Exercise-1

Marked questions are recommended for Revision.

PART - I : SUBJECTIVE QUESTIONS

Section (A) : General introduction of complex salts and definitions to be used

- A-1. K₂SO₄ solution mixed with Cr₂(SO₄)₃ solution in 1 : 1 molar ratio gives the test of Cr³⁺ ion but CuSO₄ solution mixed with aqueous ammonia in 1 : 4 molar ratio does not give the test of Cu²⁺ ion. Explain why ?
- A-2. What is the coordination number and the oxidation state of the metal in each of the following complexes?

(a) (d)æ (g)	[AgCl₂] [–] [Co(NH₃)₃(NO₂)₃] K[Pt(NH₃)Cl₅]	, , ,	(b) (e)æ	[Cr(H₂O)₅Cl]²+ [Fe(EDTA)]⁻	;	(C) (f)ක	[Co(NCS)4] ^{2–} [Cu(en)2]SO4;
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A-3. Write the name of the following ligands and classify their denticity

(A) CH ₃ NC	(B) acac ⁻¹	(Ć) N ₃ −	(D) dien	(E) edta4-
(F) edta ^{3–}	(G) ox ^{2–}	(H) dmg ⁻¹	(I) NC [_]	(J) NO ₂ -
(K) O ^{2–}	(L) O ₂ -			

A-4. Predict the different ligating sites by drawing structures in the following ligands.

- (A) $(NO_2)^-$ (B) $(SCN)^-$ (C) $(C_2O_2S_2)^{2-}$ (D) $(OCN)^-$
 - ÌÉ) ÌNOŚ)⁻
- A-5. (a) Determine the denticity of the ligands in complexes $[Fe(C_2O_4)_3]^{3-}$ and $[Pt(en)_2]^{2+}$. What are the coordination number and the oxidation number of the central metal ion?
 - (b) Designate the coordination entities and counter ions in the coordination compounds. $K_2[Ni(CN)_4]$; [Cr(en)₃] Cl₃; Fe₄[Fe₍CN)₆]₃; [PtCl₂(en)₂] (NO₃)₂.

(c)	Identify the Lewis acid a	and Lewis base components of t	he following complexes.
	(i) [HgBr ₄] ²⁻	(ii) [Ni(H ₂ O) ₆] ²⁺	(iii) [PdCl ₂ (NH ₃) ₂]
	(iv) [Al(OH) ₄] [_]	(v) [Ag(CN) ₂] ⁻	(vi) [Cr(CO) ₆]

Section (B) : Nomenclature of coordination compounds

B-1. Name the following compounds

Name	the following compounds		
(a)æ	[Co(NH ₃) ₆]Cl ₃	Prepared in 1798 by B.M. Tassaert and considered to be first complex salt prepared.	
(b)	[Rh(NH₃)₅l]l₂	A yellow colored complex obtained by heating $[Rh(NH_3)_5(H_2O)]I_3$ above 100°C.	
(C)	[Fe(CO)₅]	A highly toxic volatile liquid.	
(d)	[Fe(C ₂ O ₄) ₃] ³⁻	The ion formed when Fe ₂ O ₃ rust is dissolved in oxalic acid, H ₂ C ₂ O ₄ .	
(e)æ	[Cu(NH ₃) ₄]SO ₄	A deep blue compound obtained when CuSO ₄ is treated with excess of NH ₃ .	
(f)	Na[Cr(OH)₄]	The compound formed when $Cr(OH)_3$ precipitate is dissolved in excess of NaOH.	
(g) کھ	[Co(gly) ₃]	A complex that contains the anion of amino acid, glycine.	
(h) 🕰	[Fe(H ₂ O) ₅ (SCN)] ²⁺	The red complex ion formed in the qualitative analysis test of Fe ³⁺ ion.	
(i) کھ	K ₂ [Hgl ₄]	Alkaline solution of this complex is called Nessler's Reagent.	
(j)	Co[Hg(SCN) ₄]	Deep blue crystalline precipitate obtained in qualitative detection of Hg ²⁺ .	
(k)	Fe4[Fe(CN)6]3	Prussian blue , deep blue colored complex obtained in detection of Fe ²⁺ .	
(I)	K ₃ [Co(NO ₂) ₆]	Potassium cobaltinitrite or Fischer salt yellow precipitate obtained in detection of Co ²⁺ .	
(m)	[Ni(dmg) ₂]	Rosy red precipitate obtained in detection of Ni ²⁺ ions.	
(n)	K ₂ [PtCl ₆]	Yellow precipitate obtained in detection of potassium ions.	
(0) (0)	Na ₂ [Fe(CN) ₅ NO ⁺]	Sodium nitroprusside used for detection of sulphide ions/sulphur.	
(p) 🔊	[Fe(H ₂ O) ₅ (NO ⁺)]SO ₄	Brown ring complex, obtained in detection of Fe ⁺¹ ions.	
(q)æ	[Cu(CN) ₄] ^{3–}	Colourless stable soluble complex obtained in detection of Cu ²⁺ on adding excess of KCN solution.	
(r)	(NH ₄) ₂ [PtCl ₆]	Only few compounds of ammonium ions are precipitate this is one of	

B-2.

	these, a yellow precipitate.	
the following compounds.		
[CoBr(en) ₂ (ONO)] ⁺	(b)æ	[Co(NH ₃) ₆][Co(ONO) ₆]
[Co(NH ₃) ₅ (CO ₃)]Cl	(d)	[Pt(NH ₃) ₄ Cl ₂][PtCl ₄]
[Co(en) ₃] ₂ (SO ₄) ₃	; (f)	[(NH ₃) ₅ Co-NH ₂ -Co(NH ₃) ₄ (H ₂ O)]Cl ₅
[Cr(CO)₅(PPh₃)]	(h)	[(CO) ₅ Mn-Mn(CO) ₅]
$Cr(n^{6}-C_{6}H_{6})_{2}$	(i)	[Co(NH ₃) ₄ (OH ₂) ₂][BF ₄] ₃
Ba[Zr(OH)2(ONO)2(ox)]	(ľ) 🏊	[Co(NH ₃) ₆][Co(C ₂ O ₄) ₃]
	[CoBr(en) ₂ (ONO)] ⁺ [Co(NH ₃) ₅ (CO ₃)]Cl [Co(en) ₃] ₂ (SO ₄) ₃ [Cr(CO) ₅ (PPh ₃)] Cr(η ⁶ -C ₆ H ₆) ₂	$ \begin{array}{c c} the following compounds. \\ [CoBr(en)_2(ONO)]^+ & (b) \\ [Co(NH_3)_5(CO_3)]Cl & (d) \\ [Co(en)_3]_2(SO_4)_3 & (f) \\ [Cr(CO)_5(PPh_3)] & (h) \\ Cr(\eta^6-C_6H_6)_2 & (j) \end{array} $

B-3. Write down the formulae of the following compounds

(a)	Tetraamminezinc(II) Nitrate	The compound formed when zinc nitrate is treated with an
		excess of ammonia
(b)	TetracarbonyInickel(0)	The first metal carbonyl(prepared in 1888) and an important
		compound in the industrial refining of nickel metal
(C) 🔊	Potassium	A compound that contains a square planar anion
	amminetrichloridoplatinate(II)	
(d)	Dicyanidoaurrate(I) ion	An ion important in the extraction of gold from its ores
(e)	Sodium hexafluoridoaluminate(III)	Called cryolite, used in the electrolytic refining of aluminium
(f) 🔊	Diamminesilver(I) ion	Ion formed when AgCI is dissolved in excess of ammonia

B-4. Write down the formulae of the following compounds

(a)	diamminetriaquahydroxidochromium (III) nitrate		
(b)	tetrakis(pyridine)platinum(II) tetraphenylborate(III)		
(c)	dibromidotetracarbonyliron (II)		
(d)	ammonium diamminetetrakis(isothiocyanato)chromate(III).		
(e)	pentaamminedinitrogenruthenium(II) chloride		
(f)	barium dihydroxidodinitrito-O-oxalatozirconate(IV)		
(g)	tetrapyridineplatinum(II) tetrachloridonickelate(II)		

Section (C) : Werner's Theory

(Initial bonding theories and EAN rule)

C-1. Werner conducted many experiments to establish the formula of complexes, one of these were conductivity measurements. On the basis of the experiments performed he obtained the following values of conductivity for different type of complexes.

Type of complex	Electrical Conductivity
Nonelectrolyte	0 – 10 (due to impurities)
1:1 Electrolyte	90 – 130
1:2 or 2:1 Electrolyte	230 – 290
1:3 or 3:1 Electrolyte	390 – 450
1:4 Electrolyte	500 – 550

On the basis of above table Match the following two columns.

	COLUMN A		COLUMN B
	Formula of compound	Conductivity	Correct Werner's Representation
(a)	PtCl ₄ .2NH ₃	6.99	(i) [Cr(NH ₃) ₅ Cl]Cl ₂
(b)	PtCl ₄ .NH ₃ .KCl	106.8	(ii) [Co(NH ₃)₅Br]Br ₂
(C)	CrCl ₃ .5NH ₃	260.2	(iii) [Cr(NH ₃) ₆]Cl ₃
(d)	PtCl ₄ .2KCl	256.8	(iv) [Pt(NH ₃) ₂ Cl ₄]
(e)	CrCl ₃ .6NH ₃	441.7	(v) [Pt(NH ₃) ₆]Cl ₄
(f)	PtCl ₄ .6NH ₃	522.9	(vi) [Pt(NH ₃) ₃ Cl ₃]Cl
(g)	CoBr ₃ .5NH ₃	257.6	(vii) K ₂ [PtCl ₆]
(h)	PtCl ₄ .3NH ₃	96.8	(viii) K[Pt(NH₃)Cl₅]

C-2. 1 g of complex [Cr(H₂O)₅Cl]Cl₂.H₂O was passed through a cation exchanger to produce HCl. The acid liberated was diluted to 1 litre. What will be the molarity of acid solution [Molecular weight of complex = 266.5] ?

	dination Compounds
C-3.æ	$ \begin{array}{llllllllllllllllllllllllllllllllllll$
C-4.æ	Arrange the following compounds in order of increasing molar conductivity.(i) $K[Co(NH_3)_2(NO_2)_4]$ (ii) $[Cr(NH_3)_3(NO_2)_3]$ (iii) $[Cr(NH_3)_5(NO_2)]_3[Co(NO_2)_6]_2$ (iv) $[Cr(NH_3)_6]Cl_3$
	 (D): Valence Bond Theory + Crystal Field Theory (Part-I) A metal complex having composition Cr(NH₃)₄Cl₂Br has been isolated in two forms A and B. The form reacts with AgNO₃ to give a white precipitate readily soluble in dilute aqueous ammonia, whereas gives a yellow precipitate soluble in concentrated ammonia. (i) Write the formulae of A and B. (ii) State hybridisation of chromium in each. (iii) Calculate their magnetic moments for each (spin-only value). (iv) Calculate the EAN for both. (v) Will they conduct electricity or not. (vi) Write the formula of the complexes formed when the precipitates dissolve in aqueous ammonia the concentrated ammonia respectively.
D-2 .	Predict the hybridisation and geometry of the following complexes. (a) $[NiBr_4]^{2-}$ (b) $[AuCl_4]^{-}$ (c) $[Pt(NH_3)_4]^{2+}$
Section E-1.	 ion (E) : Valence Bond Theory + Crystal Field Theory (Part-II) For the complex K₂[Cr(NO)(NH₃)(CN)₄]; μ = 1.73 BM. (i) Write IUPAC name. (ii) What will be structure? (iii) How many unpaired electrons are present in the central metal ion? (iv) Is it paramagnetic or diamagnetic? (v) Calculate the EAN of the complex. (vi)What will be the hybridisation of the complex?
-2. a	Predict the hybridisation and geometry of the following complexes.(a) $[Fe(CN)_6]^{3-}$ (b) $[MnBr_4]^{2-}$ (c) $[Fe(H_2O)_6]^{2+}$ (d) $[Co(SCN)_4]^{2-}$
Ξ-3.	$[Co(NH_3)_6]^{3+}$ & $[CoF_6]^{3-}$ both are complexes of Co(III), but $[Co(NH_3)_6]^{3+}$ is diamagnetic while $[CoF_6]^{3-}$ paramagnetic with $\mu = 4.90$ B.M. Explain.
E-4.a	Arrange the following in increasing order as directed. (a) (i) [CoCl ₃ (NH ₃) ₃], (ii) [CoCl(NH ₃) ₅]Cl ₂ , (iii) [Co(NH ₃) ₆]Cl ₃ , (iv) [CoCl ₂ (NH ₃) ₄]Cl - Molar conductance (b) C, N, O, F (halogen) - tendency of σ donation. (c) Br [–] , S ^{2–} , NO ₂ [–] , CO, H ₂ O, CN [–] , NH ₃ , NO ₃ [–] - strength of ligands.
E-5.	$ \begin{array}{lll} \mbox{For each of the following complexes, draw a crystal field energy-level diagram, assign the electrons to orbitals, and predict the number of unpaired electrons: (a) [CrF_6]^{3-} (b) [V(H_2O)_6]^{3+} (c) [Fe(CN)_6]^{3-} (d) [Cu(en)_3]^{2+} (e) [FeF_6]^{3-} \end{array} $
Secti	on (F) : Applications of crystal field theory
-1.	(Magnetic moment of complex, Colour of complex, Limitation, Stability of complex) Cobalt (II) is stable in aqueous solution but in the presence of complexing reagents(strong field ligands it is readily oxidised. Why ?
F-2.	The value of Δ_0 for $[Ti(H_2O)_6]^{3+}$ is found to be 240 kJ mol ⁻¹ then predict the colour of the complex usin the following table. (h = 6 × 10 ⁻³⁴ J-sec, N _A = 6 × 10 ²³ , c = 3 × 10 ⁸ m/sec)

$M_{\rm A} = 0 \times 10^{-3} = 3 \times 10^{-3} \times 10$			
Absorbed light	λ (nm) (absorbed)	Colour exhibited	
Blue	435 – 480	Yellow	
green-blue	480 - 490	Orange	
blue-green	490 – 500	Red	

green	500 - 560	purple
yellow-green	560 - 580	violet
Yellow	580 – 595	blue
Red	605 – 700	blue green

F-3. (a) [Ti(H₂O)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 form a complex absorbing light of wavelength higher than 5000 Å.

