## **COORDINATION COMPOUNDS**

## CHEMISTRY

|             | Single Correct  | Answer Type                    |                         |
|-------------|---|--------------------------------|-------------------------|
| 1.          | The IUPAC name of $Na_3[Co(ONO)_6]$ is:   |                                |                         |
| 1.          | a) Sodium cobaltinitrite  |                                |                         |
|             | b) Sodium hexanitritocobaltate(III)   |                                |                         |
|             | c) Sodium hexanitrocobalt(III)  |                                | $\sim$                  |
|             | d) Sodium hexanitritocobaltate(II)  |                                |                         |
| 2.          | $CuSO_4$ decolourises on addition of KCN, the product   | t is:                          |                         |
|             | a) $Cu(CN)_4^{2-}$ b) $[Cu(CN)_4]^{3-}$   | c) Cu(CN) <sub>2</sub>         | d) CuCN                 |
| 3.          | Exchange of coordination group by a water molecul   |                                | llts in:                |
|             | a) Ionization isomerism   |                                | $\circ$                 |
|             | b) Ligand isomerism   | <b>^</b>                       | $\mathbf{X}$            |
|             | c) Hydration isomerism  |                                |                         |
|             | d) Geometrical isomerism  |                                |                         |
| 4.          | The type of isomerism found in urea molecule is   |                                |                         |
|             | a) Chain  |                                |                         |
|             | b) Position   |                                |                         |
|             | c) Tautomerism  |                                |                         |
|             | d) None of these  |                                |                         |
| 5.          | The IUPAC name of the compoundO is  | $\mathbf{\nabla}^{\mathbf{r}}$ |                         |
|             | a) Butane-2-aldehyde b) 2-methyl butanal  | c) 2-ethyl propanal            | d) None of the above    |
| 6.          | Anisol is a product obtained from phenol by the rea   | , , , , ,                      |                         |
| •           | a) Coupling b) Etherification   | c) Oxidation                   | d) Esterification       |
| 7.          | Which of the following is diamagnetic in nature?  | ,                              | ,<br>,                  |
|             | a) $[Fe(CN)_6]^{3-}$ b) $[NiCl_4]^{2-}$   | c) [Ni(CO) <sub>4</sub> ]      | d) $[MnCl_4]^{2-}$      |
| 8.          | Which is the strongest field ligand?  |                                |                         |
|             | a) $CN^-$ b) $NO_2^-$   | c) NH <sub>3</sub>             | d) en                   |
| 9.          | Nitrobenzene on reduction with Zn and $aq$ . NH <sub>4</sub> Cl g   | ives:                          |                         |
|             | a) Aniline  |                                |                         |
|             | b) Nitrosobenzene   |                                |                         |
|             | c) N-phenyl hydroxylamine   |                                |                         |
|             | d) Hydrazobenzene   |                                |                         |
| 10.         | The IUPAC name of $[Co(NH_3)_5 ONO]^{2+}$ ion is  |                                |                         |
|             | a) Pentaammine nitrito cobalt (IV) ion  | b) Pentaammine nitro co        |                         |
|             | c) Pentaammine nitrito cobalt (III) ion   | d) Pentaammine nitro co        | balt (IV) ion           |
| 11.         | The compound which does not show paramagnetism  |                                |                         |
| 12          | a) $NO_2$ b) NO   | c) $[Ag(NH_3)_2]Cl$            | d) $[Cu(NH_3)_4Cl_2]$   |
| 12.         | Which of the following is expected to undergo nitration corresponding nitro derivatives employing the usual                             |                                | y to furnish the        |
|             |   | c) $C_6H_5CH_3$                |                         |
| 12          | a) C <sub>6</sub> H <sub>6</sub> b) C <sub>6</sub> H <sub>5</sub> NO <sub>2</sub><br>The number of unpaired electrons calculated in [Co |                                | d) $C_6H_5 \cdot CCl_3$ |
| 13.         | a) 4 and 4 b) 0 and 2   | c) 2 and 4                     | d) 0 and 4              |
| 14          | The IUPAC name of   | cj 2 anu 1                     | aj o ana 1              |
| <b>I</b> 1. |   |                                |                         |
|             | HO-N COOH is  |                                |                         |

- a) 4-hydroxy amino benzene carboxylic acid
- b) 4-(N-hydroxy) imino benzene carboxylic acid
- c) 4-hydroxy imino cyclohexanoic acid
  - d) 4-(N-hydroxy) imino cyclohexane-1
     -carboxylic acid
- 15. The IUPAC name of the coordination compound  $K_2[Zn(OH)_4]$  is
  - a) Potassium tetrahydroxozine (II)
  - c) Potassium tetrahydroxozincate (II)
- 16. Arrange in order of decreasing trend towards  $S_E$  reactions,

Chlorobenzene, Benzene, Anilium chloride, Toluene: I. (II) (III) (IV)

a) 
$$II > I > III > IV$$
 b)  $III > I > IV > IV$ 

b) Dipotassium tetrahydroxo(II)

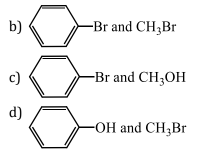
d) Potassium tetrahydroxozincate (III)

- d) I > II > III > IV
- 17. Toluene is nitrated and the resulting product is reduced with tin and hydrochloric acid. The product so obtained is diazotised and then heated with cuprous bromide. The reaction mixture so formed contains:a) Mixture of *o* and *m*-bromotoluenes
  - b) Mixture of *o* and *p*-bromotoluenes
  - c) Mixture of *o* and *p*-dibromobenzenes
  - d) Mixture of *o* and *p*-bromoanilines
- 18. A positive carbylamine test is given by:
  - a) N, N-dimethylaniline
  - b) 2,4-dimethylaniline
  - c) *N*-methyl-*o*-methylaniline
  - d) p-methyl benzylamine
- 19.  $CN^-$  is strong field ligand. This is due to the fact that
  - a) It carries negative charge
  - b) It is a pseudohalide
  - c) It can accept electrons from metal species
  - d) It forms high spin complexes with metal species.
- 20. Which of the following is not true for ligand metal complex?
  - a) Highly charged ligand forms strong bond
  - b) Greater the ionization potential of central metal, the stronger is the bond
  - c) Larger the permanent dipole moment of ligand, the more stable is the bond
  - d) Larger the ligand, the more stable is the metal-ligand bond
- 21. The nitration of nitrobenzene with fuming  $HNO_3$  will give:
- a) TNB b) 1,3-dinitrobenzene c) Picric acid d) 1,4-dinitrobenzene
- 22. A ligand can also be regarded asa) Lewis acidb) Bro
  - b) Bronsted base c) Lewis base
- d) Bronsted acid
- 23. The correct statement with respect to the complexes Ni(CO)<sub>4</sub> and [Ni (CN)<sub>4</sub>]<sup>2-</sup> is
  a) Nickel is in the same oxidation state in both
  - b) Both have terahedral geometry
  - c) Both have square planar geometry
  - d) Have tetrahedral and square planar geometry respectively
- 24. Which one of the following has lowest value of paramagnetic behaviour?
- a)  $[Cr(CN_6)_4]^{3-}$  b)  $[Mn(CN)_6]^{3-}$  c)  $[Fe(CN)_6]^{3-}$  d)  $[Co(CN)_6]^{3-}$
- 25. In the reaction;

$$\bigcirc$$
 OCH<sub>3</sub>—HBr

the products are:

