Onwards (MR-20)	MALLA REDDY ENGINEERING COLLEGE (Autonomous)	I B.Tech		
Code: A0B17	<b>Engineering Chemistry</b>	L	T	P
Credits: 4	(Common for CSE, IT, ECE, EEE, CE, ME and Min.E)		1	-

#### **Course objectives:**

The purpose of this course is to emphasize the relevance of fundamentals of chemical sciences in the field of engineering and to provide basic knowledge on atomic- molecular orbital's, electrochemistry, batteries, corrosion and the role of water as an engineering material in domestic-industrial use. They will also impart the knowledge of stereochemistry, understanding the chemical reaction path way mechanisms and synthesis of drugs. Listing out various types of fuels and understanding the concept of calorific value and combustion.

#### **Module I: Water and its treatment**

[10 Periods]

Introduction to water, hardness of water, causes of hardness, expression of hardness, units and types of hardness-Numerical Problems. Alkalinity of water, specifications of potable water (BIS); Estimation of temporary & permanent hardness of water by EDTA method. Boiler troubles - Scale & Sludge, Priming and foaming, caustic embrittlement and boiler corrosion; Treatment of boiler feed water - Internal treatment (colloidal, phosphate, carbonate and calgon conditioning). External treatment - Lime Soda process (cold & hot) and ion exchange process, Numerical Problems. Disinfection of water by chlorination and ozonization. Desalination by Reverse osmosis and its significance.

#### **Module II: Molecular structure and Theories of Bonding:**

[10 Periods]

[7 Periods]

Introduction to Molecular orbital Theory. Linear Combination of Atomic Orbital's (LCAO), significance of bonding and anti-bonding molecular orbital, Conditions for the formation of molecular orbital's. Molecular orbital energy level diagrams of diatomic molecules -,  $N_2$ ,  $O_2$  and  $F_2$ . Introduction to coordination compounds-ligand-coordination number (CN) - spectrochemical series. Salient features of crystal field theory, Crystal field splitting of transition metal complexes in octahedral (  $[CoF_6]^{3-}$  and  $[Co(CN)_6]^{3-}$ ) and tetrahedral ( $[NiCl_4]^{2-}$  and  $[Ni(CO)_4]$ ) fields - magnetic properties of complexes. Band structure of solids and effect of doping on conductance.

#### **Module III: Electrochemistry and Corrosion**

## A. Electrochemistry:

Introduction to Electrochemistry-Conductance (Specific and Equivalent) and units. Types of cells-electrolytic & electrochemical cells (Galvanic Cells)-Electrode potential- cell potential (EMF). Electrochemical series and its applications, Nernst equation its applications and numerical problems. Reference electrodes - Calomel Electrode and Glass electrode-determination of pH using glass electrode. Batteries: Primary (dry cells) and secondary (Lead-Acid cell, Ni-Cd cell) - applications of batteries. Fuel cells: Hydrogen - Oxygen fuel cell and its applications.

B. Corrosion: [7 Periods]

Causes and effects of corrosion: Theories of corrosion - Chemical & Electrochemical corrosion, Pilling-Bedworth rule, Types of corrosion: Galvanic and Water-line corrosion. Factors affecting rate of corrosion-Nature of metal and Nature of Environment, Corrosion control methods - Cathodic protection (Sacrificial anodic and impressed current cathodic methods). Surface coatings: Methods of metallic coatings - hot dipping (Galvanization), Electroplating (Copper) and Electroless plating (Nickel).

# Module IV: Stereochemistry, Reaction mechanism & synthesis of drug molecules and NMR spectroscopy: [12 Periods]

Introduction to Isomers - classification of isomers - structural (chain, positional & functional) and stereoisomerism-geometrical (cis-trans & E-Z system) - characteristics of geometrical isomerism, optical isomerism (chirality - optical activity, specific rotation, enantiomers and diastereomers) of tartaric acid and lactic acid. Conformational isomerism of n-Butane. Introduction to bond cleavage (homo & hetero cleavage) - reaction intermediates and their stability. Types of organic reactions - Mechanism of substitution (SN $^1$  & SN $^2$ ) and (E $_1$ &E $_2$ ) reactions with suitable example. Ring opening (Beckmann rearrangement), oxidation and reduction (Cannizaro reaction), cyclization (Components of Diels-Alder reaction-Mechanism of Diels-Alder reaction with suitable example) reactions. Synthesis of Paracetamol, Aspirin and their applications.

Introduction to Spectroscopy, Basic concepts of nuclear magnetic resonance spectroscopy, chemical shift and spin-spin splitting.

### **UNIT-V** Fuels and Combustion

[08 Periods]

**Fuels:** Classification- solid fuels: coal – analysis of coal – proximate and ultimate analysis and their significance. Liquid fuels – petroleum and its refining, cracking – types – moving bed catalytic cracking. Knocking – octane and cetane rating, synthetic petrol - Fischer-Tropsch's process; Gaseous fuels – composition and uses of natural gas, LPG and CNG. **Combustion:** Definition, Calorific value of fuel – HCV, LCV; Calculation of air quantity required for combustion of a fuel. Determination of calorific value by Junkers gas calorimeter-Numerical problems on combustion.

#### **Text Books:**

- 1. P.C.Jain and Monica Jain, "A Text Book of Engineering Chemistry", DhanpatRai Publications, New Delhi, 16th Edition 2014.
- 2. S.S. Dara and S.S. Umare, "A Text Book of Engineering Chemistry", S Chand Publications, New Delhi, 12th Edition 2010.
- 3. A.Jaya Shree, "Text book of Engineering Chemistry", Wiley, New Delhi, 2018.

#### **Reference Books:**

- 1. B.Rama Devi, Ch.VenkataRamana Reddy and PrasanthaRath, "Text Book of Engineering chemistry", Cengage Learning India Pvt.Ltd,2016.
- 2. M.G. Fontana and N. D. Greene, "Corrosion Engineering", McGraw Hill Publications, New York, 3<sup>rd</sup> Edition, 1996.
- 3. K. P. C. Volhardt and N. E. Schore, "Organic Chemistry: Structure and Function", 5<sup>th</sup> Edition, 2006.

#### e-Resources:

#### a) Concerned Website links:

- 1) https://books.google.co.in/books?isbn=0070669325 (Engineering chemistry by Sivasankar).
- 2) https://www.youtube.com/watch?v=yQUD2vzfgh8 (Hot dipping Galvanization).
- 3)https://archive.org/stream/VollhardtOrganicChemistryStructureFunction6th/Vollhardt\_Organic\_C hemistry\_Structure\_Function\_6th\_djvu.txt.

## b) Concerned Journals/Magazines links:

- 1) http://americanhistory.si.edu/fuelcells/sources.htm (Fuel Cell Information Sources)
- 2) https://www.abctlc.com/downloads/courses/WaterChemistry.pdf (Water Chemistry)

#### c) NPTEL Videos:

- 1) nptel.ac.in/courses/113108051/ (corrosion & electrochemistry web course)
- 2) https://www.youtube.com/watch?v=V7-8EOfZKeE (Stereochemistry)

#### **Course Outcomes:**

After completion of the course students will be able to:

- 1. Understand water treatment, specifically hardness of water and purification of water by various methods.
- 2. Analyze microscopic chemistry in terms of atomic and molecular orbital's splitting and band theory related to conductivity.
- 3. Acquire knowledge on electrochemical cells, fuel cells, batteries and their applications.
- 4. Acquire basic knowledge on the concepts of stereochemistry, reaction mechanisms and interpretation of NMR in organic molecules.
- 5. Examine various fuels based on the calorific value and identify a better fuel source of less pollution.

#### **Co-Po Mapping:**

СО	PO	PO	PO	PO	PO	PO	PO	PO	PO	PO	PO	PO
CO	1	1 2 3	3	3 4 5		6	7	8	9	10	11	12
CO1	3	3	1	2			3					
CO2	2	1	1	2								
CO3	3	3	2	2			3					
CO4	3	2	1	1								
CO5	3	3	3	3			3					

# Module-I Water and its treatment

## **Introduction:**

Water is a natural gift on the earth is essential for human animal and plants. Human being used for all domestic purposes like drinking, cooking bathing, cleaning and washing. Water is not only essential for the survival of life, but it is also used for the operation in a large number of industries as coolant, solvent, for steam generation, for air conditioning, fire-fighting etc. Without food human can survive for a number of days, but water is such an essential that without it one cannot survive.

It has a wide role in industries also. Probably, its most important use as an engineering material is in the "steam generation".

Chemically, water has two atoms of hydrogen and one atom of oxygen having the molecular formula  $H_2O$ . In several chemical reactions, water is formed along with the main product, e.g. acid reacts with base to give salt and water. Alcohols and organic acids react to give ester and  $H_2O$ 

Water molecule is a bent triatomic molecule with the bond angle  $104.5^{0}$  the oxygen atom present in the water molecule has  $sp^{3}$  – hybridization, state having two bond pairs and two lone pairs.

Water is found in three physical states, liquid (water), solid (ice) and gas (vapour). The freezing point of water is  $0^{0}$ C and the boiling point is  $100^{0}$ C. Several special behaviors found in water due to hydrogen bond present in it. In nature, water is present 3//4 of the earth surface.

The process of removing of all types of impurities from water and making fit for domestic or industrial purposes is called **Water Technology** or **Water Treatment**.

The main sources of water are,

- 1. Rain water
- 2. Surface water (Rivers and lakes)
- 3. Underground water (wells and springs)
- 4. Lake water
- 5. Sea water

**Rainwater**: It is the purest form of natural water. But unfortunately it dissolves the toxic gases like  $CO_2$ ,  $SO_2$ ,  $NO_2$  etc. and other solids, which pollute the atmosphere.

**Sea water:** It is the most impure form of water containing about 3.5% dissolved salts of which about 2.6% is sodium chloride. Other salts present include sulphates, bicarbonates, bromides of sodium, potassium, magnesium etc. Other impurities are carried to sea through rivers.

**River water:** The sources of river water are the springs and the rainwater. River water while flowing through the land collects lots of organic matters from falling trees and nearby habitats and also other soluble and suspended matters from the lands, soils etc. The dissolved matters

include the salts like sulphates, bicarbonates and bromides of sodium, potassium and magnesium.

**Lake water:** It is much purer than river water, dissolved impurities are less but contains lots of organic matter.

**Underground water**: The rainwater and other surface water percolate down through the soil and rocks and get filtered and finally collected on rocky surface or again come out as spring. Though it contains less suspended matter but the dissolved mineral content is quite high and is of high organic purity.

## **Impurities in Water**

Impurities in water are of three types

- 1. Physical impurities:
  - A. Suspended impurities:
  - B. Colloidal impurities
- 2. Chemical impurities
  - A. Dissolved salts
- 3. Biological impurities
  - A. Microorganisms

<u>HARDNESS OF WATER:</u> Most of the inorganic salts are soluble in water. The  $Ca^{+2}$  and  $Mg^{+2}$  salts are dissolved in the water causes hardness of water. When water passes through the rocks (dolomite  $MgCO_3$  and limestone  $CaCO_3$ ) and flows on the ground. The  $Ca^{+2}$  and  $Mg^{+2}$  salts are present in the rocks are dissolved in water in the presence of atmospheric  $CO_2$  and it make hard.

$$MgCO_3+CO_2+H_2O \longrightarrow Mg(HCO_3)_2$$
  
 $CaCO_3+CO_2+H_2O \longrightarrow Ca(HCO_3)_2$ 

#### **Types of water:**

a) Hard Water b) Soft Water

**Hard Water:** Water, which does not produce lather with soap solution, but produces white precipitate is called **Hard water.** This is due to presence of dissolved calcium and magnesium salts. A soap is a sodium salt or potassium salt of higher fatty acid such as **stearic acid** ( $C_{17}H_{35}COOH$ ). When soap is contact with hard water, sodium stearate will react with dissolved  $Ca^{+2}$  and  $Mg^{+2}$  salts and produce insoluble Ca-stearate (or) Mg-stearate precipitates.

**Soft water:** Water, which produce lather with soap solution is called **Soft water**. This is due to the absence of Ca & Mg salts.

$$C_{17}H_{35}COONa + H_2O \longrightarrow C_{17}H_{35}COOH+NaOH$$
  
Soap (water soluble)  
(Water soluble)  
 $C_{17}H_{35}COONa + C_{17}H_{35}COOH \longrightarrow Foam Formation$   
(Sodium stearate ) (Stearicacid)

## **Types of Hardness:**

Hardness of water is of two types

- (a) Temporary hardness (carbonate hardness)
- (b) Permanent hardness (non carbonate hardness)

## (a) Temporary hardness (carbonate hardness)

**Temporary Hardness** is due to the presence of magnesium and calcium bicarbonates [Ca(HCO<sub>3</sub>)<sub>2</sub> and Mg(HCO<sub>3</sub>)<sub>2</sub>]. It is also called **carbonate hardness** (CH). It can be removed by **Boiling**. During boiling, the soluble Mg(HCO<sub>3</sub>)<sub>2</sub> is converted into insoluble Mg(OH)<sub>2</sub> and Ca(HCO<sub>3</sub>)<sub>2</sub> is changed to insoluble CaCO<sub>3</sub>. These insoluble precipitates can be removed by filtration. Filtrate thus obtained will be soft water.

Ca(HCO<sub>3</sub>)<sub>2</sub>

Boiling

CaCO<sub>3</sub> 
$$\downarrow$$
H<sub>2</sub>O + CO<sub>2</sub>

Calcium bicarbonate

Calcium carbonate (Insoluble)

Mg(HCO<sub>3</sub>)<sub>2</sub>

Mg(HCO<sub>3</sub>)<sub>2</sub>

Mg(OH)<sub>2</sub>  $\downarrow$  + 2CO<sub>2</sub>

Magnesium bicarbonate

Mg(OH)<sub>2</sub>  $\downarrow$  + 2CO<sub>2</sub>

## Permanent (or) Non - carbonate hardness:

**Permanent Hardness** is due to the presence of soluble salts of magnesium and calcium in the form of chlorides, nitrates and sulphates in water (CaCl<sub>2</sub>, CaSO<sub>4</sub>, MgCl<sub>2</sub>, Ca(NO<sub>3</sub>)<sub>2</sub>, Mg(NO<sub>3</sub>)<sub>2</sub>and MgSO<sub>4</sub>). Permanent hardness is not removed by boiling. It is also called **non-carbonate hardness** (NCH). Only chemical treatment method can remove this hardness.

Total Hardness: Temporary hardness and permanent hardness constitute the total hardness which is also expressed as the sum of the concentration of calcium and magnesium ions

Total Hardness = Temporary Hardness + Permanent Hardness

Total Hardness =  $[Ca^{2+}] + [Mg^{2+}]$ 

**Disadvantages of Hard Water:** The following are the disadvantages of hard water:

In domestic use: (a) Hard water is not useful for various domestic purposes, viz. washing, bathing and drinking. The hardness producing ions convert soluble soaps to insoluble precipitates. This results in wastage of soaps in washing and bathing.

- (b) Due to the presence of dissolved hardness producing salts, the boiling point of water is elevated. Hence, more fuel and time are required for cooking.
- (c) Hard water causes bad effect on our health. Calcium form oxalate crystals in urinary tract.

In industrial use: (a) The dissolved Ca<sup>+2</sup>, Mg<sup>+2</sup> and Iron salts in hard water will react with dyes to form undesirable precipitates which gives improper shade.

- (b) In sugar refineries must be free from sulphates, nitrates, alkali, carbonates and bacteria, these impurities sugar may not crystalline well.
- (c) If hard water is fed directly in to the boilers for steam generation, it results in various boiler troubles viz. Scales, sludges, priming, foaming and caustic embrittlement.
- (d) Hard water is not suitable for laboratory analysis, because hardness produces ions interferes (disturbs) in various reactions.
- (e) Any water sample with hardness less than 150 ppm is good and portable while beyond 350 ppm is not suitable for consumption.

**Degree of Hardness:** The total hardness of water is caused by **8** different salts of Ca<sup>2+</sup> and Mg<sup>2+</sup>, Ca(HCO<sub>3</sub>)<sub>2</sub>, Mg(HCO<sub>3</sub>)<sub>2</sub>, CaCl<sub>2</sub>, CaSO<sub>4</sub>, MgCl<sub>2</sub>, Ca(NO<sub>3</sub>)<sub>2</sub>, Mg(NO<sub>3</sub>)<sub>2</sub> and MgSO<sub>4</sub>. The hardness of water is expressed in terms of CaCO<sub>3</sub> equivalents. CaCO<sub>3</sub> is selected for the expression of degree of hardness is due to (a) Its molecular weight is 100, which is easy for calculation (b) It is most in soluble in water.

Hardness causing salt (H.C.S)

In terms of CaCO<sub>3</sub> equivalents = Weight of the hardness causing salt x Molecular weight of CaCO<sub>3</sub> Molecular weight of the hardness causing salt

OR

Hardness causing salt (H.C.S)

In terms of CaCO<sub>3</sub> equivalents = Weight of the hardness causing salt x Eqivalent weight of CaCO<sub>3</sub> Equivalent weight of the hardness causing salt

Substance	Molecular weight	Valency	Equivalent weight
Calcium bicarbonate	162	2	81
Magnesium bicarbonate	146	2	73
Calcium sulphate	136	2	68
Calcium chloride	111	2	55.5
Magnesium sulphate	120	2	60
Magnesium chloride	95	2	47.5
Calcium carbonate	100	2	50
Calcium nitrate	164	2	82
Magnesium nitrate	148	2	74

Problem 1: Calculate  $CaCO_3$  equivalent of hardness causing salts present in 1000 liters of sample of water containing 16.2 mg/l of  $Ca(HCO_3)_2$ , 11.1 mg/l of  $CaCl_2$ , 60mg/l of  $MgSO_4$  and 19 mg/l of  $MgCl_2$ .

***************************************						
salt	Quantity	M.W	Equivalent to	Equivalent hardness causing salt in		
	of the salt		CaCO <sub>3</sub> (mg/l)	1000 L of water		
Ca(HCO <sub>3</sub> ) <sub>2</sub>	16.2	162	<u>16.2X100</u> =10	10X1000=10,000 mg		
			162			
CaCl <sub>2</sub>	11.1	111	<u>11.1X100</u> =10	10X1000=10,000 mg		
			111			
MgSO <sub>4</sub>	60	120	<u>60X100</u> =50	50X1000=50,000 mg		
G			120			
MgCl <sub>2</sub>	19	95	<u>19X100</u> =20	20X1000=20,000 mg		
<i>S</i> -			95			

**UNITS OF HARDNESS:** The following are the common units used in hardness measurements.

[1] Parts per million (ppm): It is defined as the number of parts of  $CaCO_3$  equivalent hardness causing salts present in million parts ( $10^6$ ) of water.

1 ppm = 1 part of  $CaCO_3$  equivalent hardness in  $10^6$  parts of water.

[2] **Milligram per litre:** It is defined as the number of milligrams of CaCO<sub>3</sub> equivalent hardness causing salts present in one litre of water.

 $1 \text{ mg/}l = 1 \text{ mg of CaCO}_3$  equivalent hardness in one litre of water

Since weight of 1 litre of water = 
$$1 \text{ kg} = 1000 \text{ g}$$

$$= 1000 \times 1000 = 10^6 \text{ mg}$$

1 mg/l = 1 mg of CaCO<sub>3</sub> equivalent hardness causing salts per 10<sup>6</sup> mg of water = 1 part of CaCO<sub>3</sub> per 10<sup>6</sup> parts of water = 1 ppm

Thus, mathematically both units are equal.

[3] Degree Clarke (°Cl): I t is the number of parts of CaCO<sub>3</sub> equivalent hardness per 70,000 parts of water.

 $1^{0}$  clarke ( $1^{0}$ Cl) = 1 grain of CaCO<sub>3</sub> eq hardness per gallon of water or  $1^{0}$ Cl = 1 part of CaCO<sub>3</sub> equivalent hardness per 70,000 parts of water.

[4] Degree French (°Fr): It is the number of parts of CaCO<sub>3</sub> per 10<sup>5</sup> parts of hard water thus,

 $1^{0}$ Fr = 1 part of equivalent CaCO<sub>3</sub> per  $10^{5}$  parts of hard water.

## [5] Equivalent per Million : EPM [ mill equivalent per litre(meq/l)]:

One Milligram equivalent of CaCO<sub>3</sub> per litre of hard water is called equivalent per million (or) Mill equivalent per litre is the number of milli equivalents of hardness present per litre Thus,

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1Meq/L = 1 Meq of CaCO<sub>3</sub> per Litre of water

= 10<sup>-3</sup> × 50g of CaCO<sub>3</sub> eq per litre

= 50 mg of CaCO<sub>3</sub> eq per litre

= 50 mg / l of CaCO<sub>3</sub> eq

1mg = 50 ppm

(or) 1 epm = 50 ppm of CaCO<sub>3</sub>
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The hardness of water can converted in to all the 4 units by making use of the following interconversion.

#### Relation between various units of hardness

**PROBLEM 1:** A sample of hard water contains 120 mg/l of hardness. Express hardness of water in ppm, degree of French and degree of clark?

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Hardness of water sample= 120mg/l
1 ppm = 1 mg/l = 0.1° Fr = 0.07° Cl
Hardness of water in degree of clark (°Cl)= 120X0.07=8.4 °Cl
Hardness of water in ppm=120 ppm
Hardness of water in degree of french (°Fr)= 120X0.1=12 °Fr
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#### **PROBLEM:2**

A sample on water on analysis has been found to contain the following:

$$Ca(HCO_3)_2 = 10.5 \; ppm. \; \; Mg \; (HCO_3)_2 = 12.5 \; ppm \\ CaSO_4 = 7.5 \; ppm \; \; \; CaCl_2 = 8.2 \; ppm \; \; \; MgSO_4 = 2.6 \; ppm.$$

Calculate the temporary and permanent hardness in degree French.

Sol. 
$$Ca(HCO_3)_2 = 10.5 \text{ ppm} = \frac{10.5 \times 100}{162} = 6.481 \text{ ppm}$$
 $Mg(HCO_3)_2 = 12.5 \text{ ppm} = \frac{12.5 \times 100}{146} = 8.562 \text{ ppm}$ 
 $CaSO_4 = 7.5 \text{ ppm} = \frac{7.5 \times 100}{136} = 5.515 \text{ ppm}$ 
 $CaCl_2 = 8.2 \text{ ppm} = \frac{8.2 \times 100}{111} = 7.387 \text{ ppm}$ 
 $MgSO_4 = 2.6 \text{ ppm} = \frac{2.6 \times 100}{120} = 2.167 \text{ ppm}$ 
 $\therefore$  Temporary hardness =  $(6.481 + 8.562) = 15.043 \text{ ppm}$ 
 $= 15.043 \times 0.1^{\circ} \text{ Fr} = 1.504^{\circ} \text{ Fr}$ 

Permanent hardness =  $(5.515 + 7.387 + 2.167) \text{ ppm}$ 
 $= 15.069 \text{ ppm}$ 
 $= 1.5069^{\circ} \text{ Fr}$ .

**PROBLEM 3:** A sample of hard water contains the following dissolved salts per litre. Ca(HCO<sub>3</sub>)<sub>2</sub>=16.4mg, Mg(HCO<sub>3</sub>)<sub>2</sub>=14.6mg, CaCl<sub>2</sub>=111mg, MgSO<sub>4</sub>=12mg, CO<sub>2</sub>=44mg, CaSO<sub>4</sub>=13.6mg. Calculate temporary and permanent hardness of water in ppm and degree of clark?

salt	Quantity	M.W	Equivalent to CaCO <sub>3</sub> (mg/l)
	of the salt		
Ca(HCO <sub>3</sub> ) <sub>2</sub>	16.4	162	<u>16.4X100</u> =10.12
			162
Mg(HCO <sub>3</sub> ) <sub>2</sub>	14.6	146	<u>14.6X100</u> =10
			146
CaCl <sub>2</sub>	111	111	<u>111X100</u> =100
			111
MgSO <sub>4</sub>	12	120	12X100 = 10
			120
CO <sub>2</sub>	44	44	44X100 = 100
			44
CaSO <sub>4</sub>	13.6	136	<u>13.6X100</u> =10
			136

$$\label{eq:camporary} \begin{array}{l} \text{Temporary hardness= $Ca(HCO_3)_2$+ $Mg(HCO_3)_2$} \\ &= 10.12 + 10 = 20.123 \, mg/l \\ &= 20.123 X 0.07 = 1.408 \ ^{\circ}\text{C1} \\ \text{Permanent hardness= $CaCl_2$+ $MgSO_4$+ $CaSO_4$} \\ &= 100 + 10 + 10 = 120 \ mg/l \\ &= 120 X 0.07 = 8.4 \ ^{\circ}\text{C1} \end{array}$$

**PROBLEM 4:** A sample of hard water contains the following dissolved salts per unit.  $CaCl_2=111mg$ ,  $CaSO_4=1.36$  mg,  $Ca(HCO_3)_2=16.2$  mg,  $Mg(HCO_3)_2=14.6$  mg, silica=40 mg, turbidity=10 mg. Calculate the total , permanent and temporary hardness of water in ppm, degree of clark and degree of French?

salt	Quantity	M.W	Equivalent to CaCO <sub>3</sub> (mg/l)
	of the salt		
Ca(HCO <sub>3</sub> ) <sub>2</sub>	16.2	162	<u>16.2X100</u> =10
			162
Mg(HCO <sub>3</sub> ) <sub>2</sub>	14.6	146	<u>14.6X100</u> =10
			146
CaCl <sub>2</sub>	111	111	<u>111X100</u> =100
			111
CaSO <sub>4</sub>	1.36	136	<u>1.36X100</u> =1
			136

Temporary hardness= 
$$Ca(HCO_3)_2 + Mg(HCO_3)_2$$
  
=  $10+10=20$  mg/l, 20 ppm  
=  $20X0.07=1.4$  °Cl  
= $20X0.1=2$  °Fr

Total hardness=Temporary hardness + Permanent hardness =20+101=121 mg/l, 121 ppm =121X0.07=8.47°Cl =121X0.1=12.1 °Fr

**Problem 4.** A sample of water has been found to contain the following salts:  $Ca(HCO_3)_2 = 10.5 \ ppm; \ Mg(HCO_3)_2 = 12.5 \ ppm; \ CaCl_2 = 8.2 \ ppm; \ MgSO_4 = 2.6 \ ppm; \ CaSO_4 = 7.5 \ ppm.$ 

Calculate (i) temporary and permanent hardness and (ii) the vol. of  $\frac{M}{100}$  EDTA required for titration of the 100 ml of the sample to determine the total hardness of the sample.

Sol. (i) Ca(HCO<sub>3</sub>)<sub>2</sub> = 10.5 ppm = 
$$10.5 \times \frac{100}{162}$$
 = 6.481 ppm CaCO<sub>3</sub>

$$Mg(HCO_3) = 12.5 ppm = 12.5 \times \frac{100}{146} = 8.562 ppm CaCO_3$$

$$CaCl2 = 8.2 ppm = 8.2 \times \frac{100}{111} = 7.387 ppm CaCO3$$

$$MgSO4 = 2.6 ppm = 2.6 \times \frac{100}{120} = 2.167 ppm CaCO3$$

$$CaSO4 = 7.5 ppm = 7.5 \times \frac{100}{136} = 5.515 ppm CaCO3$$
∴ Temporary hardness = (6.481 + 8.562) ppm
$$= 15.043 ppm$$
Permanent hardness = (7.387 + 2.167 + 5.15) ppm = 15.069 ppm
∴ Total hardness = (15.043 + 15.069) = 30.112 ppm
(ii) The volume of sample taken = 100 ml.
∴ Volume of M/100 EDTA required =  $\frac{30.112 \times 100}{1000} = 3.11 ml$ .

#### **BOILER TROUBLES**

A boiler is a closed vessel, in which water under pressure is transformed in to steam by heat to generate power. The water should be free from impurities and softened before being used in boilers. When hard water is fed into boiler the following problems arise,

- (1) Priming and Foaming
- (2) Caustic embrittlement
- (3) Boiler corrosion
- (4) Scale and Sludge

#### (1) Priming and Foaming

**Priming:** It may defined as the violent or rapid boiling of water in the boiler which results in carrying out of water droplets pass with steam in in the form of a spray. This process of wet steam formation is called **priming**. Priming in boilers is due to

- (1) The presence of suspended and dissolved salts
- (2) High level of water in boilers
- (3) Improper design of boiler
- (4) High velocities of steam in the boiler

Priming can be controlled by: (a) Maintain low water levels in boilers

- (b) Filtering water before feed into boilers
- (c) Avoid high-velocities of the steam
- (d) Proper evaporation and using adequate heating surfaces

**Foaming:** Formation of stable, persistent foam (or) bubbles at the surface of water in the boiler is called foaming. The oil and alkali react with water to form soap, as a result the surface tension of water decreases. Foaming is due to the presence of oil drops, grease and suspended solids.

Foaming can be removed by adding antifoaming agents like polyamide and castor oil and removing oil from boiler water by adding compounds like NaAlO<sub>2</sub>.

#### (2) Caustic embrittlement

This is a type of boiler corrosion, caused by using highly alkaline water. The source of alkali in the boiler is due to presence of residual  $Na_2CO_3$  in water by softening process by lime soda process. This undergo hydrolysis at high temperature and pressure to give NaOH.

$$Na_2CO_3+H_2O \longrightarrow 2 NaOH+CO_2$$

"The term caustic embrittlement used for the cracks in boiler water due to the increased concentration of caustic alkali". Accumulation of NaOH in the walls of the boiler parts becomes brittle, known as caustic embrittlement. Caustic embrittlement can be prevented by:

(a) Using Na<sub>3</sub>PO<sub>4</sub> as softening reagent instead of Na<sub>2</sub>CO<sub>3</sub> in external treatment of boiler water. (b) Adding tannins, lignin to boiler water, which blocks the cracks in the boiler walls. (c) Neutralizing alkali with very small quantity of acid.



#### 3. Boiler corrosion

The decay of boiler material by chemical (or) electrochemical attack its environment is called **boiler corrosion**. Main reasons for the boiler corrosion are (1) Dissolved oxygen (2) Dissolved  $CO_2$  (3) Acids produced from dissolved salts.

(1) **Removal of Dissolved O<sub>2</sub>:** Dissolved oxygen in water is mainly responsible for boiler corrosion. At high temperatures, dissolved O<sub>2</sub> in water attacks boiler material

$$2Fe + 2H_2O + O_2$$
  $\longrightarrow$   $2Fe(OH)_2$   
 $4Fe(OH)_2 + O_2 + 2H_2O$   $\longrightarrow$   $2[Fe_2O_3 \cdot 3H_2O]$  (rust)

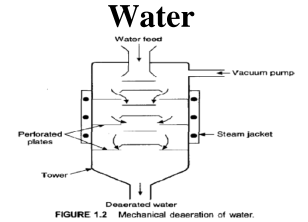
Dissolved oxygen can be removed from water by **chemical** and **mechanical** methods.

**Chemical method:** In this method Sodium sulphite (Na<sub>2</sub>SO<sub>3</sub>), hydrazine (NH<sub>2</sub>NH<sub>2</sub>) are added in small quantity (5-10 ppm) is converted to sodium sulphate and N<sub>2</sub>, respectively.

$$2Na_2SO_3 + O_2 \longrightarrow 2Na_2SO_4$$
  
 $N_2H_4 + O_2 \longrightarrow N_2 + 2H_2O$ 

Hydrazine is found to be an ideal compound for removing dissolved oxygen since the products are water and inert  $N_2$  gas. It removes oxygen without increasing the concentration of dissolved salts.

Mechanical method: Mechanical de-aeration is another method to degasify is dissolved water.



In this method, water is allowed to flow down a perforated plate fitted tower. Vacuum is applied to this tower and the sides of the tower are heated. High temperature and low pressure and large exposed area reduce the quantity of dissolved oxygen in water.

(2) **Removal of CO<sub>2</sub>:** Dissolved CO<sub>2</sub> in water produces carbonic acids which is corrosive nature.  $CO_2 + H_2O \longrightarrow H_2CO_3$ 

Carbon dioxide can be removed from water by chemical or mechanical means. In the chemical method  $CO_2$  is removed from water by the addition of a calculated quantity of  $NH_4OH$ 

$$2NH_4OH + CO_2 \longrightarrow (NH_4)_2CO_3 + H_2O$$

(3) Removal of Acids: Acids produced from salts dissolved in water are also mainly responsible for corrosion of boilers. Certain salts like MgCI<sub>2</sub> and CaCl<sub>2</sub> on hydrolysis at higher temperatures produce hydrochloric acid which corrodes the boiler.

$$MgCl_2 + 2H_2O \longrightarrow Mg(OH)_2 + 2HCl$$

The liberated acid can produce rust in the following way.

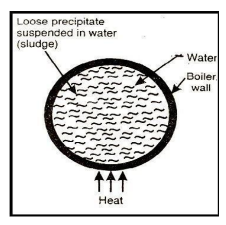
Fe + 2HCI 
$$\longrightarrow$$
 FeCl<sub>2</sub> + H<sub>2</sub>  
FeCl<sub>2</sub> + 2H<sub>2</sub>O  $\longrightarrow$  Fe(OH)<sub>2</sub> + 2HCI  
4Fe(OH)<sub>2</sub> + O<sub>2</sub> + 2H<sub>2</sub>O  $\longrightarrow$  2[Fe<sub>2</sub>O<sub>3</sub>·3H<sub>2</sub>O] (rust)

The presence of a small amount of  $MgCl_2$  can cause a amount extent of corrosion. Corrosion by acids can be avoided by (1) remove  $MgCl_2$  from water (2) adding corrosion inhibitors, sodium silicates, sodium phosphate.

#### (4) Scales and sludge formation:

In industries, boilers are continuously used for stream generation. When water is continuously converted into steam in boilers, the concentration of dissolved salts in water increases progressively .When the concentration of salts reaches to their saturation point, they are thrown out in the form of precipitation on the inner walls of the boilers. The least soluble one gets precipitate first.

**Sludge:** A loose, soft and slimy precipitate formed within the boiler is called **slude**. Sludes can be formed by substances which have greater solubility in hot water and less solubility in cold water. Salts like MgCO<sub>3</sub>, MgCl<sub>2</sub>, MgSO<sub>4</sub> and CaCl<sub>2</sub> etc are responsible for slude formation in boilers



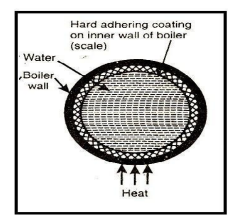
#### **Disadvantages of sludge formation:**

- (1) Sludges are poor conductor of heat, so they tend to waste a portion of heat generated.
- (2) If sludges are formed along with scales, then former gets entrapped in the lather and both get deposited as scales.
- (3) Excessive sludge formation disturbs the working of the boiler. It settles in the regions of poor water circulation in pipes.

## **Prevention of sludge formation:**

- (1) By using soft water which is free from salts like MgCO<sub>3</sub>, MgCl<sub>2</sub>, CaCl<sub>2</sub> and MgSO<sub>4</sub> can prevent sludge formation.
- (2) By frequently blow down operation.

**Scale:** On the other hand, if the precipitate is hard and sticky and forms adherent coating on the inner walls of the boiler, it is called **scale**. Scales are formed by substance like Ca (HCO<sub>3</sub>)<sub>2</sub>, CaSO<sub>4</sub> & Mg (OH)<sub>2</sub>.



Scales may be formed inside the boiler due to:

(1) Decomposition of calcium bicarbonate

$$Ca(HCO_3)_2 \longrightarrow CaCO_3 \downarrow + H_2O + CO_2$$
  
Scale

However scale composed chiefly of calcium carbonate is soft and is the main cause of scale formation in low – pressure boilers but in high pressure boilers, CaCO<sub>3</sub> is soluble

$$CaCO_3 + H_2O \longrightarrow Ca(OH)_2 + CO_2$$
  
Soluble

## (ii) Deposition of calcium sulphate:

The solubility of CaSO<sub>4</sub> in water decreases with rise of temperature. Thus, solubility of CaSO<sub>4</sub> is 3200 ppm at 15<sup>0</sup>C and it reduce to 55 ppm at 30<sup>0</sup>C and 27 ppm at 32<sup>0</sup>C. In other words, CaSO<sub>4</sub> is soluble in cold water, but almost completely insoluble in super heated water. Consequently, CaSO<sub>4</sub> gets precipitated as hard scale on the boiler. This is the main cause of scales in high – pressure boilers.

## (iii) Hydrolysis of Magnesium salts.

Dissolved Mg salts undergo hydrolysis forming magnesium hydroxide precipitate, which forms a soft type of scale.

$$MgCl_2 + 2H_2O \longrightarrow Mg (OH)_2 \downarrow + 2 HCl$$
  
Scale

## (iv) Presence of Silica (SiO<sub>2</sub>)

Silica present in small quantities, deposits as calcium silicate  $(CaSiO_3)$  and / or Magnesium silicate  $(MgSiO_3)$ , these deposits stick very firmly on the inner side of the boiler surface and are very difficult to remove. One important source of silica in water is the sand filter.

#### Disadvantage of scale formation

- (i) Wastage of Fuel: Scales have a poor thermal conductivity so the rate of heat transfer from boiler to inside water is greatly reduced. In order to provide a steady supply of heat to water excessive or over heating is done and this causes increase in fuel consumption.
- (ii) Lowering of boiler safety: Due to scale formation over heating of boilers is done in order to maintain a steady supply of steam. It makes the boiler material softer and weaker. This causes distortion of boiler tube and also makes the boiler unsafe to bear the pressure of the steam especially in high pressure boilers.
- (iii) **Decrease in efficiency:** Deposition of scale in the valves and condensers of the boiler, choke them partially. This results in decrease in efficiency of the boiler.
- (iv) **Danger of explosion**: When thick scale cracks due to uneven expansion, the water comes suddenly in contact with over heat portion and large amount of steam is formed instantaneously. This results in development of sudden high pressure which may cause explosion of the boiler.

**Prevention of sludge formation:** Scales can be removed by mechanical and chemical methods. (1) Mechanical methods: They can be removed with a scraper (or) a piece of wood (or) wire brush. They can also be removed by blow down.

(2) Chemical methods: If scales are hard and adhering, they can be removed by dissolving in chemicals.

CaCO<sub>3</sub> scales are dissolved in 5-10% HCl

CaSO4 complex is highly soluble in adding EDTA.

#### SOFTENING OF WATER

To minimize boiler troubles, the water must be perfectly soft. So the process of removing the hardness causing salts from water called softening of water. There are two methods are available for softening of water.

- 1) Internal methods
- 2) External methods
- (1) <u>Internal treatment methods:</u> In this method raw water is treated inside the boiler and addition of suitable chemicals to reduce scale and sludges formation. Some important internal treatments are (a) Collodial conditioning (b) Phosphate conditioning (c) Calgon conditioning (d) Carbonate conditioning.
  - (a) The addition of colloidal substances, like Kerosene, tannin, agar-agar added to low pressure boilers. These substances get adsorbed over the scale forming precipitates and yield non-sticky, loose deposits, which can be easily removed by blow down.
  - (b) Phosphate conditioning is generally to high pressure boilers. When Sodium phosphate is added to boiler water, it react with Mg and Ca salts forming soft sludges of Ca<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub> and Mg<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub>.

$$3CaCl_2 + 2 Na_3PO_4 \longrightarrow Ca_3(PO_4)_2 + 6NaCl$$
  
 $3MgCl_2 + 2 Na_3PO_4 \longrightarrow Mg_3(PO_4)_2 + 6NaCl$ 

Sodium phosphate is used when the alkalinity of boiler water PH 9.5-10.5. If the alkalinity of boiler water is too high, NaH<sub>2</sub>PO<sub>4</sub> (dihydrogen phosphate) is used.

(c) Sodium hexametaphosphate is called CALGON (Na<sub>2</sub>[Na<sub>4</sub>(PO<sub>3</sub>)<sub>6</sub>] (or) (NaPO<sub>3</sub>)<sub>6</sub>. It reacts with scale forming CaSO<sub>4</sub> and forms a water soluble compound.

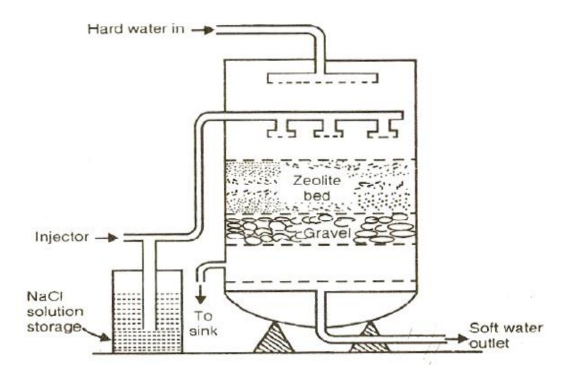
$$Na_2 [Na_4 (PO_3)_6] \longrightarrow 2Na + [Na_4P_6O_{18}]^{-2}$$
 $2CaSO_4 + [Na_4P_6O_{18}]^{-2} \longrightarrow [Ca_2P_6O_{18}]^{-2} + 2Na_2SO_4$ 

(d) The hard and strong bind scales formed due to CaSO<sub>4</sub> are avoided by the addition of Na<sub>2</sub>CO<sub>3</sub> to boiler water. The CaSO<sub>4</sub> is converted to CaCO<sub>3</sub>, which is loose sludge and it can be removed by blow down operation.

$$CaSO_4 + Na_2CO_3 \longrightarrow CaCO_3 + Na_2SO_4$$

(2) External treatment methods: The treatment given to water for the removal of hardness causing salts before into the boiler is called External treatment. Three External methods are: (a) Zeolite process (or) Permutit process (b) Ion exchange process (or) Deionisation (or) Demineralisation (c) Lime-soda process.

## (a) Zeolite process (or) Permutit process



The chemical composition of Zeolites is hydrated aluminium silicate, represented as  $Na_2O$ ,  $Al_2O_3$   $xSiO_2$  y  $H_2O$  where x=2-10 and y=2-6. It can be represented as Sodium Zeolite ( $Na_2Ze$ ). Zeolites are capable of exchange reversible its sodium ion for hardness causing  $Ca^{+2}$  and  $Mg^{+2}$  in water. Hence Zeolites are cation exchangers.

Zeolites are mainly 2 types:

- 1. Natural zeolites are non-porous, eg. Natrolite- Na2O.Al2O3.4SiO2.2H2O.
- 2. **Synthetic Zeolites** are porous and possess gel structure. Synthetic Zeolites possess higher exchange capacity.

**PROCESS:** The hard water is passed through a Zeolite bed fixed in a cylinder at a specific rate. The hardness causing ions Ca<sup>+2</sup> and Mg<sup>+2</sup> are retained by the Zeolite as CaZe and MgZe, respectively.

**REGENERATION:** After some time the Zeolite bed is completely converted to Calcium and Magnesium Zeolites and no purification of raw water takes place. i.e., the

Zeolite bed got exhausted. At this stage the purification of hard water is stopped and the Zeolite bed is regenerated by treating with 10% brine (NaCl) solution.

$$CaZe + 2 NaCl \longrightarrow Na2Ze + CaCl2$$

$$MgZe + 2 NaCl \longrightarrow Na2Ze + MgCl2$$

## **Advantages:**

- **a**. It removes the hardness completely and water of about 10 ppm hardness is produced.
- **b**. The equipment used is compact, occupying a small space.
- **c**. No impurities are precipitated, so there is no danger of sludge formation in the treated water at a later stage.
- **d**. The process automatically adjusts itself for variation in hardness of incoming water.
- e. It is quite clean
- **f**. It requires less time for softening.
- **g**. It requires less skill for maintenance as well as operation.

## **Disadvantages:**

- **a.** The treated water contains more sodium salts than in lime-soda process.
- **b**. Hard water containing acids destroys the Zeolite bed.
- **c**. If suspended particles (turbidity) are present the pores of the Zeolites are blocked and softening is not possible.
- **d**. If raw water contains large quantities of coloured ions such as Mn+2 and Fe+2 they must be removed first, because these ions produce magnesium and Ferrous zeolites, which cannot be regenerated.

## (b) Ion exchange process (or) Deionisation (or) Demineralisation

Ion exchange process also known as demineralization or de-ionization process. Ion-Exchange resins are insoluble and cross linked long chain organic polymers with a micro porous structure, and the "functional Groups" attached to the chains are responsible for the ion-exchanging properties.