(b) Calculate the magnetic moments (spin only) of the following complexes (i) [PtCl₆]²⁻

(ii) [Cr(CO)₆]

(iii) [Ir(NH₃)₆]³⁺

(iv) [Pd(en)2]2+

Section (G) : Isomerism in Coordination compounds

(Structural Isomerism, Stereoisomerism, Geometrical Isomerism, Optical Isomerism)

G-1. What type of isomers are the following :

(i)	[Mn(CO)₅SCN]	and	[Mn(CO)₅NCS]
(ii)	[Co(en) ₃] [Cr(CN) ₆]	and	[Cr(en)3] [Co(CN)6]
(iii)	[Co(NH ₃) ₅ NO ₃]SO ₄	and	[Co(NH ₃) ₅ SO ₄]NO ₃
(iv)	[Co(H ₂ O) ₂ Cl ₂ (py) ₂]Cl	and	$[Co(H_2O)Cl_3(py)_2]H_2O$

(a) Draw all possible constitutional isomers of the compound Ru(NH₃)₅(NO₂)Cl. Label the isomers as G-2. 🖎 linkage isomers or ionization isomers.

(b) There are six possible isomers for a square planar palladium(II) complex that contains two NH₃ and two SCN⁻ ligands. Sketch the structures of all six, and label them according to the classification.

How many geometrical isomers are possible for each of the following complexes? G-3.

(a) [Pt(NH₃)₂(SCN)₂] (c) $[Co(NH_3)_3(NO_2)_3]$

(e) $[Cr Br_2 (en)_2]^+$

(b) [CoCl₂Br₂]²⁻ (tetrahedral) (d) $[Pt(en)Cl_2]$ (f) [Rh(en)₃]³⁺

G-4. Which of the following complexes can exist as enantiomers? Draw their structures (a) cis-[Co(NH₃)₄Br₂]⁺ (b) cis-[Cr(H₂O)₂(en)₂]³⁺ (c) $[Cr(g|y)_3]$ (d) [Cr(en)₃]³⁺ (e) cis- $[Co(NH_3)Cl(en)_2]^{2+}$ (f) trans-[Co(NH₃)₂(en)₂]²⁺

Section (H) : Organometallic Compounds

- Draw the structures of the following metal carbonyls H-1. (b) [Cr(CO)₆]
 - (a) [V(CO)₆] (d) [Fe(CO)5]

(e) [Ni(CO)₄]

CE

H-2.a

A-1.

The figure represents the synergic bonding interaction in metal carbonyl complex. On the basis of this explain the following :

(i) Strength of Metal-ligand bond

(ii) Bond order of CO in carbonyl complex as compared to bond order in carbon monoxide.

PART - II : ONLY ONE OPTION CORRECT TYPE

Section (A): General introduction of complex salts and definitions to be used

- Ethylene diamine is an example of a ligand :
- (A) monodentate (B) bidentate
- A-2. The donor sites of (EDTA)^{4–} are ?
 - (A) O atoms only
 - (C) Two N atoms and four O atoms
- (B) N atoms only

(C) tridentate

(D) Three N atoms and three O atoms

(D) hexadentate

(c) [Mn₂(CO)₁₀]

A-3.	Some salts although c Such salts are :	containing two different	metallic elements give to	est for one of them in solution.
	(A) complex salt	(B) double salt	(C) normal salt	(D) none
A-4.	Ligands are : (A) Lewis acids	(B) Lewis bases	(C) neutral	(D) none
A-5.	The oxidation state of N (A) +2	Mo in its oxido-complex s (B) +3	species [Mo ₂ O ₄ (C ₂ H ₄) ₂ (H (C) +4	₂ O) ₂] ^{2–} is (D) +5
A-6.	Co-ordination number (A) 4	of platinum in [Pt(NH ₃) ₄ C (B) 2	l ₂] ²⁺ ion is: (C) 8	(D) 6
A-7.a	Which of the following (A) [Cu(H ₂ O) ₄] ²⁺	s copper(I) compound ? (B) [Cu(CN)4] ^{3–}	(C) [Cu(NH ₃) ₄] ²⁺	(D) All of these
A-8.	In the complex [CoCl ₂ ((A) 6 and +3	en)₂]Br, the co-ordinatior (B) 3 and +3	number and oxidation s (C) 4 and +2	tate of cobalt are : (D) 6 and +1
A-9.১	What is the charge on t (A) +3	the complex [Cr(C ₂ O ₄) ₂ (H (B) +1	H ₂ O) ₂] formed by Cr(III) ((C) +2	? (D) –1
Section B-1.	A complex cation is for coordination number of (A) Diammineethyleneo (B) Diammineethyleneo (C) Diammineethyleneo		oxidation state) with liga of the following can be i platinum (II) ion platinate (IV) ion platinum (IV) ion	ands (in proper number so that ts correct IUPAC name ?
B-2.	Which of the following (A) Potassium tetrafluo (C) Dichlorobis(urea)co	ridooxidochromate (VI)	(B) Barium tetrafluorido (D) All are impossible.	bbromate (III)
B-3.১	The formula of the com (A) [Co(en) ₂ SO ₄]	plex tris(ethylenediamin (B) [Co(en)₃SO₄]	e)cobalt(III) sulphate is : (C) [Co(en) ₃] ₂ SO ₄	(D) [Co(en) ₃] ₂ (SO ₄) ₃
B-4.	 (A) Tetraamminechlorid (B) Chloridonitrito-O-te (C) Dichloridonitrito-O- 	ne for the compound [Cc donitrito–N–cobalt(III) ch etraamminecobalt(II) chl -tetraamminecobalt(III) donitrito–O–cobalt(III) ch	loride	
B-5.	The hypothetical comp (A) [CoCl(NH ₃) ₃ (H ₂ O) ₂] (C) [Co(NH ₃) ₃ (H ₂ O) ₂ Cl]		ridocobalt(III) chloride ca (B) [Co(NH ₃) ₃ (H ₂ O)Cl ₃] (D) [Co(NH ₃) ₃ (H ₂ O) ₃]Cl	
Section	on (C) : Werner's T			
C-1.	(Initial bonding theories EAN of a metal carbon (A) 4		c number of metal M is 20 (C) 5	6, what is the value of x? (D) 6
C-2.	The EAN of platinum in (A) 90	potassium hexachlorido (B) 86	pplatinate(IV) is (Atomic r (C) 76	number of Pt = 78) : (D) 88
C-3.	aqueous solution of the	e compound showed 4 pa		olutions in the ratio of 1 : 3. The ereas molar conductivity reveals (D) K[Co(NO ₂) ₄]

	\A/biob of the full .					
C-4 .	(A) K ₄ [Fe(CN ₆]	will exhibit maximum ion (B) [Co(NH ₃) ₆] Cl ₃	ic conductivity? (C) [Cu(NH ₃) ₄] Cl ₂	(D) [Ni (CO)4]		
C-5.2a	Which of the following (A) [Co(NH ₃) ₆]Cl ₃	shows maximum molar o (B) [Co(NH ₃) ₃ Cl ₃]	conductance ? (C) [Co(NH ₃) ₄ Cl ₂] Cl	(D) [Co(NH₃)₅Cl]Cl₂		
C-6.	The complex [Cr(H₂O). (A) Br⁻	4Br₂]Cl gives the test for (B) Cl⁻	: (C) Cr ³⁺	(D) Br⁻ and Cl⁻ both		
C-7.	Which of the following complexes will be dehydrated to relatively minimum extent by conc. H_2SO_4 uidentical condition.(A) [Cr(H_2O)_5Cl]Cl_2.H_2O(B) [Cr(H_2O)_4Cl_2]Cl.2H_2O(C) [Cr(H_2O)_6]Cl_3(D) all of these					
C-8.2	precipitated is:		,	ercentage of total chloride io		
	(A) 100	(B) 75	(C) 50	(D) 25		
C-9.	structure consistent with	th the observation is:		er molecule in the solution. Th		
	(A) [Pt(NH ₃) ₄]Cl ₄	(B) [Pt(NH ₃) ₂ Cl ₄]	(C) [Pt(NH ₃) ₅ Cl]Cl ₃	(D) [Pt(NH ₃) ₄ Cl ₂]Cl ₂		
		n has no. 'd' electrons in t (B) [Co(NH ₃) ₆] ³⁺				
)-2 .	requires the measurem (A) molar conductance	nent of	(B) optical activity	the valence bond theory ofte		
	(C) magnetic moment		(D) dipole moment			
D-3.	The magnitude of c considerably less than (A) There are only four (B) The direction of th	that in the octahedral fie ligands instead of six so he orbitals does not coin n energy (Δ) by further 2 d (B) are correct	energy (CFSE of Δ_1 ld. Because the ligand fild is only 2/3 cide with the direction of) in tetrahedral complexes 3 in tetradedral complex of the ligands. This reduces th		
D-3. D-4.	The magnitude of c considerably less than (A) There are only four (B) The direction of th crystal field stabilizatio (C) Both points (A) and (D) Both points (A) and Which of the following (A) Higher ionic radius	that in the octahedral fie ligands instead of six so he orbitals does not coin n energy (Δ) by further 2 d (B) are correct	energy (CFSE of Δ1 ld. Because the ligand fild is only 2/3 cide with the direction of /3 the stability of metal ion (B) Higher charge/size	3 in tetradedral complex of the ligands. This reduces th complexes ? ratio of the metal ion		
	The magnitude of c considerably less than (A) There are only four (B) The direction of th crystal field stabilizatio (C) Both points (A) and (D) Both points (A) and Which of the following (A) Higher ionic radius (C) Lower ionisation po	that in the octahedral fie ligands instead of six so ne orbitals does not coin n energy (Δ) by further 2 d (B) are correct d (B) are wrong factors tends to increase of the metal ion otential of the metal ion	energy (CFSE of Δ1 Id. Because the ligand fild is only 2/3 cide with the direction of /3 the stability of metal ion (B) Higher charge/size (D) Lower basicity of th	3 in tetradedral complex of the ligands. This reduces th complexes ? ratio of the metal ion		
D-4.	The magnitude of c considerably less than (A) There are only four (B) The direction of th crystal field stabilizatio (C) Both points (A) and (D) Both points (A) and (D) Both points (A) and Which of the following (A) Higher ionic radius (C) Lower ionisation points The crystal field splitting related as:	that in the octahedral fie ligands instead of six so ne orbitals does not coin n energy (Δ) by further 2 d (B) are correct d (B) are wrong factors tends to increase of the metal ion otential of the metal ion	energy (CFSE of Δt ld. Because the ligand fild is only 2/3 cide with the direction of /3 the stability of metal ion (B) Higher charge/size (D) Lower basicity of th I complex (Δ_0) and that	3 in tetradedral complex of the ligands. This reduces the complexes ? ratio of the metal ion ne ligand for tetrahedral complex (Δ_t) an		
D-4.	The magnitude of c considerably less than (A) There are only four (B) The direction of the crystal field stabilizatio (C) Both points (A) and (D) Both points (A) and (D) Both points (A) and (D) Both points (A) and (C) Lower ionic radius (C) Lower ionisation point The crystal field splitting related as: (A) $\Delta_t = \frac{4}{9} \Delta_0$	that in the octahedral fie ligands instead of six so ne orbitals does not coin n energy (Δ) by further 2. d (B) are correct d (B) are wrong factors tends to increase of the metal ion otential of the metal ion ng energy for octahedra (B) $\Delta_t = 0.5 \Delta_0$ ains $t_{2g}^6 e_g^0$ configuration	energy (CFSE of Δt ld. Because the ligand fild is only 2/3 cide with the direction of /3 the stability of metal ion (B) Higher charge/size (D) Lower basicity of th I complex (Δ_0) and that (C) $\Delta_t = 0.33 \Delta_0$	3 in tetradedral complex of the ligands. This reduces the complexes ? ratio of the metal ion ne ligand for tetrahedral complex (Δ_t) an		
D-4. D-5. D-6.	The magnitude of c considerably less than (A) There are only four (B) The direction of th crystal field stabilizatio (C) Both points (A) and (D) Both points (A) and (C) Lower ionisation points (C) Lower ionisation points	that in the octahedral fie ligands instead of six so ne orbitals does not coin n energy (Δ) by further 2. d (B) are correct d (B) are wrong factors tends to increase of the metal ion otential of the metal ion ng energy for octahedra (B) $\Delta_t = 0.5 \Delta_0$ ains $t_{2g}^6 e_{g}^0$ configuration (B) K ₄ [Fe(CN) ₆] ond Theory + Crysta	energy (CFSE of Δ_1 ld. Because the ligand fild is only 2/3 cide with the direction of /3 the stability of metal ion (B) Higher charge/size (D) Lower basicity of th I complex (Δ_0) and that (C) $\Delta_t = 0.33 \Delta_0$ s. Which of the following (C) [Co(NH ₃) ₆]Cl ₃ al Field Theory (Par ound involving :	B in tetradedral complex of the ligands. This reduces the complexes ? ratio of the metal ion he ligand for tetrahedral complex (Δ_t) and (D) $\Delta_t = \frac{9}{4} \Delta_0$ complex will be paramagnetic? (D) [Fe(CN) ₅ (O ₂)] ⁻⁵		
)-4.)-5.)-6. Sectio	The magnitude of c considerably less than (A) There are only four (B) The direction of th crystal field stabilizatio (C) Both points (A) and (D) Both points (A) and (C) Lower ionisation points (C) Lower ionisation points (A) $\Delta_t = \frac{4}{9} \Delta_0$ All the metal ions conta (A) [FeCl(CN) ₄ (O ₂)] ⁴⁻	that in the octahedral fie ligands instead of six so ne orbitals does not coin n energy (Δ) by further 2. d (B) are correct d (B) are wrong factors tends to increase of the metal ion otential of the metal ion ng energy for octahedra (B) $\Delta_t = 0.5 \Delta_0$ ains $t_{2g^6} e_{g^0}$ configuration (B) K_4 [Fe(CN) ₆] ond Theory + Crysta	energy (CFSE of Δ_1 ld. Because the ligand fild is only 2/3 cide with the direction of /3 the stability of metal ion (B) Higher charge/size (D) Lower basicity of th I complex (Δ_0) and that (C) $\Delta_t = 0.33 \Delta_0$ s. Which of the following (C) [Co(NH ₃) ₆]Cl ₃ al Field Theory (Par	B in tetradedral complex of the ligands. This reduces the complexes ? ratio of the metal ion ne ligand for tetrahedral complex (Δ_t) and (D) $\Delta_t = \frac{9}{4} \Delta_0$ complex will be paramagnetic? (D) [Fe(CN) ₅ (O ₂)] ⁻⁵		
)-4.)-5.)-6. Sectio	The magnitude of c considerably less than (A) There are only four (B) The direction of th crystal field stabilizatio (C) Both points (A) and (D) Both points (A) and (C) Lower ionisation points (C) Lower ionisation poi	that in the octahedral fie ligands instead of six so ne orbitals does not coin n energy (Δ) by further 2. d (B) are correct d (B) are wrong factors tends to increase of the metal ion otential of the metal ion ng energy for octahedra (B) $\Delta_t = 0.5 \Delta_0$ ains $t_{2g}^6 e_{g}^0$ configuration (B) K ₄ [Fe(CN) ₆] ond Theory + Crysta	energy (CFSE of Δ_1 ld. Because the ligand fild is only 2/3 cide with the direction of /3 the stability of metal ion (B) Higher charge/size (D) Lower basicity of th I complex (Δ_0) and that (C) $\Delta_t = 0.33 \Delta_0$ s. Which of the following (C) [Co(NH ₃) ₆]Cl ₃ al Field Theory (Par ound involving : (C) d ² sp ³	B in tetradedral complex of the ligands. This reduces the complexes ? ratio of the metal ion ne ligand for tetrahedral complex (Δ_t) and (D) $\Delta_t = \frac{9}{4} \Delta_0$ complex will be paramagnetic? (D) [Fe(CN) ₅ (O ₂)] ⁻⁵		

