Chapter-1. Some Basics Concepts Chemistry

Some Basic concepts of chemistry

Basic S.I Units

S.N.	Physical Quantity	Symbol	Unit	Symbol Of Unit
1	Length	l	meter	m
2	Mass	m	kilogram	kg
3	Time	T	second	S
4	Electric current	I	ampere	A
5	Temperature	T	kelvin	К
6	Amount of Substance	N	mole	mol
7	Luminous Intensity	I	candela	cd

Unit	Kilogram	Kelvin	Second	Siemens
Symbol	kg	K	S	S

Derived units: The units of all other physical quantities are derived out of these of the basic physical quantities.

The units thus obtained are called the derived units.

S.	Physical	Mathematical	Name of unit	In terms of
N.	Quantity	Relation		basic SI Units
1	Volume	Length cube	Cubic meter	m ³
2	Force	Mass x	Newton(N)	kg m s ⁻²
		Acceleration		
3	Pressure	Force/Area	Pascal (Pa)	kg m ⁻¹ s ⁻²
4	Energy or	Force x Distance	joule(J)	kg m ² s ⁻²
	Work			
5	Heat	Quantity of heat	Joule/kelvsin	JK ⁻¹ mol ⁻¹ or
	Capacity	per degree rise		$m^2kgs^{-2}k^{-1}$
		in temp.		

6	Concentrati	Mol./vol.	Mol/L	mol dm ⁻³ (M)
	on			
7	E.C.E.	Eq.wt. /Faraday	Kg/C	kgC ⁻¹
8	Area	Length Square	Square meter	m ²
9	Velocity	Distance /Time	meter/sec	ms ⁻¹
10	Acceleration	Velocity	meter per	ms ⁻²
		change/Time	square	
			second	
11	Power	Energy/Time	Watt(W)	kg m ² s ⁻³ or
				J.s ⁻¹
12	Density	Mass/Volume	Kilogram per	kg m ⁻³
			cubic meter	
13	Frequency	Cycles/Second	hertz(Hz)	S-1
14	Electric	$Current \times Time$	coulomb(C)	A.s
	Charge			
15	Potential	Force/ Charge	volt(V) or	kg m ²
	difference	(force per unit	$(J.A^{-1}s^{-1})$	s-3.A-1 or
		charges)		(charge)

Some other units in SI system

Physical	Unit	Relationship with basic
Quantity		SI unit
	kilometer (km)	1 km = 1000 meter-
	meter (m)	100 cm = 1 meter
	centimeter (cm)	1000 mm = 1 meter
Longth	millimeter (mm)	$10^6 \mu m = 1 meter$
Length	micrometer or micron	$10^9 \text{ nm} = 1 \text{ meter}$
	(μm)	$10^{12} \text{pm} = 1 \text{meter}$
	nanometer (nm)	
	pico meter (pm)	

	kilogram (kg)	1 kg = 1000 g
	gram (kg)	-
Mass	milligram (mg)	1000 mg = 1 g
	microgram (μg)	$10^6 \mu g = 1 g$
	Nanogram (ng)	$10^9 \text{ng} = 1 \text{g}$
	milliter (mL)	1000 mL = 1 liter
Volume	cubic centimeter	$1000 \text{ cm}^3 = 1 \text{ liter}$
voiume	(cm ³ ,mL)	$10^6 \mu l = 1 liter$
	microliter(μL)	

Determination of Atomic Mass:

- 1 amu=(1/12) mass of an atom of 12 C = 1.66×10^{-24} g= 1.66×10^{-27} kg
- Average Atomic Mass = $\frac{(m_1 \times x_1) + (m_2 \times x_2)}{x_1 + x_2} = \frac{\% \text{ of element}}{\text{Atomic mass of element}}$ Where, m₁& m₂ are atomic masses of isotopes and x₁& x₂ are their fractions present.
- One Gram Atom = 6.023×10^{23} atoms
- Masses-Volume relation:

•
$$\frac{\text{Mass}}{\text{Molar mass}} = \frac{\text{V(dm}^3)}{22.4}$$

• n =
$$\frac{\text{Mass}}{\text{Molar mass}} = \frac{\text{No.of particles}}{6.022 \times 10^{23}}$$

- Determination of Molar Mass:
- One Gram Molecule (one mole) = 6.023 X 10²³ particles
- One mole = 6.023 X 10²³ atoms, molecules, electrons etc.,=22.4dm³ at S.T.P.(for gases)
- Molecular Mass = 2 X Vapour density
- Vapour density = $\frac{\text{Mass of certain volume of a gas or vapour}}{\text{Mass of equal volume of a } H_2 \text{ at same temp.}}$
- STP = Standard temperature & pressure, T = 273.15 K, and P = 1 atm.
- Eq. mass = $\frac{\text{Atomic Mass}}{Valency}$
- Eq. mass of the same element may be different in different compounds if its valency changes:

1) Hydrogen Displacement Method

Eq. mass of metal =
$$\frac{\text{Mass of Metal}}{\text{Mass of H}_2 \text{ displaced}} = \frac{\text{Mass of metal}}{V_0 \text{ (dm}^3) \text{ X 9 X } 10^{-2}}$$

