SOLVED EXAMPLES

Ex.1	Mixture of volatile compo	onents A and B has total va	apour pressure (in torr) :		
	$P = 254 - 119 x_{A}$				
	Where x _A is mol fraction o	f A in mixture.			
	Hence P_A° and P_B° are (in t	orr)			
	(A) 254, 119	(B) 119, 254	(C) 135, 254	(D) 154, 119	
Sol.	(C)				
	when $x_A = 0, x_B = 1$ $\therefore P = P_B^{\circ}$ $\therefore P_B^{\circ} = 254,$ when $x_B = 1, x_B = 0$				
	$\therefore P = P_{p}^{o}$				
	$\therefore P_{p}^{\circ} = 254.$				
	when $x_{1}^{B} = 1, x_{2} = 0$				
	when $x_A = 1, x_B = 0$ $\therefore P_A^\circ = P = 254 - C$	119=135			
Ex.2	When mango is placed in	-	-		
_	(A) Shrinks	(B) Swells	(C) Brusts	(D) Nothing happens	
Sol.	(A)				
				on present in the mango juice, therefore	
				pric acid. Therefore, H_2O will be passes	
	out from the mango into a	queous hydrochloric acid	. Therefore mango shrin	ks.	
Ex.3	Heptane and octane form	deal solution. At 373 K, th	e vapour pressures of the	e two liquids are 105.2 kPa and 46.8 kPa	
	respectively. What will be	the vapour pressure, in b	ar, of a mixture of 25g of	f heptane and 35 g of octane?	
Sol.	(A) Heptane $C_7 H_{16} m_A = 1$	00			
	(B) Octane $C_8 H_{18}$	$m_{\rm B}^{=114}$			
	$n_{A} = \frac{w_{A}}{m_{A}} = \frac{25}{100} = 0.25;$	$n_{\rm B} = \frac{32}{114} = 0.3$			
	$x_{A} = \frac{0.25}{0.25 + 0.30} = 0.45$	$x_{p} = \frac{0.3}{0.3} = 0.55$	5		
	0.20 0.00	^B 0.25 + 0.30			
	=0.45				
	$p = p_{\mathrm{A}}^{0} x_{\mathrm{A}} + p_{\mathrm{B}}^{0} x_{\mathrm{B}}$				
	$=105.2 \times 0.45 + 46.8 \times 0.45$	0.55			
	=47.34+25.74=73.081	Pa			
Ex.4	A certain substance 'A' tet	ramerises in water to the ex	xtent of 80%. A solution	of 2.5 g of A in 100 g of water lowers the	
	freezing point by 0.3°C. T				
	(A) 122	(B) 31	(C) 244	(D) 62	
Sol.	(D)		(0)	(2) 02	
	(-)				
	1-i 🔺 🔺				
	$\alpha = \frac{1-i}{1-\frac{1}{n}}$				
	1 n				
	$1 = -\frac{1}{1 = 1}$ $0.8 = \frac{1 - i}{1 = 1}; i = 0.4$				
	$1 - \frac{1}{1 - \frac{1}{2}}$				
	n				
	$\Delta T = iK_f \times m$				
		00			
	$0.3 = 0.4 \times 1.86 \times \frac{\mathrm{W}_{\mathrm{B}} \times 100}{\mathrm{m}_{\mathrm{B}} \times \mathrm{W}}$				
	$0.3 = 0.4 \times 1.86 \times \frac{2.5 \times 100}{m_{\rm B} \times 100}$	<u>00</u>			
	$m_{\rm B} \times 10^{-100}$	0			
	$m_{\rm B} = 62$				

Ex.5 Which of the following solutions will exhibit highest boiling point ?

(A) $0.01 \text{ M} \text{ Na}_2 \text{SO}_4$ (B) $0.01 \text{ M} \text{ KNO}_3$ (C) 0.015 M urea

Sol. (A)

 $\Delta T = i \times K_h \times m$

 $i \times m$ of Na₂SO₄ is highest, hence its boiling point will also be highest.

Na_2SO_4	$i \times m = 3 \times 0.01 = 0.03$
KNO3	$i \times m = 2 \times 0.01 = 0.02$
Urea	$i \times m = 1 \times 0.015 = 0.015$
Glucose	$i \times m = 1 \times 0.015 = 0.015$

 $\pi V = ST \{ \rightarrow n=1 \}$

- Ex.6 What is the osmotic pressure of 12% solution of can sugar (mol. wt. 342) at 17°C.
- Sol. 12 g. sugar is dissolved in 100 mL thus 342 g. sugar is dissolved in

8 litres

Now,

$$\pi = \frac{ST}{V} = \frac{0.0821 \times 290}{2.85} = 8.35 \text{ atm}$$

Ex.7 A dilute solution contains m mol of solute A in 1 kg of a solvent with molal elevation constant K_b . The solute dimerises in solution as $2A \ddagger \uparrow \uparrow \uparrow A_2$. Show that equilibrium constant for the dimer formation is

(D) 0.015 M glucose

$$K = \frac{K_{b}(K_{b}m - \Delta T_{b})}{(2\Delta T_{b} - K_{b}m)^{2}}$$

where $\Delta T_{\rm b}$ is the elevation of the boiling point for the given solution. Assume molarity = molality

after dimerization

initially

Sol. $2A \stackrel{\circ}{+} \stackrel{\circ}{-} \stackrel{\circ}{+} A_2$ m 0 $(m-m\alpha) \frac{m\alpha}{2}$

$$m(1-\alpha)$$

Where α is the degree of dimerization and m is molality which is also molarity (given). Hence due to dimerization, final molality = m

$$m(1-\alpha) + \frac{m\alpha}{2} = m\left(1-\frac{\alpha}{2}\right)$$

$$i = 1 + (n-1)\alpha$$

$$= 1 + \left(\frac{1}{2} - 1\right)\alpha$$

$$= \left(1 - \frac{\alpha}{2}\right)$$

$$\Delta T_{b} = K_{b} \times m \times i = K_{b} \times m\left(1 - \frac{\alpha}{2}\right)$$

$$\therefore \qquad \alpha = \frac{2(K_{b}m - \Delta T_{b})}{K_{a}m}$$

mα

2

equilibrium constant K for the dimer formation is



$$K = \frac{[A_2]}{[A]^2} = \frac{\frac{m\alpha}{2}}{m^2(1-\alpha)^2} = \frac{\alpha}{2m(1-\alpha)^2}$$
$$K = 2\left[\frac{K_b m - \Delta T_b}{K_b m}\right] / 2m \left[1 - \frac{2(K_b m - \Delta T_b)}{K_b m}\right]^2$$
$$K = \frac{K_b (K_b m - \Delta T_b)}{(K_b m - 2K_b m + 2\Delta T_b)^2}$$
$$K = \frac{K_b (K_b m - \Delta T_b)}{(2\Delta T_b - K_b m)^2}$$
Proved

- **Ex.8** Distribution coefficient of an organic acid between water and benzene is 4.1 in favour of C_6H_6 . If 5 g of acid is distributed in between 50 mL of benzene and 100 mL of water, calculate the concentration of acid in two solvents.
- Sol. Let the amount of organic acid in

$$C_6H_6$$
 layer = a g

volume of
$$C_6 H_6 = 50 \text{ mL}$$

- \therefore concentration of acid in $C_6 H_6 = \frac{a}{50} \text{ g mL}^{-1}$
- since total amount of acid = 5 g
- : Amount of acid in H_2O layer = (5 a) g
- and volume of $H_2O = 100 \text{ mL}$

$$\therefore$$
 Conc. of acid in H₂O = $\left(\frac{5-a}{100}\right) \frac{g}{mL}$

Now,
$$K = \frac{\text{Conc. of acid in } C_6 H_6}{\text{Conc. of acid in } H_2 O}$$

$$=\frac{a}{50}\times\frac{100}{(5-a)}$$

$$\therefore \quad 4.1 \quad = \frac{a}{50} \times \frac{100}{(5-a)}$$

- or a = 3.361 g.
- : Amount of acid in 50 mL

$$C_6 H_6 = 3.361 g$$

:. acid concentration in $C_6 H_6 = \frac{3.361}{50} \times 1000 = 67.22 \text{ g/L}$

Also, amount of acid in 100 mL $H_2O = 5 - a = 5 - 3.361 = 1.639$ g

$$\therefore$$
 acid concentration in H₂O = $\frac{1.639}{100} \times 1000 = 16.39 \text{ g/L}$



- **Ex.9** The freezing point depression of 0.001 m $K_x[Fe(CN)_6]$ is 7.10 × 10⁻³ K. Determine the value of x. Given, $K_f = 1.86 \text{ K kg mol}^{-1}$ for water.
- Sol.

$$\Delta x = i \times K_{f} \times m$$

$$7.10 \times 10^{-3} = i \times 1.86 \times 0.001$$

$$i = 3.817$$

$$\alpha = \frac{i-1}{n-1}$$

$$1 = \frac{3.817 - 1}{(x+1) - 1}$$

$$x = 2.817 \approx 3$$

- \therefore Molecular formula of the compound is K₃[Fe(CN)₆].
- **Ex.10** The composition of vapour over a binary ideal solution is determined by the composition of the liquid. If x_A and y_A are the mole fractions of A in the liquid and vapour, respectively find the value of x_A for which $(y_A x_A)$ has maximum. What is the value of the pressure at this composition ?
- Sol. Since

$$\boldsymbol{y}_{A} = \frac{\boldsymbol{x}_{A}\boldsymbol{P}_{A}^{\circ}}{\boldsymbol{P}_{B}^{\circ} + (\boldsymbol{P}_{A}^{\circ} - \boldsymbol{P}_{B}^{\circ})\boldsymbol{x}_{A}}$$

Substracting x_A from both the sides, we get

$$y_{A} - x_{A} = \frac{x_{A}P_{A}^{\circ}}{P_{B}^{\circ} + (P_{A}^{\circ} - P_{B}^{\circ})x_{A}} - x_{A}$$

Differentiating this with respect to x_A , we get

$$\frac{d(y_{A} - x_{A})}{dx_{A}} = \frac{P_{A}^{\circ}}{P_{A}^{\circ} + (P_{A}^{\circ} - P_{B}^{\circ})x_{A}} - \frac{x_{A}P_{A}^{\circ}(P_{A}^{\circ} - P_{B}^{\circ})}{\left\{P_{B}^{\circ} + (P_{A}^{\circ} - P_{B}^{\circ})x_{A}\right\}^{2}} -$$

The value of x_A at which $y_A - x_A$ has a maximum value can be obtained by setting the above differential equal to zero. Thus, we have

$$\frac{P_{A}^{\circ}}{P_{A}^{\circ} + (P_{A}^{\circ} - P_{B}^{\circ})x_{A}} - \frac{x_{A}P_{A}^{\circ}(P_{A}^{\circ} - P_{B}^{\circ})}{\left\{P_{B}^{\circ} + (P_{A}^{\circ} - P_{B}^{\circ})x_{A}\right\}^{2}} - 1 = 0$$

Solving for x_A , we get $x_A = \frac{\sqrt{P_A^\circ P_B^\circ} - P_B^\circ}{P_A^\circ - P_B^\circ}$

The value of P at this composition is

$$P = x_{A} P_{A}^{\circ} + x_{B} P_{B}^{\circ}$$

or
$$P = P_{B}^{\circ} + (P_{A}^{\circ} - P_{B}^{\circ}) x_{A}$$

or
$$P = P_{B}^{\circ} + (P_{A}^{\circ} - P_{B}^{\circ}) \left(\frac{\sqrt{P_{A}^{\circ} P_{B}^{\circ}} - P_{B}^{\circ}}{P_{A}^{\circ} - P_{B}^{\circ}} \right)$$

or
$$P = \sqrt{P_{A}^{\circ} P_{B}^{\circ}}$$



E	xercise # 1	[Sin	ngle Correct Choice	Type Questions]
1.	We have 100 mL of 0.1 M KCl solution (A) evaporate 50 mL water (C) add 0.1 mol KCl	on. To make it 0.	2 M, (B) evaporate 50 mL so (D) add 0.01 mol KCl	lution
2.	Persons are medically considered to h lead per decilitre of blood. Concentra (A) 1000 (B) 100	-		ration greater than 10 micrograms of (D) 1
3.	Which of the following concentration solution of HCl are known?			
4.	 (A) Molality (B) Molar (B) Which statement best explains the m (A) A Solute will easily dissolve a sol (B) A solvent and solute with similar (C) The only true solutions are form (D) The only true solutions are form 	eaning of the pl ute of similar n intermolecular ed when water	aass forces will readily form a s dissolves a non-polar solu	olution
5.	An ionic compound that attracts atm (A) Dilute (B) Hygre	-	so strongly that a hydrate (C) Immiscible	is formed is said to be : (D) Miscible
6.	The vapour pressure of water depend (A) Surface area of container (C) Temperature	ds upon :	(B) Volume of container (D) All	
7.	A liquid is kept in a closed vessel. If surface, then the vapour pressure of (A) More than what would be if the g (B) Same as what would be if the gla (C) Less than what would be if the gl (D) Cannot be predicted	the liquid in the class plate were ss plate were re	e vessel is : removed moved	all hole is kept on top of the liquid
8.	At higher altitudes, water boils at ten (A) temperature of higher altitudes is (C) the proportion of heavy water in	low	 °C because (B) atmospheric pressur (D) atmospheric pressur 	
9.	Among the following substances, the (A) Water (B) Merce	· ·	r pressure is exerted by : (C) Kerosene	(D) Rectified spirit
10.	When a liquid that is immiscible with contained 1.25 g of the liquid per gran mass of liquid? (A) 7.975 g/mol (B) 166 g	n of water. The v		1
11.	Two liquids X and Y are perfectly in pressure of a mixture of X and Y prep (A) 600 torr (B) 400 to	ared in weight 1		
12.	An ideal solution contains two volatile 4 mole of B then total vapour pressur (A) 150 (B) 180		· · · · · ·	r). If mixture contain 1 mole of A and (D) 198.88

