#### KARPAGAM ACADEMY OF HIGHER EDUCATION DEPARTMENT OF CHEMISTRY

# Semester-I 17CHU101 INORGANIC CHEMISTRY I: Atomic structure and chemical bonding 5H 5C Instruction Hours/week:L: 5 T:0 P:0 Marks: Internal: 40 External: 60 Total:100

#### Scope

The course presents the knowledge about the basics of atomic structure and chemical bonding. Quantum mechanical treatment of atomic structure involving quantum numbers and electronic configuration of elements were given. It further presents knowledge about ionic and covalent bonding. The covalent bonding has been explained based on Valence bond theory and molecular orbital theory.

#### **Program Outcome**

- 1. To understand and write the electronic configuration of elements
- 2. To provide a versatile knowledge of different types of bonding like ionic and covalent bonding.
- 3. To provide a knowledge about the valence Bond Theory and Molecular Orbital Theory of covalent bonding

#### **Atomic Structure:**

#### UNIT I

Review of: Bohr's theory and its limitations, dual behaviour of matter and radiation, de Broglie's relation, Heisenberg Uncertainty principle. Hydrogen atom spectra. Need of a new approach to atomic structure. What is Quantum mechanics? Time independent Schrodinger equation and meaning of various terms in it. Significance of  $\psi$  and  $\psi^2$ , Schrödinger equation for hydrogen atom. Radial and angular parts of the hydrogenic wave functions (atomic orbitals) and their variations for 1s, 2s, 2p, 3s, 3p and 3d orbitals (Only graphical representation). Radial and angular nodes and their significance. Radial distribution functions and the concept of the most probable distance with special reference to 1s and 2s atomic orbitals.

#### UNIT II

Significance of quantum numbers, orbital angular momentum and quantum numbers ml and ms. Shapes of s, p and d atomic orbitals, nodal planes. Discovery of spin, spin quantum number (s) and magnetic spin quantum number (ms). Rules for filling electrons in various orbitals, Electronic configurations of the atoms. Stability of half-filled and completely filled orbitals, concept of exchange energy. Relative energies of atomic orbitals, Anomalous electronic configurations.

# Chemical Bonding and Molecular Structure UNIT III

*Ionic Bonding:* General characteristics of ionic bonding. Energy considerations in ionic bonding, lattice energy and solvation energy and their importance in the context of stability and solubility of ionic compounds. Statement of Born-Landé equation for calculation of lattice energy, Born-Haber cycle and its applications, polarizing power and polarizability. Fajan's rules, ionic character in covalent compounds, bond moment, dipole moment and percentage ionic character

# UNIT IV

*Covalent bonding:* VB Approach: Shapes of some inorganic molecules and ions on the basis of VSEPR and hybridization with suitable examples of linear, trigonal planar, square planar, tetrahedral, trigonal bipyramidal and octahedral arrangements. Concept of resonance and resonating structures in various inorganic and organic compounds.

# UNIT V

MO Approach: Rules for the LCAO method, bonding and antibonding MOs and their characteristics for *s*-*s*, *s*-*p* and *p*-*p* combinations of atomic orbitals, nonbonding combination of orbitals, MO treatment of homonuclear diatomic molecules of  $1^{st}$  and  $2^{nd}$  periods (including idea of *s*-*p* mixing) and heteronuclear diatomic molecules such as CO, NO and NO<sup>+</sup>. Comparison of VB and MO approaches.

#### **Suggested Readings:**

#### **Text Books:**

- 1. Lee, J.D. (2010). *Concise Inorganic Chemistry*. ELBS.
- 2. Cotton, F.A., Wilkinson, G. & Gaus, P.L. (2008). *Basic Inorganic Chemistry*. 3rd ed. Hohn Wiley & sons.

#### **Reference Books:**

- 1. Douglas, B.E., McDaniel, D.H. & Alexander, J.J. (2010). *Concepts and Models in Inorganic Chemistry*. John Wiley & Sons.
- 2. Huheey, J.E., Keiter, E.A., Keiter, R.L. & Medhi, O.K. (2006). *Inorganic Chemistry: Principles of Structure and Reactivity*. Pearson Education India.

# **Program Learning Outcome**

- 1. Writing the electronic configuration of elements have been learnt
- 2. Different types of bonding like ionic and covalent bonding have been learnt .
- 3. Studied ad learnt about valence Bond Theory and Molecular Orbital Theory of covalent bonding



Karpagam Academy of Higher Education (Deemed University Established Under Section 3 of UGC Act 1956) COIMBATORE-21 DEPARTMENT OF CHEMISTRY

Name of the Staff	:	Dr. J. Balaji
Department	:	CHEMISTRY
Title of the Paper	:	Atomic Structure and Chemical Bonding
Paper Code	:	17CHU102
Class	:	I- B.Sc Chemistry 'A' Section
Year and Semester	:	2017-18 and I <sup>st</sup> Semester
Total Lecture hours	:	75 Hrs

#### Unit I

# **HOURS REQUIRED -14**

S.No	Lecture hour	Topics to be covered	Support material
1.	1	Bohr's theory and its limitations	T2 - 3-7
2.	1	Dual behaviour of matter and radiation, de Broglie's relation	T2 10-11.
3.	1	Heisenberg Uncertainty principle	T2-15-16
4.	1	Hydrogen atom spectra. Need of a new approach to Atomic structure.	T2-7-9
5.	1	Quantum mechanics Introduction.	T2-21
6.	1	Time independent Schrodinger equation and meaning of various terms in it. Significance of $\psi$ and $\psi^2$ .	T2-22-24
7.	1	Schrödinger equation for hydrogen atom.	T2-5-7
8.	1	Radial and angular parts of the hydrogenic wavefunctions (atomic orbitals)	T2-13-14
9.	1	Variations for 1 <i>s</i> , 2 <i>s</i> , 2 <i>p</i> , 3 <i>s</i> , 3 <i>p</i> and 3 <i>d</i> orbitals (Only graphical representation).	T2-15-16
10.	1	Radial and angular nodes and their significance.	T1-13
11.		Radial distribution functions .	T1-16
12.	1	Concept of the most probable distance with special reference to $1s$ and $2s$ atomic orbitals.	T2-17-21
13.	1	Concept of the most probable distance with special reference to $1s$ and $2s$ atomic orbitals.	T2-17-21
14.	1	Recapitulations and discussion of important questions	

# **References:**

**Text Books:** 

T1 Lee, J.D. (1991). Concise Inorganic Chemistry. ELBS.

T2-R.L. Madan(2015) . Chemistry for Degree students B.Sc., First year. S.Chand & Company Private Ltd., New Delhi.

# **HOURS REQUIRED 14**

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S.No	Lecture hour	Topics to be covered	Support material
1.	1	Significance of quantum numbers	R2-37-39
2.	1	Orbital angular momentum quantum numbers <i>ml</i> and <i>ms</i> .	R2-39
3.	1	Shapes of $s$ , $p$ and $d$ atomic orbitals, nodal planes	T2-20-22
4.	1	Discovery of spin, spin quantum number (s)	T2-9-11
5.	1	Magnetic spin quantum number (ms).	T2-9-11
6.	1	Rules for filling electrons in various orbitals.	T2-25-32
7.	1	Rules for filling electrons in various orbitals.	T2-25-32
8.	1	Electronic configurations of the atoms.	T2-30-32
9.	1	Stability of half-filled and completely filled orbitals.	T2-30-32
10.	1	Stability of half-filled and completely filled orbitals.	T2-30-32
11.	1	Concept of exchange energy	T2-30-32
12.	1	Relative energies of atomic orbitals	T2-167, R2-662
13.	1	Anomalous electronic configurations.	T2-167
14.	1	Recapitulations and discussion of important questions	

#### **References:**

#### **Text Books:**

T2-R.L. Madan(2015) . Chemistry for Degree students B.Sc., First year. S.Chand & Company Private Ltd., New Delhi.

#### **Reference Books:**

R2- B.R.Puri, L.R. Sharma, K.C. Kalia (2002). *Principles of Inorganic Chemistry*. Vallabh Publications, Delhi.

# **HOURS REQUIRED -14**

S.No	Lecture hour	Topics to be covered	Support material
1.	1	General characteristics of ionic bonding.	T2-141-142
2.	1	Energy considerations in ionic bonding	T2-148-150
3.	1	Lattice energy and solvation energy	T2-148-150
4.	1	Importance in the context of stability	T2-148-149
5.		Solubility of ionic compounds	T2-150
6.	1	Born-Landé equation	T2-150-151
7.		calculation of lattice energy	T2-150
8.		Statement of Born-Landé equation for calculation of lattice energy	T2-151
9.	1	Born-Haber cycle and its applications	T2-152-155
10.	1	Polarizing power and polarizability	T2-157
11.	1	Fajan's rules	T2-157-159
12.		Ionic character in covalent compounds.	T2-158
13.	1	Bond moment, dipole moment and percentage ionic character	T2-159
14.	1	Recapitulations and discussion of important questions	

# **References:**

# **Text Books:**

T2-R.L. Madan(2015) . Chemistry for Degree students B.Sc., First year. S.Chand & Company Private Ltd., New Delhi.

#### **HOURS REQUIRED -15**

S.No	Lecture hour	Topics to be covered	Support material
1.	1	Covalent bonding: VB Approach	T2-75
2.	1	Shapes of some inorganic molecules and ions on the basis of VSEPR	R2-187-188
3.	1	Hybridization with suitable examples of linear, trigonal planar arrangements.	R2-187-188
4.	1	Hybridization with suitable examples of linear, trigonal planar arrangements.	R2-187-188
5.	1	VSEPR and hybridization of square planar arrangements.	R2-188
6.	1	VSEPR and hybridization of tetrahedral arrangements	R2-188-189
7.	1	VSEPR and hybridization of tetrahedral arrangements	R2-188-189
8.	1	VSEPR and hybridization of trigonal bipyramidal arrangements	R2-189-190
9.	1	VSEPR and hybridization of octahedral arrangements.	R2-190-191
10.	1	VSEPR and hybridization of octahedral arrangements.	R2-190-191
11.	1	Concept of resonance	T2-79
12.	1	Concept of resonance	T2-79
13.	1	Resonating structures in various inorganic and organic compounds	R1-143-146
14.	1	Resonating structures in various inorganic and organic compounds	R1-143-146
15.	1	Recapitulations and discussion of important questions	

# **References:**

#### **Text Books:**

T1- Lee, J.D. (1991). Concise Inorganic Chemistry. ELBS.

T2-R.L. Madan(2015) . Chemistry for Degree students B.Sc., First year. S.Chand & Company Private Ltd., New Delhi.

# **Reference Books:**

R1 - Huheey, J.E., Keiter, E.A., Keiter, R.L. & Medhi, O.K. (2006). *Inorganic Chemistry: Principles of Structure and Reactivity*. Pearson Education India.

R2- B.R.Puri, L.R. Sharma, K.C. Kalia (2002). *Principles of Inorganic Chemistry*. Vallabh Publications, Delhi.

#### Unit IV

Unit	V
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S.No	Lecture hour	Topics to be covered	Support material
1.	1	MO Approach: Rules for the LCAO method	T1-97-98
2.	1	Bonding and antibonding MOs	T1-112
3.	1	Characteristics for <i>s-s</i> , <i>s-p</i> , <i>p-p</i> combinations of atomic orbitals	T1-90-94
4.	1	Characteristics for <i>s-s</i> , <i>s-p</i> , <i>p-p</i> combinations of atomic orbitals	T1-90-94
5.	1	Nonbonding combination of orbitals	T1-96-97
6.	1	MO treatment of homonuclear diatomic molecules of 1 <sup>st</sup> periods	T1-98-107
7.	1	MO treatment of homonuclear diatomic molecules of 2 <sup>nd</sup> periods	T1-98-107
8.	1	MO treatment of homonuclear diatomic molecules of 1 <sup>st</sup> periods	T1-98-107
9.	1	MO treatment of homonuclear diatomic molecules of 2 <sup>nd</sup> periods	T1-98-107
10.	1	Heteronuclear diatomic molecule such as CO.	T1-109-112
11.	1	Heteronuclear diatomic molecule such as NO.	T1-109-112
12.	1	Heteronuclear diatomic molecule such as NO <sup>+</sup> .	T1-109-112
13.	1	Comparison of VB and MO approaches	T1-128-129
14.	1	Recapitulations and discussion of important questions	
15.	1	Discussion of Previous year ESE Question Paper	
16.	1	Discussion of Previous year ESE Question Paper	
17.	1	Discussion of Previous year ESE Question Paper	
18.	1	Discussion of Previous year ESE Question Paper	

# **References:**

# **Text Books:**

T1- Lee, J.D. (1991). Concise Inorganic Chemistry. ELBS.

# UNIT – I ATOMIC STRUCTURE

Review of: Bohr's theory and its limitations, dual behaviour of matter and radiation, de Broglie's relation, Heisenberg Uncertainty principle. Hydrogen atom spectra. Need of a new approach to atomic structure. What is Quantum mechanics? Time independent Schrodinger equation and meaning of various terms in it. Significance of  $\psi$  and  $\psi$ 2, Schrödinger equation for hydrogen atom. Radial and angular parts of the hydrogenic wave functions (atomic orbitals) and their variations for 1s, 2s, 2p, 3s, 3p and 3d orbitals (Only graphical representation). Radial and angular nodes and their significance. Radial distribution functions and the concept of the most probable distance with special reference to 1s and 2s atomic orbitals.

#### Introduction

If you want to have a language, you will need an alphabet, so alphabets are building blocks of language. Similarly atoms join together to create matter. Atoms are the basis of chemistry, rather, for everything that exists in the universe. These atoms create the elements, molecules and the world in large. The word 'atom' has been derived from the Greek word '**atomos'** which means 'uncut-able' or indivisible. The existence of atom was known to Greek and Indian philosophers as early as 400 B.C. They were of the view that continued subdivision of matter would ultimately yield an atom. According to them atoms were the building blocks of matter and could not be further divided. But they could provide no proof to their hypothesis that "all matter is said to be composed of small particles called atoms". This was proposed by John Dalton, a British school teacher in 1805. His theory called 'Daltons Atomic Theory' regarded the atom as the ultimate particle of matter.

At the end of nineteenth century enough experimental evidence (discharge tube experiment by

J.J. Thomson, E Goldstein) was accumulated to show that an atom is made up of still smaller particles. These sub atomic particles are called the fundamental particles. Although the number of subatomic particles now known is very large, the three most important among them are **proton, neutron** and **electron**. The Central part of the atom is highly dense and is called NUCLEUS. Protons which are positively charged particles and neutrons which are neutral particles reside in the nucleus. Electrons which are negatively charged particles are present in the orbits around the nucleus. The important properties of these fundamental particles are given in Table 1.

Name	Symbol	Year of Discovery	Absolute charge/C	Relative charge	Mass/kg	Mass/u	Approx mass/u
Electron	e	1896	-1.6022x10 <sup>-19</sup>	-1	9.10939x10 <sup>-31</sup>	0.00054	0
Proton	р	1886	+1.6022x10 <sup>-19</sup>	+1	1.67262x10 <sup>-27</sup>	1.00727	1
Neutron	n	1932	0	0	1.67493x10 <sup>-27</sup>	1.00867	1

#### **Table 1: Properties of Fundamental Particles**

The atoms of all known elements contain all the three fundamental particles : electrons, protons and neutrons. The only known exception is hydrogen atom (it contains one electron, one proton but no neutron. The main milestones in the evolution of atomic structures are :

- 1808 Dalton's atomic theory.
- 1896 J.J. Thomson's discovery of electron and proton.
- 1909 Rutherford's nuclear atom.
- 1913 Mosley's determination of atomic number.
- 1913 Bohr Atom
- 1921 Bohr Bury scheme of electronic arrangement
- 1924 De Broglie's wave equation
- 1932 Chadwick's discovery of neutron

With the discovery of sub atomic particles, there arose a need to know how these particles are arranged in the atom. Therefore, different atomic models giving different pictures of the structure of atom have been proposed from time to time. Some of the earlier models are discussed here.

#### **Thomson Model of Atom**

J.J. Thomson, the discoverer of electron in 1898 proposed the **'raisin pudding'** model of atom. He assumed that an atom consists of a uniform sphere of positive charge with electrons embedded into it in such a way as to give it a stable configuration. In this model, the atom is visualized as a pudding or cake of positive charge with raisins (electrons) embedded into it hence, the name 'raisin pudding' model. Mass of the atom was considered to be evenly spread over the atom.

Thomson proposed that electrons in an atom were not at rest, but were vibrating about their equilibrium position. A vibrating electron would emit electromagnetic radiations (spectral lines) having the same frequency as that of vibrating electrons. Zeeman's effect could also be explained on this basis.

#### Limitations of Thomson model:

1. Spectral lines obtained even in the simplest case could not be explained by Thomson model. Hydrogen atom is known to have several lines in ultra violet region. However, according to this model only one spectral line is possible.

2. Rutherford's scattering experiments could not be explained by this model.

3. The mass of the electron has been found to be 1/1837 times that of hydrogen atom. This implied that solid spherical atom proposed by Thomson, especially in case of heavier atoms, must contain thousands of electrons, which is not possible.



Fig. 1. Picture of atom as viewed by Thomson

#### **Rutherford's Experiment**

Lord Rutherford performed an experiment for testing the Thomson model. He bombarded thin foils of gold with high speed  $\alpha$ - particles.

A radioactive substance like polonium which can be a source of  $\alpha$ - particles carries two units of positive charge (+2 charge) and mass equal to about 4 times that of hydrogen atom. The radio

active substance emits  $\alpha$  - particles in all directions. These rays are made to fall on a lead plate with a hole, so that only thin stream of  $\alpha$ -particles comes out of the hole, and the remaining particles are absorbed by the lead plate. These  $\alpha$ -particles bombard a thin gold foil (thickness 4 x 10<sup>-5</sup> cm) layer. A movable screen coated with zinc sulphide is placed next to gold foil. When  $\alpha$ - particles hit the screen, it produces a flash of light which could be counted. The course of  $\alpha$  - particles striking a metallic sheet is represented in Fig 2.



Fig. 2. Rutherford's scattering experiment

#### **Results:**

1) Most of the particles (about 99%) passed straight through the foil and struck the screen at the centre.

2) A small fraction of  $\alpha$ - particles deflected from their original path through varying angles (Mean scattering was about 0-87<sup>0</sup>), small number of  $\alpha$ -particles underwent large deflections and approximately one out of 20,000 made a right angle deflection).

3) Hardly one out of 2000  $\alpha$  - particles bounced back. On the basis of these observations Rutherford come to the following conclusions:



Fig: 3: schematic molecular view of gold foil

1. Atom is extraordinarily hollow i.e. most of the space in the atom is empty as a good number of  $\alpha$ - particles passed through the foil.

2.A few positively charged  $\alpha$  - particles had deflected. The deflection was attributed to enormous repulsive force showing that the positive charge of the atom is not spread throughout the atom as suggested by Thomson. The positive charge has to be concentrated in a very small volume that repelled and deflected the positively charged  $\alpha$ - particles. This very small portion of the atom was called **nucleus** by Rutherford.

3. Calculations from scattering experiment indicated that nucleus of an atom has a diameter of approximately one to six fermis (1 fermi =  $10^{-13}$  cm) and atoms have diameter about 100000 times the size of the nucleus i.e. of the order of  $10^{-8}$  cm.

4. Due to the rigidness of the nucleus, some  $\alpha$ - particles on colliding with it turn back on their original path.

#### **Rutherfod's Model**

Rutherford gave his first model in 1912. He suggested that "an atom consists of a central nucleus of small dimension within which resides the positive charge and most of the mass. Outside this nucleus are electrons to make the atom neutral"

But Rutherford himself realized that his model could not explain the stability of the atom. If the electrons are assumed to be at rest in the atom, the electrons would be attracted by the nucleus and fall inside it. To remove this limitation, he gave another model which states that "the atom consists of a central nucleus surrounded by electrons which are not at rest, but revolve round the nucleus in closed paths like the planets revolving round the sun".

Rutherford called 'electrons' as 'planetary electrons' due to analogy with the planets revolving round the sun. An atom is neutral as the number of electrons equals the number of protons in the nucleus. Later Rutherford concluded that mass number is twice the nuclear charge. He then modified his own model as;

"Atomic nucleus consists of protons and enough electrons to reduce the positive charge to about half the mass number and remaining positive charge on the nucleus was balanced by planetary electrons".

**Effective Radius of Nucleus:** In the scattering experiment the alpha particles bounce back at a point where the kinetic energy is fully converted into potential energy:

Kinetic energy =  $1/2 mv^2$ 

Potential energy of  $\propto$  particle =  $2kZe^2$ (Charge e unit and distance r from nucleus charge Ze)

where  $k = 9 \times 10^9 \text{ Nm}^2 \text{ C}^{-2}$ .

Equating KE. With P.E

$$\frac{1}{2} mv^2 = \frac{2kZe^2}{r}$$

when  $r=2x10^{7}m s^{-1}$ 

mass of  $\propto$  particle (2p+2n) = 6.694x10<sup>-27</sup>kg, Z = 79 for gold atom

r, the effective radius of gold is

$$r = \frac{4kZe^2}{mv^2} = \frac{4 \times 9 \times 10^9 (1.6 \times 10^{-19})^2 \times 79}{6.694 \times 10^{-27} \times (2 \times 10^7)^2}$$

 $r = 2.72 \text{ x } 10^{-14} \text{m}.$ 

The radius found is in agreement with that suggested by Rutherford.

**Drawbacks:** Rutherford had described the atomic model with electrons rapidly revolving around the nucleus as the planets revolve around the sun. When classical mechanics (based on Newtons laws of motion) was applied to the solar system it was found that planets revolve in well defined orbits around the sun. The planetary orbits calculated on the basis of classical mechanics were in agreement with the experimental measurements. Whenever a body is moving in an orbit, it undergoes acceleration. Thus, an electron moving round the nucleus should also undergo acceleration. In accordance with Maxwell's electromagnetic theory, charged particles when accelerated emit electromagnetic radiations and thus lose energy. In other words the orbiting electrons of the Rutherford's model, would continuously emit radiation, and in the process would lose energy, and come closer to the nucleus. Ultimately, the electron would fall into the nucleus and destroy the atom. Niels Bohr, a Danish physicist calculated that an atom would collapse in hundred millionth of a second. This was contrary to the fact that atoms are stable. Another limitation of Rutherford's model was that it was silent about the electronic structure of the atom i.e distribution of electrons around the nucleus and the energies associated with them.

#### **Bohr's Model of Atom**

In 1913, Bohr put forward a theory based on quantization of energy to improve upon the Rutherford's model of the structure of atom. This theory also satisfactorily explained the line spectrum of hydrogen atom. He postulated that:

1. The electrons move around the nucleus in one of the several fixed circular orbits called energy levels. These energy levels are arranged concentrically around the nucleus, and are characterized by an integer n, the lowest level being given the number 1. The energy level corresponding to n=1, 2, 3, 4... are also known as K, L, M, N ... shells.

2. Electrons can move about only in certain orbits which have specific energies. Their movement is possible in only those orbits for which its angular momentum is an integral multiple of  $h/2 \prod$  or mvr=nh/2  $\prod$  where 'n' is any integer 1, 2, 3, 4...n.

3. The energy level nearer to the nucleus has low energy, where as that farthest from it has maximum energy. An electron is said to be in ground state, when it moves in energy level having lowest energy. The ground state is the most stable state of the atom. The electron has a definite energy which is characteristic of the orbit in which it is moving. As long as the electron remains in an orbit, it does not lose energy. These orbits are hence called 'stationary orbits'.

4. Energy is emitted or absorbed when an electron moves from one level to another. Thus by absorbing one particular quantum of energy, the electron will jump from a energy level 1 to 2 or 2 to 3. It is then said to be in excited state. The quantum of energy absorbed in each case is equal to the difference in energies of the two levels. An electron cannot have an energy that would place it in between the two permissible orbits.

5. When an electron moves from a higher energy (E2) orbit to a lower energy (E1) orbit, the

energy ( $\Delta E = E2-E1$ ) is emitted in the form of a photon of frequency v such that  $\Delta E=E2-E1=hv$ 

It is also possible to calculate the radius of each circular orbit using the equation rn = 0.529<sup>o</sup>A (n<sup>2</sup>) where n = 1,2,3 From the above equation it is clear that with an increase in the value of 'n' the value of 'r' will increase i.e. the distance between the nucleus and election increases. The radius of first orbit r1 is called the BOHR RADIUS, -i.e. n = 1; r = 0.529 ° A. Bohr's model is also applicable to other hydrogen like atoms, eg, He<sup>+</sup>, Li<sup>+2</sup> which contain only

one electron but higher number of protons, i.e.  $Z \neq 1$  but may have values 2, 3 .

The total energy of the electron is the sum of the potential and kinetic energies. The kinetic energy of the electron is given by:

K E =  $\frac{1}{2}$  mv<sup>2</sup>, where v is the velocity of the electron and m its mass

 $P E = -k Ze^2/r$  (potential energy of the electron carrying charge -e at a distance r from the nucleus, with charge Ze).

$$\therefore \qquad \frac{kZe^2}{r^2r} = \frac{mv^2}{r^2}$$
$$mv^2 = \frac{kZe^2}{r} \qquad -(2)$$

Putting value of mv<sup>2</sup> from equation 2 in equation

1 The total energy of electron in n<sup>th</sup> orbit will be

En 
$$= \frac{kZe^2}{2r} - \frac{kZe^2}{r} = \frac{kZe^2}{2r}$$

The value of r from Bohr's Quantum theory of hydrogen atom is

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 $r = n^{2}h^{2}$  $4\pi^{2}mZe^{2}$  Where n = 1,2,3 ------

∴ En = -

$$\frac{2\pi^2 k^2}{mZ^2 e}$$
$$\frac{4}{2} n^2 h^2$$

Since for hydrogen Z = 1

$$En = -\frac{2\pi^2 k^2 m e^4}{n^2 h^2}$$

Substituting the values of constants  $\pi = 3.1416$ , k = 9 x 10<sup>9</sup> nm2 C-2, m = 9.1 x 10<sup>-31</sup>kg, e = 1.602 x 10<sup>-19</sup>C and h = 6.626 x 10<sup>-34</sup> Js in above equation

$$E = -\frac{2(3.1416)^2}{(9x10^9)^2} \frac{(9.1x10^{-31})}{(9.1x10^{-31})} \frac{(1.602x10^{19})^4}{(1.602x10^{19})^4}$$
$$= -\frac{2.18 \times 10^{-18}}{n^2} \frac{Z^2}{n^2} = \frac{AZ^2}{n^2}$$

Where A, the constant is  $2.18 \times 10^{18}$  J

$$E_n = \frac{\begin{array}{c} -18 & -1 \\ 2.18 \times 10 - \times Z \text{ atom} \end{array}}{n^2}$$

For example, in case of He<sup>+</sup>, Z= 2, the energy in the 3<sup>rd</sup> orbit will be:

$$E_{3} = \frac{2.18 \times 10 \ 2}{3^{2}} J$$

The above equation shows that the energy of the electron is inversely proportional to the square of n. Thus high the value of n, less negative will be the energy of the electron in it or it can be said that energy of the electron will have more positive value.

Significance of Negative Value of Energy electron at infinity is arbitrarily assumed to be zero. This state is called zero-energy state. When an electron moves and comes under the influence of nucleus, it does some work and sp The energy of an ends its energy in this process. Thus the energy of the electron decreases and it becomes less than zero i.e. it acquires a negative value.

The entitled radiation can have energy conditioned by E = hv

# Limitations of Bohr's Model:

1. It could not explain the spectra of atoms containing more than one electron. It could not be applied to even simple helium atom, which has two electrons.

2. It failed to account for splitting of lines into groups of finer lines as observed by means of spectroscopes of high resolving power.

3. It failed to account for splitting of spectral lines in presence of a magnetic field (Zeeman's effect) or an electric field (Stark effect).

#### **Bohr- Sommerfeld Theory**

When the spectra were examined with spectrometers of high resolving power, each line was found to consist of a group of several finer lines. To explain this limitation of Bohr's model, Sommerfeld extended Bohr's theory. While retaining the first of Bohr's circular orbit as such, he added one elliptical orbit to Bohr's second orbit, two elliptical orbits to Bohr's third orbit and so on. He inferred that like any periodic motion under the influence of central force, it will form an elliptical orbit with the force located at one of the foci.



Fig.4. Sommerfeld orbits in hydrogen atom.

The angular momentum of an electron moving in an elliptical orbit is supposed to be quantized. Thus, only a definite set of values is permissible. Angular momentum is further assumed to be

an integral part of  $\overline{2^{h_{\Pi}}}$  units

 $\therefore$  angular momentum =  $\frac{1}{2^{kh}}\Pi$ 

here 'h' Planck's constant k is azimuthal quantum number

It should be noted here that the quantum number used in Bohr's theory is principal quantum number (n). The two quantum numbers n and k are related by the expression  $n = \underline{lenglh of major axis}$ k lengh of min or axis

When n=k, the orbit will be circular and when k<n, orbit is elliptical.

The values of k for a given value of n are k = n - 1, n-2, n-3 and so on. In other words k will have n possible values (n to 1) for a given value of n.

Although this model successfully explained fine line structure of atomic spectra and introduced the basic concept of stationary states and electronic transitions between them, yet it could not provide a satisfactory picture of the structure of atom.

The structure of atom could only be fully understood by considering the wave mechanical concepts. Two important developments which contributed towards the development of a model free from the limitations of Bohr's model were;

- a) Dual behaviour of matter.
- b) Heisenberg's uncertainty principle.

#### Wave Mechanics

The classical 'Mechanical Theory' of matter considers matter to be made up of discrete particles (electrons, protons etc.), where as the 'wave theory interpreted the nature of radiations like x-ray and light. According to the wave theory, radiations such as x-rays and light consisted of continuous collection of waves traveling in space. This theory however could not explain the photoelectric effect i.e. the emission of electrons from metal surfaces by the action of light. In an attempt to explain the same, Planck and Einstein (1905) proposed that energy radiations, including those of heat and light, are emitted as small packets called quanta or photons. This is referred to as 'Wave Mechanical Theory'. It states that light exhibits both a wave and particle nature, under suitable conditions.

The distinction between particles and waves was very hazy. In 1924, Louis de Broglie suggested in his doctoral thesis that dual character – wave and particle – may not be confined to radiations alone but should be extended to matter as well: i.e matter also possessed particle and wave character. In other words electrons and protons, when in motion, bear wave properties having characteristics of waves such as wave amplitude, frequency and wave length. This concept is

#### called the wave mechanical theory of matter.

Louis de Broglie was born in an aristocratic family and began to study history at the Sorbonne in Paris, as preparation for a career in diplomacy. But after serving as a radio-operator in the First World War, he followed the lead of his brother Maurice and went into physics. He was awarded Nobel Prize in 1929.

#### **De Broglie's Equation**

This gives a relationship between the magnitude of the wave length associated with mass m of a moving body and its velocity.

According to Planck, the photon energy 'E' is given by the equation

E = hv....(i)

Where h is the Planck's constant and v the frequency of radiation.

According to Einsteins mass-energy relationship

 $E = mc^{2}....(ii)$ 

Where E is the energy associated with photon of mass m and c is the velocity of radiation, Comparing equations (i) and (ii).

mc<sup>2</sup> = hv  
Q v = 
$$\lambda^{c}$$
  
mc<sup>2</sup> = h $\lambda^{c}$   
or mc =  $\frac{h}{\lambda}$ ......(iii)....de BROGLIE 'S EQUATION  
 $\lambda$ .

 $mass \times velocity = momentum$  (p)

 $\therefore$  momentum (p) =  $\lambda^{\underline{h}}$ 

The de Broglie s equation can be stated as: the momentum of a particle in motion is inversely proportional to its wave length.

The de Brodlie's equation is true for all particles, but wave characteristic is of significance only for small particles, such as electrons. Large particles in motion do have wavelength, but it is not measurable and hence of no significance.

**Applications:** de Broglie's prediction was confirmed experimentally when it was found that an electron beam undergoes diffraction, a phenomenon associated with waves. This fact has been applied in making electron microscope which is based on wave- like behaviour of electrons. An electron microscope can magnify objects to about 15 million times.

a) For a body with large mass, the value of  $\lambda$  is found to be too small, to be measured by any instrument. Let us consider a body of mass 10 grams (10x10<sup>-3</sup>kg) moving with a velocity of

1000 ms<sup>-1</sup>. The de Broglie s wavelength  $\lambda$  is given as follows:

$$\lambda = \frac{\frac{-3}{5}}{10 \times 10^{-3} \times 1000} = 6.626 \times 10^{-34} \text{ m}$$

This wavelength is too small to be measured and hence is considered inconsequential.

b) A body with small mass eg subatomic particle such as electron, has a significant value of  $\lambda$ , which can be measured with a suitable device. Let us consider an electron of mass 9.1091x10<sup>-31</sup>kg and moving with a velocity of 1.2x10<sup>-8</sup> cm/sec i.e. 1.2 x 10<sup>-10</sup> m/sec. The de-Broglie s wave length  $\lambda$  is given by

$$\lambda = \frac{6.6256 \times 10^{-34}}{9.1091 \times 10^{-31} \times 1.2 \times 10} = 6.0613 \times 10^{-6} m$$

The above value is comparable to the wavelength of x-rays and hence detectable. It can be concluded from the foregoing discussion that everything in nature possesses properties of both the particles and of waves. The properties of large objects are best described by considering the particulate nature, while properties of extremely small bodies like electrons can be described by considering the wave nature.

#### Relationship between de-Broglie's and Bohr's theory

Bohr considered electrons to be moving in circular paths. Since electron has wave characteristic it is assumed that electron wave moves in a circle. The wave motion is considered to be in phase only when two ends of the wave meet to give a regular series of crests and troughs. This condition is possible when the circumference of the circular orbit is equal to the integral multiple of its wavelength.

 $2\prod r = n\lambda$ .....(i).

where r is the radius of the orbit,  $\lambda$  is the wavelength of the electron wave.