2) Oxide Formation method:

Eq. mass of metal =
$$\frac{8 \text{ X Mass of metal taken}}{\text{mass of O }_{2}\text{combined with metal}}$$

$$= \frac{5.6 \text{ X Mass of metal}}{\text{V of O}_2 \text{ combined with metal}}$$

3) Chloride Formation Method:

Eq. mass of metal =
$$\frac{35.5 \text{ X mass of metal taken}}{\text{mass of Cl}_2 \text{ combined with metal}}$$

4) Metal Displacement Method:

$$\frac{\text{Mass of metal A}}{\text{mass of metal B}} = \frac{\text{Eq.Mass of metal A}}{\text{Eq.mass of metal B}}$$

5) Electrolysis Method:

$$\frac{\text{Mass of A discharged}}{\text{mass of B discharged}} = \frac{\text{Eq. Mass of A}}{\text{Eq. mass of B}}$$

CHAPTER 2. STATES OF MATTER (GASEOUS & LIQUID STATES)

Gas laws:

1. Boyle's law:

- a) PV = constant
- b) $\frac{p}{d}$ = constant (at constant T & n), Where d-density of gas

2. Charle's law:

a)
$$V/T = constant (at constant P \& n),$$

b)
$$V_t = V_o + \frac{V_0 t}{273.15} = v_0 \left(1 + \frac{t}{273.15}\right)$$

3. Gay Lussac law:

P/T = constant (at constant P & n)

4. Avogadro's law:
$$V \propto \text{Number of molecules (n), } \\ (at constant P & T) or $\frac{V}{n} = \text{constant}$$$

5. Kinetic energy: K.E./Mol.=
$$\frac{3}{2}$$
RT

Average K.E. or K.E./Molecule
$$=\frac{3}{2}\frac{RT}{N} = \frac{3}{2}kT$$

(K=Boltzman constant)

6. Van der Waal's equation:

$$\begin{split} &\left[P+\frac{a}{V^2}\right][V-b]=RT \quad \text{ (For 1 mole)} \\ &\left[P+\frac{n^2a}{V^2}\right][V-nb]=nRT \quad \text{(For n mole)} \end{split}$$

a-'a' is vander Waal's constant for attraction between gas molecules.

b-'b'is vander Waal's constant for volume occupied by gas particles.

b=4.N.v (v is volume of one molecule in rest) and $a = P'V^2$

CHAPTER 3. REDOX REACTIONS

- **3.1. Oxidation:** Oxidation or de-electronation is a process which liberates electronsor increase in the oxidation number of oxidized species.
- **3.2 Reduction**: Reduction or electronation is a process which gains electrons or decreases in the oxidation number of reduced species.

Steps involved in balancing of redox equation by oxidation number method:

- **Step 1**. Find out oxidation number of each element and identify the elementWhich undergoes change in oxidation number.
- **Step 2.** Find out increase and decrease in oxidation number per atom. Multiply the increase or decrease in oxidation number with number of atoms undergoing the change.
- **Step 3.** Multiply the formula with suitable integer to equalise the increase & decrease in oxidation number.
- **Step 4.** Balance atoms other than 0 & H.
- **Step 5.** To balance O atoms, add H₂O molecules to the side containing less O atoms.
- **Step 6.** In case of ionic reaction:
 - a) If the reaction is acidic medium, the proper number of H⁺ ions are added to the side containing less number of H atoms.
 - b) If the reaction is basic medium, then proper number of H_2O molecules are added to the side containing less number of H atoms, while an equal o OH^- ions are added to the other side.

Mechanical Engineering

Scope:

Mechanical Engineers are solution providers for development of processes and products that range from tiny component designs to mega manufacturing plants. Likewise, automobile engineers design, facilitate and develop designs of vehicles like cars and motorbikes. Coming from very diverse disciplines of engineering, mechanical/automobile engineers can work right from inception stage, R&D of their product to its final commissioning. They take care of the mechanical, electronic, electric, software and safety elements during the manufacturing.

Professional Opportunities

Design

Mechanical Engineers draft technical drawings, manually or with the aid of computers.

Production

Mechanical Engineers supervise the manufacturing of mechanical components and machines.

Analysis & Testing

Mechanical Engineers analyze and test different types of machines and their parts to ensure that they function flawlessly.

Installation

Mechanical Engineers install machines and mechanical parts at their client's location.

Maintenance

Mechanical Engineers primarily ensure that machinery is working as per specifications.

Maintenance of Aircraft or missiles

Mechanical Engineers with specialization in Aircraft technology, test new aerodynamic and aviation systems and maintain them.

Electronics & Telecommunication Engineering

Scope:

Telecommunications engineering is an engineering discipline centered on electrical and computer engineering which seeks to support and enhance telecommunication systems. The work ranges from basic circuit design to strategic mass developments. A telecommunication engineer is responsible for designing and overseeing the installation of telecommunications equipment and facilities, such as complex electronic switching systems, and other plain old telephone service facilities, optical fiber cabling, IP networks, and microwave transmission systems. Telecommunication engineering also overlaps with broadcast engineering.