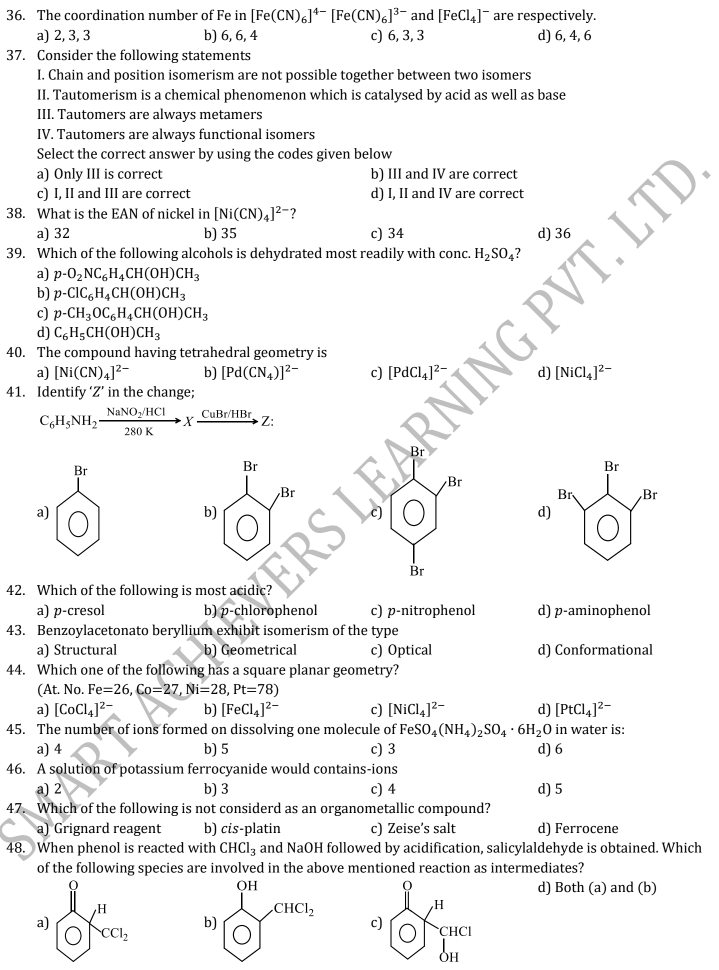
a) Br OCH<sub>3</sub> and  $H_2$ 



26. An octahedral complex is formed when central metal atom undergoes hybridization amongst the....orbitals.

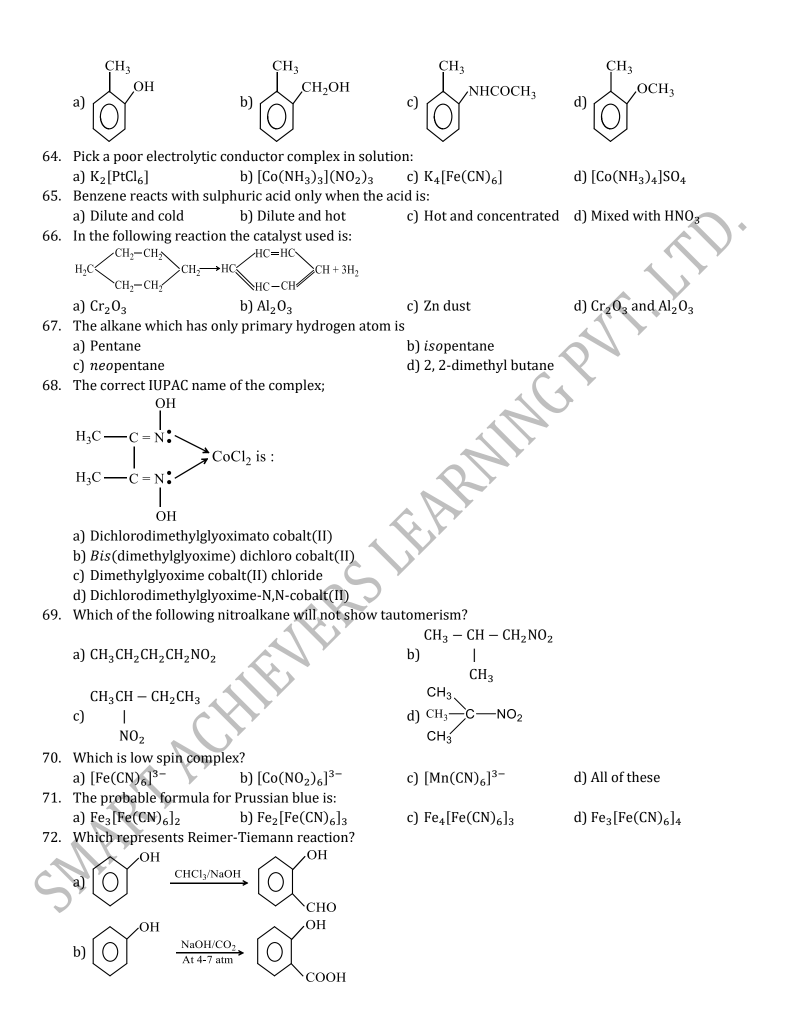
|     | theorbitals.   |  | -                             |   |
|-----|--|--|-------------------------------|---|
|     | a) <i>sp</i> <sup>3</sup>  | b) $dsp^2$   | c) $sp^3d$                    | d) $sp^3d^2$  |
| 27. | ONa  |  |                               |   |
|     | $\left[ \bigcirc \right]^{\text{OTVa}} + \text{CO}_2 \frac{-390 \text{ K}}{P}$ | $\rightarrow X \xrightarrow{\text{HCl}}$   |                               |   |
|     |  |  |                               |   |
|     | ОН   |  |                               |   |
|     |  | Let $X$ in the reaction is:  |                               | X   |
|     | СООН   |  |                               | * *   |
|     | ONa  |  |                               | <b>S</b>  |
|     | a)   |  |                               |   |
|     | СООН   |  |                               |   |
|     | COONa  |  |                               |   |
|     | b) [ ( ) ]   |  |                               |   |
|     | $\sim$   |  |                               |   |
|     | OCOONa   |  |                               |   |
|     | c) [ ( )   |  | NY III                        |   |
|     |  | C  | Y                             |   |
|     | d) OH  |  | )                             |   |
|     | COONa  |  |                               |   |
| 28. | Biological oxidation of  | C <sub>6</sub> H <sub>6</sub> taking place in body                                     | of dog, gives:                |   |
|     | a) Benzoic acid  | b) Toluic acid   | c) Maleic acid                | d) Muconic acid   |
| 29. |  |  | ith copper ions in the alkali | ne solutions but not in acidic                            |
|     | solutions .What is the r   |  |                               |   |
|     |  | ydration protects copper i   |                               |   |
|     | ni   | rotons coordinate with am  | monia molecules forming N     | $H_4^+$ ions and $NH_3$ molecules                         |
|     | are not available  |  |                               |   |
|     |  |  | pitated which is soluble in e | excess of any alkali                                      |
| 20  |  | an amphoteric substance  | ductivity in colution?        |   |
| 30. |  | has the highest molar cond<br>b) [Pt(NH <sub>3</sub> ) <sub>5</sub> Cl]Cl <sub>3</sub> | -                             | d) [Pt(NH <sub>3</sub> ) <sub>3</sub> Cl <sub>3</sub> ]Cl |
| 31. |  | is not <i>meta</i> directing grou  |                               |   |
|     | a) $-SO_3H$  | b) —NO <sub>2</sub>  | c) —CN                        | d) —NH <sub>2</sub>                                       |
| 32. | - 0  | is an organometallic comp  | ,                             |   |
|     | a) Lithium methoxide   | 0p   | b) Lithium acetate            |   |
|     | c) Lithium dimethylam  | ine  | d) Methyl lithium             |   |
| 33. |  | wing is very strong o-, p-di   | , ,                           |   |
|     | a) —Cl   | b) —0 <i>R</i>   | c) $-\mathrm{NH}_2$           | d) —NH <i>R</i>   |
| 34. | The type of hybridisati  | on in tetrahedral complexe   | es of metal atom is           |   |
|     | a) $dsp^2$   | b) <i>d</i> <sup>2</sup> <i>sp</i>   | c) <i>sp</i> <sup>3</sup>     | d) $sp^2$   |
| 35. | Chlorobenzene on heat  | ing with NaOH at 300°C un  | der pressure gives:           |   |
|     | a) Phenol  | b) Benzaldehyde  | c) Chlorophenol               | d) None of these  |