 $\lambda = \frac{2\Pi}{n}$ .....(ii).

From de Broglie equation we know that

$$\lambda = \overline{mv^h}$$

Thus

$$\frac{2\Pi_n}{n} = \lambda = \frac{h}{mv}$$
or
$$2\Pi \gamma = \frac{nh}{Mv}$$
nh

$$or \quad mvr = \frac{nn}{2\Pi}$$
(*iii*).



The above equation is same as Bohrs second postulate.

It can be said that electrons can move only in such orbits as have angular momentum an integral multiple of  $h/2\Pi$ . If the circumference is bigger than the value obtained by equation (iii), the wave will not be in phase (Fig. 5b). Thus, de Broglie's equation provides a theoretical basis for Bohr's second postulate.

Three years after de Broglie asserted that particles of matter could possess wavelike properties, the diffraction of electrons from the surface of a solid crystal was experimentally observed by C.J. Davisson and L.H. Germer of the Bell Telephone Laboratory. In 1927 they reported their investigation of the angular distribution of electrons scattered from nickel. They showed that the electron beam was scattered by the surface atoms on the nickel at the exact angles predicted for the diffraction of x-rays according to Bragg's formula, with a wavelength given by the de

Broglie equation,  $\lambda = h / mv$ .



Fig. 6

In 1927, G.P. Thomson, son of J.J. Thomson, reported his experiments, in which a beam of energetic electrons was diffracted by a thin foil. Thomson found patterns that resembled the x-ray patterns made with powdered (polycrystalline) samples. This kind of diffraction, by many randomly oriented crystalline grains, produced rings. When the wavelength of the electrons was changed by altering their incident energy, the diameters of the diffraction rings changed proportionally, as expected from Bragg's equation.

These experiments by Davisson and Germer and by Thomson proved that de Broglie's waves are not simply mathematical conveniences, but have observable physical effects. The 1937 Nobel Prize in Physics was awarded to these scientists for their pioneering work. Just as Compton showed that waves could act like particles, Davisson and Germer showed that particles could act as waves.

#### Heisenberg's Uncertainty Principle

Werner Heisenberg (1927) a German physicist, stated 'uncertainty principle', which is the consequence of dual behaviour of matter and radiation. It states that 'It is impossible to determine simultaneously the exact position and exact velocity of an electron'. This principle is an important feature of wave mechanics and discusses the relationship between a pair of conjugate properties i.e those properties that are interdependent. For example, the position and momentum of a moving particle are interdependent and thus conjugate properties. If the momentum (or velocity) is measured very accurately, the measurement of the position of the particle correspondingly becomes less precise. On the other hand, if the position is determined with accuracy, the measurement of momentum becomes less accurate. Thus, certainty of determination of one property introduces uncertainty in determination of the other. The uncertainty in measurement of position  $\Delta x$  and the uncertainty of determination of momentum  $\Delta p$  (or  $\Delta mv$ ) are related by Heisenberg s relationship as

$$x \cdot p\Delta \Delta^{h} > \text{ constant}$$

$$-\frac{1}{4}$$

This principle applies to all bodies but becomes significant when applied to bodies with small masses and is negligible in case of large objects.

S

The momentum  $\Delta p$  should be along the direction x for the principle to hold good. If the product of uncertainty in position  $\Delta x$  and momentum along the y direction is considered, it will be zero.

i.e 
$$\Delta x \cdot \Delta py = 0$$

This principle is regarded as the fundamental law of nature and is not a limitation of the measuring instrument. However, sophisticated and accurate an instrument may be, it can not give the position and momentum simultaneously with accuracy.

The following example illustrates the principle. If one is asked to measure the thickness of a sheet of paper with an unmarked metrestick, the result obtained would obviously be inaccurate and meaningless. In order to obtain accuracy, one would have to use an instrument graduated in units smaller than the thickness of the sheet. A substance can be seen only if it could reflect light or any other radiation from its surface. Since the size of electron is too small, its position may be

determined by employing light of very small wavelength such as x-rays or  $\gamma$  rays. A photon of radiation of small wavelength has large energy and also momentum. As one such photon strikes an electron, it immediately changes the momentum of the electron. Now as position of

the electron is being determined, the momentum gets changed. Thus it is impossible to determine the exact position of an electron which is moving with a definite velocity.

Experimental verification – A high powered microscope with proper lighting arrangement is set up



Fig: 7

When photons from sources collide with the electrons, some of the electrons bounce into the microscope and enable the observer to see the flash of light and thus to measure both position and momentum of the electron at a particular instant of time. The position of the electron is determined using the following equation of optics, where the resolving power of microscope is given by

$$\Delta = \frac{\lambda}{2\sin\theta}$$

 $\Delta x$ - is minimum distance between two points which can be distinguished as separate. Radiation of short wavelength (x-rays or gamma ray) are preferred to make  $\Delta x$  value very small. Thus  $\Delta x$  is the error or uncertainty in measurement or the position of electron.

 $\lambda$  - wavelength of photon

 $\theta$  - semivertical angle of the cone light coming from illuminated electron.

#### **Quantum Mechanical Model of Atom**

The motion of all macroscopic objects such as falling stone, planets and their satellites having particle like behaviour is best described by classical mechanics based on Newton's laws of motion. However, it fails when applied to microscopic objects like atoms, electrons etc. The reason lies in the fact that classical mechanics ignores the dual behaviour of matter and the uncertainty principle.

Quantum mechanics was developed by Werner Heisenberg and Erwin Schrodinger. It is a theoretical science which deals with the study of the motions of the microscopic objects that have both wave and particle like properties. It specifies the laws of motion that these objects

obey. When applied to macroscopic objects (for which wave like properties are insignificant), the results are the same as those obtained from classical mechanics.

#### **Schrodinger's Wave Equation**

The Schrodinger equation has two 'forms', one in which time explicitly appears, and so describes how the wave function of a particle will evolve in time. In general, the wave function behaves like a wave, and so the equation is often referred to as the time- dependent Schrödinger wave equation. In the other equation time dependence has been 'removed' and it is known as the time- independent Schrödinger wave equation. These are not two separate, independent equations. The time- independent equation can be derived readily from the time-dependent equation.

The time independent equation is written as

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

Here  $\hat{H}$  is the energy operator known as Hamiltonian,  $\Psi$  is the wave function of the system and E is the total energy of the system

$$\hat{H}\Psi = \check{T} + V$$

Hamiltonian operator is the sum of kinetic energy operator  $(\check{T})$  and potential energy operator (V)

Therefore, Schrodinger equation can be written as

$$(\check{T} + V) \Psi = E \Psi$$

By writing the explicit mathematical forms of  $\check{T}$  and V the above equation can be solved to get E and  $\Psi$  for the system.

For any system, the quantum mechanical study consists of following three steps;

- a. writing Schrödinger equation for the system
- b. solving Schrödinger equation to get meaningful solutions of the wave functions and corresponding energies.
- c. calculation of all the observable properties of the system.

The results obtained by solving the Schrodinger equation for a system are in agreement with the experimental findings.

#### Some Comments on Schrodinger Equation:

1. The Schrödinger's wave equation is a second degree differential equation. It has several solutions some of which are imaginary and are not valid. If the potential energy term is known, the total energy 'E' and corresponding wave function  $\psi$  can be evaluated.

- 2.  $\psi$  is called the probability amplitude and is known as the wave function. It describes the behaviour of a particle as a function of x and t. This concept is known as Born interpretation of  $\psi$ ' and refers to a system in which the probability of finding the particle at any point is independent of time. Such a state is called stationary state.
- 3. Inspite of  $\psi$  being complex,  $\psi$  can always be chosen to be real for a 1-D bound state.
- 4. At infinity the value of  $\psi$  is zero. It means that at infinity the electron is not influenced by the attractive force of the nucleus.
- 5.  $\psi$  has only one finite value at any particular point. It means that  $\psi$  is single-valued.
- 6. For any value of potential energy, there will be many solutions (frequently an infinite number) each with a corresponding energy E. But very few solutions describe the known behaviour of electrons. The permitted solutions for  $\psi$  are called eigen functions and corresponding values of E are called eigen values.

**Physical significance of**  $\psi$  - The wave function  $\psi$  gives the amplitude of the wave. The intensity of wave property is known to be proportional to the square of the amplitude of the wave function. Therefore, the intensity of the electron in the atom is proportional to the square of the amplitude  $\psi$ . By intensity of the electron is meant the electron density, i.e if  $\psi^2$  is more near one point than another point in the vicinity of the atom s nucleus, it will be found more often near the point with larger  $\psi^2$  than near the point with less value of  $\psi^2$ . Thus it can be said that  $\psi$  in itself has no physical significance, and all that can be known about the internal structure of matter is through  $\psi^2$ . It helps in locating the relative probability of finding the electron. This uncertainty in finding the electron is in conformity with Heisenberg's uncertainty principle. The exact energy can be found out with the help of equation and hence there would be a great degree of uncertainty in finding the position of the electron.

#### Important Features of the Quantum Mechanical Model of Atom

The picture of the structure of the atom which emerges from the application of Schrodinger's equation is given by Quantum Mechanical model of atom. Its main features are:

1. The electrons in atoms can have only certain specific values, hence it is said that energy of electrons in atoms is quantized.

2. The existence of quantized electronic energy levels is a direct consequence of the wave like properties of electrons.

3. From Heisenberg's uncertainty priniciple, it is established that both the exact position and exact velocity of an electron in an atom cannot be determined simultaneously. Thus the path of an electron in an atom can never be determined, hence the concept of probability of finding the electron at different points in an atom arises.

4. An atomic orbital is the wave function  $\psi$  for an electron in an atom. An electron is said to occupy an orbital, whenever it is described by a wave function. As many such wave functions

are possible for an electron, there are many atomic orbitals in an atom. An orbital can contain a maximum of two electrons. In a multi-electron atom, the electrons are filled in various orbitals in the order of increasing energy. For every electron, there is a orbital wave function characteristic of the orbital it occupies. All the information about the electron in an atom is stored in its orbital wave function  $\psi$  and quantum mechanics makes it possible to extract this information from  $\psi$ .

5. The probability of finding an electron at a point within an atom is proportional to the square of the orbital wave function i.e  $\psi^2$  at that point. This  $\psi^2$  is the probability density and is always positive.

#### **Difference between Orbit and Orbital**

As postulated by Bohr, orbit is a definite circular path around the nucleus in which an electron revolves, whereas an orbital is a three dimensional region in space around the nucleus with in which there is maximum probability of finding an electron having certain energy. An orbit specifies a definite trajectory of an electron in an atom (eg. Bohr suggested that electron of hydrogen atom will always be at a distance of 0.53 <sup>0</sup>A from the nucleus) whereas according to wave mechanics, it is impossible to describe the trajectory of an electron in an atom at any time, and only probability or relative chance of finding an electron with a probability velocity can be started. Thus an orbital does not specify a definite trajectory of an electron in an atom as electron may be found at any distance from the nucleus in the orbital. The total probability of finding the electron in a given volume can be calculated by the sum of all the products  $\psi^2$  and the corresponding volume elements. It is thus possible to get the probable distribution of an electron in an orbital. In an orbit, there is certainty about movement of electron which is in violation of Heisenberg's uncertainty principle, where as the orbital follows the uncertainty principle of Heisenberg.

#### **Quantum Mechanical Treatment of the Hydrogen Atom**

Hydrogen atom is the simplest chemical system as it consists of one proton and one electron. In a coordinate system, nucleus being stationary is taken as the origin and it is assumed that electron moves at a distance r around the stationary nucleus. Schrodinger equation for the hydrogen atom can be written in terms of the Cartesian coordinates (x,y,z) or in terms of the spherical polar

coordinates (r,  $\theta$ ,  $\phi$ ) of the electron with respect to the nucleus.



Fig. 8

Since an atom has spherical symmetry it is more convenient to write Schrodinger equation in terms of polar coordinates. The coordinates x, y and z of electron in terms of polar coordinates are given by:

x =  $r \sin \theta \cos \phi$ y =  $r \sin \theta \sin \phi$ z =  $r \cos \theta$ x<sup>2</sup> + y<sup>2</sup> + z<sup>2</sup> =  $r^2$ 

=

En

For Hydrogen atom, the possible energy states and the corresponding wave function [ $\Psi$ (r,  $\theta$ ,  $\phi$ )] can be obtained on solving Schredinger equation in polar coordinates [ $\Psi$ (r,  $\theta$ ,  $\phi$ )] are known as <u>atomic orbitals</u> The atomic orbital is the product of two factors:

- (1) Radial part dependent on r, and
- (2) Angular part dependent on  $\theta$  and  $\varphi$

The quantized electronic states of the hydrogen atom are given by

 $\frac{\text{mee}^4}{8\epsilon 0^2 \text{h}^2 \text{n}^2}$ 

where me is the mass of electron in kg, e its charge in C, h the Planck s constant, n the principal quantum number and  $\varepsilon 0$  is the permittivity of free space i.e. the ease with which electromagnetic radiation pass through it.

The quantised energy states and corresponding wave functions which are characterized by a set of three quantum numbers (principal 'n', azimuthal 'l' and magnetic 'm') arise as a natural consequence in the solution of Schrodinger equation. The restrictions on the values of three quantum numbers also come naturally from this solution. Thus all aspects of the hydrogen atom spectrum and other phenomena that could not be explained by Bohr model are successfully predicted by quantum mechanical solution of the hydrogen atom.

#### **Shapes of Atomic Orbitals**

The quantum-mechanical treatment of electrons in atoms gives a clear picture of the energy levels associated with every atomic orbital which can be defined by a set of four quantum numbers.

The location of an electron can be described in two ways using the quantum-mechanical results. If the electron is visualized as a very small object moving very rapidly, then the space it occupies can be described in terms of the probability of finding the electron at a given point or within a given space at any instant. If the desired probability is set at 99%, or 95%, a physical space occupied by the electron can be calculated on this time-average basis. If on the other hand, the electron is visualized as an electromagnetic wave, then the amplitude of that wave or the wave function will be greater at some locations than at others. The space the electron occupies can be considered to be the space within which the amplitude of its wave function is greater than 1%, or 5%, of its maximum amplitude.

The electron can be described equally well in either way or the three-dimensional spaces defined as "containing" the electron as a wave and as a particle are the same. Chemists find it

convenient to describe the location of electrons in atoms and molecules in terms of this type of shape. These representations of orbital shapes are those which are said to contain 99% or 95% of the electron density of the orbital.

**s** - **Orbitals** : Orbitals with subshell quantum number l = 0 are called s orbitals. These orbitals are spherical in shape and have spherical symmetry. This means that the wave function will depend only on the distance from the nucleus and not on the direction or on its angular position. The designation 's' refers to the lines in the spectrum as sharp. There can be only one s orbital or sub shell for any specific main energy level and diagrammatically portrayed as a sphere centered at the origin of three mutually perpendicular cartesian axis as depicted in fig 1.8. The figure shows wave function  $(\psi)^2$  against radial distance from the nucleus for hydrogen s orbitals. This plot, which represents the relative probability of finding the electron per unit volume, is difficult to visualize physically because there is much less volume close to the nucleus than further out. It is easier to visualize where the electron may be by plotting the radial probability density  $4(\prod)r^2(\psi)^2$ , rather than  $(\psi)^2$ , against the distance from the nucleus as shown on the right of the Figure.



Fig. 9

The size of the s orbital increases as the principal quantum number of the orbital increases but the geometry remains spherical. The electron density also tends to extend further. Other orbitals behave in the same way as the principal quantum number of the orbitals increase.



Fig.10. s- orbital

**p- Orbitals:** Orbitals with subshell quantum number l = 1 are called p orbitals. Since the magnetic quantum number m can be -1, 0 or +1 when the value of the subshell quantum number l is one, the p orbitals come in sets of three. One of these orbitals is aligned along each of the three mutually perpendicular axes of the atom which are traditionally designated x, y, and z. The three 2p orbitals are correspondingly designated 2px , 2py, and 2pz. The p orbitals, either as a set or individually, do not have spherical symmetry and so a simple plot of radial probability density cannot be made for them. If, however, the distance from the nucleus is taken along any one of the three axes and the orbital is along the same axis, then a suitable plot can be made as shown in the Figure 11 below.



Fig.11

A p orbital has two lobes like a dumb-bell. p refers to designation of spectral lines as 'principal'. The diagrammatic representation of the three separate p orbitals is along the three Cartesian axes X, Y and Z. However, the three individual orbitals collectively considered as superimposed may be totally visualized as concentrically spherical around the origin of the Cartesian axes (fig.12).



Fig. 12

**d- Orbitals :** Orbitals with subshell quantum number l=2 are called d orbitals. Since m can be 2, -1, 0, +1, or +2 when l is two, the d orbitals come in sets of five. These have the same radial function but differ in angular distribution. Four out of five i.e dxy, xyz, dzx and dx2-y2 are similar in shape. These are formed by insertion of two mutually perpendicular planes into spherical pattern dxy lies along the plane XY, dyz lies along the plane YZ and dzx lies along the plane ZX. Their lobes lie in between the axis, making an angle of  $45^{\circ}$  with axes. The orbital dx2 -y2 lies along the plane XY, just as dxy orbital but is related by  $45^{\circ}$  around Z axis, so that the lobes are along X and Y axes and the nodal planes lie midway between the nodal planes of the dxy orbital. The fifth orbital, dz<sup>2</sup> is symmetrical around Z-axis and has two cone shaped nodal surface with the angle between the elements of the cone and Z-axis approximately equal to  $45^{\circ}$ .

The designation 'd' refers to the lines in the spectrum as 'diffuse'



Fig. 13.

**f- Orbitals :** Orbitals with subshell quantum number l=3 are called f orbitals. Since m can be -3, -2, -1, 0, +1, +2, or +3 when 1 has the value 3, f orbitals come in sets of seven. Out of the seven orbitals, two f orbitals possess nodal planes separated by  $60^{\circ}$ ; two others have three mutually perpendicular nodal surfaces and remaining three have two conical and one planar surface each.



Fig. 14

The f orbitals are rarely of direct chemical interest because they tend to be buried deep within the electronic cloud of an atom, but they do play a role in the spectroscopy of the lanthanides and actinides. These are the most complex orbitals with which most chemists have to deal.

#### **Radial Probability Distribution**

The probability of finding an electron within a small radial space around the nucleus is termed as radial probability distribution. The main feature of radial distribution plots is that they determine the spatial extent of the orbit.

Consider the space around the nucleus to be divided into a large number of uniform concentric shells. In between radii r and r+dr the volume of the spherical shell would be  $4\prod r^2 dr$ . Thus the probability of finding an electron within this spherical shell would be  $4\prod^2 \psi r^2$ . This value is the radial probability at a distance r from the nucleus. In the function  $4\prod r^2 dr\psi^2$ , the probability factor  $\psi^2$  decreases whereas the volume factor  $4\prod r^2 dr$  increases with increase in the value of r. The function  $4\prod r^2 dr\psi^2$  when plotted against the distance or from the nucleus gives the radial probability distribution of the electron.





Fig.15 shows the radial distribution functions for 1s, 2s and 3s states of the hydrogen atom, Fig.16 shows the same for 2p, 3p and 3d states of the hydrogen atom.



**Fig. 16** 

It can be noted from the Fig.15 and 16 that:

1. When r=0, the probability of finding the electron is zero, i.e at nucleus, the probability of finding an electron is zero.

2. For a larger value of principal quantum number n, the electron will be farther from the nucleus.

3. The value r0 = 0.529 <sup>0</sup> *A* of maximum radial probability in the radial distribution curve from the nucleus has been found to be in close agreement with the Bohr's calculated value for the radius of first circular orbit.

4. Radial distributions overlap, revealing that the outer electrons penetrate the region occupied by inner electrons in an atom.

#### **Possible Questions**

#### PART A (Objective type each questions carries one mark)

1. The wave associated with particles is called------

a) De Broglie wave b) electromagnetic c) radiowaves d) infrared wave

 Position and velocity of a particle cannot be measured simultaneously with certainty is called------

a)De Broglie relation b) Einstein mass energy relationship c) **Heisenberg uncertainity principle** d) Hund's rule

- 3. Among the conditions which one is correct for the wavefunction  $\psi$  to satisfy and give meaningful solution
  - a)  $\psi$  must be single valued b)  $\psi$  must be discontinuous at allpoints in space c)  $\psi$  must become unity at infinity d)  $\psi$  should not be normalize
- 4. The azimuthal quantum number specifies
  - a) subshell to which an electron belongs to b) orientiation of orbitals in space c)spin of an electrond)The shell to which electron belongs to
- De Broglie wave is associated with----- a)electron b)atom c)molecule d) particles
- 6.  $\lambda = h/p$  is called

a)**De Broglie relation** b) Einstein mass energy relationship c) Heisenberg uncertainity principle

In Schrodinger wave equation, the electron motion is considered to be a-----a)superimposed wave b) stationary wave c)interfringes d)polarized

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- 8. Among the conditions which one is correct for the wavefunction  $\psi$  to satisfy and give meaningful solution
  - a)  $\psi$  must be multiple valued b)  $\psi$  must becontinuous at allpoints in space c)  $\psi$ mustbecome unity at infinity d)  $\psi$  should not be normalize
- 9. The number of radial nodes for 3p orbital is \_\_\_\_\_.
  - a) 3 b) 4 c) 2 d) 1
- 10. The principal quantum number specifies the
- a) subshell to which an electron belongs to b) orientiation of orbitals in space c)spin of an electron d)The shell to which electron belongs to
- 11. The wavelength of the wave associated with any material particle was calculated by analogy with photon is ------

a)**De Broglie relation** b) Einstein mass energy relationship c) Heisenberg uncertainity principle

- 12. Orbital angular momentum depends on \_\_\_\_\_\_.
  - a) **l** b) n and l c) n and m d) m and s
- 13. Photon-----
- a) Value is 4 for N shell b) Probability density c) Always positive value d) Exhibits both
   momentum and wavelength
- 14. The number of radial and angular nodes in 3p subshells are \_\_\_\_\_
- a) **1 and 1 respectively** b)1 and 2 respectively c)2 and 1 respectively d)1 only
- 15. The number of spherical nodes in 3p orbitals is ------
- 16. a)**1** b)2 c)3 d)zero

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- 17. Electron----
  - a) Value is 4 for N shell b) Probability density c) Exhibits both momentum and wavelength d)wavelength only
- 18. Which orbital does not possess angular node
  - a)s b)p c)d d)f
- 19. Among the conditions which one is correct for the wavefunction  $\psi$  to satisfy and give meaningful solution

a) $\psi$  must not be single valued b))  $\psi$  must not be discontinuous at all points in space c) $\psi$ must become zero at infinity d) $\psi$  should not be normalize

- 20. Among the conditions which one is correct for the wavefunction  $\psi$  to satisfy and give meaningful solution
  - a)  $\psi$  must be single valued b)  $\psi$  must be discontinuous at all points in space c)  $\psi$  must become unity at infinity d)  $\psi$  should not be normalize

#### PART B (each questions carries two marks)

- 21. What is the significance of  $\psi$  and  $\psi^2$ .
- 22. Write the de Broglie equation.
- 23. Draw radial probability distribution curves for 2s,3s orbitals.
- 24. Define node.
- 25. What do you understand by the term nodal plane
# **ATOMIC STRUCTURE (2017-18)**

### PART C (each questions carries six marks)

- 26. What do you understand by dual character of matter? Derive the de Broglie relation. How was it verified.
- 27. Explain Radial and angular nodes and their significance.
- 28. State the postulates put forth by Bohr to describe the theory of H-atom.How did Bohr derive the expression for the energy and the velocity of an electron moving in a particular orbit of H-atom.
- 29. Discuss briefly how the radial wave function and angular wave function yield information about the shapes of orbitals.
- 30. Write a note on Heisenberg uncertainity principle
- 31. Draw the radial probability distribution curves for 2s and 2p electrons and explain the difference
- 32. Explain Heisenberg uncertainity principle. Show that this principle is valid only for small particles and not for large objects.
- 33. Derive the time independent Schrodinger equation.

# KARPAGAM ACADEMY OF HIGHER EDUCATION UNIT I (ATOMIC STRUCTURE)

Questions	Opt-1	Opt-2	Opt-3	opt-4	Answer
The wave associated with particles is called	De Broglie wave	electromagnetic wave	radiowaves	infrared wave	De Broglie wave
The wave associated with particles is called	matter wave	electromagnetic wave	radiowaves	infrared wave	matter wave
De Broglie wave is associated with	electron	atom	molecule	particles	particles
Einstein mass energy relationship is	E = mc/2	E = mc2	E/2 = mc	E = m2c	E = mc2
		Einstein mass energy			Einstein mass energy
$E = mc^2$ is called	De Broglie relation	relationship	Heisenbergs principle	Hund's rule	relationship
		Einstein mass energy			
$\lambda = h/p$ is called	De Broglie relation	relationship	Heisenbergs principle	Hund's rule	De Broglie relation
The wavelength of the wave associated with					
any material particle was calculated by		Einstein mass energy			
analogy with photon. This is	De Broglie relation	relationship	Heisenbergs principle	Hund's rule	De Broglie relation
		Einstein mass energy			
$\Delta x.\Delta v = h/4\pi m$ The equation is called	De Broglie relation	relationship	Heisenbergs principle	Hund's rule	Heisenbergs principle
Position and velocity of a particle cannot be					
measured simuntaneously with certainty.		Einstein mass energy			
This is	De Broglie relation	relationship	Heisenbergs principle	Hund's rule	Heisenbergs principle
De Broglie equation has significance only to	microscopic particles	macroscopic particles	rocks	big stones	microscopic particles
	The wavelength	The wavelength			The wavelength
De Broglie equation is applicable and	produced by a larger	produced by a larger	Planck's constant	momentum is not	produced by a larger
siginificant only to microscopic particles	particle is too small to	particle is too large to	changes for a bigger	constant for larger	particle is too small to
because	be observed	be observed	particle	molecules.	be observed
In Schrodinger wave equation, the electron					
motion is considered to be a	superimposed wave	stationary wave	intrifringes	polarised	stationary wave
The number of radial nodes for 3p orbital					
is	3	4	2	1	1
Among the conditions which one is correct		ψ must be			
for the wavefunction $\psi$ to satisfy and give	ψ must be single	discontinuous at all	ψ must become unity	ψ should not be	ψ must be single
meaningful solution	valued	points in space	at infinity	normalise	valued
Among the conditions which one is correct		· · ·			
for the wavefunction $\psi$ to satisfy and give	ψ must be multiple	ψ must be continuous	ψ must become unity	$\psi$ should not be	ψ must be continuous
meaningful solution	valued	at all points in space	, at infinity	normalise	at all points in space

Among the conditions which one is correct		ψ must be			
for the wavefunction $\psi$ to satisfy and give	$\psi$ must not be single	discontinuous at all	$\psi$ must become zero	$\psi$ should not be	$\psi$ must become zero
meaningful solution	valued	points in space	at infinity	normalise	at infinity
Among the conditions which one is correct		ψ must be			
for the wavefunction $\psi$ to satisfy and give	$\boldsymbol{\psi}$ must not be single	discontinuous at all	$\psi$ must become unity	ψ should be	ψ should be
meaningful solution	valued	points in space	at infinity	normalise	normalise
The significant values of the wave function $\psi$					
are called	Eigen functions	Radial function	angular function	Heisenberg function	Eigen functions
Number of angular nodes for 4d orbital is					
	4	3	2	1	2
Orbital angular momentum depends on					
	I	n and I	n and m	m and s	1
					Exhibits both
			Always positive		momentum and
Photon	Value is 4 for N shell	Probability density	value		wavelength
				Exhibits both	Exhibits both
				momentum and	momentum and
Electron	Value is 4 for N shell	Probability density		wavelength	wavelength
There is no difference 2p and a 3p					- C
orbital regarding	shape	size	Energy	value of a	shape
Which orbital does not possess angular	·				
node	s	р	d	f	S
The number of radial and angular nodes					
in 3p subshells are	1 and 1 respectively	1 and 2 respectively	2 and 1 respectively	1 only	1 and 1 respectively
In the atomic spectrum of hydrogen the					
series of lines observed in the visible					
region is	Paschen series	Balmer series	Brackett series	lymann series	Balmer series
	independent of its	Depends on its		equal to square of its	independent of its
The velocity of a photon	wavelength	wavelength	Depends on its source	magnitude	wavelength
Which of the following is not the					
property of the photons	velocity	Energy	Momentum	Rest mass	Rest mass
The Planck's constant has a unit of	work	energy	Linear momentum	angular momentum	angular momentum
Of the following transitions in hydrogen					
atom, the one which gives an absorption					
line of lowest frequency	n=1 to n=2	n=3 to n=8	n=2 to n=1	n=8 to n=3	n=3 to n=8
Transition of electron from $n=3$ to $n=1$	-		-		
level results in	emission spectrum	band spectrum	IR spectrum	X-ray spectrum	emission spectrum

The number of spherical nodes in 3p					
orbitals is	1	2	3	zero	1
Which of the following is expressions					
gives the de Broglie relationship	h/mv=p	$\lambda = h/mv$	$\lambda = h/mp$	λm = v/p	$\lambda = h/mv$
In hydrogen atom when electron jumps				size of the atom	
from 1s to 2s orbital	energy is released	energy is absorbed	atom becomes cation	decreases	energy is absorbed
For azimuthal quantum number l=3, the					
maximum number of electrons will be					
-	2	14	6	10	14
Transition from n=4,5,6 to n=3					
level in hydrogen spectrum gives	Paschen series	Balmer series	Brackett series	lymann series	Paschen series
For which species Bohr's theory does not					
apply	н	He+	Li2+	Ве	Ве
Which represents the correct set of four					
quantum numbers	4,3,2,+1/2	4,2,1,0	4,3,-2,+1/2	4,2, 1,+1/2	4,2, 1,+1/2
Which of the following relates to photon					
both as wave motion and a stream of					
particles	interference	E=mc2	diffraction	E=hv	E=hv
Which electronic level will allow the					
hydrogen atom to absorb photons but not					
to emit	1s	2s	2р	3d	1s
De broglie equation describes the					
relationship of wavelength associated				distance from the	
with motion of an electron and	Mass only	mass and velocity	position	nucleus	mass and velocity
		different energy			
Each subshell of a shell possesses in		levelincreasing energy		decreasing energy	
H atom.	same energy level	level		level	same energy level
The energy of electron at infinite distance					
from the nucleus in Bohr's model is taken					
as	negative	positive	zero	any value	zero
	an electron from a	an electron from an	A proton from a	A neutron from a	an electron from a
Energy will be absorbed in separating	proton	electron	proton	neutron	proton
In the Schrodinger wave equation, $\psi$					
represents	amplitude function	orbital	wave function	all	all
In hydrogen spectrum least energetic					
transition are found in	Pfund series	Balmer series	Brackett series	lyman series	Pfund series

In hydrogen spectrum most energetic					
transition are found in	Pfund series	Balmer series	Brackett series	lyman series	lyman series
Outer orbits have energy than inner					
orbits	greater	smaller	same	equal	greater
	pauli's exclusion	plancks quantum	Heisenberg		Heisenberg
Bohr model of atom is contradicted by?	principle	theory	uncertainity principle	all the three	uncertainity principle
Heisenberg's uncertainity principle is					
applicable to	atoms only	electrons only	nucleus only	any moving object	any moving object
Bohr model of atom explains	zeeman effect	photoelectric effect	stark effect	none of the above	none of the above
Bohr model of atom cannot explains	zeeman effect	stark effect	fine spectrum	all of these	all of these
The de Broglie equation treats an			both a particle and	particle, wave not	both a particle and
electron to be	a particle	a wave	wave	explained	wave
The probability of finding the electron in		zero for some orbitals			
the nucleus is	zero for all objects	and finite for others	finite for all orbitals	infinite	zero for all objects
The electronic configuration of H+ is	1s0	1s1	1s2	1s2 2s1	1s1
The electronic density between 1s and 2s					
orbital is	high	low	zero	two	zero
The uncertainity principle and the					
concept of wave nature of matter was					
proposedbyand respectively	Heisenberg, de Broglie	de Broglie, Heisenberg	Heisenberg, Planck	Planck, Heisenberg	Heisenberg, de Broglie
The probability of finding an electron					
residing in a px orbital is zero	in the yz plane	in the xy plane	in the y direction	in the z direction	in the z direction
		spectrum of any atom			spectrum of any atom
		or ion having one	spectrum of hydrogen		or ion having one
Bohr's model can explain	spectrum of H only	electron only	molecule	solar spectrum	electron only
The transition of electron from					
level results in emission spectrum	n=3 to n=1	n=4 to n=1	n=1 to n=4	n=5 to n=1	n=3 to n=1

# UNIT II

# **QUANTUM NUMBERS**

Significance of quantum numbers, orbital angular momentum and quantum numbers ml and ms. Shapes of s, p and d atomic orbitals, nodal planes. Discovery of spin, spin quantum number (s) and magnetic spin quantum number (ms). Rules for filling electrons in various orbitals, Electronic configurations of the atoms. Stability of half-filled and completely filled orbitals, concept of exchange energy. Relative energies of atomic orbitals, Anomalous electronic configurations.

#### **Quantum Numbers**

The Bohr model was a one-dimensional model that used one quantum number to describe the distribution of electrons in the atom. The only information obtained was the size of the orbit, which was described by principal quantum number 'n'. This could hardly explain the hydrogen spectrum adequately and the spectra of other elements that are quite complex also remained unexplained by this concept. Many single lines of spectra are found to consist of a number of closely related lines when studied with the help of sophisticated instruments of high resolving power. Spectral lines split up when the source of radiation is placed in a magnetic field (Zeeman effect) or in an electric field (Stark effect). Schrödinger's model allowed the electron to occupy three dimensional space. It, therefore, required three coordinates or three QUANTUM NUMBERS, to describe the orbitals in which the electrons can be found. The three coordinates that come from Schrödinger's wave equation are the principal (n), angular (l) and magnetic (m) quantum numbers. In all, there are four identification numbers called quantum numbers which fully describe an electron in an atom. The word quantum is used to signify that all the energy levels which are available to an electron are governed by the laws of quantum mechanics.