Professional Opportunities

Telecom equipment engineer

A telecom equipment engineer is an electronics engineer that designs equipment such as routers, switches, multiplexers, and other specialized computer/electronics equipment designed to be used in the telecommunication network infrastructure.

Network engineer

A network engineer is a computer engineer who is in charge of designing, deploying and maintaining computer networks. In addition, they oversee network operations from a network operations center, designs backbone infrastructure, or supervises interconnections in a data center.

Central-office engineer

A central-office engineer is responsible for designing and overseeing the implementation of telecommunications equipment in a central office. A CO engineer is responsible for integrating new technology into the existing network, assigning the equipment's location in the wire center, and providing power, clocking (for digital equipment), and alarm monitoring facilities for the new equipment. The CO engineer is also responsible for providing more power, clocking, and alarm monitoring facilities if there are currently not enough available to support the new equipment being installed.

- Cancel any duplication that is observed on both the sides of Step7 equation.
- Step 8 Check the equation is balanced for both, the atoms and the charges.

CHAPTER 4. CHEMICAL EQUILIBRIUM

Ostwald's dilution law, pH & pOH

- $\alpha = \frac{\text{Number of moles ionised}}{\text{Total number of moles}}$
- Percentage dissociation = $(\% \alpha) = \alpha \times 100$ b)
- c) $\alpha = \frac{\% \text{ dissociation}}{100}$
- d) For weak electrolyte $\alpha = \sqrt{\frac{K}{c}}$ \therefore $(K = \alpha^2 \times C)$ e) For WA, $\alpha = \sqrt{\frac{K_a}{c}}$ $(K_a = \alpha^2 \times C)$, \because $(\alpha < 0.005)$ f) For WB, $\alpha = \sqrt{\frac{K_b}{c}}$ $(K_b = \alpha^2 \times C)$, \because $(\alpha < 0.005)$

Dissociation constant of an acid HA, $K_a = \frac{[H^+][A^-]}{[HA]}$

Dissociation constant of a base BOH, $K_b = \frac{[B^+][OH^-]}{IBOHI}$

- For two solutions of an electrolyte $\frac{\alpha_i^2}{\alpha_{r_i}^2} = \frac{C_{ii}}{C_r}$ g)
- h) $[H^+]$ =Basicity x α x C (for WA)
- [OH-]=Acidity x α x C (for WB) i)
- For S.A./S.B. $\rightarrow \alpha = 1$ i)
- k) pH=-log[H+]
- pOH=-log [OH-] 1)
- m) $[H^+]$ =antilog (-pH)
- n) $[OH^{-}]$ = antilog (-pOH)
- o) pH + pOH = 14(at 298K)
- p) $[H^+][OH^-]=K_w=1X10^{-14}(at 298K)$
- q) Alternately $[H^+] = \sqrt{K_a \times C}$
- r) $[OH^-] = \sqrt{K_b \times C}$ s) $C = \frac{\text{No.of Moles}}{\text{dm}^3}$

t)
$$n = \frac{Mass}{Molar Mass}$$

To calculate pH of buffer: (Henderson Hasselbalch equation)

a) for acidic buffer:

$$pH = pK_a + log \frac{[Salt]}{[Acid]}$$
 where, $(pK_a = -log k_a)$

- b) For basic buffer: $pOH = pK_b + log \frac{|Salt|}{|Base|} (pK_b = -log k_b)$
- 1) Salt of strong acid & weak base: $pH = \frac{1}{2}(pK_w pK_b \log C)$
- 2) Salt of weak base & strong acid: $pH = \frac{1}{2}(pK_w + pK_b + \log C)$
- 3) Salt of weak acid & weak base: $pH = \frac{1}{2}(pK_w + pK_a pK_b)$
- 4) Salt of strong acid & strong base: pH=7

Where,
$$pK_a = -logK_a$$
,
 $pK_b = -logK_b$
 $pK_w = -logK_w$
 $pK_a + pK_b = pK_w = 14$

CHAPTER 5.SOLID STATE

- a) To calculate formula of compound:
 - 1. In a cube three are 8 corners, 6 face centers, 12 edge centers & one body centers.
 - 2. Contribution towards unit cell:
 - 3. Corner= $\frac{1}{8}$, face centre= $\frac{1}{2}$, edge centre= $\frac{1}{4}$, body centre=1 Formula of the compound is same as the ratio of atoms in the unit cell.
- b) For cubic crystals of ionic compounds:

$$\rho = \frac{Z \times M}{a^3 \times N_A} \text{ gcm}^{-3}$$

Where Z=Number of formula unit present in one unit cell. M=Formula mass

c) Relation between distance and edge length:

SCC	BCC	FCC
d=a	$d = \frac{\sqrt{3}a}{2}$	$d = \frac{a}{\sqrt{2}}$

d=distance between nearest neighbors & 'a' edge length of cubic unit cell

CHAPTER 6. SOLUTION AND COLLIGATIVE PROPERTIES

1. Relation between molarity & molality:

a) Molarity =
$$\frac{\text{molality} \times \text{Mass of solvent in kg}}{\text{volume of solution in dm}^3}$$