Add. 41-42A, Ashok Park Main, New Rohtak Road, New Delhi-110035 +91-9350679141

13	The vapour pressure of two pure liquids A and B, that form an ideal solution are 100 and 900 torr respectively a temperature T. This liquid solution of A and B is composed of 1 mole of A and 1 mole of B. What will be the pressure when 1 mole of mixture has been vaporized ?				
	-	C) 300 torr	(D) None of these		
14.	For a binary ideal liquid solution, the total pressure of t (A) $P_{total} = P_A^o + (P_A^o - P_B^o) X_B$ (C) $P_{total} = P_B^o + (P_B^o - P_A^o) X_A$ (C)	he solution is given as : B) $P_{total} = P^o_B + (P^o_A - P^o_B)$ D) $P_{total} = P^o_B + (P^o_B - P^o_A)$	X _A X _B		
15.	Given at 350 K $p_A^{\circ} = 300$ torr and $p_B^{\circ} = 800$ torr, the composition (A) $X_A^{\circ} = 0.08$ (B) $X_A^{\circ} = 0.06$ (C)	position of the mixture having C) $X_A = 0.04$	g a normal boiling point of 350 K is (D) $X_A = 0.02$		
16.	Two liquids A and B have P_{A}^{o} and P_{B}^{o} in the ratio of 1 : 3 are 1 : 3 then mole fraction of 'A' in vapour phase in equ (A) 0.1 (B) 0.2 (6)				
17.	2 0	B) HNO ₃ and water D) C_6H_6 and $C_6H_5CH_3$			
18.	 A maxima or minima obtained in the temperature compo (A) an azeotropic mixture (B) an eutectic formation (C) that the liquids are immiscible with one another (D) that the liquids are partially miscible at the maximum 		of two liquids indicates		
19.		B) Positive deviations fromD) The applicability of He			
20.	For chloroform and acetone or for a solution of chloroform and acetone if p_s (observed (actual)) is compared wit p_s (Theoretical (Raoult)) then which of the following is /are true ?				
	(A) $p_{s(actual)} < p_{s(raoult)}$ (B) $\begin{array}{l} \text{Limit} \\ \chi_{\text{chloroform}} \longrightarrow 0 \end{array} (p^{o}_{acet})$	$t_{tone} - p_{sactual}) = 0$		
	(C) $\underset{X_{acetone}}{\text{Limit}} \longrightarrow 0 (p^{\circ}_{chloroform} - p_{s(actual)}) = 0$ (1)	D) $p^{o}_{acetone} > p^{o}_{chloroform} n$	ear room temperature		
21.	 The vapour pressure of the solution of two liquids A(p° = x_A = 0.4. The result shows that (A) solution exhibits ideal behaviour (B) solution shows positive deviations (C) solution shows negative deviations (D) solution will show positive deviations for lower concernance 				
22.		4 the vapour pressure of so B) $C_6H_5Cl - C_6H_5Br$ D) $nC_6H_{14} - nC_7H_{16}$	lution is 580 torr then the mixture		
23.	The solubility of $N_2(g)$ in water exposed to the a 5.3 × 10 ⁻⁴ M. Its solubility at 760 mm and at the same ten		partial pressure is 593 mm is (10) 2400 M		

(A) 4.1×10^{-4} M (B) 6.8×10^{-4} M (C) 1500 M (D) 2400 M



24.	The solubility of gases in liquids : (A) increases with increase in pressure and temperature (B) decreases with increase in pressure and temperature (C) Increases with increase in pressure and decrease in temperature (D) decreases with increase in pressure and increase in temperature					
25.	 Select correct statements : (A) Gases which have high value of Van der Waals constant 'a' are easily liquefied (B) Easily liquefiable gases are water soluble (C) Ions forming gases in a solvent are soluble in that solvent (D) Under same conditions, NH₃ has low solubility than that of CO₂. 					
26.	Some of the following g I: CO_2 ; Water insoluble gases ca (A) I, IV, V	$II: NH_3; III: HCl;$	e to formation of their ions IV : CH ₄ ; (C) I, II, III	: $V:H_2$ (D) IV, V		
27.	(A) 0.05 M K ₄ [Fe(CN) ₆] (B) 0.10 M K ₄ [Fe(CN) ₆] (C) 0.20 M NaCl and 0.10	and 0.10 M FeSO_4 and $0.05 \text{ M FeSO}_4 (\text{NH}_4)_2 \text{SO}_4$		tor be the same?		
28.	One mole of a solute A is	-	e of solvent. The association	of the solute take place as follows:		
	-	$nA \rightleftharpoons A_n$				
	(A) $i = 1 - \alpha$	(B) $i = 1 + \frac{\alpha}{n}$	(C) $i = \frac{1-\alpha+\frac{\alpha}{n}}{1}$	(D) i = 1		
29.	mass of the complex in t	he solu <mark>tion will</mark> be :	egree of ionization of K_3 [Fe((C) $M_{normal} (1 + \alpha)^{-1}$	 (D) equal to M_{normal} 		
30.		erature. If the molecular wei		95% of the vapour pressure of the solute,s the molecular weight of the solute,(D) none of these		
31.		ir pressures of a solvent and olatile solute respectively, (B) $P = P_0 N_1$		and N_1 and N_2 are the mole fractions (D) $P = P_0(N_1/N_2)$		
32.	A complex of iron and cy $(K_b = 0.52^\circ \text{ mol}^{-1} \text{ kg}):$ (A) $K_3[\text{Fe}(\text{CN})_6]$	ranide ions is 100% ionised a	at 1 m (molal). If its elevatior (C) K ₄ [Fe(CN) ₆]	n in b.p. is 2.08. Then the complex is (D) Fe(CN) ₄		
33.		sucrose in 100 grams of wa any grams of ice would hav (B) 20 grams		ce separates the freezing point goes(D) 23 grams		
34.			$MgCl_2$ by weight of solutio on of $MgCl_2$ ($K_b(H_2O) = 0.5$ (C) $T_b = 108.5^{\circ}C$	n. Calculate its normal boiling point 1 kgmol ⁻¹ K). (D) $T_b = 110.3^{\circ}C$		

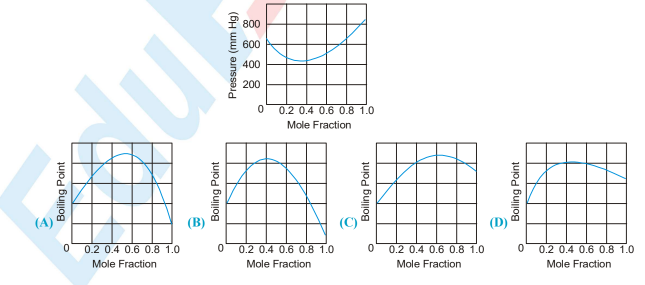


35.	$\begin{array}{ll} PtCl_4 . \ 6H_2O \ can \ exist \ as \ a \ hydrated \ complex \ 1 \ molal \ aq. \ solution \ has \ depression \ in \ freezing \ point \ of \ 3.72^\circ. \ Assume \ 100\% \ ionisation \ and \ K_f(H_2O) = 1.86^\circ \ mol^{-1} \ kg, \ then \ complex \ is \ - \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $
36.	How many moles of sucrose should be dissolved in 500 gms of water so as to get a solution which has a difference of 104°C between boiling point and freezing point. ($K_f = 1.86 \text{ K Kg mol}^{-1}$, $K_b = 0.52 \text{ K Kg mol}^{-1}$) (A) 1.68 (B) 3.36 (C) 8.40 (D) 0.840
37.	Which of the following has been arranged in order of decreasing freezing point? (A) $0.05 \text{ M KNO}_3 > 0.04 \text{ M CaCl}_2 > 0.140 \text{ M sugar} > 0.075 \text{ M CuSO}_4$ (B) $0.04 \text{ M BaCl}_2 > 0.140 \text{ M sucrose} > 0.075 \text{ M CuSO}_4 > 0.05 \text{ M KNO}_3$ (C) $0.075 \text{ M CuSO}_4 > 0.140 \text{ M sucrose} > 0.04 \text{ M BaCl}_2 > 0.05 \text{ M KNO}_3$ (D) $0.075 \text{ M CuSO}_4 > 0.05 \text{ M NaNO}_3 > 0.140 \text{ M sucrose} > 0.04 \text{ M BaCl}_2$
38.	Consider following cases : I : 2M CH ₃ COOH solution in benzene at 27°C where there is dimer formation to the extent of 100% II : 0.5 M KCl aq. solution at 27°C, which ionises 100% Which is/are true statements(s) : (A) both are isotonic (B) I is hypertonic (C) II is hypotonic (D) none is correct
39.	 Select correct statement : (A) Osmosis, like all colligative properties, results from an increase in entropy as pure solvent passes through the membrane and mixes with the solution (B) Desalination of sea-water is done by reverse osmosis (C) Both are correct statements (D) None is correct statement
40.	Osmotic pressure of 30% solution of glucose is 1.20 atm and that of 3.42% solution of cane sugar is 2.5 atm. The
	osmotic pressure of the mixture containing equal volumes of the two solutions will be(A) 2.5 atm(B) 3.7 atm(C) 1.85 atm(D) 1.3 atm.
41.	
41.	 (A) 2.5 atm (B) 3.7 atm (C) 1.85 atm (D) 1.3 atm. The dissolving process is exothermic when : (A) The energy released in solvation exceeds the energy used in breaking up solute-solute and solvent-solvent interactions. (B) The energy used in solvation exceeds the energy released in breaking up solute-solute and solvent-solvent interactions. (C) The energy released in solvation is about the same as the energy used in breaking up solute-solute and solvent-solvent interactions. (D) The energy used in solvation is about the same as the energy used in breaking up solute-solute and solvent-solvent interactions.



44.	original volume of the sa	ample ?	-	of NaCl was obtained. What was the
	(\mathbf{A}) 30 mL	(B) 333 mL	(C) 750 mL	(D) 1000 mL
45.	A 20.0 mL sample of Cu of the original solution ? (A) 48.4 M		ted to dryness, leaving 0.967	7 g of residue. What was the molarity (D) 0.303 M
46.	When KCl dissolves in (A) $\Delta H = + ve$, $\Delta S = + ve$ (C) $\Delta H = + ve$, $\Delta S = + ve$	-	ic dissolution), then : (B) $\Delta H = + ve$, $\Delta S = -ve$ (D) $\Delta H = -ve$, $\Delta S = -ve$	
47.		1		89°C under a reduced pressure of cent of chlorobenzene in the distillate (D) 38.46
48.	If two liquids A ($P_A^o = 100$) independently of the oth (A) less than 100 torr (C) between 100 to 200 to	ner, are present in a closed	r) are completely immiscible I vessel. The total vapour pr (B) greater than 200 ton (D) 300 torr	with each other, each one will behave ressure of the system will be: r
49.	the following is not a po			0 Torr respectively at 20°C. Which of hary/ternary solution of these at 20°C (D) $53\frac{1}{2}$
50.	the boiling point of a mix	(B) 1	zene	benzene chlorobenzene 90 100 110 120 $t(c^9) \rightarrow$
51	Given P y curve for a no	n ideal liquid mixture (Fig) Identify the correct T x cu	irve for the same mixture

51. Given P-x curve for a non-ideal liquid mixture (Fig.). Identify the correct T-x curve for the same mixture.





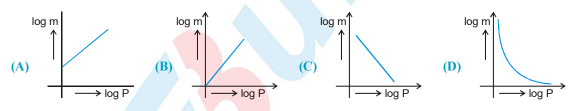
52. If vapour pressures of pure liquids 'A' & 'B' are 300 and 800 torr respectively at 25°C. When these two liquids are mixed at this temperature to form a solution in which mole percentage of 'B' is 92, then the total vapour pressure is observed to be 0.95 atm. Which of the following is true for this solution.

(A) $\Delta V_{mix} > 0$ (B) $\Delta H_{mix} < 0$ (C) $\Delta V_{mix} = 0$ (D) $\Delta H_{mix} = 0$

- 53. At 323 K, the vapour pressure in millimeters of mercury of a methanol-ethanol solution is represented by the equation $p = 120 X_A + 140$, where X_A is the mole fraction of methanol. Then the value of $\lim_{x_A \to 1} \frac{p_A}{X_A}$ is : (A) 250 mm (B) 140 mm (C) 260 mm (D) 20 mm
- 54. Consider two liquids A & B having pure vapour pressures $P_A^{\circ} \& P_B^{\circ}$ forming an ideal solution. The plot of $\frac{1}{X_A}$ v/s $\frac{1}{Y_A}$ (where X_A and Y_A are the mole fraction of liquid A in liquid and vapour phase respectively) is linear with slope and Y intercepts respectively :

(A)
$$\frac{P_A^{\circ}}{P_B^{\circ}}$$
 and $\frac{(P_A^{\circ} - P_B^{\circ})}{P_B^{\circ}}$
(B) $\frac{P_A^{\circ}}{P_B^{\circ}}$ and $\frac{(P_B^{\circ} - P_A^{\circ})}{P_B^{\circ}}$
(C) $\frac{P_B^{\circ}}{P_A^{\circ}}$ and $\frac{(P_A^{\circ} - P_B^{\circ})}{P_B^{\circ}}$
(D) $\frac{P_B^{\circ}}{P_A^{\circ}}$ and $\frac{(P_B^{\circ} - P_A^{\circ})}{P_B^{\circ}}$

- 55. Which of the following is less than zero for ideal solutions ? (A) ΔH_{mix} (B) ΔV_{mix} (C) ΔG_{mix} (D) ΔS_{mix}
- 56. Which of the following curves represents the Henry's law?