**1. Principal Quantum Number:** This quantum number denotes the principal shell to which the electron belongs. This is also referred to as major energy level. This number is designated as 'n' and is identical with Bohr's integers to describe stationary states in an atom. Thus it gives the number of principal shell in which the electron revolves around the nucleus. It designates the average distance of the electron from the nucleus hence specifies the location. As the value of 'n' increases, the energy levels get further away from the nucleus. The value of E increases or becomes less negative.

$$En = = \frac{313.3}{n^2} k \ cals$$

The value of energy of orbit increases with increase in value of n and becomes zero at  $n=\infty$ . Beyond this, the energy is no longer quantized and the electron is not bound to the nucleus and can have any arbitrary amount of kinetic energy. The spectrum in this region does not contain

lines but is continuous and is called continuum. The difference in energy between En-1 and  $En-\alpha$  is the energy needed to ionize the atom and is called the ionization energy.

The principal quantum number 'n' can have non- zero positive integral values from 1 to  $\propto$ . The energy levels 1,2,3,4 etc are also designated by letters K, L, M, N, etc. Although the quantum number 'n' may theoretically assume any integral value from 1 to  $\propto$ , only values

from 1 to 7 have so far been established for atoms of known elements in their ground states. As the first energy level is nearest to the nucleus, it binds strongly to the nucleus. The next energy level is farther from the nucleus and hence binds to the nucleus with a lesser force and so on. Therefore, the principal quantum number signifies:

- a) The size of the electron orbit i.e distance between the nucleus and the electron.
- b) The binding force between the nucleus and the electron.

**2**. **Azimuthal Quantum Number:** This quantum number is also known as subsidiary, angular momentum quantum number (l). Each shell is made up of subshells. The number of subshells in a shell is given by its principal quantum number. Thus 1<sup>st</sup> shell or K has only one sub shell. 2<sup>nd</sup> shell or L has two sub shells. 3<sup>rd</sup> shell or M has 3 subshells. These subshells differ from one another in their azimuthal quantum number (l) values, and hence differ in their shapes.

Values of l=0, 1,2,3..... are denoted by the symbols :s, p, d, f which are derived from the names of the spectral lines sharp, principal, diffuse and fundamental respectively. The s subshell is spherical, p the dumb-bell shaped, d and f have complicated shapes. Thus azimutual quantum number determines the shape of the orbital (electron cloud) i.e whether the cloud is spherical, dumb-bell shaped or has some other complicated shape. In a particular energy level the energies of its subshells are in the order

$$s$$

Thus, s electrons will shield p electrons from the attraction of the nucleus and p electrons will shield d electrons from the nucleus. Therefore, the sequence of filling the various subshells is s, p, d, f and so on.

**3. Magnetic Quantum Number:** This quantum number was introduced to describe the Zeeman's effect i.e splitting of spectral lines under the influence of an applied magnetic field. A shell is made up of one or more subshells and each subshell is made up of one or more orbitals. The number of orbitals in a shell is  $n^2$  where n is the principal quantum number. Each orbital has a characteristic magnetic quantum number and can accommodate up to two electrons.

When an atom is placed in a magnetic field (Zeeman's effect), the orbitals orient themselves relative to the magnetic field. This is due to the presence of negatively charged electrons in it. The orientation in the magnetic field is determined by the magnetic quantum number (m).

Magnetic quantum number is designated as 'm' and can have values from +1 to –1 through zero. Every value of m, for a given value of 1, corresponds to one orbital in that subshell designated by 1. Thus:

- 1) s will have one orbital as m has only one value, m = 0
- 2) p will have three orbitals as m has three values +1, 0, -1.
- 3) d will have five orbitals as m has five values +2, +1, 0, -1, -2).
- 4) f will have seven orbitals as m has seven values +3, +2, +1, 0, -1, -2, -3.

It means that two orbitals in the same subshell can have identical n & l values but must differ in 'm' values.

Thus p subshell is made up of three identical orbitals. But in a magnetic field, the three orbitals take up different positions with respect to the lines of force of field and thus have slightly different energy levels. Therefore, the three 2p orbitals are designated as 2px, 2py and 2pz. Similarly, in a magnetic field, the 3d orbitals are designated as  $dx^2-y^2$ ,  $dz^2$ , dxz, dyz, dxy.

**4. Spin Quantum Number:** The fact that an electron in its motion around the nucleus also rotates or spins about its own axis, gives birth to spin quantum number. This quantum number is designated as 's'. It indicates the direction in which the electron is spinning. Spin can be either clockwise or anticlockwise. Clockwise spin is represented by an arrow pointing upwards ( $\uparrow$ ), while anti-clockwise spin is represented by an arrow pointing downwards ( $\downarrow$ ). The spin of electron contributes to its overall angular momentum and since it can be either clockwise or anticlockwise relative to the direction of the path of electron, it may cause angular momentum to increase or decrease.



Fig. 16

This results in two sub levels of slightly different energy for each value of m, thus corresponding to two possibilities of s. It is assumed that the total energy difference between two sub levels is one quantum i.e. s level will have values  $+\frac{1}{2}$  and  $-\frac{1}{2}$ .

An orbital can accommodate two electrons provided they have opposite spins. Such electrons are known as paired electrons. When an orbital has only one electron, it is known as odd or unpaired electron.

The spin quantum number gives rise to magnetic properties of materials. When the electrons are paired, they have opposite spins and their magnetic moments will be cancelled. Presence of magnetic properties indicates the presence of one or more unpaired electrons.

athematically this quantum number is defined by the equation.

#### **Pauli's Exclusion Principle**

The four quantum numbers ascribed to an electron fully describe its nature, position and energy. Wolfgang Pauli (1925) put forward an ingenious principle which controls the assignment of values of four quantum numbers to an electron. It states that "*No two electrons in a single atom can have all the quantum numbers identical*".

The above statement implies that even if two electrons have the same values for n, l and m, they must have different values of s. It is clear that electrons having the same value of n, l and m i.e. entering the same orbital should have opposite spin. More so, state of opposite spin gives a lower energy state as compared to that parallel spins. According to the thermodynamics, a system having lower energy has maximum stability. Hence the preferred state is the one with electrons having opposite spins. Pauli's principle can also be stated as 'Two electrons can occupy the same orbital only if their spins are opposed'.

#### **Energy Distribution and Orbitals :**

a) One electron atom - Hydrogen is the simplest of all atoms as it has only one electron, which in the ground state occupies the first energy level (n=1). On absorbing energy, it may jump to higher energy levels like 2s, 2p, 3s, 3p etc. Then  $n^2$  levels for any value of 'n' in hydrogen atom have the same energy as shown in the energy level scheme for hydrogen atom.

The spectral study of hydrogen has revealed that its spectral lines correspond to the major energy levels only. Thus the energy value of an electron having a particular quantum is fixed, irrespective of the orbital to which it may belong. It implies that energy associated with



electrons in s, p, d and f orbitals of a particular principal quantum number is the same, e.g. – the energy level of 3s, 3p and 3d orbitals is equal.

Fig. 17

b) Poly-electron atom – In a two electron atom, the second electron differs from the first electron in spin only. It is also accommodated in 1s orbital thereby completing the K shell. This completed K shell affects the energy of the electrons occupying subsequent energy levels. For atoms having more than two electrons, the nuclear charge is shielded from the outer electrons by the two K shell electrons. This makes the energy level of any orbital in a principal level 'n' dependent upon the value of orbital quantum number l. The dependence of energy of orbitals of a shell on l(s, l = 0; p, l = 1) is because of the fact that s orbital electrons (1=0), penetrate near the nucleus and are, therefore, less effectively shielded from the nuclear charge. The s electron being less shielded are drawn inwards and possess lesser energy than porbital electron (l=1). The same argument can be extended for other values of l. Within each energy level 'n', the various sub levels (different l value orbitals) exhibit slightly different energies. The orbitals at a principal level 'n' get split up and come to possess different energies, which increase in the same order as the values of l. Thus, for a particular principal level, the energy of the sub levels is in the order s . The energy levels of 4s, 4p, 4dand 4f orbitals are different even though they belong to the same shell n = 4. However, the energy of electrons in the same orbital is same. Thus all the 4d orbitals (4dxy, 4dyz, 4dzx,  $4dz^2$ ,  $4dx^2-y^2$  or 3p orbitals (3px, 3py and 3pz.) are at the same level of energy, irrespective of their orientation.

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### Hund's Rule of Maximum Multiplicity

On the basis of magnetic measurements, Hund(1925) put forward an empirical rule known after his name as 'Hund's Rule of Maximum Multiplicity'. This states that 'Electrons are distributed among the orbitals of a sub shell in such a way as to give the maximum number of unpaired electrons and hence the same direction of spin'. It implies that the pairing of electrons in any orbital of a sub shell will occur only when all the available orbitals of it (sub shell) have one electron each.

**Examples :** The following examples will illustrate the applications of Hund's rule.

1. Nitrogen has atomic number 7 and hence will have the following configuration.  $1 s^2 2s^2 2p^3$ 



and not



The above configuration also explains the covalency of nitrogen which is three, as three electrons are required for complete pairing of the three electrons present in p-sub shell.

2. Oxygen has atomic number 8 and the following configuration

 $1s^2$ ,  $2s^2$ ,  $2p^4$ 



### 3. Carbon has atomic number 6

In ground state its electronic configuration is 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>2</sup>



4. In Eu (At no = 63) the electronic configuration is  $1s^2$ ,  $2s^2$ ,  $2p^6$ ,  $3s^2$ ,  $3p^6$ ,  $3d^{10}$ ,  $4s^2$ ,  $4p^6$ ,  $4d^{10}$ ,  $4f^7$ ,  $5s^2$ ,  $5p^6$ ,  $6s^2$ .

Here seven f orbitals have to accommodate seven electrons. Hence, each f orbital is singly filled i.e there is no pairing.

5) In Tb (At no = 65) the electronic configuration is  $1s^2$ ,  $2s^2$ ,  $2p^6$ ,  $3s^2$ ,  $3p^6$ ,  $3d^{10}$ ,  $4s^2$ ,

4p<sup>6</sup>, 4d<sup>10</sup>, 4f<sup>9</sup>, 5s<sup>2</sup>, 5p<sup>6</sup>, 6s<sup>2</sup>.

Here seven f orbitals have to accommodate nine electrons. Hence two of these are doubly filled while the remaining five are singly filled. Thus, in these atoms pairing starts only after all the seven f orbitals have been singly filled first.

In excited state the energies of 2s and 2p come further close and hence the four electrons (all in 2s and 2p) divide themselves between four orbitals. Thus covalency of carbon is four.

 Iron has atomic number 26. Electronic configuration according to Aufbau principle will be: 1s<sup>2</sup>, 2s<sup>2</sup>, 2p<sup>6</sup>, 3s<sup>2</sup>, 3p<sup>6</sup>, 4s<sup>2</sup>, 3d<sup>6</sup>

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But empty ( $p^0 d^0$ ), half filled ( $p^3 d^5$ ) and completely filled ( $p^6 d^{10}$ ) orbitals of a sub shell provide symmetry and are more stable as compared to other arrangements. Thus, the other stable configuration for iron will be:  $1s^2$ ,  $2s^2$ ,  $2p^{-6}$ ,  $3s^2$ ,  $3p^6$ ,  $4s^2$ ,  $3d^5$ ,  $4p^1$ 

In former case, the valency of iron will be two and in the latter case, it will be three.

**Explanation:** The three orbitals px, py and pz have equal energy but are concentrated in different regions of space at right angles to each other. Electrons can minimize the repulsive forces between themselves by occupying different orbitals and having parallel spins. This explains their preference to enter different orbitals as long as possible.

Cr (At no = 24)  $1s^2 2s^2 2p^6 3s^2 3p^6 4d^5 4s^1$ 

Mn (At no = 25)  $1s^2 s2^2 2p^6 3s^2 3p^6 4d^5 4s^2$ 

The expected electronic configuration of copper (atomic no. 29) is

Cu At no = 29

 $1s^2 2s^2 2p^6 3s^2 3p^6 3d^9 4s^2$ 

However it is observed that preferred configuration is

 $1s^2 2s^2 2p^6 3s^2 3p^6 3d^{10} 4s^1$ 

The stability of half filled and completely filled orbitals may be explained in terms of lowering of energy due to the exchange energy for a set of two electrons with parallel spins.

According to Hund's rule, electrons avoid entering the same orbital in the given sub shell as far as possible. This reduces coulombic repulsion by enlarging the distance between electrons. Also, the electrons entering the different orbitals of a sub shell have parallel spins. This keeps them further apart and lowers the energy through electron exchange and resonance.

The paramagnetic and diamagnetic character of substances especially transition and rare earth elements can be interpreted with the help of Hund's rule.

### Aufbau Principle

This is also known as 'building up principle', (Aufbau is a German word meaning building up or construction). The principle mainly deals with the building up process in which the extra nuclear electrons are added to various available orbitals so as to balance the nuclear charge. It broadly states that in the ground state of an atom, the electrons tend to occupy the available orbitals in the increasing order of energy, the orbitals of lower energy being filled first. The main postulates of this principle are:

(1) Lowest Energy Rule – When an atom in ground or unexcited state, it has minimum energy. The electrons in this state tend to occupy the lowest energy orbitals available. This implies that:

a) The orbitals with lower n-values are filled prior to orbitals with higher n-values.

b) For any given principal quantum number n, the order of filling is s, p, d and f.

(2) Order of Energy Levels – The different orbitals are filled up in the ascending order of energy. The order is

1s<2s<2p<3s<3p<4s<3d<4p<5s<4d<5p<5s<4f<5d

The correct order of filling of energy can be obtained by considering the sum of principal and subsidiary quantum number (n+1) for the various possible orbitals. The orbital with lowest (n+1) is filled first e.g. -1s(n+1 = 1) is filled before 2s(n+1 = 2). When two orbitals have the same (n+1) value, the one with lower n value is filled first i.e. 3p(n+1 = 4) before 4p(n+1 = 4)

<sup>(3)</sup> Degenerate orbitals – The orbitals having equal energy are called degenerate orbitals. So p has three equivalent energy orbitals or three fold degeneracy, d orbital has 5 fold degeneracy and f has seven fold degeneracy. While filling the orbitals, no differentiation is made between three p, five d or seven f level

(4) Screening Rule – When the inner shells are completely filled with electrons, the outer electrons are screened against attraction by the nucleus. This effect is known as screening effect. As a result of the screening effect, every ns orbital is filled earlier than the corresponding (n-1) d orbital. Hence 4s is filled before 3d.

# **Deviations from Aufbau Principle :**

In some cases, the electronic configuration predicted by Aufbau Principle is different from actual arrangement of electrons as evidenced by spectroscopic methods. This deviation from regularity involves the placing of one or two ns electrons in an (n-1) d level, the energy difference between s or (n-1) d levels being non-significant.

The deviations from regularity may occur due to one of the following reasons:

- a) half filled and completely filled orbitals are more stable than partially filled orbitals,
- b) diamagnetic nature of atom,
- c) to explain the observed properties of elements like lanthanides and actinides.

### **Electronic configuration of elements**

On the basis of Aufbau principle and deviations observed, the electronic configuration of various elements is given in following table:

# **Effective Nuclear Charge**

A hydrogen like atom is an atom that possesses only one electron. Multi electron atoms are characterized by electron- electron repulsion. Unlike a hydrogen like atom, where analytical solutions of Schrödinger equation are available, this equation cannot be solved analytically for multielectron atom owing to the electron-electron repulsion. Interestingly, it is possible to obtain a crude solution for a multi electron atom by making use of a relatively simple construct employing the "effective" nuclear charge in the equations for a hydrogen like atom. The nuclear charge, Z, is the charge of the nucleus. The effective nuclear charge Z\* is the amount of positive charge on the nucleus perceived by an electron. Electrons intervening between the nucleus and an outer electron are said to shield or screen the outer electron from the nucleus so that the outer electron does not experience the full nuclear charge.

The effective nuclear charge  $Z^*$  which is felt by an electron is not the actual charge of the nucleus (Z). J.C. Slater pointed that the amount of nuclear charge actually experienced by an electron depends on the type of orbital and on the ability of other electrons in more penetrating orbitals to screen the electron in question from the nucleus.

When among the orbitals, the value of principal quantum number is same, s orbital is said to be most stable. An orbital with appreciable electron density between the nucleus and a second orbit is said to penetrate the region of space occupied by the second orbital. Thus, the electron is more stable in the more penetrating orbital. The relative extent to which various orbitals penetrate the electron clouds of other orbitals is s>p>d>f. Thus an electron will experience the highest value of effective nuclear charge in a s orbital than a p orbital and so on for any particular value of n i.e. principal quantum number.

It has been observed that when screening of an electron is poor, the effective nuclear charge is correspondingly high. It implies that electrons occupy degenerate set of orbitals in order to experience maximum effective nuclear charge. The above fact is illustrated by the following example. Boron (at no-5) has electron configuration  $1s^2 2s^22p^1$ .

The next element in the periodic table is carbon with atomic no.6 and electronic configuration  $1s^2 2s^2 2p^2$ . Here the new electron is added in accordance with Hund's rule,

Also each new electron will have a new and different effective nuclear charge. For Boron, if the new electron is housed in px orbital, the new electron for C will be housed in another of 2p orbital, say the py orbital. py orbital is perpendicular to the px orbital and is poorly screened from the nuclear charge by px orbital. Thus the effective nuclear charge for the last electron in Carbon is higher than that for Boron. It is the geometry and orientation of the p-orbital that makes them poor at shielding one another from the nucleus. Next to Carbon in the periodic table is Nitrogen (atomic no7) with electronic configuration  $1s^2 2s^2 2p^3$ .

The third p-electron is further poorly screened from the growing nuclear charge because the other two p electrons in px and py orbitals lie at 90°C to this last one. Thus the effective nuclear charge for Carbon is higher than that of Boron and that of Nitrogen is higher than that of Carbon.

# Slater's Rules :

In 1930, J. C. Slater devised a simple set of guidelines for shielding or screening constant (S). He proposed that this empirical constant represents the cumulative extent to which the other electrons of an atom shield (or screen) any particular electron from the nuclear charge. Thus:

$$Z^* = Z$$
-S

Where Z is the atomic number of the atom and  $Z^*$  the effective nuclear charge. The value of effective nuclear charge  $Z^*$  is smaller than Z, since the electron is shielded from Z by an amount equal to S. Thus when screening is small, the effective nuclear charge  $Z^*$  is large.

The value of S for an electron in a given electronic configuration i.e. in presence of other electrons of the atom is calculated using a set of empirical rules given by Slater known as Slater's rules.

If the electron is housed in s or p orbital then

a) All electrons in principal shells higher than the electron in question contribute zero to S.

- b) Each electron in the same principal shell contributes 0.35 to S.
- c) Electrons in the (n-1) shell contribute 0.85 to S
- d) Electrons in deeper shells each contribute 1.00 to S

If the electron is housed in d or f orbital then,

a) All electrons in principal shells higher than the electron in question contribute zero to S.

- b) Each electron in the same principal shell contributes 0.35 to S.
- c) All inner shell electrons i.e.(n-1) and lower uniformly contribute 1.0 to S Thus for 1 s electron S = 0.3

For electrons in an s or p orbital with n > 1, the screening constant is given by

S = 1.00 N2 + 0.85 N1 + 0.35 N0

Where N0 is the number of other electrons in the same shell. N1 is the number of electrons in the next smaller shell (n-1) and N2 is the number of electrons in the next smaller shells (n-2 and smaller).

n	Z	S	$\mathbf{Z}^*$
1 (H)	1	0	1.00
(He)	2	0.35	1.65
2(Li)	3	1.70	1.30
(Be)	4	2.05	1.95
(B)	5	2.40	2.60
(C)	6	2.75	3.25
(N)	7	3.10	3.90
(O)	8	3.45	4.55
(F)	9	3.80	5.20
(Ne)	10	4.15	5.85

# Calculation of effective Nuclear Charge Z\* According to Slater's Equation.

Trends in Z\* become less reliable for heavier elements or in comparisons down a group in periodic table. The Slater rules indicate that electrons in the same shell as the electron of interest reduce the effective nuclear charge by 0.35 even though each electron possesses a full-1 charge. Electrons in the next inner shell (n-1) are more effective in shielding the nucleus, reducing Z\* by 0.85. Electrons closer to the nucleus are completely effective in screening the nucleus (each electron shields a full –1 charge)

# **Spin- Orbit Coupling**

An electron has spin angular momentum and since moving charges generate magnetic fields. The electron has a magnetic moment that arises from its spin. An electron in an orbital with 1>0 is in effect a circulating current and possesses a magnetic moment that arises from its orbital momentum. The interaction of the spin and orbital magnetic moments is called <u>spin-orbit coupling</u>. The strength of the coupling and its effect on the

energy levels of the atom depends on relative orientations of the spin and orbital magnetic moments.

The spin-orbit interaction on the relative orientation of the spin and orbital momenta depends on total angular momentum of the electron, the vector sum of its spin and orbital momenta. Thus when spin and orbital angular momenta are parallel, the angular momentum is high. When the two angular momenta are opposed, the total angular momentum is low.

#### **Possible Questions**

# PART A (Objective type each questions carries one mark)

- 1. The azimuthal quantum number specifies
  - a) subshell to which an electron belongs to b) orientiation of orbitals in spacec)spin of an electron d)The shell to which electron belongs to
- 2. The magnetic quantum number specifies
  - a) subshell to which an electron belongs to b) orientiation of orbitals in spacec)spin of an electron d)The shell to which electron belongs to
- 3. The principal quantum number 'n' can have values
  - a)  $n = 0, 1, 2, 3, \dots, 7$  b)  $n = 1, 2, 3, \dots, n$  infinity c) $n = 0, 1, 2, \dots, n$  infinity d)2l+1
- 4. The magnetic quantum number 'm' can have values
  - a)  $\mathbf{m} = -\mathbf{l}, \dots, \mathbf{0}, \dots, \mathbf{l}$  b) $\mathbf{m} = -2\mathbf{l}, \dots, 0, \dots, +2\mathbf{l}$  c)  $\mathbf{m} = -\mathbf{l}, \dots, \dots, +\mathbf{l}$  d) $\mathbf{m} = -2\mathbf{l}, \dots, +2\mathbf{l}$
- 5. Quantum number gives the orientiation of orbitals in space
  - a) Principal quantum number b) Azimuthal quantum number c) magnetic quantum number d) spin quantum number
- 6. The azimuthal quantum number specifies
  - a) **The angular momentum of an electron** b)orientiation of orbitals in space c)spin of an electron d)The shell to which electron belongs to
- J. Balaji Department of Chemistry, KAHE

- 7. The quantum number which identifies the shell to which electron belongs
  - a) **Principal quantum number** b) Azimuthal quantum number c) magnetic quantum number d) spin quantum number
- 8. If the azimuthal quantum number l = 0, it specifies
  - a) s orbital b) p orbital c) d orbital d) f orbital
- 9. If the azimuthal quantum number l = 1, it specifies
  - a) s orbital b) **p** orbital c) d orbital d) f– orbital
- 10. S orbital have how many orientations
  - a) **One** b) Three c) Five d) Seven
- 11. Principal quantum number represents ------
  - a) orientation of the orbital b) energy and size of orbital c) spin of electron d) shape of the orbital
- 12. If the azimuthal quantum number l = 0, it specifies
  - a) s orbital b) p orbital c) d orbital d) f– orbital
- 13. The number of possible orientiations of an orbital is given by
  - a) **2l + 1** b) 2l 1 c) l2 + 1 d) l2 + 1
- 14. p orbital have how many orientations
  - a) One **b**) **Three** c) Five d) Seven
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15. d – orbital have how many orientations

a)one b)Three c)**five** d)seven

- 16. The magnetic quantum number specifies
  - a) subshell to which an electron belongs to b) orientiation of orbitals in spacec)spin of an electron d)The shell to which electron belongs to
- 17. The principal quantum number 'n' can have values
  - a)  $n = 0, 1, 2, 3, \dots, 7$  b)  $n = 1, 2, 3, \dots, n$  infinity c) $n = 0, 1, 2, \dots, n$  infinity d)2l+1
- 18. The electron has ------ probability of being located at a node.
  - a) **0** b)1 c)2 d)3
- 19. The d orbital with the orientation along X and Y axes is called

a)  $dxy b) dz^2 c) dxz d) dx^2-y^2$ 

- 20. The principal quantum number 'n' can have values
  - a)  $n = 0, 1, 2, 3, \dots, 7$  b)  $n = 1, 2, 3, \dots, n$  infinity c) $n = 0, 1, 2, \dots, n$  infinity d)2l+1

### PART B (each questions carries two marks)

- 21. What are quantum numbers?
- 22. What do you understand by anomalous electronic configurations.
- 23. Give the significance of l and m.
- 24. What is n+1 rule.

- 25. What do you understand by the Quantum number.
- 26. Define spin orbit coupling

# PART C (each questions carries six marks)

- 27. Explain the terms.Principal quantum number,angular momentum number.Explain the significance of each of these numbers.
- 28. Write short notes on concept of exchange energy.
- 29. Explain the terms.magnetic quantum number, spin quantum number. Explain the significance of each of these numbers.
- 30. Write a note on rules for filling electron and electronic configuration of given atom.
- 31. How do the  $d_{x2-y2}$  and  $d_{xy}$  orbitals differ in their orientation in space?
- 32. Explain why in the filling up of atomic orbitals 4s orbital is filled up before 3d orbital.
- 33. Write short notes on Azimuthal quantum number and magnetic quantum number.
- 34. Explain the following giving appropriate reasons:
  - i) Half-filled and completely filled orbitals are more stable.
  - ii)The valence shell configuration of Cr-atom is  $3d^5.4s^1$  and not  $3d^4.4s^2$

### KARPAGAM ACADEMY OF HIGHER EDUCATION UNIT II (QUANTUM NUMBERS)

Questions	Opt-1	Opt-2	Opt-3	opt-4	Answer
The quantum number which identifies the shell to	Principal quantum	Azimuthal quantum	magnetic quantum		Principal quantum
which electron belongs	number	number	number	spin quantum number	number
The quantum number which specifies the subshell	Principal quantum	Azimuthal quantum	magnetic quantum		Azimuthal quantum
to which an electron belongs to	number	number	number	spin quantum number	number
Quantum number gives the orientiation of orbitals	Principal quantum	Azimuthal quantum	magnetic quantum		magnetic quantum
in space	number	number	number	spin quantum number	number
The quantum number which gives the spin of an	Principal quantum	Azimuthal quantum	magnetic quantum		
electron	number	number	number	spin quantum number	spin quantum number
	subshell to which an	orientiation of orbitals in		The shell to which	The shell to which
The principal quantum number specifies the	electron belongs to	space	spin of an electron	electron belongs to	electron belongs to
	subshell to which an	orientiation of orbitals in		The shell to which	subshell to which an
The azimuthal quantum number specifies	electron belongs to	space	spin of an electron	electron belongs to	electron belongs to
	subshell to which an	orientiation of orbitals in		The shell to which	orientiation of orbitals in
The magnetic quantum number specifies	electron belongs to	space	spin of an electron	electron belongs to	space
	subshell to which an	orientiation of orbitals in		The shell to which	
The spin quantum number specifies	electron belongs to	space	spin of an electron	electron belongs to	spin of an electron
	The angular momentum	orientiation of orbitals in		The shell to which	The angular momentum
The azimuthal quantum number specifies	of an electron	space	spin of an electron	electron belongs to	of an electron
The principal quantum number 'n' can have values	n= 0,1,2,37	n= 1,2,3 infinity	n= 0,1,2,infinity	2 +1	n= 1,2,3 infinity
The azimuthal quantum number 'l' can have values	l = 0,1,2,3(n-1)	l = 1,2,3(n-1)	l = 0,1,2,3infinity	l = 1,2,3infinity	l = 0,1,2,3(n-1)
The magnetic quantum number 'm' can have					
values	m = -l,0, +l	m = -2l,0, +2l		m = -l, +l	m = -l,0, +l
The number of possible orientiations of an orbital is					
given by	2l + 1	21 – 1	12 + 1	12 – 1	2l + 1
Principal quantum number represents	orientation of the	energy and size of			energy and size of
	orbital	orbital	spin of electron	shape of the orbital	orbital
	orientation of the	energy and size of			
Azimuthal quantum number tells	orbital	orbital	spin of electron	shape of the orbital	shape of the orbital
^	orientation of the	energy and size of		-	orientation of the
Magnetic quantum number represents	orbital	orbital	spin of electron	shape of the orbital	orbital
	orientation of the	energy and size of		1	
Spin quantum number represents	orbital	orbital	spin of electron	shape of the orbital	spin of electron
The electron has probability of			-r		-r
haing loosted at a node			_		
being located at a node.	0	1	2	3	0
lifethe action that an anti-	a subtact	D. subtal	al a sheka l	f	a sub-ta-l
If the azimuthal quantum number I = 0, it specifies	s – orbital	P-orbital	a- orbital	t-orbital	s – ordital

If the azimuthal quantum number I = 1, it specifies	s – orbital	P-orbital	d- orbital	f-orbital	P-orbital
If the azimuthal quantum number $l = 2$ it specifies	s – orbital	P-orbital	d- orbital	f-orbital	d- orbital
	5 010101			1 Of bitai	
If the azimuthal quantum number I = 3, it specifies	s – orbital	P-orbital	d- orbital	f-orbital	f-orbital
For the S- orbital the azimuthal quantum number					
value is	=1	l = 2	l = 0	= 4	l = 0
For the p- orbital the azimuthal guantum number					
value is	= 1	l = 2	l = 0	= 4	= 1
For the d- orbital the azimuthal guantum number					
value is	= 1	l = 2	l = 0	= 4	= 2
For the S- orbital thequantum number value					
is  =0	azimuthal	principal	magnetic	spin	azimuthal
S – orbital have how many orientations	one	Three	five	seven	one
p - orbital have how many orientations	one	Three	five	seven	Three
d – orbital have how many orientations	one	Three	five	seven	five
f = orbital have how many orientations	one	Three	five	seven	seven
The shape of the s- orbital	Spherical	dumb bell	double dumb bell	square	Spherical
The shape of the p- orbital	Spherical	dumb bell	double dumb bell	square	dumb bell
The shape of the f- orbital	Spherical	dumb bell	double dumb bell	square	double dumb bell
	Spherical			Square	
In an atom no two electrons can have the same					
values for all the quantum numbers proposed by	nauli	hund	aufhau	dalton	nauli
	pauli			daiton	pauli
For a given valus for azimuthal quantum number I,					
the total number of values for m are given by	l+1	2l+1	2l-1	l+2	2l+1
Which shape is associated with orbital designated					
by n=2. l=1	Spherical	tetrahedral	dumb bell	pyramidal	dumb bell
The angular momentum of electro in nth orbit is					
given by	nh	nh/2π	$n^{2}h/2\pi$	h/2 πn	nh/2π
The order of filling of electrons in the orbital of an		,			,
atom will be	3d,4s,4p,4d,5s	4s,3d,4p,5s,4d	5s,4p,3d,4d,5s	3d,4p,4s,4d,5s	4s,3d,4p,5s,4d
The number of vacant d-orbitals in completely				,,,,,,	, , , , ,
excited Cl atom is	2	3	1	5	2
The number of unpaired electron present in Cr3+	3	2	5	1	3
How many electrons can be accomadated in a					
subshell with n=3 and l=1	8	6	18	32	6
The total number of orbitals in a shell with principal		_	_		
quantum number n is	2n	$2n^2$	$n^2$	n+1	$n^2$
The quantum number not obtained from the					
schrodinger's wave equation is	n	I	m	s	S

When 3p orbtals are completely filled then the					
newely entering electron goes in to	4p	3d	4s	4d	4s
The quantum number which determines the shape					
of the orbital is	Principal	Azimuthal	magnetic	spin	Azimuthal
The number of spherical nodes in 3p orbitals are	one	three	four	two	one
Electrons are filled in orbitals according to	n+l rule	Hund's rule	Aufbau's principle	Pauli's exclusion principle	Hund's rule
An electron has magnetic quantum number as -3					
,what is its principal quantum number	1	2	3	4	4
	greater than that of 2px	less than that of 2px		same as that of 2pz	same as that of 2pz
The energy of an electron of 2py orbital is	orbital	orbital	equal to that of 2s orbital	orbital	orbital
The d-orbitals invo;ved in sp3d hybridisation is	dxy	dz2	dx2-y2	dxz	dz2
			equal work -function but	equalenergy but different	
Degenerate orbitals have	equal energy	equal work function	different energy	function	equal energy
		two spherical and one	two non spherical and		
3p orbitals has	two non spherical lobes	non spherical lobes	one spherical lobe	two spherical lobes	two spherical lobes
Which orbital notation does not have spherical					
node	n=2;l=0	n=2;l=1	n=3;l=0	n=4	n=2;l=1
The maximum number of 3d electrons having spin					
quantum number s=+1/2 are	10	14	5	any number from 1to 10	5
The d orbital with the orientation along X and Y					
axes is called	dxy	dzy	dyz	dx2-y2	dx2-y2
What is the number of subshells in principal					
quantum number 3	3	5	9	14	3
The first energy level that can contain d-orbital is	2	3	4	5	3
How many electrons with I=1 will be there in an					
atom having atomic number	2	3	4	5	3
Dumb-bell shape is associated with orbital					
designated by	n=2, l=1	n=3, l=1	n=4, l=2	n=5, l=1	n=2, l=1
Which of the following quantum number does not	Principal quantum	Azimuthal quantum	magnetic quantum		
follow from schrodinger wave equation	number,n	number,l	number,m	spin quantum number,s	spin quantum number,s
The principle which gives a way to fill the electrons					
in the available energy is	Hund's rule	Autbau's principle	Pauli's exclusion principle	n+l rule	Autbau's principle
The orbital with maximum number of possible					_
orientations	s	p	d	f	f

# Unit-III

# **Chemical Bonding – I**

**Ionic Bonding:** General characteristics of ionic bonding. Energy considerations in ionic bonding, lattice energy and solvation energy and their importance in the context of stability and solubility of ionic compounds. Statement of Born-Landé equation for calculation of lattice energy, Born-Haber cycle and its applications, polarizing power and polarizability. Fajan's rules, ionic character in covalent compounds, bond moment, dipole moment and percentage ionic character

# General characteristics of ionic bonding

Ionic compounds have ionic bonds which are electrostatic forces of attraction, they can decompose into respective ions.