- b)
- i) if 'd' of solvent is given then M=m X d (for dilute solution)
- ii) If'd' of solution is given then

$$\label{eq:Molarity} \text{Molarity} = \frac{\text{Molarity} \times 1000}{(\text{d} \times 1000) - (\text{Molarity} \times \text{Molar Mass})}$$

2. Relation Between Molarity(m) and Mole fraction of solute(X_2) in water is

$$X_2 X_2 = \frac{m}{55.56+m}$$
 (Where moles of water in 1 dm³ is 55056)

Henry's law

1.
$$S = K \times P$$
, $K \rightarrow Henry's constant (atm-1)$

$$2. m = K \times P, \ (S \propto M)$$

$$3. x_{gas} = K \times P, (M \propto x_{gas})$$

4. If $K \to Henry's$ constant in atm then $p = K \times x_{gas}$

Osmotic pressure (II):

Expression for osmotic pressure: osmotic pressure=h d g Where, h=height of column, d=density of solution in the column, g=acceleration due to $gravity=9.8ms^2$

Van't Hoff solution equation:

Determination of molar mass of solute from osmotic pressure:

$$PV = nRT \quad \text{or} \quad P = \frac{W_2RT}{M_2V} \quad \text{or} \quad M_2 = \frac{W_2RT}{\pi V}$$

Units of R, Π & V

To calculate van't Hoff factor (i):

$$\begin{split} \text{I)} \quad & \text{i} = \frac{\text{CP}_{(obs)}}{\text{CP}_{(theor)}} = \frac{\text{CP}_{(expt)}}{\text{CP}_{(normal)}} \\ \text{II)} \quad & \text{i} = \frac{\pi_{(obs)}}{\pi_{(theor)}} = \frac{\Delta P_{(obs)}}{\Delta P_{(theor)}} = \frac{\Delta T_{b(obs)}}{\Delta T_{b(theor)}} = \frac{\Delta T_{f(obs)}}{\Delta T_{f(theor)}} \\ \text{III)} \quad & \text{i} = \frac{\text{theoretical molar mass of solute}}{\text{Observed molar mass of solute}} \end{split}$$

To calculate degree van't Hoff factor (i):

Hoff factor for the substances that undergo dissociation or association:

- I) $\pi = iCRT$
- II) $\Delta T_b = iK_b m$
- III) $\Delta T_f = iK_f m$

CHAPTER 7.CHEMICAL THERMODYNAMICS AND ENERGETICS

A] First law of thermodynamics:

$$\Delta U_{(J)} = q_{(J)} + W_{(J)} = q + P\Delta V$$

Note:

- a) In this statement, q is the heat absorbed and W is the work done on the System. $\Delta U = q + W$
- b) In this statement, q is the heat absorbed and W is the work done by the System. $\Delta U = q W$

Sign convention for q,W, Δ U

- q=+ve, heat absorbed by system from surrounding & q=-ve, heat evolved by system into surrounding.
- 2. $\Delta U = +ve (U_2 > U_1)$, I.E. of system increases & $\Delta U = -ve (U_2 < U_1)$, I.E. of system decreases.
- 3. W=+ve $(V_2>V_1)$, work done on the system by the surrounding $\&W=-ve(V_2<V_1)$, work done by the system on the surrounding

B] Change in enthalpy: $\Delta H = \Delta U + P\Delta V$

- Sign convention of ΔH :
- 1. If ΔH =-ve,heat is evolved, it is called exothermic reaction.
- 2. If $\Delta H = +ve$, heat is absorbed, it is called endothermic reaction.
- 3. $\Delta H = \Sigma \Delta H_{(products)} \Sigma \Delta H_{(reactants)}$
- 4. Enthalpy change in reaction=(Sum of enthalpies of formation or (heat of combustion) of products)-(sum of enthalpies of formation of reactants)

Note:

- a) Enthalpy of element in their physical state and at standard condition is zero.
- b) Enthalpy of diamond is not considered as an elementary state of C hence

Its enthalpy is not zero.

- 1. $\Delta H = \Delta U + \Delta nRT$
- 2. $q_p = q_v + \Delta nRT$

Where,

 ΔH =Heat of reaction at a constant pressure in J.

 ΔU =Heat of reaction at a constant volume in J.

 Δn =Difference between number of moles of gaseous products and number ofmoles of gaseous reactants.

 $R=gas\ constant\ (8.314JK^{-1}\ mol^{-1})$

T=Absolute temperature (K)

 $q_p {=} \, \Delta H {=} Heat$ absorbed at a constant pressure

 $q_v \!\!= \Delta U \!\!=\! Heat$ absorbed at a constant volume

P=Atmospheric volume (Nm-2)

V=Volume (m³)

Hint: generally ΔH & ΔU are given in kJ, convert them in J as R is given $(1kJ=10^3J)$

- 1. The value of Δn may be zearo, positive or negative.
- 2. For gases:
 - a) If $\Delta n=0$ then $\Delta H = \Delta U$
 - b) If $\Delta n < 0$ then $\Delta H < \Delta U$

- c) If $\Delta n > 0$ then $\Delta H > \Delta U$
- 3. For solids and liquids, $\Delta n=0$

D] Hess's law:

- 1. $\Delta H = \Delta H_1 + \Delta H_2 + \Delta H_3$
- 2. To solve the examples on Hess's law:
- a) First step: write the data in the form of thermo chemical equation.
- b) Second step: write the required equation and balance it.
- c) Third step: obtain the required equation by adding, subtracting, multiplying or dividing the data equation.
- d) Fourth step: carry out the same operation in the sequential manner on the enthalpy values of the equations and obtain the total enthalpy change (ΔH) inthe required equation.

