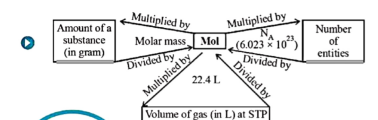


PHYSICAL CHEMISTRY



SOME BASIC CONCEPTS OF CHEMISTRY

Molecular mass

Average relative mass of one molecule $\frac{1}{12}$ × mass of C-12 atom

- Molecular mass = 2 × VD
 Eq. wt. of metal

$$= \frac{\text{wt. of metal}}{\text{wt. of H}_2 \text{ displaced}} \times 1.008$$

Eq. wt. of metal = $\frac{\text{wt. of metal}}{\text{wt. of oxygen combined}} \times 8$

$$= \frac{\text{wt. of metal}}{\text{wt. of chlorine combined}} \times 35.5$$

- Molecular formula = (Empirical formula)_n
 - Energy of electron in species with one

ATOMIC STRUCTURE

$$E_{n} = \frac{-2\pi^{2} m e^{4} Z^{2}}{n^{2} h^{2}}$$

For energy in SI syste

$$E_{n} = \frac{-2\pi^{2} me^{4} Z^{2}}{n^{2} h^{2} (4\pi\epsilon_{0})^{2}}$$

$$E_{n} = \frac{-13 \cdot 2Z^{2}}{n^{2}} kJ \text{ mol}^{-1}$$

$$mvr = \frac{nh}{2\pi}$$

- $r = \frac{n^2 h^2}{4\pi^2 m Z e^2} = 0.529 \left(\frac{n^2}{Z}\right) A$
- Total energy of electron in the nth shell $= K.E. + P.E. = kZ \frac{e^2}{2r} + \left(-\frac{kZe^2}{r}\right) = -\frac{kZe^2}{2r}$

 $\overline{\upsilon} = \frac{1}{\lambda} = RZ^2 \left[\frac{1}{n^2} - \frac{1}{n^2} \right], [R = 1.0968 \times 10^7 \,\text{m}^{-1}]$

$$E = hv = \frac{hc}{\lambda}$$

$$\lambda = \frac{h}{\sqrt{2m \times K.E.}}$$

- No. of spectral lines produced when an electron drops from n^{th} level to ground level = $\frac{n(n-1)}{2}$
- Heisenberg's Uncertainty Principle (Δx) (Δp) $\geq h/4\pi$
- Nodes (n-1) = total nodes, ℓ = angular nodes, $(n-\ell-1)$ = Radial nodes
- Orbital angular momentum : $\sqrt{\ell (\ell+1)} \frac{h}{2\pi} = \sqrt{\ell(\ell+1)}h$

CHEMICAL **BONDING**

- (i) % ionic character
- $= \frac{\text{Actual dipole moment}}{\text{Calculated dipole moment}} \times 100$
- (ii) Dipole moment is helpful in predicting geometry and polarity of molecule.
- Fajan's Rule: Following factors are helpful in increasing covalent character in ionic compounds
 - (i) Small cation
- (ii) Big anion
- (iii) High charge on cation/anion
- (iv) Cation having pseudo inert gas configuration (ns²p⁶d¹⁰) e.g. Cu⁺, Ag⁺, Zn⁺², Cd⁺²
- M.O. theory:
 - (i) Bond order = $\frac{1}{2}(N_b N_a)$
 - (ii) Higher the bond order, higher is the bond dissociation energy, greater is the stability, shorter is the bond length.
- Formal charge (F.C.) on an atom in a Lewis structure
 - = [total number of valence electrons in the free atoms]
 - -[total number of non-binding (lone pair) electrons]
 - $-\frac{1}{2}$ [total number of bonding (shared) electrons]
- Relative bond strength: $sp^3d^2>dsp^2>sp^3>sp^2>sp>p-p$ (Co-axial) > s - p > s - s > p - p (Co-lateral)



- VSEPR theory
 - (i) (LP-LP) repulsion > (LP-BP) > (BP-BP)
 - (ii) NH₂ \rightarrow Bond Angle 106° 45' because (LP-BP) repulsion > (BP-BP) H₂O \rightarrow 104° 27'because (LP-LP) repulsion >(LP-LB)>(BP-BP)
- Hybridisation:

$$= \frac{1}{2} \begin{cases} \text{number of valence electrons of central atom} \\ + \text{number of monovalent atoms attached to it} \\ + \text{negative charge if any - positive charge if any} \end{cases}$$

CHEMICAL EQUILIBRIUM

- $K_p = K_c (RT)^{\Delta n_g}$ where $\Delta n_{g} = n_P n_R$
- Free Energy Change (ΔG)
 - (a) If $\Delta G = 0$ then reversible reaction would be in equilibrium, $K_c = 0$
 - (b) If $\Delta G = (+)$ ve then equilibrium will be displaced in backward direction; $K_c < 1$
- (c) If $\Delta G = (-)$ ve then equilibrium will shift in forward direction; $K_c > 1$
- (a) K_c unit \rightarrow (moles/lit) $^{\Delta n}$,
 - (b) K_n unit $\rightarrow (atm)^{\Delta n}$
- Reaction Quotient and Equilibrium Constant

Consider the following reversible reaction

$$A+B \rightleftharpoons C+D$$

$$\therefore Q_c = \frac{[C][D]}{[A][B]}$$

Case I: If $Q_c < K_c$ then: [Reactants] > [Products]

then the system is not at equilibrium

Case II: If $Q_c = K_c$ then: The system is at equilibrium.