Ionic compounds are ionic in nature and good conductor of electricity.

They are usually found in solid state like sodium chloride etc. In ionic compounds, the constituent atoms or ions show strong attractions with other ions.

Ionic compounds generally have high melting points, boiling points and found in solid state.

Due to presence of ions, ionic compounds are polar in nature and easily soluble in polar solvents like water.

Ionic bonding can result from a redox reaction when atoms of an element (usually metal), whose ionization energy is low, give some of their electrons to achieve a stable electron configuration. In doing so, cations are formed. The atom of another element (usually nonmetal), whose electron affinity is positive, then accepts the electron(s), again to attain a stable electron configuration, and after accepting electron(s) the atom becomes an anion. Typically, the stable electron configuration is one of the noble gases for elements in thes-block and the p-block, and particular stable electron configurations for d-block and f-block elements. The electrostatic attraction between the anions and cations leads to the formation of a solid with a crystallographic lattice in which the ions are stacked in an alternating fashion. In such a lattice, it is usually not possible to distinguish discrete molecular units, so that the compounds formed are not molecular in nature. However, the ions themselves can be complex and form molecular ions like the acetate anion or the ammonium cation.

For example, common table salt is sodium chloride. When sodium (Na) and chlorine (Cl) are combined, the sodium atoms each lose anelectron, forming cations (Na<sup>+</sup>), and the chlorine atoms each gain an electron to form anions (Cl<sup>-</sup>). These ions are then attracted to each other in a 1:1 ratio to form sodium chloride (NaCl).

$$Na + Cl \rightarrow Na^+ + Cl^- \rightarrow NaCl$$

However, to maintain charge neutrality, strict ratios between anions and cations are observed so that ionic compounds, in general, obey the rules of stoichiometry despite not being molecular compounds. For compounds that are transitional to the alloys and possess mixed ionic and metallic bonding, this may not be the case anymore. Many sulfides, e.g., do form nonstoichiometric compounds.

Many ionic compounds are referred to as **salts** as they can also be formed by the neutralization reaction of an Arrhenius base like NaOH with an Arrhenius acid like HCl

 $NaOH + HCl \rightarrow NaCl + H_2O$ 

The salt NaCl is then said to consist of the acid rest Cl<sup>-</sup> and the base rest Na<sup>+</sup>.

The removal of electrons from the cation is endothermic, raising the system's overall energy. There may also be energy changes associated with breaking of existing bonds or the addition of more than one electron to form anions. However, the action of the anion's accepting the cation's valence electrons and the subsequent attraction of the ions to each other releases (lattice) energy and, thus, lowers the overall energy of the system.

Ionic bonding will occur only if the overall energy change for the reaction is favorable. In general, the reaction is exothermic, but, e.g., the formation of mercuric oxide (HgO) is endothermic. The charge of the resulting ions is a major factor in the strength of ionic bonding, e.g. a salt  $C^+A^-$  is held together by electrostatic forces roughly four times weaker than

 $C^{2+}A^{2-}$  according to Coulombs law, where C and A represent a generic cation and anion respectively.

### Statement of Born-Landé equation for calculation of lattice energy:

For a solid crystalline ionic compound the enthalpy change in forming the solid from gaseous ions is termed the lattice energy. The experimental value for the lattice energy can be determined using the Born-Haber cycle. It can also be calculated (predicted) using the Born-Landé equation as the sum of the electrostatic potential energy, calculated by summing interactions between cations and anions, and a short-range repulsive potential energy term. The electrostatic potential can be expressed in terms of the inter-ionic separation and a constant (Madelung constant) that takes account of the geometry of the crystal. The further away from the nucleus the weaker the shield. The Born-Landé equation gives a reasonable fit to the lattice energy of, e.g., sodium chloride, where the calculated (predicted) value is -756 kJ/mol, which compares to -787 kJ/mol using the Born-Haber cycle.

### The Born-Lande' equation

The Born-Landé equation is a concept originally formulated in 1918 by the scientists Born and Lande and is used to calculate the lattice energy (measure of the strength of bonds) of a compound. This expression takes into account both the Born interactions as well as theCoulomb attractions.

### Introduction

Due to its high simplicity and ease, the Born-Landé equation is commonly used by chemists when solving for lattice energy. This equation proposed by Max Born and Alfred Landé states that lattice energy can be derived from ionic lattice based on electrostatic potential and the potential energy due to repulsion. To solve for the Born-Landé equation, you must have a basic understanding of lattice energy:

- Lattice energy decreases as you go down a group (as atomic radii goes up, lattice energy goes down).
- Going across the periodic table, atomic radii decreases, therefore lattice energy increases.

# Equation

The Born-Landé equation was derived from these two following equations. the first is the electrostatic potential energy:

 $\Delta U = -LA|Z + ||Z - |e24\pi\epsilon or$ 

with

- L is Avogadro's constant (6.022×1023)
- A is the Madelung Constant (a constant that varies for different structures)
- e is the charge of an electron  $(1.6022 \times 10 19 \text{ C})$
- Z+ is the cation chargeZ- is the anion charge
- $\epsilon$ o is the permittivity of free space
- The second equation is the repulsive interaction:

 $\Delta U=LBrn$  with

- B is the repulsion coefficient and
- N is the Born Exponent (typically ranges between 5-12) that is used to measure how much a solid compresses

These equations combine to form:

- $\Delta U(0K) = LA|Z + ||Z |e24\pi \epsilon oro(1-1n)$  with
- $r_0$  = closest ion distance

# **Calculate Lattice Energy**

Lattice energy, based on the equation from above, is dependent on multiple factors. We see that the charge of ions is proportional to the increase in lattice energy. In addition, as ions come into closer contact, lattice energy also increases.

# The Born-Haber cycle

Thermodynamics may be defined as the branch of science which deals with the quantitative relationship between heat and other forms of energies. **Chemical thermodynamics deals with the change in heat during chemical processes. The energy change during the chemical reaction at constant temperature and volume is given by internal energy change.** However chemical reactions are generally involved some change in volume, therefore in order to study the heat changes of chemical reaction at constant temperature and pressure, a new function enthalpyis introduced.

Enthalpy is the total energy associated with any system which includes its internal energy and also energy due to environmental factor such as pressure-volume conditions. Mathematically enthalpy can be written as

### $\mathbf{H} = \mathbf{U} + \mathbf{P}\mathbf{V}$

Where U stands for internal energy and PV is additional energy possesses by the substance. Enthalpy is a state function and an extensive property. It is also known as heat content of the system whose value depends upon amount and chemical nature of substance as well as on pressure and temperature.

### **Born Haber Cycle Example**

Born Haber cycle is a simplified method developed by Max Born and Fritz Haber in 1919 to correlate the lattice energies of ionic solids to other thermodynamic data. Lattice enthalpy or

lattice energy is defined as enthalpy change which occurs when one mole of ionic solid is formed by close packing of constituent ions in gaseous state. Or it may be defined as energy released when mole of ionic solid dissociates one into its gaseous state. For example, the formation of sodium chloride from sodium and chloride ion involves 788 kj/mol of energy and dissociation of sodium chloride into it constituent ions requires same amountof energy.

NaCl(s) 
$$\rightarrow$$
 Na+(g) + Cl- (g)  $\triangle$ LH°= 788 kj/mol  
Na+(g) + Cl-(g)  $\rightarrow$  NaCl(s)  $\triangle$ LH° = -788 kj/mol

When oppositely charged ions interact to form ionic solid, a large amount of energy is released and dissociation of crystal also require a lot of energy which makes melting point and boiling points of crystal very high. Born Haber cycle is used to determine this lattice energy by using other energy values like ionization energy, electron affinity, dissociation energy, sublimation energy and heat of formation.

**1. Ionization Energy:** It is the energy required to remove an electron from a neutral gaseous atom or an ion.

**2. Electron Affinity:** It is the energy released when an electron is added to an isolated neutral gaseous atom or an ion.

**3. Dissociation energy:** The energy required to dissociate a compound is called as dissociation energy. Dissociation of a compound is always an endothermic process and requires an input of energy.

**4. Sublimation energy:** The energy required to change the phase from solid to gas, by passing the liquid phase is called as sublimation energy.

**5. Heat of formation:** The energy change during the formation of a compound from its elements is known as heat of formation.

Born Haber cycle is based on Hess law which states that the standard enthalpy change of reactions is the sum of algebraic sum of the standard enthalpies of reactions into which the overall reaction may be splitted or divided. **For example,** the formation of C from A and B can take place in two ways. Both ways involve same initial and final states, but one is single step and other one is multi-step reaction which involves intermediates, D and E. Both ways has an enthalpy change of 160 kJ.



The formation of ionic compounds from alkali metals like sodium, potassium with halogen such as chlorine, bromine to form alkali halide can be represent by using Born Haber cycle. Remember the heat of formation of ionic halide is equals to the sum of other energies involves in various steps.


# Born - Haber Cycle

 $\Delta H_{f^{\circ}} = \Delta H_{sub} + IE + \Delta H_{diss} + EA + U$ 

# Lattice Energy: The Born-Haber cycle

Ionic solids tend to be very stable compounds. The enthalpies of formation of the ionic molecules cannot alone account for this stability. These compounds have an additional stability due to the lattice energy of the solid structure. However, lattice energy cannot be directly measured. The Born-Haber cycle allows us to understand and determine the lattice energies of ionic solids.

# Lattice Energy

Lattice Energy is a type of potential energy that may be defined in two ways. In one definition, the lattice energy is the energy required to break apart an ionic solid and convert its component atoms into gaseous ions. This definition causes the value for the lattice energy to always be positive, since this will always be an endothermic reaction. The other definition says that lattice energy is the reverse process, meaning it is the energy released when gaseous ions bind to form

an ionic solid. As implied in the definition, this process will always be exothermic, and thus the value for lattice energy will be negative. Its values are usually expressed with the units kJ/mol. Lattice Energy is used to explain the stability of ionic solids. Some might expect such an ordered structure to be less stable because the entropy of the system would be low. However, the crystalline structure allows each ion to interact with multiple oppositely charge ions, which causes a highly favorable change in the enthalpy of the system. A lot of energy is released as the oppositely charged ions interact. It is this that causes ionic solids to have such high melting and boiling points. Some require such high temperatures that they decompose before they can reach a melting and/or boiling point.

### Fajans' Rules:

Rules formulated by Kazimierz Fajans in 1923, can be used to predict whether a chemical bond is expected to be predominantly ionic or covalent, and depend on the relative charges and sizes of the cation and anion. If two oppositely charged ions are brought together, the nature of the bond between them depends upon the effect of one ion on the other.



Covalent	Ionic
Small cation (< ~100	Large cation (> ~100
pm)	pm)
Large anion	Small anion
High positive charges	Low positive charges

Although the bond in a compound like  $X^+Y^-$  may be considered to be 100% ionic, it will always have some degree of covalent character. When two oppositely charged ions (X<sup>+</sup> and Y<sup>-</sup>) approach each other, the cation **attracts** electrons in the outermost shell of the anion but **repels** the positively charged nucleus. This results in a distortion, deformation or polarization of the anion. If the degree of polarization is quite small, an ionic bond is formed, while if the degree of polarization is large, a covalent bond results.

The ability of a cation to distort an anion is known as its polarization power **and the tendency of the anion to become polarized by the cation is known as its** polarizability.

The polarizing power and polarizability that enhances the formation of covalent bonds is favoured by the following factors:

**Small cation:** the high polarizing power stems from the greater concentration of positive charge on a small area. This explains why LiBr is more covalent than KBr (Li<sup>+</sup> 90 pm cf. K<sup>+</sup> 152 pm).

**Large anion:** the high polarizability stems from the larger size where the outer electrons are more loosely held and can be more easily distorted by the cation. This explains why for the common halides, iodides, are the most covalent in nature (I<sup>-</sup> 206 pm).

Large charges: as the charge on an ion increases, the electrostatic attractions of the cation for the outer electrons of the anion increases, resulting in the degree of covalent bond formation increasing.

Large cations are to be found on the bottom left of the periodic table and small anions on the top right. The greater the positive charge, the smaller the cation becomes and the *ionic potential* is a measure of the charge to radius ratio.

Polarization will be increased by:

High charge and small size of the cation

Ionic potential Å Z+/r+ (= polarizing power)

High charge and large size of the anion

The polarizability of an anion is related to the deformability of its electron cloud (i.e. its

"softness")

An incomplete valence shell electron configuration

Noble gas configuration of the cation produces better shielding and less polarizing power

e.g.  $Hg^{2+}$  (r+ = 102 pm) is more polarizing than  $Ca^{2+}$  (r+ = 100 pm)



On the left, the cation charge increases (size decreases) and on the right, the anion size increases, both variations leading to an increase in the covalency. Thus covalency increases in the order:

 $[Na^{+} Cl^{-}, NaCl] < [Mg^{2+} 2(Cl)^{-}, MgCl_{2}] < [Al^{3+} 3(Cl)^{-}, AlCl_{3}] \text{ and}$  $[Al^{3+} 3(F)^{-}, AlF_{3}] < [Al^{3+} 3(Cl)^{-}, AlCl_{3}] < [Al^{3+} 3(Br)^{-}, AlBr_{3}]$ 

Electronic configuration of the cation: for two cations of the same size and charge, the one with a pseudo noble-gas configuration (with 18 electrons in the outer-most shell) will be more polarizing than that with a noble gas configuration (with 8 electrons in the outermost shell). Thus zinc (II) chloride (Zn(II)  $1s^2 2s^2 2p^6 3s^2 3p^6 3d^{10}$  and Cl<sup>-</sup>  $1s^2 2s^2 2p^6 3s^2 3p^6$ ) is more covalent than magnesium chloride (Mg(II)  $1s^2 2s^2 2p^6$ ) despite the Zn<sup>2+</sup> ion (74 pm) and Mg<sup>2+</sup> ion (72 pm) having similar sizes and charges.

From an MO perspective, the orbital overlap disperses the charge on each ion and so weakens the electrovalent forces throughout the solid, this can be used to explain the trend seen for the melting points of lithium halides.

LiF = 870 °C, LiCl = 613 °C, LiBr = 547 °C, LiI = 446 °C

It is found that the greater the possibility of polarization, the lower is the melting point and heat of sublimation and the greater is the solubility in non-polar solvents.

Example: The melting point of KCl is higher than that of AgCl though the crystal radii of  $Ag^+$  and  $K^+$  ions are almost the same.

Solution: When the melting points of two compounds are compared, the one having the lower melting point is assumed to have the smaller degree of ionic character. In this case, both are chlorides, so the anion remains the same. The deciding factor must be the cation. (If the anions were different, then the answer could be affected by the variation of the anion.) Here the significant difference between the cations is in their electronic configurations.  $K^+=$  [Ar] and  $Ag^+ = [Kr] 4d^{10}$ . This means a comparison needs to be made between a noble gas core and pseudo noble gas core, which as noted above holds that the pseudo noble gas would be the more polarizing.

### Percentage of ionic character and charge distribution

Based on Fajan's rules, it is expected that every ionic compound will have at least some amount of covalent character. The percentage of ionic character in a compound can be estimated from dipole moments.

The bond dipole moment uses the idea of electric dipole moment to measure the polarity of a chemical bond within a molecule. It occurs whenever there is a separation of positive and negative charges. The bond dipole  $\mu$  is given by:

 $\mu = \delta d$ 

A bond dipole is modeled as  $+\delta - \delta$ - with a distance d between the partial charges. It is a vector, parallel to the bond axis and by convention points from minus to plus . The SI unit for an electric dipole moment is the coulomb-meter, (C m). This is thought to produce values too large to be practical on the molecular scale so bond dipole moments are commonly measured in Debye, represented by the symbol, D.

Historically the *Debye* was defined in terms of the dipole moment resulting from two equal charges of opposite sign and separated by 1 Ångstrom ( $10^{-10}$  m) as 4.801 D. This value arises from ( $1.602 \times 10^{-19} * 1 \times 10^{-10}$ ) /  $3.336 \times 10^{-30}$ 

where D =  $3.336 \times 10^{-30} \text{ C m}$  (or 1 C m =  $2.9979 \times 10^{29} \text{ D}$ ).

Typical dipole moments for simple diatomic molecules are in the range of 0 to 11 D.

# The % ionic character = $\mu_{observed}$ / $\mu_{calculated}$ (assuming 100% ionic bond) \* 100 % Bond character based on electronegativity differences

It is possible to predict whether a given bond will be non-polar, polar covalent, or ionic based on the electronegativity difference, since the greater the difference, the more polar the bond.



Electronegativity difference, ΔχΡ	Bond
$\Delta\chi < 0.4$	covalent
$0.4 < \Delta \chi < 1.7$	polar
	covalent
$\Delta \chi > 1.7$	ionic

Linus Pauling proposed an empirical relationship which relates the percent ionic character in a bond to the electronegativity difference.

# percent ionic character= $(1-e^{-(\Delta \chi/2)^{2}})^{*}$ 100

The magnitude of polarization or increased covalent character depends upon a number of factors. These factors are,

# (1) Small size of cation:

Smaller size of cation greater is its polarizing power i.e. greater will be the covalent nature of the bond.

# (2) Large size of anion:

Larger the size of anion greater is its polarizing power i.e. greater will be the covalent nature of the bond.

# (3) Large charge on either of the two ions:

As the charge on the ion increases, the electrostatic attraction of the cation for the outer electrons of the anion also increases with the result its ability for forming the covalent bond increases.

### (4) Electronic configuration of the cation:

For the two ions of the same size and charge, one with a pseudo noble gas configuration (i.e. 18 electrons in the outermost shell) will be more polarizing than a cation with noble gas configuration (i.e., 8 electron in outer most shell).

Anomalous behaviour of the 2nd row elements: Li, Be, B, C, N, O, F

For the elements in the 2nd row, as the atomic number **increases**, the atomic radius of the elements decreases, the electronegativity increases, and the ionization energy increases. The 2nd row has two metals (lithium and beryllium), making it the least metallic period and it has the most nonmetals, with four. The elements in the 2nd row often have the most extreme properties in their respective groups; for example, fluorine is the most reactive halogen, neon is most lithium the inert noble gas, and is the least reactive alkali metal. These differences in properties with the subsequent rows are a result of:

- the smaller size of the atoms
- an outer shell with a maximum of 8 electrons (2s and 2p) and an underlying shell with just 2 electrons
- no acessible d-orbitals energy too high for use in bonding

Apart from the 2nd row (ignoring H/He 1st row) the later rows all end with inert gases but these do not have completed quantum levels. The 2nd row elements in general can only use the 2s and 2p electrons for bonding restricting the total number of bonds to 4.

So N is not expected to have more than 4 bonds and 3 is common, while for P 5 and 6 bonded species are quite common.

# **Reactivity of metals and metalloids**

For Lithium, compared to other alkali metals

# **Reaction with water:**

Li reacts slowly with water at 25 °C

Na reacts violently and K in flames

 $2M(s) + 2H_2O(l) \rightarrow 2M^+(aq) + 2OH^- + H_2(g)$ 

In general Li, Be, B, C, N, O, F are less reactive towards water than their heavier congeners.

# **Reaction with oxygen:**

In conditions of excess oxygen, only Li forms a simple oxide, Li<sub>2</sub>O. Other metals form peroxides and superoxides

# **Reaction with nitrogen:**

Li reacts directly with  $N_2$  to form  $Li_3N$ 

 $6\text{Li}(s) + N_2(g) \rightarrow 2\text{Li}_3N(s).$ 

No other alkali metal reacts with N2

# Solubility:

LiF, LiOH and Li<sub>2</sub>CO<sub>3</sub> are less soluble than the corresponding Na and K compounds

For **Beryllium** compared to the other alkaline earth metals:

With								water:
All	Group	2	metals	except	Be, r	eact	with	water
M(s)	+	2H <sub>2</sub> O(l)	$\rightarrow$	$M^{2+}(aq)+$	20H	(aq)	+	$H_{2}\left(g\right)$
With				oxygen				(air):
Be only	reacts with	h air abo	ve 600 °C	if it is finely	powdered.	The Be	O that	is formed
is ampho	oteric (other		Group	2	oxides	ar	e	basic).
Of the Group 2 elements only Be reacts with NaOH or KOH to liberate $H_2$ and form $[Be(OH)_4]^2$ .								

### Chemical Bonding - I (2017 – 18) batch

Li and Be are metals but are less conducting than the higher members of Group 1 and 2 elements IEs due to their high (electrons close nucleus). are to Ionization of **Boron** to  $B^{3+}$  requires a large input of energy and B adopts a covalent polymeric structure with semi-metallic properties. The other elements of Group 14 become increasingly metallic as the group is descended due to ionization the decrease in energies. Crystalline Boron is chemically inert - unaffected by boiling HCl and only slowly oxidized by concentrated HNO<sub>3</sub> when finely hot powdered. Covalent character

Li<sup>+</sup> and Be<sup>2+</sup> are small and have strong polarizing abilities. Their compounds are more covalent heavier than those of the elements in their groups.  $BeCl_2$  is covalent while  $MCl_2$  (M = Mg-Ba) are ionic. The conductivity of fused beryllium chloride is only 1/1000 that of sodium chloride under similar conditions. Catenation

Catenation is the linkage of atoms of the same element into longer chains. Catenation occurs most readily in carbon, which forms covalent bonds with other carbon atoms to form longer chains and structures. This is the reason for the presence of the vast number of organic compounds in nature.

The ability of an element to catenate is primarily based on the bond energy of the element to itself, which decreases with more diffuse orbitals (those with higher azimuthal quantum number) overlapping to form the bond. Hence, carbon, with the least diffuse valence shell 2p orbital is capable of forming longer p-p sigma bonded chains of atoms than heavier elements which bond via higher valence shell orbitals.

**Department of Chemistry, KAHE** 

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Hetero-catenation is quite common in Inorganic Chemistry. Phosphates and silicates with P-O-P-

O and Si-O-Si-O linkages are examples of this. Multiple Bonds

C, N and O are able to form multiple bonds (double and/or triple). In Group 14, C=C double bonds are stable (134 pm) but Si=Si double bonds (227 pm) are uncommon. The diagram below shows how multiple bonds are formed involving  $\pi$  overlap of 2p orbitals. By comparison the 3p orbitals of the corresponding third row elements Si, P, and S are more diffuse and the longer bond distances expected for these larger atoms would result in poor  $\pi$  overlap. C=C bond length 134 pm and Si=Si bond length 227 = = pm Oxidizing ability oxygen fluorine of and Due to the high electron affinities and electronegativities of oxygen and fluorine, they tend to form strong ionic bonds with other elements. They even react with noble gases to form compounds XeO<sub>3</sub>, XeO<sub>4</sub>. XeF<sub>4</sub> and XeF<sub>6</sub>. such as

In 1962 Neil Bartlett at the University of British Columbia reacted platinum hexafluoride and xenon, in an experiment that demonstrated the chemical reactivity of the noble gases. He discovered the mustard yellow compound, xenon hexafluoroplatinate, which is perhaps now best formulated as a mixture of species,  $[XeF^+][PtF_5]^-$ ,  $[XeF^+][Pt_2F_{11}]^-$ , and  $[Xe_2F_3]^+[PtF_6]^-$ .

A few hundred compounds of other noble gases have subsequently been discovered: in 1962 for radon, radon difluoride (RnF<sub>2</sub>), and in 1963 for krypton, krypton difluoride (KrF<sub>2</sub>). The first stable compound of argon was reported in 2000 when argon fluorohydride (HArF) was formed at a temperature of 40 K (-233.2 °C). Neutral compounds in which helium and neon are involved in

# Chemical Bonding - I (2017 – 18) batch

chemical	bonds	have	still	not	been	formed.
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Noble gas compounds have already made an impact on our daily lives.  $XeF_2$  is a strong fluorinating agent and has been used to convert uracil to 5-fluorouracil, one of the first anti-tumor agents.

# **Possible Questions**

# PART A (Objective type each questions carries one mark)

1. Which force is strongest

a)dipole-dipole forces b)**ion-ion forces** c)ion -dipole forces d) ion-induced dipole forces

2. Greater the bond order, greater is

a)**bond dissociation energy** b) covalent nature c)bond length d)paramagnetism

3. The molecule having largest dipole moment among the following is

a)CH<sub>4</sub> b)CHCl<sub>3</sub> c)CCl<sub>4</sub> d)CHI<sub>3</sub>

- 4. The molecule which have zero dipole moment
  a)CH<sub>2</sub>Cl<sub>2</sub> b)BF<sub>3</sub> c)ClO<sub>2</sub> d)NF<sub>3</sub>
- 5. Which of the following is most polar

a)C-O b)C-F c)O-F d)N-F

- 6. According to Fajan'srule, covalent bond is favoured by
  a)large cation and small anion b)large cation and large anion c)small cation and small anion d)small cation and large anion
- 7. Which of the following compound is non polar

 $a) CHCl_3 \quad b) \textbf{SiCl}_4c) SnCl_2 \quad d) NH_3$ 

8. Which of the following is least ionic

a)AgCl b) KCl c)BaCl<sub>2</sub> d)COCl<sub>2</sub>

9. Which of the following compound has least dipole moment

a)NH<sub>3</sub>b)NF<sub>3</sub> c)SO<sub>2</sub> d)H<sub>2</sub>O

10. Which one of the following has least polarity

a)H-F b)H-Cl c) H-O d)H-S

- 11. Which of the following is most polara)C-O b)C-F c)O-F d)N-F
- 12. As the s-character of hybridised orbital increases, the bond anglea)increases b)decreases c)does not change d)equal
- 13. The weakest among the following isa)ionic bond b)covalent bond c)metallic bond d)vanderwaal's forces
- 14. The electron has ----- probability of being located at a node.a)0 b)1 c)2 d)3
- 15. Ionic compounds n general possess both
  - a) high m.pt and non-directional bondsb) high m.pt and low-boiling pointsc) directional bonds and low boiling pointsd) high solubilities in polar and non-polarsolvents
- 16. The magnitude of the lattice energy of a solid increases ifa)the ions are large b)**the ions are small** c)the ions are of equal size d)charges on the ions are small
- 17. Which of the following bonds has the highest bond dissociation energya)CH<sub>3</sub>-H b)CH<sub>3</sub>-Br c)CH<sub>3</sub>-Cl d)CH<sub>3</sub>-F
- 18. Whice of the following forces between atoms or ions or molecules is the strongest
  - a) ionic-bond b)ion dipole c)dipole-dipole d)london-dispersive

### PART B (each questions carries two marks)

- 19. State Fajan's rules.
- 20. What is an ionic bond.
- 21. Write the Born-Lande equation.
- 22. What is dipole moment.
- 23. What are the applications dipole moment.

# PART C (each questions carries six marks)

- 24. What is lattice energy? How does the solubility of an ionic solid depend upon its lattice energy.
- 25. Write a note on polarization and polarizing power
- 26. Explain why melting point of NaCl is higher than that of AlCl<sub>3</sub>.
- 27. The lattice energy of silver halides are almost the same as that of alkali metal halides, yet they are insoluble in water. Explain.
- 28. Explain how Born-Haber cycle is used to calculate the lattice energy of NaCl.
- 29. Give an account on solvation energy and importance in context of stability and solubility of ionic compounds.
- 30. What is an ionic bond? List four characteristic properties of ionic compounds.
- 31. Discuss the Fajan rules.
- 32. Derive Born –Lande equation for the lattice energy of sodium chloride.
- 33. What do you understand by the terms polarizing power and polarizability. How do these determine ionic character of a compound.

#### KARPAGAM ACADEMY OF HIGHER EDUCATION

Questions	Opt-1	Opt-2	Opt-3	opt-4	Answer
Unit-III (Chemical Bonding – I)					
Which of the following has zero dipole moment	PH <sub>3</sub>	CH <sub>4</sub>	$PH_4$	$CH_2Cl_2$	CH <sub>4</sub>
Which of the following is least ionic	AgCl	ксі	BaCl <sub>2</sub>	COCl <sub>2</sub>	COCl <sub>2</sub>
Which of the following compound has least dipole moment	NH <sub>3</sub>	NF <sub>3</sub>	SO <sub>2</sub>	H <sub>2</sub> O	NF <sub>3</sub>
Which one of the following has least polarity	H-F	H-Cl	H-O	H-S	H-S
Which of the following is most polar	C-0	C-F	O-F	N-F	C-F
According to Esian's rule covalent hand is favoured by	large cation and small	large cation and large	small cation and small	small cation and large	small cation and large
	anion	anion	anion	anion	anion
Which contains a co-ordinate and covalent bond	BaCl <sub>2</sub>	NH <sub>4</sub> Cl	HCl	H <sub>2</sub> O	NH <sub>4</sub> Cl
Which has a net dipole moment	BF <sub>3</sub>	CCl <sub>4</sub>	BeF <sub>2</sub>	SO <sub>2</sub>	$SO_2$
Which of the following compound is non polar	CHCl <sub>3</sub>	SiCl <sub>4</sub>	SnCl <sub>2</sub>	NH <sub>3</sub>	SiCl <sub>4</sub>
The angle between the overlapping of one s-orbital and one p- orbital is	180°	120°	109°28,	120060'	180°
Which force is strongest	dipole-dipole forces	ion-ion forces	ion -dipole forces	ion-induced dipole forces	ion-ion forces
Greater the bond order,greater is	bond dissociation energy	covalent nature	bond length	paramagnetism	bond dissociation energy
The molecule having largest dipole moment among the following is	$CH_4$	CHCl <sub>3</sub>	CCl <sub>4</sub>	CHI <sub>3</sub>	CHCl <sub>3</sub>
The molecule which have zero dipole moment	CH <sub>2</sub> Cl <sub>2</sub>	BF <sub>3</sub>	ClO <sub>2</sub>	NF <sub>3</sub>	BF <sub>3</sub>
Which has the lowest bond angle	NH <sub>3</sub>	BeF <sub>2</sub>	H <sub>3O+</sub>	CH <sub>4</sub>	NH <sub>3</sub>
The shape of a molecule which has three bonds and one lone pair is	Octahedral	Pyramidal	Triangular planar	Tetrahedral	Pyramidal
The type of hybridisation in SO2 and SO3 is respectively	sp, sp <sup>2</sup>	$sp^2$ , $sp^2$	$sp^2$ , $sp^3$	sp, sp <sup>3</sup>	$sp^2$ , $sp^2$
As the s-character of hybridised orbital increases, the bond angle	increases	decreases	does not change	equal	increases
The bond order is maximum in	H <sub>2</sub>	H <sub>2+</sub>	He <sub>2</sub>	He2+	H <sub>2</sub>
Strongest bond is in	NaCl	CsCl	Both a and b	LiF	CsCl
The weakest among the following is	ionic bond	covalent bond	metallic bond	vanderwaal's forces	vanderwaal's forces
Which has zero dipole moment	ClF	PCl <sub>3</sub>	SiF <sub>4</sub>	CFCl <sub>3</sub>	SiF <sub>4</sub>
The molecules having dipole moment are	2,2 dimethyl propane	trans-3-hexene	trans-2-pentene	2,2,3,3 tetramethyl butane	trans-2-pentene
Stability of ionic compound is due to	electronegativity	lattice energy	sublimation energy	electron affinity	lattice energy
Ionic compoundsin general possess both	high m.pt and non- directional bonds	high m.pt and low- boiling points	directional bonds and low boiling points	high solubilities in polar and non-polar solvents	high m.pt and non- directional bonds
The magnitude of the lattice energy of a solid increases if	the ions are large	the ions are small	the ions are of equal size	charges on the ions are small	the ions are small

Which of the following bonds has the highest bond dissociation energy	СН <sub>3</sub> -Н	CH <sub>3</sub> -Br	CH <sub>3</sub> -Cl	CH <sub>3</sub> -F	CH <sub>3</sub> -F
Whice of the following forces between atoms or ions or molecules is the strongest	ionc-bond	ion dipole	dipole-dipole	london-dispersive	ionc-bond
The unequal sharing of the bonded pair of electrons between two atoms in a molecule causes	dipole	radical formation	covalent bond	decomposition of molecule	dipole
All chemical bonds are the result of	interaction of nuclei	interaction of electrons	difference in electronegativity	interaction of the electrons and nuclei	interaction of the electrons and nuclei
CCl4 is insoluble in water because	H <sub>2</sub> O is polar	CCl4 is non-polar	they do not form intermolecular H- bonding	they do not form intramolecular H- bonding	CCl4 is non-polar
Which has the minimum bond energy	H-Br	H-I	-	н-н	1-1
The compound with maximum dipole moment is	p-dichlorobenzene	m-dichlorobenzene	o-dichlorobenzene	CCl <sub>4</sub>	o-dichlorobenzene
Dipole-dipole attractive forces are strongest between the molecule of	Не	CH <sub>4</sub>	CO <sub>2</sub>	H <sub>2</sub> O	H <sub>2</sub> O
Whice of the following molecules does not possess a permanent electric dipole	H <sub>2</sub> O	SO <sub>2</sub>	SO <sub>3</sub>	CS <sub>2</sub>	CS <sub>2</sub>
The C-H bond distance in the largest in	$C_2H_2$	$C_2H_4$	$C_2H_4Br_2$	$C_2H_6$	$C_2H_4Br_2$
Which is most ionic	P <sub>2</sub> O <sub>5</sub>	MnO	CrO <sub>3</sub>	Mn <sub>2</sub> O <sub>7</sub>	MnO
The compound which contains both ionic and covalent bond	CH <sub>4</sub>	H <sub>2</sub>	KCN	KCl	KCN
Which of the following is polar	BF <sub>3</sub>	C <sub>2</sub> H <sub>5</sub> -F	CO <sub>2</sub>	CH <sub>4</sub>	C <sub>2</sub> H <sub>5</sub> -F
Which of the following compound has dipole movement zero	CCl <sub>4</sub>	CHCl <sub>3</sub>	HF	NH <sub>3</sub>	CCl <sub>4</sub>
Which of the following does not conduct electricity	Molten NaOH	Molten KOH	Solid NaCl	Aqueous NaCl	Solid NaCl
Which bond is most polar	Cl-F	Br-F	I-F	F-F	I-F
When sodium and chlorine react, energy is	Released and ionic bond is formed	Released and covalent bond is formed	Absorbed and covalent bond is formed	Absorbed and ionic bond is formed	Released and ionic bond is formed
According to Fajan's rule, ionic character increases when	large cation and small anion	small cation and small anion	large cation and large anion	large cation and small anion	large cation and small anion
AICI3 is covalent, but AIF3 is ionic. This is justified by	Crystal structure	Valence bond theory	Fajan rule	Lattice energy	Fajan rule
In the case of stable molecule the value of bond order be	negative	positive	zero	all	positive
Which bond has the highest bond energy	sigma bond	covalent bond	multiple bond	polar covalent bond	multiple bond
Both ionic and covalent bonds are present in	CH <sub>4</sub>	KCl	$SO_2$	NaOH	NaOH
Which is least soluble in water	AgCl	AgF	Agl	AgBr	Agl
Greater the dipole moment	Greater is the ionic in nature	lesser is the polarity	smaller the ionic nature	covalent nature	Greater is the ionic in nature
Ionic compounds have melting and boiling points.	high	low	very low	neither high nor low	high
Which of the following has maximum polar character	C-I	C-Br	C-F	C-S	C-F
A substance is soluble in water if	lattice energy> hydration energy	Hydration energy> lattice energy	hydration energy=lattice energy	no relation	Hydration energy> lattice energy

An electrovalent compound is made up of	electrically charged particles	neutral molecules	neutral atoms	only neutral	electrically charged particles
NaCl crystal is made up of	NaCl molecules	Na+ and Cl - ions	Na and Cl atoms	dimers of NaCl	Na+ and Cl - ions
The stability of an ionic solid is measured in terms of its	lattice energy	hydration energy	Haber	Fajan	lattice energy
A salt dissolves in water if hydration energy is than lattice energy	more	low	very low	equal	more
In elecrovalency of electrons takes place.	transfer	sharing	electrons shared by one atom only	no electrons takes place	transfer
For a given cation ,as the size of the anion increases, the covalent characterand m.pt decreases.	decreases	increases	decreases with increasing	increasing with decreases	decreases
The rule to determine the extent of polarization of an anion by a cation was given by	Pauling	Born	Haber	Fajan	Fajan

# Unit-IV

# **Chemical Bonding – II**

*Covalent bonding:* VB Approach: Shapes of some inorganic molecules and ions on the basis of VSEPR and hybridization with suitable examples of linear, trigonal planar, square planar, tetrahedral, trigonal bipyramidal and octahedral arrangements. Concept of resonance and resonating structures in various inorganic and organic compounds.