E-4.a	Match Column-I with Column-II and select the	correct a	nswer with respect to hybridisation usir
	codes given below : Column - I		Column - II
	(Complex)		(Hybridisation)
	(I) [Au F ₄] ⁻	(p)	dsp ² hybridisation
	$\frac{(1)}{(II)} \frac{[7(U+4)]}{[Cu(CN)_4]^{3-}}$	(q)	sp ³ hybridisation
			sp ³ d ² hybridisation
		(r)	
	(IV) [Fe(H ₂ O) ₅ NO] ²⁺	(s)	d ² sp ³ hybridisation
	Codes : (I) (II) (III) (IV)	((I) (II) (III) (IV)
	(I) (II) (III) (IV) (A) q p r s	· ·	(I) (II) (III) (IV) p q s r
	(C) p q r s		q p s r
E-5.	The hybridisation and unpaired electrons in [Fe (A) sp ³ d ² ; 4 (B) d ² sp ³ ; 3		on are :
E-6.	The number of unpaired electrons in d ⁶ , low spi (A) 4 (B) 2	n, octahed (C) 1	Iral complex is : (D) 0
E-7.	$ \begin{array}{ll} \mbox{Which of the following is a high spin complex ?} \\ \mbox{(A) } [Co(NH_3)_6]^{3+} & (B) \ [Fe(CN)_6]^{4-} \end{array} $	(C) [Ni(C	CN)₄] ^{2−} (D) [FeF ₆] ^{3−}
E-8.১	$ \begin{array}{llllllllllllllllllllllllllllllllllll$	(C) [Mn(ł	H ₂ O) ₆] ²⁺ (D) [Fe(CN) ₆] ^{4–}
E-9.	The number of unpaired electrons present in co (A) 5 (B) 4	mplex ion (C) 6	[FeF ₆] ³⁻ is : (D) 0
E-10.	Which of the following complexes has a geome(A) [Ni Cl4]2-(B) Ni (CO)4	ry differen (C) [Ni(C	
E-11.	Select the correct statement. (A) Complex ion [MoCl ₆] ^{3–} is paramagnetic. (C) Both (A) and (B) are correct.		plex ion [Co(en)₃]³+ is diamagnetic. e of correct.
E-12.≿	Amongst Ni(CO) ₄ , [Ni(CN) ₄] ^{2–} and NiCl ₄ ^{2–} : (A) Ni(CO) ₄ and NiCl ₄ ^{2–} are diamagnetic and [N (B) NiCl ₄ ^{2–} and [Ni(CN) ₄] ^{2–} are diamagnetic and (C) Ni(CO) ₄ and [Ni(CN) ₄] ^{2–} are diamagnetic an (D) Ni(CO) ₄ is diamagnetic and NiCl ₄ ^{2–} and [Ni	Ni(CO)4 is d NiCl4 ²⁻ i	s paramagnetic. s paramagnetic.
	on (F) : Applications of crystal field the etic moment of complex, Colour of complex, I The compound which does not show paramagn (A) [Cu(NH ₃) ₄ Cl ₂] (B) [Ag(NH ₃) ₂]Cl	imitation , etism ?	
		(C) NO	(D) NO ₂
F-2.	Among the following ions, which one has the high (A) $[Cr(H_2O)_6]^{3+}$ (B) $[Fe(H_2O)_6]^{2+}$	ghest para (C) [Cu(F	
	on (G) : Isomerism in Coordination co ural Isomerism, Stereoisomerism, Geometrical Is The complexes [Pt(NH ₃) ₄] [PtCl ₆] and [Pt(NH ₃) ₄ (A) linkage isomers (C) co-ordination isomers	omerism, (Cl ₂] [PtCl ₄] (B) optica	Optical Isomerism)

G-2.🔈	[Co(NH ₃) ₅ NO ₂]Cl ₂ and [_ Co(NH₃)₅ONO]Cl₂ are ı	related to each other as :	
	(A) geometrical isomers(C) coordination isomers		(B) linkage isomers(D) ionisation isomers	
G-3.	The number of geometr (A) 0	ical isomer of [Co(NH ₃) (B) 2)₃(NO₃)₃] are : (C) 3	(D) 4
G-4.	Geometrical isomerism (A) 2	is found in coordinatior (B) 3	n compounds having coor (C) 4 (tetrahedral)	dination number : (D) 6
G-5.	Cis-trans isomerism is monodentate ligands) : (A) Ma ₄	found in square pla (B) Ma ₃ b	anar complexes of mole (C) Ma ₂ b ₂	ecular formula ('a' and 'b' are (D) Mab ₃
G-6.	Geometrical isomerism (A) [Ag(NH ₃)(CN)]	-	(C) [PtCl4l2]	(D) [Pt(NH ₃) ₃ Cl][Au(CN) ₄]
Section H-1.	on (H) : Organometa Which one is not an org (A) RMgX		? (C) (CH₃)₄Sn	(D) C ₂ H₅ONa
H-2.	Formula of ferrocene is (A) [Fe(CN) ₆] ^{4–}		(C) [Fe(CO)₅]	(D) [Fe(C₅H₅)₂]

- 1.2 Match the column : Column-I Column-II [Fe(en)₃]³⁺ (A) d²sp³ hybridisation of central metal (p) (B) [Co(ox)₃]³⁻ (q) sp³d² hybridisation of central metal (C) [Cr(CN)₆]³⁻ (r) paramagnetic [NiCl₆]⁴⁻ (D) (s) diamagnetic (t) metal ion has +3 oxidation state
- 2. There are some coordination compounds given in column-I which may exist in different isomeric forms as given in column-II. Select the correct option(s) for the coordination compounds and their respective isomeric forms.

	Column-I		Column-II
(A)	[Co(en)2NH3Cl]SO4	(p)	Enantiomer
(B)	[Co(NH ₃) ₄ (NO ₂) ₂](NO ₃) ₂	(q)	Geometrical isomer
(C)	[Co(en)(pn)(NO ₂) ₂]Cl	(r)	Ionization isomer
(D)	[Co(gly) ₃]	(s)	Linkage isomer

Exercise-2

 $\mathbf{\hat{s}}$ Marked questions are recommended for Revision.

PART - I : ONLY ONE OPTION CORRECT TYPE

- A complex anion is formed by Osmium (in some oxidation state) with ligands (in proper number so that coordination number of osmium becomes six). Which of the following can be its correct IUPAC name?

 (A) pentachloridonitridoosmium(VI)
 (B) pentachloridonitridoosmate(VI)
 (C) azidopentachloridoosmate(VI)
 (D) None of these
- 2.
 The EAN of metal atoms in [Fe(CO)₂(NO⁺)₂] and Co₂(CO)₈ respectively are :

 (A) 34, 35
 (B) 34, 36
 (C) 36, 36
 (D) 36, 35
- **3.** Which of the following is inner orbital complex as well as diamagnetic in nature ? (A) $[Ir(H_2O)_6]^{3+}$ (B) $[Ni(NH_3)_6]^{2+}$ (C) $[Cr(NH_3)_6]^{3+}$ (D) $[Co(NH_3)_6]^{2+}$

4.	Which of the following statement is correct ? (A) The oxidation state of iron in sodium nitro prusside Na ₂ [Fe(CN) ₅ (NO))] is +3 (B) [Ag(NH ₃) ₂] ⁺ is linear in shape (C) In [Fe(H ₂ O) ₆] ³⁺ , Fe is d ² sp ³ hybridized (D) In Ni(CO) ₄ , the oxidation state of Ni is 1				
5.24	The complexK4[Zn(CN)4(O2)2] is oxidised intoK2[Zn(CN)4(O2)2], then which of the following is correct?(A) Zn(II) is oxidised into Zn(IV)(B) Paramagnetic moment decreases(C) O-O bond length increases(D) Paramagnetic moment increases				
6.	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $				
7.24	 The complex [Fe(H₂O)₅NO]²⁺ is formed in the brown ring test for nitrates when freshly prepared FeSO solution is added to aqueous solution of NO₃⁻ ions followed by addition of conc. H₂SO₄. Select correct statement about this complex. (A) Hybridisation of iron is sp³d². (B) Iron has +1 oxidation state. (C) It has magnetic moment of 3.87 B.M. confirming three unpaired electrons in Fe. (D) All the above are correct statements. 				
8.	Which of the following statements is not correct?(A) TiCl4 is a colourless compound.(B) [Cr(NH3)6)]Cl3 is a coloured compound.(C) K3[VF6] is a colourless compound.(D) [Cu(NCCH3)4][BF4] is a colourless compound.				
9.	Among TiF ₆ ²⁻ , CoF ₆ ³⁻ , Cu ₂ Cl ₂ and NiCl ₄ ²⁻ the colourless species are: (A) CoF ₆ ³⁻ and NiCl ₄ ²⁻ (B) TiF ₆ ²⁻ and CoF ₆ ³⁻ (C) NiCl ₄ ²⁻ and Cu ₂ Cl ₂ (D) TiF ₆ ²⁻ and Cu ₂ Cl ₂				
10.	The number of geometrical isomers for octahedral [Co(NH ₃) ₂ Cl ₄] ⁻ , square planar AuCl ₂ Br ₂ ⁻ are : (A) 4, 2 (B) 2, 2 (C) 3, 2 (D) 2, 3				
11.24					
12.	Both geometrical and optical isomerism are shown by : (A) $[Co(en)_2Cl_2]^+$ (B) $[Co(NH_3)_5Cl]^{2+}$ (C) $[Co(NH_3)_4Cl_2]^+$ (D) $[Cr(OX)_3]^{3-}$				
	Among the following, metal carbonyls, the C–O bond is strongest :				

PART - II : SINGLE AND DOUBLE VALUE INTEGER TYPE

- 1. Sum of denticity of following ligands are Glycinate ion, Oxalate ion, o-phenathroline, 2,2-bipyridyl, diethylenetriamine, ethylenediamine
- **2.** How many total sodium ions are present in one formula unit of sodium ethane-1,2diaminetetraacetatochromate (II) and sodium hexanitrito cobaltate (III) ?
- A blue colour complex is obtained in the analysis of Fe⁺³ having formula Fe₄[Fe(CN)₆]₃ Let a = oxidation number of Iron in the coordination sphere b = no. of secondary valencies of central iron ion. c = Effective atomic number of Iron in the coordination sphere. Then find the value of (c + a - 2b)
- **4.** Coordination number of Cr in $CrCl_3.5H_2O$ as six. The possible volumes of 1 M Ag NO₃ needed to precipitate the chlorine inoutersphere in 200 ml of 0.1 M solution of the complex is/are. Write your answer as $V_1 + V_2 + V_3 + \dots$

- 5. Ni⁺² form a complex ion in water having the formula $[Ni(H_2O)_6]^{+2}$. How many of the following statements are true for the complex ion ?
 - (i) The complex is octahedral in shape.
 - (iii) Ni⁺² has incompletely filled 3d subshell.
- (ii) The complex is diamagnetic in nature.
- (iv) Secondary valency of Ni⁺² is 6.

(v) All the bonds (metal-ligand) are perpendicular to each other.

- (vi) All the 3d orbitals of Ni+2 are degenerate
- (vii) Total spin of the complex is 1.
- (ix) The complex is more stable than $[Ni(en)_3]^{+2}$
- (viii) The hybridisation of Ni⁺² is d²sp³
 (x) Effective atomic number of Ni⁺² is 36.

6. How many of the following is correctly matched complex ?

	Complex	Oxidation no. on central metal	Electronic configuration
(a)	K ₃ [Co(C ₂ O ₄) ₃]	+3	t_{2g}^6
(b)	(NH ₄) ₂ [CoF ₄]	+2	$t_{2g}^5 e_{g}^2$
(C)	Cis - [Cr(en) ₂ Cl ₂]Cl	+3	$t_{2g}^{3} = e_{g}^{0}$
(d)	[Mn(H ₂ O) ₆]SO ₄	+2	$t_{2g}^{3} e_{g}^{2}$

8. The number of coordination isomers possible for $[Fe(NH_3)_6]^{3+}$ $[Cr(C_2O_4)_3]^{3-}$ is _____

- **9.** Find the sum of number of geometrical isomers for following complexes.
 - (a) $[CoCl_2Br_2]^{2-}$ (b) $[Rh(en)_3]^{3+}$ (c) $[Cr(en)_2 Br_2]^+$
 - (d) [Pt en Cl₂] (e) [Co(NH₃)₃(NO₂)₃]
- **10.** What is the sum of bond order of Fe–C bond and C–O bond in $Fe(CO)_5$?
- 11. How many isomeric forms are possible for the octahedral complex, [Rh(en)₂(NO₂)(SCN)]⁺?

PART - III : ONE OR MORE THAN ONE OPTIONS CORRECT TYPE

- 1. Which of the following statement(s) are incorrect ?
 - (A) Those additional compounds which lose their identity in solution are called double salts.
 - (B) In K_3 [Fe(CN)₆] Fe²⁺ and CN⁻ ion can give quantitative identification test.
 - (C) $[KAI(SO_4)_2.12H_2O]$ is a coordination compound.
 - (D) All acids are lewis acids and σ donors.
- The effective atomic number of Co(CO)₄ is 35 and hence is less stable. It attains stability by
 (A) oxidation of Co
 (B) reduction of Co
 (C) dimerization
 (D) none
- Select the correct statements ;
 (A) Potassium ferrocyanide and potassium ferricyanide can be differentiated by measuring the solid state magnetic moment.