#### Ion exchangers are two types

- 1. Cation Exchange Resins
- 2. Anion Exchange Resins
- 1 Cation Exchange Resins: Cation exchangers are capable of exchanging their H<sup>+</sup> ions

- replaced by cations such as  $Mg^{+2}/Ca^{+2}$ . The general formula of cation exchangers are RH. These are made by copolymers of **styrene** and **divinyl benzene** containing the functional groups **COOH** (or)  $-SO_3H$ .
- 2 **Anion Exchange Resins:** Anion exchange resins are **styrene-divinyl benzene** copolymers which contain **amino**, **quaternary ammonium group** an internal part of the resins, which exchange OH ion in the dissolved salts. The anion exchangers are represented by the formula R'OH.

**Softening Process:** The hard water is first passed through cation exchange resins, which remove  $Ca^{+2}$  and  $Mg^{+2}$  ions in the following way.

$$2RH + Ca(HCO_3)_2 \longrightarrow R_2Ca + 2H_2CO_3$$

$$2RH + Mg(HCO_3)_2 \longrightarrow R_2Mg + 2H_2CO_3$$

$$2RH + MgCl_2 \longrightarrow R_2Mg + 2HCl$$

$$2RH + CaCl_2 \longrightarrow R_2Ca + 2HCl$$

$$2RH + MgSO4 \longrightarrow R_2Mg + H_2SO_4$$

$$2RH + CaSO4 \longrightarrow R_2Ca + H_2SO_4$$

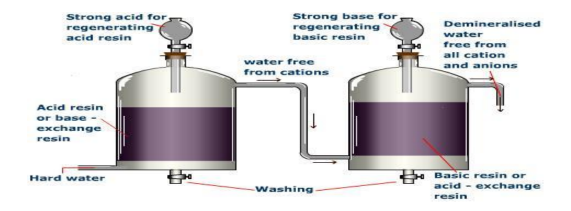
$$2RH + Mg(NO_3)_2 \longrightarrow R_2Mg + 2HNO_3$$

$$2RH + Ca(NO_3)_2 \longrightarrow R_2Ca + 2HNO_3$$

The  $\mathrm{Mg^{+2}}$  and  $\mathrm{Ca^{+2}}$  are retained by the cation exchangers as CaR and MgR releasing H+ into water. The water coming out of the resin is highly acidic. Then the water is passed through the anion exchange resin, the acids present in water are removed by the exchanger releasing  $\mathrm{OH^-}$  into water. The  $\mathrm{H^+}$  and  $\mathrm{OH^-}$  are combined and produce  $\mathrm{H_2O}$ .

$$\begin{array}{ccc} R'OH + HCI & \longrightarrow & R'CI + H_2O \\ 2 R'OH + H_2SO_4 & \longrightarrow & R'_2SO_4 + 2H_2O \\ 2 R'OH + H_2CO_3 & \longrightarrow & R'HCO_3 + H_2O \\ 2 R'OH + HNO_3 & \longrightarrow & R'NO_3 + H_2O \end{array}$$

Thus the water coming out from exchangers is free from all ions known as deionized (or) demineralized water.



**Regeneration:** After deionization the cation and anion exchangers will be exhausted. Regeneration of cation exchanger is carried out by passing dil. HCl (or)  $H_2SO_4$ .

$$R_2Ca + 2HCl \longrightarrow 2RH + CaCl_2 \text{ (washings)}$$
  
 $R_2Ca + H_2SO_4 \longrightarrow 2RH + MgSO_4 \text{ (washings)}$ 

Similarly the anion exchanger is treated with dil.NaOH solution.

$$R'Cl + NaOH \longrightarrow R'OH + NaCl (washings)$$
  
 $R_2'SO4 + NaOH \longrightarrow 2R'OH + Na_2SO_4 (washings)$   
 $R'NO_3 + NaOH \longrightarrow R'OH + NaNO_3 (washings)$ 

The washings are discarded into sink. The regenerated ion-exchanger is used for softening. **Advantages:** (1) Highly acidic (or) alkaline water samples can be purified by this process. (2) The hardness possessed by the deionized water is 2 ppm. **Disadvantages:** (1) The resins are expensive, the cost of purification is high. (2) The raw water should contain turbidity below 10 ppm, otherwise pores in the resin will be blocked and output of the process is reduced.

## (C) Lime (L)-Soda (S) Process

The basic principle of this process is to chemically convert all the soluble hardness causing impurities into insoluble precipitates which may be removed by settling and filtration.

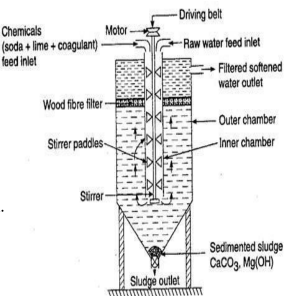
In this process Lime  $[Ca(OH)_2]$  and Soda  $[Na_2CO_3]$  are the reagents used to precipitate the dissolved salts of  $Ca^{+2}$  and  $Mg^{+2}$  as  $CaCO_3$  and  $Mg(OH)_2$ . The precipitates are filtered off. Lime soda process mainly two types, they are

- 1. Cold Lime-soda process
- 2. Hot Lime-soda process.

#### 1. Cold Lime-soda process

In this method, water is treated with lime [Ca(OH)<sub>2</sub>] and Soda [Na<sub>2</sub>CO<sub>3</sub>] at room temperature. Precipitate and filtration cannot be done easily. Hence small amount of alum is added. Sodium aluminate (NaAlO<sub>2</sub>) can act as a coagulant and help to removal of silica and oil.

Raw water along with calculated amount of chemicals (Lime+ Soda+ Coagulant) are fed from the top into the Chamber and sludge setting at the bottom is drawn off. The water sample contains residual hardness 50-60 ppm.

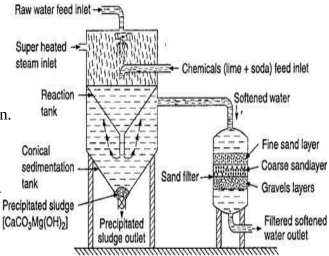


## 2. Hot Lime-soda process

In this process the raw water is treated with Softening chemicals at 80-150°C. At higher Precipitation takes place and coagulant is not Required.

Advantages of hot L-S process:

- 1. Precipitation is rapid and completes in 15 min.
- 2. No coagulant is required
- 3. Bacteria are killed in hot L-S process.
- 4. The process is economical
- 5. The residual hardness of water is 15-30 ppm.



Lime reacts with temporary hardness causing salts, magnesium permanent hardness, CO<sub>2</sub>, acids, bicarbonates, which could be removed by soda.

## Amount of Lime required for softening:

74/100 [temporary  $Ca^{+2}$  hardness+ (2 x temporary  $Mg^{+2}$ ) + permanent  $Mg^{+2}$  hardness+  $CO_2$ +  $\frac{1}{2}$  HCl+H<sub>2</sub>SO<sub>4</sub>+  $\frac{1}{2}$  NaHCO<sub>3</sub>+I<sub>2</sub>KHCO<sub>3</sub>+FeSO<sub>4</sub> + (3 x Al<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub>-1/2 NaAlO<sub>2</sub> hardness] all the hardness in terms of CaCO3 equivalent.

## Amount of soda required for softening:

106/100 [permanent  $Ca^{+2}$  hardness + permanent  $Mg^{+2}$  hardness + ½  $HCl+H_2SO_4$ -½ NaHCO<sub>3</sub>- 1/2KHCO<sub>3</sub>+ (3 x Al<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub> + FeSO<sub>4</sub> hardness] all the hardness in terms of CaCO<sub>3</sub> equivalent.

In this process, calculated amount of lime and soda is added to the hard water. Lime

(a) Removal of Dissolved CO<sub>2</sub> and H<sub>2</sub>S

$$Ca(OH)_2 + CO_2 \longrightarrow CaCO_3 \downarrow + H_2O$$

(b) Neutralization of free mineral acids

$$Ca(OH)_2 + 2HCI \longrightarrow CaCl_2 + 2H_2O$$
  
 $Ca(OH)_2 + H_2SO_4 \longrightarrow CaSO_4 + 2H_2O$ 

(c) Removal of temporary hardness

$$Ca(OH)_2 + Ca(HCO_3)_2$$
 2  $CaCO_3 \downarrow + 2H_2O$   
2 $Ca(OH)_2 + Mg(HCO_3)_2$  Mg  $(OH)_2 \downarrow + 2CaCO_3 + 2H_2O$ 

(d) Removal of permanent magnesium hardness

$$Ca(OH)_2 + MgCl_2 \longrightarrow Mg(OH)_2 \downarrow + CaCl_2$$

(e) Removal of dissolved Iron and Aluminium salts.

$$Ca(OH)_2 + FeSO_4 \longrightarrow Fe(OH)_2 + CaSO_4$$
Unstable
$$2Fe(OH)_2 + H_2O + O_2 \longrightarrow 2Fe(OH)_3$$

(f) Removal of bicarbonate salts.

$$2NaHCO_3 + Ca(OH)_2$$
  $\longrightarrow$   $CaCO_3 + Na_2CO_3 + 2H_2O$   
 $2KHCO_3 + Ca(OH)_2$   $\longrightarrow$   $CaCO_3 + K_2CO_3 + 2H_2O$ 

NaHCO<sub>3</sub> present in hard water reacts with soda and produce more sodium carbonate.

(f) Reactions of Soda (Na<sub>2</sub>CO<sub>3</sub>). Soda Remove all the Ca<sup>2+</sup> permanent hardness

$$Na_{2}CO_{3} + CaCl_{2} \longrightarrow CaCO_{3} \downarrow + 2 NaCl$$

$$Na_{2}CO_{3} + CaSO_{4} \longrightarrow CaCO_{3} \downarrow + Na_{2}SO_{4}$$

$$Al_{2} (SO_{4})_{3} + 3Ca(OH)_{2} \longrightarrow 2Al(OH)_{3} + 3CaSO_{4} + H_{2}O$$

$$NaAlO_{2} + 2H_{2}O \longrightarrow Al (OH)_{3} + NaOH$$

$$2NaOH + CaCl2 \longrightarrow Ca(OH)_{2} + 2NaCl$$

$$2NaOH = Ca(OH)_{2}$$

NaOH produced reacts with hardness causing salts and produces lime. Hence, its hardness must be subtracted from lime required.

#### Advantages of lime-soda process:

- 1. It is very economical
- 2. If this process is combined with sedimentation with coagulation, lesser amounts of coagulants shall be needed.
- 3. The process increases the pH value of the treated-water; thereby corrosion of the distribution pipes is reduced.
- 4. Besides the removal of harness, the quantity of minerals in the water is reduced.
- 5. To certain extent, iron and manganese are also removed from the water.
- 6. Due to alkaline nature of treated-water, amount of pathogenic bacteria in water is considerably reduced.

## Dis-advantages of lime-soda process:

- 1. For efficient and economical softening, careful operation and skilled supervision is required.
- 2. Disposal of large amounts of sludge or insoluble precipitates poses a problem. However, the sludge may be disposed off in raising low-lying areas of the city.

	Cold L-S Process	Hot L - S Process	
Sl.No			
1	It done at room temp $(25^{\circ} - 30^{\circ}C)$	It is done at elevated temp (90° – 100°C)	
2	It is a slow process	It is a rapid process	
3	The use of coagulants is must	Coagulants not needed	
4	Filtration is not easy	Filtration is easy as the viscosity of wate	
		becomes low at elevated temperature	
5	Softened water has residual	Softened water has residual hardness of	
	hardness around 60ppm	15-20 ppm	
6	Dissolved gases are not removed	Dissolved gases such as CO <sub>2</sub> are removed	
	_	to some extent	

LIME-SODA PROCESS	ION-EXCHANGE/RESIN
	Water treatment plant occupies less
Water treatment plant occupies more area or place.	area.
Water after treatment has lesser dissolved	Water after treatment free from
solids.	dissolved solids
This method of water treatment plants is not	This method of water treatment
much	plants is more
expensive and material used is cheap.	expensive and material used for softening is expensive.
Operation expenses are higher	Operation expenses are higher
The cost incurred on softening of water is	The cost incurred on softening of
high.	water is high
It cannot operate in under pressure.	It can even operate in under pressure.
	It can be used for treating acidic and
It can be used for treating acidic water also.	alkaline water also.
There is a problem of settling, coagulation and	There is problem of turbidity.
removal of sludge.	
It is not possible.	This method can be made automatic.
In order to meet the changing hardness of	Control test comprises only in
incoming	checking the
water, frequent control and adjustment of reagents is needed.	hardness of treated-water.
11.6	Residual hardness is low about 0 to 2
Residual hardness is low about 15 to 50 ppm	ppm
	It is very good for treating water for
It is not good for boilers	use in high pressure boilers.
	It required less skill for maintenance
Skilled persons required	as well as operation

## Estimation of temporary & permanent hardness of water by EDTA method:

The hardness of water indicates the quality in terms of Calcium and Magnesium expressed in terms of CaCO<sub>3</sub>. The analysis is done by complexometric titration using standard EDTA and EBT as an indicator.

**Principle:** In this complex metric Titration, the water sample is titrated with standard solution of Di sodium salt of EDTA using EBT indicator.

This method give more accurate results than the soap solution method.

$$\begin{array}{c} \text{HOOC - H}_2\text{C} \\ \text{N - CH}_2\text{ - CH}_2\text{ - N} \\ \text{HOOC -H}_2\text{C} \end{array} \\ \begin{array}{c} \text{CH}_2\text{ - COOH} \\ \text{CH}_2\text{ - COOH} \\ \end{array}$$

Ethylene diamine tetra acetic acid (EDTA) forms complexes with  $Ca^{2+}$  and  $Mg^{2+}$  as well as with many other metal cations, in aqueous solution. These complexes have the general formula given below

**Reactions involved**: EBT indicator when added to hard water at pH = 10, forms weak complexes with calcium and magnesium present in hard water. It results in the formation of Ca-EBT or Mg-EBT complexes which is wine-red, these are unstable.

$$\begin{array}{c} (Ca^{2+} \, / \, Mg^{2+} \, ) + EBT \longrightarrow [Ca - EBT \, ] \, / \, [Mg - EBT] \\ blue & wine-red \ color \ complex \end{array}$$

During titration with EDTA, EDTA extracts metal ions from the above complex to give a colorless and stable complex releasing the free indicator (blue). The color changes from wine-red to blue at the endpoint.

Various steps involved in this method are...

1. **Preparation of standard hard water:** Dissolve 1g of pure, dry CaCO<sub>3</sub> in minimum quantity of dil. HCl and then evaporate the solution to dryness on a water bath. Dissolve the residue in distilled water to make 1 Litre solution. Each mL of this solution thus contains 1mg of CaCO<sub>3</sub> equivalent hardness.

1 mL hard water solution = 1mg of CaCO<sub>3</sub> equivalent hardness.

- 2. **Standardization of EDTA solution:** Rinse and fill the burette with EDTA solution. Pipette out 50 mL of standard hard water in a conical flask. Add 10-15 mL of buffer solution and 4 to 5 drops indicator. Titrate with EDTA solution till wine-red colour changes to clear blue. Let volume used by V1 mL.
- 3. **Titration of Unknown Hard water:** Rinse and fill the burette with EDTA solution. Pipette out 50 mL of unknown sample hard water in a conical flask. Add 10-15 mL of buffer solution and 4 to 5 drops indicator. Titrate with EDTA solution till wine-red colour changes to clear blue. Let volume used be V2 mL.
- 4. **Titration of Permanent hardness:** Take 100 mL of the water sample in a large beaker. Boil it. Till the volume is reduced to about 50 mL, filter it, makeup the volume to 100 mL with distilled water. Then, titrate 50 mL of boiled water sample just as in Step (2). Let volume used by V3 mL.

## **CALCULATION:**

Total hardness of water = 
$$\frac{1000 \times V_2}{V_1}$$
 ppm

$$Permanent\ hardness = \frac{1000\ X\ V_3}{V_1} \quad ppm$$

Temporary hardness = [Total hardness – Permanent hardness]

$$Temporary\ hardness = \underbrace{1000\ X\ (V_{\underline{2}^{\text{-}}}\ V_{\underline{3}})}_{V1}\ ppm$$

**Impurities in water**: The impurities present in water may be broadly classified as follows. [1] *Dissolved impurities*: The dissolved impurities are mainly the carbonates, bicarbonates, chlorides and sulphates of calcium, magnesium, iron, sodium and potassium. The presence of these salts imparts hardness to water. The dissolved impurities also include dissolved gases like oxygen and carbon

- dioxide. [2] Suspended impurities: The following are the types of suspended impurities:
- (a) Inorganic: Clay and sand (b) Organic: Oil globules, vegetable and animal matter. The above suspended impurities impart turbidity, colour and odour to water.
- [3] *Colloidal impurities*: They are finely-divided silica and clay, organic waste products, complex protein amino acids, etc.
  - [4] *Microorganisms*: They are algae, fungi and bacteria

**Potable Water (water for domestic supply):** "Water free from contaminants or water that is safe for human consumption is called potable water". Municipalities have to supply potable water, i.e., It is a water of sufficiently high quality that it can be consumed or used without risk of immediate or long term harm. The following are *characteristics* of *potable water*.

- 1) Water should be clear, colorless and odorless.
- 2) Free from pathogenic bacteria and gases like H<sub>2</sub>S
- 3) Hardness of water must be 125 ppm.
- 4) PH at 7-8
- 5) Terbidity in drinking water should not exceed 25 ppm
- 6) Free from heavy metals like Pb, As, Cr and Mn.

**Desalination:** The process of removal of dissolved salts (NaCl, KCl) present in water is known as desalination. Water is divided into 3 categories on the basis of salinity.

- (1) Sea water: The salinity is greater than 35,000 mg/l
- (2) Brakish water: The salinity is in the range of 1000 -35,000 mg/l.
- (3) Fresh water: The salinity of water is less than 1000 mg/l.

  Brakish water and sea water are not fit for drinking as well as industrial purposes.

  They can be subjected to desalination to make them suitable for drinking. Important

desalinations methods are (1) Reverse Osmosis (2) Electrodialysis (3) Distillation

Various stages involved in the purification of water for domestic purposes. Sources of water ->> Screening ->> Aeration ->> Sedimentation ->> Filtration ->> Sterilization (or) Disinfection ->> Storage and Distribution

#### **Screening**

Screening is the process of removing floating materials like wood pieces and leaves from water. Raw water is allowed to pass through a screen having a large number of holes which removes the small and large floating matter.

#### Sedimentation

Sedimentation is the process of removing suspended impurities by allowing the water to stand undisturbed for 2-6 hours in big tanks. Due to force of gravity, most of the suspended particles settled down at the bottom and they are removed. Sedimentation removes only 70-75% of the suspended matter.

## Coagulation

Finely-divided silica, clay, etc. do not settle down easily and hence cannot be removed by sedimentation. Most of these are in colloidal form and are negatively charged and hence do not coalesce because of mutual repulsion. Such impurities are removed by coagulation method. Here, certain chemicals like alum and  $Al_2(SO_4)_3$  are added to water. When  $Al_2(SO_4)_3$  is added to water, it hydrolyzes to form a gelatinous precipitate of  $Al(OH)_3$ ' The gelatinous precipitate of  $Al(OH)_3$  settles to the bottom and can be removed easily.

#### **Filtration**

For removing bacteria, colour, taste, odour, fine suspended particles, etc. and to produce clear water, filtration is used. In this process, water is passed through beds of fine sand, coarse sand and other granular material. The porous material used is the filtering medium and the equipment used for filtration is known as filter, e.g. slow sand filter.

A typical slow sand filter is shown in Figure 1.5. It consists of a tank containing thick beds of fine sand (at the top), coarse sand, fine gravel and coarse gravel (at the bottom). When the water passes through the filtering medium, it flows through the various beds slowly due to gravity. The rate of filtration slowly decreases due to the blockage of impurities in the pores of the sand bed. When the rate of flow becomes very slow, filtration is stopped and the bed is

cleaned by scraping of a smaller layer of the sand bed (top layer) and replacing it with the clean sand. Bacteria are partly removed by this filtration process.

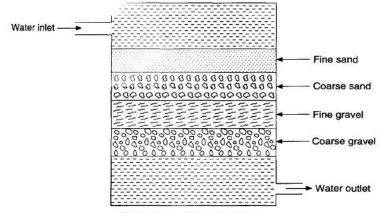


FIGURE 1.5 Slow sand filter.

#### **Disinfection / Sterilization:**

**Disinfection**: The process of destroying/killing the disease producing Bacteria, microorganisms from the water and making it safe for use, is called disinfection.

**Disinfectants:** The chemicals or substances which are added to water for killing the Bacteria. The disinfection of water can be carried out by following methods:

- A) **Boiling**: Water for 10-15 minutes, all the disease-producing bacteria are killed and water becomes safe for use. This method is not possible for the municipal supply of water.
- B) **Ozone:** Ozone is a powerful disinfectant and is readily absorbed by water. Ozone is highly unstable and decomposes to give **nascent oxygen** which is capable of destroying the bacteria.  $O_3 \longrightarrow O_2 + [O]$ . This process is relatively expensive and ozone is unstable and cannot be stored for long time.
- C) UV: UV light is used as a disinfectant for swimming pool water. No chemical are used and it is safe for skin. In this process, water is exposed to UV rays, which are generated from an electric mercury vapor lamp.
- D) **Chloramines**: When added chloramine in water, which generates hypochlorous acid, a powerful disinfectant, that kills bacteria.

$$ClNH_2 + H_2O \longrightarrow HOCl + NH_3$$

HOCl inactivates the enzymes of bacteria and kills. Chloramine is useful for disinfection in swimming pools.

E) **Bleaching powder**: Calculated amount of bleaching powder is added to water and allowed to stand for several hours. Bleaching powder contains 80% chlorine.

$$CaOCl_2 + H_2O \longrightarrow Ca(OH)_2 + Cl_2$$
  
 $Cl_2 + H_2O \longrightarrow HCI + HOCI$ 

HOCI + Bacteria → Bacteria are killed

Disadvantages: 1) Bleaching powder introduces Ca<sup>+2</sup> hardness in water and adds lime residue. 2) Excess of it gives bad smell and bad taste. 3) Excess chlorine is irritating to mucous membrane.

F) **Chlorine**: Chlorination is the process of purifying the drinking water by producing a powerful Germicide like Hypochlorous acid. When this Chlorine is mixed with water it produces Hypochlorous acid which kills the germs present in water.

Chlorine is basic (means pH value is more than 7) disinfectant and is much effective over the germs. Hence Chlorine is widely used all over the world as a powerful disinfectant. Chlorinator is an apparatus, which is used to purify the water by chlorination process.

### **Advantages**

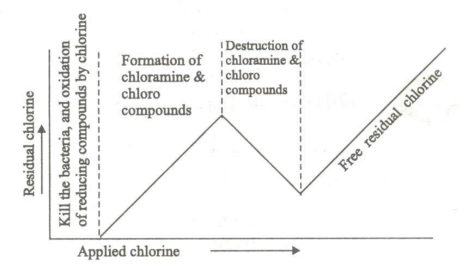
- 1. It is effective and economical
- 2. It requires very little space for storage
- 3. It is stable and does not deteriorate on keeping
- 4. It can be used at low as well as high temperatures
- 5. It introduces no salt impurities in the treated water.
- 6. It is the most ideal disinfectant.

#### **Disadvantages**

- 1. Excess chlorine if added, produces unpleasant taste and odour
- 2. Its excess produces irritation on mucous membrane

**Break-point chlorination (or free-residual chlorination)** involves the addition of sufficient amount of chlorine to oxidize (a) organic matter (b) reducing substances (c) Free ammonia; leaving behind free chlorine which possesses disinfecting action against disease-producing bacteria.

A known volume of sample of water is treated with an increasing amount of chlorine and then analyzed for the remaining chlorine. When a graph is plotted between the amounts of residual chlorine against the added chlorine, a curve is obtained as shown below. The addition of Chlorine at the dip or break is called break point chlorination. This indicates the point at which free chlorine begins to appear and resulting in water free from bad tastes and odor.



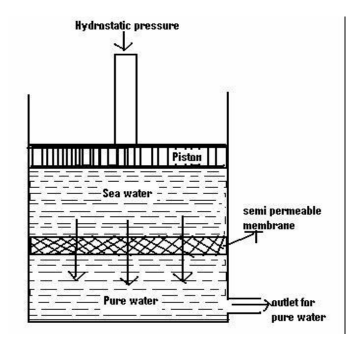
**Desalination:** The process of removing common salt from the water is known as desalination. The water containing dissolved salts with a peculiar salty taste, is called brackish water. Sea water, containing on an average about 3.5% salts, comes under this category. Brackish water is totally unfit for drinking purpose.

Commonly used methods for the desalination of brackish water is:

- 1. Reverse Osmosis
- 2. Electrodyalias
- 3. Distillation

**Reverse Osmosis:** When two solutions of unequal concentrations are separated by a semi permeable membrane, flow of solvent takes place from dilute to concentrated sides, due to osmosis. If, however, a hydrostatic pressure in excess of osmotic pressure is applied on the concentrated side, the solvent flow reverse, ie. Solvent is forced to move from concentrated side to dilute side across the membrane. This is the principle of reverse osmosis. Thus, in reverse osmosis (RO) methods, pure solvent is separated from contaminants, rather than removing contaminants from the water.

This membrane filtration is also called 'super-filtration' or 'hyper – filtration'.



Process: in Reverse osmosis process, pressure (15 to 40 kg cm<sup>-2</sup>) is applied to the sea water or impure water to force its pure water out through the semi-permeable

membrane, leaving behind the dissolved solids. The principle of reverse osmosis, as applied for treating saline or sea water. The membrane consists of very thin films of cellulose acetate, affixed to either side of a perforated tube. However, mover recently superior membranes made of polymethacrylate and polyamide polymers have come into use.

#### **Advantages:**

- a. Reverse osmosis possesses a distinct advantage of removing ionic as well as non-ionic, colloidal and high molecular weight organic matter.
- b. It removes colloidal silica, which is not removed by demineralization.
- c. The maintenance cost is almost entirely on the replacement of the semi permeable membrane.
- d. The life time of membrane is quite high, about 2 years.
- e. The membrane can be replaced within a few minutes, thereby providing nearly uninterrupted water supply.
- f. Due to low capital cost, simplicity, low operating cost and high reliability, the reverse osmosis is gaining ground at present for converting sea water into drinking water and for obtaining water for very high pressure boilers.

## **Alkalinity of water:**

Alkalinity is a measure of the capability of water to absorb  $H^+$  ions without significant change of pH. In other words, alkalinity is a measure of the acid buffering capacity of water. The determination of alkalinity of water is necessary for controlling the corrosion, to calculate the amount of lime and soda needed for water softening; in conditioning the boiler feed water, etc. Alkalinity of a sample of water is due to the presence of  $OH^-$  (hydroxide ion),  $HCO_3^-$  (bicarbonate ion) and  $CO_3^{\ 2^-}$  (carbonate ion) or the mixture of two ions present in water. The possibility of  $OH^-$  and  $HCO_3^-$  ions together is not possible since they combine together to form  $CO_3^{\ 2^-}$  ions.

$$OH^- + HCO_3^- \longrightarrow CO_3^{2-} + H_2O$$

The alkalinity due to different ions can be estimated separately by titration against standard acid solution, using selective indicators like phenolphthalein and methyl orange.

i) OH
$$^-$$
 + H $^+$   $\longrightarrow$  H<sub>2</sub>O  
ii) CO<sub>3</sub>  $^{2-}$  + H $^+$   $\longrightarrow$  HCO<sub>3</sub>  $^-$   
iii) HCO<sub>3</sub>  $^-$  + H $^+$   $\longrightarrow$  H<sub>2</sub>O + CO<sub>2</sub>

The neutralization reaction up to **phenolphthalein** end point shows the completion of reactions (i) and (ii) (OH $^-$  and CO $_3$  $^{2-}$ ) and (CO $_3$  $^{2-}$  and HCO $_3$  $^{-}$ ) only. The amount of acid used thus corresponds to complete neutralization of OH $^-$  plus half neutralization of CO $_3$  $^{2-}$ . This is also called partial alkalinity.

$$P = OH^{-} + \frac{1}{2}CO_{3}^{2}$$
  
 $M = OH^{-} + CO_{3}^{2} + HCO_{3}^{2}$ 

The titration of water sample using methyl **orange** indicator marks the completion of the reactions (i), (ii) and (iii). The amount of acid used after phenolphthalein end point corresponds to one half of normal carbonate and all the bicarbonates. Total amount of acid used represent the total alkalinity due to all ions present in water sample. This is also called total alkalinity.

The table1 below shows the type and amount of alkalinity in water

S.No	Result of Titration	OH <sup>-</sup> ion	CO <sub>3</sub> <sup>2-</sup>	HCO <sub>3</sub>
1	P=0	nil	nil	M
2	P=M	P or M	nil	nil
3	P=1/2M(V1=V2)	nil	2P	nil
4	P>1/2M(V1>V2)	2P-M	2(M-P)	nil
5	P<1/2M(V <v2)< td=""><td>nil</td><td>2P</td><td>M-2P</td></v2)<>	nil	2P	M-2P

- i) Phenolphthalein alkalinity (P) = 0; that means the volume of acid used till the completion of reaction (i) and (ii) is 0. This can only happen when both OH<sup>-</sup> and CO<sub>3</sub> <sup>2-</sup> ions are not present in water. Alkalinity is present due to HCO<sub>3</sub> <sup>-</sup> ion only which can be determined using methyl orange indicator and called methyl orange alkalinity (M).
- ii)  $P = \frac{1}{2} M$ ; indicates that only  $CO_3^{2-}$  ions are present. Using phenolphthalein indicator neutralization reaches up to  $HCO_3^{-}$  but using methyl orange indicator the complete neutralization of  $HCO_3^{-}$  takes place.
- iii)  $P > \frac{1}{2}M$ ; implies  $OH^-$  ions are also present along with  $CO_3^{\ 2^-}$  ions. Up to phenolphthalein alkalinity  $OH^-$  ions will be neutralized completely, whereas  $CO_3^{\ 2^-}$  will be neutralized up to  $HCO_3^{\ -}$  ion. But using methyl orange indicator  $HCO_3^{\ -}$  will be completely neutralized along with  $OH^-$  and  $CO_3^{\ 2^-}$ .
- iv)  $P < \frac{1}{2}$  M; indicates that beside  $CO_3^{2-}$  ions  $HCO_3^{-}$  ions are also present. The volume of acid required for the neutralization up to phenolphthalein end point correspond half neutralization of  $CO_3^{2-}$  (equation ii). Neutralization using methyl orange indicator corresponds to  $HCO_3^{-}$  obtained from  $CO_3^{2-}$  and  $HCO_3^{-}$  originally present in the water sample.
- v) P = M; indicates only  $OH^-$  ions are present.

## Specifications for Drinking water: (BIS and WHO standards):

Water is an incredibly important aspect of our daily life. Every day we drink water, cook with water, bath in water and participate in many activities involving water. It is essential for all dimensions of life. The water may contain many harmful constituents, there are no universally recognized and accepted international standard for drinking water.

Many developed and developing countries specify standards to be applied in their own. The physical parameters (temperature and turbidity) of water have significant influence on aquatic life. The chemical parameters (PH, TDS, total alkalinity, total hardness, chlorides, phosphates and nitrates) of water before attempting to study the animals and plants living in it.

**TEMPERATURE:** Temperature measurements are very useful in understanding the trend of physical, chemical and biological activities which are enhanced/ retarded by the variation of temperature. The WHO (1993) did not recommend any definite temperature value for drinking water. A temperature of about 40°C is permissible limit for drinking water (BIS 1991).

**TURBIDITY:** The acceptable limit of BIS (1991) and recommendation is 10 NTU. The WHO recommendation of turbidity is 5-25 NTU.

NTU=Nephelometric Turbidity Unit

**PH:** The pH was found ranging between 7.0 and 8.3. pH as such has no adverse effect on health, however a lower value below 4 will produce sour taste; and a higher value above 8.5 an alkaline taste. pH range from 6.5 to 7.5 is most favorable for production in a water body. The WHO and BIS recommendation of pH is 6.5- 8.5. In the current study the pH range is a safe for fish production and drinking water.

**Total dissolved salts (TDS):** The BIS (1991) has set desirable limit of TDS value to be 500 mg/l in potable water. However the permissible limit is 2000 mg/l in the absence of any alternative source in water. According to WHO (1993), the standard permissible limit for TDS is 1000 mg/l. Water at a TDS level of above 500 mg/l is unsuitable for flora and tastes unpleasant to drink.

**Alkalinity:** BIS has set the desirable limit of alkalinity for drinking water to be 200 mg/l and the permissible value has been prescribed to be 600 mg/l in the absence of any alternative source; however according to WHO (1993) it is to be 200 mg/l.

**Total hardness:** The standard permissible limit of total hardness value of drinking water set by BIS (1991) is 300 mg/l. According to WHO (1993) its value is 500 mg/l.

**Chlorides:** BIS (1991) have set a desirable limit of chloride in drinking water to be 250 mg/l and permissible value has been prescribed to be 1000 mg/l in the absence of any alternative source. Whereas according to WHO (1993), it is 250 mg/l.

**Phosphates:** There is no BIS standard permissible limit for phosphate for drinking water, while WHO (1993) has fixed it to be 0.1 mg/l. If level becomes too high, plant growth can accelerate resulting in the dense growth of algae and plants in the water body.

**Nitrate:** WHO (1993) has fixed the value of nitrate in drinking water to be 50 mg/l and BIS (1991) has set a desirable limit of nitrate in drinking water to be 45 mg/l and permissible value has been prescribed to be 100 mg/l in the absence of any alternative source.

#### COMPARISION OF DIFFERENT PHYSICO-CHEMICAL PARAMETERS

#### WITH SUGGESTED BY WHO AND BIS FOR DRINKING WATER SUPPLY

S.NO.	Parameters	Range Values	Limit for d	Permissible rinking water uality
	Temperature		BIS	WHO
1	(°C)	16.0 – 27.8	40	-
2	Turbidity (NTU)	6 -22	10	5-25
3	рН	7.0 - 8.3	6.5-8.5	6.5-8.5
4	Total Dissolved solids (mg/l)	290 - 328	500	1000
5	Total Alkalinity (mg/l)	138 - 198	200	200
6	Total Hardness (mg/l)	132- 187	300	500
7	Chloride (mg/l)	20 - 64	250	250
8	Phosphate (mg/l)	0.021 - 0.071	-	0.1
9	Nitrate (mg/l)	0.21 – 1.28	45	50

## PROBLEMS ON LIME-SODA PROCESS

-	(1) calculate the amount	of 12me and soda resulted for the
	10 000 18408	at you water containing the
		PEN LIME : CILCON = 50 Myh . Calegram
	1 0287 05	Time & source le dois cope of acres s
	R& 40/821 kg, and soda	18 R& 90 Per Kg.
	R8 40 (BB) kg, and source	7 12 1 11 0
į	.,-	

* convol. the H.C.S	HCS Pro text	100 Molemal Wolemal ws of (aws	equivalents.  50  11.1 × 100 = 10
3) Mg 504	12 mg/L	120	$\frac{12 \times 100}{120} = 10$
4) NaHco3 5) 5918ca	4·25 mg从 10 mg从	84	Does not yequire treatment.

Amount lime required:

$$= \frac{74}{100} \left( 50 + 10 + 4.31 \right)$$

$$= \frac{74}{100} \times 64.31 = 47.589 \text{ mg} \right) \text{ little}$$

$$= \frac{74}{100} \times 64.31 = 47.589 \text{ mg} \right) \text{ little}$$

amount of time required for 10,000 letres of water = 47.589 × 10,000 = 4,75890 mgs 10,000 letres

= 475.89 3ms 10,000 letre

= 0.4759 Kg] 10,000 letre.

Total cost of Irme = 40 × 0.4759 = 19.036 / = R8 = 19.04

```
Amount of soda required:
                106 ( cad2+ Mg SO4 - 1 NaHcO3 hardness convolted
                                                 to cacez equivale
              = 106 (10+10 + 4.31)
              = 106 × 15.69
               = 16.6314 mg) let
         soda required for 10,000 tetres of water
                     = 16.6314 × 10,000
                      = 166314 mg | 10,000 GHres
                       = 0. 1663 Kg (10,000 letrey.
 cost of goda = R$ 90-1- per kg
  Total cost of soda = 0.1663×90 = R$ 14.97/
Total expenditure on 12me & soda = 19.04 + 14.97 = R8 34.0/
 (3) calculate the amount of time and sala regulard of
  stor tensing 10,000 letters of water containing the following salk
 per little.
   calleon = 162 mg casey = 136 mg
                                      Mgds = 95 mg and
  Nad = 56-1 mg. purity of time is 13% and 20da is 99%.
                                          caces equivalent of
                           Amout of
    H.cs.
                 MIW
(1) ca(Hery)
                 162
                                           162×100 = 100mg/1
                              162
                                            136 ×100 = 100 mg/2
                        136
(2)
     Casal 136
                                              136
                                            95 X 100 = 95 mg/
              95
                              95
     Made
(3)
               Tross not require time soda treatment.
      Nacl
0
 Amount of time required:
      74 (Temporary cate hardness + permanent Mgtz Handness)
                                    12 12 17 17 17
       74 (100 + 100) = 74 x 200 = 148 mg/l
    Amount of time required for the treatment 10,000 let reg
                           = 14-8 × 10,000 = 1A-8,0000 mgx 10,000l
                         = 1,48,000 = 14-80 gm/10,000l
                           =\frac{1480}{1000} = 1.48 \text{ Kg} / 10,000l
       :. Purity of time is 93%
           Total amount of lime regulized = 1.48 ×100
                                       = 1.59 kg / 10,000l
```

Amount of soda required:

 $= \frac{106}{100} \left( 100 + 100 \right) = \frac{106 \times 200}{100} = 212 \text{ mg/l}$ 

treatment of 10,000 lit = 212 × 10000

= 2.12 kg | 10,000lit

purity of soda is 99%.

-. Total amount of soda required

= 2-14 Kg / 10,000l

A water sample gave the following constituents on analytis (B)

mgs (it. My (Acos) = 73

ca 304 - 68

Mgds = 95

Mg 304 - 12

ca (4 cog) = 81

Nacl - 4.8 (No hardness)

calculate the cost of the chemicals required for saftening 20,000 litres of water. The purity of time and soda are 95% and 90%. The cept per looky . Each of time and one Rx 75 and Rx 2480 respectively.

ANS: Rs: 3:04 +

Example 6.4

A water sample contains the following impurities:  $Ca^{2+} = 20$  ppm,  $Mg^{2+} = 18$  ppm,  $HCO_3^- = 180$  ppm, and  $SO_4^{2-} = 24$  ppm. Calculate the amount of lime and soda needed for softening.

Solution
$$CaCO_{3} \text{ equivalent of } Ca^{2+} = \left[20 \times \frac{50}{20}\right] \text{ mg/l} = 50 \text{ ppm}$$

$$CaCO_{3} \text{ equivalent of } Mg^{2+} = \left[18 \times \frac{50}{12}\right] \text{ mg/l} = 75 \text{ ppm}$$

$$CaCO_{3} \text{ equivalent of } HCO_{3}^{-} = \left[180 \times \frac{50}{61}\right] \text{ mg/l} = 147.54 \text{ ppm}$$

$$Lime \text{ requirement for sofening} = \frac{74}{100} \left[\text{Prem. } Mg^{2+} + HCO_{3}^{-}\right]$$

$$= \frac{74}{100} \left(75 + 147.54\right)$$

$$= 164.68 \text{ mg/l}$$

$$Soda \text{ requirement for sofening} = \frac{106}{100} \left[\text{Prem. } \{Ca^{2+} + Mg^{2+}\} - HCO_{3}^{-}\right]$$

$$= \frac{106}{100} \left(50 + 75 - 147.54\right)$$

$$= -22.54 \text{ mg/l}$$

The value is negative. This implies that soda requirement is nil.

### Example 8.5

Calculate the amount of lime and soda needed for softening 50000 l of water containing following salts per liter:  $Ca(HCO_3)_2 = 10$  mg, Mg ( $HCO_3$ )<sub>2</sub> = 15 mg,  $MgSO_4 = 20$  mg,  $CaSO_4 = 18$  mg,  $CaCl_2 = 8$  mg, and NaCl = 4 mg.

### Solution

Lime requirement for sofening

$$= \frac{74}{100} [\text{Temp. } \{\text{Ca}^{2+} + 2 \times \text{Mg}^{2+}\} + \text{Perm. } (\text{Mg}^{2+})] \times \text{Volume of water}$$

$$= \frac{74}{100} \left[ 10 \times \frac{50}{81} + 2 \times 15 \times \frac{50}{73} + 20 \times \frac{50}{60} \right] \times 50000 \text{ mg}$$

$$= 1605430 \text{ mg} = 1.605 \text{ kg}$$

Soda requirement for sofening = 
$$\frac{106}{100}$$
 [Temp.  $\{Ca^{2+} + Mg^{2+}\} \times H^{+}\}$ ] × Volume of water
$$= \frac{106}{100} \left[ 20 \times \frac{50}{60} + 18 \times \frac{50}{68} + 8 \times \frac{50}{55.5} \right] \times 50,000 \text{ mg}$$

$$= 1967360 \text{ mg} = 1.967 \text{ kg}$$

Example 8.6

Example 2 (alculate the amount of lime and soda needed for softening 50000 l of water with the following (alculate the amount of sodium aluminate as coagulant. Analysis result:  $Ca^{2+} = 100$ ,  $Mg^{2+} = 35$ ,  $O_2 = 50$ ,  $HCO_3 = 250$ ,  $H^+ = 2$  ppm.

Solution

Lime requirement for sofening (mg/l) = 
$$\frac{74}{100}$$
 [Mg<sup>2+</sup> + CO<sub>2</sub> + H<sup>+</sup> + HCO<sub>3</sub> - NaAlO<sub>2</sub>]

Soda requirment for sofening (mg/l) = 
$$\frac{106}{100}$$
 [Ca<sup>2+</sup> + Mg<sup>2+</sup> + H<sup>+</sup> - HCO<sub>3</sub>]

Substitute the values after converting into CaCO<sub>3</sub> equivalent as per Example 8.5.

### Problem 3.11 :

A sample of water contains the following dissolved salts in mgs/litre. CanO<sub>4</sub> = 6.8, MgCO<sub>4</sub> = 8.4, Al<sub>4</sub>(SO<sub>4</sub>)<sub>4</sub> = 34.2, CO<sub>2</sub> = 4.4, HCl = 3.65. Calculate the amount of lime and soda required for the treatment of 5000 litres of water. The purity of lime 80% and soda is 90%, 10% of excess chemicals were added.

Not: Conversion of the amounts of hardness causing salts into CaCO, equivalents.

7. 70.	Hardness causing salt (H.C.S.)	Amount of H.C.S. mg/l	Mol. wt. of H.C.S.	CaCO, equivalents
1 1	CaSO,	6.8	136	$\frac{6.8 \times 100}{136} = 5$
2.	месо,	8.4	84	$\frac{8.4 \times 100}{84} = 10$
3.	AI,(SO,),	34.2	342	$\frac{34.2 \times 100}{342} = 10$
4.	co,	4.4	44	$\frac{4.4 \times 100}{44} = 10$
5.	нсі	3.65	36.5	$\frac{3.65 \times 100}{36.5} = 10$

Amount of lime required =  $\frac{74}{100} \left[ (2 \times \text{MgCO}_3) + (3 \times \text{Al}_2(\text{SO}_4)_3) + \text{CO}_2 + \frac{1}{2} \text{HCl hardness in} \right]$ 

terms of CaCO3 equivalents

$$= \frac{74}{100} \times \left( (2 \times 10) + (3 \times 10) + 10 + \frac{1}{2} \times 10 \right) = \frac{74}{100} \times 65 = 48.1 \text{ mg/l}$$

Lime required for 5,000 litres of water =  $48.1 \times 5,000$ 

= 240500 mg/5000 lit. = 0.2405 kg/5000 lit.

Purity of lime = 80%.

Out of 100 kgs of raw time used only 80 kgs is pure lime

? raw lime should be used if 0.2405 kg of pure lime required

the amount of raw time required =  $\frac{0.2405 \times 100}{80}$  = 0.3006 kg/5000 lit.