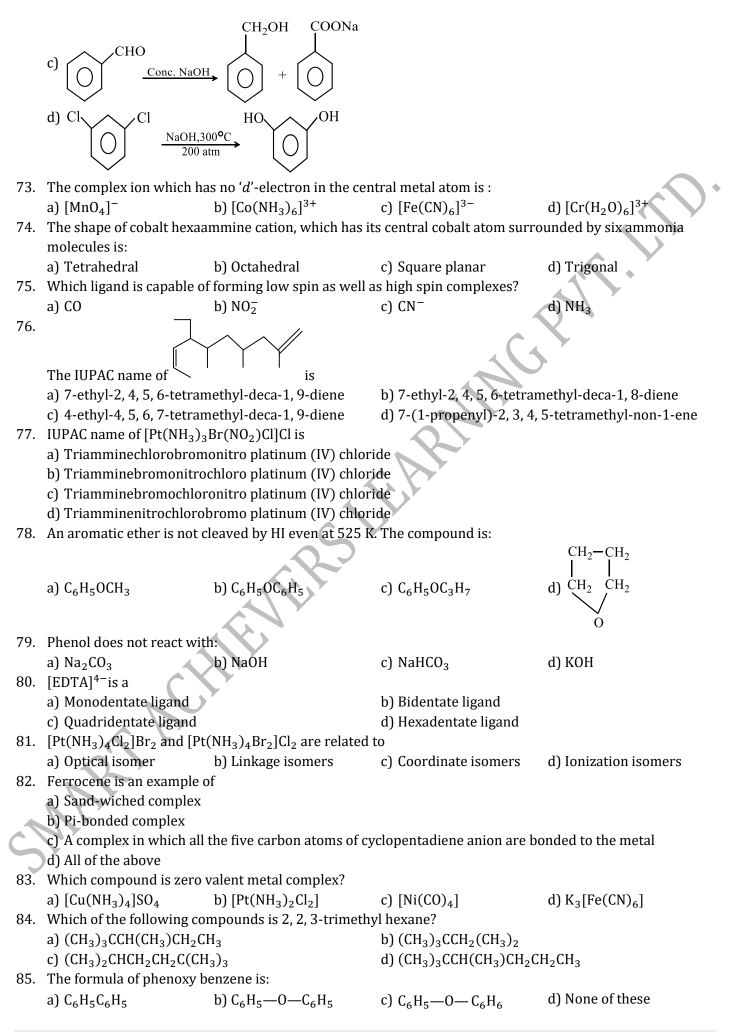
•



49. Number of geometrical isomers for the molecule

|     | c = c < R  |   |   |                             |
|-----|--|---|---|-----------------------------|
|     |  | are   |   |                             |
|     | a) 2   | b) 3  | c) 6  | d) 5                        |
| 50  | •  | oordination number of a ca  | ,   |                             |
| 001 |  | it only a single characterist   |   |                             |
|     |  |   | of ligands bonded to the m  | etal atom                   |
|     |  | =   | the tendency to surround  |                             |
|     | -  | ons as one of the rare gase   | -   |                             |
|     |  |   | nds on the size, structure ar   | nd charge of the ligands    |
| 51. | Among the following, the                         |   |   |                             |
|     | a) C <sub>6</sub> H <sub>5</sub> NH <sub>2</sub> | -   | c) <i>m</i> -NO <sub>2</sub> —C <sub>6</sub> H <sub>4</sub> NH <sub>2</sub> | d) $C_6H_5CH_2NH_2$         |
| 52. | General formula for aren                         |   |   |                             |
|     | a) C <sub>n</sub> H <sub>2n+6</sub>              | b) $C_n H_{2n+6\nu}$  | c) C <sub>n</sub> H <sub>2n</sub>   | d) $C_n H_{2n-6y}$          |
| 53. |  | pesn't have a metal-carbon  | bond?   |                             |
|     | a) Al $(0C_2H_5)_3$                              | b) C <sub>2</sub> H <sub>5</sub> MgBr                                       | c) K[Pt(C <sub>2</sub> H <sub>4</sub> )Cl <sub>3</sub> ]                    | d) Ni(CO) <sub>4</sub>      |
| 54. | How many isomers are p                           | ossible in [Co(en) <sub>2</sub> Cl <sub>2</sub> ]?                          | c) $K[Pt(C_2H_4)Cl_3]$  |                             |
|     | a) 2   | b) 4  | c) 6  | d) 1                        |
| 55. | How many carbon atoms                            | in the molecule $HOOC - (0)$  | $CHOH)_2 - COOH$ are asymm  | netric?                     |
|     | a) 1   | b) 2  | c) 3  | d) None of these            |
| 56. | In benzene, there is a del                       | ocalisation of $\pi$ -electrons. I  | Hence, each $\pi$ -electron is at   | tached bycarbon nuclei.     |
|     | a) 2   | b) 3  | c) 6  | d) 4                        |
| 57. |  | tinguish C <sub>6</sub> H <sub>5</sub> NH <sub>2</sub> and C <sub>6</sub> H |   |                             |
|     |  | l with coupling with pheno  |   |                             |
|     | b) Carbylamine reaction                          |   |   |                             |
|     | c) Reimer-Tiemann react                          | tion  | Y   |                             |
| 50  | d) None of the above                             |   |   |                             |
| 58. |  | e used in Friedel-Craft's re  |   |                             |
|     | a) <i>Cl</i> +                                   | b) RCOCl  | c) $_{RCO}^{+}$   | d) <i>R</i> +               |
| 59. | Thiophene is separated f                         |   |   |                             |
|     | a) Chlorination of thioph                        |   |   |                             |
|     | b) Sulphonation of thiopl                        |   |   |                             |
|     | c) Nitration of thiophene                        |   |   |                             |
