# **Coordination Compounds**

### Addition Compounds :

They are formed by the combination of two or more stable compounds in stoichiometric ratio.



### Double salts :

Those addition compounds which lose their identity in solution are called double salts.

$$\begin{split} & \mathsf{K}_2\mathsf{SO}_4 + \mathsf{Al}_2(\mathsf{SO}_4)_3 + 24\mathsf{H}_2\mathsf{O} \rightarrow \mathsf{K}_2\mathsf{SO}_4 \ . \ \mathsf{Al}_2(\mathsf{SO}_4)_3 \ . \ 24\mathsf{H}_2\mathsf{O} \underbrace{\overset{(\mathrm{aq.})}{\longleftarrow}} 2\mathsf{K}^+ \ (\mathrm{aq.}) + 2\mathsf{Al}^{+3} \ (\mathrm{aq.}) + 4\mathsf{SO}_4^{\ 2^-} \ (\mathrm{aq.}) \\ & \mathsf{Other examples are carnallite} \ (\mathsf{KCl.} \ \mathsf{MgCl}_2 \ . \ \mathsf{6H}_2\mathsf{O}), \ \mathsf{Mohr's salt} \ [\mathsf{FeSO}_4 \ . \ (\mathsf{NH}_4)_2\mathsf{SO}_4 \ . \ \mathsf{6H}_2\mathsf{O}], \\ & \mathsf{Potash alum} \ [\mathsf{KAl}(\mathsf{SO}_4)_2 \ . \ 12\mathsf{H}_2\mathsf{O}] \ \mathsf{etc.} \end{split}$$

## **Coordination Compounds :**

Those addition compounds which retain their identity (i.e. doesn't lose their identity) in solution are called coordination compounds.

 $Fe(CN)_2 + 4KCN \longrightarrow Fe(CN)_2$ . 4KCN or  $K_4 [Fe(CN)_6]$  (aq.)  $\implies 4K^+$  (aq.) +  $[Fe(CN)_6]^{4-}$ (aq.)

Other examples are,  $[Cu(NH_3)_4]SO_4$  (aq.)  $\equiv$   $[Cu(NH_3)_4]^{2+}$  (aq.) +  $SO_4^{2-}$  (aq.)

 $K_2[Zn(CN)_4]$  (aq.)  $\implies 2K^+$  (aq.) +  $[Zn(CN)_4]^{2-}$  (aq.)

Coordination compound is defined as a species in which metal atom or ion is attached to group of neutral molecules / ions by coordinate covalent bonds.

## Solved Examples.

**Ex.1** Explain that  $Fe(NH_4)_2(SO_4)_2$  is a double salt but  $K_3Fe(CN)_6$  is a complex.

**Sol.**  $Fe(NH_4)_2(SO_4)_2$  is a double salt because its solution gives tests of  $Fe^{2+}$ ,  $NH_4^+$  and  $SO_4^{2-}$ .  $K_3Fe(CN)_6$  is not a double salt but it is a complex because the solution of this compound renders  $K^+$  and  $[Fe(CN)_6]^{3-}$  ions and further  $[Fe(CN)_6]^{3-}$  does not give the tests of  $Fe^{3+}$  and  $CN^-$ .

## **Coordination Entity/Coordination Sphere :**

A coordination entity constitutes a central atom/ion, usually of a metal, to which are attached a fixed number of other atoms or groups each of which is called a ligand. It may be neutral or charged. Examples being :  $[Co(NH_3)_6]^{3+}$ ,  $[PtCl_4]^{2-}$ ,  $[Fe(CN)_6]^{3-}$ ,  $[NiCl_2(OH_2)_4]$  and  $(NH_3)$ , (CI-), (CN-),  $(H_2O)$  are the ligands.

The central atom/ion and the ligands attached to it are enclosed in square bracket and is collectively called as coordination sphere.

**Note :** The remaining ions apart from complex ions i.e. outside the coordination sphere are called counter ions, free ions or ionisable ions. For example, in  $K_4[Fe(CN)_6]$ , the potassium (K<sup>+</sup>) ion is counter ion of coordination entity  $[Fe(CN)_6]^4$ .

## Central Atom/Ion :

In a coordination entity-the atom/ion to which are bound a fixed number of ligands in a definite geometrical arrangement around it, is called the central atom or ion. For example, the central atom/ion in the coordination entities :  $[NiCl_2(OH_2)_4]$ ,  $[CoCl(NH_3)_5]^{2+}$  and  $[Fe(CN)_6]^{3-}$  are Ni<sup>2+</sup>, Co<sup>3+</sup> and Fe<sup>3+</sup>, respectively. These central atoms/ions are also referred to as Lewis acids.

### Ligands :

The neutral molecules, anions or cations which are directly linked with central metal atom or ion in the coordination entity are called ligands.

These may be simple ions such as  $Br^-$ , small molecules such as  $H_2O$  or  $NH_3$ , larger molecules such as  $H_2NCH_2CH_2NH_2$  or  $N(CH_2CH_2NH_2)_3$  or even macromolecules such as proteins.

When a ligand is attached to a metal atom ion through a single donor atom, as with Cl-,  $H_2O$  or NH<sub>3</sub>, the ligand is said to be unidentate. Similarly when a ligand is bound through two donor atoms, as in  $H_2NCH_2CH_2NH_2$  (ethane-1, 2-diamine) or  $C_2O_4^{2-}$  (oxalate), the ligand is said to be bidentate and when several donor atoms are present in a single ligand as in N (CH<sub>2</sub>CH<sub>2</sub>NH<sub>2</sub>)<sub>3</sub> or ethylenediaminetetraacetic acid (EDTA), the ligand is said to be polydentate.

## Ambidentate Ligand :

Ligands which can ligate through two different atoms present in it are called ambidentate ligands. Examples of such ligands are the  $CN^-$ ,  $NO_2^-$  and  $SCN^-$  ions.  $NO_2^-$  ion can coordinate through either the nitrogen or the oxygen atoms to a central metal atom/ion. Similarly,  $SCN^-$  ion can coordinate through the sulphur or nitrogen atom. Such possibilities give rise to linkage isomerism in coordination compounds. For example,

$M \leftarrow N \leq_{O}^{O}$	nitrito-N
M ← O—N=O	nitrito-O
$M \leftarrow SCN$	thiocyanato or thiocyanato-S
$M \leftarrow NCS$	isothiocyanato or thiocyanato-N

#### Chelate ligand :

Chelate ligand is a di or polydentate ligand which uses its two or more donor atoms to bind a single metal ion producing a ring. The complex formed is referred to as a chelate complex and the process of chelate formation is called chelation. The number of such ligating groups is called the denticity of the ligand.

## Denticity and Chelation :

	IUPAC Name	Formula
triphenyl phosphine	triphenyl phosphine/triphenyl phosphane	PPh <sub>3</sub>
pyridine	pyridine	C <sub>5</sub> H <sub>5</sub> N (py)
ammonia	ammine	NH <sub>3</sub>
water	aqua or aquo	H <sub>2</sub> O
carbonyl	carbonyl	СО
nitrosyl	nitrosyl	NO
fluoro	fluoro or fluorido*	F <sup>-</sup>
chloro	chloro or chlorido*	Cl⁻
bromo	bromo or bromido*	Br⁻
iodo	iodo or iodido*	I_
cyano	cyanido or cyanido-C* (C-bonded)	CN⁻
isocyano	isocyanido or cyanido-N* (N-bonded)	NC <sup>-</sup>
thiocyano	thiocyanato-S(S-bonded)	SCN⁻
isothiocyano	thiocyanato-N(N-bonded)	NCS <sup>-</sup>
hydroxo	hydroxo or hydroxido*	OH⁻
nitro	nitrito–N (N–bonded)	NO <sub>2</sub> <sup>-</sup>
nitrito	nitrito–O (O–bonded)	ONO <sup>-</sup>
nitrate	nitrato	NO <sub>3</sub> <sup>-</sup>
amido	amido	$NH_2^-$
nitride	nitrido	N <sup>3–</sup>
azido	azido	$N_3^-$
hydride	hydrido	H⁻
oxide	oxido	O <sup>2-</sup>
peroxide	peroxido	02 <sup>2-</sup>
superoxide	superoxido	0 <sub>2</sub> <sup>-</sup>
sulphate	sulphato	SO4 <sup>2-</sup>
sulphide	sulphido or thio	S <sup>2-</sup>
nitrosylium	nitrosylium or nitrosonium	NO <sup>+</sup>
nitronium	nitronium	NO <sub>2</sub> <sup>+</sup>

Table : 1Common Monodentate Ligands

Chelating Points	Common Name	IUPAC Name	Abbreviation	Formula
bidentate	ethylenediamine	1,2-ethanediamine/ ethane-1,2-diamine	en	NH <sub>2</sub> ,CH <sub>2</sub> CH <sub>2</sub> NH <sub>2</sub>
bidentate	propanediamine	1,2-propanediamine	pn	NH <sub>2</sub> -CH-CH <sub>2</sub> -NH <sub>2</sub>   CH <sub>3</sub>
tridentate	diethylenetriamine	[N-(2-aminoethyl)-1 2-ethanediamine or diethylenetriamine	dien	NH <sub>2</sub> CH <sub>2</sub> CH <sub>2</sub> NHCH <sub>2</sub> CH <sub>2</sub> NH <sub>2</sub>
tetradenate	triethylenetetraamine	[N, N'-bis-(2-aminoethyl)-1, 2-ethanediamine	trien	NH2CH2CH2NHCH2CH2NHCH2CH2NH2
	triaminotriethylamine	or triethylenetetraamine , , -tris(2-aminoe- thyl) amine.	tren	NH <sub>2</sub> CH <sub>2</sub> CH <sub>2</sub> NCH <sub>2</sub> CH <sub>2</sub> NH <sub>2</sub>   CH <sub>2</sub> CH <sub>2</sub> NH <sub>2</sub>
pentadentate	tetraethylenepentaamine	1,4,7,10 pentaazatridecan or tetraethylenepentaamin	าe าe	NH <sub>2</sub> CH <sub>2</sub> CH <sub>2</sub> NHCH <sub>2</sub> CH <sub>2</sub> NHCH <sub>2</sub> CH <sub>2</sub> NHCH <sub>2</sub> CH <sub>2</sub> NH <sub>2</sub>
hexadentate	ethylenediaminetetraacetate	1,2-ethanediyl (dinitrilo) tetraacetate or ethylenediaminetetraa	EDTA	OOCH <sub>2</sub> C NCH <sub>2</sub> CH <sub>2</sub> N CH <sub>2</sub> COO <sup>-</sup> CH <sub>2</sub> COO <sup>-</sup>
bidentate	oxalato		ох	

Table : 2Common Chelating Amines

## Coordination Number :

The coordination number of the central atom/ion is determined by the number of sigma bonds between the ligands and the central atom/ions i.e. the number of ligand donor atoms to which the metal is directly attached. Pi-bonds, if any, between the ligating atom and the central atom/ion are not considered for the determination of coordination number. The sigma bonding electrons may be indicated by a pair of dots, preceding the donor atom in the ligand formula as in :  $[Co(NH_{2})_{2}]^{3+}$ ,  $[Fe(CN)_{6}]^{3-}$ ,  $[Ni(CO)_{4}]$ ,  $[Co(CI_{4})]^{2-}$ .

