28. CO-ORDINATION COMPOUNDS

1. INTRODUCTION

The concept of coordination compounds originates from the tendency for complex formation of transition elements.

Molecular or Addition Compounds

When solutions of two or more simple stable compounds in molecular proportion are allowed to evaporate, crystals of new substances, called molecular or addition compounds, are obtained. Some common examples are given below:

$CuSO_4 + 4NH_3$	\rightarrow	CuSO ₄ ·4NH ₃
AgCN + KCN	\rightarrow	KCN·AgCN
4 KCN + Fe (CN) ₂	\rightarrow	Fe (CN) ₂ .4KCN
$K_2SO_4 + Al_2(SO_4)_3$	$+24H_2O \rightarrow K_2$	$SO_4.AI_2(SO_4)_3.24H_2O$ Alum
Simple stable		Addition or
Compounds		Molecular compounds

There are two types of molecular or addition compounds: (1) double salts or lattice compounds and (2) coordination or complex compounds.

Double Salts or Lattice Compounds

The addition compounds which are stable only in solid state but are broken down into their individual constituents when dissolved in water are called double salts or lattice compounds.

Nomenclature of Double Salts

(a) A hyphen (-) is used between the compounds while writing the names of double salts with the number of molecules appearing inside brackets at the end.

Example: $KCl \cdot MgCl_2 \cdot 6H_2O - potassium chloride-magnesium chloride-water (1/1/6)$

 K_2SO_4 ·Cr₂ (SO₄)₃·24H₂O – potassium sulphate-chromium sulphate-water (1/1/24)

(a) Cations and anions with the same oxidation number are represented with English alphabets (cation first). If more than one type of cations are present, they are arranged in the increasing order (ascending order) of oxidation numbers. Anions of different oxidation numbers are arranged in the following order:

Oxide (O_2^{-}) , hydroxide (OH^{-}) , inorganic anion, organic anion, hydride (H^{-}) .

The above rules are further clarified by following examples:

- (i) KNaCO₃ Potassium sodium carbonate (mixed salt)
- (ii) KHCO₃ Potassium hydrogen carbonate (mixed salt)

Note: Hydrogen is an exception and is written at the end.

- (iii) NH4·MgPO₄·6H₂O Ammonium magnesium phosphate-6-hydrate or water
- (iv) NaCl·NaF·2Na₂SO₄ (Hexa) sodium chloride fluoride (bis) sulphate

Coordination or Complex Compounds: Those molecular compounds which retain their identity in solid as well as in solution, are known as coordination or complex compounds. A part (or whole compound) of these compounds is not dissociated in solution and its behavior is different from its constituents.

Example:
$$K_4 \left[Fe(CN)_6 \right] \rightarrow 4K^+ + \left[Fe(CN)_6 \right]_{Complex ion}^{4-}$$

Illustration 1: Aqueous solution of potassium ferrocyanide does not give the test of Fe (II) and it is not poisonous like potassium cyanide. Why? (JEE MAIN)

Sol: Being a complex salt, it ionizes to K+ and [Fe (CN)₆]^{4–} ions. Due to absence of Fe (II) it does not give the test of Fe (II). Absence of free CN[–] makes it nonpoisonous.

Note: An imperfect complex compound may be too unstable to exist and may be completely dissociated in solution, and then becomes a double salt.

2. TYPES OF COORDINATION COMPOUNDS

- (a) Based on the nature of cations and anions, coordination compounds are classified into four types:
 - (i) Simple cation and complex anion: K_4 [Fe (CN)₆], K_2 [PtCl₆], etc.
 - (ii) Complex cation and simple anion: $[Cu (NH_3)_4] SO_4$ [Ni $(NH_3)_6] Cl_2$ etc.
 - (iii) Complex cation and complex anion: [Pt (NH₃)₄] [PtCl₄], etc.
 - (iv) Neutral complex compounds: Ni $(CO)_{4'}$ Fe $(CO)_{5'}$ etc.
- (b) Based on stability, coordination compounds are of two types:
 - (i) Perfect or penetrating complexes: These are compounds in which the complex ion is feebly dissociated in solution and is highly stable.
 - E.g. K_4 [Fe (CN)₆] \rightarrow 4K⁺ + [Fe (CN)₆]⁴⁻

$$[Fe (CN)_c]^{4-} \rightleftharpoons Fe^{2+} + 6CN^{-}$$
 (Feebly dissociated)

(ii) Imperfect or normal complexes: These are compounds in which the complex ion is appreciably dissociated and is less stable.

E.g. $K_4 [Cd (CN)_4] \rightarrow 2K^+ + [Cd (CN)_4]^{2-}$ $[Cd (CN)_4]^{2-} \rightleftharpoons Cd^{2+} + 4CN^-$

(Reversible dissociation)

3. TERMINOLOGIES IN COORDINATION COMPOUNDS

3.1 Central Metal Atom/Ion

Metal atom of complex ion which coordinates with atoms or group of atoms/ions by accepting

their lone pair of electrons. Here Cu is the central metal atom. Some complex ions may also have more than one metal atom.



3.2 Ligands

The anions, cations or neutral molecules, which form coordinate bonds with the central metal atom by donating an electron pair (lone pair) are ligands. These electron pair donors are also known as Lewis bases. Thus, a complex ion is formed as: $M^{n+} + xL \rightarrow [ML_{\nu}]^{n+}$

In the above example, NH_3 is the ligand.

Classification of Ligands

Ligands are classified as follows:

(a) On the basis of the charge of ligand:

- (i) Anionic ligands: These are negatively charged and are the most common type of ligand, such as, F⁻, Cl⁻, Br⁻, OH⁻, CN⁻, SO₃²⁻, S²⁻, SO₄²⁻, etc.
- (ii) Neutral ligands: These are uncharged and are the electron pair donor species such as H₂ Ö,RÖH, NH₃ RNH₂, R₂NH, R₃N etc.
- (iii) Cationic ligands: They are positively charged and are rare such as NO⁺, etc.
- (b) On the basis of denticity: The number of donations accepted by a central atom from a particular ligand is known as the denticity of the ligand. Based on this, ligands are classified as follows:
 - (i) Monodentate: when only one donation is accepted from the ligand. For example,

 H_2O , NO, CO, $NH_{3'}$, CO^{2-} , CI^- , etc.

(ii) Bidentate: when two donations are accepted from the ligand. For example,

(1) En: Ethylenediamine







- (iii) Tridentate: when three donations are accepted from the ligand. For example,
 - (1) Dien: Diethylenetriamine





(iv) Tetradentate: when four donations are accepted from the ligand. For example,

(1) Trien: Trie thylenetetraamine

(2) NTA: Nitrilotriacetate





(v) **Pentadentate:** when five donations are accepted from the ligand. For example, EDTA: ethylenediamine



(vi) Hexadentate: when six donations are accepted from the ligand. For example, EDTA: Ethylenediamine tetraacetate



Some other types of ligands:

(a) Flexidentate: Some ligands exhibit variable denticity and are called flexidentate ligands. For example,



(b) **Chelating:** A ligand which forms a ring structure with the central atom is called a chelating ligand. All polydentate ligands are chelating ligands.

Chelated complex compounds are more stable than similar complexes with monodentate ligands because dissociation of the complex involves breaking down two bonds rather than one. However, it must be noted that NH_2NH_2 and $H_2N(-CH_2-CH_2-)NH_2$ cannot act as chelating ligands due to their three-member ring and locked structure, respectively.

- (c) Ambidentate ligand: A ligand with more than one kind of donor sites but only one site is utilized at a time is called an ambidentate ligand. Ambidentate ligands are of two types:
 - (i) Monodentate and ambidentate:



(ii) Bidentate and ambidentate:



(d) Classification of ligands on the basis of bonding interaction between the central metal atom and ligand:

- (i) Classical or simple donor ligand: These ligands donate their lone pair of electrons to the central atom. For example, O₂⁻, OH⁻, F⁻, NH₂⁻, NH₃, N³⁻, etc.
- (ii) Nonclassical or π -acceptor ligand: These ligands donate the lone pair of electrons to the central atom and accept the electron cloud from the central atom in their low-lying vacant orbital. This kind of back donation is known as 'synergic effect' or 'synergic bonding'. For example, CO, CN⁻, NO⁺, PF₃, PR₃ (R = H, Et, Ph...), C₂H₄, C₂H₂, CO₂, etc.
 - In the case of CO, the back donation to the π^* orbital of the central atom can be depicted as:



Figure 28.1: Back bonding in metal carbonyl

As per valence bond or molecular orbital theory, it is implicit that the bond order of C–O bond decreases but the C–O bond length increases due to synergic effect. Similarly, since CN⁻ and NO⁺ are isoelectronic with CO, back donation takes place in these species also in the π^* orbitals and the same conclusion can be drawn for the bond order and bond lengths.

