UNIT 1

SOME BASIC CONCEPTS OF CHEMISTRY

Chemistry: Chemistry is the branch of science that deals with the composition, structure and properties of matter. Chemistry is called the science of atoms and molecule

Branches of Chemistry

- Organic Chemistry -This branch deals with study of carbon compounds especially hydrocarbons and their derivatives.
- Inorganic Chemistry-This branch deals with the study of compounds of all other elements except carbon. It largely concerns itself with the study of minerals found in the Earth's crust.
- Physical Chemistry-The explanation of fundamental principles governing various chemical phenomena is the main concern of this branch. It is basically concerned with laws and theories of the different branches of chemistry.
- Industrial Chemistry-The chemistry involved in industrial processes is studied under this branch.
- Analytical Chemistry-This branch deals with the qualitative and quantitative analysis of various substances.
- Biochemistry-This branch deals with the chemical changes going on in the bodies of living organisms; plants and animals.
- Nuclear Chemistry-Nuclear reactions, such as nuclear fission, nuclear fusion, transmutation processes etc. are studied under this branch.

PROPERTIES OF MATTER AND THEIR MEASUREMENT--Every substance has unique or characteristic properties. These properties can be classified into two categories – **physical properties** and **chemical properties**.

Physical properties are those properties which can be measured or observed without changing the identity or the composition of the substance. E.g. colour, odour, melting point, boiling point, density etc.

The measurement or observation of **chemical properties** requires a chemical change to occur. e.g. Burning of Mg-ribbon in air

Chemical properties are characteristic reactions of different substances; these include acidity or basicity, combustibility etc.Many properties of matter such as length, area, volume, etc., are quantitative in nature.

Metric System was based on the decimal system.

The International System of Units (SI)

The International System of Units (in French Le Systeme International d'Unitesabbreviated as SI) was established by the 11th General Conference on Weights and Measures (CGPM from *ConferenceGenerale des Poids at Measures*). The SI system has seven *base units*

Quantity	Unit	Symbol
Length	metre	m
Mass	kilogram	kg
Time	second	S
Temperature	kelvin	К
Amount of substance	mole	mol
Electric current	ampere	А
Luminous intensity	candela	Cd

Prefixes in SI system

Multiple	Prefix	Symbol
10⁻¹²	pico	р
10 ⁻⁹	nano	n
10 ⁻⁶	micro	μ
10 ⁻³	milli	m
10 ⁻²	centi	c
10⁻¹	deci	d
10	deca	da
10^{2}	hecto	h
10^{3}	kilo	k
10 ⁶	mega	Μ
10 ⁹	giga	G
10 ¹²	tera	Т

Mass and Weight-- Mass of a substance is the amount of matter present in it while weight is the force exerted by gravity on an object. The mass of a substance is constant whereas its weight may vary from one place to another due to change in gravity. The mass of a substance can be determined very accurately by using an analytical balance

Volume-- Volume has the units of $(length)^3$. So volume has units of m^3 or cm^3 or dm^3 . A common unit, litre (L) is not an SI unit, is used for measurement of volume of liquids. 1 L = 1000 mL, 1000 cm³ = 1 dm³

Density: Density of a substance is its amount of mass per unit volume.SI unit of density = SI unit of mass/SI unit of volume = kg/m^3 or $kg m^{-3}$ This unit is quite large and a chemist often expresses density in g cm⁻³.

Temperature--There are three common scales to measure temperature — $^{\circ}C$ (degree celsius), $^{\circ}F$ (degree Fahrenheit) and K (kelvin). Here, K is the SI unit.

$$^{\circ}F = \frac{9}{5}(^{\circ}C) + 32$$

K = °C + 273.15

Note—Temperature below 0 $^{\circ}$ C (i.e. negative values) are possible in Celsius scale but in Kelvin scale, negative temperature is not possible.

Scientific Notation

In which any number can be represented in the form $N \times 10^{n}$ (Where n is an exponent having positive or negative values and N can vary between 1 to 10).

e.g. We can write 232.508 as 2.32508 $\times 10^2$ in scientific notation. Similarly, 0.00016 can be written as 1.6×10^{-4} .

Precision refers to the closeness of various measurements for the same quantity. **Accuracy** is the agreement of a particular value to the true value of the result

Significant Figures

The reliability of a measurement is indicated by the number of digits used to represent it. To express it more accuratelywe express it with digits that are known with certainty. These are called as Significant figures. They contain all thecertain digits plus one doubtful digit in a number.

Rules for Determining the Number of Significant Figures

- All non-zero digits are significant. For example, 6.9 has two significant figures, while 2.16 has three significant figures. The decimal place does not determine the number of significant figures.
- A zero becomes significant in case it comes in between non zero numbers. For example, 2.003 has four significant figures, 4.02 has three significant figures.
- Zeros at the beginning of a number are not significant. For example, 0.002 has one significant figure while 0.0045 has two significant figures.
- All zeros placed to the right of a number are significant. For example, 16.0 has three significant figures, while 16.00has four significant figures. Zeros at the end of a number without decimal point are ambiguous.
- In exponential notations, the numerical portion represents the number of significant figures. For example, 0.00045 is expressed as 4.5 x 10⁻⁴ in terms of scientific notations. The number of significant figures in this number is 2, while inAvogadro's number (6.023 x 10²³) it is four.
- The decimal point does not count towards the number of significant figures. For example, the number 345601 has sixsignificant figures but can be written in different ways, as 345.601 or 0.345601 or 3.45601 all having same number of significant figures.

Retention of Significant Figures - Rounding off Figures

The rounding off procedure is applied to retain the required number of significant figures.

- 1. If the digit coming after the desired number of significant figures happens to be more than 5, the precedingsignificant figure is increased by one, 4.317 is rounded off to 4.32.
- 2. If the digit involved is less than 5, it is neglected and the preceding significant figure remains unchanged, 4.312 isrounded off to 4.31.
- 3. If the digit happens to be 5, the last mentioned or preceding significant figure is increased by one only in case ithappens to be odd. In case of even figure, the

preceding digit remains unchanged. 8.375 is rounded off to 8.38 while 8.365 is rounded off to 8.36.

Dimensional Analysis During calculations generally there is a need to convert units from one system to other. This is called **factor label method** or **unit factor method** or **dimensional analysis**.

For example- 5 feet and 2 inches (height of an Indian female) is to converted in SI unit

1 inch =
$$2.54 \times 10^{-2} \text{ m}$$

1 = $\frac{2.54 \times 10^{-2} \text{ m}}{1 \text{ inch}}$ then, 5 feet and 2 inch = 62 inch
= 62 inch $\times \frac{2.54 \times 10^{-2} \text{ m}}{1 \text{ inch}} = 1.58 \text{ m}$

Physical Classification of Matter

Properties	Solid	Liquid	Gas	
1. volume	Definite	Definite	Indefinite	
2. Shape	Definite	Indefinite	Indefinite	
3. Inter molecular	Very high	Moderate	Negligible / Very	
force of attraction			low	
4. arrangement of	Orderly arranged	Free to move	Free to move every	
molecules		within the volume	where	
5. Inter molecular	Very small	Slightly greater	Very great	
space				
7. Compressibility	Not compressible	Not compressible	Highly	
			compressible	
8. Expansion on	Very little	Very little	Highly expand	
heating				
9. Rigidity	Very rigid	Not rigid knownas	Not rigid and	
		fluid	known as fluid	
9. Fluidity	Can't flow	Can flow	Can flow	
10. Diffusion	They can diffuse	Can diffuse And	Can diffuse And	
	due to kinetic	rate of diffusion is	rate of diffusion is	
	energy of	very fast	very fast	
	liquid/gases			

Chemical Classification of matter---



Elements

An element is the simplest form of matter that cannot be split into simpler substances or built from simpler substances by any ordinary chemical or physical method. There are 114 elements known to us, out of which 92 are naturally occurring while the rest have been prepared artificially.

Elements are further classified into metals, non-metals and metalloids.

Compounds

A compound is a pure substance made up of two or more elements combined in a definite proportion by mass, which could be split by suitable chemical methods.

Characteristics of compound

- Compounds always contain a definite proportion of the same elements by mass.
- The properties of compounds are totally different from the elements from which they are formed.
- Compounds are homogeneous.
- Compounds are broadly classified into inorganic and organic compounds. Inorganic compounds are those, which areobtained from non-living sources such as minerals. For example, common salt, marble and limestone. Organiccompounds are those, which occur in living sources such as plants and animals. They all contain carbon. Commonorganic compounds are oils, wax, fats etc.

Mixtures

A mixture is a combination of two or more elements or compounds in any proportion so that the components do not lose their identity. Air is an example of a mixture Mixtures are of two types, homogeneous and heterogeneous.

Homogeneous mixtures have the same composition throughout the sample. The components of such mixtures cannot be seen under a powerful microscope. They are also called solutions. Examples of homogeneous mixtures are air, seawater, gasoline, brass etc.

Heterogeneous mixtures consist of two or more parts (phases), which have different compositions. These mixtures have visible boundaries of separation between the different constituents and can be seen with the naked eye e.g., sand and salt, chalk powder in water etc.

LAWS OF CHEMICAL COMBINATIONS

Law of Conservation of Mass (Given by Antoine Lavoisier in 1789).

It states that matter (mass) can neither be created nor destroyed.

Law of Definite Proportions or Law of Constant Composition:

This law was proposed by Louis Proust in 1799, which states that:

'A chemical compound always consists of the same elements combined together in the same ratio, irrespective of the method of preparation or the source from where it is taken'.

Law of Multiple Proportions Proposed by Dalton in 1803, this law states that:

'When two elements combine to form two or more compounds, then the different masses of one element, which combine with a fixed mass of the other, bear a simple ratio to one another'.

Gay Lussac's Law of Gaseous Volumes (Given by Gay Lussac in 1808.)

According to this law when gases combine or are produced in a chemical reaction they do so in a simple ratio by volume provided all gases are at same temperature and pressure.

e.g.H₂(g) + Cl₂(g) --- \rightarrow 2HCl(g) 1V 1V 2V

All reactants and products have simple ratio 1:1:2.

Avogadro Law (In 1811, Given by Avogadro)

According to this law equal volumes of gases at the same temperature and pressure should contain equal number of molecules.

Dalton's Atomic Theory

- All substances are made up of tiny, indivisible particles called atoms.
- Atoms of the same element are identical in shape, size, mass and other properties.
- Atoms of different elements are different in all respects.
- Atom is the smallest unit that takes part in chemical combinations.
- Atoms combine with each other in simple whole number ratios to form compound atoms called molecules.
- Atoms cannot be created, divided or destroyed during any chemical or physical change.

Atoms and Molecules

The smallest particle of an element, which may or may not have independent existence is called an atom, while the smallest particle of a substance which is capable of independent existence is called a molecule.

Molecules are classified as homoatomic and heteroatomic. Homoatomic molecules are made up of the atoms of the same element and heteroatomic molecules are made up of the atoms of the different element have different atomicity (number of atoms in a molecule of an element) like monoatomic, diatomic, triatomic and polyatomic.

Atomic Mass Unit

One atomic mass unit is defined as a mass exactly equal to one twelfth the mass of one carbon -12 atom. And 1 amu = 1.66056×10^{-24} g.

Today, 'amu' has been replaced by 'u' which is known as unified mass.

Atomic Mass

Atomic mass of an element is defined as the average relative mass of an atom of an element as compared to the mass of an atom of carbon -12 taken as 12.

Atomic mass = _____ mass of an atom

 $1/12 \text{ massofa carbon atom}(^{12} \text{ C})$

Gram Atomic Mass

The quantity of an element whose mass in grams is numerically equal to its atomic mass. In simple terms, atomic mass of an element expressed in grams is the gram atomic mass or gram atom.

For example, the atomic mass of oxygen = 16 amu

Therefore gram atomic mass of oxygen = 16 g

Molecular Mass

Molecular mass of a substance is defined as the average relative mass of its molecule as compared to the mass of an atom of C-12 taken as 12. It expresses as to how many times the molecule of a substance is heavier than 1/12th of the mass of an atom of carbon.

For example, a molecule of carbon dioxide is 44 times heavier than 1/12th of the mass of an atom of carbon. Therefore the molecular mass of CO2 is 44 amu.

It is obtained by adding the atomic masses of all the atoms present in one molecule.

Gram Molecular Mass

A quantity of substance whose mass in grams is numerically equal to its molecular mass is called gram molecular mass. In simple terms, molecular mass of a substance expressed in grams is called gram molecular mass.

e.g., the molecular mass of oxygen = 32 amu

Therefore, gram molecular mass of oxygen = 32 g

Formula Mass-

Sum of atomic masses of the elements present in one formula unit of a compound. It is used for the ionic compounds.

Mole Concept.

Mole is defined as the amount of a substance, which contains the same number of chemical units (atoms, molecules, ions or electrons) as there are atoms in exactly 12 grams of pure carbon-12.

A mole represents a collection of 6.022 $\times 10^{23}$ (Avogadro's number) chemical units..

The mass of one mole of a substance in grams is called its molar mass.

Molar Volume

The volume occupied by one mole of any substance is called its molar volume. It is denoted by Vm. One mole of all gaseous substances at 273 K and 1 atm pressure occupies a volume equal to 22.4 litre or 22,400 mL. The unit of molar volume is litre per mol or millilitre per mol

PERCENTAGE COMPOSITION—

The mass percentage of each constituent element present in any compound is called its percentage composition

Mass % of the element=<u>Mass of element in 1 molecule of the compound</u> x 100 Molecular mass of the compound

Empirical Formula and Molecular Formula—

An **empirical formula** represents the simplest whole number ratio of various atoms present in a compound. E.g. CH is the empirical formula of benzene.

The **molecular formula** shows the exact number of different types of atoms present in a molecule of a compound. E.g. C6H6 is the molecular formula of benzene.

Relationship between empirical and molecular formulae

The two formulas are related as Molecular formula = n x empirical formula

n=<u>Molecular mass</u>

empirical formula mass

Chemical Equation-

Shorthand representation of a chemical change in terms of symbols and formulae of the substances involved in the reaction is called chemical equation..

The substances that react among themselves to bring about the chemical changes are known as reactants, whereas the substances that are produced as a result of the chemical change, are known as products.

Limiting Reagent- The reactant which gets consumed first or limits the amount of product formed is known as **limiting reagent**

Reactions in Solutions-- The concentration of a solution can be expressed in any of the following ways.

1. Mass Percent is the mass of the solute in grams per 100 grams of the solution. Mass % of the solute = $\frac{Mass of the solute}{Mass of the solution} \times 100$

A 5 % solution of sodium chloride means that 5 g of NaCl is present in 100g of the solution.

2. Volume percent is the number of units of volume of the solute per 100 units of the volume of solution.

Volume % of the solute = $\frac{\text{Volume of the solute}}{\text{Volume of the solution}} \times 100$

A 5 % (v/v) solution of ethyl alcohol contains 5 cm^3 of alcohol in 100 cm^3 of the solution

3. Molarity of the solution is defined as the number of moles of solute dissolved per litre (dm³) of the solution. It isdenoted by the symbol M. Measurements in Molarity can change with the change in temperature because solutionsexpand or contract accordingly.

Molarity of the solution = $\underline{No. of moles of the solute}_{Volume of the solution in litre} = \underline{n}_{Volume of the solution in litre}$

The Molarity of the solution can also be expressed in terms of mass and molar mass

Molarity of the solution = <u>Mass of the solute</u> Molar mass of the solute X volume of the solution in liter

In terms of weight, molarity of the substance can be expressed as:

Molarity =
$$\frac{W_g}{M g \text{ mol}^{-1} \times V \text{ litre}} = \frac{W}{M \times V} \text{ mol/L}$$

Molarity equation

To calculate the volume of a definite solution required to prepare solution of other molarity, the following equation is used:

M1V1 = M2V2, where M1= initial molarity, M2= molarity of the new solution, V1= initial volume and V2= volume of the new solution.

4. Molality- Molality is defined as the number of moles of solute dissolved per 1000 g (1 kg) of solvent. Molality is expressed as 'm'.

Molality = <u>Moles of the solute</u> × 1000 Wt. of Solvent (in gm) 5. Mole Fraction is the ratio of number of moles of one component to the total number of moles (solute and solvents) present in the solution. It is expressed as 'x'.

Mole fraction of the solute = Moles of the solute Moles of solute + Moles of solvent

Mole fraction of the solvent = Moles of the solvent Moles of solute + Moles of solvent

Mole fraction of the solute + Mole fraction of solvent = 1 One Mark questions with answers

1. What is the significant figures in 1.050×10^4 ?

Ans. Four

2. What is the S.I. unit of Density?

Ans. Kg m⁻³

3. What do mean by Mole fraction?

Ans. Mole Fraction is the ratio of number of moles of one component to the total number of moles (solute and solvents) present in the solution. It is

expressed as 'x'.

4. Round off up to 3 significant figure (a) 1.235 (b) 1.225

Ans. (a) 1.24 (b) 1.22

5. What is AZT?

Ans. Azidothymidine.

6. What is limiting reagent?

Ans.The reactant which gets consumed first or limits the amount of product formed is known as **limiting reagent**

7. What is the relation between temperature in degree Celsius and degree fahrenheit?

Ans.

$$^{\circ}\mathsf{F} = \frac{9}{5}(^{\circ}\mathsf{C}) + 32$$

8. Define one mole?

Ans. One mole is the amount of a substance that contains as many particles as there are atoms in exactly 12 g of the carbon-12.

9. Calculate the formula mass calcium chloride.

Ans. Formula mass of $CaCl_2 = 40+2 \times 35.5 = 40+71 = 111 \text{ u}$

10. What is the law called which deals with the ratios of the volumes of the gaseous reactants and products?

Ans. Gay Lussac's law of gaseous volumes.

Two Marks questions with answers

1. Give the two points of differences between homogeneous and heterogeneous mixtures.

Ans.

Homogeneous mixture	Heterogeneous mixture
1. Homogeneous mixtures	1. Heterogeneous mixtures consist of
have the same composition	two or more parts (phases), which
throughout the sample.	have different compositions.
	2. These mixtures have visible
2. The components of such	boundaries of separation between the
mixtures cannot be seen	different constituents and can be seen
under a powerful	with the naked eye
microscope.	

2. Copper oxide obtained by heating copper carbonate or copper nitrate contains copper and oxygen in the same ration by mass. Which law is illustrated by this observation? State the law.

Ans.Law of Definite Proportions This law states that: A chemical compound always consists of the same elements combined together in the same ratio, irrespective of the method of preparation or the source from where it is taken.

3. Write the empirical formula of the following:

(a) N_2O_4 (b) $C_6H_{12}O_6$ (c) H_2O (d) H_2O_2 Ans. (a) NO_2 (b) CH_2O (c) H_2O (d) HO

4. Briefly explain the difference between precision and accuracy.

Ans. Precision refers to the closeness of various measurements for the same quantity. However, accuracy is the agreement of a particular value to the true value of the result.

5. Define the law of multiple proportions. Explain it with one example. Ans.When two elements combine to form two or more compounds, then the different masses of one element, which combine with a fixed mass of the other, bear a simple ratio to one another. For example- carbon combines with oxygen to form two compounds CO and CO_2

-	
CO	CO_2
12	12
16	32
	CO 12 16

Masses of oxygen which combine with a fixed mass of carbon (12g) bear a simple ratio of 16:32 or 1:2.

6. Chlorine has two isotopes of atomic mass units 34.97 and 36.97. The relative abundance of the isotopes is 0.755 and 0.245 respectively. Find the average atomic mass of chlorine.

Ans. Average atomic mass = 34.97 x 0.755 + 36.97 x 0.245 = 35.46 u

7. Calculate the percentage composition of water.

Ans. Mass % of an element = $\underline{\text{mass of that element in the compound} \times 100}$ molar mass of the compound

Molar mass of water = 18.02 g

Mass % of hydrogen = $\frac{2 \times 1.008 \times 100}{18.02}$

$$= 11.18$$

Mass % of oxygen = $\frac{16.00 \times 100}{18.02}$ = 88.79

8. State the number of significant figures in each of the following:

(i) 208.91 (ii) 0.00456 (iii) 453 (iv) 0.346

Ans.

- (i) 208.91 has five significant figures.
- (ii) 0.00456 has three significant figures.
- (iii) 453 has three significant figures.
- (iv) 0.346 has three significant figures.
- 8. Express the results of the following calculations to the appropriate number of significant figures.

(i)
$$\frac{3.24 \times 0.08666}{5.006}$$
 (ii) $\frac{(1.36 \times 10^{-4})(0.5)}{2.6}$

Ans.

(i)
$$\frac{3.24 \times 0.08666}{5.006} = 0.05608 = 0.0561$$

(ii) $\frac{(1.36 \times 10^{-4})(0.5)}{2.6}$

$$= 0.2615 \text{ x } 10-4 = 0.3 \text{ x } 10^{-4}$$

9. How are 0.50 mol Na2CO3 and 0.50 M Na2CO3 different?

Ans. Molar mass of Na2CO3= $2 \times 23 + 12 + 3 \times 16 = 106 \text{ g} / \text{mol}$

0.50 molNa2CO3means 0.50 x 106 = 53 g

0.50 M Na2CO3 means 0.50 mol i.e. 53 g of Na2CO3 are present in I L of the solution.

Three Marks questions with answers-

1. What is unit factor method? Express the following in SI units - 93 million miles (distance between earth and sun)

Ans. Method to convert units from one system to other is called unit factor method.

93 million miles = 93×10^6 miles

1 mile = 1.60934 km = 1.60934 x 10³ m
1 =
$$\frac{1.60934 \times 10^3 \text{ m}}{1 \text{ mile}}$$

93 million mile = 93 × 10⁶ mile × $\frac{1.60934 \times 10^3}{1 \text{ mile}}$

$$= 1.5 \text{ x } 10^{11} \text{ m}$$

2. Write the three points of difference between compound and mixture.

Compound	Mixture
Constituents are always present in a	Constituents may be present in any
fixed ratio by mass	ratio
May or may not be homogeneous in	Always homogeneous in nature
nature	
Constituents can be easily separated	Constituents cannot be easily

m

by simple mechanical means	separated by simple mechanical means
Properties are midway between those	Properties are entirely different from
of its constituents.	those of its constituents.

3. What do mean bygram atomic mass. One million silver atoms weigh 1.79 x 10^{16} g. Calculate the gram atomic mass of silver.

Ans.atomic mass of an element expressed in grams is the gram atomic mass Number of silver atoms = 1 million = 1×10^{6} Mass of one million Ag atoms = 1.79×10^{16} g

Mass of 6.023 x
$$10^{23}$$
 atoms of silver = $\frac{1.79 \times 10^{16} \text{ g}}{1 \times 10^{6}}$ x 6.023 x 10^{23}
= 107.8 g

4. What is the percentage of carbon, hydrogen and oxygen in ethanol?

Ans. Molecular formula of ethanol is : C_2H_5OH Molar mass of ethanol is : (212.01 + 61.008 + 16.00) g = 46.068 gMass per cent of carbon = $(24.02g / 46.068g) \times 100 = 52.14\%$ Mass per cent of hydrogen = $(6.048g / 46.068g) \times 100 = 13.13\%$ Mass per cent of oxygen = $(16.00 g / 46.068g) \times 100 = 34.73\%$

- 5. What do mean by molarity .Calculate the molarity of NaOH in the solution prepared by dissolving its 4 g in enough water to form 250 mL of the solution. Ans.The number of moles of solute dissolved per litre (dm³) of the solution is called molarity Since molarity (M) = No. of moles of solute /Volume of solution in litres =(Mass of NaOH/Molar mass of NaOH)/0.250 L =(4 g / 40 g 0.1 mol)/0.250L =0.1 mol/0.250 L = 0.4 mol L⁻¹
 - $= 0.4 \mod L$
 - = 0.4 M
- 6. Classify the following as pure substances or mixture-

(a) ethyl alcohol (b) oxygen (c) blood (d) carbon (e) steel (f) distilled water

Ans. Pure substance- ethyl alcohol ,oxygen , carbon, distilled water Mixture- blood, steel

7. What are the rules for rounding off?

Ans.1. If the digit coming after the desired number of significant figures happens to be more than 5, the precedingsignificant figure is increased by one,

2. If the digit involved is less than 5, it is neglected and the preceding significant figure remains unchanged,

3.If the digit happens to be 5, the last mentioned or preceding significant figure is increased by one only in case it happens to be odd. In case of even figure, the preceding digit remains unchanged.

- 8. Define –(a) Average atomic mass (b) Molecular mass (c) Formula mass Ans. (a) Average atomic mass- Atomic mass of an element is defined as the average relative mass of an atom of an element as compared to the mass of an atom of carbon -12 taken as 12.
 - (b) Molecular mass- it is sum of atomic masses of the elements present in a molecule.
 - (c) Formula mass- it is sum of atomic masses of the elements present in a formula unit of a compound.
- 9. Express the following in the scientific notation with 2 significant figures-(a) 0.0048 (b) 234,000 (c) 200.0

Ans. (a) 4.8×10^{-3} (b) 2.3×10^{5} (c) 2.0×10^{2} 10. Calculate the number of atoms in each of the following (i) 52 moles of Ar (ii) 52 u of He (iii) 52 g of He

Ans. (i) 1 mole of Ar = 6.022×10^{23} atoms of Ar \therefore 52 mol of Ar = $52 \times 6.022 \times 10^{23}$ atoms of Ar

 $= 3.131 \times 10^{25}$ atoms of Ar

(ii)1 atom of He = 4 u of He

Or.

4 u of He = 1 atom of He

1 u of He = 1/4 atom of He

52u of He = 52/4 atom of He

= 13 atoms of He

(iii) Molar mass of He = 4 g/mol4 g of He contains = 6.022×10^{23} atoms of He

52 g of He contains = $6.022 \times 10^{23} \text{ x } 52 = 78.286 \text{ x } 10^{23} \text{ atoms of He}$ Δ

Five Marks questions with answers-

1. What is the difference between empirical and molecular formula? A compound contains 4.07 % hydrogen, 24.27 % carbon and 71.65 % chlorine. Its molar mass is

98.96 g. What are its empirical and molecular formulas?

Ans. An empirical formula represents the simplest whole number ration of various atoms present in a compound whereas the molecular formula shows the exact number of different types of atoms present in a molecule of a compound.

Name of	Percentage	Step-1Conversion of	Step 2. number	Step 3. Divide the mole
element	of elements	mass per cent to	moles of each	value by the smallest
		grams.	element	number
С	24.27%	24.27g	24.27/12 = 2.0225	2.0225/2.018 = 1
H	4.07%	4.07g	4.07/1= 4.07	4.07/2.018=2
Cl	71.65%	71.65g	71.65/35.5 = 2.018	2.018/2.018 = 1

The empirical formula of the above compound is CH2Cl. empirical formula mass is 12 + (1x2) + 35.5 = 49.5

n= molecular mass/ empirical formula mass =98.96/49.5 = 2 Hence molecular formula is $C_2H_4Cl_2$

2. Dinitrogen and dihydrogen react with each other to produce ammonia according to the following chemical equation:

 $N_{2(g)} + H_{2(g)} \rightarrow 2NH_{3(g)}$

(i) Calculate the mass of ammonia produced if 2.00×10^3 g dinitrogen reacts with 1.00×10^3 g of dihydrogen.

(ii) Will any of the two reactants remain unreacted?

(iii) If yes, which one and what would be its mass?

Ans. (i) Balancing the given chemical equation, $N_{2(g)} + 3H_{2(g)} \longrightarrow 2NH_{3(g)}$

From the equation, 1 mole (28 g) of dinitrogen reacts with 3 mole (6 g) of dihydrogen to give 2 mole (34 g) of ammonia.

 $\Rightarrow 2.00 \times 10^{3} \text{ g of dinitrogen will react with } \frac{6 \text{ g}}{28 \text{ g}} \times 2.00 \times 10^{3} \text{ g} \text{ dihydrogen i.e.,}$

 2.00×10^3 g of dinitrogen will react with 428.6 g of dihydrogen.

Given,

Amount of dihydrogen = 1.00×10^3 g

Hence, N_2 is the limiting reagent.

 \therefore 28 g of N₂ produces 34 g of NH_{3.}

Hence, mass of ammonia produced by 2000 g of N₂ = $\frac{34g}{28 g} \times 2000 g$

= 2428.57 g

(ii) N_2 is the limiting reagent and H_2 is the excess reagent. Hence, H_2 will remain unreacted.

(iii) Mass of dihydrogen left unreacted = 1.00×10^3 g – 428.6 g

= 571.4 g

3. A welding fuel gas contains carbon and hydrogen only. Burning a small sample of it in oxygen gives 3.38 g carbon dioxide, 0.690 g of water and no other products. A volume of 10.0 L (measured at STP) of this welding gas is found to weigh 11.6 g. Calculate (i) empirical formula, (ii) molar mass of the gas, and (iii) molecular formula.

Ans. (i) 1 mole (44 g) of CO_2 contains 12 g of carbon.

```
=\frac{12 \text{ g}}{44 \text{ g}} \times 3.38 \text{ g}
        3.38 g of CO<sub>2</sub> will contain carbon
                                         = 0.9217 \text{ g}
        18 g of water contains 2 g of hydrogen.
                                                            =\frac{2 \text{ g}}{18 \text{ g}} \times 0.690
        0.690 g of water will contain hydrogen
                                                  = 0.0767 \text{ g}
        Since carbon and hydrogen are the only constituents of the compound,
the total mass of the compound is:= 0.9217 \text{ g} + 0.0767 \text{ g} = 0.9984 \text{ g}
                                                        =\frac{0.9217 \text{ g}}{0.9984 \text{ g}} \times 100
                Percent of C in the compound
                                                  = 92.32\%
                                                                 =\frac{0.0767 \text{ g}}{0.9984 \text{ g}} \times 100
                         Percent of H in the compound
                                                  = 7.68\%
                                                             = 92.32
                                                              12.00
                Moles of carbon in the compound
                                                          = 7.69
                                                                            7.68
                        Moles of hydrogen in the compound =
                                                                              1
                                                                  = 7.68
                Ratio of carbon to hydrogen in the compound = 7.69: 7.68= 1: 1
Hence, the empirical formula of the gas is CH.
(ii) Given,
Weight of 10.0L of the gas (at S.T.P) = 11.6 g
                                                 =\frac{11.6 \text{ g}}{10.0 \text{ L}} \times 22.4 \text{ L}
        Weight of 22.4 L of gas at STP
                = 25.984 g≈ 26 g
        Hence, the molar mass of the gas is 26 g.
```

(iii) Empirical formula mass of CH = 12 + 1 = 13 g

```
n = \frac{\text{Molar mass of gas}}{\text{Empirical formula mass of gas}}= \frac{26 \text{ g}}{13 \text{ g}}n = 2: \text{Molecular formula of gas} = (CH) = C
```

 \therefore Molecular formula of gas = (CH)_n= C₂H₂

HOTS (Higher Order Thinking Skills)

1. What is the difference between 160 cm and 160.0 cm

Ans. 160 has three significant figures while 160.0 has four significant figures. Hence, 160.0 represents greater accuracy.

2. In the combustion of methane, what is the limiting reactant and why?

Ans. Methane is the limiting reactant because the other reactant is oxygen of the air which is always present in excess. Thus, the amounts of CO_2 and H_2O formed depend upon the amount of methane burnt.

- 3. A compound made up of two elements A and B has A=70 %, B=30 %. Their relative number of moles in the compound are 1.25 and 1.88. calculate
 - a. Atomic masses of the elements A and B
 - b. Molecular formula of the compound , if its molecular mass is found to be 160

Ans. Relative no. of moles of an element = $\frac{\% \text{ of the element}}{100 \text{ moles}}$

```
Atomic mass
```

Or atomic mass = $\frac{\% \text{ of the element}}{\text{Relative no. of moles}}$ = $\frac{70}{1.25}$ = 56 Atomic mass of B = $\frac{30}{1.88}$ = 16 Calculation of Empirical formula

Element	Relative	no.	of	Simplest	molar	Simplest	whole	no.
	moles			ratio		molar ratio		
А	1.25			1.25/1.25	= 1	2		
В	1.88			1.88/1.25	= 1.5	3		

Empirical formula = A_2B_3

Calculation of molecular formula-

Empirical formula mass = $2 \times 56 + 3 \times 16 = 160$

n= molecular mass / Empirical formula mass = 160/160 = 1Molecular formula = A_2B_3

CHAPTER 2 STRUCTURE OF ATOM

• Atom is the smallest indivisible particle of the matter. Atom is made of electron, proton and neutrons.

PARTICLE	ELECTRON	PROTON	NEUTRON	
Discovery	Sir. J. J. Thomson	Goldstein (1886)	Chadwick (1932)	
	(1869)			
Nature of charge	Negative	Positive	Neutral	
Amount of charge	1.6 x 10 ⁻¹⁹ Coloumb	1.6 x 10 ⁻¹⁹ Coloumb	0	
Mass	9.11 x 10 ⁻³¹ kg	1.672614 x 10 ⁻²⁷ kg	$1.67492 \text{ x}10^{-27} \text{kg}$	

• Electrons were discovered using cathode ray discharge tube experiment.

• Nucleus was discovered by Rutherford in 1911.

• Cathode ray discharge tube experiment: A cathode ray discharge tube madeof glass is taken with two electrodes. At very low pressure and high voltage, current starts flowing through a stream of particles moving in the tube fromcathode to anode. These rays were called cathode rays. When a perforated anode was taken, the cathode rays struck the other end of the glass tube atthe fluorescent coating and a bright spot on the coating was developed

Results:

a. Cathode rays consist of negatively charged electrons.

b. Cathode rays themselves are not visible but their behavior can be observed with

help of fluorescent or phosphorescent materials.

c. In absence of electrical or magnetic field cathode rays travel in straight lines

d. In presence of electrical or magnetic field, behaviour of cathode rays is similar to that shown by electrons

e. The characteristics of the cathode rays do not depend upon the material of the electrodes and the nature of the gas present in the cathode ray tube.

- Charge to mass ratio of an electron was determined by Thomson. The chargeto mass ratio of an electron as 1.758820 x 10¹¹ C kg⁻¹
- Charge on an electron was determined by R A Millikan by using an oil dropexperiment. The value of the charge on an electron is -1.6×10^{-19} C.
- The mass on an electron was determined by combining the results of Thomson's experiment and Millikan's oil drop experiment. The mass of anelectron was determined to be 9.1094×10^{-31} kg.
- **Discovery of protons and canal rays:** Modified cathode ray tube experimentwas carried out which led to the discovery of protons.
- Characteristics of positively charged particles: a. Charge to mass ratio of particles depends on gas from which these originate

b. The positively charged particles depend upon the nature of gas present in the cathode ray discharge tube

c. Some of the positively charged particles carry a multiple of fundamental of electrical charge.

d. Behaviour of positively charged particles in electrical or magnetic field is opposite to that observed for cathode rays

- Neutrons were discovered by James Chadwick by bombarding a thin sheet of beryllium by α particles. They are electrically neutral particles having a mass slightly greater than that of the protons.
- Atomic number (Z) : the number of protons present in the nucleus (Moseley1913).
- Mass Number (A) :Sum of the number of protons and neutrons present in thenucleus.
- <u>Thomson model of an atom</u>: This model proposed that atom is considered asa uniform positively charged sphere and electrons are embedded in it. An important feature of Thomson model of an atom was that mass of atom isconsidered to be evenly spread over the atom. Thomson model of atom is also called as Plum pudding, raisin pudding orwatermelon modelThomson model of atom was discarded because it could not explain certainexperimental results like the scattering of α particles by thin metal foils.

Observations from α- particles scattering experiment by Rutherford: a. Most of the α- particles passed through gold foil un deflected b. A small fraction of α- particles got deflected through small angles

c. Very few α - particles did not pass through foil but suffered large deflection nearly180°

• <u>Conclusions Rutherford drew from *a*- particles scattering experiment</u>:

- a. Since most of the α -particles passed through foil undeflected, it means most of the space in atom is empty
- b. Since some of the α -particles are deflected to certain angles, it means that there is positively mass present in atom
- c. Since only some of the α -particles suffered large deflections, the positively charged mass must be occupying very small space
- d. Strong deflections or even bouncing back of α -particles from metal foil were due to direct collision with positively charged mass in atom
- <u>**Rutherford's model of atom</u>:** This model explained that atom consists ofnucleus which is concentrated in a very small volume. The nucleus comprises of protons and neutrons. The electrons revolve around the nucleus in fixedorbits. Electrons and nucleus are held together by electrostatic forces of attraction.</u>
- Drawbacks of Rutherford's model of atom:
 - a. According to Rutherford's model of atom, electrons which are negatively charged particles revolve around the nucleus in fixed orbits. Thus,

- b. theelectrons undergo acceleration. According to electromagnetic theory of Maxwell, a charged particle undergoing acceleration should emitelectromagnetic radiation. Thus, an electron in an orbit should emitradiation. Thus, the orbit should shrink. But this does not happen.
- c. The model does not give any information about how electrons are distributed around nucleus and what are energies of these electrons
- **Isotopes:** These are the atoms of the same element having the same atomic number but different mass number.e g $_1H^1$, $_1H^2$, $_1H^3$
- **Isobars:** Isobars are the atoms of different elements having the same massnumber but different atomic number.e g $_{18}Ar^{40}$, $_{20}Ca^{40}$
- **Isoelectronic species**: These are those species which have the same number of electrons.
- Electromagnetic radiations: The radiations which are associated withelectrical and magnetic fields are called electromagnetic radiations. When anelectrically charged particle moves under acceleration, alternating electricaland magnetic fields are produced and transmitted. These fields aretransmitted in the form of waves. These waves are called electromagneticwaves or electromagnetic radiations.

• Properties of electromagnetic radiations:

- a. Oscillating electric and magnetic field are produced by oscillating charged particles. These fields are perpendicular to each other and both are perpendicular to the direction of propagation of the wave.
- b. They do not need a medium to travel. That means they can even travel in vacuum.

• Characteristics of electromagnetic radiations:

- a. Wavelength: It may be defined as the distance between two neighbouring crests or troughs of wave as shown. It is denoted by λ .
- b. **Frequency** (v): It may be defined as the number of waves which passthrough a particular point in one second.
- c. Velocity (v): It is defined as the distance travelled by a wave in onesecond. In vacuum all types of electromagnetic radiations travel with thesame velocity. Its value is $3 \times 10^8 \text{m sec}^{-1}$. It is denoted by v

d. Wave number: Wave number (\bar{v}) is defined as the number of wavelengths per unit length.

• Velocity = frequency x wavelength $c = v\lambda$

• Planck's Quantum Theory-

- The radiant energy is emitted or absorbed not continuously but discontinuously in the form of small discrete packets of energy called 'quantum'. In case of light, the quantum of energy is called a 'photon'
- The energy of each quantum is directly proportional to the frequency of the radiation, i.e. $E \alpha v$ or E = hv where $h = Planck's constant = 6.626 x 10^{-27} Js$
- Energy is always emitted or absorbed as integral multiple of this quantum. E=nhv Where n=1,2,3,4,.....

- **Black body**: An ideal body, which emits and absorbs all frequencies, is calleda black body. The radiation emitted by such a body is called black bodyradiation.
- **Photoelectric effect**: The phenomenon of ejection of electrons from the surface of metal when light of suitable frequency strikes it is called photoelectric effect. The ejected electrons are called photoelectrons.
- Experimental results observed for the experiment of Photoelectric effect-
 - When beam of light falls on a metal surface electrons are ejectedimmediately.
 - Number of electrons ejected is proportional to intensity or brightness of light
 - Threshold frequency (vo): For each metal there is a characteristic minimum frequency below which photoelectric effect is not observed. This is called threshold frequency.
 - If frequency of light is less than the threshold frequency there is noejection of electrons no matter how long it falls on surface or how high isits intensity.
- Photoelectric work function (Wo): The minimum energy required to ejectelectrons is called photoelectric work function.Wo= hvo
- Energy of the ejected electrons : $h(v-v_0) = \frac{1}{2}m_ev^2$
- Dual behavior of electromagnetic radiation- The light possesses both particle and wave like properties, i.e., light has dual behavior . whenever radiation interacts with matter, it displays particle like properties.(Black body radiation and photoelectric effect) Wave like properties are exhibited when it propagates(interference an diffraction)
- When a white light is passed through a prism, it splits into a series of coloured bands known as spectrum.
- Spectrum is of two types: continuous and line spectrum
 - a. The spectrum which consists of all the wavelengths is called continuous spectrum.
 - b. A spectrum in which only specific wavelengths are present is known as a line
 - spectrum. It has bright lines with dark spaces between them.
- Electromagnetic spectrum is a continuous spectrum. It consists of a range ofelectromagnetic radiations arranged in the order of increasing wavelengths ordecreasing frequencies. It extends from radio waves to gamma rays.
- Spectrum is also classified as emission and line spectrum.
 - Emission spectrum: The spectrum of radiationemitted by a substance that has absorbed energy is called an emissionspectrum.
 - Absorption spectrum is the spectrum obtained when radiation is passed through a sample of material. The sample absorbs radiation of

certainwavelengths. The wavelengths which are absorbed are missing and comeas dark lines.

- The study of emission or absorption spectra is referred as spectroscopy.
- Spectral Lines for atomic hydrogen:

Series	n ₁	n 2	Spectral Region
Lyman	1	2, 3, 4, 5	Ultraviolet
Balmer	2	3, 4, 5	Visible
Paschen	3	4, 5	Infrared
Brackett	4	5, 6	Infrared
Pfund	5	6, 7	Infrared

• Rydberg equation

$$\overline{v} = 109,677 \left(\frac{1}{n_1^2} - \frac{1}{n_2^2} \right) \text{ cm}^{-1}$$

 $R = Rydberg's constant = 109677 cm^{-1}$

- Bohr's model for hydrogen atom:
 - a. An electron in the hydrogen atom can move around the nucleus in a circular path of fixed radius and energy. These paths are called orbits orenergy levels. These orbits are arranged concentrically around thenucleus.
 - b. As long as an electron remains in a particular orbit, it does not lose or gain energy and its energy remains constant.
 - c. When transition occurs between two stationary states that differ inenergy, the frequency of the radiation absorbed or emitted can becalculated

$$v = \frac{\Delta E}{h} = \frac{E_2 - E_1}{h}$$

d. An electron can move only in those orbits for which its angular momentum is an integral multiple of $h/2\pi$

$$m_{e} vr = n. \frac{h}{2\pi}$$
 $n = 1, 2, 3....$

- The radius of the *n*th orbit is given by $r_n = 52.9 \text{ pm x} \frac{n^2}{7}$
- energy of electron in *n*th orbit is :

$$E_n = -2.18 \times 10^{-18} \left(\frac{Z^2}{n^2} \right) J$$

- Limitations of Bohr's model of atom:
 - a. Bohr's model failed to account for the finer details of the hydrogen spectrum.
 - b. Bohr's model was also unable to explain spectrum of atoms containing more than one electron.
- **Dual behavior of matter:** de Broglie proposed that matter exhibits dualbehavior i.e. matter shows both particle and wave nature. de Broglie's relation is

$$\lambda = \frac{h}{mv} = \frac{h}{p}$$

• Heisenberg's uncertainty principle: It states that it is impossible to determine simultaneously, the exact position and exact momentum (or velocity) of an electron. The product of their uncertainties is always equal to or greater than $h/4\pi$.