(B) The complex $[Co(NH_3)_5Br]SO_4$ and $[Co(NH_3)_5SO_4]Br$ can be differentiated by adding aqueous solution of barium chloride

(C) The complex $[Co(NH_3)_5CI]Br$ and $[Co(NH_3)_5Br]CI$ can be differentiated by adding aqueous solution of silver nitrate.

(D) the complex $[\text{Co}(\text{NH}_3)_6]\text{Cl}_3$ and $[\text{Co}(\text{NH}_3)_5\text{Cl}]\text{Cl}_2$ can be differentiated by measuring molar conductance

- 4. Consider the following statements : S1: [MnCl6]³⁻, [FeF6]³⁻ and [CoF6]³⁻ are paramagnetic having four, five and four unpaired electrons respectively. S_2 : Low value of formation constant of a complex indicates its high thermodynamic stability. S_3 : The crystal field splitting Δ_0 , depends upon the field produced by the ligand and charge on the metal ion. and arrange in the order of true/ false. (D) T F F (A) T T T(B) T F T (C) FTF 5.2 Which of the following is/are correctly matched ? (A) $[Ni(CO)_4]$ - dsp² and diamagnetic. (B) $[Ni(en)_3](NO_2)_2 - sp^3d^2$ and two unpaired electrons. (C) $[V(NH_3)_6]CI_3 - sp^3d^2$ and two unpaired electrons. (D) $[Mn(NO^+)_3(CO)]$ - sp³ and diamagnetic. 6. Which of the following statement(s) is/are correct with respect to the crystal field theory ? (A) It considers only the metal ion d-orbitals and gives no consideration at all to other metal orbitals. (B) It cannot account for the π bonding in complexes. (C) The ligands are point charges which are either ions or neutral molecules (D) The magnetic properties can be explained in terms of splitting of d- orbitals in different crystal field. 'Spin only' magnetic moment of Ni in [Ni(dmg)₂] is same as that found in : 7.2 (A) Ni in [NiCl₂ (PPh₃)₂] (B) Mn in [MnO₄]-(C) Co in [CoBr₄]²⁻ (D) Pt in $[Pt(H_2O)_2Br_2]$ 8. Which complex of the following pairs has the larger value of Δ_0 ? (i) $[Co(CN)_6]^{3-}$ and $[Co(NH_3)_6]^{3+}$ (ii) [Co(H₂O)₆]³⁺ and [Rh(H₂O)₆]³⁺ (iii) [Co(H₂O)₆]³⁺ and [Co(H₂O)₆]²⁺ (iv) [Co(NH₃)₆]³⁺ and [CoF₆] ³⁻ Select the correct one (A) $[Co(CN)_6]^{3-} > [Co(H_2O)_6]^{3+}$ (B) $[Co(H_2O)_6]^{2+} < [Co(H_2O)_6]^{3+}$ (D) $[Co(NH_3)_6]^{3+} < [CoF_6]^{3-}$ (C) $[Co(H_2O)_6]^{3+} > [Rh(H_2O)_6]^{3+}$ 9. Which of the following isomerisms is/are shown by the complex [CoCl₂(OH₂)₂(NH₃)₂]Br ? (D) optical (A) Ionization (B) Linkage (C) Geometrical Three arrangements are shown for the complex [Co(en)(NH₃)₂Cl₂]⁺. Pick up the wrong statement. 10.2 CI CI NH H₂N NH ŃН. ċι ĊI (I) (II) (III) (A) I and II are geometrical isomers (B) II and III are optical isomers (C) I and III are optical isomers (D) II and III are geometrical isomers 11. Consider the following complexies $[V(CO)_6]^-$, $[Cr(CO)_6]$ and $[Mn(CO)_6]^+$. Then incorrect statement (s) about metal carbonyls is /are. (A) 'C–O' bond is strongest in the cation and weakest in the anion (B) 'C-O' bond order is less in the cation than in anion. (C) 'C–O' bond longer in the cation than in anion or neutral carbonyl. (D) 'M-C' bond order is higher in the cation than in anionic or neutral carbonyl.
- **12.**Following Sidwick's rule of EAN, $Co(CO)_x$ will be :
(A) $Co_2(CO)_4$ (B) $Co_2(CO)_3$ (C) $Co_2(CO)_8$ (D) $Co_2(CO)_{10}$

PART - IV : COMPREHENSION

	Read the following	passage carefully and	answer the questior	1S.	
Comp	One method involve formula of the compl concentrated H ₂ SO ₄	s treating the complex version of the second s	with known reagents isomer of the comple uffers loss in weight a	o find out the structure of complexes. and from the nature of reaction, the x Co(en) ₂ (H ₂ O)Cl ₂ Br, on reaction with and on reaction with AgNO ₃ solution it	
1.	The <u>correct</u> formula (A) [CoClBr(en) ₂]H ₂ C (C) [Co Br(en) ₂ (H ₂ O))	(B) [CoCl(en) ₂ (H ₂ (D) [CoBrCl(en) ₂]	/-	
2.		e coordination sphere o lex ion (due to spin only) (B) 5.9 BM		be replaced by F⁻, then the magnetic (D) 1.73 BM	
3.24		nds in the coordination s to spin only) will be : (B) 0.0 BM	sphere be replaced by (C) 4.9 BM	y NO₂ ⁻ , then the magnetic moment of (D) 5.9 BM	
4.		al complex is treated wit PbCl ₂) formed will be : (B) 1.0	th excess Pb(NO ₃) ₂ s (C) 0.0	solution, then the number of moles of (D) 3.0	
5.	The number of geor complex): (A) 2	(B) 3	formula of the above (C) 4	e original complex are (including the (D) 1	
Comp	Ni ²⁺ (aq.) + Dimethyl	q.) \longrightarrow Complex (X). glyoxime $\xrightarrow{NH_4OH}$ C ber of cobalt and nickel		Y are four.	
6.24	The IUPAC names of the complexes (X) and (Y) are respectively : (A) tetrathiocyanato-S-cobalt(II) and bis(dimethylglyoximate) nickel(II). (B) tetrathiocyanato-S-cobaltate (II) and bis(dimethylglyoximato)nickel (II). (C) tetrathiocyanato-S-cobaltate (II) and bis(dimethylglyoximato)nickelate(II). (D) tetrathiocyanato-S-cobaltate(III) and bis(dimethylglyoximato)nickel(II).				
7.24	The geometry of con (A) tetrahedral and s (B) both tetrahedral. (C) square planar an (D) both square plan	d tetrahedral	espectively :		
8.24	(A) (X) is paramagne (B) (Y) is diamagneti	atement for the complexe tic with two unpaired ele c and shows intermolecu tic with three unpaired e	ctrons. Ilar H-bonding.	amagnetic.	

(D) (X) and (Y) both are diamagnetic.

Comprehension # 3

Q.9, Q.10 and Q.11 by appropriately matching the information given in the three columns of the following table.

Let us consider following columns				
Column 1	Column 2	Column 3		
μ (in B.M>)	Hybridisation state	No. of geometrical isomers		
(l) μ = 2.83 B.M.	(i) sp ³	(P) 2		
(II) μ = 5.93 B.M.	(ii) sp ³ d ²	(Q) 3		
(III) μ = 3.88 B.M.	(iii) d²sp³	(R) 4		
(IV) $\mu = 0$ B.M.	(iv) dsp ²	(S) 5		
[Nate: Atomic Number of $O_{2} = O(4)/(-O_{2} - O(4))$]				

[Note: Atomic Number of Cr = 24, V = 23, Pt = 78)]

9.	About (CrCl₃(NH₃)₃) wł (A) (III), (iii), P	nich of following combina (B) (II), (iv), Q	tion is correct ? (C) (IV), (i), R	(D) (I), (ii), S
10.	Correct combination fo (A) (II), (i), P	r [VCl₂(NO₂)₂(NH₃)₂]⁻. (B) (I), (iii), S	(C) (III), (ii), R	(D) (IV), (iv), Q
11.	Correct combination fo (A) (II), (iii), Q	r [PtCl ₂ (NH ₃) ₂] is : (B) (I), (iv), S	(C) (IV), (iv), P	(D) (III), (ii), R

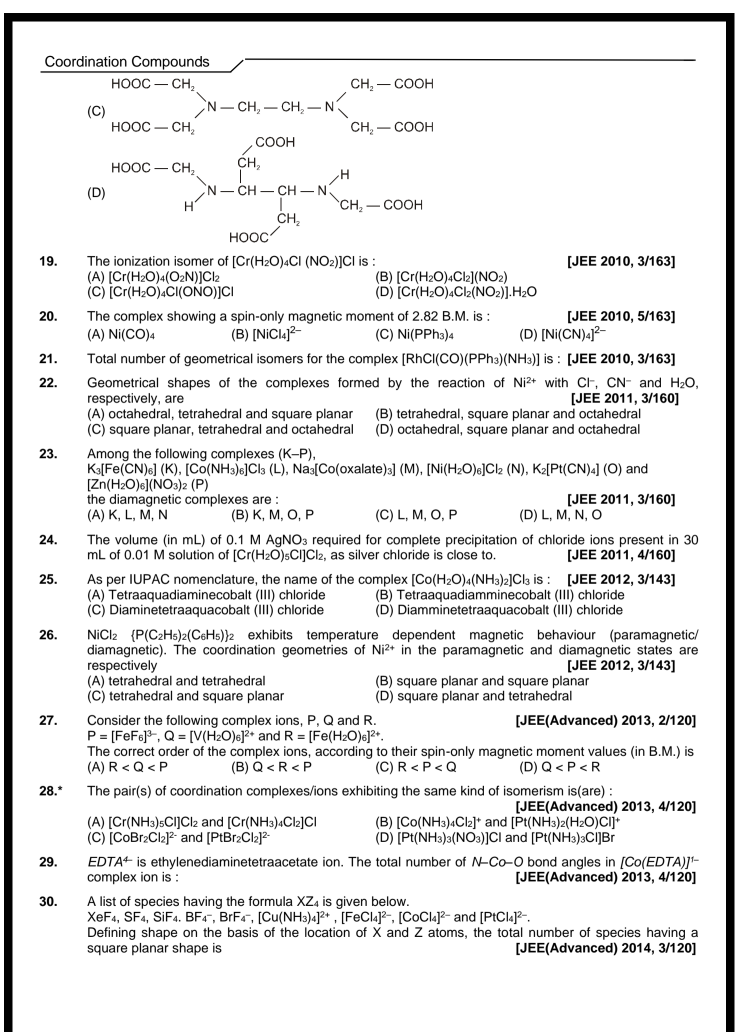
Exercise-3

* Marked Questions may have more than one correct option.

PART - I : JEE (ADVANCED) / IIT-JEE PROBLEMS (PREVIOUS YEARS)

			•
The species having tetra (A) [PdCl ₄] ²⁻		(C) [Pd(CN) ₄] ^{2–}	[JEE 2004, 3/84] (D) [NiCl4] ²⁻
The spin magnetic mom (A) $\sqrt{3}$	thent of cobalt in the composition (B) $\sqrt{8}$		s : [JEE 2004, 3/84] (D) √24
ammonia solution, a bri (a) Draw the structure o (b) Write the oxidation s	ght red coloured precipit f bright red substance. tate of nickel in the subs	ate is obtained. tance and hybridisation.	[JEE 2004, 4/60]
(A) Geometrical and ion	ization	Iral [Co(NH ₃) ₄ Br ₂]Cl ? (B) Geometrical and opt (D) Geometrical only	[JEE 2005, 3/84] tical
The bond length in CO i (A) 1.158 Å	s 1.128 Å. What will be t (B) 1.128 Å	he bond length of CO in (C) 1.178 Å	Fe(CO) ₅ ? [JEE 2006, 5/184] (D) 1.118 Å
a b a b a b a b a b a b a b a b a b a b			
	name of the complex an The species having tetra (A) $[PdCl_4]^{2-}$ The spin magnetic mom (A) $\sqrt{3}$ When dimethyl glyoxim ammonia solution, a brig (a) Draw the structure of (b) Write the oxidation s (c) State whether the su Which kind of isomerism (A) Geometrical and ionization (C) Optical and ionization The bond length in CO in (A) 1.158 Å	name of the complex and draw the structure of a The species having tetrahedral shape is : (A) [PdCl ₄] ²⁻ (B) [Ni(CN) ₄] ²⁻ The spin magnetic moment of cobalt in the comp (A) $\sqrt{3}$ (B) $\sqrt{8}$ When dimethyl glyoxime is added to the aque ammonia solution, a bright red coloured precipita (a) Draw the structure of bright red substance. (b) Write the oxidation state of nickel in the subs (c) State whether the substance is paramagnetic Which kind of isomerism is exhibited by octahed (A) Geometrical and ionization (C) Optical and ionization The bond length in CO is 1.128 Å. What will be t (A) 1.158 Å (B) 1.128 Å	(A) $[PdCl_4]^{2-}$ (B) $[Ni(CN)_4]^{2-}$ (C) $[Pd(CN)_4]^{2-}$ The spin magnetic moment of cobalt in the compound, Hg $[Co(SCN)_4]$ is(A) $\sqrt{3}$ (B) $\sqrt{8}$ (C) $\sqrt{15}$ When dimethyl glyoxime is added to the aqueous solution of nickel(II ammonia solution, a bright red coloured precipitate is obtained.(a) Draw the structure of bright red substance.(b) Write the oxidation state of nickel in the substance and hybridisation.(c) State whether the substance is paramagnetic or diamagnetic.Which kind of isomerism is exhibited by octahedral $[Co(NH_3)_4Br_2]CI$?(A) Geometrical and ionization(B) Geometrical and op(C) Optical and ionization(D) Geometrical onlyThe bond length in CO is 1.128 Å. What will be the bond length of CO in