Some common co-ordination number of important metals are as given below.

Metal	C.N.	Metal	C.N.
Cu⁺	2, 4	Ni <sup>2+</sup>	4, 6
Ag⁺	2	Fe <sup>2+</sup>	6
Au⁺	2, 4	Fe <sup>3+</sup>	6
Cu <sup>2+</sup>	4, 6	Co <sup>2+</sup>	6
Ag <sup>2+</sup>	4	Co <sup>3+</sup>	6
Pt <sup>2+</sup>	4	Pt <sup>4+</sup>	6
Pd <sup>2+</sup>	4	Pd <sup>4+</sup>	6

## **Coordination Polyhedron :**

The spatial arrangement of the ligand atoms which are directly attached to the central atom/ion defines a coordination polyhedron about the central atom. Figure below shows the shapes of tetrahedral, square planar, octahedral, square pyramidal and trigonal bipyramidal coordination polyhedra.  $[Co(NH_3)_6]^{3+}$  has an octahedral geometry, while  $[PtCl_4]^{2-}$  and  $Ni(CO)_4$ , are square planar and tetrahedral, respectively.

## **Oxidation number of Central Atom :**

The oxidation number of the central atom is defined as the charge it would carry if all the ligands are removed along with the electron pairs that are shared with the central atom. Metal oxidation number is represented by a Roman numeral in parentheses following the name of the coordination entity. For example oxidation number of iron in  $[Fe(CN)_{e}]^{3-}$  is +3 and it is written as Fe(III).

#### Homoleptic and heteroleptic complexes

Complexes in which a metal is bound to only one type of donor groups, e.g.,  $[Cr(NH_3)_6]^{3+}$ , are known as homoleptic. Complexes in which a metal is bound to more than one type of donor groups. e.g.,  $[Co(NH_3)_4Br_2]^+$ , are known as heteroleptic.

## -Solved Examples

- **Ex.2** Give the coordination entities and counter ions in the coordination compounds :  $[Cr(NH_3)_6]Cl_3$ ;  $K_4[Fe(CN)_6]$ ,  $K_2[PtCl_4]$ ;  $[Ni(CO)_4]$ ;  $K_2[Ni(CN)_4]$ .
- Sol. The respective coordination entities are : [Cr(NH<sub>3</sub>)<sub>6</sub>]<sup>3+</sup>; [Fe(CN)<sub>6</sub>]<sup>4-</sup>; [PtCl<sub>4</sub>]<sup>2-</sup>; [Ni(CO)<sub>4</sub>]; and the counter ions are Cl<sup>-</sup>, K<sup>+</sup>, K<sup>+</sup>, (no counter ion) and K<sup>+</sup>, respectively in the given coordination compounds.
- **Ex.3** Explain why  $[Fe(en)_3)^{3+}$  is less stable than  $[Fe(EDTA)]^-$ ?
- **Sol.** [Fe(en)<sub>3</sub>)<sup>3+</sup> has three, five membered rings in the structure. On the other hand [Fe(EDTA)]<sup>-</sup> has five, six membered ring in the structure.



The five membered rings are less stable than six membered rings ; so  $[Fe(en)_3)^{3+}$  is less stable than  $[Fe(EDTA)]^{-}$ 

**Ex.4** Draw the structure of the complexes  $[Fe(C_2O_4)_3]^{3-}$  and  $[Pt(en)_2]^{2+}$ . Determine the dentisity of the ligands and identify any chelate rings. What are the coordination number and the oxidation number of the central metal ion ?



The ligands, oxalate and ethylenediamine are bidentate as each ligand has to donor atoms. So in 1st case the number of chelate rings (five membered) are three where as in 2nd case the number of chelate rings (five membered) are two.

The coordination number and oxidation state of iron are six and +3 respectively and the coordination number and oxidation state of platinum are four and +2 respectively.

## **Nomenclature of Coordination Compounds**

Writing the formulas of Mononuclear Coordination Entities :

The following rules are followed while writing the formulas :

- (i) The central atom is placed first.
- (ii) The ligands are then placed in alphabetical order. The placement of a ligand in the list does not depend on its charge.
- (iii) Polydentate ligands are also placed alphabetically. In case of abbreviated ligand, the first letter of the abbreviation is used to determine the position of the ligand in the alphabetical order.
- (iv) The formula for the entire coordination entity, whether charged or not, is enclosed in square brackets. When ligands are polyatomic, their formulas are enclosed in parentheses. Ligands abbreviations are also enclosed in parentheses.
- (v) There should be no space between the ligands and the metal within a coordination sphere.
- (vi) When the formula of a charged coordination entity is to be written without that of the counter ion, the charge is indicated outside the square brackets as a right superscript with the number before the sign. For example, [Co(H<sub>2</sub>O)<sub>6</sub>]<sup>3+</sup>, [Fe(CN)<sub>6</sub>]<sup>3-</sup> etc.
- (vii) The charge of the cation(s) is balanced by the charge of the anion(s).

#### Writing the name of Mononuclear Coordination Compounds :

The following rules are followed when naming coordination compounds :

(i) Like simple salts the cation is named first in both positively and negatively charged coordination entities.

**Examples** : [Ag(NH<sub>3</sub>)<sub>2</sub>]Cl, diamminesilver(l)chloride. K<sub>3</sub>[Fe(CN)<sub>6</sub>],potassium hexacyanidoferrate(III).

(ii) The ligands are named in an alphabetical order (according to the name of ligand, not the prefix) before the name of the central atom/ion.

#### Examples :

 $[Pt(NH_3)BrCl(CH_3NH_2)]$ , amminebromidochloridomethylamineplatinum(II).  $[Co(H_2O)_2(ox)_2]^-$ , diaquabis(oxalato)cobaltate(III).

(iii) Names of the anionic ligands end in –o and those of neutral ligands are the same except aqua for  $H_2O$ , ammine for  $NH_3$ , carbonyl for CO, thiocarbonyl for CS and nitrosyl for NO. But names of cationic ligands end in–ium. The neutral an cationic are placed within enclosing marks ().

Some more important examples of neutral and cationic ligands are :

tetraphosphorus	—	$P_4$
dioxygen		O <sub>2</sub>
octasulphur		$S_8$

(iv) Prefixes mono, di, tri, etc., are used to indicate the number of the one kind of ligands in the coordination entity. When the names of the ligands include a numerical prefix are complicated or whenever the use of normal prefixes creates some confusion, it is set off in parentheses and the second set of prefixes is used.

di	bis
tri	tris
tetra	tetrakis
penta	pentakis
hexa	hexakis
hepta	heptakis
	di tri tetra penta hexa hepta

**Examples**;  $[CoCl_2(NH_2CH_2NH_2)_2]^+$ , dichloridobis(ethane-1,2-diamine)cobalt(III).

 $[NiCl_2(PPh_3)_2]$ , dichloridobis(triphenylphosphine)nickel(II).

(v) Oxidation state of the metal in cation, anion or neutral coordination entity is indicated by Roman numeral in the parentheses after the name of metal.

(vi) If 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. If the complex ion is an anion, the name of the metal ends with the suffix - ate. For example, Co in a complex anion,  $[Co(SCN)_4]^{2-}$  is called cobaltate. For some metals, the Latin names are used in the complex anions.

iron (Fe)	ferrate	lead (Pb)	plumbate
silver (Ag)	argentate	tin (Sn)	stannate
gold (Au)	aurate		

**Examples**; [Co(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>]<sup>+</sup>, pentaamminechloridocobalt(III).

 $(NH_4)_2$  [Co(SCN)<sub>4</sub>], ammonium tetrathiocyanato-S-cobaltate(II).

- (vii) The neutral complex molecule is named similar to that of the complex cation.
- **Example**; [CrCl<sub>3</sub>(py)<sub>3</sub>], trichloridotris(pyridine)chromium(III).
- (viii) The prefixes cis- and trans- designate adjacent and opposite geometric locations. For examples,

[Pt(NH<sub>2</sub>)<sub>2</sub>Cl<sub>2</sub>], cis- and trans-diamminedichloridoplatinum(II),

 $[CoCl_2(NH_3)_4]^+$ , cis- and trans-tetraamminedichloridocobalt(III).



## Solved Examples

Ex.5 Write the structural formula corresponding to each of the following IUPAC names :

- (a) tetraamminedichoridocobalt(III) chloride
- (b) dicvanidoargentate(I) ion (c) tris(ethane-1,2-diamine)copper(II) sulphate (d) sodium diaguatetrahydroxidoaluminate(III)

Sol. According to IUPAC nomenclature the structural formula are as follows :

(a) [Co <sup>III</sup> (NH <sub>3</sub> ) <sub>4</sub> Cl <sub>2</sub> ]Cl	(b)	[Ag <sup>I</sup> (CN) <sub>2</sub> ] <sup>−</sup>
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 $Na[AI^{III}(H_2O)_2(OH)_4]$ (c)  $[Cu^{II}(en)_3]SO_4$ (d)

Write IUPAC names of the following Ex.6

(a)	Na <sub>2</sub> [Fe(CN) <sub>5</sub> NO]	(b)	[Co(NH <sub>3</sub> ) <sub>4</sub> Cl(NCS)] <sup>+</sup>
(c)	K₄[FeO₄]	(d)	$K_2[Co(N_3)_4]$

According to IUPAC nomenclature the IUPAC names are as follows : Sol.

