Revision Notes on Solution:

A solution is a homogeneous mixture of two (or more) substances, the composition of which may vary between certain limits. A solution consisting of two components is called *binary solution*. The component which is present in large quantity is called *solvent* and the component which is small in quantity is called *solute*. If both components are in same physical state.

Type of Solutions

All the three states of matter (gas, liquid or solid) may behave either as solvent or solute. Depending on the state of solute or solvent, mainly there may be the following seven types of binary solutions.

S.No.	Solute	Solvent	Example
1	Gas	Gas	Air
2	Gas	Liquid	Aerated water $(CO_2 + H_2O)$
3	Gas	Solid	Hydrogen in palladium
4	Liquid	Liquid	Alcohol in water, benzene in toluene
5	Liquid	Solid	Mercury in zinc amalgam
6	Liquid	Gas	CO ₂ dissolved in water
7	Solid	Liquid	Sugar in water, common salt in water
8	Solid	Gas	Smoke
9	Solid	Solid	Various alloys

Vapour Pressure:

- The pressure exerted by vapours over the liquid surface at equilibrium is called vapour pressure of the liquid.
- If solute is non volatile solid or liquid the vapour pressure of solution is equal to partial vapour pressure of solvent in the solution.
- If the solute is volatile solid or liquid, then vapour pressure will be equal to the sum of partial vapour pressure of solute and that of solvent.

Raoult's Law:

"The partial vapour pressure of any component in the solution is directly proportional to its mole fraction".				
For a binary solution of two components A and B,				
$P_A = X_A$				

$$P_B = X_B$$

Where

 P^0_A = vapour pressure of component A in pure state.

 P_A = vapour pressure of component A in the solution.

 P^{0}_{B} = vapour pressure of component B in pure state.

 P_B = vapour pressure of component B in the solution

Limitations of Raoult's Law

- Raoult's law is applicable only to very dilute solutions.
- It is applicable to solutions containing non-volatile solute only.
- It is not applicable to solutes which dissociate or associate in a particular solution

Raoult's Law in Combination with Dalton's Law of Partial Pressure:

$$P_T = X_A P_A^0 + X_B P_B^0 = P_B^0 + (P_A^0 - P_B^0) X_A$$

Where

 P_T = Total Vapour Pressure of the Solution.

Ideal and Non-Ideal Solutions:

• Ideal Solution:

?These solutions obey Raoult's law at all compositions of solute in solvent at all temperature

Two liquids A and B form and ideal solution when A – A and B–B molecular attractions will be same and hence A–B molecular attraction will be almost same as A–A and B–B molecular attraction.

For Ideal Solution:

- 1. $dH_{mixing} = 0$, i.e. no heat should be absorbed or evolved during mixing
- 2. $dV_{mixing} = 0$, i.e. no expansion or contraction on mixing

Examples, Ethyl chloride and ethyl bromide, n-hexane and n-heptane, CCl₄ and SiCl₄

• Non-Ideal Solution:

?These solutions deviate from ideal behaviour and do not obey Raoult's law over entire range of composition.

For non ideal solutions,

1.
$$dH_{mixing} \neq 0$$

2.
$$dH_{mixing} \neq 0$$

Here we may have two cases

A) Positive Deviation:

$$1. P_A > X_A \& P_B > X_B$$

$3. \mathrm{dV}_{\mathrm{mix}} > 0$		
Example: Cyclohexane and Ethanol		7
B) Negative Deviation:		
1. $P_A > X_A \& P_B > X_B$		
$2. dH_{mix} < 0$		
$3. dV_{\text{mix}} < 0$		
Colligative Properties		
The properties of dilute solutions which care called <i>colligative properties</i> (denoted)		particles of solute present in the solution and not on their identity ellection).
Lowering of Vapour Pressure b	y a Non-Volatile S	Solute
The relative lowering of vapour pressure present in the solution.	of a solution containing	g a non-volatile solute is equal to the mole fraction of the solute
Elevation of Boiling Point by a I	Non-Volatile Solu	ite:
Since the addition of a non-volatile solute	e lowers the vapour pre	essure of the solvent, the vapour pressure of a solution is always

lower than that of the pure solvent, and hence it must be heated to a higher temperature to make its vapour pressure equal to

Depression of Freezing Point by a Non-Volatile Solute:

where M_1 = molecular weight of solute and w_2 and w_1 are weights of solute and solvent

atmospheric pressure.

 $2. dH_{mix} > 0$

whore $M = molecular$	ular weight of solute and w_2 and w_1 are weights of solute and solvent
-	
Osmosis and Os	mone Pressure:
	he phenomenon of the passage of pure solvent from a region of lower concentration (of the solution) to a higher concentration through a semi-permeable membrane.
	ressure: Excess pressure which must be applied to a solution in order to prevent flow of solvent into the ugh the semi-permeable membrane.
V = nRT	
where	
= Osmotic p	pressure
V = volume of s	solution
n = no. of mole	es of solute that is dissolved
R = Gas consta	nt
T = Absolute to	emperature
sotonic Solutions:	A pair of solutions having same osomotic pressure is called isotonic solutions.
Abnormal N	lolecular Weight and Van't Hoff Factor:
Van't Hoff Facto	or:
Van't Hoff, in order	r to account for all abnormal cases introduced a factor i known as the Van't Hoff factor, such that
Degree of Assoc	ciation:
	total number of molecules which combine to form bigger molecule
Let a be the degree	e of association, then,
The number of una	ssociated moles = 1-
The number of asso	ociated moles =/n
Total number of eff	fective moles = 1- /n
Obviously, $i < 1$	
Degree of Disso	ciation
The fraction of the	total number of molecules which dissociates in the solution, that is, breaks into simpler molecules or ions.
$KCl \leftrightarrow K^+ + Cl^-$	
1	

Thus, the total number of moles after dissociation = 1-

Hence,
$$i = (1 + 1)/1$$

$$i = 1 +$$
 = $1 + (2-1)$

In general,
$$i = 1 + (n-1)$$
,

Where, n = number of particles (ions) formed after dissociation

From the above formula, it is clear that i > 1