Coordination Compounds 7. The IUPAC name of complexes 'A' & 'B' are respectively : [JEE 2006, 5/184] (A) Potassium tetracyanonickelate(II) and Potassium tetrachloronickelate(II) (B) Potassium tetracyanonickel(II) and Potassium tetrachloronickel(II) (C) Potassium cyanonickelate(II) and Potassium chloronickelate(II) (D) Potassium cyanonickel(II) and Potassium chloronickel(II) 8. The hybridisation of both complexes are : [JEE 2006, 5/184] (A) dsp² (B) $sp^2 \& dsp^2$ (C) $dsp^2 \& sp^3$ (D) both sp³ What are the magnetic nature of 'A' & 'B' ? [JEE 2006, 5/184] 9. (A) Both diamagnetic. (B) 'A' is diamagnetic & 'B' is paramagnetic with one unpaired electrons. (C) 'A' is diamagnetic & 'B' is paramagnetic with two unpaired electrons. (D) Both are paramagnetic. 10. Among the following metal carbonyls, the C-O bond order is lowest in : [JEE 2007, 3/162] (D) [Fe(CO)5] (A) [Mn(CO)₆]⁺ (B) [V(CO)₆]⁻ $(C) [Cr(CO)_6)]$ 11. Match the complexes in Column-I with their properties listed in Column-II. [JEE 2007, 6/162] Column-I Column-II Geometrical isomers (A) [Co(NH₃)₄(H₂O)₂]Cl₂ (p) (B) $[Pt(NH_3)_2Cl_2]$ (q) Paramagnetic (C) [Co(H₂O)₅Cl]Cl (r) Diamagnetic Metal ion with +2 oxidation state (D) [Ni(H₂O)₆]Cl₂ (s) 12. The IUPAC name of [Ni(NH₃)₄] [NiCl₄] is : [JEE 2008. 3/163] (A) Tetrachloronickel(II) tetraamminenickel (II) (B) Tetraamminenickel(II) tetrachloronickel (II) (C) Tetraamminenickel(II) tetrachloronickelate (II) (D) Tetraamminenickel(II) tetrachloronickelate (0) 13. Both [Ni(CO)₄] and [Ni(CN)₄]²⁻ are diamagnetic. The hybridisation of nickel in these complexes, [JEE 2008, 3/163] respectively, are : (A) sp^3 , sp^3 (B) sp³, dsp² (C) dsp^2 , sp^3 (D) dsp^2 , sp^2 14. Statement-1: The geometrical isomers of the complex [M(NH₃)₄Cl₂] are optically inactive, and Statement-2: Both geometrical isomers of the complex [M(NH₃)₄Cl₂] possess axis of symmetry. [JEE 2008, 3/163] (A) Statement-1 is true, statement-2 is true; statement-2 is a correct explanation for statement-1. (B) Statement-1 is true, statement-2 is true; statement-2 is NOT a correct explanation for statement-1. (C) Statement-1 is true, Statement-2 is false. (D) Statement-1 is false, Statement-2 is true. 15. Statement-1: [Fe(H₂O)₅NO]SO₄ is paramagnetic, and **Statement-2**: The Fe in [Fe(H₂O)₅NO]SO₄ has three unpaired electrons. [JEE 2008, 3/163] (A) Statement-1 is true, statement-2 is true; statement-2 is a correct explanation for statement-1. (B) Statement-1 is true, statement-2 is true; statement-2 is NOT a correct explanation for statement-1. (C) Statement-1 is true, Statement-2 is false. (D) Statement-1 is false, Statement-2 is true. 16.* The compound(s) that exhibit(s) geometrical isomerism is(are) : [JEE 2009, 4/160] (A) $[Pt(en)Cl_2]$ (B) $[Pt(en)_2]Cl_2$ (C) $[Pt(en)_2Cl_2]Cl_2$ (D) [Pt(NH₃)₂Cl₂] 17. The spin only magnetic moment value (in Bohr magneton units) of Cr(CO)₆ is : [JEE 2009, 3/160] (A) 0 (B) 2.84 (C) 4.90 (D) 5.92 The correct structure of ethylenediaminetetraacetic acid (EDTA) is : 18. [JEE 2010, 3/163] $HOOC - CH_2$ $CH_2 - COOH$ ÌΝ — CH = CH — Ν Ĵ (A) HOOC - CH, CH₂ — COOH COOH HOOC N — CH, — CH, — N 🤇 (B) HOOC COOH



31. Match each coordination compound in List-I with an appropriate pair of characteristics from List-II and select the correct answer using the code given below the lists. **[JEE(Advanced) 2014, 3/120]** $\{en = H_2NCH_2CH_2NH_2 : atomic numbers : Ti = 22; Cr = 24; Cp = 27; Pt = 78\}$

1011							
	List-I				List-II		
Ρ.	[Cr(N⊦	13)4Cl2)Cl]	1.	Paramagnetic and exhibits ioni	sation iso	merism
Q.	[Ti(H ₂ C	D)₅CI](NC) 3)2	2.	Diamagentic and exhibits cis-tr	ans isome	erism
R.	[Pt(en)	(NH ₃)Cl]	NO ₃	3.	Paramagentic and exhibits cis-	trans ison	nerism
S.	[Co(NH	H ₃) ₄ (NO ₃)2]NO3	4.	Diamagentic and exhibits ionis	ation isom	erism
Code	e :						
	Р	Q	R	S	P Q	R	S
(A)	4	2	3	1	(B) 3 1	4	2
(C)	2	1	3	4	(D) 1 3	4	2

- 32. For the octahedral complexes of Fe³⁺ in SCN⁻ (thiocyanato-S) and in CN⁻ ligand environments, the difference between the spin-only magnetic moments in Bohr magnetons (when approximated to the nearest integer) is : [Atomic number of Fe = 26] [JEE(Advanced) 2015, 4/168]
- 33. In the complex acetylbromidodicarbonylbis(triethylphosphine)iron(II), the number of Fe–C bond(s) is [JEE(Advanced) 2015, 4/168]
- **34.** Among the complex ions, $[Co(NH_2-CH_2-CH_2-NH_2)_2Cl_2]^+$, $[CrCl_2(C_2O_4)_2]^{3-}$, $[Fe(H_2O)_4(OH)_2]^+$, $[Fe(NH_3)_2(CN)_4]^-$, $[Co(NH_2-CH_2-CH_2-NH_2)_2$ (NH₃)Cl]²⁺ and $[Co(NH_3)_4(H_2O)Cl_2]^{2+}$, the number of complex ion(s) that show(s) *cis-trans* isomerism is : **[JEE(Advanced) 2015, 4/168]**
- **35.** Among [Ni(CO)₄], [NiCl₄]²⁻, [Co(NH₃)₄Cl₂]Cl, Na₃[CoF₆], Na₂O₂ and CsO₂, the total number of paramagnetic compounds is : (A) 2 (B) 3 (C) 4 (D) 5
- **36.** The number of geometric isomers possible for the complex $[CoL_2Cl_2]^-$ (L = H₂NCH₂CH₂O⁻) is [JEE(Advanced) 2016, 3/124]

37. The geometries of the ammonia complexes of Ni²⁺, Pt²⁺ and Zn²⁺, respectively, are

[JEE(Advanced) 2016, 3/124]

(A) octahedral, square planar and tetrahedral(C) tetrahedral, square planar and octahedral

(B) square planar, octahedral and tetrahedral (D) octahedral, tetrahedral and square planar

38.* Addition of excess aqueous ammonia to a pink coloured aqueous solution of MCl₂.6H₂O (**X**) and NH₄Cl gives an octahedral complex **Y** in the presence of air. In aqueous solution, complex **Y** behaves as 1 : 3 electrolyte. The reaction of **X** with excess HCl at room temperature results in the formation of a blue coloured complex **Z**. The calculated spin only magnetic moment of **X** and **Z** is 3.87 B.M., whereas it is zero for complex **Y**. [JEE(Advanced) 2017, 4/122]

Among the following options, which statement(s) is (are) correct?

- (A) The hybridization of the central metal ion in \mathbf{Y} is d^2sp^3
- (B) Addition of silver nitrate to Y gives only two equivalents of silver chloride
- (C) When **X** and **Z** are in equilibrium at 0°C, the colour of the solution is pink
- (D) **Z** is a tetrahedral complex

39.* The correct statement(s) regarding the binary transition metal carbonyl compounds is (are) (Atomic numbers: Fe = 26, Ni = 28) [JEE(Advanced) 2018, 4/128] (A) Total number of valence shall electrons at motal centre in Eq.(CO) or Ni(CO), is 16

- (A) Total number of valence shell electrons at metal centre in $Fe(CO)_5$ or $Ni(CO)_4$ is 16
- (B) These are predominantly low spin in nature
- (C) Metal-carbon bond strengthens when the oxidation state of the metal is lowered
- (D) The carbonyl C-O bond weakens when the oxidation state of the metal is increased

40. Among the species given below, the total number of diamagnetic species is

[JEE(Advanced) 2018, 3/120]

H atom, NO₂ monomer, O₂⁻ (superoxide), dimeric sulphur in vapour phase, Mn₃O₄, (NH₄)₂[FeCl₄], (NH₄)₂[NiCl₄], K₂MnO₄, K₂CrO₄

41. The ammonia prepared by treating ammonium sulphate with calcium hydroxide is completely used by NiCl₂.6H₂O to form a stable coordination compound. Assume that both the reactions are 100% complete. If 1584 g of ammonium sulphate and 952 g of NiCl₂.6H₂O are used in the preparation, the combined weight (in grams) of gypsum and the nickel-ammonia coordination compound thus produced is _____. (Atomic weights in g mol⁻¹: H = 1, N = 14, O = 16, S = 32, Cl = 35.5, Ca = 40, Ni = 59)

[JEE(Advanced) 2018, 3/120]

[JEE(Advanced) 2018, 4/120]

42.* The correct option(s) regarding the complex [Co(en)(NH₃)₃(H₂O)]³⁺

- $(en = H_2NCH_2CH_2NH_2)$ is (are)
- (A) It has two geometrical isomers
- (B) It will have three geometrical isomers if bidentate 'en' is replaced by two cyanide ligands

(1)

(2)

(3)

(4)

(5)

(6)

LIST-II

[FeF6]4-

[FeCl₄]²⁻

Ni(CO)₄ [Ni(CN)4]2-

Ti(H₂O)₃Cl₃

[Cr(NH₃)₆]³⁺

- (C) It is paramagnetic
- (D) It absorbs light at longer wavelength as compared to [Co(en)NH₃)₄]³⁺

43. Match each set of hybrid orbitals from LIST-I with complex(es) given in LIST-II. [JEE(Advanced) 2018, 3/120]

- LIST-I dsp²
- (P) (Q) sp³
- sp³d² (R)
- (S) d²sp³

The correct option is :

(A) $P \rightarrow 5$; $Q \rightarrow 4$, 6; $R \rightarrow 2$, 3; $S \rightarrow 1$ (C) $P \rightarrow 6$; $Q \rightarrow 4, 5$; $R \rightarrow 1$; $S \rightarrow 2, 3$ (B) $P \rightarrow 5, 6; Q \rightarrow 4; R \rightarrow 3; S \rightarrow 1, 2$ (D) $P \rightarrow 4, 6; Q \rightarrow 5, 6; R \rightarrow 1, 2; S \rightarrow 3$

PART - II : JEE (MAIN) / AIEEE PROBLEMS (PREVIOUS YEARS)

JEE(MAIN) OFFLINE PROBLEMS

One mole of Co(NH₃)₅Cl₃ gives 3 moles of ions on dissolution in water. One mole of this reacts with two 1. moles of AgNO3 to give two moles of AgCI. The complex is : [AIEEE 2003, 3/225] (1) [Co(NH₃)₄Cl₂]Cl.NH₃ (2) [Co(NH₃)₄Cl]Cl₂.NH₃ (3) [Co(NH₃)₅Cl]Cl₂ (4) [Co(NH₃)₃Cl₃].2NH₃ 2. Ammonia forms the complex $[Cu(NH_3)_4]^{2+}$ with copper ions in alkaline solution but not in acid solution. The reason for it is : [AIEEE 2003, 3/225] (1) in alkaline solution Cu(OH)₂ is precipitated which is soluble in excess of alkali. (2) copper hydroxide is amphoteric. (3) in acidic solution hydration protects Cu²⁺ ions. (4) in acidic solution protons coordinates with ammonia molecule forming NH₄⁺ ions and NH₃ molecules are not available. In the coordination compound K₄[Ni(CN)₄], the oxidation state of nickel is : 3. [AIEEE 2003, 3/225] (1) - 1(3) + 1(2) 0(4) + 2The co-ordination number of a central metal atom in a complex is determined by : [AIEEE 2004, 3/225] 4. (1) the number of only anionic ligands bonded to metal ion (2) the number of ligands around a metal ion bonded by pi bonds (3) the number of ligands around a metal ion bonded by sigma and pi bonds (4) the number of ligands around a metal ion bonded by sigma bonds 5. Which one is an outer orbital complex ? [AIEEE 2004, 3/225] (1) [Ni(NH₃)₆]²⁺ (2) [Mn(CN)₆]^{4–} (3) [Co(NH₃)₆]³⁺ (4) [Fe(CN)₆]⁴⁻ Co-ordination compounds have great importance in biological systems. In this context, which statement 6. is incorrect? [AIEEE 2004, 3/225] (1) Carboxypeptidase-A is an enzyme and contains zinc. (2) Haemoglobin is the red pigment of blood and contains iron. (3) Cyanocobalmin is B₁₂ and contains cobalt. (4) Chlorophylls are green pigments in plants and contain calcium. 7. Which one has largest number of isomers ? [AIEEE 2004, 3/225] (3) [Ir(PhR₃)₂H(CO)]²⁺ (4) [Ru(NH₃)₄Cl₂]⁺ (1) $[Co(en)_2Cl_2]^+$ (2) [Co(NH₃)₅Cl]²⁺