Total amount of lime along with 10% excess chemicals =  $\frac{0.3006 \times 110}{100}$ = 0.33066 kg/5000 litre

Amount of soda required

$$= \frac{106}{100} (\text{CaSO}_4 + (3 \times \text{Al}_2(\text{SO}_4)_3) + \frac{1}{2} \text{HCl hardness in CaCO}_3 \text{ equivalents}$$

$$= \frac{106}{100} [5 + (3 \times 10) + 5]$$

$$= \frac{106}{100} \times 40 = 42.4 \text{ mg/l}.$$

Lime required for 5000 litres of water =  $42.4 \times 5000$ 

= 212000 mg/1000 lit = 0.212 kg/5000 li

Purity of soda = 90%

Amount of raw soda to be required =  $\frac{0.212 \times 100}{90}$  = 0.2356 kg/5000 litre

10% excess soda is added

Total amount of soda along with 10% excess chemicals

$$= \frac{0.2356 \times 110}{100} = 0.2591 \text{ kg/5000 litre}$$

### Problem 3.12:

Calculate the amount of lime and soda required for softening 10,000 litres of water containing the following salts per litre.

 $Ca(HCO_3)_2 = 162$  mgs,  $CaSO_4 = 136$  mgs,  $MgCl_2 = 95$  mgs and NaCl = 56.1 mgs. Purity of lime is 93% and that of soda is 99%. (BIT Mesra I.B.E. 1993)

Sol:

S.No.	Hardness causing salt (H.C.S.)	Amt. of H.C.S. in mg/l	Mol.wt.of H.C.S.	CaCO <sub>3</sub> equivalent of H.C.S.
1.	Ca(HCO <sub>3</sub> ) <sub>2</sub>	162	162	$\frac{162 \times 100}{162} = 100  \text{mgs//}$
2.	CaSO <sub>4</sub>	136	136	$\frac{136 \times 100}{136} = 100 \mathrm{mgs/l}$
3.	MgCl <sub>2</sub>	95	95	$\frac{95 \times 100}{95} = 100 \text{mgs//}$
4.	NaCl	Does	not require lin	ne soda treatment

Amt. of lime required =  $\frac{74}{100}$  (temporary hardness of Ca(HCO<sub>3</sub>)<sub>2</sub> + permanent hardness of MgCl<sub>2</sub>)

$$=\frac{74}{100}\times200 = 148 \text{ mg/l}.$$

Amount of lime required for the treatment 10,000 litres

= 
$$148 \times 10000 = 14,80,000 \text{ mgs/}10000 l$$
  
=  $\frac{1480000}{1000} = 1480 \text{ gms/}10000 l$   
=  $\frac{1480}{1000} = 1.48 \text{ kg/}10000 l$ 

Purity of lime is 93%

Total amount of lime required =  $\frac{1.48 \times 100}{93} = 1.59 \text{ kg}/10,000 \text{ l}$ 

Amount of soda required =  $\frac{106}{100} \times 200 = 212 \,\text{mgs/}l$ 

for the treatment of 10,000 litres =  $\frac{212 \times 10000}{1000 \times 1000} = 2.12 \text{ kg}/10000 l$ 

Purity of soda is 99%

Total amount of soda required =  $\frac{2.12 \times 100}{99}$  = 2.14 kg/10,000 *l* 

problem : 3.13

A water works has to supply 1000 litres of water, that contains  $Mg^{2+}$  36 ppm  $_{HCO_3^-} = 18.3$  ppm and  $H^+ = 1.5$  ppm. Calculate the amount of lime and soda required  $_{50}$ !

S.No.	(H.C.S)	Amt. in ppm	Mol.wt.	CaCO <sub>3</sub> equivalents
1.	Mg <sup>2+</sup>	36	24	$\frac{36 \times 100}{24} = 150 \text{ ppm}$
2.	HCO <sub>3</sub>	18.3	122*	$\frac{18.3 \times 100}{122} = 15  \text{ppm}$
3.	H <sup>+</sup>	1.5	2*	$\frac{1.5\times100}{2} = 75 \mathrm{ppm}$

Amount of lime required 
$$= \frac{74}{100} \left[ Mg^{2+} + HCO_3^- + H^+ \right]$$
$$= \frac{74}{100} (150 + 15 + 75)$$
$$= \frac{74}{100} (240) = 177.6 \,\text{mg/l}$$

Lime required for the treatment of 1000 litres of water

$$= \frac{177.6 \times 1000}{1000 \times 1000} = 0.1776 \,\mathrm{kg}$$

Soda required for the treatment = 
$$\frac{106}{100} (Mg^{2+} + H^{+} - HCO_{3}^{-})$$
  
=  $\frac{106}{100} (150 + 75 - 15)$   
=  $\frac{106}{100} \times 210 = 222.6 \text{ mg/}l$ 

Amount of soda required for the treatment of 1000 litres of water

$$=\frac{222.6\times1000}{1000\times1000}=0.2226\,\mathrm{kg}$$

[\* Note:  $\frac{1}{2}$  HCO<sub>3</sub> and  $\frac{1}{2}$  H<sup>+</sup> hardness must be considered. That is why the molecular

weights of HCO<sub>3</sub> and H<sup>+</sup> are taken as 122 and 2 instead of 61 and 1 respectively Problem 3.14:

Analysis of water gave the following results:  $H_2SO_4 = 196 \text{mg/l}$ ,  $MgSO_4 = 24 \text{mg/l}$ CaSO<sub>4</sub> = 272 mg/l and NaCl = 25 mg/l. Water is to be supplied to the/town of the population one lakh only. The daily consumption of water is 50 litres per head Calculate the cost of lime and soda required for softening of hardwater for town for the month April 2002, if the cost of lime is Rs. 5.00 per kg and cost of soda is Rs. 8.00per kg. (MREC 2002)

Sol:

S.No.	(H.C.S)	Amt. in mg/l	Mol. wt.	CaCO <sub>3</sub> equivalents	
1.	H <sub>2</sub> SO <sub>4</sub>	196	98	$\frac{196 \times 100}{98} = 200$	
2.	MgSO <sub>4</sub>	24	120	$\frac{24 \times 100}{120} = 20$	
3.	Cache	272	136	$\frac{272 \times 100}{136} = 200$	
4.	NaCl	No treatment with lime & soda required			

Amount of lime required =  $\frac{74}{100}$  (Hardness of  $H_2SO_4 + MgSO_4$  in terms of CaCO<sub>3</sub> equivalents)  $= \frac{74}{100} \times 220 = 162.8 \,\text{mg/l}$ 

$$=\frac{74}{100}\times220=162.8\,\mathrm{mg/l}$$

Daily consumption of water = 50 l per head. Population of the town = 1,00,000

Lime required for the treatment for the population =  $\frac{162.8 \times 50 \times 1,00,000}{1000 \times 1000} = 814 \text{kg}$ 

For April 2002 (30 days) total amount of lime required  $= 814 \times 30 = 24420 \text{ kgs}.$ 

Given cost of lime = 5/- per kg.

Total cost of lime in Rs. =  $24420 \times 5 = 1,22,100/$ -

Amount of soda required =  $\frac{106}{100}$  (hardness of H<sub>2</sub>SO<sub>4</sub> + MgSO<sub>4</sub> + Cast in terms of CaCO<sub>3</sub> equivalents)

$$=\frac{106}{100}(420)=445.2\,\mathrm{mg/}l$$

Soda required for the treatment for the population

$$= \frac{445.2 \times 50 \times 1,00,000}{1000 \times 1000} = 2226 \,\mathrm{kg}.$$

For April 2002 (30 days) the amount of soda required =  $2226 \times 30 = 66780$  kg Cost of soda = Rs 8/- per kg.

Total cost of soda =  $66780 \times 8 = Rs.5,34,240/$ 

### Problem 3.15:

Calculate the amount of lime and soda required for softening 10,000 litres of water, which was analysed as follows:

Analysis of raw water:

 $Ca^{2+} = 380 \text{ ppm}$ , Dissolved  $CO_2 = 120 \text{ ppm}$ 

 $Mg^{2+} = 144 \text{ ppm}, \text{ FeSO}_4 .7H_2O = 278 \text{ ppm}$ 

 $HCO_3^- = 1500 \text{ ppm}$ 

Analysis of treated water

$$OH^- = 34 \text{ ppm}$$
,  $CO_3^{2-} = 32 \text{ppm}$ .

(MREC 2000)

S.No.	H.C.S	Amt. (ppm)	At.wt/mol. wt	CaCO <sub>3</sub> equivalents
Raw water		: · · · · · · · · · · · · · · · · · · ·	· • • • • •	
$\overline{}$	a <sup>2+</sup>	380	40	$\frac{380 \times 100}{40} = 950$
2. N	Ig <sup>2+</sup>	144	24	$\frac{144 \times 100}{24} = 600$

		♥ WAIER AND IIS II	SEALWENT Y	
S.No.	H.C.S	Amt. (ppm)	At.wt/mol. wt	CaCO <sub>3</sub> equivalent
Raw w	ater		L	
3.	HCO <sub>3</sub>	1500	122	$\frac{1500 \times 100}{122} = 1229$
4.	Dissolved CO <sub>2</sub>	120	44	$\frac{1420 \times 100}{44} = 272.7$
5.	FeSO <sub>4</sub> .7H <sub>2</sub> O	278	278	$\frac{278 \times 100}{278} = 100$
reated	water		2	
6.	OH-	34	34	$\frac{34 \times 100}{34} = 100$
7.	CO <sub>3</sub> <sup>2-</sup>	32	60	32×100

Amount of lime required for raw water =  $\frac{74}{100}$  (Hardness of Mg<sup>2+</sup> + HCO<sub>3</sub> + CO<sub>2</sub>+ FeSO<sub>4</sub>.7H<sub>2</sub>O converted in terms of CaCO<sub>3</sub> equivalents)

$$= \frac{74}{100} (600 + 1229.5 + 272.7 + 100) = 1629.6 \text{ mg/l}.$$

Amt. of lime required for 10,000 litres of raw water

$$= \frac{1629.6 \times 10,000}{1000 \times 1000} = 16.296 \,\mathrm{kg}/10,000 \,l$$

Amt. of soda required for raw water =  $\frac{106}{100}$  (Hardness of Ca<sup>2+</sup> + Mg<sup>2+</sup> + FeSO<sub>4</sub>.7H<sub>2</sub>0 - HCO<sub>3</sub> in terms of CaCO<sub>3</sub> equivalents)  $=\frac{106}{100}(950+600+100-1229.5)$  $= \frac{106}{100} \times 420.5 = 445.73 \,\mathrm{mg/l}$ 

Amt. of soda required for 10,000 litres of raw water

$$= \frac{445.73 \times 10000}{1000 \times 1000} = 4.4573 \,\mathrm{kg}/10,000 \,l$$

Amount of lime required for excess OH- in treated water

$$= \frac{74}{100} \times 100 = 74 \,\mathrm{mg/l}$$

Amount of lime required for excess OH- in 10,0001 of treated water

$$=\frac{74\times10000}{1000\times1000}=7.4\,\mathrm{kg}/10,000\,l$$

Total amount of lime required for the treatment of raw + treated 10,000 litres of water = 16.296 + 7.4 = 23.696 kg/10,000 l.

Amt. of soda required for excess of OH- and HCO3 in treated water

$$= \frac{106}{100} (100 + 53.3) = \frac{106}{100} \times 153.3 = 162.498 \,\mathrm{mg/l}$$

Amount of soda required for excess OH- and HCO3 in 10,000 l of treated water

$$= \frac{162.498 \times 10000}{1000 \times 1000} = 1.625 \,\mathrm{kg}/10,000 \,l$$

Total amount of soda required for the treatment raw + treated 10,000 l of water = 4.4573 + 1.625 = 6.0823 kg/10000 l.

### Problems for Practise:

1. A water sample gave the following constituents on analysis in mgs/litre. Mg(HCO<sub>3</sub>)<sub>2</sub> = 73, CaSO<sub>4</sub> = 68, MgCl<sub>2</sub> = 95, MgSO<sub>4</sub>=12, Ca(HCO<sub>3</sub>)<sub>2</sub> = 81, NaCl = 4.8. Calculate the cost of the chemicals required for softening 20,000 litres of water. The purity of lime and soda are 95% and 90%. The costs per 100 kg. each of lime and soda are Rs. 75 and Rs. 2480 respectively. (Nagpur IBE 1997)

[Ans: Rs. 
$$3.04 + 93.50 = 96.54/-$$
]

2. Calculate the amount of lime and soda required to soften 1 million litres of a water sample containing the following dissolved salts (calcium permanent hardness) Ca<sup>2+</sup> = 20 ppm, Mg<sup>2+</sup> = 24 ppm, HCO<sub>3</sub> = 150 ppm, and CO<sub>2</sub> = 30 ppm. The purity of lime is 87% and soda is 91%. 10% of excess chemicals were added.

[Ans: time = 
$$272.46 \text{ kg}$$
, Soda =  $34.6 \text{ kg}$ ]

#### ♦ WATER AND ITS TREATMENT

3. A sample of raw water contains the following dissolved salts.  $Ca(HCO_3)_2 = 8.1$ ppm,  $Mg(HCO_3)_2 = 14.6$  ppm,  $HCO_3 = 91.5$  ppm,  $CaCl_2 = 33.3$  ppm,  $MgCl_2 = 38$ ppm,  $Al_2(SO_4)_3 = 17.1$  ppm. Calculate amounts of lime (90% pure) and soda (JNTU 1988) (98% pure) for the treatment of 10,00,000 litres of water.

[Ans : Lime 214.05 kg, Soda = 10.82 kg]

- 4. Calculate the quantity of lime and soda required for softening 60,000 litres of water containing  $CO_2 = 20$  ppm,  $CaCO_3 = 45.45$  ppm, Mg  $(HCO_3)_2 = 25$  ppm, HCl = 8.4ppm,  $Al_2 (SO_4)_3 = 40$  ppm and  $MgCl_2 = 12$  ppm. 10% of excess chemicals were [Ans : Lime = 14.07 kg, Soda = 11.39 kg] added.
- 5. A water works has to supply 1 m<sup>3</sup>/s of water. The raw water contain Mg(HCO<sub>1</sub>), = 219 ppm,  $Mg^{2+} = 36$  ppm,  $HCO_3 = 18.3$  ppm and  $H^+ = 1.5$  ppm. Calculate the cost of treating water per day, if lime (90% pure) and soda (95% pure) has the cost Rs. 500/- per tonne and Rs. 7000/- per tonne respectively.
  - (Hint : Amount of water purified per day = 1  $\text{m}^3/\text{s} \times 60 \times 60 \times 24 \text{ s/day} = 8.64 \times 10^4 \text{m}^3 / \text{day}$ =  $8.64 \times 10^7$  litres / day.1 tonne =  $10^3$  kg =  $(10^9$  mgs))

[Ans: Rs.1,60,894/-]

Calculate the amount of lime and soda required for softening 10,000 litres of hard water containing Ca(HCO<sub>3</sub>)<sub>2</sub> = 81 mgs, CaSO<sub>4</sub> = 13.6 mgs, MgCl<sub>2</sub> = 95 mgs and NaCl = 56 mgs. Purity is 93% and Soda is 99%.

[Ans : Lime = 1.19 kg, Soda = 1.178 kg

# Module-II Molecular structure and Theories of Bonding

# II. Molecular structure and Theories of Bonding

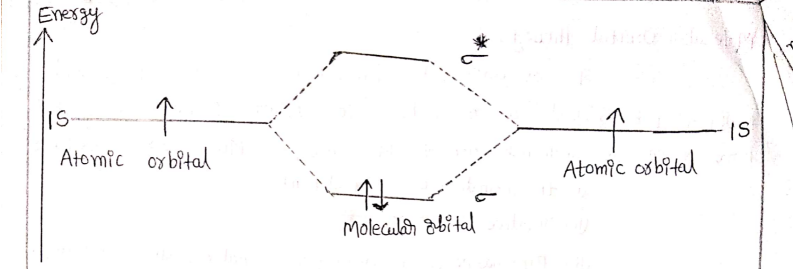
## Molecular orbital theory:

To overcome the limitations of VBT, Hund and Mulliken put forward a new theory to explain formation of covalent bonds known as molecular orbital (MO) theory. MO theory to explain:

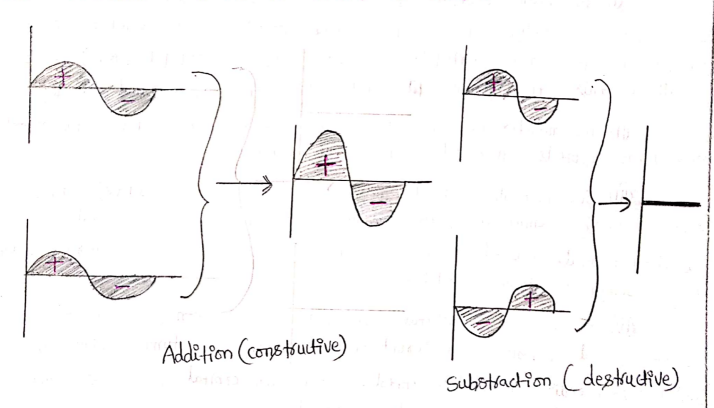
- 1) the formation of chemical bond
- (ii) relative bond strengths
- (iii) paramagnetic (or) diamagnetic nature, etc of molecules.

Salient features of MO theory are:

- The atomic orbitals (with same energy (or) nearly same energy and appropriate symmetry under (mean combination) of the combing atoms overlap to form new orbitals, called molecular orbitals (M.O). As a result of this, the atomic porbitals lose their individual identity.
- (i) The number of molecular orbitals formed is equal to the number of atomic orbitals participating in linear combination.
- atomic oxbitals remain in phase overlap of atomic oxbitals is called "Bonding molecular oxbital". Such an oxbital possesses less energy than the combining atomic oxbitals.
- (iv) The molecular expital formed by substraction (destructive) (when atomic orbitals remain in different phase) everlap of atomic expitals is called "Anti bonding molecular expital". Such an expital possesses more energy than the combining atomic dibitals.
- in unlike atomic orbitals, molecular abital is polycentric region in space, defined by its size and shape. Thus, in a molecular orbital, electrons are revolving in the field of more than one nucleus.
- oxbitals from which they are generated.
- (VII) Filling of molecular abitals follows the same rules used for filling of atomic orbitals.



Linear Combination of Atomic orbitals (LCAO):



According to wave mechanics the atomic oxbitals can be expressed by wave functions (4/8), which represent by the amplitude of electron waves. Their values can be calculated from the solutions of Schrodinger's wave equation. However, it is difficult to solve the above exactly for many electron systems. Therefore, an approximate method known as Linear combination of atomic oxbitals (LCAO) has been adapted.

According to this method, the molecular orbitals are formed by the Linear combination (addition (constructive) or substraction (destructive) of atomic orbitals of the constituent atoms of the

Let us consider this method to simplest homonuclean diatomic molecule, hydrogen which consists of two hydrogen atoms represented by HA and HB. The atomic orbitals of these atoms are represented by the wave functions of and UB. when these atoms approach each other come two possibilities.

(1) Mole auton Erbital is formed by the addition of wave functions atomic orbitals. It can be represented by 4(MO) = 4A+4B - (constructive interaction)

The M.O formed is called "Bonding molecular orbital". It lowers the energy than that constituent overlapping atomic orbitals. The difference in energy between combining atomic orbitals and the bond molecular Bubital formed, is called stabilization emergy. Thus, bonding m.o stabilizes the molecule.

# characteristics of bonding molecular orbital:

(1) It possesses lower emergy than that of the combining the atomic orbitals. (ii) It possesses high electron-density in the

region between the two nuclei

(iii) It imports stability to the molecule.

(V) Every electron in it makes contribution to the

attraction of two combining atoms.

(i) If is only formed, when the lobes of the combining atomic orbitals possess same sign.

Molecular oxbital is formed by the substraction of wave functions of atomic orbitals. It can be represented by

 $\Psi_{(MO)}^* = \Psi_A - \Psi_B - (destructive interaction)$ 

The M.O formed is called anti-bonding molecular orbital. This type of M.O corresponds to higher energy state. The difference in energy between the anti-bonding molecular orbital and the combining atomic orbitals, excalled de stabilization emergy. Thus, antibonding M.O destabilizes the molecule.

characteristics of anti-bonding molecular orbital:

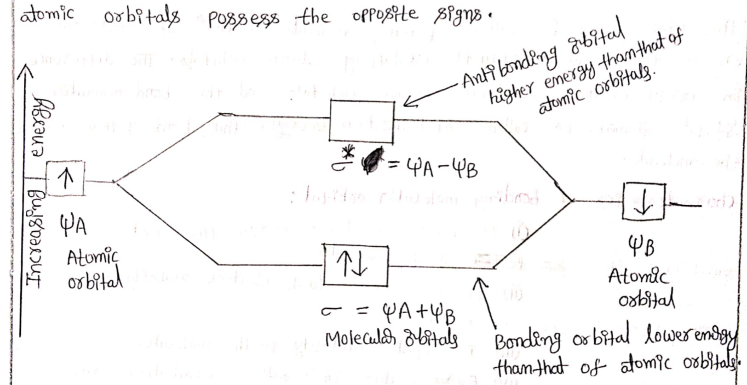
domic orbitals. (ii) It possesses low-electron density in the region between the two nucles.

in It imports instability to the molecule

(iv) Every electron in it makes contribution to the repulsion

of two combining atoms.

is It is formed, when the lobes of the combining



Formation of bonding (=) and antibonding (=\*) molecular substals by the linear combination of atomic orbitals (PA and PB centered on two atoms A and B respectively.

Significance of bonding and anti-bonding molecular oxbital:

The probability density in bonding and anti-bonding molecular subitals, the bonding molecular subital filmation (4) greater than that of anti-bonding molecular subital (4\*2) formation. This is because we known that probability is gives by the secure of amplitude (4), therefore

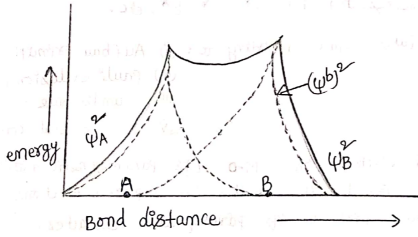
$$\psi_{(M,0)}^{2} = (\psi_{A} + \psi_{B})^{2} = \psi_{A}^{2} + \psi_{B}^{2} + 2 \psi_{A} \psi_{B}$$

$$\psi_{(M,0)}^{*2} = (\psi_{A} - \psi_{B})^{2} = \psi_{A}^{2} + \psi_{B}^{2} - 2 \psi_{A} \psi_{B}$$

Taking sociates of the bonding MO wave function (4b), we get  $(4b)^2 = 4^2 + 4^2 + 24A4B$ 

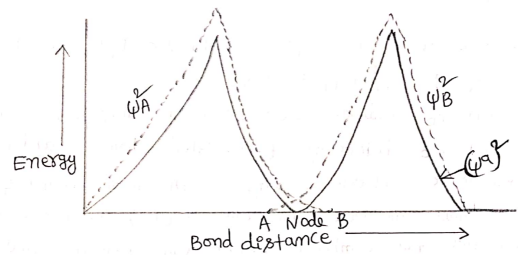
The terms who and yis indicate electronic change densities of cave functions (yA and yB belonging to isolated atoms A and B, respectively.

(yb) is the electronic change density of the wave function yb, i.e., the bonding M.O. Thus, the more is the value of (yb), the more is the change density between the two combining nuclei. Again from the equation is also evident that (yb) > ya + yB by a term, a ya yB. This term a ya yB results from the overlap of Ao's and is called overlap integral. Larger is the overlap integral more is the change density between the nuclei and stable bond is formed. The plot of electron change density as a function of inter-nuclear distance for bonding M.O. is shown below.



: Again, taking square of the anti-bonding M.O wave-function ( $\psi^{a}$ )
we get  $(\psi^{a})^{2} = \psi^{a}_{A} + \psi^{b}_{B} - 2\psi_{B}\psi_{A}$ 

It is evident from the above equation that  $(\psi^b)^2 < \psi^2 + \psi^3 = by$  a term  $2 \psi_A \psi_B$ . Thus, in anti-bonding M.O., there is less thange density between the nuclei. It is, therefore, the energy of the anti-bonding m.o. is more than the sum of the energies of the two interacting atoms A and B. The plot of electron change density is shown below.



- a) M.O. gives the electron probability distribution around a group of mudei Just as an A.O. gives the electron probability distribution around the mudeus.
- (b) the shape of the M.O formed depends on the type of the combining atomic orbitals.
- antibonding M.O are represented by -, TT, 8 etc., where as
- d) The filling of M.O takes place according to: in Aufbau primiple
  - (i) pauli exclusion primaple
  - (III) Hund's rule
  - (iv) If a molecule contains

one (or) more unpaised electrons in it M.O, it is paramagnetic nature; otherwise diamagnetic. Greater the number of unpaired electrons in M.O's of a substance, high is its paramagnetic character.

## Conditions for the formation of molecular orbitals:

Any two atomic exbitals on combination donot form molecular oxbitals. In fact, there are certain lemitations to the combination of atomic oxbitals. They are:

This is true only for homonuclean diatomic amolecules.

This is true only for homonuclean diatomic amolecules.

The combining atomic 8451tals must be overlap to the maximum extent.

Greater the extent of overlap, the greater the nude of a molecular 8451tal.

about the molecular axis.

By convention z-axis is taken as the molecular axis.

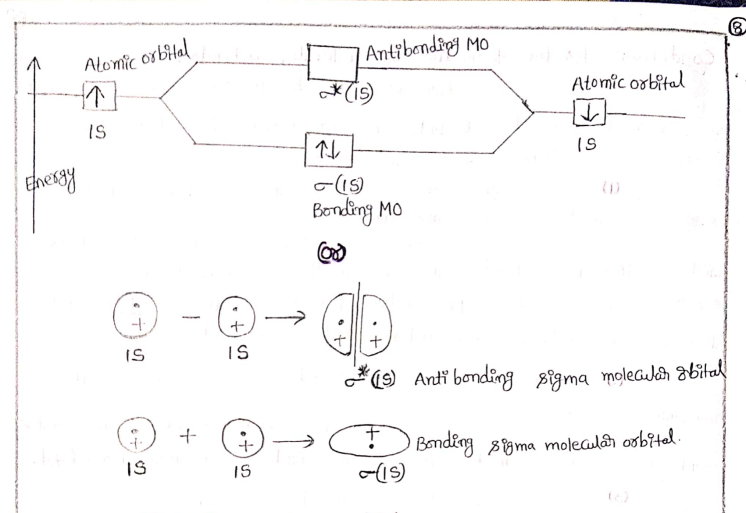
By convention z-axis is taken as the molecular axis.

Axis. It is important to note that atomic oxbitals having same (ox) nearly

the same emergy will not combine it they do not have the same symmetry. For example, are orbital of one of atom can combine with are substal of another atom but not with are (or) ary stritals because of their different symmetries.

## Formation of M.O'S:

From S-domic orbitals: when two 15 orbitals combine two new molecular dibitals are formed. one of these pertains to the bonding molecular dibital with lower energy while the other corresponds to higher energy as compared to both the atomic orbitals concerned in the process. Thus we find that the number of molecular dibitals formed from atomic orbitals is equal to the number of atomic orbitals responsible for their formation and are shown below.



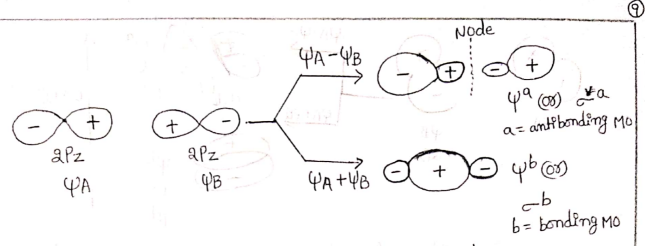
### Molecular orbitals from P-atomic orbitals:

Combination of p-oxbitals is slightly complicated.

There being three types of 2p-oxbitals namely 2px, 2py and 2pz directed along the cartesian cooldinates of, y and z, respectively. By convertion we assume that z-axis is the intermolecular axis.

(i) Combination of 2Pz atomic oxbitals: The linear combination of two 2Pz oxbitals on two atoms approach each other along the internuclear axis, the combine by the addition of electrons wave.

This results in the formation of bonding - (2Pz) oxbital while combination pubstraction of electron waves gives rise to - (2Pz). It is represented as follows.

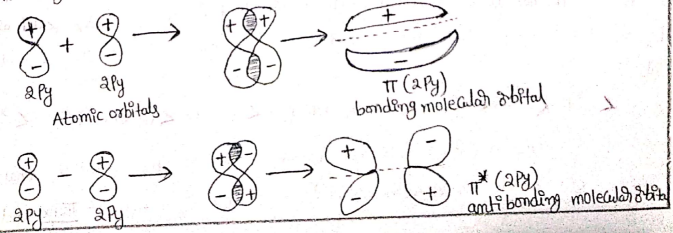


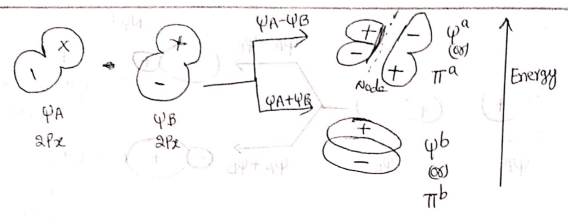
Molecular osbitals formed from 2Pz osbitals

# (1) Combination of apy and apx atomic orbitals:

when two apy and apa orbitals on two atoms are approaching in a manner that their exes one mutually parallel, they interact to give rise to the formation of molecular orbitals that are not symmetrical about the internuclear axis and are called molecular orbitals represented by TT (2Py) (08) TT (2Px) bonding orbitals. These T(2Py) and T(2Px) bonding Mo's have zero electron demosity on the plane that contains the nuclear axis (nodal plane), while the electron demosty is concentrated in two regions above and below the modal plame.

The TT\* (2Py) and TT\* (2Px) anti-bonding Mos have higher energy than their corresponding molecular and atomic orbitals. Since apy and apx atomic orbitals on an atom are degenerate (one of equal emongy), the TT molecular orbitals assising out of them Will also be degenerate le; TT (2Pg) and TT (2Px) bonding molecular osbitals possess equal energy and the same situation prevails in antibording molecular orbitals TX (2Px) And TX (2Px).





Molecular orbitals formed from 2Px and 2Py orbitals

## Energy level diagram for Molecular orbitals:

we have seen that 15 atomic orbitals on two atoms form two molecular orbitals designated as <u>-15</u> and <u>+15</u>. In the same manner, the <u>25</u> and <u>2P</u> atomic orbitals (eight atomic orbitals on two atoms) give rise to the following eight molecular orbitals.

Antibonding Mo's: #(5), #QS, #QPZ), #QPY), #QPY)

The energy levels of these molecular subitals have been determined exparimentally from spectroscopic data for homonuclear distance molecules of second row elements of parodic table.

The increasing order of energies of various molecular abitals for 02 and F2 is given below.

-15 L = 15 L = 25 L = 12 Px = 112 Py) < (1 2 Px = 112 Py)

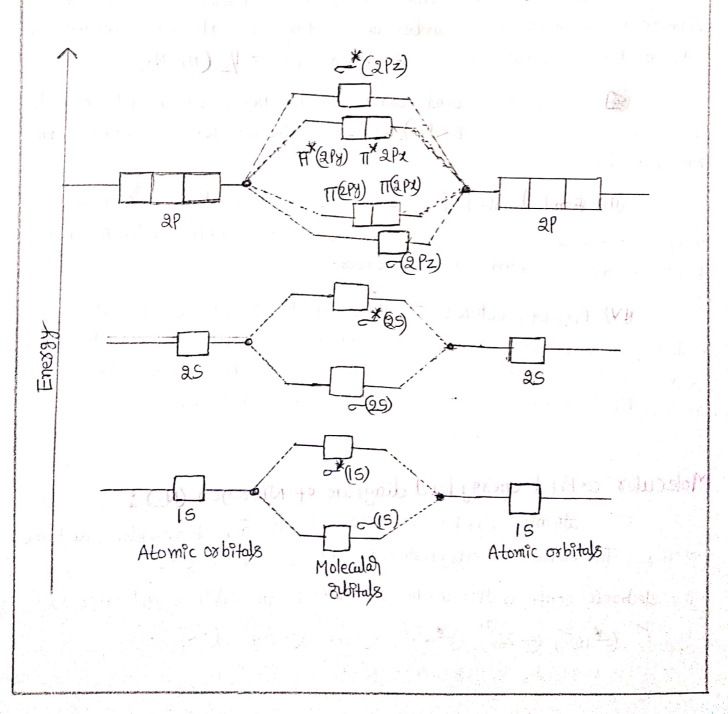
However, this sequence of energy levels of molecular abitals is not correct for the remaining molecules Lia, Be2, B2, C2, N2. For instance, it has been observed experimentally that for molecules such as B2, C2, N2 etc. the increasing order of energies of various molecular abitalis

-15 6 -15 (-25 < -25 < (Tapz=TTapy) < -2 PZ < (TTapz=TTapy) < -2 PZ < (TTapz=T

The 9mportant characteristics feature of this order 98 that the energy of -2Pz molecular orbital is higher than that of TIZPX and

TT 2 Py molecular abitals.

For diatomic molecules obtained from atom having more than one electronic shell, the inner shells do not appreciably affect the bonding and may be omitted. Such inner electrons are called non-bonding electrons. Electronic structures of simple molecules can be wisked out by feeding in the molecular inditals in the coder of increasing energy. Here also Aufball primarle feeding electrons in the increasing order of enagy of subitals repeat it self. These order shown is figure.



### Electronic configuration and molecular behaviour:

The distribution of electrons armong various molecular Sibitals is called electronic configuration of the molecule. From the electronic configuration of the molecule, It is possible to get important information about the molecule as discussed below.

- d) Stability of molecule: If Nb is the number of electrons occupying orbitals and Na the number occupying the anti-bonding subitals the (a) the molecule is stable if Nb is greater than Nb, and (b) the molecule 95 unstable 97 Nb 95 less than Na.
- (11) Bond order: Bond order (B.O) is defined as one half the difference between the number of electrons present in the bonding and anti-bonding orbitals i.e; Bond order (B.O) = 1/2 (Nb-Na)
- A positive bond order (i.e Nb>Na) means a stable molecule whele a negative (i.e Nb < Na) (08) zero (i.e Nb=Na) bond order means am umstable molecule.
- (11) Bond-length: Bond order between two atoms in a molecule may be taken as an approximate measure of the bond length. The bond length decreases as bond order Proxeases.
- (iv) Magnetic nature: It all the molecular Bloitals in a molecule are doubly occupied, the substance is diamagnetic (repelled by magnetic field). However if one (ex) more molecular orbitals are singly occupied it is paramagnetic (attracted by magnetic field). Eg. 02 molecule.

# Molecular orbital energy level diagram of Nitrogen (N2):

The atomic number of Nitrogen is 7, Homonudean, diatomic molecule. The electronic configuration of N 98 1525 253.

The electronic configuration of N2 according to molecular sibital approach is (=15), (=15), (=25), (=25), (T2PX), (T2PX), (=2P2).

Total number of electrons N+N = 7+7 = 14

Number of electrons in bonding molecular orbitals (Nb) =10

Number of electrons in anti-bonding molecular orbitals (Na) = 4

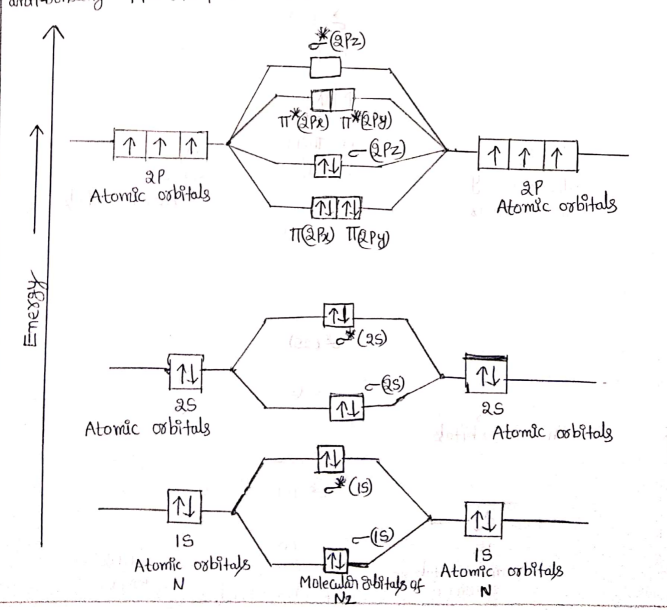
Total number of electrons = 14

Bond order =  $\frac{1}{2} \left[ \frac{10-4}{100} \right]$   $= \frac{1}{2} \left[ \frac{10-4}{100} \right]$   $= \frac{1}{2} \left[ \frac{10-4}{100} \right]$ 

bonded with three covalent bonds. i.e., a triple bond.

High value of bond order shows that Nz Contains highest bond dissociation energy.

Magnetic nature: N2 9% diamagnetic, since all electrons in bonding as well as anti-bonding Mo. She paixed and it has no unpaired electrons.



Presence of N2 paired electrons (all paired electrons) shows that N2 98 dia magnetic. The bond length 98 Provinsely proportional to the bond order.

### Molecula orbital Energy level diagram of 02:

Oxygen molecule is homonuclean, diatomic molecule with following

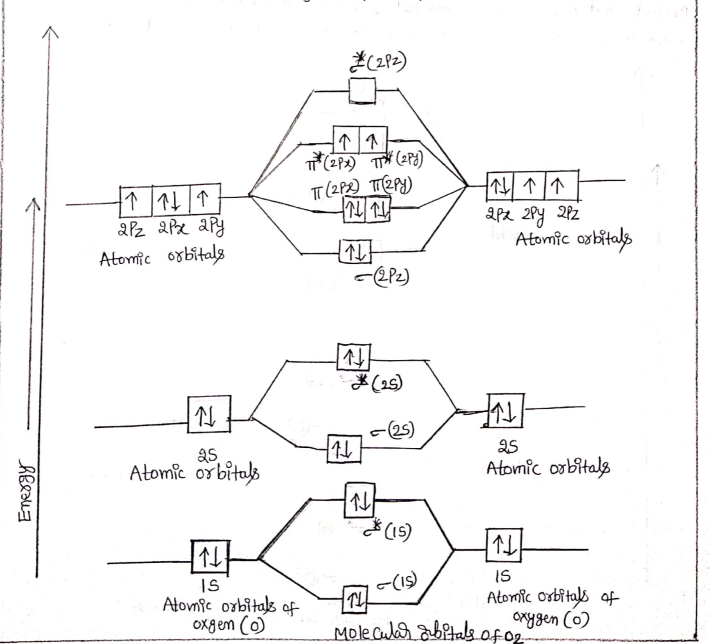
configuration. Atomic number of oxygen (0) =8

The electronic configuration is = 15, 25, 2px, 2py, 2pz

The electionic configuration of  $O_2$  according to molecular orbitals 98 given as

-(15), -(25), -(25), π(219), π(219), π(219), π(219), π(219), σ(219), σ(219)

The molecular orbital energy diagram of 02 is given as follows.



Number of Bonding electrons (Nb) = 10

Number of anti-bonding electrons (Na) = 6

Bond order = 
$$\frac{1}{2}[Nb-Na]$$

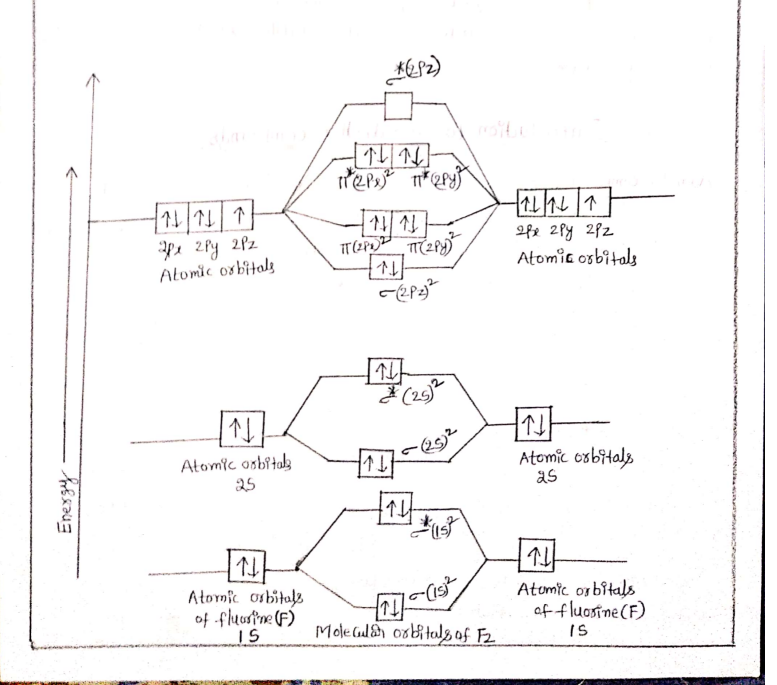
=  $\frac{1}{2}[10-6]$ 
=  $\frac{1}{2}x4 = 2.0$ 

The bond Ender & Justiffes the presence of a double bond in on molecule.

The presence of two umpaired electrons show that exygen molecule is paramagnetic, because it contains two umpaired electrons in Tr (2px) and Tr (2py)

molecular substal.

## Molecular energy diagram of Fluorine (F2):



The electronic configuration of fluorine molecule orbitals is

(=15), (=15), (-25), (=25), (=27), (=

Number of electrons in bonding molecular stortals (Nb) = 10 Number of electrons in anti-bonding molecular stortals (Na) = 8

Bond order = 
$$\frac{1}{2}[Nb-Na]$$
  
=  $\frac{1}{2}[10-8]$   
=  $\frac{1}{2}x2 = 1.0$ 

The molecule (F2) is very stable. The presence of all paired electrons in both bonding and anti-bonding molecular states shows that fluorine molecule is diamagnetic.

# Introduction to coordination compounds

Addition compounds: The compounds obtained upon evaporation of solutions containing two (or more simple compounds in molecular proportions are known as addition (or) molecular compounds, for example

$$K_{2}S_{04} + Al_{2}(S_{04})_{3} + 24 H_{20} \rightarrow K_{2}S_{04} + Al_{2}S_{04})_{3} \cdot 24 H_{20} \rightarrow Potash alum Potash alum double salts 
$$(NH_{4})_{2}S_{04} + FeS_{04} + 6 H_{20} \rightarrow (NH_{4})_{2}S_{04} \cdot FeS_{04} \cdot 6 H_{20} \rightarrow Moh 8/8 \text{ salt}$$$$

cooxdination compounds

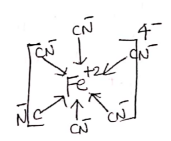
The addition compounds can be classified into two types on the basis of their behaviour in solution phase.

Pouble salts: The addition compounds which one stable in solid state but dissociate to provide all of their constituent ions upon dissolution in water (or) any other polar solvents are known as double salts. Therefore, such a salt behave similar to simple salts in solution. For example, potash alum upon dissolution in water provides kt, Alts, so42 ions.

coordination compounds: The addition compounds which do not provide all of their constituent sons in solution are known as coordination (or) complex compounds. In such compounds, some of the constituent sons lose their sample, potassium ferrocyanide upon dissolution in water provides only kt and [Fe(N)] 4. Here Fet2 and cn sons lose their sample and a complex son [Fe(N)] 4. Here Fet2 and cn sons lose their sample.

### Termenologies:

Ligand: The molecules (or) ions which remain directly attached to the central metal in a complex compound through coordinate linkages are known as ligands. The atom of the ligand which actually donates the electron pain for coordination is known as donor (or) coordination atoms. For example, in ferro gamide ion, cn ions are ligands and "c" is the coordinating atom.



Ligands are classified usually on the basis of their number of donor sites into the following two categories.

(1) Monodentate ligands: such a ligand carries only one donor, site.

For example, NH3, ph3P, H2O:, CgH3N, 10, F, NH2 etc

(2) polydentate ligands: such a ligand comies more than one donor sites for example: EDTA, oxalate etc.

## co-ordination number:

Co-ordination number of the central metal atom/Pon

is the number of donor sites of the ligands which remain attached to it through coordinate linkage. For example, [Fe (N)]: coordination number is 6; [Cu (en)2]: coordination number is 4; [Co (143)54] Biz: coordination of number of some common transition metals are given in the following table.

Metal 90n	coardination number	Metal Pon	coordin	ation number
Fet2 cot2	6 4,6	Aut Alt <sup>3</sup>	2,4	Sparit Species
Nº +2	4,6	Cst3	6	
$ \begin{array}{c} \text{cut}^2 \\ \text{zn}^2 \\ \text{Pt}^2 \end{array} $	4	cot3 Aut3	6	yre is
cu <sup>†</sup> Ag <sup>†</sup>	2,4	pt+4	6	z Laung

Coordination Sphere: The central metal atom/ for along with the ligands attached to it are placed within square brackets and is known as inner sphere (or) coordination sphere. The patforn outside the square brakets is known as ionization (or) outersphere. The species in the inner sphere are non ionizable, whereas those in the outer sphere are ionizable.

