**Exercise-1** 

# **PART - I : SUBJECTIVE QUESTIONS**

## Section (A) : General Introduction & types of solution

A-1. What are the characteristics of the supersaturated solution.

A-2. What do you mean by Hygroscopic compound ?

A-3. Why are some solution processes exothermic whereas others are endothermic ?

# Section (B) : Concentration terms (Revision of mole)

Commit to mem	nory :	
Molarity =	no. of moles of solute volume of solution (in L);	Molality = $\frac{\text{no. of moles of solute}}{W(g) \text{ of solvent}} \times 1000$
	volume of solution (in L) '	W(g) of solvent
Molality –	molarity ×1000	(where d is density of solution in a/ml)
100 anty = 1	000d - molarity × m.wt. of solu	— (where d is density of solution in g/ml) ute
$\% \frac{w}{w} = \frac{m_{so}}{m_{sol}}$	$\frac{x_{\text{solute}}(g)}{x_{\text{solute}}(g)} \times 100$ ; $x_{\text{solute}} = x_{\text{solute}}$	$\frac{n}{n+N}$ (where, n is moles of solute, N is moles of solvent)
Xsolute + Xso	lvent = 1	

**B-1.** Calculate the concentration of NaOH solution in g/ml which has the same molarity as that of a solution of HCl of concentration 0.0365 g/ml.

B-2. ★ The density of 3M solution of sodium thiosulphate (Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>) is 1.58 g/ml. Calculate (i) amount of Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> in % w/w
(ii) mole fraction of Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>
(iii) molality of Na<sup>+</sup> and S<sub>2</sub>O<sub>3</sub><sup>2-</sup> ions.

**B-3.** Calculate the molality and molarity of a solution made by mixing equal volumes of 30% by weight of  $H_2SO_4$  (density = 1.20 g/mL) and 70% by weight of  $H_2SO_4$  (density = 1.60 g/mL).

# Section (C) : Vapour Pressure

#### Commit to memory :

- \* The partial pressure of vapours of X in equilibrium with X at a given temperature is called as its vapour pressure. (X is given pure solid or pure liquid)
- \* A gas is said to be saturated with vapours of a liquid if the partial pressure of liquid vapours is equal to its (saturated) vapour pressure.
- \* Partial pressure of vapours of X (pure solid or liquid) will remain equal to its vapour pressure till the vapours of X are in equilibrium with X.
- C-1.> The vapour pressure of water at 80°C is 355 torr. A 100 ml vessel contained water-saturated oxygen at 80° C, the total gas pressure being 760 torr. The contents of the vessel were pumped into a 50.0 ml, vessel at the same temperature. What were the partial pressures of oxygen and of water vapour and the total pressure in the final equilibrium state? Neglect the volume of any water which might condense.
- **C-2.** A vessel has nitrogen gas and water vapours in equilibrium with liquid water at a total pressure of 1 atm. The partial pressure of water vapours is 0.3 atm. The volume of this vessel is reduced to one third of the original volume, at the same temperature, then find total pressure of the system. (Neglect volume occupied by liquid water)

# Section (D) : Solutions of Solid and Gases in Liquids

## Commit to memory :

- Henry's law: The solubility of gas in a liquid at a given temperature is directly proportional to its partial pressure above liquid in which it is dissolved.
  - $P = K_{HX}$  (where x is mole fraction of unreacted, dissolved gas and P is its partial pressure above liquid.)
- The partial pressure of ethane over a solution containing  $6.56 \times 10^{-3}$  g of ethane is 1 bar. If the solution D-1. contains  $5.00 \times 10^{-2}$  g of ethane, then what shall be the partial pressure of the gas ?
- D-2.2 If N<sub>2</sub> gas is bubbled through water at 293 K, how many millimoles of N<sub>2</sub> gas would dissolve in 300 mole of water, if N<sub>2</sub> exerts a partial pressure of 1 bar. Given that Henry's law constant for N<sub>2</sub> at 293 K is 75.00 kbar.

# Section (E) : Immiscible Liquids

## Commit to memory :

- If A and B are volatile immiscible liquids, then above their mixture.
  - $P_T = P_A^o + P_B^o$  ( $P_T = Total pressure above mixture of A and B, <math>P_A^o = vapour pressure of A$ ,  $P^{0}_{B}$  = vapour pressure of B)

 $\frac{P_A^{\circ}}{P_D^{\circ}} = \frac{n_A}{n_B} \quad (n_A \text{ and } n_B \text{ are moles of } A \text{ and } B \text{ in distillate})$ 

 $\frac{w_{A}}{w_{B}} = \frac{P_{A}^{\circ}M_{A}}{P_{B}^{\circ}M_{B}}$  (w<sub>A</sub> and w<sub>B</sub> are masses of A and B in distillate, M<sub>A</sub> and M<sub>B</sub> are molar masses of A and B)

- E-1. A mixture of an organic liquid A and water distilled under one atmospheric pressure at 99.2°C. How many grams of steam will be condensed to obtain 1.0 g of liquid A in the distillate ? (Vapour pressure of water at 99.2°C is 739 mm Hg. Molecular weight of A = 123)
- Boiling point of a mixture of water and nitrobenzene is 99°C, the vapour pressure of water is 733 mm of E-2. Hg and the atmospheric pressure is 760 mm of Hg. The molecular weight of nitrobenzene is 123. Find the ratio of weights of the components of the distillate.

# Section (F) : Completely miscible liquids : Raoult's law

## Commit to memory :

Statement of Raoult's law (for volatile liq. mixture) : In solution of volatile liquids, the partial vapour pressure of each component is directly proportional to its mole fraction in solution.

 $\mathbf{D}_{\mathrm{A}} \propto \mathbf{X}_{\mathrm{A}}$ 

..... (1)

 $p_A = x_A P_A^0$ where  $p_A$  = Partial vapour pressure of component A,  $x_A$  = Mole fraction of component 'A' in solution,  $P_{A^{o}}$  = Vapour pressure of pure component 'A' at given temperature

Hence  $P_T = x_A P_A^0 + x_B P_B^0$ ..... (2)

( $P_T$  = Total pressure of vapour above solution,  $P_{B^0}$  = vapour pressure of pure component 'B' at given temperature.)

Thus,  $\frac{1}{P_T} = \frac{y_A}{P_A^\circ} + \frac{y_B}{P_B^\circ}$  (for complete derivation, refer page no. 12 of sheet)

 $(y_A = mole fraction of A in vapour phase above the solution and y_B = mole fraction of B in vapour phase$ above the solution)

- F-1.3 Two liquids A and B form an ideal solution. At 300 K, the vapour pressure of a solution containing 1 mole of A and 3 moles of B is 550 mm of Hg. At the same temperature, if one mole of B is added to this solution, the vapour pressure of the solution increases by 10 mm of Hg. Determine the vapour pressure of A and B in their pure states.
- F-2. Two liquids, A and B, form an ideal solution. At the specified temperature, the vapour pressure of pure A is 200 mm Hg while that of pure B is 75 mm Hg. If the vapour over the mixture consists of 50 mol percent A, what is the mole percent A in the liquid ?
- F-3. Two solutions of A and B are available. The first is known to contain 1 mole of A and 3 moles of B and its total vapour pressure is 1.0 atm. The second is known to contain 2 moles of A and 2 moles of B; its vapour pressure is greater than 1 atm, but it is found that this total vapour pressure may be reduced to

1 atm by the addition of 6 moles of C. The vapour pressure of pure C is 0.80 atm. Assuming ideal solutions and that all these data refer to 25°C, calculate the vapour pressure of pure A and of pure B.

- F-4. At 80°C, the vapour pressure of pure benzene is 753 mm Hg and of pure toluene 290 mm Hg. Calculate the composition of a liquid in mole per cent which at 80°C is in equilibrium with the vapour containing 30 mole per cent of benzene.
- **F-5.** Vapour pressure of C<sub>6</sub>H<sub>6</sub> and C<sub>7</sub>H<sub>8</sub> mixture at 50°C is given P (mm Hg) =  $180X_B + 90$ , where X<sub>B</sub> is the mole fraction of C<sub>6</sub>H<sub>6</sub>. A solution is prepared by mixing 12 mol benzene and 8 mol toluene and if vapours over this solution are removed and condensed into liquid and again brought to the temperature 50°C, what would be mole fraction of C<sub>6</sub>H<sub>6</sub> in the vapour state. (At. wt. of C = 12, H = 1)

## Section (G) : Non-ideal Solutions

+ve deviation	-ve deviation
$P_{T.exp} > (x_A P^o_A + x_B P^o_B)$	$P_{T.exp} < (X_A P_A^0 + X_B P_B^0)$
(where PT.exp is experimental tot	al pressure above mixture of volatile liquids A and B).
$\Delta H_{mix} = +ve$	$\Delta H_{mix} = -ve$
$\Delta V_{mix} = +ve$	$\Delta V_{mix} = -ve$
$\Delta S_{mix} = +ve$	$\Delta S_{mix} = +ve$
$\Delta G_{mix} = -ve$	$\Delta G_{mix} = -ve$

**G-1.** A non ideal solution was prepared by mixing 30 ml chloroform and 50 ml acetone. Comment on volume of mixture.

G-2. ➤ Total vapour pressure of mixture of 1 mole of volatile component A (P<sup>o</sup><sub>A</sub> = 100 mm Hg) and 3 mole of volatile component B (P<sup>o</sup><sub>B</sub> = 80 mm Hg) is 90 mm Hg. Find out nature of solution and sign of entropy of solution.

#### Section (H) : Degree of Ionisation/Dissociation for Weak Electrolytes

#### Commit to memory :

 For dissociation/association i = 1 + (n - 1)α (where n is total number of particles produced per solute particle after association/dissociation, i = vant Hoff factor, α = degree of dissociation / association.
 Moles of solute particles after association/dissociation of X
 Observed value of colligative property

- $\frac{X}{T} = \frac{\text{Observed value of colligative property}}{\text{Theoretical value of that colligative property}}$
- Moles of X without association / dissociation
- $i = \frac{\text{Theoretical molar mass of solute}}{2}$ 
  - Observed molar mass of solute

#### H-1.2 Complete the following table.

Solute	Dissociation / association reaction	Degree of dissociation / association	n	i
KCI		1		
H <sub>2</sub> SO <sub>4</sub>		1		
CH <sub>3</sub> COOH (in water)		0.2		
CH <sub>3</sub> COOH (in benzene)		0.5		
Urea				
NaBr		0.8		
A	$3A \rightarrow A_3$	1		

**H-2.** Calculate the percentage degree of dissociation of an electrolyte XY<sub>2</sub> (Normal molar mass = 164) in water if the observed molar mass by measuring elevation in boiling point is 65.6.

#### Section (I) : Relative lowering of vapour pressure

#### Commit to memory :

\*  $RLVP = \frac{P^{o} - P_{S}}{P^{o}} = x_{solute};$   $\frac{P^{o} - P_{S}}{P_{S}} = \frac{n}{N}$ 

(where  $P^{o}$  = vapour pressure of pure solvent,  $P_{s}$  = partial pressure of vapour above solution, n = dissolved moles of solute, N = moles of solvent.)

 $\frac{P^{o} - P_{s}}{P_{s}} = \frac{m_{solute(g)}}{M_{solute}} \times \frac{M_{solvent}}{m_{solvent(g)}}$ (where m = given mass in solution, M = molar mass.)

$$\frac{P^{\circ} - P_{s}}{P_{s}} = (\text{ molality }) \times \frac{M_{solvent}}{1000}$$

If solution is of single solute and it gets associated or dissociated;

$$\frac{P^{\circ} - P_{s}}{P_{s}} = \frac{i.n}{N} \text{ ; that is } \frac{P^{\circ} - P_{s}}{P_{s}} = i \times \text{(molality)} \times \frac{M_{\text{solvent}}}{1000}$$

- I-1.> Twenty grams of a solute are added to 100 g of water at 25°C. The vapour pressure of pure water is 23.76 mmHg; the vapour pressure of the solution is 22.41 Torr.
  - (a) Calculate the molar mass of the solute.