- 57. According to Henry's law, the solubility of a gas in a given volume of liquid increases with increase in : (A) Temperature (B) Pressure (C) Both (A) and (B) (D) None of these
- 58. The degree of dissociation of an electrolyte is α and its van't Hoff factor is i. The number of ions obtained by complete dissociation of 1 molecule of the electrolyte is:

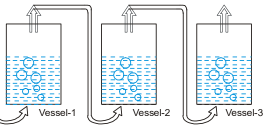
(C) $\frac{i-1}{\alpha}$

(A)
$$\frac{i+\alpha-1}{\alpha}$$
 (B) $i-\alpha-1$

(D)
$$\frac{i+1+\alpha}{1-\alpha}$$

59. Dry air is slowly passed through three solutions of different concentrations, c_1 , c_2 and c_3 ; each containing (non volatile) NaCl as solute and water as solvent, as shown in the Fig. If the vessel 2 gains weight and the vessel 3 loses weight, then :







60.	Pressure cooker reduce (A) the heat is more even (B) a large flame is used (C) boiling point of wat (D) whole matter is con	enly distributed inside the l er is elevated	cooker	
61.	its 0.1 molal solution w solute which neither u trimerization?	as found to be identical to ndergoes association nor	the boiling point elevation dissociation. To what po	solvent. The boiling point elevation of n in case of a 0.08 molal solution of a ercent had the solute 'S' undergone
	(A) 30%	(B) 40%	(C) 50%	
62.			. If that complex is suppose a number of Co ²⁺ in the co (C) Four	ed to be 75% ionised in water with vant mplex can be : (D) Six and Four both
63.		AgNO ₃ , this solution will f	orm(assuming no change i	es [Cu(NH ₃) ₃ Cl] Cl in 3 moles H ₂ O is n degree of ionisation of substance on
	(A) 1 mol AgCl	(B) 0.25 mol AgCl	(C) 0.5 mol AgCl	(D) 0.40 mol AgCl
64.	In the following aqueou (A) 1 m sucrose (B) 1 m potassium ferric (C) 1 m potassium sulpl maximum value of vapo (A) A	zyanide	hat of : (C) C	(D) equal
65.		(no volume change on mi ing points remain same become lower		
66.	The freezing point of aq 1.86 and assume molarit (A) 290 K		ns 3% urea, 7.45% KCl and (C) 267.42 K	19% of glucose is (given K_f of water = (D) 250 K
67.	x mole of KCI and y mo		ved in 1 kg of water. Given	that $x + y = 0.1$ and K_f for water is 1.85 (D) 0.37^0 to 0.93^0
68.	same volume of 0.03 M	aqueous solution of NaCl, equation is attached ? Volu	are placed side by side in a	
69.	injection that is to have	the same osmotic pressur	e as blood is :	to be used per litre for an intravenous
	(A) 0.3	(B) 0.2	(C) 0.1	(D) 0.4
70.	HgCl ₂ is 34.3. What is the (A) as Hg ₂ Cl ₂ molecules	he state of mercurous chlor s	ide in HgCl ₂ ? (Hg – 200, 6 (B) as HgCl molecule	S
	(C) as Hg ⁺ and Cl ⁻ ions		(D) as Hg_2^{2+} and Cl^-	ions



Add. 41-42A, Ashok Park Main, New Rohtak Road, New Delhi-110035 +91-9350679141

71.	 At a constant temperature, ΔS will be maximum for which of the following processes : (A) Vaporisation of a pure solvent (B) Vaporisation of solvent from a solution containing nonvolatile and nonelectrolytic solute in it (C) Vaporisation of solvent from a solution containing nonvolatile but electrolytic solute in it (D) Entropy change will be same in all the above cases
72.	FeCl ₃ on reaction with $K_4[Fe(CN)_6]$ in aq. solution gives blue colour. These are separated by a semipermeable membrane PQ as shown. Due to osmosis there is- (A) blue colour formation in side X (B) blue colour formation in both of the sides X and Y (D) no blue colour formation
73.	 The solubility of gases in liquids : (A) increases with increase in pressure and temperature (B) decreases with increase in pressure and temperature (C) Increases with increase in pressure and decrease in temperature (D) decreases with increase in pressure and increase in temperature
74.	Colligative properties have many practical uses, some of them may be : I : Melting of snow by salt II : Desalination of sea water III : Determination of molar mass IV : Determination of melting point and boiling point of solvent Actual practical uses are : (A) I, II (B) III, IV (C) I, II, III (D) II, III, IV
75.	 Select correct statement(s): (A) When solid CaCl₂ is added to liquid water, the boiling temperature rises (B) When solid CaCl₂ is added to ice at 0°C, the freezing temperature falls (C) Both (A) and (B) (D) None of the above
76.	The concentration of pollutant in ppm (w/w), that has been measured at 450 mg per 150 kg of sample is :(A) 3 ppm(B) 6 ppm(C) 3000 ppm(D) 330 ppm
77.	Available are $1L ext{ of } 0.1 ext{ M vaCl}$ and $2L ext{ of } 0.2 ext{ M CaCl}_2$ solutions. Using only these two solutions what maximum volume of a solution can be prepared having $[Cl^-] = 0.34 ext{ M exactly}$. Both electrolytes are strong (A) $2.5 ext{ L}$ (B) $2.4 ext{ L}$ (C) $2.3 ext{ L}$ (D) None of these
78.	Mole fraction of $C_3H_5(OH)_3$ in a solution of 36 g of water and 46 g of glycerine is :(A) 0.46(B) 0.36(C) 0.20(D) 0.40
79.	A complex containing K^+ , Pt(IV) and Cl ⁻ is 100% ionised giving i = 3. Thus, complex is :(A) $K_2[PtCl_4]$ (B) $K_2[PtCl_6]$ (C) $K_3[PtCl_5]$ (D) $K[PtCl_3]$
80.	In which case, van't Hoff factor i remains unchanged ? [Assume common complexes of these ions] (A) PtCl ₄ reacts with KCl (B) aq. ZnCl ₂ reacts with aq. NH ₃ (C) aq. FeCl ₃ reacts with aq. K ₄ [Fe(CN)] ₆ (D) KMnO ₄ reduced to MnO ₂ in alkaline medium (MnO ₂ a black ppt)
81.	If $pK_a = -\log K_a = 4$, and $K_a = Cx^2$ then Van't Hoff factor for weak monobasic acid when $C = 0.01$ M is : (A) 1.01 (B) 1.02 (C) 1.10 (D) 1.20



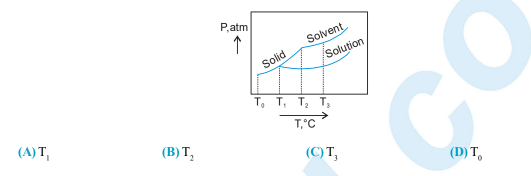
1 atm P,atm solver B D T°C (A)A **(B)**B (C) C **(D)**D 83. Select correct statement? (A) Heats of vaporisation for a pure solvent and for a solution are similar because similar intermolecular forces between solvent molecules must be overcome in both cases (B) Entropy change between solution and vapour is smaller than the entropy change between pure solvent and vapour (C) Boiling point of the solution is larger than that of the pure solvent (D) All are correct statements 84. Ratio of $\Delta T_{\rm h}/K_{\rm h}$ of 6% AB, and 9% A,B (AB, and A,B both are non-electrolytes) is 1 mol/kg in both cases. Hence, atomic masses of A and B are respectively : (A) 60, 90 **(D)** 10, 40 **(B)** 40, 40 (C) 40, 10 85. Consider following terms (m = molality): $\frac{\Delta T_b}{\cdot}$ **III** : II: mK, i IV:K $I: mK_{L};$ Terms which can be expressed in degree (temperature) are **(D)** I, III (A) III, IV **(B)** I, II (C) I, II, IIII Elevation in b.p. of an aqueous urea solution is 0.52° . (K_k = 0.52° mol⁻¹ kg) Hence, mole-fraction of urea in this 86. solution is : (A) 0.982 **(B)** 0.567 (C) 0.943 **(D)** 0.018 87. Insulin $(C_2H_{10}O_5)_n$ is dissolved in a suitable solvent and the osmotic pressure π of the solution of various concentration (in kg/m³) is measured at 20°C. The slope of a plot of π against c is found to be 8.134 × 10⁻³ (SI units) The molecular weight of the insulin (in kg/mol) is : (D) 8.314×10^{5} (A) 4.8×10^5 **(B)** 9×10^{5} (C) 293×10^3 88. An aqueous solution of a solute AB has b.p. of 101.08°C (AB is 100% ionised at boiling point of the solution) and freezes at -1.80° C. Hence, AB (K_b / K_f = 0.3) (A) is 100% ionised at the f.p. of the solution (B) behaves as non-electrolyte at the f.p. of the solution (C) forms dimer (D) none of the above 89. Density of 1M solution of a non-electrolyte $C_6H_{12}O_6$ is 1.18 g/mL. If $K_r(H_2O)$ is 1.86° mol⁻¹ kg, solution freezes at : $(A) - 1.58^{\circ}C$ **(B)** $-1.86^{\circ}C$ **(C)**−3.16°C **(D)** 1.86°C **90**. Mole fraction of a non-electrolyte in aqueous solution is 0.07. If K_c is 1.86° mol⁻¹ kg, depression in f.p., ΔT_c is: (A) 0.26° **(B)** 1.86° (C) 0.13° **(D)** 7.78° 91. Total vapour pressure of mixture of 1 mol of volatile component A ($p_A^\circ = 100 \text{ mmHg}$) and 3 mol of volatile component B $(p_p^{\circ} = 60 \text{ mmHg})$ is 75 mm. For such case : (A) there is positive deviation from Raoult's low (B) boiling point has been lowered (C) force of attraction between A and B is smaller than that between A and A or between B and B (D) all the above statements are correct

What is the normal boiling point of the solution represented by the phase diagram ?

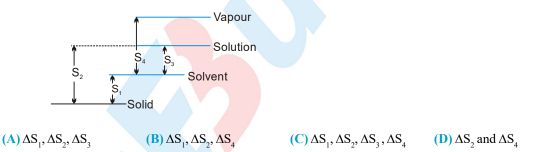


82.

- 92. Water and chlorobenzene are immiscible liquids. Their mixture boils at 90°C under a reduced pressure of 7.82 × 10⁴ Pa. The vapour pressure of pure water at 90°C is 7.03 × 10⁴ Pa. On weight percent basis, chlorobenzene in the distillate is equal to (mol. wt. of chlorobenzene is 112.5 g mol⁻¹)
 (A) 50
 (B) 60
 (C) 70
 (D) 80
- 93.Relative decrease in vapour pressure of an aqueous NaCl is 0.167. Number of moles of NaCl present in 180g of H2O is(A) 2 mol(B) 1 mol(C) 3 mol(D) 4 mol
- 94. What is the normal freezing point of the solution represented by the phase diagram ?



- **95.** Select correct statement :
 - (A) Solution has more molecular randomness than a pure solvent has, the entropy change between solution and solid is larger than the entropy change between pure solvent and solid
 - (B) Heats of fusion of solution and solvent are similar since similar forces of intermolecular forces are involved
 - (C) Sugar containing solution freezes at a lower temperature than pure water
 - (D) All are correct statements
- **96.** Some entropy change are represented in figure. Select correct entropy change.



- 97. Which statement comparing solutions with pure solvent is not correct
 - (A) A solution containing a non-volatile solute has a lower vapour pressure than pure solvent
 - (B) A solution containing a non-volatile solute has a lower boiling point than pure solvent
 - (C) A solution containing a non-volatile solute has a lower freezing point than pure solvent
 - (D) A solution will have a greater mass than an equal volume of pure solvent if the solute has a molar mass greater than the solvent
- 98. A colligative property of a solution depends on the :
 (A) arrangement of atoms in solute molecule
 (C) number of molecules of solute in solution
- 99. Which has maximum freezing point ?(A) 6g urea solution in 100 g H₂O
 - (C) 6g sodium chloride in 100 g H₂O

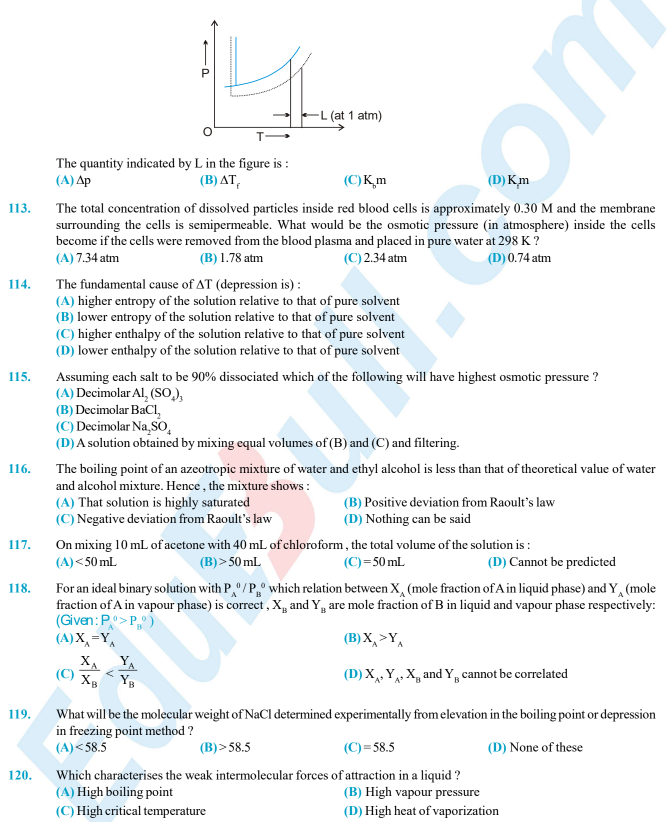
- (B) total number of molecules of solute and solvent
- (D) mass of the solute molecules
- (B) 6g acetic acid solution in 100g H₂O
- (**D**) All have equal freezing point



100.	Mole fraction of the toluene in the vapour phase whi and toluene ($p^\circ = 80$ Torr) having 2.0 mol of each is :	-	olution of benzene (p° = 120 Torr)
	(A) 0.50 (B) 0.25	(C) 0.60	(D) 0.40
101.	The vapour pressure of pure benzene, C_6H_6 at 50°C is benzene is required to prepare a solution of benzene (A) 0.377 (B) 0.605		
102.	 An azeotropic solution of two liquids has a boiling p (A) shows negative deviation from Raoult's law (B) shows positive deviation from Raoult's low (C) shows ideal behaviour (D) is saturated 	point lower than either of th	em when it :
103.	Van't Hoff factors of aqueous solutions of X, Y, Z are all three cases) (A) b.p. : X < Y < Z (C) osmotic pressure : X = Y = Z	 e 1.8, 0.8 and 2.5. Hence, the (B) f. p. Z < X < Y (D) v. p. : Y < X < Z 	eir (assume equal concentrations in
104.	 Select correct statements : (A) The fundamental cause of all colligative proper pure solvent (B) The freezing point of hydrofluoride solution is laterative (C) 1M glucose solution and 0.5 M NaCl solution and (D) All are correct statements 	arger than that of equimolal	hydrogen chloride solution
105.	The vapour pressure of a pure liquid A is 40 mmHg liquid B is 32 mmHg. Mole fraction of A in the soluti (A) 0.8 (B) 0.5		-
106.	Depression of freezing point of 0.01 molal aq. CH ₃ C 1.86°C. Assuming molality equal to molarity, pH of C (A) 2 (B) 3		. 1 molal urea solution freezes at –
107.	If relative decrease in vapour pressure is 0.4 for a solu (A) 60% (B) 50%	tion containing 1 mol NaCl i (C) 100%	n 3 mol H_2O , NaCl is % ionised. (D) 40%
108.	Which of the following azeotropic solutions has the (A) CHCl ₃ and CH ₃ COCH ₃ (C) CH ₃ CH ₂ OH and CH ₃ COCH ₃	 b.p. less than b.p. of the c (B) CS₂ and CH₃COCH₃ (D) CH₃CHO and CS₂ 	onstituents A and B ?
109.	Vapour pressure of CCl ₄ at 25°C is 143 mmHg. 0.5 g of mL of CCl ₄ (density = 1.538 g mL ⁻¹). Vapour pressur (A) 141.9 mmHg (B) 94.4 mmHg		mass = 65 mol^{-1}) is dissolved in 100 (D) 144.1 mmhg
110.	A 0.50 molal solution of ethylene glycol in water is u 1.86° per molal, at which temperature will the mixture (A) 1.56°C (B) -0.93 °C		freezing point constant of water is (D) 0.93°C
111.	The depression of freezing points of 0.05 molal aqua1. NaCl $2. K_2 SO_4$ Which one of the above compounds will exhibit the solution of the above compound solution of the solution of	3. $C_6 H_{12} O_6$	4. $Al_2(SO_4)_3$