Case III : If $Q_c > K_c$ then : [Products] > [Reactants]

The system is not at equilibrium.

A relationship between the equilibrium constant K_C, reaction quotient and Gibb's energy.

 $\Delta G = \Delta G^{\circ} + RT \ln Q$

At equilibrium $\Delta G = 0$ and Q = K then

- $\Delta G^{\circ} = -RT \ln K_c$
- ∴ $\Delta G^{\circ} = -RT \ln K_p$ Le-Chatelier's principle
 - (i) Increase of reactant conc. (Shift reaction forward)
 - (ii) Decrease of reactant conc. (Shift reaction backward)
 - (iii) Increase of pressure (from more moles to less moles)
 - (iv) Decrease of pressure (from less moles to more moles)
 - For exothermic reaction decrease in temp. (Shift
 - forward)
 - (vi) For endothermic increase in temp. (Shift backward)

IONIC **EQUILIBIRIUM**

- Lewis Acid (e⁻ pair acceptor) → CO₂, BF₃, AlCl₃, ZnCl₂, normal
- (ii) Lewis Base (e⁻ pair donor) → NH₃, ROH, ROR, H₂O, RNH₂, normal anions
- Dissociation of Weak Acid and Weak Base

 Weak Acid, $K_a = Cx^2/(1-x)$ or $K_a = Cx^2$; x << 1Weak Base, $K_b = Cx^2/(1-x)$ or $K_b = Cx^2$; x << 1

- Buffer solution {Henderson equation}:
 - Acidic, $pH = pK_a + log \{Salt/Acid\}$. For maximum buffer action $pH = pK_a$ Range of buffer $pH = pK_a \pm 1$
 - Alkaline \rightarrow pOH = pK_b + \log {Salt/Base} for max. buffer action pH = $14 - pK_b$ Range pH = $14 - pK_b \pm 1$
 - (iii) Buffer Capacity = $\frac{\text{Moles / lit of Acid or Base Mixed}}{\text{Moles / lit of Acid or Base Mixed}}$ Change in pH
- Relation between ionisation constant (K_i) and degree of ionisation(α):-

$$K_i = \frac{\alpha^2}{(1-\alpha)V} = \frac{\alpha^2 C}{(1-\alpha)}$$
 (Ostwald's dilution law)

It is applicable to weak electrolytes for which $\alpha \ll 1$ then

$$\alpha = \sqrt{K_i V} = \sqrt{\frac{K_i}{C}} \text{ or } V \uparrow C \downarrow \alpha \uparrow$$

Common ion effect: By addition of X mole/L of a common ion, to a weak acid (or weak base) a becomes equal to

$$\frac{K_a}{X}$$
 (or $\frac{K_b}{X}$) [where α = degree of dissociation]

- (i) If solubility product > ionic product then the solution is 0 unsaturated and more of the substance can be dissolved in it. (ii) If ionic product > solubility product the solution is super saturated (principle of precipitation).
- Salt of weak acid and strong base:

pH = 0.5 (pK_w + pK_a + log c); h =
$$\sqrt{\frac{K_h}{c}}$$
; $K_h = \frac{K_w}{K_a}$
(h = degree of hydrolysis)

Salt of weak base and strong acid:

$$pH = 0.5 (pK_w - pK_b - \log c); h = \sqrt{\frac{K_w}{K_b \times c}}$$

Salt of weak acid and weak base:

$$pH = 0.5 (pK_w + pK_a - pK_b); h = \sqrt{\frac{K_w}{K_a \times K_b}}$$
Unit of rate constant:

- - $k = mol^{1-n} lit^{n-1} sec^{-1}$

CHEMICAL **KINETICS**

- Order of reaction It can be fraction, zero or any whole number.
- Molecularity of reaction is always a whole number. It is never more than three. It cannot be zero.
- First Order Reactions:

$$k = \frac{2.303}{t} \log_{10} \frac{a}{(a-x)} \& t_{1/12} = \frac{0.693}{k}$$

 $[A]_{t} = [A]_{0}e^{-kt}$

Second Order Reactions:

When concentration of A and B taking same.

$$k_2 = \frac{1}{t} \left(\frac{x}{a(a-x)} \right)$$

When concentration of A and B are taking different -

$$k_2 = \frac{2.303}{t(a-b)} \log \frac{b(a-x)}{a(b-x)}$$



Zero Order Reaction: x = kt and $t_{1/2} = \frac{a}{2k}$

The rate of reaction is independent of the concentration of the reacting substance.