(a) sodium pentacyanidonitrosoniumferrate(II); It is sodium nitroprusside which is diamagnetic

(b) tetraamminechloridothiocynato-N-cobalt(III) ion

(c) potassium tetraoxidoferrate(IV) (d) potassium tetraazidocobaltate(II);  $N_3^-$  is azide.

## Effective Atomic Number Rule given by Sidgwick :

Effective Atomic Number (EAN) = No. of electron present on the metal atom/ion + No. of electrons donated by ligands to it.

OR

Effective Atomic Number (EAN) = Atomic no. of central metal - Oxidation state of central metal + No. of electrons donated by ligands.

The complexes in which the EAN of the central atom equals the atomic number of the next noble gas, are found to be extra stable.

Note: The EAN rule is generally found to be not valid in case of most of the complexes but in case of metal carbonyls this rule is found to be valid in all cases except one or two exceptions.

## -Solved Examples -

Determine the E.A.N. of the metal atom in the following : Ex.7 (a)  $K_{3}[Cr(C_{2}O_{4})_{3}]$ (b)  $K_4[Fe(CN)_6]$ 

Sol.	V⁻ (Z = 23)	∴ E.A.N. = 23 + 1 +	12 = 36 which is atomi	ic number of Kr.
Ans.	(3)			
	(1) $[Co(SCN)_4]^{2-}$	(2) [Cr(NH <sub>3</sub> ) <sub>6</sub> ] <sup>3+</sup>	(3) [V(CO) <sub>6</sub> ] <sup>-</sup>	(4) [Ni(Cl) <sub>4</sub> ] <sup>2–</sup>
Ex.8	In which of the follow	ving complexes, the eff	fective atomic number of	of the metal atom equals the atomic
	(b) $Fe^{2+}(Z = 26)$	∴ E.A.N. = 24 + 12	= 36 (more stable)	
Sol.	(a) Cr <sup>3+</sup> (Z = 24)	∴ E.A.N. = 21 + 6 ×	<pre>&lt; 2 = 33 (Less stable)</pre>	

#### Werner's Theory :

Werner in 1983 presented a theory known as Werner's coordination theory. More important postulates of this theory are :

Most element exhibit two types of valencies : (a) Primary valency and (b) Secondary valency.

#### (a) Primary valency :

This corresponds to oxidation state of the metal ion. This is also called principal, ionisable or ionic valency. It is satisfied by negative ions and its attachment with the central metal ion is shown by dotted lines.

#### (b) Secondary or auxiliary valency :

It is also termed as coordination number (usually abbreviated as CN) of the central metal ion. It is nonionic or non-ionisable (i.e. coordinate covalent bond type). This is satisfied by either negative ions or neutral molecules having lone pair of electrons (e.g., H2O, NH3 etc.) or even sometimes by some positive groups. The ligands which satisfy the coordination number are directly attached to the metal atom or ion and shown by thick lines.

Every element tends to satisfy both its primary and secondary valencies. In order to meet this requirement a negative ion may often show a dual behaviour, i.e. it may satisfy both primary and secondary valencies since in every case the fulfillment of coordination number of the central metal ion appears essential. This dual behaviour is represented by both thick and dotted lines. For example,  $[CoCl(H_2O)_5]Cl_2$  is represented as



The ions/groups bound by the secondary valencies have characteristic spatial arrangements corresponding to different coordination number. In the modern terminology, such spatial arrangements are called coordination polyhedra and various possibilities are

C.N. = 2	linear	C.N. = 3	Triangular
C.N. = 4	tetrahedral or square planar	C.N. = 6	octahedral.

To distinguish between the two types of valencies, Werner introduced the square brackets [] to enclose those atoms making up the coordination complex and which are, therefore, not ionized.

On the basis of the above postulates Werner formulated the coordination compounds, CoCl<sub>3</sub>. 6NH<sub>4</sub>, CoCl<sub>3</sub>. 5NH<sub>3</sub> and CoCl<sub>3</sub>. 4NH<sub>3</sub> as : [Co(NH<sub>3</sub>)<sub>6</sub>]Cl<sub>3</sub>, [Co(NH<sub>3</sub>)<sub>5</sub>Cl]Cl<sub>2</sub> and [Co(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>]Cl respectively; the species within the square brackets being the coordination entitles (complexes) and the ions outside the square brackets the counter ions. He further postulated that octahedral, square, planar and tetrahedral geometrical shapes are more common in coordination compounds of transition metals. Thus,  $[Co(NH_3)_{\epsilon}]^{3+}$ ,  $[CoCl(NH_3)_{5}]^{2+}$ ,  $[CoCl_2(NH_3)_{4}]^{+}$  are octahedral entities, while  $[Ni(CO)_{4}]$  and  $[PtCl_{4}]^{2-}$  are tetrahedral and square-planar, respectively.

S.No.	Werner complex	Modern notation	Ionisation	Secondary valency satisfied by	Primary valency satisfied by
1	CoCl <sub>3.</sub> 6NH <sub>3</sub>	[Co(NH <sub>3</sub> ) <sub>6</sub> ]Cl <sub>3</sub>	[Co(NH <sub>3</sub> ) <sub>6</sub> ] <sup>3+</sup> +3Cl <sup>−</sup>	six (NH <sub>3</sub> )	three (CΓ)
2	CoCl <sub>3.</sub> 5NH <sub>3</sub>	[Co(NH <sub>3</sub> ) <sub>5</sub> Cl]Cl <sub>2</sub>	[Co(NH <sub>3</sub> ) <sub>5</sub> Cl] <sup>2+</sup> +2Cl <sup>−</sup>	five (NH <sub>3</sub> ) and one (Cl <sup>-</sup> )	three (CΓ) including one (CΓ) with dual nature
3	CoCl <sub>3.</sub> 4NH <sub>3</sub>	[Co(NH <sub>3</sub> ) <sub>4</sub> Cl <sub>2</sub> ]Cl	[Co(NH <sub>3</sub> ) <sub>4</sub> Cl <sub>2</sub> ] <sup>+</sup> + Cl <sup>−</sup>	four (NH <sub>3</sub> ) and two (CГ)	three (Cl <sup>−</sup> ) including two (Cl <sup>−</sup> ) with dual nature
4	CoCl <sub>3.</sub> 3NH <sub>3</sub>	[Co(NH <sub>3</sub> ) <sub>3</sub> Cl <sub>3</sub> ]	[Co(NH <sub>3</sub> ) <sub>3</sub> Cl <sub>3</sub> ]	three (NH <sub>3</sub> ) and three (CI <sup><math>-</math></sup> )	three (Cl <sup>-</sup> ) all with dual nature

## -Solved Examples ————

**Ex.9** Match the pairs of complexes listed in column-I with the method(s) used for their differentiation listed in column-II.

		Column – I	Colum	n–II				
	(1) [Cr(H <sub>2</sub> O) <sub>6</sub> ]Cl	$_{_3}$ and Cr(H $_2$ O) $_5$ Cl] Cl $_2$ .H $_2$ O	(p) Only neutral molecu secondary valences in	le(s) satisfiy(es) all both complexes.				
	(2) [Co(NH <sub>3</sub> ) <sub>5</sub> Br	$]SO_4$ and $[Co(NH_3)_5SO_4]Br$	(q) Negative ion(s) satis well as secondary valer	sfiy(es) both primary as nces in both complexes.				
	(3) [Co(NH <sub>3</sub> ) <sub>5</sub> Cl]	$[Cl_2 \text{ and } [Co(NH_3)_6]Cl_3$	(r) Both complexes give aqueous solution.	same number of ions in				
	(4) [Cu(H <sub>2</sub> O) <sub>4</sub> ]S	$O_4.H_2O$ and $[Cu(H_2O)_6](NO_3)_2$	(s) Both complexes give	e white precipitate with $AgNO_3$ .				
Ans.	(1) − s ;	; (2) – q, r ; (3) – s ; (4) – p						
Sol.	(1)	$[Cr(H_2O)_6]Cl_3 (aq) \rightleftharpoons [Cr(H_2O)_6]Cl_3 (aq) \models [Cr(H_2O)_6]Cl_4 (aq) \models [Cr($	)) <sub>6</sub> ]³+ (aq) + 3Cl⁻ (aq)					
		$[Cr(H_2O)_5Cl] Cl_2.H_2O (aq) \Longrightarrow$	$[Cr(H_2O)_5Cl]^{2+}$ (aq) + 2Cl	⊢ (aq)				
		$Ag^{+} + CI^{-} \longrightarrow AgCI \downarrow (white)$						
	(2)	$[Co(NH_3)_5Br]SO_4 (aq) \implies [Co$	$(NH_3)_5Br]^{2+}$ (aq) + SO <sub>4</sub> <sup>2-</sup>	(aq)				
			1:1 electrolyte					
		$[Co(NH_3)_5SO_4]Br (aq) \longrightarrow [Co$	o(NH <sub>3</sub> )₅SO₄]⁺ (aq) + Br⁻ (a	aq)				
			1 : 1 electrolyte					
		Br <sup>-</sup> and $SO_4^{2-}$ satisfy both prima	ry as well as secondary	valences.				
	(3)	$[Co(NH_3)_5CI]CI_2 (aq) \rightleftharpoons [Co(NH_3)_5CI]CI_2 (aq) = [Co(NH_3)_5CI]CI$	NH <sub>3</sub> )₅Cl]²+ (aq) + 2Cl⁻ (ac	() (				
		$[Co(NH_3)_6]Cl_3 (aq) \rightleftharpoons [Co(NH_3)_6]Cl_3 (aq)$	H <sub>3</sub> ) <sub>6</sub> ]³+ (aq) + 3Cl⁻ (aq)					
	(4)	$[Cu(H_2O)_4]SO_4.H_2O (aq)   [$	$Cu(H_2O)_4]^{2+}(aq) + SO_4^{2-}$ $u(H_2O)_2]^{2+}(aq) + 2NO_2^{-}(aq)$	(aq) q)				
Ex.10	Which of the fol	lowing has the highest molar con	ductivity in solution ?					
	(A) [Pt(NH <sub>3</sub> ) <sub>6</sub> ]Cl	(B) [PtCl(NH <sub>3</sub> ) <sub>5</sub> ]Cl <sub>3</sub>	(C) $[PtCl_2(NH_3)_4]Cl_2$	(D) [PtCl <sub>3</sub> (NH <sub>3</sub> ) <sub>3</sub> ]Cl				
Ans.	А							
Sol.	(A) [Pt(NH <sub>3</sub> ) <sub>6</sub> ]Cl	₄ [Pt(NH <sub>3</sub> ) <sub>6</sub> ] <sup>4+</sup> (aq) + 4Cl⁻ (a	aq).					
	(B) [PtCl(NH <sub>3</sub> ) <sub>5</sub> ]	$Cl_3 \stackrel{aq.}{\longleftarrow} [PtCl(NH_3)_5]^{3+}$ (aq) + 30	Cl⁻ (aq).					
	(C) [PtCl <sub>2</sub> (NH <sub>3</sub> ) <sub>4</sub>	$]\mathrm{Cl}_{_2} \xleftarrow{\mathrm{aq.}} [\mathrm{Pt}\mathrm{Cl}_{_2}(\mathrm{NH}_{_3})_{_4}]^{_2+} (\mathrm{aq}) +$	2Cl⁻ (aq).					
	(D) [PtCl <sub>3</sub> (NH <sub>3</sub> ) <sub>3</sub>	$]CI \stackrel{aq.}{\longleftarrow} [PtCI_3(NH_3)_3]^+ (aq) + C$	Cl⁻ (aq).					
	Since [Pt(NH <sub>3</sub> ) <sub>e</sub>	]Cl <sub>4</sub> gives maximum number of	ions in the solution and	d as number of ions increases				
	molar conductiv	vity in solution increases.						