Spectrochemical series: An arrangement of ligands according to their ability to increase magnitude (a) for a given metal center is called spectrochemical series.

Weak - I, Bo, SCN, cl, NJ, F, HN CONID, OH, OX2, O2, H20, NCS, Py, NH3, en, bpy, phen, NJ, CH3, CH5, CN, CO-strong

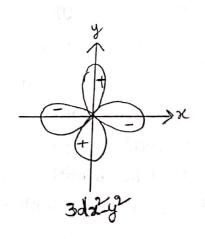
A ligand that produces a large \$ 980 strong-field (or low spin) ligand.
A ligand that produces a small \$ 980 weak-field (or high spin) ligand.

Based on the orientation of the lobes with respect to coordinates, five d-subitals have been grouped as follows:

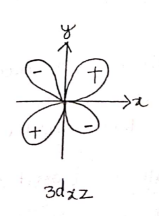
(b) eg set of orbitals: This group has the orbitals which have their lobes along the axes and hence one called axial subitals. Quite obviously these one dry and dz orbitals. Group theory alls these obviously these one dry and dz orbitals. Group theory alls these of orbitals in which "e" refers to doubly degenerate set.

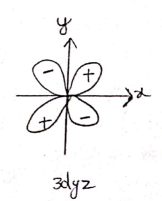
(i) tog set of orbitals: This group includes the orbitals whose lobes lie between the axes and whe called non-axial orbitals. Group theory author obviously these are day, dyz and dza orbitals. Group theory calls these tog orbitals, in which "t" refers to triply degenerate set-

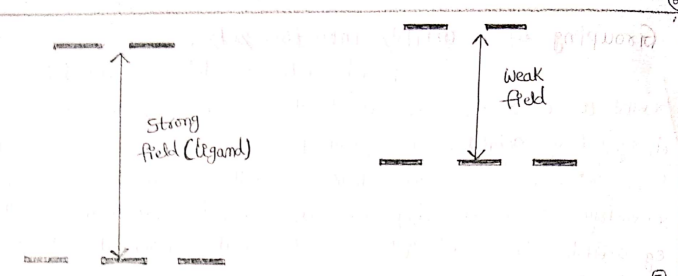
eg set:



tog set: + + > 2







The CN ligands has maximum spilitting power whereas the ligands I has least splitting power.

Salient features of Crystal field theory (CFT): This theory was developed by Brethe and Van Vieck. The salient features of this theory one as follows. (i) The central metal cation is surrounded by ligands, which contain one or more lone pain of electrons.

(ii) The legands are treated as point changes.

(11) The Ponic (Egands (eg: F, d, cN etc) one regarded as

negative point changes, while neutral ligands (eg: H20, NH3 etc) are regarded as dipoles (i.e dipolar). In metal complexes, the negative end of the metal cation.

(IV) There is no interaction between metal expitals

and legand orbitals.

is purely electroptatic (or) coulombic attraction between anion (Positively charged) and negatively charged anion (or) megative end of neutral depole molecule. (ii) All the d-orbitals on the metal have the same energy (i.e degenerate) in the free atom. However, when a complex is formed, the legand destroy the degeneracy of these orbitals, i.e. the orbitals now have the different energies.

# Coystal field splitting of d-orbitals in octahedral complexes:

In case of free metal for all the five d-subitals one degenerate, i.e., these have the same energy. Now let us consider an octahedral complex [ML6] "In which the central metal cation, m" 18 placed at the centre of the octahedron and its surrounded by six ligands which reside at the corners of the octahedron as shown in fig.

(1) The ligands on each of the three axes are allowed to approach towards the metal cation, mit from both the ends of the exes. In this process the electrons in d-sibilate of the metal cation one repelled by negative point charge (as) by the negative, end of the dipole of the ligands.

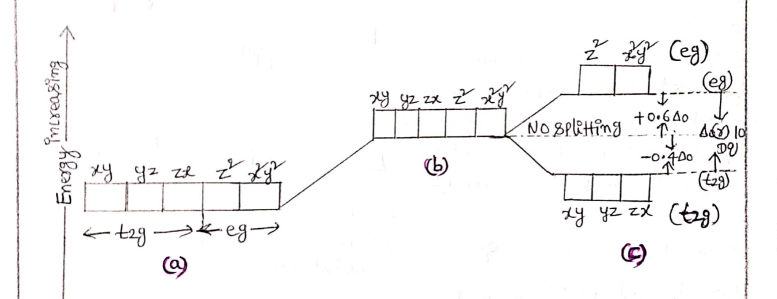
possition of the central metal cation, (11) The lobes of the two eg orbitals mit and sex legands, L's in an

(i.e dz and dxy), lie directly 9n

the path of the approaching ligands, the electrons in these or bitals exponence greater force of repulsion than those in three tog orbitals (dry, dyz and dzi), whose lobes are directed in space between the path of the approaching ligands. So the energy of eg orbitals is increased while that tog orbitals is decreased. (greater is the repulsion, greater is Increase the energy).

So under the influence of approaching ligands the five d-abitals which were split into two levels. (1) tog level, which is triply degenerate and is of lower energy and iii eg level, which Ps doubly degenerate and Ps of higher emergy. The separation of five d-abitals of the metal for into 2 sets having different emerses is called crystal field splitting (or) energy level splitting.

This enough difference asizes because of the electrostatic field exerted by the legands on tag and eg sets of orbitals of the central metal cation is called crystal field stabilization enough. It is denoted by  $\Delta_0$  (a) to  $\Delta_0$ . Thus, we find that tag set losses an emonsy equal to  $0.4\Delta_0$  (= 4DV), while eg set gains an emonsy equal to  $0.6\Delta_0$  (= 6DV). The loss and gain in emonsion of tag and ey orbitals is shown by negative and positive signs, respectively and ey orbitals is shown by negative and positive signs, respectively



(a) five degenerate d-sibitals of free metal cation which are free from any ligand field. (b) Hypothetical degenerate d-sibitals at a higher energy level under spherically ligand field. (c) splitting of d-sibitals is under influence of approaching ligands in to tag and eg sets.

Distribution of d-electrons in tog and eg substals in octahedral complexes: The distribution of d-electrons in tog and eg oxbitals takes place on the basis of the nature of the ligands. i.e. whether the ligands are weak (or) strong.

is when the ligands are weak: under the influence of weak ligands the energy difference, so between tzg and eg sets is relatively small and hence all the five d-orbitals of these two selfs may be supposed to be degenerate, i.e. in the presence of weak ligands all the d-orbitals have the same energy and consequently the distribution of d-electrons in tzg and eg sets takes place according to Hund's rule, which states that electrons will pairul only when each of the five-d-orbitals is at least singly filled.

when the ligands are strong: In octahedral complexes containing strong ligands, distribution of d-electrons in tzg and eg sets does not obey Hund's rule. Thus in stronger field the first six electrons numbered as 1-6 will go tzg-set and the remaining four electrons numbered as 7-10 enter eg-set.

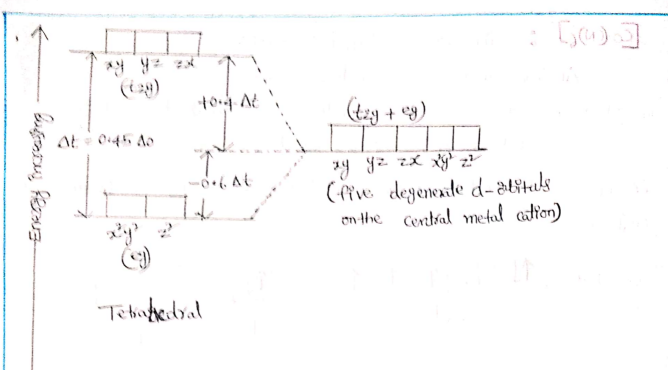
dn	Stronger field (	Low-spin c	omplexes)	weak field (hig	h-spin con	nplexes)
9095	tegeg configuo	d n	5	tag eg configurdio	n n	S
d'	t28 e8°	1	1/2	tag ego	1.	1/2
d2	tag eg	2	-1	t292 ego	2	1,
$d^3$	t293 eg°	3	3/2	t293 ego	3	3/2
d4	tagt ego	2	1	+293 eg	4	2
d <sup>5</sup>	+295 eg	1	1/2	+293 eg2	5	5/2
d6	tagé ego	0	0	t2g4 eg2	4 1	2
d <sup>7</sup>	t296 eg 1	1 1	1/2	t295 cg2	3	3/2
dg	t28 eg2	2	1	t296 eg2	9_	
d9	£296 eg3	1	1/2	+296 eg3		1/2
d10	t2\$ eg4	О	0	t296 e94	О	0

n = No. of unpaised electrons S = Regultant spin = 1/2 xn

# Crystal field splitting of d-obstals in Tetrahedral complexes:

A tetrahedron placed in a cube. The centre of the cube is the centre of the tetrahedron at which is placed central metallic cation MMT. Four corners of the cube one the four corners of the tetrahedron at which one placed the four nagative ligands which have been shown by costles with negative

In order to understand the splitting of d-dblitals in testahedral complexes it is convienient to picture a testahedron placed inside a cube. The four negative ligands placed at the four corners of the testahedron are located at the four corners of the cube. Thus, these four ligands are lying between the three axes viz: x, y and z axes. The lobes of tag distrib (day, dyz and dzx) are lying between the axes, i.e are lying directly in the patr of the ligands, these orbitals will experience greater force of repulsion from the ligands than those of eg oxbitals (dz² and dx²y²) whose lobes are lying along the axes, i.e are lying in space between the ligands. Thus the energy of tag oxbitals will be increased while that of eg oxbitals will be decreased. Consequently the d-switals are again split into two sets as shown below. The order of energy of tag and eg sets is the revose of that seen in odahabal complexes. The energy difference between tag and eg sets for testahedral complex is represented as  $\Delta E$ .



# Distribution of d-electrons in tetrahedral complexes:

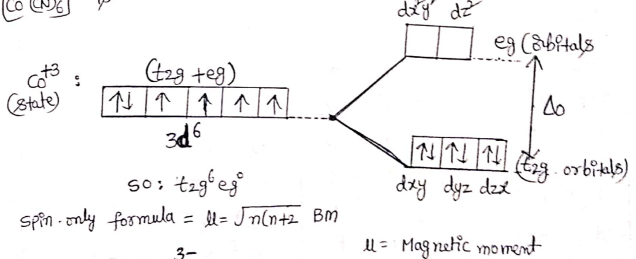
1	Annual Control of the	The Section of the Contract of	management descriptions and the same				
cl <sup>m</sup> con	weak field (HS-complexes)			Strong Ateld (LS-complexes)			
Banapi	tog eg configuration	m	S	tog eg configuration	প	S	
d	t29° e9'	1	1/2	£29° eg'	1 ,	Y2_	
$d^2$	429° eg2	2	1	teg eg2	2	1	
$d^3$	t29 eg2	3	4/2	-129° 093	1	V <sub>2</sub>	
14	t292 eg2	4	થ	tog egt	0	0	
d5	tzg3 eg2	5	5/2	tog egt	1	1/2_	
db	t293 eg3	4	2	tg2 egt	2	1 -	
d7	-6293 eg4	3	3/2	1283 est	3	3/2	
q <sub>8</sub>	t294 e94	2	1	t28+ 99	2	1	
9	t295 egt	1 1	1/2	t295 e94	1	1/2	
d <sup>10</sup>	t296 eg4	0	0	t296 e97	0	0	

n= number of un flured electrops

S= nx1.

CFT is helpful in determining the number of unpulsed electrons in a given HS (high spin) - and LS (Low spin) - tetral heard and octahedral complexes, and consequently with the help of "spin only" formula. It = In(n+2 BM.

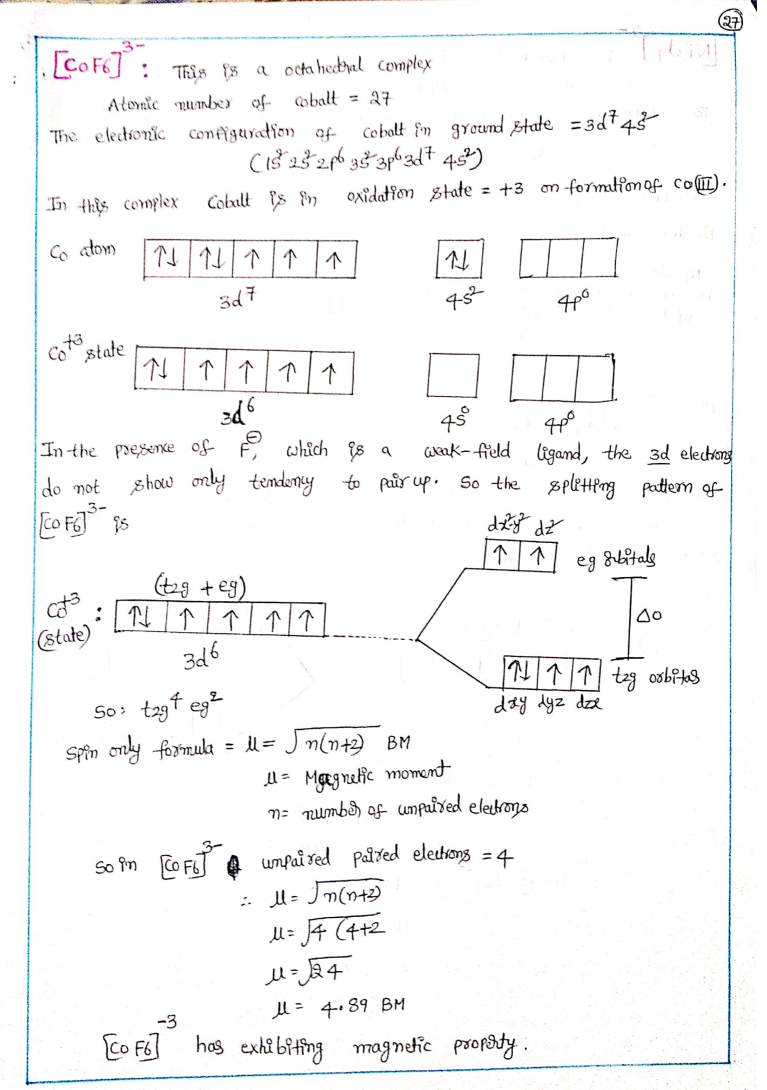
[Co (CN) ? This is a octahedral complex Atomic number of cobalt is = 27 The electronic configuration of cobalt in ground state = 3d 45° (15252p6353p6453d7) In this complex cobalt is in exidation state = +3 on formation of Co (III). 11/11 Co atom 3d7 cot3 state In the presence of CN, which is a strong-field ligand, the 3d electrons show tendency to pain up. so the splitting pattern of (co (N)6]3- 95 dxy dz eg (Sibitals



So in [Co(N)] number of unpaired electrons

S= 11=0 BM

So [Co(N)6] has no magnetic property.



[Nid4]2-: This is a tetrahedral complex

Atomic number of Nickel is = 28

The electronic configuration of Nickel in ground state = 3d 15

(18282p6383p6483d8)

In this complex Nickel Ps in oxidation = +2, so on-formation of Ni (II) complex.

Ni atom
in ground 11 11 11 1 1

State
3d8

13 40

Nitestate [] 1] 1] 1 1

4.8 AP

In the presence of it, which is weak-field legand, it does not have the tendancy to pair up the electrons in the 3d level. so the splitting pattern of [Ni d4] is

N9+2: (+28 + eg)
state [1] 1] 1] 1 1 1

50: £294 eg4

SPM only formula = Il= m(n+2) BM 1)

1 = Magnetic moment

n = number of unpaired electrons

So in [NPU4] = 1 = Jm(n+2)

M= J2(2+2)

U= \[ \begin{aligned}
8 \\
\end{aligned}

从= 2,82 BM

[NICN4]?: This is a Tetrahedral complex. In this case again NI is in (+2) exidation state. The electronic configuration of NI atom is 3d8 45 (15252pb 353pb 753d8) in ground state which changes to 3d8 45 on the formation of NICII) with the loss
of two electrons.  No atom in 11 11 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Nº (II) state 11 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Since cho is a very strong field ligand, it has the tendency to pair up electrons in the 3d level so as to set up space for its own electrons to be accommodated. Now, the configuration changes as shown
N9+2: 3d8 48 4P°  ASP hybridization
Thus, empty 3d-3rbitals, 45 and two 4P grobitals hybridize to
from cyanide ligands occupy these orbitals to form semare planar and diamagnetic complex.  [Ni (CN)4] = Pn this complex unpaired electrons = 0 80 8Pin
INT (CN)4]  The formula $U = \int m(n+2)$

= [p(y)) M

## BAND Structure of solids:

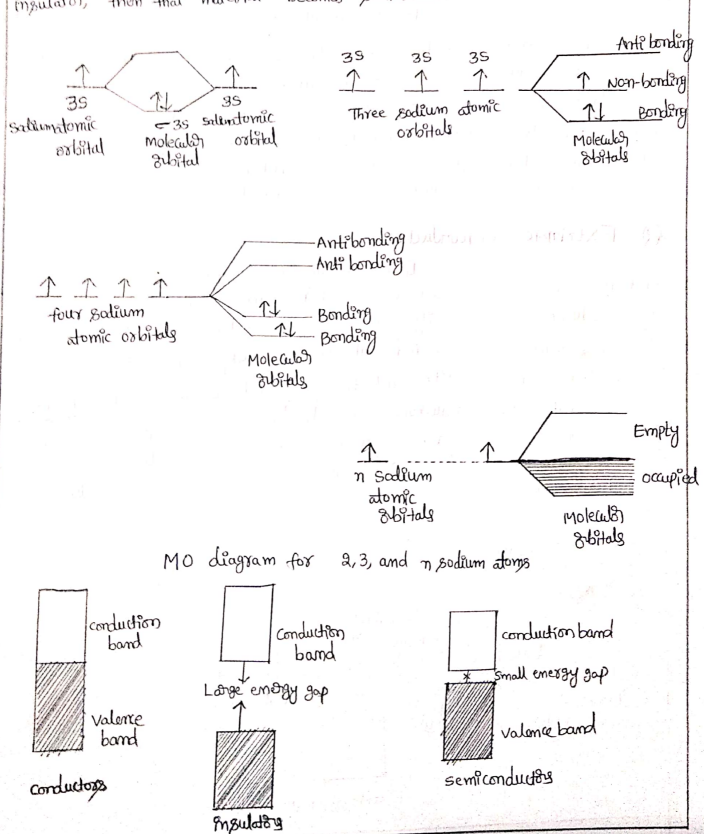
The molecular orbital theory of metals is known as band model. Since the smallest perceptible piece of metal would contain about 1020 atoms. A large number of molecular orbital are formed from a large number of atomic orbitals.

Let us consider the example of a piece of sodium metal. Each sodium atom has single 35 valence electron. When two atoms are closer together, the atomic orbitals will combine to form two molecular orbitals, 35 and 35. The two elections will occupy the bonding molecular arbital, while the antibording orbital will remain empty. Similarly for three atoms, three molecular orbitals are formed - bonding, non-bonding and antibonding. when four sodium atoms combine, we two 35 orbitals and 35 th orbitals. The quantum mechanical considerations do not permit these orbitals to be degenerate but they are closely placed, being alongst similar in energy. The concept can be extended to n number of sodium atoms present in a prece of sodium metal. In this multi-atom system, the number of molecular subital states will be equal to the number (1) of the atomic orbitals combining. Since the number of molecular orbitals is large, the spacing between them decreases to become almost negligable and we get a band of continuous emergy levels". These molecular orbitals extend in all three dimensions over all the atoms in the metal piece. The was energy levels Known as bands. The molecular orbital theory is known as Band model.

The molecular orbital which contains the pair of electrons is called valence orbitals and the band formed it is called valence band the molecular orbitals and the band formed it is called valence band the molecular orbitals, which is formed by the band is called conducting orbitals, which is formed by the band is called conducting band. Now the emongy of between the valence band and conducting band plays an important role. If the fermi gap is more the material becomes insulate, if the fermi energy gap is least

the material becomes conductor because when the election in valunce bound are exposed to heat, light, emongy, they absorb the enougy and exilted to higher emongy level. when fermi energy gap is more, the electrons cannot be excited to the conducting band, and the material becomes insulation.

Engulator, then that material becomes semiconductor.



# Role of doing in band structures:

Types of semiconductors: semiconductors are of

two types (1) Intrinsic semiconductors

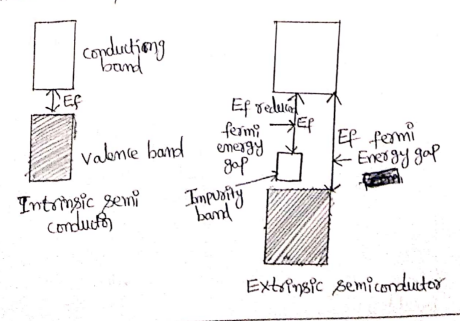
(2) Extrinsic Semi conductos

## W Intrinsic Semicondudios:

If the fearmi energy gap is small between the valence band and conducting band, the electron from valence band is exited to conducting band, it is called intrinsic semiconductor both the hole left in valence band and the excited electron to the conduction band contribute tawards conductivity. As the temperature rises, the number of electrons promoted to the conduction band increases contributing to the increase in conductivity of semiconductor.

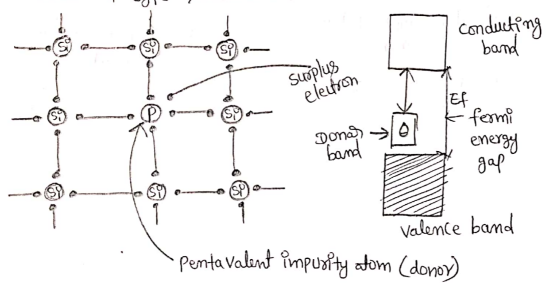
# @ Exterimenc semiconductoss:

If the fermi energy gap is more, where the electrons cannot be excited to conduction band, conductance can be improved by introducing an impurity into the semiconducting material which is called doping. Doping can be corried out by introducing an electron deficient (05) electron rich impurity into the material which produces p-type and n-type semiconductors respectively. The impurity forms an impurity band called donor band between Valence band and conducting band of the material, whereby the formi energy gap is reduced and conduction takes place.



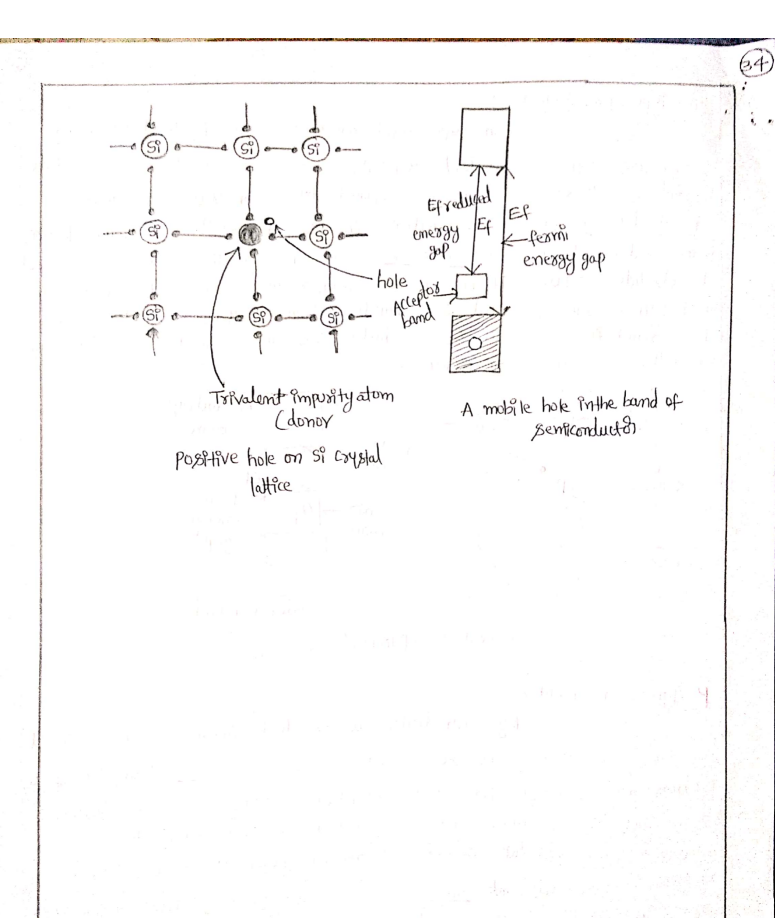
## n - Type semi conductor:

n-type seme conductor's were produced by doping Si (os) Gie with pentavalent impurity atoms like P, As etc. A minute amount of Si 60 Gie atoms are replaced by P (ox) Gie atoms are leutrons from covalent bonds with Si (ox) Gie and the fifth eleutrons are absolute zero. At the normal temperature some of the fifth eleutrons of impurity are promoted to conduction band, causing conduction. Since the conductivity of such semiconductors are due to negative eleutrons, there are called n-type semiconductors.



P- Type semicondutor:

By introducing a trivalent impusity atom like, B, Al, Gia. Pinto Si (or) Gie the replacement of these Si (or) Gie atoms by impusity produces an incomplete bond in the structure producing a positive hole. The positive holes are localized around trivalent impusity atom at low temperatures (or) absolute zero. At normal temperatures the valence electrons on the adjacent Si (or) Gie atom may gain sufficient energy to move into the hole, thus creating a new hole on the Si (or) Gie. By a series of hops, the positive hole can migrate across the crystall, thuy current is carried out by the migration of positive centres.



# Module-III Electrochemistry and Corrosion

#### **MODULE-III**

#### ELECTRO CHEMISTRY AND CORROSION

#### Q.1. Define Specific, Equivalent and Molecular conductance and their units?

**Specific conductance:** The resistance offered by a conductor to the passage of electricity through it is directly proportional to length (l) and inversely proportional to the area of cross section (a). The resistance R

R 
$$\alpha l/a$$
  
R =  $\rho l/a$   
 $\rho$ = specific resistance  
R =  $\rho l/a$   
 $\rho$  =Ra/l

Units: Resistance = R = Ohm

Where  $\rho$  (rho) is a constant known as "specific resistance". Specific conductance (Kv) is the reciprocal of specific resistance. Kv=  $1/\rho$  Kv=1/Ra

Specific conductance is expressed in reciprocal Ohm<sup>-1</sup>Cm<sup>-1</sup>(or) SCm<sup>-1</sup>S= siemens

**Equivalent conductance:** "If one gram equivalent weight of an electrolyte is dissolved in "V" ml of the solvent, the conductivity of all ions produced at the dilution V is known as equivalent conductance, which is denoted by  $\lambda v$ ".

The equivalent conductance is equal to the product of specific conductance (Kv) and the volume (V).  $\lambda v = Kv X V$  V=1000/C or 1000/normality

$$\lambda \mathbf{v} = 1/\text{Ra X Kv}$$
  
 $\lambda \mathbf{v} = 1/\text{Ra X } 1000/\text{N}$ 

**Molecular conductance:** "If one gram molecular weight of an electrolyte is dissolved in "V" ml of the solvent, the conductivity of all ions produced at the dilution V is known as equivalent conductance, which is denoted by  $\mu v$ ".

$$\mu v$$
=Kv X V V=1000/C or 1000/molarity

$$\lambda \mathbf{v} = 1/\text{Ra X Kv}$$
  
 $\lambda \mathbf{v} = 1/\text{Ra X } 1000/\text{Moles}$ 

$$\lambda$$
v=Ohm<sup>-1</sup> cm<sup>2</sup> mol<sup>-1</sup> (or) S cm<sup>2</sup> mol<sup>-1</sup>

#### Cell constant (x)

It is a constant, characteristic of the cell in which the electrolytic is taken and its value depends on the distance between the electrodes and area of cross section of the electrodes

Cell Constant = Distance between the electrodes

Area of cross section of each electrodes

$$X = \frac{l}{a}$$

(1) The equivalent conductance of 0.005 N NaOH solution is 240 mho cm<sup>2</sup> equi<sup>-1</sup>. What is the specific conductance and resistance the electrodes are 1 cm apart and each have a surface area of 1cm<sup>2</sup>.

$$V = \frac{1000}{C}$$

$$C$$

$$\lambda v = Kv \ X \ V$$

$$240 = Kv \ X \ \frac{1000}{0.005}$$

$$Kv = \frac{240 \ X \ 0.005}{1000}$$

$$Kv = 1.2 \ X10^{-3} \ ohm^{-1} \ cm^{-1}$$

$$Specific conductance of 0.005 \ N \ NaOH \ is = 0.0012 \ ohm^{-1} \ cm^{-1}$$

$$Kv = I/Ra \qquad I = 1 \ cm \ and \ a = 1 \ cm2$$

$$R = 1/Kv$$

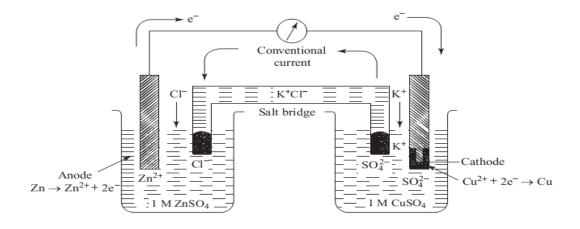
$$R = 1/0.0012$$

$$R = 833.3 \ ohm$$

#### Q.2. Write a short note on galvanic cell and cell potential.

**Galvanic Cell:** An electrochemical cell is a device for converting chemical energy into electrical is called galvanic cell. A galvanic cell is made of two half cells. One is oxidation or anodic half cell and the other one is reduction or cathodic half cell.

Daniel cell is an example of galvanic cell having zinc and copper electrodes. The first half cell consists of zinc electrode dipped in ZnSO4 solution and the second half cell is made of copper electrode dipped in copper sulphate solution. Both half cells are connected externally by metallic conductor (connecting wire) and internally by a bent glass tube having saturated solution of a strong electrolyte (KCl) called salt bridge. It acts as a bridge between the two half cells.



The chemical reaction that takes place in a galvanic cell is the redox reaction. One electrode acts as anode in which oxidation takes place and the other acts as the cathode in which reduction takes place. In electrochemical cells anode has negative polarity and cathode has positive polarity.

#### **Representation of Galvanic cell:**

A zinc or a copper galvanic cell can be represented as

The double bar shows a salt bridge, i.e. electrolyte—electrolyte junction. The chemical reactions taking place at both electrodes may be written as follows:

At anode: Oxidation takes place with the liberation of two electrons.

$$Zn \rightarrow Zn^{+2} + 2e^{-}$$
 (oxidation)

At cathode: Reduction occurs and cuprous ion is reduced to metallic copper.

$$Cu^{+2} + 2e^{-} \rightarrow Cu$$
 (reduction)

The overall reaction is

$$Zn + Cu^{+2} Zn^{+2} + Cu$$

- The electrode showing oxidation reaction is anode and the other electrode where reduction occurs is cathode. As per IUPAC convention, the anode is always represented on the left and the cathode always represented on the right side of the cell.
- As the connection is complete, the flow of electrons will be externally from anode to cathode and internally from cathode to anode through the salt bridge. The flow of current is due to the difference in electrode potentials of both the electrodes. The potential difference in the cell is called the EMF and is measured in volts. It can be measured by the potentiometer. The flow of current becomes slow after using the electrodes for a long time because of the polarization of the electrodes.

Salt bridge: Salt bridge is a U-shaped tube containing concentrated solution of an inert electrolyte such as KCl, KNO<sub>3</sub>, and  $K_2SO_4$  or a paste of inert electrolyte (whose ions do not take part in redox reaction and do not react with the electrolyte) in agar—agar medium or gelatin.

Functions of salt bridge:

- Salt bridge helps to complete the circuit by allowing the ions to flow from one solution to the other without mixing the two solutions.
- It helps to maintain electrical neutrality of the solution in the two half cells.

#### **Cell Potential:**

The potential difference between the two electrodes in a galvanic cell is called a cell potential or emf of the cell. It is measured in volts. The difference of potential, which causes flow of electrons from the electrode of higher potential to the electrode of lower potential, is called electromotive force (EMF). The emf of galvanic cell is calculated from the reduction half-cell potentials using the following relation.

$$E(cell) = E(right) - E(left)$$

emf of the cell = Potential of the half cell on the right hand side (Cathode) - Potential of the half cell on the left hand side (Anode)

$$E_{cell} = E_{cathode} - E_{anode}$$

$$OR$$

$$E(cell) = E(right) - E(left)$$

Where E(cell) = e.m.f of cell E(right) = reduction potential of right hand side electrode E(left) = reduction potential of left hand side electrode

The difference in potentials of the two half – cells of a cell arises due to the flow of electrons from anode to cathode and flow of current from cathode to anode.

#### Q.3. what is Single Electrode potential and how to measure single electrode potential?

Potential difference is developed between the metal (electrode) ions from metal to the solution (electrolyte solution) (or) from solution to the metal is called Single Electrode potential. The tendency of an electrode to lose electrons is a direct measure of its tendency to get oxidized: and this tendency is called, oxidation potential, similarly, the tendency of an electrode to gain electrons is a direct measure of its tendency to get reduced and this tendency is known as reduction potential.

It is quite obvious that the value of reduction potential negative of its oxidation potential and vice versa. Thus, the oxidation potential of an electrode is - x volt, then its reduction potential will have a value of + x volt.

(1) Calculate the e.m.f. of the following reaction at 25  $^{\circ}$ C.

$$Cu^{+2}+Zn \longrightarrow Zn^{+2}+Cu$$
  
 $E^0_{Zn}(oxi)=0.763 \text{ volts}$   
 $E^0_{Cu}(oxi)=-0.337 \text{ volts}$ 

#### **Measurement of Single Electrode potential:**

Single electrode potential cannot be directly measured. The single electrode potential of a half cell depends on

- 1. Concentration of the ions in the solution
- 2. Tendency to form ions
- 3. Temperature

The single electrode potential is conveniently measured by combining the half cell with a standard electrode or reference electrode and measuring the total EMF of a cell

$$E(cell) = E(right) - E(left)$$

If the standard cell acts as Anode the equation becomes

$$E(cell) = E(right) - E^0$$

If the standard cell acts as Cathode the equation becomes

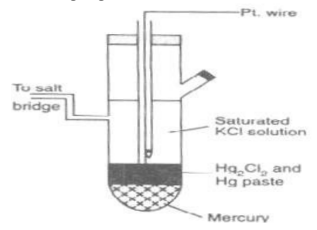
$$E(cell) = E^0 - E(left)$$

Where  $\mathbf{E}^0$  = Standard electrode potential

**Standard electrode potential**: The potential exhibited by a single electrode at unit concentration of the concerned metal ion or non-metal ion solution at 25° C is called **standard electrode potential**. This is expressed by 'E<sup>0</sup>'

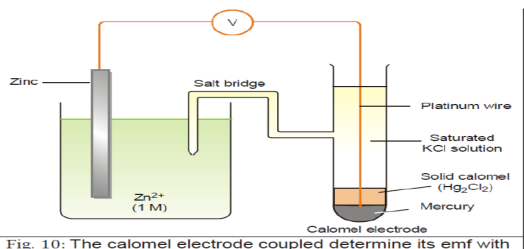
## Q.4. Explain the construction and working of Calomel Electrode and mention its advantages.

The second type of reference electrode employed for measuring the single electrode potential is Calomel electrode. Calomel electrode is particularly very simple to construct, free from surface sensitivity and accurate to use even in a very normal laboratory. It consists of a glass tube having side tube on each side. High purity of mercury is placed at the bottom of the tube and connected to the circuit by a platinum wire, sealed in a glass tube. The surface of mercury is covered with a paste of mercurous chloride (Hg<sub>2</sub>Cl<sub>2</sub>) (commercially known as calomel) and mercury in KCl solution. KCl solution is used as electrolyte. The electrode is represented as Hg, Hg<sub>2</sub>Cl<sub>2</sub>/KCl. The standard electrode potential of this electrode at 25°C is



0.1M KCl | Hg<sub>2</sub>Cl<sub>2</sub>(s) | Hg, E<sup>0</sup>=0.3338 V 1.0M KCl | Hg<sub>2</sub>Cl<sub>2</sub> (s) | Hg, E<sup>0</sup>= 0.2800 V Saturated KCl | Hg<sub>2</sub>Cl<sub>2</sub> (s) | Hg, E<sup>0</sup>= 0.2422V The corresponding electrode reaction is; Hg<sub>2</sub>Cl<sub>2</sub>+2e- $\rightarrow$ 2Hg + 2Cl<sup>-</sup>

Measurement of single electrode potential of Zn using saturated calomel electrode.



To measure the electrode potential of Zn, a cell is made by combining the saturated calomel electrode with Zn electrode, the potential to determine. The reduction potential of Zn electrode is less than E<sup>0</sup> of calomel electrode (0.2422 v), the calomel electrode will act as cathode and Zn electrode act as anode. The following cell reaction will occur in the above cell.

 $Zn \rightarrow Zn^{+2} + 2e^{-}$  (oxidation) At anode:  $\begin{array}{ccc} \underline{Hg2Cl2+2e-} & \underline{2Hg+2Cl-} \\ Zn+Hg2Cl2 & \overline{\longrightarrow} ZnCl2+2Hg \end{array}$ At cathode:

e.m.f=1.0052 volts Cell reaction:

The electrode potential is calculated from the e.m.f of the cell as follows.

$$\begin{split} E(cell) &= E(right) - E(left) \\ E(cell) &= E^0(calomel) - E(left) \\ E(cell) &= E^0(calomel) - E(Zn) \\ E(Zn) &= E^0(calomel) - Ecell \\ &= 0.2422\text{-}1.0052 \\ &= -0.763 \text{ volts} \end{split}$$

The reduction potential of Zn electrode is = -0.763 volts

#### Q. 5. Write Nernst equation? Explain its terms and its applications.

Nernst equation is a thermodynamic equation which relates the change in free energy  $(\Delta G)$  and cell potential with concentration,  $M^{n+}$ 

The decrease in free energy in the reversible reaction will produce electrical energy i.e.

$$-\Delta$$
 G = nFE and

$$\Delta G^0 = -nFE^0$$
-----Eq-(1)

Where E = Electrode potential

 $E^0$  = Standard electrode potential

F = Faraday (96,500 coulombs)

Consider the following redox reaction

$$M^{n+} + ne^- \leftrightarrow M$$

For such a redox reversible reaction, the free energy change (D G) and its equilibrium constant (K) are related as:

$$\Delta G = \Delta G^0 + RT \ln [Product] / [Reactant] ------Eq. (2)$$

Where  $\Delta G^0$  = standard free energy change

The above equation is known as Van't Hoff Isotherm.

Comparing equation 1 & 2 
$$\begin{split} -nFE &= -nFE^0 + RT \; ln[M]/[M^{n+]} \\ &= -nFE^0 + RT \; ln1/[M^{n+]} \end{split}$$
 Where, concentration of the metal is unity or 
$$-nEF &= -nFE^0 - RT \; ln \; [M^{n+]} \\ Dividing \; the \; equation \; by \; -nF \\ E &= E^0 + RT/nFln \; [M^{n+]} \end{split}$$

$$E = E^0 + 2.303RT/nF log [M^{n+1}]$$
 R= 8.314J/K/mole, T= 298 K (25  $^{0C}$ )

$$E = E^0 + 0.0591/n \log [M^{n+}]$$
-----Eq-(3)

This equation-3 is known as "Nernst Equation" for reduction electrode potential. For oxidation potential  $E=E^0$  - 0.0591/n log [M<sup>n+</sup>]

#### **Applications of Nernst Equation**

- 1. It can be used to study the effect of electrolyte concentration on electrode potential.
- 2. It can be used for the calculation of the potential of a cell under non-standard conditions.
- 3. Determination of unknown concentration of one of the ionic species in a cell is possible with the help of Nernst equation, provided  $E_{\text{cell}}$  and concentration of other ionic species are known.
- 4. The pH of a solution can be calculated from the measurement of EMF and Nernst equation.
- 5. It can be also be used for finding the valency of an ion or the number of electrons involved in the electrode reaction.
- (1) What is the potential of a lead electrode that is in contact with a solution of 0.015 M Pb<sup>+2</sup>ions. Standard electrode potential (E0) for Pb  $\longrightarrow$  Pb<sup>+2</sup> + 2e- is 0.13 volts

The Nernst equation for oxidation potential of Pb is  $E = E^0 - 0.0591/n \log [Pb^{+2}]$ 

E=E<sup>0</sup>-0.05991/2 log [0.015] E=0.13-0.05991/2 log [0.015]

E=0.13-0.02955 log [-1.824]

E = 0.13 + 0.0539

Oxidation potential of Pb electrode is E=0.184 volts

(2) Calculate the reduction potential of Cu+2/Cu=0.5M at 25°C.E° of Cu+2/Cu=0.337 volts.

Concentration of [Cu+2]=0.5M

 $E^{0}$  Cu+2/Cu=0.337

n=2

The Nernst equation for reduction potential of Cu+2 is

 $E = E^0 + 0.0591/n \log [Cu^{+2}]$ 

 $E = 0.337 + 0.0591/2 \log [0.5]$ 

E =0.3337+0.02955[-0.3010]

E = 0.337 - 0.0089

E=0.328 volts

Reduction potential of Cu =0.328 volts

(3) Find the Oxidation potential of  $Zn/Zn^{+2}=0.2M$  electrode at  $25^{\circ}C$ . Standard oxidation potential of  $Zn/Zn^{+2}$  is 0.763 volts.

Concentration of [Zn<sup>+2</sup>]=0.2M

 $E^0 Zn/Zn^{+2} = 0.763 \text{ volts}$ 

The Nernst equation for reduction potential of Zn is

 $E = E^0 - 0.0591/n \log [Zn^{+2}]$ 

 $E = 0.763 - 0.0591/2 \log [0.2]$ 

E =0.763-0.02955[-0.69897]

E = 0.763 + 0.02065

Oxidation potential of Zn E = 0.78365 volts

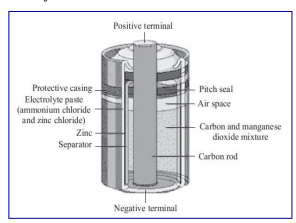
#### Q. 6. Explain the primary batteries with suitable example.

**Primary batteries or Primary Cell:** These batteries can convert chemical energy in to electrical energy. These are also called non-rechargeable batteries. Eg., Voltaic cells, Leclanche cell, and lithium primary cell.

#### Dry cell or Leclanche cell or Zn-MnO<sub>2</sub> cell:

Dry cell is one important primary cell. The cell contains a cylindrical Zinc container that acts

as an Anode. A graphite rod placed in the centre of the cell and acts as a cathode. The space between the anode and cathode is packed with the paste of NH<sub>4</sub>Cl, ZnCl<sub>2</sub> and the graphite rod is surrounded by powdered MnO<sub>2</sub> and carbon as shown in figure. This cell is called Dry cell because of the absence of any liquid phase, even the electrolyte consists of NH<sub>4</sub>Cl, ZnCl<sub>2</sub>and MnO<sub>2</sub>to which starch is added to make a thick paste which prevents leakage. The graphite rod is fitted with a metal cap and the cylinder is sealed at the top with a pitch.



Anode: Zinc container

Cathode: Graphite rod

Electrolyte: NH<sub>4</sub>Cl+ZnCl<sub>2</sub>+MnO<sub>2</sub>

At anode: 
$$Zn \rightarrow Zn^{2+} + 2e$$
- (oxidation)

At cathode:  $2MnO_2+H_2O+2e-\rightarrow Mn_2O_3+2OH^-$  (reduction)

The net cell reaction is 
$$Zn+2MnO_2+H_2O$$
  $Zn^{2+}+Mn_2O_3+2OH^{-}$ 

The resulting OH-ions react with NH<sub>4</sub>Cl to produce NH<sub>3</sub>, which is not liberated as gas but immediately combines with  $Zn^{2+}$  and Cl<sup>-</sup> ions to form a complex [ $Zn(NH_3)_2Cl_2$ ] (diamminedichloro zinc).

$$2NH_4Cl + 2OH^- \longrightarrow 2NH_3 + 2Cl^- + 2H_2O$$

$$Zn^{2+}+2NH_3+2Cl^- \longrightarrow [Zn(NH_3)_2Cl_2]$$

The obtained zinc complex is disrupts the current flow. The cell exhibits a potential of nearly 1.5 volts. When the reactants have been completed (or) converted in to products, no electricity is produced and the cell becomes dead, and cannot be used after that.

#### **Advantages:**

1) These cells have voltage ranging from 1.25v to 1.50v.

- 2) Primary cells are used in the torches, radios, transistors, hearing aids, pacemakers, watches etc.
- 3) Price is low.