 \bigcap Time of n^{th} fraction of first order process,

$$t_{1/n} = \frac{2.303}{k} \log \left(\frac{1}{1 - \frac{1}{n}} \right)$$

- Amount of substance left after 'n' half lives = $\frac{[A]_0}{2^n}$
- Arrhenius equation: $k = Ae^{-E_a/RT}$, slope = $\frac{-E_a}{2.303R}$ and Temperature Coefficient

$$\log\left(\frac{k_2}{k_1}\right) = \frac{E_a}{2.303} \left(\frac{T_2 - T_1}{T_1 T_2}\right)$$

It has been found that for a chemical reaction with rise in temperature by 10 °C, the rate constant gets nearly doubled.

$$k = PZ_{AB}e^{-E_a/RT}$$

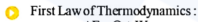


- Oxidant itself is reduced (gives O_2)
 Or Oxidant $\longrightarrow e^-(s)$ Acceptor
 Reductant itself is oxidised (gives H_2)
 Or reductant $\longrightarrow e^-(s)$ Donor
- (i) Strength of acid ∞ O.N (ii) Strength of base ∞ 1/O.N
- (i) Electrochemical Series:- Li, K, Ba, Sr, Ca, Na, Mg, Al, Mn, Zn, Cr, Fe, Cd, Co, Ni, Sn, Pb, H₂, Cu, Ag, Pt, Au.
- (ii) As we move from top to bottom in this series
- (a) Standard Reduction Potential 1
- (b) Standard Oxidation Potential ↓
- (c) Reducing Capacity ↓
- (d) IP ↑

THERMO-

DYNAMICS

(e) Reactivity ↓



 $\Delta E = Q + W$

Expression for pressure volume work $W = -P\Delta V$

Maximum work in a reversible expansion:

$$W = -2.303 \text{ n RT log } \frac{V_2}{V_1}$$

= -2.303 nRT log
$$\frac{P_1}{P_2}$$

$$W_{rev} \ge W_{irr}$$

 $q_v = c_v \Delta T = \Delta U, q_p = c_p \Delta T = \Delta H$

Enthapy changes during phase transformation

- (i) Enthalpy of Fusion
- (ii) Heat of Vapourisation
- (iii) Heat of Sublimation
- Kirchoff's equation :

$$\Delta E_{T_2} = \Delta E_{T_1} + \Delta C_V (T_2 - T_1) [constant V]$$

$$\Delta H_{T_2} = \Delta H_{T_1} + \Delta C_P (T_2 - T_1)$$
 [constant P]

Entropy(s): Measure of disorder or randomness $\Delta S = \Sigma S_p - \Sigma S_R$

$$\Delta S = \frac{q_{rev}}{T} = 2.303 \text{ nR log } \frac{V_2}{V_1} = 2.303 \text{ n R log } \frac{P_1}{P_2}$$

Free energy change: $\Delta G = \Delta H - T\Delta S$, $\Delta G^o = -nFE^o_{cell}$ $-\Delta G = W(maximum) - P\Delta V$, $\Delta G_{system} = -T\Delta S_{total}$

Ο ΔΗ	ΔS	ΔG	Reaction characteristics
-	+	Always negative	Reaction is spontaneous at all temperature.
+	-	Always positive	Reaction is nonspontaneous at all temperature
_	_	Negative at low	Spontaneous at low temp. &
		temperature but positive at high temperature	non spontaneous at high temperature
+	+	Positive at low temp. but negative at high	Non spontaneous at low temp. & spontaneous at high temp.
		temperature	comp.



- m = Z.I.t
- Degree of dissociation : $\alpha = \frac{\lambda_{eq}}{\lambda_{eq}^0}$
- Specific conductance

$$\kappa = \frac{1}{\rho} = \frac{\ell}{R.a} = G \times \frac{\ell}{a} = G \times \text{cell constant (G*)};$$

$$\Lambda_{\rm m} = \frac{\kappa \times 1000}{M}, \ \Lambda_{\rm eq} = \frac{\kappa \times 1000}{N}$$

- Nohlrausch's law: $\Lambda_m^0 = x\lambda_A^0 + y\lambda_B^0$
- Nernst Equation

ELECTRO-

CHEMISTRY

$$E = E^{0} - \frac{0.0591}{n} \log_{10} \frac{[Products]}{[Reactants]}$$

&
$$E^{o}_{Cell} = E^{o}_{right} + E^{o}_{left}$$
 & $K_{eq.} = antilog \left[\frac{nE^{o}}{0.0591} \right]$
 $\Delta G = -nFE_{cell}$ & $\Delta G^{o} = -nFE^{o} cell = -2.303 RT log K_{cell}$

&
$$W_{\text{max}} = +nFE^{\circ} \& \Delta G = \Delta H + T \left(\frac{\partial \Delta G}{\partial T} \right)_{P}$$

Calculation of pH of an electrolyte by using a calomel

electrode:
$$pH = \frac{E_{cell} - 0.2415}{0.0591}$$

Thermodynamic efficiency of fuel cells :

$$\eta = \frac{-\Delta G}{\Delta H} = \frac{-nFE_{cell}^{o}}{\Delta H}$$

For H₂-O₂ fuel cells it is 95%.