3. Note:

- a) Substracting is conveniently done by adding the reversed equation and changing the sign of ΔH while reversing it. $\Delta H_{direct \, reaction} = \Delta H_{reverse \, reaction}$
- b) In thermo chemical equation the symbol 'aq' when used alone indicates a very large quantity of water. While multiplying and equation containing 'aq' by any number, the term 'aq' need not be multiplied.

H] Change in entropy:

a)
$$\Delta S = \frac{q_{rev}}{T}$$

b)
$$\Delta S_{\rm sys} = \frac{q_{\rm sys}}{T}$$
 and $\Delta S_{\rm surr} = \frac{-q_{\rm surr}}{T}$

c)
$$\Delta S_{universe} = \Delta S_{total} = \Delta S_{sys} + \Delta S_{surr}$$

Standard molar energy (ΔS^0):

$$\Delta S^0 = \Sigma \Delta S^0_{\text{(products)}} - \Sigma \Delta S^0_{\text{(reactants)}}$$
 or $\Delta S^0 = (cS^0C + dS^0D) - (aS^0A + bS^0B)$

 ΔG^0 and equilibrium constant (K):

$$\Delta G^0 = -2.303 RT log_{10} K_C$$
 or $\Delta G^0 = -2.303 RT \ log_{10} K_p$

Where
$$K_C = \frac{(Products)}{(Reactants)} = \frac{[C]^c[D]^d}{[A]^a[B]^b}$$
 or $K_p = \frac{(P_c)^c(P_d)^d}{(P_A)^a(P_B)^b}$

C=concentrations & P=Partial pressure

CHAPTER 8 ELECTROCHEMISTRY

a) To calculate Λ_0 of weak acid

1. Λ (weak acid)= Λ_0 (salt of WA and SB)+ Λ_0 (SA)- Λ_0 (salt of SA and SB) OR

Ex:
$$\Lambda_0(CH_3COOK) + \Lambda_0(HBr) - \Lambda_0(HBr)$$

b) Faraday's 1st law:

- a) $W(kg)=Z(kg/C) \times Q(C)$
- b) $W(kg)=Z(kg/C) \times i(amp) \times t(sec)$
- c) $E=F \times Z$
- d) $E(kg) = 96500(C) \times Z(kg/C)$

Avogadro's law:

1mole=Molar mass=6.023x10²³molecules=22.4dm³ of a gas at N.T.P.

Form Avogadro's law and faraday we can calculate the volume of different gases liberated by 1 faraday.

1mole gas (mass) = Molar volume = (mass/eq.mass) number of equivalents= number of Faraday

Relation between quantity and amount of substance deposited:

Quantity of electricity to discharge one mole of ions=n x F (Where n=charge on the ion)

CHAPTER 8. CHEMICAL KINETICS

A) To calculate rate constant for first order reaction:

$$k = \frac{2.303}{t} log_{10} \frac{[A]_0}{[A]_t}$$

Where, $[A]_0$ =Initial concentration of reaction A,

 $[A]_t$ =concentration of reaction A left after time t.

B) To calculate rate constant (k) according to Arrhenius equation:

- a) $k=Ae^{-Ea/RT}$
- b) $\log_{10} = \log_{10} A \frac{E_a}{2.303RT}$

Where, $A \rightarrow$ Frequency factor or pre-exponential factor

 $E_a\!\to\! Activation\ energy,\ R\text{-}Gas\ constant,\ T\text{-}Absolute\ temperature}$

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c)
$$E_a \Rightarrow \frac{E_a}{RT} \Rightarrow \frac{E_a}{RT} \Rightarrow e^{-\frac{E_a}{RT}} \Rightarrow k \Rightarrow rate$$

decreases decreases increases increases increases dE_a =Threshold energy - Average energy

C) Arrtenius equation and temperature variation or to calculate Ea:

a)
$$\log_{10} \frac{k_2}{k_1} = \frac{E_a}{2.303R} \left[\frac{T_2 - T_1}{T_1 \times T_2} \right]$$

 $k_1 \rightarrow \text{rate constant at temperature } T_{1,}$

 $k_2 \rightarrow rate constant at temperature T_2$

b) Rate either doubled or tripled for every degree 10^{0} rise in the temperature as $\frac{k_{T}+10}{k_{T}}=2$ to 3

Relation between [A] and rate and half life:

 a) Time for the completion of 'x' fraction of the reaction may be evaluated by the equation.

$$t_{x} = \frac{2.303}{k_{1}} \log \frac{1}{1-x}$$

b) Time taken for the completion of 100% reaction:

$$a - x = 0$$
, $t_x = \frac{2.303}{k} = \log \frac{a}{o} = \infty$

c) Amount left after n half lives:

$$[A] = [A]_0 \, 2^{\cdot n} \text{ or } [A]_t = [A]_0 \, 2^{\cdot n} \quad \text{where } n = \frac{\text{total time}}{t_0 \, 5}$$