#### **Disadvantages:**

This cell does not have a long life, because the acidic NH<sub>4</sub>Cl corrodes the container even when the cell is not in use.

## Q. 7. Describe the construction and working of lead-acid battery along with reactions involved during discharging and charging.

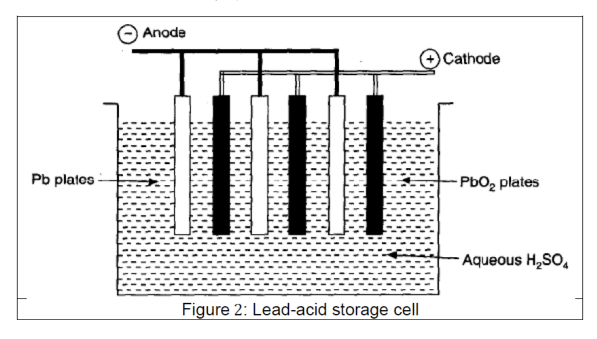
These batteries are rechargeable. A good secondary cell undergoes a number of **discharging** and **charging** cycles. In these cells both "voltaic cell" and "electrolytic reaction" takes place. When operating voltaic cell, it supplies electrical energy and it becomes "run down", when being charged the cell operates as an electrolytic cell.

#### **Examples**

- 1. Lead-acid battery
- 2. Nickel-cadmium battery
- 3. Lithium-ion cell battery
- 1. **Lead-acid batteries:** Lead-acid battery consists of **Pb** anode and lead dioxide (Pbo<sub>2</sub>) cathode. A number of Pb plates (-Ve) are connected in parallel and Pbo<sub>2</sub> (+Ve) plates are also connected in parallel. The plates are separated by one insulators like strips of wood, rubber or glass fibre and entire combination was immersed in dil H<sub>2</sub>SO<sub>4</sub> (38%), which is worked as an electrolyte.

**Anode:** Sponge metallic Lead (Pb) **Cathode:** Lead-dioxide (PbO<sub>2</sub>) **Electrolyte:** 5M H<sub>2</sub>SO<sub>4</sub> (38%)

**EMF:** 2V



**Discharging:** When the battery is in use (discharging) the reactions at the electrodes are as follows.

At anode:  $Pb + SO_4^{2-} \rightarrow PbSO_4 + 2e- (0.356V)$ 

At cathode:  $PbO_2 + SO_4^{2-} + 4H^+ + 2e^- \rightarrow PbSO_4 + 2H_2O$  (1.658V)

The overall reaction is:  $Pb(s) + PbO_2(s) + 2H_2SO_4 \rightarrow 2PbSO_4 + 2H_2O$  (2.041 V)

During discharging of battery,  $H_2SO_4$  (d=1.84 g/cm<sup>3</sup>) is consumed for the oxidation of Pb, and as a hence the density of  $H_2SO_4$  falls. When it is falls below 1.20g/cm<sup>3</sup>, the battery needs recharging. In discharging, the cell acts as a voltaic cell.

**Charging:** In recharging, the cell operates like an electrolytic cell and an external EMF greater than 2V is passed, the electrode reactions are the reverse of those that occur during discharge.

$$PbSO_4 + 2e - \rightarrow Pb + SO_4^{2-}$$
  
 $PbSO_4 + 2H_2O \rightarrow PbO_2 + SO_4^{2-} + 4H^+ + 2e$ 

The overall reaction is

$$2PbSO_4 + 2H_2O \rightarrow Pb(s) + PbO_2(s) + 2H_2SO_4$$

During this process, lead (Pb) is deposited at the cathode and PbO<sub>2</sub> is deposited at the anode and H<sub>2</sub>SO<sub>4</sub> is regenerated. A 12 volts Lead-acid battery is generally used, which consists of six cells.

**Applications:** Automobiles and construction equipment, standby/backup systems. Used mainly for engine batteries, these cells represent over half of all battery sales. Some advantages are their low cost, long life cycle, and ability to withstand mistreatment. They also perform well in high and low temperatures.

#### **Advantages**

- These cells have a low cycle life, a quick self discharge, and low energy densities. However, with a nominal voltage of 2V and power densities of up 600 W/kg, the lead-acid cell is an adequate, if not perfect, design for car batteries.
- Rechargeable, portable constant potential at low cost.

#### **Disadvantages**

• Use of H<sub>2</sub>SO<sub>4</sub> is dangerous its usage is fragile.

# Q. 7. Describe the construction and working of Ni-Cd battery along with reactions involved during discharging and charging.

**Nickel-Cadmium** (**Nicad**) **Cells:** It is a rechargeable secondary cell. It consists of a Cadmium (Cd) as the anode and Nickel oxyhydroxide (NiO(OH) acts as cathode. KOH is used as an electrolyte. The cell reaction during discharging and charging are as follows.

**Anode:** Cadmium

Cathode: Nickel oxy-hydroxide (NiO(OH))

**Electrolyte**: Aqueous potassium hydroxide (KOH)

**EMF**: 1.4V

#### **Discharging:**

At anode:  $Cd + 2OH^{-} \rightarrow Cd(OH)_2 + 2e$ - (during discharging)

At cathode:  $2NiO(OH) + 2H_2O + 2e \rightarrow 2Ni(OH)_2 + 2OH^{-}(during discharging)$ 

\_\_\_\_\_

Net reaction: Cd +2NiO(OH)+2H<sub>2</sub>O  $\rightarrow$  2Ni(OH)<sub>2</sub>+Cd(OH)<sub>2</sub>

**Charging:** 
$$Cd(OH)_2 + 2e - \rightarrow Cd + 2OH^- (during charging)$$

 $2Ni(OH)_2 + 2OH \rightarrow 2NiO(OH) + 2H_2O + 2e$ - (during charging)

The overall reaction :  $Cd(OH)_2 + 2Ni(OH)_2 \rightarrow 2H_2O + Cd + 2NiO(OH) + 2H_2O$ 

**Applications:** Calculators, digital cameras, pagers, laptops, tape recorders, flashlights, medical devices (defibrillators), electrical vehicles, space applications **Advantages** 

- Good performance in high-discharge and low-temperature applications.
- They also have long shelf and use life.
- Small size
- High rate of charge/discharge capacity
- Higher energy efficiency

**Disadvantages** are that they cost more than the lead –acid battery and have lower power densities. Possibly it's most well-known limitation is a memory effect, where the cell retains the characteristics of the previous cycle. This term refers to a temporary loss of cell capacity, which occurs when a cell is recharged without being fully discharged. This can cause cadmium hydroxide to passive the electrode, or the battery to wear out.

# Q. 8. What are fuel cells? Explain the construction and working of $H_2$ - $O_2$ fuel cell and its applications.

A fuel cell is an electrochemical cell which converts chemical energy contained in readily available fuel oxidant system into electrical energy.

**Principle:** The basic principle of the fuel cell is same as that of an electrochemical cell. The fuel cell operates like a galvanic cell. The only difference is that the fuel and the oxidant are stored outside the cell. Fuel and oxidant are supplied continuously and separately to the electrodes at which they undergo redox reactions. Fuel cells are capable of supplying current as long as reactants are replenished.

Fuel+Oxidant→ OxidationProducts+Electricity

Examples: (1) H<sub>2</sub>- O<sub>2</sub>fuel cell

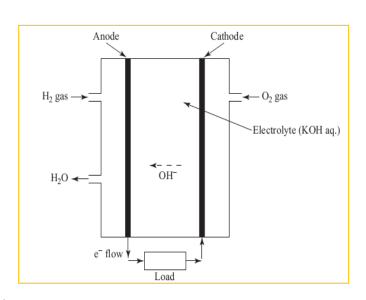
(2) Propane – O<sub>2</sub>fuel cell

(3) CH<sub>3</sub>OH – O<sub>2</sub>fuel cell

#### Hydrogen-oxygen fuel cell:

The best example of fuel cell is Hydrogenoxygen fuel and in which hydrogen gas is used as a fuel and oxygen as oxidant. A schematic diagram of H<sub>2</sub>-O<sub>2</sub> fuel cell is shown below;

The cell consists of a two inert porous electrodes (made of graphite impregnated with finely divided platinum or Nickel) and an



electrolyte 2.5% KOH solution. The hydrogen gas fuel is continuously supplied at the anode and oxygen gas is supplied at the cathode. As hydrogen gas diffuses through the anode, it is adsorbed on the electrode surface and reacts with hydroxyl ions to form water. At the cathode oxygen diffusing through the electrode is adsorbed and reduced to hydroxyl ions. These electrode reactions are summarized below;

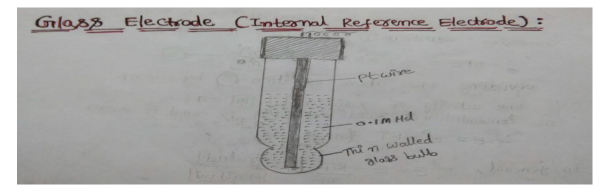
At the anode: 
$$2H_2\rightarrow 4H^++4e^ 4H^++4OH^-\rightarrow 4H_2O$$
 $2H_2+4OH^-\rightarrow 4H_2O+4e^-$ 
At the cathode:  $O_2+2H_2O+4e^-\rightarrow 4OH^-$ 
The net cell reaction is,  $2H_2+O_2\rightarrow 2H_2O$ 

The product discharged is  $H_2O$  and standard e.m.f of the cell is  $E^0=1.23$  volts. A number of such fuel cells are connected together in series to make a fuel battery.

#### **Advantages of Fuel Cells:**

- No emission of toxic gases and chemical wastes are in safe limits.
- They have less weight, so these cells are preferred for space crafts and the product of H<sub>2</sub>O is a valuable water source for astronauts.
- The reactants and products are environment friendly and only we have to bother about disposal of cell material.
- High efficiency (75-85%) of energy conversion from chemical energy to electrical energy. So offer an excellent use of our renewable energy resources.
- No noise pollution like in generators and low thermal pollution.
- Low maintenance costs, fuel transportation costs, cell parts are modular and exchangeable.
- The fuels and electrolyte materials are available in plenty and inexhaustible unlike fossil fuels.
- Unlike acid cells used in automotives the fuel cells are far less corrosive.
- Unlike nuclear energy, fuel cell energy is economical and safe.

# Q. 9. Explain the construction, working of glass electrode and the determination of pH using glass electrode.



Glass electrode is widely used for pH measurements by combining with calomel electrode. The glass electrode is made of a special glass, with the composition Na<sub>2</sub>O (22%), CaO (6%) and SiO<sub>2</sub> (72%). The glass electrode consists of thin-walled glass bulb and relatively low melting point and high electrical conductivity. Glass electrode containing a Pt wire in a 0.01 M HCl. The glass electrode is represented as

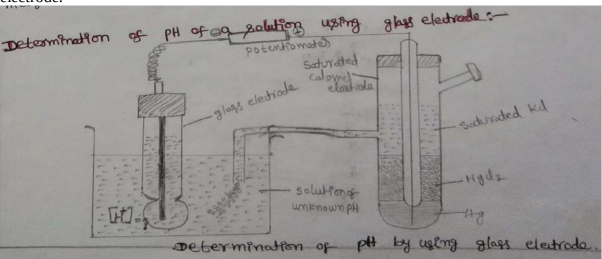
#### Pt, 0.1M HCl/Glass

HCl in the bulb furnishes a constant  $H^+$  ion concentration. Glass electrode is used as the "internal reference electrode". When glass electrode is immersed in another solution determined for pH values especially coloured solutions containing oxidizing (or) reducing agents. The glass membrane of the glass electrode undergoes an ion exchange reaction between the  $Na^+$  ions of glass and  $H^+$  ions in solution. The potential difference varies with the  $H^+$  ion concentration and its e.m.f is given by the expression.

Glass membrane 
$$^-$$
—Na $^+$  + H $^+$  — Glass membrane  $^-$ —H $^+$  + Na $^+$ 

$$E_G = E^0G - 0.0592 log [H]^+$$
(or)
$$E_G = E^0G + 0.0592 pH \qquad \text{so} \quad [-log H^+ = pH]$$

From the equation it is clear that the pH of a solution is a direct measure of e.m.f of a glass electrode.



The glass electrode is placed in the solution under test and is coupled with saturated calomel electrode as shown in the figure above.

The e.m.f of the cell reaction is measured. From the e.m.f , the pH of the solution is calculated as follows.  $E_{cell} = E_{(Right)} - E_{(Left)}$ 

$$\begin{split} E_{cell} &= E_{(calomel)} - E_{(glass~electrode)} \\ E_{cell} &= E_{(calomel)} - E(E^0G~+0.0592~pH~) \\ pH &= \underbrace{E(Calomel) - E^0G - E_{cell}}_{0.0592} \end{split}$$

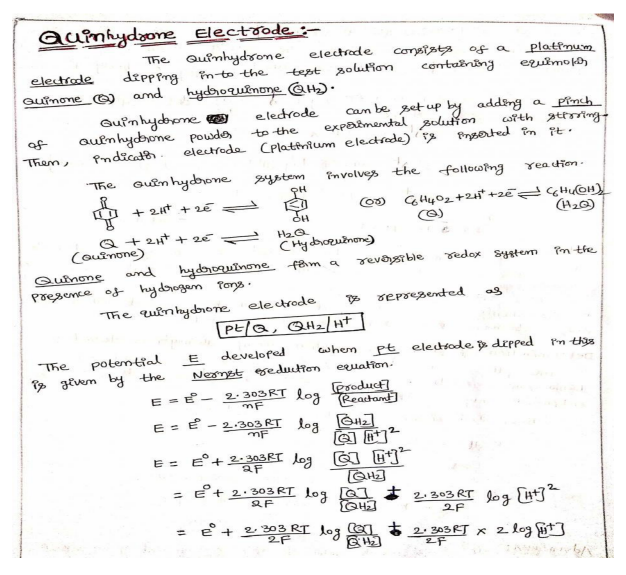
Where E<sup>0</sup>G is a constant, which is determined by using a solution of known pH.

Advantages: (a) It can be easily constructed and readily used

- (b) The results are accurate
- (c) Equilibrium is rapidly achieved.
- (d) Glass electrode is easy to operate.

Disadvantages: (a) The resistance is quite high, special electronic potentiometers are required for measurements. (b) The glass electrode can be used in solutions only pH range of 0 to 10. However, above the pH 12 (high alkalinity), cations of the solution affect the glass and make it the electrode useless.

Q10. Explain the construction and working principle of Quinhydrone electrode and to determination of PH of solution by using quinhydrone electrode.



```
[ log [it] = 2 log [it] and hydroquione one usually taken in equinolar
     Quinone
                                                                      0
          i.e., [] = [] the quantity [] pg unity =1
 mantities.
 E= E + 2.303RT log 1 $ 2.303RT de log [HT]
E= E + 0 + 2.303RT x2 log [HT]
                                                                      0
              C
               E = E° + 0,0592 log [#] at 25°C
          (00)
E = E° ★ 0.0592 PH
                                                - log 田丁 = PH
                                                                      C
          E° = Standard electrode potential of quentydrone electrode
                                                                      0
                   at 25°C. But reduction potential 98 = +0.69962
                                                                      0
  =-0.6996 Volt
                                                                      C
             E = 0.6996 - 0.0592 PH
* The quinkydrome electrode depends upon the concentration of hydrogen
        Pt can be used for the determination of PH value
                                                                      0
Determination of pH of a solution using Quinhydrome electrode?
                                                                      C
    The PH of a solution is determined by connecting the
                                                                      0
quentydowne electrode with a saturated calomel electrode. This
                                                                      0
Combination may be represented as
                                                                      6
  Pt/OHz, Q, Ht (unknown) / Kd (saturated), Hgz Uz(5) / Hgt
                                                                      0
                                                                      6
       : ECell = Eright-Elett
                                                                      6
             Ecel = Ecolomel) - Equintydopre
              Eccu = 0.2422 - (0.6994 -0.0592 PH)
               Ecell + 6.6994 -0.2422 = 0.0592 PH
                                                                      0
                                                                      0
                     pH = Ecell + 0.6994 - 0.2422
                                                                      (Z
                                  0.0592
                It & useful putte tetration of an aid by an alkali.
            ů)
Advantages:
                 Equilibrium is sapidly attained
                 The pt value obtained is very accurate.
```

Dis advantages: (1) It gives very good result in solution only when pH is less than 8. In more alkaline solution equilibrium between a and H2Q is disturbed.

Q11. Explain the electrochemical series and its applications.

### Electrochemical Sexies:

when the metals one of an in the order of the respect to one motor solutions of in soies that the middle electrochomical soies. In this soies, hydrogen is situated in the middle having zero, electrode potential. Any metal above hydrogen will displace hydrogen from dilute aid solution. For the Na reads with water to likewate hydrogen because the Eoof Nat/Ng (-2.714 V) is less tham Eoof Ht/Hz (zero).

In the socies will be reduced by hydrogen.

The higher page was the home	they undergo <u>corrosion</u> eagly.	ps to be oxidized
	undand <u>oxidation</u> potentials at	25°c
Electrode	HalfCell reaction	E Volts
12/12+	B→ E+E	+3.045
K(K+	K→ K+E	+ 2.925
	ca -> cat2+2E	+2.87
ca/cat2		+2.714
Nalnat	Na -> Nat +e	+2.37
mg/mg+2	$Tng \rightarrow mg^{+2} + 2e^{-}$	+0.763 AUM
zn zntz	2m -> 2n+2+2e-	amo
	Fe-> Fe+2+2e-	40.440
Fe/ Fet2	Fe Fe + 72	40.403
cd/cd+2	$cd \rightarrow cd^{+2} + 2e^{-}$	+0.126
Pb/Pb+2	Pb-> Pb+2+2e	0.000
Pt Hzg Ht (im)	H2(1atm) -> 2H+ +2E-	0.000
, Clother Cas	1 2+ . 25	-0°337->
culat2	Cupo cu2++2E	-0.799
Agl Ag+7	Ag -> Ag+1 +e-	7
1 -1 2+	20 -> U2 + 2E	-1.36 \ Nob
Pt/u-/u2+	$Au \rightarrow Aut^3 + 3e^-$	-1.50 / Catho
ANY Act 3	Au -> Au. 7 50	

The higher a metal is in the soils, the greater tendency to be oxidized. The metals highly in the soiles are strong reducing agents, and their rons are stable whereas those mean the bottom of the soiles are inactive, and their rons are easily reduced to metals.

Applications of electrochemical socies: (1) The relative corrosion tendencies of the metals and allays. (2) Relative ease of oxidation (05) reduction of metals. (3) Replacement tendency to metals. (4) alculating the equilibrium constant as given below.

E = RT In Kee)

E = 2.303 RT log 10 Kee)

Log 10 Kee) MEE

2.303 RT

Log 10 Kee) ME

2.303 RT

Acg 10 Kee) - ME

0.0591

Dysedicting spontamenty of redox reautisms.

#### **CORROSION**

#### **Types of corrosion (Theories of corrosion):**

The corrosion process proceeds in two types (1) **chemical (Dry corrosion)** and (2) **electrochemical (wet corrosion)** attack of environment.

## Q.1. EXPLAIN THE MECHANISM OF CHEMICAL CORROSION (DRY CORROSION).

This type of corrosion occurs mainly through the direct chemical action of environment/atmospheric gases such as oxygen, halogen, hydrogen sulphides, Sulphur dioxide, nitrogen or anhydrous inorganic liquids.

There are three main types of chemical corrosion: (a) oxidation corrosion

- (b) corrosion by other gases
- (c) liquid metal corrosin

#### a. Oxidation corrosion:

"Direct action of oxygen at high or low temperatures on metals in the **absence of moisture** is called oxidation corrosion". Metals such as Ag, Au, and Pt are not oxidised as they are noble metals.

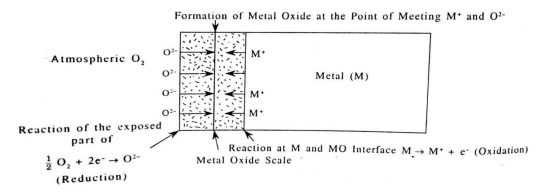
$$2 \text{ M} \longrightarrow 2 \text{ M}^+ + 2 \text{ e}^- \text{ (Loss of electron: Oxidation)}$$

$$O_2 + 2 \text{ e}^- \longrightarrow 2 \text{ O}^{2-} \text{ (Gain of electron: Reduction)}$$

$$2 \text{ M} + O_2 \longrightarrow 2 \text{ M}^+ + 2O^{2-} \longrightarrow 2\text{MO(metal oxide layer)}$$

**Mechanism**: The oxidation of the metal occurs at the surface first, resulting the formation of metal oxide scale, which restricts further oxidation.

Further oxidation to continue, either the metal must diffuse outwards through the scale to the surface or the oxygen must diffuse inwards through the scale to the underlying metal. Of the two types of diffusions, the diffusion of the metal is rapid because the size of the metal ion is smaller than oxygen ion, hence higher mobility to metal ion.



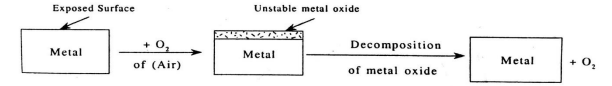
Nature of the oxide formed plays an important part in oxidation corrosion process. The following types of films are there:

1. **Stable metal oxide**: If the formed metal oxide is stable, it prevents further corrosion due to tightly bind to the metal surface is impermeable to the further attacking of oxygen. Fe (iron) corrodes

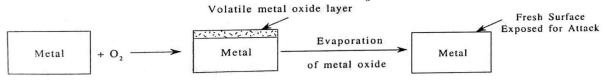
faster than Al (aluminium) because Al forms tightly adhering protective Al<sub>2</sub>O<sub>3</sub> film on the Al metal surface.



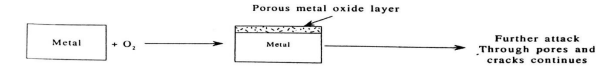
**2. Unstable metal oxide**: The oxide layer formed decomposes back into the metal and oxygen. Consequently, oxidation corrosion is not possible in such a case. Thus, Ag, Au, and Pt do not undergo oxidation corrosion.



**3. Volatile oxide layer:** The oxide layer formed evaporates as soon as it is formed, leaving the underlying metal surface exposed for further attack. This causes rapid and continuous corrosion, leading to excessive corrosion. Ex: Molybdenum oxide (MoO<sub>3</sub>) is volatile.



4. **Porous oxide layer**: If the formed metal oxide layer is porous, it allows further corrosion. A porous layer of metal oxide contains cracks, gaps etc. provides access to oxygen to reach the underlying surface of the metal. Eg. Alkaline earth metals and alkaline metals.



**Pilling Bedworth rule**: According to Pilling-Bedworth rule, a metal which possess high specific volume ratio does not undergo further corrosion (or) prevent further corrosion.

## Specific volume ration=Volume of metal oxide Volume of metal

If the volume of the metal oxide layer is at least as great as the volume of metal, which is formed non-porous and protective layer by tightly adhering to the base metal.

Specific volume ratio is more the rate of corrosion is less. For example, the specific volume ratios of W, Cr and Ni are 3.6, 2.0 and 1.6 respectively. Hence, the rate of corrosion is least in Tungsten (W).

#### 2. Corrosion by other gases:

The gases like  $SO_2$ ,  $CO_2$ ,  $Cl_2$ ,  $H_2S$  and  $F_2$  are causes for chemical corrosion and these corrosion mainly depends on the reactivity the gas. Eg:  $Cl_2$  attacke the metal (Sn) forming volatile  $SnCl_4$  and Ag metal produces adherent ,non porous oxide film (AgCl).

#### 3. Liquid metal corrosion:

The corrosion reaction involves either dissolution of a solid metal by a liquid metal or internal penetration of the liquid metal into the solid metal. Both these modes of corrosion cause weakening of the solid metal. Eg: Liquid metal mercury dissolves most metals by forming amalgums, there by corroding them.

## Q. 2. EXPLAIN THE MECHANISM OF ELECTROCHEMICAL CORROSION (WET CORROSION) TAKING IRON (Fe) AS AN EXAMPLE.

Wet or electrochemical corrosion takes place under wet or moist conditions through the formation of short circuited electrochemical cells. According to this theory, there is formation of galvanic cell on the surface of metal. Some of the metal surface act as anode and rest act as cathode. The chemical in the environment and humidity acts as an electrolyte. Oxidation takes place at anode and it results in corrosion, while reduction takes place at cathode.

Electrochemical corrosion involves flow of electron-current between the anodic and cathodic areas. The anodic reaction involves in dissolution of metal as corresponding metallic ions with the liberation of free electrons.

At anodic area: 
$$M \longrightarrow M^{n+} + n e^{-}$$
 (Oxidation)

On the other hand, the cathodic reaction either H<sup>+</sup>, O<sub>2</sub>, H<sub>2</sub>O consumes the electrons generating from anode to produce non-metallic ions like OH<sup>-</sup> or O<sup>2-</sup> either by evolution of hydrogen or absorption of oxygen, depending on the nature of corrosive environment.

Wet corrosion takes place by the following ways.

#### a) Evolution of hydrogen at cathode:

This type of corrosion occurs usually in acidic environments. The Rusting of iron takes place in acidic medium. Considering metal like Fe the anodic reaction is dissolution of iron as ferrous ions with the liberation of electrons.

Fe 
$$\longrightarrow$$
 Fe<sup>2+</sup> + 2e (Oxidation)

These electrons flow through the metal, from anode to cathode, where  $H^{+}$  ions are eliminated as hydrogen gas.

$$2H^{+} + 2e^{-} \longrightarrow H_{2}$$
 (Reduction)  
The net overall reaction is: Fe + 2H<sup>+</sup>  $\longrightarrow$  Fe<sup>2+</sup> + H<sub>2</sub>

In Hydrogen Evolution type of corrosion the anodes are usually large and the cathodes are small areas.

In Neutral medium at the cathode formation of OH- takes place along with the liberation of hydrogen gas.

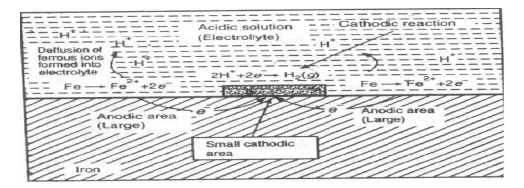
$$2H_2O + 2e^{-} \longrightarrow 2OH_1 + H_2$$
 (Reduction)

The Fe<sup>2+</sup> and OH<sup>-</sup> diffuse towards each other through the conducting medium forming the corrosion product rust in between the cathodic and anodic areas.

$$Fe^{2+} + 2OH^{-} \longrightarrow Fe(OH)_{2} (Ferrous Hydroxide)$$

$$4 Fe(OH)_{2} + O_{2} + 2H_{2}O \longrightarrow 4 Fe(OH)_{3} \longrightarrow 2(Fe_{2}O_{3}. 3H_{2}O) \text{ rust}$$

$$Ferric Hydroxide \qquad Hydrated ferrous oxide$$



In this case, metals react with acidic and neutral conditions to release H<sub>2</sub> gas. All metals above hydrogen in electrochemical series can show this type of corrosion.

**b) Absorption of oxygen at cathode**: This type of corrosion takes place in neutral and basic medium in the presence of oxygen. The oxide of iron covers the surface of the iron. The small scratch on the surface creates small anodic area and rest of the surface acts as cathodic area.

The following chemical reactions occur at anode and cathode.

Fe 
$$\longrightarrow$$
 Fe<sup>2+</sup> + 2e (Oxidation)

The liberated electrons flow from anodic to cathodic areas, through iron metal, where electrons are intercepted by the dissolved oxygen as:

$$O_2 + H_2O + 2e^{-} \longrightarrow 4OH$$
 (Reduction)

The Fe<sup>2+</sup> ion at anode and OH ions at cathode diffuse and when they meet, ferrous hydroxide is precipitated.

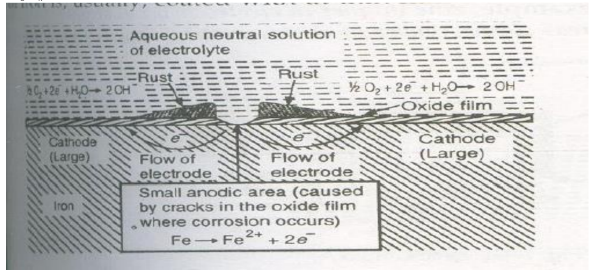
$$Fe^{2+} + 2OH$$
 Fe(OH)<sub>2</sub>

If enough oxygen is present, ferrous hydroxide is easily oxidized to ferric hydroxide.

$$4 \operatorname{Fe(OH)}_{2} + \operatorname{O}_{2} + 2\operatorname{H}_{2}\operatorname{O} \longrightarrow \operatorname{Fe(OH)}_{3} \longrightarrow 2(\operatorname{Fe}_{2}\operatorname{O}_{3}.3\operatorname{H}_{2}\operatorname{O})$$

This product is called yellow rust, actually corresponds to 2(Fe<sub>2</sub>O<sub>3</sub>. 3H<sub>2</sub>O)

If the supply of oxygen is limited the corrosion product may be even black anhydrous magnetite,  $\mathrm{Fe_3O_4}$ 



#### Q. 3. Explain the Factors influencing rate of corrosion.

Since corrosion is a process of destruction of metal surface by its environment, the two factors that govern the corrosion process are:

(i) Nature of Metal and (ii) Nature of Environment.

#### 1. Nature of the metal:

#### (i)Position in galvanic series:

When the oxidation potentials are arranged in the decreasing order of activity in a series of metals called as galvanic series.

Activ	e 1.	Mg
(or anod	lic) 2.	Mg alloys
1 1	3.	Zn
	4.	Al
	5.	Cd
	6.	Al alloys
1 1 1	7.	Mild steel
1 1 1	8.	Cast iron
	9.	High Ni cast iron
1 11	10	. Pb-Sn solder
1 11	11	. РЬ
	12	. Sn
1 11	13	. lconel
1 1 1	14	. Ni - Mo - Fe alloys
1 1	15	. Brasses
1 11	16	. Monel (7 = Ni, 30 = Cu, rest = Fe)
1 1 1	17	. Silver solder
1 1 1	18	. Cu
1 1 1	19	. Ni
1 1 1	20	Cr stainless steel
1 1 1	21	18 – 8 stainless steel
	22	18 – 8 Mo stainless steel
	23	Ag
	24	Ti
	25	Graphite
Noble	26	Au
(or catho	lic) 27	. Pt

In the galvanic series series, a metal high in the series is more anodic and undergoes corrosion faster than the metal below it. For example, **Li** corrodes faster than **Mg**; **Zn** corrodes faster than **Fe**; **Fe** corrodes faster than **Sn**; **Cu** corrodes faster than **Ag** and so on. Also more difference in the position of galvanic series will cause faster corrosion at anodic metal.

(ii) **Relative areas of the anodic and cathodic parts:** When two dissimilar metals (or) alloys are in contact, the corrosion of the anodic part is directly proportional to the areas of the cathodic and anodic parts. Smaller the area of anode compared to cathode will lead to faster corrosion of anode. Because the larger cathodic area demands more electrons, which will be fulfilled by fast corrosion at anode (oxidation).

Corrosion a <u>Area of cathode</u> Area of anode (iii) **Purity of metal: Impurities in a metal**: Pure metal resists corrosion, while impurities in a metal form a local galvanic cell (metal as anode and impurity as cathode) and result in the corrosion of metal.

Purity less: more corrosion Purity more: corrosion less

(iv) **Physical state of metal**: The rate of corrosion is influenced by physical state of the metal. The smaller the grain-size of the metal or alloy, the greater will be its solubility and hence, greater will be its corrosion.

More stress: more corrosion Less stress: less corrosion

- (v) **Nature of surface film**: The ratio of the volumes of the metal oxide to the metal is known as a "specific volume ratio." Greater the specific volume ratio, lesser is the oxidation corrosion rate. According to Pilling- Bedworth rule the volume of oxide film is greater than metal from which metal oxide formed, then the film is protective.
- (vi) **Solubility and volatility of corrosion products**: In both the cases, the corrosion will be faster. MoO<sub>3</sub>, SnCl<sub>4</sub> are volatile, so faster is corrosion of **Sn** in Cl<sub>2</sub> atmosphere. In case of soluble corrosion product by water and metal surface will be exposed to further corrosion.
- **2. Nature of the environment:** The role of environment in the corrosion of a metal is very important. Environmental parameters like temperature, humidity, PH etc. play important role.
- (i) **Temperature and humidity**: The rate of diffusion increase by rise in temperature, hence the rate of corrosion is also increased. But higher temperature reduces the concentration of  $O_2$  and hence corrosion is reduced. In humidity gases like CO2, SO2, NOx are dissolved which form electrolytes. It will cause galvanic corrosion.
- (ii) **Influence of pH**: PH value means concentration of H<sup>+</sup> (acidic nature). In acidic medium (less than 7), corrosion is faster. In basic medium, some metal such as Pb, Zn, Al, etc. form complexes and hence they corrode.

Eg: Zn corrodes minimum at P<sup>H</sup> 11, but at higher P<sup>H</sup> (more than 11) it corrodes faster. At P<sup>H</sup> 5.5, Al corrodes minimum.

- (iii) **Nature of ions present in vicinity**: Cu<sup>+2</sup> ions present in the vicinity (medium) of Fe, accelerate corrosion, while silicates present in the medium resist corrosion.
- (iv) Conductance of corroding medium: Due to presence of salts and water in earth, it is of conducting nature. More conductance leads to more stray current and hence fast corrosion. Dry sandy soil is less conducting and hence less corrosion, while mineralised clay soil is more conducting hence more corrosion occurs.
- (v) Oxygen concentration: Oxygen is one of the important element responsible for corrosion. The % of  $O_2$  in atmosphere increases, the rate of corrosion also increases due to formation of Oxygen concentration cell on the surface of metal (iron rod half dipped in water corrodes due to this effect).

Anode Fe 
$$\longrightarrow$$
 Fe<sup>2+</sup> + 2e (Oxidation)  
Cathode  $\frac{1}{2}$  O<sub>2+</sub> H<sub>2</sub>O + 2e  $\longrightarrow$  2OH  $\longrightarrow$  Fe +  $\frac{1}{2}$  O<sub>2+</sub> H<sub>2</sub>O  $\longrightarrow$  Fe<sup>2+</sup> + 2OH  $\longrightarrow$ 

$$Fe^{2+} + 2OH \longrightarrow Fe(OH)$$

If enough oxygen is present, ferrous hydroxide is easily oxidized to ferric hydroxide.

$$4 \operatorname{Fe(OH)}_{2} + \operatorname{O}_{2} + 2\operatorname{H}_{2}\operatorname{O} \longrightarrow \operatorname{Fe(OH)}_{3} \longrightarrow 2(\operatorname{Fe}_{2}\operatorname{O}_{3}.3\operatorname{H}_{2}\operatorname{O})$$

This product is called yellow rust, actually corresponds to 2( Fe<sub>2</sub>O<sub>3</sub>. 3H<sub>2</sub>O)

- (vi) **Polarisation of the electrode:** More the polarisation at electrodes, less current passed and hence less corrosion.
- (vii) **Flow of process stream:** The velocity of the diffusion of ions in the medium increases, the rate of corrosion increases.

#### **CORROSION CONTROL METHODS:**

The following some of the important corrosion control methods are described:

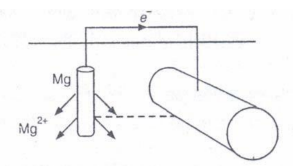
- ✓ Cathodic Protection
- ✓ Protective coatings or metallic coatings
- ✓ Using Pure metal & alloys
- ✓ Modifying the Environment
- ✓ Use of Inhibitors
- ✓ Proper Designing

✓

#### Q. 4. Explain the Cathodic protection:

"The principle involved in this method is that the metal is forced to behave like cathode, thereby preventing corrosion". There are two types of cathodic protection.

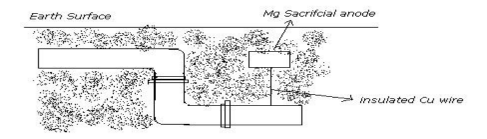
- (a) Sacrificial anodic protection
- (b) Impressed current cathodic protection
- **(a) Sacrificial anodic protection**: In this method of protection, the metallic structure to be protected called "base metal" is connected to more anodic metal through a wire, so that all the corrosion is concentrated at the more anodic metal. As the more anodic metal is sacrificed in the process of saving base metal from corrosion. Hence, it is known as *sacrificial anode*. Metals commonly used as sacrificial anodes are Zn, Al,Mg.
- Zn (or) Mg metals are more positive than Fe and, hence in electrochemical it will act as an anode and the Fe will act as cathode. According to the principle of galvanic cell, anode that undergo corrosion and saving cathode from corrosion. Thus, Mg sacrifies itself for saving the Fe.



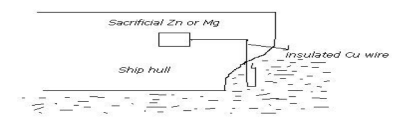
In cathodic protection, an anode of a more strongly reducing metal is sacrificed to maintain the integrity of the protected object (e.g. a pipeline, bridge, ship hull or boat).

Applications: Important applications of this technique are

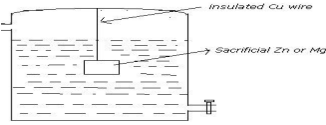
a. Protection of underground cables and pipelines from soil corrosion



b. Protection of ships and boat hulls from marine corrosion. Sheets of Mg or Zn are hung around the ship hull, these sheets being anodic to corrosion so these sheets get corroded, when consumed completely, these are replaced by fresh one.



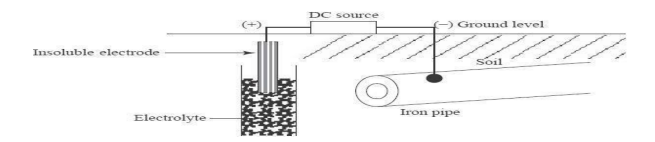
c. The formation of rusty water is prevented by the insertion of Mg sheets or rods into domestic water boilers or tanks.



**Advantages:** (1) It is a simple method (2) It has low maintainance and installation cost (3) It does not required external power

**Disadvantages**: (1) Some times more than one anode is required (2) Current out put is less (3) It does not work properly in high corrosive environment (4) The sacrificial anode must be replaced periodically, when it is consumed.

#### (b) Impressed current cathodic protection:



In this method the metal to be protected (the metal is exposed to soil (or) other electrolyte) is made anode by connecting to an external battery. The battery contains an inert graphite anode, which is buried in a back fill to maintain electric contact. A known quantity of direct current is passed through the electrolyte, when the applied current is equal to the current, the corrosion of the metal stops.

Impressed current cathodic protection is given to (i) water tanks (ii) transmission line towers (iii) buried pipe lines and (iv) marine pipes.

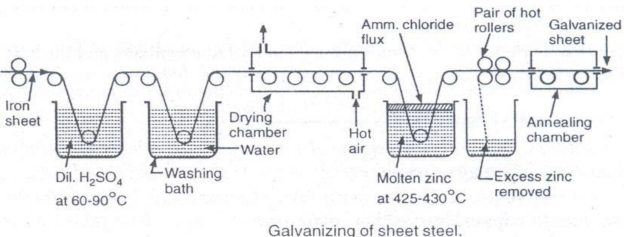
#### Q. 5. Explain the protective coatings (or) metallic coatings.

Protective coatings act as a barrier between the coated surface and its environment. Some of the methods used for imparting coating of one metal over another. Metallic coating are applied by

- (1) Hot dipping
- (2) Electroplating (Cu)
- (3) Electrolessplating (Ni)
- (4) Paints
- (5) Metal Cladding
- (6) Cementation
- (1) Hot dipping: It is one of the common methods for applying metallic coatings on the surface of base metal. In this method, molten metals such as Zn (m.p.=419°C), Sn (m.p.=232°C) Pb, Al, etc., having low melting points than the coated metals such as on iron (Fe), steel and copper (Cu) which have relatively higher melting points.

Galvanizing: "The process of coating Fe (iron) with Zn (zinc) is known as galvanization". The method of coating iron (Fe) or any base metal with Tin (Sn) is called Tinning.

The base metal iron or steel sheet is cleaned by acid pickling method with dil.H<sub>2</sub>SO<sub>4</sub> for 15-20 minutes at 60-90 °C. The sheet is then washed well and dried. It is dipped in a bath of molten zinc maintained at 425-435 °C. The surface of the bath is kept covered with ammonium chloride (NH<sub>4</sub>Cl) flux to prevent oxide formation. The sheet is taken out and excess Zn is removed by passing it between a pair of hot rollers. Then the sheet is subjected to annealing process at cooled slowly.



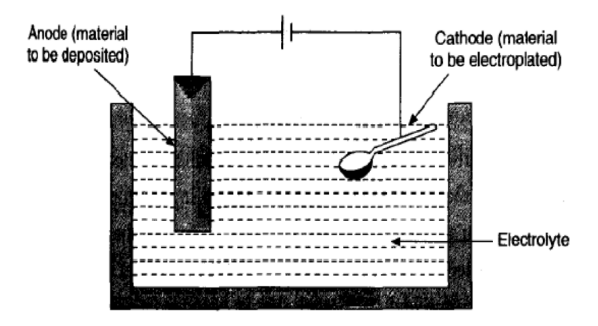
Applications: It is mostly used to protect iron used for roofing sheets, wires, pipes, nails, bolts, screws, buckets and tubes.

Galvanizing utensils cannot be used for preparing and storing food stuffs especially acidic in nature, because zinc dissolves to form highly toxic or poisonous compounds.

(2) Electroplating (or) Electrodeposition: "The process of depositing the coating metal on the surface of base metal/non metal by electrolysis is called electroplating".

It is an electrochemical process in which a base metal is coated by Zn, Ag, Cr, Au, Sn etc. to protect it from corrosion and also to make it shine and decorative. The base metal is made cathode, dipped in a suitable electrolyte, and the metal to be deposited is made anode (or) an inert material good electrical conductivity material is also used as an anode. A direct current (DC) is passed through the electrolytic solution.

The article to be electroplated to solvent cleaning by trichloroethylene to remove oil, grease etc., Then it is subjected to acid pickling with dil. HCl (or) dil. H<sub>2</sub>SO<sub>4</sub> to remove any oxides etc., The cleaned article is made cathode of the electrolytic cell and electrodes were dipped in the salt solution of the coating metal, which acts as an electrolyte. When direct current is passed, the coating metal ions migrate to the cathode and deposit on the base metal article in the form of a thin layer.



For eg: electroplating of copper on iron articles, the following are maintained.

At anode:  $M \longrightarrow M^{n+} + ne^{-}$  (dissolution of metal)

At cathode:  $M^{n+}+ne^{-} \longrightarrow M(deposition on substance)$ 

1. Electrolytic bath solution: copper sulphate

2. Temperature maintained: 40-70 °C

3. Current density: 20-30 mA/cm<sup>2</sup>

4. Cathode: base metal

If inert electrode used as anode copper sulphate to replenish it.

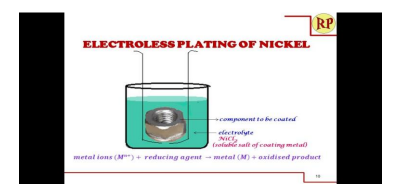
**Application:** Electroplating is a most important and frequently used technique in industries to produce metallic coatings. Both metals and non metals can be electroplated. In metals the electroplating increase resistance to corrosion, chemical attack, hardness, wear resistance and surface properties. In non metals electroplating increases strength and decorates the surface of non metals like plastics, wood, glass etc.

(3) Electrolessplating (Ni): In this method of depositing a metal from its salt solution on a catalytically active surface by a suitable reducing agent without using electrical energy is called electroless plating.

$$M^{n+}$$
 + Reducing agent  $\rightarrow M$  + oxidized product.

The reducing agent supplies electrons for the reduction of metal ions to metal. The metal gets deposited on the surface of the base metal there by producing a uniform thin coating. Eg: Ni, Co, Cu and some precious metals can be deposited by electroless plating.

The base object, a plastic material like acrylonitrile butadiene styrene (ABS) can be coated with nickel. The base metal surface is activated by stannous chloride (SnCl<sub>2</sub>) followed by palladium chloride (PdCl<sub>2</sub>) to get a layer of palladium and the surface is dried. The base metal is dipped in a solution of NiCl<sub>2</sub> (nickel chloride), sodium hypophosphite (reducing agent), sodium acetate buffer, sodium succinate (complexing agent) PH of the bath maintained is 4.5 and temperature is 93 <sup>0</sup>C. The following reactions takes place and nickel get plated electrolessly on the surface of the base object.



$$Ni^{+2} + H_2PO^{2-} + H_2O$$
  $\longrightarrow$   $Ni + H_2PO^{3-} + 2H^+$ 

Applications: (i) They are used in electronic industry for fabricating printed circuits and diodes. (ii) ABS plastic coated objects are used for decorative applications automotives, electronics etc.