112. The phase diagrams for the pure solvent (solid lines) and the solution (non-volatile solute, dashed line) are recorded below :





121.	 A teacher one day pointed out to his students the preculia fact that water is unique liquid which freezes exactly at 0° C and boils exactly at 100° C. He asked the students to find the correct statement based on this fact : (A) Water dissolves anything however sparingly the dissolution may be (B) Water is a polar molecule (C) Boiling and freezing temperatures of water were used to define a temperature scale (D) Liquid water is denser than ice 				
122.	 When an ideal binary solution is in equilibrium with its vapour , molar ratio of the two components in the solution and in the vapour phase is : (A) Same (B) Different (C) May or may not be same depending upon volatile nature of the two components (D) None of the above 				
123.		e lower than its usual melti 1 water			
124.	 Which of the following liquid pairs shows a positive deviation from Raoult's law ? (A) Acetone – chloroform (B) Benzene – methanol (C) Water – nitric acid (D) Water – hydrochloric acid 				
125.	The relationship between dissolved in 250 mL of wa (A) $P_1 > P_2 > P_3$	ater is :	when 10 g glucose (P_1) , 10 g (C) $P_2 > P_1 > P_3$	(D) $P_2 > P_3 > P_1$	
126.	The amount of ice that wi 9.3°C is : $(K'_f = 1.86 \text{ K mo})$ (A) 38.71 g		a solution containing 50 g of (C) 42 g	ethylene glycol in 200 g water to – (D) 42 mg	
127.	A liquid is in equilibrium equal : (A) Potential energy	with its vapours at its boil (B) Total energy	ing point. On the average the (C) Kinetic energy	e molecules in the two phases have (D) Intermolecular forces	
128.	On the basis of intermole (A) $CH_3OH > H_2 > CH_4$ (C) $CH_4 > CH_3OH > H_2$	cular forces predict the co	rrect order of decreasing bot (B) $CH_3OH > CH_4 > H_2$ (D) $H_2 > CH_4 > CH_3OH$	iling points of the compounds:	
129.	During depression of free (A) Liquid solvent, solid (B) Liquid solvent, solid (C) Liquid solute, solid so (D) Liquid solute, solid so	solvent solute blute	following are in equilibriun	n :	



Exercise # 2 Part # I [Multiple Correct Choice Type Questions]

For the given electrolyte $A_x B_y$, the degree of dissociation ' α ' can be given as 1.

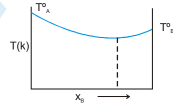
(A)
$$\alpha = \frac{i-1}{x+y-1}$$
 (B) $i = (1-\alpha) + x\alpha + y\alpha$ (C) $\alpha = \frac{1-i}{1-x-y}$ (D) None

A graph plotted between $\frac{P}{d}$ vs d (where P is osmotic pressure of solution of a solute of mol. wt. m and d is its 2. density at temperature T. Pick out the correct statements about the plots :

(A)
$$\left[\frac{P}{d}\right]_{d\to 0} = \frac{ST}{m}$$
 (B) The intercept of the plot $\frac{ST}{m}$
(C) The intercept of the plot = $\left[\frac{P}{d}\right]_{d\to 0}$ (D) $\left[\frac{P}{d}\right]_{d\to 0}$ is independent of temperature

(C) The intercept of the plot =
$$\left\lfloor \frac{P}{d} \right\rfloor_{d \to 0}$$

- Which of the following is/are correct for an ideal binary solution of two volatile liquids (eg. benzene & toluene)? 3. (A) Its vapour is always richer in the more volatile component (compared to the liquid).
 - (B) The liquid will gradually become richer in the less volatile component if such a mixture is boiled (distilled).
 - (C) The P_T (i.e. the total pressure) above the solution will be the sum of the vapor pressures of the two pure components.
 - (D) The boiling point of the solution will be less than the boiling points of the two components.
- The diagram given below represents boiling point composition 4. diagram of solution of component A and B, which is/are incorrect among the following? T(k) (A) The solution shows negative deviation
 - (B) A-B-interactions are stronger than A-A and B-B
 - (C) The solution is ideal solution
 - (D) The solution shows positive deviation.



- According to Henry's law, the partial pressure of gas (P'_{o}) is directly proportional to mole fraction of gas in 5. dissolved state , i.e., $P_{gas} = K_{H}$. X_{gas} where K_{H} is Henry's constant. Which are correct? (A) K_{H} is characteristic constant for a given gas-solvent system

 - (B) Higher is the value of $K_{\rm H}$, lower is solubility of gas for a given partial pressure of gas
 - (C) K_{H} has temperature dependence
 - **(D)** K_{H} increases with temperature
- Which of the following concentration factors can be calculated if the mole fraction and density of an aqueous 6. solution of HCl are known?

(A) Molality (B) Molarity

(C) Percent by mass

(D) Normality

- 7. Consider following solutions :
 - I: 1 M aqueous glucose solution
 - II: 1M aqueous sodium chloride solution
 - **III**: 1M aqueous ammonium phosphate solution
 - **IV**: 1M benzoic acid in benzene
 - Select correct statements for the above solutions :
 - (A) All are isotonic solutions
 - (B) III is hypertonic of I, II and IV
 - (C) IV is hypotonic of I, II and III
 - (D) II is hypotonic of III but hypertonic of I and IV



8. Which combination of (I) vapour pressure, (II) intermolecular forces and (III) ΔH_{vap} (latent heat of vaporisation) is matched correctly

	Ι	II	Ш
(A)	high	weak	small
(B)	high	strong	large
(C)	low	weak	large
(D)	low	strong	small

- 9. Which is/are true about ideal solutions ?
 - (A) The volume of mixing is zero
 - (B) The enthalpy of mixing is zero
 - (C) The entropy of mixing is zero
 - **(D)** The enthalpy of mixing is negative
- **10.** Freezing point lowering expression is
 - $\Delta T_{f} = K_{f} m \text{ (molality)}$

Which of the following assumptions are considered for the validity of above equation ?

- (A) The solution is dilute
- (B) The ΔH_f (latent heat of fusion of solvent) is independent of temperature between the actual and normal freezing point
- (C) The solid-phase consists of pure solvent when solution is allowed to cool
- (D) ΔT_{f} is not equal to $3K_{f}$ for 3 mol L⁻¹ solution
- 11. At 35°C, the vapour pressure of CS_2 is 512 mmHg, and of acetone is 344 mmHg. A solution of CS_2 and acetone in, which the mole fraction of CS_2 is 0.25, has a total vapour pressure of 600mmHg. Which of the following statements is/are correct?
 - (A) A mixture of 100 mL of acetone and 100 mL of CS, has a volume of 200 mL
 - (B) When acetone and CS₂ are mixed at 35°C, heat must be absorbed in order to produce a solution at 35°C
 - (C) Process of mixing is exothermic
 - (D) Entropy of mixing is zero
- **12.** Which is/are correct statement(s) ?
 - (A) When mixture is more volatile, there is positive deviation from Raoult's law
 - (B) When mixture is less volatile, there is negative deviation from Raoult's law
 - (C) Ethanol and water form ideal solution
 - (**D**) CHCl₃ and water form ideal solution
- 13. At 40°C, vapour pressure in Torr of methanol and ethanol solution is P = 119x + 135 where x is the mole fraction of methanol. Hence
 - (A) vapour pressure of pure methanol is 119 Torr
 - (B) vapour pressure of pure ethanol is 135 Torr
 - (C) vapour pressure of equimolar mixture of each is 127 Torr
 - (D) mixture is completely immiscible
- 14. Which one of the statements given below concerning properties of solutions, describes a colligative effect ?
 - (A) Boiling point of pure water decreases by the addition of ethanol
 - (B) Vapour pressure of pure water decreases by the addition of nitric acid
 - (C) Vapour pressure of pure benzene decreases by the addition of naphthalene
 - (D) Boiling point of pure benzene increases by the addition of toluene



15. For a dilute solution having molality m of a given solute in a solvent of mol.wt. M, b.pt. T_b and heat of vaporisation

per mole
$$\Delta H$$
; $\left[\frac{\partial T_b}{\partial m}\right]_{m \to 0}$ is equal to

(A) Molal elevation constant of solvent

(B)
$$\frac{RT_b^2M}{\Delta H_{vap}}$$
; where M in kg ΔH_{vap} and R in J mol⁻¹

(C)
$$\frac{RT_b^2M}{\Delta S_{vap}}$$
; where M in kg; ΔS_{vap} and R in J mol⁻

- (D) $\frac{RT_b^2M}{1000\Delta H_{vap}}$; where M in g; R and ΔH_{vap} expressed in same unit of heat.
- 16. Which facts are true when we use van't Hoff equation PV = CST for osmotic pressure P of dilute solutions ?
 - (A) The equation is identical to that of ideal gas equation
 - (B) The solute particles in solution are analogous to the gas molecules and the solvent is analogous to the empty space between the gas molecules
 - (C) Solute molecules are dispersed in the solvent the way the gas molecules are dispersed in empty space
 - (D) The equation is not identical to that of ideal gas equation
- 17. Which statements are correct about antifreeze mixtures to use to melt ice or snow on roads ?
 - (A) Antifreeze mixture of CaCl₂ + water (f.pt. -50° C) is preferred over KCl + water (f.pt. -10° C)
 - (B) The low freezing point of aq. CaCl, solution is due to its vant Hoff factor i = 3.
 - (C) The use of antifreeze for salt solutions causes major problems of corrosion of steel car bodies and reinforcement bars in concrete road structures.
 - (D) More is the amount of salt spreaded on road, easier is melting of ice.
- 18. The vapour pressure of a dilute solution of a solute is influenced by :
 - (A) Temperature of solution
- (B) Mole fraction of solute
- (C) M.pt. of solute (D) Degree of dissociation of solute
- 19. In the depression of freezing point experiment, it is found that the :
 - (A) Vapour pressure of the solution is less than that of pure solvent
 - (B) Vapour pressure of the solution is more than that of pure solvent
 - (C) Only solute molecules solidify at the freezing point
 - (D) Only solvent molecules solidify at the freezing point
- 20. Which of the following statements are correct for van't Hoff factor 'i' for a solution of weak electrolyte $A_x B_y$?
 - (A) i = 1 a + Xa + Ya
 - **(B)** i > 1 at normal dilution
 - (C) i increases more rapidly with dilution and attains a limiting value of (x + y) at infinite dilution
 - (D) The increase in 'i' with dilution is due to increase in molality of solution with dilution



	Part # II	>>	[Assertion & Reason Type Q	uestions]
	Directions : H	Each question	has 5 choices (A), (B), (C), (D) and (E) o	out of which only one is correct.
	(B) Statemen(C) Statemen	t-1 is True, Sta tt-1 is True, St tt-1 is False, S	atement-2 is True; Statement-2 is a corre atement-2 is True; Statement-2 is NOT a c atement-2 is False. tatement-2 is True. alse.	1
1.	Statement -1	The molar m	ass obtained for benzoic acid in benzene	e is found to be nearly 244.
	Statement -2	Benzoic acid	I has the formula HOOC	Соон
2.	Statement-1 :	The differen is decreased		tion of HCl and HF decreases as their molarity
	Statement-2 :	The extent o	f dissociation decreases steadily with ine	creasing dilution.
3.		boiling point if 'a' mL of 0. from the boi	t of the solution is no different from the be	another 'b' mL of 0.1 molal glucose solution, the oiling points of the samples prior to mixing but olal HF the boiling point of the mixture is different
4.			ions of urea and sucrose will freeze at s int of a solution is inversely proportion	-
5.			uric iodide is added to the aqueous solu vith KI forming complex ion [HgI ₄] ²⁻ .	tion of KI, the freezing point is raised.
6.			n of Glauber's salt is isotonic with 1 M s	solution of KNO ₃ . may or may not have same osmotic pressure.
7.	Statement-I :	0.1 M soluti temperature.		e than 0.1 M solution of glucose at same
	Statement-II	In solution,	NaCl dissociates to produce more numb	per of particles.
8.			vering of vapour pressure is equal to mo wering of vapour pressure is a colligative	
9.				solvent. g point when 1 mole of the solute is dissolved
10.		will also boi	l at the same temperature. olal concentration of aqueous solutions	°C, then decimolal solution of calcium chloride of electrolytes, the elevation of boiling point



Exercise # 3 Part # I [Matrix Match Type Questions]

1.

Column-I

Assuming all the solutes are non volatile and all solutions are ideal and neglect the hydrolysis of cation and anion.

- (A) 10 ml 0.1 M NaOH aqueous solution is added to 10 ml 0.1 M HCl aqueous solution
- (B) 10 ml 0.1 M NaOH aqueous solution is added to 10 ml 0.1 M CH₃COOH aqueous solution
- (C) 10 ml 0.1 M HCl aqueous solution is added to 10 ml 0.1 M NH₃ aqueous solution
- (D) 10 ml 0.1 M HCl aqueous solution is added to 10 ml 0.1 M KOH aqueous solution

2. Column I

(A) Acetone + $CHCl_3$ (B) Ethanol + Water (C) $C_2H_5Br + C_2H_5I$ (D) Acetone + Benzene

3.

Column-I (Properties)

(A) Relative lowering of vapour pressure

- (B) Elevation in boiling point
- (C) Freezing point
- (D) Osmotic pressure

Column-II

- (p) Osmotic pressure of solution increases
- (q) Vapour pressure of solution increases
- (r) Boiling point of solution increases
- (s) Freezing point of solution increases

Column II

(p) $\Delta S_{mix.} > 0$ (q) $\Delta V_{mix.} > 0$ (r) $\Delta H_{mix.} < 0$ (s) Maximum boiling azeotropes (t) Minimum boiling azeotropes

Column-II (Affecting factors)

- (p) Directly proportional to van't Hoff factor, i
- (q) Directly proportional to molality
- (r) Directly proportional to molarity
- (s) Indirectly proportional to lowering of vapour pressure

Part # II

[Comprehension Type Questions]

Comprehension #1

Addition of non–volatile solute to a solvent always increases the colligative properties such as osmotic pressure, ΔP , ΔT_{b} and ΔT_{f} . All these colligative properties are directly proportional to molality if solutions are dilute. The increases in colligative properties on addition of non–volatile solute is due to increase in number of solute particles.