- e) From the above equation we can arrive at:
 - a) For 75% of the reaction; $t_{0.75} = 2x t_{0.5}$
 - b) For 87.5% of the reaction; $t_{0.875} = 2x t_{0.5}$
 - c) For 99.9% of the reaction; $t_{0.999} = 10x t_{0.5}$

CHAPTER 9.CO-ORDINATION COMPOUNDS

- I) Structural Isomerism:
- a) Ionisation Isomerism:

Ion-Ion exchange. Ex.: $[Co(NH_3)_5SO_4]Br$ and $[Co(NH_3)_5Br]SO_4$

- **b) Linkage Isomerism:**Linkage of atoms of lingands. Ex.:[Co(NH₃)₅NO₂|Cl₂ and [Co(NH₃)₅.ONO]Cl₂
- c) Co-ordination Isomerism:

 Exchange of ligands.Ex.: [Cu(NH₃)₄][PtCl₄] and[Pt(NH₃)₄][CuCl₄]
- d) Hydrated Isomerism:

Number of H_2O molecules inside and outside of coordination sphere.

 $Ex.:[Cr(H_2O)_6]Cl_3, [Cr(H_2O)_5Cl]Cl_2.H_2O$ and $[Cr(H_2O)_4Cl_2]Cl_2.H_2O$

II) Stereo-Isomerism:

Geometrical Isomerism:

CN=4

- a) Tetrahedral geometry: do not show
- b)Square planner geometry: (Cis-same and trans opposite)

CHAPTER 10.HALOARENES

1) Friedel-Craft Alkylation:

2) Friedel-Craft Acetylation:

3) Wurtz – Fitting Reaction:

CHAPTER 11 - PHENOL

Preparation:

- 1) From chlorobenzene:
 - a) Dow's process(1928):

b) Rasching process (Industrial Method 1943):

2) From Aniline or aminobenzene (Diazotisation):

3) From Cumene (Isopropyl Benzene):

4) From benzene sulphonic acid:

ii)
$$SO_3 Na \qquad OH$$

$$2 \qquad + H_2SO_4 \rightarrow 2 \qquad + Na_2SO_4$$
dil

5) Kolbe's Reaction: (Formation of salicylic acid)

Salicylic acid (2 - Hydroxybenzoic Acid)

6) Reimer - Tiemann reaction: (formation of salicylaldehyde):

CHAPTER 12 - ETHERS

1) Preparation of ether

By intermolecular dehydration of alcohols (Continuous etherification)

2) By methylation of alcohol: (from diazomethane and alcohol)

$$CH_{3} - OH + CH_{2}N_{2}$$
 $\xrightarrow{HBF_{4}}$ $CH_{3} - O - CH_{3} + N_{2}$
 $C_{2}H_{5} - OH + CH_{2}N_{2}$ $\xrightarrow{HBF_{4}}$ $C_{2}H_{5} - O - CH_{3} + N_{2}$

3) By Williamson's synthesis: (from sodium alkoxide and alkyl halide)

$$C_2H_5$$
 - ONa + I - C_2H_5 \longrightarrow C_2H_5 - O - C_2H_5 + NaI
$$C_2H_5$$
 - ONa + I - CH_3 \longrightarrow C_2H_5 - O - CH_3 + NaI

4) By Williamson's synthesis:

ONa + I -
$$CH_3$$
 O - CH_3 + NaI

- A) Electrophilic substitution reaction
 - 1) Friedel Craft alkylation:

2) Friedel – Craft Acetylation:

- B) Preparation of aromatic aldehydes only:
 - 1) By formylation of benzene (Gattermann Koch reaction):

C) Preparation of aromatic ketones only:

Friedel - Craft alkylation:

From GR and phenyl cyanide

$$C \equiv N \qquad MgBr \qquad \qquad C \qquad \qquad + 2H_2O \qquad \qquad dil. \ HCl \qquad \qquad O \qquad \qquad \\ C \qquad \qquad C \qquad \qquad + NH_3 + Mg(Br)OH \qquad \qquad C \qquad \qquad C$$

CHAPTER 13. COMPOUND CONTAINING NITROGEN

A) Chemical Properties

1) Hydrolysis:

(Distinguishing test between 1°,2°, 3° nitroalkane)

$$R- CH_2-NO_2+H_2O \xrightarrow{HCI/\triangle} R- COOH + NH_2OH$$

$$2R_2CH-NO_2 \xrightarrow{HCI/\triangle} 2R- CO-R+N_2O+H_2O$$

$$R_3C-NO_2 \xrightarrow{HCI/\triangle} No reaction$$

2) Halogenation (Alkaline bromination):

$$R - CH_2NO_2 + 2Br_2 + 2NaOH \longrightarrow R - C(Br)_2 - NO_2 + 2NaBr + 2H_2O$$

$$R_2 - CH - NO_2 + Br_2 + NaOH \longrightarrow R_2C(Br) - NO_2 + NaBr + H_2O$$

$$R_3C - NO_2 + Br_2 + NaOH \longrightarrow No \ reaction$$

$$CH_3 - NO_2 - 3CI_2 + 3NaOH \longrightarrow CCI_3 - NO_2 + 3NaCI + 3H_2O$$

B) Amines

Preperation

1) By alkylation of pthalimide (Gabriel)

Ring substitution (ESR):

a) Halogenation:

b) Nitration:

c) Sulphonation:

d) Friedel - Craft reaction: Do not given by benzoic acid.