#### Q. Explain the Types of Corrosions.

Galvanic Corrosion: When two dissimilar metals (eg., zinc and copper) are electrically connected and exposed to an electrolyte, the metal higher in electrochemical series undergoes corrosion. In this process, the more active metal (with more negative electrode potential) acts as a anode while the less active metal (with less negative electrode potential) acts as cathode. In the above example, zinc (higher in electrochemical series) forms the anode and is attacked and gets dissolved; whereas copper (lower in electrochemical series or more noble)acts as cathode.

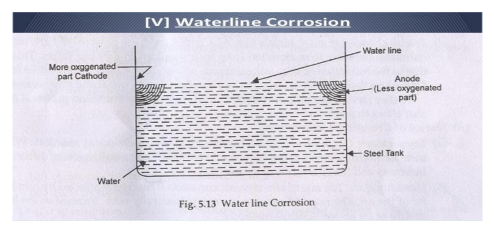
**Mechanism:** In acidic solution, the corrosion occurs by the hydrogen evolution process; while in neutral or slightly alkaline solution, oxygen absorption occurs. The electron-current flows from the anode metal, zinc to the cathode metal, copper.

At anode: 
$$Zn \rightarrow Zn^{2+} + 2^{e-}$$
 (Oxidation) corrosion

Thus it is evident that the corrosion occurs at the anode metal; while the cathodic part is protected from the attack.

Prevention of galvanic corrosion: (i) coupling of metals close to the electrochemical series (ii) fixing insulating material between two metals (iii) using larger anodic metal and smaller cathodic metal.

**Waterline corrosion:** It has been observed in the case of an iron tank containing water, that the portion of iron tank just below the water level undergoes corrosion. It is due to the difference in oxygen concentration. Corroding portion is poor in oxygen and acts as anode.



A distinct brown line is formed just below the water line due to the deposition of rust. The reactions may be represented as follows:

#### **Cell reactions:**

At anode:  $M \rightarrow Mn+ + ne- (Oxidation of metal M)$ 

At cathode: O2 + 2H2O + 4 e→ 4OH- (Reduction of oxygen)

Other example: Ships which remain partially immersed in sea water for a long time undergo water line corrosion.

# **Module-IV**

Stereochemistry, Reaction mechanism & synthesis of drug molecules and NMR spectroscopy

0

Stevenchemistry, a sub discipline of chemistry, that deals with the relative special anxingement of atoms that form the structure of molecules and their manipulation. Stevenchemistry is a part of chemistry which deals with structure in three dimensions ("steven" means three dimensional), one appeal of stevenchemistry is stevenizomerism.

"I somers are different compounds, having some molecular formula but different proporties (physical, chemical, or both) are known as isomers, and the phenomenon is known as isomersm".

Types of isomerism: Isomerism is of two types.

- (a) Structural Promesism (or constitutional Promesism
- (b) Steven someofern (or Spatial Esomeofern.

## (a) Structural iso merism (ds) constitutional isomerism:

The compounds having same molecular formula but possess different structural arrangement of atoms. This difference may be due to for the nature of carbon chain (chain isomerism) or due to the position of the substituent (position isomerism) or due to the difference in the nature of the functional group (functional isomerism).

Chain (somerism: It auxes due to the different in the nature of carbon chain. Example:

- (1) n-butane (C4-H10) CH3-CH2-CH3-CH3
  PROBULTANE (C4-H10) CH3-CH-CH3
  CH3
- (2) m- Perntame (C5 H12) CH3-CH2-CH2-CH3

  Iso Perntame (C5 H12) CH3-CH2-CH3

  CH3

  Neo-Perntame (C5 H12) CH3-CH3

  CH3

  CH3

  CH3

  CH3

This type of isomerism is due to the difference in the nature of the corbon chain (i.e straight con branched) which forms the nucleus of the molecule.

C4 H10 and C5 H12 can exist in 2 and 3 isomeric from respectively.

DECISEO Chemiator SINME

substituent atom (or) group (or) an unsaturated linkage in the carbon chain. Examples: (1) N-propyl iodide (C3H7I) CH3-CH2-CH2-I

Isopropyl Podlde (C3H7I) CH3-CH2-CH3-I

(2) 1- butene (C4H8) CH2=CH-CH-CH3 2- butene (C4H8) CH3-CH=CH-CH3

(3) F/4080 propane (C3H7F) CH3-CH2-CH2-F
2-f/4080 propane (C3H7F) CH3-CH-CH3 maj x = m

(4) N-propyl alcohol (CgH80) CH3-CH2-CH2-OH
Isopropyl alcoho (C3H80) CH3-CH-CH3

(iii) Functional Promerism: compounds having same molecular formula but possess different functional groups.

Examples: (1) Ethyl alcohol (C2H60) CH3-CH2-OH (alcoholic group)

Dimethyl ether (C2H60) CH3-O-CH3 (ether group)

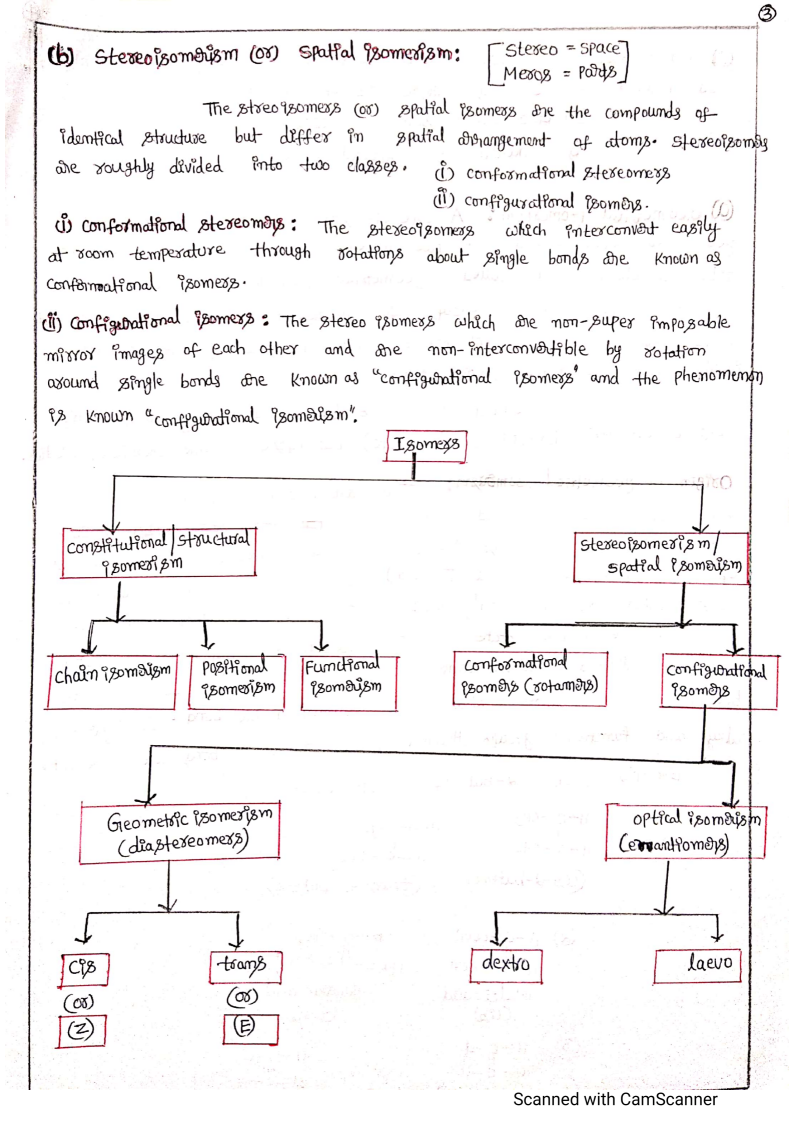
(2) propanaldehyde (C3H60) CH3-CH2-CH0 (althyde group)
Acetone (C3H60) CH3-E-CH3 (Ketone group)
Allyl alcohol (C3H60) CH2=CH-CH2-OH (alcohol)

(3) propromic and (c3 H602) CH3-CH2-COOH (and group)
Methyl acetate (C3 H602) CH3-E-O-CH3 (ester group)

Methyl ethyl amine (C3HqN) CH3-CH2-CH2-NH2 (i amine)
Methyl ethyl amine (C3HqN) CH3-NH-CH2-CH3 (2° amine)
Trimethyl amine (C3HqN) CH3-N-CH3 (3° amine)
CH3

Thus type of isomerism is due to difference in the nature of fundional

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- (a) Gaometrical geometre (Diastereomers)
- (b) Emantformers (optical isomer)

O Geometrical Isomerism: A type of Isomerism due to the different geometrical arrangement of two different groups about the airbon-carbon double bond is called geometrical isomorism on as—trans Isomorism: when similar groups are attached to the same side of molecula it is called the cis (or syn) isomer, while in trans (or and) isomer similar groups lie on the opposite side.

Geometrical Promers differ in their physical proporties (such as M.P., B.P., density, solubility etc.) but possess same chemical proporties.

Oxigin of geometrical isomorism: Two corbon atoms Toined by a single bond can be freely votated around it. Ext other two corbon atoms. But when two corbon atoms one Toined by a double bond (which consists of a sigma-bond and a  $\pi$ -bond), the free votation about around the corbon-corbon axis is hardly possible. The two corbon atoms in element compounds one in the state of sp hybridization. In this way, the atoms (or) groups attached to corbon atoms in compounds containing c=c double bond become fixed up in space. Now alkeness contain double bond. If hydrogen, alkyl and functional groups attached to corbon-corbon bonds one different.

Examples: (1) 2-butene CH3-CH=CH-CH3

Conditions for geometrical isomousm: (1) The molecule must contain a double bond (2) Each of two carbon atoms of double bond must remain attached to different substituent.

Characteristics of Diastereomers: (i) They exhibit similar, but somewhat different chemical proposties (ii) Their rates of reaction one different.

(iii) The physical proposties (Eq: melting point, boiling point, density, solubility, refractive index etc) one different. (iv) They can be separated by I fractional distillation, frational crystallization and chromatography. The relative physical constants of cis and trans gromers are mentioned in the following table.

Physical constants	la a	CL3	1	d	trans	
Melting point	squit-	Lower	Ø,	5	Higher	
Boiling point	11	Higher	Hog	- 3-	Lower	0-3-4
Solubility	3-4	Higher	}	>5\ 500	Lower	* -'- H
Density	. X. k.	Higher	kinnj	dinnit	Lower	· LA
Dipolemoment		માંમાન			Lower	
Refrautive Index	II.	Higher		1-5-	Lower	
- Heat of combustion		Higher	arohi i	b- R.J- c	Lower	
le .	meleti		90	ethyli		

Eg: Cis isomers have higher dipole moments than toans isomers which may have zero dipole moments. For example, the dipole moment of cis-maleic and is quite high while that of fumblic and is zero because dipole moments of C-cook bonds cancel out the effects of each other as these one in opposite directions.

similarly in 1,2-dichloroetheme, the dipole moment of c-d bond is cancelled out in the trans grower.

### Nomenclature:

a cla-trum system: The isomer having the same groups on the same side of the double bondisknown as cla-form, whereas the one in which same group remain on the opposite side of the double bond is known as transform.

to the carbon atoms of a double bond, then cis and teams designations cannot be employed

C=c

6 d

In such cases, cahn, Ingold and prelog developed E and Z
system of nomenclature, which is based on a priority system. In this
system, the two atoms froups attached to each coulon atom of the
doubly bond colons one put in odds of precedence based on sequence
rules.

Of the symbol "E" (from German word Entgegen =
across (m) opposite) is assigned to the isomers in which the atoms/

7

groups of higher precedence one on the opposite side.

(ii) The symbol "z" (from serman word zusammen = together)

Ps assigned to the isomer in which the atoms groups of higher precedence
one on the same side. Thus

Sequence sules: (1) Higher the atomic number, higher is the priority assigned to the atom. For example

Precedence order: I > Br > d > H

Atomic number: 53 35 17 1

(2) In case of groups, the precedence order is decided on the basis of atomic number of first atom of the group. For example precedence order: -d> -503H> -OH> -NHCH3> -COOH

Atomic number of first atom: 17 160 8 Hz 7

(3) If the procedence order of groups cannot be settled on the basis of atomic number of first atom, then second atom (or) the subsequent groups are considered. For example.

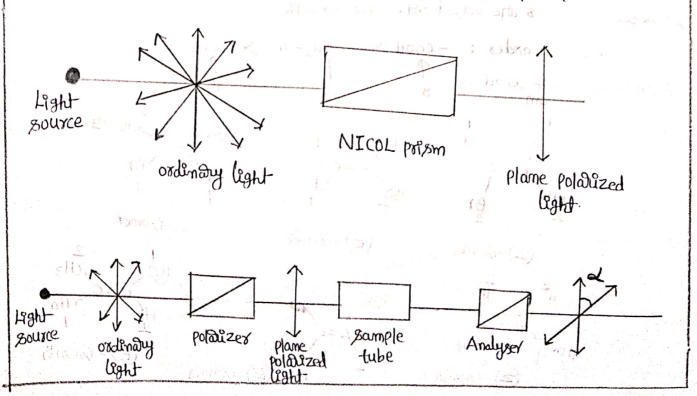
Geometrical Examplify in compounds with more than one double bond.

(a) when a compound contains I'm" number of dizzimilarly
substituted double bonds, then the number of geometrical isomers is 27.

$$\frac{1}{1}$$
  $\frac{1}{1}$   $\frac{1}$ 

## optical activity (optical geomorism):

plane polarized light: Ordinary light corplists of wars vibrating in all planes perpendicular to in the propagation, when such a light is allowed to pass through NICOL prism, its vibrations in all planes one cut of except one plane. This light having wave motion in one plane only, is called plane polarized light and the plane perpendicular to the plane of vibration, is called the plane of polarisation.



A mexture of these two varieties in erual proportions will be optically inactive and it is called "racemic form".

#### Polorimeter:

The Pristrument for Studying interaction between planepolatized light and chiral molecules is known as a polatineter. (Fig.)

Intially, a beam of unpolatized light is passed through a polatical sheet, called polatizer, then through a sample tute. By rotating the analyzer, another polatical sheet, minimal light transmission can be achieved. Next the sample tube is filled with a solution containing am optically active compound. As the plane-polatized light passes through the sample tube this time, its plane of polatization is rotated either to be right to the left by certain amount depending on whether the optical isomer is in the "d" (08) "L"-form. This rotation can be measured readily by turning the analyser in the appropriate in the appropriate direction, until minimal light transmission is again achieved. The angle of rotation in degrees, &, depends not only on the nature of the molecules, but also on the concentration of the solution and the length of sample tube.

Specific votation: optical activity is retailed in terms of specific votation; which is characteristics of a particular compound. It is defined as the rotation in degrees observed when a plane polarized is passes through I dm of solution having concentration of I ginfind at a specified temperature and cavelength. This is usually expressed as

where Dis the D-line of sodium having wavelength 589.3 tis the temperature (2)

I is the length of polarimeter cube,

C is the concentration of solution (9/ml).

The specific rotation of pure liquid is given by  $\mathbb{Z}_{p}^{t} = \frac{\mathbb{Z}_{p}^{t}}{14^{-3}}$ 

If the substance is in solution from then:  $[x]_0^t = \frac{1}{10}$ The extent of votation [x] of the plane of polarization depends on

is nature of substance

(li) 94s temperature

(III) length "1" through which (ight passes

(iv) density "d" of the substance and

(V) wavelength of the light employed.

Requirement of optical activity: (00) cause of optical activity:

Compound possessing identical molecular and structural formulae and identical physical and chemical proporties, but differing only optical activity (i.e; their action on plane polarized (19th) are called optical isomers (ox) emantioness and the phenomenon is called optical isomerism (ox) enanthomerism.

Example: Fluoro chiorobromo methame, CH Fd Br, contains an assymetric central contains atom; and 9t exists in two optical active forms, which the related to each other as mirror images and the the two the non-super pmpossible; ! H CHO : CHO

Pmpossible if F F By

Mins

CHO

H-C-OH

CH2OH

CH2OH

CH2OH

Glycoladdehydai Glgwraddehyda

(O)

mirris (O)

Course of optical autivity: The only factor responsible for the optical activity (or soldien of the plane of polarized light) is Mokewan Assymetry. A plane which biseds the object into super-impossible mirror image halves, is called the plane of symmetry. "An object which how no plane of symmetry is asymmetric (or) chiral". Thus the reflection of an asymmetric object in a mirror is, not super-impossible on the object. Asymmetric object and its

misson image one called "enautioners".

(Symmetric)

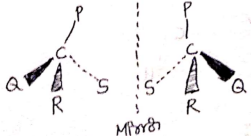
A A

(Asymmetric)

Symmetric and assymetric object

From the above discussion, it is clear that for an organic compound to show optical activity, it should be in have an assymmetric (or) chiral contamination and in not have a plane of symmetry.

Asymmetric (on chiral carbon atom: A carbon atom bonded to four different atoms (or) groups, is called "asymmetric carbon atom". Thus, if a molecule has am asymmetric carbon atom, it is no longer super improssible on its mirror image and will, therefore, exhibit optical activity.



Plane of symmetry: If a molecule cambe divided by an imaginary plane on which a way that points on one side of plane form a mirror

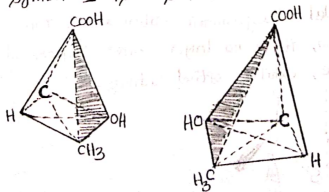
material parmets

Centre of Symmetry: This is an imaginary point in a molecule from which, when equal lengths are drawn on both sides, then they meet exactly at similar points in the molecule. If a compound has a centre of symmetry, it is "optically imartive".

optical isomerism of Lauric and: [CH3 CH OH. COOH]

Latte and contains an asymmetric control colon atom, and it exists in two optical active forms, which are related to each other as mirron images and the two one non-superimpossable.

A third form called racemic mixture is obtained, when equimolar quantities of d- and 1-enantioners are mixed. Such a mixture is optically inactive as a result of the cancellation of the equal and opposite rotations of two isomers. The symbol  $\pm$  is used to denote a racemic mixture.



optical promerism of Taxtaric and:

Tartanic and has two asymmetric coulon atoms, each linked to the four different groups, i.e., -OH, -H, -COOH and -CHOH) COOH.

The spatfal assungements of various groups in tartain and camba represented in four ways, correspondingly, tortails and can exists in four different forms given below.

B Dextro totalic and 9% on which rotates the plane polatization of light to the right. This votation due to upper half is strengthened by one due to lower half.

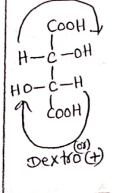
(i) Levotantaic and is a missor image of the above and rotates the plane of polarization of the light to the left.

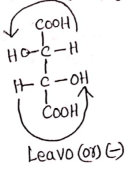
(ii) Racemic trottable and 1/2 a 50-50 mixture of the above two valieties and is hence, madive. It can be repolved into the adive constituents, Viz; dextro and levo-vouletie. It is therefore, inactive by external compensation.

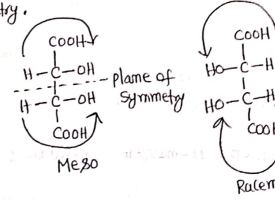
(1V) Meso-tartalic and 9s the grantive variety as the solution of upper-half as compensated by the rotation due to the lower-half (being in opposite direction). It cannot be resolved into active constituents.

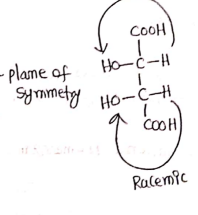
It 98, therefore, martine by internal compunsation, unlike dextro- and levo-

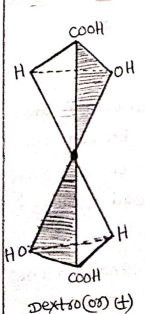
varieties, it has a plane of symmetry.

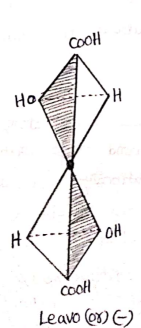


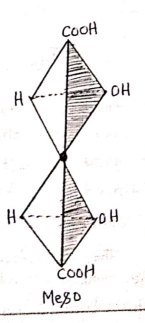


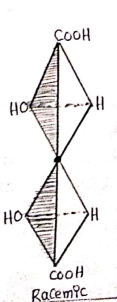












characteristics of enantionness: (1) They have identical physical proporties such as metting points boiling points, densities and refractive indices.

except their behaviour toward optically affive compounds.

(3) Earthorneys have different biological

proposites. (3)

(4) They differ in their action toward

plane polarized light. They rotate the plane polarized light to the same extent but in opposite disections.

(5) They cannot be separated by methods such as fractional constallization and fractional distillation, chromatography.

#### Conformational isomerism:

Conformational isomerism refers to any one of the infinite number of spatial abhangements of the atoms of a molecule that can be from rotation about a single bond. The term should not be confused with configuration isomerisms, which are spatial abhangements of the atoms of a molecule that can be changed only by breaking and making of bonds.

The steepersomers which are non-super impossible but abe easily inter convertible by rotation about the single bond are known as conformational isomerism.

## conformational isomerism of n-tidanc:

the sigma bond Johning the corbon atoms a and 3 of butane, is eylindsically symmetrical about the nuclear axis.

HS-C-13-CH3

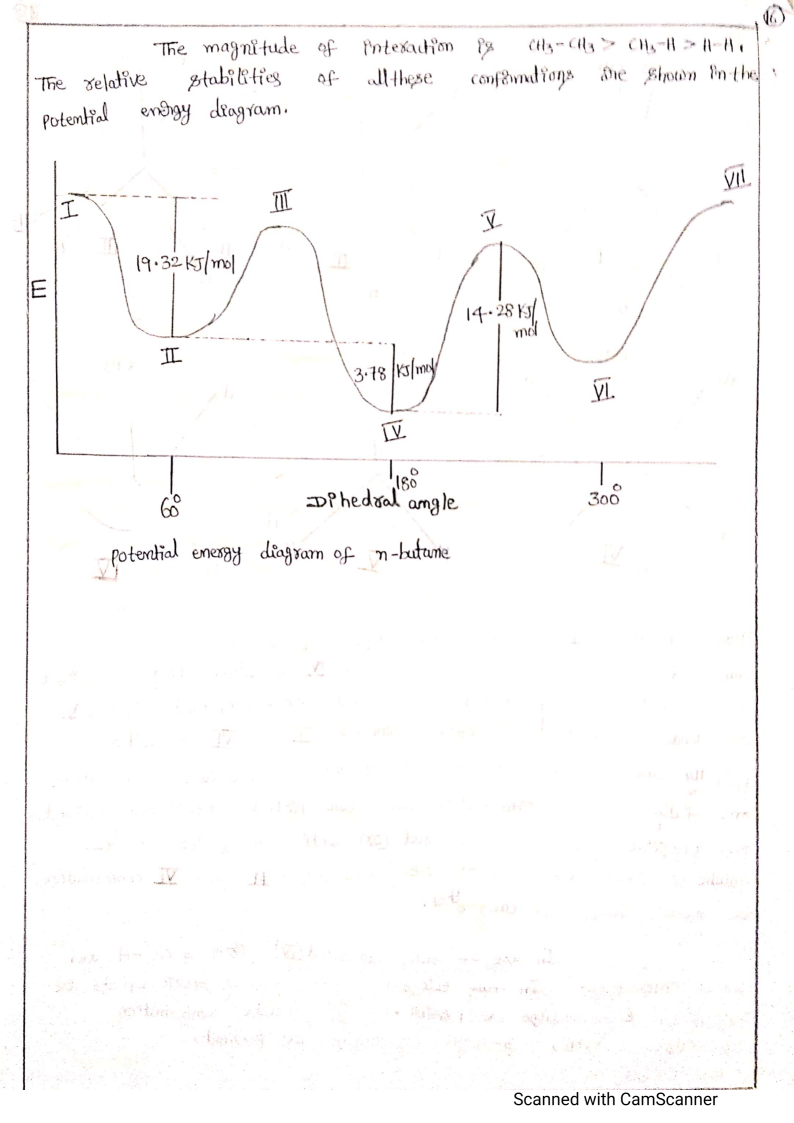
This in-turn permits the free rotation of coulon atoms number 2 and 3 with respect to each other along their bond axis without breaking the bond. If one CH3-CH2- group is kept stationary and other CH3-CH2- group is allowed to rotate through 368 in six steps (60 each time), then following bix conformations of n-batane one obtained.

that results from the free rotation about C2-C3 bond the called conformational isomers of n-butane and this phenomena

The conformation I in which two methyl groups are very much close go called fully eclipsed, and the conformation IV in which they are failtiest apart is called fully staggered (or) anti-firm. conformation III and II are called see known as partially eclipsed, where as II and II are called partially staggered (or) gauche (or) skew. In the case of n-butane one fully eclipsed conformation and two partially staggered conformations are observed. one fully staggered (or) anti conformation and two gauche (or) skew conformations are present. II and III conformations she minds in and III conformations.

In case of fully staggered (IV) from 4 cH3-H and 2H-H interactions. In fully eclipsed form greater steric effects are alting, i.e.; 6 interactions are possible. In southe conformation 1 cH3-CH3, 2-CH3-H, 3-H-H interactions are present.

CH3



(A) eHEMICAL SHIFT: The number of signals in an NMR spectrum tells the number of the sets of equivalent protons in a molecule. The position of the signals in the spectrum help ys to know the nature of protons.

If the resonance frequency of all protons in a molecule were the same, them the NMR spectrum would show only one peak for the compound. The induced field opposing the applied field. The field felt by the proton, is dismissed and the proton is said to be "shellded". The induced field reinforces the applied field, the proton feels a higher field strength and thus, such a proton is said to be "deshielded." "Shielding shifts" the absortion "upfield" and "deshielding shifts" the absortion "down field".

Such shifts in the positions of NMR absorbing which orise due to the shielding (or) destilling of protons by the electrons are called "chemical shifts". (6). For measuring chemical shifts of various protons in a molecule, the original for tetrametryl solane (Tms) is taken as a reference, due to city in the low electronegativity of solicon the Si-city.

(i) the low electronegativity of states the confounds.

protons with the same chemical shifts are called equivalent protons. Non-equivalent protons are have different chemical shifts. & (delta) (08)

(tau) scales one commonly used.

Shielded signals (upfield) Tims

deshielded signals (downfield)

10 9 8 7 6 5 4 3 2

7 1 1 1 1 1 1 1 Tau

The values of 8 fb a substance with respect to TMS can be obtained by measuring

where 
$$v_3 = \text{Rezonance frequency of the zample}$$

$$\delta = \frac{\Delta v}{\text{operating frequency in mega cycles}}$$

The value of 8 is expressed in parts permillion (ppm). most chemical shifts have 8 values between 0 and 10.

Type of protons	chemical shift in ppm
CH4	0.23
CH3-I	2.16
CH3- BY	2.68
CH3-CL	3.25 4.26
-4.5	

(3) Spin - Spin splitting (spin-spin coupling): "The splitting of signal lines in the spectrum into two (or) more components is called spin-spin splitting".

consider a molecule of CHz-CHz-Br (ethyl bromide).

This molecule has two kinds of protons in it and thus, two signals one excepted in its own NMR spectrum. For "a" kind of protons (CHz), a triple i.e.: a group of three Peaks is observed and a quotet is noticed for "b" kind of protons (CHz).

The spin of two protons (-CH2-) can couple with the adjacent methyl group (-CH3-) in three different ways relative to the external field. The three different ways of allignment one:

Thus, a triplet of peaks results with the intensity ratio of 1:2:1. Similarly, the spin of three protons (-cHz) can couple with the adjacent muthylene group (-CHz) in four different ways. Thus, quartet of peaks results with an intensity ratio of 1:3:3:1.

(\_AMA\_)

Ü 117

(i) TTT TTT 3

(ii) TTT TTT 3

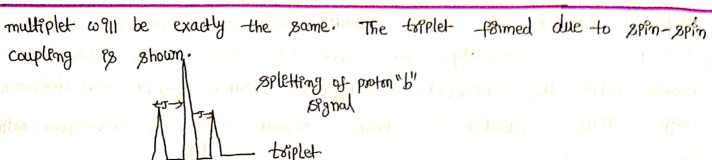
(v) 111

The multiplicity of the signal for any group of equivalent protogs is clearly related to the number of protogs of the adjacent atoms. As a semple rule the multiplicity of a given group is (n+1), where m is the number of protogs of the adjacent atoms. Accordingly m-propyl indide (CH3-CH2-CH2-I) has three different types of protogs. The NMR spectrum of 3H (triplet) (1:2:1), 2H (sextet) (1:5:10:10:5:1) and 2H (triplet) (1:2:1).

# (a) Coupling complant (J):

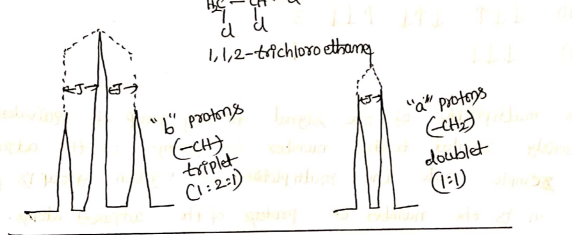
"The distance between the centires of the two advacent peaks in a multiplet is usually constant and is called the coupling constant". The value of the coupling constant 9s independent of the external field. It is measured in thest (Hz)(os) in cps (cycles per second). It is denoted by the letter J. The value of "J" generally lies between o and so Hz.

Now let us consider a compound >CH-CH2In this compound two signals one expected in the NMR spectrum. The
influence of two equivalent protons "a" the signal for proton "b" will
appear as a triplet. The distance between any two adjacent peaks in a



coupling constant in the multiplets are observed.

NMR spentrum of 1,1,2-trichloro ethane two



coupling constant in a-butene:

How the constant 
$$C = c$$
 the constant  $C = c$  the

percently has beducen a und at the consider a compound Sandy.

The talk percent as expended to the compound Sandy and the considered and the constants and the constants. The talk are constants and the constants and the constants and the constants. The

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bungan K at

# Module: V: Reaction Mechanism and Synthesis of drugs

# (B)-1. Explain homo & heterolytic cleavage with suitable example.

The breaking of a covalent bond between two atoms can take place mainly in two alternate ways, viz., homolytic and heterolytic fissions depending upon the relative electronegatives of the two central atoms.

Homolytic fission (or) homolytic bond fission, one electron of the bonding pair goes with each of the departing atom resulting en (6) two electrically neutral fragments (or) atoms known as "free radicals".

Homolytic fission takes place when the two atoms are usually of similar electronegativity. Free radical reactions are catalysed by light, peroxides (H202, benzoyl peroxide) and high termperature, these are said to be the free radical producing agents.

Heterolytic fixsion (n) he boolysis: In this type of fixsion the electron pair forming the covalent bond goes a single atom and thus electrically changed fragments (ions) are formed. The heterolytic fixsion of the covalent bond can occur in two ways. Heterolytic fixsion takes place when the two atoms are usually of different electronegativities.

such organic species has only six paired of electrons and a positive at Pts corbon centre is known as "corbonium for". corbonium fors one generally symbolized as R.

(i) when the electron pair between "c" and "x" the organic group and the substituent "x" is devoid of its bonding electron and thus gets positive while the organic group is negatively charged.

Such organic species which has eight paired electrons and a negative charge on one of its corbon center known as "carbanion". carbangons are generally symbolized as Romannia

### (3). a. what are reaction intermediates? Explain their formation and stability with examples.

Reaction intermediates one generally short-lived and highly reactive species and they react with reagent to form the products. Several intermediates during analysis of organic realions. They one (i) Free radicals

(ii) carbocations

(III) Carbani ons

(i) Free radicals: Free radicals may be defined as an odd electron neutral species a which the rosides on a corbon atom. These one formed by the homolysis of avalent bonds.

A free radical may have a sp2 hybridized carbon in which the odd electron remains in the unused, patital; the shape of this type of 128 ) Sp2 hybridized free vadical will be planar.

stability: Alkyl free radicals are classified into primary (1), secondary (2) and textiary (3) depending on nature of "c" atom that cornies the odd electron. The stability of corbon (alkyl) free radicals is not influenced by inductive effect because they have no charge. However, they are stabilised by hyperconjugation (non-bond resonance). This explains stability of alkyl free radials which

of the corbin wide is

follows the order 3>2>1> methyl. More the number of possible hyperconjugative structures more is delocalization of add electron and therefore more is the stability. In age of 3, nine hyperconjugative structures are possible, while in 2° and i, six and three structures are possible, respectively, as shown below. This explains the order of Stability.

H-C. > No hyperconjugative structure

Methyl free radical

How the Hold of ethyl radical i) i freezadial

H-C+13

2º free radical

# (1) corpo conflows (corporation long):

Carbo cation is positively changed species

containing a corbon atom having an incomplete octet, i.e., only sixelectrons In three covalent bonds. These one formed by hetereolysis of covalent bonds.

carbocations are classified as 1, 2 and 3 depending upon the nature of the iii) carbonions

Proproplastoction methylcoloation ethyl colocation

t-butyl corbocation

The assum atom in consocation is sith hybridized and the system is plant at the cationic "c" center and bond angle of 128.

Stability: The stability of carbocation cambe explained by either through resonance effect and Inductive effect (electron pushing effect). (a) Resonance effect: Resonance also helps in delocalizing the +ve charge and there by explains the stability of carbocations, e.g., in case of benzyl and allyl carbocations.

(b) Inductive effect: The alkyl group releases electrons to positive corbon and thus reduces its charge and electrons to positive corbon and thus reduces its charge and its persal of its turn itself becomes some what positive. This dispersal of the charge stabilizes the corbocation. Therefore, a tertiary corbocation charge stabilizes the corbocation. Therefore, a tertiary corbocation (with a alkyl (with 3 alkyl groups) is more stable than a secondary (with 1 alkyl groups) which in turn is more stable than a primary (with 1 alkyl group), muthyl codion (with no makeyl group) is the least stable.

CH3 
$$\rightarrow$$
 CH3  $\rightarrow$  CH3

(iii) carbanions: carbanions are negatively changed species containing a carbon atom with six electrons in three bonds and in addition an unshared pair of electrons.

H3C-CO

The corbon atom that coories the negative change is sp3 hybridizally corbanions are classified as i, of and 3° depending upon the nature of the corbon atom bearing the negative change.

Stability: The stability of carbanions mainly responsible for the dispersal of the charge are "Industive effect" and "resonance" The electron - releasing substituents tend to intensity the charge at the electron rich carbon (I) and thoreby making it less stable. Electron withdrawing groups tends to decrease the negative charge dampsity on the electron-orch carbon (II) and have stabilize it.

Inductive effect: I II

(c) The stability of alkyl combanions which follows the order.

(b) Resonance: Resonance also helps in decolalizing the -ve change and these by explains stability of carbanions.

Electrophiles: These are electron seeking (or loving species (a) positive electrophiles: Ht, Brt, Not

(b) Neutral electrophiles: BF3, Ald3, znd2, So3

Nucleophiles: These are electron sufficient species.

(a) Negative nucleophiles: X, oH, NH2, CN

(b) Neutral nucleophiles: H20, NH3, ROH, RNH2.

(Q) 3. Define Substitution readings and Explain the mechanismos 5N' and SN' reactions with suitable example.

Those substitution reactions which involve the attack of a nucleophile one known as nucleophilic substitution reactions. These are usually written as SN ("s" stands for substitution and "N' 48 Nucleophilic) and one more common in diphatic compounds Hydrolysis of alkyl halides by aqueous KOH 98 a common example of nucleophilic substitution.

a) Bimolecular mudeophilic substitution (sn) mechanism:

Let us consider the nudeophilic substitution

reaction of a methyl bromide in aqueous Nath solution.

CH3 BY + OH -> CH3 OH + BY
Substrate Nudeophile product Lea

Mechanism: This is a one-step mechanism and since this step Provolues two molecules, It is known as bimolecular. Kinetics data reveals that the rate depends on concentration of both substrate and nucleophile.

Rate of Eliser [OH]

In this attacking nucleophile OH, attacks at corbon in CH3BV, forming a high energy transition state in which: (a) formation of bond between OH-carbon (b) bond By-carbon 98 9m process breaking with simultaneous of the leaving group, Br:

In this reaction, the molar concentrations of substrate (CH3BY) and Nucleophile (off) one changed and hence, the reaction is of 2nd order (or bimolecular) called SN reaction. In SN reaction, the nucleophile attacks a c-atom from the side opposite to that of leaving group.

In transition state (TS) both nucleophile and leaving group (Br.) carry partial negative charges. "c" is spr hybridized in this state. energy required to break the "c-Br" bond is compensated by energy released during making of the "c-off" bond. the

(1) Unimpleated nucleophilic substitution (SN) mechanism:

Let us consider the nucleophilic substitution

reaction of a t-butyl bromide in aqueous NaoH solution.

This is a two-steps mechanism. (a) The first step is slow and it is rate determining step, old bond break and (b) The second step Ps the attack of a nudeophile (OH) on the carbonium for and it is a fast step, new bonds one formed.

testiony butyl

The rate-determing step is first step and this step does not involve the attacking nucleophile. The molen concentration of one readant ((CH3)3-C-Br) is only changed, so the overall reaction is of first order, because the time taken by the reaction depends mainly on step-I. such reactions one SN reaction (which means substitution, nucleophilic and unimolecular).

These reactions may proceed either by bimolecular (Ez) (or) by unimolecular (Ei) mechanism.

(1) Ezmechanism: Let us consider the elimination reaction of a primary alkyl halide, say ethyl bromide by a strong base sodium ethoxide.

CH3-CH2-BY + C2H5ONQ -> CH2=CH2 + C2H5OH + BS (base) Product

(substrate) The vale of an elimination reactions depend upon the concentration of the substrate and the base, the reaction is said to be of

concentration of the substrate and the ast, the seather is such that second order and is represented as E2 (bimolecular elimination). The common example of E2 elimination reaction is the dehydrohalogenation of privately halides to form alkeres. Here the abstraction of the proton (by alkali) from the B-carbon atom and the expulsion of the halide ion from the d-carbon occurs simultaneously.

on the concentration of the substrate, the readfon is said to be of first one and designated as EI (unimolecular elimination). EI readfons are also two step processes. The common example of EI readfon is dehydrohalogenation of t-belog halides. In the first step, under the influence of solvation, the nudeophile goes away along with the bording electrons to form a contraction. This is the rate determining step and slow. Subsequently, the contraction loss a proton to solvent (or) some other proton acceptor.

Orientation: The elimination reactions of unsymmetrical substrates usually yield a mixture of all possible products.

double bond in either side of the chain prefer to yield most substituted double bond in either side of the chain prefer to yield most substituted alkene, i.e., the loss of the β-hydrogen causs preferably from the substituted position.

CH3

CH3

H3C-C-C-CH2-CH3

H3C-C-C-CH2-CH3

H3C-C-C-CH2-CH3

CH3-CH-CH3 (mays)

CH3-CH-CH3 (mays)

CH3-CH-CH3 (mays)

(B) 5. What the addition readings? Explain the addition readings with examples and mechantsms.

Reactions in which atoms (00) groups of atoms added to a molecule one known as addition reactions. Addition reaction occurs in compounds containing double (or triple bonds.

Addition reactions may be instituted by electrophiles, nucleophiles (08) free radicule. Hence addition reactions may also be of three types depending upon the mechanism, namely (a) Electrophilic addition reactions (AdE)

(b) Nucleophilic

(C) Free radical addition reactions.

(AdE) with suitable example:

Addition reactions brought about by electrophiles are called electrophilic addition realthous. They are characteristics of alkenes altyries and occur by the formation of carbocation as intermediates.

Eg: consider the addition of hydrogen halldes (H-X) to alkemes: Inthis addition reaction, the addendum first associates it self to electronofth TT-center to form a TT-complex. The TT-complex subsequently gets transferred into the intermediate corboration. In the next step, the nucleophile, i.e., the halide for attaches It self to the intermediate to form the final addition product.

$$c=c(+++-x)$$
  $c=c(--c)$ 

Alkene (addendum)

 $c=c(--c)$ 
 $c=c(--c)$ 

Electrophilic addition to asymmetrical alkenes takes place in accordance to the "Markownikoff's" rule. According to this rule, during addition across unsymmetrical double bond, the negative part of the attacking reagent attaches it self to the carbon atom carrying lesser number of hydrogen atoms, while positive part goes to the carbon atom with more number of hydrogen atoms.

As the 2° carbocation is more stable, it is formed predominantly and the latter adds the negative ion to form the mass product.

# (b) Explain the Mechanism of Nucleophilic addition reactions with suitable example:

Addition reactions which one brought about by nucleophiles one called nucleophilic addition reactions. common examples of compounds undergoing nucleophilic addition reactions one additions, Ketones etc. In additional additions one pulled towards oxygen (as it is more electronegative than carbon) making the carbonyl carbon electron-deficient and the carbonyl oxygen electron-ofch. The reaction can be catalysed by an aid (Ox) by a base.

Eg: Mechanism can be explained by considering the addition of HCN to a Ketone to form cyanohydown.

# C) Explain the Free radial addition reaction with suitable example:

Addition reaction brought about by free radicals one called free radical addition reactions. The addition of HBY to alkene Por the presence of peroxide is an example of free radical addition reaution.

The free radical addition of HBY to an unsymmetrical alkeme in the presence controlly to the Markorounekofs's rule and is known of peroxpde occurs as the Anti-Markownikoff's addition (Or) peroxide effect. This phenomenon gs also known as "kharasch effeit". It states that im case of addition of HBY to alkeme Pm the prosonce of peroxide, the megative post of the attacking we reasent (Br) will be attack the carbon atom coorying longer number of hydrogen atoms.

a-promo

Mechanism: In the presence of peroxide, the addition of HBr to okefings

takes place by free radical mechanism.

3) In Hadron: R-0-0-R homolytic a RO. (R18 generally GH500) RO. +HBY ROH + BY.

(ii) propagation: Br. free radical them attacks the propene to form i and CH3-CH=CH2 + Br CH3-CH-CH2 (1 free radical)

CH3-CH-CH2 (1 free radical)

CH3-CH-CH2 BY

CH3-CH-CH2 BY 2° free radicals. (less stable)

(2º free radical)

Sonce a free radical is more stable than of free radical, hence the final product is obtained from the secondary free radical.

(HU) Termination:

## (a). 6. Explain the mechanism of Beckmann reasonangement suitable example.

It is the conversion of Keteximes Caliphatic, anomatic, diaryl, alicyclic, heterolyclic) to amides (or) N-substituted amides by intermolecular rearrangement with Lewis aids like H2504, pcl5, P205, Societe.

Mechanism: (i) conversion of -OH group to a better leaving group.

bonding electron pair from carbon to nitrogen.

CH3 
$$C=0$$
 + HJN-OH  $\xrightarrow{-H_2O}$  CH3  $C=N$ :

CH3 Hydroxyl acetorie amine

CH3  $C=0$  + HJN-OH  $\xrightarrow{-H_2O}$  CH3  $C=N$ :

 $CH_3$   $C=N$ 

HC = N = migration of alkyl group

The readrangement is highly stereospecific in which the group anti to off group.

The reaction ps utilized for sing expansion. An important example is the conversion of cyclohexamone exime into capsolactum, which has considerable synthetic importance in manufacture of person (a textile polymer).

(a) 8. Explain the mechanism of commizarys reaction with suitable example.

Aldehydes containing no a-hydrogen atom (es Hetto, (formaldehyde)), C6 H5CHO (benzaldehyde)) on treatment with conc. NaoH/ KOH undergo self oxidation and reduction to yield an alcohol and a salt of corbosylic aid. One molecule of an aldehyde is reduced to an alcohol and another molecule is oxidized to a corbosylic aid.

Me changem: Following three steps are involved:

Step-I: Attack of of Pons (of NaOH/ KOH) on the cobonyl combon atom

$$O \rightarrow C \rightarrow H \rightarrow O \rightarrow C \rightarrow H$$

Benzaldehyde

(A)

Step-III: Transfer of hydride for (OH) to another molecule of aldehyde.

Step-III: Acid-base reaction between (B) and (C)

(a) 9. Explain the Diel's-Alder mechanism with suitable example.

Diets-alder reaction consists in addition of a conjugated diene to an unsaturated molecule called a dienophile, resulting in the formation of a six membered ring. The example is the reaction between butadiene and ethylene.