• In PR₃, the back donation can be depicted as:



Figure 28.2: Back bonding in case of phosphine ligand

• In C_2H_4 , the back donation is clearly depicted with the example of Zeise's salt.



Figure 28.3: Back donation in case of ethylene ligand

Here, back donation is received in the p* orbital of the C–C bond. Hence, the bond order of C–C bond decreases and the bond length increases as compared to the free C_2H_4 molecule. Due to backbonding, the C_2H_4 molecule loses its planarity and likewise the C_2H_2 molecule loses its linearity.

Illustration 2: What is meant by the denticity of a ligand? Give examples of a unidentate and a bidentate ligand. (JEE MAIN)

Sol: Denticity indicates the number of donor sites in a ligand. It can be 1, 2, 3, 4 and 6 for unidentate, bidentate, terdentate, tetradentate and hexadentate ligands respectively. Unidentate ligands: Cl, H_2O , NH_3 , Bidentate ligands: Ethylene diamine, Propylene, diamine.

Illustration 3: Although NH₂·NH₂ possesses two electron pairs for donation, it does not act as a chelating agent. Why? (JEE MAIN)

Sol: The coordination by $NH_2 \cdot NH_2$ leads to a three-member strained ring which is highly unstable and hence it does not act as chelating agent.

3.3 Coordination Number

- (a) The number of atoms in a ligand that directly bond to the central metal atom or ion by coordinate bonds is called the coordination number of the metal atom or ion.
- (b) In other words, it is the number of coordinate covalent bond which the ligands form with the central metal atom or ion.
- (c) Some common coordination numbers exhibited by metal ions are 2, 4 and 6. The light transition metals exhibit 4 and 6 coordination numbers while heavy transition metals exhibit coordination numbers above 6.
- (d) For example, the coordination number of Ni in the complex $[Ni (NH_3)_4] Cl_2$ is 4 and that of Pt in the complex $K_2 [PtCl_6]$ is 6.

3.4 Coordination Sphere



The central metal atom/ion and the ligands directly attached to it are collectively termed as the coordination sphere. Coordination sphere is represented inside square brackets, e.g. $\left[Ni (NH_3)_a \right]^{2+}$

3.5 Oxidation Number

The actual charge that a metal atom experiences in a complex is known as its oxidation number. In other words, oxidation number of a metal atom will be equal to the total charge on this atom if all the ligands are removed without their electron pair.

Calculation of oxidation number: Algebraic sum of oxidation numbers of all the atoms of a molecule/ion is equal to the charge on it.

For example, for $M_a [M'_b (L)_x]$ [a× (O.N. of M)] + [b × O.N. of M'] + [x × (O.N. of L)] = 0 For ion $[M_a (L)_x]^{y+}$ a × (O.N. of M) + x × (O.N. of L) = y e.g:- Oxidation number of Co (let it be x) in ion [Co (CN) (H₂O) (en)₂]²⁺ can be calculated as: x + (O.N. of CN) + (O.N. of H₂O) + 2 (O.N. of en) = +2

 $x + (-1) + (0) + (2 \times 0) = +2$ $\therefore x = +3$

MASTERJEE CONCEPTS

- (a) Transition metals exhibit variable oxidation states.
- (b) Oxidation number of different species:
 - (i) Alkali metals (Li, Na, K, Rb, Cs) = +1
 - (ii) Alkaline earth metals (Be, Mg, Ca, Sr, Ba, Ra) = +2
 - (iii) Oxidation number of ion = charge on ion
 - (iv) Oxidation number of neutral molecule = 0

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Illustration 4: Specify the oxidation number of the central metals in the following coordination entities:

(a) [Co (CN) (H ₂ O) (en) ₂] ²⁺	(b) [PtCl ₄] ²⁻	(c) [CrCl ₃ (NH ₃) ₃]	(d) [CoBr ₂ (en) ₂] ⁺
(e) K ₂ [Fe (CN) ₆]			(JEE MAIN)
Sol:			

(a) +3 (b) +2 (c) +3 (d) +3 (e) +3

3.6 Effective Atomic Number

- (a) Effective atomic number was first proposed by Sidgwick in order to explain the stability of the complex.
- (b) It can be defined as the resultant number of electrons present in the metal atom or ion after accepting electrons from the donor atoms of the ligands.
- (c) In some cases, the effective atomic number coincides with the atomic number of the next noble gas.
- (d) Effective atomic number is calculated as follows:

EAN = atomic number of the metal – number of electrons lost in ion formation + number of electrons gained from the donor atoms of the ligands.

MASTERJEE CONCEPTS

Calculation of EAN: Effective atomic number = Atomic number (Z) – Electron donated (Equal to O.N.) + Electrons accepted from ligands $(2 \times No. of coordinate bonds formed)$

Or $EAN = Z - O.N. + 2 \times (C.N.)$

Note: EAN and stability: An ion with central metal atom possessing EAN equal to next inert gas will be more stable.

Vaibhav Krishnan (JEE 2009, AIR 22)

Illustration 5: Metal carbonyls having formula M (CO)_x where the number of carbonyl units coordinated to metal M are formed by Fe, Cr and Ni. If effective atomic number of each metal is 36, write the formulas of these metal carbonyls. (JEE MAIN)

Sol: M (CO), In Fe (CO) EAN = At. No. of Fe + $2 \times$ No. of ligands.

(:: O.N. of Metals in Metal carbonyls is zero as these compounds are neutral species formed by neutral ligand CO).

i.e. CO \therefore 36 = 26+2x; \therefore x = 5

:. Formula of iron carbonyl is Fe (CO)₅, Similarly, we get Cr (CO)₆ and Ni (CO)₄.

4. FORMULA AND IUPAC NOMENCLATURE OF COORDINATION COMPOUNDS

4.1 Formula of a Complex

- (a) In formulas of both simple and complex salts, cation precedes the anion. Nonionic compounds are written as single units.
- (b) Complex ions are written inside square brackets without any space between the ions.
- (c) Metal atom and ligands are written in the following order:
 - (i) In the complex part, the metal atom is written first followed by ligands in the order, anionic \rightarrow neutral \rightarrow cationic.
 - (ii) If more than one ligand of one type (anionic, neutral or cationic) are present, then they are arranged in English alphabetical order, e.g. between H_2O and $NH_{2'}$, H_2O should be written first. Similarly, order of NO_2^- , SO_3^{2-} and OH^- will be NO_2^-
 - (iii) When ligands of the same type have similar name for the first atom, then the ligand with less number of such atoms is written first. Sometimes the second atom may be used to decide the order. When number of atoms are also same e.g., Out of NO_2^- , NH_2^- will be written first. In H₃ and \ddot{N}_2 , $\ddot{N}H_3$ will be written first as it contains only one N-atom.
 - (iv) Polyatomic ligands and abbreviations for ligands are always written in lower case letters. e.g. (en), (py), etc.
 - (v) Charge of a complex ion is represented as over script or square bracket.

E.g.

- K_4 [Fe (CN)₆] First cation and then anion [Rule 1 and 2]
- [CrCl₂ (H₂O)₄] Br—Cl– (negative ligand) before H₂O (neutral ligand [Rule 3-(i)]

4.2 Nomenclature of Coordination Compounds

Mononuclear coordination compounds are named by following these rules:

- (a) In both the positively and negatively charged coordination compounds, the cation is named first followed by the anion.
- (b) The ligands are named in alphabetical order before the name of the central atom/ion. (This procedure is reversed in writing its formula).

(c) Names of the anionic ligands end in –o. E.g.

Symbol	Name as ligand	Symbol	Name as ligand
N ³⁻	Azido	OH⁻	Hydroxo
CI-	Chloro	CO ₃ ²⁻	Carbonato
O-	Peroxo	C ₂ O ₄ ⁻²	Oxalato
Br	Bromo	SO ₄ ⁻²	Sulphato
O₂H⁻	Perhydroxo	NO ₃	Nitrato
CN-	Cyano	SO ₃ ²⁻	Sulphito
S ²⁻	Sulphido	CH₃COO-	Acetato
0 ₂ -	Охо	NO ₂	(Bonded through oxygen) nitrite
NH ₂	Amido		(Bonded through nitrogen) nitro

(d) Names of neutral and cationic ligands are the same except for aqua for H₂O, ammine for NH₃, carbonyl for CO and nitrosyl for NO. These are placed within parentheses ().