Mathematically
$$\Delta x \times \Delta p \ge \frac{h}{4\pi}$$

where $\Delta x =$ uncertainty in position,
 $\Delta p =$ uncertainty in momentum

- Heisenberg's uncertainty principle rules out the existence of definite pathsor trajectories of electrons and other similar particles
- Failure of Bohr's model:
 - a. It ignores the dual behavior of matter.
 - b. It contradicts Heisenberg's uncertainty principle.
- Classical mechanics is based on Newton's laws of motion. It successfully describes the motion of macroscopic particles but fails in the case of microscopic particles.

Reason: Classical mechanics ignores the concept of dual behaviour of matter especially for sub-atomic particles and the Heisenberg's uncertainty principle.

- **Quantum mechanics** is a theoretical science that deals with the study of themotions of the microscopic objects that have both observable wave like andparticle like properties.
- Quantum mechanics is based on a fundamental equation which is calledSchrodinger equation.
- Schrodinger's equation: For a system (such as an atom or a molecule whoseenergy does not change with time) the Schrödinger equation is written as:

$$\hat{H}\Psi = E\Psi$$

 \hat{H} is the Hamiltonian operator E is the total energy of the system Ψ represents the wave function which is the amplitude of the electron Wave

- When Schrödinger equation is solved for hydrogen atom, the solution gives the possible energy levels the electron can occupy and the corresponding wave function(s) of the electron associated with each energy level. Out of the possible values, only certain solutions are permitted. Each permitted solution is highly significant as it corresponds to a definite energy state. Thus, we can say that energy is quantized.
- ψ gives us the amplitude of wave. The value of ψ has no physical significance.
- Ψ^2 gives us the region in which the probability of finding an electron ismaximum. It is called probability density.

- Orbital: The region of space around the nucleus where the probability offinding an electron is maximum is called an orbital.
- Quantum numbers: There are a set of four quantum numbers which specifythe energy, size, shape and orientation of an orbital. To specify an orbital only three quantum numbers are required while to specify an electron all four quantum numbers are required.
- Principal quantum number (n): It identifies shell, determines sizes and

N	1	2	3	4
Shell no.:	К	L	М	Ν
Total number of orbitals in a shell = n ²	1	4	9	16
Maximum number of electrons = 2n ²	2	8	18	32

energy of orbitals

• Azimuthal quantum number (l): Azimuthal quantum number. 'l' is also known as orbital angular momentum or subsidiary quantum number. l. It identifies sub-shell, determines the shape of orbitals, energy of orbitals in multi-electron atoms along with principal quantum number and orbital angular

momentum, *i.e.*, $\sqrt{l(l+1)} \frac{h}{2\pi}$ The number of orbitals in a subshell = 2l + 1. For a given value of *n*, it can have *n* values ranging from 0 to n-1. Total number of subshells in a particular shell is equal to the value of n.

Subshell	S	р	d	f	g
notation					
Value of ' <i>l</i> '	0	1	2	3	4
Number of	1	3	5	7	9
orbitals					

- Magnetic quantum number or Magnetic orbital quantum number (ml): Itgives information about the spatial orientation of the orbital with respect tostandard set of co-ordinate axis.For any sub-shell (defined by 'l' value) 2l+1 values of ml are possible.For each value of l, $m_l = -1, -(1-1), -(1-2)...0, 1...$ (1-2), (l-1), l
- Electron spin quantum number (ms): It refers to orientation of the spin of theelectron. It can have two values +1/2 and -1/2. +1/2 identifies the clockwisespin and -1/2 identifies the anti- clockwise spin.
- The region where this probability density function reduces to zero is callednodal surfaces or simply nodes.
- Radial nodes: Radial nodes occur when the probability density of wave function for the electron is zero on a spherical surface of a particular radius. Number of radial nodes = n 1 1
- Angular nodes: Angular nodes occur when the probability density wavefunction for the electron is zero along the directions specified by a particularangle. Number of angular nodes = 1
- Total number of nodes = n 1

- Degenerate orbitals: Orbitals having the same energy are called degenerateorbitals.
- Shape of p and d-orbitals



• Shielding effect or screening effect: Due to the presence of electrons in theinner shells, the electron in the outer shell will not experience the full positivecharge on the nucleus.

So, due to the screening effect, the net positive charge experienced by theelectron from the nucleus is lowered and is known as <u>effective nuclear</u> <u>charge</u>. Effective nuclear charge experienced by the orbital decreases with increase of azimuthal quantum number (1).

• Aufbau Principle: In the ground state of the atoms, the orbitals are filled inorder of their increasing energies

- n+l rule-Orbitals with lower value of (n+l) have lower energy. If two orbitals have the same value of (n+l) then orbital with lower value of nwill have lower energy.
- The order in which the orbitals are filled isas follows:
 - 1s, 2s, 2p, 3s, 3p, 4s, 3d, 4p, 5s, 4d, 5p, 4f, 5d, 6p, 7s...
- **Pauli Exclusion Principle**: No two electrons in an atom can have the same setof four quantum numbers. Only two electrons may exist in the same orbitaland these electrons must have opposite spin.
- Hund's rule of maximum multiplicity: Pairing of electrons in the orbitalsbelonging to the same subshell (p, d or f) does not take place until eachorbital belonging to that subshell has got one electron each i.e., it is singlyoccupied.
- Electronic configuration of atoms: Arrangement of electrons in different orbitals of an atom. The electronic configuration of differentatoms can be represented in two ways.
 - a. $s^a p^b d^c$ notation.

b. Orbital diagram:, each orbital of the subshell is represented by a box and the electron is represented by an arrow (\uparrow) a positive spin or an arrow (\downarrow) a negative spin.

• Stability of completely filled and half filled subshells:

a. Symmetrical distribution of electrons- the completely filled or half filled sub-shells have

symmetrical distribution of electrons in them and are more stable.

b. Exchange energy-The two or more electrons with the same spin present in the degenerate orbitals of a sub-shell can exchange their position and the energy released due to this exchange is called exchange energy. The number of exchanges is maximum when the subshell is either half filled or completely filled. As a result the exchange energy is maximum and so is the stability.

ONE MARK QUESTIONS

1. Neutrons can be found in all atomic nuclei except in one case. Which is this atomic nucleus and what does it consists of?

Ans. Hydrogen atom. It consists of only one proton.

2. Calculate wave number of yellow radiations having wavelength of 5800 A^0 . Ans. Wave number = 1/ wavelength

Wavelength = $5800 \text{ A}^{0} = 5800 \text{ x} 10^{-10} \text{ m}$

Wave number = $1/5800 \times 10^{-10} \text{ m} = 1.72 \times 10^{6} \text{ m}^{-1}$

3. What are the values of n and 1 for 2p orbital?

Ans. n=2 and l=1

4. Which of the following orbitals are not possible? 1p, 2s, 3f and 4d Ans. 1p and 3f are not possible.

5. Write the electronic configuration of the element having atomic number 24. Ans. $1s^2 2s^2 2p^6 3s^2 3p^6 3d^5 4s^1$

6. What atoms are indicated by the following electronic configurations?

a. $1s^2 2s^2 2p^1$

b.
$$[Ar]4s^2 3d^1$$

Ans. a. Boron b. Scandium

7. What is the relationship between frequency and wavelength of light? Ans. velocity of light = frequency x wavelength. Frequency and wavelength are inversely proportional to each other.

8. State Pauli Exclusion Principle.

Ans. No two electrons in an atom can have the same set of four quantum numbers or an orbital can have maximum two electrons and these must have opposite spin.

9. When α - rays hit a thin foil of gold, very few α - particles is deflected back. What does it prove?

Ans. There is a very small heavy body present within the atom.

10. What is the difference between a quantum and a photon?

Ans. The smallest packet of energy of any radiation is called a quantum whereas that of light is called photon.

TWO MARKS QUESTIONS

- 1. Write the complete symbol for the atom with the given atomic number (Z) and mass number(A). (a) Z = 17, A = 35 (b) Z = 92, A = 233Ans. (a) ${}^{35}_{17}Cl$ (b) ${}^{233}_{92}U$
- 2. Using s,p,d and f notation, describe the orbital with the following quantum numbers-

(a) n=1,l=0 (b) n=3, l=1 (c) n=4, l=2 (d) n=4, l=3

Ans. (a) 1s (b) 3p (c)4d (d) 4f

3. How many electrons in an atom have the following quantum numbers?

a. n=4, $m_s=-1/2$ b. n=3, l=0

Ans. (a) 16 electrons (b) 2 electrons.

4. An element with mass number 81 contains 31.7 % more neutrons as compared to protons. Assign the atomic symbol.

Ans. Mass number = 81, i.e., p + n = 81

If protons = x, then neutrons = $x + \frac{31.7}{100}$ X x = 1.317 x

- x+1.317x = 81 or 2.317x = 81 x=35 Thus proton = 35, i.e., atomic no. = 35 Hence symbol is ${}^{81}_{35}$ Br
- 5. (i) The energy associated with the first orbit in the hydrogen atom is -2.18 x 10^{-18} J/atom. What is the energy associated with the fifth orbit
 - (ii) Calculate the radius of Bohr's fifth orbit for hydrogen atom.

Ans. (i) $E_n = -2.18 \times 10^{-18} / n^2$ (ii) For H atom, $r_n = 0.529 \times n^2$ $E_5 = -2.18 \times 10^{-18} / 5^2 = -8.72 \times 10^{-20} J$ $r_5 = 0.529 \times 5^2 = 13.225 A^0 = 1.3225 nm$

6. Explain, giving reasons, which of the following sets of quantum numbers are not possible.

(a) n=0, l=0; $m_l = 0$, $m_s = +\frac{1}{2}$ (c) n=1, l=0; $m_l = 0$, $m_s = -\frac{1}{2}$

(b) n=1, l=1; m_l =- 0, m_s= + $\frac{1}{2}$ (d) n=2, l=1; m_l = 0, m_s= + $\frac{1}{2}$

Ans. (a) Not possible because $n \neq 0$ (c) Not possible because when $n=1, l \neq 1$

(b) Possible

(d) Possible

- 7. (a)What is the lowest value of n that allows g orbitals to exist?
 (b)An electron is in one of the 3d orbitals, Give the possible values of n,l and m_l for this electron.
- Ans.(a) minimum value of n=5

 $(b)n=3, l=2, m_l = -2, -1, 0, +1, +2$

- 8. Calculate the total number of angular nodes and radial nodes present in 30 orbitals.
- Ans. For 3p orbitals, n=3, l=1

Number of angular nodes = l = 1

Number of radial nodes = n-l-1 = 3-1-1=1

9. Mention the draw backs of Rutherford's atomic model.

Ans. 1. It could not explain the stability of an atom.

2. It could not explain the line spectrum of H- atom.

10. State de-Broglie concept of dual nature of matter. How do dual nature of electron verified?

Ans. Just as light has dual nature, every material particle in motion has dual nature (particle nature and wave nature). The wave nature has been verified by Davisson and Germer's experiment whereas particle nature by scintillation experiment.

THREE MARKS QUESTIONS

1. State (a)Hund's Rule of maximum Multiplicity (b) Aufbau Principle (c) n+l rule

Ans.(a) Pairing of electrons in the orbitals belonging to the same subshell (p, d or f) does not takeplace until each orbital belonging to that subshell has got one electron each i.e., it is singly occupied.

(b)In the ground state of the atoms, the orbitals are filled inorder of their increasing energies

(c) Orbitals with lower value of (n+1) have lower energy. If two orbitals have the same value of (n+1) then orbital with lower value of n will have lower energy.

2. Write down the quantum numbers n and 1 for the following orbitals

a. 2p b. 3d c. 5f

3. Write the 3 points of difference between orbit and orbital.

Ans.

Ans.

Orbit		orbita	1
1.	An orbit is a well defined	1.	An orbital is the three dimensional
	circular path around the		space around the nucleus within
	nucleus in which the		which the probability of finding an
	electrons revolve		electron is maximum(upto 90 %)
2.	It represents the planar	2.	It represents the three dimensional
	motion of an electron around		motion of an electron around the
	the nucleus		nucleus
3.	All orbits are circular and	3.	Different orbitals have different
	disc like		shapes, i.e., s-orbitals are spherically
			symmetrical, p-orbitals are dumb-bell
			shaped and so on.

4. State Heisenberg's uncertainty principle.calculate the uncertainty in the position of an electron if the uncertainty in its velocity is 5.7×10^5 m/s.

Ans. It states that it is impossible to determine simultaneously, the exact position and exact momentum (or velocity) of an electron. The product of their uncertainties is always equal to or greater than $h/4\pi$.

$$\Delta x \quad x (m \ x \ \Delta v) = h/4 \square$$

$$\Delta x = h/4 \square \ x \ m \ x \ \Delta v = \frac{6.6 \ x \ 10^{-34}}{4 \ x \ 3.14 \ x \ 9.1 \ x \ 10^{-31} \ x \ 5.7 \ x \ 10^5} = 1.0 \ x \ 10^{-10} \ m$$

5. Write 3 points of differences between electromagnetic waves and matterwaves.

Electromagnetic waves	Matter waves
1. These are associated with	1. These are not associated with
electric and magnetic	electric and magnetic field.
fields	
2. They do not require any	2. They require medium for
medium for propagation.	propagation
3. They travel with the same	3. They travel with lower speeds
speed a that of light	not constanr for all matter
	waves

6. (i) Calculate the number of electrons which will together weigh one gram.(ii) Calculate the mass and charge of one mole of electrons

Ans. (i) Mass of one electron = 9.10939×10^{-31} kg

 \therefore Number of electrons that weigh 9.10939 \times 10⁻³¹ kg = 1

Number of electrons that will weigh 1 $g = (1 \times 10^{-3} \text{kg})$

$$=\frac{1}{9.10939\times10^{-31} \text{ kg}}\times(1\times10^{-3} \text{ kg})$$

 $= 0.1098 \times 10^{-3+31}$

 $= 0.1098 \times 10^{28}$

- $= 1.098 \times 10^{27}$
- (ii) Mass of one electron = 9.10939×10^{-31} kg

Mass of one mole of electron = $(6.022 \times 10^{23}) \times (9.10939 \times 10^{-31} \text{ kg})$

 $= 5.48 \times 10^{-7} \text{ kg}$

Charge on one electron = 1.6022×10^{-19} coulomb

Charge on one mole of electron = $(1.6022 \times 10^{-19} \text{C}) (6.022 \times 10^{23})$

 $= 9.65 \times 10^4 \,\mathrm{C}$

7. Find energy of each of the photons which
(i) correspond to light of frequency 3× 10¹⁵Hz.
(ii) have wavelength of 0.50 Å.

Ans.(i) Energy (E) of a photon is given by the expression,

E = hvWhere, h = Planck's constant = 6.626×10^{-34} Js v = frequency of light = 3×10^{15} Hz Substituting the values in the given expression of *E*: $E = (6.626 \times 10^{-34}) (3 \times 10^{15})$ $E = 1.988 \times 10^{-18}$ J

(ii) Energy (*E*) of a photon having wavelength (λ) is given by the expression,

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

h = Planck's constant = 6.626×10^{-34} Js c = velocity of light in vacuum = 3×10^8 m/s Substituting the values in the given expression of *E*:

$$E = \frac{\left(6.626 \times 10^{-34}\right) \left(3 \times 10^{8}\right)}{0.50 \times 10^{-10}} = 3.976 \times 10^{-15} \text{ J}$$

$$\therefore E = 3.98 \times 10^{-15} \text{ J}$$

8. What is the wavelength of light emitted when the electron in a hydrogen atom undergoes transition from an energy level with n = 4 to an energy level with n = 2?

Ans.The n_i = 4 to n_f = 2 transition will give rise to a spectral line of the Balmer series. The energy involved in the transition is given by the relation,

$$E = 2.18 \times 10^{-18} \left[\frac{1}{n_i^2} - \frac{1}{n_f^2} \right]$$

Substituting the values in the given expression of E:

$$E = 2.18 \times 10^{-18} \left[\frac{1}{4^2} - \frac{1}{2^2} \right]$$
$$= 2.18 \times 10^{-18} \left[\frac{1-4}{16} \right]$$
$$= 2.18 \times 10^{-18} \times \left(-\frac{3}{16} \right)$$

 $E = -(4.0875 \times 10^{-19} \text{ J})$

The negative sign indicates the energy of emission.

Wavelength of light emitted $(\lambda) = \frac{hc}{E}$

 $\left(\text{since } E = \frac{\text{hc}}{\lambda} \right)$

Substituting the values in the given expression of λ :

$$\lambda = \frac{\left(6.626 \times 10^{-34}\right) \left(3 \times 10^{8}\right)}{4.0875 \times 10^{-19}}$$
$$\lambda = 4.8631 \times 10^{-7} \text{ m}$$
$$= 486.3 \times 10^{-9} \text{ m}$$
$$= 486 \text{ nm}$$

9. An atom of an element contains 29 electrons and 35 neutrons. Deduce (i)the number of protons and (ii) the electronic configuration of the element (iii) Identify the element.

Ans.(i)For an atom to be neutral, the number of protons is equal to the number of electrons.

: Number of protons in the atom of the given element = 29 (ii) The electronic configuration of the atom is $1s^22s^2 2p^6 3s^2 3p^64s^2 3d^{10}$ (iii) Copper

10. Give the number of electrons in the species H_2^+ , H_2 and O_2^+

Ans. Number of electrons present in hydrogen molecule $(H_2) = 1 + 1 = 2$

: Number of electrons in $H_2^+ = 2 - 1 = 1$

Number of electrons in $H_2 = 1 + 1 = 2$

Number of electrons present in oxygen molecule $(O_2) = 8 + 8 = 16$

 \therefore Number of electrons in $O_2^+ = 16 - 1 = 15$

FIVE MARKS QUESTIONS WITH ANSWERS

1. What are the draw backs of Bohr's atomic model? Show that the circumference of the Bohr orbit for the hydrogen atom is an integral multiple of the de Broglie wavelength associated with the electron revolving around the orbit.

Ans.1.Bohr's model failed to account for the finer details of the hydrogen spectrum.

- 2. Bohr's model was also unable to explain spectrum of atoms containing more than one electron.
- 3. Bohr's model was unable to explain Zeeman effect and Stark effect i
- 4. Bohr's model could not explain the ability of atoms to form molecules by chemical bonds

Since a hydrogen atom has only one electron, according to Bohr's postulate, the angular momentum of that electron is given by:

$$mvr = n \frac{h}{2\pi}$$
.....(1)
Where, $n = 1, 2, 3, ...$
According to de Broglie's equation:
 $\lambda = \frac{h}{mv}$

or
$$mv = \frac{h}{\lambda}$$
.....(2)

Substituting the value of 'mv' from expression (2) in expression (1):

$$\frac{\mathrm{h}r}{\lambda} = n\frac{\mathrm{h}}{2\pi}$$

or $2\pi r = n\lambda$(3)

Since $2\pi r$ represents the circumference of the Bohr orbit (*r*), it is proved by equation (3) that the circumference of the Bohr orbit of the hydrogen atom is an integral multiple of de Broglie's wavelength associated with the electron revolving around the orbit.

2. State photo electric effect. The work function for caesium atom is 1.9 eV. Calculate (a) the threshold wavelength and (b) the threshold frequency of the radiation. If the caesium element is irradiated with awavelength 500 nm, calculate the kinetic energy and the velocity of the ejected photoelectron. Ans. Photoelectric effect: The phenomenon of ejection of electrons from the surface of metal when light of suitable frequency strikes it is called photoelectric effect. The ejected electrons are called photoelectrons. It is given that the work function (W_0) for caesium atom is 1.9 eV.

(a) From the expression,
$$W_0 = \frac{hc}{\lambda_0}$$
, we get:
 $\lambda_0 = \frac{hc}{W_0}$

Where,

 λ_0 = threshold wavelength h = Planck's constant c = velocity of radiation Substituting the values in the given expression of (λ_0):

$$\lambda_0 = \frac{\left(6.626 \times 10^{-34} \text{ Js}\right) \left(3.0 \times 10^8 \text{ ms}^{-1}\right)}{1.9 \times 1.602 \times 10^{-19} \text{ J}}$$
$$\lambda_0 = 6.53 \times 10^{-7} \text{ m}$$

Hence, the threshold wavelength λ_0 is 653 nm.

(b) From the expression, $W_0 = h v_0$, we get:

$$v_0 = \frac{W_0}{h}$$
Where,
 v_0 = threshold frequency
 h = Planck's constant
Substituting the values in the given expression of v_0 :
 $v_0 = \frac{1.9 \times 1.602 \times 10^{-19} \text{ J}}{6.626 \times 10^{-34} \text{ Js}}$
 $(1 \text{ eV} = 1.602 \times 10^{-19} \text{ J})$
 $v_0 = 4.593 \times 10^{14} \text{ s}^{-1}$
Hence, the threshold frequency of radiation (v_0) is $4.593 \times 10^{14} \text{ s}^{-1}$.
(c) According to the question:

Wavelength used in irradiation (λ) = 500 nm

Kinetic energy = $h(v - v_0)$

$$= hc \left(\frac{1}{\lambda} - \frac{1}{\lambda_0}\right)$$

= $(6.626 \times 10^{-34} \text{ Js})(3.0 \times 10^8 \text{ ms}^{-1}) \left(\frac{\lambda_0 - \lambda}{\lambda \lambda_0}\right)$
= $(1.9878 \times 2^{-26} \text{ Jm}) \left[\frac{(653 - 500)10^{-9} \text{ m}}{(653)(500)10^{-18} \text{ m}^2}\right]$
= $\frac{(1.9878 \times 10^{-26})(153 \times 10^9)}{(653)(500)}$ J
= 9.3149×10^{-20} J

Kinetic energy of the ejected photoelectron = 9.3149×10^{-20} J

Since K.E

$$= \frac{1}{2}mv^{2} = 9.3149 \times 10^{-20} \text{ J}$$

$$v = \sqrt{\frac{2(9.3149 \times 10^{-20} \text{ J})}{9.10939 \times 10^{-31} \text{ kg}}}$$

$$= \sqrt{2.0451 \times 10^{11} \text{ m}^{2} \text{s}^{-2}}$$

$$v = 4.52 \times 10^{5} \text{ms}^{-1}$$

Hence, the velocity of the ejected photoelectron (v) is $4.52 \times 10^5 \text{ms}^{-1}$. 3. (a)The quantum numbers of six electrons are given below. Arrange them in order of increasing energies. If any of these combination(s) has/have the same energy lists: 1. n=4, l=2, $m_l=-2$, $m_s=-1/2$ 2. n=3, l=2, $m_l=1$, $m_s=+1/2$ 3. n=4, l=1, $m_l=0$, $m_s=+1/2$ 4. n=3, l=2, $m_l=-2$, $m_s=-1/2$ 5. n=3, l=1, $m_l=-1$, $m_s=+1/2$

6. n = 4, l = 1, $m_l = 0$, $m_s = +1/2$

(b)Among the following pairs of orbitals which orbital will experience the larger effective nuclearcharge? (i) 2s and 3s, (ii) 4d and 4f, (iii) 3d and 3p

Ans.(a)For n = 4 and l = 2, the orbital occupied is 4d.

For n = 3 and l = 2, the orbital occupied is 3*d*.

For n = 4 and l = 1, the orbital occupied is 4p.

Hence, the six electrons i.e., 1, 2, 3, 4, 5, and 6 are present in the 4d, 3d, 4p, 3d, 3p, and 4p orbitals respectively.

Therefore, the increasing order of energies is 5(3p) < 2(3d) = 4(3d) < 3(4p) = 6(4p) < 1 (4*d*).

(b)Nuclear charge is defined as the net positive charge experienced by an electron in the orbital of a multi-electron atom. The closer the orbital, the greater is the nuclear charge experienced by the electron (s) in it.

(i) The electron(s) present in the 2s orbital will experience greater nuclear charge (being closer to the nucleus) than the electron(s) in the 3s orbital. (ii) 4d will experience greater nuclear charge than 4fsince 4d is closer to the nucleus.

(iii) 3*p* will experience greater nuclear charge since it is closer to the nucleus than 3*f*.

- 4. (i) The unpaired electrons in Al and Si are present in 3p orbital. Which electrons will experience more effective nuclear charge from the nucleus? (ii) Indicate the number of unpaired electrons in: (a) P, (b) Si, (c) Cr, (d) Fe Ans. (i) the electrons in the 3p orbital of silicon will experience a more effective nuclear charge than aluminium.
 - (ii) (a) Phosphorus (P):

Atomic number = 15 The electronic configuration of P is: $1s^2 2s^2 2p^6 3s^2 3p^3$ The orbital picture of P can be represented as:

1s 2s 2p 3s 3p

From the orbital picture, phosphorus has **three** unpaired electrons. **(b)** Silicon (Si):

Atomic number = 14

The electronic configuration of Si is: $1s^2 2s^2 2p^6 3s^2 3p^2$

The orbital picture of Si can be represented as:

1s 2s 2p 3s 3p

From the orbital picture, silicon has **two** unpaired electrons.

(c) Chromium (Cr):

Atomic number = 24

The electronic configuration of Cr is: $1s^2 2s^2 2p^6 3s^2 3p^6 4s^1 3d^5$ The orbital picture of chromium is: 1s 2s 2p 3s 3p 4s 3d

From the orbital picture, chromium has six unpaired electrons.

(d) Iron (Fe):

Atomic number = 26 The electronic configuration is: $1s^2 2s^2 2p^6 3s^2 3p^6 4s^2 3d^6$ The orbital picture of chromium is: $\underbrace{\texttt{ft}}_{1s} \underbrace{\texttt{ft}}_{2s} \underbrace{\texttt{ft}}_{2p} \underbrace{\texttt{ft}}_{3s} \underbrace{\texttt{ft}}_{3p} \underbrace{\texttt{ft}}_{4s} \underbrace{\texttt{ft}}_{3d} \underbrace{\texttt{ft}}_{3d} \underbrace{\texttt{ft}}_{3d}$ From the orbital picture, iron has **four** unpaired electrons.

HOTS QUESTIONS WITH ANSWERS

- 1. Give the name and atomic number of the inert gas atom in which the total number of d-electrons is equal to the difference between the numbers of total p and total s electrons.
- Ans. electronic configuration of Kr (atomic no.=36) = $1s^2 2s^2 2p^6 3s^2 3p^6 3d^{10} 4s^2 4p^6$ Total no. of s-electrons = 8, total no. of p-electrons = 18. Difference = 10 No. of d- electrons = 10
- 2. What is the minimum product of uncertainty in position and momentum of an electron?

Ans.h/ 4π

3. Which orbital is non-directional ?

Ans. s- orbital

4. What is the difference between the notations *l* and L ?

Ans. l represents the sub-shell and L represent shell.

5. How many electrons in an atom can have n + l = 6? Ans. 18

6. An anion $A^{3+}_{.}$ has 18 electrons. Write the atomic number of A.

Ans.15

7. Arrange the electron (*e*), protons (*p*) and alpha particle (α) in the increasing order for the values of *e/m* (charge/mass).

Ans.. α

CHAPTER-3

CLASSIFICATION OF ELEMENTS AND PERIODICITY IN PROPERTIES OF ELEMENTS

Mandeleev's Periodic Law:- The properties of the elements are the periodic function of their atomic masses.

Moseley, the English physicist showed that atomic number is more fundamental property of an element than its atomic mass. Therefore, the position of an element in the periodic table depends on its atomic number than its atomic mass.

Modern Periodic Law: The physical and chemical properties of elements are the periodic functions of their atomic numbers.

Types of Elements: s-, p-, d- and f- blocks.

MAIN GROUP ELEMENTS/ REPRESENTATIVE ELEMENTS:

The s- and p- block elements are called main group elements or representative elements.

s- block elements: Group-1 (Alkali metals) and Group-2 elements (Alkaline earth metals) which respectively have ns¹ and ns² outermost electronic configurations.

p-Block elements: They belongs to group- 13 to 18. The outer most electronic configuration is $ns^2 np^{1-6}$. He $(1s^2)$ is a s- block element but is positioned with the group 18 elements $(ns^2 np^6)$ because it has completely filled valence shell and as a result, exhibits properties characteristic of other noble gases.

d- block elements (Transition elements) are the elements of group 3 to 12 having outer electronic configuration $(n-1) d^{1-10} ns^{1-2}$. Four transition series are 3d, 4d, 5d and 6d. The 6d- series is incomplete. Atomic radius generally decreases across a period and increases as we descend the group.

f-Block elements (Inner- transition Series)

Lanthanoids charecterised by the filling of 4 f-orbitals, are the elements following lanthanum from ${}_{58}$ Ce to ${}_{71}$ Lu. Actinoids characterised by filling of 5f-orbitals, are the elements following actinium from ${}_{70}$ Th to ${}_{103}$ Lr. Characteristic outer electronic configuration is (n-2) f¹⁻¹⁴ (n-1) d⁰⁻¹ ns².

Noble Gases: The gaseous elements of group 18 are called noble gases. The general outermost electronic configuration of noble gases (except He) is $ns^2 np^6$. He exceptionally has $1s^2$ configuration. Thus the outermost shell of noble gases is completely filled.

PERIODICITY: The repetition of similar properties after regular intervals is called periodicity.

Cause of Periodicity: The properties of elements are the periodic repetition of similar electronic configuration of elements as the atomic number increases.

ATOMIC PROPERTIES: The physical characteristics of the atom of an element are called atomic properties. The properties such as atomic radius, ionic radius, ionisation energy, electro-negativity, electron affinity and valence etc., called atomic properties.

ATOMIC RADIUS- The distance from the centre of the nucleus to the outermost shell of the electrons in the atom of any element is called its atomic radius.

Periodicity- (a) In period- Atomic radius of elements decreases from left to right in a period.

(b) In Group- Atomic radius of elements increases on moving top to bottom in a group.

COVALENT RADIUS- Half the inter-nuclear distance between two similar atoms of any element which are covalently bonded to each other by a single covalent bond is called covalent radius.

VAN DER WAALS' RADIUS: Half the inter-nuclear separation between two similar adjacent atoms belonging to the two neighbouring molecules of the same substance in the solid state is called the van der waals' radius of that atom.

METALLIC RADIUS: Half the distance between the nuclei of the two adjacent metal atoms in a close packed lattice of the metal is called its metallic radius.

Van der Waals' radius > Metallic radius > Covalent radius

IONIC RADIUS: The effective distance from the centre of the nucleus of an ion upto which it has an influence on its electron cloud is called its ionic radius.
A cation is smaller but the anion is larger than the parent atom. In case of isoelectronic species, the cation with greater positive charge has smaller radius but anion with greater negative charge has the larger radii.

IONISATION ENTHALPY: The ionisation enthalpy is the molar enthalpy change accompanying the removal of an electron from a gaseous phase atom or ion in its ground state. Thus enthalpy change for the reaction; $M_{(g)} \rightarrow M^+_{(g)} + e^-$

Is the ionisation enthalpy of the element M. Like ionisation energies for successive ionisation, the successive ionisation enthalpy may also be termed as 2^{nd} ionisation enthalpy ($\Delta_r H_2$), third ionisation enthalpy ($\Delta_r H_3$) etc. The term ionisation enthalpy is taken for the first ionisation enthalpy, ($\Delta_r H_1$) is expressed in kg mol⁻ or in eV.

Periodicity:

i) Generally the ionisation enthalpies follow the order (there are few exceptions):

 $(\Delta_r H_1)$ < $(\Delta_r H_2)$ < $(\Delta_r H_3)$

- ii) The ionisation enthalpy decreases on moving top to bottom in a group.
- iii) The ionisation enthalpy increases on moving from left to right in a period.

ELECTRON GAIN ENTHALPY: The electron gain enthalpy $((\Delta_{eg}H)$ is the molar enthalpy change when an isolated gaseous atom or ion in its ground state adds an electron to form the corresponding anion thus the enthalpy change for the reaction; $X_{(g)} + e^- \rightarrow X_{(g)}^-$

Is called the electron gain enthalpy $(\Delta_{eg} H)$ of the element X. The $\Delta_{eg} H$ may be positive or negative.

The successive values for the addition of second, third etc. Electron, these are called second, third etc. electron gain enthalpies. For example,

 $X_{(g)} + e^{-} \rightarrow X^{-}_{(g)}$ $\Delta H = \Delta_{eg} H_1$ is called first electron gain enthalpy $X^{-}_{(g)} + e^{-} \rightarrow X^{2^{-}}_{(g)}$ $\Delta H = \Delta_{eg} H_2$ is called second electron gain enthalpy

 $X^{2-}_{(g)} + e^{-} \rightarrow X^{3-}_{(g)}$ $\Delta H = \Delta_{eg} H_3$ is called third electron gain enthalpy Usually the term electron gain enthalpy ($\Delta_{eg}H$) means the first electron gain enthalpy.

Periodicity:

- (i) In period- The electron gain enthalpy increases from left to right in a period.
- (ii) In group- The electron gain enthalpy decreases from top to bottom in a group.

ELECTRONEGATIVITY: "The relative tendency of an atom in a molecule to attract the shared pair of electrons towards itself is termed as its electronegativity."

Periodicity:

- (i) In period- The electro-negativity increases from left to right in a period.
- (ii) In group- The electro-negativity decreases from top to bottom in a group.

VALENCE ELECTRONS: The electrons present in outermost shell are called as valence electron. Because the electrons in the outermost shell determine the valency of an element.

VALENCY OF AN ELEMENT: The number of hydrogen or halogen atom or double the number of oxygen atom, which combin with one atom of the element is taken as its valency. According to the electronic concept of valency, " the number of electrons which an atom loses or gains or shares with other atom to attain the noble gas configuration is termed as its valency."

Periodicity:

- (i) In period- The valency first increases then decreases from left to right in a period.
- (ii) In group- The valency remains constant from top to bottom in a group.

ELECTROPOSITIVE OR METALLIC CHARACTER: The tendency of an element to lose electrons and forms positive ions (cations) is called electropositive or metallic character. The elements having lower ionisation energies have higher tendency to lose electrons, thus they are electropositive or metallic in their behaviour.

Alkali metals are the most highly electropositive elements.

Periodicity: In period- The electropositive or metallic characters decreases from left to right in a period.

In group- The electropositive or metallic characters increases from top to bottom in a group.

ELECTRO-NEGATIVE OR NON- METALLIC CHARACTERS: the

tendency of an element to accept electrons to form an anion is called its non metallic or electronegative character. The elements having high electro-negativity have higher tendency to gain electrons and forms anion. So, the elements in the upper right hand portion of the periodic table are electro-negative or non-metallic in nature.

Periodicity:

- (i) In period- The electro-negative or non- metallic characters increases from left to right in a period.
- (ii) In group- The electro-negative or non-metallic characters decreases from top to bottom in a group.

REACTIVITY OF METALS:

Periodicity:

(i) In period- The tendency of an element to lose electrons decreases in a period. So the reactivity

of metals decreases from left to right in a period.

(ii) In group- The tendency of an element to lose electrons increases in a period. So the reactivity

of metals increases from top to bottom in a group.

REACTIVITY OF NON- METALS:

 (i) In period- The tendency of an element to gain electrons increases in a period. So the reactivity

of non-metals increases from left to right in a period.

(ii) In group- The tendency of an element to gain electrons decreases in a group. So the reactivity

of non-metals increases from top to bottom in a group.

SOLUBILITY OF ALKALI METALS CARBONATES AND BICARBONATES:

PERIODICITY IN GROUP: The solubility of alkali metal carbonates and bicarbonates in water increases down the group (From Lithium to Caesium).

SOLUBILITY OF ALKALINE EARTH METAL HYDROXIDES AND SULPHATES:

PERIODICITY IN GROUP: The solubility of alkaline earth metal hydroxide and sulphates in water increases down the group (From Beryllium to Barium).

BASIC STRENGTH OF ALKALINE EARTH METAL HYDROXIDES:

PERIODICITY IN GROUP: The basic strength of alkaline earth metal hydroxide in water increases down the group (From Beryllium to Barium), i.e.,

 $Be(OH)_2 \qquad < Mg(OH)_2 < Ca(OH)_2 < Sr(OH)_2 < Ba(OH)_2$

Basic strength increases

THERMAL STABILITY OF CARBONATES OF ALKALI AND ALKALINE EARTH METALS:

Except lithium carbonate, (LiCO₃), the carbonates of all other alkali metals are stable towards heat, i.e., carbonates of alkali metals (except LiCO₃) do not decompose on heating. LiCO₃ decomposes on heating to give lithium oxide (LiCO₃).

The carbonates of alkaline earth metals are relatively less stable. On heating, they decompose to give corresponding oxide and CO_2 gas. The decomposition temperature for alkaline earth metal carbonates increases as we go down the group.

Anomalous Properties of Second Period Elements

Their anomalous behaviour is attributed to their small size, large charge/radius ratio, high electro negativity, non- availability of d- orbitals in their valence shell. the first member of each group of p-Block elements displays greater ability to form pp-pp multiple bonds to itself (e.g. C=C, C=C O=O, N=N) and to other second period elements (e.g. C=O, C=N, N=O) compared to subsequent member of the group.

ONE MARK QUESTIONS

Q1. Select the species which are iso-electronic (same number of electron) with each other. (1) Ne (2) Cl^{-} (3) Ca^{2+} (4) Rb^{+}

Ans-The Cl⁻ and Ca²⁺. Both have $18 e^{-}$ each.

Q.2. What the elements of a group have common among them?

Ans- They have same number of electrons in the valence shell.

Q.3. What the s- and p- block elements are collectively called?

Ans- Representative elements.

Q.4. Define atomic radius.

Ans- The one-half the distance between the nuclei of two covalently bonded atoms of the same element in a molecule is called as atomic radius.

Q.5. State the modern periodic law.

Ans- The physical and chemical properties of the elements are the periodic function of their atomic numbers.

Q.6. Name the groups of elements classified as s-, p- and d- blocks.

Ans- s- block= 1,2 (including He), p- block= 13 to 18 (except He), d- block= 3 to 12.

Q.7. Define the term ionisation enthalpy.

Ans- The energy required to remove the outer most electron from the valence shell of an isolated gaseous atom is called as ionisation enthalpy.

Q.8.In how many groups and periods the elements in modern periodic table are classified?

Ans- In 18 groups and 7 periods.

Q.9. What do you mean by electronic configuration of the elements?

Ans- The systematic distribution of the electrons among the orbitals of an atom of an element according to increasing order of their energies is called as electronic configuration of that element.

TWO MARKS QUESTIONS

Q.1. Describe the two merits of long form periodic table over the Mendeleev's periodic table?

Ans- 1. It removed the anomalies about the position of isotopes which existed in the Mendeleev's table.

2. It relates the position of an element in the periodic table with its electronic configuration.

Q.2. What is a period in the periodic table? How do atomic sizes change in a period with an increase in atomic number?

Ans- The horizontal rows in periodic table are called as periods. The atomic sizes decrease in a period with an increase in atomic number.

Q.3. The outer electronic configuration of some elements are:

(a) $3s^2 3p^4$ (b) $3d^{10}4s^2$ (c) $3s^2 3p^6 4s^2$ (d) $6s^2 4f^3$

To which block of elements in the periodic table each of these belongs?

Ans- (a) p- Block (b) d- Block (c) s- Block (d) f- Block

Q.4. What is meant by periodicity in properties of elements? What is the reason behind this?

Ans- The repetition of similar properties after regular intervals is called as periodicity. It is due to the similarity in the outer electronic configurations which gives rise to the periodic properties of the elements.

Q.5. How do atomic radii vary in a group and a period?

Ans- In group- Atomic size increases on moving from top to bottom.

In period- Atomic size decreases on moving left to right in a period.

Q.6. Arrange the following in the order of increasing radii:

(a) I, I^+ , I^- (b) F, Cl, Br

Ans- (a) $I^+ < I < I^+$ (b) O < N < P

Q.7. Name the factors which affect the ionisation enthalpy of an element.

Ans- (i) Size of atom or ion (ii) Nuclear charge (iii) Electronic configuration

(iv) Screening effect (v) Penetration effect of the electrons

Q.8. How does ionisation enthalpy vary in a group and a period?

Ans- In Period- It increases from left to right

In group- It decreases down the group.

Q.9. Noble gases have zero electron gain enthalpy values. Explain.

Ans- Because the outer most shell of noble gases is completely filled and no more electrons can be added.

Q.10. Elements in the same group have equal valency. Comment on it.

Ans- Because the general outer most electronic configurations of the elements of a group remain same and they contain equal number of electrons in their respective outer most shells.

THREE MARKS QUESTIONS

Q.1. The first ionisation enthalpy of magnesium is higher than that of sodium. On the other hand, the second ionisation enthalpy of sodium is very much higher than that of magnesium. Explain.

Ans- The 1st ionisation enthalpy of magnesium is higher than that of Na due to higher nuclear charge and slightly smaller atomic radius of Mg than Na. After the loss of first electron, Na⁺ formed has the electronic configuration of neon (2,8). The higher stability of the completely filled noble gas configuration leads to very high second ionisation enthalpy for sodium. On the other hand, Mg⁺ formed after losing first electron still has one more electron in its outermost (3s) orbital. As a result, the second ionisation enthalpy of magnesium is much smaller than that of sodium.

Q.2. What are the major differences between metals and non- metals? Ans-

Property	Metal	Non- Metal	
Nature	Electropositive	Electronegative	
Type of ion	Cation (Positively Charged)	Anion (Negatively	
formed		Charged)	
Reaction with	Active metals displace	Do not displace hydrogen	
acids	hydrogen		
Oxides	Basic	Acidic	

Q.3. Among the elements of the second period Li to Ne pick out the element:

(i) with the highest first ionisation energy(ii) with the highest electronegativity

(iii) with the largest atomic radius Give the reason for your choice.

Ans- (i) The ionisation energy increases on going from left to right. Therefore, the element with the highest ionisation energy is Ne.

(ii) The electro negativity is electron- accepting tendency. This increases on going from left to right and decreases down the group. Therefore, the element with the highest electro- negativity is F.

(iii) The atomic radius decreases across a period on going from left to right. Thus, the first element of any period should have the largest atomic radii. Here, Li has the largest atomic radii.