Dies alder reaction belongs to special class of reactions called policyclic reactions, which one characterised by the making and breaking of two (or) more bonds in a single concerted step through cyclic transition states. These reactions are known as cycloaddition reactions. The above example involves addition between a 4TT electron system (diene) and att

7 (0)-CH20H

electron system (dienophik), 9t 9s known as [4+2] cycloadditton.

Mechanism: Both dieme and demophile underogo electrometric shifts simultaneously, thereby producing their respective excited state. Then, these excited states condense forming an adduct in 1,4-positions of the dieme.

The vate of the above readfon go promoted by the presence of:

- (i) Electron-donating group in the diene (like-cHz somp).
- (ii) Electron-withdrawing groups on the demophile (or) the alterne (like (40, (00), (00), (00)).

## (a). 10. Write the Synthesis and pharmaceutical applications of paracetamol.

paracetamol is also known as "acetaminophen" is a medicine and was discovered in 1877. The structure of paracetamol is given below.

It is prepared by the netration of phenol with sodium netrite, which gives two isomers of orthonitro phenol and para nitrophenol. The ortho isomer is removed by steam distillation and the para isomer is reduced to amine in the presence of sodium borohydrida (Na BH4) to give p-aminophenol.

Finally the armine is acetylated with acetic arrhydride to produce acetaminaphen widely known as paracetamol.

Applications: (i) paracetamol 98 used for reducing fever in people of all ages under the trade names Tylenol, paradol etc.

(ii) paracetamol is used for the relief of mild to moderate pain.

(iii) It has analgesic proporties. paracetamol can relieve orthritis
i.e., pain of the hip, hand (or) knee.

geveral first line therapies for treatment of tempion of migrame headache. (i) paracetamol is used for controlling dental pain dusing due to dental procedures.

(vi) Although it causes lesser gastrointentinal side effects as compared to aspirin, but overdose of paracetamol causes damage to lever.

## (3)-11. Write the Synthesis and pharma central applications of Ibuprafen.

Ibuprofem, 2-(4-isobutyl phend) propronic aid, cambe synthesized by various method. To the synthesize of ibuprofem consists of the chloromethylation of iso-butyl benzene, giving 4-iso-butyl benzylchioride. This product is reacted with sodium cyanide (Nacn), making 4-iso-butyl benzyl cyanide, which is alkylated in the presence of sodium avaide by methyl iodide into 2-(4-isobutyl benzyl) propronitiske.

Hydrolysis of the resulting product in the presence of a base produced thuprofem.

Applications: (i) Ibuprofen exhibits analgesic, fever reducing and antifenfammatory action. (ii) It is used in treating rheumatorid arthritis, in various forms of articular and non afficular diseases (iii) It is used to pain relieving from inflammatory posipheral nerve system (iv) It is used for head aches and toothaches.

monocipied to emiliarly a hostology beauty between the contract of the contrac

concepts and theory:

"Nuclear magnetic resonance, spectroscopy (NMR) is an absorption spectroscopy in which certain magnetic nuclei under the influence of a static external magnetic field one subjected to a second oscillating electromagnetic field in the form of radiofresumy radiation which induces the nucleus to reponate."

NMR spectroscopy is the study of NMR phenomenon to examine the physical and chemical proportion of a molecule.

Many stormic mucle" have mechanical spin and angular momentum which is described in terms of "nuclear spin quantum muber I". It has values of 0, 1/2, 1, 3/2 and so on (I=0 despotes mo spin). For the nucled to be magnetic, they should have alther odd number of protons (0x) add number of meutrons (0x) both only the nucled that have spin quantum number greater than zero can exhibit phenomenon (Eg: 14, 11B, 13c, 14N, 15N, 170, 19, 31, etc).

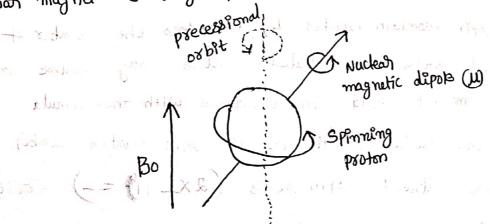
The spin quantum number I determines the number of quantum mechanical states an isolated nucleus may assume in external tuniform magnetic field, in accordance with the formula (2I+1). Hence the nucleus of I has the spin quantum number I = I/2 and has allowed spin states  $(2X_1^1+1)^2=2$  for its nucleus which are -I/2 and +I/2. In the absence of applied magnetic field, the embry of all the spin states of a nucleus one equivalent (degenerate).

Element H H E B B H 16 19 F Nuclear spin (I) 1/2 1 0 1/2 1 0 1/2 1 0 1/2 1 0 1/2 1 0 1/2 1 Number of spin 2 3 0 2 3 0 2

Because of the spin and change, a nucleus can behave like a magnet. NMR spectroscopy involves application of magnetic field to mudel and them measuring the amount of energy necessary to put vorious nuclei of the sample into resonance. Nuclei in different environments (shielded and unshielded) require different amount of energy to bring into resonance. Am NMR spectrum provides a signal (or) peak representing the energy necessary to bring each nucleus into resonance.

### Phenomenon of Nuclean Magnetic Resonance:

The magnetic nucles (H, 13cdc. under the influence of external magnetic field behave as a spirming both magnet as they possess both electric change and mechanical spin.

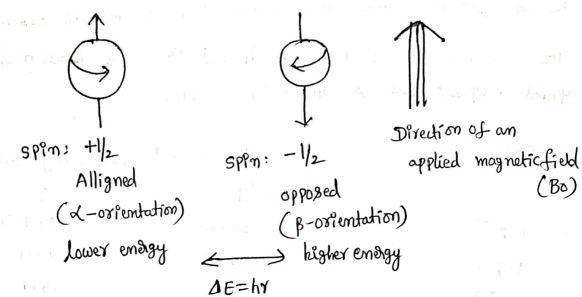


Spinneng proton en a magnetic field Bo

The proton like a magnet tends to align itself with the external magnet (parallel) with the magnetic field, called &- orienties (low energy state) or opposed (anti-parallel) to the field, called B-orientiation (higher energy state). In these & and B-states, the proton also

Concepts and theory:

spins around the axis of an applied magnetic field. This is called precessional motion.



pre cessional frequency: A transition from the lower emergy state to the higher emergy state can be brought about by applying radiation of exactly the required frequency (in the radiofrequency region) for a given stationary magnetic field of strength Bo. The fundamental NMR equation correlates to a particular value of the applied radiofrequency known as Lamor frequency v, which is is directly proportional to the magnetic field strength. Thus

V & Bo

V X Po

$$V = \left(\frac{3}{2\pi}\right) B_0$$

SPrice 
$$\Delta E = hV$$

DEX BO (: h, ) and IT

De constati

h is plank's constant

y is gyromagnetic ratio and

Bo P8 the strength of the magnetic field.

The radio frequency v is typically in the order of megahertz (mHz).

A frequency of 300 MHz is needed at a magnetic field strength

BO of 7.05 T (tesla) for proton. If an external field of strength

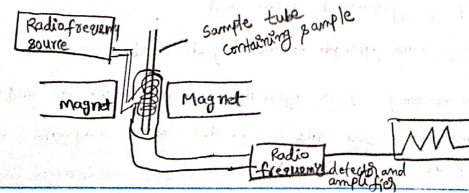
1.4 T will precess 60 MHz. At this ratio, the system is said to be in resonance, hence the name "nuclear magnetic reponance".

The frequences of protons in H and 13 as a function of field of applied magnetic field is shown in Table.

Во Стена)	Precessional	frequency (mHz)
1.4 T	60 MHz	15.1 MHz
1.9 T	15 M 120 1 1 20 80 - 11	ा । १५ १, १००।
2·3T	1100	25.1
•	200	50.3
4.77	200	75.5000000
7.1 T	300	125.7
11.7 T	milianshur an 500 prouve	1 HILLEY 151 1-1000
14.14	600	the specifical prophers

## Instrumentation of NMR spectroscopy:

The early NMR instruments for proton magnetic resonance were built using permanent magnets (85) electromagnets, generally with field strengths of 1.4, 1.87, 2.20.(67) 2.35 tests and corresponding frequencies of 60, 80, 90 (67) 100 MHz, respectively. The basic components, a detection system to measure the energy being alosabled by the nuclei from the radio frequency beam and a display monitor. These is shematically represented as



The sample under investigation is dissolved in deuterated solvent and the solution taken in a glass tube is placed in the probe with a spinner that sping the tube so that the non-homogenous components of the magnetic fields one averaged out. The transmitter and received coils also form part of the probe. The computer interface acquired the data and further processes to delive the NMR spectrum, which appears as a solies of peaks (corresponding to protong in different chemical environment) whose area are proportional to the number of protons they represent. The NMR spectrum obtained is useful for identification of compounds as well as quantitative analysis and purity of the sample.

CHEMICAL SHIFT: The number of signals in an NMR spectrum tells the number of the sets of equivalent protons in a molecule. The position of the signals in the spectrum help us to know the nature of posterns. Viz: aromatic, aliphatic, a cetylinic, vinylic etc. Each of these type of protons will have different electronic environments and thus, they absoluted applied field strengths.

If the resonance frequency of all protons in a molecule were the same, then NMR spectrum would show only one peak for the compound. The field strength experienced by the protons in the sample is not same as the strength of the applied field. "A magnetic field induces electron circulations in the change cloud in a plane perfendicular to the applied field, and in such a direction so as to produce a field opposing the applied field. The field felt by the proton, is dismissed and the proton is said to be "shielded".

The Produced field reinforces reinforces the applied field, the proton feels a higher field strength and thus, such a proton is said to be "deshielded?

the abantifun down field.



Applied field.

"Buch shifts in the positions of NMR absortions which dise due to the shielding (a) deshielding of protons by the electrons the called "chemical shifts"(6) 13 measuring chemical shifts of various protons in a molecule, the signal for tetrametryl silome (TMS) is taken as a reference, due to cus

i) the Low electromegativity of silicon. CH3-59-CH3

(ii) the shielding of equivalent protons on TMS
TMS

1/2 greater than most of the organic compounds.

protons with the same chemical shift are called equivalent protons. Non-equivalent protons are have different chemical shifts. 8 (delta) or

T(tau) & cales one commenty used.

Shielded signals (upfield) Tins

deshielded signals (downfield)

1 1 1 1 1 1 0 6



obtained by measuring Us - VTMS

operating frequency in megacycles

where  $\Delta v = P8$  the frequency 8 hift.

The value of 8 is expressed pn posts per million (ppm). Most chemical shifts have 8 values between 0 and 10. In the 7 scale, signal for the standard reference, This is taken as 10 ppm.

The value of 8 is expressed pn posts per million (ppm). Most chemical shifts have 8 values between 0 and 10. In the 7 scale, signal for the standard reference, This is taken as 10 ppm.

NMR signal ps usually plotted with magnetic field strength increasing to the right. Thus the signal for TMS (highly shielded) appears at the extreme right of spectrum with S=0 ppm. Greater the deshielding of protons, larger will be the value of S. The values of chemical shifts for protons in different environments one:

2.7*	Types of protons	Chemial shefts in ppm
£		8
1	R-CH2	0.9
- bt		1 12-3.5
3	c=c-H	6-9.0
3	Ar-H	
9		Jan 1 1 23
4	CH4	2.16
3	CH3-I	2.68
6	CH3-BY	3.25
<b>(P)</b>	cH3-cl	4.26
8	CH3-F	works to the standard be soon where soon

 $\oplus$ 

## (3) Spin - Spin spletting ( spin-spin coupling);

"The splitting of signal lines in the spectrum into two (as) more components is called spin-spin splitting".

consider a molecule of CH3-CH2-Br (ethylbromid).

This molecule has two kinds of protons in it and thus, two signals are excepted in its own NMR spectrum. For "a" kind of protons (CH3), a triple i.e. a group of three peaks is observed and q quartet is noticed for "b" kind of protons (CH2).

The spin of two protons (CHz) can couple with the adjacent methyl group (-CHz) in three different ways relative to the external field. The three different ways of allignment are:

ii) 17 (Remaind)

iii) 11 11 (Anterso) | External field triplet

Thus, a triplet of peaks results with the intensity ratio of 1:2:1 which corresponds to the distribution ratio of alignment.

Similarly, the spin of three protons (-cH3-) can couple with the adjacent methylene group (-cH2) in four different ways thus, quartet of peaks results with an intensity ratio of 1:3:3:1.

(i) 111 111 3 (ii) 111 111 3 (iii) 111 111 3

\* The multiplicity of the signal for any group of equivalent protons is clearly related to the number of protons of the advacent atoms. As a simple rule the multiplicity of a given group is (n+1), where

m is the number of protons of the adjacent atoms. Accordingly, n-propyl rodide (CH3-CH2-CH2-I) has three different types of protons. The NMR spectrum of 3H (triplet) (1:2:1), 2H (sextet) (1:5:10:10:5:1) and 2H (triplet) (1:2:1).

The peak Pontensities of the various multiplets can be described as

Peak area ratio

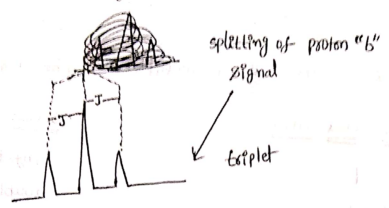
Singlet	doublet
1:1	Triplet
1:3:3:1	quartet
1:4:6:4:1	sextet

# (J):

"The distance between the centres of the two adjacent peaks in a multiplet is usually constant and is called the coupling constant". The value of the coupling constant is independent of the external field. It is measured in Hertz (Hz) con in cos (cycles per second). It is denoted by the letter I.

The spectrum of a particular compound at different radio-frequencies, the separation of signals due to different chemical shifts change but the separation of two adjacent peaks in a multiplet remains always constant. The value of "I" generally lies between 0 and 20 Hz.

Now let us consider a compound ) CH-CH2-In this compound two signals are expected in the NMR spectrum. The furthernce of two equivalent protons "a" the signal for proton "b" will appeals as a triplet. The distance between any two adjacent peaks in a multiplet will be exactly the same. The triplet formed due to spin-spin coupling is shown



sprinkaly In the NMR speatrum of 1,1,2 tolchioro ethane two multiples are observed.

Coupling constant in 2-buteme:

How the constant 
$$C = c$$
  $CH3$   $C = c$   $CH3$   $C = c$   $CH3$   $C = c$   $CH3$   $CH3$ 

# **Module-V**Fuels and Combustion

#### **Chapter 4: Fuels and Combustion**

**Fuel:** "A fuel can be defines as a combustible substance containing carbon and releasing a large amount of heat, that can be used for domestic and industrial needs".

"Combustion is the process of chemical reaction between fuel and oxygen. The combustion process is an exothermic chemical reaction".

$$Fuel + O_2 \xrightarrow{high \ temp} Combustion \ products + Heat$$

As most of the fuels contain carbon or carbon and hydrogen, the combustion involves the oxidation of carbon to carbon dioxide and hydrogen to water. Sulphur, if present, is oxidised to sulphur dioxide while the mineral matter forms the ash.

**Classification fuels**: Fuels can be classified into 2 types based on two factors

- 1. Their Occurrence
- 2. Physical state of aggregation.
- 1. Based on the occurrence, fuels can be further divided into 2 types (a) natural or primary fuels: Fuels which are found in nature are called natural (or) primary fuels. e.g., wood, peat, coal, petroleum, natural gas etc.
- (b) Artificial or secondary fuels: Fuels, which are prepared from the primary fuels. For example, charcoal, coke, kerosene, diesel, petrol, coal gas, oil gas, producer gas etc.
- 2. Based on the physical state of aggregation the fuels are divided in to solids, liquids and gases.

S.NO	Characteristic	Solid fuels	Liquid fuels	Gaseous fuels
	property of a fuel			
1	example	Coal	Crude oil	Coal gas
2	Cost	Cheap	Costlier than solid fuels	Costly
3	Storage	Easy to store	Closed containers should be used for storing	Storage space required is huge and should be leak proof.
4	Risk towards fire hazards	Less	More	Very high, since these fuels are highly inflammable
5	Combustion rate	It is a slow process	Fast process	Very rapid and efficient
6	Combustion control	Cannot be controlled	Cannot be controlled or stopped when necessary	Controlled by Regulating the supply of air
7	Handling cost	High since labour is required in their storage & transport.	be transported through pipes	fuels, these can be transported through pipes
8	Ash	Ash is produced and its disposal also possess problems	No problem of ash	No problem of ash
9	Smoke	Produce smoke invariably	Clean, but liquids associated with high carbon and aromatic fuels produce smoke	Smoke is not produced
10	Thermal efficiency	Least	High	Highest
11	Calorific value	Least	High	highest
12	Use in internal combustion engine	Cannot be used	Can be used	Can be used

#### Characteristics of a Good/ideal fuel:

While selecting an ideal fuel for domestic or industrial purpose we should keep in mind that the fuel selected must possess the following characteristic properties.

- 1. It should possess high calorific value. (Liberation of large amount of heat per unit mass/volume of the fuel)
- 2. It should have proper ignition temperature. The ignition temperature of the fuel should neither be too low nor too high.

- 3. It should not produce poisonous products during combustion. In other words, it should not cause pollution o combustion.
- 4. It should have moderate rate of combustion.
- 5. Combustion should be easily controllable i.e., combustion of fuel should be easy to start or stop as and when required.
- 6. It should not leave behind much ash on combustion.
- 7. It should be easily available in plenty.
- 8. It should have low moisture content.
- 9. It should be cheap.
- 10. It should be easy to handle and transport.

**Solid fuels:** The main solid fuel is wood. Wood contains higher percentage of carbon in the form of cellulose, lignocellulose. The process of conversion of wood into coal is called as **coalification**. The calorific value of wood is 4000-4500 Kcal/kg and thus is used as a domestic fuel. Wood can be converted into charcoal by a process is called **carbonization** (destructive distillation of wood). Charcoal is used as absorbent of gases and for decolourisation of sugars.

**Coal** is highly carbonaceous matter and is regarded as a fossil fuel produced from large accumulations of vegetable debris and alternation of vegetable matter like plants etc. under certain favourable conditions by the action of heat and pressure over a millions of years. Coal is mainly composed of carbon, oxygen, hydrogen and nitrogen.

Its formation can be explained by the following theories.

- i) In-situ theory: this theory states that the coal formation took place at the same area where vegetation grew and accumulated originally. The great purity of coal appear more reasonable on the basis of this theory.
- ii) Drift theory: according to this theory, trees when uprooted due climating conditions, in the deeper parts of the soil under transformation to coal due to high temperature, pressure absence of oxygen and presence of bacteria.

**Coal** has been classified in several ways. The most common method of classification is on the basis of rank. From the origin of coal it is clear the wood, after a long interval of time and under certain conditions was converted into coal. The successive stages in the transformation of vegetable matter into coal are wood, peat, lignite, bituminous coal and anthracite coal.

Coal is classified based on the carbon content. The following is sequence of conversion.

Peat → Lignite → Bituminous → Anthracite −

← Moisture content, H, O, S, N, Volatile matter

Carbon content, calorific value, hardness → −

**Peat**: peat is regarded as the first stage in the transformation of wood into coal. Brown, fibrous, jelly like mass. Un-economical fuel and contains 80-90% of  $H_2O$ . Composition C = 57%, H = 6%, O = 35%, ash 2.5 to 6%. Calorific value = 5400 kcal/kg.

**Lignite:** (Brown coal) soft, brown, coloured lowest rank coal moisture content is 20 to 60%. Composition: C = 60%, O = 20%, Calorific value = 6,500 to 7,100 k.cal/kg

**Bituminous coal:** Bituminous coal (common coal) Black to dark colored. This coal is largely used in industries for making metallurgical coke, coal gas and for domestic heating. It has laminated structure it is sub classified based on carbon content. Composition is % of C = 78 to 90%, VM = 20 to 45%, CV = 8000 to 8500 kcal/kg.

**Anthracite:** Highest rank of coal. These coals have very low volatile matter, ash & moisture. This coal is very hard, dense and lustrous in appearance. % of C = 98 % has lowest volatile matter hardest, dense, lustrous. CV = 8650 to 8700 k.cal/kg.

**Grading:** Coal is graded as caking coal and coking coal.

The coal which on heating becomes soft, plastic and fuse together are known as caking coal.

The coal which on heating gives porous, hard and strong residues are called coking coals.

#### Analysis of coal

The composition of coal varies widely and hence it is necessary to analyse the coal samples so that types of coal can be selected for a particular industrial use. The following methods of analysis can be utilized for the selection of coal.

- 1. Proximate analysis: This analysis records moisture, volatile matter, ash and fixed carbon as percentages of the original weight of the coal sample. Proximate analysis is of significance in commercial classification and industrial utilization of coal.
- 2. Ultimate analysis: This consists of determination of C, H, S, N and O. The ultimate analysis is essential.

#### **Proximate Analysis of Coal**

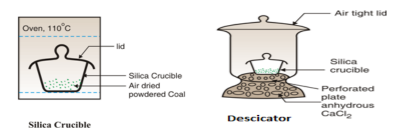
Proximate analysis is the study or analysis of coal sample in which

- (i) moisture %
- (ii) volatile matter %
- (iii) ash %
- (iv) fixed carbon %, are found out.

#### (i) Moisture %

- (a) Principle: All moisture in coal escapes on heating coal at 110°C for 1 hour.
- (b) Method: A known weight of powdered and air dried coal sample is taken in a crucible and it is placed in an oven for 1 hour at  $110^{\circ}$ C. Then the coal is cooled in a desiccator and weighed out. If the initial weight of the coal is **m** gms and final weight is **m**<sub>1</sub> gms. The loss in weight (m m<sub>1</sub>) corresponds to moisture in coal.

(c) Formula: Moisture % = 
$$\frac{\text{Loss in weight}}{\text{Weight of coal sample}} \times 100 = \frac{\text{m} - \text{m}_1}{\text{m}} \times 100$$



- (ii) Volatile Matter (V.M.) %
- (a)Principle: At 925°C, coal molecules undergo thermal degradation to produce volatile matter.
- (b)Method: Moisture free coal left in the crucible in first experiment  $(m_1)$  is covered with a lid loosely. Then it is heated at 925°C in a muffle furnace for 7 minutes. The crucible is taken out and cooled in a descicator. Then it is weighed  $(m_2 \text{ gms})$ . The loss in weight  $(m_1 m_2)$  is due to loss of volatile matter in the m gms of the coal sample. (Volatile matter is the thermally decomposed coal during burning of coal, that escapes without combustion, in the form of smoke).

(c) Formula: Volatile matter 
$$\% = \frac{\text{Weight of volatile matter}}{\text{Weight of air dried coal}} \times 100 = \frac{m_1 - m_2}{m} \times 100$$

The volatile matter % can also be determined by taking the fresh weight of the air dried coal but the loss in weight at 925°C, will be due to loss of moisture and volatile matter. If w is the weight of air dried coal and  $w_1$  is the mass of coal left at 925°C heating, then

Volatile matter % = 
$$\frac{\text{Loss in weight due to moisture and V.M.} \times 100}{\text{Weight of coal sample}} - \text{moisture \%}$$

$$= \frac{(w - w_1) \times 100}{w} - \text{moisture \%}$$

#### (iii) Ash %

(a)Principle: Inorganic matter in the coal gets oxidised to form metal oxides and silica, which is non-combustible and left as ash.

(b)Method: The residual coal in the above experiments is heated and burnt in a open crucible at above 750°C for half hour. The coal gets burnt. The ash left in crucible is cooled in a descicator and weighed (m<sub>3</sub> gm).

(c) Formula: Ash % = 
$$\frac{\text{weight of ash}}{\text{weight of coal}} \times 100 = \frac{m_3}{m} \times 100$$

#### (iv) Fixed carbon %

Fixed carbon % = 100 - (% Moisture + % V. M. + % ash)

#### **Significance (Importance of Proximate Analysis)**

- (i) High % of moisture is undesirable because it reduces the calorific value, increase the cost of transportation and causes wastage of heat.
- (ii) The volatile matter present in coal may be due to coumbustible gases (H<sub>2</sub>, CO, CH<sub>4</sub> etc) and non-coumbustible gases (CO<sub>2</sub>, N<sub>2</sub>). A high volatile matter containing coals give long flames, high smoke and low calorific value.
- (iii) Ash is formed in the presence of non-coubustible elements like Fe, Mn, and Co. Many Indian coals have high ash content. High ash content in coal normally leads to (a) reduce the calorific value (b) ash if present as clinkers
- (iv) The higher % of fixed carbon in a coal, greater is its calorific value and better the quality of coal.

#### **Ultimate Analysis of Coal**

Definition: The analysis of coal in which percentages of C, H, N, S and O elements are found out, is known as ultimate analysis.

#### Carbon and Hydrogen.

Method for determination:

- A known weight of powdered and air dried coal sample is burnt in the presence of pure oxygen, in a combustion apparatus. C and H are converted to CO<sub>2</sub> and H<sub>2</sub>O vapours respectively.
- The gaseous products are allowed to pass through first the preweighed U-tube containing anhydrous  $CaCl_2$  or magnesium per chlorate (absorbing  $H_2O$  vapours) and then through the KOH solution in a preweighed U-tube (absorption of  $CO_2$ ).
- The increase in weight of U-tube containing anhydrous CaCl<sub>2</sub> corresponds to weight of water formed and increase in weight of U-tube containing KOH solution corresponds to CO<sub>2</sub> formed, by combusting the coal sample.

#### Reactions:

$$C + O_2 \longrightarrow CO_2$$
;  $2 \text{ KOH} + CO_2 \longrightarrow K_2CO_2 + H_2O$   
 $12 \qquad 44$   
 $2 \text{ H} + \frac{1}{2}O_2 \longrightarrow H_2O$ ;  $CaCl_2 + 7 \text{ H}_2O \longrightarrow CaCl_2 \cdot 7 \text{ H}_2O$   
 $2 \qquad 18$ 

Formulae for calculation:

$$C \% = \frac{\text{Weight of CO}_2 \text{ formed } \times 12 \times 100}{\text{Weight of coal sample } \times 44}$$

$$H \% = \frac{\text{Weight of H}_2 \text{ O formed } \times 2 \times 100}{\text{Weight of coal sample } \times 18}$$

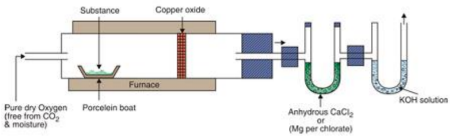


Fig. 4: Combustion apparatus

#### Sulphur:

Principle: Sulphur present in coal converts to first SO<sub>3</sub> which is soluble in water forming H<sub>2</sub>SO<sub>4</sub>. H<sub>2</sub>SO<sub>4</sub> is then converted to BaSO<sub>4</sub> precipitate when treated with BaCl<sub>2</sub>.

Sulphur is determined from the washings obtained from the known mass of coal, used in a bomb calorimeter for determination of calorific value. The washings are treated with barium chloride solution, barium sulphate is precipitated. The precipitate is filtered, washed and dried.

% 
$$S = \frac{wt. of\ BaSO4\ formed\ X\ 32\ X\ 100}{wt. of\ coal\ sample\ taken\ X\ 233}$$

#### Nitrogen:

#### Principle:

N in coal gets converted to ammonium sulphate, by action of hot concentrated H<sub>2</sub>SO<sub>4</sub> and then on treatment with alkali solution, equivalent amount of NH<sub>3</sub> is liberated.

#### Method:

A known weight of powdered and air dried coal is heated with concentrated H<sub>2</sub>SO<sub>4</sub> alongwith K<sub>2</sub>SO<sub>4</sub> catalyst in a long necked Kjeldahl flask.

After the contents become clear, it is treated with alkali solution in a round bottom flask. The ammonia (basic gas) liberated is passed in known volume of standard acid solution.

The unused acid is determined by back titration with NaOH solution.

The unused acid is determined by back titration with NaOH solution.

$$\begin{array}{c}
N \\
\text{(in coal)}
\end{array}$$
 $\begin{array}{c}
H_2SO_4, \text{ heat} \\
K_2SO_4
\end{array}$ 
 $\begin{array}{c}
NH4)_2SO_4
\end{array}$ 
 $\begin{array}{c}
\text{alkali, heat} \\
\text{NH}_3
\end{array}$ 
 $\begin{array}{c}
\text{Basic} \\
\text{acid solution}
\end{array}$ 

- i) Mass of coal = m gm.
- ii) $V_2$  ml = (Blank titration reading).
- iii) $V_1$  ml = (Back titration reading, after passing NH<sub>3</sub>)
- iv) Volume of the acid consumed by  $NH_3 = (V_2 V_1)$  ml.

#### Formula:

$$N \% = \frac{\text{Volume of acid consumed} \times \text{normality of NaOH} \times 1.4}{\text{Weigth of coal sample}}$$

OR

N % = 
$$\frac{\text{Volume of acid} \times \text{change in normality of acid} \times 1.4}{\text{Weight of coal sample}}$$

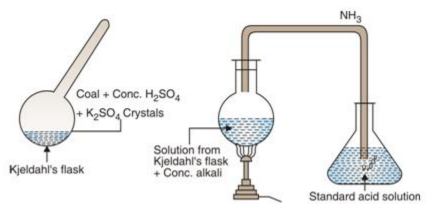


Fig. 5: Estimation of nitrogen in coal

Significances of Ultimate analysis: (1) Higher the % of carbon and hydrogen, the better is the quality of coal and its calorific value. (2) Nitrogen does not have any calorific value. It has no significance. (3) Sulphur increases calorific value. But the products of SO<sub>2</sub> and SO<sub>3</sub> are forms H2SO4 leading to corrosive effect on equipment. (4) Oxygen present in coal, it combined with moisture which causes low calorific value. An increases 1% of oxygen calorific value decreases by 1.7%. Hence a good quality of coal should be low % of oxygen.

**Ex. 1**) 0.25 gm of a coal sample on burning in a combustion chamber in the current of pure oxygen was found to increase weight of U-tube with anhydrous CaCl2 by 0.075 gm and of KOH U-tube by 0.52gm. Find C and H percentages in coal.

Given: Weight of coal = w = 0.25 gm

Increase in weight of U - tube containing  $CaCl_2$  = Weight of moisture formed = 0.075 gm Increase in weight of KOH U-tube = Weight of  $CO_2$  formed = 0.52 gm

$$2H + \frac{1}{2}O_2 \rightarrow H_2O$$
  
 $(2 \times 1)$  (18)  
 $C + O_2 \rightarrow CO_2$   
(12) (44)  
% hydrogen =  $\frac{\text{Weight of moisture formed}}{\text{Weight of coat}} \times \frac{2}{18} \times 100$   
=  $\frac{2}{18} \times \frac{0.075}{0.25} \times 100 = 0.33$  %  
% carbon =  $\frac{12}{44} \times \frac{\text{Weight of CO}_2}{\text{Weight of coal}} \times 100$   
=  $\frac{12}{44} \times \frac{0.52}{0.25} \times 100 = 56.73$  % ...Ans.

**Ex. 2)** One gram of coal sample was burnt in oxygen. Carbon Dioxide was absorbed in KOH and water vapour in CaCl2. The increase in weight of KOH and CaCl2 was 3.157 and 0.504 gm respectively. Determine % C and % H in the sample

Given: Weight of  $CO_2$  absorbed in KOH = 3.157 gm

Weight of H<sub>2</sub>O absorbed in CaCl<sub>2</sub> = 0.504 gm

Weight of coal burnt = 1.0 gm

$$C\% = \frac{12}{44} \times \frac{\text{Weight of CO}_2}{\text{Weight of coal}} \times 100 = \frac{12}{44} \times \frac{3.157}{1} \times 100 = 86.1\%$$
 ...Ans.

$$\label{eq:H_M_sight} H \,\% \ = \ \frac{2}{18} \ \times \frac{\mbox{Weight of } H_2\mbox{O}}{\mbox{Weight of coal}} \ \times \ 100 = \frac{2}{18} \ \times \frac{0.504}{1} \ \times \ 100 = 5.6 \,\% \qquad \mbox{...Ans.}$$

**Ex. 3)** Find the % of C and H in coal sample from the following data- 0.20 gm of coal on burning in a combustion tube in presence of pure oxygen was found to increase in the weight of CaCl2 tube by 0.08 gm and KOH tube by 0.12 gm.

Given: W = Weight of coal = 0.2 gm

Weight of CO<sub>2</sub> = increase in weight of KOH = 0.12 gm

Weight of H<sub>2</sub>O formed = increase in weight of CaCl<sub>2</sub> = 0.08 gm

**Ex. 4)** 2.4 gm of coal sample was weighed in silica crucible. After heating for one hour at 110°C, the residue weighed as 2.25 gm. The crucible was then covered with a vented lid and strongly heated for exactly 7 minutes

at 950°C. The residue weighed as 1.42 gm. The crucible was further heated without lid until a constant weight was obtained. The last residue was found to be 0.22 gm. Calculate the % results of the above analysis.

Given: W = weight of coal = 2.4 gm

Moisture % = 
$$\frac{\text{Weight of moisture}}{\text{Weight of coal}} \times 100 = \frac{0.15}{2.4} \times 100 = 6.25 \%$$

Weight Volatile matter = Weight of coal after 110°C - Weight of residue at 950°C

$$= 2.25 - 1.42 = 0.83 \text{ gm}$$

VM % = 
$$\frac{\text{Weight of VM}}{\text{Weight of coal}} \times 100 = \frac{0.83}{2.4} \times 100 = 34.58 \%$$

Weight of ash residue = 0.22 gm

Ash % = 
$$\frac{\text{Weight of ash}}{\text{Weight of coal}} \times 100 = \frac{0.22}{2.4} \times 100 = 9.17\%$$

Fixed carbon % = 
$$100 - (Moisture \% + VM \% + ash \%)$$
  
=  $100 - (6.25 + 34.58 + 9.17) = 50 \%$ 

The given coal contains,

**Ex. 5)** 0.5 gm of a coal sample on burning in a combustion chamber in the current of pure oxygen was found to increase weight of U tube with anhydrous CaCl2 by 0.145 gm and of KOH U tube by 0.90 gm. Find 'C' and 'H' percentage in coal.

Soln.:

Weight of coal burnt = 0.5 gm

Weight of CO<sub>2</sub> formed = Increase in weight of U-tube containing KOH = 0.9 gm

Weight of H<sub>2</sub>O formed = Increase in weight of U-tube containing anhydrous CaCl<sub>2</sub> = 0.145 gm

$$C\% \text{ in coal } = \frac{12}{44} \times \frac{\text{Weight of CO}_2}{\text{Weight of coal}} \times 100 = \frac{12}{44} \times \frac{0.9}{0.5} \times 100 = 49.99 \%$$

$$\text{H \% in coal } = \frac{2}{18} \times \frac{\text{Weight of H}_2\text{O}}{\text{Weight of coal}} \times 100 = \frac{2}{18} \times \frac{0.145}{0.5} \times 100 = 3.22 \ \%$$

#### LIQUID FUELS

Petroleum is one of the best primary liquid fuels. It is also known as crude oil. Petrol, diesel, kerosene are secondary liquid fuels and derived from petroleum. (i) The thermal efficiency of liquid fuels is higher than solid fuels. Low and high boiling fractions of petroleum are used in internal combustion of petrol and diesel engines, respectively. (ii) Liquid fuels possess higher calorific value per unit than solid fuels. (iii) Liquid fuels does not produce ash, dust, clinkers etc. during combustion.

**PETROLEUM (OR) CRUDE OIL:** The crude oil (or) petroleum also known as rock oil (or) mineral oil. The unpleasant odour of petroleum is due to the presence of some foul smelling sulphur compounds. Petroleum does not have definite composition. It is a complex mixture of various hydrocarbons and small quantity optically active compounds of S, N, O and traces of Fe, Cu, V, etc.

#### Composition of crude oil

Elemental composition

Liemental composition	
Element	Percent range
Carbon	80 to 87%
Hydrogen	11 to 15%
Nitrogen	0.4 to 0.9%
Oxygen	0.1 to 0.9%
Sulfur	0.1 to 3.0%
Metals	< 0.1%

#### Molecular composition

Open chain Alkanes
Cycloalkanes
Aromatics
Asphaltenes
Resins

Origin petroleum: There are 2 theories to explain the origin of petroleum.

(a) Carbide theory (or) Mendeleev's theory: This theory is also called inorganic theory. Metals inside the earth react with carbon to form metal carbides. These carbides are converted into hydrocarbons in the presence of moisture (or) steam, which on further hydrogenation, polymerise to give a complex mixture of paraffin's, olefins and aromatic hydrocarbons.

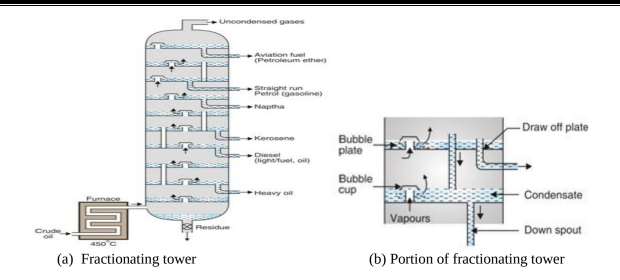
Drawbacks: This theory was unable to explain the presence of N, S and optically active compounds in petroleum.

**(b) Engler's theory:** According to this theory, organic matter, animals, vegetation and marine accumulated in sea. They were decomposed under high temperature and pressure by anaerobic bacteria to a dark viscous liquid called petroleum. This theory is better accepted and presence of optically active compounds in petroleum favours Engler's theory.

#### REFINING OF PETREOLUEM

The crude oil is a mixture of several hydrocarbons and is not fit to be marketed. "The process of separating various fractions of crude oil is called **refining** and hence where the industry refining of crude oil is takes place is called **oil refinery**. Refining of petroleum involves the following 3 steps.

- (a) Separation of water (Cottrelle's process): The crude oil is an emulsion of oil and water. This mixture was passed between two highly charged electrodes the emulsion films will destroy and the colloidal water droplets separated into bigger drops. They can be separated out from the oil.
- **(b) Removal of sulphur compounds:** In order to remove sulphur compounds from crude oil, it is treated with copper oxide. The sulphur compounds converted to insoluble copper sulphide, which can be removed filtration.
- (c) Fractional distillation: Crude oil obtained after (a) and (b) steps is then heated at about 400 °C in an iron retort. All volatile constituents are evaporated leaving behind some residue. The hot vapours are passed through different fractionating columns. The apparatus is a long cylindrical vessel with several trays, each having chimney with a loose cap.



The vapours go upward, they gradually become cool and their fractional condensation takes place at different levels in the column. The uncondensed gases escape from top of the fractionating column. These gases are liquefied and used as **LPG**. Different boiling fractions are condensed at different trays. Higher boiling fraction is condensed first and then gradually the lower ones.

Sr. No	Name of fraction	Boiling range	Composition of hydrocarbon	Uses
1.	Uncondensed gases	Below 40°C	C <sub>1</sub> to C <sub>4</sub>	Domestic and industrial fuel under 'LPG' name.
2.	Aviation fuel or petroleum ether	40° – 70°C	C <sub>5</sub> to C <sub>7</sub>	Fuel for aeroplane, Helicopters, as solvent
3.	Petrol or gasoline	60° C – 120°C	C <sub>5</sub> to C <sub>8</sub>	Fuel for petrol engines, dry cleaning, as solvent.
4.	Naphtha or solvent spirit	120° C – 180°C	C <sub>7</sub> to C <sub>10</sub>	As solvent and for dry cleaning, for chemicals.
5.	Kerocene	180° C – 250°C	C <sub>10</sub> to C <sub>16</sub>	For illumination, domestic fuel, for oil gas and fuel of jet engines.
6.	Diesel	250° C – 320° C	C <sub>15</sub> to C <sub>18</sub>	Diesel engine fuel.
7.	Heavy oil	320° C – 400° C	C <sub>17</sub> to C <sub>30</sub>	For making petrol by cracking.

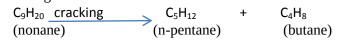
Heavy oil on re-fractionation produces

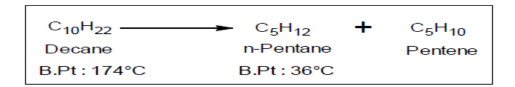
- (a) Lubricating oil (as a lubricant)
- (b) Petroleum jelly (in cosmetics, medicines and as a lubricant)
- (c) Grease (as a lubricant)
- (d) Paraffin wax (preparing wax papers, shoe polishes and candles)

Residue: Above 400 <sup>o</sup>C residue the following products are obtained (>30 carbons) (a) Asphalt (or) tar (for laying roads and water proofing houses of the roofs (b0 Petroleum coke (for moulding the rods of arc lights and as a fuel)

**PETROL (OR) GASOLINE:** Petrol is highly volatile, inflammable secondary liquid fuel and used in the internal combustion engines of automobiles. Only 20% of the petrol is coming from petroleum fractionating column and it is called **straight run petrol**. 50% of the petrol is coming by process called **cracking** and remaining 30% of the petrol is produced by synthesis from coal and water, which is called **synthetic petrol**.

CRACKING: Cracking is familiar process of breaking of bigger hydrocarbons of high molecular weight, high boiling to simple, low boiling, low molecular weight hydrocarbons. For eg: 1

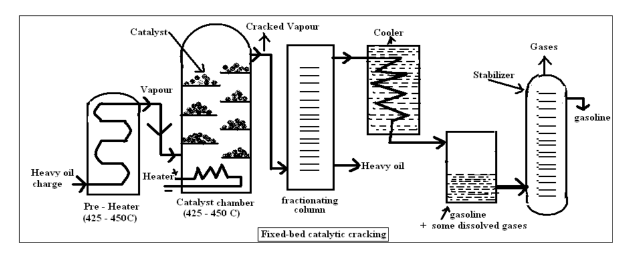




The process of cracking is mainly of two types. (1) Thermal cracking (2) Catalytic cracking

- (1) **Thermal cracking:** Breaking heavier hydrocarbon molecules into simpler, low boiling, lower molecular weight hydrocarbons by use of temperature and pressure is called "thermal cracking". Thermal cracking is carried out by two ways. (a) Liquid phase thermal cracking (b) Vapour phase thermal cracking. The liquid phase cracking takes place at 475°C to 530°C at a pressure 100kg/cm². While the vapour phase cracking occurs at 600 to 650°C at a low pressure of 10 to 20 kg/cm².
- (2) Catalytic cracking: In this type of cracking catalysts are used. The best one is aluminosilicate with some metal oxides (oxides of Ca, Fe, Mg, Cr, Na). This process completes at lower temperatures compared to the thermal cracking (300-400 °C., 1-5 Kg/cm² pressure). Catalytic cracking is also of two types.
  - (a) Fixed-bed catalytic cracking
  - **(b)** Moving-bed catalytic cracking

#### (a) Fixed-bed catalytic cracking:



(i) In this method, vapours are heated in a pre-heater to cracking temperature 425-450 °C (ii) The hot vapours are forced to catalytic chamber (containing artificial clay + ZrO<sub>2</sub>) maintained at 425-450 °C and 1.5 kg/cm<sup>2</sup> pressure. About 40% carbon is converted into gasoline and 2-4% carbon is deposited

on catalytic bed. (iii) The vapours produced are then passed through a fractionating column, where heavy oil fractions are condensed. (iv) The vapours are then led through a cooler, where some of the gases are condensed along with gasoline. The uncondensed gases move on to stabilizer, where the dissolved gases are removed and pure gasoline is obtained. (v) The catalyst after 8-10 hr stop functioning due to deposition of carbon. This is reactivated by burning off the deposited carbon. During reactivation, the oil vapours are diverted through another catalytic chamber.

## Q. a) What is knocking? b) What is octane number & cetane number c) What is leaded petrol? Discuss its advantages and disadvantages.

In internal combustion engines, diesel or petrol mixed with air is used as fuel and ignited in the cylinder. In **petrol engines** the ignition brought by an electric spark (spark engines) and compressing air (compression engines) in **diesel engines**.

Premature and instantaneous ignition of petrol – air (fuel-air) mixture in a petrol engine, leading to production of an explosive violence is known as knocking. The resistance offered by gasoline to knocking cannot be defined in absolute terms. It is generally expressed on an arbitrary scale known as Octane rating.

#### Knocking causes loss of efficiency of the engine, increase fuel consumption and damage to spark plug.

The tendency of fuel constituents to knock in the following order. Straight – chain paraffins > Branched-chain paraffins (i.e.,iso paraffins)> Olefines> Cycloparaffins (i.e.,naphthalenes)> aromatics. Thus, olefins of the same carbon chain length possess better anti knock properties than the corresponding paraffins and so on.