(b) What mass of this solute is required in 100 g of water to reduce the vapour pressure to one-half the value for pure water ?

I-2. ➤ The degree of dissociation of Ca(NO<sub>3</sub>)<sub>2</sub> in a dilute aqueous solution containing 7 g salt per 100 g of water at 100°C is 70%. If the vapour pressure of water at 100°C is 760 mm of Hg, calculate the vapour pressure of the solution.

#### Section (J) : Elevation of Boiling Point & Depression of Freezing Point

#### Commit to memory :

 $\Delta T_b = i \times K_b \times m$  (where  $\Delta T_b =$  elevation in boiling point of solution,  $K_b =$  ebullioscopic constant, m = molality of single solute,  $T_b =$  boiling point of solvent (in K).

$$K_{b} = \frac{RT_{b}^{2}M}{1000 \times \Delta H_{vap}} = \frac{RT_{b}^{2}}{1000 \times L_{vap}}$$
 (M = Molar mass of solvent)

 $\Delta T_f = i \times K_f \times m$  (where  $\Delta T_f$  = depression in freezing point of solution,  $K_f$  = cryoscopic constant,  $T_f$  = freezing point of solvent (in K).

$$K_{f} = \frac{RT_{f}^{2} M}{1000 \times \Delta H_{fusion}} = \frac{RT_{f}^{2}}{1000 \times L_{fusion}}$$

J-1. (a) A solution containing 0.5 g of naphthalene in 50 g CCl<sub>4</sub> yield a boiling point elevation of 0.4 K, while a solution of 0.6 g of an unknown solute in the same mass of the solvent gives a boiling point elevation of 0.65 K. Find the molar mass of the unknown solute.

(b) The boiling point of a solution of 0.1 g of a substance in 16 g of ether was found to be  $0.100^{\circ}$ C higher that of pure ether. What is the molecular mass of the substance.  $K_b$ (ether) = 2.16 K kg mol<sup>-1</sup>.

- J-2.∞ The amount of benzene that will separate out (in grams) if a solution containing 7.32 g of triphenylmethane in 1000 g of benzene is cooled to a temperature which is 0.2°C below the freezing point of benzene ? (K<sub>f</sub> = 5.12 K-Kg/mol)
- J-3. The boiling point of a solution of 5 g of sulphur in 100 g of carbon disulphide is 0.474°C above that of pure solvent. Determine the molecular formula of sulphur in this solvent. The boiling point of pure carbon disulphide is 47°C and its heat of vaporisation is 84 calories per gram.

[Hint : 
$$K_b = \frac{RT_b^2}{1000L_v} = \frac{2 \times (320)^2}{1000 \times 84} = 2.438$$
]

J-4. A 0.01 molal solution of ammonia freezes at  $-0.02^{\circ}$ C. Calculate the van't Hoff factor, i and the percentage dissociation of ammonia in water. ( $K_{f(H,O)}$ ) = 1.86 deg molal<sup>-1</sup>.

## Section (K) : Osmotic Pressure

## Commit to memory :

- \*  $\pi = CRT = \frac{n}{V} RT$  (n = Total moles of solute particles in solution, V = Total volume of solution in L)
- K-1. (a) Predict the osmotic pressure order for the following (assume salts are 100% dissociated).
  - I 0.1 M urea
- II 0.1 M NaCl IV 0.1 M Na<sub>3</sub>PO<sub>4</sub>

 $\begin{array}{cccc} III & 0.1 \ M \ Na_2SO_4 & IV & 0.1 \ M \ Na_3PO_4 \\ \textbf{(b)} \ If \ equal \ volumes \ of \ all \ these \ solutions \ are \ mixed \ then \ calculate \ the \ osmotic \ pressure \ of \ the \ net \ resultant \ solution \ obtained \ at \ 300K. \end{array}$ 

- **K-2.** A solution containing 3.00 g of calcium nitrate in 100 c.c. of solution had an osmotic pressure of 11.2 atmosphere at 12°C. Calculate the degree of ionisation of calcium nitrate at this dilution and temperature.
- **K-3.** 17.4% (w/v) K<sub>2</sub>SO<sub>4</sub> solution at 27°C is isotonic with 5.85% (w/v) NaCl solution at 27°C. If NaCl is 100% ionised, what is % ionisation of K<sub>2</sub>SO<sub>4</sub> in aq. solution ?
- K-4.★ At 2°C the osmotic pressure of a urea solution is found to be 500 mm of Hg. The solution is diluted and the temperature is raised to 27°C, when the osmotic pressure is found to be 109.09 mm of Hg. Determine the extent of dilution.

# PART - II : ONLY ONE OPTION CORRECT TYPE

## Section (A) : General Introduction & types of solution

- A-1. Which statement best explains the meaning of the phrase "like dissolves like "?
  - (A) A Solute will easily dissolve a solute of similar mass
  - (B) A solvent and solute with similar intermolecular forces will readily form a solution
  - (C) The only true solutions are formed when water dissolves a non-polar solute
  - (D) The only true solutions are formed when water dissolves a polar solute
- A-2. An ionic compound that attracts atmospheric water so strongly that a hydrate is formed is said to be : (A) Dilute
  (B) Hygroscopic
  (C) Immiscible
  (D) Miscible

## Section (B) : Concentration terms (Revision of mole)

B-1. Persons are medically considered to have lead poisoning if they have a concentration greater than 10 micrograms of lead per decilitre of blood. Concentration in parts per billion is :

 (A) 1000
 (B) 100
 (C) 10
 (D) 1

## Section (C) : Vapour Pressure

- C-1. A liquid is kept in a closed vessel. If a glass plate (negligible mass) with a small hole is kept on top of the liquid surface, then the vapour pressure of the liquid in the vessel is :
  - (A) More than what would be if the glass plate were removed
  - (B) Same as what would be if the glass plate were removed
  - (C) Less than what would be if the glass plate were removed
  - (D) Cannot be predicted
- C-2.> The vapour pressure of water depends upon :
  - (A) Surface area of container (
    - (C) Temperature

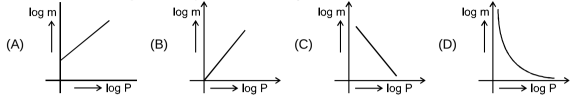
- (B) Volume of container (D) All
- C-3. Among the following substances, the lowest vapour pressure is exerted by : (A) Water (B) Mercury (C) Acetone (D) Ethanol

C-4. At higher altitudes, water boils at temperature < 100°C because

- (B) atmospheric pressure is low
- (A) temperature of higher altitudes is low(C) the proportion of heavy water increases
- (D) atmospheric pressure becomes more.

## Section (D) : Solutions of Solid and Gases in Liquids

- **D-1.** 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
- D-2. Which of the following curves represents the Henry'slaw?



D-3. 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

**D-4.** Some of the following gases are soluble in water due to formation of their ions : I:  $CO_2$ ; II:  $NH_3$ ; III: HCI; IV:  $CH_4$ ; V:  $H_2$ Water insoluble gases can be : (A) I, IV, V (B) I, V (C) I, II, III (D) IV, V

**D-5.** The solubility of N<sub>2</sub>(g) in water exposed to the atmosphere, when its partial pressure is 593 mm is  $5.3 \times 10^{-4}$  M. Its solubility at 760 mm and at the same temperature is : (A)  $4.1 \times 10^{-4}$  M (B)  $6.8 \times 10^{-4}$  M (C) 1500 M (D) 2400 M

## Section (E) : Immiscible Liquids

E-1. When a liquid that is immiscible with water was steam distilled at 95.2°C at a total pressure of 748 torr, the distillate contained 1.25 g of the liquid per gram of water. The vapour pressure of water is 648 torr at 95.2°C, what is the molar mass of liquid?
(A) 7.975 g/mol
(B) 166 g/mol
(C) 145.8 g/mol
(D) None of these

## Section (F) : Completely miscible liquids : Raoult's law

(B) 140 mm

- F-1. For a binary ideal liquid solution, the total pressure of the solution is given as : (A)  $P_{total} = P^{o}_{A} + (P^{o}_{A} - P^{o}_{B}) X_{B}$  (B)  $P_{total} = P^{o}_{B} + (P^{o}_{A} - P^{o}_{B}) X_{A}$ (C)  $P_{total} = P^{o}_{B} + (P^{o}_{B} - P^{o}_{A}) X_{A}$  (D)  $P_{total} = P^{o}_{B} + (P^{o}_{B} - P^{o}_{A}) X_{B}$
- F-2.28An ideal solution contains two volatile liquids A (p° = 100 torr) and B (p° = 200 torr). If mixture contain 1<br/>mole of A and 4 mole of B then total vapour pressure of the distillate is:<br/>(A) 150(B) 180(C) 188.88(D) 198.88
- **F-3.** 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

F-4.∞ Given at 350 K p<sub>A</sub>° = 300 torr and p<sub>B</sub>° = 800 torr, the composition of the mixture having a normal boiling point of 350 K is :

(C) 260 mm

(D) 20 mm

(A) 
$$X_A = 0.08$$
 (B)  $X_A = 0.06$  (C)  $X_A = 0.04$  (D)  $X_A = 0.02$ 

- **F-5.** Two liquids A and B have  $P^{o}_{A}$  and  $P^{o}_{B}$  in the ratio of 1 : 3 and the ratio of number of moles of A and B in liquid phase are 1 : 3 then mole fraction of 'A' in vapour phase in equilibrium with the solution is equal to:
  - (A) 0.1 (B) 0.2 (C) 0.5 (D) 1.0

## Section (G) : Non-ideal Solutions

G-1.æ	<ul> <li>The vapour pressure of the solution of two liquids A(p<sup>o</sup> = 80 mm) and B(p<sup>o</sup> = 120 mm) is found to be 100 mm when x<sub>A</sub> = 0.4. The result shows that</li> <li>(A) solution exhibits ideal behaviour</li> <li>(B) solution shows positive deviations</li> <li>(C) solution shows negative deviations for lower concentration and negative deviations for higher concentrations.</li> </ul>				
G-2.≿	Consider a binary mix then the mixture could (A) CHCl <sub>3</sub> – CH <sub>3</sub> COCH (C) C <sub>6</sub> H <sub>6</sub> – C <sub>6</sub> H <sub>5</sub> CH <sub>3</sub>	be $(p_A^0 = 300 \text{ torr}, p_B^\circ)$			,
G-3.	A solution of sulphuric (A) Negative deviations (C) Ideal properties			ons from Raoult's law y of Henry's law	
G-4.	When KCI dissolves in (A) $\Delta H = + ve$ , $\Delta S = + ve$ (C) $\Delta H = + ve$ , $\Delta S = + ve$	ve, $\Delta G = + ve$	thermic dissolution), the (B) $\Delta H = + ve, \Delta S =$ (D) $\Delta H = - ve, \Delta S =$	$= - ve, \Delta G = - ve$	
G-5.	<ul> <li>The dissolving process is exothermic when :</li> <li>(A) The energy released in solvation exceeds the energy used in breaking up solute-solute and solvent-solvent interactions.</li> <li>(B) The energy used in solvation exceeds the energy released in breaking up solute-solute and solvent-solvent interactions.</li> <li>(C) The energy released in solvation is about the same as the energy used in breaking up solute-solute and solvent-solvent-solvent interactions.</li> <li>(D) The energy used in solvation is about the same as the energy used in breaking up solute-solute and solvent-solvent interactions.</li> </ul>				)
G-6.১	Which of the following (A) $\Delta H_{mix}$	is less than zero for ide (B) $\Delta V_{mix}$	eal solutions ? (C) $\Delta G_{mix}$	(D) ∆S <sub>mix</sub>	
	on (H) : Degree of I			ectrolytes	

H-1. One mole of a solute A is dissolved in a given volume of solvent. The association of the solute take place as follows :  $nA \rightleftharpoons A_n$ 