**Molecular geometry** is the three-dimensional arrangement of the atoms that constitute a molecule. It determines several properties of a substance including its reactivity, polarity, phase of matter, color, magnetism, and biological activity. The angles between bonds that an atom forms depend only weakly on the rest of molecule, i.e. they can be understood as approximately local and hence transferable properties.

# VALENCE SHELL ELECTRON PAIR REPULSION (VSEPR) THEORY

In order to predict the geometry of molecules, Nyholm and Gillespie developed a qualitative model known as **Valence Shell Electron Pair Repulsion Theory (VSEPR Theory)**. The basic assumptions of this theory are summarized below.

1) The electron pairs in the valence shell around the central atom of a molecule repel each other and tend to orient in space so as to minimize the repulsions and maximize the distance between them.

2) There are two types of valence shell electron pairs viz., i) Bond pairs and ii) Lone pairs

**Bond pairs** are shared by two atoms and are attracted by two nuclei. Hence they occupy less space and cause less repulsion.

**Lone pairs** are not involved in bond formation and are in attraction with only one nucleus. Hence they occupy more space. As a result, the lone pairs cause more repulsion.

The order of repulsion between different types of electron pairs is as follows:

Lone pair - Lone pair > Lone Pair - Bond pair > Bond pair - Bond pair

Note: The bond pairs are usually represented by a solid line, whereas the lone pairs are represented by a lobe with two electrons.

3) In VSEPR theory, the multiple bonds are treated as if they were single bonds. The electron pairs in multiple bonds are treated collectively as a single super pair.

The repulsion caused by bonds increases with increase in the number of bonded pairs between two atoms i.e., a triple bond causes more repulsion than a double bond which in turn causes more repulsion than a single bond.

4) The shape of a molecule can be predicted from the number and type of valence shell electron pairs around the central atom.

When the valence shell of central atom contains only bond pairs, the molecule assumes symmetrical geometry due to even repulsions between them.

However the symmetry is distorted when there are also lone pairs along with bond pairs due to uneven repulsion forces.

5) Primary & Secondary effects on bond angle and shape:

i) The **bond angle** decreases due to the presence of lone pairs, which cause more repulsion on the bond pairs and as a result the bond pairs tend to come closer.

ii) The repulsion between electron pairs increases with increase in electronegativity of central atom and hence the bond angle increases. The bond pairs are closer and thus by shortening the distance between them, which in turn increases the repulsion. Hence the bonds tend to move away from each other. The bond pairs tend to move away from each other since the distance between them is shortened as they are more localized on more electronegative central atom.

However the bond angle decreases when the electronegativities of ligand atoms are more than that of central atom. There is increase in the distance between bond pairs since they are now closer to ligand atoms. Due to this, they tend to move closer resulting in the decrease in bond angle.

The bond pairs tend to come closer since the distance between them is increased as they are more localized on more electronegative ligand atoms.

iii) The bond angle decreases with increase in the size of central atom.

On smaller central On bigger central atoms, atoms the bond pairs the bond pairs are more are closer and hence distant from each other tend to move away and hence there is less from each other so as repulsion. Hence they to minimize repulsion. tend to move closer and Hence bond angle will be more. bond angle. However the bond angle increases with increase in the size of ligand atoms, which surround the central atom.

There is less repulsion	
	These is more repulsion
between smaller	
	between bigger ligand
ligand atoms and they	
	atoms and hence they
can move closer to	. 1 .
anah other and thus	tend to move away
each other and thus	from each other. Thus
decrease the bond	nom cach other. Thus
deerease the bolid	bond angle increases
angle.	oona angie mereuses.

iv) The bond angles are also changed when multiple bonds are present. It is due to uneven repulsions.

6) When there are two or more resonance structures, the VSEPR theory is applicable to any of such contributing structure.

Types of molecular structure

A bond angle is the geometric angle between two adjacent bonds. Some common shapes of simple molecules include:

- Linear: In a linear model, atoms are connected in a straight line. The bond angles are set at 180°. For example, carbon dioxide and nitric oxide have a linear molecular shape.
- **Trigonal planar:** Molecules with the trigonal planar shape are somewhat triangular and in one plane (flat). Consequently, the bond angles are set at 120°. For example,boron trifluoride.

- Bent: Bent or angular molecules have a non-linear shape. For example, water (H<sub>2</sub>O), which has an angle of about 105°. A water molecule has two pairs of bonded electrons and two unshared lone pairs.
- Tetrahedral: Tetra- signifies four, and -hedral relates to a face of a solid, so "tetrahedral" literally means "having four faces". This shape is found when there are four bonds all on one central atom, with no extra unshared electron pairs. In accordance with the VSEPR (valence-shell electron pair repulsion theory), the bond angles between the electron bonds are arccos(-1/3) = 109.47°. For example, methane (CH<sub>4</sub>) is a tetrahedral molecule.
- Octahedral: Octa- signifies eight, and -hedral relates to a face of a solid, so "octahedral" means "having eight faces". The bond angle is 90 degrees. For example, sulfur hexafluoride (SF<sub>6</sub>) is an octahedral molecule.
- **Trigonal pyramidal:** A trigonal pyramidal molecule has a pyramid-like shape with a triangular base. Unlike the linear and trigonal planar shapes but similar to the tetrahedral orientation, pyramidal shapes require three dimensions in order to fully separate the electrons. Here, there are only three pairs of bonded electrons, leaving one unshared lone pair. Lone pair bond pair repulsions change the bond angle from the tetrahedral angle to a slightly lower value. For example, ammonia (NH<sub>3</sub>).

### VSEPR table

The bond angles in the table below are ideal angles from the simple VSEPR theory, followed by the actual angle for the example given in the following column where this differs. For many cases, such as trigonal pyramidal and bent, the actual angle for the example differs from the ideal angle, but all examples differ by different amounts. For example, the angle in  $H_2S$  (92°) differs from the tetrahedral angle by much more than the angle for  $H_2O$  (104.48°) does.

Bonding electron pairs	Lone pairs	Electron domains (Steric #)	Shape	Ideal bond angle (example's bond angle)	Example	Image
2	0	2	linear	180°	CO <sub>2</sub>	
3	0	3	trigonal planar	120°	BF <sub>3</sub>	<b>"</b>
2	1	3	bent	120° (119°)	SO <sub>2</sub>	<b>~</b>
4	0	4	tetrahedral	109.5°	CH <sub>4</sub>	<b>"</b>
3	1	4	trigonal pyramidal	109.5 (107.8°)	NH <sub>3</sub>	
2	2	4	Bent	109.5° (104.48°) <sup>[10][11]</sup>	H <sub>2</sub> O	<b>~</b> >
5	0	5	trigonal bipyramidal	90°, 120°, 180°	PCl <sub>5</sub>	⊶ <b>}</b> c
4	1	5	seesaw	ax-ax 180° (173.1°), eq-eq 120° (101.6°),	SF <sub>4</sub>	್ರ್ಯಾಂ

Bonding electron pairs	Lone pairs	Electron domains (Steric #)	Shape	Ideal bond angle (example's bond angle)	Example	Image
				ax-eq 90°		
3	2	5	T-shaped	90° (87.5°), 180° < 180°)	ClF <sub>3</sub>	***
2	3	5	linear	180°	XeF <sub>2</sub>	
6	0	6	octahedral	90°, 180°	SF <sub>6</sub>	sign of the second s
5	1	6	square pyramidal	90° (84.8°)	BrF5	ي <mark>ل</mark> و
4	2	6	square planar	90°, 180°	XeF <sub>4</sub>	÷
7	0	7	pentagonal bipyramidal	90°, 72°, 180°	IF7	3 george
6	1	7	pentagonal pyramidal	72°, 90°, 144°	XeOF5 <sup>-</sup>	ages
5	2	7	planar pentagonal	72°, 144°	XeF5 <sup>-</sup>	<b>→</b> *3
8	0	8	square antiprismatic		XeF <sub>8</sub> <sup>2–</sup>	×

Bonding electron pairs	Lone pairs	Electron domains (Steric #)	Shape	Ideal bond angle (example's bond angle)	Example	Image
9	0	9	tricapped trigonal prismatic		ReH9 <sup>2-</sup>	≽

# RELATION BETWEEN NUMBER & TYPE OF VALENCE ELECTRON PAIRS WITH THE SHAPE OF MOLECULE

\* The shape of molecule and also the approximate bond angles can be predicted from the number and type of electron pairs in the valence shell of central atom as tabulated below.

In the following table the molecule is represented by "AXE" notation, where

A = Central atom

X = Ligand atom bonded to the central atom either by a single bond or by multiple bond; indicating a bond pair. E = Lone pair

\* The sum of number of ligand atoms (X) and number of lone pairs (E) is also known as steric number.

\* The bond pairs are shown as green colored thick lines, whereas the lone pairs are shown as point charges using green colored lobes.

Steric number	Number of Bond pairs	Number of Lone pairs	Formula	Shape of molecule	Approximate Bond angles	Examples
1	1	0	AX	Linear	180°	ClF, BrF, BrCl, HF, O <sub>2</sub>
2	2	0	$AX_2$	Linear	180°	BeCl <sub>2</sub> , HgCl <sub>2</sub> , CO <sub>2</sub>
3	3	0	AX3	Trigonal planar	120°	BF <sub>3</sub> , CO <sub>3</sub> <sup>2-</sup> , NO <sub>3</sub> <sup>-</sup> , SO <sub>3</sub>
	2	1	AX2E	Angular	120°	SO <sub>2</sub> , SnCl <sub>2</sub> , O <sub>3</sub> , NSF, $NO_2^-$
4	4	0	$AX_4$	Tetrahedral	109°28'	CH4, SiCl4, NH4 <sup>+</sup> , PO4 <sup>3-</sup> , SO4 <sup>2-</sup> , ClO4 <sup>-</sup>
	3	1	AX3E	Trigonal pyramidal	around 109°28'	NH <sub>3</sub> , PCl <sub>3</sub> , XeO <sub>3</sub>
	2	2	AX <sub>2</sub> E <sub>2</sub>	Angular	around 109°28'	$H_2O$ , $SCl_2$ ,

						Cl <sub>2</sub> O,
						OF <sub>2</sub>
5	5	0	AX5	Trigonal bipyramidal	120° & 90°	PCl5, SOF4
	4	1	AX₄E	See saw or distorted tetrahedral	-	SF4, TeCl4
	3	2	AX <sub>3</sub> E <sub>2</sub>	T-Shape	90°	ClF <sub>3</sub> , BrF <sub>3</sub> , BrCl <sub>3</sub>
	2	3	AX <sub>2</sub> E <sub>3</sub>	Linear	180°	$XeF_2$ , $I_3^-$
6	6	0	$AX_6$	Octahedral	90°	SF <sub>6</sub>
	5	1	AX5E	Square pyramidal	90°	ClF5, BrF5, ICl5
	4	2	AX <sub>4</sub> E <sub>2</sub>	Square planar	90°	XeF <sub>4</sub>
7	7	0	AX <sub>7</sub>	Pentagonal bipyramidal	72° & 90°	IF7
	6	1	AX6E	Pentagonal pyramidal	around 72° & 90°	XeOF <sub>5</sub> <sup>-</sup> , IOF <sub>5</sub> <sup>2-</sup>

# STEPS INVOLVED IN PREDICTING THE SHAPES OF MOLECULES USING VSEPR THEORY

\* The first step in determination of shape of a molecule is to write the Lewis dot structure of the molecule.

\* Then find out the number of bond pairs and lone pairs in the valence shell of central atom.

While counting the number of bond pairs, treat multiple bonds as if they were single bonds. Thus electron pairs in multiple bonds are to be treated collectively as a single super pair.

\* Use the above table to predict the shape of molecule based on steric number and the number of bond pairs and lone pairs.

# **APPLICATIONS & ILLUSTRATIONS OF VSEPR THEORY**

# 1) Methane (CH<sub>4</sub>):

\* The Lewis structure of methane molecule is:

\* There are 4 bond pairs around the central carbon atom in its valence shell. Hence it has tetrahedral shape with 109°28' of bond angles.

# 2) Ammonia (NH<sub>3</sub>):

\* The Lewis structure of ammonia indicates there are three bond pairs and one lone pair around the central nitrogen atom.

\* Since the steric number is 4, its structure is based on tetrahedral geometry. However, its shape is pyramidal with a lone pair on nitrogen atom.

\* The bond angle is decreased from 109°28' to 107°48' due to repulsion caused by lone pair on the bond pairs.

### 3) Water (H<sub>2</sub>O):

\* It is evident from the Lewis structure of water molecule, there are two bond pairs and two lone pairs in the valence shell of oxygen. Hence its structure is based on tetrahedral geometry. However its shape is angular with two lone pairs on oxygen.

\* The bond angle is decreased to 104°28' due to repulsions caused by lone pairs on bond pairs. It can be noted that the bond angle decreases with increase in the number of lone pairs on the central atom.

### 4) Sulfur tetrachloride (SCl<sub>4</sub>):

Since there are four bond pairs and one lone pair around sulfur in its valence shell, the structure of SCl<sub>4</sub> is based on trigonal bipyramidal geometry. It has seesaw shape with a lone pair occupying the equatorial position.

The angles between P-Cl<sub>axial</sub> and P-Cl<sub>equatorial</sub> are less than 90° due to repulsion exerted by the lone pair. The angle between P-Cl<sub>equatorial</sub> bonds also decreases from its usual value, 120°.

The lone pair occupies the equatorial position to minimize the repulsions.

Usually the lone pairs, bulky groups and less electronegative atoms tend to occupy equatorial position to minimize repulsions. This is because they experience repulsion only from two groups at 90°, when they occupy the equatorial positions. However the repulsion will be more when they occupy axial positions, since they encounter three groups at 90°.

### 5) **PF<sub>3</sub>Cl<sub>2</sub>:**

There are only 5 bond pairs on phosphorus atom. Hence it has trigonal bipyramidal shape. The chlorine atoms occupy the equatorial positions to minimize the repulsions since they are not only bulkier and also less electronegative than fluorine atoms.

The bond pair of P-Cl is slightly more closer towards the P atom when compared to the bond pair of P-F, since the chlorine atoms are comparatively less electronegative than fluorine atoms. Hence there is comparatively more negative charge accumulation towards P atom, which makes the P-Cl bonds to experience more repulsion than P-F bonds. Hence they orient in equatorial positions at 120° to minimize repulsions.

Note that, here we are comparing the polarity of P-Cl bond with P-F bond. But one should keep in mind that the bond pair of P-Cl bond is still closer to Cl, since it is more electronegative than P atom.

#### 6) Formaldehyde (HCHO):

There are three bond pairs around the central carbon atom. The double bond between C and O is considered as a single super pair. Hence the shape of the molecule is trigonal planar and the bond angles are expected to be equal to 120°.

However, the C=O exerts more repulsion on the C-H bond pairs. Hence the  $\angle$ H-C-H bond angle will be less than 120° and the  $\angle$ H-C-O is greater than 120°.

### Resonance

Resonance is a mental exercise and method within the Valence Bond Theory of bonding that describes the delocalization of electrons within molecules. It compares and contrasts two or more possible Lewis structures that can represent a particular molecule. Resonance structures are

used when one Lewis structure for a single molecule cannot fully describe the bonding that takes place between neighboring atoms relative to the empirical data for the actual bond lengths between those atoms. The net sum of valid resonance structures is defined as a resonance hybrid, which represents the overall delocalization of electrons within the molecule. A molecule that has several resonance structures is more stable than one with fewer. Some resonance structures are more favorable than others.

Electrons have no fixed position in atoms, compounds and molecules but have probabilities of being found in certain spaces (orbitals). Resonance forms illustrate areas of higher probabilities (electron densities). This is like holding your hat in either your right hand or your left. The term Resonance is applied when there are two or more possibilities available. Resonance structures do not change the relative positions of the atoms like your arms in the metaphor. The skeleton of the Lewis Structure remains the same, only the electron locations change.

Most arrows in chemistry cannot be used interchangeably and care must be given to selecting the correct arrow for the job.

- $\longleftrightarrow$ : A double headed arrow on both ends of the arrow between Lewis structures is used to show their inter-connectivity
- $\overrightarrow{}$ : Double harpoons are used to designate equilibria
- \_\_\_\_: A single harpoon on one end indicate the movement of **one** electron
- $\longrightarrow$ : A double headed arrow on one end is used to indicate the movement of **two** electrons

# **Example 1: Ozone**

Consider ozone (O<sub>3</sub>)



An animation of how one can do a resonance with ozone by moving electrons

# **Delocalization and Resonance Structures Rules**

In resonance structures, the electrons are able to move to help stabilize the molecule. This movement of the electrons is called delocalization.

- 1. Resonance structures should have the same number of electrons, do not add or subtract any electrons. (check the number of electrons by simply counting them).
- 2. All resonance structures must follow the rules of writing Lewis Structures.
- 3. The hybridization of the structure must stay the same.
- 4. The skeleton of the structure can not be changed (only the electrons move).
- 5. Resonance structures must also have the same amount of lone pairs.

# **Formal Charge**

Even though the structures look the same, the formal charge (FC) may not be. Formal charges are charges that are assigned to a specific atom in a molecule. If computed correctly, the overall formal charge of the molecule should be the same as the oxidation charge of the molecule (the charge when you write out the empirical and molecular formula). We want to choose the resonance structure with the least formal charges that add up to zero or the charge of the overall molecule. The equation for finding Formal Charge is:

Formal Charge = (number of valence electrons in free orbital) - (number of lone-pair electrons) - (12 number bond pair electrons)

The formal charge has to equal the molecule's overall charge,e.g., the CNS– has an overall charge of -1, so the Lewis structure's formal charge has to equal -1.

# **Example 2: Thiocyanate Ion**

Consider the thiocyanate (CNS-) ion.

1. Find the Lewis Structure of the molecule. (Remember the Lewis Structure rules.)

$$\left[:N \equiv C - \vdots:\right]^{-}$$

2. Resonance: All elements want an octet, and we can do that in multiple ways by moving the terminal atom's electrons around (bonds too).



3. Assign Formal Charges

Formal Charge = (number of valence electrons in free orbital) - (number of lone-pair electrons) -

(12 number bond pair electrons)

Remember to determine the number of valence electron each atom has before assigning Formal

# Charges

C = 4 valence  $e^{-}$ , N = 5 valence  $e^{-}$ , S = 6 valence  $e^{-}$ , also add an extra electron for the (-1) charge. The total of valence electrons is 16.


4. Find the most ideal resonance structure. (Note: It is the one with the least formal charges that adds up to zero or to the molecule's overall charge.)



5. Now we have to look at electronegativity for the "Correct" Lewis structure.

The most electronegative atom usually has the negative formal charge, while the least electronegative atom usually has the positive formal charges.



# **Resonance Hybrids**

Resonance Structures are a representation of a Resonance Hybrid, which is the combination of all resonance structures. The resonance structure with the Formal Charge closest to zero is the most accepted structure, however, the correct Lewis structure is actually a combination of all the resonance structures and is not solely describe as one.

- 1. Draw the Lewis Structure & Resonance for the molecule (using solid lines for bonds).
- 2. Where there **can** be a double or triple bond, draw a dotted line (-----) for a bond.

3. Draw only the lone pairs found in all resonance structures, do not include the lone pairs that are not on all of the resonance structures.

The correct Lewis structure is actually a combination of all the resonance structures and hence is not solely described as one.

### **Example 3: Carbonate Ion**

Consider the carbonate ion: CO<sub>3</sub><sup>2-</sup>



Step 1: Draw the Lewis Structure & Resonance.

$$co_{3}^{2-} \\ \begin{bmatrix} \ddot{c} - c = \ddot{c} \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} = c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ I \\ : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - \ddot{c} : \\ \vdots : \dot{c} : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \hline \vdots : \end{bmatrix}^{2-} \\ \leftarrow \begin{bmatrix} \ddot{c} - c - c : \\ \hline$$

Step 2: Combine the resonance structures by adding (dotted) bonds where other resonance bonds can be formed.



Step 3: Add only the lone pairs found on ALL resonance structures.



The bottom is the finished resonance hybrid for  $CO_3^{2-}$ .

# Rules for estimating stability of resonance structures

- 1. The **greater the number of covalent bonds**, the greater the stability since more atoms will have complete octets
- 2. The structure with the least number of formal charges is more stable
- 3. The structure with the least separation of formal charge is more stable

- 4. A structure with a negative charge on the more electronegative atom will be more stable
- 5. Positive charges on the least electronegative atom (most electropositive) is more stable
- Resonance forms that are equivalent have no difference in stability and contribute equally (eg. benzene)

#### **Example 4: Multiple Resonance of other Molecules**

Molecules with multiple resonance forms



Some structural resonance conformations are the major contributor or the dominant forms that the molecule exists. For example, if we look at the above rules for estimating the stability of a molecule, we see that for the third molecule the first and second forms are the major contributors for the overall stability of the molecule. The nitrogen is more electronegative than carbon so, it can handle the negative charge more than carbon. A carbon with a negative charge is the least favorable conformation for the molecule to exist, so the last resonance form contributes very little for the stability of the Ion.



The different resonance forms that the molecule has helps and directs the reactivity to specific sites

The Hybrid Resonance forms show the different Lewis structures with the electron been delocalized. This is very important for the reactivity of chlorobenzene because in the presence of an electrophile it will react and the formation of another bond will be directed and determine by resonance. The long pair of electrons delocalized in the aromatic substituted ring is where it can potentially form a new bond with an electrophile, as it is shown there are three possible places that reactivity can take place, the first to react will take place at the para position with respect to the chloro substituent and then to either ortho position.

# **Possible Questions**

# PART A (Objective type each questions carries one mark)

- Which of the following be octahedral
   a)SF<sub>6</sub> b)BF<sup>4-</sup> c)PCl<sub>5</sub> d)H<sub>2</sub>
- When the hybridisation state of carbon atom changes from sp3 to sp2 to sp ,the angle between the hybridised orbitals
   a)decreases gradually b)increases gradually c)decreases with increasing d)decreases considerably
- 3. .According to VSEPR theory the shape of water molecule is a)octahedral b)**distorted tetrahedral** c)planar triangle d)linea
- 4. In ------ the atoms retain their individuality in the molecule.
  - a) Valence bond theory b)Molecular orbital theory c)Crystal field theory d)coordinate compounds
- Resonance energy is equal to actual bond energy minus energy of the pure ----- a)ionic bond b)Covalent bond c)Coordinate bond d)isomorphous compound
- 6. Bond order in a molecule of NO is ----
  - a)**1** b)2.5 c)3 d) 3.5
- 7. Which one of the following compound is covalent

a)H2b)CaO c)KCl d)Na2S

- 8. The shape of the sulphate ion isa)tetrahedralb)square planar c)trigonalbipyramidal d)hexagonal
- 9. In a double bond connecting two atoms, there is a sharing of a)two electrons b)**four electrons** c)one electrons d)all electrons
- 10. Which of the following compound has not sp3 hybridisation. a)H<sub>2</sub>O b)NH<sub>3</sub> c)CH<sub>4</sub> d)SF<sub>4</sub>
- 11. BF<sub>3</sub> involves ------ hybridisation and the shape of the molecule is triangular planar.
  a)spb)sp<sup>2</sup>c)sp<sup>3</sup> d)d<sup>2</sup>sp<sup>3</sup>
- 12. Directed bond in water form an angle of ------a)90°C b)120°C c)105°C d)60°C

- 13. Which one of the following is strongest bonda)Cl-F b)F-F c)Br-F d)Br-Cl
- 14. Decreasing order of size of various hybrid orbitals is
  a)sp>sp<sup>2</sup>>sp<sup>3</sup> b)sp<sup>3</sup>>sp<sup>2</sup>>sp c)sp<sup>2</sup>>sp>sp<sup>3</sup> d)sp>sp<sup>3</sup>>sp<sup>2</sup>
- 15. Bonded electron pairs present in octahedral SF<sub>6</sub> moleculea)3 b) 4 c)6 d)8

### PART B (each questions carries two marks)

- 16. What do you understand by resonance.
- 17. What shapes are associated with the following hybrid orbitals  $sp^2$ ,  $sp^3$ ,  $sp^3d$ ,  $sp^3d^2$
- On the basis of hybridization, discuss the geometry of the following molecules SF<sub>6</sub>, IF<sub>7</sub>, H<sub>2</sub>O, NH<sub>3</sub>.
- 19. What is meant by hybridization.
- 20. What is sigma and pi bonds.

### PART C (each questions carries six marks)

- 21. What is valence bond theory of bonding? How does valence bond theory account for bonding in a hydrogen molecule.
- 22. Account for the following:
  - i) All the bond lengths in  $SF_6$  are equal but these are not equal in  $IF_5$  molecule.

ii)The geometries of  $H_2O$  and  $H_2S$  are same, but their bond angles are  $H_2O>H_2S$ .

iii)BH<sub>3</sub> is trigonal planar while NH<sub>3</sub> is pyramidal.

- 23. Give examples and explain the concept of hybridisation with reference to : i)sp<sup>3</sup> ii)sp<sup>3</sup>d
- 24. What do you understand by the term resonance energy? What are the essential rules for writing resonating structure.

- 25. With suitable examples explain the hybridization and geometry of molecules in octahedral complexes.
- 26. What is resonance? What are the rules which have restricted our choice of resonating structures?
- 27. Explain the hybridization and geometry of molecules in tetrahedral complexes by giving suitable examples.
- 28. What do you mean by resonance? Write the resonance structure of CO and cause of its low dipole moment.
- 29. .i) What are the essential conditions for writing resonating structures.
  - ii) Draw all the resonating structures of nitrous oxide.
- 30. State the geometrical arrangements possible in sp<sup>3</sup>d and sp<sup>3</sup>d<sup>2</sup>hybridisations.Name the dorbitals involved in these.

#### KARPAGAM ACADEMY OF HIGHER EDUCATION Unit-IV (Chemical Bonding – II)

	Unit-IV (Chemical Bo	nung – 11)			
Questions	Opt-1	Opt-2	Opt-3	opt-4	Answer
In the atoms retain their individuality in the molecule.	Valence bond theory	Molecular orbital theory	Crystal field theory	coordinate compounds	Valence bond theory
Resonance energy is equal to actual bond energy minus energy of the pure	ionic bond	Covalent bond	Coordinate bond	isomorphous compound	Covalent bond
Carbon dioxide has dipole moment equal to	one	two	zero	three	zero
Bond order in a molecule of NO is	1	2.5	3	3.5	2.5
Which one of the following compound is covalent	H <sub>2</sub>	CaO	KCl	Na <sub>2</sub> S	H <sub>2</sub>
BF3 involves hybridisation and the shape of the molecule is	sp	sp <sup>2</sup>	sp <sup>3</sup>	d <sup>2</sup> sp <sup>3</sup>	sp <sup>2</sup>
triangular planar. The change of the culobate ion is	totuch o duo l		tuis su al hiny manai dal	havecorol	totuch o duci
In a double hand connecting two stores, there is a charing of		square planar			four electrons
Which of the following compound has not on? hybridication					
which of the following compound has not sp3 hybridisation.		IN F13		564	
The shape of SICI4	planar triangle	octahedral	tetrahedral	square planar	tetrahedral
BF3 involves sp2 hybridisation and the shape of the molecule is	tetrahedral	square planar	triangular planar	linear	triangular planar
Which of the following be octahedral	SF <sub>6</sub>	BF4-	PCl <sub>5</sub>	H <sub>2</sub>	SF <sub>6</sub>
When the hybridisation state of carbon atom changes from sp3 to sp2 to	dooroogoo gradually	increases gradually	depropage with increasing	dooroogoo considerably	ingraage gradually
sp ,the angle between the hybridised orbitals	decreases gradually	increases gradually	decreases with increasing	decreases considerably	increases gradually
According to VSEPR theory the shape of water molecule is	octahedral	distorted tetrahedral	planar triangle	linear	distorted tetrahedral
Directed bond in water form an angle of	90∘C	120°C	105°C	60∘C	105∘C
Which of the following involves sp2 hybridisation	$CO_2$	$SO_2$	N <sub>2</sub> O	СО	$SO_2$
The structure of ICl2- is	trigonal	trigonal bipyramidal	octahedral	square planar	trigonal bipyramidal
A sp3 hybrid orbital contains	1/4 s-character	1/2 s-character	2/3 rd s-character	3/4 s-character	1/4 s-character
In which of the following the angle between the covalent bonds is maximum	SO <sub>2</sub>	CH <sub>4</sub>	NH <sub>3</sub>	H <sub>2</sub> O	CO <sub>2</sub>
Compounds formed by sp3d hybridisation will have structure	planar triangle	pyramidal	angular	trigonal bipyramidal	trigonal bipyramidal
Which one of the following is most polar	CCl <sub>4</sub>	CHCl <sub>3</sub>	CH <sub>3</sub> Cl	CH <sub>3</sub> OH	CH <sub>3</sub> OH
Which of the following possess strong covalent bond	H-Cl	CI-CI	C-Cl	B-Cl	H-Cl
Which one of the following is strongest bond	Cl-F	F-F	Br-F	Br-Cl	Br-F
Decreasing order of size of various hybrid orbitals is	sp>sp2>sp3	sn3>sn2>sn	sn2>sn>sn3	sp>sp3>sp2	sn3>sn2>sn
Bonded electron pairs present in octahedral SF6 molecule	3	4	6	8	6 6
In PCI5 molecule, P is	sp3 hybridised	dsp2 hybridised	ds3p hybridised	sp3d hybridised	sp3d hybridised
The equilateral triangle shape has	sp hybridisation	sp2 hybridisation	sp3 hybridisation	sp3d hybridisation	sp2 hybridisation
Which hybrid orbitals are used for bonding in a square planar molecule or ion	sp3	dsp2	sp	sp3d	dsp2
The molecule which has pyramidal shape	PCl <sub>3</sub>	SO3	SF <sub>6</sub>	NO3-	PCl <sub>3</sub>
Which molecule is not linear	CO <sub>2</sub>	SO <sub>3</sub>	C <sub>2</sub> H <sub>2</sub>	MgCl <sub>2</sub>	SO <sub>3</sub>
Which of the following molecule does not have a linear arrangement of molecules	C <sub>2</sub> H <sub>2</sub>	H <sub>2</sub> S	BeCl <sub>2</sub>	CS <sub>2</sub>	H <sub>2</sub> S
Which hybridisation results in non-planar orbitals	sn	sn?	sn3	dsp2	sn3
How many bonded electron pairs are present in IE7 molecule	зр 6	3p2 7	8	5	7
In compound X all the bond angles are exactly $109\circ 28'$ . The X is	chloromethane	iodoform	Carbon tetrachloride	Chloroform	Carbon tetrachloride
Which is expected to have linear structure	SO <sub>2</sub>	CO <sub>2</sub>	$CO_3^{2}$	SO4 <sup>2-</sup>	CO <sub>2</sub>
Which is not linear	CO <sub>2</sub>	HCN	C <sub>2</sub> H <sub>2</sub>	H <sub>2</sub> O	H <sub>2</sub> O
Which compound does not contain double or triple bond	C <sub>2</sub> H <sub>4</sub>	H <sub>2</sub> O	N <sub>2</sub>	HCN	H <sub>2</sub> O
H20 is notmolecule	linear	nyramidal	angular	aatabadral	linear
Thehybridisation of Ag in the linear compley [Ag(NH2)2]+ is	en3	pyrannuar en?	en	den?	en
The estat rule is not valid for the male!	sho	sp2	o bh	usp2	94 140
The octet rule is not valid for the molecule	$CO_2$	n <sub>2</sub> O	$O_2$	0	n <sub>2</sub> O

Which of the following molecule /ion has triangular pyramidal shape	BF <sub>3</sub>	NO <sup>3-</sup>	H3O+	CO <sub>3</sub> <sup>2-</sup>	H3O+
Which of the following does not have a covalent bond	O <sub>3</sub>	H3O+	PC15	HNO <sub>3</sub>	PC15
The unequal sharing of the bonded pair of electrons between two atoms in a molecule gives rise to	ionic bond	polar covalent bond	non-polar covalent bond	co-ordinate bond	polar covalent bond
Which of the following is not tetrahedral	BF <sup>4-</sup>	NH4+	CO3 <sup>2-</sup>	SO4 <sup>2-</sup>	CO <sub>3</sub> <sup>2-</sup>
The shape of SO42- ion is	tetrahedral	square planar	trigonal	trigonal planar	tetrahedral
A molecule with four bonded electron pairs on the central atom and no lone pair is likely to be	linear	tetrahedral	octahedral	trigonal planar	tetrahedral
The shape of CIO4- ion is	square planar	square pyramidal	tetrahedral	trigonal bipyramidal	tetrahedral
The bond angle in NH3 molecule is	109°28'	90°	107°	105°	107°
Shape of molecules is decided by	sigma bonds	Pi bonds	both sigma and pi bonds	neither sigma nor pi bonds	sigma bonds
Which one of the following is not tetrahedral	H <sub>2</sub> O	$C_2Cl_6$	CH <sub>4</sub>	diamond	H <sub>2</sub> O
Metallic sodium is	insulator	semiconductor	conductor of electricity	conductor only in molten state	conductor of electricity
For which of the following hybridisation the bond angle is maximum	sp	sp2	sp3	dsp2	sp
Aqueous solution contains	H+	OH+	H <sub>3</sub> O+	H2	H <sub>3</sub> O+
Which of the following has square planar geometry	HgCl2	CO3 <sup>2-</sup>	NH4+	Ni(CN) <sub>4</sub> <sup>2-</sup>	Ni(CN)4 <sup>2-</sup>
Phosphorous tri fluoride molecule is	square planar	trigonal bipyramidal	tetrahedral	trigonal pyramidal	trigonal pyramidal
Which is the highest melting halide	NaCl	NaBr	NaF	NaI	NaF
The hybridisation in SiCl4	sp3	sp2	sp	dsp2	sp3
The greater the number of in a contributing structure ,greater is the importance of that structure	covalent bonds	ionic bond	metallic bond	co-ordinate bond	covalent bonds
The resonance energy of carbon dioxide molecule is	142	138.2	156.5	162.2	138.2
The resonating structure fail to explain the paramagnetic character of	oxygen molecule	nitric oxide molecule	carbon dioxide molecule	nitrate ion	oxygen molecule

# Unit-V

# **Molecular Structures**

MO Approach: Rules for the LCAO method, bonding and antibonding MOs and their characteristics for s-s, s-p and p-p combinations of atomic orbitals, nonbonding combination of orbitals, MO treatment of homonuclear diatomic molecules of 1st and 2nd periods (including idea of s-p mixing) and heteronuclear diatomic molecules such as CO, NO and NO+.