The knocking of petrol can be measured by a test is called **octane rating** (or) **octane number**. The **n-heptane** knocks very badly and hence, anti-knocking value has been arbitrarily fixed as zero and **isooctane** (2,2,4-trimethy pentane) gives very little knocking, so its anti-knock value fixed as 100.

$$\begin{array}{c} CH_3 \\ CH_3 - C - CH_2 - CH_2 - CH_3 \\ CH_3 & CH_3 \\ 2,2,4 \text{- trimethyl pentane} \\ \text{(isooctane) octane number 100 (good fuel)} \end{array} \qquad \begin{array}{c} CH_3 - (CH_2)_5 \text{- } CH_3 \\ \text{n-heptane} \\ \text{octane number zero (bad fuel)} \end{array}$$

Several mixtures of n-heptane and isooctane were prepared like **isooctane: n-heptane** -40:60, 50:50, 60:40, 70:30 etc. Each mixture is taken into an internal combustion engine and its knocking is observed and matched with the knocking characteristic of the petrol sample under examination.

For eg: Octane number of the petrol sample is matches with 70:30 mixture, the octane number of the test sample is 70. So "octane number can be defined as the percentage of isooctane in a mixture of isooctane and n-heptane and whose knocking of the petrol sample under examination".

**LEAD petrol:** The variety of petrol in which tetra ethyl lead is added, it is leaded petrol.

$$C_2H_5$$
  
 $C_2H_5 - Pb - C_2H_5$  Tetra ethyl lead (TEL)  
 $C_2H_5$ 

To improve the anti-knock of the petrol sample, TEL and diethyl telluride  $[(C_2H_5)_2Te]$  are added. These materials are extremely poisonous and about 1 to 1.5 ml of TEL is added per litre of petrol. When TEL is fed into the internal combustion engine along with petrol, TEL undergoes and converted to lead peroxide. This lead peroxide reacts with any hydrocarbon molecule, there by slowing down the reaction of pre-ignition. So TEL acts as a negative catalyst for pre-ignition of the petrol sample. This lead peroxide is deposited in I/C engine which is harmful. To eliminate lead peroxide, ethylene-dibromide is added, which converts lead peroxide to lead dibromide. The lead dibromide is volatile and easy escapes out along with other flue gases.

**Cetane number (knocking of diesel):** Diesel fraction is obtained between 250-320 °C during fractional distillation of petroleum. The oil contains 85% C, 12% H and calorific value is 11,000 Kcal/mole.

"Diesel knock can be defined as the rattling sound produced due to ignition-lag (the delay in the ignition of the fuel)". The structure of diesel is responsible for knocking. The hydrocarbons possess antidisel knock property in the order.

n-alkanes > naphthalenes > alkenes > branched alkanes > aromatics.

"The diesel knock is measured by cetane number. For this **hexadecane** is selected which is having **100%** antiknock property. **2-methyl naphthalene** is aromatic compound having anti knock diesel property is **zero**".





Antiknock: 100 antiknock: 0

The mixtures containing **n-hexadecane** and **2-methylnaphthalene** like 40:60, 50:50, 60:40, 70:30 etc. respectively are prepared and their anti-diesel knock property is measured by taking in a diesel engine. For eg: the diesel knock of the test sample matches with 60:40 mixtures, the cetane number of the diesel is 60. "So the cetane number can be defined as the % of hexadecane in a mixture of 2-methyl naphthalene whose diesel knock matches with the diesel knock of the test sample under examination". For low speed diesel engines the cetane number is 25, for high speed diesel engines the cetane number is 45 and 35 for medium speed diesel engines.

#### **Gaseous Fuels**

Natural gas is the primary gaseous fuel. A variety of secondary fuels are obtained from coal (or) petroleum. They include coal gas, producer gas, water gas derived from coal and LPG, CNG derived from petroleum and natural gas.

(a) Natural Gas: Natural gas is primarily methane gas and it is a fossil fuel. Commercially natural gas produced from oil fields and natural gas fields. It is also called **mars** gas. It consists of methane and other saturated hydrocarbons. The average composition of natural gas is as follows.

Composition: Average composition of a natural gas is;

 $CH_4 = 70-90\%$ 

 $C_2H_6 = 5-10\%$ 

 $H_2 = 3\%$ 

CO+CO<sub>2</sub>=0.7%

Its calorific value varies from 12,000 to 14,000 kcal/m<sup>3</sup>. If natural gas contains lower hydrocarbons like methane and ethane it is called **lean or dry gas.** In the natural gas contains higher hydrocarbons like propane, butane along with methane it is called **rich or wet gas.** 

Appreciable quantities of H<sub>2</sub>S can be removed from natural gas by scrubbing with monomethanol amine.

Applications: 1. It is used as a very good domestic fuel.

- 2. It is used in the preparation of ammonia (used for urea manufacturing).
- 3. It is used to prepare carbon-black which is used as filler for rubber industry.
- 4. A large number of chemicals are synthesised from natural gas.
- 5. It is also used for the generation of electricity by using it in fuel cells.
  - (b) LPG (liquefied Petroleum Gas): LPG is a common fuel for domestic and industries. The main components of LPG are n-butane, isobutane, butylene and propane. LPG is dehydrated and desulphurised. LPG is highly inflammable and consists of hydrocarbons can be readily liquefied under

pressure and exists as gas at room temperature. Its Calorific Value is 27,800 kcals/m3 and LPG is supplied under different trade names like Indane, H.P etc.

#### Composition

Its approximate composition is

n-Butane = 70 %

Isobutane = 17 %

n-Propane = 11 %

Butylene and Ethane = rest.

#### Uses

- i. It is used as a fuel for domestic cooking.
- ii. Used for heating industrial furnaces.
- iii. Used as an alternate for Gasoline in automobiles.

#### **Disadvantages**

- 1. It is difficult to handle as fuel.
- 2. Engines working at low compression ratio cannot use LPG as fuel.
  - (c) CNG (Compressed Natural Gas): Natural gas contains mainly CH<sub>4</sub>. When natural gas is compressed at high pressure (1000 atm) or cooled to –160°C, it is converted into CNG. It is now replacing gasoline as it releases less pollutant during its combustion. It is environmentally clean alternative to those fuels which produce toxic pollutants. In some of the metro cities, CNG-vehicles are used to reduce pollution. LNG (liquified natural gas) is different from CNG. LNG is costlier than CNG. Its calorific value is about 13000 kcal/m3

#### **Advantages of CNG**

- 1. Due to higher temperature of ignition, CNG is better fuel than petrol and diesel.
- 2. Operating cost of CNG is less. Cost of production is less. It can be easily stored.
- 3. It releases least pollutants like CO and unburnt hydrocarbons.
- 4. It undergoes regular combustion.

#### Disadvantages

- 1. Faint odour; leakage cannot be detected easily.
- 2. CNG tanks require a large tank space.
- 3. Refueling network for CNG is very expensive.

#### **COMBUSTION**

Combustion is an exothermic chemical reaction. It is defined as "the process of oxidation of the fuel by oxygen and subsequent liberation of energies like heat, light etc". For example, combustion of carbon in oxygen:

$$C_{(s)} + O_{2(s)} \longrightarrow CO_{2(g)} + 97 \text{ kcal}$$

#### Factors affecting the rate of combustion

The rate of combustion depends on the following factors:

- 1. The concentration of the fuel and air.
- 2. The nature of the combustible substance
- 3. The temperature
- 4. With increase in pressure or surface area of the fuel the rate of combustion can be increased.
- 5. It increases with increase in pressure of air.
- 6. It Increases with preheating of fuel and air.

**Calorific value of a fuel:** "Calorific value can be defined as the amount of heat produced by the combustion of unit mass (or) unit volume of a fuel. The higher the calorific value, the better will be the quality of fuel. There are different units for measuring the quantity of heat. They are

- (a) Calorie: It is the amount of heat required to increase the temperature of 1 gm of water through one degree centigrade.
- **(b) Kilocalorie:** The amount of heat required to increase the temperature of **1kg** of water through **one degree centigrade** (i.e. 15  $^{0}$ C to 16  $^{0}$ C).
- (c) **British thermal unit (BTU):** The amount of heat required to raise the temperature of one pound of water by 1°F (Fahrenheit) (60 °F to 61 °F) is BTU.

$$1 \text{ BTU} = 252 \text{ cal} = 0.252 \text{ kcal}$$
  
 $1 \text{ kcal} = 3.968 \text{ BTU}$ 

(d) Centigrade heat unit (C.H.U): It is the amount of heat required to raise the temperature of one pound of water through 1°C.

$$1 \text{ kcal} = 3.968 \text{ BTU} = 2.2 \text{ CHU}$$

Inter conversion of various units of heat

1 
$$kcal = 1000 cal = 3.968 BTU = 2.2 CHU$$

Types of calorific values: (1) Gross calorific value (G.C.V.) or High calorific value

- (2) Net calorific value (N.C.V.) or lower calorific value
- (1) Gross calorific value (G.C.V.) or High calorific value: The higher calorific value can be defined as the total heat generated when a unit quantity of fuel is completely burnt and the products of combustion are cooled down to cooled down to  $60^{\circ}$  For  $15^{\circ}$  C.

For example, when a fuel containing hydrogen is burnt, it under goes combustion and will be converted to steam. If the combustion product is cooled to room temperature, the steam gets condensed into water and the latent heat is evolved. Therefore the latent heat of combustion of condensation of 'steam' so liberated is included in gross calorific value.

(2) Net (or) Lower Calorific Value (NCV or LCV): Lower the calorific value is defined as the net heat produced when a unit quantity of fuel is completely burnt and the products of combustion are allowed to escape into the atmosphere. LCV does not include latent heat of steam (or) water vapour.

#### Relationship between HCV (GCV) and LCV (NCV)

(The energy required in calories to completely convert 1 gm of H<sub>2</sub>O to steam without increase the temperature is called latent heat of steam). **The latent heat of steam is 587 cal/gm.** 

Since 1 part by weight of H<sub>2</sub> produces 9 parts by weight of H<sub>2</sub>O as given by the equation below

$$H_2$$
 +  $\frac{1}{2}$   $O_2$   $\longrightarrow$   $H_2O$   
2gms 16gms 18gms  
1 8 9

Hence, LCV = HCV-(mass of hydrogen X 9 X latent heat of steam)

$$NCV = GCV - \frac{9}{100}H \times 587 \text{ kcal/kg}$$

$$NCV = GCV - 0.09H \times 587 \text{ kcal/kg}$$

where, 
$$H = \%$$
 of  $H_2$  in the fuel.

#### **Dulong's formula (Theoretical calculation)**

Dulong's formula for the theoretical calculation of calorific value is

GCV (or) HCV = 
$$\frac{1}{100}$$
 (8080  $C + 34500 [H - \frac{O}{8}] + 2240 S) kcal/kg$ 

where, C, H, O & S represent the % of the corresponding elements in the fuel.

It is based on the assumption that the calorific values of C, H & S are found to be 8080, 34500 and 2240 kcal, when 1 kg of the fuel is burnt completely. However, all the oxygen in the fuel is assumed to be present in combination with hydrogen in the ratio H: O as 1:8 by weight. So the surplus hydrogen available

for combustion is 
$$H - \frac{O}{8}$$
.

"The amount of heat produced when 1 gm molecule of the reactants are burnt in the presence of oxygen" for example  $C + O_2 \longrightarrow CO_2 + 96960$  cal

12 gm of carbon liberates 96960 calories of heat

1 gm of carbon liberates = 96960/12=8080 cal.

Similarly H<sub>2</sub> burns in the presence of O<sub>2</sub>, resulting in the formation of

$$H_2 + 1/2O_2 \longrightarrow H_2O + 69000 \text{ cal}$$

2gm 16gm 18 gm

2 gm of  $H_2$  liberates 69000 calories of heat, so 1gm of  $H_2$  liberates = 69000/2= 34500 cal.

$$S + O_2 \longrightarrow SO_2 + 71680 \text{ cal}$$
  
32 gm 32 gm 96 gm

1 gm of Sulphur produces = 71680/32= 2240 cal.

1) Calculate the gross and net calorific value of a coal sample having the following composition C= 80%, H=7%, O=3%, S=3.5 %, N=2% and ash 5%.

GCV (or) HCV = 
$$\frac{1}{100}$$
 (8080  $C + 34500 [H - \frac{O}{8}] + 2240 S$ ) kcal/kg

=1/100 (8080x34500 (7-3/8)+2240x3.5 HCV=8828 Kcal/kg

> And LCV=HCV-0.09HX587 8828-0.09x7587 LCV=8458 Kcal/kg

```
net colorafic value of coal containing
                                 C= 85%, H2=8%, S= 1%, N2=2%,
(2) calculate the gross and
  02 = 2%, agh = 2% and latent heat of steam Ps 587 cal/gm.
  the following composition.
  :. The gross adortific value of the fuel 98 calculated by making
            Dulong's formula.
         GeV= 100 [8080xc+34500 (H-0)+2240 XS] callym
  use of
              = \frac{1}{100} \left[ 8080 \times 85 + 34500 \left( 8 - \frac{2}{8} \right) + 2240 \times 1 \right]
              = 100 [68 6800 + 34500 (7.75) + 2240]
                = 956415 = 9565.15 callgon
   2 Net calorific value = GICV - (0.09 XH X587)
                        = 9564.15 - (0.09 x8 x587)
                         = 9564.15 - 422.64 = 9141-51 cal) gm.
```

A coal has the following composition by weight C = 90%, 0 = 3%, S = 0.5%, N = 0.5% and agh = 2.5%. The net caloritie 8490.5 Kcal/ kg. calculate value of the fuel was found to be percentage of hydrogen and HCV of the fuel. : Higher calorific value = 100 [8080xc +34500 (H-0)+2240xs] alg  $= \frac{1}{100} \left[ (8080 \times 90) + 34500 \left( H - \frac{3}{8} \right) + 2240 \times 0.5 \right]$ = 7272 + (345H -129.4) + 11.2 callgm = [7153.8 + 345 H] callom Lower coloristic value = HCV - (0.09 XH X587) (LCV) HCV = LCV + 0.09# X587 = 8490.5 + 52.8 H callgm 7153°8 + 345 H = 8490,5 + 52.8 H 292.2 H = 8490.5 - 7153.8 H = 8490.5 - 7153.8 29212 H= 4.575%. o/o of H= 4.58%

HCV = 8490 + (0.09 x4.58 x 587) = 8732.5 cal gm.

1. Calculate the Gross and Net calorific values of a coal having the following compositions, C = 80 %,  $H_2 = 08 \%$ ,  $O_2 = 08 \%$ , S = 2 % and ash=2. Latent heat of steam is = 587 cal/gm.

#### Solution

(i) Gross Calorific Value (GCV)

GCV = 
$$\frac{1}{100}$$
(8080  $C + 34500 [H - \frac{O}{8}] + 2240 S$ )  
=  $\frac{1}{100}$ (8080 × 80 + 34500 [8 -  $\frac{8}{8}$ ] + 2240 × 2 ) kcal/kg  
=  $\frac{1}{100}$ (646400 + 241500 + 4480) kcal/kg  
=  $\frac{1}{100}$ (892380) kcal/kg  
= 8923.8 kcal/kg.

(ii) Net Calorific Value (NCV)

= 
$$GCV - \frac{9}{100}H \times 587$$
 kcal/kg  
=  $8923.8 - \frac{9}{100} \times 8 \times 587$  kcal/kg  
=  $8923.8 - 422.64$  kcal/kg  
=  $8501.16$  kcal/kg

2. Calculate the Gross and Net calorific values of a coal having the following compositions, C = 63%,  $H_2 = 19\%$ ,  $O_2 = 03\%$ , S = 13% and ash=2. Latent heat of steam is = 587 cal/gm.

#### **Solution**

(i) Gross Calorific Value (GCV)

GCV = 
$$\frac{1}{100}$$
(8080 C + 34500 [ $H - \frac{O}{8}$ ] + 2240 S) kcal/kg  
=  $\frac{1}{100}$ (8080 × 63 + 34500 [19- $\frac{3}{8}$ ] + 2240 ×13 ) kcal/kg  
=  $\frac{1}{100}$ (509040 + 64562 + 29120) kcal/kg  
=  $\frac{1}{100}$ (1180722) kcal/kg

 $= 11807.22 \ kcal / kg.$ 

(ii) Net Calorific Value (NCV)

= 
$$GCV - \frac{9}{100}H \times 587$$
 kcal/kg  
=  $11807.22 - \frac{9}{100} \times 19 \times 587$  kcal/kg  
=  $11807.22 - 1003.77$  kcal/kg  
=  $10803.45$  kcal/kg

3. Calculate the Gross and Net calorific values of a solid fuel having 80% of carbon & 20% of hydrogen. Latent heat of steam is = 587 cal/gm.

#### Solution

(i) Gross Calorific Value (GCV)

GCV = 
$$\frac{1}{100}$$
 (8080 C + 34500 [H- $\frac{O}{8}$ ] + 2240 S) kcal/kg

Here, the % of  $H_2$  and S are Zero.

$$= \frac{1}{100} (8080 \times 80 + 34500 [20 - \frac{0}{8}] + 2240 \times 0) \text{ kcal/kg}$$

$$= \frac{1}{100} [646400 + 690000] \text{ kcal/kg}$$

$$= \frac{1}{100} [1336400] \text{ kcal/kg}$$

 $= 13364 \, kcal / kg.$ 

(ii) Net Calorific Value (NCV)

$$= GCV - \frac{9}{100}H \times 587 \text{ kcal/kg}$$

$$= 13364 - \frac{9}{100} \times 20 \times 587 \text{ kcal/kg}$$

$$= 13364 - 1056.6 \text{ kcal/kg}$$

$$= 12307.4 \text{ kcal / kg}$$

4. A coal sample on analysis gives C = 75%,  $H_2 = 6\%$ ,  $O_2 = 3.5\%$  S = 03% and the rest ash. Calculate the Gross and Net calorific values of the fuel. Latent heat of steam is = 587 cal/gm

#### **Solution**

(i) Gross Calorific Value (GCV)

GCV = 
$$\frac{1}{100}$$
 (8080  $C + 34500$  [ $H - \frac{O}{8}$ ] + 2240  $S$ ) kcal/kg  
=  $\frac{1}{100}$  (8080 × 75 + 34500 [ $6 - \frac{3.5}{8}$ ] + 2240 ×3 ) kcal/kg  
=  $\frac{1}{100}$  [606000 + 191906 + 6720] kcal/kg  
=  $\frac{1}{100}$  [804626] kcal/kg  
=  $80462.6$  kcal/kg.

(ii) Net Calorific Value (NCV)

$$= GCV - \frac{9}{100}H \times 587 \text{ kcal/kg}$$

$$= 80462.6 - \frac{9}{100} \times 6 \times 587 \text{ kcal/kg}$$

$$= 80462.6 - 316.98 \text{ kcal/kg}$$

$$= 80145.62 \text{ kcal / kg}$$

5. On analysis, a coal sample has the following composition by weight; C = 75 %,  $O_2 = 04 \%$ , S = 05 %, and ash = 3%. Net calorific value of the fuel is 9797.71kcal / kg. Calculate the percentage of hydrogen and gross calorific value of coal.

#### Solution

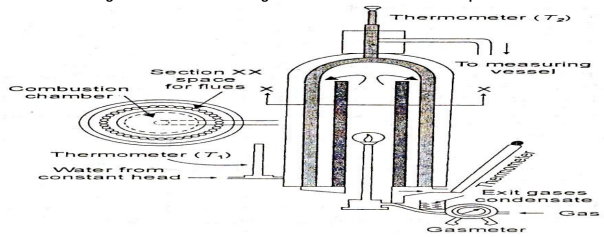
(i) Gross Calorific Value (GCV)

GCV = 
$$\frac{1}{100}$$
(8080 × 75 + 34500 [13  $-\frac{4}{8}$ ] + 2240 ×5 ) kcal/kg  
GCV =  $\frac{1}{100}$ [606000 + 431250 + 11200] kcal/kg  
=  $\frac{1}{100}$ [1048450] kcal/kg  
= 10484.5 kcal/kg.

#### (Q) Determination of calorific value of gaseous fuel by Junker's gas Calorimeter?

The calorific value a fuel is determined by the combustion of the fuel in a special type apparatus called calorimeter. Junker's gas calorimeter is used for measuring the calorific values of gaseous and liquid fuels which get vaporised easily. This calorimeter works on junker's principle. According to this principle

#### Calorific value of gaseous fuel X volume of gas=volume of water X rise in temperature of water



Calorific value of gaseous fuel can be determined by using Junker's calorimeter; it consists of a vertical cylindrical combustion chamber where combustion of gaseous fuel can be carried out with the help of Bunsen burner. The supply of gaseous fuel is regulated with the help of pressure governor. The volume of gas, flowing in a particular time, is measured with the help of gasometer. The combustion chamber is surrounded by an annular water space. Inside the outer flues, heat exchanger coil are also fitted. Radioactive and convective heat loss from the calorimeter is prevented with the help of outer jacket which is chromium plated. Moreover, the outer jacket contains air which is very good heat insulator. Around the combustion chamber, there is an annular space where water is made to circulate. At the appropriate places there are the openings where thermometers arplaced for measuring the temperatures of the inlet and outlet water.

A known volume of gas is burned in excess of air at a constant rate in combustion chamber in such a manner that all the heat produced is absorbed in water. Water is flowing at a constant rate in annular space around the combustion chamber. The increase in the temperature of the water is measured and the heat evolved from the burning of the gas can be readily calculated. The weight of water flowing is also recorded for the calculation of calorific value of gaseous fuel.

Experimental calculations and observations: Let "V" be the volume of gaseous fuel burnt at a given temperature and pressure in a certain time "t"

Volume of water used for cooling combustion products in the "t" = w

Weight of steam condensed in time "t"=m

Temperature of inlet water= t<sub>1</sub><sup>0</sup>C

Temperature out let water= t<sub>2</sub><sup>0</sup>C

Rise in temperature=  $(t_2-t_1)$ 

Mass of water condensed in the out let water in time "t" is "m" kg and hence =m/v

If L kcal/volume is the calorific value of gaseous fuel, then Heat produced by fuel=V x L

Heat absorbed by water (w kg) = Wx  $(t_2-t_1)$ 

By junker's principle  $V \times L = W \times (t_2-t_1)$  (gross calorific value)

HCV or calorific value of gas  $L = W \times (t_2-t_1)/V$ 

If latent heat of steam is 587 kcal/kg, then

 $HCV=(M \times 587)$ 

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So, Net calorific value (NCV) =  $(L-(M \times 587) \text{ kacl/m} 3)$ 

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#### RENEWABLE ENERGY SOURCES

The term energy means capacity to do work. Energy can neither be created nor destroyed, but transformed one form to another. Energy is present in a number of forms such as mechanical, thermal, biological energy etc. There are two types of energy sources namely

- (1) Renewable energy sources
- (2) Non-renewable energy sources

Energy exists freely in nature. Some of them exists infinitely (never run out called renewable). The important renewable energy sources are (1) solar energy (2) wind energy (3) hydropower energy (4) biomass energy

(1) Solar energy: The energy is derived from the sun is known as solar energy. It can be used for direct heating (or) sun's heat is converted into electricity. Solar cells (or) photovoltaic cells are devices that convert light energy directly into electrical energy. Solar cells consist of two semiconductor materials joined together. Silicon is "doped" with phosphorous to create an "n" type semiconductor, which is joined to silicon "doped" with boron "p" type semiconductor to create a p-n junction.

When the solar cell is exposed to sunlight, energy from sunlight excites electron from n-type silicon to the holes of p-type silicon. From the p-type silicon, electrons flow through the external circuit as an electric current. The voltage of solar cell is not large, so a series of solar cells are connected to get a solar panel.

#### Advantages:

Solar cells make absolutely no noise at all.

Solar cells create absolutely no pollution (oil burning release harmful greenhouse gases, carcinogens and CO<sub>2</sub> into air).

Very little maintenance is required to keep solar cells running.

Solar panels may quite expensive, long you run can use.

A number of solar equipment's have been developed to utilize to heat water, to cook food, to pump water, to certain machines and used for street lighting, railway signals etc.

#### **Disadvantages:**

The major disadvantage is solar energy does not produce during cloudy weather, night time, rains and other natural conditions.

(2) Wind energy: Moving air is called wind. Energy recovered from the force of wind is called wing energy. The energy possessed by wind is of its high speed. The earth's surface has both land and water. When the sun fall down, the air over the land heats up quicker than that over water. The heated air is lighter and its rises.

The moving air (wind) has huge amounts of kinetic energy (K.E), and this can be transferred into electrical energy using wind turbines and connects to a generator.

#### Advantages:

- (a) It does not cause any air pollution.
- (b) Reduce fossil fuel consumption.
- (c) Environmentally friendly (not cause for any pollution).
- (d) Low operating/maintenance costs.
- (e) High net energy yield.

#### Disadvantages:

- (a) Public resists for locating the wind forms in populated areas due to noice generated by machines and loss of aesthetic appearance.
- (b) Wind forms located on the migratory routes of birds will cause hazards.
- (c) Wind forms produce unwanted sound.
- (d) Wind turbines interface with electromagnetic signals (TV, Radio signals).
- (e) Low energy density of wind (Must use large areas of land).
- (f) Wind turbines cannot work, if there is no wind (or) if wind speed is so high it would damage them.
  - The Largest wind turbine in the world is located in Hawaii, and has blades length of a football court.
  - The average wind speed of 14 miles per hour is needed to convert wind energy into electricity.
  - One wind turbine can produce enough electricity to power up to 300 homes.
- (3) Hydropower: Hydropower is one of the oldest methods of producing power. Hydropower is energy is obtained from flowing water. Electrical power is generated by hydro-electric projects in which dams are constructed across the river. The kinetic energy of flow water is converted into mechanical energy by means of turbines and in turn, the mechanical energy transferred into electrical energy by generators.

#### Advantages:

- 1. Cheap to operate.
- 2. Long life and lower operating costs than all other power plants.

- 3. Renewable.
- 4. High yield of energy obtained.
- 5. Lower energy cost than any other method.
- 6. Some countries depend almost entirely on it.
- 7. Not intermittent (if reservoir is large enough).
- 8. Reservoirs have multiple uses.
- 9. Flood control, drinking water, aquaculture, recreation.
- 10. Less air pollution than fossil fuel combustion.

#### **Disadvantages:**

- 1. Human population displacement.
- 2. Reduces availability of water downstream

#### (a) Ecosystem impacts:

- 1. Barriers to migrating fish
- 2. Loss of biodiversity both upstream and downstream
- 3. Coastal erosion
- 4. Reduces nutrient flow (dissolved and particulate)

#### (b) Water pollution problems:

- 1. Low dissolved oxygen (DO)
- 2. Increased H<sub>2</sub>S toxicity; other DO-related problems
- 3. Siltation a big problem (also shortens dam life).
- (4) Biomass energy: Bio-mass is an organic material from living beings or its residues. It is a renewable sources of energy derived from the waste of various human and natural activities. The bio-mass energy sources include wood, animal manure, sugar cane waste, agriculture crops, house hold waste, roots of plants, garbage etc.

The simplest way of using bio-mass energy sources is to allow them to dry out in the sun and burn them. Liquid fuels including ethanol, methanol, biodiesel and gaseous fuels such as methane (CH<sub>4</sub>) derived from bio-mass feed stocks. Ethanol is made from starch (or) sugar.

Bio gas is a mixture of methane (CH<sub>4</sub>) (68%), CO<sub>2</sub> (31%) and N<sub>2</sub> (1%). Bio gas produced from natural organic wastes of cattle dung, human excretia, poultry waste, plant leaves, paddy husk. Bio gas used as cooking gas whose calorific value is 4400-6200 kcal. Heat value of bio gas can be improved by reducing its  $CO_2$  content. "Bio gas another name is also called gobar gas".

#### **Advantages:**

- (a) It is renewable energy.
- (b) No harmful emissions.
- (c) It is clean energy source.
- (d) Reduce dependency on fossil fuels
- (e) Reduce land fills (waste is harmful to the environment and occupy the land. So waste be burned to create useful biomass energy)
- (f) Prepare different products from biofuels.
- (g) Emits less SO<sub>2</sub> and NO<sub>x</sub> than fossil fuel.

#### **DISADVANTAGES:**

- (a) Low energy density/yield: In some cases (eg, corn-derived bioethanol) may yield no net energy
- (b) Land conversion:
- (i) Biodiversity loss.
- (ii) Possible decrease in agricultural food productivity
- (iii)Usual problems associated with intensive agriculture, (a) Nutrient pollution (b) Soil depletion (c) Soil erosion

**Biodiesel**: Biodiesel is renewable and clean burning fuel that is made from waste vegetable oils, animal fats and restaurant wastes. "Biodiesel can be defined as a fuel comprised of mono-alkyl esters of long chain fatty acids derived from vegetable oils (or) animal fats, designated as B100".

Biodiesel is made through a chemical process is called "trans esterification". There are 3 basic routes to biodiesel production oils and fats. (a) Base catalysed trans- esterification of the oil.

- (b) Acid catalysed trans- esterification of the oil.
- (c) Conversion of the oil to its fatty acids and then bio diesel.

Reaction for Biodiesel Formation: Generally base catalysed trans-esterification method widely used for the synthesis of biodiesel.

During the chemical conversion of vegetable oil to biodiesel we get water soluble glycerol and a small amount of sodium soaps. The water soluble part can be easily separated from biodiesel by washing the mixture with water. The alkaline sodium methoxide catalyst, saponifies some small amount of oil to give soap.

Compounds present in biodiesel are like,

methyl palmitate  $H_3C$  -  $(CH_2)_{14}$  -  $COOCH_3$ 

methyl stearate  $H_3C$  -  $(CH_2)_{16}$  -  $COOCH_3$ 

methyl oleate  $H_3C - (CH_2)_7 - CH = CH - (CH_2)_7 - COOCH_3$ 

methyl linoleate  $H_3C - (CH_2)_5 - (CH = CH)_2 - (CH_2)_7 - COOCH_3$ 

Biodiesel produces less toxic pollutants and greenhouse gases than petroleum diesel. It can be used pure form (B100) or blended with diesel in the form of B2 (2% biodiesel, 98% diesel), B5 (5% biodiesel, 95% diesel), B20 (20% biodiesel, 80% diesel). Biodiesel can be used in any diesel engine with little (or) no modification to the engine (or) fuel system.

#### **Advantages of Biodiesel:**

- (i) Biodiesel is renewable unlike petroleum based diesel.
- (ii) Biodiesel is safer to handle compared to standard diesel.
- (iii) Biodiesel can be easily blended with standard diesel.
- (iv) Biodiesel can help cut our reliance on fossil fuels.
- (v) Biodiesel emitting significantly less harmful carbon emission compared to standard diesel.
- (vi) Biodiesel has very good lubricating properties, significantly better than standard diesel.
- (vii) Biodiesel has shorter ignition delay compared to standard diesel.
- (viii) Biodiesel has no sulphur content, and it does not acid formation.

#### **Limitations of Biodiesel:**

- (i) Biodiesel is currently mostly produced from corn, which could lead to food shortages and increased fodd prices.
- (ii) Biodiesel is 20 times more susceptible to water contamination compared to standard diesel. This could lead to corrosion.
- (iii)Cloud and pour points of biodiesel are higher than diesel and can cause problem in fuel flow line. So it cannot be used in cold regions.
- (iv) Biodiesel is significantly more expensive compared to around 11% to standard diesel.
- (v) Biodiesel can release nitrogen oxide, which can lead to the formation smog.

#### **Calculation of Air for Combustion**

Substances always combine in definite proportions. These proportions are determined by their molecular masses.

 $C + O_2 \rightarrow CO_2$  (12:32:44)

- (i) 22.4 L of any gas at 0°C and 760mm pressure (STP) has a mass equal to its 1 mol.
- (ii) Air contain 21% of oxygen by volume and 23% of oxygen by mass.
- (iii) 28.94 g/mol is taken as molar mass of air.
- (iv)  $O_2$  required for combustion = theoretical  $O_2$  required  $O_2$  present in the fuel.
- (v) Dry flue gases means products of combustion except moisture.

For solid or Liquid fuels:

$$O_2$$
 quantity =  $\left(\frac{32}{12}C + 8H + S - O\right)$  kg.

Where C, H, S and O are amounts of elements in Kg.

Air quantity = 
$$\frac{oxygen\ quantity\ X\ (100+\%\ excess\ air)}{23}$$
 Kg.

For Gaseous fuels:

 $O_2$  volume required = volume gas component in m<sup>3</sup> x volume of  $O_2$  per volume of gas.

Air volume required = 
$$\frac{Oxygen\ volume\ X\ (100+\%\ excess\ air)}{21}$$
 m<sup>3</sup>.

Ex 10) A sample of a fuel contains the following by weight C=81%, H=4%, O2=2%, N2=10%, S=1% the reminder is ash. Calculate the volume of air required for the perfect combustion of 1kg of fuel assuming STP conditions.

S.No.	Percentage composition	Actual composition in 1 kg of fuel	Combust reaction	Wt. of O <sub>2</sub> required for combustion
1.	C = 81	0.81 kg	$\begin{array}{ccc} C + O_2 \rightarrow CO_2 \\ 12 & 32 & 44 \end{array}$	$\frac{0.81 \times 32}{12} = 2.16  \text{kg}$
2.	H = 4%	0.04 kg	$H_2 + \frac{1}{2}O_2 \rightarrow H_2O$ 2 16 18	$\frac{0.04 \times 16}{2} = 0.32 \text{k}$
3.	O <sub>2</sub> = 2%	0.02 kg	Oxygen present in the fuel must be subtracted from the wt. of O <sub>2</sub> required	- 0.02
4.	$N_2 = 10$	0.10 kg	$N_2$ is non-combustible does not require $O_2$ for combustion	
5.	S = 1%	0.01	$S + O_2 \rightarrow SO_2$ 32 32 64	$\frac{0.01 \times 32}{32} = 0.01$

Total Weight of  $O_2$  required for combustion = 2.47 kg

```
Volume of Oxygen required: 32 gms of O_2 at STP occupies a volume of 22.4 litres 2470 gms of O_2 at STP occupies a volume of =\frac{2470\times22.4}{32}=1729l Wt. of O_2 required for combustion: 100 kg of air contains 23 kg of O_2? of air contains 2.47 kg of O_2 Wt. of O_2 required for combustion =\frac{2.47\times100}{23}=10.74 kg Vol. of air required for combustion: 100 litres of air contains 21 litres of O_2? of air contains 1729 I of O_2
Volume of air required for combustion =\frac{1729\times100}{21}=8233.3I
```

Ex.11) A petrol sample contains 14 % H and 86 % carbon. Calculate the quantity of air required for complete combustion of 1 kg petrol.

Soln.:

Given: Weight of petrol fuel = 1 kg. 
$$C \% = 86$$
 and  $H \% = 14$ ,

$$C = Weight \ of \ carbon \ in \ 1 \ kg \ petrol = \frac{86}{100} \ \times \ 1 = 0.86 \ kg$$

H = Weight of hydrogen in 1 kg petrol = 
$$\frac{14}{100} \times 1 = 0.14$$
 kg.

$$O_2$$
 quantity =  $\left(\frac{32}{12}C + 8H + S - O\right)$  kg.

$$= \left(\frac{32}{12} \times 0.86 + 8 \times 0.14 + 0 + 0\right) \text{ kg}.$$

Excess air used = zero

Air quantity = 
$$\frac{\text{Oxygen quantity} \times (100 + 0 \text{ excess})}{23}$$
  
=  $\frac{3.413 \times 100}{23}$  = 14.839 kg air ...Ans.

Ex. 12) Volumetric analysis of producer gas is,  $H_2 = 20 \%$  CO = 22 %  $N_2 = 50 \%$ ,  $CH_4 = 2 \%$  and  $CO_2 = 6 \%$ . Find volume of air required for complete combustion of 1 m<sup>3</sup> of the gas

Soln.:

Volume of 
$$H_2$$
 in 1 m<sup>2</sup> gas =  $\frac{20}{100} \times 1 \text{ m}^2 = 0.2 \text{ m}^3$ 

Similarly, Volume of 
$$CO = 0.22 \text{ m}^3$$

Volume of 
$$CH_4 = 0.02 \text{ m}^3$$

(N2 and CO2 are not combustible)

Reaction	Volume of O <sub>2</sub> required
$H_2 + \frac{1}{2} O_2 \rightarrow H_2 O$	$\frac{1}{2}$ × volume of $H_2 = \frac{1}{2}$ × 0.2 = 0.1 m <sup>3</sup>
1 vol. 1/2 vol.	
$CO + \frac{1}{2} O_2 \rightarrow CO_2$	$\frac{1}{2} \times 0.22 = 0.11 \text{ m}^3$
$CH_4 + 2 O_2 \rightarrow CO_2 + 2 H_2 O$	$2 \times \text{volume of CH}_4 = 2 \times 0.02 = 0.04 \text{ m}^3$

Total volume of 
$$O_2$$
 required =  $0.25 \text{ m}^3$ 

Volume of air required = 
$$\frac{\text{Volume of O}_2 \times (100 + 0)}{21}$$
$$= \frac{0.25 \times 100}{21} = 1.19 \text{ m}^3$$

**Ex. 13**) A gas has following composition by volume, H2 = 20 %, CH4 = 6 %, CO = 18 %, O2 = 5 %, N2 = 43 %. If 25 % excess air is used. Find volume of air actually supplied per m3 of the gas.

Soln.: 1 m<sup>3</sup> of the gaseous fuel contains,

$$H_2 = \frac{20}{100} \times 1 = 0.2 \text{ m}^3, CH_4 = \frac{6}{100} \times 1 = 0.06 \text{ m}^3,$$

$$CO \; = \; \frac{18}{100} \, \times 1 = 0.18 \; m^3, \, O_2 = \frac{5}{100} \, \times 1 = 0.05 \; m^3.$$

Reaction	Volume of O2 required
$H_2 + \frac{1}{2} O_2 \rightarrow H_2 O$	Volume of gas $\times \frac{1}{2} = 0.2 \times \frac{1}{2} = 0.1 \text{ m}^3$
1 vol. $\frac{1}{2}$ vol.	
$CH_4 + 2 O_2 \rightarrow CO_2 + H_2 O$ 1 vol. 2 vol.	Volume of gas $\times$ 2 = 0.06 $\times$ 2 = 0.12 m <sup>3</sup>
$CO + \frac{1}{2} O_2 \rightarrow CO_2$	Volume of gas $\times \frac{1}{2} = \frac{0.18}{2} = 0.09 \text{ m}^3$
1 vol. ½ vol.	
	Total $O_2$ required = $0.1 + 0.12 + 0.09$
	$= 0.31 \text{ m}^3$
	Less O <sub>2</sub> present in fuel = -0.05
	Net O <sub>2</sub> required = 0.26 m <sup>3</sup>

Volume of 
$$O_2 = 0.26 \text{ m}^3$$

Volume of air with 25 % excess

$$= \frac{\text{Volume of oxygen} \times (100 + 25)}{21} = \frac{0.26 \times 125}{21}$$

$$= 1.548 \text{ m}^3. \dots \text{Ans.}$$

 $\therefore$  Actual volume of air supplied with 25 % excess for combusting 1 m<sup>3</sup> of the gaseous fuel is 1.548 m<sup>3</sup>.

**Ex. 14)** A gas has following composition by volume :  $H_2 = 20 \%$ ;  $CH_4 = 6\%$ ; CO = 22%; = CO2 O = 4% 2 N = 4% 2 = 44% Find the volume of air actually supplied per m3 of this gas.

Soln.:

Volume of 
$$H_2$$
 in  $1m^3$  gas =  $\frac{20}{100} \times 1 = 0.2 \text{ m}^3$   
Volume of  $CH_4$  in  $1m^3$  gas =  $\frac{6}{100} \times 1 = 0.06 \text{ m}^3$   
Volume of  $CO$  in  $1m^3$  gas =  $\frac{22}{100} \times 1 = 0.22 \text{ m}^3$   
Volume of  $O_2$  in  $1m^3$  gas =  $\frac{4}{100} \times 1 = 0.04 \text{ m}^3$ 

(CO2 and N2 do not burn and do not require air).

(eeg and 1,2 do not built and do not require air).		
Reaction	Volume of O <sub>2</sub> required = Volume of gas × Volume of O <sub>2</sub> per molecule	
$H_2 + \frac{1}{2}O_2 \longrightarrow H_2O$	$0.2 \times \frac{1}{2} = 0.1 \mathrm{m}^3$	
$CH_4 + 2 O_2 \longrightarrow CO_2 + 2 H_2O$	$0.06 \times 2 = 0.12 \text{ m}^3$	
$CO + \frac{1}{2}O_2 \longrightarrow CO_2$	$0.22 \times \frac{1}{2} = 0.11 \mathrm{m}^3$	
	Total $O_2 = 0.33 \text{ m}^3$	
	Less $O_2$ in fuel = $0.33 - 0.04 = 0.29$	

**Ex. 15**) A sample of coal was found to have the following composition:

H 
$$C = 75\%$$
, 2 O = 5.2%, 2 N  $S = 1.2\%$ , = 12.8%, 2 ash = 2.1% = 3.7%,

Calculate the minimum amount of air necessary for complete combustion of 1 kg of coal

Soln.:

Amount of C in 1 kg coal = 
$$\frac{75}{100} \times 1$$
 kg = 0.75 kg  
Amount of H in 1 kg coal =  $\frac{5.2}{100} \times 1$  kg = 0.052 kg  
Amount of O in 1 kg coal =  $\frac{12.8}{100} \times 1$  kg = 0.128 kg  
Amount of S in 1 kg coal =  $\frac{1.2}{100} \times 1$  kg = 0.012 kg

(N and ash do not get burnt, do not require air, during combustion of coal).

**Ex. 16**) A sample of coal requires 20% excess air for complete combustion. Calculate weight of air for 250 gm of the coal, if its composition is C = 81%, H = 4%, N = 1.5%, S = 1.2%, O = 3%, ash = 9.35.May 2010

Soln.:

Weight of carbon in 250 gm coal = 
$$\frac{81}{100} \times 250$$
 gm = 202.5 gm

Weight of hydrogen in 250 gm coal =  $\frac{4}{100} \times 250$  gm = 10 gm

Weight of sulphur in 250 gm coal =  $\frac{1.2}{100} \times 250$  gm = 3gm

Weight of oxygen in 250 gm coal =  $\frac{3}{100} \times 250 = 7.5$  gm

(N and ash do not burn, do not require oxygen)

$$O_2 \text{ quantity } = \left[\frac{32}{12} \text{ C} + 8 \text{ H} + \text{S} - \text{O}\right] \text{ gm}$$

(C, H, S, O are the weights of elements in gms)

$$\therefore O_2 \text{ quantity } = \frac{32}{12} \times 202.5 + 8 \times 10 + 3 - 7.5 \text{ gm}$$

$$= 615.5 \text{ gm oxygen}$$

$$Quantity \text{ of air } = \frac{O_2 \text{ quantity } (100 + \% \text{ excess air})}{23}$$

$$= \frac{615.5 (100 + 20)}{23}$$

$$= 3211.3 \text{ gm air}$$

$$= 3.2113 \text{ kg air}$$

Weight of coal = 250 gm

Ex. 17) A gas used in internal combustion engine contains,  $H_2=45$  %, CO=15%,  $CH_4=35$ %  $N_2=5$  % Find the minimum quantity (volume ) of air required per 1 m<sup>3</sup> of the gas for complete combustion.

Soln.:

Volume of 
$$H_2$$
 in 1  $m^3$  gas =  $45 \times 1$   $m^3$  /  $100 = 0.45$   $m^3$ 

Similarly, Volume of 
$$CH_4 = 35 \times 1 / 100 = 0.35 \text{ m}^3$$

Volume of CO = 
$$15 \times 1 / 100 = 0.15 \text{ m}$$
3

(N<sub>2</sub> is not combustible)

Reaction	Volume of O <sub>2</sub> required
$H_2 + \frac{1}{2} O_2 \rightarrow H_2 O$	$\frac{1}{2}$ × volume of H <sub>2</sub> = $\frac{1}{2}$ × 0.45= 0.225 m <sup>3</sup>
1 vol. 1/2 vol.	
$CO + \frac{1}{2} O_2 \rightarrow CO_2$	$\frac{1}{2}$ × 0.15= 0.075 m <sup>3</sup>
$CH_4 + 2 O_2 \rightarrow CO_2 + 2 H_2 O$	$2 \times \text{volume of CH}_4 = 2 \times 0.35 = 0.7 \text{ m}^3$

Total volume of O2 required = 1 m3

Volume of air required = 
$$\frac{\text{Volume of O}_2 \times (100 + 0)}{21}$$
$$= 1 \times 100 / 21 = 4.762 \text{ m}^3$$

...Ans.