| (0  | d) Oxidation of thiophen                         |   |   |                             |
| 60. |  |   |   | of 3 ions when dissolved in |
|     | a) 3   | b) 1  | secondary valencies in this<br>c) 4   | d) Zero                     |
| 61  |  | of alcohol [(CH <sub>3</sub> ) <sub>2</sub> CH] <sub>3</sub> COF            |   | uj zero                     |
| 01. | a) Tri isopropyl carbinol                        |   | b) 2, 4-dimethyl-3-isopro   | ppvl pentan-3-ol            |
|     | c) 2,4-dimethyl-3-(1-me                          | thvl) ethvl pentan-3-ol   | d) None of the above  |                             |
| 62. |  | l complexes can be explain  | -   |                             |
| 5   | a) Completely filled <i>d</i> -or                |   | -   |                             |
| ¥   | b) Vacant <i>d</i> -orbitals                     |   |   |                             |
|     | c) $d - d$ transition                            |   |   |                             |
|     | d) None of the above                             |   |   |                             |
| 63. | Which is most reactive to                        | wards electrophilic reagen  | its:  |                             |



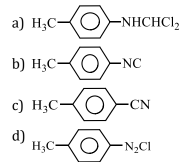


86. Ziegler-Natta catalyst is an organometallic compound containing b) Titanium c) Rhodium d) Zirconium a) Iron 87. Ziegler-Natta catalyst is a)  $(Ph_3P)_3RhCl$ b)  $K[PtCl_3(C_2H_4)]$ c)  $[Al_2(C_2H_6)_6 + TiCl_4]$ d)  $[Fe(C_2H_5)_2]$ 88. The tendency to show complex formation is maximum in ....elements. a) s-block b) p-block c) *d*-block d) f-block 89. EDTA has coordination number a) 3 b) 4 c) 5 d) 6 90. Which of the following is used in Friedel-Craft's acylation reaction? CH<sub>3</sub>CO b) CH<sub>3</sub>CH<sub>2</sub>Cl c) CH<sub>3</sub>COOCH<sub>3</sub> d) CH<sub>3</sub>Cl a) CH<sub>3</sub>CO 91. The correct IUPAC name of  $Mn_3(CO)_{12}$  is a) Dodacacarbonyl maganate (0) b) Dodacacarbonyl maganate (II) c) Didacacarbonyl trimaganese (0) d) Manganic dodecacarbanyl (0) 92. The  $\pi$  –bonded organometallic compound which has ethene as one of its component is a) Zeise's salt b) Ferrocene c) Dibenzene chromium d) Tetraethyl tin 93. IUPAC name of the compound CH<sub>3</sub>--COOC<sub>2</sub>H<sub>5</sub> is b) Ethyl-2-methyl-2-(o-nitro) phenyl propanoate a) Ethyl-2-methyl-2-(*m*-nitro) phenyl propanoate c) Ethyl-2-methyl-2-(3-nitro) phenyl propanoate d) Ethyl-2-methyl-2-(3-nitro) phenyl propanoic acid 94. What is the product obtained in the following reaction: NH  $NH_2$ d) a) 95.  $[Co(NH_3)_6]Cl_3$  is called: a) Hexaammine cobalt (III) chloride b) Amino cobalt chloride (III) c) Cobalt chloride hexaammine d) Hexaammine tricobalt chloride 96. The complexes  $[PtCl_2(NH_3)_4]Br_2$  and  $[PtBr_2(NH_3)_4]Cl_2$  are example for isomerism a) Geometrical c) Ionization b) Optical d) Linkage 97. Geometrical shapes of the complexes formed by the reaction of  $Ni^{2+}$  with  $Cl^{-}$ ,  $CN^{-}$  and  $H_2O$ , respectively, are a) Octahedral, tetrahedral and square planar b) Tetrahedral, square planar and octahedral c) Square planar, tetrahedral and octahedral d) Octahedral, square planar and octahedral 98. Identify the correct order of reactivity in electrophilic substitution reactions of the following compounds: a) 1 > 2 > 3 > 4b) 4 > 3 > 2 > 1c) 2 > 1 > 3 > 4 d) 2 > 3 > 1 > 4