Comparison of VB and MO approaches.

The molecular orbital (MO) theory is a powerful and extensive approach which describes electrons as delocalized moieties over adjacent atoms. The applications of the MO theory extend beyond the limitations of the Valence Shell Electron Pair Repulsion (VSEPR) model and the Valence Bond theory. These previous theories provide substantial information on predicting the structure and bonding properties of a molecule by portraying electrons as localized "balloons" of high density.

#### Introduction

Although VSEPR and the Valence Bond theory accurately predict bond properties, they fail to fully explain some molecules. The MO theory incorporates the wave character of electrons in developing MO diagrams. MO diagrams predict physical and chemical properties of a molecule such as shape, bond energy, bond length and bond angle. They also provide information in predicting a molecule's electronic spectra and paramagnetism. The objective of this wiki is to provide readers with the fundamental steps in constructing simple homonuclear and heteronuclear diatomic molecular orbital diagrams. These steps may then be extrapolated to construct more difficult polyatomic diagrams.

#### **Molecular Orbitals**

The region an electron is most likely to be found in a molecule. A MO is defined as the combination of atomic orbitals.

# **Homonuclear Diatomics**

Molecules consisting of two identical atoms are said to be homonuclear diatomic, such as:  $H_2$ ,  $N_2$ ,  $O_2$ , and  $F_2$ .

# **Heteronuclear Diatomics**

Molecules consisting of two non-identical atoms are said to be heteronuclear diatomic, such as: CO, NO, HF, and LiF.

# **Bonding and Antibonding Orbitals**

Orbitals that are out-of-phase with one of another are "antibonding" orbitals because regions with dense electron probabilities do not merge which destabilizes the molecule. "Bonding" orbitals are less energetic than antibonding atomic orbitals and are in-phase, as depicted in the figure below. Note how the bonding orbitals come together constructively, while the antibonding orbitals do not. According to Valence bond theory:

A covalent bond is formed by the overlapping of partially filled orbitals of two
 batch atoms.

• Overlapping orbitals must have electrons with opposite spin.

Atoms involved in bond formation should have unpaired electrons.

The number of covalent bonds formed by an atom would be equal to the number of half filled orbital.

Resulting molecular orbital is obtained by the combination of the two wave functions (AOs) of two unpaired electrons.

• Atoms which are involved in bond formation maintain their identity.

#### **Molecular Orbital Theory :**

A covalent bond is formed by the overlapping of atomic orbitals which form molecular orbital. Bonding electrons occupy molecular orbital not atomic orbital.

An electron in a molecular orbital is polycentric because it is influenced by more than one nuclei.

Formation of molecular orbital is based on the linear combination of atomic orbitals (LCAO).

There are two kinds of molecular orbitals:

- 1. BONDING MOLECULAR ORBITAL
- 2. ANTI-BONDING MOLECULAR ORBITAL

### Sigma bond:

- 1. Sigma bond is formed by the linear or head to head or end on overlapping of orbitals.
- 2. Sigma bonds are the strong bonds due to maximum overlapping of orbitals.
- 3. Electron density is maximum around the bond axis.

4. Compounds having sigma bonds are stable and hence less reactive.

5. First bond formed between two atoms is always a sigma bond.

6. s-s overlap, s-p overlap and p-p overlap give rise to sigma bond.

Examples of Sigma bond :

### s-s overlap in H<sub>2</sub> molecule

Hydrogen molecule consists of two H-atoms. Each atom contains one electron in 1s-orbital.

 $E.C = 1s^1$ 

According to molecular orbital theory two  $1s^1$  orbitals of two H-atoms overlap linearly to produce two types of molecular orbitals.

- 1. Bonding molecular orbital (□-orbital)
- 2. Anti-bonding molecular orbital ( $\Box^*$ -orbital)

Bonding molecular orbital ( $\Box$ -orbital) has lower energy and results in the covalent bond formation which is a sigma bond between two H-atoms while the anti-bonding molecular orbital possess high energy remains unoccupied.

### s-p overlap in HF molecule

s-p type overlap occurs in HF molecule when one **1s-orbital** of hydrogen atom overlaps **2p-** orbital of fluorine to form two types of molecular orbitals.

- 1. Bonding molecular orbital (□-orbital)
- 2. Anti-bonding molecular orbital ( $\Box^*$ -orbital)

Bonding molecular orbital ( $\Box$ -orbital) has lower energy and results in the covalent bond formation (sigma bond) between H-atom and F-atomwhile the anti-bonding molecular orbital ( $\Box$ \*-orbital) possess high energy remains unoccupied.

### p-p overlap in HF molecule

p-p type overlap occurs in  $F_2$  molecule when one  $2p_z$ -orbital of one fluorine atom overlaps  $2p_z$ orbital of other fluorine atom form two types of molecular orbitals.

1. Bonding molecular orbital ( $\Box$ -orbital)

2. Anti-bonding molecular orbital ( $\Box^*$ -orbital)

Bonding molecular orbital ( $\Box$ -orbital) has lower energy and results in the covalent bond formation (sigma bond) between two F-atoms while the anti-bonding molecular orbital ( $\Box^*$ -orbital) possess high energy remains unoccupied.

# Strength of sigma bond:

The relative strength of a sigma bond is related to the extent of overlap of the atomic orbitals.

This is known as the 'principle of maximum overlap'.

Due to spherical charge distribution in s-orbital, generally s-s overlapping is not so effective as s-p and p-p overlapping.

Where p-orbitals have directional charge distribution and longer lobes which cause more effective overlapping. Thus s-s sigma bond is relatively weak.

Order of the strength of sigma bonds is as follows:

nature of sigma bond	S-S	s-p	р-р
bond strength	1.0	1.73	3.0

# Pi bond:

A Pi bond is formed by the lateral or side ways or parallel overlapping of P-orbital of the atoms which are already bonded by a sigma bond and their axes are coplanar. This type of overlap generates two types of molecular orbitals:

# (a) Pi-bonding molecular orbital ( -orbital)

# (b) Pi-antibonding molecular orbital ( $\Box^*$ -orbital)

A pi-bonding orbital has two regions of electron density below and above the nodal plane. The electron contained in it are called pi-bonding electrons which form the pi bond. It is not linearly symmetrical with respect to the bond axis, rather it has a nodal plane.

Pi-bonds are weaker than sigma bonds.

In Pi-bonds, electron density lies in the regions above and below the nuclei.

Compounds having pi bonds are more reactive.

Pi bond is formed when two atoms already bonded by a sigma bond.



#### Phases and nodes

Phases are designated either (+) or (-) relative to their wave "up" or wave "down" displacements. A node occurs if the phase signs change from (+) to (-) or vice versa. It is important to notice that the phase signs do NOT symbolize charges. Nodes are regions where the probability of finding an electron is ZERO.

# Sigma and Pi Bonds

A sigma-bond is an "end-to-end" bond formed from symmetric atomic orbitals. A pi-bond is formed from a "sideways" overlap.

### Valence bond theory



**Molecular Orbital Diagrams** 



 The Y-axis of a MO diagram represents the total energy (not potential nor Gibbs Energy) of the orbitals.

- 2. Individual atomic orbitals (AO) are arranged on the far left and far right of the diagram.
- Overlapping atomic orbitals produce molecular orbitals located in the middle of the diagram. These MO overlap with either a sigma or pi bond and are designated in bonding, nonbonding, or antibonding orbitals with respect to their phases.
- Electrons from the atomic orbitals are assigned molecular orbitals in accordance with the Pauli Exclusion Principle. Lower energy MOs are filled first, followed by consecutively increasing orbitals.

### FUNDAMENTAL STEPS IN DERIVING MO DIAGRAMS

There are several steps common in all MO diagrams. Understanding these basic steps to derive simple homonuclear and heteronuclear MOs will enable us to construct more complicated, polyatomic diagrams.

#### STEP 1

Find the valence electron configuration of each atom in the molecule. The valence electrons will be placed on the atomic orbital for that atom. Do this for **each** atom. Ex) Boron (B) =  $(He)2s^2 2p^1 = 3$  valence electrons

#### STEP 2

Decide if the molecule is homonuclear of heteronuclear. If the molecule is homonuclear, the AOs will be symmetric. Heteronuclear AOs will be slightly different because the more electronegative atom will be placed lower on the diagram. This is due to lone pairs of electrons being more

stable on more electronegative elements leading them to be lower in energy. Ex) HF = "F" will be placed lower in the diagram than "H".

### **STEP 3**

Fill molecular orbitals using energy and bonding properties of the overlapping atomic orbitals. Keep in mind the energy of the atomic orbitals and molecular orbitals! The following factors contribute to the position of one MO with respect to other MOs.

- More nodes = more energetic = higher MOs
- Sigma orbitals are stronger than pi bonds
- Antibonding MOs are higher in energy than bonding MOs
- Constructive overlap = fewer nodes = more stable (less energetic)
- Destructive overlap = more nodes = less stable (more energetic)

### **STEP 4**

Use the diagram to predict properties of the molecule. Remember: the number of individual atomic orbitals should equal the number of MOs! Ex) Bond order, bond angle, paramagnetism, etc.

### Building Molecular Orbital Diagrams for Homonuclear and Heteronuclear

### **Diatomic Molecules**

Due to symmetry of the molecule, homonuclear MO's are less difficult to derive than heteronuclear molecules and polyatomic molecules. (Such as H<sub>2</sub>O, NH<sub>3</sub>, and CH<sub>4</sub>.)

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#### Molecular Structures (2017 – 18) batch

However, **notice** the difference between orbitals of homonuclear diatomics for elements with an atomic number less than or equal to 7 versus more than 7. In general,  $B_2$ ,  $C_2$ , and  $N_2$  have the MO diagram depicted on the left.  $O_2$  and  $F_2$  have the MO on the right. Why does this difference exist?



Constructing MO diagrams for heteronuclear molecules require the same 4 steps as above. However, recall that the more electronegative atom will be lower on the diagram. The best way to learn how to draw MO diagrams is to work on practice problems. Try the following MO's on your own, and then check with the answers provided

#### **MO diagram for N2:**

**STEP 1:**  $N_2 = (He) 2s^2 2p^3$  (5 valence electrons for each atom)

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**STEP 2:** Homonuclear molecule; each "N" AO will be equivalent in energy.





Figure 2: MD. Diagram of Na

### **STEP 4**

BOND ORDER: 1/2 (Number of Bonding orbitals – Number of Antibonding orbitals)

$$=1/2(8-2)=3.$$

MO for O<sub>2</sub>:

**STEP 1:**  $O_2 = (He) 2s^2 2p^4$  (6 valence electrons for each atom)

**STEP 2:** Homonuclear molecule; each "O" AO will be equivalent in energy.

**STEP 3:** Fill the MOs with electrons:

**STEP 4:** Check. Do the number of AO's = number of MO's? If so, calculate the bond order.

BOND ORDER: 1/2 (#Bonding orbitals - #Antibonding orbitals)

=1/2(8-4)=2. Consistent with Oxygen's double bond.

# MO diagram for HF

**STEP 1:**  $H = 1s^1$  (1 valence electron) F= (He)  $2s^2 2p^5$  (7 valence electrons)

STEP 2: Heteronuclear molecule; "F" will be lower on the diagram.

**STEP 3:** Fill the MOs with electrons:



# Bonding and antibonding orbitals

Molecular orbital theory is concerned with the combination of atomic orbitals to form new molecular orbitals. These new orbitals arise from the linear combination of atomic orbitals to form bonding and antibonding orbitals. The bonding orbitals are at a lower energy than the antibonding orbitals, so they are the first to fill up. By figuring out the molecular orbitals, it is easy to calculate bond order.

#### Introduction

The valence bond theory is an extension of the Lewis Structures that considers the overlapping of orbitals to create bonds. The valence bond theory is only limited in its use because it does not explain the molecular geometry of molecules very well. This is where hybridization and the molecular orbital theory comes into place.

### Hybridization

Hybridization is a simple model that deals with mixing orbitals to from new, hybridized, orbitals. This is part of the valence bond theory and helps explain bonds formed, the length of bonds, and bond energies; however, this does not explain molecular geometry very well.

#### sp:

An example of this is acetylene ( $C_2H_2$ ). This combines one s orbital with one p orbital. This means that the s and p characteristics are equal.

 $sp^2$ 

An example of this is ethylene ( $C_2H_4$ ). This is the combination of one s orbital and two p orbitals.

### sp<sup>3</sup>

An example of this is methane (CH<sub>4</sub>). This is the combination of one s orbital and three p orbitals.

If you add the exponents of the hybridized orbitals, you get the amount of sigma bonds associated with that bond. The  $sp^2$  hybridized orbital has one p orbitals that is not hybridized and so it can form a pi bond. This means that  $sp^2$  orbitals allow for the formation of a double bond. Also, sp hybridized orbitals form a triple bond.

# **Antibonding vs. Bonding Orbitals**

Electrons that spend most of their time between the nuclei of two atoms are placed into the bonding orbitals, and electrons that spend most of their time outside the nuclei of two atoms are placed into antibonding orbitals. This is because there is an increasing in electron density between the nuclei in bonding orbitals, and a decreasing in electron density in antibonding orbitals (Chang 459). Placing an electron in the bonding orbital stabilizes the molecule because it is in between the two nuclei. Conversely, placing electrons into the antibonding orbitals will decrease the stability of the molecule. Electrons will fill according to the energy levels of the orbitals. They will first fill the lower energy orbitals, and then they will fill the higher energy

orbitals. If a bond order of zero is obtained, that means that the molecule is too unstable and so it will not exist.

Below are a few examples of bonding and antibonding orbitals drawn out:

- Hydrogen Example (homonuclear):
- Oxygen Example (homonuclear):
- Hydrogen Flouride Example (heteronuclear)

# **Bond Order**

Bond order is the amount of bonds formed between two atoms. For example, two bonds are formed between oxygen atoms, so the bond order is 2. The following is the equation to find bond order.

1/2(electrons in bonding molecular orbitals - electrons in antibonding molecular orbitals)

Bond order gives information about bond length and strength. Generally, higher bond order correlates to a shorter bond length. This is due to the greater number of bonds between the atoms. In addition, because of the greater number of bonds between the atoms, the strength should also be greater as bond order increases.

# Molecular orbital

Molecular orbital theory and Molecular orbital diagram

Complete acetylene (H–C=C–H) molecular orbital set. The left column shows MO's which are occupied in the ground state, with the lowest-energy orbital at the top. The white and grey line visible in some MO's is the molecular axis passing through the nuclei. The orbital wave functions are positive in the red regions and negative in the blue. The right column shows virtual MO's which are empty in the ground state, but may be occupied in excited states.

In chemistry, a **molecular orbital** (**MO**) is a mathematical function describing the wave-like behavior of an electron in a molecule. This function can be used to calculate chemical and physical properties such as the probability of finding an electron in any specific region. The term orbital was introduced by Robert S. Mulliken in 1932 as an abbreviation for one-electron orbital wave function.<sup>[1]</sup> At an elementary level, it is used to describe the region of space in which the function has a significant amplitude. Molecular orbitals are usually constructed by combining atomic orbitals or hybrid orbitals from each atom of the molecule, or other molecular orbitals from groups of atoms. They can be quantitatively calculated using the Hartree–Fock or self-consistent field (SCF) methods.

A molecular orbital (MO) can be used to represent the regions in a molecule where an electron occupying that orbital is likely to be found. Molecular orbitals are obtained from the combination of atomic orbitals, which predict the location of an electron in an atom. A molecular orbital can specify the electron configuration of a molecule: the spatial distribution and energy of one (or one pair of) electron(s). Most commonly an MO is represented as a linear combination of atomic orbitals (the LCAO-MO method), especially in qualitative or very approximate usage. They are invaluable in providing a simple model of bonding in molecules, understood through molecular orbital theory. Most present-day methods incomputational chemistry begin by calculating the MOs of the system. A molecular orbital describes the behavior of one electron in the electric field generated by the nuclei and some average distribution of the other electrons. In the case of two electrons occupying the same orbital, the Pauli principle demands that they have opposite spin. Necessarily this is an approximation, and highly accurate descriptions of the molecular electronic wave function do not have orbitals (see configuration interaction).

#### Formation of molecular orbitals

Molecular orbitals arise from allowed interactions between atomic orbitals, which are allowed if the symmetries (determined from group theory) of the atomic orbitals are compatible with each other. Efficiency of atomic orbital interactions is determined from the overlap (a measure of how well two orbitals constructively interact with one another) between two atomic orbitals, which is significant if the atomic orbitals are close in energy. Finally, the number of molecular orbitals that form must equal the number of atomic orbitals in the atoms being combined to form the molecule.

In the molecular structure, the molecular orbitals can be obtained from the "Linear combination of atomic orbitals molecular orbital method" ansatz. Here, the molecular orbitals are expressed as linear combinations of atomic orbitals.

#### Linear combinations of atomic orbitals (LCAO)

Molecular orbitals were first introduced by Friedrich Hund and Robert S. Mulliken in 1927 and 1928. The linear combination of atomic orbitals or "LCAO" approximation for molecular orbitals was introduced in 1929 by Sir John Lennard-Jones. His ground-breaking paper showed how to derive the electronic structure of thefluorine and oxygen molecules from quantum

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principles. This qualitative approach to molecular orbital theory is part of the start of modern quantum chemistry. Linear combinations of atomic orbitals (LCAO) can be used to estimate the molecular orbitals that are formed upon bonding between the molecule's constituent atoms. Similar to an atomic orbital, a Schrödinger equation, which describes the behavior of an electron, can be constructed for a molecular orbital as well. Linear combinations of atomic orbitals, or the sums and differences of the atomic wavefunctions, provide approximate solutions to the Hartree–Fock equations which correspond to the independent-particle approximation of the molecular Schrödinger equation. For simple diatomic molecules, the wavefunctions obtained are represented mathematically by the equations

$$\Psi = c_a \psi_a + c_b \psi_b$$
$$\Psi^* = c_a \psi_a - c_b \psi_b$$

where  $\Psi$  and  $\Psi^*$  are the molecular wavefunctions for the bonding and antibonding molecular orbitals, respectively,  $\psi_a$  and  $\psi_b$  are the atomic wavefunctions from atoms a and b, respectively, and  $C_a$  and  $C_b$  are adjustable coefficients. These coefficients can be positive or negative, depending on the energies and symmetries of the individual atomic orbitals. As the two atoms become closer together, their atomic orbitals overlap to produce areas of high electron density, and, as a consequence, molecular orbitals are formed between the two atoms. The atoms are held together by the electrostatic attraction between the positively charged nuclei and the negatively charged electrons occupying bonding molecular orbitals.

# Bonding, antibonding, and nonbonding MOs

When atomic orbitals interact, the resulting molecular orbital can be of three types: bonding, antibonding, or nonbonding.

# **Bonding MOs:**

- Bonding interactions between atomic orbitals are constructive (in-phase) interactions.
- Bonding MOs are lower in energy than the atomic orbitals that combine to produce them.

# **Antibonding MOs:**

- Antibonding interactions between atomic orbitals are destructive (out-of-phase) interactions, with a nodal plane where the wavefunction of the antibonding orbital is zero between the two interacting atoms
- Antibonding MOs are higher in energy than the atomic orbitals that combine to produce them.

# Nonbonding MOs:

- Nonbonding MOs are the result of no interaction between atomic orbitals because of lack of compatible symmetries.
- Nonbonding MOs will have the same energy as the atomic orbitals of one of the atoms in the molecule.

#### Sigma and pi labels for MOs

The type of interaction between atomic orbitals can be further categorized by the molecular-orbital symmetry labels  $\sigma$  (sigma),  $\pi$  (pi),  $\delta$  (delta),  $\varphi$  (phi),  $\gamma$  (gamma) etc. paralleling the symmetry of the atomic orbitals s, p, d, f and g. The number of nodal planes containing the internuclear axis between the atoms concerned is zero for  $\sigma$  MOs, one for  $\pi$ , two for  $\delta$ , etc.

#### **σ** symmetry

A MO with  $\sigma$  symmetry results from the interaction of either two atomic s-orbitals or two atomic p<sub>z</sub>-orbitals. An MO will have  $\sigma$ -symmetry if the orbital is symmetric with respect to the axis joining the two nuclear centers, the internuclear axis. This means that rotation of the MO about the internuclear axis does not result in a phase change. A  $\sigma^*$ orbital, sigma antibonding orbital, also maintains the same phase when rotated about the internuclear axis. The  $\sigma^*$  orbital has a nodal plane that is between the nuclei and perpendicular to the internuclear axis.

#### $\pi$ symmetry

A MO with  $\pi$  symmetry results from the interaction of either two atomic  $p_x$  orbitals or  $p_y$  orbitals. An MO will have  $\pi$  symmetry if the orbital is asymmetric with respect to rotation about the internuclear axis. This means that rotation of the MO about the internuclear axis will result in a phase change. There is one nodal plane containing the internuclear axis, if real orbitals are considered.

A  $\pi^*$  orbital, pi antibonding orbital, will also produce a phase change when rotated about the internuclear axis. The  $\pi^*$  orbital also has a second nodal plane between the nuclei.<sup>[11][12][13][14]</sup>

#### δ symmetry

A MO with  $\delta$  symmetry results from the interaction of two atomic  $d_{xy}$  or  $d_{x}^{2}$ ,  $y^{2}$  orbitals. Because these molecular orbitals involve low-energy d atomic orbitals, they are seen intransition-metal complexes. A  $\delta$  bonding orbital has two nodal planes containing the internuclear axis, and a  $\delta^{*}$  antibonding orbital also has a third nodal plane between the nuclei.

#### **φ** symmetry



Suitably aligned f atomic orbitals overlap to form phi molecular orbital (a phi bond)

Theoretical chemists have conjectured that higher-order bonds, such as phi bonds corresponding to overlap of f atomic orbitals, are possible. There is as of 2005 only one known example of a molecule purported to contain a phi bond (a U–U bond, in the moleculeU<sub>2</sub>).

### Gerade and ungerade symmetry]

For molecules that possess a center of inversion (centrosymmetric molecules) there are additional labels of symmetry that can be applied to molecular orbitals. Centrosymmetric molecules include:

- Homonuclear diatomics, X<sub>2</sub>
- Octahedral, EX<sub>6</sub>
- Square planar, EX<sub>4</sub>.

Non-centrosymmetric molecules include:

- Heteronuclear diatomics, XY
- Tetrahedral, EX<sub>4</sub>.

If inversion through the center of symmetry in a molecule results in the same phases for the molecular orbital, then the MO is said to have gerade (g) symmetry, from the German word for even. If inversion through the center of symmetry in a molecule results in a phase change for the molecular orbital, then the MO is said to have ungerade (u) symmetry, from the German word for odd. For a bonding MO with  $\sigma$ -symmetry, the orbital is  $\sigma_g$  (s' + s" is symmetric), while an antibonding MO with  $\sigma$ -symmetry the orbital is  $\sigma_u$ , because inversion of s' – s" is antisymmetric. For a bonding MO with  $\pi$ -symmetry the orbital is  $\pi_u$  because inversion through the center of symmetry for would produce a sign change (the two p atomic orbitals are in phase with each other but the two lobes have opposite signs), while an antibonding MO with  $\pi$ -symmetry is  $\pi_g$  because inversion through the center of symmetry for would not produce a sign change (the two p orbitals are antisymmetric by phase).<sup>[16]</sup>

#### **Bonding in molecular orbitals**

#### **Orbital degeneracy**

Molecular orbitals are said to be degenerate if they have the same energy. For example, in the homonuclear diatomic molecules of the first ten elements, the molecular orbitals derived from the  $p_x$  and the  $p_y$  atomic orbitals result in two degenerate bonding orbitals (of low energy) and two degenerate antibonding orbitals (of high energy).

#### **Ionic bonds**

When the energy difference between the atomic orbitals of two atoms is quite large, one atom's orbitals contribute almost entirely to the bonding orbitals, and the others atom's orbitals contribute almost entirely to the antibonding orbitals. Thus, the situation is effectively that some electrons have been transferred from one atom to the other. This is called an (mostly) ionic bond.

### **Bond order**

The bond order, or number of bonds, of a molecule can be determined by combining the number of electrons in bonding and antibonding molecular orbitals. A pair of electrons in a bonding orbital creates a bond, whereas a pair of electrons in an antibonding orbital negates a bond. For example, N<sub>2</sub>, with eight electrons in bonding orbitals and two electrons in antibonding orbitals, has a bond order of three, which constitutes a triple bond.

Bond strength is proportional to bond order—a greater amount of bonding produces a more stable bond—and bond length is inversely proportional to it—a stronger bond is shorter.

There are rare exceptions to the requirement of molecule having a positive bond order. Although Be<sub>2</sub> has a bond order of 0 according to MO analysis, there is experimental evidence of a highly unstable Be<sub>2</sub> molecule having a bond length of 245 pm and bond energy of 10 kJ/mol.
# **Possible Questions**

## PART A (Objective type each questions carries one mark)

1. HCl involves

a)s-s overlapping b)**sp-overlapping** c)p-p overlapping d) $P\pi$ - $P\pi$  overlapping

2. Which shows non directional bonding

a)BCl<sub>3</sub> b)CsCl c)NCl<sub>3</sub> d)BeCl<sub>3</sub>

- Which species has lone pair of electrons on central atom
   a)CCl<sub>4</sub> b)CH<sub>4</sub> c)NH<sup>4+</sup> d)H<sub>2</sub>O
- 4. Which atomic orbital is always involved in sigma bonding onlya)s b)p c)d d)f
- The molecule having bond order 3 is
   a)H<sub>2</sub>b)N<sub>2</sub>c)O<sub>2</sub> d) He<sup>2+</sup>
- 6. The number of antibonding electron pairs in  $O_2^{2-}$  molecular ion on the basis of MOT is (Atomic number of O=8)

a)**4** b)3 c)2 d)5

7.  $O_2$  molecule is

a)**paramagnetic** b)diamagnetic c)ferromagnetic d)ferromagnetic

8. Which one of the following molecule is paramagnetic

a)CO<sub>2</sub> b)SO<sub>2</sub> c)NO d)H<sub>2</sub>O

9. Lateral overlapping is expected in

a)sigma bonds b)pi bonds c)metallic bonds d)coordinate bonds

10. The energy of antibonding molecular orbital is

a)**greater than the bonding M.O** b)smaller than the bonding M.O c)equal to that of bonding orbital d)very low

11. Strongest bond is formed by the overlapping of

a)**2s and 2p orbitals** b)2p and 2p orbitals c)2s and 2s orbitals d)is orbital

12. Greater than the bonding M.O

a)Pi bond b)increases bond lengthc)**decrease s bond length**d)Distorts the geometry of molecule

13. A pi bond is formed by sideway overlapping of

a)s-s orbitals b)**p-p orbitals**c)s-p orbitals d)s-p-s orbitals

- 14. Sigma bond is formed bya)orbitalsb)p-p orbitalsc)end to end overlapping of orbitals d) shortens the internuclear distance
- 15. Fluorine molecule is formed by the overlap of

a)s-p orbitalsb)s-s c)**p-p orbitals by end to end manner**d)p-p orbitals by side to side manner

16. Linear combination of two hybridised orbitals , belonging to two atoms and each having one electron leads to

a)**sigma bond**b)double bondc)co-ordinate covalent bond d)pi bond

17. When two atomic orbitals combine they form

a)one molecular orbital b)**two molecular orbitals** c)two bonding molecular orbital

d)two antibonding molecular orbitals

18. Which has the largest bond anglea)H<sub>2</sub>O<sub>2</sub> b)NH<sub>3</sub> c)H<sub>2</sub>S d)H<sub>2</sub>O

19. Which of the following species has the longest bond length

a)NO+ b)O<sup>2-</sup>c)O<sup>2+</sup> d)N<sup>2+</sup>

20. Hybridisation involves

a)addition of an electron pair b)**mixing up of atomic orbitals** c)separation of orbitals

d)removal of electron pair

# PART B (each questions carries two marks)

- 21. What is LCAO molecular orbital approach
- 22. What is bond order?
- 23. What is meant by bonding molecular orbital.
- 24. What is meant by non bonding molecular orbital
- 25. What is the bond order of  $N_2$  and  $O_2$  molecule.

# PART C (each questions carries six marks)

- 26. What are the main postulates of VSEPR theory.
- 27. Distinguish between molecular orbital theory and valence bond theory.
- 28. Distinguish two aspects of the following:
  - i) atomic and molecular orbitals

- ii) bonding and antibonding orbitals
- 29. Draw MO diagram of CO and calculate its bond order.
- 30. What is the meaning of overlap of atomic orbitals? Explain s-s,s-p and p-p overlap with examples.
- 31. With the help of M.O. diagrams explain why
  - i) the bond order in  $N_2^+$  ion is less than that in  $N_2$  molecule whereas the bond order in  $O_2^+$  is greater than that in  $O_2$  molecule.
  - ii) the bond energy of  $NO^+$  is higher than that of NO.
- 32. Explain how the atomic orbitals combine to form bonding and antibonding M.O. What are the limitations to such combination.
- 33. Draw the molecular orbital diagram of CO molecule and explain.
- 34. Sketch the shapes of Mo obtained by the overlap of
  - i) two s-orbitals ii)end on overlap of two p-orbitals iii)sidewise overlap of two p-orbitals
- 35. Write the electronic configuration of NO molecule.
  - iii) What is meant by bond order
  - iv) Will the bond length be shorter or longer than in NO<sup>+</sup>
- 36. How many unpaired electrons will be present.