## Bonding in coordination compounds

#### Valence bond theory :

The valence bond theory, VBT, was extended to coordination compounds by Linus Pauling in 1931. The formation of a complex involves reaction between a lewis base (ligand) and a lewis acid (metal or metal ion) with the formation of a coordinate-covalent (or dative) bonds between them. The model utilizes hybridisation of (n-1) d, ns, np or ns, np, nd orbitals of metal atom or ion to yield a set of equivalent orbitals of definite geometry to account for the observed structures such as octahedral, square planar and tetrahedral, and magnetic properties of complexes. The number of unpaired electrons, measured by the magnetic moment of the compounds determines which d-orbitals are used.

These hybrid orbitals are allowed to overlap with ligand orbitals that can donate electron pairs for bonding.

Following table provides the types of hybridisation with different coordination number.

Coordiantion number of metal	Type of hybridisation	Shape of complex
4	sp <sup>3</sup>	Tetrahedral
4	dsp <sup>2</sup>	Square planer
5	sp <sup>3</sup> d	Trigonal bipyramidal
6	sp <sup>3</sup> d <sup>2</sup>	Octahedral
6	d <sup>2</sup> sp <sup>3</sup>	Octahedral

It is to be noted that the type of hybridisation of metal and shape of complex involved can be predicted conveniently, if some characteristic of the complex like magnetic nature, geometry or whether exhibits isomerism or not, etc., be known.

#### **Coordination Number Six.**

In the diamagnetic octahedral complex,  $[Co(NH_3)_6]^{3+}$ , the cobalt ion is in +3 oxidation state and has the electronic configuration represented as shown below.



Thus, the complex has octahedral geometry and is diamagnetic because of the absence of unpaired electron. Since in the formation of complex the inner d-orbital (3d) is used in hybridisation, the complex is called an inner orbital or low spin or spin paired complex.

The complex  $[FeF_6]^{4-}$  is paramagnetic and uses outer orbital (4d) in hybridisation (sp<sup>3</sup>d<sup>2</sup>) ; it is thus called as outer orbital or high spin or spin free complex. So :



Six pairs of electrons from six F- ions.

#### **Coordination Number Four:**

In the paramagnetic and tetrahedral complex [NiCl<sub>4</sub>]<sup>2-</sup>, the nickel is in +2 oxidation state and the ion has the electronic configuration 3d8. The hybridisation scheme is as shown in figure.



The compound is paramagnetic since it contains two unpaired electrons.

Similarly complex [Ni(CO),] has tetrahedral geometry and is diamagnetic as it contains no unpaired electrons. The hybridisation scheme is as shown in figure.



[PtCl<sub>4</sub>]<sup>2-</sup>

dsp<sup>2</sup> hybrid orbitals Four pairs of electrons from four CI-

Similarly the hybridisation scheme for  $[Ni(CN)_4]^{2-}$  is as shown in figure.

	3d				<u>4s</u>			4p		
Ni <sup>2+</sup> , [Ar]3d <sup>8</sup>	16	11	11	1	1					
[Ni(CN) <sub>4</sub> ] <sup>2–</sup>	11	11	11	11	<u>.</u>			1	4 <u>1</u>	

dsp<sup>2</sup> hybrid orbitals

Four pairs of electrons from four CN-

While the valence bond theory, to a large extent, explains the formation, structures and magnetic behaviour of coordination compounds, it suffers from the following shortcomings :

- 1. A number of assumptions are involved.
- 2. There is no quantitative interpretation of magnetic data.
- 3. It has nothing to say about the spectral (colour) properties of coordination compounds.
- 4. It does not give a quantitative interpretation of the thermodynamic or kinetic stabilities of coordination compounds.
- 5. It does not make exact predictions regarding the tetrahedral and square-planar structures of 4coordinate complexes.
- 6. It does not distinguish between strong and weak ligands.

#### Magnetic Properties of Coordination Compounds : (NCERT-Ref.)

Additional information for understanding the nature of coordination entities is provided by magnetic susceptibility measurements. We have noted that coordination compounds generally have partially filled d orbitals and as such they are expected to show characteristic magnetic properties depending upon the oxidation state, electron configuration, coordination number of the central metal and the nature of the ligand field. It is experimentally possible to determine the magnetic moments of coordination compounds which can be utilized for understanding the structures of these compounds.

The number of unpaired electrons in any complex can be easily calculated from the configuration of the metal ion, its coordination number and the nature of the ligands involved( strong or weak from the spectrochemical series) and after that the magnetic moment of the complexes can be easily calculated using

### Magnetic Moment = $\sqrt{n(n + 2)}$ Bohr Magneton

For metal ions with upto three electrons in the d-orbitals like  $Ti^{3+}$ ,  $(d^1)$ ;  $V^{3+}$   $(d^2)$ ;  $Cr^{3+}$   $(d^3)$ ; two vacant d-orbitals are easily available for octahedral hybridisation. The magnetic behaviour of these free ions and their coordination entities is similar. When more than three 3d electrons are present, like in  $Cr^{2+}$  and  $Mn^{3+}$   $(d^4)$ ;  $Mn^{2+}$  and  $Fe^{3+}$   $(d^5)$ ;  $Fe^{2+}$  and  $Co^{3+}$   $(d^6)$ ; the required two vacant orbitals for hybridisation is not directly available (as a consequence of Hund's rules). Thus, for  $d^4$ ,  $d^5$  and  $d^6$  cases, two vacant d-orbitals are only available for hybridisation as a result of pairing of 3d electrons which leaves two, one and zero unpaired electrons respectively.

The magnetic data agree with maximum spin pairing in many cases, especially with coordination compounds containing  $d^6$  ion. However, there are complications with the coordination compounds / species having  $d^4$  and  $d^5$  ions.

 $[Mn(CN)_6]^{3-}$  has a magnetic moment equal to two unpaired electrons while  $[MnCl_6]^{3-}$  has a magnetic moment equal to four unpaired electrons.

Similarly  $[Fe(CN)_6]^{3-}$  has magnetic moment of a single unpaired electron while  $[FeF_6]^{3-}$  has a magnetic moment of five unpaired electrons.  $[CoF_6]^{3-}$  is paramagnetic with four unpaired electrons while  $[Co(C_2O_4)]^{3-}$  is diamagnetic.

This anomalous behaviour is explained by valence bond theory in terms of formation of inner orbitals and outer orbitals complexes.

 $[Mn(CN)_6]^{3-}$ ,  $[Fe(CN)_6]^{3-}$  and  $[Co(C_2O_4)_2]^{3-}$  are inner orbital complexes involving d<sup>2</sup>sp<sup>3</sup> hybridisation, the former two are paramagnetic and the latter diamagnetic.  $[MnCl_6]^{3-}$ ,  $[FeF_6]^{3-}$  and  $[CoF_6]^{3-}$  are outer orbital complexes involving sp<sup>3</sup>d<sup>2</sup> hybridisation and are paramagnetic having four, five and four electrons respectively.

Solved Examples

- **Ex.11** How would you account for the following ?
  - (a)  $[Fe(CN)_6]^{3-}$  is weakly paramagnetic while  $[Fe(CN)_6]^{4-}$  is diamagnetic
  - (b) Ni(CO)<sub>4</sub> possesses tetrahedral geometry while  $[Ni(CN)_4]^{2-}$  is square planar
  - (c)  $[Cr(NH_3)_6]^{3+}$  is paramagnetic while  $[Ni(CN)_4]^{2-}$  is diamagnetic. Explain why?
- **Sol.** (a)  $[Fe(CN)_6]^{3-}$  involves  $d^2sp^3$  hybridization.

	3d	<u>4s</u> 4p
Fe³+, [Ar] 3d⁵	$\uparrow \uparrow \uparrow \uparrow \uparrow \uparrow$	
[Fe(CN) <sub>6</sub> ]³−		
	Rearrangement	d <sup>2</sup> sp <sup>3</sup> hybridization

One d-orbital is singly occupied, hence it is weakly paramagnetic in nature.  $[Fe(CN)_6]^{4-}$  involves also d<sup>2</sup>sp<sup>3</sup> hybridization but it has Fe<sup>2+</sup> ion as central ion.

	3d	<u>4s</u> 4p	
Fe <sup>2+</sup> , [Ar] 3d <sup>6</sup>	$\uparrow \downarrow \uparrow \uparrow \uparrow \uparrow \uparrow$		
	$\uparrow\downarrow\uparrow\downarrow\uparrow\downarrow$		•
[Fe(CN) <sub>6</sub> ] <sup>4–</sup>	Rearrangement	d <sup>2</sup> sp <sup>3</sup> hybridizatic	⁄

All electrons are paired, hence it is diamagnetic in nature.

(b) In the formation of  $Ni(CO)_4$  nickel atom undergoes sp<sup>3</sup> hybridization, hence it is tetrahedral in shape.



In the formation of  $[Ni(CN)_4]^{2-}$ ,  $Ni^{2+}$  ion undergoes dsp<sup>2</sup> hybridization, hence it is square planar in shape.