1. For different aqueous solutions of 0.1 N NaCI, 0.1 M urea, 0.1 N Na₂SO₄ and 0.1 N Na₃PO₄ solution at 27° C. The correct statements are :

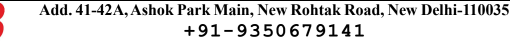
1. The order of osmotic pressure is, $NaCl > Na_2SO_4 > Na_3PO_4 > urea$

2.
$$\pi = \frac{\Delta T_b}{K_b} \times ST$$
 for urea solution

3. Addition of salt on ice increases its melting point

4. Addition of salt on ice brings in melting of ice earlier

(A) 2, 3, 4 (B) 2, 4 (C) 1, 2, 3 (D) 3, 4



- 2. 1 g mixture of glucose and urea present in 250 mL aqueous solution shows the osmotic pressure of 0.74 atm at 27°
 - C. Assuming solution to be dilute , which are correct ?
 - 1. Percentage of urea in mixture is 17.6.
 - 2. Relative lowering in vapour pressure of this solution is 5.41×10^{-4} .
 - 3. The solution will boil at 100.015° C, if K, of water is 0.5 K molality⁻¹.
 - 4. If glucose is replaced by same amount of sucrose, the solution will show higher osmotic pressure at 27°C.
 - 5. If glucose is replaced by same amount of NaCl, the solution will show lower osmotic pressure at 27°C.

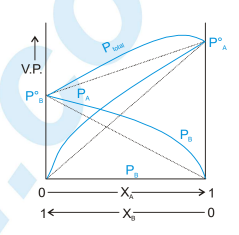
(A) 1, 2, 3 (B) 1, 2, 3, 5 (C) 2, 4, 5 (D) 1, 4, 5

Comprehension #2

Answer the questions (given below) which are based on the following diagram.

Consider some facts about the above phase diagram :

Vapour pressure diagram for real solutions of two liquids A and B that exhibit a positive deviation from Raoult's law. The vapour pressure of both A and B are greater than predicted by Raoult's law. The dashed lines represented the plots for ideal solutions.



- A: This is observed when A...B attractions are greater than average of A...B and B...B attraction: B: ΔH_{mix} = +ve, ΔV_{mix} = +ve C: Boiling point is smaller than expected such that vaporisation is increased D: Mixture is called azeotropic mixture Select correct facts (A) A, B, C
 (B) B, C, D
 (C) A, C, D
 (D) A, B, C, D
 (
- 2. Total vapour pressure of mixture of 1 mol of volatile component A ($p_{A}^{\circ} = 100 \text{ mmHg}$) and 3 mol of volatile component B ($p_{B}^{\circ} = 60 \text{ mmHg}$) is 75 mm. For such case :
 - (A) There is positive deviation from Raoult's law
 - (B) Boiling point has been lowered
 - (C) Force of attraction between A and B is smaller than that between A and A or between B and B.
 - (D) All the above statements are correct.

Comprehension #3

Following passage explains effect of temperature on the vapour pressure of liquid. Answer the questions given at the end.

Effect of temperature on Vapour pressure

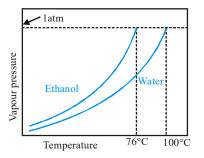
The quantity of heat required to evaporate a given liquid at constant temperature is defined as the heat of vaporisation. Variation of vapour pressure with temperature is given by

Clausius-Clapeyron equation.

$$\log_{e} P = -\frac{\Delta H_{vap}}{RT} + \log_{e} A$$



A liquid is said to be at its boiling temperature if its vapour pressure is equal to external pressure. Therefore, the boiling point of water in particular and of liquids in general decreases as altitude of a place increases where the external pressure is less than 1 atmosphere (normal b.p. of water is 373.15 K at 1 atmosphere)



On top of Mount Everest, for example, where the atmospheric pressure is only about 260 mm Hg, water boils at approximately 71°C. Conversely, if the external pressure on a liquid is greater than 1 atm., the vapour pressure necessary for boiling than normal boiling is reached later, and the liquid boils at a temperature greater than normal boiling point.

1. Clausius-Clapeyron equation can be written in the following form :

(A)
$$P = Ae^{-\Delta H_{vap}/RT}$$

(B) $\frac{dlog_{10}P}{dT} = \frac{\Delta H_{vap}}{2.303RT^2}$
(C) $\frac{dlog_eP}{dT} = -\frac{\Delta H_{vap}}{RT^2}$
(D) $P = Ae^{\Delta H_{vap}/RT}$

2. For a given liquid at a given temperature vapour pressure is given by :

(B) –9 mm

$$\log_{10} P \text{ (mm)} = -\frac{400(K)}{T} + 10$$

Vapour pressure of the liquid at 400 K is :

(A) 9 mm

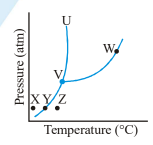
(C) 10⁹ mm

(D) 10^{-9} mm

3. Latent heat of vaporisation of the above case in the given temperature range is :

(A) -400 R		(B) 400 R
(C) -400×2.303 R		(D) 400×2.303 R

Comprehension #4



The phase diagram for a pure substance is shown above. Use this diagram and your knowledge about changes of phase to answer the following questions.

- 1. What does point V represent :
 - (A) point of equilibrium (B) point of fusion



⁽C) point of vaporisation (D) Triple point

- 2. What characteristics are specific to the system only at point V?
 - (A) Liquid ‡ˆˆ† Solid
 - (B) Solid ‡ˆˆ⁺ Vapour
 - (C) Liquid ‡[^]↑ ★ Vapour
 - (**D**) Solid [↑][↑][↑] Liquid [↑][↑][↑] Vapour
- 3. What happens if temperature is increased from X to Y at 1.0 atm ?
 - (A) solid is competely vaporised
 - (B) solid and vapour are in equilibrium
 - (C) solid and liquid are in equilibrium
 - (D) liquid and vapour are in equilibrium
- 4. Select correct statement (s) :
 - (A) curve VU is solid-liquid equilibrium curve
 - (B) curve VU has a positive slope
 - (C) curve VW is vapour pressure curve for liquid substance
 - (D) In the solid liquid mixture of the substance, solid will float
- 5. If the given substance is water then :
 - (A) curve VU would have negative slope
 - **(B)** in ice $\hat{\ddagger} \hat{}^{\dagger}$ water liquid mixture, ice will float
 - (C) as the temperature increases, pressure at which solid and liquid are in equilibrium, decreases
 - (D) increase in pressure at constant temperature causes ice to be converted to liquid water
- 6. If the triple point pressure of a substance is greater than 1 atm, we expect :
 - (A) the solid to sublime without melting
 - (B) the boiling point temperature to be lower than the triple point temperature
 - (C) the melting point of the solid to come at a lower temperature than the triple point
 - (D) that the substance cannot exist as a liquid
- 7. In a phase change (say solid to liquid or liquid to solid) $\Delta G = \Delta H T\Delta S$ where :
 - (A) ΔH is the enthalpy change associated with making or breaking the intermolecular attractions that hold solid and liquid together and ΔS is associated with change in disorder between the various phases.
 - (B) ΔH is associated with change in disorder while ΔS is associated with energy change
 - (C) both are associated with change in disorder
 - (D) both are associated with change in energy



Exercise # 4[Subjective Type Questions]1. (a) The vapour pressure of n-hexane and n-heptane at 273 K are 45.5 mm Hg and 11.4 mm Hg,

- respectively. What is the composition of a solution of these two liquids if its vapour pressure at 273 K is 37.3 mm Hg.
 - (b) The mole fraction of n-hexane in the vapour above a solution of n-hexane and n-heptane is 0.75 at 273 K. What is the composition of the liquid solution.
- A solution containing 30 g of a nonvolatile solute in exactly 90 g water has a vapour pressure of 21.85 mm Hg at 25°C. Further 18 g of water is then added to the solution. The resulting solution has vapour pressure of 22.18 mm Hg at 25°C. Calculate (a) molar mass of the solute, and (b) vapour pressure of water at 25°C.
- 3. The freezing point of ether was lowered by 0.60°C on dissolving 2.0 g of phenol in 100 g of ether. Calculate the molar mass of phenol and comment on the result. Given : $K_{f}(ether) = 5.12 \text{ K kg mol}^{-1}$.
- 4. The vapour pressure of ethanol and methanol are 44.5 mm Hg and 88.7 mm Hg, respectively. An ideal solution is formed at the same temperature by mixing 60 g of ethanol and 40 g of methanol. Calculate the total vapour pressure of the solution and the mole fraction of methanol in the vapour.
- 5. The vapour pressure of pure benzene at a certain temperature is 640 mm Hg. A non-volatile solid weighing 2.175 g is added to 39.0 g of benzene. The vapour pressure of the solution is 600 mm Hg. What is the molar mass of the solid substance?
- 6. Addition of 0.643 g of a compound to 50 mL of benzene (density : 0.879 g mL⁻¹) lower the freezing point from 5.51°C to 5.03°C. If K_f for benzene is 5.12 K kg mol⁻¹, calculate the molar mass of the compound.
- 7. A solution contains 3.22 g of $HClO_2$ in 47.0 g of water. The freezing point of the solution is 271.10 K. Calculate the fraction of $HClO_2$ that undergoes dissociation to H^+ and ClO_2^- . Given : K_f (water) = 1.86 K kg mol⁻¹.
- 8. A 0.1 molar solution of NaCl is found to be isotonic with 1% urea solution. Calculate (a) Van't Hoff factor, and (b) degree of dissociation of sodium chloride. Assume density of 1% urea equal to 1 g cm⁻³.
- 9. The addition of 3 g of a substance to 100 g CCl_4 (M = 154 g mol⁻¹) raises the boiling point of CCl_4 by 0.60°C. If K_b (CCl₄) is 5 K mol⁻¹ kg, calculate (a) the freezing point depression (b) the relative lowering of vapour pressure (c) the osmotic pressure at 298 K and (d) the molar mass of the substance. Given : K_f (CCl₄) = 31.8 K kg mol⁻¹ and ρ (solution) = 1.64 g cm⁻³.
- 10. A solution containing 0.122 kg of benzoic acid in 1 kg of benzene (b. pt. 353 K) boils at 354.5 K. Determine the apparent molar mass of benzoic acid (which dimerizes) in the solution and the degree of dimerization. Given : $\Delta_{van}H_{1m}$ (benzene) = 394.57 J g⁻¹.
- 11. A solution containing 0.011 kg of barium nitrate in 0.1 kg of water boils at 100.46°C. Calculate the degree of ionization of the salt. K_{b} (water) = 0.52 K kg mol⁻¹.
- 12. When 3.24 g of mercuric nitrate Hg (NO₃)₂ dissolved in 1 kg of water, the freezing point of the solution is found to be -0.0558°C. When 10.84 g of mercuric chloride HgCl₂ is dissolved in 1 kg of water, the freezing point of the solution is -0.0744°C. K_f = 1.86 mol⁻¹ K kg. Will either of these dissociate into ions in an aqueous solution ?
- 13. To 500 cm³ of water 3.0×10^{-3} kg of acetic acid is added. If 23% of acetic acid is dissociated, what will be the depression of freezing point? K_r and density of water are 1.86 K kg mol⁻¹ and 0.997 g cm⁻³, respectively.



- 14. A 0.01 m aqueous solution of $K_3[Fe(CN)_6]$ freezes at -0.062°C. What is the apparent percentage of dissociation? [K_r for water = 1.86]
- 15. The degree of dissociation of Ca $(NO_3)_2$ in a dilute aqueous solution containing 7 g of the salt per 100 g of water at 100°C is 70%. If the vapour pressure of water at 100°C is 760 mm, calculate the vapour pressure of the solution.
- 16. The vapour pressure of solution containing 6.69 g of $Mg(NO_3)_2$ dissolved in 100 g of water is 747 Torr at 373 K. Calculate the degree of dissociation of the salt in the solution.
- 17. At 353 K, the vapour pressure of pure ethylene bromide and propylene bromide are 22.93 and 16.93 k Nm⁻², respectively, and these compounds form a nearly ideal solution. 3 mol of ethylene bromide and 2 mole of propylene bromide are equilibrated at 553 K and a total pressure of 20.4 k Nm⁻².
 - (a) What is the composition of the liquid phase?
 - (b) What amount of each compound is present in the vapour phase?
- 18. The vapour pressure of two pure liquids, A and B, that form an ideal solution are 300 and 800 torr, respectively, at temperature T. A mixture of the vapour of A and B for which the amount fraction of A is 0.25 is slowly compressed at temperature T. Calculate :
 - (a) The composition of the first drop of the condensate,
 - (b) The total pressure when this drop is formed,
 - (c) The composition of the solution whose normal boiling point is T.
 - (d) The pressure when only the last bubble of vapour remains.
 - (e) The composition of the last bubble.
- 19. A certain mass of a substance, when dissolved in 100 g C_6H_6 , lowers the freezing point by 1.28°C. The same mass of solute dissolved in 100 g water lowers the freezing point by 1.40°C. If the substance has normal molecular weight in benzene and is completely ionised in water, into how many ions does it dissociate in water? K_f for H_2O and C_6H_6 are 1.86 and 5.12 K kg mol⁻¹.
- 20. The cryoscopic constant for acetic acid is 3.6 K kg/mol. A solution of 1 g of a hydrocarbon in 100 g of acetic acid freezes at 16.14°C instead of the usual 16.60°C. The hydrocarbon contains 92.3% carbon. What is the molecular formula?
- 21. A radiator was filled with 10 L of water to which 2.5 L of methanol (density = 0.8 g.mL^{-1}) were added. At 9 : 00 pm, the vehicle is parked outdoors where the temperature is 0°C. The temperature is decreasing at a uniform rate of 0.5° C / min. Upto what time will there be no danger to the radiator of the car. K_f(water) = 1.86 kg.mo^{-1} K. Assume methanol to be non-volatile.
- 22. Sea water is found to contain 5.85% NaCl and 9.50% MgCl₂ by weight of solution. Calculate its normal boiling point assuming 80% ionisation for NaCl and 50% ionisation of MgCl₂[K_{k} (H₂O) = 0.51 kg mol⁻¹K].
- 23. Find the freezing point of a glucose solution whose osmotic pressure at 25°C is found to be 30 atm. $K_{f}(water) = 1.86 \text{ kg.mol}^{-1}$. K.
- 24. The latent heat of fusion of ice is 80 calories per gram at 0°C. What is the freezing point of a solution of KC● in water containing 7.45 grams of solute 500 grams of water, assuming that the salt is dissociated to the extent of 95%?
- 25. At 300 K, two solutions of glucose in water of concentration 0.01 M and 0.001 M are separated by semipermeable membrane. Pressure needs to be applied on which solution, to prevent osmosis? Calculate the magnitude of this applied pressure?
- 26. At 10°C, the osmotic pressure of urea solution is 500 mm. The solution is diluted and the temperature is raised to 25°C, when the osmotic pressure is found to be 105.3 mm. Determine extent of dilution.