Q.4. Arrange the following as stated:

	(i) N_2 , O_2 , F_2 , Cl_2			(Increasing order of bond dissociation energy)				
	(11) F, Cl, Br, I			(Increasing order of electron gain enthalpy)				
	(111)	$T_2, IN_2,$	CI_2, O	ν_2 (1	ncreas	sing ore	ler of	bond length)
Ans-	(i)	F_2	<	Cl_2	<	O_2	<	N_2
	(ii)	Ι	<	Br	<	F	<	Cl
	(iii)	N_2	<	O ₂	<	F_2	<	Cl_2

Q.5. Why does the first ionisation enthalpy increase as we go from left to right through a given period of the periodic table?

Ans- In a period, the nuclear charge (the number of protons) increases on going from left to right. The electron added to each element from left to right enters the same shell. This results in an increase of the effective nuclear charge across the period on moving from left to right. As a result, the electron get more firmly bound to the nucleus. This causes an increase in the first ionisation enthalpy across the period.

Q.6. Use the periodic table to answer the following questions.

(i) Identify the element with five electrons in the outer sub-shell.

(ii) Identify an element that would tend to lose two electrons.

(iii) Identify an element that would tend to gain two electrons.

Q.7. Explain why are cations smaller and anions larger in size than their parent atoms?

Ans- (a) The cations are smaller than their parent atoms due to the following reasons:

(i) Disappearance of the valence shell.

(ii) Increase of effective nuclear charge

(b) The anions are larger than their parent atoms due to the following reason:

An increase in the number of electrons in the valence shell reduces the effective nuclear charge due to greater mutual shielding by the electrons. As a result, electron cloud expands, i.e., the ionic radius increases.

Q.8. Describe the theory associated with the radius of an atom as it

(a) gains an electron (b) loses an electron

Ans- (a) When an atom gains an electron, its size increases. When an electron is added, the number of electrons goes up by one. This results in an increase in repulsion among the electrons. However, the number of protons remains the same. As a result, the effective nuclear charge of the atom decreases and the radius of the atom increases.

(b) When an atom loses an electron, the number of electrons decreases by one while the nuclear charge remains the same. Therefore, the interelectronic repulsions in the atom decrease. As a result, the effective nuclear charge increases. Hence, the radius of the atom decreases.

Q.9. How does atomic radius vary in a period and in a group? How do you explain the variation?

Ans- Atomic radius generally decreases from left to right across a period. This is because within a period, the outer electrons are present in the same valence shell and the atomic number increases from left to right across a period, resulting in an increased effective nuclear charge. As a result, the attraction of electrons to the nucleus increases.

On the other hand, the atomic radius generally increases down a group. This is because down a group, the principal quantum number (n) increases which results in an increase of the distance between the nucleus and valence electrons.

Q.10. Consider the following species:

 $N^{3-}\!\!,\,O^{2-}\!\!,\,F^{-}\!\!,\,Na^{+}\!\!,\,Mg^{2+}$ and Al^{3+}

(a) What is common in them?

(b) Arrange them in the order of increasing ionic radii.

Ans- (a) the same number of electrons (10 electrons). Hence, the given species are isoelectronic.

(b) $Al^{3+} < Mg^{2+} < Na^+ < F^- < O^{2-} < N^{3-}$

FIVE MARKS QUESTIONS

Q.1. What is the cause of the periodicity in the properties of the elements? How do the following properties vary in (a) a group and (b)in a period

(i) electronegativity (ii) ionisation enthalpy (iii) Atomic size

Ans- It is due to the similarity in the outer electronic configurations which gives rise to the periodic properties of the elements.

(a) In a group:

- (i) Electronegativity- It decreases down the group.
- (ii) Ionisation enthalpy- It decreases down the group.
- (iii) Atomic size- It increases down the group.

(b) In a period:

- (i) Electronegativity- Increases
- (ii) Ionisation enthalpy- Increases
- (iii) Atomic size- Dereases.

Q.2. The first $(\Delta_i H1)$ and the second $(\Delta_i H)$ ionization enthalpies (in kJ mol⁻¹) and the $(\Delta_{eg} H)$ electron gain enthalpy (in kJ mol⁻¹) of a few elements are given below:

Elements	$\Delta_i H$	$\Delta_i H$	$\Delta_{ m eg} H$
Ι	520	7300	-60
II	419	3051	-48
III	1681	3374	-328
IV	1008	1846	-295
V	2372	5251	+48
VI	738	1451	-40

Which of the above elements is likely to be :

(a) the least reactive element. (b) the most reactive metal.

(c) the most reactive non-metal. (d) the least reactive non-metal.

(e) the metal which can form a stable binary halide of the formula MX_2 , (X=halogen).

(f) the metal which can form a predominantly stable covalent halide of the formula MX (X=halogen)?

Ans- (a) Element V is likely to be the least reactive element. This is because it has the highest first ionization enthalpy $(\Delta_i H_1)$ and a positive electron gain enthalpy $(\Delta_{eg} H)$.

(b) Element II is likely to be the most reactive metal as it has the lowest first ionization enthalpy $(\Delta_i H_1)$ and a low negative electron gain enthalpy $(\Delta_{eg} H)$.

(c) Element III is likely to be the most reactive non-metal as it has a high first ionization enthalpy ($\Delta_{i}H_{1}$) and the highest negative electron gain enthalpy ($\Delta_{eg}H$).

(d) Element V is likely to be the least reactive non-metal since it has a very high first ionization enthalpy ($\Delta_i H_2$) and a positive electron gain enthalpy ($\Delta_{eg} H$).

(e) Element VI has a low negative electron gain enthalpy (Δ_{eg} H). Thus, it is a metal. Further, it has the lowest second ionization enthalpy (Δ_i H₂). Hence, it can form a stable binary halide of the formula MX₂(X=halogen).

(*f*) Element I has low first ionization energy and high second ionization energy. Therefore, it can form a predominantly stable covalent halide of the formula MX (X=halogen).

CHAPTER-4

CHEMICAL BONDING AND MOLECULAR STRUCTURE

OCTET RULE- During a chemical reaction the atoms tend to adjust their electronic arrangement in such a way that they achieve 8 e⁻ in their outermost electron. This is called octet rule.

CHEMICAL BOND- the chemical force which keeps the atoms in any molecule together is called a chemical bond.

IONIC BOND- The columbic force of attraction which holds the appositively charged ions together is called an ionic bond. An ionic bond is formed by the complete transfer of one or more electrons from the atom of a metal to an atom of non- metal.

LATTICE ENTHALPY- The molar enthalpy change accompanying the complete separation of the constituent particles that compose of the solids (such as ions for ionic solid, molecules for molecular solids) under standard conditions is called lattice enthalpy ($\Delta_l H^o$). The lattice enthalpy is a positive quantity.

ELECTRO VALENCY: The number of electrons lost or gain by an atom of an element is called as electrovalency.

The element which give up electrons to form positive ions are said to have positive valency, while the elements which accept electrons to form negative ions are said to have negative valency.

FORMATION OF AN IONIC BOND: It is favoured by, (i) the low ionisation enthalpy of a metallic element which forms the cations, (ii) High electron gain enthalpy of non- metallic element which forms the anions, (iii) Large lattice enthalpy i.e; the smaller size and the higher charge of the atoms.

COVALENCY: The number of electrons which an atom contributes towards mutual sharing during the formation of a chemical bond called its covalency in that compound.

SINGLE COVALENT BOND: A covalent bond formed by the mutual sharing of one pair of electrons is called a single covalent bond, or simply a single bond. A single covalent bond is represented by a small line (–) between the two atoms.

DOUBLE COVALENT BOND: A covalent bond formed by the mutual sharing of two pair of electrons is called a double covalent bond, or simply a double bond. A double covalent bond is represented by two small horizontal lines (=) between the two atoms. E.g. O=O, O=C=O etc.

TRIPLE COVALENT BOND: A covalent bond formed by the mutual sharing of three pair of electrons is called a triple covalent bond, or simply a triple bond. A triple covalent bond is represented by three small horizontal lines (\equiv) between the two atoms. E.g. N \equiv N, H-C \equiv C-H etc.

FORMATION OF A COVALENT BOND: Formation of a covalent bond is favoured by

- (i) High ionisation enthalpy of the combining elements.
- (ii)Nearly equal electron gain enthalpy and equal electro-negativities of combining elements.
- (iii) High nuclear charge and small atomic size of the combining elements.

POLAR COVALENT BOND: The bond between two unlike atoms which differ in their affinities for electrons is said to be polar covalent bond. E.g. H-Cl

COORDINATE BOND: The bond formed when one sided sharing of electrons take place is called a coordinate bond. Such a bond is also known as dative bond. It is represented by an arrow (\rightarrow) pointing towards the acceptor atom. E.g. $H_{3N} \rightarrow BF_3$

Bond Length: Bond length is defined as the equilibrium distance between the nuclei of two bonded atoms in a molecule

Bond Angle: It is defined as the angle between the orbitals containing bonding electron pairs around the central atom in a molecule/complex ion

Bond Enthalpy: It is defined as the amount of energy required to break one mole of bonds of a particular type between two atoms in a gaseous state.

Bond Order: In the Lewis description of covalent bond, the Bond Order is given by the number of bonds between the two atoms in a molecule

Resonance:whenever a single Lewis structure cannot describe a molecule accurately, a number of structures with similar energy, positions of nuclei, bonding and non-bonding pairs of electrons are taken as the canonical structures of the hybrid which describes the molecule accurately

Dipole moment : The product of the magnitude of the charge and the distance between the centres of positive and negative charge. It is a vector quantity and is represented by an arrow with its tail at the positive centre and head pointing towards a negative centre. Dipole moment (μ) = charge (Q) × distance of separation (r) **SIGMA BOND:** A covalent bond formed due to the overlapping of orbitals of the two atoms along the line joining the two nuclei (orbital axis) is called sigma (σ) bond. For example, the bond formed due to s-s and s-p, p-p overlapping along the orbital axis are sigma bonds.

Pi-BOND: A covalent bond formed by the side wise overlapping of p- or dorbitals of two atoms is called as pi (π) bond. For example, the bond formed due to the sideways overlapping of the two p- orbitals is a pi- bond.

HYDROGEN BOND: The bond between the hydrogen atom of one molecule and a more electro- negative element of same or another molecule is called as hydrogen bond.

HYBRIDIZATION: The process of mixing of the atomic orbitals to form new hybrid orbitals is called hybridization. All hybrid orbitals of a particular kind have equal energy, identical shapes and are symmetrically oriented in shape.

The hybrid orbitals are designed according to the type and the atomic orbitals merging together, e.g.,







Q.1. What is the total number of sigma and pi bonds in the following molecules?

(a) C₂H₂ (b) C₂H₄

Ans- there are three sigma and two pi-bonds in C_2H_2 .

there are five sigma bonds and one pi-bond in C_2H_4 .

Q.2. Write the significance of a plus and a minus sign shown in representing the orbitals.

Ans- Molecular orbitals are represented by wave functions. A plus sign in an orbital indicates a positive wave function while a minus sign in an orbital represents a negative wave function.

Q.3. How do you express the bond strength in terms of bond order?

Ans- Bond strength represents the extent of bonding between two atoms forming a molecule. The larger the bond energy, the stronger is the bond and the greater is the bond order.

Q.5. Define the bond length.

Ans- Bond length is defined as the equilibrium distance between the nuclei of two bonded atoms in a molecule.

Q.6. Arrange the bonds in order of increasing ionic character in the molecules: LiF, K_2O , N_2 , SO_2 and ClF_3 .

Ans- $N_2 < SO_2 < ClF_3 < K_2O < LiF$.

Q.7. The skeletal structure of CH_3COOH as shown below is correct, but some of the bonds are shown incorrectly. Write the correct Lewis structure for acetic acid.

Ans- The correct Lewis structure for acetic acid is as follows:

Q.8. Define octet rule.

Ans- The elements tend to adjust the arrangement of their electrons in such a way that they (except H and He) achieve eight electrons in their outermost shell. This is called octet rule.

Q.9. Define lattice enthalpy.

Ans- The energy required when one mole of an ionic compound in crystalline form is split into the constituent ions is called lattice enthalpy.

Q.10. Which type of bond is formed when the atoms have zero difference in electronegativity?

Ans- Covalent bond.

TWO MARKS QUESTIONS

Q.1. Define hydrogen bond. Is it weaker or stronger than the van der Waals forces?

Ans- A hydrogen bond is defined as an attractive force acting between the hydrogen attached to an electronegative atom of one molecule and an electronegative atom of a different molecule (may be of the same kind).

There are two types of H-bonds:

- (i) Intermolecular H-bond e.g., HF, H₂O etc.
- (ii) Intramolecular H-bond e.g., o-nitrophenol



Hydrogen bonds are stronger than Van der Walls forces since hydrogen bonds are regarded as an extreme form of dipole-dipole interaction.

Q.2. Write the favourable factors for the formation of ionic bond.

Ans-(i) Low ionization enthalpy of metal atom.

(ii) High electron gain enthalpy $(\Delta_{eg} H)$ of a non-metal atom.

(iii) High lattice energy of the compound formed.

Q.3. Although geometries of NH_3 and H_2O molecules are distorted tetrahedral, bond angle in water is less than that of ammonia. Discuss.

Ans- The molecular geometry of NH_3 and H_2O can be shown as:



The central atom (N) in NH_3 has one lone pair and there are three bond pairs. In H_2O , there are two lone pairs and two bond pairs.

The two lone pairs present in the oxygen atom of H_2O molecule repels the two bond pairs. This repulsion is stronger than the repulsion between the lone pair and the three bond pairs on the nitrogen atom.

Since the repulsions on the bond pairs in H_2O molecule are greater than that in NH_3 , the bond angle in water is less than that of ammonia.

Q.4. Explain the important aspects of resonance with reference to the CO_3^{2-} ion.

Ans- According to experimental findings, all carbon to oxygen bonds in $^{CO_3^{2-}}$ are equivalent. Hence, it is inadequate to represent $^{CO_3^{2-}}$ ion by a single Lewis structure having two single bonds and one double bond.

Therefore, carbonate ion is described as a resonance hybrid of the following structures:



Q.5. H_3PO_3 can be represented by structures 1 and 2 shown below. Can these two structures be taken as the canonical forms of the resonance hybrid representing H_3PO_3 ? If not, give reasons for the same.



Ans- The given structures cannot be taken as the canonical forms of the resonance hybrid of H_3PO_3 because the positions of the atoms have changed.

Q.6. Use Lewis symbols to show electron transfer between the following atoms to form cations and anions: (a) Ca and O (c) Al and N.

Ans(a) <u>Ca and O:</u>

The electronic configurations of Ca and O are as follows:

Ca: 2, 8, 8, 2 O: 2, 6

Oxygen requires two electrons more to complete its octet, whereas calcium has two electrons more than the nearest noble gas i.e., Argon. Hence, the electron transfer takes place as:

Ca: $Ca^{2+}_{2,8,8,2}$ $Ca^{2+}_{2,8,8}$ $Ca^{2+}_{2,8,8} \equiv Ca^{2+}o^{2-}$ (b) Al and N: The electronic configurations of Al and N are as follows:

Al: 2, 8, 3 N: 2, 5

Nitrogen is three electrons short of the nearest noble gas (Neon), whereas aluminium has three electrons more than Neon. Hence, the electron transference can be shown as:

$$Al^{3+}_{2,8,3} \xrightarrow{N}_{2,5} Al^{3+}_{2,8,2,8} \xrightarrow{N}_{2,8}^{3-} \equiv Al^{3+} N^{3-}_{2,8}$$

Q.7. Although both CO_2 and H_2O are triatomic molecules, the shape of H_2O molecule is bent while that of CO_2 is linear. Explain this on the basis of dipole moment.

Ans- According to experimental results, the dipole moment of carbon dioxide is zero. This is possible only if the molecule is linear so that the dipole moments of C-O bonds are equal and opposite to nullify each other.

$$: \overset{\leftarrow}{\circ} = c = \overset{\leftarrow}{\circ} :$$

Resultant $\mu = 0$ D

 H_2O , on the other hand, has a dipole moment value of 1.84 D (though it is a triatomic molecule as CO_2). The value of the dipole moment suggests that the structure of H_2O molecule is bent where the dipole moment of O–H bonds are unequal.



Q.8. Write the significance/applications of dipole moment.

Ans- Dipole moment is the measure of the polarity of a bond. It is used to differentiate between polar and non-polar bonds since all non-polar molecules (e.g. H_2 , O_2) have zero dipole moments. It is also helpful in calculating the percentage ionic character of a molecule.



Q.9. Use molecular orbital theory to explain why the Be_2 molecule does not exist.

Ans- The electronic configuration of Beryllium is $1s^2 2s^2$.

The molecular orbital electronic configuration for Be₂ molecule can be written as:

$$\sigma_{1s}^2 \sigma_{2s}^{\star 2} \sigma_{2s}^{\star 2} \sigma_{2s}^{\star 2}$$
Hence, the bond order for Be₂ is $\frac{1}{2} (N_b - N_a)$.

Where,

 N_b = Number of electrons in bonding orbitals N_a = Number of electrons in anti-bonding orbitals

 $\therefore \text{Bond order of Be}_2 = \frac{1}{2} (4-4)_{=0}$

A negative or zero bond order means that the molecule is unstable. Hence, Be_2 molecule does not exist.

Q.10. Distinguish between a sigma and a pi bond.

Ans- The following are the differences between sigma and pi-bonds:

Sigma (σ) Bond	Pi (π) Bond
(a) It is formed by the end to end overlap of orbitals.	It is formed by the lateral overlap of orbitals.
(b) The orbitals involved in the overlapping are <i>s</i> – <i>s</i> , <i>s</i> – <i>p</i> , or <i>p</i> – <i>p</i> .	These bonds are formed by the overlap of p -porbitals only.
(c) It is a strong bond.	It is weak bond.
(d) The electron cloud is symmetrical about the line joining the two nuclei.	The electron cloud is not symmetrical.
(e) It consists of one electron cloud, which is symmetrical about the internuclear axis.	There are two electron clouds lying above and below the plane of the atomic nuclei.
(f) Free rotation about σ bonds is possible.	Rotation is restricted in case of pi- bonds.

Q.11. Explain with the help of suitable example polar covalent bond.

Ans- When two dissimilar atoms having different electronegativities combine to form a covalent bond, the bond pair of electrons is not shared equally. The bond pair shifts towards the nucleus of the atom having greater electronegativity. As a result, electron distribution gets distorted and the electron cloud is displaced towards the electronegative atom.

As a result, the electronegative atom becomes slightly negatively charged while the other atom becomes slightly positively charged. Thus, opposite poles are developed in the molecule and this type of a bond is called a polar covalent bond.

HCl, for example, contains a polar covalent bond. Chlorine atom is more electronegative than hydrogen atom. Hence, the bond pair lies towards chlorine and therefore, it acquires a partial negative charge.

$$\mathbf{H} \underbrace{\mathbf{O}}_{\mathbf{H}} := \mathbf{H} \underbrace{\mathbf{O}}_{\mathbf{H}} \cdot \mathbf{C}^{\mathbf{h}}$$

Bond pair attracted more toward

THREE MARKS QUESTIONS

Q.1. Write Lewis dot symbols for atoms of the following elements: Mg, Na, B, O, N, Br.

Ans-Mg: Na Br Br

Q.3. Draw diagrams showing the formation of a double bond and a triple bond between carbon atoms in C_2H_4 and C_2H_2 molecules.

Ans- C_2H_4 : The electronic configuration of C-atom in the excited state is:

 $_{6}C = 1s^{2}2s^{1}2p_{x}^{1}2p_{y}^{1}2p_{z}^{1}$

In the formation of an ethane molecule (C₂H₄), one sp^2 hybrid orbital of carbon overlaps a sp^2 hybridized orbital of another carbon atom, thereby forming a C-C sigma bond. The remaining two sp^2 orbitals of each carbon atom form a sp^2 -s sigma bond with two hydrogen atoms. The unhybridized orbital of one carbon atom undergoes sidewise overlap with the orbital of a similar kind present on another carbon atom to form a weak π -bond.



$C_2H_2:$

In the formation of C_2H_2 molecule, each C-atom is *sp* hybridized with two 2*p*-orbitals in an unhybridized state.

One *sp* orbital of each carbon atom overlaps with the other along the internuclear axis forming a C–C sigma bond. The second *sp* orbital of each C–atom overlaps a half-filled 1*s*-orbital to form a σ bond.

The two unhybridized 2p-orbitals of the first carbon undergo sidewise overlap with the 2p orbital of another carbon atom, thereby forming two pi (π) bonds between carbon atoms. Hence, the triple bond between two carbon atoms is made up of one sigma and two π -bonds.







Q.4. Explain the formation of H_2 molecule on the basis of valence bond theory.

Ans- Let us assume that two hydrogen atoms (A and B) with nuclei (N_A and N_B) and electrons (e_A and e_B) are taken to undergo a reaction to form a hydrogen molecule. When A and B are at a large distance, there is no interaction between them. As they begin to approach each other, the attractive and repulsive forces start operating.

Attractive force arises between:

(a) Nucleus of one atom and its own electron i.e., $N_A - e_A$ and $N_B - e_{B.}$

(b) Nucleus of one atom and electron of another atom i.e., N_{A} – e_{B} and N_{B} – $e_{A.}$

Repulsive force arises between:

(a) Electrons of two atoms i.e., $e_A - e_{B_{-}}$

(b) Nuclei of two atoms i.e., $N_A - N_{B_{-}}$

The force of attraction brings the two atoms together, whereas the force of repulsion tends to push them apart.



Repuisive Forces

The magnitude of the attractive forces is more than that of the repulsive forces. Hence, the two atoms approach each other. As a result, the potential energy decreases. Finally, a state is reached when the attractive forces balance the repulsive forces and the system acquires minimum energy. This leads to the formation of a dihydrogen molecule.

Q.5. Write the important conditions required for the linear combination of atomic orbitals to form molecular orbitals.

Ans- The given conditions should be satisfied by atomic orbitals to form molecular orbitals:

(a) The combining atomic orbitals must have the same or nearly the same energy. This means that in a homonuclear molecule, the 1*s*-atomic orbital of an atom can combine with the 1*s*-atomic orbital of another atom, and not with the 2*s*-orbital.(b) The combining atomic orbitals must have proper orientations to ensure that the overlap is maximum.

(c) The extent of overlapping should be large.

Q.6. Describe the hybridisation in case of PCl₅. Why are the axial bonds longer as compared to equatorial bonds?

Ans- The ground state and excited state outer electronic configurations of phosphorus (Z = 15) are:

Ground state:

† ↓	1	
35	3 <i>p</i>	3 <i>d</i>

Excited state:

1	1	†
35	3 <i>p</i>	3 <i>d</i>

Phosphorus atom is sp^3d hybridized in the excited state. These orbitals are filled by the electron pairs donated by five Cl atoms as:

PCl₅

†∔	14 14 14	11	
1	<u>t</u> t t t	† ;	
Ċl	CI CI CI	C1 ¦	

The five sp^3d hybrid orbitals are directed towards the five corners of the trigonal bipyramidals. Hence, the geometry of PCl₅ can be represented as:



There are five P–Cl sigma bonds in PCl₅. Three P–Cl bonds lie in one plane and make an angle of 120° with each other. These bonds are called equatorial bonds. The remaining two P–Cl bonds lie above and below the equatorial plane and make an angle of 90° with the plane. These bonds are called axial bonds. As the axial bond pairs suffer more repulsion from the equatorial bond pairs, axial bonds are slightly longer than equatorial bonds.

Q.7. What is meant by the term bond order? Calculate the bond order of: N_2 , O_2 , O_2^+ and O_2^- .

Ans- Bond order is defined as one half of the difference between the number of electrons present in the bonding and anti-bonding orbitals of a molecule.

Bond order = $\frac{1}{2}(N_{\rm b} - N_{\rm a})$

Bond order of N_2

 $[\sigma(1s)]^{2}[\sigma^{*}(1s)]^{2}[\sigma(2s)]^{2}[\sigma^{*}(2s)]^{2}[\pi(2p_{s})]^{2}[\pi(2p_{s})]^{2}[\sigma(2p_{z})]^{2}$

Number of bonding electrons, $N_{\rm b} = 10$

Number of anti-bonding electrons, $N_a = 4$

Bond order of nitrogen molecule $=\frac{1}{2}(10-4)$ = 3

Bond order of O₂ $[\sigma - (1s)]^2 [\sigma^*(1s)]^2 [\sigma(2s)]^2 [\sigma(2s)]^2 [\sigma(1p_z)]^2 [\pi(2p_x)]^2 [\pi(2p_y)]^2 [\pi^*(2p_x)]^1 [\pi^*(2p_y)]^1$

Bond order $=\frac{1}{2}(N_{\rm b}-N_{\rm a}) =\frac{1}{2}(8-4) = 2$

Hence, the bond order of oxygen molecule is 2.

Similarly, the electronic configuration of O_2^* can be written as:

 $\mathrm{KK}[\sigma(2s)]^{2}[\sigma^{*}(2s)]^{2}[\sigma(2p_{z})]^{2}[\pi(2p_{x})]^{2}[\pi(2p_{y})]^{2}[\pi^{*}(2p_{x})]^{1}$

Bond order of $O_2^+ = \frac{1}{2}(8-3) = 2.5$

The electronic configuration of O_2^- ion will be:

 $\mathrm{KK}[\sigma(2s)]^{2}[\sigma^{*}(2s)]^{2}[\sigma(2p_{z})]^{2}[\pi(2p_{x})]^{2}[\pi(2p_{y})]^{2}[\pi^{*}(2p_{x})]^{2}[\pi^{*}(2p_{y})]^{1}$

Bond order of $O_2^- = \frac{1}{2}(8-5) = 1.5$

Q.8. Discuss the shape of the following molecules using the VSEPR model:

BeCl₂, BCl₃, SiCl₄, AsF₅, H₂S, PH₃

Ans- $\underline{BeCl_2}$: Cl: Be: Cl The central atom has no lone pair and there are two bond pairs. i.e., $BeCl_2$ is of the type AB₂. Hence, it has a linear shape.

<u>BCl_{3:} Cl:B:Cl</u> The central atom has no lone pair and there are three bond pairs.

Hence, it is of the type AB₃. Hence, it is trigonal planar.

CI: Si CI

SiCl_{4:} The central atom has no lone pair and there are four bond pairs. Hence, the shape of SiCl₄ is tetrahedral being the AB_4 type molecule.



<u>AsF_{5:}</u> F The central atom has no lone pair and there are five bond pairs. Hence, AsF₅ is of the type AB₅. Therefore, the shape is_trigonal bipyramidal.

H₂S: H:S:H The central atom has one lone pair and there are two bond pairs. Hence, H₂S is of the type AB_2E . The shape is Bent.

н: Р :н

<u>PH_{3:}</u> The central atom has one lone pair and there are three bond pairs. Hence, PH₃ is of the AB₃E type. Therefore, the shape is trigonal bipyramidal.

Q.9. Write the resonance structures for SO₃, NO₂ and $^{NO_3^-}$.

Ans- The resonance structures are:

(a) SO₃:





Q.10. What do you understand by bond pairs and lone pairs of electrons? Illustrate by giving example.

Ans- The shared pairs of electrons present between the bonded atoms are called **bond pairs**. All valence electrons may not participate in bonding. The electron pairs that do not participate in bonding are called**lone pairs** of electrons.

In H₂O, there are two bond pairs and two lone pairs on the central atom (oxygen).



FIVE MARKS QUESTIONS

Q.1. Define octet rule. Write its significance and limitations.

Ans-- The octet rule or the electronic theory of chemical bonding was developed by Kossel and Lewis. According to this rule, atoms can combine either by transfer of valence electrons from one atom to another or by sharing their valence electrons in order to attain the nearest noble gas configuration by having an octet in their valence shell.

O :: C:: O Or O = C = O

The octet rule successfully explained the formation of chemical bonds depending upon the nature of the element.

Limitations of the octet theory:

The following are the limitations of the octet rule:

(a) The rule failed to predict the shape and relative stability of molecules.

(b) It is based upon the inert nature of noble gases. However, some noble gases like xenon and krypton form compounds such as XeF_2 , KrF_2 etc.

(c) The octet rule cannot be applied to the elements in and beyond the third period of the periodic table. The elements present in these periods have more than eight valence electrons around the central atom. For example: PF_5 , SF_6 , etc.



(d) The octet rule is not satisfied for all atoms in a molecule having an odd number of electrons. For example, NO and NO_2 do not satisfy the octet rule.

$$N = 0$$
 $0 = N - 0$

(e) This rule cannot be applied to those compounds in which the number of electrons surrounding the central atom is less than eight. For example, LiCl, BeH₂, AlCl₃ etc. do not obey the octet rule.

Li:Cl Cl:Al:Cl H:Be:H

Q.2. Which hybrid orbitals are used by carbon atoms in the following molecules?

CH₃-CH₃; (b) CH₃-CH=CH₂; (c) CH₃-CH₂-OH; (d) CH₃-CHO (e) CH₃COOH

Ans- (a)



Both C_1 and C_2 are sp^3 hybridized.

(b)



 C_1 is sp^3 hybridized, while C_2 and C_3 are sp^2 hybridized.



Both C_1 and C_2 are sp^3 hybridized.



 C_1 is sp^3 hybridized and C_2 is sp^2 hybridized.





 C_1 is sp^3 hybridized and C_2 is sp^2 hybridized.

Q.3. Compare the relative stability of the following species and indicate their magnetic properties;

 O_2, O_2^+, O_2^- (superoxide), $O_2^{2^-}$ (peroxide)

Ans- There are 16 electrons in a molecule of dioxygen, 8 from each oxygen atom. The electronic configuration of oxygen molecule can be written as:

$$[\sigma - (1s)]^{2}[\sigma^{*}(1s)]^{2}[\sigma(2s)]^{2}[\sigma^{*}(2s)]^{2}[\sigma(1p_{z})]^{2}[\pi(2p_{x})]^{2}[\pi(2p_{y})]^{2}[\pi^{*}(2p_{x})]^{1}[\pi^{*}(2p_{y})]^{1}[\pi^{*}$$

Since the 1s orbital of each oxygen atom is not involved in boding, the number of bonding electrons = $8 = N_b$ and the number of anti-bonding orbitals = $4 = N_a$.

Bond order
$$= \frac{1}{2} (N_b - N_a)$$
$$= \frac{1}{2} (8 - 4)$$
$$= 2$$

Similarly, the electronic configuration of O_2^+ can be written as:

$$\mathrm{KK}[\sigma(2s)]^{2}[\sigma^{*}(2s)]^{2}[\sigma(2p_{z})]^{2}[\pi(2p_{x})]^{2}[\pi(2p_{y})]^{2}[\pi^{*}(2p_{x})]^{1}$$

 $N_{\rm b} = 8$

 $N_{\rm a} = 3$

Bond order of $O_2^+ = \frac{1}{2}(8-3)$

= 2.5

Electronic configuration of O_2^- ion will be:

 $\mathrm{KK}[\sigma(2s)]^{2}[\sigma^{*}(2s)]^{2}[\sigma(2p_{z})]^{2}[\pi(2p_{x})]^{2}[\pi(2p_{y})]^{2}[\pi^{*}(2p_{x})]^{2}[\pi^{*}(2p_{y})]^{1}$

 $N_{\rm b}=8$

 $N_{\rm a} = 5$

Bond order of $O_2^- = \frac{1}{2}(8-5)$

Electronic configuration of O_2^{2-} ion will be:

$$\mathrm{KK}[\sigma(2s)]^{2}[\sigma^{*}(2s)]^{2}[\sigma(2p_{z})]^{2}[\pi(2p_{x})]^{2}[\pi(2p_{y})]^{2}[\pi^{*}(2p_{x})]^{2}[\pi^{*}(2p_{y})]^{2}[\pi^{*}(2p_{$$

 $N_{\rm b} = 8$

 $N_{\rm a}=6$

Bond order of $O_2^{2-} = \frac{1}{2}(8-6)$

= 1

Bond dissociation energy is directly proportional to bond order. Thus, the higher the bond order, the greater will be the stability. On this basis, the order of stability is $O_2^+ > O_2 > O_2^- > O_2^{2-}$.

HOTS

Q.1. Apart from tetrahedral geometry, another possible geometry for CH_4 is square planar with the four H atoms at the corners of the square and the C atom at its centre. Explain why CH_4 is not square planar?

Ans- Electronic configuration of carbon atom: ${}_{6}C: 1s^{2} 2s^{2} 2p^{2}$

In the excited state, the orbital picture of carbon can be represented as:

† ↓	1	1	1	1	
15	2 <i>s</i>	$2p_x$	2p	,2p	1

Hence, carbon atom undergoes sp^3 hybridization in CH₄ molecule and takes a tetrahedral shape.



For a square planar shape, the hybridization of the central atom has to be dsp^2 . However, an atom of carbon does not have *d*-orbitalsto undergo dsp^2 hybridization. Hence, the structure of CH₄ cannot be square planar.

Moreover, with a bond angle of 90° in square planar, the stability of CH₄ will be very less because of the repulsion existing between the bond pairs. Hence, VSEPR theory also supports a tetrahedral structure for CH₄.

Q.2. Explain why BeH₂ molecule has a zero dipole moment although the Be–H bonds are polar.

Ans- The Lewis structure for BeH₂ is as follows: H:Be:H

There is no lone pair at the central atom (Be) and there are two bond pairs. Hence, BeH_2 is of the type AB₂. It has a linear structure.

 $H \xrightarrow{++} Be \xrightarrow{++} H$

Dipole moments of each H–Be bond are equal and are in opposite directions. Therefore, they nullify each other. Hence, BeH₂ molecule has zero dipole moment.

Q.3. Which out of NH₃ and NF₃ has higher dipole moment and why?

Ans- In both molecules i.e., NH_3 and NF_3 , the central atom (N) has a lone pair electron and there are three bond pairs. Hence, both molecules have a pyramidal shape. Since fluorine is more electronegative than hydrogen, it is expected that the net dipole moment of NF_3 is greater than NH_3 . However, the net dipole moment of NH_3 (1.46 D) is greater than that of NF_3 (0.24 D).

This can be explained on the basis of the directions of the dipole moments of each individual bond in NF_3 and NH_3 . These directions can be shown as:



Thus, the resultant moment of the N–H bonds add up to the bond moment of the lone pair (the two being in the same direction), whereas that of the three N – F bonds partly cancels the moment of the lone pair.

Hence, the net dipole moment of NF_3 is less than that of NH_3 .

Q.4. What is meant by hybridisation of atomic orbitals? Describe the shapes of sp, sp^2 , sp^3 hybrid orbitals.

Ans- Hybridization is defined as an intermixing of a set of atomic orbitals of slightly different energies, thereby forming a new set of orbitals having equivalent energies and shapes.

For example, one 2*s*-orbital hybridizes with two 2*p*-orbitals of carbon to form three new sp^2 hybrid orbitals.

These hybrid orbitals have minimum repulsion between their electron pairs and thus, are more stable. Hybridization helps indicate the geometry of the molecule.

Shape of *sp* **hybrid orbitals:** *sp* hybrid orbitals have a linear shape. They are formed by the intermixing of *s* and *p* orbitals as:



Shape of sp^2 hybrid orbitals:

 sp^2 hybrid orbitals are formed as a result of the intermixing of one *s*-orbital and two 2*p*-orbitals. The hybrid orbitals are oriented in a trigonal planar arrangement as:



Shape of sp^3 hybrid orbitals:

Four sp^3 hybrid orbitals are formed by intermixing one s-orbital with three *p*-orbitals. The four sp^3 hybrid orbitals are arranged in the form of a tetrahedron as:



Q.5. Describe the change in hybridisation (if any) of the Al atom in the following reaction.

 $AlCl_3 + Cl^- \longrightarrow AlCl_4^-$

Ans- The valence orbital picture of aluminium in the ground state can be represented as:

$$\begin{array}{c|c} \hline \hline \hline \\ 3s^2 & 3p \end{array}$$

The orbital picture of aluminium in the excited state can be represented as:

1	1	1	
35	$3p_x$	3p)	,3pz

Hence, it undergoes sp^2 hybridization to give a trigonal planar arrangement (in AlCl₃).

To form AlCl₄⁻, the empty $3p_z$ orbital also gets involved and the hybridization changes from sp^2 to sp^3 . As a result, the shape gets changed to tetrahedral.
<u>CHAPTER 5</u>

STATES OF MATTER

- 1. Change in state : It is over all effect of Intermolecular forces, molecular Interactional energy & thermal energy:
- 2. Measurable properties of gases : P,V, T, n, Viscosity, specific heat are some measurable properties.
- **3. Gas Laws** : The quantitative relationship b/w any two of the variables (V, n, P,T) when other two are constant.
- 4. Boyle's Law : The pressure of fixed msss of gas varies inversely with the volume at constant T. P α 1/V(n,T const.)

 $P_1V_1 = P_2V_2$

5. Charle's Law : At constant P, the volume of fixed amount of gas varies directly with its absolute temperature.

V
$$\alpha$$
 T or $\frac{V}{T} = cons \tan t, \ \frac{V_1}{T_1} = \frac{V_2}{T_2}$

6. Gay lussac's Law : At constant V, The pressure of fixed amount of gas varies directly with its absolute temperature.

P
$$\alpha$$
 T or $\frac{P}{T} = const$, $\frac{P_1}{T_1} = \frac{P_2}{T_2}$

- 7. Ideal gas equation : The relationship b/w P, V and T by Gas Laws PV= nRT.
- 8. Avogadro's Law : At given T and P, the volume of gas varies directly to the amount of gas . V α n (P, T constant)
- **9.** Dalton's Law of partial persure : The pressure enerted by a mixture of non reacting gases is equal to the sum of their partial pressure at constant (V,T)

 $P(total) = P1 + P2 + P3 + \dots (T, V, constant)$

- **10.** Kinetic Molecular theory :
 - a. Gases consist of large number of identical particles (atoms or molecules) that are so small that the actual volume of the molecules is negligible in comparison to the empty space between them.
 - b. There is no force of attraction between the particles of a gas at ordinary temperature and pressure
 - c. Particles of a gas are always in constant and random motion
 - d. Pressure is exerted by the gas as a result of collision of the particles with the walls of the container
 - e. Collisions of gas molecules are perfectly elastic
 - f. At any particular time, different particles in the gas have different speeds and hence different kinetic energies
 - g. Average kinetic energy of the gas molecules is directly proportional to the absolute temperature
- **11.** Real Gases : The gases which deviates from Ideal behavior at higher pressure and low temperature b/c of force of attraction b/w molecules increases .
- **12.** Compressibility factor (Z) : It determine extent of devation

of real gases from Ideal gas behavior : $Z = \frac{PV}{n.R.T.}$ for ideal

gas Z=1, for Nonideal gas Z< 1, Z > 1

13. Vander waal's Equation :
$$\left(P + \frac{n^2 a}{V^2}\right) \Psi - nb = nRT.$$

- 14. Critical Temperature : (Tc) The temperature above which a gas cannot be liquefied whatever high pressure may be
- **15.** Critical Pressure : The minimum pressure required to liquity a gas at its critical temperature.
- **16.** Critical Volume : The volume of 1 mole of gat at Tc, Pc.
- Super cooled liquids : The liquids which are cooled to a temperature below its freezing point without freezing.

- **18.** Elastic Collision : The collisions in which no loss of K.E. only there is transfer of energy.
- **19.** Vapour pressure : The equilibrium pressure by vapour of liquid in a container at given temperature (T)
- **20.** At higher altitude : The b.p. of water decreases b/c the atmospheric pressure is less than one atmosphere.
- **21.** Surface Tension (V) : It is force acting per unit length perpendicular to the line drawn on the surface : (Nm^{-1}) : It decreases with increases in T, it increases with increase in external pressure, b/c of it falling drops of liquid are spherical, liquid in capillary tube rises.
- 22. Viscosity (η) : It is resistance offered to the flow of liquid due to friction b/w layer of fluids . $F = n.A.\frac{dv}{dn}$
- 23. Effect of T & P on viscosity : It decreases with increase in T, and increases with increase in P.
- **24.** Low M.P. & B.P. of molecular liquids is due to low magnitude of molecular interaction energy.

One mark questions

- 1. At what condition surface tension vanishes ?
- 2. Why Helium is used in balloons in place of hydrogen?
- 3. At what temperature below which a gas does not obey ideal gas law?
- 4. At what temperature the volume of a gas is supposed to be zero?
- 5. What is the molar volume at 0° C and 1 bar pressure?

- 6. Name the temperature above which a gas cannot be liquefied by any amount of pressure.
- 7. What is the effect of increase of temperature on surface tension and viscosity in a liquid?
- 8. How is the partial pressure of a gas in a mixture is related to the total pressure of the gaseous mixture?
- 9. Why vegetables are cooked with difficulty at hill station.
- 10. What is the value of Z (compressibility factor) for an Ideal gas?

Answers to One mark questions

- Ans 1 At critical temperature, meniscus b/w liquid and vapours disappears.
- Ans 2. He is incombustible, though heavier than H_2 .
- Ans 3. Below Boyle Temperature.
- Ans 4. At absolute zero (O K) temperature.
- Ans 5. It 22400 ml.
- Ans 6. It is critical temperature (Tc)
- Ans 7. Both decreases with increase in temperature
- Ans 8. $P_1 = X_1 \times P_{(total)}$
- Ans 9. At hill station the atmospheric pressure is less and so boiling point decreases.
- Ans10. For ideal gas Z=1.

Two Marks Questions

Q.1 How do you convert pressure in atmosphere in to SI unit.

- Q.2 What type graph will you get when PV is plotted against P at constant temperature.
- Q.3 What would have happened to the gas if the molecular collisions were not elastic?
- Q.4 At a particular temperature, why vapour pressure of acetone is less than of ether?
- Q.5 Why liquids diffuse slowly as compared to gases?
- Q.6 What would be the SI unit for quantity $\frac{P.V^2.T^2}{n}$?
- Q.7 In terms of Charle's law explain why -273° C is the lowest temperature?
- Q.8 For real gases the relation b/w P,V,T is given by vander Waal's equation, write it for n moles?
- Q.9 What correction is applied to obtain pressure of dry gas with the total pressure & aqueous tension?
- Q.10 Name two phenomena that can be explained on the basis of surface tension.

Answers to two marks questions

- Ans 1. $1 \text{ atm} = 101325 \text{ Pa or } \text{Nm}^{-2}, 1 \text{ bar} = 10^5 \text{ Pa}.$
- Ans 2 .A straight line parallel to pressure axis.
- Ans 3. On every collision there is loss of energy, so molecules would have slowed down & settled down in vessel and pressure reduce to O.
- Ans 4. b/c molecular force of attraction in acetone is stronger than those present in ether.

Ans 5. In liquids the molecules are less free than in a gas. i. e intermolecular forces in liquid are greater than in gas.

Ans 6.
$$\frac{P.V^2.T^2}{n} = \frac{(N.m^{-2})m^3}{mal} = N.m^4 k^2 mal^{-1}$$

Ans 7 At -273° C, the Volume of gas becomes equal to zero i. e. gas ceases to exist.

