa (1atm)
Example 4.7 – Determine the number of theoretical plates required!

i ) Find flow rates of the feed and top product – Conservation of mass!

F = B + D = 25 + D $x_F F = x_B B + x_D D$ 

Substituting in the numerical values,

$$0.79F = (0.05)(25) + 0.98D = 1.25 + 0.98(F - 25)$$

Solving for the feed stream flow rate,

$$F = \frac{(24.5 - 1.25)}{(0.98 - 0.79)} = 122.37 \text{ mol/s}$$
$$D = F - B = 122.37 - 25 = 97.37 \text{ mol/s}$$

#### **Example 4.7 – Determine the number of theoretical plates required!**

ii) Applying the First law of thermodynamics to the entire column,

 $\dot{\mathbf{Q}}_{\mathrm{B}} = \dot{\mathbf{Q}}_{\mathrm{D}} + \mathbf{h}_{\mathrm{D}}\mathbf{D} + \mathbf{h}_{\mathrm{B}}\mathbf{B} - \mathbf{h}_{\mathrm{F}}\mathbf{F}$ 

$$\label{eq:h} \begin{split} h &= 1050 \, \text{J/mol} \, \left( \text{saturated liquid} \right) = h_D = h_B \\ H &= 6916 \, \text{J/mol} \, (\text{saturated vapor}) \end{split}$$

The enthalpy of the feed stream(two-phase mixture) is,

 $h_F = (0.831)(1050) + (1 - 0.831)(6916) = 2041 \text{ J/mol}$ 

Using the values in the First law equation,

$$\dot{Q}_B = 1,071,000 + (1050)(97.37) + (1050)(25) - (2041)(122.37)$$
  
 $\dot{Q}_B = 949,731 \text{ W} = 949.731 \text{ kW}$ 

#### **Example 4.7 – Determine the number of theoretical plates required!**

iii) Determine the operating line equations

$$\frac{D}{V_{\rm n}} = \frac{6916 - 1050}{(1,071,000/97.37) + 1050 - 1050} = 0.5336$$

The reflux ratio,

$$\frac{L_{n+1}}{V_n} = 1 - 0.5336 = 0.4664$$

Operating line for the upper section,

$$y_n = 0.4664x_{n+1} + (0.5336)(0.98) = 0.4664x_{n+1} + 0.5229$$

## **4.7 Theoretical Plate Calculations for Columns**

**Example 4.7 – Determine the number of theoretical plates required!** 

$$\frac{B}{V_{\rm m}} = \frac{6916 - 1050}{(949,731/25) - 1050 + 1050} = 0.1544$$

The reflux ratio,

$$\frac{L_{m+1}}{V_m} = 1 + 0.1544 = 1.1544$$

Operating line for the lower section,

$$y_{m} = 1.1544x_{m+1} - (0.05)(0.1544) = 1.1544x_{m+1} - 0.00772$$

## **4.7 Theoretical Plate Calculations for Columns**

#### **Example 4.7 – Determine the number of theoretical plates required!**

The equation for the feed line,

$$q = \frac{6916 - 2041}{6916 - 1050} = 0.831$$

We obtain feed line,

$$y = \frac{0.79}{1 - 0.831} - \frac{0.831}{1 - 0.831} x = 4.6746 - 4.9172x$$

## **4.8 Minimum Number of Theoretical Plates**



**Rectification column** 

## **4.8 Minimum Number of Theoretical Plates**



System used in the development of the theoretical-plate equations. The flow rate of vapor *leaving* the *n*th plate is denoted by  $V_n$  and the flow rate of the liquid *leaving* the *n*th plate is denoted by  $L_n$ 

# **4.8 Minimum Number of Theoretical Plates**

Minimum number of theoretical plates  $\rightarrow$  when the slope of the operating lines approach unity



Upper operating line  $y_n = \frac{L_{n+1}}{V_n} x_{n+1} + \frac{D}{V_n} x_D$ 

Lower operating line  $y_m = \frac{L_{m+1}}{V_m} x_{m+1} - \frac{B}{V_m} x_B$ 

 $\begin{array}{c} \mbox{Minimum refrigeration} \\ Q_D \downarrow \\ \mbox{Requirement of an infinite} \\ \mbox{number of theoretical plates} \end{array}$ 

# **4.9 Types of Rectification Columns**

### Packed column

- Cylinders filled with a packing material such as metallic rings and wire gauze
- The packing material provides a large surface area
- This type of column is not so frequently used in cryogenic systems ex) diffuser

### Plate column

- Several plates to allow the vapor to bubble -Small holes
- Fine mesh
- Bubble cap

### **4.9 Types of Rectification Columns**

#### (a) Perforated plate





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### **4.9 Types of Rectification Columns**

(a) Bubble-cap plate plate





Bubble-cap trays

## 4.10 Linde Single-Column System



Fig. Linde single-column gas-separation system

# **4.11 Linde Double-Column System**



Fig. Linde double-column gas-separation system. If the product were desired in the liquid phase, the heat exchanger would be a two-channel heat exchanger, and The stream marked "pure oxygen gas out" would not be present

## **4.11 Linde Double-Column System**

The Linde single-column system has two serious disadvantages.

- 1. Only pure oxygen can be produced
- 2.Large quantities of oxygen are wasted in the impure nitrogen exhaust gas

## 4.11 Linde Double-Column System



Fig. Heylandt gas-separation system

## 4.12 Heylandt System

Schematics and pictures of Air separation systems



Fig. Real pictures of air separation systems