Ni<sup>2+</sup>, [Ar] 3d<sup>8</sup>

[Ni(CN)]2-

3d	4s	4р	
$\uparrow\downarrow\uparrow\downarrow\uparrow\downarrow\uparrow\downarrow\uparrow\uparrow\uparrow\uparrow\uparrow$			
$\uparrow\downarrow\uparrow\downarrow\uparrow\downarrow\uparrow\downarrow\uparrow\downarrow$	•	••••	
Rearrangement		hybridiza	/ tion

(c) Chromium in +3 oxidation i.e., [Ar]<sup>18</sup> 3d<sup>3</sup> electron configuration always have three unpaired electrons with strong as well as with weak field ligands. So,

				C			 		 5
[Cr(NH <sub>3</sub> ) <sub>6</sub> ] <sup>3+</sup>	1	1	1						
	d²sp	o³ hỵ	ybri	d or	bita		 	,	
[Ni(CN) <sub>4</sub> ] <sup>2-</sup>	11	11	11	11		40			

dsp<sup>2</sup> hybrid orbitals

It is diamagnetic as all electrons are paired in 3d<sup>8</sup> configuration of Ni<sup>2+</sup>.

**Ex.12** Sodium nitroprusside,  $Na_2[Fe(CN)_5NO]$  is a diamagnetic substance and a important laboratory reagent for the testing of sulphide ions. The metal involved in the complexation in this is present in which of the following hybridisation state :



- **Ex.13** When the d-orbitals involved in the hybridisation are inner (n–1) d-orbitals ; the complex formed are referred to as :
  - (1) high spin-complexes
- (2) low spin complexes
- (3) zero spin complexes (4) positive spin complexes
- Ans. (2)

- Sol. When in the formation of complex the inner d-orbital, (n 1)d is used in hybridisation, the complex is called an inner orbital or low spin or spin paired complex.
   When in the formation of complex the outer d-orbital, nd is used in hybridisation, the complex is called a outer orbital or high spin or spin free complex.
- **Ex.14** Explain  $[Co(NH_3)_6]^{3+}$  is an inner orbital complex whereas  $[Ni(NH_3)_6]^{2+}$  is an outer orbital complex.
- **Sol.** In the presence of  $NH_3$ , the 3d electrons pair up leaving two d-orbitals empty to be involved in  $d^2sp^3$  hybridisation forming inner orbital complex in case of  $[Co(NH_3)_6]^{3+}$ .

In Ni(NH<sub>3</sub>)<sub>6</sub><sup>2+</sup>, Ni is in +2 oxidation state and has d<sup>8</sup> configuration, the hybridisation involved is sp<sup>3</sup>d<sup>2</sup> forming outer orbital complex.

### **Crystal Field Theory :**

The drawbacks of VBT of coordination compounds are, to a considerable extent, removed by the Crystal Field Theory.

The crystal field theory (CFT) is an electrostatic model which considers the metal-ligand bond to be ionic arising purely from electrostatic interaction between the metal ion and the ligand. Ligands are treated as point charges in case of anions or dipoles in case of neutral molecules. The five d orbitals is an isolated gaseous metal atom/ion have 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 due to ligands (either anions or the negative ends of dipolar molecules like  $NH_3$  and  $H_2O$ ) in a complex, it becomes asymmetrical and the degeneracy of the d orbitals is lost. It results in splitting of the d orbitals. The pattern of spitting depends upon the nature of the crystals field.

#### (a) Crystal field splitting in octahedral coordination entities :

In an octahedral coordination entity with six ligands surrounding the metal atom/ion, there will be repulsion between the electrons in metal d orbitals and the electrons (or negative charges) of the ligands. Such a repulsion is more when the metal d orbitals is directed towards the ligand than when it is away from the ligand. Thus, the  $d_{x^2-v^2}$  and  $d_{z^2}$  orbitals (axial orbitals) which point towards the axis

along the direction of the ligand will experience more repulsion and will be raised in energy ; and the d<sub>xy</sub> , d<sub>yz</sub> and d<sub>zx</sub> orbitals (non-axial) orbitals which are directed between the axis will be lowered in energy relative to the average energy in the spherical crystal field. Thus, the degeneracy of the d orbitals has been removed due to ligand electron-metal electron repulsions in the octahedral complex to yield three orbitals of lower energy, t<sub>2g</sub> set and two orbitals of higher energy, e<sub>g</sub> set. This splitting of the degenerate levels due to the presence of ligands in a definite geometry is termed as crystal field splitting and the energy separation is denoted by  $\Delta_0$  (the subscript o is for octahedral). Thus, the energy of the two e<sub>g</sub> orbitals will increase by (3/5) $\Delta_0$  and that of the three t<sub>2g</sub> will decrease by (2/5)  $\Delta_0$ .



Ι-

The crystal field splitting,  $\Delta_0$ , depends upon the fields produced by the ligand and charge on the metal ion. Some ligands are able to produces strong fields in which case, the splitting will be large whereas others produce weak fields and consequently result in small splitting of d orbitals. In general, ligands can be arranged in a series in the orders of increasing field strength as given below :

$$\frac{\langle Br^{-} \langle SCN^{-} \langle Cl^{-} \langle S^{2-} \langle F^{-} \langle OH^{-} \langle C_{2}O_{4}^{2-} \langle H_{2}O \rangle}{\text{weak} \text{ ligand}} < \underbrace{NCS^{-} \langle edta^{4-} \langle NH_{3} \langle en \langle NO_{2}^{-} \langle CN^{-} \langle CO \rangle }{\text{strong} \text{ ligand}}$$

Such a series is termed as spectrochemical series. It is an experimentally determined series based on the absorption of light by complexes with different ligands. For d<sup>4</sup> configuration, the fourth electron will singly occupy  $e_g$  orbital (according to Hund's rule) or will undergo pairing in  $t_{2g}$  orbital, which of these possibilities occurs, depends on the relative magnitude of the crystal field splitting,  $\Delta_0$  and the pairing energy, P (P represents the energy required for electron pairing in a single orbital). The two possibilities are :

(i) If  $\Delta_0 < P$ , the fourth electron enters one of the  $e_g$  orbitals giving the configuration  $t_{2g}^3 e_g^1$ . Ligands for which  $\Delta_0 < P$  are known as weak field ligands and form high spin complexes.

(ii) If  $\Delta_0 > P$ , it becomes more energetically favourable for the fourth electron to occupy a  $t_{2g}$  orbital with configuration  $t_{2g}^4 e_g^0$ . Ligands which produce this effect are known as strong field ligands and form low spin complexes.

#### (b) Crystal field splitting in tetrahedral coordination entities :

In tetrahedral coordination entity formation, the d orbital splitting is inverted and is smaller as compared to the octahedral field splitting. For the same metal, the same ligands and metal-ligand distances, it can be shown that  $\Delta_t = (4/9)\Delta_0$ . This may attributes to the following two reasons.

(i) there are only four ligands instead of six, so the ligand field is only two thirds the size ; as the ligand field spliting is also the two thirds the size and (ii) the direction of the orbitals does not concide with the

direction of the ligands. This reduces the crystal field spliting by roughly further two third. So  $\Delta_t = \frac{2}{3}$ 

$$\times \frac{2}{3} = \frac{4}{9} \Delta_{o}.$$

Consequently, the orbital splitting energies are not sufficiently large for forcing pairing and, therefore, low spin configurations are rarely observed.



Since  $\Delta_t < \Delta_o$ , crystal field spliting favours the formation of

octahedral complexes.

**Ex.15** The magnetic moments of following, arranged in increasing order will be (atomic number of Co = 27) (A)  $[Co(en)_3]^{3+}$  (B)  $[CoF_6]^{3-}$  (C)  $[Co(SCN)_4]^{2-}$ 

(1) 
$$A > B > C$$
 (2)  $B > C > A$  (3)  $C > B > A$  (4)  $B > A > C$   
Ans. (B)



**Ex.16** Amongst the following ions which one has the highest magnetic moment value? (i)  $[Cr(H_2O)_{e}]^{3+}$ (ii)  $[Fe(H_2O)_{e}]^{2+}$ (iii)  $[Zn(H_2O)_6]^{2+}$ .

[Fe(H<sub>2</sub>O)<sub>2</sub>]<sup>2+</sup> Ans.

- Sol. (i) [Cr(H<sub>2</sub>O)<sub>6</sub>]<sup>3+</sup> contains 3 electrons (ii) [Fe(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup> contains 4 electrons (iii) In [Zn(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup> all electrons are paired
- Ex.17 Arrange the following in increasing order as directed. (a) Mn<sup>3+</sup>, V<sup>3+</sup>, Cr<sup>3+</sup>, Fe<sup>3+</sup> – magnetic moment in aqueous solutions (octahedral complex).
  - (b)  $Br^-$ ,  $S^{2-}$ ,  $NO_2^-$ ,  $CO_1$ ,  $H_2O_1$ ,  $CN^-$ ,  $NH_3$ ,  $NO_3^-$  strength of ligands.
- (a) V<sup>3+</sup> < Cr<sup>3+</sup> < Mn<sup>3+</sup> < Fe<sup>3+</sup> (b)  $Br^- < S^{2-} < NO_3^- < H_2O < NH_3 < NO_2^- < CN^- < CO$ Ans.
- (a)  $Mn^{3+}$ ,  $3d^4 4$  unpaired electrons;  $V^{3+}$ :  $3d^2 2$  unpaired electrons;  $Cr^{3+}$ ,  $3d^3 3$  unpaired electrons; Sol. Fe<sup>3+</sup>,  $3d^5 - 5$  unpaired electrons.
  - (b) According to spectrochemical series (values of  $\Delta$  have been experimentally determined).

#### **COLOUR IN COORDINATION COMPOUNDS:**

According to the crystal field theory the colour is due to the d-d transition of electron under the influence of ligands. The mechanism of light absorption in coordination compounds is that photons of appropriate energy can excite the coordination entity from its ground state to an excited state. Consider the Ti(III) ion in solution, that is [Ti(H2O)]3+. This is a violet colour octahedral complex, where in the ground state of the complex a single electron is present in t<sub>2q</sub> level. The next higher state available for the transition is the empty e, level. If the light corresponding to the energy of yellow-green is absorbed by the complex, it would excite the electron from  $t_{2g}$  level to  $e_{g}$  level. Consequently the complex appears violet in colour.