If  $\alpha$  is the degree of association of A, the van't Hoff factor i is expressed as :

(A) 
$$i = 1 - \alpha$$
 (B)  $i = 1 + \frac{\alpha}{n}$  (C)  $i = \frac{1 - \alpha + \frac{\alpha}{n}}{1}$  (D)  $i = 1$ 

H-2. The degree of dissociation of an electrolyte is  $\alpha$  and its van't Hoff factor is i. The number of ions obtained by complete dissociation of 1 molecule of the electrolyte is :

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

H-3. If  $M_{normal}$  is the normal molecular mass and  $\alpha$  is the degree of ionization of K<sub>3</sub>[Fe(CN)<sub>6</sub>], then the abnormal molecular mass of the complex in the solution will be : (A)

$$M_{normal} (1 + 2\alpha)^{-1}$$
 (B)  $M_{normal} (1 + 3\alpha)^{-1}$  (C)  $M_{normal} (1 + \alpha)^{-1}$  (D) equal to  $M_{normal}$ 

H-4. A complex containing K<sup>+</sup>, Pt(IV) and Cl<sup>-</sup> is 100% ionised giving i = 3. Thus, complex is : (A) K<sub>2</sub>[PtCl<sub>4</sub>] (B) K<sub>2</sub>[PtCl<sub>6</sub>] (C) K<sub>3</sub>[PtCl<sub>5</sub>] (D) K[PtCl<sub>3</sub>]

I-1.a		vering of vapour pro		spectively and $N_1$ and $N_2$ are the
1-1.23		olvent and non-volatile s (B) $P = P_0 N_1$		
I-2.	Relative decrease in v in 180g of H <sub>2</sub> O is : (A) 2 mol	rapour pressure of an ac (B) 1 mol	ueous NaCl is 0.167. I (C) 3 mol	Number of moles of NaCl presen (D) 4 mol
I-3.æ	The vapour pressure c	of pure benzene, $C_6H_6$ at	50°C is 268 Torr. How	many moles of non-volatile solut ng a vapour pressure of 167 Tor
	(A) 0.377	(B) 0.605	(C) 0.623	(D) 0.395
I-4.	% ionised.		-	1 mol NaCl in 3 mol H <sub>2</sub> O, NaCl i
	(A) 60%	(B) 50%	(C) 100%	(D) 40%
I-5.æ	pressure of the solven		re. If the molecular wei	solvent A is 95% of the vapou ight of the solvent is 0.3 times the solute. (D) none of these
I-6.æ	moles H <sub>2</sub> O is 0.50. Of ionisation of substance	n reaction with AgNO <sub>3</sub> , e on adding AgNO <sub>3</sub> )	this solution will form(a	ning 2 moles [Cu(NH <sub>3</sub> ) <sub>3</sub> Cl]Cl in a assuming no change in degree c
	(A) 1 mol AgCl	(B) 0.25 mol AgCl	(C) 0.5 mol AgCl	(D) 0.40 mol AgCl
	• •	f Boiling Point & D	•	
	An aqueous solution of glucose in the same vo	containing 1g of urea bo	•	
J-1. 🖎	An aqueous solution of glucose in the same vo (A) 100.75 °C	containing 1g of urea bo blume will boil at - (B) 100.5 °C	ils at 100.25°C. The ad (C) 100°C	queous solution containing 3 g c (D) 100.25°C
	An aqueous solution of glucose in the same vo (A) 100.75 °C	containing 1g of urea bo blume will boil at - (B) 100.5 °C bint was 0.52 °C when	ils at 100.25°C. The ad (C) 100°C	queous solution containing 3 g c
J-1.≿ J-2.	An aqueous solution of glucose in the same vo (A) 100.75 °C Elevation in boiling po Molecular weight of x i (A) 120 A solute'S' undergoes elevation of its 0.1 mo 0.08 molal solution of	containing 1g of urea bo blume will boil at - (B) 100.5 °C bint was 0.52 °C when s : (K = 0.52 kg mol <sup>-1</sup> ) (B) 60 a reversible trimerization blal solution was found to a solute which neither u	ils at 100.25°C. The ad (C) 100°C 6 g of a compound x v (C) 100 n when dissolved in a o be identical to the bo	queous solution containing 3 g c (D) 100.25°C was dissolved in 100 g of water
J-1.≿ J-2.	An aqueous solution of glucose in the same vo (A) 100.75 °C Elevation in boiling po Molecular weight of x i (A) 120 A solute'S' undergoes elevation of its 0.1 mo	containing 1g of urea bo blume will boil at - (B) 100.5 °C bint was 0.52 °C when s : (K = 0.52 kg mol <sup>-1</sup> ) (B) 60 a reversible trimerization blal solution was found to a solute which neither u	ils at 100.25°C. The ad (C) 100°C 6 g of a compound x v (C) 100 n when dissolved in a o be identical to the bo	queous solution containing 3 g c (D) 100.25°C was dissolved in 100 g of water (D) 342 certain solvent. The boiling poin oiling point elevation in case of a
J-1.≿ J-2.	An aqueous solution of glucose in the same vo (A) 100.75 °C Elevation in boiling po Molecular weight of x i (A) 120 A solute'S' undergoes elevation of its 0.1 mo 0.08 molal solution of had the solute 'S' under (A) 30% A complex of iron and the complex is ( $K_b = 0$ )	containing 1g of urea bo plume will boil at - (B) 100.5 °C pint was 0.52 °C when s : (K = 0.52 kg mol <sup>-1</sup> ) (B) 60 a reversible trimerization blal solution was found to a solute which neither to ergone trimerization? (B) 40% cyanide ions is 100% io .52° mol <sup>-1</sup> kg) :	ils at 100.25°C. The ad (C) 100°C 6 g of a compound x v (C) 100 on when dissolved in a o be identical to the bo indergoes association in (C) 50% onised at 1m (molal). If	queous solution containing 3 g c (D) 100.25°C was dissolved in 100 g of water (D) 342 certain solvent. The boiling poin oiling point elevation in case of a nor dissociation. To what percen (D) 60% its elevation in b.p. is 2.08. The
J-1. & J-2. J-3. &	An aqueous solution of glucose in the same vo (A) 100.75 °C Elevation in boiling po Molecular weight of x i (A) 120 A solute'S' undergoes elevation of its 0.1 mo 0.08 molal solution of had the solute 'S' under (A) 30% A complex of iron and	containing 1g of urea bo blume will boil at - (B) 100.5 °C bint was 0.52 °C when s : (K = 0.52 kg mol <sup>-1</sup> ) (B) 60 a reversible trimerization blal solution was found t a solute which neither u ergone trimerization? (B) 40% cyanide ions is 100% io	ils at 100.25°C. The ac (C) 100°C 6 g of a compound x o (C) 100 on when dissolved in a o be identical to the bo indergoes association o (C) 50%	queous solution containing 3 g c (D) 100.25°C was dissolved in 100 g of water (D) 342 certain solvent. The boiling poin oiling point elevation in case of a nor dissociation. To what percer (D) 60%
J-1. & J-2. J-3. &	An aqueous solution of glucose in the same vol (A) 100.75 °C Elevation in boiling por Molecular weight of x i (A) 120 A solute'S' undergoes elevation of its 0.1 mo 0.08 molal solution of had the solute 'S' under (A) 30% A complex of iron and the complex is ( $K_b = 0$ . (A) $K_3$ [Fe(CN) <sub>6</sub> ] PtCl <sub>4</sub> .6H <sub>2</sub> O can exist a	containing 1g of urea bo plume will boil at - (B) 100.5 °C point was 0.52 °C when s : (K = 0.52 kg mol <sup>-1</sup> ) (B) 60 a reversible trimerization plal solution was found to a solute which neither to ergone trimerization? (B) 40% cyanide ions is 100% ion (B) 40% cyanide ions is 100% ion (B) Fe(CN) <sub>2</sub> as a hydrated complex onisation and K <sub>f</sub> (H <sub>2</sub> O) =	(C) 100°C (C) 100°C 6 g of a compound x v (C) 100 on when dissolved in a o be identical to the bo indergoes association in (C) 50% onised at 1m (molal). If (C) K₄[Fe(CN) <sub>6</sub> ] 1 molal aq. solution ha	queous solution containing 3 g o (D) 100.25°C was dissolved in 100 g of wate (D) 342 certain solvent. The boiling poir oiling point elevation in case of a nor dissociation. To what percer (D) 60% its elevation in b.p. is 2.08. The (D) Fe(CN)4 as depression in freezing point of mplex is - 2H <sub>2</sub> O
J-1.2 J-2. J-3.2	An aqueous solution of glucose in the same vo (A) 100.75 °C Elevation in boiling po Molecular weight of x i (A) 120 A solute'S' undergoes elevation of its 0.1 mo 0.08 molal solution of had the solute 'S' under (A) 30% A complex of iron and the complex is ( $K_b = 0$ . (A) K <sub>3</sub> [Fe(CN) <sub>6</sub> ] PtCl <sub>4</sub> .6H <sub>2</sub> O can exist a 3.72°. Assume 100% i (A) [Pt(H <sub>2</sub> O) <sub>6</sub> ]Cl <sub>4</sub> (C) [Pt(H <sub>2</sub> O) <sub>3</sub> Cl <sub>3</sub> ]Cl . 3 How many moles of so	containing 1g of urea bo containing 1g of urea bo plume will boil at - (B) 100.5 °C point was 0.52 °C when s : (K = 0.52 kg mol <sup>-1</sup> ) (B) 60 a reversible trimerization plal solution was found to a solute which neither to regone trimerization? (B) 40% cyanide ions is 100% ionomous (B) 40% cyanide ions is 100% ionomous (B) Fe(CN)2 as a hydrated complex onisation and K <sub>f</sub> (H <sub>2</sub> O) = H <sub>2</sub> O ucrose should be dissolved tween boiling point and formula to the top of t	<ul> <li>ils at 100.25°C. The ad</li> <li>(C) 100°C</li> <li>6 g of a compound x v</li> <li>(C) 100</li> <li>on when dissolved in a</li> <li>o be identical to the boundergoes association in</li> <li>(C) 50%</li> <li>onised at 1m (molal). If</li> <li>(C) K4[Fe(CN)6]</li> <li>1 molal aq. solution ha</li> <li>1.86° mol<sup>-1</sup> kg, then cor</li> <li>(B) [Pt(H<sub>2</sub>O)<sub>4</sub>Cl<sub>2</sub>]Cl<sub>2</sub>.</li> <li>(D) [Pt(H<sub>2</sub>O)<sub>2</sub>Cl<sub>4</sub>]. 4H</li> <li>red in 500 g of water so</li> </ul>	queous solution containing 3 g c (D) 100.25°C was dissolved in 100 g of water (D) 342 certain solvent. The boiling poin oiling point elevation in case of a nor dissociation. To what percen (D) 60% its elevation in b.p. is 2.08. Ther (D) Fe(CN)4 as depression in freezing point c mplex is - 2H <sub>2</sub> O