- $P = K_H \cdot x$
- Normality (N) = $\frac{\text{number of equivalents}}{\text{volume of the solution in litres}}$
- $Molarity(M) = \frac{number of moles}{volume of the solution in litres}$





SOLUTION AND COLLIGATIVE **PROPERTIES**

- P=p_A+p_B=p°_AX_A+p°_BX_B
 Characteristics of an ideal solution: (i) $\Delta_{\text{sol}} V = 0$ (ii) $\Delta_{sol}H=0$
- Relative lowering of vapour pressure

$$= \frac{P_A^o - P_A}{P_A^o}$$

$$\frac{P_A^o-P_A}{P_A^o}=X_B=\frac{n_B}{n_A+n_B}$$

- Colligative ∞ Number of particles/ ions/ moles of solute properties
- Depression of freezing point, $\Delta T_f = K_f m$
- Elevation in boiling point with relative lowering of vapour

$$\Delta T_b = \frac{1000 K_b}{M_1} \left(\frac{p^o - p}{p^o} \right) (M_1 = \text{mol. wt. of solvent})$$

Osmotic pressure (P) with depression in freezing point ΔT_f

$$P = \Delta T_f \times \frac{dRT}{1000K_f}$$

- Relation between Osmotic pressure and other colligative properties:
 - (i) $\pi = \left(\frac{p_A^0 p_A}{p_A^0}\right) \times \frac{dRT}{M_B}$ Relative lowering of vapour pressure
 - (ii) $\pi = \Delta T_b \times \frac{dRT}{1000K_b}$ Elevation in boiling point
 - (iii) $\pi = \Delta T_f \times \frac{dRT}{1000K_f}$ Depression in freezing point
- $i = \frac{Normal\ molar\ mass}{Observed\ molar\ mass} = \frac{Observed\ colligative\ property}{Normal\ colligative\ property}$
- Degree of association $a = (1 i) \frac{n}{n-1}$

& degree of dissociation (α) = $\frac{1-1}{n-1}$

GASEOUS STATE

In 1 Ideal gas equation: PV = nRT(i) R = 0.0821 liter atm. deg^{-1} mole⁻¹ (ii) R = 2 cals. $deg.^{-1}$ mole⁻¹ (iii) R = 8.314 JK⁻¹ mole⁻¹ Velocities related to gaseous state

RMS velocity =
$$\sqrt{\frac{3PV}{M}} = \sqrt{\frac{3RT}{M}} = \sqrt{\frac{3P}{d}}$$

Average speed =
$$\sqrt{\frac{8RT}{M}}$$
 & Most probable speed = $\sqrt{\frac{2RT}{M}}$

Average speed = $0.9213 \times RMS$ speed RMS speed = $1.085 \times \text{Average speed}$ $MPS = .816 \times RMS$; RMS = 1.224 MPSMPS: A.V. speed: RMS = 1:1.128:1.224

- Rate of diffusion $\propto \frac{1}{\sqrt{\text{density of gas}}}$
- van der Waal's equation

$$\left(P + \frac{n^2 a}{V^2}\right) (V - nb) = nRT \text{ for } n \text{ moles}$$

Z (compressibility factor) = $\frac{PV}{nPT}$; Z = 1 for ideal gas

$$T_C = \frac{8a}{27Rb}$$
, $P_C = \frac{a}{27b^2}$, $V_C = 3b$, $T_b = \frac{a}{bR}$

Available space filled up by hard spheres (packing fraction):

SOLID AND LIQUID STATE

Simple cubic =
$$\frac{\pi}{6}$$
 = 0.52

$$bcc = \frac{\pi\sqrt{3}}{8} = 0.68 \quad fcc = \frac{\pi\sqrt{2}}{6} = 0.74$$

$$hcp = \frac{\pi\sqrt{2}}{6} = 0.74$$

diamond =
$$\frac{\pi\sqrt{3}}{6}$$
 = 0.34

Radius ratio and co-ordination number (CN)

Limiting radius ratio	CN	Geometry
[0.155-0.225]	3	[Plane triangle]
[0.255–0.414]	4	[Tetrahedral]
[0.414–0.732]	6	[Octahedral]
[0.732–1]	8	[bcc]

Atomic radius r and the edge of the unit cell: Pure elements:

Simple cubic =
$$r = \frac{a}{2}$$
; $bcc r = \frac{\sqrt{3}a}{4}$; $fcc = \frac{\sqrt{2}a}{4}$

- Relationship between radius of void (r) and the radius of the sphere (R): r (tetrahedral) = 0.225 R; r (octahedral) =0.414R
- Paramagnetic: Presence of unpaired electrons [attracted by magnetic field]
- Ferromagnetic: Permanent magnetism [↑↑↑↑]
- Antiferromagnetic: Net magnetic moment is zero $[\uparrow \downarrow \uparrow \downarrow]$
 - Ferrimagnetic : Net magnetic moment is three $[\uparrow \downarrow \downarrow \uparrow \uparrow \uparrow]$ Emulsion: Colloidal soln. of two SURFACE
 - **CHEMISTRY &** immiscible liquids [O/W emulsion, COLLOIDAL W/O emulsion] STATE Emulsifier : Long chain

hydrocarbons are added to

- stabilize emulsion. Lyophilic colloid: Starchy gum, gelatin have greater affinity for solvent.
- Lyophobic colloid: No affinity for solvent, special methods are used to prepare sol. [e.g. As₂S₃, Fe(OH)₃ sol]
- Preparation of colloidal solution: (i) Dispersion methods (ii) Condensation method.
- Coagulating power $\propto \frac{1}{\text{Flecculating value}}$
- Properties of colloidal solution: (i) Tyndall effect (ii) Brownian movement (iii) Coagulation (iv) Filtrability.