Cool	rdination Compounds	_/		
8.	The correct order of ma (1) Fe(CN)6 ⁴⁻ > [CoCl4] (3) [Fe(CN)6] ⁴⁻ > [MnC		oin value in BM) among i (2) [MnCl₄]²- > [Fe(CN) (4) [MnCl₄]²- > [CoCl₄]²	$ _{6} ^{4-} > [CoCl_{4}]^{2-}$
9.	The oxidation state of ((1))	Cr in [Cr(NH ₃) ₄ Cl ₂]+ is : (2) +1	(3) +2	[AIEEE 2005, 1½/225] (4) +3
10.	The IUPAC name of K (1) Potassium hexacya (3) Potassium hexacya	noferrate(II)	(2) Potassium hexacya (4) Tripotassium hexac	
11.	Which of the following (1) [Cu(NH ₃) ₄] ²⁺	will show optical isomeris (2) [ZnCl ₄] ^{2–}	sm ? (3) [Cr(C ₂ O ₄) ₃] ^{3–}	[AIEEE 2005, 3/225] (4) [Co(CN) ₆] ³⁻
12.	Which one of the follow	ving complexes would ex	hibit the lowest value of	paramagnetic behaviour ?
	(1) [Co(CN) ₆] ³⁻	(2) [Fe(CN) ₆] ^{3–}	(3) [Mn(CN)6] ³⁻	[AIEEE 2005, 3/225] (4) [Cr(CN) ₆] ³⁻
13.	The value of 'spin only one is: (Assuming octa (1) d ⁴ (in strong field lig (3) d ³ (in weak as well	hedral complex) gand)	ne of the following config (2) d ⁴ (in weak field liga (4) d ⁵ (in strong field lig	
14.	Nickel (Z = 28) combines with a uninegative monodentate ligand X ⁻ to form a paramagnetic cc [NiX ₄] ²⁻ . The number of unpaired electron(s) in the nickel and geometry of this complex ion respectively : (1) one, tetrahedral (2) two, tetrahedral (3) one, square planar (4) two, square planar			
15.	The IUPAC name for the IUPAC name for the (1) Nitrito-N-pentaamm (3) Pentaamminenitrito			
16.	In Fe(CO)₅, the Fe – C (1) π-character only (3) ionic character only		[AIEEE 2006, 3/165] (2) both σ and π characters (4) σ -character only	
17.	How many EDTA (et complex with a Ca ²⁺ io (1) Six		acid) molecules are re (3) One	equired to make an octahedral [AIEEE 2006, 3/165] (4) Two
18.		tic moment (in units of B	. ,	 ²⁺ in aqueous solution would be [AIEEE 2006, 3/165] (4) 1.73
19.	Which one of the follow (1) [NiCl₄] ^{2–} (At. no. Co = 27, Ni = 2	ving has a square planar (2) [PtCl₄] ^{2–} 28, Fe = 26, Pt = 78)	geometry? (3) [CoCl ₄] ^{2–}	[AIEEE 2007, 2/120] (4) [FeCl ₄] ²⁻
20.	The coordination number and the oxidation state of the element 'E' in the complex [E(en)2(C2O4)](when 'en' is ethylene diamine) are, respectively,[AIEEE 2008, 3/10](1) 4 and 2(2) 4 and 3(3) 6 and 3(4) 6 and 2			[AIEEE 2008, 3/105]
21.	In which of the follow highest? (1) [Co(C ₂ O ₄) ₃] ^{3–}	ing octahedral complex (2) [Co(H ₂ O) ₆] ³⁺	es of Co (at no. 27), w (3)[Co(NH ₃) ₆] ³⁺	vill the magnitude of ∆₀ be the [AIEEE 2008, 3/105] (4) [Co(CN) ₆] ^{3–}
22.	Which of the following (1) [Co (en) (NH ₃) ₂] ²⁺	has an optical isomer ? (2) [Co(H ₂ O) ₄ (en)] ³⁺	(3) [Co (en) ₂ (NH ₃) ₂] ³⁺	[AIEEE 2009, 4/144] (4) [Co (NH ₃) ₃ Cl] ⁺
23.	(1) [Pd(PPh ₃) ₂ (NCS) ₂]			[AIEEE 2009, 4/144]

	-			
24.	exchanger. The chlorid		on were treated with exe	$10l^{-1}$) is passed through a cation cess of AgNO ₃ to give 4.78 g of nass of Ag = 108 u) [AIEEE 2010, 8/144]
	(1) [Co(NH ₃) ₆] Cl ₃	(2) [CoCl ₂ (NH ₃) ₄] Cl	(3) [CoCl ₃ (NH ₃) ₃]	(4) [CoCl(NH ₃) ₅] Cl ₂
25.	Which one of the follov (1) [Zn(en)(NH ₃) ₂] ²⁺ (en = ethylenediamine)	ving has an optical isome (2) [Co(en)₃]³+)	er ? (3) [Co(H ₂ O) ₄ (en)] ³⁺	[AIEEE 2010, 4/144] (4) [Zn(en) ₂] ²⁺
26.	(1) The complex involv(2) The complex is part(3) The complex is an offer the		nd is octahedral in shape	
27.	The magnetic moment (1) 1.82 BM	(spin only) of [NiCl ₄] ^{2–} is (2) 5.46 BM	: (3) 2.82 BM	[AIEEE 2011, 4/144] (4) 1.41 BM
28.	Which among the follo	owing will be named as di	ibromidobis (ethylene dia	amine) chromium (III) bromide?
	(1) [Cr (en) ₃]Br ₃	(2) [Cr(en) ₂ Br ₂]Br	(3) [Cr(en)Br ₄] [_]	[AIEEE 2012, 4/144] (4) [Cr(en)Br ₂]Br
29.	., , _	complex species is not e	xpected to exhibit optical	l isomerism ?
	(1) [Co(en) ₃] ³⁺	(2) [Co(en) ₂ Cl ₂] ⁺	(3) [Co(NH ₃) ₃ Cl ₃]	[JEE(Main) 2013, 4/120] (4) [Co(en) (NH ₃)₂ Cl ₂] ⁺
30.	wavelengths in the reg strength of the four liga	gion of red, green, yellow	v and blue, respectively	gands L ₁ , L ₂ , L ₃ and L ₄ absorb . The increasing order of ligand [JEE(Main) 2014, 4/120] (4) L ₁ < L ₂ < L ₄ < L ₃
31.	The number of geom (py = pyridine): (1) 2	(2) 3	exist for square plana	ar [Pt(Cl)(py)(NH ₃)(NH ₂ OH)]⁺ is [JEE(Main) 2015, 4/120] (4) 6
32.		me magnetic moment is :		
	(1) [Cr(H₂O) ₆] ²⁺ and [Fe (3) [CoCl₄] ^{2−} and [Fe(H	e(H ₂ O) ₆] ²⁺	(2) [Mn(H ₂ O) ₆] ²⁺ and [C (4) [Cr(H ₂ O) ₆] ²⁺ and [C	[JEE(Main) 2016, 4/120] Cr(H ₂ O) ₆] ²⁺
33.	Which one of the follow (1) <i>cis</i> [Co(en) ₂ Cl ₂]Cl (en = ethylenediamine)	ving complexes shows op (2) <i>trans</i> [Co(en) ₂ Cl ₂]Cl)		[JEE(Main) 2016, 4/120] (4) [Co(NH ₃) ₃ Cl ₃]
34.	On treatment of 100 precipitated. The comp (1) [Co(H ₂ O) ₃ Cl ₃].3H ₂ C (3) [Co(H ₂ O) ₅ Cl]Cl ₂ .H ₂ d	olex is :)	CoCl ₃ .6H ₂ O with exce (2) [Co(H ₂ O) ₆]Cl ₃ (4) [Co(H ₂ O) ₄ Cl ₂]Cl.2H	ss AgNO ₃ ; 1.2 × 10 ²² ions are [JEE(Main) 2017, 4/120] ₂ O
35.	[Co(NH ₃) ₄ Br ₂] ⁺ + Br ⁻ \rightarrow (I) Two isomers are pro (II) Two isomers are pr (III) Only one isomer is	oduced if the reactant cor oduced if the reactant co produced if the reactant produced if the reactant produced if the reactant	mplex ion is a cis-isomer mplex ion is a <i>tran</i> -isomer complex ion is a <i>trans</i> -is	er. somer.
36.	The oxidation states of	^E Cr in [Cr(H ₂ O) ₆]Cl ₃ , [Cr(C ₆ H ₆) ₂], and K ₂ [Cr(CN) ₂ ($O_2(O_2)(NH_3)$] respectively are :
	(1) +3, 0, and +6	(2) +3, 0, and +4	(3) +3, +4, and +6	[JEE(Main) 2018, 4/120] (4) +3, +2, and +4

Coor	rdination Compounds			
		JEE(MAIN) ONL	INE PROBLEMS	
1.	is:	-		blved in the formation of the comlex a) 2014 Online (09-04-14), 4/120]
2.	(1) sp ³ d ² (2)The correct statement about	dsp² t of the magnetic pr	perties of [Fe(CN) ₆] ³⁻	
	 (1) both are paramagnetic (2) both are diamagnetic (3) [Fe(CN)₆]³⁻ is diamagnet (4) [Fe(CN)₆]³⁻ is paramagnet 		nagnetic.	2014 Online (09-04-14), 4/120]
3.	Which of the following name	e formula combinati		n <u>) 2014 On</u> line (11-04-14), 4/120]
	Formula	Name		
	(1) $K_2[Pt(CN)_4]$		yanoplatinate (II)	
	(2) [Mn(CN) ₅] ^{2–} (3) K[Cr(NH ₃) ₂ Cl ₄]	Pentacyanoma	gnate (II) ion imine tetrachlorochror	mate (III)
	$(4) [Co(NH_3)_4(H_2O)I]SO_4$		quaiodo cobalt (III) sul	
4.	Consider the coordination c	ompound, [Co(NH ₃)6]Cl ₃ . In the formatior	n of the complex, the species which
	acts as the Lewis acid is :		[JEE(Mair	n) 2014 Online (11-04-14), 4/120]
	(1) $[Co(NH_3)_6]^{3+}$ (2)	Cŀ	(3) Co ³⁺	(4) NH ₃
5.	Among the following specie		[JEE(Mair	n) 2014 Online (12-04-14), 4/120]
	(1) CN ⁻ (2)	NH ₃	(3) F [_]	(4) CO
6.	Which one of the following of (At nos. Sc = 21, Ti = 22, V (1) $[Sc(H_2O)_6]^{3+}$ (2)			ight ? n) 2014 Online (12-04-14), 4/120] (4) [Zn(NH ₃) ₆] ²⁺
7.	An octahedral complex with molecular composition M.5NH ₃ .Cl.SO ₄ has two isomers, A and B. T solution of A gives a white precipitate with AgNO ₃ solution and the solution of B gives white precipit with BaCl ₂ solution. The type of isomerism exhibited by the complex is:			solution of B gives white precipitate
	(1) Linkage isomerism(3) Coordinate isomerism		(2) Ionisation isome (4) Geometrical isor	rism
8.	The hybridisation involved respectively:		of unpaired electro	orm a diamagnetic complex [NiL₄] ^{2–} ons present in the complex are n) 2014 Online (19-04-14), 4/120] (4) sp ³ , zero
9.	The correct statement on th	e isomerism associ		
	[JEE(Main) 2015 Online (10-04-15), 4/120 (a) [Ni(H ₂ O) ₅ NH ₃] ²⁺ , (b) [Ni(H ₂ O) ₄ (NH ₃) ₂] ²⁺ and (c) [Ni(H ₂ O) ₃ (NH ₃) ₃] ²⁺ is : (1) (a) and (b) show only geometrical isomerism. (2) (b) and (c) show geometrical and optical isomerism (3) (b) and (c) show only geometrical isomerism (4) (a) and (b) show geometrical and optical isomerism			
10.	Which molecule/ion among	the following canno		
	(1) CH ₄ (2)	CN-	[JEE(Mair (3) Br−	n) 2015 Online (10-04-15), 4/120] (4) CO
11.	orbitals?	nplex ions has ele [Mn(CN)6] ^{4_}		metrically filled in both t _{2g} and e _d a) 2015 Online (11-04-15), 4/120] (4) [Co(NH ₃) ₆] ²⁺

Coo	ordination Compounds	
12.	Identify the correct trend given below: (Atomic	No.: Ti = 22, Cr = 24 and Mo = 42) [JEE(Main) 2016 Online (09-04-16), 4/120]
	(1) Δ_0 of [Cr(H ₂ O) ₆] ²⁺ < [Mo(H ₂ O) ₆] ²⁺ and Δ_0 of	
	(2) Δ_0 of [Cr(H ₂ O) ₆] ²⁺ > [Mo(H ₂ O) ₆] ²⁺ and Δ_0 of	
	(3) Δ_0 of [Cr(H ₂ O) ₆] ²⁺ > [Mo(H ₂ O) ₆] ²⁺ and Δ_0 of	
	(4) Δ_0 of [Cr(H ₂ O) ₆] ²⁺ < [Mo(H ₂ O) ₆] ²⁺ and Δ_0 of	$[Ti(H_2O)_6]^{3+} > [Ti(H_2O)_6]^{2+}$
13.	Which one of the following complexes will cons	sume more equivalents of aqueous solution of Ag(NO ₃) ? [JEE(Main) 2016 Online (09-04-16), 4/120]
	(1) Na ₃ [CrCl ₆] (2) [Cr(H ₂ O) ₅ Cl]Cl ₂	(3) [Cr(H ₂ O) ₆]Cl ₃ (4) Na ₂ [CrCl ₅ (H ₂ O)]
14.	Which of the following is an example of homole	eptic complex ?
	(1) [Co(NH ₃) ₄ Cl ₂] (2) [Co(NH ₃) ₆]Cl ₃	[JEE(Main) 2016 Online (09-04-16), 4/120] (3) [Co(NH ₃) ₅ Cl]Cl ₂ (4) [Pt(NH ₃) ₂ Cl ₂]
15.	sp ³ d ² hybridization is not displayed by : (1) PF_5 (2) SF_6	[JEE(Main) 2017 Online (08-04-17), 4/120] (3) [CrF ₆] ^{3–} (4) BrF₅
16.	[Co ₂ (CO) ₈] displays : (1) one Co–Co bond, four terminal CO and fou (2) one Co–Co bond, six terminal CO and two (3) no Co–Co bond, four terminal CO and four (4) no Co–Co bond, six terminal CO and two b	[JEE(Main) 2017 Online (09-04-17), 4/120] Ir bridging CO bridging CO bridging CO
17.	The correct combination is : (1) [NiCl ₄] ²⁻ – square-planar; [Ni(CN) ₄] ²⁻ – para (2) [Ni(CN) ₄] ²⁻ – tetrahedral; [Ni(CO) ₄] – param (3) [NiCl ₄] ²⁻ – paramagnetic; [Ni(CO) ₄] – tetrah (4) [NiCl ₄] ²⁻ – diamagnetic; [Ni(CO) ₄] – square-	nagnetic edral
18.	(1) $[ZnCl_4]^{2-} > [NiCl_4]^{2-} > [CoCl_4]^{2-} > [MnCl_4]^{2-}$	nts among the following is : (n = 30) [JEE(Main) 2018 Online (15-04-18), 4/120] (2) [CoCl ₄] ²⁻ > [MnCl ₄] ²⁻ > [NiCl ₄] ²⁻ > [ZnCl ₄] ²⁻ (4) [MnCl ₄] ²⁻ > [CoCl ₄] ²⁻ > [NiCl ₄] ²⁻ > [ZnCl ₄] ²⁻
19.	The total number of possible isomers for squar	re-planar [Pt(Cl)(NO₂)(NO₃) (SCN)]²- is :
		[JEE(Main) 2018 Online (15-04-18), 4/120]
	(1) 8 (2) 12	(3) 16 (4) 24
20.		nd point is estimated spectrophotometrically (through light ht and only 'L' absorbs, then the titration plot between
	$(1)^{A} \xrightarrow{(2)^{A}} (2)^{A} \xrightarrow{V (Ligand) \rightarrow} V (Ligand) \rightarrow$	$(3)^{A} (4)^{A} (4)^{A}$
21.	In Wilkinson's catalyst, the hybridization of cen	
21.	(1) sp ³ d, trigonal bipyramidal (3) dsp ² square planar	 (2) d²sp³, octahedral (4) sp³ tetrahedral