J-7.2		has been arranged in or 04 M CaCl <sub>2</sub> > 0.140 M su	5	ng point?
	(C) 0.075 M CuSO <sub>4</sub> >	140 M sucrose > 0.075 M 0.140 M sucrose > 0.04 0.05 M NaNO <sub>3</sub> > 0.140 N	M BaCl <sub>2</sub> > 0.05 M KNO3	3
J-8.		of sucrose in 100 grams ).25ºC. How many grams (B) 20 grams		<ul><li>P°C. As ice separates the freezing arated?</li><li>(D) 23 grams</li></ul>
J-9.2			<b>°</b>	nt of solution. Calculate its normal of MgCl <sub>2</sub> ( $K_b(H_2O) = 0.51 \text{ kgmol}^{-1}$
	(A) T <sub>b</sub> = 101.9°C	(B) T <sub>b</sub> = 102.3°C	(C) T <sub>b</sub> = 108.5°C	(D) T <sub>b</sub> = 110.3°C
Section	on (K) : Osmotic P	ressure		
K-1.	sucrose (P <sub>3</sub> ) are disso	lved in 250 mL of water i	s :	ose (P <sub>1</sub> ), 10 g urea (P <sub>2</sub> ) and 10 g
	(A) $P_1 > P_2 > P_3$	(B) $P_3 > P_1 > P_2$	(C) $P_2 > P_1 > P_3$	(D) $P_2 > P_3 > P_1$
K-2.	•	blood is 7.40 atm at 27°C hat is to have the same c (B) 0.2		plucose to be used per litre for an od is : (D) 0.4
K-3.	membrane surroundin	ig the cells is semiperme	eable. What would the o	is approximately 0.30 M and the osmotic pressure (in atmosphere) sma and placed in pure water at
	(A) 7.34 atm	(B) 1.78 atm	(C) 2.34 atm	(D) 0.74 atm
K-4.æ		tion in benzene at 27°C v tion at 27°C, which ionise		mation to the extent of 100%
	(A) both are isotonic	(B) I is hypertonic	(C) II is hypotonic	(D) none is correct
		PART - III : MAT		N
1.	It at a particular tempe	erature, the density of 18	M H <sub>2</sub> SO <sub>4</sub> is 1.8 g cm <sup><math>-3</math></sup> .	Then :

1. If at a particular temperature, the density of  $18 \text{ M H}_2\text{SO}_4$  is  $1.8 \text{ g cm}^{-3}$ . Then :

	Column – I		Column – II
(A)	Molality	(p)	0.1
(B)	% concentration by wt. of solute	(q)	0.9
(C)	mole fraction of H <sub>2</sub> SO <sub>4</sub>	(r)	500
(D)	mole fraction of H <sub>2</sub> O	(s)	98

#### 2. Match the following column-

	Column – I		Column – II
(A)	Acetone + CHCl <sub>3</sub>	(p)	$\Delta S_{mix.} > 0$
(B)	Ethanol + Water	(q)	$\Delta V_{mix.} > 0$
(C)	$C_2H_5Br + C_2H_5I$	(r)	ΔH <sub>mix.</sub> < 0
(D)	Acetone + Benzene	(s)	Maximum boiling azeotropes
		(t)	Minimum boiling azeotropes

# **Exercise-2**

# **PART - I : ONLY ONE OPTION CORRECT TYPE**

1. All of the water in a 0.20 M solution of NaCl was evaporated and a 0.150 mol of NaCl was obtained. What was the original volume of the sample ? (A) 30 mL (B) 333 mL (C) 750 mL (D) 1000 mL A 20.0 mL sample of CuSO<sub>4</sub> solution was evaporated to dryness, leaving 0.967 g of residue. What was 2.2 the molarity of the original solution ? (Cu = 63.5) (B) 0.0207 M (C) 0.0484 M (D) 0.303 M (A) 48.4 M The vapour pressure of water at 20°C is 17.54 mmHg. What will be the 3.2 vapour pressure of the water in the apparatus shown after the piston is lowered, decreasing the volume of the gas above the liquid to one half of its Water vapour initial volume (assume temperature constant). (A) 8.77 mmHg (B) 17.54 mmHa Liquid water (C) 35.08 mmHg (D) between 8.77 and 17.54 mmHg A sample of air is saturated with benzene (vapor pressure = 100 mm Hg at 298 K) at 298K, 750mm Hg 4.2 pressure. If it is isothermally compressed to one third of its initial volume, the final pressure of the system is (C) 2050 torr (A) 2250 torr (B) 2150 torr (D) 1950 torr 5. Water and chlorobenzene are immiscible liquids. Their mixture boils at 89°C under a reduced pressure of 7.7 x 10<sup>4</sup> Pa. The vapour pressure of pure water at 89°C is 7 x 10<sup>4</sup> Pa. Weight percent of chlorobenzene in the distillate is: (C) 78.3 (D) 38.46 (A) 50 (B) 60 If two liquids A ( $P_A^0 = 100$  torr) and B ( $P_B^0 = 200$  torr) are completely immiscible with each other, each 6. one will behave independently of the other, are present in a closed vessel. The total vapour pressure of the system will be: (B) less than 200 torr (C) between 100 to 200 torr (D) 300 torr (A) less than 100 torr Given P-x curve for a non-ideal liquid mixture (Fig.). Identify the correct T-x curve for the same mixture. 7.2 ମ୍ମି 800 mm) 600 Pressure 400 200 0 0204060810 Mole Fraction Point Point Point Point Boiling F (A) Boiling Boiling ' Boiling (D) (B) (C) 0 0.2 0.4 0.6 0.8 0 02040608 1.0 0 0204060810 0 02040608 1.0 1.0 Mole Fraction Mole Fraction Mole Fraction Mole Fraction

8. 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)  $\Delta H_{mix} = 0$ 

9. Barium ions, CN<sup>-</sup> and Co<sup>2+</sup> form an ionic complex. If that complex is supposed to be 75% ionised in water with vant Hoff factor 'i' equal to four, then the coordination number of Co<sup>2+</sup> in the complex can be: (A) Six
 (B) Five
 (C) Four
 (D) Six and Four both

- In the following aqueous solutions (a) 1 m sucrose, (b) 1 m potassium ferricyanide and (c) 1 m potassium sulphate Maximum value of vapour pressure of solution is that of :
   (A) a
   (B) b
   (C) c
   (D) equal
- 11.> When only a little quantity of HgCl₂(s) is added to excess Kl(aq) to obtain a clear solution, which of the following is true for this solution? (no volume change on mixing). The reaction is 4Kl(aq.) + HgCl₂(s) → K₂[Hgl₄] (aq.) + 2KCl (aq.)
   (A) Its boiling and freezing points remain same (B) Its boiling point is lowered
  - (C) Its vapour pressure become lower (D) Its boiling point is raised
  - (E) Its freezing point is lowered.
- **12.** x mole of KCI and y mole of BaCl<sub>2</sub> are both dissolved in 1 kg of water. Given that x + y = 0.1 and K<sub>f</sub> for water is 1.85 K/molal, what is the observed range of  $\Delta T_f$ , if the ratio of x to y is varied ? (A) 0.37° to 0.555° (B) 0.185° to 0.93° (C) 0.56° to 0.93° (D) 0.37° to 0.93°
- **13.** FeCl<sub>3</sub> on reaction with K4[Fe(CN)<sub>6</sub>] 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 side Y
  - (C) blue colour formation in both of the sides X and Y
  - (D) no blue colour formation
- **14.** Two beakers, one containing 20 ml of a 0.05 M aqueous solution of a non volatile, non electrolyte and the other, the same volume of 0.03 M aqueous solution of NaCl, are placed side by side in a closed enclosure. What are the volumes in the two beakers when equilibrium is attained ? Volume of the solution in the first and second beaker are respectively.

(A) 21.8 ml and 18.2 mL

(C) 20 mL and 20 mL

(B) 18.2 mL and 21.8 mL (D) 17.1 mL and 22.9 mL

# PART - II : SINGLE AND DOUBLE VALUE INTEGER TYPE

- **1.** What volume of 98% sulphuric acid (in ml) should be mixed with water to obtain 200 mL of 15% solution of sulphuric acid by weight ? Given density of  $H_2O = 1.00$  g cm<sup>-3</sup>, sulphuric acid (98%) = 1.88 g cm<sup>-3</sup> and sulphuric acid (15%) = 1.12 g cm<sup>-3</sup>.
- **2.** At 300 K, 40 mL of O<sub>3</sub> (g) dissolves in 100 g of water at 1.0 atm. What mass of ozone (in gram) dissolved in 1600 g of water at a pressure of 4.0 atm at 300 K?
- **3.** An ideal aqueous solution containing liquid A(M.Wt. = 128) 64% by weight has a vapour pressure of 145 mm Hg. If the vapour pressure of A is x mm of Hg and that of water is 155 mm Hg at the same temperature. Then find x/5. The solutions is ideal.
- **4.** A and B form ideal solutions; at 50°C, P<sub>A</sub>° is half P<sub>B</sub>°. A solution containing 0.2 mole of A and 0.8 mole of B has a normal bolling point of 50°C. Find 18 × P<sub>B</sub>°. (P<sub>B</sub>° is in atm)
- **5.** The vapour pressure of pure liquid A at 300 K is 577 Torr and that of pure liquid B is 390 Torr. These two compounds form ideal liquid and gaseous mixtures. Consider the equilibrium composition of a mixture in which the mole fraction of A in the vapour is 0.35. Find the mole % of A in liquid.
- 6. How many facts related to CHCl<sub>3</sub> + ethylmethylketone solution are correct ?
   (a) It shows negative derivation.
   (b) It forms maximum boiling azeotropic mixture
  - (b) (c) it forms maximum boining azeotropic (d)  $\Delta G < 0$
  - (c)  $\Delta S > 0$
  - (e) Components can be separated by fractional distillation.
- 7. How many of the following solutions show negative deviation from Raoult's Law ?

	Liquid A	+	Liquid B
(i)	(CH <sub>3</sub> ) <sub>2</sub> CO	+	$CS_2$
(ii)	CCl <sub>4</sub>	+	$C_6H_6$
(iii)	CCl <sub>4</sub>	+	CHCl₃
(iv)	H <sub>2</sub> O	+	C₂H₅OH

F	2			
0.1M	0.01M			
K₄Fe(CN) <sub>6</sub>	FeCl <sub>3</sub>			
F = = = = = =	= = = = = = =			
= ∶Side X =	i⊑ Side Y⊑ ⊒			
= = = = = = =	_= = = = = = =			

(v)	(C <sub>2</sub> H <sub>5</sub> ) <sub>2</sub> O	+	(CH <sub>3</sub> ) <sub>2</sub> CO
(vi)	CH <sub>3</sub> COOH	+	C₅H₅N (pyridine)
(vii)	$C_6H_5NH_2$	+	(CH <sub>3</sub> ) <sub>2</sub> CO
(viii)	C <sub>6</sub> H₅CI	+	C <sub>6</sub> H₅Br
(ix)	Cyclohexane	+	Ethanol

- 0.1 mole XY<sub>2</sub> is dissolved in 2L water, where it ionizes to give X<sup>2+</sup> and Y<sub>2</sub><sup>2-</sup>. Observed osmotic pressure 8.2 is 3 atm. Molar mass of X is 24 and Y is 32. Find Mobserved + 2i (where Mobserved is observed molar mass of XY<sub>2</sub>). (Use R = 1/12 L-atm/mol.K and temperature is 87°C)
- 9.2 How many grams of sucrose (C12H22O11) must be dissolved in 90 g of water to produce a solution over which the relative humidity is 80%? Assume the solution is ideal. Give your answer after dividing by 10.
- 10.ര. 1.22 g of a monobasic acid is dissolved in 100 g of benzene. Boiling point of solution increases by 0.13°C with respect to pure benzene. Find the molecular mass of acid in benzene solvent (in u). Report your answer after dividing it by 100 and Round it off to nearest integer. (K<sub>b</sub> of benzene = 2.6 K kg mol<sup>-1</sup>).
- 11.2 1 g of a monobasic acid dissolved in 200 g of water lowers the freezing point by 0.186°C. On the other hand when 1 g of the same acid is dissolved in water so as to make the solution 200 mL, this solution requires 125 mL of 0.1 M NaOH for complete neutralization. Calculate % dissociation of acid ? (K<sub>f</sub> K - kg

2.

- 12. At 27°C, a 1.2% solution (wt./vol.) of glucose is isotonic with 4.0 g/litre of solution of solute X. Find the molar mass of X, if the molar mass of glucose is 180. (R = 0.082 L atm mol<sup>-1</sup> K<sup>-1</sup>, Molar mass of qlucose = 180 q/mole)
- 10 g of solute A and 20 g of solute B both are dissolved in 500 ml. of water. The solution has the same 13. osmotic pressure as 6.67 g of A and 30 g of B are dissolved in the same volume of water at the same temperature. If the ratio of molar masses of A and B is x/v, find x + v.