Ans 8. Vander waal's equation

$$\left(P+\frac{n^2a}{v^2}\right)\Psi-nb=n.R.T.$$

Ans 9. $P_{(dryygas)} = P_{(Total)} - Aqueous tension$

Ans 10. Surface tension can explain (i) capillary action. (ii) Spherical shape of small drops of liquid.

Three Marks Questions

Q.1 Calculate temperature of 4.0 moles of gas occupying 5dm^3 volume at 3.32 bar. (R= 0.083 bar dm³ k⁻¹ mol⁻¹)

Ans PV = nRT, or $T = \frac{PV}{nR}$

Or
$$T = \frac{3.32 \ bar \ x \ 5 \ dm^3}{4.0 \ mol \ x \ 0.083 \ bar \ \ dm^3 \ k^{-1} \ mol^{-1}} = 50. \ Ans$$

Q.2 34.05 ml of phosphorus vapours weight 0.0625g at 543° C and 1 bar pressure what is molar mass of phosphorous?

Ans.
$$\frac{P_1V_1}{T_1} = \frac{P_2V_2}{T_2}, \Rightarrow \frac{1x34.05}{(54b+273)} = \frac{1xv_2}{273} at 0^\circ C$$

Or V₂ = 11.35 ml
Now 11.35 ml of vapours at 0°C and 1 bar pressure weight 0.0625 gm

Now 22700 ml of vapors at 0° C and 1 bar pressure weight $\frac{0.0625}{11.35}x22700$

- Q.3 A mixture of dihydrogen and dinitrogen at 1 bar pressure contains 20% by weight of H_2 . Calculate partial pressure of H_2 .
- Ans The mixture of H_2 and O_2 contain 20% H_2 means 20 gm H_2 & 80gm O_2

$$n_{2} (H_{2}) = \frac{20}{2} = 10 \text{moles}, \ n(O_{2}) = \frac{80}{32} = 2.5 \text{contest}.$$
$$P_{(H2)} = \frac{n(H_{2})}{n(H_{2}) + n(V_{2})} \quad x \ P_{(total)} = \frac{10}{10 + 2.5} \ x \ 1bar = 0.8bar$$

- Q.4 What will be minimum pressure required compressing 500dm^3 of air at 1 bar to 200 dm^3 at 30° C?
- Ans. $P_1 = 1bar$. $V_1 = 500 dm^3$ $P_2 = ?$ $V_2 = 200 dm^3$ & temp. constant $30^{\circ}C$ So by Boyle's low $P_1V_1 = P_2 V_2$

1 x 500 = P₂ x 200 or P₂
$$\frac{500}{200}$$
 = 2.5 bar

Q.5 Calcutate the total pressure in mixture of 4g of O_2 and 2gm. Of H₂ can fixed to a total volume of IL at 0° C. (R = 0.0821) Latm. Mol⁻¹

Q.6 Account for the following :

(i) The size of weather balloon becomes longer and larger as it ascend up into higher altitudes.

Ans (i) At higher altitudes atmospheric pressure decreases, but inside the balloon gas exerts pressure and size becomes larger.

- Q.8 What do you mean by ideal gas and real gas? Why do real gases deviate from ideal behaviors?
- Ans. Ideal gas follows gas laws at all P & TReal gas donot " gas laws at all P & T.Real gas deviate from ideal behaviour due to force of attraction and because of volume of molecules of gases are not negligible.
- Q.9 Using the equation of state PV=nRT Show that at given T, density of gas proportional to gas pressure P.

Ans
$$PV = \underline{n}RT \Longrightarrow P = \frac{n}{v}.RT$$

 $\Rightarrow P = \frac{m}{M}, \frac{1}{V}, RT. = \frac{m}{V}.\frac{RT}{M}$
 $\Rightarrow P = P.\frac{RT}{M}$ (Where $P = \frac{m}{v}$)
 $\Rightarrow P \times P\left(\because \frac{RT}{M} = const.\right)$

- Q.10 Critical temperature of CO_2 and CH_4 gases are $31.1^{\circ}C$ and $81.9^{\circ}C$ respectively. Which of these has strong intermolecular forces & why?
- Ans CO_2 has strong vander waal's forces of attraction than CH_4 . b/c of greater polarity and high molecular wt.

Hots Questions with Answers.

- (a)Why aerated water bottles kept under water during summer? Q.1
 - Which property of liquid is responsible for spherical (b) shape of drop?
 - Why is moist air lighter than dry air? (c)
 - Define aqueous tension? (d)
 - What is units of a and b which are vander waal's (e) constant?
- To reduce temperature so to reduce pressure, other wise Ans (a) bottle may burst.

(b) Surface Tension.

(c) Moist air has water vapours which lowers vapour density, so it is lighter.

(d) It is pressure of water vapours at given T. (e)Unit of a in atm. $L^2 \text{ mol}^{-2}$, b is L mol^{-1.}

Why does sharpened edge becomes smooth on heating up to Q.2 melting point?

On heating the glass it melts and take up rounded shape at Ans edges which has minimum surface area. b/c of surface tension. Q.3 Arrange following in order of increasing density:

$$d = \frac{PM}{RT}$$
, O_2 at 25° C, 1atm, O_2 at O^0C , 2atm, O_2 at 273° C, 1atm.

Ans $d = \frac{PM}{RT}$, R, M are constant, so d depends upon $\frac{P}{T}$ so at 25°C,

 $1 \operatorname{Atm} \frac{P}{T} = \frac{1}{298}$, At 273° C, 1atm $\frac{P}{T} = \frac{1}{546}$ Hence increasing order of

density will be: O_2 at 273^o C, 1atm < O_2 at 25^o C, 2atm.

An O₂ cylinder has 10L O₂ at 200 atm. It patient takes 0.50ml Q.4 of O_2 at 1 atm in one breath at 37^{0} C. How many breaths are possible? 10L at 200 atm = ? L at 1 atm $\Rightarrow P, V, = P_2 V_2 \Rightarrow 200 \times 10 = 1 \times V_2$ or Ans

$$V_2 = 2000 \text{ L}$$
, No of breathes = $\frac{Total Volume}{Volume \text{ for 1 breath}} = \frac{2000 \text{ L}}{0.5 \times 10^{-3} \text{ L}} = 4 \times 10^6$

<u>CHAPTER 6</u> <u>THERMODYNAMICS</u>"

Brief Summary of the chapter:

- 1. Thermodynamics: Science which deals with study of different forms of energy and quantitative relationship.
- 2. System & Surroundings: The part of universe for study is called system and remaining portion is surroundings.
- State of system & state function: State of system is described in terms of T,
 P, V etc. The property which depends only on state of system not upon path is called state function eg. P, V, T, E, H, S etc.
- 4. Extensive & Intensive Properties: Properties which depends on quantity of matter called extensive prop. eg. mass, volume, heat capacity, enthalpy, entropy etc. The properties which do not depends on matter present depends upon nature of substance called Intensive properties. eg. T,P, density, refractive index, viscosity, bp, pH, mole fraction etc.
- 5. Internal energy: The total energy with a system.

i.e. $U = E_e + E_n + E_c + E_p + E_k + \dots$

 $\Delta U = U_2 - U_1$ or $U_P - U_R$ & U is state function and extensive properly. If $U_1 > U_2$ energy is released.

- 6. Heat (q): It I a form of energy which is exchanged between system and surrounding due to difference of temperature. Unit is Joule (J) or Calorie (1 Calorie = 4.18μ J).
- 7. First Law of Thermodynamics: It is law of conservation energy. Energy can neither be created not destroyed, it may be converted from one from into another.

Mathematically $\Delta U = q + w$, w = -p. ΔV (work of expansion)

 $\Delta U = q - p$. ΔV or $q = \Delta U + p$. ΔV , q,w are not state function.

But ΔU is state function.

8. Enthalpy (H): At constant volume
$$\Delta V = 0, \rightarrow q_v = \Delta U$$

So $H = U + p$. ΔV , $q_p = H_2 H_1 = \Delta H$
 $\Rightarrow \Delta H = \Delta U + P$. ΔV .

- 9. Relationship between q_p , q_v i.e. $\Delta H \& \Delta U$ It is $\Delta H = \Delta U + \Delta ng.RT$ or $q_p = q_{v+} \Delta ng.RT$
- 10. Exothermic and Endothermic reactions: $\Delta H = -Ve$ for exothermic and $\Delta H = +Ve$ for endothermic reaction i.e. evolution and absorption of heat. Eg C+O₂ \rightarrow CO₂ + 393.5 KJ, $\Delta H = -393.5$ KJ (exothermic) N₂ + O₂ \rightarrow 2NO - 180.7 KJ, $\Delta H = 180.7$ KJ (Endothermic)
- 11. Enthalpy of reaction (Δ_r H): The amount of heat evolved or absorbed when the reaction is completed.
- 12. Standard Enthalpy of reaction $(\Delta_r H^0)$ at 1 bar pressure and specific temp. (290K) i.e. standard state.
- 13. Different types of Enthalpies of reactions:
 - (i) Enthalpy of combustion ($\Delta_c H$), (ii) Enthalpy of formation ($\Delta_f H$)
 - (iii) Enthalpy of neutralization (iv) Enthalpy of solution
 - (v) Enthalpy of atomization($\Delta_a H$), (vi)Enthalpy of Ionisation ($\Delta_i H$)
 - (vii) Enthalpy of Hydration ($\Delta_{hyol.}$ H) (viii) Enthalpy of fusion ($\Delta_{fus.}$ H)
 - (ix) Enthalpy of vaporization $(\Delta_{vap.}H)$ (x)Enthalpy of sublimation $(\Delta_{sub.}H)$ $(\Delta_{sub.}H) = \Delta_{fus.}(H) - \Delta_{vap}(H)$
- 14. Hess's Law of constant heat summation: The total amount of heat change is same whether the reaction takes place in one step or in several steps.

i.e. $\Delta H = \Delta H_1 + \Delta H_2 + \Delta H_3 + \cdots$

- 15. Bond enthalpy: It is amount of energy released when gaseous atoms combines to form one mole of bonds between them or heat absorbed when one mole of bonds between them are broken to give free gaseous atoms. Further $\Delta_r H = \Sigma B.E.$ (Reactants) - $\Sigma B.E.$ (Products)
- 16. Spontaneous & Non Spontaneous Processes: A process which can take place by itself is called spontaneous process. A process which can neither take place by itself or by initiation is called non Spontaneous.
- Driving forces for spontaneous process: (i) Tendency for minimum energy state. (ii) Tendency for maximum randomness.
- Entropy (S): It is measure of randomness or disorder of system.
 i.e. Gas>Liquid>Solid.

Entropy change (ΔS) = $\frac{q_{(rev.)}}{T} J.K^{-1}.mol^{-1}$

19. Spontaneity in term of (ΔS)

 $\Delta S_{(total)} = \Delta S_{(universe)} = \Delta S_{(system)} + \Delta S_{(surrounding)}$

If $\Delta S_{(total)}$ is +ve, the process is spontaneous.

If $\Delta S_{(total)}$ is -ve, the process is non spontaneous.

- 20. Second Law of thermodynamics: In any spontaneous process, the entropy of the universe always increases. A spontaneous process cannot be reversed.
- 21. Gibb's free energy (G): defined as $G = H T.S & \Delta G = \Delta H T. \Delta S$ (Gibb's Helmholts equation) it is equal useful work i.e. $-\Delta G = W_{(useful)} = W_{(max.)}$

If ΔG = ve, process is spontaneous.

- 22. Effects of T on spontaneity of a process: $\Delta G = \Delta H T$. ΔS .
 - (i) For endothermic process may be non spontaneous at law temp.

- (ii)For exothermic process may be non spontaneous at high temp. and spontaneous at law temp.
- 23. Calculation of $(\Delta_r G^0)$

$$\Delta_{\mathbf{r}}\mathbf{G}^{0} = \Sigma \Delta_{\mathbf{f}}\mathbf{G}^{0} (\mathbf{p}) - \Sigma \Delta_{\mathbf{f}}\mathbf{G}^{0} (\mathbf{r})$$

24. Relationship between $(\Delta_r G^0)$ & equilibrium constant (k)

 $\Delta G = \Delta G^{0} + RT lnQ \& \Delta G^{0} = -2.303RT \log k.$

25. Calculation of entropy change:

 $\Delta_{\rm r} {\rm S}^0 = \Sigma \Delta {\rm S}^0 ({\rm p}) - \Sigma {\rm S}^0 ({\rm r})$

ONE MARK QUESTIONS:

- 1. State First Law of thermodynamics.
- 2. What is a thermodynamic state function?
- 3. Give enthalpy (H) of all elements in their standard state.
- 4. From thermodynamic point to which system the animals and plants belong?
- 5. Predict the sign of ΔS for the following reactions.

 $CaCO_{3(s)} + CO_{2(g)} \xrightarrow{heat} CaO_{(s)} + CO_{2(g)}$

- 6. For the reaction $2Cl_{(g)} \rightarrow Cl_{2(g)}$, What will be the sign of ΔH and ΔS ?
- 7. State Hess's Law for constant heat summation?
- 8. What is Gibb's Helmhaltz equation?
- 9. Define extensive properties.
- 10. Give relationship between ΔH , ΔU for a reaction in gaseous state.

ANSWERS FOR ONE MARK QUESTIONS

- 1. Energy can neither be created nor destroyed. The energy of an isolated system is constant. $\Delta U = q + w$.
- 2. A function whose value is independent of path. eg. P, V, E, H
- 3. In standard state enthalpies of all elements is zero.
- 4. Open system.
- 5. ΔS is positive (entropy increases)
- 6. Δ H: (-ve) b/c energy is released in bond formation and

 ΔS : (-ve) b/c atoms combines to form molecules.

- 7. The change of enthalpy of reaction remains same, whether the reaction is carried out in one step or several steps.
- 8. $\Delta G = \Delta H T. \Delta S$
- 9. Properties which depends upon amount of substance called extensive properties. Volume, enthalpy, entropy.
- 10. $\Delta H = \Delta U + ng. RT.$

TWO MARKS QUESTIONS:-

- Q.1 In a process, 701J heat is absorbed and 394J work is done by system. What is change in Internal energy for process?
- Q.2 Given: $N_{2(g)} + 3H_{2(g)} \rightarrow 2NH_{3(g)}, \Delta_r H^0 = -92.4 \text{KJ.mol}^{-1}$. What is the standard enthalpy of formation of $NH_{3(g)}$.
- Q.3 Calculate entropy change in surroundings when 1.0 mol of H₂O(*l*) is formed under standard conditions? Given $\Delta H^0 = -286 \text{KJmol}^{-1}$.
- Q.4 Give relationship between entropy change and heat absorbed/evolved in a reversible reaction at temperature T.
- Q.5 What is spontaneous change? Give one example.

- Q.6 A real crystal has more entropy than an Ideal Crystal. Why?
- Q.7 Under what condition, the heat evolved/absorbed in a reaction is equal to its free energy change?
- Q.8 Predict the entropy change in-
 - (i) A liquid crystallizes into solid
 - (ii) Temperature of a crystallize solid raised from OK to 115K
- Q.9 What is bond energy? Why is it called enthalpy of atomization?
- Q.10 Calculate entropy change for the following process.

 $H_2O_{(s)} \Leftrightarrow H_2O_{(l)}$, is 6.0 KJ mol⁻¹ at 0⁰C.

ANSWER FOR TWO MARKS QUESTIONS:

1. q = 701J, w = 394J, so $\Delta U = q + w = 701 - 394 = 307J$.

2.
$$\Delta_{\rm f} {\rm H}^{\Theta} {\rm NH}_{3({\rm g})} = -\frac{92.4}{2} = 46.2 {\rm KJ.mol}^{-1}$$

3.
$$q_{(rev.)} = -\Delta H^0 = -286 \text{ KJmol}^{-1} = 286000 \text{Jmol}^{-1}$$

$$\Delta S = \frac{q_{(rev.)}}{T} = \frac{286000 \text{ J.mol}^{-1}}{298 \text{ K}} = 959 \text{ J.K}^{-1} \text{ mol}^{-1}$$

- 4. $\Delta S = \frac{q_{(rev)}}{T}$
- A process which can take place of its own or initiate under some condition.
 eg. Common salt dissolve in water of its own.
- 6. A real crystal has some disorder due to presence of defects in their structural arrangement, and Ideal crystal does not have any disorder.
- 7. In $\Delta G = \Delta H T$. ΔS , when reaction is carried out at OK or $\Delta S = 0$, then $\Delta G = \Delta H$.

- 8. (i) Entropy decreases b/c molecules attain an ordered state.
 (ii)entropy increase b/c from OK to 115K particles begin to move.
- 9. It is the amount of energy required to dissociate one mole of bonds present b/w atoms in gas phase. As molecules dissociates into atoms in gas phase so bond energy of diatomic molecules is called enthalpy of atomization.
- 10. $H_2O_{(s)} \Leftrightarrow H_2O_{(l)}$ at 0^0C , $\Delta_{fus}H = 6KJ \text{ mol}^{-1}$

 $= 6000 \text{J mol}^{-1}$

 $\Delta T_f = 0^0 C = 0 + 273 = 273 K$

Do $\Delta_{fus}s = \frac{\Delta_{fus}H}{T} = \frac{6000 \text{ J.mol}^{-1}}{273 \text{ K}} = 21.98 \text{ J.K}^{-1} \text{mol}^{-1}$

THREE MARKS QUESTIONS:

- Q.1 For oxidation of iron, $4Fe_{(s)} + 3O_{2(g)} \rightarrow 2Fe_2O_{3(s)}$ ΔS is $-549.4J.K^{-1}$ mol⁻¹, at 298K. Inspite of –ve entropy change of this reaction, Why the reaction is spontaneous? ($\Delta_r H^0 = -1648 \times 10^3 \text{ J.mol}^{-1}$)
- Q.2 Using the bond energy of $H_r = 435 \text{ KJ mol}^{-1}$, $Br_2 = 192 \text{ KJ mol}^{-1}$, $HBr = 368 \text{ KJmol}^{-1}$. Calculate enthalpy change for the reaction $H_{2(g)} + Br_{2(g)} \rightarrow 2HBr_{(g)}$
- Q.3 Enthalpies of formation of $CO_{(g)}$, $CO_{2(g)}$, $N_2O_{(g)}$ and $N_2O_{4(g)}$ and -110, -393, 81 and 9.7 KJ mol⁻¹ respectively. Find value $\Delta_r H$ for the reaction $N_2O_{4(g)} + 3CO_{(g)}$
- Q.4 For the reaction at 298K, $2A+B \rightarrow C$, $\Delta H = 400 \text{ KJ mol}^{-1}$, $\Delta S = 0.2 \text{ KJ mol}^{-1} \text{ K}^{-1}$. At what temperature will the reaction become spontaneous, considering ΔH , ΔS be constant at the temp.
- Q.5 The equilibrium constant for a reaction is 10. What will be the value of ΔG^{0} ? R = 8.314J.K⁻¹mol⁻¹ T = 300K.
- Q.6 What do you understand by state function? Neither q nor w is a state function but q + w is a state function? Explain.
- Q.7 Justify the following statements:

- (i) An endothermic reaction is always thermodynamically spontaneous.
- (ii) The entropy always increases on going from liquid to vapour state at any temperature T.
- Q.8 Calculate the temperature above which the reduction reaction becomes spontaneous: $PbO_{(s)} + C_{(s)} Pb_{(s)} + CO_{(g)}$, given [$\Delta H = 108.4 \text{ KJ mol}^{-1}$, $\Delta S = 190 \text{J.K}^{-1} \text{ mol}^{-1}$].
- Q.9 From the data given below at 298K for the reaction:

$$\begin{split} CH_{4(g)} &+ 2O_{2(g)} \, CO_{2(g)} + 2H_2 O_{(l)} \, Calculate \, enthalpy \, of \, formation \, of \, CH_{4(g)} \, at \\ 298K. \, Given: [\Delta_r H = -890.5 \, KJ, \\ [\Delta_r H_{(CO_2)} &= -393.5 \, KJ. mol^{-1}, \Delta_r H_{(H_2O)} = -286.0 \, KJ. mol^{-1}] \end{split}$$

Q.10 For the reaction $NH_4Cl_{(s)} NH_{3(g)} + HCl_{(g)}$ at 25^0C enthalpy change $\Delta H = 177KJ.mol^{-1}$ and $\Delta S = 285J.K^{-1} mol^{-1}$. Calculate free energy change ΔG at 25^0C and predict whether the reaction is spontaneous or not.

ANSWER TO THREE MARKS QUESTIONS:

Ans.1 $\Delta_{(surr.)} = -\frac{\Delta_r H^0}{T} = -\frac{C - 1648 \text{ x} 10^3 \text{ J.mol}^{-1}}{298 \text{ K}} = 5530 \text{ JK}^{-1} \text{mol}^{-1}.$

& $\Delta S_{(system)} = -549.4 \text{QJK}^{-1} \text{mol}^{-1}$.

 $\Delta_r S_{(total)} = 5530 - 5494 = 4980.6 \text{ J.K}^{-1} \text{mol}^{-1}$

Since $\Delta_r S_{(total)}$ is +ve, so the reaction is spontaneous.

Ans.2 $\Delta_r H^0 = \Sigma$ bond enthalpies_(rect.) - Σ bond enthalpies_(prod.)

$$= \mathbf{H}_{H-H}^{0} + \mathbf{H}_{Br-Br}^{0} + \mathbf{H}_{H-Br}^{0}$$
$$= [435 - 192] - [2 \times 368] \text{ KJ mol}^{-1}$$
$$= 627 - 736 = -109 \text{ KJ. Mol}^{-1} \text{ Ans.}$$

Ans.3 $\Delta_r H = \Delta_f H^0_{(\text{prod.})} \Delta_r H^0_{(\text{rect})}$

$$= \mathbf{h}_{f} H^{0}_{(N_{2}O)} + 3\Delta_{f} H^{0}_{(CO_{2})} - \mathbf{h}_{f} H^{0} N_{2}O_{4} + 3\Delta_{f} H^{0}_{(CO)}$$
$$= [81 + 3(-393)] - [9.7 + (-110)]$$

= [81 + 1179] - [9.7 - 330] = -777.7 KJAns.4 $\Delta H = 400 \text{ KJ mol}^{-1}, \Delta S = 0.2 \text{ KJK}^{-1} \text{ mol}^{-1}.$ $\Delta G = \Delta H - T. \Delta S$ $O = 400 - 0.2 \text{ x T} (\Delta G = 0 \text{ at equilibrium})$ $T = \frac{400}{0.2} = 2000 \text{K}$, so reaction will be spontaneous above 2000 K.
Ans.5 $\Delta_r G^0 = -2.303 \text{ RT log K}$ = -2.303 x 8.314 x 300 x log 10 = -19.147 x 300 x 1 = -5744.1 J $\Delta_r G^0 = -5.7441 \text{KJ.mol}^{-1}$

Ans.6 The property whose value depends upon state of system and is independent of path. $q + w = \Delta U$, which is a state function as value of ΔU does not depends upon path.

Ans.7 (a) It is false, exothermic reaction is not always spontaneous. If $\Delta S = +ve$ and T. $\Delta S > \Delta H$. The process will be non spontaneous even it. It is endothermic. (b) The entropy of vapour is more than that of liquid, so entropy increases during

vaporization.

Ans.8 $\Delta G = \Delta H - T$. ΔS , at equilibrium $\Delta G = 0$, $\Rightarrow \Delta H = T$. ΔS

$$\Rightarrow T = \frac{\Delta H}{\Delta S} = \frac{108.4 \times 10^{3} \text{ J.mol}^{-1}}{190 \text{ J.K}^{-1} \text{ mol}^{-1}} = 570.526 \text{ K}$$

So the reaction will be spontaneous above 570.52K, as above this temperature ΔG will be -ve.

Ans.9 $\Delta_{\rm r} H = \Delta_{\rm f} H_{\rm (CO_2)} + 2\Delta_{\rm f} H C H_{\rm 4(g)} - \Delta_{\rm f} H_{\rm (O_2)}$

 $-890.5 \text{KJ} = -393.5 \text{KJ} + 2 \text{v} - 286 - \Delta H_f(\text{CH}_4) - \text{O}$

 $= \Delta H_{\rm f}({\rm CH_4}) = -75.0 \text{ KJ.mol}^{-1}.$

 $= \Delta H_{f(CH_{c})} = -75.0 \text{KJ.mol}^{-1}$.

Ans.10 $\Delta H = 177 \text{ KJ mol}^{-1}$, $\Delta S = 285 \text{ JK}^{-1} \text{ mol}^{-1}$

$$\Delta G = \Delta H - T. \ \Delta S = 177 \text{KJ} - \frac{298 \times 285}{1000} \text{KJ}$$

 $= 177 \text{ KJ} - 84.93 \text{ KJ} = 92.07 \text{ KJ.mol}^{-1}$.

Since ΔG is +ve, so the reaction is non spontaneous.

FIVE MARKS QUESTIONS:-

- Q.1 What is entropy? Why is the entropy of a substance taken as zero at 0K? Calculate the $\Delta_r G$ for the reaction? $N_{2(g)} + 3H_{2(g)} \implies 2NH_{3(g)}$ at 298K The value of equilibrium constant (K) is 6.6×10^5 , R = $8.314 \text{ JK}^{-1} \text{ mol}^{-1}$.
- Ans: It is measure of randomness or disorder of system.

Because at O K there is complete order in the system.

$$\Delta G^{0} = -2.303 \text{ RT} \log K = -2.303 \text{ x } 8.314 \text{ x } 298 \text{ x } \log 6.6 \text{ x } 105$$
$$= -5705.8[\log 6.6 + \log 10^{5}]$$
$$= -5705.8[0.8195 + 5.0] = -5705.8 + 5.8195\text{J}$$
$$= -33204.903\text{J}$$
$$\Delta G^{0} = -33.205 \text{ KJ mol}^{-1}.$$

Q.2 (i) What are extensive property and intensive properties?(ii)Calculate the value of equilibrium constant (K) at 400K for

2 NOCl_(g) →2NO_(g) + Cl_{2(g)}.

$$\Delta H^0 = 77.2$$
KJ.mol⁻¹, $\Delta S^0 = 122$ J.K⁻¹mol⁻¹ at 400K, R = 8.314 J.K⁻¹mol⁻¹.

Ans. (i) An **extensive property** is a property whose value depends on the quantity or size of matter present in the system Those properties which do not depend on the quantity or size of matter present are known as **intensive properties**

(ii) $\Delta H^0 = \Delta H^0 - T. \ \Delta S^0 = 77.2 \text{KJ} - \frac{400 \text{x} 122}{1000} \text{KJ.mol}^{-1}$ = 77.2 - 48.8 = 28.4 KJ mol⁻¹ and $\Delta G^0 = -2.303 \text{ RT} \log \text{K}.$ 28400 = -2.303 x 8.314 x 400 log K.

$$\log K = \frac{7.1}{2.303 \times 8.314} = -3.7081 = \overline{4}.2919 K$$

K – antilog ($\overline{4}$.2919) = 1.95 x10⁻⁴ Ans.

Q.3 Define standard enthalpy of formation. Calculate the enthalpy of formation of benzene from data

$$C_{6}H_{6(1)} + \frac{15}{2}O_{2(g)} \rightarrow 6CO_{2(g)} + 3H_{2}O_{(1)}, \Delta_{c}H^{0} = -3266.0KJ$$

$$C_{(s)} + O_{2(g)} \rightarrow CO_{2(g)} \qquad \Delta_{f}H^{0} = -393.1KJ$$

$$H_{2(g)} + \frac{1}{2}O_{2(g)} \rightarrow H_{2}O_{(1)} \qquad \Delta_{f}H^{0} = -286.0KJ$$

Ans. Ans. The standard enthalpy change for the formation of one mole of a compound from its elements in their most stable states of aggregation (also known as reference states) is called Standard Molar Enthalpy of Formation.

$$\Delta_{c}H^{0} = 6\Delta_{f}H^{0}_{(co2_{2})} + 3\Delta_{f}H^{0}_{(H_{2}O)} - \Delta_{f}H^{0}_{(C_{6}H_{6})} - \frac{15}{2}\Delta H^{0}_{f(O_{2})}$$
$$= -3266\text{KJ} = 6 \text{ x} - 393.1 + 3\text{ x} - 286 - \Delta_{f}H^{0}_{(C_{6}H_{6})} - 0$$

= -3218 kJ/mol

HOTS QUESTIONS

 Why standard entropy of an elementary substance is not zero whereas standard enthalpy of formation is taken as zero? Ans. A substance has a perfectly ordered arrangement only at absolute zero.

Hence, entropy is zero only at absolute zero. Enthalpy of formation is the heat change involved in the formation of one mole of the substance from its elements. An element formed from it means no heat change.

2. The equilibrium constant for a reaction is one or more if ΔG° for it is less than zero. Explain

Ans. $-\Delta G^{\circ} = RT \ln K$, thus if ΔG° is less than zero. i.e., it is negative, then ln K will be positive and hence K will be greater than one.

3. Many thermodynamically feasible reactions do not occur under ordinary conditions. Why? Ans. Under ordinary conditions, the average energy of the reactants may be

Ans. Under ordinary conditions, the average energy of the reactants may be less than threshold energy. They require some activation energy to initiate the reaction.

CHAPTER-7

<u>EQUILIBRIUM</u>

Equilibrium state- When rate of formation of a product in a process is in competition with rate of formation of reactants, the state is then named as "Equilibrium state".

solid \rightleftharpoons liquid \rightleftharpoons gas

- ➢ Equilibrium in physical processes:
 H₂O_{(s}) ⇒ H₂O_(l) ⇒ H₂O_(vap)
- Law of chemical equilibrium: At a given temperature, the product ofconcentrations of the reaction productsraised to the respective stoichiometriccoefficient in the balanced chemicalequation divided by the product ofconcentrations of the reactants raised totheir individual stoichiometric coefficientshas a constant value. This is known asthe Equilibrium Law or Law of ChemicalEquilibrium.

 $aA +bB \rightleftharpoons cC + dD$ $K_c = [C]^c [D]^d / [A]^a [B]^b$

Chemical equation		Equilibriumconstant
aA + b B ≓ c C + D		K
$cC + dD \rightleftharpoons aA + bB$		K'c = (1/Kc)
na A + nb B ⇒ncC + ndD		$K'''c = (Kc^n)$
	C	

Concentrations or partial pressure of pure solids orliquids do notappear in the expression of the equilibriumconstant. In the reaction,

 $Ag_2O(s) + 2HNO_3(aq) \rightleftharpoons 2AgNO_3(aq) + H_2O(l)Kc = \frac{[AgNO_3]^2}{[HNO_3]^2}$

- > If Qc > Kc, the reaction will proceed in the direction of reactants (reverse reaction). If Qc < Kc, the reaction will proceed in the direction of the products (forward reaction)
- Kp is equilibrium constant in terms of partial pressure of gaseous reactants and products.
- Kc is equilibrium constant in terms of molar concentration of gaseous reactants and products.
- ► Kp =Kc $(RT)^{\Delta n}$ here **R** is gas constant, **T** is temperature at which the process is carried out & Δn is no. of moles of gaseous product minus no. of moles of gaseous reactants.
- > If $Kc > 10^3$; Kc is very high i.e. the reaction proceeds nearly to completion.
- > If $Kc < 10^3$; Kc is very small i.e. the reaction proceeds rarely.
- If Kcis ranging in the range of 10³ to 10⁻³; i.e. reactants and products are just in equilibrium.
- $\blacktriangleright \Delta G^0 = -RT \ln K$ or $\Delta G^0 = -2.303RT \log K$
- Factors affecting equilibrium constant:- temperature, pressure, catalyst and molar concentration of reactants and products.

- Le Chatelier's principle:- It states that a change in any of the factors thatdetermine the equilibrium conditions of asystem will cause the system to change such a manner so as to reduce or tocounteract the effect of the change.
- \blacktriangleright Arrhenius acids are the substances that ionize in water to form \mathbf{H}^+ .
- > Arrhenius bases are the substances that ionize in water to form **OH**⁻.
- Lewis acids are lone pair (of e-) accepters while Lewis bases are lone pair donators.
- Proton donor are acids while proton accepters are bases(Bronsted-Lowry concept).
- The acid-base pair thatdiffers only by one proton is called a conjugateacidbase pair. IfBrönsted acid is a strong acid then itsconjugate base is a weak base and viceversa.
- > Ionic product of water. $Kw = [H^+][OH^-]$
- > $\mathbf{pH} = -\log [\mathbf{H}^+]$; here $[\mathbf{H}^+]$ is molar concentration of hydrogen ion.
- ➢ pH + pOH =14
- ➢ pKa + pKb =14
- \blacktriangleright Ka x Kb = Kw = ionic product of water=1 x 10⁻¹⁴
- Buffer solution :The solutions which resist change in pH on dilution or with the addition of small amounts of acid or alkali are called Buffer Solutions.
- common ion effect: It can be defined as a shift in equilibrium on adding a substance that provides more of an ionic species already present in the dissociation equilibrium.
- Hydrolysis of Salts: process of interaction between water andcations/anions or both of salts is calledhydrolysis.
- The cations (e.g., Na⁺, K⁺, Ca²⁺, Ba²⁺, etc.) of strong bases and anions(e.g., Cl⁻, Br⁻, NO³⁻, ClO⁴⁻ etc.) of strong acids simply get hydrated but do not hydrolyse, andtherefore the solutions of salts formed fromstrong acids and bases are neutral i.e., theirpH is 7.
- Salts of weak acid and strong base e.g., CH₃COONa are basic in nature.
- Salts of strong acid and weak base e.g.,NH₄Cl, are acidic
- Salts of weak acid and weak base, e.g., CH₃COONH₄. The pH is determined by the formula $pH = 7 + \frac{1}{2} (pK_a pK_b)$
- Solubility product- product of the molar concentrations of the ions in a saturated solution, each concentration term raised to the power equal to the no. of ions produced.

ONE MARK QUESTIONS

Q.1. Mention the factors that affect equilibrium constant.

Ans. Temperature, pressure, catalyst and molar concentration of reactants and products.

Q.2. What is ionic products of water? Ans. $Kw = [H^+] [OH^-]$

Q.3. Write conjugate acids of H_2O & NH_3 . Ans. H_3O^+ & NH_4^+ .

Q.4. Define Arrhenius acids. Ans. Arrhenius acids are the substances that ionize in water to form \mathbf{H}^+ .

Q.5. Define the term degree of ionization. Ans.Extent up to which an acid/base/salt ionize to form ions.

Q.6. What are Buffer solutions?

Ans.The solutions which resist change in pH on dilution or with the addition of small amounts of acid or alkali are called Buffer Solutions.

Q.7. Write Kc for the gaseous reaction- $N_2 + 3H_2 \rightleftharpoons 2NH_3$ Ans. Kc=[NH₃]²/[N₂] [H₂]³

Q.8. Out of $H_2O \& H_3O^+$ which is stronger acid? Ans. H_3O^+ .

Q.9. What is common ion effect?

Ans. Shift in equilibrium on adding a substance that provides more of an ionic species already present in the dissociation equilibrium.

Q.10. Write relationship between Kp and Kc for the gaseous reaction - N₂ + O₂ \rightleftharpoons 2NO

Ans. $Kp = Kc as\Delta n$ is zero for the above said reaction.

TWO MARKS QUESTIONS

1. What is effect of catalyst on equilibrium constant 'Kc'?

Ans . A catalyst does not affect equilibrium constant because it speeds up both forward and backward reactions to the same extent.

- State Le Chatelier'r principle. Ans.It states that a change in any of the factors thatdetermine the equilibrium conditions of asystem will cause the system to change such a manner so as to reduce or tocounteract the effect of the change.
- 3. What is meant by conjugate acid –base pairs? Explain.

Ans:- H_2O + $HCl \rightleftharpoons H_3O^+$ + Clbase acid conjugate acid conjugate base

- Classify the following bases as strong and weak bases: NaHCO₃, NaOH, KOH, Ca(OH)₂, Mg(OH)₂. Ans:-strong base NaOH, KOH ; weak bases NaHCO₃,Ca(OH)₂, Mg(OH)₂.
- 5. The concentration of hydrogen ion in a sample of soft drink is 3.8×10^{-3} M. What is its pH ? Ans:-pH = $-\log[3.8 \times 10^{-3}]$ = $-\{\log[3.8] + \log[10^{-3}]\}$ = $-\{(0.58) + (-3.0)\} = -\{-2.42\} = 2.42$ Therefore, the pH of the soft drink is 2.42and it is acidic.
- 6. The species: H₂O, HCO₃⁻, HSO₄⁻ and NH₃can act both as Bronsted acids and bases.For each case give the corresponding conjugate acid and conjugate base. Ans:-

Species	Conjugate acid	Conjugate base
H ₂ O	H_3O^+	OH ⁻
HCO ₃ ⁻	H_2CO_3	CO_{3}^{2}
HSO_4^-	H_2SO_4	SO_4^{2-}
NH ₃	$\mathrm{NH_4}^+$	NH ₂

 Explain Lewis acids and bases with suitable examples. Ans:-Lewis acids are lone pair (of e-) accepters while Lewis bases are lone pair donators.

AlCl₃ is a Lewis acid while NH₃ is a Lewis base.

- 8. What is difference between alkali and bases? Give examples. Ans:- An alkali is a water soluble base. All the alkalis are bases but all the bases are not alkali.
 Ex- NaOH is an alkali/base. Ca(OH)₂ is a base but not an alkali.
- 9. Explain homogeneous and heterogeneous equilibrium giving examples.

Ans:- If all the reactants and products present in an equilibrium mixture are in same phase→homogeneous equilibrium.

If all the reactants and products present in an equilibrium mixture are in different phase \rightarrow heterogeneous equilibrium.

 $N_{2(g)} + 3H_{2(g)} \rightleftharpoons 2NH_{3(g)}$ homogeneous equilibrium

 $CaCO_{3(s)} \rightleftharpoons CaO_{(s)} + CO_{2(g)}$ heterogeneous equilibrium

THREE MARK QUESTIONS

1. The pH of some common substances is given bellow. Classify the substances as acidic/basic

Name of fluid	pH
Lime water	10
Milk of magnesia	10
Human saliva	6.4
Lemon juice	2.2
Sea water	7.8
Vinegar	3
milk	6.8

Ans .:- acidic-Human saliva, Lemon juice, milk, vinegar

Basic- Lime water, sea water, milk of magnesia.

2. Explain general characteristics of acids and bases.

Ans.:- Most of the acids taste sour.Acids are known to turn blue litmus paper into red and liberate dihydrogen on reacting with some metals.

Bases are known to turn red litmus paper blue, tastebitter and feel soapy.

3. Water is amphoteric in nature. Explain.

Ans.:- Water can react with acid as well as base

$H_2O + HCl \rightarrow H_3O^+$	$+C1^{-}$	water is basic
$H_2O + NH_3 \rightarrow OH^-$	$+ NH_4^+$	water is acidic

4. Describe the effect of :

a) addition of H₂

b) addition of CH ₃ OH	
c) removal of CO	
d) removal of CH ₃ OH	
on the equilibrium of the r	reaction:
$2H_{2(g)} + CO_{(g)} \rightleftharpoons CH_3OH_{(g)}$	
Ans.:- a) addition of H_2	equilibrium will shift on H

$Ans.:-a$) addition of H_2	equilibrium will shift on RHS
b) addition of CH ₃ OH	equilibrium will shift on LHS
c) removal of CO	equilibrium will shift on LHS
d) removal of CH ₃ OH	equilibrium will shift on RHS

5. Classify the following species into Lewisacids and Lewis bases and show how these act as such:

(a) $HO^{-}(b)F^{-}(c) H^{+}(d) BCl_{3}$

Solution

(a) Hydroxyl ion is a Lewis base as it candonate an electron lone pair (:OH⁻).

(b) Flouride ion acts as a Lewis base asit can donate any one of its fourelectron lone pairs.

(c) A proton is a Lewis acid as it canaccept a lone pair of electrons frombases like hydroxyl ion and fluorideion.

(d) BCl_3 acts as a Lewis acid as it canaccept a lone pair of electrons fromspecies like ammonia or aminemolecules.

6. For the equilibrium,2NOCl(g) \rightleftharpoons 2NO(g) + Cl₂(g)the value of the equilibrium constant, *Kc* is 3.75 × 10⁻⁶ at 1069 K. Calculate the *Kp* for the reaction at this temperature?

Solution

We know that, $Kp = Kc(RT)^{\Delta n}$ For the above reaction, $\Delta n = (2+1) - 2 = 1$ $Kp = 3.75 \times 10^{-6} (0.0831 \times 1069)$ Kp = 0.033.

7. Hydrolysis of sucrose gives, Sucrose + H₂O \rightarrow Glucose + Fructose Equilibrium constant *Kc* for the reaction is 2 ×10¹³ at 300K. Calculate ΔG^0 at 300K.

Solution

$$\Delta G^{0} = -RT \ln Kc$$

$$\Delta G^{0} = -8.314 \text{J} \text{ mol}^{-1} \text{K}^{-1} \text{J} \text{ x } 300 \text{K} \times \ln(2 \times 10^{13})$$

$$\Delta G^{0} = -7.64 \times 10^{4} \text{ J} \text{ mol}^{-1}$$

8. Explain the following :

(i) Common ion effect (ii) solubility products (iii) pH

Ans. (i) Suppression of ionization of weak electrolyte by adding a strong electrolyte having an ion common.

(ii) Product of the molar concentrations of the ions in a saturated solution, each concentration term raised to the power equal to the no. of ions produced.

(iii) Negative logarithm of hydrogen ion concentration.

9. The values of Ksp of two sparingly solublesalts Ni(OH)₂ and AgCN are 2.0 $\times 10^{-15}$ and 6×10^{-17} respectively. Which salt ismore soluble? Explain.

Solution

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AgCN⇒Ag<sup>+</sup> + CN<sup>-</sup>

Ksp = [Ag<sup>+</sup>][CN<sup>-</sup>] = 6 × 10<sup>-17</sup>

Ni(OH)<sub>2</sub>⇒Ni<sup>2+</sup> + 2OH<sup>-</sup>

Ksp = [Ni<sup>2+</sup>][OH<sup>-</sup>]<sup>2</sup> = 2 × 10<sup>-15</sup>

Let [Ag<sup>+</sup>] = S<sub>1</sub>, then [CN-] = S<sub>1</sub>

Let [Ni<sup>2+</sup>] = S<sub>2</sub>, then [OH<sup>-</sup>] = 2S<sub>2</sub>

S<sub>1</sub><sup>2</sup> = 6 × 10<sup>-17</sup>, S<sub>1</sub> = 7.8 × 10<sup>-9</sup>

(S<sub>2</sub>)(2S<sub>2</sub>)<sup>2</sup> = 2 × 10<sup>-15</sup>, S<sub>2</sub> = 0.58 × 10<sup>-4</sup>

Ni(OH)<sub>2</sub> is more soluble than AgCN.
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FIVE MARKS QUESTIONS

1. At 473 K, equilibrium constant *Kc* for decomposition of phosphorus pentachloride, PCl₅ is 8.3×10^{-3} . If decomposition is depicted as,

PCl₅ (g) \rightleftharpoons PCl₃ (g) + Cl₂ (g) $\Delta r H^0 = 124.0 \text{ kJ mol}^{-1}$ a) Write an expression for *Kc* for the reaction.

b) What is the value of *Kc* for the reverse reaction at the same temperature?

c)what would be the effect on Kc if (i) more PCl₅ is added (ii) pressure is increased(iii) the temperature is increased ?