- 27. When cells of the skeletal vacuole of a frog were placed in a series of NaCl solutions of different concentration at 25°C, it was observed microscopically that they remained unchanged in 0.7% NaCl solution, shrank in more cocentrated solutions, and swelled in more dilute solutions. Water freezes from the 0.7% salt solution at -0.406°C. What is the osmotic pressure of the cell cytoplasm at 25°C ? $K_r = 1.86 \text{ kg mol}^{-1} \text{ K}.$
- 28. An ideal solution of two volatile liquid A and B has a vapour pressure of 402.5 mmHg, the mole fraction of A in vapour & liquid state being 0.35 & 0.65 respectively. What are the vapour pressure of the two liquid at this temperature.
- 29. Dry air was drawn through bulbs containing a solution of 40 grams of urea in 300 grams of water, then through bulbs containing pure water at the same temperature and finally through a tube in which pumice moistened with strong H_2SO_4 was kept. The water bulbs lost 0.0870 grams and the sulphuric acid tube gained 2.036 grams. Calculate the molecular weight of urea.
- **30.** A 0.1 M solution of potassium ferrocyanide is 46% dissociated at 18°C. What will be its osmotic pressure?
- **31.** At 100°C, benzene & toluene have vapour pressure of 1375 & 558 Torr respectively. Assuming these two form an ideal binary solution that boils at 1 atm & 100°C. What is the composition of vapour issuing at these conditions?
- 32. Two beaker A and B present in a closed vessel. Beaker A contains 152.4 g aqueous solution of urea, containing 12 g of urea. Beaker B contains 196.2 g glucose solution, containing 18 g of glucose. Both solutions allowed to attain the equilibrium. Determine wt. % of glucose in it's solution at equilibrium:
- 33. The vapour pressure of two pure liquids A and B, that form an ideal solution are 100 and 900 mm Hg respectively at temperature T. This liquid solution of A and B is composed of 1 mole of A and 1 mole of B. What will be the pressure, when 1 mole of mixture has been vaporized ?
- 34. The addition of 3 g of substance to 100 g CCl_4 (M = 154 g mol⁻¹) raises the boiling point of CCl_4 by 0.60°C of K_b (CCl_4) is 5.03 kg mol⁻¹ K. Calculate :
 - (a) the freezing point depression
 - (b) the relative lowering of vapour pressure
 - (c) the osmotic pressure at 298 K
 - (d) the molar mass of the substance
 - Given $K_f(CCl_4) = 31.8 \text{ kg mol}^{-1} \text{ K}$ and ρ (density) of solution = 1.64 g/cm³.
- 35. A one litre solution is prepared by dissolving some solid lead-nitrate in water. The solution was found to boil at 100.15°C. To the resulting solution 0.2 mole NaCl was added. The resulting solution was found to freeze at -0.83°C. Determine solubility product of PbCl₂. Given K_b= 0.5 and K_f = 1.86. Assume molality to be equal to molarity in all case.
- 36. A protein has been isolated as sodium salt with their molecular formula $Na_x P$ (this notation means that xNa^+ ions are associated with a negatively charged protein P^{-x}). A solution of this salt was prepared by dissolving 0.25 g of this sodium salt of protein in 10 g of water and ebulliscopic analysis revealed that solution boils at temperature 5.93×10^{-3} °C higher than the normal boiling point of pure water. K_b of water 0.52 kg mol⁻¹. Also elemental analysis revealed that the salt contain 1% sodium metal by weight. Deduce molecular formula and determine molecular weight of acidic form of protein H_P.



- 37. The vapour pressure of two miscible liquids (A) and (B) are 300 and 500 mm of Hg respectively. In a flask 10 mole of (A) is mixed with 12 mole of (B). However, as soon as (B) is added, (A) starts polymerising into a completely insoluble solid. The polymerisation follows first-order kinetics. After 100 minute, 0.525 mole of a solute is dissolved which arrests the polymerisation completely. The final vapour pressure of the solution is 400 mm of Hg. Estimate the rate constant of the polymerisation reaction. Assume negligible volume change on mixing and polymerisation, and ideal behaviour for the final solution.
- 38. If 20 mL of ethanol (density = 0.7893 g/mL) is mixed with 40 mL water (density = 0.9971 g/mL) at 25°C, the final solution has density of 0.9571 g/mL. Calculate the percentage change in total volume of mixing. Also calculate the molality of alcohol in the final solution.
- 39. Mixture of two liquids A and B is placed in cylinder containing piston. Piston is pulled out isothermally so that volume of liquid decreases but that of vapour increases. When negligibly small amount of liquid was remaining, the mole fraction of A in vapour is 0.4. Given $P_{A}^{\circ} = 0.4$ atm and $P_{B}^{\circ} = 1.2$ atm at the experimental temperature. Calculate the total pressure at which the liquid has almost evaporated. (Assume ideal behaviour)
- 40. 1.5 g of monobasic acid when dissolved in 150 g of water lowers the freezing point by 0.165°C. 0.5 g of the same acid when titrated, after dissolution in water, requires 37.5 mL of N/10 alkali. Calculate the degree of dissociation of the acid (K_f for water = 1.86° C mol⁻¹).
- 41. 30 mL of CH₃OH (d = 0.7980 g cm⁻³) and 70 mL of H₂O (d = 0.9984 g cm⁻³) are mixed at 25°C to form a solution of density 0.9575 g cm⁻³. Calculate the freezing point of the solution. $K_t(H_2O)$ is 1.86 kg mol⁻¹ K. Also calculate its molarity.
- 42. Vapour pressure of C_6H_6 and C_7H_8 mixture at 50°C is given by P (mm Hg) = 179 $X_B + 92$, where X_B is the mole fraction of C_6H_6 . A solution is prepared by mixing 936 g benzene and 736 g toluene and if the vapours over this solution are removed and condensed into liquid and again brought to the temperature of 50°C, what would be mole fraction of C_6H_6 in the vapour state ?
- 43. When the mixture of two immiscible liquids (water and nitrobenzene) boils at 372 K and the vapour pressure at this temperature are 97.7 kPa (H_2O) and 3.6 kPa ($C_6H_5NO_2$). Calculate the weight % of nitrobenzene in the vapour.
- 44. The molar volume of liquid benzene (density = 0.877 g mL^{-1}) increase by a factor of 2750 as it vaporizes at 20°C and that of liquid toluene (density = 0.867 g mL^{-1}) increases by a factor of 7720 at 20°C solution has a vapour pressure of 46.0 torr. Find the mole fraction of benzene in the vapour above the solution.
- 45. Calculate the boiling point of a solution containing 0.61 g of benzoic acid in 50 g of carbon disulphide assuming 84% dimerization of the acid. The boiling point and K_{h} of CS, are 46.2°C and 2.3 K kg mol⁻¹, respectively.
- 46. At 25°C, 1 mol of A having a vapour pressure of 100 torr and 1 mol of B having a vapour pressure of 300 torr were mixed. The vapour at equilibrium is removed, condensed and the condensate is heated back to 25°C. The vapour now formed are again removed, recondensed and analyzed. What is the mole fraction of A in this condensate ?
- 47. The vapour pressure of a certain liquid is given by the equation :

 $Log_{10}P = 3.54595 - \frac{313.7}{T} + 1.40655 log_{10} T$ where P is the vapour pressure in mm and T = Kelvin Temperature.

Determine the molar latent heat of vaporisation as a function of temperature. Calculate the its value at 80 K.



- **48.** A very dilute saturated solution of a sparingly soluble salt A_3B_4 has a vapour pressure of 20 mm of Hg at temperature T, while pure water exerts a pressure of 20.0126 mm Hg at the same temperature. Calculate the solubility product constant of A_3B_4 at the same temperature.
- 49. The molar volume of liquid benzene (density = 0.877 g mL^{-1}) increases by a factor of 2750 as it vaporises at 20°C while in equilibrium with liquid benzene. At 27°C when a non volatile solute (that does not dissociate) is dissolved in 54.6 cm³ of benzene vapour pressure of this solution, is found to be 98.88 mm Hg. Calculate the freezing point of the solution.

Given : Enthalpy of vaporization of benzene (l) = 394.57 J/gMolal depression constant for benzene = $5.12 \text{ K kg. mol}^{-1}$ Freezing point of benzene = 278.5 K.

- 50. The freezing point of 0.02 mol fraction solution of acetic acid (A) in benzene (B) is 277.4 K. Acetic acid exists partly as a dimer $2A = A_2$. Calculate equilibrium constant for the dimerisation. Freezing point of benzene is 278.4 K and its heat of fusion ΔH_f is 10.042 kJ mol⁻¹.
- 51. Tritium, T (an isotope of H) combines with fluorine to form weak acid TF, which ionizes to give T⁺. Tritium is radioactive and is a β-emitter. A freshly prepared aqueous solution of TF has pT (equivalent of pH) of 1.5 and freezes at -0.372°C. If 600 mL of freshly prepared solution were allowed to stand for 24.8 years, calculate (i) ionization constant of TF. (ii) Number of β-particles emitted.

(Given : K_f for water = 1.86 kg mol K^{-1} , $t_{1/2}$ for tritium = 12.4 years.)

- 52. An ideal solution was prepared by dissolving some amount of cane sugar (non-volatile) in 0.9 moles of water. The solution was then cooled just below its freezing temperature (271 K), where some ice get separated out. The remaining aqueous solution registered a vapour pressure of 700 torr at 373 K. Calculate the mass of ice separated out, if the molar heat of fusion of water is 96 kJ.
- 53. The freezing point depression of a 0.109 M aq. solution of formic acid is -0.21°C. Calculate the equilibrium constant for the reaction,

HCOOH (aq) $\frac{1}{2}$ $\hat{}$ $\hat{}$ H⁺ (aq) + HCOO^{Θ} (aq)

 K_{f} for water = 1.86 kg mol⁻¹ K

54. 10 g of NH_4Cl (mol. weight = 53.5) when dissolved in 1000 g of water lowered the freezing point by 0.637°C. Calculate the degree of hydrolysis of the salt if its degree of dissociation of 0.75. The molal depression constant of water is 1.86 kg mol⁻¹ K.



E	xercise # 5	Part # I [Prev	ious Year Questions]	[AIEEE/JEE-MAIN]
1.	6.02×10^{20} molecules	of urea are present in 100 ml	of its solution. The concen	tration of urea solution is : [AIEEE-2004]
	(A) 0.001 M	(B) 0.01 M	(C) 0.02 M	(D) 0.1 M.
2.	Which one of the follo (A) $0.01 \text{ M Na}_2\text{SO}_4$	(B) 0.01 M KNO ₃	exhibit highest boiling poin (C) 0.015 M urea	nt ? [AIEEE-2004] (D) 0.015 M glucose
3.	-	the same solvent have : but different freezing point ame freezing points	(B) same freezing point(D) different boiling and	[AIEEE-2005] but different boiling point d freezing points
4.		tance (non electrolyte) are mix lution. What is the molarity o (B) 1.50 M	<u> </u>	: 480 ml of 1.5 M first solution + 520 [AIEEE-2005] (D) 2.70 M
5.				re of benzene is 75 torr and that of n containing 78 g of benzene and 46 [AIEEE-2005] (D) 53.5
6.	If α is the degree of dis	ssociation of Na ₂ SO ₄ , the var	t Hoff's factor (i) used for	calculating the molecular mass is :
	(A) $1 + \alpha$	(B) $1 - \alpha$	(C) $1 + 2\alpha$	[AIEEE-2005] (D) 1-2α.
7.	Density of a 2.05 M so (A) 3.28 mol Kg ⁻¹	lution of acetic acid in water (B) 2.28 mol Kg ⁻¹	is 1.02 g/mL. The molality (C) 0.44 mol Kg ⁻¹	of the solution is [AIEEE-2006] (D) $1.14 \mod \text{Kg}^{-1}$
8.	-			m at 300 K. The vapour pressure of apour pressure (in mm) at the same [AIEEE-2007] (D) 300
9.	A 5.25% solution of a	substance is isotonic with a	1.5% solution of urea (mo	blar mass = 60g mol^{-1}) in the same n ⁻³ , molar mass of the substance will [AIEEE-2007] (D) 15.0 g mol ⁻¹
10.		water at 20° C is 17.5 mm Hg of the resulting solution will (B) 16.500 mm Hg		0 ₆) is added to 178.2 g of water at 20° [AIEEE-2008] (D) 17.675 mm Hg
11.	solution of 'A' and 'B' bo	ressure of pure liquid 'A' is 52 bils at 80° C and 1 atm pressur	•	quid 'B' is 1000 mm Hg. If a mixture nixture is (1atm = 760 mm Hg) [AIEEE-2008]
	(A) 34 mol percent	(B) 48 mol percent	(C) 50 mol percent	(D) 52 mol percent
12.	correct regarding the b(A) The solution is nor(B) The solution in nor	ehaviour of the solution ? h-ideal, showing +ve deviatio h-ideal, showing -ve deviatio ve deviation while ethanol sh	n from Raoult's Law. n from Raoult's Law.	h one of the following statement is [AIEEE-2009] aoult's Law.