Transition of an electron in [Ti(H<sub>2</sub>O)<sub>6</sub>]<sup>3+</sup>

Table below gives the relationship of the wavelength absorbed and the colour observed. Relationship between the wavelength of light absorbed and the colour observed In some coordination entitles

Coordination entity	Wavelength of light absorbed (nm)	Colour of light absorbed	Colour of coordination entity		
[CoCl(HN <sub>3</sub> ) <sub>5</sub> ] <sup>2+</sup>	535	Yellow	Violet		
[Co(NH <sub>3</sub> ) <sub>5</sub> (H <sub>2</sub> O)] <sup>3+</sup>	500	Blue Green	Red		
[Co(NH <sub>3</sub> ) <sub>6</sub> ] <sup>3+</sup>	475	Blue	Yellow Orange		
[Co(CN) <sub>6</sub> ] <sup>3-</sup>	310	Ultraviolet	Pale Yellow		
[Cu(H <sub>2</sub> O) <sub>4</sub> ] <sup>2+</sup>	600	Red	Blue		
[Ti(H <sub>2</sub> O) <sub>6</sub> ] <sup>3+</sup>	498	Blue Green	Purple		

**Note :** (a) In absence of ligand, crystal field splitting does not occur and as a consequence the substance appears colourless. For example (i) removal of water from violet coloured complex  $[Ti(H_2O)_6]CI_3$  on heating makes it colourless. (ii) Similarly anhydrous copper sulphate  $(CuSO_4)$  is white, but hydrated copper sulphate  $(CuSO_4.5H_2O)$  is blue coloured.

(b) The nature of the ligand and the molar ratio of metal : ligand also influence the colour of the complex for example in the pale green complex of  $[Ni(H_2O)_6]$ , the colour change is absorbed when ethylenediamine is progressively added to it.

Molar ratio of en : Ni	Coloured observed
1:1	Pale blue
2:1	Blue/Purple
3 : 1	Violet

**Note** : Ruby is  $AI_2O_3$  in which 0.5–1% Cr<sup>3+</sup> ions (d<sup>3</sup> electron system) are randomly distributed in the positions normally occupied by  $AI^{3+}$ . We may consider Cr(III) species as octahedral Cr(III) complexes incorporated into the alumina lattice ; d-d transition of electron at these centres give rise to the colour (red).

Emerland is the mineral beryl  $(Be_3Al_2Si_6O_{18})$  in which  $Cr^{3+}$  ions occupy octahedral sites, but in this case low energy corresponding to yellow red and blue is absorbed and light corresponding to green region is transmitted.

#### Limitations of crystal field theory

- (1) It considers only the metal ion d-orbitals and gives no consideration at all to other metal orbitals (such as s,  $p_x$ ,  $p_y$  and  $p_z$  orbitals).
- (2) It is unable to account satisfactorily for the relative strengths of ligands. For example it gives no explanation as to why H<sub>2</sub>O is a stronger ligand than OH<sup>-</sup> in the spectrochemical series.
- (3) Account to this theory, the bond between the metal and ligands are purely ionic. It gives no account on the partly covalent nature of the metal ligand bonds.
- (4) The CFT cannot account for the  $\pi$ -bonding in complexes.

## Metal Carbonyls :

Compounds of metals with CO as a ligand are called metal carbonyls.

They are of two types.

(a) Monomeric : Those metal carbonyls which contain only one metal atom per molecule are called monomeric carbonyls. For examples :[Ni(CO)<sub>4</sub>] (sp<sup>3</sup>, tetrahedral) ; [Fe(CO)<sub>5</sub>] (dsp<sup>3</sup>, trigonal bipyramidal)

 $[Cr(CO)_{e}]$  (d<sup>2</sup> sp<sup>3</sup>, octahedral);  $[V(CO)_{e}]$  (d<sup>2</sup> sp<sup>3</sup>, octahedral, only carbonyl which is paramagnetic having one unpaired electron. This is least stable among all the four carbonyls)



(b) Polymeric : Those metal carbonyl which contain two or more than two metal atom per molecule and they have metal-metal bonds are called polymeric carbonyl. For example : Mn<sub>2</sub> (CO)<sub>10</sub> , Co<sub>2</sub>(CO)<sub>8</sub>, etc.



The metal-carbon bond in metal carbonyls possess both s and p character. The M-C  $\sigma$  bond is formed by the donation of lone pair of electrons on the carbonyl carbon (CO is a weak base) into a vacant orbital of the metal. The M — C  $\pi$  bond is formed by the donation of a pair of electron from a filled d orbital of metal into the vacant antibonding  $\pi^*$  orbital of carbon monoxide. Thus carbon monoxide acts as  $\sigma$  donor (OC  $\rightarrow$  M) and a  $\pi$  acceptor (OC  $\leftarrow$  M), with the two interactions creating a synergic effect which strengthens the bond between CO and the metal as shown in figure.



#### **STABILITY OF COORDINATION COMPOUNDS :**

The stability of a coordination compound [ML] is measured in terms of the stability constant (equilibrium constant) given by the expression,

 $\beta_n = [ML_n]/[M(H_2O)_n][L]^n$ 

for the overall reaction :

 $M(H_2O)_n + nL \Longrightarrow ML_n + nH_2O$ 

By convention, the water displaced is ignored, as its concentration remains essentially constant. The above overall reaction takes place in steps, with a stability (formation) constant, K1, K2, K3, ..... K for each step as represented below :

$$M(H_2O)_n + L \rightleftharpoons ML(H_2O)_{n-1} + H_2O$$
  
$$K_4 = [ML(H_2O)_{n-1}] / \{[M(H_2O)_n][L]\}$$

 $ML_{n-1} (H_2O) + L \Longrightarrow ML_n + H_2O$ 

$$K_{n} = [ML_{n}] / \{[ML_{n-1} (H_{2}O)] [L]\}$$

 $M(H_2O)_n + nL \Longrightarrow ML_n + nH_2O$ 

$$\beta_n = K_1 \times K_2 \times K_3 \times \dots \times K_n$$

 $\beta_{\text{n}}\!\!,$  the stability constant, is related to thermodynamic stability when the system has reached equilibrium. Most of the measurements have been made from aqueous solutions, which implies that the complex is formed by the ligand displacing water from the agua complex of the metal ion. Ignoring the charge and taking L as an unidentate ligand, the stepwise formation of the complex is represented as shown above.  $K_1, K_2, K_3, \dots, K_n$  representing the stepwise stability (or formation) constants.

The above is thermodynamic stability criteria, there can be another kind of stability called Kinetic Stability, which measures the rate of ligand replacement.

Some important generalisation regarding stability constants :

For a given metal and ligand the stability is generally greater when the charge on the metal ion is greater. Thus, stability of coordination entities of ions of charge 3+ is greater than the entities of 2+ ions. Further, for the divalent ions of the first row transition elements, irrespective of the ligand involved, the stabilities vary in the Irving-Williams order :

Mn" < Fe" < Co" < Ni" < Cu" > Zn"

This order is according to the size of the ions, smaller the size of the ion or greater the charge density on the metal greater is the stability of the complex.

In F<sup>-</sup>, Cl<sup>-</sup>, Br<sup>-</sup>, I<sup>-</sup>; F<sup>-</sup> forms strongest complexes due to small size & hence high charge density.

The stability also depends on the formation of chelate rings. If L is an unidentate ligand and L–L, a didentate ligand and if the donor atoms of L and L–L are the same element, then L–L will replace L. The stabilisation due to chelation is called the chelate effect. It is of great importance in biological systems and analytical chemistry. The chelate effect is maximum for the 5- and 6-membered rings. In general, rings provide greater stability to the complex.

## Solved Examples.

Ex.18	What will be the correct order for the wavelength of absorption in the visible region for the following : $[Ni(NO_2)_{\beta}]^{4-}$ , $[Ni(NH_3)_{\beta}]^{2+}$ , $[Ni(H_2O)_{\beta}]^{2+}$ ?							
Sol.	(i) The order of the ligand in the spectrochemical series : $H_2O < NH_3 < NO_2^{-}$ .							
	Hence the wavelength of the light observed will be in the order : $[Ni(H_2O)_6]^{2+} < [Ni(NH_3)_6]^{2+} < [Ni(NO_2)_6]^{4-}.$							
Ex.19	Thus, wavelengths absorbed (E = hc/ $\lambda$ ) will be in the opposite order. A solution of [Ni(H <sub>2</sub> O) <sub>6</sub> ] <sup>2+</sup> is green but a solution of [Ni(CN) <sub>4</sub> ] <sup>2-</sup> is colourless. Explain.							
301.	of the complex. In $[Ni(CN)_4]^{2-}$ there is no unpaired electron for d-d transition ; so it is colourless. In $[Ni(H_2O)_6]^{2+}$							
	there are two unpaired electrons available for d-d transition; so it is coloured.							
Ex.20 Sol.	<ul> <li>[Fe(CN)<sub>6</sub>]<sup>4-</sup> and [Fe(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup> are of different colours in dilute solutions. Why?</li> <li>The colour of the complex in the dilute solution depends on the nature of ligands.</li> <li>(i) The order of the ligand in the spectrochemical series : H<sub>2</sub>O &lt; CN<sup>-</sup></li> </ul>							
	Hence the wavelength of the light observed will be in the order : $[Fe(H_2O)_6]^{2+} < [Fe(CN)_6]^{4-}$ .							
	Thus, wavelengths absorbed (E = $hc/\lambda$ ) will be in the opposite order. Therefore they so different colours in dilute solutions.							
Ex.21	Calculate the overall complex dissociation equilibrium constant for the $[Cu(NH_3)_4]^{2+}$ ion, given that $\beta^4$ for this complex is 2.1 × 10 <sup>13</sup> .							
Sol.	The overall dissociation constant is the reciprocal of overall stability constant i.e. $\frac{1}{\beta^4} = 4.7 \times 10^{-14}$ .							
	– Self Fractice Froblem —							
1.	Aqueous copper sulphate solution (blue in colour) gives :							
	(i) a green precipitate with aqueous potassium fluoride and							

(ii) a bright green solution with aqueous potassium chloride. Explain these experimental results.

- **2.** Amongst the following, the most stable complex is (i)  $[Fe(H_2O)_a]^{3+}$  (ii)  $[Fe(NH_2)_a]^{3+}$  (iii)  $[Fe(C_2O_4)_2]^{3-}$  (iv)  $[FeCl_a]^{3-}$
- **3.**  $[Ti(H_2O)_6]^{3+}$  absorbs light of wavelength 5000 Å. Name one ligand which would form a titanium(III) complex absorbing light of lower wavelength than 5000 Å and one ligand which would a complex absorbing light of wavelength higher than 5000 Å.