99. The centric formula for benzene was proposed by:

| a) Dewar  |   |   |
|---|---|---|
| b) Armstrong and Baeyer   |   |   |
| c) Ladenberg  |   |   |
| d) Kekule   |   |   |
| 100. Which is the correct statement?                                  |   |   |
| a) Benzyl alcohol is more acidic than phenc                           | bl  |   |
| b) Ethanol is a powerful oxidizing agent                              |   |   |
| c) Phenol is more acidic than propanol                                |   |   |
| d) Ethane has high boiling point than ethan                           | ol  |   |
| 101. Phenol on sulphonation gives:                                    |   |   |
| a) o-phenol sulphonic acid  |   |   |
| b) <i>p</i> -phenol sulphonic acid                                    |   |   |
| c) <i>m</i> -phenol sulphonic acid                                    |   |   |
| d) Mixture of <i>o</i> -and <i>p</i> -phenol sulphonic acid           | ds  |   |
| 102. Which of the following organometallic com                        | -   |   |
| a) $Fe(CH_3)_3$ b) $[Co(CO)_5NH_3]$                                   | ] <sup>2+</sup> c) [Fe( $\eta^5 - C_5 H_5$ ) <sub>2</sub> ] | d) K[PtCl <sub>3</sub> ( $\eta^2 - C_2H_4$ )]                                   |
| 103. The number of double bonds in BHC (gamm                          | nexane) is:   |   |
| a) 1 b) 2   | c) 3  | d) Zero   |
| 104. Given the molecular formula of the hexa co                       | ordinated complexes (A) $CoCl_3$                            | $\cdot$ 6NH <sub>3</sub> ( <i>B</i> )CoCl <sub>3</sub> $\cdot$ 5NH <sub>3</sub> |
| (C) $CoCl_3 \cdot 4NH_3$ . If the number of coordination              | ted $NH_3$ molecules in A, B and C                          | C respectively are 6, 5 and 4,  |
| primary valency in (A), (B) and (C) are                               |   |   |
| a) 0, 1, 2 b) 3, 2, 1   | c) 6, 5, 4  | d) 3, 3, 3  |
| 105. Type of isomerism shown by [Cr(NH <sub>3</sub> ) <sub>5</sub> NO | $[2]Cl_2$ is  |   |
| a) Optical b) Ionisation  | c) Geometrical  | d) Linkage  |
| 106. $[Sc(H_2O)_6]^{3+}$ ion is                                       |   |   |
| a) Colourless and diamagnetic   | b) Coloured and octa  | hedral  |
| c) Colourless and paramagnetic  | d) Coloured and para  | magnetic  |
| 107. Which one of the following octahedral com                        | plexes will not show geometrica                             | ll isomerism? ( <i>A</i> and <i>B</i> are                                       |
| monodentate ligands)  |   |   |
| a) $[MA_4B_2]$ b) $[MA_5B]$   | c) $[MA_2B_4]$  | d) $[MA_3B_3]$  |
| 108. The IUPAC name of the following compound                         | d is  |   |
| $O = C - CH - CH_2$   |   |   |
| $ $ $ $ $ $ $ $ OH NH <sub>2</sub> OH                                 |   |   |
| a) 3-amino-2-hydroxy propanoic acid                                   | b) 2-aminopropan-3-   | ol-1-oic acid   |
| c) 2-amion-3-hydroxy propanoic acid                                   | d) Aminohydroxy pro   |   |
| 109. Which of the following complex ion is not e                      |   |   |
| a) [Ni(CN) <sub>4</sub> ] <sup>2-</sup>                               | xpected to absorb visible light:                            |   |
| b) $[Cr(NH_3)_6]^{3+}$  |   |   |
| c) $[Fe(H_2O)_6]^{2+}$  |   |   |
|   |   |   |
| d) $[Ni(H_2O)_6]^{2+}$  | fin hin   |   |
| 110. The correct sequence of activating power o                       | r a group in benzene is:                                    |   |
| a) $-NH_2 > -NHCOCH_3 > -CH_3$<br>b) $-NH_2 < -NHCOCH_3 < -CH_3$      |   |   |
|   |   |   |
| c) $-NH_2 > -NHCOCH_3 < -CH_3$  |   |   |
| d) $-NH_2 < -NHCOCH_3 > -CH_3$  |   |   |
| 111. The pair of compounds having metals in the                       |   |   |
| a) $MnO_2$ , FeCl <sub>3</sub>  | b) $[MnO_4]^-$ , $CrO_2Cl_2$                                | _   |
| c) $[Fe(CN)_6]^{3-}$ , $[Co(CN)_3]$                                   | d) $[NiCl_4]^{2-}$ , $[CoCl_4]^{-}$                         |   |
| 112. Total number of geometrical isomers for th                       |   |   |
| a) 1 b) 2   | c) 3  | d) 4  |

113. The reaction of chloroform with alc. KOH and *p*-toluidine forms:



114. Which order is correct in spectrochemical series of ligands?

a)  $Cl^- < F^- < [C_2O_4]^{2-} < NO_2^- < CN^-$ 

b)  $CN^- < [C_2O_4]^{2-} < Cl^- > NO_2^- < F^-$ 

c)  $[C_2O_4]^{2-} < F^- < Cl^- > NO_2^- < CN^-$ 

d)  $F^- < Cl^- < NO_2^- < CN^- < [C_2O_4]^{2-}$ 

- 115. The IUPAC name of compound  $K_3$ [Fe(CN)<sub>5</sub>NO] is
  - a) Pentacyano nitrosyl potassium ferrate(II)
  - c) Potassium pentacyanonitrosyl ferrate (III)

116. The colour of  $[Ti(H_20)_6]^{3+}$  is due to:

- a) Transfer of an electron from one Ti to another
- b) Presence of water molecule
- c) Excitation of electrons from d d

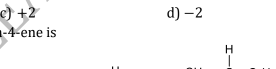
d) Intramolecular vibration

117. The oxidation number of Fe in  $K_4[Fe(CN)_6]$  is

118. Correct structures of [E][S]-5-bromo-2,7-dimetyl, non-4-ene is

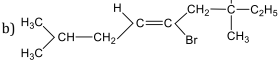
a)  $\overset{H_3C}{\underset{H_3C}{\longrightarrow}}CH - CH_2 \overset{H}{\underset{CH_2}{\longrightarrow}}C = C \overset{Br}{\underset{CH_2}{\overset{L}{\underset{CH_2}{\longrightarrow}}}C} \overset{CH_3}{\underset{H_3}{\overset{L}{\underset{CH_2}{\longrightarrow}}}C}$ 

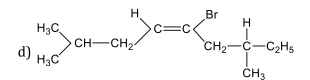
c) 
$$\overset{H_3C}{\underset{H_3C}{\longrightarrow}}$$
 cH-CH<sub>2</sub> CH-CH<sub>2</sub> CH-CH<sub>2</sub> CH<sub>3</sub>  
 $\overset{CH_3}{\underset{Br}{\longrightarrow}}$  CH-CH<sub>2</sub> CH-CH<sub>2</sub> CH-CH<sub>3</sub>



b) Potassium cyano pentanitrosyl ferrate(II)

d) Potassium pentacyanonitrosyl ferrate (II)