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#### Unit-V (Molecular Structures)

Questions	Opt-1	Opt-2	Opt-3	opt-4	Answer
The molecule having bond order 3 is	H <sub>2</sub>	N <sub>2</sub>	0 <sub>2</sub>	He2+	N <sub>2</sub>
The number of antibonding electron pairs in O22- molecular ion on the	1	2	2	5	4
basis of MOT is (Atomic number of O=8)	+	5	2		4
O2 molecule is	paramagnetic	diamagnetic	ferromagnetic	ferrimagnetic	paramagnetic
Which of the following is paramagnetic	0 <sub>2</sub>	CN-	со	NO+	O <sub>2</sub>
The bond order of O2+ is	2	2.5	1.5	3	2.5
Which one of the following molecule is paramagnetic	$CO_2$	SO <sub>2</sub>	NO	H <sub>2</sub> O	NO
Which of the following is not paramagnetic	NO	S <sup>2-</sup>	O <sup>2-</sup>	N <sup>2-</sup>	S <sup>2-</sup>
The molecular species having highest bond order is	0 <sub>2</sub>	O <sup>2-</sup>	$O^{2+}$	H2+	$O^{2^+}$
Which species is paramagnetic	O <sup>2-</sup>	CH <sup>3-</sup>	со	NO3-	O <sup>2-</sup>
The linear combination of two hybridised orbitals each having one electron and belonging two different atoms leads to a	sigma bond	Pi bonds	double bond	coordinate bond	sigma bond
Strongest bond is formed by the overlapping of	2s and 2p orbitals	2p and 2p orbitals	2s and 2s orbitals	is orbital	2s and 2p orbitals
The bond order of CO molecule on the basis of M.O. theory is	0	2	3	1	3
Which can not exist on the basis of MO theory	0 <sub>2</sub>	He2+	NO3-	He2	He2
When two atomic orbitals combine they form	one molecular orbital	two molecular orbitals	two bonding molecular orbital	two antibonding molecular orbitals	two molecular orbitals
Which has the largest bond angle	H2O2	NH <sub>3</sub>	H <sub>2</sub> S	H <sub>2</sub> O	NH <sub>3</sub>
Which of the following species has the longest bond length	NO+	O <sup>2-</sup>	02+	N2+	02+
HCI involves	s-s overlapping	sp-overlapping	p-p overlapping	Ρπ-Ρπ overlapping	sp-overlapping
Which shows non directional bonding	BCI3	CsCl	NCI3	BeCl3	
Which species has lone pair of electrons on central atom	CCI4	CH4	NH4+	H <sub>2</sub> O	H <sub>2</sub> O
Which atomic orbital is always involved in sigma bonding only	s	р	d	f	s
Lateral overlapping is expected in	sigma bonds	pi bonds	metallic bonds	coordinate bonds	pi bonds
The energy of antibonding molecular orbital is	greater than the bonding M.O	smaller than the bonding M.O	equal to that of bonding orbital	very low	greater than the bonding M.O
Pi bond	increases bond length	decrease s bond length	Distorts the geometry of molecule	makes homoatomic molecules more reactive	decrease s bond length
The example of the p-p orbital overlapping is the formation of	H2 molecule	Cl2 molecule	Hydrogen chloride	Hydrogen bromide molecule	
Maximum covalency of an atom of an element is equal to	No. of unpaired lectrons in the s and p-orbitals	No.of unpaired electrons in the p-orbitals	total number of s and p- electrons in the outermost shell	total number of electrons in the p-orbitals	total number of s and p- electrons in the outermost shell
A pi bond is formed by sideway overlapping of	s-s orbitals	p-p orbitals	s-p orbitals	s-p-s orbitals	p-p orbitals
Sigma bond is formed by	orbitals	p-p orbitals	end to end overlapping of orbitals	shortens the internuclear distance	end to end overlapping of orbitals
Fluorine molecule is formed by the overlap of	s-p orbitals	S-S	p-p orbitals by end to end manner	p-p orbitals by side to side manner	p-p orbitals by end to end manner
Linear combination of two hybridised orbitals , belonging to two atoms and each having one elctron leads to	sigma bond	double bond	co-ordinate covalent bond	pi bond	sigma bond
Which is not characteristic of pi bond	pi bond is formed when asigma bond already exists	Pi bonds are formed from hybrid orbitals	pi bond may be formed by the overlapping of p and or d orbitals	pi bond results from lateral overlap of atomic orbitals	pi bonds are formed from hybrid orbitals
Which of the following has maximum bond energy	02	02+	O <sub>2</sub> <sup>2-</sup>	O <sup>2-</sup>	02+
Hybridisation involves	addition of an electron pair	mixing up of atomic	separation of orbitals	removal of electron pair	mixing up of atomic
The bond strength increases	with increasing bond order	with increasing extent of overlapping of orbitals	with decreasing diference between energies of overlapping orbitals	with decreasing order	with increasing bond order

The HCN molecule contains	3 sigma and 2 pi bonds	2 sigma and 2 pi bonds	3 sigma and 3 pi bonds	2 sigma one pi bond	2 sigma and 2 pi bonds	
The formation of N2+ from N2 , the electron is removed from	a σ orbital	π orbital	σ* orbital	π* orbital	a σ orbital	
MOT put forwarded by	Hund and Mullikan	Heisenberg	debroglie	Aufbau	Hund and Mullikan	
The valence shell configuration of O-atom has unpaired						,
electrons	1	. 2	4	. 3	3	-
Hydrogen molecule is formed by	s-s overlap	s-p overlap	p-p head to head overlap	p-p side to side overlap	s-s overlap	
Hydrogen fluoride is formed by	s-s overlap	s-p overlap	p-p head to head overlap	p-p side to side overlap	s-p overlap	
The example of ammonia molecule belongs to	s-p overlap	p-p head to head overlap	p-p side to side overlap	s-s overlap	s-p overlap	
Oxygen molecule is formed by the overlap of	s-p overlap	s-s overlap	p-p head to head overlap	p-p overlap	p-p head to head overlap	
p-p head to head overlap is otherwise called as	end to end overlap	sideway overlap	lateral overlap	sidewise	end to end	overlap
Nitrogen molecule is an example for	p-p head to head overlap	s-p overlap	s-s overlap	p-p overlap	p-p head to head overlap	
p-p side to side overlap is otherwise called as	lateral overlap	end to end overlap	head on overlap	linear overlap	lateral overlap	
Two s-orbitals on two different atom is	s-s overlap	s-p overlap	p-p head to head overlap	p-p side to side overlap	s-s overlap	
The electron density in this bond is distributedabout the	summatrical	ungummetrical	parallal	nornondicular	our manatrical	
nuclear axis	symmetrical	unsymmetrical	parallel	perpendicular	symmetrical	
Atomic orbitals are	polycenteric	monocentric	bicentric	zero	monocentric	
Bonding MO is formed by the combination of electron waves of the	same	opposite	positive and negative	negative and positive	same	
sign						
Molecular orbitals are	polycenteric	monocentric	bicentric	zero	polycenteric	
Antibonding MO possessesenergy than the atomic orbitals	lossor	higher	very less	low and high	higher	
from which it is formed						
In antiboding MO the electron density in between the nuclei is	low	high	very low	very low	low	
and hence the repulsion between the nuclei high						
Bonding MO possessesenergy than the atomic orbitals from	higher	lower	very high		lower	
which it is formed	inglici	lower	verynign		lowel	
The degenerate molecular orbitals obeyed by	Hund's rule	Heisenberg	debroglie	Aufbau	Hund's rule	
The maximum number of electrons that a MO can contain two. These	pauli exclusion principle	Heisenberg	Hund's rule	Aufbau	pauli exclusion principle	
two electrons must have opposite spin is called						
Resonance plays an important role in	VBT	MOT	CFT	Metallic	VBT	
Thestarts with the nuclei of the constituent atoms	VBT	MOT	CFT	Metallic	MOT	
Bond order of Boron molecule is equal to	1	. 2	3	4	l 1	L
The neon molecule has the bond order of	0	1	. 2		3 C	)
The NO molecule is	paramagnetic	diamagnetic	ferromagnetic	ferrimagnetic	paramagnetic	
In Nitrosyl ion (No+) it hasin character.	paramagnetic	diamagnetic	ferromagnetic	ferrimagnetic	diamagnetic	

[17CHU101] (For the candidates admitted from 2017 & onwards) **B.Sc. DEGREE EXAMINATION First Semester** Chemistry **INTERNAL TEST - I** ATOMIC STRUCTURE AND CHEMICAL BONDING

Maximum: 50 marks

Time: 2 Hours

#### PART-A

#### Answer All the Questions (20 x 1 = 20 Marks)

1. The wave associated with particles is called------

a)De Broglie wave b) electromagnetic c) radiowaves d) infrared wave 2.  $\lambda = h/p$  is called

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a)De Broglie relation b) Einstein mass energy relationship c) Heisenberg uncertainity principle 3. The principal quantum number specifies the

a) subshell to which an electron belongs to b) orientiation of orbitals in space c)spin of an electron d)The shell to which electron belongs to

4. The azimuthal quantum number specifies

a) subshell to which an electron belongs to b) orientiation of orbitals in space c)spin of an electron d)The shell to which electron belongs to

5.In Schrodinger wave equation, the electron motion is considered to be a------

a)superimposed wave b) stationary wave c)interfringes d)polarized

6.Among the conditions which one is correct for the wavefunction w to satisfy and give meaningful solution

a)  $\psi$  must be multiple valued b)  $\psi$  must be ontinuous at all points in space c)  $\psi$  must become unity at infinity d) w should not be normalize

7. Einstein mass energy relationship is----

a)E = mc/2 b)  $E = mc^2$ c)E/2 = mc d)  $E = m^2 c$ 8.Photon-----

a) Value is 4 for N shell b) Probability density c) Always positive value d) Exhibits both momentum and wavelength

9. The number of radial and angular nodes in 3p subshells are

a)1 and 1 respectively b)1 and 2 respectively c)2 and 1 respectively d)1 only

10. Among the conditions which one is correct for the wavefunction  $\psi$  to satisfy and give meaningful solution

a)  $\psi$  must be single valued b)  $\psi$  must be discontinuous at all points in space c)  $\psi$  must become unity at infinity d) w should not be normalize 11.d - orbital have how many orientations

a)one

b)Three c)five d)seven 12. The magnetic quantum number specifies

a) subshell to which an electron belongs to b) orientiation of orbitals in space c)spin of an electron d)The shell to which electron belongs to

13. The principal quantum number 'n' can have values

a) n=0,1,2,3,...,7 b) n=1,2,3,...,n infinity c)n=0,1,2,...,n infinity d)2|+1. 14. The electron has ------ probability of being located at a node. a)0 b)1 c)2 d)3 15. The number of possible orientiations of an orbital is given by a)21 + 1b)21 - 1c) 12 + 1d)12 - 116. p - orbital have how many orientations a)one b)Three c)five d)seven 17. The shape of the s- orbital a)Spherical b)dumb bell c) doubledumb bell d)square 18. Principal quantum number represents -a) orientation of the orbital b) energy and size of orbital c) spin of electron d) shape of the orbital 19. Quantum number gives the orientiation of orbitals in space a) Principal quantum number b) Azimuthal quantum number c) magnetic quantum number d) spin quantum number 20. The azimuthal quantum number specifies a)The angular momentum of an electron b)orientiation of orbitals in space c)spin of an electron d)The shell to which electron belongs to

PART-B (3x2=6 Marks) ANSWER ALL THE QUESTIONS:

21. Draw the shapes of the orbitals. 22. What is the significance of  $\psi$  and  $\psi^2$ . 23. What do you understand by the term nodal plane.

PART-C (3x8=24 Marks) ANSWER ALL THE OUESTIONS:

24. a.What do you understand by dual character of matter? Derive the de Broglie relation. How was it verified.

(Or)b.Explain Radial and angular nodes and their significance.

25. a.Explain the terms:magnetic quantum number, spin quantum number. Explain the significance of each of these numbers.

(Or)b.Write a note on rules for filling electron and electronic configuration of given atom.

26. a.Write a note on Heisenberg uncertainity principle. (Or)

b.Explain why in the filling up of atomic orbitals 4s orbital is filled up before 3d orbital.

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### [16CHU101]

# (For the candidates admitted from 2016 & onwards) B.Sc. DEGREE EXAMINATION, August 2016. Third Semester Chemistry INTERNAL TEST - I ATOMIC STRUCTURE AND CHEMICAL BONDING

### **Answer Key**

#### PART-A

- 1. De Broglie wave
- 2. De Broglie relation
- 3. The shell to which electron belongs to
- 4. subshell to which an electron belongs to
- 5. stationary wave
- 6. ψ must becontinuous at allpoints in space
- 7. **E** =  $mc^2$
- 8. Exhibits both momentum and wavelength
- 9. 1 and 1 respectively
- 10.  $\psi$  must be single valued
- 11. **five**
- 12. orientiation of orbitals in space
- 13. **n**= **1,2,3** ..... infinity
- 14. **0**
- 15. **2l** + **1**
- 16. Three
- 17. Spherical
- 18. energy and size of orbital
- 19. magnetic quantum number
- 20. The angular momentum of an electron

### PART-B

21. Draw the shapes of the orbitals.

**S** - **Orbitals** : Orbitals with subshell quantum number 1 = 0 are called s orbitals.



**p- Orbitals:** Orbitals with subshell quantum number l = 1 are called p orbitals. A p orbital has two lobes like a dumb-bell. The diagrammatic representation of the three separate p orbitals is along the three Cartesian axes X, Y and Z. However, the three individual orbitals collectively considered as superimposed may be totally visualized as concentrically spherical around the origin of the Cartesian axes



**d- Orbitals :** Orbitals with subshell quantum number l=2 are called d orbitals. Since m can be 2, -1, 0, +1, or +2 when l is two, the d orbitals come in sets of five.

22. What is the significance of  $\psi$  and  $\psi^2$ .

The wave function  $\psi$  gives the amplitude of the wave. The intensity of wave property is known to be proportional to the square of the amplitude of the wave function. Therefore, the intensity of the electron in the atom is proportional to the square of the amplitude  $\psi$ . By intensity of the electron is meant the electron density, i.e if  $\psi 2$  is more near one point than another point in the vicinity of the atoms nucleus, it will be found more often near the point with larger  $\psi 2$  than near the point with less value of  $\psi 2$ . Thus it can be said that  $\psi$  in itself has no physical significance, and all that can be known about the internal structure of matter is through  $\psi 2$ . It helps in locating the relative probability of finding the electron. This uncertainty in finding the electron is in conformity with Heisenbergs uncertainty principle. The exact energy can be found out with the help of equation and hence there would be a great degree of uncertainty in finding the position of the electron.

### 23. What do you understand by the term nodal plane.

The orbital dx2-y2 lies along the plane XY, just as dxy orbital but is related by 450 around Z axis, so that the lobes are along X and Y axes and the nodal planes lie midway between the nodal planes of the dxy orbital. The fifth orbital,  $dz^2$  is symmetrical around Z-axis and has two cone shaped nodal surface with the angle between the elements of the cone and Z-axis approximately equal to 450.

### PART-C

24. a.What do you understand by dual character of matter? Derive the de Broglie relation. How was it verified.

This gives a relationship between the magnitude of the wave length associated with mass 'm' of a moving body and its velocity.

According to Planck, the photon energy 'E' is given by the equation

E = hv.....(i)

Where h is the Planck s constant and v the frequency of radiation. According to Einstein s massenergy relationship

 $E = mc^{2}$ .....(ii)

Where E is the energy associated with photon of mass 'm' and c is the velocity of radiation, Comparing equations (i) and (ii).

$$mc2 = hv$$
$$v=c/\lambda$$
$$mc2 = h c/\lambda$$
or mc= h/  $\lambda$ 

This equation is known as de BROGLIE 'S EQUATION

b. Explain Radial and angular nodes and their significance.

The probability of finding an electron within a small radial space around the nucleus is termed as radial probability distribution. The main feature of radial distribution plots is that they determine the spatial extent of the orbit. Consider the space around the nucleus to be divided into a large number of uniform concentric shells. In between radii r and r+dr the volume of the spherical shell would be  $4\Pi r^2 dr$ . Thus the probability of finding an electron within this spherical

shell would be  $4\Pi 2\psi r^2$ . This value is the radial probability at a distance r from the nucleus. In the function  $4\Pi r^2 dr \psi^2$ , the probability factor  $\psi^2$  decreases whereas the volume factor  $4\Pi r^2 dr$  increases with increase in the value of r. The function  $4\Pi r^2 dr \psi^2$  when plotted against the distance or from the nucleus gives the radial probability distribution of the electron.

### Significance

- 1. When r=0, the probability of finding the electron is zero, i.e at nucleus, the probability of finding an electron is zero.
- 2. For a larger value of principal quantum number n, the electron will be farther from the nucleus.
- 3. The value r0 = 0.529 of maximum radial probability in the radial distribution curve
- 4. from the nucleus has been found to be in close agreement with the Bohr's calculated value for the radius of first circular orbit.
- 5. Radial distributions overlap, revealing that the outer electrons penetrate the region occupied by inner electrons in an atom.

25. a.Explain the terms:magnetic quantum number, spin quantum number. Explain the significance of each of these numbers.

### Magnetic Quantum Number:

This quantum number was introduced to describe the Zeeman's effect i.e splitting of spectral lines under the influence of an applied magnetic field. A shell is made up of one or more subshells and each subshell is made up of one or more orbitals. The number of orbitals in a shell is n2 where n is the principal quantum number. Each orbital has a characteristic magnetic quantum number and can accommodate up to two electrons. When an atom is placed in a magnetic field (Zeeman's effect), the orbitals orient themselves relative to the magnetic field. This is due to the presence of negatively charged electrons in it. The orientation in the magnetic field is determined by the magnetic quantum number (m). Magnetic quantum number is designated as 'm' and can have values from +l to –l through zero.

Every value of m, for a given value of l, corresponds to one orbital in that subshell designated by l. Thus:

- 1. s will have one orbital as m has only one value, m = 0
- 2. p will have three orbitals as m has three values +1, 0, -1.

- 3. d will have five orbitals as m has five values +2, +1, 0, -1, -2).
- 4. f will have seven orbitals as m has seven values +3, +2, +1, 0, -1, -2, -3.
- It means that two orbitals in the same subshell can have identical n & l values but must differ in 'm' values.

Thus p subshell is made up of three identical orbitals. But in a magnetic field, the three orbitals take up different positions with respect to the lines of force of field and thus have slightly different energy levels. Therefore, the three 2p orbitals are designated as 2px, 2py and 2pz.

#### **Spin Quantum Number:**

The fact that an electron in its motion around the nucleus also rotates or spins about its own axis, gives birth to spin quantum number. This quantum number is designated as 's'. It indicates the direction in which the electron is spinning. Spin can be either clockwise or anticlockwise. Clockwise spin is represented by an arrow pointing upwards ( $\uparrow$ ), while anticlockwise spin is represented by an arrow pointing downwards ( $\downarrow$ ). The spin of electron contributes to its overall angular momentum and since it can be either clockwise or anticlockwise relative to the direction of the path of electron, it may cause angular momentum to increase or decrease.



This results in two sub levels of slightly different energy for each value of m, thus corresponding to two possibilities of s. It is assumed that the total energy difference between two sub levels is one quantum i.e. s level will have values +1/2 and -.1/2

An orbital can accommodate two electrons provided they have opposite spins. Such electrons are known as paired electrons. When an orbital has only one electron, it is known as odd or unpaired electron. The spin quantum number gives rise to magnetic properties of materials. When the electrons are paired, they have opposite spins and their magnetic moments will be cancelled. Presence of magnetic properties indicates the presence of one or more unpaired electrons.

b. Write a note on rules for filling electron and electronic configuration of given atom.

On the basis of magnetic measurements, Hund(1925) put forward an empirical rule known after his name as Hund's Rule of Maximum Multiplicity'. This states that Electrons are distributed among the orbitals of a sub shell in such a way as to give the maximum number of unpaired electrons and hence the same direction of spin'. It implies that the pairing of electrons in any orbital of a sub shell will occur only when all the available orbitals of it (sub shell) have one electron each. **Examples :** The following examples will illustrate the applications of Hund's rule.

1. Nitrogen has atomic number 7 and hence will have the following configuration. 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>3</sup>



and not as



The above configuration also explains the covalency of nitrogen which is three, as three electrons are required for complete pairing of the three electrons present in p-sub shell.

26. a. Write a note on Heisenberg uncertainity principle.

Werner Heisenberg (1927) a German physicist, stated uncertainty principle, which is the consequence of dual behaviour of matter and radiation. It states that "It is impossible to

determine simultaneously the exact position and exact velocity of an electron". This principle is an important feature of wave mechanics and discusses the relationship between a pair of conjugate properties i.e those properties that are interdependent. For example, the position and momentum of a moving particle are interdependent and thus conjugate properties. If the momentum (or velocity) is measured very accurately, the measurement of the position of the particle correspondingly becomes less precise. On the other hand, if the position is determined with accuracy, the measurement of momentum becomes less accurate. Thus, certainty of determination of one property introduces uncertainty in determination of the other. The uncertainty in measurement of position  $\Delta x$  and the uncertainty of determination of momentum  $\Delta p$  (or  $\Delta mv$ ) are related by Heisenberg's relationship as

### $\Delta x \cdot \Delta p > h/4\pi$ where h is Planck's constant

This principle applies to all bodies but becomes significant when applied to bodies with small masses and is negligible in case of large objects. The momentum  $\Delta p$  should be along the direction x for the principle to hold good. If the product of uncertainty in position  $\Delta x$  and momentum along the y direction is considered, it will be zero.

### i.e $\Delta x \cdot \Delta py = 0$

The following example illustrates the principle. If one is asked to measure the thickness of a sheet of paper with an unmarked metrestick, the result obtained would obviously be inaccurate and meaningless. In order to obtain accuracy, one would have to use an instrument graduated in units smaller than the thickness of the sheet. A substance can be seen only if it could reflect light or any other radiation from its surface. Since the size of electron is too small, its position may be determined by employing light of very small wavelength such as x-rays or  $\gamma$  rays. A photon of radiation of small wavelength has large energy and also momentum. As one such photon strikes an electron, it immediately changes the momentum of the electron. Now as position of the electron is being determined, the momentum gets changed. Thus it is impossible to determine the exact position of an electron which is moving with a definite velocity.



When photons from sources collide with the electrons, some of the electrons bounce into the microscope and enable the observer to see the flash of light and thus to measure both position and momentum of the electron at a particular instant of time. The position of the electron is determined using the following equation of optics, where the resolving power of microscope is given by

$$\Delta x = \frac{\lambda}{2\sin\theta}$$

 $\Delta x$ - is minimum distance between two points which can be distinguished as separate. Radiations of short wavelength (x-rays or gamma ray) are preferred to make  $\Delta x$  value very small. Thus  $\Delta x$  is the error or uncertainty in measurement or the position of electron.

 $\lambda$  - wavelength of photon

 $\theta$  - semivertical angle of the cone light coming from illuminated electron.

b. Explain why in the filling up of atomic orbitals 4s orbital is filled up before 3d orbital.

The three orbitals  $p_x$ ,  $p_y$  and  $p_z$  have equal energy but are concentrated in different regions of space at right angles to each other. Electrons can minimize the repulsive forces between themselves by occupying different orbitals and having parallel spins.

This explains their preference to enter different orbitals as long as possible.

Cr (At no = 24)  $1s^2 2s^2 2p^6 3s^2 3p^6 4d^5 4s^1$ Mn (At no = 25)  $1s^2 2s^2 2p^6 3s^2 3p^6 4d^5 4s^2$ 

The expected electronic configuration of copper (atomic no. 29) is  $C_u At no = 29 \ 1s^2 \ 2s^2 \ 2p^6 \ 3s^2 \ 3p^6 \ 3d^9 \ 4s^2$ 

However it is observed that preferred configuration is 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup> 3s<sup>2</sup> 3p<sup>6</sup> 3d<sup>10</sup> 4s<sup>1</sup>

The stability of half filled and completely filled orbitals may be explained in terms of lowering of energy due to the exchange energy for a set of two electrons with parallel spins.

According to Hund's rule, electrons avoid entering the same orbital in the given sub shell as far as possible. This reduces coulombic repulsion by enlarging the distance between electrons. Also, the electrons entering the different orbitals of a sub shell have parallel spins. This keeps them further apart and lowers the energy through electron exchange and resonance.

[17CHU101] (For the candidates admitted from 2017 & onwards) B.Sc. DEGREE EXAMINATION, August 2017 First Semester Chemistry INTERNAL TEST - II ATOMIC STRUCTURE AND CHEMICAL BONDING Maximum: 50 marks

Reg. No. : -----

No of Copies : 110

Time: 2 Hours

Answer All the Questions

PART-A  $(20 \times 1 = 20 \text{ Marks})$ 1. Greater the bond order, greater is a)bond dissociation energy b) covalent nature c)bond length d)paramagnetism 2. Which of the following is most polar a)C-O b)C-F c)O-F d)N-F 3. Which of the following is least ionic a)AgC1 b) KCl c)BaCl2 d)COCl2 4. Which of the following compound has least dipole moment a)NH<sub>3</sub>b)NF<sub>3</sub> c)SO<sub>2</sub> d)H<sub>2</sub>O 5. Which one of the following has least polarity a)H-F b)H-Cl c) H-O d)H-S 6. BF<sub>3</sub> involves sp<sup>2</sup> hybridisation and the shape of the molecule is-----a)tetrahedralb)square planar c)triangular planar d)linear 7. The magnitude of the lattice energy of a solid increases if a)the ions are large b)the ions are small c)the ions are of equal size d)charges on the ions are small 8. Which of the following bonds has the highest bond dissociation energy a)CH<sub>3</sub>-H b)CH<sub>3</sub>-Br c)CH<sub>3</sub>-Cl d)CH<sub>3</sub>-F 9. Whice of the following forces between atoms or ions or molecules is the strongest a)ionic-bond b)ion dipole c)dipole-dipole d)london-dispersive 10. The molecule which have zero dipole moment a)CH2Cl2 b)BF3 c)ClO2 d)NF3 11. The shape of a molecule which has three bonds and one lone pair is a)Octahedral b)Pyramidal c)Triangular planar d)Tetrahedral 12.Bonded electron pairs present in octahedral SF6 molecule a)3 b) 4 c)6 d)8 13.Decreasing order of size of various hybrid orbitals is

a)sp>sp<sup>2</sup>>sp<sup>3</sup> b)sp<sup>3</sup>>sp<sup>2</sup>>sp c)sp<sup>2</sup>>sp>sp<sup>3</sup> d)sp>sp<sup>3</sup>>sp<sup>2</sup> 14. Resonance energy is equal to actual bond energy minus energy of the pure -----a)ionic bond b)Covalent bond c)Coordinate bond d)isomorphous compound 15. Bond order in a molecule of NO is ----

a)1 b)2.5 c)3 d) 3.5 16.Which of the following compound has not  $sp^3$  hybridisation.

......

18. The weakest among the following is
a) ionic bond b) covalent bond c) metallic bond d) vanderwaal's forces
19. Directed bond in water form an angle of ------a) 90°C
b) 120°C
c) 105°C
d) 60°C
20. Which one of the following is strongest bond
a) Cl-F b) F-F c) Br-F d) Br=Cl

PART-B (3x2=6 Marks) ANSWER ALL THE QUESTIONS:

21. What is an ionic bond.22. What do you understand by the term resonance energy?23. What is meant by hybridization.

PART-C (3x8=24 Marks) ANSWER ALL THE QUESTIONS:

24. a.Give an account on solvation energy and importance in context of stability and solubility of ionic compounds.

b.Describe Born-Haber cycle for calculating the lattice energy of sodium chloride.

25. a.With suitable examples explain the hybridization and geometry of molecules in octahedral complexes.

b. Derive the Born-Lande equation.

26. a. Discuss the Fajan rules.

Or

b. a.Explain the hybridization and geometry of molecules in tetrahedral complexes by giving suitable examples.

Reg. No. : -----

[16CHU101]

### (For the candidates admitted from 2016 & onwards) B.Sc. DEGREE EXAMINATION, September 2016.

### Chemistry INTERNAL TEST - II ATOMIC STRUCTURE AND CHEMICAL BONDING

**Answer Key** 

Time: 2 Hours

Maximum: 50 marks

### PART- A

- 1. bond dissociation energy
- 2. **C-F**
- 3. COCl<sub>2</sub>
- 4. NF<sub>3</sub>
- 5. **H-S**
- 6. triangular planar
- 7. the ions are small
- 8. CH<sub>3</sub>-F
- 9. ionic-bond
- 10. **BF**<sub>3</sub>
- 11. Pyramidal
- 12. 6
- 13. **sp<sup>3</sup>>sp<sup>2</sup>>sp**
- 14. Covalent bond
- 15.1
- 16. SF<sub>4</sub>
- 17. **sp**<sup>2</sup>
- 18. vanderwaal's forces
- **19. 105°C**
- 20. Br-F

### PART-B

21. What is an ionic bond.

Ionic compounds have ionic bonds which are electrostatic forces of attraction, they can decompose into respective ions. Ionic compounds are ionic in nature and good conductor of electricity. They are usually found in solid state like sodium chloride etc. In ionic compounds, the constituent atoms or ions show strong attractions with other ions. Ionic compounds generally have high melting points, boiling points and found in solid state. Due to presence of ions, ionic compounds are polar in nature and easily soluble in polar solvents like water.

For example, common table salt is sodium chloride. When sodium (Na) and chlorine (Cl) are combined, the sodium atoms each lose anelectron, forming cations (Na+), and the chlorine atoms each gain an electron to form anions (Cl–). These ions are then attracted to each other in a 1:1 ratio to form sodium chloride (NaCl).

$$Na + Cl \rightarrow Na^+ + Cl^- \rightarrow NaCl$$

22. What do you understand by the term resonance energy?

Resonance is a mental exercise and method within the Valence Bond Theory of bonding that describes the delocalization of electrons within molecules. It compares and contrasts two or more possible Lewis structures that can represent a particular molecule. Resonance structures are used when one Lewis structure for a single molecule cannot fully describe the bonding that takes place between neighboring atoms relative to the empirical data for the actual bond lengths between those atoms. The net sum of valid resonance structures is defined as a resonance hybrid, which represents the overall delocalization of electrons within the molecule. A molecule that has several resonance structures is more stable than one with fewer.

23. What is meant by hybridization.

Hybridization is a simple model that deals with mixing orbitals to from new, hybridized, orbitals. This is part of the valence bond theory and helps explain bonds formed, the length of bonds, and bond energies; however, this does not explain molecular geometry very well.

### PART-C

24. a.Give an account on solvation energy and importance in context of stability and solubility of ionic compounds.

The role of the two quantities is discussed as under:

- 1. Greater is the lattice energy, greater the stability of the ionic solid. Because more energy will be needed to pull apart the positive and negative ions.
- 2. Salts with high lattice energy required a greater input of thermal energy to break down the crystal lattice. Consequently, salts have high melting points.
- 3. The magnitude of lattice energy of the solid may give us an idea about the solubility of that substance in different solvents. Ionic solids in general are insoluble in non polar solvents such as CCl4, where as these are soluble in polar solvents, water.
- 4. For a solid to dissolve in water, the strong force of attraction between its ions must be over come. This energy can be over come by ion-solvent interaction. The solvation of ions is reffered to interms of solvation energy, which is always negative, i.e, energy is released in the process.
- 5. The solvation energy increases if the solvent has high dipole moment/high polarizability.
- 6. The solid that dissolves in water its hydration energy is greater than the lattice energy of that solid.

Hydration energy > Lattice energy Hydration energy < Lattice energy b. Describe Born-Haber cycle for calculating the lattice energy of sodium chloride.

Born Haber cycle is a simplified method developed by Max Born and Fritz Haber in 1919 to correlate the lattice energies of ionic solids to other thermodynamic data. Lattice enthalpy or lattice energy is defined as enthalpy change which occurs when one mole of ionic solid is formed by close packing of constituent ions in gaseous state. Or it may be defined as energy released when one mole of ionic solid dissociates into its gaseous state.

**For example,** the formation of sodium chloride from sodium and chloride ion involves 788 kj/mol of energy and dissociation of sodium chloride into it constituent ions requires same amount f energy.

NaCl(s)  $\rightarrow$  Na+(g) + Cl- (g)  $\triangle$ LH° = 788 kj/mol Na+(g) + Cl-(g)  $\rightarrow$  NaCl(s)  $\triangle$ LH° = -788 kj/mol

When oppositely charged ions interact to form ionic solid, a large amount of energy is released and dissociation of crystal also require a lot of energy which makes melting point and boiling points of crystal very high. Born Haber cycle is used to determine this lattice energy by using other energy values like ionization energy, electron affinity, dissociation energy, sublimation energy and heat of formation.

**1.Ionization Energy:** It is the energy required to remove an electron from a neutral gaseous atom or an ion.

**2.Electron Affinity:** It is the energy released when an electron is added to an isolated neutral gaseous atom or an ion.

**3.Dissociation energy:** The energy required to dissociate a compound is called as dissociation energy. Dissociation of a compound is always an endothermic process and requires an input of energy.

**4.Sublimation energy:** The energy required to change the phase from solid to gas, by passing the liquid phase is called as sublimation energy.

**5.Heat of formation:** The energy change during the formation of a compound from its elements is known as heat of formation.

Born Haber cycle is based on Hess law which states that the standard enthalpy change of reactions is the sum of algebraic sum of the standard enthalpies of reactions into which the

overall reaction may be splitted or divided.

**For example,** the formation of C from A and B can take place in two ways. Both ways involve same initial and final states, but one is single step and other one is multi-step reaction which involves intermediates, D and E. Both ways has an enthalpy change of 160 kJ.

25. a.With suitable examples explain the hybridization and geometry of molecules in octahedral complexes.

The basic principles, which are involved in the valence bond treatment of coordination compounds are:

(a) Hybridization of valance orbitals of the central metal/ ion

(b) Bonding between ligand and the metal ion/atom.