# PART - III : ONE OR MORE THAN ONE OPTIONS CORRECT TYPE

- We have 100 mL of 0.1 M KCl solution. To make it 0.2 M, 1.2 (A) evaporate 50 mL water (B) evaporate 50 mL solution (C) add 0.1 mol KCl
  - (D) add 0.01 mol KCI Which of the following concentration factors can be calculated if the mole fraction and density of an
  - aqueous solution of HCl are known ? (A) Molality (B) Molarity
- (C) Percent by mass (D) Normality
- The vapour pressure of a dilute solution of a solute is influenced by : 3.2 (A) Temperature of solution (B) Mole fraction of solute (C) M.pt. of solute
  - (D) Degree of dissociation of solute
- 4.2 According to Henry's law, the partial pressure of gas  $(P'_g)$  is directly proportional to mole fraction of gas in dissolved state , i.e.,  $P_{gas}' = K_H$ .  $X_{gas}$  where  $K_H$  is Henry's constant. Which are correct ?
  - (A) K<sub>H</sub> is characteristic constant for a given gas-solvent system
  - (B) Higher is the value of  $K_H$ , lower is solubility of gas for a given partial pressure of gas
  - (C) K<sub>H</sub> has temperature dependence
  - (D) K<sub>H</sub> increases with temperature
- Select correct statements : 5.2
  - (A) Gases which have high value of van der Waals constant 'a' are easily liquefied
  - (B) Easily liquefiable gases are water soluble
  - (C) Gases which forms ions in a solvent are soluble in that solvent
  - (D) Under same conditions,  $NH_3$  has low solubility in water than that of  $CO_2$ .
- 6.2 Two liquids X and Y are perfectly immiscible. If X and Y have molecular masses in ratio 1 : 2, the total vapour pressure of a mixture of X and Y prepared in weight ratio 2 : 3 should be ( $P_x^0 = 400$  torr,  $P_v^0 = 200 \text{ torr}$ ) (D) 1000 torr
  - (B) 400 torr (A) 600 torr (C) 800 torr

- 7.2 Which is/are true about ideal solutions ? (A) The volume change on mixing is zero (C) The entropy of mixing is zero
- (B) The enthalpy of mixing is zero
- (D) The enthalpy of mixing is negative
- 8. 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
- 9. Which is/are correct statement(s) ?

(A) When mixture is more volatile than both pure components, there is positive deviation from Raoult's law.

(B) When mixture is less volatile than both pure components, there is negative deviation from Raoult's law.

(C) Ethanol and water form ideal solution

(D) CHCl<sub>3</sub> and water form ideal solution

At 35°C, the vapour pressure of CS<sub>2</sub> is 512 mm Hg, and of acetone is 344 mm Hg. A solution of CS<sub>2</sub> 10.2 and acetone in which the mole fraction of  $CS_2$  is 0.25, has a total vapour pressure of 600 mm Hq. Which of the following statements is/are correct?

(A) A mixture of 100 mL of acetone and 100 mL of CS<sub>2</sub> has a volume of 200 mL

(B) When acetone and CS<sub>2</sub> 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
- 11. For the given electrolyte  $A_x B_y$ , the degree of dissociation ' $\alpha$ ' can be given as

(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

- 12.2 In which of the following pairs of solutions will the values of the vant Hoff factor be the same? (A) 0.05 M K<sub>4</sub> [Fe(CN)<sub>6</sub>] and 0.10 M FeSO<sub>4</sub>
  - (B) 0.10 M K<sub>4</sub>[Fe(CN)<sub>6</sub>] and 0.05 M FeSO<sub>4</sub> (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. 6H<sub>2</sub>O
  - (C) 0.20 M NaCl and 0.10 M BaCl<sub>2</sub>
  - (D) 0.05 M FeSO<sub>4</sub> (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>.6H<sub>2</sub>O and 0.02 M KCI.MgCl<sub>2</sub>.6H<sub>2</sub>O
- 2 g of non-volatile hydrocarbon solute dissolved in 100 g of hypothetical organic solvent (molar mass = 13. 50) was found to lower vapour pressure from 75.50 to 75 mm of Hg at 20°C. Given that hydrocarbon contains 96% of C. Then which of the following are true ?
  - (A) molecular wt of solute = 150
- (B) molecular formula =  $C_{12}H_6$
- (C) molecular wt. of solute = 132
- (D) none of these
- 14.2 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
- 15.2 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

- **16.** Which facts are true when we use van't Hoff equation PV = nST 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

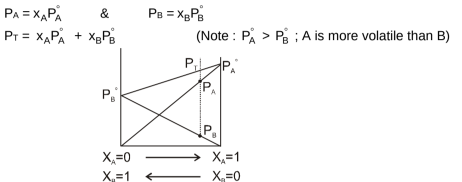
(C) Solute molecules are dispersed in the solvent the way the gas molecules are dispersed in empty space

(D) The equation is not identical to that of ideal gas equation

# **PART - IV : COMPREHENSIONS**

#### Read the following passage carefully and answer the questions.

#### Comprehension #1



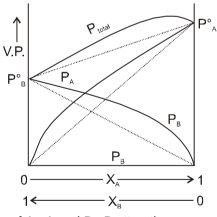
Vapour pressure of mixtures of Benzene (C<sub>6</sub>H<sub>6</sub>) & toluene (C<sub>7</sub>H<sub>8</sub>) at 50°C are given by  $P_M = 179 X_B + 92$ , where  $X_B$  is mole fraction of C<sub>6</sub>H<sub>6</sub>.

(D) none of these

- 1. What is vapour pressure of pure liquids ?
  - (A)  $P_B^{\circ} = 92 \text{ mm}, P_T^{\circ} = 179 \text{ mm}$  (B)  $P_B^{\circ} = 271 \text{ mm}, P_T^{\circ} = 92 \text{ mm}$
  - (C)  $P_B^{\circ} = 180 \text{ mm}, P_T^{\circ} = 91 \text{ mm}$
- 2.2Vapour pressure of liquid mixture obtained by mixing 936 g C6H6 & 736 g tolene is :(A) 300 mm Hg(B) 250 mm Hg(C) 199.4 mm Hg(D) 180.6 mm Hg

#### 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...A and B...B attraction: B : ΔH<sub>mix</sub> = +ve, ΔV<sub>mix</sub> = +ve C : Boiling point is smaller than expected such that vaporisation is increased D : Mixture can form azeotropic mixture Select correct facts (A) A, B, C (B) B, C, D (C) A, C, D (D) A, B, C, D

- 4.∞ Total vapour pressure of mixture of 1 mol of volatile component A (p°<sub>A</sub> = 100 mm Hg) and 3 mol of volatile component B (p°<sub>B</sub> = 60 mm Hg) 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

Addition of non-volatile solute to a solvent always increases the colligative properties such as osmotic pressure,  $\Delta P$ ,  $\Delta T_b$  and  $\Delta T_f$ . 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.

- **5.** For different aqueous solutions of 0.1 M NaCl, 0.1 M urea, 0.1 M Na<sub>2</sub>SO<sub>4</sub> and 0.1 M Na<sub>3</sub>PO<sub>4</sub> solution at 27°C, the correct statements are : (Consider the solutions to be dilute)
  - 1. The order of osmotic pressure is, NaCl > Na<sub>2</sub>SO<sub>4</sub> > Na<sub>3</sub>PO<sub>4</sub> > 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 or	n ice brings in melti	ng of ice earlier	

(A) 2, 3, 4 (B) 2, 4	(C) 1, 2, 3	(D) 3, 4
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- **6.** 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  $5.41 \times 10^{-4}$ .
  - 3. The solution will boil at 100.015°C, if K<sub>b</sub> of water is 0.5 K molality<sup>-1</sup>.
  - 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 # 4

Let us consider a binary solution of two volatile liquids 'A' and 'B', when taken in a closed container. Both the components would evaporate and an equilibrium would be established between vapour phase and liquid phase. Let the total vapour pressure at this stage be  $p_{total}$  and  $p_A$  and  $p_B$  are partial pressures of A and B. Mole fractions of these components in liquid solution are  $x_A$  and  $x_B$ , that of vapour phase are  $y_A$  and  $y_B$  respectively  $p_A^\circ \& p_B^\circ$  are vapour pressure of pure A & pure B.

Column-I	Column-II	Column-III
(i) Ideal solution	(a) $p_T = p_A^{\circ} x_A + p_B^{\circ} x_B$	(P) Hole fraction of A
(ii) Non ideal solution (Positive deviation)	A—A (b) interactions B—B interactions≠ A—B interactions	(Q) $P_T = P^{\circ}_A + P^{\circ}_B$
(iii) Non ideal solution (negative deviation)	(c) Form azeotropic mixture	(R) $\Delta S_{surrounding} = +ve$

Solu	tion & Colligative Prope	erties				
	(iv) Immiscible liquids	(d) solution having vap	oours of	() Vapour Pressure of A	Mole fraction of A	P <sub>A</sub> °
7.	For Hexane + Heptane (A) (i) (a) (R)	$e \rightarrow$ solution correct set (B) (ii) (b) (P)	is : (C) (ii) (d)	(R)	(D) (i) (a) (P)	
8.	For water + ethanol —; (A) (ii) (b) (Q)	Solution correct set is : (B) (ii) (c) (S)	(C) (iv) (d)	(S)	(D) (iv) (c) (R)	
9.	For water + $H_2SO_4 \rightarrow$ (A) (i) (a) (P)	solution correct set is (B) (ii) (b) (Q)	(C) (iii) (b)	(R)	(D) (iv) (c) (P)	
10.	For a mixture of water (A) (i) (a) (P)	and chlorobenzene corr (B) (ii) (b) (Q)	rect set is (C) (iii) (b)	(R)	(D) (iv) (d) (Q)	
	Exercise	-3				
* Ma	rked Questions may hav		ect option.			
<u> </u>	PART - I : JEE (AD	VANCED) / IIT-J	EE PROI	BLEMS	(PREVIOUS YE	EARS)
 1.	In the depression of fro I. The vapour pressure II. The vapour pressur III. Only solute molecu	eezing point experiment of the solution is less the of the solution is more les solidify at the freezing cules solidify at the freezing	, it is found th nan that of pu than that of p ng point.	at : ire solvent.	[JEE 1999	
	In the depression of from I. The vapour pressure II. The vapour pressure III. Only solute molecu IV. Only solvent molecu (A) I, II	eezing point experiment e of the solution is less the e of the solution is more les solidify at the freezir	, it is found th nan that of pu than that of p ng point. ing point. (C) I, IV	at : ire solvent. pure solvent	[JEE 1999 t. (D) I, II, III. f dissociation is :	9, 3/80]
1.	In the depression of from I. The vapour pressure II. The vapour pressure III. Only solute molecu IV. Only solvent molecu (A) I, II	eezing point experiment e of the solution is less th e of the solution is more les solidify at the freezir cules solidify at the freez (B) II, III	, it is found th nan that of pu than that of p ng point. ing point. (C) I, IV	at : ire solvent. pure solvent	[JEE 1999 t. (D) I, II, III.	9, 3/80]
1.	In the depression of fr I. The vapour pressure II. The vapour pressure III. Only solute molecu IV. Only solvent molecu (A) I, II The van't Hoff factor fo (A) 91.3% To 500 cm <sup>3</sup> of water, 3 be the depression in respectively	eezing point experiment e of the solution is less the e of the solution is more les solidify at the freezin cules solidify at the freez (B) II, III or 0.1 M Ba(NO <sub>3</sub> ) <sub>2</sub> soluti (B) 87% 3.0 × 10 <sup>-3</sup> kg of acetic a freezing point? k <sub>f</sub> and o	, it is found th nan that of pu than that of p ing point. (C) I, IV on is 2.74. Th (C) 100% cid is added. density of wa	at : pure solvent. pure solvent ne degree of If 23% of ac iter are 1.86	[JEE 1999 t. i dissociation is : [JEE 1999 (D) 74% cetic acid is dissociate 5 K kgmole <sup>-1</sup> and 0 [JEE 2000	<b>9, 3/80]</b> <b>9, 3/80]</b> ed, what will .997 g cm <sup>-3</sup>
1. 2.	In the depression of fr I. The vapour pressure II. The vapour pressure III. Only solute molecu IV. Only solvent molecu (A) I, II The van't Hoff factor fo (A) 91.3% To 500 cm <sup>3</sup> of water, 3 be the depression in respectively (A) 0.186 K	eezing point experiment e of the solution is less th e of the solution is more les solidify at the freezin cules solidify at the freez (B) II, III or 0.1 M Ba(NO <sub>3</sub> ) <sub>2</sub> soluti (B) 87% 3.0 × 10 <sup>-3</sup> kg of acetic a freezing point? k <sub>f</sub> and o (B) 0.228 K reezing point in a solution d solvent	, it is found th nan that of pu than that of p ing point. (C) I, IV on is 2.74. Th (C) 100% cid is added. density of wa (C) 0.372 on, the followi (B) Liquid	nat : pure solvent. pure solvent ne degree of lf 23% of ac tter are 1.86	[JEE 1999 (D) I, II, III. dissociation is : [JEE 1999 (D) 74% cetic acid is dissociate (D) 74% cetic acid is dissociate (D) 74% (D) 0.556 K (D) 0.556 K guilibrium : [JEE 2003 d solute	<b>9, 3/80]</b> <b>9, 3/80]</b> ed, what will .997 g cm <sup>-3</sup> <b>), 3/35]</b>
1. 2. 3.	In the depression of fr I. The vapour pressure II. The vapour pressure III. Only solute molecu IV. Only solvent molecu (A) I, II The van't Hoff factor fo (A) 91.3% To 500 cm <sup>3</sup> of water, 3 be the depression in respectively (A) 0.186 K During depression of f (A) Liquid solvent-solid (B) Liquid solute-solid A 0.004 M solution of	eezing point experiment e of the solution is less th e of the solution is more les solidify at the freezin cules solidify at the freez (B) II, III or 0.1 M Ba(NO <sub>3</sub> ) <sub>2</sub> soluti (B) 87% 3.0 × 10 <sup>-3</sup> kg of acetic a freezing point? k <sub>f</sub> and o (B) 0.228 K reezing point in a solution d solvent	, it is found th nan that of pu- than that of p ing point. (C) I, IV on is 2.74. Th (C) 100% cid is added. density of wa (C) 0.372 on, the followi (B) Liquid (D) Liquid	at : pure solvent. pure solvent ne degree of If 23% of ac ter are 1.86 K ng are in ec solvent-solid	[JEE 1999 (D) I, II, III. dissociation is : [JEE 1999 (D) 74% cetic acid is dissociate (D) 74% cetic acid is dissociate (D) 74% (D) 74	<b>9, 3/80]</b> <b>9, 3/80]</b> ed, what will .997 g cm <sup>-3</sup> <b>0, 3/35]</b> <b>3, 3/84]</b> erature. The