119. Name the metal *M* which is extracted on the basis of following reactions,  $4M + 8CN^- + 2H_2O + O_2 \rightarrow 4[M(CN)_2]^- + 4OH^-$ 

 $2[M(CN)_2]^- + Zn \longrightarrow [Zn(CN)_4]^{2-} + 2M:$ a) Nickel b) Silver c) Copper d) Mercury 120. EAN of Cr in [Cr(NH<sub>3</sub>)<sub>6</sub>]Cl<sub>3</sub> is: a) 32 c) 34 d) 35 b) 33 121. The complex  $[Pt(NH_3)_6]Cl_4$  furnishes: Ja) 5 ions b) 6 ions c) 4 ions d) 2 ions 122. Ammoniacal solution of Ni(CN)<sub>2</sub> reacts with C<sub>6</sub>H<sub>6</sub> to produce a light violet coloured crystalline compound of the formula: a) Ni(CN)<sub>2</sub>  $\cdot$  C<sub>6</sub>H<sub>5</sub> b)  $C_6H_5CH_3$ c) Ni(CN)<sub>2</sub>C<sub>6</sub>H<sub>6</sub> d) Ni(CN)<sub>2</sub>NH<sub>3</sub>  $\cdot$  C<sub>6</sub>H<sub>6</sub>

123. Ammonia forms the complex ion  $[Cu(NH_3)_4]^{2+}$  with copper ions in alkaline solution but not in acidic solution. What is the reason for it?

a) In acidic solutions, hydration protects copper ions

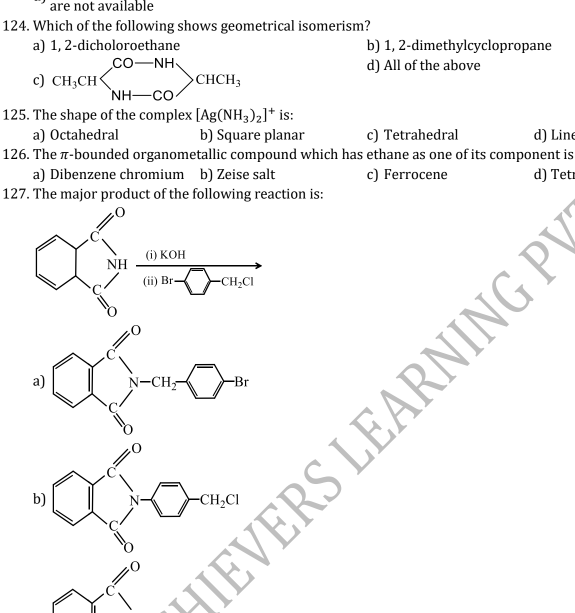
b) In alkaline solution, insoluble Ci(OH)<sub>2</sub> is precipited which in excess of any alkali

- c) Copper hydroxide is an amphoteric substance
- In acidic solutions, protons coordinate with ammonia molecules forming  $NH_4^+$  ions and  $NH_3$  molecules are not available

d) Linear

d) Tetraethyl tin

124. Which of the following shows geometrical isomerism?



128. Which is true in the case of  $Ni(CO)_4$  complex?

- a) Hybridization of Ni is  $sp^3$
- b) Tetrahedral shape of the molecule
- c) Diamagnetic

c)

d)

d) All are correct

129. The reaction,  $C_6H_5N_2Cl \xrightarrow{Cu_2Cl_2/HCl} C_6H_5Cl + N_2$  is called:

CH<sub>2</sub>Cl

a) Etard's reaction b) Sandmeyer's reaction c) Wurtz-Fittig reaction d) Perkin's reaction

| 130. Which of the following does not show optical iso               | merism?                     |                                   |
|---|-----------------------------|-----------------------------------|
|   | c) $[Co(NH_3)_3Cl_3]^0$     | d) $[Co(en)Cl_2(NH_3)_2]^+$       |
| 131. CH <sub>3</sub>  |                             |                                   |
|   |                             |                                   |
|   |                             |                                   |
| CH <sub>3</sub>   |                             |                                   |
| Having the IUPAC name as  |                             |                                   |
| a) 1, 2-dimethyl cyclobutane  | b) 2, 3-dimethyl cyclo      |                                   |
| c) 2, 3-dimethyl butane   | d) 1, 2-dimethyl cyclo      |                                   |
| 132. Which of the following ions is produced when we                | e prepare nitrating mixture | e by mixing together              |
| concentrated $HNO_3$ and concentrated $H_2SO_4$ ?                   |                             |                                   |
| a) $NO_2^-$ b) $NO_2^+$   | c) NO <sub>3</sub>          | d) SO <sub>3</sub> <sup>+</sup> H |
| 133. The correct IUPAC name of                                      |                             |                                   |
| F<br>  D.   |                             |                                   |
| Br  |                             |                                   |
| is  |                             |                                   |
|   |                             |                                   |
| a) 1-brmo-2-chloro-6-fluoro-4-iodobenzene                           | -                           | 2-fluoro-4-iodobenzene            |
| c) 2-bromo-1-chloro-3-floro-5-iodobenzene                           | d) 2-bromo-3-chloro-        | 1-floro-5-odobenzene              |
| 134. $[Co(NH_3)_4(NO_2)_2]$ Cl exhibits:                            |                             |                                   |
| a) Ionization isomerism, geometrical isomerism                      |                             |                                   |
| b) Linkage isomerism, geometrical isomerism an                      |                             |                                   |
| c) Linkage isomerism, ionization isomerism and                      |                             |                                   |
| d) Linkage isomerism, ionization isomerism and                      |                             |                                   |
| 135. Which of the following complexes are not correct $\frac{1}{2}$ | tly matched with hybridisa  | tion of their central metal ion?  |
| 1. $[Ni(CO)_4]$ $sp^3$  |                             |                                   |
| 2. $[Ni(CO)_4]^{2-}$ $sp^3$   |                             |                                   |
| 3. $[CoF_6]^{3-}$ $d^2sp^3$   |                             |                                   |
| 4. $[Fe(CN)_6]^{3-}$ $sp^3d^2$                                      | ,                           |                                   |
| Select the correct answer using the codes given b                   |                             |                                   |
| a) 1 and 2 b) 1 and 3   | c) 2 and 4                  | d) 2, 3 and 4                     |
| 136. Which of the following is an explosive?                        |                             |                                   |
| a) $PCl_5$ b) $HNO_3$   | c) $C_6H_5OH$               | d) 2,4,6-trinitrophenol           |
| 137. The coordination number of Cr in $[Cr(NH_3)_3(H_2)]$           |                             |                                   |
| a) 3 b) 4   | c) 6                        | d) 2                              |
| 138. The major product obtained when 3-phenyl-1, 2-                 | -propane-diol is neated wit | $H_2 SO_4 IS:$                    |
| a) $C_6H_5$ — $CH_2$ — $CO$ — $CH_3$                                |                             |                                   |
| b) $C_6H_5$ $-CH_2$ $-CH_2$ $-CHO$                                  |                             |                                   |
| c) $C_6H_5$ — $CH_2$ — $CH = CH_2$                                  |                             |                                   |
| d) $C_6H_5 - CH_2 - CH CH_2$  |                             |                                   |
|   |                             |                                   |
| 139. Rate of substitution in phenol is:                             |                             |                                   |
| $\checkmark$ a) Slower than as in benzene                           |                             |                                   |
| b) Faster than as in benzene  |                             |                                   |
| c) Equal to that as in benzene                                      |                             |                                   |
| d) None of the above  |                             |                                   |
| 140. Magnetic moment of $[Ag(CN)_2]^-$ is zero. How ma              | any unpaired electrons are  | there?                            |
| a) Zero b) 4  | c) 3                        | d) 1                              |
| 141. Chlorophyll is a coordination compound having o                | central atom of:            | -                                 |
| a) Ca b) Mg   | c) Na                       | d) K                              |
|   | -                           | -                                 |