## Answers

- 1. (i)  $[Cu(H_2O)_4]^{2+} + 4F^- \longrightarrow [CuF_4]^{2-} + 4H_2O$ . (ii)  $[Cu(H_2O)_4]^{2+} + 4CI^- [CuCI_4]^{2-} + 4H_2O$ .
- **2.**  $[Fe(C_2O_4)_3]^{3-}$ .
- **3.** CN<sup>-</sup>, F<sup>-</sup>

## Isomerism in coordination compounds :



#### (1) Structural isomerism :

The structural isomerism arises due to different lignads within the coordination sphere.

#### (A) Ionisation Isomerism

This type of isomerism occurs when the counter ion in a coordination compound is itself a potential ligand and can displace a ligand which can then become the counter ion. For example, following complexes show ionisation isomerism.

 $[\text{Co}(\text{NH}_3)_5\text{SO}_4]\text{NO}_3$  and  $[\text{Co}(\text{NH}_3)_5\text{NO}_3]\text{SO}_4$ 

 $[\text{Co}(\text{NH}_3)_4(\text{NO}_2)\text{CI}]\text{CI} \text{ and } [\text{Co}(\text{NH}_3)_4\text{CI}_2]\text{NO}_2.$ 

[Co(NH<sub>3</sub>)<sub>4</sub>(H<sub>2</sub>O)Cl]Br<sub>2</sub> and [Co(NH<sub>3</sub>)<sub>4</sub>BrCl]Br.H<sub>2</sub>O. [Also an example of hydrate isomers.]

 $[Pt(NH_3)_4Cl_2]Br_2$ , and  $[Pt(NH_3)_4Br_2]Cl_2$ .

[CoCl(en)<sub>2</sub>(NO<sub>2</sub>)]SCN, [Co(en)<sub>2</sub>(NO<sub>2</sub>)SCN]Cl and [Co(en)<sub>2</sub>(SCN)Cl]NO<sub>2</sub>

## (B) Solvate/Hydrate Isomerism :

It occurs when water forms a part of the coordination entity or is outside it. This is similar to ionisation isomerism. Solvate isomers differ by whether or not a solvents molecule is directly bonded to the metal ion or merely present as free solvent molecules in the crystal lattice. For example,  $CrCl_3 \cdot 6H_2O$  exists in three distinct isomeric forms :  $[Cr(H_2O)_6]Cl_3$ , violet ;  $[CrCl(H_2O)_6]Cl_2 \cdot H_2O$ , blue green :  $[CrCl_2(H_2O)_4]Cl.2H_2O$ , dark green.

## (C) Linkage isomerism :

In some ligands, like ambidentate ligands, there are two possible coordination sites. In such cases, linkage

isomerism exist. e.g.,  $NO_2$  group can be bonded to metal ions through nitrogen (-NO<sub>2</sub>) or through oxygen

(-ONO). SCN too can be bonded through sulphur (-SCN) thiocyanate or through nitrogen (-NCS) isothiocyanate.

For example :  $[Co ONO (NH_3)_5] Cl_2 \& [Co NO_2 (NH_3)_5] Cl_2$ .

#### (D) Coordination Isomerism

Coordination compounds made up of cationic and anionic coordination entities show this type of isomerism due to the interchange of ligands between the cation and anion entities. Some of the examples are :

- (i)  $[Co(NH_3)_6][Cr(CN)_6]$  and  $[Cr(NH_3)_6](Co(CN)_6]$
- (ii)  $[Cu(NH_3)_4](PtCl_4]$  and  $[Pt(NH_3)_4][CuCl_4]$
- (iii)  $[Co(NH_3)_6][Cr(SCN)_6]$  and  $[Cr(NH_3)_4(SCN)_2](Co(NH_3)_2(SCN)_4]$
- (iv)  $[Pt(NH_3)_4][PtCI_6]$  and  $[Pt(NH_3)_4CI_2][PtCI_4]$

Such isomers are expected to have significant differences in their physical and chemical properties.

#### (2) Stereoisomerism

The isomers in which atoms are bonded to each other in the same order but that differ in the arrangement of these atoms in the space are called as stereoisomers and the phenomenon as stereoisomerism.

#### (A) Geometrical Isomerism

This type of isomerism arises in heteroleptic complexes due to different possible geometric arrangements of the ligands. Geometrical isomerism is common among coordination compounds with coordination numbers 4 and 6.

#### (i) Coordination Number Four :

#### (a) Tetrahedral Complex :

The tetrahedral compounds can not show geometrical isomerism as we all know that all four positions are equivalent in tetrahedral geometry.

#### (b) Square Planar Complex :

In a square planar complex of formula  $[Ma_2b_2]$  [a and b are unidentate], the two ligands 'a' may be arranged adjacent to each other in a cis isomer, or opposite to each other in a trans isomer as depicted.





Square planar complex of the type Ma<sub>2</sub>bc (where a,b,c are unidentates) shows two geometrical isomers.



Square planar complex of the type Mabcd (where a,b,c,d are unidentates) shows three geometrical isomers.



Example is  $[Pt(NH_3)BrCl(py)]$ . three isomers of the complex  $[Pt(NH_3)(NH_2OH)(py)(NO_2)]^+$  have been isolated and identified.

Square planar complex of the type  $M(AB)_2$  (where AB are unsymmetrical bidentates) shows two geometrical isomers. Example is  $[Pt(gly)_2]$  in which gly is unsymmetrical ligand.



Similarly, M(AB)(CD) also shows two geometrical isomers.

**Note :** M(AA)<sub>2</sub>, (where AA are symmetrical bidentates) does not show geometrical isomerism. e.g., [Cu(en)<sub>2</sub>]<sup>2+</sup> [Pt(ox)<sub>2</sub>]<sup>2-</sup>, etc.

#### (ii) Coordination Number Six :

Geometrical isomerism is also possible in octahedral complexes.



#### Geometrical isomers (cis and trans) of [Co(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>]\*

Number of possible isomers and the spatial arrangements of the ligands around the central metal ion for the specific complexes are given below.

(I) Complexes containing only unidentate ligands (a)  $Ma_2b_4 - 2$  (b)  $Ma_4bc - 2$  (c)  $Ma_3b_3$ 

Complexes of the formula  $Ma_3b_3$ , where a and b are monodentate ligands, may show two isomeric forms called fac– and mer–. Facial isomers have three identical ligands on one triangular face where as meridional isomers have three identical ligands in a plane bisecting the molecule. Similar isomers are possible with some chelating ligands.



The facial(fac) and meridional(mer) isomers of  $[Co(NH_3)_3(NO_2)_3]$ .

#### Unsymmetrical bidentate ligands also show fac-mer isomerism

(d)  $Ma_3b_2c - 3$  (e)  $Ma_3bcd - 4$  (f)  $Ma_2b_2c_2 - 5$ (g)  $Ma_2b_2cd - 6$  (h)  $Ma_2bcde - 9$ (i) Mabcdef, [Pt(py)(NH<sub>3</sub>)(NO<sub>2</sub>)(Cl)(Br)(l)] - 15

(II) Compounds containing bidentate ligand and unidentate ligands.

(i)  $M(AA)a_3b$  – Two geometrical isomers are possible.



(ii)  $M(AA)a_2b_2$  – Three geometrical isomers are possible.



### (B) Optical Isomerism :

A Coordination compound which can rotate the plane of polarised light is said to be optically active. When the coordination compounds have same formula but differ in their ability to rotate directions of the plane of polarised light are said to exhibit optical isomerism and the molecules are optical isomers. Optical isomers are mirror images that cannot be superimposed on one another. These are called as enantiomers. The molecules or ions that cannot be superimposed are called chiral. This is due to the absence of elements of symmetry in the complex. The two forms are called dextro(d) and laevo(l) depending upon the direction they rotate the plane of polarised light in a polarimeter (d rotates to the right, I to the left).

#### (a) Octahedral complexes :

Optical isomerism is common in octahedral complexes involving didentate ligands. For example,  $[Co(en)_3]^{3+}$  has d and I forms as given below.



Cis-isomer of  $[PtCl_2(en)_2]^{2+}$  show optical isomerism as shown below because of the absence of plane of symmetry as well as centre of symmetry.



But trans isomer of  $[PtCl_2(en_2)]^{2+}$  does not show optical isomerism.



cis-[Co(NH<sub>3</sub>)<sub>2</sub>Cl<sub>2</sub>(en)]<sup>+</sup> can show optical isomerism due to the absence of plane of symmetry as well as centre of symmetry.



#### (b) Tetrahedral Complexes.

Optical isomerism is expected in tetrahedral complexes of the type [Mabcd] analogous to tetrahedral carbon atom.



#### (c) Square Planar Complexes.

Square planar complexes are rarely found to show the optical isomerism. The plane formed by the four ligating atoms and the metal ion is considered to be a mirror plane and thus prevents the possibility of chirality.

#### Applications of coordination and organometallic compounds

- (i) Coordination compounds are of great importance in biological systems. Example being chlorophyll (the green pigment in plants); haemoglobin (the red pigment of blood, which acts as oxygen carrier) along with myoglobin (which stores oxygen and is a regulator of respiration); Vitamin B<sub>12</sub>, cyanocobalammine, the anti-pernicions anaemia factor. All of these, respectively, are the coordination compounds of magnesium, iron and cobalt with the macrocyclic porphyrin and corrin ligands.
- (ii) There are many examples of the use of coordination compounds in qualitative and quantitative chemical analysis. The familiar colour reactions given by metal ions with a number of ligands (especially the chelating ligands), as a result of formation of coordination entities, form the basis for their detection and estimation by classical and instrumental methods of analysis. Familiar examples of such reagents are :

ethylenediaminetetraaceticacid (EDTA), dimethylglyoxime,  $\alpha$ -nitroso  $\beta$ -naphthol, cupron, etc.

- (iii) Some important extraction processes of metals, like those of extraction of silver and gold, make use of complex formation. Gold, for example, combines with cyanide in the presence of oxygen and water to form the coordination entity [Au(CN)<sub>2</sub>]<sup>-</sup> in aqueous solution. Gold can be precipitated from this solution by the addition of Zinc.
- (iv) Purification of metals can be achieved through formation and subsequent decomposition of their coordination compounds. For example, impure nickel is converted to [Ni(CO)<sub>4</sub>], which is decomposed to yield pure nickel.