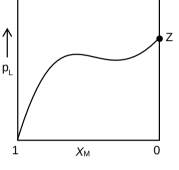
	ion & Colligative Flope			
6.	of the solution in aceto	ne increases by 0.17⁰C, is 1.7 K kg mol⁻¹ and 2.6	while that of, in the benz	enzene separately. Boiling point ene increases by 0.13°C ; K <sub>b</sub> for ilar weight of benzoic acid in two [JEE 2004, 4/60]
7.			5	kg of water using the following molal <sup>-1</sup> ) : <b>[JEE 2005, 3/84]</b> (D) 0.2
8.			issolved in 50 g of ben he van't Hoff factor (i) is (C) 2	zene (K <sub>f</sub> = 1.72 K kg mol <sup>-1</sup> ), a : <b>[JEE-2007, 3/162]</b> (D) 3
Comp	solute molecules are Application of colligative ethylene glycol and wa A solution <b>M</b> is prepare Given : Freezing point Boiling point el Boiling point el Standard freez Standard freez Standard boilin Vapour pressu Vapour pressu Molecular weig	added to get homoger /e properties are very u ter mixture as anti-freez ed by mixing ethanol and depression constant of depression constant of evation constant of water evation constant of ethan ing point of water = 273 ing point of ethanol = 15 ing point of ethanol = 351 ire of pure water = 32.8 r ire of pure ethanol = 40 r ght of ethanol = 46 g mol	neous solution. These a seful in day-to-day life. ( ing liquid in the radiator of water. The mole fraction water ( $K_f^{water}$ ) = 1.86 K kg ethanol ( $K_f^{ethanol}$ ) = 2.0 K er ( $K_b^{water}$ ) = 0.52 K kg mo nol ( $K_b^{ethanol}$ ) = 1.2 K kg m K 5.7 K 5.7 K 5.5 K mm Hg mm Hg	n of ethanol in the mixture is 0.9 g mol <sup>_1</sup> kg mol <sup>_1</sup> ol <sup>_1</sup>
	In answering the follow non-volatile and non-di		the solution to be ideal of	lilute solutions and solutes to be
9.		ssociative.	the solution to be ideal of (C) 234.2 K	lilute solutions and solutes to be [JEE 2008, 3/163] (D) 150.9 K
9. 10.	non-volatile and non-di The freezing point of th	issociative. ne solution <b>M</b> is (B) 268.5 K		[JEE 2008, 3/163]
	non-volatile and non-di The freezing point of th (A) 268.7 K The vapour pressure o (A) 39.3 mm Hg	issociative. ne solution <b>M</b> is (B) 268.5 K f the solution <b>M</b> is (B) 36.0 mm Hg solution <b>M</b> such that the	(C) 234.2 K (C) 29.5 mm Hg	[JEE 2008, 3/163] (D) 150.9 K [JEE 2008, 3/163]
10.	non-volatile and non-di The freezing point of th (A) 268.7 K The vapour pressure o (A) 39.3 mm Hg Water is added to the boiling point of this solu (A) 380.4 K The Henry's law const	issociative. ne solution <b>M</b> is (B) 268.5 K f the solution <b>M</b> is (B) 36.0 mm Hg solution <b>M</b> such that the ution is (B) 376.2 K tant for the solubility of 0.8. The number of mole	<ul> <li>(C) 234.2 K</li> <li>(C) 29.5 mm Hg</li> <li>mole fraction of water in</li> <li>(C) 375.5 K</li> <li>N<sub>2</sub> gas in water at 298</li> </ul>	[JEE 2008, 3/163] (D) 150.9 K [JEE 2008, 3/163] (D) 28.8 mm Hg n the solution becomes 0.9. The [JEE 2008, 3/163]
10. 11.	non-volatile and non-di The freezing point of th (A) 268.7 K The vapour pressure o (A) 39.3 mm Hg Water is added to the boiling point of this solu (A) 380.4 K The Henry's law const fraction of N <sub>2</sub> in air is 0 and 5 atm pressure is 3 (A) $4 \times 10^{-4}$	issociative. The solution <b>M</b> is (B) 268.5 K f the solution <b>M</b> is (B) 36.0 mm Hg solution <b>M</b> such that the ution is (B) 376.2 K tant for the solubility of D.8. The number of mole (B) 4.0 × 10 <sup>-5</sup> PC) of a solution contain	<ul> <li>(C) 234.2 K</li> <li>(C) 29.5 mm Hg</li> <li>mole fraction of water in</li> <li>(C) 375.5 K</li> <li>N<sub>2</sub> gas in water at 298 as of N<sub>2</sub> from air dissolve</li> <li>(C) 5.0 × 10<sup>-4</sup></li> </ul>	[JEE 2008, 3/163] (D) 150.9 K [JEE 2008, 3/163] (D) 28.8 mm Hg n the solution becomes 0.9. The [JEE 2008, 3/163] (D) 354.7 K 3 K is 1.0 × 10 <sup>5</sup> atm. The mole ed in 10 moles of water of 298 K [JEE 2009, 3/160]
10. 11. 12.	non-volatile and non-di The freezing point of th (A) 268.7 K The vapour pressure o (A) 39.3 mm Hg Water is added to the boiling point of this solu (A) 380.4 K The Henry's law const fraction of N <sub>2</sub> in air is 0 and 5 atm pressure is 3 (A) 4 × 10 <sup>-4</sup> The freezing point (in 0 (K <sub>f</sub> = 1.86 K kg mol <sup>-1</sup> ) is (A) - 2.3 × 10 <sup>-2</sup> For a dilute solution co elevation in boiling point	issociative. The solution <b>M</b> is (B) 268.5 K f the solution <b>M</b> is (B) 36.0 mm Hg solution <b>M</b> such that the ution is (B) 376.2 K tant for the solubility of D.8. The number of mole (B) $4.0 \times 10^{-5}$ PC) of a solution contain s: (B) $-5.7 \times 10^{-2}$ ontaining 2.5 g of a non- nt at 1 atm pressure is 2	(C) 234.2 K (C) 29.5 mm Hg mole fraction of water in (C) 375.5 K N <sub>2</sub> gas in water at 298 s of N <sub>2</sub> from air dissolve (C) 5.0 × 10 <sup>-4</sup> ing 0.1 g of K <sub>3</sub> [Fe(CN) <sub>6</sub> ] (C) $- 5.7 \times 10^{-3}$ n- volatile non- electroly <sup>o</sup> C. Assuming concentra	[JEE 2008, 3/163] (D) 150.9 K [JEE 2008, 3/163] (D) 28.8 mm Hg n the solution becomes 0.9. The [JEE 2008, 3/163] (D) 354.7 K 3 K is 1.0 × 10 <sup>5</sup> atm. The mole ed in 10 moles of water of 298 K [JEE 2009, 3/160] (D) 4.0 × 10 <sup>-6</sup> (Mol. Wt. 329) in 100 g of water [JEE 2011, 3/180]

mixed in solution

15.*	Benzene and naphthalene fe	form an ideal	solution at roor	n temperature.	For this process,	the true
	statement(s) is (are) :			[JEE(A	dvanced) 2013, 4/	<b>'120]</b>
	(A) $\Delta G$ is positive		(B) $\Delta S_{system}$	is positive		
	(C) $\Delta S_{surroundings} = 0$		(D) ∆H = 0			

- 16. MX<sub>2</sub> dissociates into M<sup>2+</sup> and X<sup>-</sup> ions in an aqueous solution, with a degree of dissociation (α) of 0.5. The ratio of the observed depression of freezing point of the aqueous solution to the value of the depression of freezing point in the absence of ionic dissociation is [JEE(Advanced) 2014, 3/120]
- 17. If the freezing point of a 0.01 molal aqueous solution of a cobalt(III) chloride-ammonia complex (which behaves as a strong electrolyte) is -0.0558°C, the number of chloride(s) in the coordination sphere of the complex is [K<sub>f</sub> of water = 1.86 K kg mol<sup>-1</sup>] [JEE(Advanced) 2015 4/168]
- 18. \* Mixture(s) showing positive deviation from Raoult's law at 35°C is(are) [JEE(Advanced) 2016, 4/124]
   (A) carbon tetrachloride + methanol
   (B) carbon disulphide + acetone
   (C) benzene + toluene
   (D) phenol + aniline

**19.\*** 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, 4/122] (A) The point Z represents vapour pressure of pure liquid M and Raoult's law is obeyed from  $x_{L} = 0$  to  $x_{L} = 1$ . (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

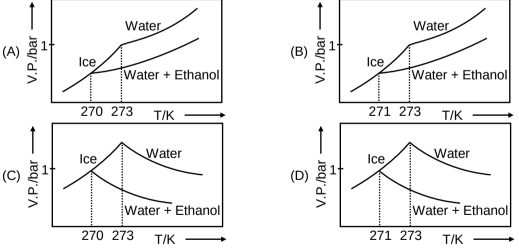


(C) The point **Z** represents vapour pressure of pure liquid **M** and Raoult's law is obeyed when  $x_{L} \rightarrow 0$ 