142. Which of the following statements is incorrect? a) In  $K_3$  [Fe(CN)<sub>6</sub>], the ligand has satisfied only the secondary valency of ferric ion. b) In  $K_3$  [Fe(CN)<sub>6</sub>], the ligand has satisfied both primary and secondary valencies of ferric ion. c) In K<sub>4</sub>[Fe(CN)<sub>6</sub>], the ligand has satisfied both primary and secondary valencies of ferrous ion. d)  $In[Cu(NH_3)_4]SO_4$ , the ligand has satisfied only the secondary valenecy of copper. 143. Maximum number of open chain isomers that an alkene can have with the molecular formula  $C_4H_8$  is a) 5 b) 4 c) 3 d) 2 144. Which one is the wrong statement? a) Open chain compounds are called aliphatic b) Unsaturated compounds contain multiple bonds in them c) Saturated hydrocarbons are called alkene d) Aromatic compounds possess a characteristic aroma 145. According to postulates of Werner's theory for coordination compounds, which of the following is true? a) Primary valencies are ionizable b) Secondary valencies are ionizable d) Primary and secondary valencies are nonc) Only primary valencies are non-ionizable ionizable 146. Atomic numbers of Cr and Fe are respectively 24 and 26. Which of the following is paramagnetic with the spin of the electron? a)  $[Cr(CO)_6]$ b)  $[Fe(CO)_5]$ c)  $[Fe(CN)_6]^{4-1}$ d)  $[Cr(NH_3)_6]^{3+}$ 147. Which of the following structures correspond to the product expected, when excess of  $C_6H_6$  reacts with CH<sub>2</sub>Cl<sub>2</sub> in presence of anhy. AlCl<sub>3</sub>? CHCl<sub>2</sub> b) · C1148. Which of the following will give a pair of enantiomorphs? a)  $[Co(en)_2Cl_2]Cl$ b)  $[Cr(NH_3)_6][Co(CN)_6]$ c)  $[Pt(NH_3)_4][PtCl_6]$ d)  $[Co(NH_3)_4Cl_2]NO_2$ 149. The crystal field splitting energy for octahedral( $\Delta_0$ ) and tetrahedral ( $\Delta_t$ ) complexes is related to b)  $\Delta_t = \frac{1}{2} \Delta_0$ c)  $\Delta_0 = 2\Delta_t$ a)  $\Delta_t = \frac{4}{9} \Delta_0$ d)  $\Delta_0 = \frac{4}{0} \Delta_t$ 150. The correct name of the compound  $[Cu(NH_3)_4](NO_3)_2$ , according to IUPAC system is: a) Cuprammonium nitrate b) Tetraamminecopper(II) dinitrate c) Tetraamminecopper(II) nitrate d) Tetraamminecopper(I) dinitrate 151. Which among the following will not show chain isomerism? b)  $C_4H_{10}$ c)  $C_5 H_{12} O$ d)  $C_5 H_{10} O$ a)  $C_3H_8$ 152. Phenol (1 mole) reacts with bromine to give s-tribromophenol. How much bromine is needed? a) 1.5 mole b) 3.0 mole c) 4.5 mole d) 6.0 mole 153. Dimethyl glyoxime forms a coloured complex with a) Ag b) Ni c) Cr d) Zn 154. Which has regular tetrahedral geometry? a)  $[Ni(CN)_4]^{2+}$ c)  $[BF_4]^$ d) XeF<sub>4</sub> b)  $SF_4$ 155. In haemoglobin the iron shows oxidation state: a) +2 b) +3c) +1 d) +4 156. For the given complex  $[CoCl_2(en)(NH_3)_2]^+$ , the number of geometrical isomers, the number of optical isomers and total number of isomers of all type possible respectively are a) 2, 2 and 4 b) 2, 2 and 3 c) 2, 0 and 2 d) 0, 2 and 2 157. Which can show aromatic character?

| 1               |
|-----------------|
| 1<br><b>) •</b> |
| 1               |
| 1<br>) •        |
| 2.              |
| 2.              |
| 2.              |
| 2.              |
| >.              |
|                 |
| F               |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
| 0 M 0           |
| one             |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
| vn              |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
|                 |
| v               |