Symbol	Name as ligand	Symbol	Name as ligand
H ₂ O	Aqua	NO	Nitrosyl
NH ₃	Ammine	CS	Thiocarbonyl
СО	Carbonyl		

(e) Positive ligands are named as:

Symbol	Name as ligand	
NH ⁺⁴		
NO ⁺	Nitrosylium	
NH ₂ NH ₃ ⁺	Hydrazinium	

- (f) Prefixes mono, di, tri, etc. are used to indicate the number of the individual ligands in a coordination compound. When the names of the ligands include a numerical prefix, then the terms, bis, tris, tetrakis are used, and the ligand to which they refer is placed in parentheses. For example, [NiCl₂ (PPh₃)₂] is named as dichlorobis (triphenylphosphine) nickel (II).
- (g) Oxidation state of the metal in a cation, anion or a neutral coordination compound is indicated by a Roman numeral in parenthesis.
- (h) When the complex ion is a cation, the metal is named same as the element. For example, Co in a complex cation is called cobalt and Pt is called platinum. In an anion, Co is called cobaltate. For some metals, their Latin names are used in the complex anions, e.g. ferrate for Fe.
- (i) Nomenclature of a neutral complex molecule is done in the similar way as that of a complex cation.

The following examples illustrate the nomenclature for coordination compounds:

- [Cr (NH₃)₃ (H₂O)₃] Cl₃ is named as: Triamminetriaquachromium (III) chloride
- [Co (H₂NCH₂CH₂NH₂)₃]₂ SO₄ is named as: Tris (ethane-1, 2-diammine) cobalt (III) sulphate
- [Ag (NH₃)₂] [Ag (CN)₂] is named as: Diamminesilver (I) dicyanoargentate (I)
- (j) Ligands which join two metals are known as 'Bridge ligands' and they are prefixed by ' μ ' (mu).

E.g.
$$\left((NH_3)_4 C \circ \left(\frac{NH_2}{NO_2} \right) \right) \right) (NO_3)_4$$
, in this complex

Here, NH_2 and NO_2 are bridge ligands and they are named μ -amido and μ -nitro, respectively.

Illustration 6: Write the formula	(JEE MAIN)		
(A) Tetraammineaquachloridocol	balt (III) chloride	(B) Potassium tetrahydroxidozincate	(II)
(C) Potassium trioxalatoaluminat	e (III)	(D) Dichloridobis (ethane-1, 2-diam	ine) cobalt (III)
Sol: (A) [Co (NH ₃) ₄ (H ₂ O) Cl] Cl ₂	(B) K ₂ [Zn (OH)]	¹] (C) $K_3 [AI (C_2O_4)_3]$ (D) $[CoCl_2 (en)_2]^+$
Illustration 7: Write the IUPAC	names of the following o	coordination compounds:	(JEE MAIN)
(A) [Pt (NH ₃) ₂ Cl (NO ₂)]	(B) K ₃ [Cr (C ₂ O ₄) ₃]	(C) $[CoCl_2 (en)_2] Cl$	
(D) [Co (NH ₃) ₅ (CO ₃)] Cl	(E) Hg [Co (SCN) ₄]	(F) [Ni (CO) ₄]	
Sol: (A) Diamminechloridonitrito	-N-platinum (II)	(B) Potassium trioxalatochro	omate (III)
(C) Dichloridobis (ethane-1, 2-diamine) cobalt (III) chlorid		de (D) Pentaamminecarbonato	cobalt (III) chloride
(E) Mercury tetrathiocyanatocobaltate (III)		(F) Tetracarbonylnickel (0)	

5. ISOMERISM IN COORDINATION COMPOUNDS

Compounds having the same molecular formula but a different arrangement of atoms and different properties are called isomers and the phenomenon is called isomerism. Types of isomerism exhibited by complex compounds are summarized below:



5.1 Structural Isomerism

Structural isomerism occurs due to the difference in chemical linkages and distribution of ligands within and outside the coordination sphere. In Structural isomerism, isomers possess dissimilar bonding pattern. Different types of isomers are discussed below:

(a) **Ionization Isomerism:** Ionization isomerism is the result of the exchange of groups or ions between the coordinating sphere and the ionization sphere.

This isomerism occurs only in compounds where counter ions act as potential ligands. Ionization isomers exhibit different physical as well as chemical properties.

 $[CoBr(NH_{3})_{5}]SO_{4} \xrightarrow{\text{Ionisation}} [CoBr(NH_{3})_{5}]^{2+} + SO_{4}^{2-}$ Pentaamminebromocobalt (III) sulphate

 $[Co(SO_{4})(NH_{3})_{5}]Br \xrightarrow{Ionisation} [Co(SO_{4})(NH_{3})_{5}]^{+} + Br^{-}$ Pentaamminesulphatocobalt (III) bromide

Here (A) and (B) are ionization isomers. (A) forms white precipitate $(BaSO_4)$ with $BaCl_2$ whereas (B) does not react with $BaCl_2$. Similarly (B) gives yellowish white precipitate (AgBr) with AgNO₃ while (A) does not react with AgNO₃. Other examples of ionization isomers are:

(i) [PtCl ₂ (NH ₃) ₄] SO ₄	and	[Pt (SO ₄) (NH ₃) ₄] Cl ₂
(ii) [CoCl ₂ (NH ₃) ₄] NO ₂	and	$[CoCl (NO_2) (NH_3)_4] Cl$
(iii) [Pt (OH) ₂ · (NH ₃) ₄] SO ₄	and	[Pt (SO ₄) (NH ₃) ₄] (OH) ₂

- (b) Hydrate Isomerism (Solvate Isomerism): In a complex compound, water molecules behave in two ways:
 - (i) Water molecules which behave as ligands are coordinated with the metal atom and are part of the complex ion, e.g. [M (H₂O)_x].
 - (ii) Water molecules act as water of crystallization and these appear outside the coordination sphere, e.g. [ML_x].nH₂O.

Isomerism which occurs due to dissimilar number of water molecules as ligands (inside the sphere) and as water of crystallization (outside the sphere), is known as hydrate isomerism. This isomerism is analogous to ionization isomerism, in which water molecules inside and outside the sphere are exchanged.

For example,

 $Cr (H_2O)_6 Cl_3$ has three possible structures:

- $[Cr (H_2O)_6] Cl_3$ (violet)
- [Cr (H₂O)₅Cl] Cl₂H₂O (green)
- [Cr $(H_2O)_4CI_2$] Cl.₂H₂O (dark green)

These complex compounds differ from one another with respect to the number of water molecules acting as ligands.

Other hydrate isomers are:

- [Co (NH₃)₄ (H₂O)Cl]Cl₂.
- [Co (NH₃)₄Cl₂] Cl H₂O

(c) Linkage or Salt Isomerism:

(i) Linkage isomerism occurs in complex compounds having ambidentate ligands like

---CN, ---NC, ---NO₂, ---ONO, ---CNO, ---CNS, ---NCS, ---SCN, etc.

- (ii) In this isomerism, an ambidentate ligand coordinates with different atoms.
- (iii) These isomers can be differentiated by IR spectroscopy.

For example,

[Co (NO ₂) (NH ₃) ₅] Cl ₂	and	[Co (ONO) (NH ₃) ₅] Cl ₂
(A)		(B)
Pentaamminenitrocobalt (III) chloride		Pentaammine nitritocobalt (II) chloride
(Yellow-red)		(Red)

(A) is not decomposed by the action of acids whereas (B) liberates HNO3 by the action of acid. Other examples of linkage isomers are:

(i) [Cr (SCN) (H ₂ O) ₅] ²⁺	and	[Cr (NCS) (H ₂ O) ₅] ²⁺
(ii) [Co (NO ₂) (py) ₂ (NH ₃) ₂] NO ₃	and	[Co (ONO) (py) ₂ (NH ₃) ₂] NO ₃

(d) **Polymerization Isomerism:** When two compounds possess stoichiometric composition but different molecular formulas, they are known as polymerization isomers of each other. Molecular formula of one isomer will be the integral multiple of the other one.

Example: [PtCl₂ (NH₃)₂] and [Pt (NH₃)₄] [PtCl₄]

(e) Coordination Isomerism:

- (i) This isomerism occurs only in those complexes in which both cation and anion are complex.
- (ii) It occurs as a result of the exchange of ligands between the cation and anion.
- (iii) It may occur in those complexes also in which both cation and anion have the same metal atoms.