(c) Relation between the type of bond and the observed magnetic behavior

Let us explain by taking simple examples such as  $[CoF_6]^{3-}$  and  $[Co(NH_3)_6]^{3+}$ . Although in both the complexes, the oxidation state of cobalt is +3, but  $[CoF_6]^{3-}$  is paramagnetic and  $[Co(NH_3)_6]^{3+}$  is diamagnetic, why? The formation of a complex may be considered as a series of hypothetical steps. First the appropriate metal ion is taken e.g.  $Co^{3+}$ . Cobalt atom has the outer electronic configuration  $3d^74s^2$ . Thus Co3+ ion will have the configuration  $3d^6$  and the electrons will be arranged as:



 $\operatorname{Co}^{+3}$  ion forms both paramagnetic (outer orbital) and diamagnetic (inner orbital) complexes depending upon the nature of ligands as illustrated below. As  $\operatorname{Co}^{3+}$  ion combines with six fluoride ligands in  $[\operatorname{CoF}_6]^{3-}$ , empty atomic orbitals are required on the metal ion to receive the coordinated lone pair of electrons. The orbitals used are one 4*s*, three 4*p* and two 4*d*. These are hybridized to give a set of six equivalent sp<sup>3</sup>d<sup>2</sup> hybrid orbitals. A ligand orbital containing a lone pair of electron forms a coordinate bond by overlapping with an empty hybrid orbital on the

metal ion. In this way a bond is formed with each ligand. The *d*-orbitals used are the 4 2 2 xy d and 4 2zd. It is shown below:



Since the outer 4d orbitals are used for bonding, this is called an outer orbital complex. The energy of these orbitals is quite high, so the complex will be reactive. This complex will be high-spin paramagnetic, because it has four unpaired electrons. An alternative octahedral arrangement in  $[Co(NH_3)_6]^{3+}$  is possible when the electrons on metal ion are rearranged as shown below:



Since inner *d*-orbitals are used this is called an inner orbital complex. There is no unpaired electron, the complex will be low-spin diamagnetic.

#### b. Derive the Born-Lande equation

The Born-Landé equation is a concept originally formulated in 1918 by the scientists Born and Lande and is used to calculate the lattice energy (measure of the strength of bonds) of a compound. This expression takes into account both the Born interactions as well as the Coulomb attractions.

Due to its high simplicity and ease, the Born-Landé equation is commonly used by chemists when solving for lattice energy. This equation proposed by Max Born and Alfred Landé states that lattice energy can be derived from ionic lattice based on electrostatic potential and the potential energy due to repulsion. To solve for the Born-Landé equation, you must have a basic understanding of lattice energy:

• Lattice energy decreases as you go down a group (as atomic radii goes up, lattice energy goes

down).

• Going across the periodic table, atomic radii decreases, therefore lattice energy increases.

The Born-Landé equation was derived from these two following equations. the first is the electrostatic potential energy with

 $\Delta U = -LA|Z + ||Z - |e24\pi \epsilon or$ 

- L is Avogadro's constant (6.022×1023)
- A is the Madelung Constant (a constant that varies for different structures)
- e is the charge of an electron  $(1.6022 \times 10 19 \text{ C})$
- Z+ is the cation charge Z- is the anion charge
- $\epsilon$ o is the permittivity of free space
- The second equation is the repulsive interaction:

 $\Delta U=LBrn$  with

• B is the repulsion coefficient and

• N is the Born Exponent (typically ranges between 5-12) that is used to measure how much a solid compresses

These equations combine to form:  $\Delta U(0K)=LA|Z+||Z-|e24\pi\epsilon oro(1-1n)$  with

• r<sub>0</sub> = closest ion distance

26. a. Discuss the Fajan rules.

Rules formulated by Kazimierz Fajans in 1923, can be used to predict whether a chemical bond is expected to be predominantly ionic or covalent, and depend on the relative charges and sizes of the cation and anion. If two oppositely charged ions are brought together, the nature of the bond between them depends upon the effect of one ion on the other.



Although the bond in a compound like X+Y- may be considered to be 100% ionic, it will always have some degree of covalent character. When two oppositely charged ions (X+ and Y-)

approach each other, the cation attracts electrons in the outermost shell of the anion but repels the positively charged nucleus. This results in a distortion, deformation or polarization of the anion. If the degree of polarization is quite small, an ionic bond is formed, while if the degree of polarization is large, a covalent bond results.

The ability of a cation to distort an anion is known as its polarization power **and the tendency of the anion to become polarized by the cation is known as its** polarizability. The polarizing power and polarizability that enhances the formation of covalent bonds is favoured by the following factors:

**Small cation:** the high polarizing power stems from the greater concentration of positive charge on a small area. This explains why LiBr is more covalent than KBr (Li+ 90 pm cf. K+ 152 pm). **Large anion:** the high polarizability stems from the larger size where the outer electrons are more loosely held and can be more easily distorted by the cation. This explains why for the common halides, iodides, are the most covalent in nature (I- 206 pm).

**Large charges:** as the charge on an ion increases, the electrostatic attractions of the cation for the outer electrons of the anion increases, resulting in the degree of covalent bond formation increasing.

Large cations are to be found on the bottom left of the periodic table and small anions on the top right. The greater the positive charge, the smaller the cation becomes and the *ionic potential* is a measure of the charge to radius ratio.

### **Electronic configuration of the cation:**

Two cations of the same size and charge, the one with a pseudo noble-gas configuration (with 18 electrons in the outer-most shell) will be more polarizing than that with a noble gas configuration (with 8 electrons in the outermost shell). Thus zinc (II) chloride (Zn(II) 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup> 3s<sup>2</sup> 3p<sup>6</sup> 3d<sup>10</sup> and Cl- 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup> 3s<sup>2</sup> 3p<sup>6</sup> ) is more covalent than magnesium chloride (Mg(II) 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup>) despite the Zn2+ ion (74 pm) and Mg2+ ion (72 pm) having similar sizes and charges. From an MO perspective, the orbital overlap disperses the charge on each ion and so weakens the electrovalent forces throughout the solid, this can be used to explain the trend seen for the melting points of lithium halides.

LiF = 870 °C, LiCl = 613 °C, LiBr = 547 °C, LiI = 446 °C

b. a. Explain the hybridization and geometry of molecules in tetrahedral complexes by giving suitable examples.

- Coordination compounds contain metal ions, in which ligands form covalent-coordinate bonds to the metal.
- Ligands must have a lone pair of electrons.
- Metal should have an empty orbital of suitable energy available for bonding.
- Atomic orbitals are used for bonding (rather than molecular orbitals)
- This theory is useful to predict the shape and stability of the metal complexes.

### Limitations:

(1) Does not explain why some complexes are colored and others are not

(2) Does not explain the temp. Dependence of the magnetic properties

The metal ion can also form 4-coordinate complexes. For such complexes two different arrangements are possible i.e. tetrahedral  $(sp^3)$ 



 $sp^3$  hybridization for Ni<sup>2+</sup>(to afford a tetrahedral geometry).

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[17CHU101]

KARPAGAM UNIVERSITY, (Under Section 3 of UGC Act 1956) COIMBATORE-641 021 (For the candidates admitted from 2017& onwards)

B.Sc. DEGREE EXAMINATION, September 2017.

First Semester

Chemistry

Internal Test-III

ATOMIC STRUCTURE AND CHEMICAL BONDING Time: 2 Hours Maximum: 50 Marks

Date :

#### PART-A (20 x 1 = 20 Marks) Answer All the Questions

1. When the hybridisation state of carbon atom changes from  ${\rm sp}^3\;$  to  $\,{\rm sp}^2\;$  to sp , the angle between the hybridised orbitals

a)decreases gradually b)increases gradually c)decreases with increasing d)decreases considerably

2.According to VSEPR theory the shape of water molecule is

a)octahedral b)distorted tetrahedral c)planar triangle d)linear

3.HCl involves

a)s-s overlapping b)sp-overlapping c)p-p overlapping d)P $\pi$ -P $\pi$  overlapping 4. Which species has lone pair of electrons on central atom

a)CCl<sub>4</sub> b)CH<sub>4</sub> c)NH<sup>4+</sup> d)H<sub>2</sub>O

5. Which atomic orbital is always involved in sigma bonding only

a)s b)p c)d d)f

6.In ------ the atoms retain their individuality in the molecule.a) Valence bond theory b)Molecular orbital theory c)Crystal field theory d)coordinate compounds

7. Resonance energy is equal to actual bond energy minus energy of the pure -----a)ionic bond b)Covalent bond c)Coordinate bond d)isomorphous compound 8. The molecule having bond order 3 is a) $H_2$  b) $N_2$  c) $O_2$  d)  $He^{2+}$ 9. The number of antibonding electron pairs in  $O_2^{2-}$  molecular ion on the basis of MOT is (Atomic number of O=8) a)4 b)3 c)2 d)5 10.O<sub>2</sub> molecule is a)paramagnetic b)diamagnetic c)ferromagnetic d)ferromagnetic 11. Which one of the following molecule is paramagnetic a)CO<sub>2</sub> b)SO<sub>2</sub> c)NO d)H<sub>2</sub>O 12. Lateral overlapping is expected in a)sigma bonds b)pi bonds c)metallic bonds d)coordinate bonds 13. The shape of the sulphate ion is a)tetrahedral b)square planar c)trigonalbipyramidal d)hexagonal 14.In a double bond connecting two atoms, there is a sharing of a)two electrons b)four electrons c)one electrons d)all electrons 15. Which of the following compound has not sp<sup>3</sup> hybridisation. a)H<sub>2</sub>O b)NH<sub>3</sub> c)CH<sub>4</sub> d)SF<sub>4</sub> 16. Hybridisation involves a)addition of an electron pair b)mixing up of atomic orbitals c)separation of orbitals d)removal of electron pair 17.A pi bond is formed by sideway overlapping of a)s-s orbitals b)p-p orbitals c)s-p orbitals d)s-p-s orbitals 18.Sigma bond is formed by a)orbitals b)p-p orbitals c)end to end overlapping of orbitals d) shortens the internuclear distance 19.Strongest bond is formed by the overlapping of a)2s and 2p orbitals b)2p and 2p orbitals c)2s and 2s orbitals d)1s orbital 20.Greater than the bonding M.O a)Pi bond b)increases bond length c)decrease s bond length d)Distorts the geometry of molecule

Reg. No. : -----

[16CHU101]

#### (For the candidates admitted from 2016 & onwards)

### **B.Sc. DEGREE EXAMINATION, October 2016.**

**First Semester** 

Chemistry

**Internal Test III** 

#### ATOMIC STRUCTURE AND CHEMICAL BONDING

### **Answer Key**

21.Define node.

Phases are designated either (+) or (-) relative to their wave "up" or wave "down" displacements. A node occurs if the phase signs change from (+) to (-) or vice versa. It is important to notice that the phase signs do NOT symbolize charges. Nodes are regions where the probability of findingan electron is ZERO.

22. What do you understand by anomalous electronic configurations.

Two ions of the same size and charge, one with a pseudo noble gas configuration (i.e. 18 electrons in the outermost shell) will be more polarizing than a cation with noble gas configuration (i.e., 8 electron in outer most shell).

23.What is dipole moment.

The percentage of ionic character in a compound can be estimated from dipole moments. The bond dipole moment uses the idea of electric dipole moment to measure the polarity of achemical bond within a molecule. It occurs whenever there is a separation of positive and negative charges.

24. What shapes are associated with the following hybrid orbitals  $sp^2,\,sp^3,\,sp^3d,sp^3d^2$ 

1. SP hybridization

Example CO<sub>2</sub>



2. SP<sup>3</sup>hybridization

Example CH4



3.  $sp^3d^2hybridization$ Example  $SF_6$ 



25. What is LCAO molecular orbital approach.

Linear combinations of atomic orbitals (LCAO) can be used toestimate the molecular orbitals that are formed upon bonding between the molecule's constituentatoms. Similar to an atomic orbital, a Schrödinger equation, which describes the behavior of anelectron, can be constructed for a molecular orbital as well. Linear combinations of atomicorbitals, or the sums and differences of the atomic wavefunctions, provide approximate solutions to the Hartree–Fock equations which correspond to the independent-particle approximation of the molecular Schrödinger equation.

#### Part C

26.a. Explain Heisenberg uncertainity principle.Show that this principle is valid only for small particles and not for large objects.

Werner Heisenberg (1927) a German physicist, stated uncertainty principle, which is the consequence of dual behaviour of matter and radiation. It states that It is impossible to determine simultaneously the exact position and exact velocity of an electron. This principle isan important feature of wave mechanics and discusses the relationship between a pair of conjugate properties i.e those properties that are interdependent. For example, the position and momentum of a moving particle are interdependent and thus conjugate properties. If themomentum (or velocity) is measured very accurately, the measurement of the position of the particle correspondingly becomes less precise. On the other hand, if the position is determined with accuracy, the measurement of momentum becomes less accurate. Thus, certainty of determination of one property introduces uncertainty in determination of the other.

The uncertainty in measurement of position  $\Delta x$  and the uncertainty of determination of momentum  $\Delta p$  (or  $\Delta mv$ ) are related by Heisenbergs relationship as  $4\pi\Delta x.\Delta p > h$  where h is Plancks constant. This principle applies to all bodies but becomes significant when applied to bodies with smallmasses and is negligible in case of large objects. The momentum  $\Delta p$  should be along the

direction x for the principle to hold good. If the productof uncertainty in position  $\Delta x$  and momentum along the y direction is considered, it will be zero.

### i.e $\Delta x \cdot \Delta py = 0$

The following example illustrates the principle. If one is asked to measure the thickness of asheet of paper with an unmarked metrestick, the result obtained would obviously be inaccurate and meaningless. In order to obtain accuracy, one would have to use an instrument graduated inunits smaller than the thickness of the sheet. A substance can be seen only if it could reflect lightor any other radiation from its surface. Since the size of electron is too small, its position may be employing light of very small wavelength such as x-rays or  $\gamma$  rays. A photon of small wavelength has large energy and also momentum. As one such photon strikes an electron, it immediately changes the momentum of the electron. Now as position of the electron is being determined, the momentum gets changed. Thus it is impossible to determine exact position of an electron which is moving with a definite velocity.



When photons from sources collide with the electrons, some of the electrons bounce into themicroscope and enable the observer to see the flash of light and thus to measure both positionand momentum of the electron at a particular instant of time. The position of the electron isdetermined using the following equation of optics, where the resolving power of microscope

$$\Delta x = \frac{\lambda}{2\sin\theta}$$

 $\Delta x$ - is minimum distance between two points which can be distinguished as separate. Radiations of short wavelength (x-rays or gamma ray) are preferred to make  $\Delta x$  value very small. Thus  $\Delta x$  is the error or uncertainty in measurement or the position of electron.

 $\lambda$  - wavelength of photon

 $\theta$  - Semivertical angle of the cone light coming from illuminated electron.

b.Derive the time independent Schrodinger equation.

The Schrodinger equation has two forms, one in which time explicitly appears, and sodescribes how the wave function of a particle will evolve in time. In general, the wave functionbehaves like a wave, and so the equation is often referred to as the time- dependent Schrödingerwave equation. In the other equation time dependence has been removed and it is known as thetime- independent Schrödinger wave equation. These are not two separate, independencequations. The time- independent equation can be derived readily from the time-dependent equation.

The time independent equation is written as

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

Here  $\hat{H}$  is the energy operator known as Hamiltonian,  $\Psi$  is the wave function of the system and Eis the total energy of the system

$$\hat{H} \Psi = \check{T} + V$$

Hamiltonian operator is the sum of kinetic energy operator ( $\check{T}$ ) and potential energy operator (V)

Therefore, Schrodinger equation can be written as

$$(\check{T} + V) \Psi = E \Psi$$

By writing the explicit mathematical forms of  $\check{T}$  and V the above equation can be solved toget E and  $\Psi$  for the system.For any system, the quantum mechanical study consists of following three steps;

a. writing Schrödinger equation for the system

b. solving Schrödinger equation to get meaningful solutions of the wave functions and corresponding energies.

c. calculation of all the observable properties of the system.

The results obtained by solving the Schrodinger equation for a system are in agreement with the experimental findings.

- 1. The Schrödingers wave equation is a second degree differential equation. It has several solutions some of which are imaginary and are not valid. If the potential energy term is known, the total energy 'E' and corresponding wave function  $\psi$  can be evaluated.
- 2.  $\psi$  is called the probability amplitude and is known as the wave function. It describes the behavior of a particle as a function of x and t. This concept is known as Born interpretation of
- 3.  $\psi$  and refers to a system in which the probability of finding the particle at any point is independent of time. Such a state is called stationary state.

- 4. Inspite of  $\psi$  being complex,  $\psi$  can always be chosen to be real for a 1-D bound state.
- 5. At infinity the value of  $\psi$  is zero. It means that at infinity the electron is not influenced by theattractive force of the nucleus.
- 6.  $\psi$  has only one finite value at any particular point. It means that  $\psi$  is single-valued.
- 7. For any value of potential energy, there will be many solutions (frequently an infinite number) each with a corresponding energy E. But very few solutions describe the known behaviour of electrons. The permitted solutions for ψ are called eigen functions and corresponding values of E are called eigen values.

27.a.Explain the terms.Principal quantum number, angular momentum number.Explain the significance of each of these numbers.

The Bohr model was a one-dimensional model that used one quantum number to describe the distribution of electrons in the atom. The only information obtained was the size of the orbit, which was described by principal quantum number 'n'. This could hardly explain the hydrogenspectrum adequately and the spectra of other elements that are quite complex also remained unexplained by this concept. Many single lines of spectra are found to consist of a number of closely related lines when studied with the help of sophisticated instruments of high resolvingpower. Spectral lines split up when the source of radiation is placed in a magnetic field (Zeemaneffect) or in an electric field (Stark effect). Schrödingers model allowed the electron to occupythree dimensional space. It, therefore, required three coordinates or three QUANTUMNUMBERS, to describe the orbitals in which the electrons can be found. The three coordinatesthat come from Schrödingers wave equation are the principal (n), angular (l) and magnetic (m)quantum numbers. In all, there are four identification numbers called quantum numbers whichfully describe an electron in an atom. The word quantum is used to signify that all the energylevels which are available to an electron are governed by the laws of quantum mechanics.

**1. Principal Quantum Number:** This quantum number denotes the principal shell to which the electron belongs. This is also referred to as major energy level. This number is designated as 'n' and is identical with Bohr's integers to describe stationary states in an atom. Thus it gives the number of principal shell in which the electron revolves around the nucleus. It designates the

average distance of the electron from the nucleus hence specifies the location. As the value of n increases, the energy levels get further away from the nucleus. The value of E increases or becomes less negative

$$En = \frac{-313.3}{n^2} k \ cals$$

 $\gamma = 0.529 \ 2 \ 0$  where 'n is the principal quantum number.

The value of energy of orbit increases with increase in value of 'n' and becomes zero at  $n=\infty$ . Beyond this, the energy is no longer quantized and the electron is not bound to the nucleus and can have any arbitrary amount of kinetic energy. The spectrum in this region does not containlines but is continuous and is called continuum. The difference in energy between En-1 and En- $\alpha$  is the energy needed to ionize the atom and is called the ionization energy. The principal quantum number 'n' can have non- zero positive integral values from 1 to  $\alpha$ . The quantum number 'n' can have non- zero positive integral values from 1 to  $\alpha$ . The quantum number n may theoretically assume any integral value from 1 to  $\alpha$ , only values from 1 to 7 have so far been established for atoms of known elements in their ground states. As the first principal level is nearest to the nucleus, it binds strongly to the nucleus. The next energy level isfarther from the nucleus and hence binds to the nucleus with a lesser force and so on. Therefore, the principal quantum number signifies:

a) The size of the electron orbit i.e distance between the nucleus and the electron.

b) The binding force between the nucleus and the electron.

b.Write short notes on concept of exchange energy.

Resonance is a mental exercise and method within the Valence Bond Theory of bonding that describes the delocalization of electrons within molecules. It compares and contrasts two or morepossible Lewis structures that can represent a particular molecule. Resonance structures areused when one Lewis structure for a single molecule cannot fully describe the bonding that takes place between neighboring atoms relative to the empirical data for the actual bond lengths between those atoms. The net sum of valid resonance structures is defined as a resonance
hybrid, which represents the overall delocalization of electrons within the molecule. A molecule that hasseveral resonance structures is more stable than one with fewer. Some resonance structures aremore favorable than others.

Electrons have no fixed position in atoms, compounds and molecules but have probabilities ofbeing found in certain spaces (orbitals). Resonance forms illustrate areas of higher probabilities(electron densities). This is like holding your hat in either your right hand or your left. The termResonance is applied when there are two or more possibilities available. Resonance structures not change the relative positions of the atoms like your arms in the metaphor. The skeleton of the Lewis Structure remains the same, only the electron locations change.

### **Delocalization and Resonance Structures Rules**

In resonance structures, the electrons are able to move to help stabilize the molecule. This movement of the electrons is called delocalization.

1. Resonance structures should have the same number of electrons, do not add or subtract any electrons.

2. All resonance structures must follow the rules of writing Lewis Structures.

3. The hybridization of the structure must stay the same.

4. The skeleton of the structure can not be changed (only the electrons move).

5. Resonance structures must also have the same amount of lone pairs.

### **Example : Carbonate Ion**

Consider the carbonate ion:  $CO_3^{2-}$ 

$$\begin{bmatrix} \ddot{c} & = c \\ \vdots & = \ddot{c} \\ \vdots & \vdots \\ \vdots & \vdots \\ \vdots & \vdots \end{bmatrix}^{2}$$

Step 1: Draw the Lewis Structure & Resonance.



Step 2: Combine the resonance structures by adding (dotted) bonds where other resonance bonds can be formed.



Step 3: Add only the lone pairs found on ALL resonance structures.



28.a.What do you understand by the terms polarizing power and polarizability.How do these determine ionic character of a compound.

The polarizing power and polarizability that enhances the formation of covalent bonds is favoured by the following factors:

**Small cation:** the high polarizing power stems from the greater concentration of positive chargeon a small area. This explains why LiBr is more covalent than KBr (Li+ 90 pm cf. K+ 152 pm).

**Large anion:** the high polarizability stems from the larger size where the outer electrons aremore loosely held and can be more easily distorted by the cation. This explains why for the common halides, iodides, are the most covalent in nature (I- 206 pm).

**Large charges:** as the charge on an ion increases, the electrostatic attractions of the cation for the outer electrons of the anion increases, resulting in the degree of covalent bond formation increasing.

Large cations are to be found on the bottom left of the periodic table and small anions on the top right. The greater the positive charge, the smaller the cation becomes and the *ionic potential* is a measure of the charge to radius ratio.

Polarization will be increased by:

- High charge and small size of the cation
- Ionic potential Å Z+/r+ (= polarizing power)
- High charge and large size of the anion

The polarizability of an anion is related to the deformability of its electron cloud (i.e. its

"softness")An incomplete valence shell electron configuration

Noble gas configuration of the cation produces better shielding and less polarizing power e.g. Hg2+ (r+ = 102 pm) is more polarizing than Ca2+ (r+ = 100 pm



On the left, the cation charge increases (size decreases) and on the right, the anion size increases, both variations leading to an increase in the covalency. Thus covalency increases in the order:

$$[Na^+ Cl^-, NaCl] < [Mg^{2+} 2(Cl)^-, MgCl_2] < [Al^{3+} 3(Cl)^-, AlCl_3] and [Al^{3+} 3(F)^-, AlF_3] < [Al^{3+} 3(Cl)^-, AlCl_3] < [Al^{3+} 3(Br)^-, AlBr_3]$$

b.DescribeBorn-Haber cycle for calculating the lattice energy of sodium chloride.

Born Haber cycle is a simplified method developed by Max Born and Fritz Haber in 1919 tocorrelate the lattice energies of ionic solids to other thermodynamic data. Lattice enthalpy orlattice energy is defined as enthalpy change which occurs when one mole of ionic solid is formedby close packing of constituent ions in gaseous state. Or it may be defined as energy released when one mole of ionic solid dissociates into its gaseous state. **For example,** the formation of sodium chloride from sodium and chloride ion involves 788 kj/mol of energy and dissociation of sodium chloride into it constituent ions requires same amount f energy.

$$NaCl(s) \rightarrow Na+(g) + Cl-(g) \Delta LH^{\circ} = 788 \text{ kj/mol}$$

 $Na+(g) + Cl-(g) \rightarrow NaCl(s) \Delta LH^{\circ} = -788 \text{ kj/mol}$ 

When oppositely charged ions interact to form ionic solid, a large amount of energy is released and dissociation of crystal also require a lot of energy which makes melting point and boilingpoints of crystal very high. Born Haber cycle is used to determine this lattice energy by usingother energy values like ionization energy, electron affinity, dissociation energy, sublimationenergy and heat of formation.

**1.Ionization Energy:** It is the energy required to remove an electron from a neutral gaseous atom or an ion.

**2.Electron Affinity:** It is the energy released when an electron is added to an isolated neutral gaseous atom or an ion.

**3.Dissociation energy:** The energy required to dissociate a compound is called as dissociation energy. Dissociation of a compound is always an endothermic process and requires an input of energy.

**4.Sublimation energy:** The energy required to change the phase from solid to gas, by passing the liquid phase is called as sublimation energy.

**5.Heat of formation:** The energy change during the formation of a compound from its elements is known as heat of formation.

Born Haber cycle is based on Hess law which states that the standard enthalpy change of reactions is the sum of algebraic sum of the standard enthalpies of reactions into which the overall reaction may be splitted or divided.

**For example,** the formation of C from A and B can take place in two ways. Both ways involvesame initial and final states, but one is single step and other one is multi-step reaction whichinvolves intermediates, D and E. Both ways has an enthalpy change of 160 kJ.



 $\triangle H_{f^{\diamond}} = \triangle H_{sub} + IE + \triangle H_{diss} + EA + U$ 

29.a.i)What are the essential conditions for writing resonating structures.

In resonance structures, the electrons are able to move to help stabilize the molecule. This

movement of the electrons is called delocalization.

- 1. Resonance structures should have the same number of electrons, do not add or subtract anyelectrons. (check the number of electrons by simply counting them).
- 2. All resonance structures must follow the rules of writing Lewis Structures.
- 3. The hybridization of the structure must stay the same.
- 4. The skeleton of the structure can not be changed (only the electrons move).
- 5. Resonance structures must also have the same amount of lone pairs.

ii)Draw all the resonating structures of nitrous oxide.

$$\underset{N}{\overset{\oplus}{=}} \overset{\Theta}{\xrightarrow{}} \overset{O}{\xrightarrow{}} \overset{O}{\xrightarrow{}} \overset{W}{\xrightarrow{}} \overset{W}{\xrightarrow{}}$$

- 1. Nitrous oxide known to be resonance hybrid of two highly polar and linear structures with opposing dipole moments
- 2. It has very low dipole moment
- 3. N<sub>2</sub>O is linear structure
- 4. N-N bond length is  $1.12 \text{ \AA}$
- 5. N-O bond length is 1.19 Å

b.State the geometrical arrangements possible in  $sp^3d$  and  $sp^3d^2hybridisations.$ Name the d-orbitals involved in these.

The basic principles, which are involved in the valence bond treatment of coordination compounds are:

- (a) Hybridization of valance orbitals of the central metal/ ion
- (b) Bonding between ligand and the metal ion/atom.
- (c) Relation between the type of bond and the observed magnetic behavior.

Let us explain by taking simple examples such as [CoF6]3- and [Co(NH3)6]3+. Although in both the complexes, the oxidation state of cobalt is +3, but [CoF6]3- is paramagnetic and  $[Co(NH_3)_6]^{3+}$  is diamagnetic, why? The formation of a complex may be considered as a series of hypothetical steps. First the appropriate metal ion is taken e.g.  $Co^{3+}$ . Cobalt atom has the outer electronic configuration  $3d^74s^2$ . Thus  $Co^{3+}$ ion will have the configuration  $3d^6$  and the electrons will be arranged as:



 $\text{Co}^{3+}$  ion forms both paramagnetic (outer orbital) and diamagnetic (inner orbital) complexes depending upon the nature of ligands as illustrated below. As  $\text{Co}^{3+}$  ion combines with six fluoride ligands in  $[\text{CoF6}]^{3-}$ , empty atomic orbitals are required on the metal ion to receive the coordinated lone pair of electrons. The orbitals used are one 4*s*, three 4*p* and two 4*d*. These are hybridized to give a set of six equivalent

 $sp^3d^2$  hybrid orbitals. A ligand orbital containing a lone pair of electron forms a coordinate bond by overlapping with an empty hybrid orbital on the metal ion. In this way a bond is formed with each ligand.



Since the outer 4d orbitals are used for bonding, this is called an outer orbital complex. The energy of these orbitals is quite high, so the complex will be reactive. This complex will be high-spin paramagnetic, because it has four unpaired electrons.

An alternative octahedral arrangement in [Co(NH3)6]3+ is possible when the electrons on metal ion are rearranged as shown below:



Since inner *d*-orbitals are used this is called an inner orbital complex. There is no unpaired electron, the complex will be low-spin diamagnetic.

30.a. Explain how the atomic orbitals combine to form bonding and antibonding M.O. What are the limitations to such combination.

A covalent bond is formed by the overlapping of atomic orbitals which form molecular orbital. Bonding electrons occupy molecular orbital not atomic orbital. An electron in a molecular orbital is polycentric because it is influenced by more than one nuclei. Formation of molecular orbital is based on the linear combination of atomic orbitals

There are two kinds of molecular orbitals:

### 1. BONDING MOLECULAR ORBITAL

# 2. ANTI-BONDING MOLECULAR ORBITAL

### Sigma bond:

1. Sigma bond is formed by the linear or head to head or end on overlapping of orbitals.

2. Sigma bonds are the strong bonds due to maximum overlapping of orbitals.

3. Electron density is maximum around the bond axis.

4. Compounds having sigma bonds are stable and hence less reactive.

5. First bond formed between two atoms is always a sigma bond.

6. s-s overlap, s-p overlap and p-p overlap give rise to sigma bond.

Examples of Sigma bond :

### s-s overlap in H<sub>2</sub> molecule

Hydrogen molecule consists of two H-atoms. Each atom contains one electron in 1sorbital.

# $E.C = 1s^1$

According to molecular orbital theory two  $1s^1$  orbitals of two H-atoms overlap linearly to produce two types of molecular orbitals.

1. Bonding molecular orbital (s-orbital)

# 2. Anti-bonding molecular orbital (s\*-orbital)

Bonding molecular orbital (s-orbital) has lower energy and results in the covalent bond formation which is a sigma bond between two H-atoms while the anti-bonding molecular orbital (s\*-orbital) possess high energy remains unoccupied.

# s-p overlap in HF molecule

s-p type overlap occurs in HF molecule when one **1s-orbital** of hydrogen atom overlaps **2p-** orbital of fluorine to form two types of molecular orbitals.

1. Bonding molecular orbital (s-orbital)

2. Anti-bonding molecular orbital (s\*-orbital)

Bonding molecular orbital (s-orbital) has lower energy and results in the covalent bondformation (sigma bond) between H-atom and F-atomwhile the anti-bonding molecularorbital (s\*-orbital) possess high energy remains unoccupied.

# p-p overlap in HF molecule

p-p type overlap occurs in F2 molecule when one **2pz**-orbital of one fluorine atomoverlaps **2pz**-orbital of other fluorine atom form two types of molecular orbitals.

- 1. Bonding molecular orbital (s-orbital)
- 2. Anti-bonding molecular orbital (s\*-orbital)

Bonding molecular orbital (s-orbital) has lower energy and results in the covalent bond

formation (sigma bond) between two F-atoms while the anti-bonding molecular orbital

(s\*-orbital) possess high energy remains unoccupied.

b.Distinguish between molecular orbital theory and valence bond theory.

# Valence bond theory

According to Valence bond theory:

- 1. A covalent bond is formed by the overlapping of partially filled orbitals of twoatoms.
- 2. Overlapping orbitals must have electrons with opposite spin.
- 3. Atoms involved in bond formation should have unpaired electrons.
- 4. The number of covalent bonds formed by an atom would be equal to thenumber of half filled orbital.
- Resulting molecular orbital is obtained by the combination of the two wavefunctions (AOs) of two unpaired electrons.
- 6. Atoms which are involved in bond formation maintain their identity.

# **Molecular Orbital Theory:**

- 1. A covalent bond is formed by the overlapping of atomic orbitals which form molecular orbital.
- 2. Bonding electrons occupy molecular orbital not atomic orbital.
- 3. An electron in a molecular orbital is polycentric because it is influenced by more than onenuclei.

#### Reg. No.....

[16CHU101]

# KARPAGAM UNIVERSITY

Karpagam Academy of Higher Education (Established Under Section 3 of UGC Act 1956) COIMBATORE - 641 021 (For the candidates admitted from 2016 onwards)

### B.Sc. DEGREE EXAMINATION, NOVEMBER 2016 First Semester

#### CHEMISTRY

### **INORGANIC CHEMISTRY I**

(Atomic Structure and Chemical Bonding) Maximum : 60 marks

Time: 3 hours

### PART – A (20 x 1 = 20 Marks) (30 Minutes) (Question Nos. 1 to 20 Online Examinations)

### PART B (5 x 2 = 10 Marks) (2 ½ Hours) Answer ALL the Questions

- 21. Find out the de-Broglie wavelength of an electron moving with a velocity of 2.2  $\times 10^{6}$  m/s.(The mass of an electron = 9.108  $\times 10^{-31}$  kg).
- 22. What is the sequence of filling electrons in sub-levels according to Aufbau principle?
- 23. Ionic compounds are high melting points. Why ?
- 24. Why borazole is also called as inorganic benzene?
- 25. How will you represent bonding and antibonding molecular orbitals on the basis of LCAO approximation?

#### PART C (5 x 6 = 30 Marks) Answer ALL the Questions

### 26. a. Write down the significance of $\psi$ and $\psi^2$

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- b. What is quantum mechanics? Explain their postulates.
- 27. a. Draw the shapes of s, p and d orbitals.

Or

b. What are the causes of stability of half filled and completely filled sublevels? Explain with one example each. 28. a. What are the factors which affect the polarisation of the ion? Or

- b. How will you determine the lattice energy of an ionic solid?
- 29. a. Based on VSEPR theory discuss the structures of  $PCl_5$  and  $SF_6$ .
  - b. Explain the concept of resonance with one example of organic and inorganic compound.

30. a. Distinguish between bonding molecular orbitals and anti bonding orbitals. Or
b. Draw the M.O diagram of NO and NO<sup>+</sup> is more stable than NO. Why?