(v) Organometallic compounds are used as catalysts. These catalysts are either of the homogeneous type (soluble in the reaction medium) or of the heterogeneous type (insoluble in the reaction medium). The catalysed polymerisation of alkenes at atmospheric pressure and ambient temperature using Ziegler-Natta catalyst (titanium tetrachloride plus triethylaluminium) is one of the important discoveries of organometallic chemistry. The first effective homogeneous catalyst chloridotris(triphenylphosphine) rhodium(I), [RhCl(PPh<sub>3</sub>)<sub>3</sub>] for hydrogenation was given by Wilkinson.

(vi) Tetra ethyl lead (TEL) is used as antiknock compound in gasoline.

- (vii)Articles can be electroplated with silver and gold much more smoothly and evenly from solutions of the complexes, [Ag(CN)<sub>2</sub>]<sup>-</sup> and [Au(CN)<sub>2</sub>]<sup>-</sup> than from a solution of simple metal ions.
- (viii) In black and white photography, the developed film is fixed by washing with hypo solution which dissolves the undecomposed AgBr to form a complex ion,  $[Ag(S_2O_3)_2]^{3-}$ .
- (ix) The chelate therapy is now a day is used in medicinal chemistry.

(a) Excess of copper and iron are removed by the chelating ligands D-penicillamine and desferrioxime B via the formation of coordination compound in plant/animal systems.

(b) EDTA is used in the treatment of lead poisoning.

(c) Coordination compounds of platinum like cis-platin and related compounds effectively inhibit the growth of tumours.

Solved Examples

- **Ex.22** Give evidence that  $[Co(NH_3)_4ClBr]Cl$  and  $[Co(NH_3)_4Cl_2]Br$  are ionisation isomers.
- **Sol.** The ionisation isomers dissolve in water to yield different ions and thus react differently to various reagents :

$$[Co(NH_3)_4ClBr]Cl \xrightarrow{aq}_{aq} [Co(NH_3)_4ClBr]^+ + Cl^-; Ag^+ + Cl^- \longrightarrow AgCl \downarrow (white)$$

$$[Co(NH_2),Cl_2]Br \stackrel{aq.}{\longrightarrow} [Co(NH_2),Cl_2]^+ + Br^- \quad ; Ag^+ + Br^- \longrightarrow AgBr \downarrow (yellow)$$

- **Ex.23** Which is incorrect statement ?
  - (A)  $[Co(en)_3] [Cr(CN)_6]$  will display coordination isomerism.
  - (B) [Mn(H<sub>2</sub>O)<sub>5</sub>(SCN)] will display linkage isomerism.
  - (C)  $[Co(NH_3)_5(NO_2)]SO_4$  will display ionization isomerism
  - (D) None of these.

#### Ans. (D)

Sol. (A) Coordination isomerism arises due to exchange of ligands between cationic and anionic complexes.
 (B) SCN<sup>-</sup> is an ambidentate ligand and thus can attach either through nitrogen or sulphur to central metal ion causing linkage isomerism.

(C) Exchange of  $NO_2^-$  (in coordination sphere) and  $SO_4^{2-}$  (in ionization sphere) produces ionization isomerism.

**Ex.24** Indicate the types of isomerism exhibited by the following complexes.

(i)  $K[Cr(H_2O)_2(C_2O_4)_2]$  (ii)  $[Co(en)_3]Cl_3$ 

(iii)  $[Co(NH_3)_4(NO_2)_2](NO_3)_2$  (iv)  $[Pt(NH_3)(H_2O)CI_2]$ .

**Sol.** (i) Both geometrical (cis-, trans-) and optical isomers for cis can exist.

(ii) Two optical isomers can exist.

(iii) There are 10 possible isomers. (Hint : There are geometrical, ionisation and linkage isomers possible).

- (iv) Geometrical (cis-, trans-) isomers can exist.
- **Ex.25** Out of the following two coordination entities which is chiral (optically active)? (a) cis- $[CrCl_2(ox)_2]^{3-}$  (b) trans- $[CrCl_2(ox)_2]^{3-}$ .
- **Sol.** The two entities are represented as



	<ul> <li>(b) [Fe<sup>II</sup>Cl<sub>4</sub>]<sup>2−</sup>; EAN = 2</li> <li>(c) [Co<sup>II</sup>(CN)<sub>6</sub>]<sup>4−</sup>; EAN =</li> <li>(d) [Ni<sup>II</sup>(NH<sub>3</sub>)<sub>6</sub>]<sup>2+</sup>; EAN =</li> </ul>	6 - 2 + 8 = 32 = 25 + 12 = 37 = 26 + 12 = 38							
6.	Consider the following (i) $K_2$ PtCl <sub>6</sub>	complexes : (ii) PtCl <sub>4</sub> . 2NH <sub>3</sub>	(iii) PtCl <sub>4</sub> . 3NH <sub>3</sub>	(iv) PtCl <sub>4</sub> . 5NH <sub>3</sub>					
Ans. Sol.	<ul> <li>(1) 256, 0, 97, 404</li> <li>(1)</li> <li>The electrical conduct</li> </ul>	tance in aqueous solution (2) 404, 0, 97, 256 ance of the complexes o	n are : (3) 256, 97, 0, 404 depends upon the numb	(4) 404, 97, 256, 0 er of ions given by them in the					
	aqueous solution. (i) $K_2[PtCl_6] \stackrel{aq}{\longleftarrow} 2K^*(aq) + [PtCl_6]^{2-}(aq)$ (ii) $[Pt(h L_1), Q L_1, aq] > [Pt(h L_1), Q L_1, qq]$								
	(iii) $[Pt(NH_3)_3Cl_3]Cl \stackrel{aq}{=}$ (iv) $[Pt(NH_3)_3Cl_3]Cl \stackrel{aq}{=}$	≥ [Pt(NH <sub>3</sub> ) <sub>3</sub> Cl <sub>3</sub> ]⁺ (aq) + Cl ≥ [Pt(NH <sub>3</sub> ) <sub>3</sub> Cl <sub>3</sub> ]⁺ (aq) + Cl ≥ [Pt(NH <sub>2</sub> ) <sub>2</sub> Cl]³⁺ + 3Cl⁻	- (aq)						
		Number of ions	Expected elec	trical conductance					
	(i) K <sub>a</sub> [PtCl <sub>a</sub> ]	3	Expected elec	256					
	(ii) [Pt(NH <sub>3</sub> ) <sub>2</sub> Cl <sub>4</sub> ]	0		0					
	(iii) [Pt(NH <sub>3</sub> ) <sub>3</sub> Cl <sub>3</sub> ]Cl	2		97					
	(iv) [Pt(NH <sub>3</sub> ) <sub>5</sub> Cl]Cl <sub>3</sub>	4		404					
	So, the correct option i	s (A).							
7.	Explain the following : (i) All the octahedral co (ii) $[CoF_6]^{3-}$ is paramag	omplexes of Ni <sup>2+</sup> must be netic but [Co(CN) <sub>6</sub> ] <sup>3–</sup> is di	outer orbital complexes. amagnetic.						
Sol.	<ul> <li>(i) Ni<sup>2+</sup> configuration 4s 4p 4d</li> <li>(ii) Ni<sup>2+</sup> configuration 4s 4p 4d</li> <li>(iii) In [CoF<sub>6</sub>]<sup>3-</sup>, Co<sup>3+</sup> undergoes sp<sup>3</sup>d<sup>2</sup> hybridization. Four d-orbitals are singly occupied. Hence, it is paramagnetic.</li> </ul>								
		rangement sp d nybridiza	tion						
	In [Co(CN) <sub>6</sub> ] <sup>3-</sup> , Co <sup>3+</sup> und [Co(CN) <sub>6</sub> ] <sup>3-</sup> `	3d     4s       ↑↓     ↑↓     ↑↓       Rearrangement     d²sp³ hybr	All electrons are paire	ed and thus it is diamagnetic.					
8.	The increasing order $(1) H_2O < OH^- < NCS^-$	of the crystal field splitting $< F^- < NH_3$	power of some commo (2) H₂O < NCS <sup>−</sup> < OH <sup>−</sup>	n ligands is : <nh₃< f<sup="">−</nh₃<>					
	(3) $CN^- < H_2O < OH^-$	$< NCS^{-} < H_2O^{-}$	(4) $F^- < OH^- < H_2O <$	$NCS^{-} < NH_{3}$					
Ans.	(4)								
Sol.	Spectro chemical series based on experimental data follows the order : $I^- < Br^- < SCN^- < CI^- < S^{2-} < F^- < OH^- < C_2O_4^{2-} < H_2O < NCS^- < edta^{4-} < NH_3 < en < NO_2^- < CN^- < CO$								

## **CHEMISTRY FOR NEET**

9.





[PtCl<sub>2</sub>(NH<sub>2</sub>)(OH<sub>2</sub>)]



dsp<sup>2</sup> hybrid orbitals

Four pairs of electrons from four CI-

Ma, bc have cis- and trans isomers.

- 16. In metal carbonyls the metal carbon bond length is found to be less than the expected bond length. Explain why?
- It is due to synergic interaction between metal and CO which develops partial double bond character Sol. between metal and CO.



Synergic bonding

- 17. (a) Draw all possible constitutional isomers of the compound Ru(NH<sub>3</sub>)<sub>5</sub>(NO<sub>2</sub>)Cl. Label the isomers as linkage isomers or ionization isomers.
  - (b) There are six possible isomers for a square planar palladium(II) complex that contains two Cl<sup>-</sup> and two SCN- ligands. Sketch the structures of all six, and label them according to the classification.
- (a) There are three constitutional isomers Ans.
  - (i) [Ru(NH<sub>3</sub>)<sub>5</sub>(NO<sub>2</sub>)]Cl
  - (ii) [Ru(NH<sub>3</sub>)<sub>5</sub>Cl](NO<sub>2</sub>) or [Ru(NH<sub>3</sub>)<sub>5</sub>Cl]ONO
  - (iii) [Ru(NH<sub>3</sub>)<sub>5</sub> ONO]Cl

18.

- (i) & (ii) are ionisation isomers
- (i) & (iii) are linkage isomers



Sol. cis- platin is cis[Pt(NH<sub>2</sub>)<sub>2</sub>Cl<sub>2</sub>]. So it contains platinum.