Example:

- (i) $[Cr (NH_3)_6] [Cr (SCN)_6]$ and $[Cr (SCN)_2 (NH_3)_4] [Cr (SCN)_4 (NH_3)_2]$ (ii) $[Co (NH_3)_6] [Cr (C_2O_4)_3]$ and $[Cr (NH_3)_6] [Co (C_2O_4)_3]$
- (f) Coordination Position Isomerism: It occurs in complexes containing bridge ligands and is the result of

dissimilar arrangement of metal atoms forming bridge, e.g. $\left(NH_{3}\right)_{4}CO\left(NH_{3}\right)_{2}Cl_{2}SO_{4}$ and

$$\left(CI \left(NH_{3} \right)_{4} Co \left(CI \left(NH_{3} \right)_{4} Co \left(CI \left(NH_{3} \right)_{3} CI \right) SO_{4} \right) \right)$$

5.2 Stereoisomerism

Stereoisomerism occurs as a result of the different arrangements of ligands around the central metal atom. It may be of two types: (1) Geometrical isomerism and (2) Optical isomerism.

5.2.1 Geometrical Isomerism

Isomerism which occurs due to different relative arrangements of ligands around the central metal atom is known as geometrical isomerism. Geometrical isomers are of two types:

- (a) **Cis-isomer:** In a disubstituted complex molecule/ion, when two similar ligands are at right angle (90°), the geometrical isomer is known as Cis-isomer.
- (b) **Trans-isomer:** When two ligands are positioned in opposite directions, i.e. at 180° to each other, the isomer formed is trans-isomer.

Cis- and Trans- positions are indicated in figures:



Cis- positions: (1, 2), (2, 3), (3, 4), (1, 4)

(1, 2), (2, 3), (3, 4), (1, 4), (1, 5), (4, 5), (3, 5), (2, 5), (1, 6), (2, 6), (3, 6) and (4, 6)

Trans- positions: (1, 3) and (2, 4)

Geometrical Isomerism and Coordination Numbers

Geometrical Isomerism with Coordination Number 4:

Tetrahedral complexes do not show geometrical isomerism as all the four valences are identical.

Square-planar complexes:

- (a) Complexes of type MA₄, MA₃B and MAB₃ do not show geometrical isomerism, where A and B are monodentate ligands.
- (b) Complexes of formula MA₂B₂ and MA₂BC types have two geometrical isomers, where A and B are monodentate ligands.

Example:

(i) $[PtCl_2 (NH_3)_2]$ resembles MA_2B_2 in formula and exists in two isomeric forms:



(ii) [PtCl (NH₃) (py)₂] resembles MA₂BC and exists in two isomeric forms:



(c) Complexes of formula MABCD exist in three isomeric forms:



- (III) and (IV) are similar.
- e.g. [Pt (NO₂) (NH₂OH) (NH₃) (py)] + exists in 3 isomeric forms.

[Pt (gly)₂] gly = glycinate ($\ddot{N}H_2$ —CH₂—COO–)

 $A = NO_{2'}$ $B = NH_2OH$, $C = NH_{3'}$ D = py

(d) If A is an unsymmetrical bidentate ligand, then compounds having formula MA₂ tend to exhibit geometrical isomerism, e.g.



(e) Bridged dinuclear complexes of formula M2A2B4 also exhibit geometrical isomerism, e.g. $|PtCl_2(P(C_6H_5)_2)|_2$



Geometrical Isomerism with Coordination Number 6:

- (a) Complexes of type MA₆ and MA₅B type do not show geometrical isomerism.
- (b) Complexes of type MA_4B_2 or MA_4BC exist in two isomeric forms, e.g. $[CoCl_2 (NH_3)_4]^+$



(c) Complexes of type MA₃B₃ exist in two geometrical forms which are named as facial (fac–) and meridonial (mer–) isomers. When three ligands of the same type are arranged in one triangular face, then isomer is facial. fac- and mer- isomers of complex MA₃B₃ are as follows:



E.g. $[Co (NO_2)_3) (NH_3)_3]$ can be represented in fac- and mer- isomeric forms as follows:



Similarly, [RhCl₃ (py)₃] also exists in fac- and mer- forms.

- (d) Complex compound of formula MABCDEF may exist in 15 isomeric forms and only one compound of this type is identified so far [Pt (Br) (Cl) (I) (NO₂) (NH₃) (py)].
- (e) Complexes of formula M(AA)₂B₂ and M(AA)₂BC also exhibit geometrical isomerism, where A is the symmetrical bidentate ligand, e.g. ethylenediamine (en), oxalate (ox), etc. [CoCl₂ (en)₂]⁺



(f) Complex of type $M(AA')_3$ also exists in Cis- and Trans- forms. Where AA' is unsymmetrical bidentate ligand, e.g. [Cr (gly)₃], gly: glycinate (NH₂CH₂COO⁻)



Illustration 8: Draw the structure of geometrical isomers of [Pt (gly),] where gly is NH₂CH₂COO⁻. (JEE ADVANCED)



5.2.2 Optical Isomerism

Optical activity: Compounds which rotate on the plane of polarized light are optically active. If the plane rotates clockwise, then the isomer is said to be dextro rotator (d or +) and if the plane rotates anticlockwise then the isomer is said to be laevo rotator (ℓ or -). Equimolar mixture of d– and isomer is optically inactive and is called racemic mixture. Optical isomers differ in optical properties.

(a) Optical isomerism in complexes with coordination number 4:

(i) Tetrahedral complexes: Like carbon compounds, complex MABCD must be optically active but due to their labile nature, such complex cannot be resolved in d or ℓ form. However, tetrahedral complexes with unsymmetrical bidentate ligand are optically active. In optically active tetrahedral compounds, the ligand must be unsymmetrical. It is not necessary whether it is chiral (asymmetric) or not, e.g. bis (benzoyl acetonato) beryllium (II)



Another example of this type is [Ni (CH₂NH₂COO)₂]—bis (glycinato) nickel (II)

$$\binom{O}{N} \times Ni \binom{O}{N} = \binom{O}{N} \times Ni \binom{O}{N}$$

Illustration 9: Draw all the optical isomers for $[(en)_2 Co < NO_2 > Co(en)_2]^{4+}$ (JEE MAIN)

Sol: Complex compound shows optical isomerism and exists in d ℓ and meso forms.



I and II d and ℓ form (mirror image of each other), III meso-form

(ii) **Square planar complexes:** Generally square planar complexes are not optically active as they have all the ligands and metal atoms in one plane. That is why there is a plane of symmetry.

Note: However some optically active square planar complexes are identified, e.g. isobutylenediaminemesostilbenediaminoplatinum (II) ion.



(a) Optical isomerism in compounds of coordination number 6 – Octahedral complexes:

- (i) Complexes of type MA₄R₂ exist in cis- and trans- forms and both forms are optically inactive due to plane of symmetry.
- (ii) Complexes of type MA₃B₂ exist in facial and meridonial forms but both are optically inactive.
- (iii) Complexes of type $MA_2B_2C_2$ are optically active, e.g. five geometrical isomers of $[PtCl_2 (NH_3)_2 (py)_2]^{2+}$ are possible. Out of these five possible isomers, three have been prepared. Their cis- form is optically active while trans- forms are optically inactive due to symmetry.



(iv) Complex MABCDEF has 15 geometrical isomers and each isomer exists as pair of enantiomers and hence total 30 optical isomers will be possible. Only one such compound has been identified so far – [Pt (Br) (Cl) (I) (NO₂) NH₂) (py)].