INORGANIC CHEMISTRY

PERIODIC TABLE

General electronic configuration

(of outer orbits) s-block ns¹⁻² s-block

 ns^2np^{1-6} p-block

Pr (L To R)

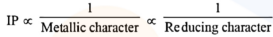
 $(n-1)d^{1-10} ns^{1-2}$ d-block $(n-2)f^{1-14}s^2p^6d^{10}$ f-block

 $(n-1)s^2p^6d^0$ or 1 ns^2

Gr(TtoB)

Property

- (i) Atomic radius (ii) Ionisation potential
- (iii) Electron affinity
- (iv) Electronegativity
- (v) Metallic character or electropositive character
- (vi) Alkaline character of hydroxides
- (vii) Acidic character
- (viii) Reducing property (ix) Oxidising property
- (x) Non metallic character



 $EA \propto \frac{1}{\text{size}} \propto \text{nuclear charge.}$

Second electron affinity is always negative.

Electron affinity of chlorine is greater than fluorine (small atomic size).

The first element of a group has similar properties with the second element of the next group. This is called diagonal relationship. The diagonal relationship disappears after IV

group.

s-BLOCK

ELEMENTS

Atomic radii: Li < Na < K < Rb < Cs

- Electronegativity: Li>Na>K>Rb>Cs
- First ionization potential: Li > Na > K > Rb>Cs
- Melting point Li > Na > K > Rb
- Colour of the flame Li Red, Na -Golden, K - Violet, Rb - Red, Cs - Blue, Ca - Brick red, Sr - Blood red, Ba-Apple green
- Rb and Cs show photoelectric effect.
- Stability of hydrides: LiH > NaH > KH > RbH > CsH
- Basic nature of hydroxides: LiOH < NaOH < KOH < RbOH <CsOH
- Hydration energy: Li>Na>K>Rb>Cs
- Reducing character: Li > Cs > Rb > K > Na

Stability of +3 oxidation state:

B>A1>Ga>In>T1

- Stability of +1 oxidation state: Ga < In < Tl
- Basic nature of the oxides and

B<Al<Ga<In<Tl

Relative strength of Lewis acid: $BF_3 < BCl_3 < BBr_3 < BI_3$

Ionisation energy: B>Al < Ga > In < Tl

Electronegativity: Electronegativity first decreases from B to Al and then increases marginally.

CARBON FAMILY

- Reactivity: C < Si < Ge < Sn < Pb
 </p>
- Metallic character : C < Si < Ge < Sn < Pb</p>
- Acidic character of the oxides: $CO_2 > SiO_2 > GeO_2 > SnO_2 > PbO_2$ Weaker acidic (amphoteric)
- Reducing nature of hydrides $CH_4 < SiH_4 < GeH_4 < SnH_4 < PbH_4$

Thermal stability of tetrahalides

 $CCl_4 > SiCl_4 > GeCl_4 > SnCl_4 > PbCl_4$

- Oxidising character of M⁺⁴ species $GeCl_4 < SnCl_4 < PbCl_4$
- Ease of hydrolysis of tetrahalides

NITROGEN FAMILY

 $SiCl_4 < GeCl_4 < SnCl_4 < PbCl_4$ Acidic strength of trioxides : $N_2O_3 >$ $P_2O_3 > As_2O_3$

Acidic strength of pentoxides $N_2O_5 > P_2O_5 > As_2O_5 > Sb_2O_5 > Bi_2O_5$ Acidic strength of oxides of nitrogen $N_2O < NO < N_2O_3 < N_2O_4 < N_2O_5$

Basic nature, bond angle, thermal stability and dipole moment of hydrides

 $NH_3 > PH_3 > AsH_3 > SbH_3 > BiH_3$

Stability of trihalides of nitrogen: NF₃ > NCl₃ > NBr₃

Lewis base strength: NF₃ < NCl₃ > NBr₃ < NI₃

Ease of hydrolysis of trichlorides $NCl_3 > PCl_3 > AsCl_3 > SbCl_3 > BiCl_3$

- Lewis acid strength of trihalides of P, As and Sb $PCl_3 > AsCl_3 > SbCl_3$
- Lewis acid strength among phosphorus trihalides $PF_{3} > PCl_{3} > PBr_{3} > PI_{3}$
- Nitrogen displays a great tendency to form $p\pi p\pi$ multiple bonds with itself as well as with carbon and oxygen.
- The basic strength of the hydrides

 $NH_{3} > PH_{3} > AsH_{3} > SbH_{3}$

The thermal stability of the hydrides decreases as the atomic size increases.