Ans: (a) $Kc=[PCl_3][Cl_2]$ [PCl₅]

(b)120.48

- (c) (i) equilibrium will shift on RHS
- (ii) equilibrium will shift on LHS
- (iii) equilibrium will shift on RHS
- 2. Dihydrogen gas is obtained from natural gas by partial oxidation with steam asper following endothermic reaction:CH₄ (g) + H₂O (g) ≓CO (g) + 3H₂ (g) (a) Write as expression for *Kp* for the above reaction.

(b) How will the values of *Kp* and composition of equilibrium mixture be affectedby(i) increasing the pressure(ii) increasing the temperature(iii) using a catalyst?

Ans. (a) $K_p = p(CO).p(H_2)^3 / p(CH_4).p(H_2O)$

(b)(i) value of Kp will not change, equilibrium will shift in backward direction.

(ii) Value of Kp will increase and reaction will proceed in forward direction.

(iii)no effect.

3. What is meant by the conjugate acid-base pair? Find the conjugate acid/basefor the following species:HNO₂, CN⁻, HClO₄, F⁻, OH⁻, CO₃²⁻, and S²⁻

Ans.The acid-base pair thatdiffers only by one proton is called a conjugate acid-base pair

	1
Species	Conjugate
	acid/base
HNO ₂	NO ₂ ⁻
CN^{-}	HCN
HClO ₄	ClO ₄ ⁻
F ⁻	HF
OH ⁻	H2O
CO_{3}^{2}	HCO_3^{2-}



1. The value of *Kc* for the reaction $2A \rightleftharpoons B + C$ is 2×10^{-3} . At a given time, the composition of reaction mixture is $[A] = [B] = [C] = 3 \times 10^{-4}$ M. In which direction the reaction will proceed?

Solution

For the reaction the reaction quotient Qc is given by, $Qc = [B][C]/[A]^2$

as $[A] = [B] = [C] = 3 \times 10^{-4} M$

 $Qc = (3 \times 10^{-4})(3 \times 10^{-4}) / (3 \times 10^{-4})2 = 1$

asQc>Kcso the reaction will proceed in the reverse direction.

2. PCl₅, PCl₃ and Cl₂ are at equilibrium at500 K and having concentration 1.59M PCl₃, 1.59M Cl₂ and 1.41 M PCl₅. Calculate *Kc* for the reaction, PCl₅ \rightleftharpoons PCl₃ + Cl₂ **Solution**

The equilibrium constant *Kc* for the abovereaction can be written as,

$$Kc = [PCl_3][Cl_2]$$
$$[PCl_5]$$

 $=(1.59)^2/1.41=1.79$

3. Why is ammonia termed as a base though it does not contain OH⁻ ions? Ans.ammona is termed as a base on the basis of Lewis concept it can donate a lone pair of electrons.

<u>CHAPTER-8</u> <u>REDOX REACTIONS</u>

oxidation	<u>reduction</u>
1. Addition of oxygen	1. Removal of oxygen
2. Removal of hydrogen	2. Addition of hydrogen
3. Addition of an electronegative	3. Removal of an electronegative
element	element
4. Removal of an electropositive	4. Addition of an electropositive
element	element
5. Loss of electron	5. Gain of electron

Oxidation number denotes theoxidation state of an element in a compound ascertained according to a set of rules formulated on the basis that electron in a covalent bond belongsentirely to more electronegative element.

Calculation of oxidation number-

- 1. O. S. of all the elements in their elemental form (in standard state) is taken as zero O. S. of elements in Cl₂, F₂, O₂, P₄, O₃, Fe(s), H₂, N₂, C(graphite) is zero.
- 2. Common O. S. of elements of group one (1st) is one. Common O. S. of elements of group two (2nd) is two.
- 3. For ions composed of only one atom, theoxidation number is equal to the chargeon the ion.
- The oxidation number of oxygen in most compounds is −2. While in peroxides (e.g., H₂O₂, Na₂O₂), eachoxygen atom is assigned an oxidationnumber of −1, in superoxides (e.g., KO₂, RbO₂) each oxygen atom is assigned anoxidation number of −(¹/₂).
- 5. In oxygendifluoride (OF₂) and dioxygendifluoride (O₂F₂), the oxygen is assigned no oxidation number of +2 and +1, respectively.
- 6. The oxidation number of hydrogen is +1 but in metal hydride its oxidation no. is-1.
- 7. In all its compounds, fluorine has an idation number of -1.
- 8. The algebraic sum of the oxidation number of all the atoms in a compound must bezero.
- 9. In polyatomic ion, the algebraic sumof all the oxidation numbers of atoms of the ion must equal the charge on the ion.

Stocknotation:the oxidationnumber is expressed by putting a Romannumeral representing the oxidation numberin parenthesis after the symbol of the metal inthe molecular formula. Thus aurous chlorideand auric chloride are written as Au(I)Cl andAu(III)Cl₃. Similarly, stannous chloride andstannic chloride are written as Sn(II)Cl₂andSn(IV)Cl₄.

Oxidation: An increase in the oxidationnumber *Reduction:* A decrease in the oxidationnumber

Oxidising agent: A reagent which canincrease the oxidation number of an elementin a given substance. These reagents are calledas **oxidants** also. *Reducing agent:* A reagent which lowers the oxidation number of an element in a given substance. These reagents are also called as **reductants**.

Redox reactions: Reactions which involvechange in oxidation number of the interactingspecies

Balancing of redox reactions:

Oxidation Number Method:

Write the net ionic equation for the reaction of potassium dichromate(VI), $K_2Cr_2O_7$ with sodium sulphite,Na2SO3, in an acid solution to give chromium(III) ion and the sulphate ion.

Step 1: The skeletal ionic equation is:

 $Cr_{2}O_{7}^{2-(aq)} + SO_{3}^{2-(aq)} \rightarrow Cr^{3+}(aq) + SO_{4}^{2-}(aq)$ Step 2: Assign oxidation numbers for Cr and S +6 -2 +4 -2 +3 +6 -2 Cr₂O₇²⁻⁽aq) + SO₃²⁻(aq) → Cr³⁺(aq)+ SO₄²⁻(aq)

Step 3: Calculate the increase anddecrease of oxidation number, and make them equal:

$$+6-2$$
 $+4-2$ $+3$ $+6$
 $Cr_2O_7^{2-(aq)} + 3SO_3^{2-}(aq) \rightarrow 2Cr^{3+}(aq) + 3SO_4^{2-}(aq)$

Step 4: Balance the charge by adding H⁺as the reaction occurs in theacidic medium,

$$\operatorname{Cr}_{2}\operatorname{O}_{7}^{2-(aq)} + 3\operatorname{SO}_{3}^{2-(aq)} 8\operatorname{H}^{+} \rightarrow 2\operatorname{Cr}^{3+}(aq) + 3\operatorname{SO}_{4}^{2-(aq)}$$

Step 5: Balance the oxygen atom by adding water molecule.

 $Cr_2O_7^{2-(aq)} + 3SO_3^{2-(aq)} 8H^+ \rightarrow 2Cr^{3+(aq)} + 3SO_4^{2-(aq)} + 4H_2O(1)$

Half Reaction Method

balance the equation showing the oxidation of Fe²⁺ ions to Fe³⁺ ions by dichromate ions $(Cr_2O_7)^{2-}$ in acidic medium, wherein, $Cr_2O_7^{2-}$ ions are reduced to Cr^{3+} ions.

Step 1: Produce unbalanced equation for thereaction in ionic form :

$$Fe^{2+(aq)} + Cr_2O_7^{2-}(aq) \rightarrow Fe^{3+}(aq) + Cr^{3+(aq)}$$

Step 2: Separate the equation into halfreactions:

+2 +3Oxidation half : Fe²⁺ (aq) \rightarrow Fe³⁺⁽aq) +6-2+3

Reduction half : $\operatorname{Cr}_2\operatorname{O}_7^{2-}(\operatorname{aq}) \to \operatorname{Cr}^{3+}(\operatorname{aq})$

Step 3: Balance the atoms other than O andH in each half reaction individually.

$$\operatorname{Cr}_2\operatorname{O_7}^{2-}(\operatorname{aq}) \to \operatorname{Cr}^{3+}(\operatorname{aq})$$

Step 4: For reactions occurring in acidicmedium, add H₂O to balance O atoms and H⁺to balance H atoms.Cr₂O₇²⁻ (aq) +14 H⁺ \rightarrow Cr³⁺⁽aq) + 7H²O (l) **Step 5:** Add electrons to one side of the halfreaction to balance the charges. If need be,make the number of electrons equal in the twohalf reactions by multiplying one or both halfreactions by appropriate coefficients. Fe²⁺ (aq) \rightarrow Fe³⁺ (aq) + e– Cr₂O₇²⁻ (aq) + 14H⁺ (aq) + 6e– \rightarrow 2Cr³⁺⁽aq) +7H₂O (l) 6Fe²⁺ (aq) \rightarrow 6 Fe³⁺ (aq) +6 e– **Step 6:** We add the two half reactions toachieve the overall reaction and cancel theelectrons on each side. This gives the net ionicequation as : 6Fe²⁺⁽aq) + Cr₂O₇²⁻⁽aq) + 14H⁺⁽aq) \rightarrow 6 Fe³⁺⁽aq) +2Cr³⁺⁽aq) + 7H₂O(l) A **redox couple** is defined as havingtogether the oxidised and reduced forms of asubstance taking part in an oxidation orreduction half reaction. Represented as Zn^{2+/}Zn and Cu²⁺/Cu.

 Electrochemical cells are the devices which are used to get electric current by using chemical reaction.



The potential associated with eachelectrode is known as **electrode potential**. If the concentration of each species taking part in the electrode reaction is unity (if any gasappears in the electrode reaction, it is confined to 1 atmospheric pressure) and further thereaction is carried out at 298K, then the potential of each electrode is said to be the **Standard Electrode Potential**.

• SHE is used to measure electrode potential and its standard electrode potential is taken as 0.00 V.

ONE MARK QUESTIONS

- 1. Define oxidation and reduction in terms of oxidation number. Ans Increase in Oxidation Number is Oxidation and decrease in Oxidation Number is called reduction.
- 2. What is meant by disproportionation? Give one example. Ans : In a disproportionation reaction an element simultaneously oxidized and reduced.

 $P_4 + 3OH^- + 3H_2O \rightarrow PH_3 + 3H_2PO_2^-$

3. What is O.N. of sulphur in H_2SO_4 ?Ans: +6

- 4. Identify the central atom in the following and predict their O.S. HNO₃
 Ans: central atom:- N; O.S. +5
- 5. Out of Zn and Cu which is more reactive? Ans: Zn.
- 6. What is galvanization? Ans: Coating of a less reactive metal with a more reactive metal e.g.- coating of iron surface with Zn to prevent rusting of iron.
- 7. How is standard cell potential calculated using standard electrode potential? Ans: $E^{0}_{cell} = E^{0}_{cathode} - E^{0}_{anode}$
- 8. What is O.S. of oxygen in H₂O₂? Ans: - -1.
- 9. The formation of sodium chloride from gaseous sodium and gaseous chloride is a redox process justify.

Ans: Na atom get oxidize and Cl is reduced.

TWO MARKS QUESTIONS

- 1. Write the balanced redox reaction .
 - (I) $\operatorname{MnO_4^-(aq)} + \operatorname{Fe}^{2+}(aq) \to \operatorname{Mn}^{2+}(aq) + \operatorname{Fe}^{3+}(aq)$ [acidic medium]
 - (II) $\operatorname{Cr}_2 \operatorname{O}_7^{2-} + \operatorname{Fe}^{2+} \rightarrow \operatorname{Cr}^{3+} + \operatorname{Fe}^{3+}$ [Acidic medium]

Ans:- (i) $MnO_{4}^{-}(aq) + 5Fe^{2+}(aq) + 8H^{+}(aq) \rightarrow Mn^{2+}(aq) + 5Fe^{3+}(aq) + 4H_{2}O_{(1)}$ (ii) $Cr_{2}O_{7}^{2-} + 6Fe^{2+} + 14H^{+} \rightarrow 2Cr^{3+} + 6Fe^{3+} + 7H_{2}O$

2. Identify the strongest & weakest reducing agent from the following metals: .Zn, Cu, Na, Ag, Sn

Ans: Strongest reducing agent: Na, weakest reducing agent: Ag.

3. Determine the oxidation no. of all the atoms in the following oxidants: $KMnO_{4,}$ $K_2Cr_2O_7$ and $KClO_4$

Ans :

In KMnO₄ K = +1, Mn = +7, O = -2In $K_2Cr_2O_7K = +1$, Cr = +6, O = -2In KClO₄K = +1, Cl = =+7, O = -2

4. Determine the oxidation no. of all the atoms in the following species: Na_2O_2 and OF_2 .

Ans: In Na₂O₂Na = +1, O = -1 InOF₂, F = -1, O = +2

5. Is it possible to store :

(i) H₂SO₄ in Al container?(ii) CuSO4 solution in Zn vessel? Ans : (i) yes. (ii) No.

- 6. Calculate the standard e.m.f. of the cell formed by the combination of $Zn/Zn^{2+} \prod Cu^{2+}/Cu$. Solution- : $E^{0}_{cell} = E^{0}_{cathode} - E^{0}_{anode}$ =0.34 - (-0.76) = 1.10V.
- 7. Identify the oxidizing and reducing agents in the following equations:

(i) $MnO_{4}^{-}(aq) + 5Fe^{2+}(aq) + 8H^{+}(aq) \rightarrow Mn^{2+}(aq) + 5Fe^{3+}(aq) + 4H_{2}O_{(1)}$ (ii) $Cr_{2}O_{7}^{2-} + 6Fe^{2+} + 14H^{+} \rightarrow 2Cr^{3+} + 6Fe^{3+} + 7H_{2}O$ Ans : (i) O.A. = MnO_{4}^{-} ; R.A.= Fe^{2+}

(ii

i)O.A.=
$$Cr_2O_7^{2-}$$
; R.A.= Fe²⁺

- 8. Predict all the possible oxidation states of Cl in its compounds. Ans:- 0, -1, +1, +3, +5, +7
- 9. Formulate possible compounds of 'Cl' in its O.S.is: 0, -1, +1, +3, +5, +7Ans: Cl₂, HCl, HOCl, HOClO, HOClO₂, HOClO₃ respectively.
- 10. List three measures used to prevent rusting of iron.
 - Ans: (i) galvanization(coating iron by a more reactive metal)
 - (ii) greasing/oiling
 - (iii) painting.

THREE MARK QUESTIONS

1. Write short notes on :

(a) Electrochemical series(b) redox reactions (c) oxidizing agents Ans :(a) Electrochemical series :- arrangement of metals(non-metals also) in increasing order of their reducing power or vice versa.

(b) Reactions in which both Oxidation and reduction take place simultaneously are REDOX REACTIONS.

(c)oxidizing agents : chemical specie which can oxidize the other one or can reduce itself.

2. Calculate O. S. of sulphur in the following oxoacids of 'S' :

H₂SO₄,H₂SO₃H₂S₂O₈and H₂S₂O₇

Ans :+6, +4, +6 and +6 respectively.

(calculate by considering x of 'S' and taking +1 of H, -2 0f "O" and -1 of "O" in peroxide bond.)

3. Explain role of salt bridge in Daniell cell.

Ans : (a) it completes the electric circuit in the cell.

(b) it maintains the electric neutrality in the cell.

- 4. Account for the followings :
 - sulphur exhibits variable oxidation states. (i)
 - Ans. Due to the presence of vacant 'd' orbitals in 'S'

(ii) Fluorine exhibits only -1 O.S.

Ans. It is most electronegative element

(iii) oxygen can't extend its valency from 2.

Ans. Small size/unavailabilityof vacant 'd' orbitals in O

 $MnO_4^- + \Gamma \rightarrow Mn^{2+} + I_2 + H_2Oby$ ion electron 5. Balance the equation method in acidic medium.

Ans :Step-I Balancing of reduction half reaction by adding protons and electrons on LHS and more water molecules on RHS:

 $8H^+ + MnO_4^- + 5e^- \rightarrow Mn^{2+} + 4H_2O$

<u>Step-II</u> Balancing of oxidation half reaction by adding electrons on RHS: $2I \rightarrow I_2 + 2e^-$

Step-III To multiply the OHR by 5; RHR by2 andto add OH & RH reactions to get overall redox reaction(cancellation of electrons of RH & OH reactions):

 $[8H^+(aq)+MnO_4^-(aq)+5e \rightarrow Mn^{2+}(aq)+4H_2O(l)] \ge 2$

$$[2I^{-} \rightarrow I_2 + 2e^{-}] \quad x \quad 5$$

 $MnO_{4}^{-}(aq) + 5Fe^{2+}(aq) + 8H^{+}(aq) \rightarrow Mn^{2+}(aq) + 5Fe^{3+}(aq) + 4H_{2}O_{(1)}$

6. complete and balance the following equations:

(i)
$$H^+ + Cr_2O_7^{2-} + Br^- \rightarrow 2Cr^{3+} + Br_2 + \cdots$$

(ii)
$$H_2O_2 + Cl \rightarrow OH + Cl_2$$

(iii) $Zn + Cu^{2+} \rightarrow ?$

Ans :(i)
$$14H^+ + Cr_2O_7^{2-} + 6Br^- \rightarrow 2Cr^{3+} + 3Br_2 + 7H_2O$$

(ii) $H_2O_2 + 2Cl \rightarrow 2OH + Cl_2$

(ii) $Zn + Cu^{2+} \rightarrow Zn^{2+} + Cu$

- 7. Identify the oxidizing and reducing agents in the following equations:
 (i) Fe + H₂SO₄→FeSO₄ + H₂
 (ii)H₂ + Cl₂ →2HCl
 (iii) MnO₂ + 4HCl→MnCl₂ + 2H₂O + Cl₂
 Ans :(i) O.A. =H₂SO₄; R.A.= Fe
 (ii) O.A. = Cl₂; R.A.=H₂
 (iii)O.A. = MnO₂; R.A. =HCl
- 8. Arrange the following in increasing order of their reducing power:

Cu, Ag, Au, Zn, Fe, Al, Na, Mg, Pt(SHE), Hg, Ca, K

Ans : Au, Hg, Ag, Cu, Pt(SHE), Fe, Zn, Al, Mg, Na, Ca, K

^{9.} Indicate O.S. of each atom present in given structure of peroxodisulphuric acid





10. What is SHE? What is its use?

Ans :Standard Hydrogen Electrode (SHE) has beenselected to have zero standard potential at alltemperatures. It consists of a platinum foilcoated with platinum black (finely divided platinum) dipping partially into an aqueous solution in which the activity (approximate concentration 1M) of hydrogen ion is unity and hydrogen gas is bubbled through the solutionat 1 bar pressure. The potential of the other half cell is measured by constructing a cell in which reference electrode is standard hydrogen electrode. The potential of the other half cell is equal to the potential of the cell.



Fig: SHE

HOTS QUESTIONS

1. Is rusting of iron an electrochemical phenomenon? How ?explain.

Ans : Yes. Rusting of iron is an electrochemical phenomenon because this is possible due to formation of a small electrochemical cell over rough surface of iron and the following redox reaction takes place there in that cell-



- Fig. 5.14 Corrosion of iron in atmosphere.
- 2. We expand croreof Rupees and even thousands of lives every year due to corrosion. How can be preventing it. Explain.

Ans : (i) By Galvanization: Coating of a less reactive metal with a more reactive metal e.g.- coating of iron surface with Zn to prevent rusting of iron.

(ii) By greasing /oiling (to keep away the object from the contact of air & moisture.)

(iii)By painting (to keep away the object from the contact of air & moisture.)
<u>Hydrogen</u>

(1) Position of hydrogen in the periodic table

Hydrogen is the first element in the periodic table. Hydrogen is placed in no specific group due to its property of giving electron (When H^- is formed) and also losing electron (When H^+ is formed).

(i) Hydrogen is placed in group I (Alkali metals) as,

(a) It has one electron in its (Outer) shell- $1s^1$ like other alkali metals which have (inert gas) ns^1 configuration.

(b) It forms monovalent H^+ ion like Li^+ , Na^+ ...

(c) Its valency is also 1.

(d) Its oxide (H_2O) is stable as Li_2O , Na_2O .

(e) It is a good reducing agent (In atomic as well as molecular state) like *Na*, *Li*...

(ii) Hydrogen also resembles halogens (Group VII A) as,

(a) It is also diatomic (H_2) like $F_2, Cl_2 \dots$

(b) It also forms anion H^- like F^- , CF^- ... by gain of one electron.

(c) H^- has stable inert gas (*He*) configuration as CH_4, C_2H_6 like halogens CCI_4, SF_2CI_2 etc.

(d) *H* is one electron short of duplet (Stable configuration) like *F*, *Cl*,... which are also one electron deficient than octet, $F-2s^22p^5$; $Cl-3s^23p^5$.

(e) (*IE*) of $H(1312 \text{ kJ mol}^{-1})$ is of the same order as that of halogens.

(iii) (*IE*) of *H* is very high in comparison with alkali metals. Also size of H^+ is very small compared to that of alkali metal ion. *H* forms stable hydride only with strongly electropositive metals due to smaller value of its electron affinity (72.8 kJ mol⁻¹).

(iv) In view of the anomalous behaviour of hydrogen, it is difficult to assign any definite position to it in the periodic table. Hence it is customary to place it in group I (Along with alkali metals) as well as in group VII (Along with halogens).

(2) **Discovery and occurrence :** It was discovered by *Henry Cavendish* in 1766. Its name hydrogen was proposed by *Lavoisier*. Hydrogen is the 9th most abundant element in the earth's crust.

Hydrogen exists in diatomic state but in triatomicstate it is called as Hyzone. Systematic name of water is oxidane.

(3) **Preparation of Dihydrogen :** Dihydrogen can be prepared by the following methods,

(i) By action of water with metals

(a) Active metals like *Na*, *K* react at room temperature

 $2M+2H_2O \rightarrow 2MOH+H_2$ [*M* = *Na*, *K* etc.]

(b) Less active metals like *Ca*, *Zn*, *Mg*, *Al* liberate hydrogen only on heating.

 $2A/+3H_2O {\rightarrow} A/_2O_3+3H_2$

(c) Metals like *Fe, Ni, Co, Sn* can react only when steam is passed over red hot metals.

$$3 Fe+4 H_2 O(\text{steam}) \rightarrow Fe_3 O_4 + 4 H_2$$

Ferrosoferric oxide

(ii) By the action of water on alkali and alkaline earth metals hydrides

 $NaH + H_2 O \rightarrow NaOH + H_2$

 $CaH_2 + 2H_2O \rightarrow Ca(OH)_2 + 2H_2$

(iii) By reaction of metals like Zn, Sn, Al with alkalies (NaOH or KOH)

$$Zn + 2 NaOH \xrightarrow{\Delta} Na_2 ZnO_2 + H_2$$

sod. zincate
$$A/+ 2 NaOH + H_2O \xrightarrow{\Delta} 2 NaA/Q_2 + 2H_2$$

Sod. meta-aluminate
$$Si + 2 NaOH + 2H_2O \xrightarrow{\Delta} Na_2 SiO_3 + 3H_2$$

Silicon
$$Sn + 2 NaOH \xrightarrow{\Delta} Na_2 SnO_2 + H_2 \uparrow$$

Sod. stannite

(iv) By action of metal with acids : All active metals which lie above hydrogen in electrochemical series, can displace hydrogen gas from dilute mineral acids like HCl, H_2SO_4 .

 $\textit{Fe+2HCl} \rightarrow \textit{FeCl}_2 + \textit{H}_2$

(v) By the electrolysis of acidified water

 $2 H_2 O \xrightarrow{H^+ / \text{Electrolyis}} 2 H_2 \uparrow + O_2 \uparrow \\ \underset{\text{At cathode}}{\text{At anode}} 2 H_2 \uparrow + O_2 \uparrow$

(vi) Laboratory method : In laboratory, it is obtained by action of granulated zinc with dilute H_2SO_4 .

 $Zn + dil.H_2SO_4 \rightarrow ZnSO_4 + H_2$

It must be noted that

(a) Pure zinc is not used for the preparation of H_2 as rate of reaction of pure Zn with dil. H_2SO_4 is quite slow.

(b) Conc. H_2SO_4 is not used because then SO_2 gas is evolved instead of H_2 .

(vii) *Preparation of pure hydrogen*: It can be obtained by

(a) The action of pure dil. H_2SO_4 on pure magnesium ribbon.

 $Mg + H_2SO_4 \rightarrow MgSO_4 + H_2$

(b) Hydrogen of high purity (> 99.95%) is obtained by electrolysing warm aqueous barium hydroxide between nickel electrodes.

> (c) By the action of water on sodium hydride. $NaH+H_2O \rightarrow NaOH+H_2\uparrow$

(d) By the action of KOH (aq.) on aluminium.

 $2\textit{AI}+2\textit{KOH}+2\textit{H}_{2}\textit{O} \!\rightarrow\! 2\textit{KAIO}_{2}+3\textit{H}_{2} \uparrow$

(viii) Commercial production of dihydrogen

(a) **Bosch process :** In this method, water gas is mixed with twice its volume of steam and passed over heated catalyst $F_{P_2}O_3$ in the presence of a promoter $C_{T_2}O_3$ or ThO_2 at 773 K when CO_2 and H_2 are obtained. CO_2 is removed by dissolving it in

water under pressure (20-25 *atm*) and H_2 left undissolved is collected.

$$C + H_2 O \xrightarrow{1270 \text{ K}} CO + H_2$$

Watergas
$$H_2 + CO + H_2 O \xrightarrow{773 \text{ K}}_{Fe_2O_3, C_2O_3} CO_2 + 2H_2$$

About 18% of the world's production of H_2 is obtained from coal.

(b) Lane's process : By passing steam over spongy iron at 773-1050 K.

 $3Fe+4H_2O \rightarrow Fe_3O_4+4H_2$

The ferrosoferric oxide (Fe_3O_4) so produced is reduced back to iron with water. this reaction is known as **Vivification reactions**

$$Fe_{3}O_{4} + 4H_{2} \rightarrow 3Fe + 4H_{2}O$$
$$Fe_{3}O_{4} + 4CO \rightarrow 3Fe + 4CO_{2}$$

(c) **By electrolysis of water** : Electrolysis of acidified water using platinum electrodes is used for the bulk preparation of hydrogen.

(d) **From hydrocarbons** : Hydrocarbons (alkanes) react with steam at high temperature to produce carbon monoxide and hydrogen, *e.g.*,

$$CH_4(g) + H_2O(g) \xrightarrow[\text{Catalyst}]{1270 \text{ K}} CO(g) + 3H_2(g)$$

The mixture of *CO* and H_2 so obtained can be converted into hydrogen as in Bosch process. About 77% of the world's production of H_2 is obtained from hydrocarbons.

(e) It is also produced as a by-product of the brine electrolysis process for the manufacture of Cl_2 and NaOH.

(4) **Physical properties of dihydrogen**: It is a colourless, tasteless and odourless gas. It is slightly soluble in water. It is highly combustible. The Physical constants of atomic hydrogen are,

Atomic radius (pm) – 37 Ionic radius of H^- ion (pm) – 210 Ionisation energy $(k J mol^{-1}) - 1312$ Electron affinity $(k J mol^{-1}) - 72.8$ Electronegativity - 2.1

(5) Chemical properties of dihydrogen : Dihydrogen is quite stable and dissociates into hydrogen atoms only when heated above 2000 K, $H_2 \xrightarrow{2000 \ \ } H + H$. Its bond dissociation energy is very high, $H_2 \rightarrow H + H$; $\Delta H = 435.9 \ \ \ Mol^{-1}$. Due to its high bond dissociation energy, it is not very reactive. However, it combines with many elements or compounds.

(i) Action with metals : To forms corresponding hydrides. $2Na + H_2 \xrightarrow{Heat} 2NaH$; $Ca + H_2 \xrightarrow{Heat} CaH_2$.

With transition metals (elements of d – block) such as *Pd*, *Ni*, *Pt* etc. dihydrogen forms interstitial hydrides in which the small molecules of dihydrogen occupy the interstitial sites in the crystal lattices of these hydrides. As a result of formation of interstitial hydrides, these metals adsorb large volume of hydrogen on their surface. This property of adsorption of a gas by a metal is called **occlusion**. The occluded hydrogen can be liberated from the metals by strong heating.

(ii) Reaction with Non-metals

$$2H_{2} + O_{2} \xrightarrow{970 \text{ K}} 2H_{2}O$$

$$N_{2} + 3H_{2} \xrightarrow{Fe,Mo} 2NH_{3}$$

$$H_{2} + F_{2} \xrightarrow{Dark} 2HF$$

$$H_{2} + CI_{2} \xrightarrow{Sunlight} (Fressure) 2HCI$$

$$H_{2} + BI_{2} \rightarrow 2HBr$$

$$H_{2} + I_{2} \xrightarrow{673 \text{ K}} 2HI$$

The reactivity of halogen towards dihydrogen decreases as, $F_2 > Cl_2 > Br_2 > l_2$

As a result, F_2 reacts in dark, Cl_2 in the presence of sunlight, Br_2 reacts only upon heating while the reaction with l_2 occurs in the presence of a catalyst.

(iii) **Reaction with unsaturated hydrocarbons** : H_2 reacts with unsaturated hydrocarbons such as ethylene and acetylene to give saturated hydrocarbons.

$$\begin{array}{l} H_{2}C = CH_{2} + H_{2} \xrightarrow{Nior Ptor Pd} CH_{3} - CH_{3} \\ Ethylene \end{array}$$

$$\begin{array}{l} HC \equiv CH + 2H_{2} \xrightarrow{Nior Ptor Pd} CH_{3} - CH_{3} \\ Acetylene \end{array}$$

This reaction is used in the *hydrogenation or hardening of oils*. The vegetable oils such as groundnut oil or cotton-seed oil are unsaturated in nature because they contain at least one double bond in their molecules. Dihydrogen is passed through the oils at about 473 *K* in the presence of catalyst to form solid fats. The vegetable ghee such as Dalda, Rath, etc. are usually prepared by this process.

Vegetableoil+
$$H_2 \xrightarrow[(liquid)]{Ni} Fat$$

(liquid)

- (6) Uses of Dihydrogen
- (i) As a reducing agent
- (ii) In the hydrogenation of vegetable oils
- (iii) As a rocket fuel in the form of liquid H_2
- (iv) In the manufacture of synthetic petrol
- (v) In the preparation of many compounds
- a) Synthesis of Ammonia by Haber's process.

$$N_{2(g)} + 3H_{2(g)} \Longrightarrow 2 NH_{3(g)}$$

b) Preparation of HCl

$$H_{2(g)} + Cl_{2(g)} \rightarrow 2HCl_{(g)} \xrightarrow{water} 2HCl_{(aq)} \xrightarrow{acid}$$

c) Synthesis of Methyl alcohol:

$$\begin{array}{c} H_2 + \begin{pmatrix} CO + H_2 \end{pmatrix} \xrightarrow{\text{2nO.CrO}_3} CH_3OH \\ \text{water gas} & (\text{catalyst}) \end{array}$$

d) In the production of vanaspathi or Margarine:

(vi) It is used in the oxy-hydrogen torch for welding if temperature around 2500°C is required. It is also used in atomic hydrogen torch for welding purposes in which temperature of the order of 4000°C is required.

vii) Synthetic petrol is prepared by Fischer-Tropsch process. Here Iron oxide is used to remove sulphur from a mixture of water gas and Hydrogen.

Different forms of hydrogen

(1) Atomic hydrogen : It is obtained by the dissociation of hydrogen molecules. The atomic hydrogen is stable only for a fraction of a second and is extremely reactive. It is obtained by passing dihydrogen gas at atmospheric pressure through an electric arc struck between two tungsten rods.

The electric arc maintains a temperature around 4000 – 4500°C. As the molecules of dihydrogen gas pass through the electric arc, these absorb energy and get dissociated into atoms as

 $H_2(g) \xrightarrow{Electric} 2H(g): \Delta H = 435.90 \text{ KJ mol}^{-1}$

This arrangement is also called atomic hydrogen torch.



(2) **Nascent hydrogen :** The hydrogen gas prepared in the reaction mixture in contact with the substance with which it has to react, is called nascent hydrogen. It is also called newly born hydrogen. It is more reactive than ordinary hydrogen. For example, if ordinary hydrogen is passed through acidified $KMnO_4$ (pink in colour), its colour is not discharged. On the other hand, if zinc pieces are added to the same solution, bubbles of hydrogen rise through the solution and the colour is discharged due to the reduction on $KMnO_4$ by nascent hydrogen.

$$\begin{split} & \textit{KMnO}_4 + \underbrace{H_2}_{\textit{Molecular}} + \underbrace{H_2SO}_4 \rightarrow \textit{No reaction} \\ & \textit{Zn} + \underbrace{H_2SO}_4 \rightarrow \textit{ZnSO}_4 + \underbrace{2[H]}_{\textit{Nascent hydrogen}} \\ & 2\textit{KMnO}_4 + 3\textit{H}_2SO_4 + \underbrace{10[H]}_{\rightarrow} \textit{K}_2SO_4 + 2\textit{MnSO}_4 + 8\textit{H}_2O \end{split}$$

(3) **Ortho and para hydrogen** : A molecule of dihydrogen contains two atoms. The nuclei of both the atoms in each molecule of dihydrogen are spinning. Depending upon the direction of the spin of the nuclei, the hydrogen is of two types,



(i) Molecules of hydrogen in which the spins of both the nuclei are in the same directions, called ortho hydrogen.

(ii) Molecules of hydrogen in which the spins of both the nuclei are in the opposite directions, called para hydrogen.

Ordinary dihydrogen is an equilibrium mixture of ortho and para hydrogen. Ortho hydrogen \rightleftharpoons Para hydrogen. The amount of ortho and para hydrogen varies with temperature as,

(a) At $0^{\circ}K$, hydrogen contains mainly para hydrogen which is more stable.

(b) At the temperature of liquefaction of air, the ratio of ortho and para hydrogen is 1:1.

(c) At the room temperature, the ratio of ortho to para hydrogen is 3:1.

(d) Even at very high temperatures, the ratio of ortho to para hydrogen can never be more than 3:1.

Thus, it has been possible to get pure para hydrogen by cooling ordinary hydrogen gas to a very low temperature (close to 20 *K*) but it is never possible to get a sample of hydrogen containing more than 75% of ortho hydrogen. i.e., Pure ortho hydrogen can not be obtained.

(4) **Hydrides :** Hydrogen forms binary hydrides of the type MH_x or M_mH_n with

(a) All main group elements except noble gases and probably indium and thallium.

(b) All lanthanoids and actinoids.

(c) Transition metals (*Sc, Y, La, Ac, Tc, Zr, Hf* and to a lesser extent *V, Nb, Ta, Cr, Cu* and *Zn*). In group 6 only *Cr* forms hydride (*CrH*).

Hydrides are classified into three main categories.

(i) Saline or ionic hydrides : Most of the s-block metals form this type of hydrides. These are non-volatile, non-conducting crystalline solids. However, BeH_2 and MgH_2 have covalent polymeric structure. These ionic hydrides have rock-salt structure. Thermal stability of 1st and 2nd group hydrides are in the order;

LiH > NaH > KH > RbH > CsH CaH₂ > SrH₂ > BaH₂

 BeH_2 , MgH_2 and *LiH* have significant covalent character.

Electrolysis of solution of saline hydride in molten alkali halide produces H_2 at anode. Saline hydrides react explosively with water.

 $NaH(s) + H_2O(aq) \rightarrow NaOH(aq) + H_2(g)$

The fire so produced cannot be extinguished by CO_2 as it gets reduced by the hot metal hydride. Only sand is useful, as it is a solid.

Alkali metal hydrides are used for making *LiAIH*, *NaBH*₄ etc. Alkali metal hydrides are also used for the removal of last traces of water from organic compounds.

(ii) *Metallic or interstitial hydrides* : Elements of groups 3, 4, 5 (*d*-block) and *f*-block elements form metallic hydrides. In group 6, only *Cr* forms hydride (*CrH*). Metals of group 7, 8, 9 do not form hydrides. This region of periodic table from group 7 to group 9 is known as hydride gap. Examples of hydrides of group 3 to 5 are, ScH_2 , YH_2 , YH_3 , LaH_2 , LaH_3 , TiH_2 , ZrH_2 , HfH_2 , VH,

VH2, NbH, NbH2, TaH

The *f*-block metals form hydrides of limiting compositions of MH_2 and MH_3 . All these hydrides are non-stoichiometric with variable composition *e.g.*,

 $ZrH_x(1.30 \le x \le 1.75)$, $TiH_x(1.8 \le x \le 2.0)$.

Most of these hydrides are good conductors of electricity in solid state.

Metallic hydrides can be used to store hydrogen especially in cars working on fuel cells.

(iii) *Molecular or covalent hydrides* : Hydrogen form molecular compounds with *p*-block elements (*B*, *C*, *N*, *O*, *F*; *Si*, *P*, *S*, *Cl*; *Ga*, *Ge*, *As*, *Sb*, *Br*; *In*, *Sn*, *Sb*, *Te*, *I*; *TI*, *Pb*, *At*). common examples of such hydrides are *CH*₄, *NH*₃, *H*₂*O*, *HF*etc. The stability of these

5

hydrides decreases down the group. For example, $NH_3 > PH_3 > ASH_3 > SbH_3 > BiH_3$. In a period the stability increases with increasing electronegativity. For example, $CH_4 < NH_3 < H_2O < HF$. Molecular hydrides are classified as electron rich, electron precise and electron deficient hydrides.

(a) *Electron rich molecular hydrides* : These hydrides have one or more lone pairs of electrons around the central more electronegative element. For example

$$\begin{array}{c} \vdots \\ H-O-H, H-N-H, H-F: \\ \vdots \\ H \end{array}$$

(b) Electron precise molecular hydrides : Elements of group 14 form such hydrides. The bond length increases on going down the group. A common example of electron precise molecular hydrides is CH_4 .

(c) Electron deficient molecular hydrides : These hydrides have lesser number of electrons than that required for writing the conventional Lewis structure. A common example of such molecular hydride is diborane, B_2H_6 .

(d) *Systematic names of molecular hydrides* : The systematic names of these hydrides are obtained from the name of the element and the suffix –ane. For example,

> PH_3 H_2O NH_3 Phosphane oxidane ozane

Isotopes of Hydrogen

Isotopes are the different forms of the same element, which have the same atomic number but different mass numbers.

Isotopes	of hydrogen
----------	-------------

Name	Sym	Ato	Mass	Relativ	Nature
	bol	mic	num	e	radioacti
		num	ber	abunda	ve or

		ber		nce	non-
					radioacti
					ve
Protiu	$^1_1\mathrm{H}~\mathrm{or}$	1	1	99.985	Non-
m or	Н			%	radioacti
Hydro					ve
gen					
Deute	2_1 H O	1	2	0.015%	Non-
rium	r D				radioacti
					ve
Tritiu	³ ₁ H O	1	3	10^{-15} %	Radioact
m	r T				ive

Physical constants of H_2 , D_2 and T_2

Property	H ₂	D ₂	T ₂
Molecular mass	2.016	4.028	6.03
Melting point (K)	13.8	18.7	20.63
Boiling point (K)	20.4	23.9	25.0
Heat of fusion (kJ mol ⁻¹)	0.117	0.197	0.250
Heat of vaporisation (kJ mol ⁻¹)	0.994	1.126	1.393
Bond energy (kJ mol ⁻¹)	435.9	443.4	446.9

Isotopic effect : In general chemical properties of isotopes are same but quantiative differences are noticed amongst them. For example, the reaction between H_2 and Cl_2 is 13.4 times faster between D_2 and Cl_2 under similar conditions. Such differences in chemical properties, which are due to difference in the mass numbers of isotopes is known as isotopic effect.

Water

Water is the oxide of hydrogen. It is an important component of animal and vegetable matter. Water constitutes about 65% of our body. It is the principal constituent of earth's surface.

(1) **Structure** : Due to the presence of lone pairs, the geometry of water is



distorted and the H-O-H bond angle is 104.5°, which is less than the normal tetrahedral angle (109.5°). The geometry of the molecule is regarded as angular or bent. In water, each O-H bond is polar because of the high electronegativity of oxygen (3.5) in comparison to that of hydrogen (2.1). The resultant dipole moment of water molecule is 1.84D.

In ice, each oxygen atom is tetrahedrally surrounded by four hydrogen atoms; **two by** covalent bonds and two by hydrogen bonds. The resulting structure of ice is open structure having a number of vacant spaces. Therefore, the density of ice is less than that of water and ice floats over water. It may be noted that water has maximum density $(1 g cm^{-3})$ at 4°C (277 K).

(2) Heavy water : Chemically heavy water is deuterium oxide (*D*₂*O*). It was discovered by *Urey*.

It is obtained as a by-product in some industries where \mathcal{H}_2 is produced by the electrolysis of water.

Heavy water (D_2O) is used (*a*) as a moderator and coolant in nuclear reactors (*b*) in the study of mechanism of chemical reactions (*c*) as a starting material for the preparation of a number of deuterium compounds, *e.g.*,

$SO_3 + D_2O \rightarrow D_{\text{Deuteric}}$	D ₂ SO ₄ osulphuricaci	d	
$A_{4}C_{3} + 12D_{2}O \rightarrow D_{D}$	$3CD_4$	+ ne	4 <i>A</i> /(<i>OD</i>) ₃
$CaC_2 + 2D_2O \rightarrow Deut$	$C_2 D_2$ terioacetylen	+ e	Ca(OD) ₂

(3) **Physical properties :** Water is colourless, odourless and tasteless liquid at ordinary temperature.

At 273*K* water is in equilibrium with ice and vapour this point is known triple point.

Some physical constants of H₂O and D₂O at 298 K

Constant	Ordinary	Heavy
	water H ₂ O	water D ₂ O

Molecular mass	18.015	20.028
Maximum density	1.000	1.106
(g cm ⁻³)		
Melting point (K)	273.2	276.8
Boiling point (K)	373.2	374.4
Heat of	6.01	6.28
fusion (kJ mol ⁻¹) at		
273 <i>K</i>		
Heat of vaporisation	40.66	41.61
(kJ mol ⁻¹) at 373K		
Heat of formation	- 285.9	- 294.6
(<i>kJ mol</i> ⁻¹)		
Ionisation constant	1.008×10^{-14}	1.95×10^{-15}

(4) Chemical properties : Water shows a versatile chemical behaviour. It behaves as an acid, a base, an oxidant, a reductant and as ligand to metals.