13.	Two liquids X and Y form an ideal solution. At 300K, vapour pressure of the solution containing 1 mol of X and 3 mol of Y is 550 mmHg. At the same temperature, if 1 mol of Y is further added to this solution, vapour pressure of the solution increases by 10 mmHg. Vapour pressure (in mmHg) of X and Y in their pure states will be, respectively : [AIEEE-2009]			
	(A) 300 and 400	(B) 400 and 600	(C) 500 and 600	(D) 200 and 300
14.	change in freezing po $(K_f = 1.86 \text{ K kg mol}^{-1})$	int of water (ΔT_f) , when	0.01 mole of sodium sulp	s and anions in aqueous solution, the hate is dissolved in 1 kg of water, is [AIEEE-2010]
15.	(heptane and octane) as	re 105 kPa and 45 kPa resp	ectively. Vapour pressure of	(D) 0.0186 K pressures of the two liquid components of the solution obtained by mixing 25.0 and of octane = 114 g mol^{-1}
16.	-		(C) 96.2 kPa e radiator holds 1.0 kg of wa of the solution lowered to – (C) 39 g	[AIEEE-2010] (D) 144.5 kPa ater, how may grams of ethylene glycol -2.8°C ? [AIEEE 2012] (D) 27 g
17.	18g glucose ($C_6H_{12}O_6$)	is added to 178.2 g water.	The vapor pressure of wat	er (in torr) for this aqueous solution is: [Jee (Mains) 2016]
18.		mer in benzene, percentage	(C) 759.0 C when 0.2 g of acetic acid is e association of acetic acid (C) 74.6 %	(D) 7.6 added to 20 g of benzene. If acetic acid
19.				show the highest freezing point ?
17.	i or i motar aqueous s		inpounds, which one will s	[Jee (Mains) 2018]
	(A) [Co(H,O),Cl]Cl,.H,	O (B) [Co(H,O),Cl,]Cl.2]	H,O (C) [Co(H,O),Cl,].3H	
20.	An aqueous solution c	contains an unknown conc ins to precipitate. The fina	entration of Ba ²⁺ . When 5	0 mL of a 1 M solution of Na_2SO_4 is blubility product of $BaSO_4$ is 1×10^{-10} . [JEE(Main) 2018]
	(A) 2×10^{-9} M	(B) 1.1×10^{-9} M	(C) 1.0×10^{-10} M	(D) $5 \times 10^{-9} \mathrm{M}$
	Part # II	Previous Year Qu	estions][IIT-JEE AD	VANCED]
1.	A 0.004 M solution of percentage dissociation		0.010 M solution of gluco	se at same temperature. The apparent [JEE 2004]
2.	(A) 25% 1.22 g of benzoic acid i in acetone increases by	(B) 50% s dissolved in 100 g of ace (0.17°C, while that of, in	the benzene increases by 0	(D) 85% separately. Boiling point of the solution 0.13° C; K _b for acetone and benzene is a two cases and justify your answer.
3.		g point of a solution of 13.4 f CuCl ₂ = 134.4 and $K_b = 0$ (B) 0.05		[JEE 2004] er using the following information, will [JEE 2005] (D) 0.2
4.		ic acid (C ₁₁ H ₈ O ₂) is disso pserved. The van't Hoff fac		$C_{\rm f} = 1.72 \text{ K kg mol}^{-1}$), a freezing point [JEE-2007]
	(A) 0.5	(B) 1	(C) 2	(D) 3
	Add. 41-4	42A, Ashok Park Ma	in, New Rohtak Road	l, New Delhi-110035

3

d. 41-42A, Ashok Park Main, New Rohtak Road, New Delhi-110035 + 91 – 9350679141

Comprehension

Properties such as boiling point, freezing point and vapour pressure of a pure solvent change when solute molecules are added to get homogeneous solution. These are called colligative properties. Application of colligative properties are very useful in day-to-day life. One of its example is the use of ethylene glycol and water mixture as anti-freezing liquid in the radiator of automobiles

A solution M is prepared by mixing ethanol and water. The mole fraction of ethanol in the mixture is 0.9

- Given: Freezing point depression constant of water $(K_f^{water}) = 1.86 \text{ K kg mol}^{-1}$
 - Freezing point depression constant of ethanol ($K_{f}^{ethanol}$) = 2.0 K kg mol⁻¹
 - Boiling point elevation constant of water $(K_{b}^{water}) = 0.52 \text{ K kg mol}^{-1}$
 - Boiling point elevation constant of ethanol ($K_b^{ethanol}$) = 1.2 K kg mol⁻¹
 - Standard freezing point of water = 273 K
 - Standard freezing point of ethanol = 155.7 K
 - Standard boiling point of water = 373 K
 - Standard boiling point of ethanol = 351.5 K
 - Vapour pressure of pure water = 32.8 mm Hg
 - Vapour pressure of pure ethanol = 40 mm Hg
 - Molecular weight of water = 18 g mol^{-1}
 - Molecular weight of ethanol = 46 g mol^{-1}

In answering the following questions, consider the solution to be ideal dilute solutions and solutes to be non-volatile and non-dissociative.

5.	The freezing point of the s	oluti <mark>on M is</mark>			[JEE 2008]
	(A) 268.7 K	(B) 268.5 K	(C) 234.2 K	(D) 150.9 K	
6.	The vapour pressure of th	e solution M is			[JEE 2008]]
	(A) 39.3 mm Hg	(B) 36.0 mm Hg	(C) 29.5 mm Hg	(D) 28.8 mm Hg	

- 7. Water is added to the solution M such that the mole fraction of water in the solution becomes 0.9. The boiling point of this solution is [JEE 2008]
 - (A) 380.4 K (B) 376.2 K (C) 375.5 K
- 8. The Henry's law constant for the solubility of N_2 gas in water at 298 K is 1.0×10^5 atm. The mole fraction of N_2 in air is 0.8. The number of moles of N_2 from air dissolved in 10 moles of water of 298 K and 5 atm pressure is :

[**JEE 2009**]

(D) 354.7 K

(A) 4×10^{-4}	(B	4.0×10^{-5}		(C) 5.0×10^{-4}	(D) 4.0×10^{-6}	
The freezing point $(K_{e} = 1.86 \text{ K kg mol}^{-1})$		f a solution	containing	0.1 g of K ₃ [F	e(CN) ₆] (Mol	. Wt. 329) in	100 g of water [JEE 2011]
$(A) - 2.3 \times 10^{-2}$		-5.7×10^{-2}		$(C) - 5.7 \times 10^{-3}$	3	(-1.2×10^{-2})	



9.

10.For a dilute solution containing 2.5 g of a non- volatile non- electrolyte solute in 100 g of water, the elevation in
boiling point at 1 atm pressure is 2°C. Assuming concentration of solute is much lower than the concentration of
solvent, the vapour pressure (mm of Hg) of the solution is (take $K_b = 0.76 \text{ K kg mol}^{-1}$)[IIT 2012]

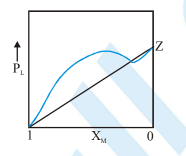
(A) 724 (B) 740 (C) 736 (D) 718

11. The mole fraction of a solute in a solution is 0.1. At 298 K, molarity of this solution is the same as its molality. Density

of this solution at 298 K is 2.0 g cm⁻³. The ratio of the molecular weights of the solute and solvent, $\left(\frac{MW_{solute}}{MW_{solvent}}\right)$, is

[JEE ADVANCED 2016]

12. For a solution formed by mixing liquids L and M, the vapour pressure of L plotted against the mole fraction of M in solution is shown in the following figure. Here x_L and x_M represent mole fractions of L and M, respectively, in the solution. The correct statement(s) applicable to this system is(are) [JEE ADVANCED 2017]



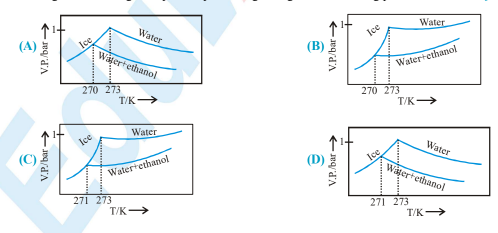
(A) The point Z represent vapour pressure of pure liquid M and Raoult's law is obeyed from $x_1 = 0$ to $x_1 = 1$

(B) Attractive intermolecular interactions between L–L in pure liquid L and M–M in pure liquid M are stronger than those between L-M when mixed in solution

(C) The point Z represents vapour pressure of pure liquid M and Raoult's law is obeyed when $x_1 \rightarrow 0$

(D) The point Z represents vapour pressure of pure liquid L and Raoult's law is obeyed when $x_L \rightarrow 1$

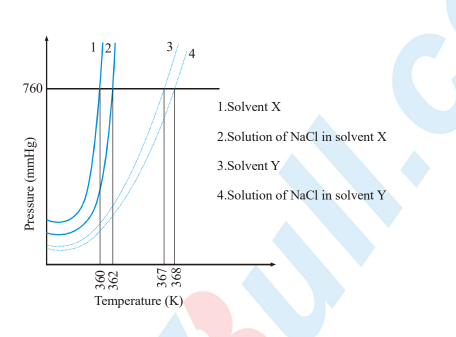
Pure water freezes at 273 K and 1 bar. The addition of 34.5 g of ethanol of 500 g of water changes the freezing point of the solution. Use the freezing point depression constant of water as 2 K kg mol⁻¹. The figures shown below represent plots of vapour pressure (V.P.) versus temperature (T). [molecular weight of ethanol is 46 g mol⁻¹] Among the following the option representing change in the freezing point is [JEE ADVANCED 2017]





[JEE(ADVANCED) 2018]

- 14. Liquids A and B form ideal solution over the entire range of composition. At temperature T, equimolar binary solution of liquids A and B has vapour pressure 45 Torr. At the same temperature, a new solution of A and B having mole fractions x_A and x_B , respectively, has vapour pressure of 22.5 Torr. The value of x_A / x_B in the new solution is _____. (given that the vapour pressure of pure liquid A is 20 Torr at temperature T) [JEE(ADVANCED) 2018]
- **15.** The plot given below shows P–T curves (where P is the pressure and T is the temperature) for two solvents X and Y and isomolal solutions of NaCl in these solvents. NaCl completely dissociates in both the solvents.



On addition of equal number of moles of a non-volatile solute S in equal amount (in kg) of these solvents, the elevation of boiling point of solvent X is three times that of solvent Y. Solvent S is known to undergo dimerization in these solvents. If the degree of dimerization is 0.7 in solvent Y, the degree of dimerization in solvent X is ____.



Т	VOC	CCT
$\mathbf{\Gamma}$		

SECTION - I : STRAIGHT OBJECTIVE TYPE

1. Three solutions are prepared by adding 'w' gm of 'A' into 1 kg of water, 'w' gm 'B' into another 1 kg of water and 'w' gm of 'C' in another 1 kg of water (A,B,C are non electrolytic). Dry air is passed from these solutions in sequence $(A \rightarrow B \rightarrow C)$. The loss in weight of solution A was found to be 2gm while solution B gained 0.5 gm and solution C lost 1 gm. Then the relation between molar masses of A,B and C is :

(A) MA: MB: MC = 4:3:5	(B) MA : MB : MC = $\frac{1}{4} : \frac{1}{3} : \frac{1}{5}$
(C) MC > MA > MB	(D)MB > MA > MC

- Barium ions, CN- and Co⁺² form an ionic complex. If this complex is 75% ionised in aqueous solution with Vant Hoff factor (i) equal to four and paramagnetic moment is found to be 1.73 BM(due to spin only) then the hybridisation state of Co(II) in the complex will be :

 (A) sp³d
 (B) d²sp³
 (C) sp³d²
 (D) dsp³
- **3.** The incorrect statement is :
 - (A) Vapour pressure of a liquid always increases by increasing the temperature.
 - (B) Vapour pressure only depends on temperature and not on the nature of the substance.
 - (C) Vapour pressure does not depend on the quality of the liquid taken and the surface area of the liquid.
 - (D) Vapour pressure is not a colligative property and is independent of the concentration of a liquid.
- 4. Relative decrease in in vapour pressure of an aqueous solution containing 2 moles [Cu(NH₂),Cl]Cl in 3 moles H₂O is 0.50. On reaction with AgNO₃, this solution will form (A) 1 mol AgCl (B) 0.25mol AgCl (D) 0.40 mol AgCl (C) 2 mol AgCl A solution of x moles of sucrose in 100gms of water freezes at -0.2° C. As ice separates the freezing point goes down 5. to 0.25°C. How many grams of ice would have separated? (C) 25 grams (A) 18 grams (B) 20 grams (D) 23 grams 6. A sample of air is saturated with benzene (vapour pressure = 100 mm Hg at 298K) at 298 K, 750mm Hg pressure. If it is isothermally compressed to one third of its initial volume, the final pressure of the system is (D) 1950 torr (A) 2250 torr **(B)** 2150 torr (C) 2050 torr Available solutions are 1L of 0.1 M NaCl and 2L of 0.2 M CaCl,. Using only these two solutions what maximum 7. volume of a solution can be prepared having [Cl] = 0.34 M exactly. Both electrolytes are strong (A) 2.5L **(B)**2.4L (C)2.3L (D) None of these

8. Calculate the osmotic pressure of the solution prepared in the above ques T = 300K, (R=0.082 L atm mol-1 K-1) (A) 10.8 atm (B)12.8 atm (C) 5.6 atm (D) None of these

9. Consider equimolal aqueous solutions of NaHSO₄ and NaCl with Δ Tb and Δ T'b as their respective boiling point

elevations. The value of $\lim_{m\to 0} \frac{\Delta T_b}{\Delta T'_b}$ will be

- 10.
- The vapour pressure of benzene, toulene and a xylene are 75 Torr, 22 Torr and 10 Torr at 20°C. Which of the following is not a possible value of the vapour pressure of an equimolar binary/ternary solution of these at 20°C? Assume all form ideal solution with each other.
 - (A) 48.5 (B) 16 (C) $35\frac{2}{3}$ (D) 53.5

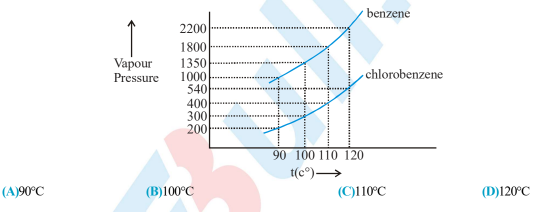


11. $3.24g \text{ of Hg(NO}_{3})_2$ (molar mass = 324) dissolved in 1000g of water constitutes a solution having a freezing point of -0.0558° C while 21.68 g of HgCl₂ (molar mass = 271) in 2000g of water constitutes a solution with a freezing point of

-0.0744°C. The Kf for the water is $1.86 \frac{K-kg}{mol}$. About the state of ionization of these two solids in water it can be

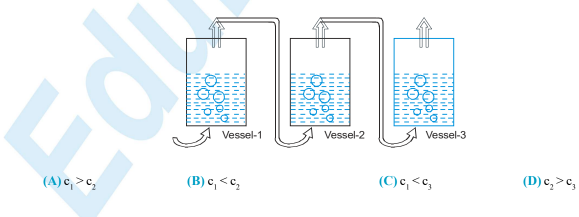
inferred that :

- (A) $Hg(NO_3)_2$ and $HgCl_2$ both are completely ionized.
- (B) $Hg(NO_3)_2$ is fully ionized but $HgCl_2$ is fully unionized.
- (C) $Hg(NO_3)_2$ and $HgCl_2$ both are completely unionized.
- (D) $Hg(NO_3)_2$ is fully unionized and $HgCl_2$ is fully ionized.
- For a solution of 0.849 g of mercurous chloride in 50 g og HgCl2 (●) the freezing point depression is 1.24°C. Kf for HgCl2 is 34.3. What is the state of mercurous chloride in HgCl2? (Hg-200, Cl-35.5).
 (A) as Hg2Cl₂ molecules
 (C)As Hg+ and Cl- ions
 (D)As Hg²⁺ and Cl- ions
- 13. Assuming that the formation of an ideal solution, determine the boiling point of a mixture containing 1560 g benzene (molar mass = 78) and 1125 g chlorobenzene (molar mass = 112.5) using the following against an external pressure of 1000 torr.