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Number of Possible Isomers for Specific Complexes

Formula	Number of Stereoisomers	Pairs of Enantiomers
MA ₆	1	0
MA _s B	1	0
MA ₄ B ₂	2	0
MA ₃ B ₃	2	0
MA ₄ BC	2	0
MA ₃ BCD	5	1
MA ₂ BCDE	15	6
MABCDEF	30	15
MA ₂ B ₂ C ₂	6	1
MA ₃ B ₂ CD	8	2
MA ₃ B ₂ C	3	0
M (AA) BCDE	10	5
M (AB) ₂ CD	11	5

Saurabh Gupta (JEE 2010, AIR 443)

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Number of Possible Isomers for Specific Complexes

Formula	Number of Stereoisomers	Pairs of Enantiomers
M (AB) (CD)EF	20	10
M (AB) ₃	4	2
M (ABA)CDE	9	3
M (ABC) ₂	11	5
M (ABBA)CD	7	3
M (ABCBA)D	7	3

Saurabh Gupta (JEE 2010, AIR 443)

6. PREPARATION AND IDENTIFICATION OF COMPLEX COMPOUNDS

Preparation of Complex Compounds

(a) By substitution reactions: Ion of a salt can be substituted by a ligand to form a complex compound, e.g.

$$CuSO_{4} + 4NH_{3} \longrightarrow [Cu(NH_{3})_{4}]SO_{4}$$

tetraamminecopper(II)sulphate

(b) By combination reaction: Various complexes can be formed by combination reactions:

 $NiCl_2 + 6\ddot{N}H_3 \longrightarrow [Ni(NH_3)_6]Cl_2$

 $AgCI + 2\ddot{N}H_{3} \longrightarrow [Ag(NH_{3})_{2}]CI$

(c) By redox reactions: Two important examples are:

$$2[Co(H_2O)_6](NO_3)_2 + 8\ddot{N}H_3 + 2NH_4NO_3 + H_2O_2 \longrightarrow 2[Co(NO_3)(NH_3)_5](NO_3)_2 + 14H_2O_3 \longrightarrow 2[Co(NO_3)(NH_3)_5](NO_3)(NH_3)_2 + 14H_2O_3 \longrightarrow 2[Co(NO_3)(NH_3)(NH_3)_2 + 14H_2O_3 \longrightarrow 2[Co(NO_3)(NH_3)(NH_3)](NH_3) + 14H_2O_3 \longrightarrow 2[Co(NO_3)(NH_3)(NH_3)(NH_3))$$

$$2\text{CoCl}_2 + 2\text{NH}_4\text{Cl} + 10\text{NH}_3 + \text{H}_2\text{O}_2 \longrightarrow 2[\text{Co(NO}_3)_6]\text{Cl}_3 + 2\text{H}_2\text{O} + 9\text{H}_2$$

Identification of Complex Compound Formation

(a) Change in solubility: Solubility of a complex compound changes abnormally when complex is formed, e.g.

 $\begin{array}{l} AgCN + KCN \longrightarrow K[Ag(CN)_{2}] \\ Partialy soluble & & \\ Soluble & \\ AgCl + 2NH_{3} \longrightarrow [Ag(NH_{3})_{2}]Cl \\ Insoluble & \\ Ni^{2+} + 2dmg \longrightarrow [Ni(dmg)_{2}]^{2+} \\ Soluble & \\ Mig^{2+} + 2dmg \longrightarrow [Ni(dmg)_{2}]^{2+} \\ Soluble & \\ Mig^{2+} + 2dmg \longrightarrow [Ni(dmg)_{2}]^{2+} \\ Mig^{2+} + 2dmg \longrightarrow [Ni(dmg)_{$

- (b) Change in conductivity: As complex formation changes the solubility, the number of ions in solution also changes and hence conductance changes suddenly.
- (c) Change in chemical properties: Change in chemical properties of metal ion also indicates the formation of complex, e.g.

Ag⁺ is precipitated by KCl solution but in the presence of NH_3 , KCl does not precipitate Ag⁺ due to formation of [Ag (NH₃)₂] Cl.

(d) Change in color: Change in color indicates complex formation, eg.

$$C_{\text{Pink}}^{O^{2+}} + 4Cl^{-} \longrightarrow [CoCl_{4}]^{2-}_{\text{Blue}}$$

$$C_{\text{Lightblue}}^{Q^{2+}} + 4\ddot{N}H_{3} \longrightarrow [Cu(NH_{3})_{4}]^{2+}_{\text{Deepblue}}$$

$$[Ni(H_{2}O)_{6}]^{2+} + 6NH_{3} \longrightarrow [Ni(NH_{3})_{6}]^{2+} + 6H_{2}O_{\text{Blue}}^{Q^{2+}}$$

(e) Change in pH, EMF, Magnetic properties or colligative properties also indicate the complex formation.

7. THEORIES OF BONDING IN COORDINATION COMPOUNDS

7.1 Werner's Theory

In 1898, Werner propounded his theory of coordination compounds. Werner proposed the concept of a primary valence and a secondary valence for a metal ion. Main postulates of his theory are:

- (a) In coordination compounds metals show two types of linkages (valencies) primary and secondary.
- (b) The primary valencies are normally ionizable and are satisfied by negative ions.
- (c) The secondary valencies are nonionizable. These are satisfied by the neutral molecules or negative ions (ligands). The secondary valency is equal to the coordination number and is constant for a metal.
- (d) The ion groups bounded by the secondary linkages to the metal have a characteristic spatial arrangement corresponding to their different numbers.

Illustration 10: $PtCl_4$ and NH_3 may form five complexes, A ($PtCl_4 \cdot 6NH_3$), B ($PtCl_4 \cdot 5NH_3$), C ($PtCl_4 \cdot 4NH_3$), D $PtCl_4 \cdot 3NH_3$ and E ($PtCl_4 \cdot 2NH_3$). One mole of each A, B, C, D and E reacts with excess of AgNO₃ to yield 4, 3, 2 and 1 mole AgCl respectively, while E gives no AgCl. The conductances of their solutions are in the order A > B > C > D > E. On the basis of Werner's theory, write their structure and give the total number of ions given by one complex.

(JEE ADVANCED)

Formula	Structural formula	Ionization	No. of ions
(A) PtCl ₄ ·6NH ₃	[Pt (NH ₃) ₆] Cl ₄	\rightleftharpoons [Pt (NH ₃) ₆] ⁴⁺ + 4Cl ⁻	5
(B) PtCl ₄ ·5NH ₃	[PtCl (NH ₃) ₅] Cl ₃	$\rightleftharpoons [PtCl (NH_3)_5]^{3+} + 3Cl^-$	4
(C) PtCl ₄ ·4NH ₃	[PtCl ₂ (NH ₃) ₄] Cl ₂	\rightleftharpoons [PtCl ₂ (NH ₃) ₄] ²⁺ + 2Cl ⁻	3
(D) PtCl ₄ ·3NH ₃	[PtCl ₃ (NH ₃) ₃] Cl	$\rightleftharpoons [PtCl_3 (NH_3)_3]^+ + Cl^-$	2
(E) PtCl ₄ ·2NH ₃	[PtCl ₄ (NH ₃) ₂]	No isonisation possible	0

Sol:

7.2 Valence Bond Theory

The salient features of the valence bond theory are summarized below:

- (a) The central metal ion has a number of empty orbitals for accommodating electrons donated by the ligands. The number of empty orbitals is equal to the coordination number of the metal ion for the particular complex.
- (b) The atomic orbitals (s, p or d) of the metal ion hybridize to form hybrid with definite directional properties. These hybrid orbitals tend to form strong chemical bonds with the ligand orbitals.
- (c) The d-orbitals involved in the hybridization may be either inner (n 1) d orbitals or outer n d-orbitals. The complexes formed in these two ways are referred to as low spin and high spin complexes, respectively.
- (d) Each ligand possesses a lone pair of electrons.
- (e) A covalent bond is formed when a vacant hybridized metal orbital and a filled orbital of the ligand overlap. The bond is also known as a coordinate bond or dative bond.
- (f) When a complex contains unpaired electrons, it is paramagnetic in nature, whereas if it does not contain unpaired electrons, it is diamagnetic in nature.
- (g) The number of unpaired electrons in the complex, determines the geometry of the complex as well as hybridization of the central metal ion and vice-versa. In practice, the number of unpaired electrons in a complex is found from magnetic moment measurements as illustrated below. $\mu = \sqrt{n(n+2)}$ where n = no. of lone pair.
- (h) Thus the knowledge of the magnetic moment can be of great help in ascertaining the type of complex.
- (i) Under the influence of a strong ligand, the electrons can be forced to pair up against the Hund's rule of maximum multiplicity.

Coordination Number	Type of Hybridization	Distribution of hybrid orbital in space
4	sp ³	Tetrahedral
4	dsp ²	Square planar
5	sp³d	Trigonal bipyramidal
6	sp ³ d ²	Octahedral
6	d ² sp ³	Octahedral

Application of Valence Bond Theory on Coordination Complexes

(a) **Complex with Coordination Number 4:** Tetra coordinated complexes have either tetrahedral or square planar geometry depending on the nature of orbitals involved in hybridization. If one ns and three np orbitals are involved in bonding, geometry will be tetrahedral and hybridization sp³. If (n - 1) d, ns and two np are involved in bonding, geometry will be square planar and hybridization dsp². Tetra coordinated complexes are common with Ni (II), Cu (II), Pt (II), etc.