Action of cold (HNO₂ +dil HCL):

a)
$$2\text{CH}_3\text{NH}_2 + 2\text{HNO}_2 \xrightarrow{\text{Cold}} \text{CH}_3 - 0 - \text{CH}_3 + 2\text{N}_2 + 3\text{H}_2\text{O}$$
 and $\text{CH}_3\text{NH}_2 + 2\text{HNO}_2 \xrightarrow{\text{Cold}} \text{CH}_3 - 0 - \text{N} = 0 + \text{N}_2 + 2\text{H}_2\text{O}$

b)
$$C_2H_5NH_2 + HO - NO \xrightarrow{Cold} C_2H_5 - OH + H_2O + N_2 \uparrow$$

c)
$$C_6H_5NH_2 + NaNO_2 + 2HCl \xrightarrow{Cold} C_6H_5N_2^+Cl^- + NaCl + 2H_2O$$

d)
$$(C_2H_5)_3NH + HO - NO \xrightarrow{Cold} (C_2H_5)_2N - N = O + H_2O$$

e)
$$(C_2H_5)_3N + HNO_2 \xrightarrow{Cold} [C_2H_5NH]NO_2$$

$$(CH_3)_2N$$
 + HO- N = O Cold $(CH_3)_2N$ N = O + H₂O

Haffmann's Carbyl amine test(Isocyanide formation):b

$$\begin{aligned} R - NH_2 + CHCL_3 + 3KOH &\stackrel{alc}{\rightarrow} R - NC + 3KCl + 3H_2O \\ C_6H_5NH_2 + CHCL_3 + 3KOH &\stackrel{alc}{\rightarrow} C_6H_5NC + 3KCl + 3H_2O \end{aligned}$$

A) Aniline:

Electrophilic substitution reaction:

1) Bromiantion

B) Benzene Diazonium Salt

Preparation:

1) By diazotization of aniline:

Chemical Properties

I) Replacement of diazonium group:

Replacement by -H(formation of arenes/ Deamination):

$$Ar - N_2^+X^- + H_3PO_2 \xrightarrow{CuCl} Ar - H + N2 \uparrow + H_3PO_3 + HX$$

 $Ar - N_2^+X^- + CH_3CH_2OH \xrightarrow{CuCl} Ar - H + N2 \uparrow + CH_3CHO + HX$

CHAPTER 14 - BIOMOLECULES

Presence of carbonyl group:

Action of HCN:

CHO

CHOH

(CHOH₄ + HCN
$$\longrightarrow$$
 (CHOH)₄

CH₂OH

CH₂OH

Vitamins:

Some vitamins, their sources and diseases due to deficiencies

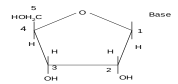
S.N.	Vitamins	Sources	Diseases due to deficiencies
	Vita-A	Milk, fish liver,	Night blindness,
1	(Retinol/	tomatoes, carrot,	retardation of growth,
1	Axerophtohl)	potatoes	dryness of skin and hair
	Vita-B	Rice, wheat, meat,	Beriberi
2	(Thiamine)	green vegetable	Delibert
	Vit-B ₂ or G	Egg yolk, fishes,	Inflammation of tongue,
	Riboflavin	yeast, liver	dryness of lips and
3	Tubonavin	y cast, 11 ver	mouth, cheilosis
			(retarding the growth and
			digesion)
	Vit-B ₃	Yeast, liver,	Dermatities, graying of
	(Panthothermic	tomatoes, egg,	hair, retard body mental
4	acid)	meat	growth, reproductive
	,		disability.
	Vit-B ₅	Barley, maize,	Pigmentation of skin
	(Nicotinamide)	wheat, rice	(pellagra), retard body
5			and mental growth,
			degeneration of spinal
			cord, mental confusion.
	Vit-B ₆	Milk, liver, maize,	Convulsion, loss of
	(Pyriodoxine or	wheat, fish, yeast	weight, mental change,
6	Pyridoxal		derangement of enzymes
0	Pyridoxamine)		(which control
			carbohydrates
			metabolism)
7	Vit-B ₁₂	Egg, liver of pig,	Degradation of spinal
	(Cyanocobalamn)	sheep	cord, anaemia
	Vit-C	Orange, grapes,	Scurvy (bleeding,
8	(Ascorbic acid)	lemon, tomatoes,	spongy, swollen) gums
		onion, cabbage	

	Vit-D	Butter, liver, egg,	Rickets, Osteomalacia
9	(Ergocalciferol)	fish oil, milk,	
9		meat, in skin cell	
		in sun	
	Vit-E	Light rice, liver of	Weakness of muscles,
10	(Tocopherol)	cattle, seed oils,	abnormal growth and of
10		wheat deposition	tissue, decrease
			reproductive power
	Vit-H	Yeast, egg, fruits,	Skin lesions, loss of
11	(Biotin)	wheat	apatite, hair fall,
			paralysis
	Vi-K	Green leaf of	Increase boold clotting
12	(Phylloquinine)	spinach, fish	time (hemorrhage), poor
		meat, cauliflower	coagulation of blood
13	Vit-P	Orange, grapes	Haemorrhagia, decrease
13			in capillary resistance.