SECTION - II : MULTIPLE CORRECT ANSWER TYPE

14. Dry air is slowly passed through three solutions of different concentration, c_1, c_2, c_3 ; each containing (non volatile) NaCl as solute and water as solvent, as shown in the figure. If the vessel 2 gains weight, then





- In which of the following pairs of solution will have the values of Vant hoff factors can be the same?
 (A) 0.05 M K₄[Fe(CN)₆] and 0.10 M FeSO₄
 (B) 0.10 M K4[Fe(CN)₆] and 0.05 M FeSO₄(NH₄)₂SO₄.6H₂O
 (C) 0.20 M NaCl and 0.10 M BaCl₂
 - (D) $0.05 \text{ M FeSO}_{4}(\text{NH}_{4})$, SO₄.6H₂O and $0.02 \text{ M KCl.MgCl}_{2}$.6H₂O
- 16. Vapour pressure of solution containing 6g of a non-volatile solute in 180 g water is 20.0 Torr. If 1 mole water is further added vapour pressure increases by 0.02 Torr. Which of the following is true?
 - (A) The molecular weight of solute is 54g/mol
 - (B) The vapour pressure of pure water is 20.22 Torr.
 - (C) Addition of more water in the solution will further raise the vapour pressure of solution.
 - **(D)** The vapour pressure of pure water is 22.22 Torr.

SECTION - III : ASSERTION AND REASON TYPE

- 17. Statement 1 When a cell is placed in hypertonic solution, it shrinks.
 - Statement 2 Reverse osmosis is used for desalination of water.
 - (A) Statement-1 is True, Statement-2 is True; Statement-2 is a correct explanation for Statement-1.
 - (B) Statement-1 is True, Statement-2 is True; Statement-2 is NOT a correct explanation for Statement-1.
 - (C) Statement-1 is True, Statement-2 is False.
 - (D) Statement-1 is False, Statement-2 is True.
- 18. Statement 1 The difference in the boiling points of equimolar solution of HCl and HF decreases as their molarity is decreased.
 - Statement 2 The extent of dissociation decreases steadily with increasing dilution.
 - (A) Statement-1 is True, Statement-2 is True; Statement-2 is a correct explanation for Statement-1.
 - (B) Statement-1 is True, Statement-2 is True; Statement-2 is NOT a correct explanation for Statement-1.
 - (C) Statement-1 is True, Statement-2 is False.
 - **(D)** Statement-1 is False, Statement-2 is True.
- **19. Statement 1** The molar mass obtained for benzoic acid in benzene is found to be nearly 244.

Statement - 2 Benzoic acid has the formula HOOC

- (A) Statement-1 is True, Statement-2 is True; Statement-2 is a correct explanation for Statement-1.
- (B) Statement-1 is True, Statement-2 is True; Statement-2 is NOT a correct explanation for Statement-1.

COOH

- (C) Statement-1 is True, Statement-2 is False.
- (D) Statement-1 is False, Statement-2 is True.
- 20. Statement -1 When 'a' mL of a 0.1 molal urea solution is mixed with another 'b' mL of 0.1 molal glucose solution, the boiling point of the solution is no different from the boiling points of the samples prior to mixing but if 'a' mL of 0.1 molal urea is mixed with 'b' mL of 0.1 molal HF the boiling point of the mixture is different from the boiling point of the separate samples.
 - Statement 2 HF is an electrolyte (weak) whereas glucose is a non electrolyte.
 - (A) Statement-1 is True, Statement-2 is True; Statement-2 is a correct explanation for Statement-1.
 - (B) Statement-1 is True, Statement-2 is True; Statement-2 is NOT a correct explanation for Statement-1.
 - (C) Statement-1 is True, Statement-2 is False.
 - (D) Statement-1 is False, Statement-2 is True.



SECTION - IV : COMPREHENSION TYPE

Read the following comprehensions carefully and answer the questions.

Ideal Solution at Fixed Temperature

Consider two liquid 'B' and 'C' that form an ideal solution. We hold the temperature fixed at some value T that is above freezing points of 'B' and 'C'. We shall plot the system's pressure P against xB, the overall mole fraction of B in the system :

$$\mathbf{xB} = \frac{\mathbf{n}_{\mathrm{B,total}}}{\mathbf{n}_{\mathrm{total}}} = \frac{\mathbf{n}_{\mathrm{B}}^{\mathrm{l}} + \mathbf{n}_{\mathrm{B}}^{\mathrm{v}}}{\mathbf{n}_{\mathrm{B}}^{\mathrm{v}} + \mathbf{n}_{\mathrm{C}}^{\mathrm{l}} + \mathbf{n}_{\mathrm{B}}^{\mathrm{l}}}$$

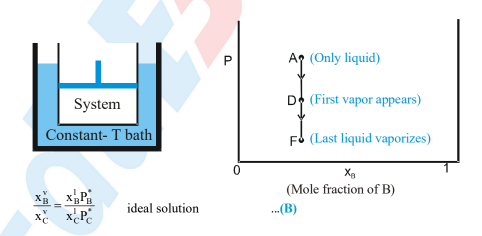
Where n_B^l and n_B^v are the number of moles of B in the liquid and vapor phases, respectively. For a closed system xB is fixed, althoud n_B^l and n_B^v may vary.

Let the system be enclosed in a cylinder fitted with a piston and immersed in a constant-temperature bath. To see what the P-versus-xB phase diagram looks like, let us initially set the external pressure on the piston high enough for the system to be entirely liquid (point A in figure) As the pressure is lowered below that at A, the system eventually reaches a pressure where the liquid just begins to vaporizes (point D). At point D, the liquid has composition x_B^1 where x_B^1 at D is equal to the overall mole fraction xB since only an infinitesimal amount of liquid has vapourized. What is the composition of the first vapour that comes off? Raoult's law, $P_B \equiv x_B^v P_B^0$ relates the vapour-phase mole fractions to the liquid composition as follows :

$$x_{B}^{0} = \frac{X_{B}^{1}P_{B}^{0}}{P}$$
 and $x_{C}^{v} = \frac{x_{C}^{1}P_{C}^{0}}{P}$...(A)

Where P_B^0 and P_C^0 are the vapour pressures of pure 'B' and pure 'C' at T, where the system's pressure P equals the

sum PB + PC of the partial pressures, where $P_B^l = \frac{n_B^l}{(n_B^l + n_C^l)}$, and the vapor is assumed ideal.

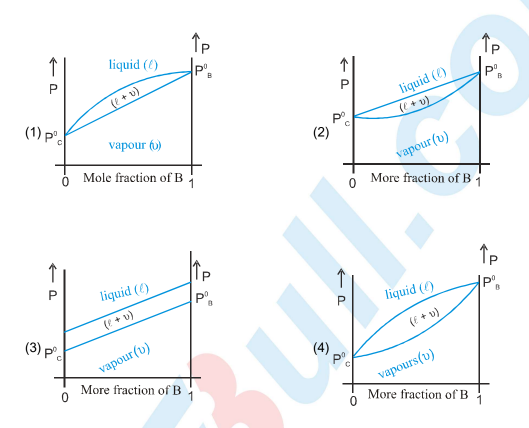


Let B be the more volatile component, meaning that $P_B^0 > P_C^0$. Above equation then shows that $x_B^v / x_C^v > x_B^l / x_C^l$. The vapor above an ideal solution is richer than the liquid in the more volatile component. Equations (A) and (B) apply at any pressure where liquid–vapor equilibrium exists, not just at point D.



Now let us isothermally lower the pressure below point D, causing more liquid to vaporize. Eventually, we reach point F in the figure, where the last drop of liquid vaporizes. Below F, we have only vapor. For any point on the line between D and F liquid and vapor phases coexist in equilibrium.

21. If the above process is repeated for all other compositions of mixture of C and B. If all the points where vapours start converting into liquid are connected and all the points where vapours get completely converted into liquid are connected obtained graph will look like.



22. The equation of the curve obtained by connecting all those points where the vapors of above mixture (all mixtures of different composition are taken) just start forming will be

(A) $P = P^{\circ}C + (P^{\circ}B - P^{\circ}C) X \bullet B$	(B) $P = P^{\circ}B + (P^{\circ}B - P^{\circ}C) X \bullet B$
(C) $P = \frac{P_B^0 P_C^0}{X_B^0 (P_C^0 - P_B^0) + P_B^0}$	(D) $P = \frac{P_B^0 P_C^0}{X_B^1 (P_C^0 - P_B^0) + P_B^0}$

23. Two liquids A and B have the same molecular weight and form an ideal solution. The solution has a vapor pressure of 700 Torrs at 80°C. It is distilled till 2/3rd of the solution (2/3rd moles out of total moles) is collected as condensate. The composition of the condensate is x'A = 0.75 and that of the residue is x''A = 0.30. If the vapor pressure of the residue at 80°C is 600 Torrs, find the original compositions of the liquid.

(A)
$$XA = 0.5$$
 (B) $XB = 0.6$ (C) $XA = 0.6$ (D) $XB = 0.3$



SECTION - V : MATRIX - MATCH TYPE

24.	Match the following :
-----	-----------------------

Column II

(A) Pyridine + CH_3COOH	(p) Δ Smixing = +Ve
(B) $C_6H_5Cl + C_6H6Br$	(q) $\Delta V mixing = +Ve$
(C) $\operatorname{CHCl}_3 + \operatorname{CCl}_4$	(r) $\Delta Gmixing = -Ve$
(D) $H_2O + HClO_4$	(s) Δ Hmixing = -Ve

25.	Match	the	foll	owing	:

Column I

Column II

(A) Relative lowering in vapour pressure (p) ne

(B) Depression in freezing point

(C) Δ Hmix < Zero

(D) Osmotic pressure

(p) negative deviation from ideal behaviour(q) Walker and Ostwald Method

- (r) Beckmann thermometer
- (s) Berkeley and Hartley's method

SECTION - VI : SUBJECTIVE TYPE

26. Barium ions, CN- and Co⁺² form a water soluble ionic complex with Ba²⁺ as free cations. For a 0.01 M solution of this complex osmotic pressure is observed to be equal to 0.984 atm and degree of dissociation is found out to be 75%. Then find the coordination number of Co(II) in this complex.

(T = 300K) (R = 0.082 L atm mol-1 K-1).

- 27. What is the osmotic pressure of a solution (in Torr) made by mixing 20mL of 0.01 M BaCl₂ with 30mL of 0.01 M NaF at 27° C? Ksp (BaF₂) = $2.4 \times 10-5$. R = 0.082 lit atm mol-1 K-1, 1 atm= 760Torr.
- 28. The composition of vapour over a binary ideal solution is determined by the composition of a liquid. If XA and YA are the mole fractions of A in the liquid and vapour, respectively, find the value of XA for which YA–XA has a maximum value. What is the value of pressure at this composition in terms of P°A and P°B.
- 29. When 2g of non volatile hydrocarbon containing 94.4 percent carbon is dissolved in 100g benzene, the vapour presure of benzene is lowered from 74.66 torr to 74.01 torr. Determine the molecular formulae for the hydrocarbon.
- 30. A saturated solution of sparingly soluble salt MCl_2 has a vapour pressure of 31.96 torr while pure water has a vapour pressure of 32.0 torr at the same temperature. Calculate the solubility of the product of MCl_2 .



ANSWER KEY

EXERCISE - 1

۲

1. A, B, D	2. B	3. A, B, C,	D 4. B 5.	В 6. С 7.	В 8. В 9.	B 10. C 11. A					
12. C 13. C	14. B 15.	A 16. A	17. A, B, C 18	. A 19. A 20	. A, B, C, D 2 1	I. C 22. A 23. B					
24. C 25. A	, C 26. D	27. B,D	28. C 29. B	30. B 31. B	32. A 33. B	34. A 35. C					
36. D 37. A	38. A 39.	C 40. C	41. A 42. C	43. B 44. C	45. D 46. C	47. D 48. D					
49. D 50. B	51. B 52.	В 53. С	54. B 55. C	56. A 57. B	58. A 59. B	,D 60. C					
61. A 62. B	63. C 64.	A 65. B	66. C 67. A	68. B 69. A	70. A 71. A	72. D 73. C					
74. C 75. C	76. A 77.	A 78. C	79. B 80. B	81. C 82. D	83. D 84. C	85. C 86. D					
87. C 88. B	89. B 90 .	D 91. D	92. C 93. B	94. A 95. D	96. C 97. B	98. C 99. A					
100. D 101. A	102.B 103	3.B 104.D	105. A 106. B	107. C 108. C	109. A 110. B	111. C 112. C					
113. A 114. A	115. A 116	.B 117.A	118.C 119.A	120. B 121. C	122. C 123. A	124. B 125. C					
126. A 127. C	128.B 129										
EXERCISE - 2 : PART # I											
1. A, B, C	2. A, B, C	3. A, E			B, C, D 6.	A, B, C, D					
7. B,C,D	8. A	9. A, E				2. A, B					
13. B	14. B,C	15. A, E	B, D 16. A,	B,C 17. A,		B. A, B, D					
19. A, D 20. A, B, C											
PART # II											
1. C 2. C	3. A 4.	В 5. А	6. D 7. A	8. D 9. B	10. D						
			EXERCISE -	3 : PART # I							
1. $A \rightarrow q, s, B$	\rightarrow q, s, C \rightarrow q	$s, D \rightarrow q, s$		s, r, $B \rightarrow p$, q, t, C	\rightarrow p, D \rightarrow p, q, t						
3. $A \rightarrow p; B \rightarrow$					1. 1. 1.						
			PART	C # 11							
Comprehension	#1: 1.	в 2.	A								
Comprehension	#2: 1.	в 2.	D								
Comprehension	#3: 1.	A, B 2.	C 3. D								
Comprehension	#4: 1.	D 2.	D 3. B 4.	A, B, C 5. A,	B, C, D 6. A	7. A					
EXERCISE - 5 : PART # I											
1 D 7 A	3 C /	C E A	6 C 7 P	9 C 0 P	10 C 11 C	17 A 12 D					
1. B 2. A	3. C 4. 16. B 17.		о. С 7. В 19. С 20. В	8. C 9. B	10. C II. C	12. A 13. B					
14. B 15. A	10. D 17.	D 10. D		с <i>щ</i> тт							
1 0 0 100			PART		0 4 10 4	11 0 1 0 D D					
		A 4. A	5. D 6. B	7. B 8. A	9. A 10. A	11. 9 12. B,D					
13. B 14. 19 15. 0.05 MOCK - TEST											
			NUUN								
1 C 2 D	3 R 1	4 5 R		8 B 0 D	10 D 11 P	17 A 13 R					
			6. C 7. A			12. A 13. B					
	B,D 16.	A,B,C 17.	6. C 7. A B 18.C 19		.B 22.A 23						