Some examples of tetra coordinated complexes are given below:

(i) Tetrahedral Complexes:

Ni (CO)₄: In Ni (CO)₄, Ni has zero oxidation state and exists as Ni (0). Four ligands (CO) are attached to central metal atom Ni and require four orbitals. The electronic configuration in
 Ni (CO) can be written as:

Ni $(CO)_4$ can be written as:



When four CO ligands are present, it is a strong ligand and the electrons pair up against "Hund's rule for maximum multiplicity".



Explanation: Four sp³ hybrid orbitals are arranged tetrahedrally making it a tetrahedral complex. Since all the electrons are paired, it is diamagnetic.

(ii) Square planar complexes:

• [Ni (CN)₄]²⁻: Here, Ni is in (II) oxidation state and the electronic arrangement is as follows:



Ni²⁺ ion in [Ni (CN) 4]²⁻ 3d⁸, 4s⁰

Explanation:

• CN– is a strong ligand and so it pairs up 3d-electrons against Hund's rule. The d-orbital thus made vacant, takes part in hybridization.

CN

CN CN CN

- Four dsp² hybrid orbitals are arranged in this manner and hence the geometry is square planar.
- Complex compounds are diamagnetic because all the electrons are paired.
- The complex makes use of the inner d-orbital, and so it is known as inner orbital or low spin or hyper ligand or spin paired complex.

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Cu [Ar]: 3d10, 4s1

Exception: Structure of $[Cu (NH_3)_4]^{2+}$ ion: It is an exceptional case which involves sp² d hybridization. Here, Cu is tetra coordinated and may exist as square planar or tetrahedral complex. Physical measurement have indicated that tetrahedral geometry for $[Cu (NH_3)_4]^{2+}$ is not possible. If square planar geometry is supposed to be correct, then the following electronic arrangement must be followed:



For dsp² hyb.3d-electron must be excited to 4-p with the following configuration.

 Cu^{2+} ion in $[Cu (NH_3)_4]^{2+}$:

 Cu^{2-} ion: \rightarrow [Ar]3d⁹, 4s⁰

ds	p² hybri	disation
11111	Nerit in the second sec	11
T NH,	↑ NH ₃	↑ ↑ NH₃ NH₃



Now if the above configuration is correct, the unpaired electrons present in higher energy, 4-p orbital should be expected to be easily lost and Cu²⁺ must be easily oxidized to Cu³⁺, but it never occurs, so the configuration is not satisfactory. To explain it Huggin suggested sp²d hybridization.

 Cu^{2+} ion in $[Cu (NH_2)_4]^{2+}$:



Note: Pt (II) and Au (III) always form square planar complexes irrespective of their ligands being strong or weak.

Neeraj Toshniwal (JEE 2009, AIR 21)

(b) **Complexes with Coordination Number 6:** Hexacoordinated complexes are of two types, inner orbital complexes and outer orbital complexes. They possess octahedral geometry.

(i) Inner orbital complexes: In this type of complexes the d-orbitals used are of lower quantum number, i.e. (n - 1). Some examples are given below:

- Complexes formed by using the inner orbitals are diamagnetic or have reduced paramagnetism.
- These are also known as low spin or spin paired complexes,

Example 1: [Fe (CN)₆]⁴⁻

 e^{-} configuration of $Fe^{26} = [Ar] 3d^{6}4s^{2}$



The above rearrangement is due to presence of the cyanide ligand.

At this stage, Fe^{2+} undergoes d^2sp^3 hybridization to form six d^2sp^3 hybrid orbitals, each of which accepts an electron pair donated by CN^- ions. The complex is Diamagnetic as it has no unpaired electron.

Example 2: [CO(NH₈)₆]³⁺



As this d^2sp^3 hybridization leads to octahedral geometry, the complex $[Cr (NH_3)_6]^{3+}$ will be octahedral in shape. Since the complex ion has 3 unpaired electrons, it is paramagnetic.

Other complexes of chromium with similar inner structure are $[Cr (CN)_6]^{3-}$ and $[Cr (H_2O)_6]^{3+}$.

(ii) Outer orbital complexes

- In these complexes s, p and d orbitals which are involved in hybridization, belong to the highest quantum number (n).
- Complex compound formed by the use of outer n and d orbitals will be paramagnetic.
- Outer orbital complexes are also known as high-spin or spin free complexes.
- The outer orbital complexes have a high number of unpaired electrons, E.g. $[CoF_6]^{3-}$



Owing to the octahedral orientation of six sp^3d^2 hybridized orbitals, shape of $[CoF_6]^{3-}$ complex ion is octahedral.

As it possesses four unpaired electrons in the 3d orbital, $[CoF_6]^{3-}$ ion is paramagnetic.

Some other examples are [FeF₆]³⁻, [Fe (NH₃)₆]²⁺, [Ni (NH₃)₆]²⁺, [Cu (NH₃)₆]²⁺, [Cr (H₂O)₆]³⁺, etc.

Limitations of valence bond theory: Even though the valence bond theory explains the formation, structures and magnetic behavior of coordination compounds to a larger extent, it suffers from the following short comings:

- It includes a number of assumptions.
- It fails to provide quantitative interpretation of magnetic data.
- It lacks explanation to the color exhibited by coordination compounds.
- It does not provide a quantitative interpretation of the thermodynamic or kinetic stabilities of coordination compounds.
- It is unable to predict the tetrahedral and planar structures of 4-coordinate complexes accurately.
- This theory does not distinguish between weak and strong ligands in compounds.

7.3 Crystal Field Splitting Theory

The important terms in Crystal Field Splitting theory are as follows:

- (a) **Degenerate orbitals:** in free state, all the d-orbitals (viz., $d_{xy'} d_{yz'} d_{xz'} d_{x^2-y^2}$ and d_{z^2}) will possess the same energy and are said to be degenerate.
- (b) t_{2g} and e_g set of orbitals: In a d-subshell, there are five d-orbitals and on the basis of orientation of lobes of these five d-orbitals with respect to coordinates, they have been grouped into two sets.
 - (i) e_g set of orbital: $d_{x^2-y^2}$, and orbitals have their lobes arranged along the axes and they constitute e_g set. These orbitals are also called axial orbital. Term e_g refers to 'doubly Degenerate', according to group theory (e = doubly degenerate set)
 - (ii) t_{2g} set of orbital: This set includes orbitals whose lobes lie between the axes and this set includes $d_{xy'}$ d_{yz} and d_{xz} orbitals. These orbitals are also known nonaxial orbitals. Group theory called these orbital t_{2g} where 't' refers to 'triply degenerate'.

Crystal Field Theory: The crystal field splitting theory (CFT) is an electrostatic model which considers the metalligand bond to be ionic occurring purely due to the electrostatic interaction between the metal ion and the ligand. Ligands are treated as point charges in case of anions and dipoles in case of neutral molecules. The five d orbitals in an isolated gaseous metal atom/ion have the same energy, i.e. they are degenerate. This degeneracy is maintained if a spherically symmetrical field of negative charges surrounds the metal atom/ion. However, when this negative field is resulted by the ligands (either anions or the negative ends of dipolar molecules like NH₃ and H₂O) in a complex, it becomes asymmetrical and the degeneracy of the d orbital is lifted. It results in splitting of the d orbitals. The pattern of splitting depends upon the nature of the crystal field. Let us discuss this splitting in different crystal fields in detail.

(a) **Crystal field splitting in octahedral field:** The orientation of d-orbital in octahedral field is represented in the diagram.

The lobes of $t_{2g'}$ set of orbital $(d_{xy'} d_{yz}$ and $d_{xz})$ point in between x, y and z axes while lobes of eg set $(d_{z'})$ and $d_{x'-y'}$ point along the x, y and z axes. Thus, energy of the eg set increases higher than that of the t_{2g} set. The splitting of orbital can be represented by Fig. 4:



Figure 28.4: Splitting of d-orbital is a octahedral crystal field

The difference in energy of t_{2g} and e_g set is known as crystal field splitting energy or crystal field stabilization energy (CFSE), which is represented by Δ_o (o stands for octahedral) or 10 Dq. The value of 10 Dq or Δ_o can be measured by UV-visible spectrum.