Melting and boiling point of hydrides $H_2O > H_2Te > H_2Se > H_2S$

Volatility of hydrides $H_2O < H_2Te < H_2Se < H_2S$

Reducing nature of hydrides $H_2S < H_2Se < H_2Te$

Covalent character of hydrides $H_2O < H_2S < H_2Se < H_2Te$

The acidic character of oxides (elements in the same oxidation

 $SO_2 > SeO_2 > TeO_2 > PoO_2$; $SO_3 > SeO_3 > TeO_3$

Acidic character of oxide of a particular element (e.g. S) $SO < SO_2 < SO_3$; $SO_2 > TeO_2 > SeO_2 > PoO_2$

BORON

FAMILY



HALOGEN FAMILY

- Bond energy of halogens: $Cl_2 > Br_2 > F_2 > I_2$
- Solubility of halogen in water:
- $F_2 > Cl_2 > Br_2 > I_2$ Oxidising power: $F_2 > Cl_2 > Br_2$
- Enthalpy of hydration of X^- ion : $F^ Cl^- > Br^- > I^-$
- Reactivity of halogens: F > Cl > Br > IIonic character of M - X bond in halides

M-F > M-Cl > M-Br > M-I

- Reducing character of X^- ion : $I^- > Br^- > Cl^- > F^-$
- Acidic strength of halogen acids: HI > HBr > HCl > HF
- Conjugate base strength of halogen acids

 $I^- < Br^- < Cl^- < F^-$

- Reducing property of hydrogen halides HF < HCl < HBr < HI
- Oxidising power of oxides of chlorine

 $Cl_2O > Cl_2O_2 > Cl_2O_6 > Cl_2O_7$

- Acidic character of oxyacids of chlorine HClO<HClO₂<HClO₃<HClO₄
- Oxidising power of oxyacids of chlorine HClO>HClO₂>HClO₃>HClO₄

NOBLE GASES

 $XeF_2 + PF_5 \longrightarrow [XeF]^+ [PF_6]$ $XeF_4 + SbF_5 \longrightarrow [XeF_3]^+ [SbF_6]$ $XeF_6 + H_2O \longrightarrow XeOF_4 + 2HF$ $XeF_6 + 2H_2O \longrightarrow XeO_2F_2 + 4HF$ $2XeF_4 + 3H_2O \longrightarrow$

$$Xe + XeO_3 + 4HF + F_2$$

 $XeF_6 + 3H_2O \longrightarrow XeO_3 + 6HF$ $2XeF_6 + SiO_2 \longrightarrow 2XeOF_4 + SiF_4$

 $2XeOF_4 + SiO_2 \longrightarrow 2XeO_2F_2 + SiF_4$

 $2XeO_2F_4 + SiO_2 \longrightarrow 2XeO_3 + SiF_4$

TRANSITION ELEMENTS (d-and f-BLOCK **ELEMENTS**

COORDINATION

COMPOUNDS

- The element with exceptional configuration are Cr²⁴[Ar 1 3d⁵4s¹, Cu²⁹[Ar 1 3d¹⁰4s¹ Mo⁴²[Kr] 4d⁵5s¹, Pd⁴⁶[Kr] 4d¹⁰5s⁰ Ag⁴⁷[Kr] 4d¹⁰5s¹, Pt⁷⁸[Xe] 4f¹⁴5d¹⁰6s⁰
- **Inner Transition Elements**
- **Electronic Configuration -** $[Xe]4f^{0-14}5d^{0-1}6s^2$
- (ii) Magnetic properties Magnetic moment is given by the formula $\mu = \sqrt{4S(S+1) + L(L+1)}$

where L=Orbital quantum number, S=Spin quantum number

- Coordination number is the number of the nearest atoms or groups in the coordination sphere.
- Ligand is a Lewis base donor of electrons that bonds to a central metal atom in a coordination compound.
- Paramagnetic substance is one that is attracted to the magnetic field, this results on account of unpaired electrons present in

the atom/molecule/ion.

- Effective atomic number EAN $= (Z - Oxidation number) + (2 \times Coordination number)$
- Factors affecting stability of complex (i) Greater the charge on the central metal ion, greater is the stability.
 - (ii) Greater the ability of the ligand to donate electron pair (basic strength) greater is the stability.
 - (iii) Formation of chelate rings increases the stability.
- Isomerism in coordination compounds:
 - Structural Isomerism (ii) Ionization Isomerism (iv) Linkage Isomerism (iii) Hydration Isomerism
 - (vii) Polymerisation Isomerism (viii) Valence Isomerism
 - (ix) Coordination Position Isomerism
 - Stereo Isomerism
 - (a) Geometrical
 - Square planar complexes of the type MA₂X₂; MABX₂; MABXY
 - Octahedral of the type: MA_4XY , MA_4X_2 , MA_3X_3 , $MA_2X_2Y_2$. $M(AA)_2X_2$ and M(ABCDEF).
 - (b) Optical isomerism

ORGANIC CHEMISTRY

GENERAL ORGANIC CHEMISTRY

- The order of decreasing electronegativity of hybrid orbitals is $sp > sp^2 > sp^3$.
- Conformational isomers are those isomers which arise due to rotation around a single bond.
- A meso compound is optically inactive, even though it has asymmetric centres (due to internal compensation of

rotation of plane polarised light)

- An equimolar mixture of enantiomers is called racemic mixture, which is optically inactive.
- Reaction intermediates and reagents: Homolyticfission → Free radicals

Heterolytic fission \rightarrow Ions (Carbonium ions, carbanions

etc.) Nucleophiles - Electron rich Two types: (i) Anions (ii) Neutral molecules with lone pair of electrons (Lewis bases)

Electrophiles: Electron deficient.