(i) **Dissociation of water** : Water is quite stable and does not dissociate into its elements even at high temperatures. Pure water has a small but measurable electrical conductivity and it dissociates as, $H_2O + H_2O \rightleftharpoons H_3O^+ + OH^-$

 $K_W = 1.0 \times 10^{-14} \text{ mol}^2 L^2 \text{ at } 298 \text{K}$

(ii) *Amphoteric nature*: Water can act both as an acid and a base and is said to be amphoteric.
However, water is neutral towards litmus and its *p*H is 7.

(iii) **Oxidising and reducing nature** : Water can act both as an oxidising and a reducing agent in its chemical reactions. e.g.

$$2 Na + 2H_2O \rightarrow 2 NaOH + H_2$$

Oxidisin g agent
$$2F_2 + 2H_2O \rightarrow 4 HF + O_2$$

Reducing agent

(iv) *Hydrolytic reactions* : Water can hydrolyse many oxides, halides, hydrides, carbides, nitrides,

phosphides, carbonates etc. to give an acid or a base or both as shown below :

$$SO_{2} + H_{2}O \rightarrow H_{2}SO_{3}$$

$$Mg_{3}N_{2} + 6H_{2}O \rightarrow 3Mg(OH)_{2} + 2NH_{3}$$

$$CaH_{2} + 2H_{2}O \rightarrow Ca(OH)_{2} + 2H_{2}$$

$$CaO + H_{2}O \rightarrow Ca(OH)_{2}$$

$$Na_{2}CO_{3} + 2H_{2}O \rightarrow 2NaOH + H_{2}CO_{3}$$

$$SiCl_{4} + 4H_{2}O \rightarrow Si(OH)_{4} + 4HCl$$

$$Ca_{3}P_{2} + 6H_{2}O \rightarrow 3Ca(OH)_{2} + 2PH_{3}$$

$$CaC_{2} + 2H_{2}O \rightarrow Ca(OH)_{2} + C_{2}H_{2}$$

(v) *Water forms hydrates with metal salts* : There are three main types of hydrates.

(a) Compounds in which water molecule are co-ordinated to the metal ion (complex compounds) $[N(OH_2)](NO_3)_2$, $Fe(OH_2)_6]CI_3$ etc.

(b) Compound in which water molecule may be hydrogen bonded to oxygen to form oxo-anion. For example in $CuSO_4.5H_2O$, 4 molecules of water are co-ordinated to Cu^{2+} while the fifth molecule is hydrogen bonded to SO_4^{2-} ion.

(c) In some compounds, water molecule occupies, interstitial sites in the crystal lattice *e.g.*, $BaCl_2.2H_2O$.

(5) Hard and Soft water

Water which produces lather with soap solution readily is called *soft water*. e.g. distilled water, rain water and demineralised water.

Water which does not produce lather with soap solution readily is called *hard water*. e.g. sea water, river water, well water and tap water.

(i) *Cause of hardness of water* : The hardness of water is due to the presence of bicarbonates, chlorides and sulphates of calcium and magnesium.

Hard water does not produce lather because the cations $(Ca^{+2} \text{ and } Mg^{+2})$ present in hard water react with soap to form insoluble precipitates, $M^{+2}_{From hard water} + 2C_{17}H_{35}COONa \rightarrow (C_{17}H_{35}COO)_2M + 2Na^+$ Sodium stearate(soap) Metalstearate(PPt)

Where *M* = *Ca* or *Mg*

Therefore, no lather is produced until all the calcium and magnesium ions are precipitated. This also results into wastage of lot of soap.

(ii) *Type of hardness of water* : The hardness of water is of two types,

(a) *Temporary hardness* : This is due to the presence of bicarbonates of calcium and magnesium. It is also called carbonate hardness.

(b) *Permanent hardness* : This is due to the presence of chlorides and sulphates of calcium and magnesium. It is also called non-carbonate hardness.

(iii) *Softening of water* : The process of the removal of hardness from water is called softening of water.

(a) *Removal of temporary hardness* : It can be removed by the following methods,

• By boiling : During boiling, the bicarbonates of *Ca* and *Mg* decompose into insoluble carbonates and give CO_2 . The insoluble carbonates can be removed by filtration.

$$Ca(HCO_3)_2 \xrightarrow{Heat} CaCO_3 + CO_2 + H_2O$$

$$Cal. bicarbonate \xrightarrow{PPt} PPt$$

$$Mg(HCO_3)_2 \xrightarrow{Heat} MgCO_3 + CO_2 + H_2O$$

$$Mag. bicarbonate \xrightarrow{PPt} PPt$$

• Clark's method : This process is used on a commercial scale. In this process, calculated amount of lime [Ca(OH)₂] is added to temporary hard water.

$$Ca(HCO_3)_2 + Ca(OH)_2 \longrightarrow 2CaCO_3 \downarrow + 2H_2O$$
Soluble Lime Mg(CO_3)_2 + Ca(OH_2) \longrightarrow MgCO_3 \downarrow + CaCO_3 \downarrow + 2H_2O
Soluble Lime (Insoluble)

(b) *Removal of permanent hardness* : Permanent hardness can be removed by the following methods, • By washing soda method : In this method, water is treated with a calculated amount of washing soda (Na_2CO_3) which converts the chlorides and sulphates of *Ca* and *Mg* into their respective carbonates which get precipitated.

$$CaCl_{2} + Na_{2}CO_{3} \longrightarrow CaCO_{3} + 2NaCl_{ppt.}$$
$$MgSO_{4} + Na_{2}CO_{3} \longrightarrow MgCO_{3} + Na_{2}SO_{4}_{ppt.}$$

• Permutit method : This is a modern method employed for the softening of hard water. hydrated sodium aluminium silicate $(Na_2AI_2Si_2O_8.xH_2O)$ is called permutit. These complex salts are also known as zeolites.

The permutit as loosely packed in a big tank over a layer of coarse sand. Hard water is introduced into the tank from the top. Water reaches the bottom of the tank and then slowly rises through the permutit layer in the tank. The cations present in hard water are exchanged for sodium ions. Therefore this method is also called ion exchange method.

$$\begin{array}{l} Na_2 Z + Ca^{+2} \longrightarrow CaZ + 2Na^+ \\ \begin{array}{c} \text{Sodium} \\ \text{colite} \end{array} \xrightarrow{(\text{From hard} \\ \text{water})} & CaI \\ \begin{array}{c} \text{zeolite} \end{array} \\ Na_2 Z + Mg^{+2} \longrightarrow MgZ + 2Na^+ \\ \begin{array}{c} \text{Sodium} \\ \text{Magnesium} \end{array} \\ \begin{array}{c} \text{Sodium} \\ \text{water} \end{array} \\ \begin{array}{c} \text{Where} \\ Z = Al_2 Si_2 O_8. \quad XH_2 O \end{array}$$

Hydrogen peroxide

Hydrogen peroxide (H_2O_2) was discovered by French chemist **Thenard**.

(1) **Preparation :** It is prepared by

(i) Laboratory method : In laboratory, H_2O_2 is prepared by Merck's process. It is prepared by adding calculated amounts of sodium peroxide to ice cold dilute (20%) solution of H_2SO_4 . $Na_2O_2 + H_2SO_4 \longrightarrow Na_2SO_4 + H_2O_2$

(ii) By the action of sulphuric acid or phosphoric acid on hydrated barium peroxide $BaO_2.8 H_2O$

(a) $BaO_2 \cdot 8H_2O + H_2SO_4 \rightarrow BaSO_4 \downarrow + H_2O_2 + 8H_2O_2$

It must be noted that anhydrous barium peroxide does not react readily with sulphuric acid (because a coating of insoluble barium sulphate is formed on its surface which stops further action of the acid). Therefore, hydrated barium peroxide, $BaQ_{2.8}H_{2.0}$ must be used.

(b) $3BaO_2 + 2H_3PO_4 \rightarrow Ba_3(PO_4)_2 + 3H_2O_2$ $Ba_3(PO_4)_2 + 3H_2SO_4 \rightarrow 3BaSO_4 + 2H_3PO_4$

Phosphoric acid is preferred to H_2SO_4 because soluble impurities like barium persulphate (from $BaO_2.8H_2O+H_2SO_4$) tends to decompose H_2O_2 while H_3PO_4 acts as preservative (negative catalyst) for H_2O_2 .

(iii) *Industrial method* : On a commercial scale, H_2O_2 can be prepared by the electrolysis of 50% H_2SO_4 solution. In a cell, peroxy disulphuric acid is formed at the anode.

$$2H_2SO_4 \xrightarrow{\text{Elecrolysi}} H_2S_2O_8(aq) + H_2(g)$$
Peroxy disulphuric acid

This is drawn off from the cell and hydrolysed with water to give H_2O_2 .

 $H_2S_2O_8 + 2H_2O \longrightarrow 2H_2SO_4 + H_2O_2$ The resulting solution is distilled under reduced pressure when H_2O_2 gets distilled while H_2SO_4 with high boiling point, remains undistilled.

(iv) **By redox process** : Industrially H_2O_2 is prepared by the auto-oxidation of 2alkylanthraquinols. The process involves a cycle of reactions. The net reaction is the catalytic union of H_2 and O_2 to give H_2O_2 .



The H_2O_2 formed (about 1%) is extracted with water and concentrated.

(2) Physical properties

(i) Pure hydrogen peroxide is a pale blue syrupy liquid.

(ii) It freezes at $-0.5^{\circ}C$ and has a density of 1.4 in pure state.

(iii) Hydrogen peroxide is diamagnetic.

(iv) It is more highly associated via hydrogen bonding than water.

(v) Although it is a better polar solvent than H_2O . However, it can't be used as such because of strong autooxidation ability.

(vi) Dipole moment of H_2O_2 is 2.1 D.

(3) Chemical properties

(i) **Decomposition** : Pure H_2O_2 is an unstable liquid and decomposes into water and O_2 either upon standing or upon heating, $2H_2O_2 \longrightarrow 2H_2O + O_2$; $\Delta H = -196.0 \text{ kJ}$

(ii) **Oxidising nature** : It is a powerful oxidising agent. It acts as an oxidising agent in neutral, acidic or in alkaline medium. *e.g.* $2K/H + H_2O_2 \longrightarrow 2KOH + I_2$ [In neutral medium]

 $2FeSO_4 + H_2SO_4 + H_2O_2 \longrightarrow Fe_2(SO_4)_3 + 2H_2O$ [In acidic medium]

 $MnSO_4 + H_2O_2 + 2NaOH \longrightarrow MnO_2 + Na_2SO_4 + 2H_2O$ [In alkaline medium]

(iii) **Reducing nature** : H_2O_2 has tendency to take up oxygen from strong oxidising agents and thus, acts as a reducing agent, $H_2O_2 + O \longrightarrow H_2O + O_2$. It can act as a reducing agent

in acidic, basic or even neutral medium.

In acidic medium, $H_2O_2 \longrightarrow 2H^+ + O_2 + 2e^-$

In alkaline medium,

 $H_2O_2 + 2OH^- \longrightarrow 2H_2O + O_2 + 2e^-$

(iv) **Bleaching action** : H_2O_2 acts as a bleaching agent due to the release of nascent oxygen.

 $H_2O_2 \longrightarrow H_2O + O$

Thus, the bleaching action of H_2O_2 is due to oxidation. It oxidises the colouring matter to a colourless product, Colouring matter $+O \rightarrow$ Colour less matter.

 H_2O_2 is used to bleach delicate materials like ivory, silk, wool, leather etc.

(v) **Acidic nature** : Anhydrous hydrogen peroxide is acidic in character ($K_a = 1.55 \times 10^{-12}$ at 298 K). its dissociation in aqueous solution may be given as

 $H_2O_2 + H_2O \rightarrow H_3O^+ + HO_2^-$

It forms two types of salts $NaOH + H_2O_2 \rightarrow NaHO_2 + H_2O$ sod. hydroperox ide (Acidicsalt) $2NaOH + H_2O_2 \rightarrow Na_2O_2 + 2H_2O$ Sod. peroxide (Normal salt)

(vi) *Addition reactions* : Hydrogen peroxide is capable of adding itself to ethylenic linkage.

$$\begin{array}{c} CH_2\\ \parallel\\ CH_2\\ CH_2\\ \text{Ethylene} \end{array} + H_2O_2 \rightarrow \begin{array}{c} CH_2OH\\ \parallel\\ CH_2OH\\ \text{Ethyleneglyco} \end{array}$$

(4) **Structure of H_2O_2 :** Hydrogen peroxide is non-linear, non-planar molecule. It has a open book structure. The -O-O- linkage is called peroxy linkage. The structure is shown below.



(5) Concentration of H_2O_2 : Dilute H_2O_2 is concentrated to about 50% by slow evaporation on a water bath. It is further concentrated to 90% in a vacuum desiccator using conc. H_2SO_4 as dehydrating agent. Further concentration to 99% is obtained by distillation under reduced pressure. Last traces of moisture in 99% of H_2O_2 are removed or anhydrous H_2O_2 is obtained by cooling it to 263 *K* in a cold bath of ether and dry ice followed by seeding with a few crystals of solid H_2O_2 when needle-shaped crystals of 100% H_2O_2 separate out. These crystals are removed, dried and melted to get 100% H_2O_2 .

(6) **Storage of H_2O_2:** H_2O_2 is not stored in glass bottles since the alkali metal oxides present in glass catalyse its decomposition. It is, therefore, stored in paraffin wax coated glass, plastic or teflon bottles. Small amounts of acid, glycerol, alcohol, acetanilide and H_3PO_4 are often used as stablizers to check its decomposition.

Uses of hydrogen peroxide

(i) For bleaching delicate articles like wool, hair, feather, ivory, etc.

(ii) For restoring colour of old lead paintings whose white lead has blackened due to formation of *PbS* by H_2S of atmosphere. Hydrogen peroxide converts the black lead sulphide to white lead sulphate

(iii) As an aerating agent in production of spong rubber.

(iv) As an antiseptic and germicide for washing wounds, teeth and ears, under the name of perhydrol.

(v) In the manufacture of sodium perborate, sodium percarbonate. These are used in high quality detergents.

(vi) As an antichlor.

(vii) As an oxidant for rocket fuel.

(viii) In the detection of *Ti*, *V* and *Cr* ions with which it forms peroxides of characteristics colours.

(ix) In the production of epoxides, propylene oxide and polyurethanes.

(x) In the synthesis of hydroquinone, pharmaceuticals (cephalosoporin) and food products like tartaric acid.

(xi) For pollution control of domestic effluents where it restores the aerobic conditions of sewage wastes. For pollution control of industrial effluents containing CN^- ions. H_2O_2 oxidises CN^- ions to harmless products.

A Hydrogen forms more compounds than even carbon.

✓ Metals like *Pd*, *Pt*, *Au* etc., have the property of absorbing large quantity of hydrogen at normal or higher temperature. Colloidal *Pd* can absorb 2950 times its own volume of hydrogen and *Pd* metal can absorb 900 times its own volume of hydrogen.

This phenomenon is known as occlusion of hydrogen. the occlusion property of these metals is in the order

Colloidal Palladium > Palladium > Platinum > Gold > Nickel.

In solids, water molecules can also be present as zeolite water and clathrate water.

∠ Ice is a good thermal insulator.

 \swarrow 30% H_2O_2 is called perhydrol. Its volume strength is 100 and molarity is 8.8.

Chapter 10 The s- Block Element

Important points

- > Groups (1 & 2) belong to the s-block of the Periodic Table.
- Group 1 consists of : lithium, sodium, potassium, rubidium, caesium and francium and collectively known as the *alkali metals*.
- Group 2 include : beryllium, magnesium, calcium, strontium, barium and radium. Except Beryllium they are known as *alkaline*
- * Physical properties-
- a) **Large atomic radii:** The atomic radii of alkali metals are the largest in their respective periods. These increase as we travel down the group.
- b) **Large ionic radii:** The ionic radii increase as we move down the group due to the addition of a new energy shell with each succeeding element.
- c) **Low ionization enthalpy:** The ionization enthalpies decrease as we move down the group. The ionization enthalpies of the alkali metals are the lowest due to loosely held s- electron.
- d) **Hydration enthalpy:** It decreases with the increase in ionic radii. The hydration enthalpy of Li ion is the maximum and the hydration enthalpy of Cs ion is the minimum.
- e) **Oxidation state:** The alkali metals exhibit oxidation state of +1 in their compounds and are strongly electropositive in character. The electropositive character increases from Li to Cs.
- f) Metallic character: The metallic character increases down the group.
- g) Melting point and boiling point:: The m p and b p of alkali metals are very low and decrease with increase in atomic number.
- h) **Nature of bonds formed:** These metals form ionic bonds. The ionic character increases as we down the group.
- i) **Flame colouration:** All the alkali metals impart a charactersistic colour to the flame.
- j) Photoelectric effect: Alkali metals (except Li) exhibits photoelectric effect.
- * Chemical features of alkali metals:
 - a) **Reducing character:** As the ionization enthalpies of the alkali metals decrease down the group their reducing character or reactivity in the gaseous state increases down the group. i.e., Li < Na < K < Rb < Cs.
 - b) **Reaction with dihydrogen:** Alkali metals react with dry hydrogen at about 673 K to form crystalline hydrides which are ionic in nature and have high melting points.

Heat

 $2 M + H_2 \longrightarrow 2M^+ H^-$

c) **Oxides and hydroxides:** Alkali metals when burnt in air form different compounds, for example the alkali metals on reaction with limited quantity of oxygen form normal oxides (M_2O) M= Li, Na, K, Rb, Cs

- d) **Reaction with halogens:** The members of the family combine with halogen to form corresponding halides which are ionic crystalline solids. Reactivity of alkali metls with particular halogen increases from Li to Cs.
- e) **Reaction with water:** Alkali metals react with water and other compounds containing acidic hydrogen atoms such as hydrogen halides, acetylene etc. to liberate hydrogen gas.
- f) **Solubility in liquid ammonia:** All alkali metals dissolve in liquid ammonia giving deep blue solutions which are conducting in nature.
- g) **Reaction with sulphur and phosphorus:** Alkali metals react with sulphur and phosphorus on heating to form sulphides and phosphides respectively.

* Diagonal relationship between Li and Al

Li resembles Mg mainly due to similarity in sizes of their atoms and ions. The main points of similarity are:

- i) Both are quite hard.
- ii) Both LiOH and $Mg(OH)_2$ are weak bases.
- iii) Carbonates of both on heating decompose to produce oxides and carbondioxide.
- iv) Both react with nitrogen to give ionic nitrides.
- v) Nitrates of both decompose on heating to give oxides.
- vi) Both Li and Mg do not form solid bicarbonates.
- vii) Because of covalent character LiCl and MgCl₂ are soluble in ethanol.
- viii) The hydroxides, bicarbonates and fluorides of both Li and Mg are sparingly soluble in water.

✤ Biological importance of Na and K

- i) Sodium ions participate in the transmission of nerve signals.
- ii) Sodium ions also regulate flow of water across the cell membranes and in transport of sugars and amino acids into the cells.
- **iii)** Potassium ions are the most abundant cations within cell fluids, where they activate many enzymes, participate in oxidation of glucose to produce ATP.
- iv) Potassium ions in combination with sodium ions are responsible for transmission of nerve signals.
- v) The functional features of nerve cells depend upon the sodium potassium ion gradient that is established in the cell.

***** Group 2 elements: Alkaline earth metals

(a) Atomic radii : The atomic radii of alkaline earth metals are fairly large though smaller than the corresponding alkali metals and they increase down the group. This is because on moving down the group, atomic radii increase primarily due to the addition of an extra shell of electrons in each succeeding element.

(b) Ionic radii: the atoms of these elements form divalent ions which show the same trend of increase in their size down the group.

(c) **Ionization enthalpy:** The alkaline earth metals have fairly low Ionizations enthalpies though greater than those of the corresponding elements of group 1 and this value decreases down the group.

(d) Hydration enthalpy: the Hydration enthalpies of alkaline earth metal ion decrease as the size of the metal ion increases down the Group

 $Be^{2+} > Mg^{2+} > Ca^{2+} > Sr^{2+} > Ba^{2+}$

(e) Oxidation State: All the members of the family exhibit +2 oxidation state in their compounded and the form divalent cations (M^{2+})

(f) Electro negativity : The electro negativity values of alkaline earth metals are quite close to those of alkali metals, though slightly more.

(g) Metallic Character : Alkaline earth metals have stronger metallic bonds as compared to the alkali metals present in the same period.

(h) Melting and boiling point : The melting and Boiling points of these metals are higher than those of alkali metals present in the same period.

(i) Colouration to the flame : With the exceptio9n of beryllium and magnesium, the rest of the elements impart characters in colour to the same flame. For example,

Be Mg Ca Sr Ba Ra

- Brick Red Crimson Grassy Green Crimson

J) Complex formation: Generally the members do not form complexes. However, smaller ions (Be & Mg Ions) form complexes with the electron donor species

- k) **Formation of organo-metallic compounds:** Both beryllium and magnesium form a number of organo-metallic compounds containing M-C bond with certain organic compounds. For example, magnesium reacts with alkyl halide in the presence of dry ether to give Grignard reagent.
- 1) **Reducing character:** Alkaline earth metals are weak reducing agent than the corresponding alkali metals which have lower ionization enthalpies and comparatively bigger atomic sizes.
- m) **Reaction with oxygen:** With the exception of Ba and Ra which form peroxides (MO₂) rest of the metals form normal oxides (MO) on heating with excess of oxygen.
- n) **Reaction with halogens**: The members of the family combine directly with halogen at appropriate temperature to form corresponding halides.
- o) **Reaction with water:** The members of this group are less reactive towards water as compared to the corresponding alkali metals because these are less electropositive in nature.
- p) **Reaction with hydrogen:** The members except Be combine with hydrogen directly upon heating to form metal hydrides.

Uses of some important compounds:-

(i) Caustic soda:

It is used: in soap, paper, textile, petroleum industry

ii) Sodium carbonate

It is used:

- a) in glass and soap industry
- b) in paper making and textile manufacturing
- c) in paint and dye stuffs
- d) in metal refining

e) in production of sodium compounds such as borax, caustic soda, sodium phosphate etc.

iii) Quick lime:

It is used:

- a. in the preparation of cement, glass and calcium carbide.
- b. In the purification of sugar
- c. In softening of hard water d. As a flux in the extraction of metal
- iv) Lime stone: It is used
- a) as building material
- b) in the manufacture of quick lime
- c) in Solvay process to prepare Na_2CO_3 as it is a source of CO_2
- d) in metallurgy for the extraction of iron
- e) in toothpaste and certain cosmetics
- v) **Cement:** It is an important building material. It is used in concrete and reinforced concrete, in plastering and in the construction of bridges, dams and buildings.
- vi) Plaster of paris: It is used
- a) in making moulds for pottery and ceramics etc.
- b) in surgical bandages for setting broken bones of the body
- c) for making statues, models, decorative materials and black board chalk.

* Biological importance of Ca and Mg

- i) Magnesium ions are concentrated in animal cells and Calcium ions are concentrated in body fluids, outside the cell.
- ii) All enzymes that utilize ATP in phosphate transfer require magnesium ion as cofactor.
- iii) In green plants magnesium is present in chlorophyll.
- iv) Calcium and magnesium ions are also essential for the transmission of impulses along nerve fibres.
- v) Calcium ions are important in blood clotting and are required to trigger the contraction of muscles.
- vi) Calcium ions also regulate the beating of the heart.

One mark questions:

1. Why are halides of beryllium polymeric?

Ans:- the halides of Be are electron deficient as their octets are incomplete. Therefore, to complete their octets, the halides polymerize.

2. Name the groups which constitute s-block elements.

Ans:- group-1 and 2

3.Arrange the alkaline earth metal carbonates in the decreasing order of thermal stability.

Ans:- $BaCO_3 > SrCO_3 > CaCO_3 > MgCO_3 > BeCO_3$

4. Write the general electronic configuration of s-block elements.

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Ans:- [Noble gas] ns<sup>1-2</sup>
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5. What is the chemical formula of Plaster of Paris?

Ans:- $CuSO_4.1/2H_2O$

6.Name the compound which can be obtained by Solvay's process.

Ans:- Sodium carbonate

7. How does the basic character of hydroxides of alkali metals vary down the group? Ans:- Increases down the group

8. Which out of $MgSO_4$ or $BaSO_4$ is more soluble in water?

Ans:- MgSO₄

9.Name radioactive elements of group 1 and 2.

Ans:- Francium and Radium.

10. Which elements of alkaline earth metals family do not give characteristic flame colouration?

Ans:- Be and Mg

Two marks questions

- 1. Among the alkali metals which has
 - (i) Highest melting point
 - (ii) Most electropositive character
 - (iii) Lowest size of ion
 - (iv) Strongest reducing character.

Ans:- (i) Li (ii) Cs (iii) Li (iv) Li

2. Complete the following reactions:

- (i) $Mg(NO_3)_2 \xrightarrow{Heat}$
- (ii) LiOH Heat

(iii) Na₂O + H₂O
$$\longrightarrow$$

(iv) Na + O₂
$$\rightarrow$$

Ans:-

(i) $2Mg(NO_3)_2$ Heat $2MgO + 4NO_2 + O_2$

- (ii) $2\text{LiOH} \xrightarrow{Heat} \text{Li}_2\text{O} + \text{H}_2\text{O}$
- (iii) $Na_2O + H_2O \longrightarrow Na_2CO_3$

(iv)
$$2Na + O_2 \longrightarrow Na_2O_2$$

3. Name the chief factors responsible for anomalous behaviour or lithium. Ans:- the anomalous behaviour of lithium is because of its:

- (i) Small size of stom and ion
 - (i) Small size of atom and ion,
 - (ii) High ionization enthalpy, and
 - (iii) Absence of d-orbitals in its Valence shell.

4. Which out of Li and Na has greater value for the following properties:

- (i) Hydration enthalpy
- (ii) Stability of hydride
- (iii) Stability of carbonate
- (iv) Basic character of hydroxide

Ans:- (i) Li (ii) Li (iii) Na (iv)Na

5. Why are alkali metals not found in nature?

Ans. Alkali metals are highly reactive in nature due to low ionization enthalpy and strong electropositive character. They do not occur in free state and are always combined with other elements. As a result alkali metals are not generally found in nature.

6. Why are lithium salts commonly hydrated and those of the other alkali ions usually anhydrous?

Ans. In the lithium salt, the Li⁺ ion due to very small size gets readily hydrated on coming in contact with moisture (water). Therefore, lithium salts are commonly hydrated. But the other alkali metal ions are comparatively big in size. They have therefore, lesser tendency to get hydrated. These salts are usually anhydrous.

7. Beryllium and magnesium do not give colour to flame whereas other alkaline earth metals do so why?

Ans: Beryllium and magnesium atoms in comparison to other alkaline earth metals are comparatively smaller and their ionisation enthalpies are very high. Hence, the energy of the flame in not sufficient to excite their electrons to higher energy levels. These elements, therefore, do not give any colour in Bunsen flame.

7. Why are alkali metals soft and have low melting points?

Ans: Alkali metals have only one valence electron per metal atom. As a result, the binding energy of alkali metal ions in the close-packed metal lattices are weak. Therefore, these are soft and have low melting point.

8. Which out of the following and why can be used to store an alkali metal? H₂O, C₂H₅OH and Benzene

Ans:- Benzene can be used to store an alkali metal because other substance react with alkali metal as:

Na + H ₂ O \longrightarrow	NaOH + $1/2H_2$
Na + C_2H_5OH	$C_2H_5ONa + 1/2H_2$

9. Why are alkali metals not found free in nature?

Ans:- alkali metals are highly reactive and therefore, are not found free in nature, they are present in the combined state in the form of halides, oxides, silicates, nitrates, etc.

Three marks questions

1. When an alkali metal dissolves in liquid ammonia the solution can acquire different colours. Explain the reasons for this type of colour change.

Ans. The dissolution of the metal in liquid ammonia is accompanied by their formation of ammoniated electrons that give rise to dark colour. This is because ammoniated electrons absorb energy corresponding to the red region of the visible light. However, if the concentration increases above 3M, the colour changes to copper-bronze and the solution acquires metallic luster due to the formation of metal ion clusters.

 $\mathbf{M} + (\mathbf{x} + \mathbf{y})\mathbf{N}\mathbf{H}_3 \quad \rightarrow \qquad [\mathbf{M}(\mathbf{N}\mathbf{H}_3)_3] + [\mathbf{e}(\mathbf{N}\mathbf{H}_3)]$

2. In what ways lithium shows similarities to magnesium in its chemical behaviour?

Ans. Li resembles Mg mainly due to similarity in sizes of their atoms and ions. The main points of similarity are:

Both are quite hard.

1 Both LiOH and $Mg(OH)_2$ are weak bases.

2 Carbonates of both on heating decompose to produce oxides and carbondioxide.

3 Both react with nitrogen to give ionic nitrides.

3. Discuss the various reactions that occur in the Solvay process.

Ans. In Solvay ammonia process.

When carbon dioxide is passed through a concentrated solution of brine saturated with NH₃, NaHCO₃ gets precipitated. NaHCO₃ on subsequent heating gives Na₂CO₃.

 $NaCl + NH_3 + CO_2 + H_2O$ $NaHCO_3 + NH_4Cl$ \rightarrow

$$2 \text{ NaHCO}_3 \rightarrow \text{Na}_2\text{CO}_3 + \text{CO}_2 + \text{H}_2\text{O}$$

CO₂ needed for the reaction is prepared by heating calcium carbonate and the quick lime, CaO is dissolved in water to form slaked lime, $Ca(OH)_2$

CaCO₃ $CaO + CO_2$ $CaO + H_2O$ Ca(OH)₂ \rightarrow

 NH_3 needed for the purpose is prepared by heating NH_4Cl and $Ca(OH)_2$

$$2 \text{ NH}_4\text{Cl} + \text{Ca}(\text{OH})_2 \rightarrow 2 \text{ NH}_3 + \text{Ca}\text{Cl}_2 + \text{H}_2\text{O}$$

4. What happen when (i) magnesium is burnt in air (ii) quick lime is heated with silica (iii) chlorine reacts with slaked lime (iv) calcium nitrate is heated?

Ans. (i) A mixture of magnesium oxide and magnesium nitride is formed

 $5Mg + O_2 + N_2$ \rightarrow 2

In air

$$2 MgO + Mg_3N_2$$

(ii) Calcium silicate is formed.

 $CaO + SiO_2$ CaSiO₃ \rightarrow

(iii) Calcium oxychloride (bleaching powder) is formed

$$Ca(OH)_2 + Cl_2 \longrightarrow CaOCl_2 + H_2C$$

(iv) Nitrogen dioxide is evolved.

Heat $Ca(NO_3)_2$ $CaO + 2 NO_2 + O_2$

5. Describe the importance of the following (i) limestone (ii) cement (iii) plaster of paris.

Ans. i) Lime stone: It is used

f) as building material

- g) in the manufacture of quick lime
- h) in Solvay process to prepare Na_2CO_3 as it is a source of CO_2
- i) in metallurgy for the extraction of iron

i) in toothpaste and certain cosmetics

ii) Cement: It is an important building material. It is used in concrete and reinforced concrete, in plastering and in the construction of bridges, dams and buildings.

iii) Plaster of paris: It is used

- d) in making moulds for pottery and ceramics etc.
- e) in surgical bandages for setting broken bones of the body
- f) for making statues, models, decorative materials and black board chalk.

6. What happens when:

a) Sodium metal is dropped in water?

b) Sodium metal is heated in free supply of air?

c) Sodium peroxide dissolves in water?

Ans. a) Sodium metal catches fire and hydrogen gas is evolved

$$2Na + 2H_2O \longrightarrow 2NaOH + H_2 + Heat$$

b) Sodium peroxide is formed

$$Na + O_2 \rightarrow Na_2O_2$$

c) (i) Sodium peroxide reacts with water at ordinary temperature to liberate oxygen gas

$$Na_2O_2 + 2H_2O \rightarrow 4 NaOH + O_2$$

ii) With ice cold water, H_2O_2 is formed
 $Na_2O_2 + 2H_2O \rightarrow 2 NaOH + H_2 O_2$
State as to why

7.

a) a solution of Na₂CO₃ is alkaline?

b) alkali metals are prepared by electrolysis of their fused chlorides?

c) sodium is found to be more useful than potassium?

Ans. (a) Sodium carbonate being a salt of strong base (NaOH) and weak acid (H_2CO_3) forms alkaline solution upon hydrolysis

 $Na_2CO_3 + 2H_2O \rightarrow 2NaOH + H_2CO_3$

(b) Since the discharge potential of alkali metals is much higher than that of hydrogen, therefore, when the aqueous solution of any alkali metal chloride is subjected to electrolysis, H_2 instead of the alkali metal is produced at the cathode. Therefore, to prepare alkali metals, electrolysis of their fused chlorides is carried out.

(c) Sodium is relatively more abundant than potassium. At the same time, it is also less reactive and its reactions with other substances can be better controlled.

8. Why are potassium and cesium, rather than lithium used in photoelectric cells?

Ans. The ionization enthalpy of lithium is quite high. The photons of light are not in a position to eject electrons from the surface of lithium metal. Therefore photoelectric effect is not noticed. However, both potassium and cesium have comparatively low ionization enthalpies. This means that the electrons can quite easily be ejected from the surface of these metals when photons of certain minimum frequency (threshold frequency) strike against their surface

9. Why is Li₂CO₃ decomposed at a lower temperature whereas Na₂CO₃ at higher temperature?

Ans. Li^+ ion is very small in size. It is stabilized more by smaller anions such as oxide ion rather than large anions such as carbonate. Therefore Li_2CO_3 decomposes into Li_2O on mild heating. On the other hand, Na^+ ion is larger in size. It is stabilized

more by carbonate ion than oxide ion. Hence, Na_2CO_3 does not undergo thermal decomposition easily.

10.Explain why can alkali and alkaline earth metals not be obtained by chemical reduction methods?

Ans. The metals belonging to both these families are very strong reducing agents. It is therefore not possible to reduce their oxides by reacting with common reducing agents like carbon (coke), zinc etc. These are normally isolated by carrying out the electrolysis of the salts of these metals in the molten state.

Five marks questions:

Compare the solubility and thermal stability of the following compounds of the alkali metals with those of the alkaline earth metals.(a) Nitrates (b) Carbonates (c) Sulphates.

Ans. Solubility:

In case of alkali metals: Nitrates, carbonates and sulphates of alkali metals are soluble in water. In alkali metals lattice energies decrease more rapidly than the hydration energies, therefore their solubility increases down the group.

In case of alkaline earth metals: Nitrates of all alkaline earth metals are soluble in water but their solubility decreases down the group because their hydration energies decrease more rapidly than their lattice energies.

Since the size of $CO_3^{2^-}$ and $SO_4^{2^-}$ anions is much larger than the cations, therefore lattice energies remain almost constant with in a particular group. Since, the hydration energies decrease as we move down the group, therefore the solubility of alkaline earth metal carbonates and sulphates decrease down the group. However, the hydration energy of Be²⁺ and Mg²⁺ ions overcome the lattice energy factor and therefore BeSO₄ and MgSO₄ are readily soluble in water while the solubility of other sulphates decreases down the group from CaSO₄ to BaSO₄.

Thermal Stability:

a) Nitrates: Nitrates of both alkali and alkaline earth metals decompose on heating. All alkaline earth metal nitrates decompose to form metal oxide, NO_2 and O_2 .

$$2M(NO_3)_2 \longrightarrow 2MO + 4NO_2 + O_2$$

M= Be, Mg, Ca, Sr, or Ba

The nitrates of Na, K. Rb and Cs decompose to form metal nitrites and O₂.

 $2MNO_3 \longrightarrow 2MNO_2 + O_2$

However, due to diagonal relationship between Li and Mg, lithium nitrate decomposes like $Mg(NO_3)_2$ to form metal oxide, NO_2 and O_2 .

4LiNO₃ Heat $2LiO_2 + 4NO_2 + O_2$

b) Carbonates: Carbonates of alkaline earth metals decompose on heating to form metal oxide and carbon di oxide.

 $2MCO_3 \longrightarrow 2MO + CO_2 M = Be, Mg, Ca, Ba$

Further as the electropositive character of the metal increases down the group the stability of these metal carbonates increases or the temperature of their decomposition increases.

c) Sulphates: Sulphates of alkaline earth metals decompose on heating to form metal oxide and SO₃.

$$MSO_4 \xrightarrow{\text{neur}} 2MO + SO_3 M = Be, Mg, Ca, Ba$$

The temperature of decomposition of these sulphates increases as the electropositive character of the metal or the basicity of the metal hydroxide increases down the group.

Among the alkali metals due to diagonal relationship, Li_2SO_4 decomposes like MgSO₄ to form the corresponding metal oxide and SO₃.

 $\begin{array}{cccc} \text{Li}_2\text{SO}_4 & \xrightarrow{\text{Heat}} & \text{Li}_2\text{O} + \text{SO}_3 \\ \\ \text{MgSO}_4 & \xrightarrow{\text{Heat}} & 2\text{MgO} + \text{SO}_3 \\ \end{array}$

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Other alkali metals are stable to heat and do not decompose easily.

- 2. Compare the alkali metals and alkaline earth metals with respect to (i) ionization enthalpy (ii) basicity of oxides and (iii) solubility of hydroxides..
- Ans.
- (i) Ionization enthalpy (I E): I E of alkaline earth metals are higher than those of alkali metals of group 1. This is because the atoms of alkaline earth metals have smaller size (due to higher nuclear charge) as compared to the alkali metals.
- (ii) Basicity of oxides: The oxides of alkali and alkaline earth metals dissolve in water to form their respective hydroxides. These hydroxides are strong bases. The hydroxides of alkaline earth metals are less basic than of alkali metals of the corresponding periods. This is due to their (i) high ionization enthalpy (ii) small ionic size and (iii) dipositive charge on the ions.

As a result M-O bond in these hydroxides is relatively stronger than that of corresponding alkali metals and therefore does not break.

- (iii) Solubility of hydroxides: Because of smaller size and higher ionic charge, the lattice enthalpies of alkaline earth metals are much higher than those of alkali metals and hence the solubility of alkali metal hydroxides is much higher than that of alkaline earth metals. However the solubility of the hydroxides of both alkali and alkaline earth metals increase down the group due to large decrease in their lattice enthalpies as compared to their hydration enthalpies.
- 3. Explain the significance of sodium, potassium, magnesium and calcium in biological fluids.
- Ans. Significance of sodium and potassium:
 - (i) Sodium ions participate in the transmission of nerve signals.
 - (ii)Sodium ions also regulate flow of water across the cell membranes and in transport of sugars and amino acids into the cells.
 - (iii) Potassium ions are the most abundant cations within cell fluids, where they activate many enzymes, participate in oxidation of glucose to produce ATP.

- (iv) Potassium ions in combination with sodium ions are responsible for transmission of nerve signals.
- (v) The functional features of nerve cells depend upon the sodium potassium ion gradient that is established in the cell.

Significance of Magnesium and Calcium:

- 1. Magnesium ions are concentrated in animal cells and Calcium ions are concentrated in body fluids, outside the cell.
- 2. All enzymes that utilize ATP in phosphate transfer require magnesium ion as cofactor.
- 3. In green plants magnesium is present in chlorophyll.
- 4. Calcium and magnesium ions are also essential for the transmission of impulses along nerve fibres.
- 5. Calcium ions are important in blood clotting and are required to trigger the contraction of muscles.
- 6. Calcium ions also regulate the beating of the heart.

HOTS QUESTIONS

1. Potassium carbonate cannot be prepared by Solvay process. Why?

Ans. This is due to the reason that potassium bicarbonate (KHCO₃) formed as an intermediate (when CO_2 gas is passed through ammoniated solution of potassium chloride) is highly soluble in water and cannot be separated by filtration.

2. The hydroxides and carbonates of sodium and potassium are easily soluble in water while the corresponding salts of magnesium and calcium are sparingly soluble in water. Explain.

Ans. All the compounds are crystalline solids and their solubility in water is guided by both lattice enthalpy and hydration enthalpy. In case of sodium and potassium compounds, the magnitude of lattice enthalpy is quite small as compared to hydration enthalpy since the cationic sizes are large. Therefore, the compounds of sodium and potassium that are mentioned, readily dissolve in water. However, in case of corresponding magnesium and calcium compounds, the cations have smaller sizes and more magnitude of positive charge. This means that their lattice enthalpies are more as compared to the compounds of sodium and potassium. Therefore, the hydroxides and carbonates of these metals are only sparingly soluble in water.

3. Why is LiF almost insoluble in water whereas LiCl soluble not only in water but also in acetone?

Ans. The low solubility of LiF in water is due to its very high lattice enthalpy (F ion is very small in size). On the other hand, in lithium chloride (LiCl) the lattice enthalpy is comparatively very small. This means that the magnitude of hydration enthalpy is quite large. Therefore lithium chloride dissolves in water. It is also soluble in acetone due to dipolar attraction. (Acetone is polar in nature)

CHAPTER 11

The p-block elements

Elements in which the last electron enters in the any one of the three p- orbital of their outermost shells – p-block elements

Gen. electronic configuration of outer shell is ns^2np^{1-6}

The inner core of e-config.may differ which greatly influences their physical & to some extent chemical properties.

- The block of elements in the periodic table consisting of the main groups :
- <u>Group 13</u> (B to Tl)
- $Group \underline{14}$ (C to Pb)
- <u>Group15</u> (N to Bi)
- <u>Group 16</u> (O to Po)
- Group 17 (F to At)
- Group <u>18</u> (He to Rn)

(1) Members at the top and on the right of the *p*-block are nonmetals (C, N, P, O, F, S, Cl, Br, I, At).

(2) Those on the left and at the bottom are metals (Al, Ga, In, Tl, Sn, Pb, Sb Bi, Po).