The crystal field splitting, $\Delta_{o'}$ depends upon the field produced by the ligand and the charge on the metal ion. Some ligands are able to produce strong fields, and correspondingly, the splitting will be large whereas others produce weak fields and these consequently result in small splitting of d orbital. Ligands can be arranged according to their order of increasing field strength as follows:

$I^{-} < Br^{-} < SCN^{-} < CI^{-} < S^{2-} < F^{-} < OH^{-} < C_{2}O_{4}^{-2-} < H_{2}O < NCH^{-} < edta^{4-} < NH_{3} < en < CN^{-} < CO^{-}$

This series is known as the spectrochemical series. It is an experimentally determined series based on the absorption of light by complex compounds with various ligands. Let us assign electrons in the d orbital of the metal ion in octahedral coordination entities. Obviously, the single d electron occupies one of the lower energy t_{2g} orbital. In d² and d³ coordination compounds, the d electrons occupy the t_{2g} orbital singly in accordance with the Hund's rule. For d⁴ ions, two possible patterns of electron distribution arise:

(a) The fourth electron could either enter the t_{2a} level and pair with an existing electron, or

(b) It could avoid paying the price of the pairing energy by occupying the eg level.

Either of these two possibilities depends on the relative magnitude of the crystal field splitting, Δ_{o} and the pairing energy, P (presents the energy required for electron pairing in a single orbital). The two options are:

- (i) If $\Delta_{o} < P$, the fourth electron enters one of the e_{g} orbital exhibiting the configuration $t_{2g}^{3}e_{g}^{1}$. Ligands for which $\Delta_{o} < P$ are known as weak field ligands, form high spin complexes.
- (ii) If $\Delta_o > P$, it becomes more energetically favorable for the fourth electron to occupy a t_{2g} orbital with configuration $t_{2g}^4 e_g^0$. Such ligands are known as strong field ligands and they form low spin complexes. It is observed from calculations that d⁴ to d² coordination entities are more stable for strong field cases compared to their weak counterparts.



Figure 28.5: Splitting of d-orbital is a octahedral crystal field

(b) Crystal field splitting in tetrahedral complexes: The orientation of ligands in a tetrahedral complex is given in fig. 5. Although none of the d-orbitals point towards axes, the t_{2g} set is close to the direction in which ligands are approaching so their energy is higher.

The magnitude of Δ , is considerably less than that in the octahedral field, which is mainly due to two reasons:

- (i) In tetrahedral complex, number of the ligands is only four instead of six.
- (ii) In tetrahedral complexes, the direction of the orbitals does not coincide with the direction of the ligands, both the factors reduce the CFSE by 2/3 and so Δ_{t} is roughly 4/9 times to Δ_{o} .
- (c) Crystal Field Splitting in Square Planar Complexes: The square planar geometry can be considered to be derived from the octahedral by removing negative charges from the z-axis. As these negative charges are removed, d_{xv} d_{xz} and d_{vz} orbital, all of which have a Z-component become more stable as shown in Fig. below.

This type of splitting can be further explained as follows:

As the lobes of point towards the ligands, this orbital has the highest energy. The lobes of dxy orbital lie between the ligands but are coplanar with them, hence this orbital has the second highest energy. The lobes of d_{x^2} orbital point out of the plane of the complex but the belt around the center of the orbital (which contains about 1/3rd of the electron density) lies in the plane. Therefore, d_{z^2} orbital is next highest in energy. The lobes of dxz and dyz orbital point out of the plane of the complex, and so they are least affected by the electrostatic field of the ligands, they degenerate and have the lowest in energy.



Figure 28.6: Splitting of d-orbital in a square planar crystal

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Weak ligands favor high spin complexes because they cannot pair up the electrons against Hund's rule while strong ligands favor low spin complexes.

Vaibhav Krishnan (JEE 2009, AIR 22)

Illustration 11: $Mn_{(aq.)}^{2+}$ ion is light pink colored while $[Mn (CN)_6]^{4-}$ is blue in color. Explain. (JEE ADVANCED)

Sol: In complexes, where Mn (II) is present, configuration of a metal ion is d⁵. There may be two types of spin arrangements in the presence of different kinds of ligands.

(A) High spin complex (with weak field ligands) and (B) Low spin complex (with strong field ligands)

The arrangement of electrons in these complexes can be depicted as:



In high spin complex compounds, it is observed that d–d transition requires reversion of spin which is against the spin selection rules and this makes them spin forbidden and the intensity of color is of only about 1/100 when the transition is allowed.

In $[Mn (CN)_6]^4$ on the other hand, d–d transitions do not have any such restrictions and are spin allowed. Intense color also is observed when transition takes place.

7.3.1 Factors Affecting CFSE

(a) Nature of ligand: The value of Δ depends upon the nature of ligands. Ligands with a small degree of crystal field splitting capacity are termed as weak field ligands and those ligands which cause large splitting are called strong field ligands. In general, ligands can be arranged in the ascending order of CFSE caused by them. This series remains practically constant for different metals and is known as spectrochemical series. It is an experimentally determined series. The order is difficult to explain due to involvement of both σ and π bonding. Some ligands in spectrochemical series are given below:

 $I^- < Br^- < S^{2-} < Cl^- < N_3^-$, $F^- < Urea$, $OH^- < Oxalate$, $O^{2-} < H_2O < NCS^- < EDTA < py$, $NH_3 < en = SO_3^{-2-} < bipy$, phen $< NO_2^- < CH_3^- < C_6H_5^- < CN^- < CO$. For strong field ligands, the order depends on the donor atom and is in the following order:

C-donor > N-donor > O-donor > Halogen donor

- (b) Geometry of the Complex: Δ_t is approximately 4/9 times of Δ° . The lower value of Δ_t is due to lesser number of ligands in tetrahedral complex. Also, in tetrahedral complexes the orbital does not point toward the axes, resulting in less interaction.
- (c) Oxidation state of metal ion: It is observed that the higher the charge on the central metal atom (or oxidation state), the higher the CFSE.

E.g., Δ_{o} for [Fe (H₂O)₆]³⁺ is greater than [Fe (H₂O)₆]²⁺, Δ_{o} for [Co (H₂O)₆]³⁺ is greater than [Co (H₂O)₆]²⁺ and Δ_{o} for [V (H₂O)₆]²⁺ is greater than [Cr (H₂O)₆]³⁺.

(d) Nature of metal ion: The value of CFSE is also determined by the transition series to which the metal belongs and the order for this is observed to be 3d < 4d < 5d. The value of Δ increases by 30% to 50% for 3d to 4d series and from 4d to 5d series. Hence, metals of 4d and 5d series have more tendency to form low spin complexes, e.g. CFSE for the given complexes follow the order:

 $[Co (NH_3)_6]^{3+} < [Rh (NH_3)_6]^{3+} < [Ir (NH_3)_6]^{3+}]$

When two metal ions possess the same charge but different number of d-electrons, the magnitude of Δ_o decreases with increase in the number of d-electrons in the central metal atom. E.g. Δ_o for [Co (H₂O)₆]²⁺ is greater than Δ_o for [Ni (H₂O)₆]²⁺ because Co²⁺ possesses 3d⁷ configuration while Ni²⁺ has 3d⁸ configuration.

7.3.2 Applications of CFSE

(a) Magnetic character of complexes: Complexes containing unpaired electrons tend to be attracted by magnetic fields and hence known as paramagnetic. In contrast, when all the electrons are paired, the complex is slightly repelled by a magnetic field and is said to be diamagnetic. The





magnetic moment of a transition metal wholly depends on the number of unpaired electrons and is equal to $\sqrt{n(n+2)}$ B.M., where n is number

of unpaired electrons. For diamagnetic substance, the magnetic moment will be zero.

Magnetic moments of coordination compounds can be experimentally determined and this data provide information to examine the nature of coordination entities further. These measurements are termed as magnetic susceptibility measurements. For example, [Fe $(H_2O)_6$]²⁺ is paramagnetic while [Fe $(CN)_6$]⁴⁻ is diamagnetic. This observation can be explained on the basis of the electronic configurations of Fe²⁺ in [Fe $(H_2O)_6$]²⁺ and [Fe $(CN)_6$]⁴⁻. H₂O is a weak field ligand while CN⁻ is a strong field ligand. So [Fe $(CN)_6$]⁴⁻ is the inner orbital low spin complex whereas [Fe $(H_2O)_6$]²⁺ is an outer-orbital high spin complex. The configurations of Fe²⁺ in both the compounds are further explained in the diagram here.

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Complexes possessing d⁰ or d¹⁰ configuration of a metal ion are always diamagnetic.

Nikhil Khandelwal (JEE 2009, AIR 94)

(b) Colour of complexes: In many complexes the d-orbital split takes place in the two sets t_{2a} and e_{a} , which

possess different energies. The difference in energies of t_{2g} and and e_g lies in the visible region of the spectrum and this helps transition metal complexes to absorb color. This makes them colored complementary to the color absorbed. This transition involves t_{2g} and e_g sets of d-orbital and is called as d–d transition. Thus d–d transition is responsible for the color of transition metal complexes. d–d transition can be represented diagrammatically as shown here.