Nuclear Acid

Nucleosides= Base + Suga

a) Ribonucleosides



CHAPTER 15 - POLYMERS

Chart of Polymers

Name	Monomer	St. of Polymer	Type/Applicati
			on
		a)LDPE Branched	Addition
Dolvathana	CH CH	473K/1000 atm:	Reaction
Polyethene	$CH_2=CH_2$	$[CH_2-CH_2]_n$	Homopolymer
		b)HDPE Linear 473/6-7	

	1		
		atm	Insulator for
		$T_iCl_4+C_2H_5)_3Al$	cables
		ÇH₃	Addition
Polypropene	CH ₃ -CH=CH ₂	{CH₂ - CH} _n	Reaction
	555, 555, 555,	_ ··	homopolymer
	0,		Addition
Teflon	$CF_2 = CF_2 \xrightarrow{O_2}$	$[CF_2-CF_2]_n$	Reaction
			homopolymer
(Pan)			Addition
Polyacryloni	$CH_2 = CH-CN$	CN	Reaction
trile	$CH_2 - CH - CN$	{CH₂ - CH} _n	homopolymer
unc			
(Orlon)			Addition
	Peroxide		Reaction
		ÇI	homopolymer
Polyvinyl	CH ₂ =CHCl	[СН ₂ - СН] _п	
Chloride		- "	Pipes
			Addition
Naammana	çı	СІ {CH ₂ - C = CH - СӉ ₁	Reaction
Neoprene	СН ₂ = С- СН - СӉ	{CH ₂ - C = CH - CH ₂ } _n	homopolymer
	O 2 →		Making
			container
	2-chloro-1,		Converyour
	3-butadiene		belt.
	CH ₂ = CH - CH = CH	{ CH₂ - CH = CH CH₂ - CH₃ - CH-1N	Addition
Buna-	•	1 2 2 3 3 3 3 3 3 3 4 4 4	Reaction
S(SBR)	1,3-Butadiene K ₂ S ₂ O ₈	 	Co-polymer
or (GNR)	₊	¢ ₆ H ₅ ·cH=CHૃ	Making
or (ONK)			Bubble gum

CHAPTER 16 - CHEMICALS IN MEDICINES:

A) Analgesic (Pain killer):

Types of analgesic:

a) Narcotic analgesic: These drugs produce depression on the central nervous system and relive pain instantly.

E.g. (1) Morphine (M.F.: C₁₇H₁₉NO₃)

(2) Heroine (M.F.: C₂₁H₂₃ NO₅)

B) Non – Narcotic analgesic: These drugs when consumed do not produce any significant depression of the central nervous system called non – narcotic analgesic.

E.g. Paracetomol, Aspirin, Ibuprofen, Methyl salicylate, etc.

Aspirin:

Acetyl salicylic acid is commercially called aspirin.

Molecular formula: C₉H₈O₄, Common Name: Acetyl salicylic acid

IUPAC name: 2 – Acetoxybenzoic acid

Preparation: from salicylic acid:

C) Antimicrobials:

The drugs used to kill or prevent the growth of the diseases forming micro-organisms are called antimicrobials. These are Antibiotics Antiseptic, Disinfectants, etc.

- Antibiotics: The term 'antibiotics' was proposed by Vuillemin in 1889. (Anti-against, biotic- life). Chemical substances which are derived from one type of micro-organisms such as bacteria and are used to inhibit the growth and destroy disease forming micro-organisms, which causes infections, are called antibiotics.
- 2) Antiseptic: The chemical substances that can kill bacteria or prevent the growth of micro-organisms in living tissues are called antiseptics.

Common antiseptics are:

Chloroxylenol (M.F:C₈H₉OCl)

1) Terineol(M.F:C₁₀H₁₈O)

- Disinfectants: Chemicals which are antimicrobial agents that are applied to non-living objects to destroy micro-organism which may cause infections are called disinfectants.
 - Ex: (1) Chlorine (Cl2)water, (2) 1%Phenol (C_6H_5OH) (3) Sulphur dioxide (SO_2)
- 2) Antifertility drugs: The drugs which are used to control the population by family planning

Ex: (1) Novestrol (Ethynylestradiol (M.F:C₂₀H₂₄O₂)

2) Norethindrone (M.F: C₂₀H₂₆O₂)

- 5) Antacids and anthistamines:
 - a) Antacids: The chemicals which are used to neutralize the
 excess acid (HCI) formedin gastric juice and raise the pH to an
 appropriate level in stomach is called antacids.
 Sweetening agents: The substances which are used for
 sweetening the food andfood material are called sweeteners.
- 1) Saccharine (M.F: C₇H₅SNO₃)