(3) Between the two, from the top left to bottom right, lie an ill-defined group of metalloid elements (B, Si, Ge, As, Te)

GROUP 13 : The boron group

- Outer Electronic Configuration:-ns²np¹
- group members: boron (B), aluminum (Al), gallium (Ga), indium (In)& thallium (Tl) . All, except boron, are metals.
- Boron show diagonal relationship with Silicon; both are semiconductors metalloids & forms covalent compounds.
- Boron compounds are electron deficient, they are lack of an octet of electrons about the B atom .
- diborane B_2H_6 , is simplest boron hydride

- Structure: three-center two-electron: the H atoms are simultaneously bonded to two B atoms the B-H bridging bond lengths are greater than B-H terminal.
- - Boron oxide is acidic (it reacts readily with water to form boric acid)
- aluminium compounds:aluminium oxide is amphoteric
- aluminum halides, e.g., AlCl₃ is dimer, an important catalyst in organic chemistry have an incomplete octet, acts as Lewic acid by accepting lone pairs from Lewic bases, forming adduct
- aluminum hydride, e.g., LiAlH₄, a reducing agent

Element	Symbol	Atomic	Electronic	Abundance in Earth's
		No.	Configuration	Crest (in ppm)
Boron	В	5	$[He]2s^2 2p^1$	8
Aluminium	Al	13	$[Ne]3s^2 3p^1$	81,300
Galium	Ga	31	$[Ar]3d^{10}4s^2 4p^1$	15
Indium	In	49	$[Kr] 4d^{10}5s^2 5p^1$	1
Thallium	T1	81	$[Xe] 5d^{10}6s^2 6p^1$	0.3

• Atomic Properties - Electronic Configurations

. Atomic and ionic radii

- The atomic and ionic radii of group 13 elements are compared to corresponding elements of group 2. From left to right in the period, the magnitude of nuclear charge increases but the electrons are added to, the same shell. These electrons do not screen each other, therefore, the electrons experience greater nuclear charge.
- In other words, effective nuclear charge increases and thus, size decreases. <u>Therefore, the elements of this group have smaller size than the corresponding</u> <u>elements of second group.</u>
- On moving down the group both atomic and ionic radii are expected to increase due to the addition of new shells. However, the observed atomic radius of Al (143 pm) is slightly more than that of Ga (135 pm).

Ionization energies

The first ionization energies of group 13 elements are less than the corresponding members of the alkaline earths.

The sharp decrease in I.E. from B to Al is due to increase in size. In case of Ga, there are ten d-electrons in its inner electronic configuration.

The very high value of 3^{rd} I. E. of thallium indicates that +3 O.N. state is not stable, rather +1 is more stable for thallium .

Electropositive (or metallic) character

the elements of group 13 are less electropositive as compared to elements of group 2. On moving down the group the electropositive (metallic) character increases because ionization energy decreases. For e.g., Boron is a non-metal white the other elements are typical metals.

Oxidation states

The common oxidation states of group 13 elements are +3 and +1. The stability of the +1 oxidation state increases in the sequence Al <Ga<In <Tl, Due to Inert pair effect.

Element	В	Al	Ga	In	T1
Oxidation state	+3	+3	+3, +1	+3, +1	+3, +1

Chemical reactivity of Gr.13 Elements

All elements in their compounds exhibit the oxidation state of + 3 and +1. <u>Hydrides</u>

- INUDUSTRIAL PREPERATION :-

 $2BF_3(g) + 6LiH(s) \rightarrow B_2H_6(g) + 6LiF(s)$

- Laboratory method:
 - (i) By the reaction of iodine with sodium borohydride in a high boiling solvent.

 $2NaBH_4 + I_2 \rightarrow B_2H_6 + 2NaI + H_2$

(ii) By reduction of BCl₃ with LiAlH₄ 4BCl₃ + 3LiAlH₄ \rightarrow 2 B₂H₆ + 3AlCl₃ + 3 LiCl

Structure of Diborane, B₂H₆



Some important characteristics of boranes:

i) Lower boranes are colourless gases while higher boranes are volatile liquids or solids.

ii) They undergo spontaneous combustion in air due to strong affinity of boron

for oxygen.

 $B_2H_6 + 3O_2 \rightarrow B_2O_3 + 3H_2O + Heat$

iii) Boranes react with alkali metal hydrides in diethyl ether to form borohydride complexes.

 $B_2H_6 + 2MH \rightarrow 2M^+[BH_4]^-$ (M= Li or Na)

Metal borohydride

• (iv) Diborane reacts with ammonia to give borazine at 450 K.

 $B_2H_6 \ + \ 6NH_3 \ \rightarrow 3B_3N_3H_6 \ + \ 12H_2$

- Borazine has a cyclic structure similar to benzene and thus is <u>called inorganic</u> <u>benzene</u>
- The other elements of this group form only a few stable hydrides. The thermal stability decreases as we move down the group.

 AlH₃ is a colourless solid polymerized via Al - H - Al bridging units. These hydrides are weak Lewis acids and readily form adducts with strong Lewis base (B:) to give compounds of the type MH₃ (M = Al or Ga). They also form complex-tetrahydrido anions, [MH4]-. The most important tetrahydrido compound is Li[AlH₄]



Dimeric structure of aluminium chloride

 Boron halides do not form dimers because the size of boron is so small that it is unable to coordinate four large-sized halide ions.

F

CI or Br



• Anomalous properties of boron

1. Boron is a non-metal & bad conductor of electricity whereas aluminium is a metal & good conductor. B is hard but Al is a soft metal.

2. Boron exists in two forms-crystalline and amorphous. But Al does not exist in different forms.

3. The melting and boiling point of boron are much higher than that of Al.

4. Boron forms only covalent compounds whereas Al forms even some ionic compounds.

5. The hydroxides and oxides of boron are acidic in nature whereas those of aluminium are amphoteric.

6. The trihalides of boron exist as monomers. On the other hand, aluminium halides exist as dimers .

7. The hydrides of boron are quite stable while those of aluminium are unstable

- Boron and silicon exhibit the typical properties of non-metals. These do not form cations. Both exist in amorphous as well as crystalline forms.
- Boron oxide (B_2O_3) and silica (SiO_2) both are acidic and dissolve in alkali solutions to form borates and silicates respectively.

 $B_2O_3 \ + \ 6NaOH \rightarrow 2Na_2BO_3 \ \ + \ 3H_2O$

 $SiO_2 + 2NaOH \rightarrow Na_2SiO_3 + H_2O$

• The chlorides of both B and Si get hydrolyzed by water to boric acid and silicic acid respectively.

 $BCl_3 + 3H_2O \rightarrow H_3BO_3 + 3HCl \qquad SiCl_4 + 3H_2O \rightarrow H_2SiO_3 + 4HCl$

- The hydrides of Boron and Silicon are quite stable. Numerous volatile hydrides are also known which catch fire on exposure to air and are easily hydrolyzed.
- Both elements are semiconductors.

Behavior in Aqueous Solutions

1 Al, Ga, In and Tl exhibit a well-defined aqueous chemistry in their tripositive states. Species like [M(OH)4]-, [M(H2O)2(OH)4]-, [M(OH2)6]3+ for M = Al, Ga, In, exist in aqueous solution.

2. Al, Ga. In and T1 ions exist as octahedral aqua ions, [M(OH2)6]3+ in aqueous solution and many salts like halides, sulphates, nitrates and perchlorates exist as hydrates.

3. Aluminiumsulphate forms double salts - called alum, having the general formula

 M_2SO_4 . $Al_2(SO_4)_3$.12 H_2O , where M=Na+ or K+.

USES OF BORON & ALUMINIUM

• Aluminium is used extensively in industry and everyday life. It forms many useful alloys with Cu. Mn, Mg, Si and Zn. Hence, aluminium and its alloys find use in packaging, utensil making, construction, aerospace and other transportation industries. It is used as a conductor for transmission of electricity. Aluminium is alsoused in the alumino-thermite process for production of chromium and manganese from their ores.

Group 14 Elements:-The Carbon Family

Group 14 includes carbon (C), silicon (Si), germanium (Ge), tin (Sn) and lead (Pb).

General electronic configuration of carbon family is ns²np².

Covalent radius:-Covalent radius expected to increase from Cto Si,

From Si to Pb small increase is found.

Ionization Enthalpy:-The first ionization enthalpies of group 14 elements are higher than those of the corresponding group 13 elements.

Electronegativity:-Group 14 elements are smaller in size as compared to group 13 elements that's why this group elements are slightly more electronegative than group 13

Chemical properties:-

Carbon and silicon mostly show +4 oxidation state. Germanium forms stable compounds in +4 state and only few compounds in +2 state.

Tin forms compounds in both oxidation states. Lead compounds in +2 state are stable and in +4 state are strong oxidizing agents.

Exception:- Pb_4 and SnF_4 are ionic in nature.

Except CCl₄ other tetrachlorides are easily hydrolysed by water.

Since carbon does not have d-orbitals and hence cannot expand its coordination number beyond 4

$CCl_4 + H_2O$	No Reaction
SiCl ₄ +4H ₂ O	Si(OH) ₄ +4HCl

Silicic acid

Allotropes of Carbon:-The three types of allotropes are -

1-Diamond

2-Graphite

3-Fullerence

Diamond:-In diamond each carbon atom undergas SP³hybridisation.

Each carbon is tetrahedrally linked to four other carbon atoms.

Graphite:-In graphite, carbon is SP²-hyberdized graphite has a two-dimensional sheet like structure consisting of a number of hexagonal rings fused together.

Graphite conducts electricity along the sheet. It is very soft and Slippery

FullerenceFullerence was discovered collectively by three scientists namely R.E Smalley, R.F Curl and H.W Kroto

SOME Important Compounds Of Carbon and Silicon

Carbon monoxide:-It I prepared by direct oxdisation of C in limited supply of oxygen.

$$2C+O_2(g) \rightarrow 2CO(g)$$

Commercially it is prepared by the passage of steam over hot coke

Carbon Dioxide:-It is prepared by complete combustion of carbon and carbon fuels in excess of air.

$$C(s) + O_2(g) \rightarrow CO_2(g)$$

Laboratory method:-

In laboratory it is prepared by the treatment of dilHCl on CaCO₃

$$CaCO_3(s) + 2HCl(aq) \rightarrow CaCl_2(aq) + CO_2(g) + H_2O(l)$$

Silicon dioxide:-Silicon dioxide is a COVALENT THREE DIMENSIONAL NETWORK SOLID.

Each silicon atom is covalently bonded in a tetrahedral manner to four oxygen atoms.

Silicones:-Silicones are the synthetic organo-siliconpolymers having general formulae $(R_2SiO)_n$ in which R = alkyl (methyl,ethyl or phenyl)

Silicates:-Silicates are exist in nature in the form of feldspar, zeolites, mica and asbestos etc.

The basic structure of silicates is SiO_4^{4-}

Zeolites:-Zeolites is aalumino-silicate of metal. Metal cations participating in formation of Zeolite are use usually Na^+ , K^+ , or Ca^{2+} .

Zeolites are used to remove permanent hardness of water.

ONE MARK QUESTIONS

1. Why is boron used in nuclear reactions?

Ans:-Because Boron can absorb neutrons.

2. By giving a balanced equation show how $B(OH)_3$ behaves as an acid in water.

Ans:-B(OH)₃ +2HOH \rightarrow [B(OH)₄]⁻ +H₃O⁺

3.Name the element of group 14 which exhibits maximum tendency for catenation?

Ans:-Carbon

4. What is the basic building unit of all silicates?

Ans:- SiO_4^{4-} is the basic unit of all silicates.

5. What happens when NaBH₄ reacts with iodine?

Ans:-2NaBH₄ + $I_2 \rightarrow B_2H_6$ +2NaI + H_2 .

6. What happens when boric acid is heated

Ans:- $4H_3BO_3 \longrightarrow 4HBO_2 \longrightarrow H_2B_4O_7$.

7. What is producer gas?

Ans:-Producer gas is a mixture of CO and N₂ in the ratio of 2:1.

8.Write the state of hybridization of 'B' in BF₃.

ANS:-Hybridisation of 'B' in BF_3 is Sp^2 .

9. Mention the state of hybridization in B in BH_4^- .

Ans:- Sp^{3} .

10. Which oxide of carbon is regarded as anhydride of carbonic acid.

Ans:-CO₂ is regarded as a hydride of carbonic acid .

Because $H_2CO_3 \rightarrow H_2O + CO_2$

TWO MARKS QUESTIONS

- 1. Give the chemical reaction as an evidence for each of the following observations.
 - (i) Tin (II) is a reducing agent where as lead (II) is not.

(ii)Gallium (I) undergoes disproportionation reaction.

Ans:- (i) Due to inert pair effect pb^{2+} is more stable than Pb^{4+} . Whereas Sn^{4+} is more stable than Sn^{2+} .

(ii) $3Ga^+ \rightarrow 2Ga + Ga^{3+}$

This is because Ga^{3+} is more stable than Ga^+ .

- 2. What happens when
 - (i) Quick lime is heated with coke?
 - (ii) Carbon monoxide reacts with Cl_2

Ans:- (i) Cao +3C \rightarrow CaC₂ +CO

(iii) $CO + Cl_2 \rightarrow COCl_2$

3. Give reason

(i) C and Si are always tetravalent but Ge,Sn,Pb show divalency.

(ii) Gallium has higher ionization enthalpy than Al. Explain.

Ans:-(i) Ge, Sn, Pb show divalency due to inert pair effect, Pb^{2+} is more stable than Pb^{4+} .

(ii) Due to poor shielding effect of d-electrons in Ga effective nuclear charge increases as compared to Al thus the I.E is higher than Al.

4. Give reason why boron and aluminium tend to form covalent compounds.

Ans:-Sumof three ionization of both the element are very high. Thus they have no tendency to lose electrons to form ionic compound.Instead they form covalent compounds.

5.If B-Cl bond has a dipole moment, Explain why BCl₃ molecule has zero dipole moment.

Ans:- B-Cl bond has dipole moment because of polarity.In BCl₃ since the molecule is symmetrical thus the polarities cancel out.

6.Suggest a reason as to why CO is poisonous.

Ans:-CO reacts with haemoglobin to form carboxy-haemoglobin which can destroy the oxygen carrying capacity of haemoglobin and the man dies of suffocation.

7. What do you understand by-

(a) Inert pair effect:-The pair of electron in the valence shell does not take part in bond formation it is called inert pair effect.

(b) Allotropy:-It is the property of the element by which an element can exists in two forms which have same chemical properties but different physical properties due to their structures.

8. How is excessive content of CO₂ responsible for global warming?

Ans:-Excess of CO_2 absorbs heat radiated by the earth.Some of it dissipated into the atmosphere while the remaining part is radiated back to the earth.Temperature of the earth increases.

9.Describe two similarities and two dissimilarities between B and Al.

Ans:-Similarities:-

- (i) Both have same number of valence electrons.
- (ii) Both have similar electronic configuration.

Dissimilarities:-

- (i) Bis a non- metal where Al is a metal
- (ii) B forms acidic oxide whereas Al forms atmospheric oxides.

10. What are fullerene? How they were prepared?

Ans:- Fullerene are the allotropes of carbon.Its structure is like a soccer ball.

They are prepared by heating graphite in electric arc in presence of inert gases such as helium or argon.

THREE MARKS QUESTIONS

1.What happens when

(a)Borax is heated strongly

(b)Boric acid is added to water

(c)Aluminium is treated with dilute NaOH

Ans:-(a) Na₂B₄O₇ . 10H₂0 \rightarrow Na₂B₄O₇ \rightarrow 2NaBO₂ + B₂O₃ (b) B(OH)₃ + H₂O \rightarrow [B(OH)₄]⁻ + H⁺ (C) 2Al + 2NaOH + H₂O \rightarrow 2NaAlO₂ + 3H₂

2.Explain the following reactions.

(a)Silicon is heated with methyl chloride at high temperature in the presence of copper.

(b)Silicon dioxide is treated with hydrogen fluoride.

(c) CO is heated with ZnO.

Ans:- (a) A mixture of mono-,di- and trimethylchlorosilianes along with a small amount of tetramethylsilane is formed.

 $CH_{3}Cl + Si \longrightarrow CH_{3}SiCl_{3} + (CH_{3})_{2}SiCl_{2} + (CH_{3})_{3}SiCl + (CH_{3})_{4}Si$

(b) The initially formed silicon tetrafluroide dissolves in HF to form hydrofluorosilicic acid

 $SiO_2 + 2HF \rightarrow SiF_4 + 2H_2O$ $SiF_4 + 2HF \rightarrow H_2SiF_6$

(c) ZnO is reduced to zinc metal

 $ZnO + CO \longrightarrow Zn + CO_2$

3. Give reasons:-

(a)Diamond is used as an abrasive.

(b) Aluminium alloys are used to make aircraft body.

(c) Aluminium utensils should not be kept in water overnight.

Ans:- (a) Diamond is used as an abrasive because it is an extremely hard substance. (b)Alloys of alumimium likeduralium is used to make aircraft body due to Someof its property . (c)Generally aluminium metal does not react with water quickly but when it is kept overnight.It reacts slowly with water in presence of air.

 $2Al(s) + O_2(g) + H_2O(l) \rightarrow Al_2O_3(s) + H_2(g)$

4.A certain salt X, gives the following results.

(i)Its aqueous solution is alkaline to litmus.

(ii)It swells up to a glassy material Y on strong heating.

(iii)when conc. H_2SO_4 is added to a hot solution of X,whitecrystalof an acid Z separates out.

Ans:- (i)
$$Na_2B_4O_7 + 10H_2O \rightarrow 2NaOH + H_2B_4O_7 + 8H_2O$$

(ii) $Na_2B_4O_7 \rightarrow 2NaBO_2 + B_2O_3$

 $(iii)Na_2B_4O_7.10H_2O + H_2SO_4 \rightarrow 4H_3BO_3 + Na_2SO_4 + 5H_2O$

5. draw structure of diborane .

Ans.



FIVE MARKS QUESTIONS

1 Explain the formation of (i) Water gas (ii) Producer gas. Give their uses. What happens when CO_2 is passed through lime water (i) for short duration (ii) folong duration.

Ans:-(i)C(s) + H₂O(g) \rightarrow CO(g) +H₂(g) (Water gas) (ii) 2C(s) + O₂ +4N₂(g) \rightarrow 2CO(g) +4N₂(g) (Producer gas) Water gas and Producer gas are used as fuel. Ca(OH)₂ +CO₂ \rightarrow CaCO₃ + H₂O

(White ppt.)

(i)
$$CaCO_3 + CO_2 + H_2O \rightarrow Ca(HCO_3)_2$$

(Soluble)

- 2 (a) Why do Boron halides from addition compound with NH₃ ?
 (b) Assign appropriate reason for each of the following observations :-
 - (i) Anhydrous AlCl₃ is used as a catalyst in many organic reactions.
 - (ii) No form of elemental silicon is comparable to graphite.

Ans:- (a) It is because BX₃ is electron deficient whereas NH₃ is electron rich.

(b) (i) It is Lewis acid.

(ii) It cannot form $p\pi - p\pi$ bond due to large size.

- 3. (i) Give reason for the following observations:-
 - (a) The tendency for catenation decreases down the group in Group 14.
 - (b) The decreasing stability of +3 oxidations state with increasing atomic number in group 13.
 - (c) PbO_2 is a stronger oxidizing agent than SnO_2 .
 - (d) Molten aluminium bromide is a poor conductor of electricity.

Ans:- (i)(a) It is due to decrease in bond dissociation energy which is due to increase in atomic size.

C-C > Si-Si > Ge-Ge > Sn-Sn > Pb-Pb.

(b) It is due to inert pair effect.

(c) PbO_2 is stronger oxidizing agent than SnO_2 because Pb^{2+} is more stable than Pb^{4+} whereas Sn^{4+} is more stable than Sn^{2+} .

(d) Molten $AlBr_3$ is poor conductor of electricity because it is covalent compound.
CHAPTER 12

Organic chemistry : Some Basic Principles and Techniques

Organic compounds are the hydrocarbons and their derivatives and organic chemistry is that branch of chemistry that deals with the study of these compounds <u>Tetravalency of carbon</u>

The atomic number of Carbon is 6 and its electronic configuration is 2,4 i.e. it has 4 valence electrons. Thus carbon is always tetracovalent, i.e. it forms 4 covalent bonds with other atoms



Due to tetravalency of carbon it has a tetrahedron shape.

<u>Catenation</u>- The self linking property of carbon is known as catenation. This is the main reason of existence of such large number of compounds <u>Classification of organic compounds</u>



Functional groups:A functional group may be defined as an atom or a group of atoms present in a molecule which largely determines the chemical properties.

CLASS OF ORGANIC COMPOUNDS	NAME OF FUNCTIONAL GROUP	STRUCTURE
Alkenes	double bond	=C=C=
Alkynes	triple bond	- C Ξ C -
Halogens	halogen	- X (F,Cl,Br,I)
Alcohols	hydroxyl	-OH
Aldehydes	aldehydic(formyl)	-CHO
Carboxylic acids	carboxyl	-COOH
Acid amides	amides	-CONH ₂
Primary amines	amino	- NH ₂

HOMOLOGOUS SERIES

Homologous series is defined as a family or group of structurally similar organic compounds all members of which contain the same functional group, show a gradation in physical and similarity in chemical properties and any two adjacent members of which differ by $-CH_2$ group. The individual members of this group are called homologues and the phenomenon is called homology.

NOMENCLATURE OF ORGANIC COMPOUNDS

Organic chemistry deals with millions of compounds. In order to clearly identify them, a systematic method of naming known as IUPAC system of nomenclature is adopted. The names are such that the listener can deduce the structure from it. The IUPAC name consists of three parts:

Prefix Word root Suffix

EX: 3 methlyoctane

NOMENCLATURE OF ALKANES

Straight chain alkanes:

The names of such compounds is based on their chain structure, and end with suffix '-ane' and carry a prefix indicating the number of carbon atoms present in the chain.

Branched chain hydrocarbons:

- 1.) The longest carbon chain in the molecule is identified.
- 2.) The numbering is done in such a way that the branched carbon atoms get the lowest possible value.
- 3.) The names of the alkyl groups attached as a branch are then prefixed to the name of the parent alkane and its position is indicated by numbers.
- 4.) The lower number is given to the first in alphabetical order.
- 5.) The carbon atom of the branch that attaches to the root alkane is numbered 1.

Organic compounds having Functional Groups:

The longest chain of carbon atoms containing the functional groups is numbered in such a way that the functional group attached to the carbon atom gets the lowest possible number in the chain.

When there are more functional groups then a priority order is followed as:

-COOH, -SO₃H, -COOR, COCl, -CONH₂, -CN, -HC=O, =C=O, -OH, -NH₂, =C=C=, -CΞ C-.

ISOMERISM

Two or more compounds having the same molecular formula but different physical and chemical properties are called isomers and this phenomenon is called isomerism.

Chain isomerism: When two or more compounds having same molecular formula but different carbon skeletons are referred to as chain isomers.



Position Isomerism : Compounds which have the same structure of carbon chain but differ in position of double or triple bonds or functional group are called position isomers and this phenomenon is called Position Isomerism. e g

 $CH_3-CH_2-CH=CH_2$ $CH_3-CH=CH-CH_3$

Functional Isomerism :Compounds which have the same molecular formula but different functional group are called functional isomers and this phenomenon is called functional Isomerism. e g

 $CH_3 - CH_2 - OH$ $CH_3 - O - CH_3$

Metamerism: It is due to the presence of different alkyl groups on either side of functional group in the molecule. Ex. $C_4H_{10}O$ represents $C_2H_5OC_2H_5$ and $CH_3OC_3H_7$.

FISSION OF COVALENT BOND

Heterolytic cleavage: In this cleavage the bond breaks in such a way that the shared pair of electron remains with one of the fragments.

 $H_3C - Br \rightarrow CH_3 + Br$

Homolytic Cleavage: In this cleavage the shared pair of electron goes with each of the bonded atom.

 $R - X \longrightarrow R^{\cdot} + X^{\cdot}$

Alkyl free radical

Nucleophiles : A reagent that brings an electron pair is called nucleophile ie nucleus seeking e g -OH , -CN

Electrophiles: A reagent that takes away electron pair is called electrophile I e electron seeking e g > C = O, $R_3C - X$

Inductive Effect: The displacement of the electron along the chain of the carbon atoms due to presence of an atom or group at the end of the chain.

$\begin{array}{c} d + + + d + + d + \\ CH_3 - \longrightarrow C H_2 & \rightarrow CH_2 & \rightarrow Cl \end{array}$

Resonance Effect : The polarity produced in the molecule by the interaction of two pi bonds or between a pi bond and lone pair of electron present on an adjacent atom.

There are two types of resonance effect:

1) Positive resonance effect : In this effect the transfer of electrons is away from an atom or substituent group attached to the conjugated system.

The atoms or groups which shows +R effect are halogens,-OH , -OR,- NH_2

2) Negative resonance effect : In this effect the transfer of electrons is towards the atom or substituent group attached to the conjugated system.

The atoms or groups which shows -R effect are -COOH, -CHO, -CN

METHODS OF PURIFICATION OF ORGANIC COMPOUNDS :

Sublimation : This method is used to separate the sublimable compounds from non sublimable compounds.

Crystallisation: This method is based on the difference in the solubilities of compound and impurities in a suitable solvent. The impure compound is dissolved in solvent at heated at higher temp .on cooling the hot and conc solution pure compounds crystallizes out.

Distillation: This method is used to separate volatile liquids from non volatile liquids and liquids having sufficient difference in their boiling points.

Fractional distillation: If the boiling points of two liquids is not much, they are separated by this method.

Distillation under reduced pressure : This method is used to purify liquids having high boiling points and decomposes at or below their boiling points.

Steam distillation : This method is used to separate substances which are steam volatile and are immiscible with water.

Differential Extraction: When an organic compound is present in an aqueous medium it is separated by shaking it with organic solvent in which it is more soluble than in water. The aqueous solution is mixed with organic solvent in a separating funnel and shaken for sometimes and then allowed to stand for some time .when organic solvent and water form two separate layers the lower layer is run out by opening the tap of funnel and organic layer is separated. the process is repeated several times and pure organic compound is separated.

Chromatography :This technique is used to separate mixtures in to their components ,purify the compounds and test the purity of compounds.It is classified as

Adsorption Chromatography : It is based on the fact that different compounds are adsorbed on an adsorbent to different degrees. Silica jel or alumina is used as adsorbents.

Partition Chromatography : It is based on the continuous differential portioning of components of a mixture between stationary and mobile phase.

QUALITATIVE ANALYSIS OF ORGANIC COMPOUNDS

Detection of Carbon and Hydrogen: The Carbon and Hydrogen present in the Organic compound is detected by heating the compound with Copper II oxide in a hard glass tube when carbon present in the compound is oxidized to CO2 which can be tested with lime Water and Hydrogenis converted to water which can be tested with anhydrous copper sulphate which turns blue.

 $\begin{array}{r} C + CuO \longrightarrow 2Cu + CO_2 \\ 2 H + CuO & Gu + H2O \\ CO_2 + Ca (OH)_2 & CaCO_3 + H_2O \\ 5H_2O + CuSO_4 & -GuSO_4.5H_2O \\ \hline \end{array}$

Sodium Fusion Extract: A small piece of dry Sodium metal is heated with a organic compound in a fusion tube for 2 -3 minutes and the red hot tube is plunged in to distilled water contained in a china dish. The contained of the china dish is boiled ,cooled and filtered. The filtrate is known as Sodium fusion extract.

Test for Nitrogen : The sodium fusion extract is boiled with iron II sulphate and then acidified with Concsulphuric acid, the formation of Prussian blue colour confirms the presence of nitrogen.

 $6CN^{-} + Fe^{2+} \longrightarrow [Fe(CN)_6]^{4-}$ $3[Fe(CN)_6]^{4-} + 4Fe^{3+} \xrightarrow{xH \ O}_{2} \qquad Fe_4[Fe(CN)_6]_{3.}xH_2O$

Test for Sulphur: the sodium fusion extract is acidified with acetic acid and lead acetate is added to it. A black precipitate of lead sulphide indicates the presence of sulphur. $S^{2-} + Pb^{2+} \longrightarrow PbS$

Black

Test for halogens: The sodium fusion extract is acidified with nitric acid and then treated with silver nitrate. A white precipitate, soluble in ammonium hydroxide shows the presence of chlorine, a yellowish ppt. sparingly soluble in ammonium hydroxide shows the presence of bromine, a yellowish ppt. insoluble in ammonium hydroxide shows the presence of iodine.

$$X^- + Ag^+ \longrightarrow AgX$$

QUANTITIVE ANALYSIS(Carbon and Hydrogen)

Let the mass of organic compound be m g. Mass of water and carbon dioxide produced be m_1 and m_2 g respectively;

% of carbon = $12 \times m_2 \times 100$ $44 \times m$ % of hydrogen = $2 \times m_1 \times 100$ $18 \times m$

Nitrogen

DUMAS METHOD: A known mass of organic compound is heated with excess of CuO in an atmosphere of CO_2 , when nitrogen of the organic compound is converted into N_2 gas. The volume of N_2 thus obtained is converted into STP and the percentage of nitrogen determined by applying the equation:

Volume of Nitrogen at $STP = P_1V_1 \times 273$

 $%N = 28 \text{ x vol of } N_2 \text{ at STP x 100}$

22400 x mass of the substance taken

KJELDAHL'S METHOD: A known mass of organic compound is heated with conc. H_2SO_4 in presence of K_2SO_4 and little $CuSO_4$ or Hg in a long necked flask called Kjeldahl's flask when nitrogen present in the organic compound is quantitatively converted into $(NH_4)_2SO_4$. $(NH_4)_2SO_4$ thus obtained is boiled with excess of NaOH solution to liberate NH_3 gas which is absorbed in a known excess of a standard acid such as H_2SO_4 or HCl.

The vol of acid unused is found by titration against a standard alkali solution. From the vol of the acid used, the percentage of nitrogen is determined by applying the equation,

%N= 1.4 x Molarity of the acid x Basicity of the acid x Vol of the acid used

Mass of the substance taken

Halogens

Carius method:

A known mass of an organic compound is heated with fuming nitric acid in the presence of silver nitrate contained in a hard glass test tube known as carius tube in a furnace. Carbon and hydrogen present in the compound are oxidized to carbon dioxide and water. The halogen present forms the corresponding silver halide. It is filtered, dried, and weighed.

Let the mass of the organic compound taken = m g Mass of AgX formed = m₁ g 1 mol of AgX contains 1 mol of of X Mass of halogen in m₁ g of AgX = ______ at mass of X xm₁ g Molecular mass of AgX % of halogen = $at mass of X xm_1 g x 100$ % Molecular mass of AgX x m Subbur

Sulphur

Let the mass of the organic compound taken = m g

Mass of $BaSO_4$ formed = m_1 g

% of sulphur = $32 \times m_1 \times 100$ %

233 x m

Phosphorous

Let the mass of the organic compound taken = m g

Mass of ammonium phosphomolydate = $m_1 g$

 $\% \ of \ phosphorous = 31 \ x \ m_1 \ x \ 100 \qquad \%$

Oxygen

Let the mass of the organic compound taken = m g

Mass of $CO_2 = m_1 g$

% of oxygen = $32 \times m_1 \times 100$ %

44 x m

One Mark Questions

Q1 Suggest a method to purify a liquid which decomposes at its boiling point.

A 1 The process Distillation Under reduced pressure is used to purify a liquid which decomposes at its boiling point.

Q 2 How will you separate a mixture of O-nitrophenol and p- nitrophenol ?

A 2 O-nitrophenol is steam volatile therefore it can be separated by Steam distillation.

Q 3 Lassaigne's test is not shown by diazonium Salt. Why?

A 3 On heating diazonium Salts loses Nitrogen and could not fuse with the Sodium metal therefore diazonium Salt do not show Positive Lassaigne's test for nitrogen.

Q 4 Alcohols are weaker acids than Water, Why?

A 4 The alkyl group in alcohols has + I effect due to which electron density is increases on Oxygen atom which makes the release of hydrogen ion more difficult from alcohol.R $\rightarrow O \rightarrow H$

Q 5 Why is nitric acid is added to Sodium extract before adding Silver nitrate for testing halogens ?

A 5 Nitric acid is added to decompose NaCN and Na₂S

 $NaCN + HNO_3 \rightarrow NaNO_3 + HCN$

 $Na_2S + 2HNO_3 \rightarrow 2NaNO_3 + H_2S$

Q 6 which of the two $O_2NCH_2CH_2^-$ or $CH_3CH_2O^-$ is expected to be more stable and why ?

A 6 NO₂ group has –I effect and disperse the negative charge on Oxygen atom

 $O_2N \leftarrow CH_2 \leftarrow CH_2O^-$

Q 7 Arrange the following in increasing Order of Stability ;

```
(CH_3)_3C^+, CH_3CH_2CH_2C^+H_2, CH_3CH_2C^+HCH_3, CH_3C^+H_2, CH_3CH_2C^+H_2
```

```
A 7 CH_3C^+H_2 < CH_3CH_2C^+H_2 < CH_3CH_2CH_2C^+H_2 < CH_3CH_2C^+HCH_3 < (CH_3)_3C^+
```

Q 8 Write the IUPAC name of the following

CH₃CH₃

A 8 2,3Dimethylpentane

Q 9 Write the hybridized state of C atoms in the following

$$CH_2 = CH - C \Xi N$$

A 9
$$sp^2sp^2sp$$

 $CH_2 = CH - C \Xi N$

Q 10 Give the IUPAC name of the following compound.



A 10 2,5Dimethylheptane

Two Marks Questions

Q 1 Draw the Structures of the following compounds.



Q 2 Explain Inductive effect with example.

A 2 Inductive Effect: The displacement of the electron along the chain of the carbon atoms due to presence of an atom or group at the end of the chain.

$$\delta + + + \delta + + \delta +$$

 $CH_3 {\rightarrow} \qquad C \; H_2 \; {\rightarrow} \quad CH_2 \; {\scriptstyle \rightarrow} Cl$

Q 3 Explain why $(CH_3)_3C^+$ is more stable than $CH_3C^+H_2$.

A 3 $(CH_3)_3C^+$ has nine alpha hydrogens and has nine hyperconjugation structures while $CH_3C^+H_2$ has three alpha hydrogens and has three hyperconjugation structures, therefore $(CH_3)_3C^+$ is more stable than $CH_3C^+H_2$

Q 4 Give the number of Sigma and pi bonds in the following molecules

a) CH₃NO₂ b)HCONHCH₃

A 4 a) 6 Sigma and 1 pi bond

b) 8 Sigma and 1 pi bond

Q 5 Write the condensed and bond line formula of 2,2,4-Trimethylpentane.



Q 6 How Sodium fusion extract is prepared ?

A 6 A small piece of dry Sodium metal is heated with a organic compound in a fusion tube for 2 -3 minutes and the red hot tube is plunged in to distilled water contained in a china dish. The contained of the china dish is boiled ,cooled and filtered. The filtrate is known as Sodium fusion extract.

Q 7 Explain the principle of paper chromatography.

A 7 Paper chromatography is based on the difference in the rates at which the components of a mixture are adsorbed. The material on which different components are adsorbed is called Stationary phase which is generally made up of alumina, silica jel or activated charcoal. The mixture to be separated is dissolved in a suitable medium and it is called moving phase. The moving phase is run on the Stationary phase , the different compounds are adsorbed on stationary phase at different rates.

Q 8 Why is an organic compound fused with Sodium for testing nitrogen, halogens and sulphur ?

A 8 On fusing with sodium metal the elements presents in an organic compounds are converted in to sodium salts which are water soluble which can be filtered and detected by the respective tests.

Q 9 It is not advisable to use sulphuric acid in place of acetic acidfor acidification while testing sulphur by lead acetate test. Give reason

A 9 Lead acetate on reacting with sulphuric acid will give a white ppt of lead sulphatewhih interfere in the detection of sulphur.

 $(CH_3COO)_2Pb + H_2SO_4 \rightarrow PbSO_4 + 2 CH_3COOH$

Q 10 Under what conditions can the process of steam distillation is used ?

A 10 Steam distillation is used to purify the liquids which are steam volatile and water and the liquid are not miscible with each other.

Three Marks Questions

Q 1 In an estimation of sulphur by carius method 0.468 g of an organic compound gave 0.668 g of barium sulphate. Find the percentage of sulphur in the compound.

A 1 Mass of the compound = 0.468 g

Mass of the barium sulphate = 0.668 g

% of sulphur = 32 X Mass of barium sulphate X 100

233 X Mass of the compound = $32 \times 0.668 \times 100$ 233 x0.468 = 19.60 %

Q 2 Which bond is more polar in the following pairs of molecules.

a) H₃C-H, H₃C-Br b) H₃C-NH₂, H₂C-OH c) H₃C-OH, H₃C-SH

A 2 a) C-Br because Br is more electronegative than H

- b) C-O because O is more electronegative than N
- c) C-O because O is more electronegative than S

Q 3 Define Isomerism.Explain position Isomerism and Functional Isomerism with examples.

A 3 Two or more compounds having the same molecular formula but different physical and chemical properties are called isomers and this phenomenon is called isomerism.

Position Isomerism : Compounds which have the same structure of carbon chain but differ in position of double or triple bonds or functional group are called position isomers and this phenomenon is called Position Isomerism. e g

 $CH_3-CH_2-CH=CH_2$ $CH_3-CH=CH-CH_3$

Functional Isomerism :Compounds which have the same molecular formula but different functional group are called functional isomers and this phenomenon is called functional Isomerism. e g

 $CH_3 - CH_2 - OH$ $CH_3 - O - CH_3$

Q 4 write the IUPAC names of the following compounds.

$$\begin{array}{c} O & O \\ || \\ A.CH3 - CH2 \stackrel{O}{-}C - CH2 \stackrel{||}{-}C - CH3 \\ B & HC \equiv C - CH = CH - CH - CH_2 \\ C & Cl CH_2CH_2CH_2CH_2Br \end{array}$$

A 4 A hexane 2,4dione

- B hexa-1,3-dien-5-yne
- C 1-bromo-4-chlorobutane

Q 5 Define Homologous series. Write the general formula of alkane, alkene and alkynes.

A 5 Homologous Series : It is defined as group of similar organic compounds which contains the similar functional groups and the two adjacent members of the series is differ by a $-CH_2$ group.

Alkanes	$C_nH_{2n+2} \\$
Alkenes	C_nH_{2n}
Alkynes	C_nH_{2n-2}
Q 6 How	many Sigma

Q 6 How many Sigma and pi bonds are present in the following molecules .

A HC Ξ CCH = CHCH₃

B $CH_2 = C = CHCH_3$

A 6 A Sigma bonds = 10 pi bonds = 3

B Sigma bonds = 9 pi bonds = 2

Q 7 Define functional groups. Write the general formula of Carboxylic acids acid chlorides.

A 7 Functional Groups : It is an atom or group of atoms bonded together in a unique manner which is usually the site of chemical reactivity in an organic molecule. e g CH_3OH

General formula of Carboxylic acids : $C_nH_{2n+1}COOH$

General formula of acid chlorides :RCOCl

Q 8 Write a shirt note on differential extraction.

A 8 When an organic compound is present in an aqueous medium it is separated by shaking it with organic solvent in which it is more soluble than in water. The aqueous solution is mixed with organic solvent in a separating funnel and shaken for sometimes and then allowed to stand for some time .when organic solvent and water form two separate layers the lower layer is run out by opening the tap of funnel and organic layer is separated. the process is repeated several times and pure organic compound is separated.

Q 9 How carbon and Hydrogen is detected in a organic compounds.

A 9 The Carbon and Hydrogen present in the Organic compound is detected by heating the compound with Copper II oxide in a hard glass tube when carbon present

in the compound is oxidized to CO2 which can be tested with lime Water and Hydrogenis converted to water which can be tested with anhydrous copper sulphate which turns blue.

 $C + CuO \rightarrow 2Cu + CO_2$

 $2 H + Cu \Theta Cu + H_2O$

 $CO_2 + Ca (OH) \rightarrow CaCO_3 + H_2O$

 $5H_2O + CuSO_4 \rightarrow CuSO_4.5H_2O$

Q 10 Write a short note on Resonance effect.

Resonance Effect : The polarity produced in the molecule by the interaction of two pi bonds or between a pi bond and lone pair of electron present on an adjacent atom.

There are two types of resonance effect:

- Positive resonance effect: In this effect the transfer of electrons is away from an atom or substituent group attached to the conjugated system. The atoms or groups which shows +R effect are halogens,-OH , -OR,-NH₂
- 2. Negative resonance effect: In this effect the transfer of electrons is towards the atom or substituent group attached to the conjugated system.

The atoms or groups which shows -R effect are -COOH, -CHO, -CN

Five Marks Questions

Q 1 Differentiate between the principle of estimation of nitrogen in an organic compound by i) Dumas method ii) Kjeldahl's method.

Ans: DUMAS METHOD: A known mass of organic compound is heated with excess of CuO in an atmosphere of CO_2 , when nitrogen of the organic compound is converted into N_2 gas. The volume of N_2 thus obtained is converted into STP and the percentage of nitrogen determined by applying the equation:

Volume of Nitrogen at $STP = P_1V_1 \ge 273$

 $%N = 28 \text{ x vol of } N_2 \text{ at STP x 100}$

22400 x mass of the substance taken

KJELDAHL'S METHOD: A known mass of organic compound is heated with conc. H_2SO_4 in presence of K_2SO_4 and little $CuSO_4$ or Hg in a long necked flask called Kjeldahl's flask when nitrogen present in the organic compound is quantitatively converted into $(NH_4)_2SO_4$. $(NH_4)_2SO_4$ thus obtained is boiled with excess of

NaOHsolution to liberate NH_3 gas which is absorbed in a known excess of a standard acid such as H_2SO_4 or HCl.

The vol of acid unused is found by titration against a standard alkali solution. From the vol of the acid used, the percentage of nitrogen is determined by applying the equation,

%N = 1.4 x Molarity of the acid x Basicity of the acid x Vol of the acid used

Mass of the substance taken

Q 2 A sample of 0.50g of organic compound was treated according to Kjeldahl's method. The ammonia evolved was absorbed in 50mL of 0.5M H_2SO_4 . The residual acid required 60mL of 0.5M solution of NaOH for neutralization. Find the percentage composition of nitrogen in the compound.

Ans: the vol ofH_2SO_4 used.

Vol of acid taken=50mL of 0.5M H₂SO₄= 25mL of 1M H₂SO₄

Vol of alkali used for neutralization of excess acid= 60 mL of 0.5m NaOH=30mL of 1M NaOH

Now 1 mole of H₂SO₄ neutralizes 2 moles of NaOH

(i.e. $H_2SO_4 + 2 NaOH \rightarrow Na_2SO_4 + 2H_2O$)

 \dots 30 mL of 1M NaOH = 15mL of 1M H₂SO₄

% of nitrogen.

1 mole of H_2SO_4 neutralizes 2 moles of $NH_3 \dots 10mL$ of $1M H_2SO_4 = 20mL$ of $1M NH_3$

But 1000mL of 1M NH₃ contain N=14g.

20 ml of 1M NH₃ will contain nitrogen = 14×20

1000

But this amount of nitrogen is present in 0.5 g of organic compound

 $\therefore \% \text{ of } N = \underbrace{14 \ x \ 20 \ x \ 100}_{1000 \ x \ 0.5} = 56.0$

Q 3 You have a mixture of three liquids A, B, C. there is a large difference in the boiling point of A and the rest two liquids. Boiling points of liquids B and C are quite close. Liquid A boils at higher temperature than B and C and boiling point of B is lower than C. How will you separate the components of the mixture.