MASTERJEE CONCEPTS

Complexes like CrO_4^{2-} , $Cr_2O_7^{2-}$, and MnO_4^{-} , etc. have d⁰ configuration of the metal ion but still exhibit intense color. Here the color is caused by the charge transfer spectra (CT) and not by the d–d transition.

Saurabh Gupta (JEE 2010, AIR 443)

7.3.3 Stability of Complexes

Complexes normally exhibit two kinds of stabilities: (i) Thermodynamic stability and (ii) Kinetic stability. Thermodynamic stability deals with the metal–ligand bond energy, stability constants, redox potentials, etc., that affect the equilibrium. On the basis of thermodynamic stability, Blitz classified the complexes into stable or penetration complexes and unstable or normal complexes.

Kinetic stability deals with the rates of reaction of complexes in a solution. On the basis of kinetic stability, Taube classified the complexes into labile and inert complexes. Ligands of labile complexes are easily replaceable while ligands of inert complexes cannot be replaced with easily.

Chelate effect: Complexes containing chelate rings are more stable, e.g. $[Ni (NH_3)_6]^{2+}$ and so is less stable than $[Ni (en)_3]^{2+}$.

Macrocylic effect: When a multidentate ligand is cyclic without any considerable steric effect, then the complex formed is more stable than acyclic ligand. This phenomenon is called the macrocyclic effect.

Illustration 12: $[Cu (CN)_4]^{2-}$ is a more stable complex than $[Cu (NH_3)_4]^{2+}$. Why? (JEE MAIN)

Sol: The higher stability constant $K = 2 \times 10^{27}$ for $Cu^{2+} + 4CN^{-} \rightarrow [Cu (CN)_4]^{2-}$ than for $[Cu (NH_3)_4]^{2+}$ (which is 4.5 × 10¹¹) explains stability. Also CN^{-} is stronger ligand than NH_3 .

8. ORGANOMETALLIC COMPOUNDS

Organometallic compounds are defined as compounds in which carbon forms a bond with an atom (metal/non-metal) which is less electronegative than carbon.

These compounds are classified into two – covalently bonded compounds and ionic organometallic compounds.

Covalently bonded compounds: In covalently bonded compounds, the metal and carbon atoms are attached to each other by a covalent bond. They can be further classified into: three groups:

(i) σ (sigma) bonded complexes: A σ -bonded complex consists of a metal atom and a carbon atom of the ligand joined together with a σ bond. In another words, the ligand contributes one electron and is called one electron donor. Tetramethyltin, (CH₃)₄Sn and trimethyl aluminum, (CH₃)₃ Al are examples of σ -bonded organometallic compounds. The latter exists as dimmer and has



a structure analogous to diborane. In this, two methyl groups bridge between two aluminum atoms.

(ii) π complexes: Organometallic compounds with π -bonds present in them are called π -complexes. Zeise's salt, ferrocene and dibenzene chromium are π -complexes. In these compounds, the π electrons interact with the metal ion and occupy one of the coordination sites. For example, in ferrocene and dibenzene chromium, the iron and chromium atoms are sandwiched between two aromatic rings.



The number of carbon atoms taking part in the

formation of π -complexes is indicated by the power of η^x (pronounced as eta). For example, ferrocene is represented as $[\text{Fe}\eta^5-\text{C}_5\text{H}_5)_2]$ indicating that five carbon atoms or cyclopentadienyl anion are involved in the π - complication with the metal. Similarly, one can write dibenzene chromium as $[\text{Cr} (\eta^6-\text{C}_6\text{H}_6)_2]$ indicating that all the six carbons of benzene are involved in π -complexation with chromium.

8.1 Bonding in Organometallic Compounds

Bonding in Metal Carbonyls: The metal–carbon bond in metal carbonyls exhibits σ as well as π characteristics.

(i) σ -overlap: The lone pair of electron is present on the bonding orbital of carbon monoxide in a σ bonded complex and it interacts with the empty d-orbital of the metal to form a metal–carbon bond.



(ii) π -overlap: Besides, the antibonding orbitals of CO also overlaps with the filled d-orbital of the metal resulting in back bonding as previously explained. Thus metal carbonyls become much more stable due to this multiple bonding.

It is important to note that the σ -bond is positioned in the nodal plane of the σ -electrons whereas π -overlap is perpendicular to the nodal plane.



Metal orbital

Anibonding orbital of carbon mono-oxide

Backbonding metal corbonyl

Bonding of Alkenes to a Transition Metal: There are two components in the bonding of alkenes to a transition metal to form complexes. First, the σ -electron density of the alkene overlaps with a π -type vacant orbital of the metal atom. Second is the backbonding resulting from the flow of electron density from a filled d-orbital on the metal into the vacant σ -antibonding molecular orbital on the carbon atom as depicted in the following diagram:



MASTERJEE CONCEPTS

As the electron density on metal atom increases, strength of backbonding from the metal to carbon increases and the metal–carbon bond length decreases. Likewise, when C–O bond order decreases, C–O bond length increases and vice versa.

Neeraj Toshniwal (JEE 2009, AIR 21)

8.2 Synthesis of Organometallic Compounds

Some important methods to generate metal-carbon bond as follows:

By the direct reaction of metals:

(a) n-Butyl lithium is prepared by the reaction of n-butyl bromide with lithium in ether.

$$\begin{array}{c} n-C_{4}H_{9}Br+2Li \xrightarrow{ \ \ E ther \ \ } n-C_{4}H_{9}Li+2LiBr\\ n-Butyl \ \ bromide \ \ n-Butyl \ \ lithium \end{array}$$

(b) Likewise, tetra ethyl lead can be prepared as follows:

$$4C_2H_5CI + 4NaPb_{\text{Sodium-lead}} \longrightarrow (C_2H_5)_4Pb + 4NaCI + 3Pb_{\text{Tetra-ethyl lead}}$$

(c) Grignard reagents are obtained by the reaction of alkyl halide (in ether) with magnesium:



By using an alkylating agent: Grignard reagent and alkyl lithium or reaction with most of the metal and non-metal halides in the presence of ether as solvent yield other organometallic compounds.

 $PCI_{3} + 3C_{6}H_{5}MgCI \xrightarrow{Ether} P(C_{6}H_{5})_{3} + 3MgCI_{2}$ Triphenyl phosphine

 $SnCl_{_4} + 4n - C_{_4}H_{_9}Li \longrightarrow (n - C_{_4}H_{_9})Sn + 4LiCl$

Preparation of Metal Carbonyls:

(a) Nickel carbonyl is obtained when finely divided nickel reacts with CO at room temperature.

 $Ni + 4CO \longrightarrow Ni(CO)_{4(g)} \xrightarrow{\Lambda} Ni + 4CO$

The nickel carbonyl so formed is in gaseous state and its subsequent thermal decomposition gives Ni and CO. This principle is used in the purification of Ni via Mond's process.

(b) Iron carbonyl is formed when iron reacts with CO at high pressure and temperature.

 $Fe+5CO \longrightarrow [Fe(CO)_5]$

MASTERJEE CONCEPTS

Carbonyls containing only metal and CO are known as homoleptic carbonyls.

Nikhil Khandelwal (JEE 2009, AIR 94)

POINTS TO REMBEMBER

Coordination compound	A compound containing central metal atom or ion bonded to a fixed number of ions or molecules (called ligands).
Ligands	The ions or molecules which donate a pair of electrons to the central metal atom or ion and form coordinate bonds.
Coordination sphere	The central metals atom or ion and the ligands attached to it is collectively called coordination sphere.
Coordination number	The number of coordinate bonds formed by the ligands with the metal atom i.e. number of unidentate ligands or double the number of bidentate ligands, etc.
Denticity	The number of coordinating or ligating atoms preset in a ligand
Chelation	When a bidentate or a polydentate ligand uses it two or more donor atoms to bind to the central metal atom or ion forming ring structure, it is called chelation
Coordination polyhedron	The spatial arrangement of the ligands which are directly attached to the central metal atom.
Homoleptic complexes	The complexes which contain only one type of ligands.
Structural isomers	The isomers having same molecular formula but different structural arrangement of atoms or group of atoms around the central metal ion.
Stereiosomers	The isomers which have the same position of atom or group of atoms but they differ in the spatial arrangement around the central metal atom.
Crystal field splitting	The conversion of five degenerate d-orbitals of the metal ion into different sets of orbitals having different energies in the presence of electrical field of ligands is called crystal field splitting.
Spectrochemical series	The arrangement of ligands in the increasing order of crystal field splitting