Two types: (i) Cations (ii) Neutral molecules with vacant orbitals (Lewis acids).

- Inductive effect is due to σ electron displacement along a chain and is permanent effect.
- +I (inductive effect) increases basicity, I effect increases acidity of compounds.
- Resonance is a phenomenon in which two or more structures can be written for the same compound but none of them actually exists.



- Pyrolytic cracking is a process in which alkane decomposes to a mixture of smaller hydrocarbons, when it is heated strongly, in the absence of oxygen.
- Ethane can exist in an infinite number of conformations. They are



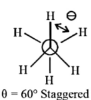






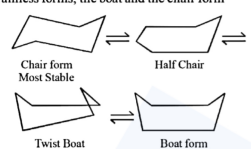








Conformations of Cyclohexane: It exists in two nonplanar, strainless forms, the boat and the chair form





ALKYNES

In dehydration and dehydrohalogenation the preferential order for removal of hydrogen is 3° > 2° > 1° (Saytzeff's rule).

(Least Stable)

The lower the ΔH_h (heat of hydrogenation) the more stable the alkene is

Alkenes undergo anti-Markonikov addition only with HBr in the presence of peroxides.

Alkynes add water molecule in presence of mercuric sulphate and dil. H₂SO₄ and form carbonyl compounds.

Terminal alkynes have acidic H-atoms, so they form metal alkynides with Na, ammonical cuprous chloride solution and

ammoniacal silver nitrate solution.

Alkynes are acidic because of H-atoms which are attached to sp 'C' atom which has more electronegativity and 's' character than sp² and sp³ 'C' atoms.



All o and p-directing groups are ring activating groups

(except - X)

They are: -OH, $-NH_2$, -X, -R, -OR, etc.

All m-directing groups are ring deactivating groups.

They are: -CHO, -COOH, $-NO_2$, -CN, $-NR_3$, etc



The order of reactivity is

(i)RI>RBr>RCI>RF

(i) Allyl halide > Alkyl halide > Vinyl halide

(iii) Alkyl halide > Aryl halide

S_N1 reaction: Mainly 3° alkyl halides undergo this reaction and form racemic mixture. S_N1 is favoured by polar solvent and low concentration of nucleophile.

S_N2 reaction: Mainly 1° alkyl halides undergo this substitution. Walden inversion takes place. S_N2 reaction is preferred by non-polar solvents and high concentration of nucleophile.

Reaction with metals:

(i)
$$R - X + Mg \xrightarrow{Dry \text{ ether}} R - Mg - X$$
Alkyl
halides
Grignard reagent

(ii) Wurtz reaction:

$$R \xrightarrow{L} X + 2 Na + X \vdash R \xrightarrow{Dry \text{ ether}} R - R + 2Na^{+}X^{-}$$
Alkane

ALCOHOLS

Alkenes are converted to alcohol in different ways as follows

Reagent dil H₂SO₄ Markovnikov
B₂H₆ and H₂O₂, OH⁻
Oxymercuration demercuration

Types of addition
Markovnikov
Anti-Markovnikov
- Markovnikov

Oxidation of

1° alcohol \longrightarrow aldehyde \longrightarrow carboxylic acid (with same no. (with same no. of of C atom) C atom)

2° alcohol \longrightarrow ketone \longrightarrow carboxylic acid (with same no. (with less no. of C atom)

3° alcohol → ketone → carboxylic acid (with less no. (with less no. of of C atom) C atom)

PHENOLS

ETHERS

CARBONYL

COMPOUNDS

Phenol ← CHCl₃/OH^Θ → Phenolic aldehyde (Reimer-Tieman reaction)

Phenol $\xrightarrow{CO_2}$ Phenolic carboxylic acid (Kolbe's reaction)

Acidity of phenols

(i) Increases by electron withdrawing substituents like

$$-NO_2$$
, $-CN$, $-CHO$, $-COOH$, $-X$, $-\stackrel{+}{N}R_3$

OR

(ii) decreases by electron releasing substituents like -R, -OH, $-NH_2$, $-NR_2$, -

 $\begin{array}{c}
2ROH \xrightarrow{Al_2O_3} R - O - R + H_2O \\
RONa + X - R' \longrightarrow ROR' + NaX \\
\text{(Williamson's synthesis)}
\end{array}$