(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 to 500 g of water changes the freezing point of the solution. Use the freezing point depression constant of water as 2 K kg mol<sup>-1</sup>. The figures shown below represent plots of vapour pressure (V.P.) versus temperature (T). [Molecular weight of ethanol is 46 g mol<sup>-1</sup>] [JEE(Advanced) 2017, 3/122]

Among the following, the option representing change in the freezing point is



21. 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 \_\_\_\_\_. [JEE(Advanced) 2018, 3/120]

(given that the vapour pressure of pure liquid A is 20 Torr at temperature T)

- 22. The plot given below shows P-T 2 curves (where P is the pressure and T is the temperature) for two solvents X and Y and isomolal 760 solutions of NaCl in these 1. Solvent X solvents. NaCl completely Pressure(mmHg) dissociates in both the solvents. 2. Solution of NaCl in solvent X On addition of equal number of 3. Solvent Y moles of a non-volatile solute S in equal amount (in kg) of these 4. Solution of NaCl in solvent Y solvents, the elevation of boiling point of solvent X is three times that of solvent Y. Solute S is known to undergo dimerization in 360 362 367 368 these solvents. If the degree of dimerization is 0.7 in solvent Y. Temperature (K) the degree of dimerization in solvent X is [JEE(Advanced) 2018, 3/120] PART - II : JEE (MAIN) / AIEEE PROBLEMS (PREVIOUS YEARS) **JEE(MAIN) OFFLINE PROBLEMS**  $6.02 \times 10^{20}$  molecules of urea are present in 100 ml of its solution. The concentration of urea solution is: 1 [AIEEE-2004, 3/225] (3) 0.02 M (1) 0.001 M (2) 0.01 M (4) 0.1 M. Which one of the following aqueous solutions will exhibit highest boiling point ? [AIEEE-2004, 3/225] 2. (1) 0.01 M Na<sub>2</sub>SO<sub>4</sub> (2) 0.01 M KNO<sub>3</sub> (3) 0.015 M urea (4) 0.015 M glucose If  $\alpha$  is the degree of dissociation of Na<sub>2</sub>SO<sub>4</sub>, the vant Hoff's factor (i) used for calculating the molecular 3. [AIEEE-2005, 1½/225] mass is : (1) 1 +  $\alpha$ (2)  $1 - \alpha$ (3)  $1 + 2\alpha$ (4)  $1 - 2\alpha$ . 4. Equimolar solutions in the same solvent have : [AIEEE-2005, 3/225] (1) same boiling point but different freezing point (2) same freezing point but different boiling point (3) same boiling and same freezing points (4) differnet boiling and freezing points 5. Two solutions of a substance (non electrolyte) are mixed in the following manner. 480 ml of 1.5 M first solution + 520 mL of 1.2 M second solution. What is the molarity of the final mixture? [AIEEE-2005, 3/225] (1) 1.20 M (2) 1.50 M (3) 1.344 M (4) 2.70 M Benzene and toluene form nearly ideal solutions. At 20°C, the vapour pressure of benzene is 75 torr 6. and that of toluene is 22 torr. The partial vapour pressure of benzene at 20°C for a solution containing 78 g of benzene and 46 g of toluene in torr is : [AIEEE-2005, 3/225] (1) 50 (2) 25 (3) 37.5 (4) 53.5 7. Density of a 2.05 M solution of acetic acid in water is 1.02 g/mL. The molality of the solution is
  - [AIEEE-2005, 3/165] (2) 2.28 mol Kg<sup>-1</sup> (3) 0.44 mol Kg<sup>-1</sup> (1) 3.28 mol Kg<sup>-1</sup> (4) 1.14 mol Kg<sup>-1</sup>
- A mixture of ethyl alcohol and propyl alcohol has a vapour pressure of 290 mm at 300 K. The vapour 8. pressure of propyl alcohol is 200 mm. If the mole fraction of ethyl alcohol is 0.6, its vapour pressure (in mm) at the same temperature will be : [AIEEE-2007, 3/120] (3) 350 (4) 300 (1) 700 (2)360
- 9. A 5.25% solution of a substance is isotonic with a 1.5% solution of urea (molar mass =  $60 \text{g} \text{ mol}^{-1}$ ) in the same solvent. If the densities of both the solutions are assumed to be equal to 1.0 g cm<sup>-3</sup>, molar mass of the substance will be [AIEEE-2007, 3/120] (1) 105.0 g mol<sup>-1</sup> (2) 210.0 g mol<sup>-1</sup> (3) 90.0 g mol<sup>-1</sup> (4) 15.0 g mol<sup>-1</sup>

- 10.The vapour pressure of water at 20°C is 17.5 mm Hg. If 18 g of glucose (C<sub>6</sub>H<sub>12</sub>O<sub>6</sub>) is added to 178.2 g<br/>of water at 20°C, the vapour pressure of the resulting solution will be :[AIEEE-2008, 3/105](1) 15.750 mm Hg(2) 16.500 mm Hg(3) 17.325 mm Hg(4) 17.675 mm Hg
- At 80°C, the vapour pressure of pure liquid 'A' is 520 mm Hg and that of pure liquid 'B' is 1000 mm Hg. If a mixture solution of 'A' and 'B' boils at 80° C and 1 atm pressure, the amount of 'A' in the mixture is (1 atm = 760 mm Hg) [AIEEE-2008, 3/105]
   (1) 34 mol percent (2) 48 mol percent (3) 50 mol percent (4) 52 mol percent
- A binary liquid solution is prepared by mixing n-heptane and ethanol. Which one of the following statement is correct regarding the behaviour of the solution ? [AIEEE-2009, 4/144]
   (1) The solution is non-ideal, showing +ve deviation from Raoult's Law.
  - (2) The solution in non-ideal, showing –ve deviation from Raoult's Law.
  - (3) n-heptane shows +ve deviation while ethanol shows -ve deviation from Raoult's Law.
  - (4) The solution formed is an ideal solution.
- 14.If sodium sulphate is considered to be completely dissociated into cations and anions in aqueous<br/>solution, the change in freezing point of water ( $\Delta T_f$ ), when 0.01 mole of sodium sulphate is dissolved in<br/>1 kg of water, is (K<sub>f</sub> = 1.86 K kg mol<sup>-1</sup>)<br/>(1) 0.0372 K[AIEEE-2010, 4/144]<br/>(3) 0.0744 K(4) 0.0186 K
- 15. On mixing, heptane and octane form an ideal solution. At 373 K, the vapour pressures of the two liquid components (heptane and octane) are 105 kPa and 45 kPa respectively. Vapour pressure of the solution obtained by mixing 25.0 g of heptane and 35 g of octane will be (molar mass of heptane = 100 g mol<sup>-1</sup> and of octane = 114 g mol<sup>-1</sup>) [AIEEE-2010, 4/144] (1) 72.0 kPa (2) 36.1 kPa (3) 96.2 kPa (4) 144.5 kPa
- 16. K<sub>f</sub> for water is 1.86 K kg mol<sup>-1</sup>. If your automobile radiator holds 1.0 kg of water, how may grams of ethylene glycol (C<sub>2</sub>H<sub>6</sub>O<sub>2</sub>) must you add to get the freezing point of the solution lowered to -2.8°C ? [AIEEE 2012, 4/120]

(3) 39 g

(4) 27 g

- Consider separate solution of 0.500 M C<sub>2</sub>H<sub>5</sub>OH(aq), 0.100 M Mg<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub>(aq), 0.250 M KBr(aq) and 0.125 M Na<sub>3</sub>PO<sub>4</sub>(aq) at 25°C. Which statement is true about these solution, assuming all salts to be strong electrolytes ?
  - (1) They all have the same osmotic pressure.
  - (2) 0.100 M Mg<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub>(aq) has the highest osmotic pressure.
  - (3) 0.125 M Na<sub>3</sub>PO<sub>4</sub>(aq) has the highest osmotic pressure.
  - (4) 0.500 M C<sub>2</sub>H<sub>5</sub>OH(aq) has the highest osmotic pressure.
- The vapour pressure of acetone at 20°C is 185 torr. When 1.2 g of a non-volatile substance was dissolved in 100 g of acetone at 20°C, its vapour pressure was 183 torr. The molar mass (g mol<sup>-1</sup>) of the substance is: [JEE(Main) 2015, 4/120]
   (1) 32
   (2) 64
   (3) 128
   (4) 488

 19.
 18 g glucosse (C<sub>6</sub>H<sub>12</sub>O<sub>6</sub>) is added to 178.2 g water. The vapor pressure of water (in torr) for this aqueous solution is.
 [JEE(Main) 2016, 4/120]

 (1) 76.0
 (2) 752.4
 (3) 759.0
 (4) 7.6

- 20. The freezing point of benzene decreases by 0.45°C when 0.2 g of acetic acid is added to 20 g of benzene. If acetic acid associates to form a dimer in benzene, percentage association of acetic acid in benzene will be : (K<sub>f</sub> for benzene = 5.12 K kg mol<sup>-1</sup>) [JEE(Main) 2017, 4/120] (1) 80.4% (2) 74.6% (3) 94.6% (4) 64.6%
- For 1 molal aqueous solution of the following compounds, which one will show the highest freezing point ? [JEE(Main) 2018, 4/120]
   (1) [Co(H<sub>2</sub>O)<sub>4</sub>Cl<sub>2</sub>]Cl.2H<sub>2</sub>O
   (2) [Co(H<sub>2</sub>O)<sub>3</sub>Cl<sub>3</sub>].3H<sub>2</sub>O

		JEE(MAIN) ONL	INE PRO	BLEMS
1.	Choose the correct stat	tement with respect to the		essure of a liquid among the following : [JEE(Main) 2014 Online (19-04-14), 4/120]
	<ul><li>(2) Increase non-linearl</li><li>(3) Decreases linearly</li></ul>	ith increasing temperatu y with increasing temper with increasing arly with increasing temp	re ature	
2.			of Van't Hot	n of $Fe(NH_4)_2(SO_4)_2$ at 25°C is 10.8 atm ff factor (i) will be respectively : (R = 0.082 L [JEE(Main) 2014 Online (19-04-14), 4/120] 3.42 (4) 3 and 5.42
3.	For an ideal solution of	two components A and I	B, which of	the following is true ? [JEE(Main) 2014 Online (19-04-14), 4/120]
	(1) $\Delta H_{\text{mixing}} < 0$ (zero)			
		tronger than A–A and B– interactions are identical		ons
4.	pure benzene and pure	e toluene at this temper	ature are 7 benzene (2) 30.5 to	3.5 mol of toluene. If the vapour pressure of 4.7 torr and 22.3 torr, respectively, then the mole fraction in equilibrium with it will be, [JEE(Main) 2015 Online (10-04-15), 4/120] orr and 0.389 orr and 0.280
5.	Determination of the m by : (1) partial ionization (3) complex formation	olar mass of acetic acid	l in benzene (2) dissoci (4) associa	
6.	The solubility of N <sub>2</sub> in v at 750 torr partial press (1) 0.02		orr partial p (3) 0.0075	pressure is 0.01 g L <sup>-1</sup> . The solubility (in g L <sup>-1</sup> )         [JEE(Main) 2016 Online (09-04-16), 4/120]         5       (4) 0.005
7.	An aqueous solution o dissociation for this solution (1) 0.67		emperature (3) 0.80	e has a van't Hoff factor of 2. The degree of [JEE(Main) 2016 Online (10-04-16), 4/120] (4) 0.50
8.	Na <sub>2</sub> SO <sub>4</sub> is 81.5% ionise	ed, the value of $x$ (K <sub>f</sub> for v	vater = 1.86	e in freezing point was found to be 3.82°C. If 5°C kg mol <sup>-1</sup> ) is approximately : [JEE(Main) 2017 Online (08-04-17), 4/120] (4) 15 g
9.		e 415 and 200 mmHg res		95 g of CHCl <sub>3</sub> . If vapour pressure of CH <sub>2</sub> Cl <sub>2</sub> he mole fraction of CHCl <sub>3</sub> in vapour form is : [JEE(Main) 2017 Online 09-04-17), 4/120] (4) 0.325
10.				electrolyte non-volotatile solute separately in nts are $M_X$ and $M_Y$ , respectively where $M_X$ =
	$\frac{3}{4}$ M <sub>Y</sub> . The relative lowe	ering of vapour pressure	of the solut	tion in X is "m" times that of the solution in Y.
	is :			omparison to that of solvent, the value of "m" [JEE(Main) 2018 Online 15-04-18), 4/120]
	(1) $\frac{4}{3}$	(2) $\frac{3}{4}$	(3) $\frac{1}{2}$	(4) $\frac{1}{4}$
11.		atile, non-electrolyte solu its vapour pressure to 7 (2) 75 g		nass = 50 g mol <sup>-1</sup> ) needed to be dissolved in [JEE(Main) 2018 Online 16-04-18), 4/120] (4) 50 g