(3) dsp², square planar

(4) sp^3 , tetrahedral

22.	Which of the following	complexes will st	now geo	metrical		18 Online (16-04-18), 4/120]
	(2) Pentaaquachloroch(3) Aquachlorobis(ethy) Potassium tris(oxalato)chromate(III)) Pentaaquachlorochromium(III)chloride) Aquachlorobis(ethylenediamine)cobalt(II) chloride) Potassium amminetrichloroplatinate(II)				
23.	Two complexes [Cr(H ₂ incorrect statement reg (1) Δ_0 value for (A) is le	arding them is :		₃]Cl₃ (B)		ellow coloured respectively. The 19 Online (09-01-19), 4/120]
	(2) both absorb energie	es corresponding	to their	•	•	
	(3) Δ_0 values of (A) and (4) both are paramagnet					d yellow light, respectively.
24.	absorb wavelengths in strength is :	the region of gre	en, blue	and rec	l respectively. Th [JEE(Main) 20	nodentate ligands L ₁ , L ₂ and L ₃ ne increasing order of the ligand 19 Online (09-01-19), 4/120]
	(1) L ₁ < L ₂ < L ₃	(2) L ₃ < L ₂ < L ₁		(3) L ₂ <	< L1 < L3	(4) $L_3 < L_1 < L_2$
25.	The complex that has h	nighest crystal fie	eld splitti	ng energ		19 Online (09-01-19), 4/120]
	(1) K ₂ [COCl ₄]	(2) [CO(NH ₃) ₅ (I	H₂O)]Cl₃	(3) [CC		(4) K ₃ [CO(CN) ₆]
26.	Wilkinson catalyst is : (1) [(Et_3P) ₃ IrCl] ($Et = C$ (3) [(Ph_3P) ₃ RhCl]	₂ H ₅)		• • • •	[JEE(Main) 20 33P)₃RhCl] n₃P)₃IrCl]	19 Online (10-01-19), 4/120]
27.	The total number of isc	mers for a squar	e planar	[.] comple		
	(1) 16	(2) 4		(3) 12	[JEE(Main) 20	19 Online (10-01-19), 4/120] (4) 8
28.	The difference in the octahedral complexes i (1) Co ²⁺			lectrons (3) Mn ²	[JEE(Main) 20	in its high-spin and low-spin 19 Online (10-01-19), 4/120] (4) Ni ²⁺
29.	products A (violet colo	oured) and B(grossomers does A a	een colo	oured). / present? (2) Linl	A can show opt	e ratio generates two isomeric tical activity, but, B is optically 19 Online (10-01-19), 4/120] s
30.	Match the metals (colu	mn I) with the co	ordinatic	on comp		
	(column I)		(colum	nn II)	[JEE(Main) 20	19 Online (11-01-19), 4/120]
	Metal		Coord	ination	compound(s)/e	nzyme(s)
	(A) Co (B) Zn		(i) (ii)	Chloro	son catalyst phyll	
	(C) Rh (D) Mg		(iii) (iv)	Vitamir	n B ₁₂ nic anhydrase	
	(1) (A)-(i);(B)-(ii);(C)-(iii) (3) (A)-(iii);(B)-(iv);(C)-((10)	(2) (A)·	-(iv);(B)-(iii);(C)-(-(ii);(B)-(i);(C)-(iv	
31.		er of Th in K₄[Th	(C ₂ O ₄) ₄ (OH ₂) ₂] is	s: [JEE(Main) 2 (019 Online (11-01-19), 4/120]
	$(C_2 O_4^{2-} = Oxalato)$ (1) 14	(2) 10		(3) 6		(4) 8
32.	The number of bridging		nd Co-Co	. ,		espetively are :
	(1) 4 and 0	(2) 0 and 2		(3) 2 a		19 Online (11-01-19), 4/120] (4) 2 and 0

Coo	rdination Compounds	/		
33.	The metal d-orbitals the	nat are directly facing th	e ligands in K₃[Co(CN)6] [JEE(Main) [are : 2019 Online (12-01-19), 4/120]
	(1) d _{xz} , d _{yz} and d _{z²}	(2) d_{xy} and $d_{x^2-y^2}$,	(4) $d_{x^2-y^2}$ and d_{z^2}
34.	$Mn_2(CO)_{10}$ is an orgar	nometallic compound du	•	2019 Online (12-01-19), 4/120]
	(1) Mn – C bond	(2) C – O bond	(3) Mn – O bond	(4) Mn – Mn bond
35.	The magnetic momer this complex is: (1) Ethylenediamine	nt of an octahedral hon (2) CN ⁻		s 5.9 BM. The suitable ligand for 2019 Online (12-01-19), 4/120] (4) CO

Answers

EXERCISE - 1

PART - I

CuSO4.4NH₃ [Cu(NH₃)₄] SO₄ $\stackrel{\text{aq.}}{=}$ [Cu(NH₃)₄]²⁺ (aq) + SO₄²⁻ (aq)

As copper (II) is present in coordination sphere it will not give the test of Cu²⁺ ion.

A-2. 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.

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.

	Complex	Coordination Number	Oxidation State
(a)	[AgCl₂]⁻	2	1
(b)	[Cr(H ₂ O) ₅ Cl] ²⁺	6	3
(C)	[Co(NCS)4] ²⁻	4	2
(d)	[Co(NH ₃) ₃ (NO ₂) ₃]	6	3
(e)	[Fe(EDTA)] [_]	6	3
(f)	[Cu(en) ₂]SO ₄	4	2
(g)	K[Pt(NH ₃)Cl ₅]	6	4

A-3. (A) methyl isocyanide, monodentate.

(B) acetylacetonato, bidentate

(C) azido, monodentate

(D) diethylenetriamine, tridentate

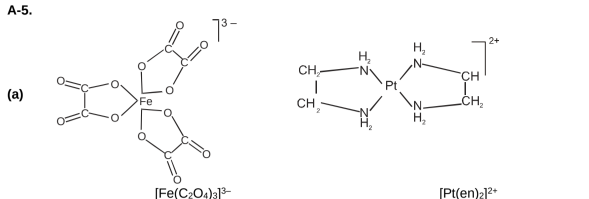
(F) ethylenediamine triacentato, pentadentate

- (E) ethylenediamine tetraacetato, hexadentate
- (G) oxalato, bidentate (I) isocyanido, monodentate
- (K) oxido, monodentate

(H) dimethylglyoximato (J) nitrito, monodentate

(L) superoxido, monodentate

A-4. (A) $M \leftarrow N \leqslant_{O}^{O}$ nitrito-N $M \leftarrow O - N = O$ nitrito-O (B) $M \leftarrow SCN$ thiocyanato or thiocyanato-S, $M \leftarrow NCS$ isothiocyanato or thiocyanato-N (C) I or I dithioxalate (D) $M \leftarrow OCN$ cyanato-O or cyanato-N, $M \leftarrow NCO$ isothiocyanato or thiocyanato-N (E) $M \leftarrow NOS$ thionitrito-N or , $M \leftarrow SON$ thionitrito-S



The ligands, oxalate and ethylenediamine are bidentate as each ligand has two 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.

(b)

 $\begin{array}{l} \mathsf{K}_{2}[\mathsf{Ni}(\mathsf{CN})_{4}] \stackrel{\text{aq.}}{\longrightarrow} 2\mathsf{K}^{+}(\mathsf{aq}) + [\mathsf{Ni}(\mathsf{CN})_{4}]^{2-}(\mathsf{aq}) \\ [\mathsf{Cr}(\mathsf{en})_{3}]\mathsf{Cl}_{3} \stackrel{\text{aq.}}{\longrightarrow} [\mathsf{Cr}(\mathsf{en})_{3}]^{3+}(\mathsf{aq}) + 3\mathsf{Cl}^{-}(\mathsf{aq}) \\ \mathsf{Fe}_{4}[\mathsf{Fe}(\mathsf{CN})_{6}]_{3} \stackrel{\text{aq.}}{\longrightarrow} 4\mathsf{Fe}^{3+}(\mathsf{aq}) + 3[\mathsf{Fe}(\mathsf{CN})_{6}]^{4-}(\mathsf{aq}) \end{array}$

 $[PtCl_2(en)_2]$ (NO₃)₂ $\stackrel{aq.}{=}$ $[PtCl_2(en)_2]^{2+}$ (aq) + 2(NO₃)⁻(aq)

So, [Ni(CN)₄]²⁻, [Cr(en)₃]³⁺, 3[Fe(CN)₆]⁴⁻ and [PtCl₂(en)₂]²⁺ are coordination entities and K⁺, C⁺, Fe^{3+} and NO_{3}^{-} are counter ions.

Coordination compounds are acid-base adduct. Cations are electron deficient, therefore, are (c) called Lewis acids where as ligands are electrons donors, therefore, are called as Lewis base.

		LEWIS ACID	LEWIS BASE
(i)	[HgBr ₄] ²⁻	Hg ²⁺	4Br−
(ii)	[Ni(H ₂ O) ₆] ²⁺	Ni ²⁺	6H ₂ O
(iii)	[PdCl ₂ (NH ₃) ₂]	Pd ²⁺	2CI- & 2NH3
(iv)	[AI(OH)4] ⁻	Al ³⁺	4OH-
(v)	[Ag(CN) ₂] ⁻	Ag⁺	2CN⁻
(vi)	[Cr(CO) ₆]	Cr ⁰	6CO

B-1.

(a)	[Co(NH ₃) ₆]Cl ₃	Hexaamminecobalt(III) chloride
(b)	[Rh(NH₃)₅I]I₂	Pentaammineiodidorhodium(III) iodide
(c)	[Fe(CO)₅]	Pentacarbonyliron(0)
(d)	[Fe(C ₂ O ₄) ₃] ³⁻	Trioxalatoferrate(III) ion OR Tris(oxalato)ferrate(III) ion
(e)	[Cu(NH ₃) ₄]SO ₄	Tetraamminecopper(II) sulphate
(f)	Na[Cr(OH)₄]	Sodium tetrahydroxidochromate(III)
(g)	[Co(gly)₃]	Triglycinatocobalt(III) OR Tris(glycinato)cobalt(III)
(h)	[Fe(H₂O)₅(SCN)] ²⁺	Pentaaquathiocyanato-S-iron(III) ion
(i)	K ₂ [Hgl ₄]	Potassium tetraiodidomercurate(II)
(j)	Co[Hg(SCN)₄]	Cobalt(II) tetrathiocyanato-S-mercurate(II)
(k)	Fe ₄ [Fe(CN) ₆] ₃	Iron(III) hexacyanidoferrate(II)
(I)	K ₃ [Co(NO ₂) ₆]	Potassium hexanitrito–N–cobaltate(III)
(m)	[Ni(dmg)₂]	Bis(dimethylglyoximato)nickel(II)
(n)	K ₂ [PtCl ₆]	Potassium hexachloridoplatinate(IV)
(o)	Na₂[Fe(CN)₅NO ⁺]	Sodium pentacyanidonitrosoniumferrate(II)
(p)	[Fe(H ₂ O) ₅ (NO ⁺)]SO ₄	Pentaaquanitrosoniumiron(I) sulphate
(q)	[Cu(CN) ₄] ^{3–}	Tetracyanidocuperate(I) ion
(r)	(NH ₄) ₂ [PtCl ₆]	Ammonium hexachloridoplatinate(IV)

B-2.

-		
(a)	[CoBr(en) ₂ (ONO)] ⁺¹	Bromidobis(ethylenediamine)nitrito-O-cobalt(III)
(b)	[Co(NH ₃) ₆][Co(ONO) ₆]	Hexaamminecobalt(III) hexanitrito-O-cobaltate(III)
(C)	[Co(NH ₃) ₅ (CO ₃)]Cl	Pentaamminecarbonatocobalt(III) chloride
(d)	[Pt(NH ₃) ₄ Cl ₂][PtCl ₄]	Tetraamminedichloridoplatinum(IV)
(u)		tetrachloridoplatinate(II)
(e)	[Co(en)3]2(SO4)3	Tris(ethylenediamine)cobalt(III) sulphate or Tris(ethane-1,
(6)		2-diamine)cobalt(III) sulphate.
(f)	[(NH3)5C0-NH2-C0(NH3)4(H2O)]Cl5	Pentaamminecobalt(III)-µ-amidotetraammineaquacobalt
(1)		(III) chloride
(g)	[Cr(CO) ₅ (PPh ₃)]	Pentacarbonyltriphenylphosphinechromium(0)
(h)	[(CO)₅Mn-Mn(CO)₅]	Decacarbonyldimanganese(0)
(i)	Cr(π-C ₆ H ₆) ₂	Bis(η ⁶ -benzene)chromium(0)
(j)	[Co(NH ₃) ₄ (OH ₂) ₂][BF ₄] ₃	Tetraamminediaquacobalt(III) tetrafluoridoborate(III)
(I)	Ba[Zr(OH) ₂ (ONO) ₂ (ox)]	Barium dihydroxidodinitrito-O-oxalatozirconate(IV)
(I)	$[Co(NH_3)_6][Co(C_2O_4)_3]$	Hexaamminecobalt(III) trioxalatocobaltate(III)

B-3.

(a)	Tetraamminezinc(II) Nitrate	[Zn(NH ₃) ₄](NO ₃) ₂
(b)	Tetracarbonylnickel(0)	[Ni(CO)4]
(C)	Potassium amminetrichloridoplatinate(II)	K[Pt(NH ₃)Cl ₃]
(d)	Dicyanidoaurrate(I) ion	[Au(CN)₂] [−]
(e)	Sodium hexafluoridoaluminate(III)	Na ₃ [AIF ₆]
(f)	Diamminesilver(I) ion	[Ag(NH ₃) ₂]+

B-4.