Ans Since the boiling point of liquid A is much higher than those of liquids B and C, therefore separate liquid A by simple distillation. Since boiling points of liquids B and C are quite close but much lower than liquid A therefore mixture of B and C will distil together leaving behind A. on further heating A will distil over.

Now place the mixture of liquids B and C in a flask fitted with fractionating column. Since the b.p. of liquid B is lower than that of C, on fractional distillation first liquid B will distil over and than liquid C.

HOTS QUESTIONS

Q 1 Explain hyperconjugation effect. How does hyperconjugation effect explain the stability of alkenes?

Ans The relative stability of various classes of carbonium ions may be explained by the number of no-bond resonance structures that can be written for them. Such structures are obtained by shifting the bonding electrons from an adjacent C-H bond to the electron deficient carbon so the positive charge originally on carbon is dispersed to the hydrogen. This manner of electron release by assuming no bond character in the adjacent C-H bond is called Hyperconjugation. Greater the hyperconjugation greater will be the stability of alkenes.

$$CH_3 - CH = CH - CH_3 < CH_3 - C = CH - CH_3$$
 $< CH_3 - C = C - CH_3$
 $| CH_3 - CH_3 - C = C - CH_3$
 $| CH_3 - CH_3 - C = C - CH_3$

Q 2In DNA and RNA nitrogen is present in the ring system. Can kjeldahl method be used for the estimation of nitrogen present in these ?give reasons

AnsIn DNA and RNA nitrogen is present in hetrocyclicrings.Kjeldahl method can not be used to estimate nitrogen present in the ring because cannot be completely converted in to $(NH_4)_2SO_4$ during digestion. Therefore Kjeldahl method can not be used to estimate nitrogen present in DNA and RNA.

Q 3 1.216 g of an organic compound was Kjeldahlised and the ammonia evolved was absorbed in 100 mL 1N H_2SO_4 . The remaining acid solution was made upto500ml by addition of water. 20ml of this dilute solution required 32mL of N/10 caustic soda solution for complete neutralization. Calculate the percentage of nitrogen in the organic compound.

Ans 20 ml of dil. Unreacted $H_2SO_4 = 32mL$ of N/10 NaOH sol.

500ml of dilunreacted $H_2SO_4 = 32 \times 500 \text{ mL of } NNaOH = 32 \times 500 \text{ ml of } 1 \text{ N NaOH}$

= 80ml 1 N NaOH

But 80ml 1 N NaOH= 80ml 1 N NaOH So, acid left unused = 80ml 1 N H_2SO_4 Acid used =(100 - 80) = 20ml 1 N H_2SO_4

%N = 1.4 x Normality of the acid x Vol of the acid used

Mass of the substance taken

= 1.4 x 1 x 20 = 23.026

1.216

CHAPTER 13

HYDROCARBON

- > <u>Hydrocarbons</u> are composed of Carbon and hydrogen.
- ➤ The important fuels like Petrol, kerosene, coal gas, CNG, LPG etc. are all hydrocarbons or their mixture.

Sources:

Petroleum and natural gas are the major sources of aliphatic hydrocarbon while coal is an important source of aromatic hydrocarbons. The oil trapped inside the rocks is known as petroleum. PETRA – ROCK, OLEUM – OIL. The oil in the petroleum field is covered with a gaseous mixture known as natural gas. The main constituents of the natural gas are methane, ethane, propane and butane.



Preparation:-

• Wurtz reaction:-

 $2CH_{3}CH_{2}Br + 2Na \xrightarrow{Dry} CH_{3}CH_{2}CH_{2}CH_{3} + 2NaBr$

- Follow mainly free radical mechanism
- Useful in preparing an alkane containing even number of carbon atoms

Stepping up reaction

Frankland reaction

 $RX + Zn + Rx \longrightarrow R - R + ZnX_2$

②From Grignard reagent (RMgX)

 $RMaX + HOH \rightarrow RH + Ma(OH)X$ $RMaX + R'OH \rightarrow RH + Ma(OR')X$ $RMgX + R'NH_2 \rightarrow RH + Mg(NHR')X$

 $R - CH = CH_2 + H_2 \xrightarrow{Ni/\Delta} R - CH_2 - CH_3$ $R - C \equiv CH + H_2 \xrightarrow{Ni/\Delta} R - CH_2 - CH_3$

4. From carboxylic acids-

Decarboxylation.-

 $CH_3COO^-Na^+ + NaOH \xrightarrow{CaO} CH_4 + Na_2CO_3$

Sodium ethanoate Kolbe's electrolytic method-

> $2CH_3COO^-Na^+ + 2H_2O$ Sodium acetate

> > ↓Electrolysis

 $CH_3 - CH_3 + 2CO_2 + H_2 + 2NaOH$

Physical Properties:-

 Nature:- Non-Polar due to covalent nature of C—C bond and C—H bond. C— C bond enrgy = 83 kj/mole and C—H bond energy = 99 kj/mole.

 C_1 — C_4 = gases, C_5 — C_{17} = colourless odourless liquid and > C_{17} = Solid.

(2) Solubility:- Like dissolve like

Viz, Polar compounds dissolve in polar solvent and Non-Polar compound dissolve in non polar solvent.

(3) Boiling point:- Low boiling point due to non polar in nature.

The molecules are held together only by weak Van der Waalls' forces.

Since we known that the magnitude of Van der Waalls' forces is directly proportional to the molecular size. Therefore, the boiling point increases with increase the molecular size i.e. with increase in number of carbon atoms.

Noted:- the boiling points of the branched chain Alkanes are less than the straight chain isomers.

This is due to the fact that branching of the chain makes the molecule more compact and thereby decreases the surface aria and consequently, the magnitudes of Van der Waalls' forces also decrease.

✤ Chemical properties

• Combustion:- $CH_4 + 2O_2 \rightarrow CO_2 + 2H_2O$ $\Delta H = -217.0 \text{ K cal/mole}$

Oxidation:-

$$CH_4 + O_2 \xrightarrow{Cu} 2CH_3OH$$

$$CH_4 + O_2 \xrightarrow{Mo_2O_3} HCHO + H_2O$$
Methanal

• Substitution:-

① Halogenation:-

 $CH_4 + Cl_2 \xrightarrow{UV} CH_3Cl + HCl$ $CH_3Cl \xrightarrow{UV} CH_2Cl_2 \xrightarrow{UV} CHCl_3 \xrightarrow{UV} CCl_4$

Noted:- Iodination is a reversible reaction. So it is carried out by heating alkane in the presence of some oxidizing agent like iodic acid (HIO_3) or nitric acid (HNO_3) or mercuric oxide (HgO) which oxidizes HI formed during the reaction.

Noted:- Fluorination of alkane takes place explosively resulting even in the rupture of C—C bond in higher alkanes.

- Features of Halogenations:-
 - (i) The reactivity of Halogens:- $F_2 > Cl_2 > Br_2 > I_2$.

(ii) The rate of replacement of Hydrogens of alkanes is: $3^{\circ} > 2^{\circ} > 1^{\circ}$

$$CH_{3}CH_{2}CH_{2}CH_{3} \xrightarrow{Cl_{2}} CH_{3}CH_{2}CH_{2}CH_{2}CH_{2}CI + CH_{3}CH_{2}CHCH_{3}$$

n - Butane

$$CH_{3} \xrightarrow{\circ} CH_{3} \xrightarrow{\circ} CH_{3} \xrightarrow{\circ} CH_{3} \xrightarrow{Cl_{2}}$$

$$CH_{3} \xrightarrow{\circ} CH_{3} \xrightarrow{\circ} CH_{3} \xrightarrow{Cl_{2}}$$

$$Isobutane$$

$$CH_{3} \xrightarrow{CH_{3}} CH_{3} CH_{3} \xrightarrow{CH_{3}} CH_{3} \xrightarrow{CH_{3}} CH_{3} \xrightarrow{CH_{3}} CH_{3} \xrightarrow{CH_{3}}$$

Mechanism:- halogenations reaction take place by free radical mechanism. The reaction proceeds in the following steps: Initiation

(i) Chain initiation step:-

 $CI - CI \xrightarrow{h_V} 2CI$

(ii) Chain Propagation step:-

$$\begin{array}{rcl} \mathsf{CH}_4 &+& \mathsf{CI}^\bullet & \longrightarrow \overset{\bullet}{\mathsf{CH}}_3 &+& \mathsf{HCI} \\ \overset{\bullet}{\mathsf{CH}}_3 &+& \mathsf{Cl}_2 & \longrightarrow & \mathsf{CH}_3\mathsf{CI} &+& \mathsf{CI} \bullet \end{array}$$

(iii) Chain Termination step:-

$$CI^{\bullet} + CI^{\bullet} \longrightarrow CI_{2}$$

$$CH_{3} + CH_{3} \longrightarrow CH_{3} - CH_{3}$$

$$CH_{3} + CI^{\bullet} \longrightarrow CH_{3}CI$$

② Nitration:-

- The reaction takes places by free radicals mechanism at high temp (450° C).
- The At high temp C—C bond is also broken so that mixture of nitroalkanes is obtained.

$$CH_{3}CH_{2}CH_{3} \xrightarrow{450^{\circ}C} CH_{3}CH_{2}CH_{2}NO_{2} + CH_{3}CHCH_{3} + CH_{3}CH_{2}NO_{2} + CH_{3}NO_{2}$$

$$NO_{2}$$

$$25\% \qquad 40\% \qquad 10\% \qquad 25\%$$

The reaction occurs as: HO-NO₂ $\xrightarrow{450\circ C}$ HO \circ + \circ NO₂ RH + \circ OH $\xrightarrow{}$ R \circ + HOH R \circ + \circ NO₂ $\xrightarrow{}$ R \circ + HOH RNO₂

③ Sulphonation:- replacement of hydrogen atom of alkane by –SO₃H group.



GAromatization:-

This method is also called dehydrogenation or hydroforming

Similarly, heptane gives toluene, n-Octane give o-xylene and 2, methyl heptane give m-xylene.

• Thermal decomposition or Pyrolysis or cracking or Fragmentation: - when higher alkanes are heated at high temp (about 700-800k) in the presence of alumina or silica catalysts, the alkanes break down to lower alkanes and alkenes.

 $\begin{array}{cccc} CH_3-CH_2-CH_3 &\longrightarrow & CH_3-CH-CH_2 + CH_3-CH_3 + C_2H_4 + CH_4 \\ \hline \bullet & \text{Action of steam:- catalyst:} & \text{nickel, alumina } Al_2O_3 \\ & 1000 \ ^0C \\ CH_4 + H_2O(\text{Steam}) & \longrightarrow CO + 3H_2 \end{array}$

This reaction is used for the industrial preparation of hydrogen from natural gas. 8. Isomerisation:-



✤ CONFORMATIONAL ISOMERISM:

The different molecular arrangements arising as a result of rotation around carbon carbon single bonds are called conformational isomers or rotational isomers and the phenomenon is called conformational isomerism.

Numerous possible arrangements of ethane are possible. Two extreme conformations are known. These are eclipsed conformation and staggered conformation.

SAWHORSE REPRESENTATION



Alkenes

- Turnaturated hydrocarbon which have double bond.
- ☞ General molecular formula C_nH_{2n}
- **☞** C−C bond hybridization 1.34 A⁰
- \sim sp² hybridization
- When we treated Alkene with chlorine, oily products are obtained. So Alkenes are also known as Olefins. (Greek olefiant meaning oil forming).
- Show chain, positional and geometrical isomerism
- Structure of double bond:-



* Preparation:-

1. From Alkynes:- Alkynes on partial reduction with Partially deactivated palladised charcoal known as *Lindlar's catalyst* give alkynes.

 $\begin{array}{ccc} \mathrm{CH}{=}\mathrm{CH}{+}\mathrm{H}_2 \xrightarrow{\mathrm{Pd/C}} \mathrm{CH}_2{=}\mathrm{CH}_2 \\ \text{Ethyne} & \text{Ethene} \end{array}$

2. From Haloalkanes: - dehydrohalogenation

(E₂ or 1,2-elimination or Bita-elimination)



3. From Dihaloalkanes: - dehalogenation



4. From Alcohols:- Dehydration (E1 - elimination)

 $CH_{3}CH_{2}CH_{2}OH \xrightarrow{Conc.H_{2}SO_{4}} CH_{3}CH = CH_{2} + H_{2}O$

 $\begin{array}{c} CH_{3}CH_{2}CH_{2}CH_{2}OH \\ Mechanism \\ CH_{3}-CH_{2}-CH_{2}-OH \end{array} \xrightarrow{AI_{2}O_{3}} CH_{3}CH_{2}CH = CH_{2} \\ \hline H^{+} \\ CH_{3}-CH_{2}-CH_{2}-OH \xrightarrow{H^{+}} CH_{3}-CH_{2}-OH_{2}^{+} \xrightarrow{-H_{2}O} CH_{3}-CH_{2}-CH_{2} \\ \hline \end{array}$ $\begin{array}{ccc} CH_3-CH-CH_2^+ & \xrightarrow{-H^+} & CH_3-CH=CH_2\\ & & 1-propene\\ & H \end{array}$ loss of H₂O

$$\begin{array}{c} OH \\ I_1 & 2 \\ H_3C - C - C - CH_2 - CH_3 \\ I \\ CH_3 \\ 3 \end{array} \xrightarrow{from 1, 2} from 1, 2 \\ Ioss of H_2O \\ from 1, 3 \\ position \end{array} \xrightarrow{CH_3 - C - CH_2 - CH_3} (Major) \\ CH_3 \\ CH_2 \\ CH_2 \\ (Minor) \end{array}$$

✤ Chemical Properties:-

• Addition Reaction:- Alkene show electrophilic addition reaction.

1. Addition of Hydrogen:-

$$RCH = CH_2 \xrightarrow{H_2/Ni} RCH_2CH_3$$

2. Addition of Halogens:-

$$CH_{2} = CH_{2} + Br_{2} \xrightarrow{CCl_{4}} Br Br$$

$$CH_{2} = CH_{2} + Br_{2} \xrightarrow{H_{2}O} Br - CH_{2} - CH_{2} - OH + HBr$$

$$(Brown colour) (Colourless)$$

3. Addition of hydrogen halides-Addition reaction of HBr to symmetrical alkenes

$$CH_2 = CH_2 + H - Br \longrightarrow CH_3 - CH_2 - Br$$

Addition reaction of HBr to unsymmetrical alkenes takes place according to Markovnikov Rule

Markownikov rule:- negative part of the addendum (adding molecule) gets attached to that carbon atom which possesses lesser number of hydrogen atoms. e g



Peroxide effect or Kharasch (Anti Markownikoff's addition):- In 1933 Kharasch and Mayo observed that when HBr is added to an unsymmetrical double bond in the presence of organic peroxide, the reaction take places opposite to the Markovnikov rule.

 $CH_{3}-CH = CH_{2} \xrightarrow{HBr} CH_{3}-CH_{2}-CH_{2}Br$ Propyl bromide $\begin{pmatrix} (1) & O & O \\ II & II \\ C_{6}H_{5}-C-O-O-C-C_{6}H_{5} \xrightarrow{Homolysis} \\ Benzoyl peroxide & O \\ II \\ 2C_{6}H_{5}-C-\dot{O} \stackrel{\cdot}{:} \rightarrow 2\dot{C}_{6}H_{5}+2CO_{2} \\ \end{pmatrix}$

CH₃-CH=CH₂ + Br (iii) Homolysis CH₃-CH-CH₂ CH₃-CH-CH₂-Br Br (b) (a) (less stable (more stable primary free secondary free radical) radical) (iv) $CH_3 - \dot{C}H - CH_2Br + H - Br Homolysis$ $CH_3 - CH_2 - CH_2Br + Br$ (major product)

Noted:- peroxide effect is applicable only to HBr and not to HF, HCl and HI. Addition of HF, HCl and HI takes place according to Markovnikov's rule even in the presence of peroxide.

4. Addition of water (Hydration):- Acid catalyzed addition of water

$$CH_{3}-CH = CH_{2} + H_{2}O \xrightarrow{65-70\% H_{2}SO_{4}} CH_{3}-CH-CH_{3}$$

$$CH_{3}-CH_{2} \xrightarrow{CH_{2}} CH_{2} \xrightarrow{H_{2}O/H} CH_{3}-CH_{3}$$

$$CH_{3}-CH_{2} \xrightarrow{CH_{3}-CH_{3}} CH_{3}$$

$$CH_{3}-CH_{3}-CH_{3}$$

$$CH_{3}-CH_{3}-CH_{3}-CH_{3}$$

$$CH_{3}-CH_{3}-CH_{3}-CH_{3}$$

$$CH_{3}-CH_{3}-CH_{3}-CH_{3}$$

$$CH_{3}-CH_{3}-CH_{3}-CH_{3}$$

$$CH_{3}-CH_{3}-CH_{3}-CH_{3}$$

$$CH_{3}-CH_{3}-CH_{3}-CH_{3}-CH_{3}-CH_{3}$$

$$CH_{3}-$$

2 Oxidation:-

① Combustion:- $CO_2 + H_2O$

O Hydroboration–oxidation:- Alkanes react with diborane to form trialkyl boranes which on oxidation with alkaline H_2O_2 give alcohols.

$$3CH_2 = CH_2 \xrightarrow{BH_3} CH_3CH_2 \xrightarrow{3} B \xrightarrow{H_2O_2/OH^-} 3CH_3CH_2OH$$

Triethylborane Triethylborane

③ Oxymercuration-demercuration:-

$$H_2C = CH_2 \xrightarrow{Hg(OAC)_2} CH_2 - CH_2 - Hg OAC \xrightarrow{NaBH_4} CH_3CH_2OH + Hg$$

④ Oxidation with potassium permanganate:-2KMnO₄ + H₂O → 2KOH + 2MnO₂ + 3[O]

- > This reaction is also called *Hydroxylation*
- Cis product I.e. cis-diol is obtained.

Noted:- The alkaline potassium permanganate solution is known as Baeyer's reagent. It has bright pink colour. It oxidizes alkenes to glycols which is colourless. This reaction is used as a test for the presence of double bond in a molecule. This is also known as Baeyer test.

$$CH_{3}-CH = CH_{2} \xrightarrow{(i) \text{ Alk.KMnO}_{4}} CH_{3} COOH + CO_{2} + H_{2}O$$
(ii) H⁺ Δ

© Oxidation with Ozone:- Ozonolysis – give carbonyls compounds



Noted:- Bromine water test and Baeyer's test are used to detect the presence of double bond while ozonolysis is used to detect the position of double bond.

Alkynes

- The Unsaturated hydrocarbon which have triple bond.
- General molecular formula C_nH_{2n-2}
- sp hybridization
- Those shows chain, positional and functional isomerism
- Preparation:-

From vicinal dihalides: - dehalogenation

$$CH_{3} - CH - CH_{2} \xrightarrow{2KOH (alc)} CH_{3} - C \equiv CH + 2KBr + 2H_{2}O$$

Br Br Br

By the action of water on calcium carbide:-

 $CaC_2 + H_2O \rightarrow HC \equiv CH + Ca(OH)_2$

- Chemical Properties:-
- Addition Reaction:- Alkyne show electrophilic addition reaction.

① Addition of Hydrogen:- Hydrogenation.

$$CH_{3}C \equiv CH + 2H_{2} \xrightarrow{NI} CH_{3}CH_{2}CH_{3}$$

Propyne

Noted:- It may be noted that the hydrogenation can be controlled at the alkene stage only. This is possible by using a *Lindlar's catalysts* or *sodium in liquid* NH_3 at 200k temp..

Noted:- It may be again noted that the catalytic reduction of alkynes in the presence of Lindlar's catalyst gives cis-alkenes while in the presence of sodium in liquid NH₃ (Birch reduction) gives trans-alkenes.



^②Addition of Halogens:-



③ Addition of hydrogen halides:-

 $\mathsf{HC} \blacksquare \mathsf{CH} \ + \ \mathsf{2HBr} \ \longrightarrow \ \mathsf{CH}_3\mathsf{CH} \ \mathsf{Br}_2$

④ Addition of water (Hydration):- Acid catalyzed addition of water

 $HC \equiv CH + H_{2}O \xrightarrow{HgSO_{4}} \left[CH_{2} \equiv CHOH \right] \xrightarrow{CH_{3}CHO} CH_{3}CHO$ Unstable $CH_{3}C \equiv CCH_{3} \xrightarrow{H_{2}O/H^{+}} CH_{3} \xrightarrow{C} CH_{2}CH_{3}$ $CH_{3}C \equiv CCH_{2}CH_{3} \xrightarrow{H_{2}O/H^{+}} CH_{3}CH_{2}CCH_{2}CH_{3} + CH_{3}CCH_{2}CH_{2}CH_{3}$ $CH_{3}C \equiv CCH_{2}CH_{3} \xrightarrow{H_{2}O/H^{+}} CH_{3}CH_{3}CH_{2}CCH_{2}CH_{3} + CH_{3}CCH_{2}CH_{2}CH_{3}$ $CH_{3}C \equiv CCH_{2}CH_{3} \xrightarrow{H_{2}O/H^{+}} CH_{3}CH_{3}CH_{2}CCH_{2}CH_{3} + CH_{3}CCH_{2}CH_{2}CH_{3}$ $CH_{3}C \equiv CCH_{2}CH_{3} \xrightarrow{H_{2}O/H^{+}} CH_{3}CH_{3}CH_{2}CCH_{2}CH_{3} + CH_{3}CCH_{2}CH_{2}CH_{3}$ $CH_{3}C \equiv CCH_{2}CH_{3} \xrightarrow{H_{2}O/H^{+}} CH_{3}CH_{3}CH_{3}CCH_{2}CH_{3} + CH_{3}CCH_{2}CH_{2}CH_{3}$ $CH_{3}C \equiv CCH_{2}CH_{3} \xrightarrow{H_{2}O/H^{+}} CH_{3}CH_{3}CH_{3}CCH_{2}CH_{3} + CH_{3}CCH_{3}CCH_{2}CH_{3}$ $CH_{3}C \equiv CCH_{2}CH_{3} \xrightarrow{H_{2}O/H^{+}} CH_{3}CH_{3}C$

- 5. Polymerisation
 - a. *Linear polymerisation*: of ethyne gives polyacetylene or polyethyne which is a high molecular weight polyene containing repeating units of (CH = CH CH = CH) and can be represented as -(CH = CH CH = CH)n CH = CH
 - b. Cyclic polymerization- results in the formation of aromatic compound.



Acidity of Alkynes- Terminal alkynes are acidic in nature.

HC ≡ CH + Na → HC ≡ C⁻Na⁺ + $\frac{1}{2}$ H₂ Monosodium ethynide CH₃ - C ≡ C - H + Na⁺NH₂⁻ ↓ CH₃ - C ≡ C⁻Na⁺ + NH₃ Sodium propynide

Alkanes, alkenes and alkynes follow the following trend in their acidic behaviour :

1) $HC \equiv CH > H_2C = CH_2 > CH_3 - CH_3$ 11) $HC \equiv CH > CH_3 - C \equiv CH >> CH_3 - C \equiv C - CH_3$ <u>AROMATIC HYDROCARBON</u>

Aromatic compounds containing benzene ring are known as **benzenoids** and those not containing a benzene ring are known as **non-benzenoids**. <u>Structure of Benzene</u>- Kekulé structure



Resonance and stability of benzene-Benzene is a hybrid of various resonating structures.



The orbital overlapping picture benzene- All the six carbon atoms in benzene are sp^2 hybridized and these hybrid orbitals form sigma bonds.



The unhybridised p orbital of carbon atoms are close enough to form a π bond by lateral overlap.



The six π electrons are thus delocalised and can move freely about the six carbon nuclei. The delocalised π electron cloud is attracted more strongly by the nuclei of the carbon atoms than the electron cloud localized between two carbon atoms. Therefore, presence of delocalised π electrons in benzene makes it more stable .

<u>Aromaticity:-</u> The compounds that follow the following features are to be considered aromatic.

(i) Planarity

(ii) Complete delocalisation of the π electrons in the ring

(iii) Presence of $(4n + 2) \pi$ electrons in the ring where *n* is an integer (n = 0, 1, 2, ...). This is often referred to as **Hückel Rule**.

Preparation of Benzene:

(i) Cyclic polymerisation of ethyne:

(ii) Decarboxylation of aromatic acids:



(iii) *Reduction of phenol:* Phenol is reduced to benzene by passing its vapours over heated zinc dust



Physical properties:

- 1. Aromatic hydrocarbons are non- polar molecules and are usually colourless liquids or solids with a characteristic aroma.
- 2. Aromatic hydrocarbons are immiscible with water but are readily miscible with organic solvents.
- 3. They burn with sooty flame.

Chemical properties

Arenes are characterised by electrophilic substitution reactions proceed via the following three steps:

- (a) Generation of the eletrophile
- (b) Formation of carbocation intermediate
- (c) Removal of proton from the carbocation intermediate



benzene ontreatment with excess of chlorine in the presence of anhydrous AlCl3 in dark yields hexachlorobenzene (C_6Cl_6)



Directive influence of a functional group in monosubstituted benzene:-

1. Ortho and para directing groups and activating- –OH, –NH₂, –NHR, – NHCOCH₃, –OCH₃, –CH₃, –C₂H₅, etc.



2. *Meta directing group and deactivating:*-NO₂, -CN, -CHO, -COR, -COOH, -COOR, -SO₃H, etc.



3. Ortho and para directing groups and deactivating- Halogens because of their strong – I effect, overall electron density on benzene ring decreases. However,

due to resonance the electron density on o- and p- positions is greater than that at the *m*-position. Hence, they are also o- and p- directing groups.

CARCINOGENICITY AND TOXICITY-Benzene and polynuclear hydrocarbons containing more than two benzene rings fused together are toxic and said to possess cancer producing (carcinogenic) property.

ONE MARK QUESTIONS

- What are hydrocarbons? Ans. Compounds of hydrogen and carbon.
- 2. What is the general formula of alkanes? Ans. C_nH_{2n+2}
- 3. Write the general formula of alkenes. Ans. C_nH_{2n}
- 4. What is the general formula of alkynes? Ans. C_nH_{2n-2}
- 5. Give the IUPAC name of lowest molecular weight alkane that contains a quaternary carbon.

Ans. 2,2dimethylpropane.

- 6. Arrange the following in the increasing order of C-C bond length- C_2H_6 C_2H_4 C_2H_2 Ans. C_2H_2 < C_2H_4 < C_2H_6
- 7. Out of ethylene and acetylene which is more acidic and why? Ans. Acetylene, due to greater electonegativity of the sp hybrid carbon.
- 8. Name two reagents which can be used to distinguish between ethene and ethyne.

Ans.Tollen's reagent and ammonical CuCl solution.

- Arrange the following in order of decreasing reactivity towards alkanes. HCl, HBr, HI, HF Ans.HI> HBr> HCl >HF
- 10. How will you detect the presence of unsaturation in an organic compound? Ans. Generally Unsaturated organic compound decolourise Bayer's reagent and Bromine water.
- 11.What is Grignard reagent? Ans. Alkyl magnesium halides

TWO MARKS QUESTIONS

1. Write the IUPAC names of the following-



a. CH₂=CH-C≡C-CH₃

Ans. a .Pent-en-3-yne

b. 2-methylphenol

- 2. Write chemical equations for combustion reaction of (i) Butane (ii) Toluene Ans.
 - (i) $2C_4H_{10(g)} + 13O_{2(g)} \longrightarrow 8CO_{2(g)} + 10H_2O_{(g)} + Heat$ Butane

(ii)
$$\overset{\text{CH}_3}{\longmapsto} +9\text{O}_{2(g)} \longrightarrow 7\text{CO}_{2(g)} + 4\text{H}_2\text{O}_{(g)} + \text{Heat}$$

Toluene

- 3. What are the necessary conditions for any system to be aromatic? Ans. A compound is said to be aromatic if it satisfies the following three conditions: (i) It should have a planar structure.
 (ii) The π-electrons of the compound are completely delocalized in the ring.
 - (iii) The total number of π -electrons present in the ring should be equal to (4n + 2), where $n = 0, 1, 2 \dots$ etc. This is known as Huckel's rule.
- 4. What effect does branching of an alkane chain has on its boiling point? Ans. As branching increases, the surface area of the molecule decreases which results in a small area of contact. As a result, the Van der Waals force also decreases which can be overcome at a relatively lower temperature. Hence, the boiling point of an alkane chain decreases with an increase in branching.
- 5. How would you convert the following compounds into benzene?
 - (i) Ethyne (ii) Ethene



(ii) Benzene from Ethene:



- 6. Suggest the name of Lewis acids other than anhydrous aluminium chloride which can be used during ethylation of benzene. Ans. anhydrous FeCl₃, SnCl₄, BF₃ etc.
- 7. Write the name of all the possible isomers of C2H2Cl2 and indicate which of them is non-polar.
 Ans.(i) cis-1,2-dichloroethene (ii) trans-1,2-dichloroethene (iii) 1,1-

dichloroethene. trans-1,2-dichloroethene is non-polar.

8. Although benzene is highly unsaturated, it does not undergo addition reactions, why?

Ans. Because of extra stability due to delocalization of π -electrons.

- 9. What are alkanes? Why are they called paraffins? Ans. Those hydrocarbons which contain single bond between carbon- carbon are called alkanes. They are called paraffins because they are very less reactive (Latin- Parum= little, affins = affinity)
- 10.How can ethene be prepared from (i) ethanol (ii) ethyl bromide? Ans. (i) Ethene from ethanol- by acidic dehydration of alcohols

$$\begin{array}{ccc} H & H \\ H - C & - C \\ | & | \\ & | \\ & | \\ & | \\ & H \\ & OH \\ & Ethanol \\ \end{array} \end{array} \xrightarrow{Conc. H_2SO_4} CH_2 = CH_2 + H_2O \\ & Ethene \\ \end{array}$$

(ii) Ethene from ethyl bromide- by dehydrohalogenation of ethyl bromide $CH_3CH_2Br + KOH (alc) \rightarrow H_2C = CH_2 + KBr + H_2O$

THREE MARKS QUESTIONS

1. What is Wurtz reaction? How can it be used to prepare butane? Ans- When alkyl halides is treated with metallic Na in presence of dry ether, alkanes are formed. This reaction is called Wurtz reaction.

Butane is prepared by the reaction of bromoethane with metallic Na in presence of dry ether

$C_2H_5Br+2Na+BrC_2H_5 \xrightarrow{dry ether} C_2H_5 - C_2H_5$ Bromoethane n-Butane

2. An alkene 'A' contains three C – C, eight C – H σ bonds and one C – C π bond. 'A' on ozonolysis gives two moles of an aldehyde of molar mass 44 u. Deduce IUPAC name of 'A'.

Ans.. The formation of two moles of an aldehyde indicates the presence of identical structural units on both sides of the double bond containing carbon atoms. Hence, the structure of 'A' can be represented as:

XC = CX

There are eight C–H σ bonds. Hence, there are 8 hydrogen atoms in 'A'. Also, there are three C–C bonds. Hence, there are four carbon atoms present in the structure of 'A'.
Combining the inferences, the structure of 'A' can be represented as:



3. In the alkane H3C – CH2 – C(CH3)2 – CH2 – CH(CH3)2, identify 1°,2°,3° carbon atoms and give the number of H atoms bonded to each one of these. Ans.

The given structure has five 1° carbon atoms and fifteen hydrogen atoms attached to it.

The given structure has two 2° carbon atoms and four hydrogen atoms attached to it.

The given structure has one 3° carbon atom and only one hydrogen atom is attached to it

FIVE MARKS QUESTIONS

4. Addition of HBr to propene yields 2-bromopropane, while in the presence of benzoyl peroxide, the same reaction yields 1-bromopropane. Explain and give mechanism

Ans. Addition of HBr to propene is an example of an electrophilic substitution reaction.

Hydrogen bromide provides an electrophile, H^+ . This electrophile attacks the double bond to form 1° and 2° carbocations as shown:



Secondary carbocations are more stable than primary carbocations. Hence, the former predominates since it will form at a faster rate. Thus, in the next step, Br^- attacks the carbocation to form 2 – bromopropane as the major product.



This reaction follows Markovnikov's rule

In the presence of benzoyl peroxide, an addition reaction takes place anti to Markovnikov's rule. The reaction follows a free radical chain mechanism as:



Secondary free radicals are more stable than primary radicals. Hence, the former predominates since it forms at a faster rate. Thus, 1 - bromopropane is obtained as the major product.

 $CH_3 - \dot{C}H - CH_2Br + H - Br$ Homolysis $CH_3 - CH_2 - CH_2Br + \dot{B}r$ 1 - Bromopropanemajor product

HOTS QUESTIONS

- How will you demonstrate that double bonds of benzene are somewhat different from that of olefins? Ans.The double bonds of olefins decolourize bromine water and discharge the pink colour of Bayer's reagent while those of benzene not
- 2. How will you separate propene from propyne? Ans. By passing the mixture through ammonical silver nitrate solution when propyne reacts while propene passes over.
- Write is the structure of the alkene which on reductive ozonolysis gives butanone and ethanol, Ans.-CH₃CH₂C(CH₃)=CHCH₃

CHAPTER 14

ENVIRONMENTAL CHEMISTRY

- Environmental chemistry deals with the study of the origin, transport, reactions, effects, fates of chemical species in the environment.
- ENVIRONMENTAL POLLUTION:-Environmental pollution is the effect of undesirable changes in our surroundings that have harmful effects on plants, animals and human beings. A substance which causes pollution is called a pollutant.they can be solid, liquid or in the gaseous state.
- ATMOSPHERIC POLLUTION:-The atmosphere that surrounds the earth is not of the same thickness at different heights.Atmospheric pollution is generally studied as tropospheric and stratospheric pollution.The ozone layer prevents about 99.5% of the sun's UV rays.
- TROPOSPHERIC POLLUTION:-Tropospheric pollution occurs due to the presence of undesirable solid or gaseous particles in the air. The following are the major gaseous and particulate pollutants present in the troposphere;
 - Gaseous air pollutants: These are oxides of sulphur, nitrogen and carbon, hydrogen sulphide, hydrocarbons, ozone and other oxidants.
 - Particulate pollutants; these are dust, mist, fumes, smoke, smog etc
- GLOBAL WARMING AND GREENHOUSE EFFECT:-About 75% of the solar energy reaching the earth is absorbed by the earth's surface,which increases it's temperature. The rest of the heat radiates back to the atmosphere. Some of the heat is trapped by the gases such as carbon dioxide, methane, ozone, CFCS and Water vapour. they add to the heating of the atmosphere causing Global warming

In a greenhouse, visible light passes through the transparent glass and heats up the soil and the plants. The warm soil and plants emit infrared rays, it partly reflects and partly absorbs these radiations, this mechanism keeps the energy of the sun trapped in the greenhouse.

ACID RAIN:When the pH of the rain water drops below5.6, it is called acid rain.Acid rain is harmful for agriculture, trees and plants as it dissolves and washes away nutrients needed for their growth. It causes respiratory ailments in human beings and animals. When acid rain falls and flows as ground water to reach rivers, lakes etc. it affects plants and animal life in aquatic ecosystem

SMOG:The word smog is derived from smoke and fog.There are two types of smog:classical and photochemical smog. Classical smog occurs in cool humid climate. It is a mixture of smoke, fog and sulphur dioxide. It is also called reducing smog. Whereas photochemical smog occurs in warm and dry sunny climate. It has high concentration of oxidizing agents and therefore ,it is also called as oxidizing smog

OZONE HOLE:Depletion of ozone layer is known as ozone hole.

EFFECTS OF DEPLETION OF THE OZONE LAYER: With the depletion of ozone layer, more UV radiation filters into troposphere. UV radiations lead to ageing of skin, cataract, sunburn, skin cancer, killing of many phytoplanktons, damage to fish productivity etc

WATER POLLUTION:-contamination of water by foreign substances which make it harmful for health of animals or plants or aquatic life and make it unfit for domestic, industrial and agriculture use.

SOURCES/ CAUSES OF WATER POLLUTION-

- Sewage and domestic wastes
- Industrial effluents
- Agriculture effluents
- Siltation-mixing of soil or rock into water
- Thermal pollutants
- Radioactive discharge

EUTROPHICATION: The process in which nutrientenriched water bodies support a dense plantpopulation, which kills animal life by deprivingit of oxygen and results in subsequent loss ofbiodiversity is known asEutrophication

BOD: The amount of oxygen required by bacteria to break down the organic matter present in a certain volume of a sample of water, is called Biochemical Oxygen Demand (BOD)

SOIL POLLUTION:Insecticides, pesticides and herbicides cause soil pollution.

GREEN CHEMISTRY:Green chemistry us a way of thinking and is about utilizing the existing knowledge and principles of chemistry and other sciences to reduce the adverse impact on environment.Green chemistry is a production process that would bring out minimum pollution or deterioration to the environment..Utilization of existing knowledge base for reducing the chemical hazards along with the development of activities is the foundation of green chemistry.

ONE MARK QUESTION

- 1. What is the name of the compound formed when CO combines with blood? Ans:-Carboxyhaemoglobin.
- 2. Which zone is known as ozonosphere? Ans:-Stratosphere.
- 3. Which main gas esis responsible for damage in ozone layer? Ans:-NO and CFCs(freons).
- 4. What is the nature of classical smog? Ans:-Reducing
- 5. Name the acids which are responsible for acid rain? Ans:-H₂SO₄,HNO₃ and HCl.
- 6. List out the gasses which are considered as major source of air pollution? Ans:-Carbon monoxide(CO),sulphur dioxide(SO₂)and oxides of nitrogen(NO₂).
- 7. What is PAN stands for? Ans:-It is peroxyacetyl nitrate.
- 8. Give the examples of insecticides? Ans:-DDT,BHC.
- 9. Which gas is mainly responsible for BHOPAL gas tragedy? Ans:-Methyl isocyanate.
- 10. What should be the tolerable limit of F^- ions in drinking water? Ans:-1ppm or 1mg dm⁻³.

TWO MARKs QUESTION

 What is 'acid rain'? How is it harmful to the environment? Ans:-Acid rain is the rain water mixed with small amount of sulphuric acid, nitric acid along with hydrochloric acid which are formed from the oxides of sulphur and nitrogen present in air as pollutants. It has a pH of 4-5. Harmful effects of acid rain:-It is toxic to vegetation and aquatic life.

It damages buildings and status

- 2. What do youmean byGreen house effect ?What is the role of CO_2 in the greenhouse effect. Ans:-It is the phenomenon in which earth's atmosphere traps the heat from the sun and prevents it from escaping in outer space. Gases such as CO_2 , methane,ozone,CFCs are believed to be responsible for this effect. Heat from the sun after being absorbed by the earth absorbed by CO_2 and then radiated back to the earth.Thus making the environment of the earth warm.
- 3. Which gases are responsible for greenhouse effect?List some of them. Ans:-CO₂ is mainly responsible for greenhouse effect.Other greenhouse gases are methane nitrous oxide, water vapours.
- 4. What is smog?How is classical smogdifferent from photochemical smog? Ans:-The word smog is a combination of smoke andfog.It is a type of air pollution that occurs in many cities throughoutthe world.Classical smog occurs in cool humid climate.It is also called reducing smog.
 Whereas photochemical smog occurs in warm and dry sunny climate. It has high concentrationof oxidizing agents and therefore ,it is also called as
- 5. What are the reactions involved for ozone layer depletion in the stratosphere? Ans:- $CF_2Cl_2(g) + UV \longrightarrow Cl(g) + CF_2Cl(g)$ $Cl(g) + O_3(g) \longrightarrow ClO(g) + O_2(g)$ $ClO(g) + O(g) \longrightarrow Cl + O_2(g)$

oxidizing smog.

- What is the full form of BOD and COD? Ans:-BOD stands for Biochemical Oxygen Demand whereas COD stands for Chemical Oxygen Demand.
- What are viable and non-viable particulates?
 Ans:-Viable particulates:-They are minute living organisms that are dispersed in the atmosphere including bacteria,fungi,moulds,algae etc.
 Non-viable particulates:-They are formed by the breakdownof larger materials or by the condensation of minute particles and droplets.
- 8. What is B.H.C?Give its IUPAC name? Ans:-B.H.C is BENZENE HEXACHLORIDE Its IUPAC name is 1,2,3,4,5,6hexa chlorocyclohexane.
- 9. What is meant by PCBs? Ans:-PCBs are polychlorinated biphenyls. They are contaminates of water. They are used as fluids in transformers and capacitors.

10. What is the compound formed when COcombines with blood? Ans:-When CO combines with blood, the following reaction occurs forming carboxyhaemoglobin:-

 $HB + CO \longrightarrow (HBCO)$ Carboxyhaemoglobin

THREE MARKS QUESTION

 What do you understand by- (i) Mist (ii) Smoke (iii) Fumes Ans(i) Mists:-Mists are produced by particles of spray liquids and the condensation of vapours in air. (ii)Smoke:- They are very small soot particles produced by burning of organic matter.

(iii)Fumes:- These are condensed vapours;fumes of metals are well known particulates of this type.

2. Define the term pesticides? What are three categories of pesticides?

Ans:-Pesticides are substances which are used to kill or block the reproductive process of unwanted organisms.

The three main categories of pesticides are:-

(i)Insecticides:-These are used to control insects and curb diseasesand protect crops.

(ii)Herbicides:-These are used to kill weeds .Example- sodium chlorate

(NaClO₃), sodium arsinite(Na₃AsO₃)

(iii)Fungicides:- These are used to check the growth of fungi. Example-methyl, mercury.

3. What do you mean by ozone hole? What are its consequences?

Ans:-Depletion of ozone hole creates some sort of holes in the blanket of ozone which surround as. This is known as ozone hole.

(i)With the depletion of the ozone layer UV radiation filters into the troposphere which leads to aging of skin, cataract, sunburn etc.

(ii)By killing many of the phytoplanktons it can damage the fish productivity.

4. What are harmful effects of photochemical smog and how can they becontrolled?

Ans:- (i)Photochemical smog causes eye irritation.

(ii)It damages plants (the leaves develop a metallic sheen)

(iii)Rubber on exposure to photochemical smog loses its elasticity and becomes inflexible and brittle.

Usually catalyticconverters are used in the automobiles, whichprevent the release of nitrogen oxide andhydrocarbons to the atmosphere. Certainplants e.g., Pinus, Juniparus, Quercus, Pyrusand Vitis can metabolise nitrogen oxide andtherefore, their plantation could help in thismatter.

5. Give three examples in which green chemistry has been applied.

Ans:-

(i) In dry cleaning , use of liquefied CO_2 in place of tetrachloroethene ($Cl_2C=CCl_2$)

(ii)In bleaching of paper using H_2O_2 in place of chlorine.

(iii)In the manufacture of chemicals like ethanal using environment-friendly chemicals and conditions.