Formation of alcohols using RMgX(a) Formaldehyde + RMgX

 $\xrightarrow{\text{Hydrolysis}} 1^{\circ} \text{ alcohol}$ (b) Aldehyde + RMgX $\xrightarrow{\text{Hydrolysis}}$ 2° alcohol

(other than HCHO)

(c) Ketone + RMgX $\xrightarrow{\text{Hydrolysis}}$ $\xrightarrow{\text{3}^{\circ}}$ alcohol



Cannizzaro reaction (Disproportionation)

Aldehyde
$$\xrightarrow{\text{Hot conc.}}$$
 Alcohol + Salt of acid

(no a H-atom)

Aldol condensation:

Carbonyl compound + dil. alkali $\longrightarrow \beta$ -hydroxy carbonyl (with α H-atom) compound

Benzoin condensation

The relative reactivities of different acid derivatives towards nucleophilic acyl substitution reaction follow the order:

CARBOXYLIC ACIDS

- The rate of esterfication decreases when alcohol, acid or both have branched substituents.
- Ortho effect: All ortho substituted benzoic acids (irrespective of type of substituent) are stronger than benzoic acid.

NITROGEN COMPOUNDS Order of basicity:

$$(R = -CH_3 \text{ or } -C_2H_5)$$

 $2^{\circ} > 1^{\circ} > 3^{\circ} > NH_3$

Hofmann degradation

Amides Br2/KOH 1° amine

The basicity of amines is

(i) decreased by electron withdrawing

groups (ii) increased by electron releasing groups

Reduction of nitrobenzene in different media gives different products

Medium

Acidic

Basic Neutral

Product

Aniline

Azoxy, Azo and finally hydrazobenzene Phenyl hydroxylamine

Carbohydrates are polyhydroxy aldehydes or ketones.

CARBOHYDRATES,
AMINO ACIDS AND
Containing three to nine carbon atoms.

AMINO ACIDS AND POLYMERS

Characteristic reactions :

Homologous series

(i) Alkanes

(ii) Alkenes and alkynes

(iii) Arenes

(iv) Alkyl halides

(v) Aldehyde and ketones

Type of reactions

Substitution

Mostly free radical

Wostry nee radical

Electrophillic addition Electrophillic substitution

Nucleophillic substitution

Nucleophillic addition

Tests to differentiate :

1°, 2° and 3° alcohols

1°, 2° and 3° amines

1°, 2° and 3° nitro compounds

Aryl halides and alkyl halides

Aldehydes and ketones

Aromatic aldehydes and Aliphatic aldehydes

(i) Lucas test

(ii) Victor meyer's test

Hinsberg test

Test with HNO₂ and KOH

Test with AgNO₃ solution Tollen's test/Fehling's test

Fehling's test

IMPORTANT REAGENT

- Dil. H₂SO₄ [or Conc. H₂SO₄ + H₂O] Use → Hydrating agent (+HOH)
- Alc. KOH or NaNH₂(Use \rightarrow -HX)

 $CH_3CH_2Cl \xrightarrow{alc.KOH} CH_2=CH_2$

- Lucas Reagent ZnCl₂ + Conc. HCl
 Use → For distinction between 1°, 2°
 & 3° alc.
- Tilden Reagent NOCl (Nitrosyl chloride)

$$C_2H_5NH_2 \xrightarrow{NOCl} C_2H_5Cl$$

Alkaline KMnO₄(Strong oxidant)

Toluene → Benzoic acid

Bayer's Reagent : 1% alkaline KMnO₄(Weak oxidant)

Use:
$$\rightarrow$$
 For test of $>$ C = C $<$ or $-$ C = C $-$

$$CH_2=CH_2+H_2O+[O] \xrightarrow{BR} CH_2OH-CH_2OH$$

- Acidic $K_2Cr_2O_7$ (Strong oxidant): $RCH_2OH \xrightarrow{[O]} RCHO$
- SnCl₂/HCl or Sn/HCl used for reduction of nitrobenzene in acidic medium.

$$C_6H_5NO_2 \xrightarrow{SnCl_2/HCl} C_6H_5NH_2$$

- Lindlar's Catalyst = Pd/CaCO₃
 - + in small quantity (CH₃COO)₂Pb

$$2 - \text{butye} + \text{H}_2 \xrightarrow{\text{"}} \text{Cis-2-butene}$$

(main product)

Ziegler –Natta Catalyst (C₂H₅)₃Al + TiCl₄

Use →In Addition polymerisation

IDENTIFICATION TESTS:

- (i) Unsaturated compound (Bayer's reagent)
 Decolourising the reagent
- (ii) Alcohols (Ceric ammonium nitrate solution) Red colouration
- (iii) Phenols (Neutral FeCl₃ solution) Violet/deep blue colouration
- (iv) Aldehydes and ketones (2, 4-D.N.P.)
 Orange precipitate
- (v) Acids (NaHCO₃ solution) Brisk effervescence (CO₂ is evolved)
- (vi) 1° amine (CHCl₃ + KOH) Foul smell (isocyanide)
- (vii) 2° amine (NaNO₂+HCl) Yellow oily liquid (Nitrosoamine)