C-1.

(a)	Diamminetriaquahydroxido	te	$[Cr(NH_3)_2(H_2O)_3(OH)](NO_3)_2$	
(b)	Tetrakis(pyridine)platinum(I	[Pt(Py) ₄][B(ph) ₄] ₂		
(C)	Dibromidotetracarbonyliron	[Fe(Br) ₂ (CO) ₄]		
(d)	Ammonium diamminetetrak	(NH ₄)[Cr(NH ₃) ₂ (NCS) ₄]		
(e)	Pentaamminedinitrogenruth	[Ru(NH ₃) ₅ N ₂]Cl ₂		
(f)	Barium dihydroxidodinitrito-	e(IV)	Ba[Zr(OH) ₂ (ONO) ₂ (ox)]	
(g)	Tetrapyridineplatinum(II) te	trachloridoplatinate	e(II)	[Pt(py) ₄][PtCl ₄]
(a) – i	v, (b) – viii,	(c) — i,	(d) – vii,	
(e) – i	ii, $(f) - v$,	(g) — ii,	(h) – vi	
0.007	5.			

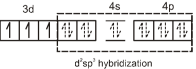
C-2.

C-3.	(a) (e)	36 36	(b) (f)		(c) (g)	36 54	(d) (h)	36 86
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C-4. ii < i < iv < iii.

(i) $[Cr(NH_3)_4Cl Br]Cl \rightleftharpoons [Cr(NH_3)_4Cl Br]^+ + Cl^-; Ag^+ + Cl^- \longrightarrow AgCl \downarrow (white) ; soluble in dilute NH_3.$ $[Cr(NH_3)_4Cl_2]Br \rightleftharpoons [Cr(NH_3)_4Cl_2]^+ + Br^-; Ag^+ + Br^- \longrightarrow AgBr \downarrow (yellow) ; soluble in conc. NH_3.$ D-1. So, $A = [Cr(NH_3)_4Cl Br]Cl$ and $B = [Cr(NH_3)_4Cl_2]Br$.

(ii) In both complexes chromium is in +3 oxidation state. Chromium with 3d³ configuration has 3 unpaired electrons with weak field as well as strong field ligand. So, the hybridisation scheme is as follow :



(iii) $\mu = \sqrt{n (n+2)} = \sqrt{15}$

(iv) EAN = 24 - 3 + 12 = 33

(v) Yes, both have two ions per formula unit.

(vi) AgCl + 2NH₃ = [Ag(NH₃)₂]Cl ; AgBr + 2NH₃ = [Ag(NH₃)₂]Br

D-2.

	Complex	Hybridization	Geometry
(a)	[NiBr ₄] ^{2–}	sp ³	Tetrahedral
(b)	[AuCl₄]⁻	dsp ²	square planar
(C)	[Pt(NH ₃) ₄] ²⁺	dsp ²	square planar

E-1. (i) potasium amminetetracyanidonitrosoniumchromate(I)

(ii) Octahedral

(iii) One unpaired electron

(v) EAN = $24 - 1 + 2 \times 6 = 35$

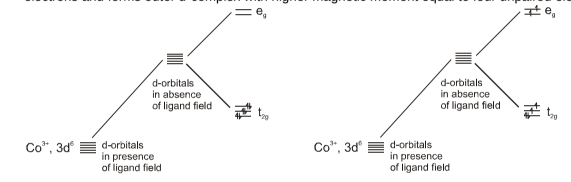
(iv) It is paramagnetic with one unpaired electron (vi) d^2sp^3

E-2.

	Complex	Hybridization	Geometry
(a)	[Fe(CN) ₆] ^{3–}	d ² sp ³	octahedral
(b)	[MnBr ₄] ^{2–}	sp ³	Tetrahedral
(C)	[Fe(H ₂ O) ₆] ²⁺	sp ³ d ²	Octahedral
(d)	[Co(SCN) ₄] ²⁻	sp ³	Tetrahedral

(b) X < O < N < C

E-3. Since ammonia is a strong field ligand so can pair up the electrons of Co(III), so will form an inner dorbital complex having zero magnetic moment while fluoride being a weak field ligand can not pair up electrons and forms outer d-complex with higher magnetic moment equal to four unpaired electrons.

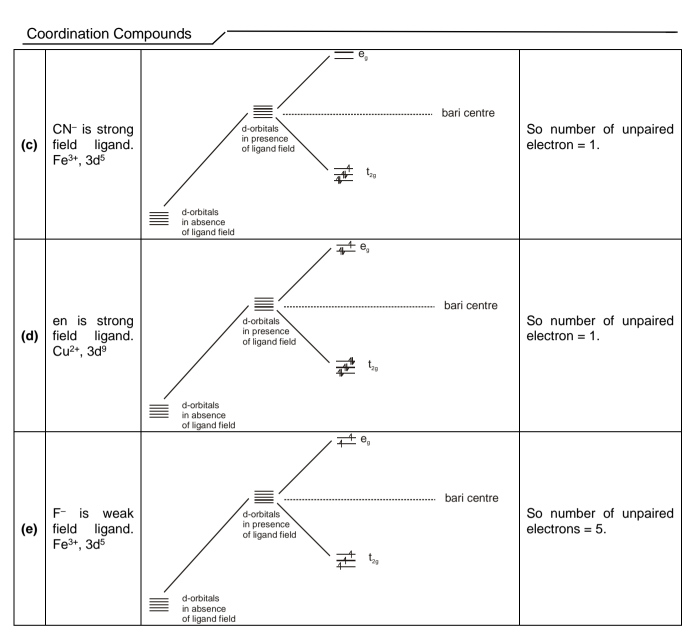


E-4. (a) i < iv < ii < iii

(c) $Br^- < S^{2-} < NO_3^- < H_2O < NH_3 < NO_2^- < CN^- < CO$

E-5.

J			
(a)	F⁻ is weak field ligand. Cr³+, 3d³	e _g d-orbitals in presence of ligand field d-orbitals in absence of ligand field	So number of unpaired electrons = 3
(b)	H ₂ O is weak field ligand. V ³⁺ , 3d ²	e _g d-orbitals in presence of ligand field d-orbitals in absence of ligand field	So number of unpaired electrons = 2

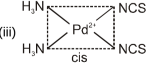


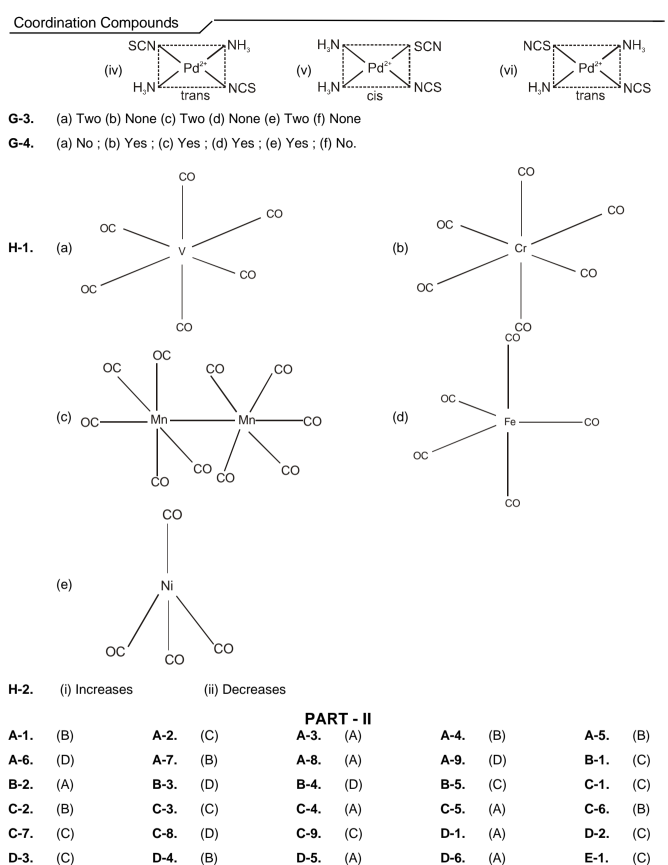
F-1. As 3d⁶ configuration has higher CFSE as compared to 3d⁷ so it gets oxidised in presence of complexing reagent to easily have d²sp³ hybridisation.

F-2. Yellow colour

G-1. (i) Linkage (ii) Coordination (iii) Ionisation (iv) Hydrate

G-2. (a) There are three constitutional isomers (i) [Ru(NH₃)₅(NO₂)]Cl (ii) [Ru(NH₃)₅Cl](NO₂) or [Ru(NH₃)₅Cl]ONO (iii) [Ru(NH₃)₅ ONO]Cl (i) & (ii) are ionisation isomers (i) & (iii) are linkage isomers >NH₃ H₃N_NSCN NCS H₃N (b) (iii) (i) (ii) H₃N⊻--SCN trans





E-2.

E-7.

E-12.

G-3.

(A)

(D)

(C)

(B)

E-3.

E-8.

F-1.

G-4.

(A)

(C)

(B)

(D)

E-4.

E-9.

F-2.

G-5.

(B)

(A)

(B)

(C)

G-6.

(C)



H-1. (D)

H-2. (D)

	PART - III										
1.	(A - p,r,t); (B -	p,s,t); (0	C - p,r,t); (D - q,r)		2. (A - p	o,q,r); (B	- q,r,s) ; (C - p,q,	r,s) ; (D	- p,q)		
EXERCISE - 2											
PART - I											
1.	(B)	2.	(C)	3.	(A)	4.	(B)	5.	(D)		
6.	(D)	7.	(D)	8.	(C)	9.	(D)	10.	(B)		
11.	(B)	12.	(A)	13.	(A)						
	PART - II										
1.	13	2.	05	3.	26	4.	(40 + 20) ml = 60 ml				
5.	4 (i, iii, iv, vii)	6.	3 (a, c, d)	7.	2 (i & v)	8.	4				
9.	4 (0 + 0 + 2 +	0 + 2 = 4	4)	10.	4	11.	12				
				PAI	RT - III						
1.	(BCD)	2.	(BC)	3.	(ABCD)	4.	(B)	5.	(BD)		
6.	(ABCD)	7.	(BD)	8.	(AB)	9.	(ACD)	10.	(BCD)		
11.	(BCD)	12.	(C)								
				PAF	RT - IV						
1.	(D)	2.	(C)	3.	(B)	4.	(A)	5.	(A)		
6.	(B)	7.	(A)	8.	(C)	9.	(A)	10.	(B)		
11.	(C)										

EXERCISE - 3

PART - I

1. IUPAC name is :

Potassium amminetatracyanidonitrosoniumchromate(I) or Potassium amminetatracyanidonitrocyliumchromate(I).

Let n is the number of unpaired electron in the chromium ion.

Since
$$\mu = \sqrt{n(n+2)}$$
 or $1.73 = \sqrt{n(n+2)}$ B.M. or

Hence n = 1.

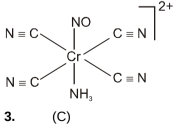
,

 $1.73 \times 1.73 = n^2 + 2n$.

As the CN⁻ and NH₃ are strong fields ligands, they compel for pairing of electrons. So, 3d 4s 4p

		U 4			
		·····	•••••	••••••	·····
$[Cr(NO)(CN)_4(NH_3)]^{2-} = [$	11 11	1 1 1			

Hence, the oxidation state of chromium is +1 (having $3d^5$ configuration). So according to charge on the complex NO should be NO⁺ and the structure of this complex is octahedral with d^2sp^3 hybridisation as given below



2.

(D)

4.

26.

31.

(3)

(2)

27.

32.

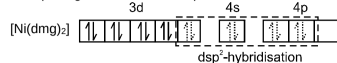
(3)

(3)

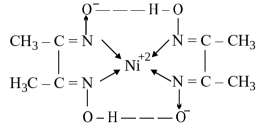
Ni²⁺ + 2dmg $\xrightarrow{NH_4OH}$ [Ni(dmg)₂] \downarrow (bright red).

It acquires stability through chelation and intra molecular H-bonding.

In [Ni(dmg)₂] the nickel is in +2 oxidaiton state and to have square planar geometry because of chelation the pairing of electrons takes place. So



As all electrons are paired, so complex is diamagnetic. Nickel with coordination number four will have the structure as given below.



rosy red ppt

5.	(A)	6.	(A)	7.	(A)	8.	(C)	9.	(C)
10.	(B)	11.	(A - p,q,s); (B -	- p,r,s); (C - q,s); (D - q,s)	12.	(C)	13.	(B)
14.	(B)	15.	(A)	16.*	(CD)	17.	(A)	18.	(C)
19.	(B)	20.	(B)	21.	3	22.	(B)	23.	(C)
24.	6	25.	(D)	26.	(C)	27.	(B)	28.*	(BD)
29.	8	30.	4	31.	(B)	32.	4	33.	3
34.	6	35.	(B)	36.	5	37.	(A)	38.	(ACD)
39.*	(BC)	40.	1	41.	2992	42.*	(ABD)	43.	(C)

PART - II JEE(MAIN) OFFLINE PROBLEMS

			- (-	_			
1.	(3)	2.	(4)	3.	(2)	4.	(4)	5.	(1)
6.	(4)	7.	(1)	8.	(4)	9.	(4)	10.	(2)
11.	(3)	12.	(1)	13.	(1)	14.	(2)	15.	(4)
16.	(2)	17.	(3)	18.	(1)	19.	(2)	20.	(3)
21.	(4)	22.	(3)	23.	(1)	24.	(1)	25.	(2)
26.	(3)	27.	(3)	28.	(2)	29.	(3)	30.	(2)
31.	(2)	32.	(1)	33.	(1)	34.	(4)	35.	(4)
36.	(1)								
			JEE	(MAIN) ON	LINE PRO	BLEMS			
1.	(3)	2.	(1)	3.	(2)	4.	(3)	5.	(4)
6.	(3)	7.	(2)	8.	(2)	9.	(3)	10.	(1)
11.	(1)	12.	(4)	13.	(3)	14.	(2)	15.	(1)
16.	(2)	17.	(3)	18.	(4)	19.	(2)	20.	(1)
21.	(3)	22.	(3)	23.	(3)	24.	(4)	25.	(4)

28.

33.

(1)

(4)

29.

34.

(4)

(1)

30.

35.

(3)

(3)