12.	Which one of the follo	wing statements regardir	0 /	rect ? 2019 Online 09-01-19), 4/120]
	<ul><li>(2) The value of K<sub>H</sub> in</li><li>(3) The partial pressu</li><li>solution.</li></ul>		w constant) values at th temperature and $K_H$ is functional to the temperature of t	e same temperature. unction of the nature of the gas the mole fraction of the gas in the
13.	A solution containing	62 g ethylene glycol in 2	50 g water is cooled to	–10ºC. If K <sub>f</sub> for water is 1.86 K kg
	mol <sup>-1</sup> , the amount of v (1) 16	water (in g) separated as (2) 32	ice is: [JEE(Main) 2 (3) 48	2019 Online 09-01-19), 4/120] (4) 64
14.	pure A and pure B a		< 10 <sup>3</sup> Pa, respectively. percent of A at this temp	At 350 K, the vapor pressures of The composition of the vapor in perature is : 2019 Online 10-01-19), 4/120]
	(1) $x_A = 0.28$ ; $x_B = 0.7$ (3) $x_A = 0.37$ ; $x_B = 0.6$		(2) $x_A = 0.76$ ; $x_B = 0.3$ (4) $x_A = 0.4$ ; $x_B = 0.6$	
15.		g point for 1 molal solution glucose in the same solv	vent is 2 k. the relation b	e depression in the freezing point between k <sub>b</sub> and K <sub>f</sub> is : 2019 Online 10-01-19), 4/120]
	(1) $K_b = 0.5 K_f$	(2) K <sub>b =</sub> K <sub>f</sub>	(3) $K_{b} = 1.5 K_{f}$	(4) $k_b = 2 k_f$
16.			en added to pure milk to	2°C, while it should have been make the diluted sample ? 2019 Online 11-01-19), 4/120]
	<ul><li>(1) 1 cup of water to 3</li><li>(3) 2 cups of water to</li></ul>		(2) 1 cup of water of 2 (4) 3 cups of water to	2 cups of pure milk
17.	K <sub>2</sub> Hgl <sub>4</sub> is 40% ionised	l in aqueous solution. The		actor (i) is : 2019 Online 11-01-19), 4/120]
	(1) 1.8	(2) 1.6	(3) 2.0	(4) 2.2
18.				t of 12% aqueous solution of Y. If 2019 Online 12-01-19), 4/120] (4) A
19.	benzene shows a dep form dimer in the solu (Given that K <sub>f</sub> = 5 kg r	pression in freezing point tion is 80, then w is : mol <sup>_1</sup> , Molar mass of ben.	equal to 2 K. If the pero [JEE(Main) 2 zoic acid = 122 g mol <sup>-1</sup> )	
	(1) 2.4 g	(2) 1.8 g	(3) 1.0 g	(4) 1.5 g

\_\_\_\_

# Answers

E

# **EXERCISE - 1**

					CI3E -					
				PA	RT - I					
A-1.	Some of the characteristics of supersaturated solution are given below (i) If a crystal of solute is added to supersaturated solution, crystallisation occurs rapidly. (ii) Supersaturated solutions contain more solute than they should have at a particular temperature.									
A-2.	hydrates. Suc	Certain compounds combine with the moisture of atmosphere and are converted into hydroxides or hydrates. Such substances are called hygroscopic. e.g., anhydrous CuSO <sub>4</sub> , quick lime (CaO), anhydrous Na <sub>2</sub> CO <sub>3</sub> etc.								
A-3.	solute, solvent	-solven	ange associate t and solute–sc han disrupting t	olvent inte	eractions. Th	e process is	exothermi	c if the new int	eracti	
B-1.	0.04 g/mL		<b>B-2.</b> (i) 3	0%, (ii) 0	.046 (iii) Na	+ = 5.42 m, S	$S_2O_3^{2-} = 2.7$	71 m		
B-3.	e e	4 m. Ma	plarity = 7.55 M	. ,						
C-1.	-		<sub>o</sub> = 355 mm Hg		= 1165 mm l	⊣g	C-2.	2.4 atm		
D-1.	7.62 bar	D-2.	4 mmol.	E-1.	5.15 g		E-2.	$\frac{W_W}{W_N} = 3.973$		
F-1.	p <sub>A</sub> <sup>0</sup> = 400 mm	of Hg, p	$_{\rm B^0} = 600 \rm mm of$	f Hg			F-2.	27.3 mole %		
F-3.	$P_{B^0} = 0.7 \text{ atm}$	$8 P_{A^0} =$	1.9 atm	F <b>-4</b> .	14.16 mo	le percent be	enzene			
F-5.			V < 80ml.	G-2.				Raoult's law, $\Delta$	S > 0	
н-1.	1 0 - 0.002.	•	V COOM	0 2.					0 - 0	
· · · · ·	Solute	Dis	sociation/assoc	iation	Degree of d	issociation /a	ssociation	n	i	
	KCI		$\frac{\text{reaction}}{\text{KCI} \rightarrow \text{K}^+ + \text{CI}}$	-		1		2	2	
	$H_2SO_4$	H	$I_2SO_4 \rightarrow 2H^+ + S$			1		3	3	
	H <sub>3</sub> COOH (in water)		$COOH \rightarrow H^+ + CH$			0.2		2	1.2	
CH	3COOH (in benzene		$_{3}COOH \rightarrow (CH_{3}COOH)$		No oppo	0.5	aiatian	0.5	0.75	
	Urea NaBr	INU as	ssociation or diss NaBr $\rightarrow$ Na <sup>+</sup> + E		NU 8550	ciation or disso 0.8	JCIALION	Not defined	1 1.8	
	A		$3A \rightarrow A_3$	-		1		0.33	0.33	
H-2.	75%	I-1.	(a) 60 g/mol	, (b) 333.	3 g		I-2.	746.2 mm of H	la	
J-1.	(a) M = 94.52		., .	J-2.	232			S <sub>8</sub>	5	
J-4.	1.075, 7.5.	<b>K-1.</b>	(a) Urea < N			⊃O₄ (h) 6 ′	15 atm	0.		
к-2.		K-3.	. ,			nust have be		timoo		
<b>~-</b> 2.	α = 0.81	<b>к-</b> э.	50% K2SO4	K-4.		nust nave be		umes		
	(D)	A 2	(D)		RT - II	<b>C</b> 1	<b>(D)</b>	<b>C</b> 2		
^ 1	(B)	A-2.	(B)	B-1.	(B)	C-1.	(B)	C-2.	(C)	
		• •				D-2.		D-3.	(B)	
C-3.	(B)	C-4.	(B)	D-1.	(C)		(A)			
C-3. D-4.	(D)	D-5.	(B)	E-1.	(C)	F-1.	(B)	F-2.		
C-3. D-4.								F-2. G-2.		
C-3. D-4. F-3.	(D)	D-5.	(B)	E-1.	(C)	F-1.	(B)		(A)	
C-3. D-4. F-3. G-3.	(D) (C)	D-5. F-4.	(B) (A)	E-1. F-5.	(C) (A)	F-1. G-1.	(B) (C)	G-2.	(A) (C)	
C-3. D-4. F-3. G-3. H-2.	(D) (C) (A)	D-5. F-4. G-4.	(B) (A) (C)	E-1. F-5. G-5.	(C) (A) (A)	F-1. G-1. G-6.	(B) (C) (C)	G-2. H-1.	(A) (C) (B)	
C-3. D-4. F-3. G-3. H-2. I-3.	(D) (C) (A) (A)	D-5. F-4. G-4. H-3.	(B) (A) (C) (B)	E-1. F-5. G-5. H-4.	(C) (A) (A) (B)	F-1. G-1. G-6. I-1.	(B) (C) (C) (B)	G-2. H-1. I-2.	(A) (C) (B) (D)	
A-1. C-3. D-4. F-3. G-3. H-2. I-3. J-2. J-7.	(D) (C) (A) (A) (B)	D-5. F-4. G-4. H-3. I-4.	(B) (A) (C) (B) (C)	E-1. F-5. G-5. H-4. I-5.	(C) (A) (A) (B) (B)	F-1. G-1. G-6. I-1. I-6.	(B) (C) (C) (B) (A)	G-2. H-1. I-2. J-1.	<ul> <li>(C)</li> <li>(A)</li> <li>(C)</li> <li>(B)</li> <li>(D)</li> <li>(D)</li> <li>(A)</li> </ul>	

<u>Solut</u>	ion & Colligativ	e Prop	erties						
1.	(A) – (r); (B) –	- (s); (C)	– (q); (D) – (p)	PA	RT - III 2.	(A) – (p,s,r); (E	B) – (p,q,t); (C	) – (p); (C	0) – (p,q,
			E	XER	CISE ·	- 2			
				PA	RT - I				
1.	(C)	2.	(D)	3.	(B)	4.	(C)	5.	(D)
6.	(D)	7.	(B)	8.	(B)	9.	(B)	10.	(A)
11.	(B)	12.	(A)	13.	(D)	14.	(B)		
				PA	RT - II				
1.	18	2.	5	3.	21	4.	20	5.	27
6.	4 (A,B,C,D)	7.	2	8.	48	9.	43	10.	2
11.	60	12.	60	13.	4				
					RT - III				
1.	(ABD)	2.	(ABCD)	3.	(ABD)	4.	(ABCD)	5.	(AC)
6.	(A)	7.	(AB)	8.	(B)	9.	(AB)	10.	(B)
11.	(ABC)	12.	(BD)	13.	(AB)	14.	(AD)	15.	(BCD)
16.	(ABC)								
					RT - IV				
1.	(B)	2.	(C)	3.	(B)	4.	(D)	5.	(B)
6.	(A)	7.	(D)	8.	(B)	9.	(C)	10.	(D)
			E	XER	CISE ·	- 3			
				PA	RT - I				
1.	(C)	2.	(B)	3.	(B)	4.	(A)	5.	(C)
6.	122, 224	7.	(A)	8.	(A)	9.	(D)	10.	(B)
11.	(B)	12.	(A)	13.	(A)	14.	(A)	15.	(BCD)
16.	2	17.	(1)	18.	(AB)	19.	(BD)	20.	(A)
21.	19	22.	0.05						
					RT - II				
4	( <b>2</b> )	n				OBLEMS	(2)	F	(2)
1. c	(2)	2.	(1)	3.	(3) ( <b>2</b> )	4.	(3)	5.	(3)
6.	(1)	7.	(2)	8.	( <b>3</b> )	9.	(2)	10.	(3)
11.	(3)	12.	(1)	13.	(2)	14.	(2)	15.	(1)
16.	(2)	17.	(1)	18.	(2)	19.	(2)	20.	(3)
21.	(2)		JEE(MA	AIN) ON		OBLEMS			
1.	(2)	2.	(1)	3.	(4)	4.	(3)	5.	(4)
6.	(2)	7.	(4)	8.	(1)	9.	(4)	10.	(2)
11.	(Bonus)	12.	(4)	13.	(4)	14.	(1)	15.	(4)