b) Tautomerism occurring in phenol

c) The fact that the electronegativity of oxygen is more than that of hydrogen

d) None of the above

175. In triethylenediamine cobalt(III) chloride the coordination number of cobalt is:



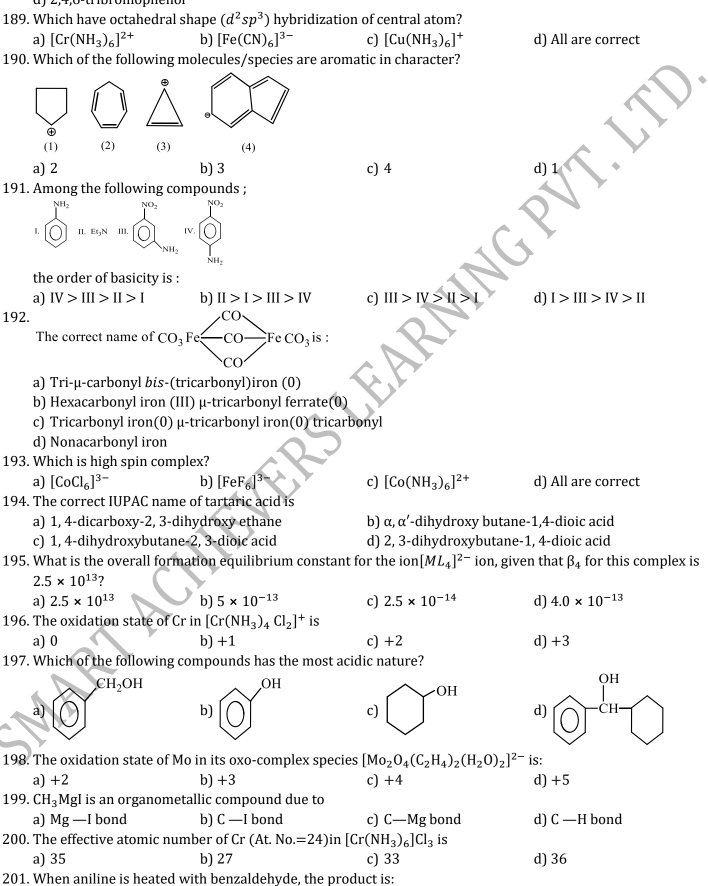
188. When phenol is treated with excess bromine water, it gives:

a) *m*-bromophenol

b) *o*-and *p*-bromophenol

c) 2,4-dibromophenol

d) 2,4,6-tribromophenol

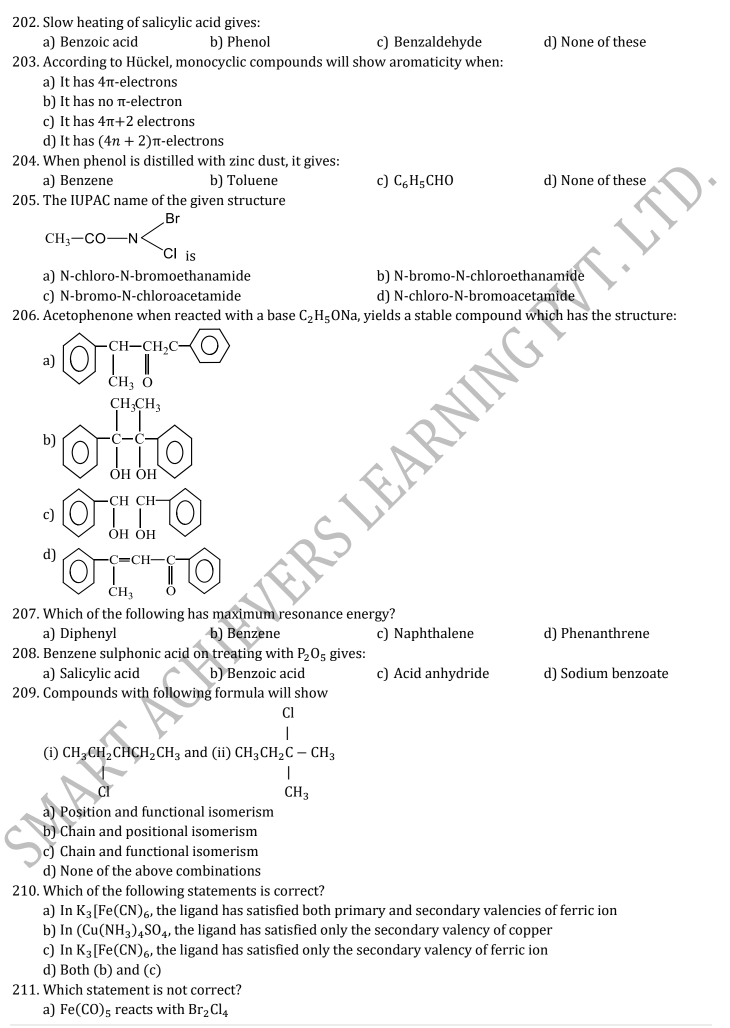


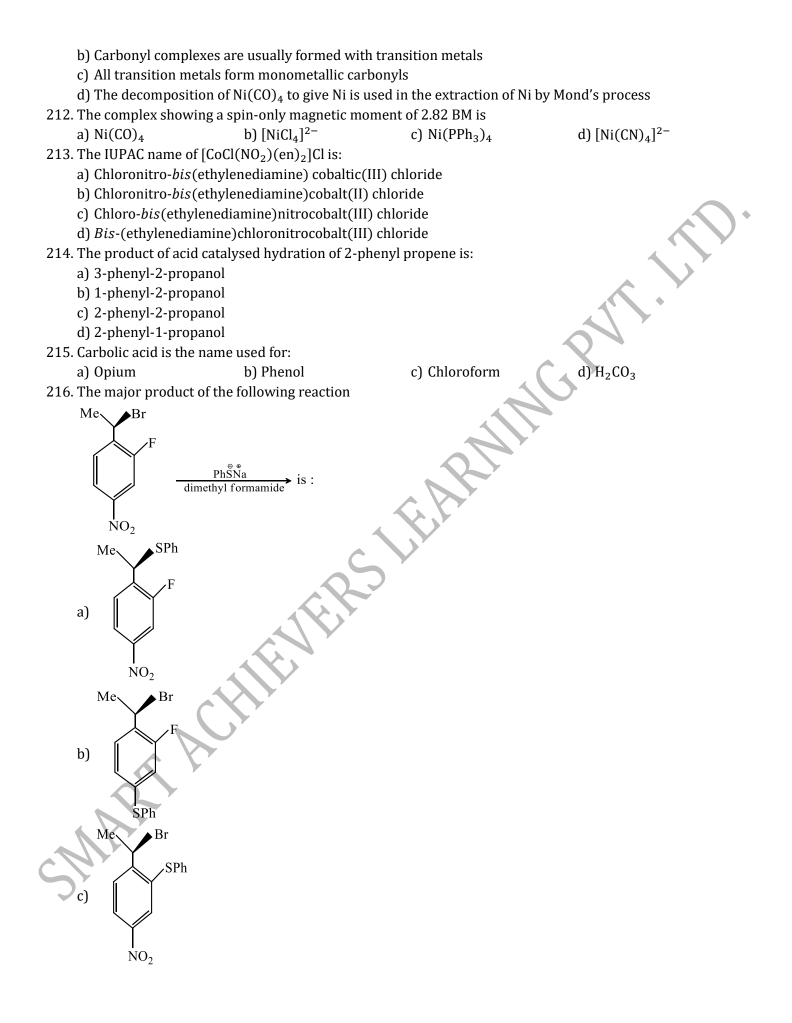
c) Unsaturated acid

b) Schiff's base

a) Benzoin

d) Azoxy benzene

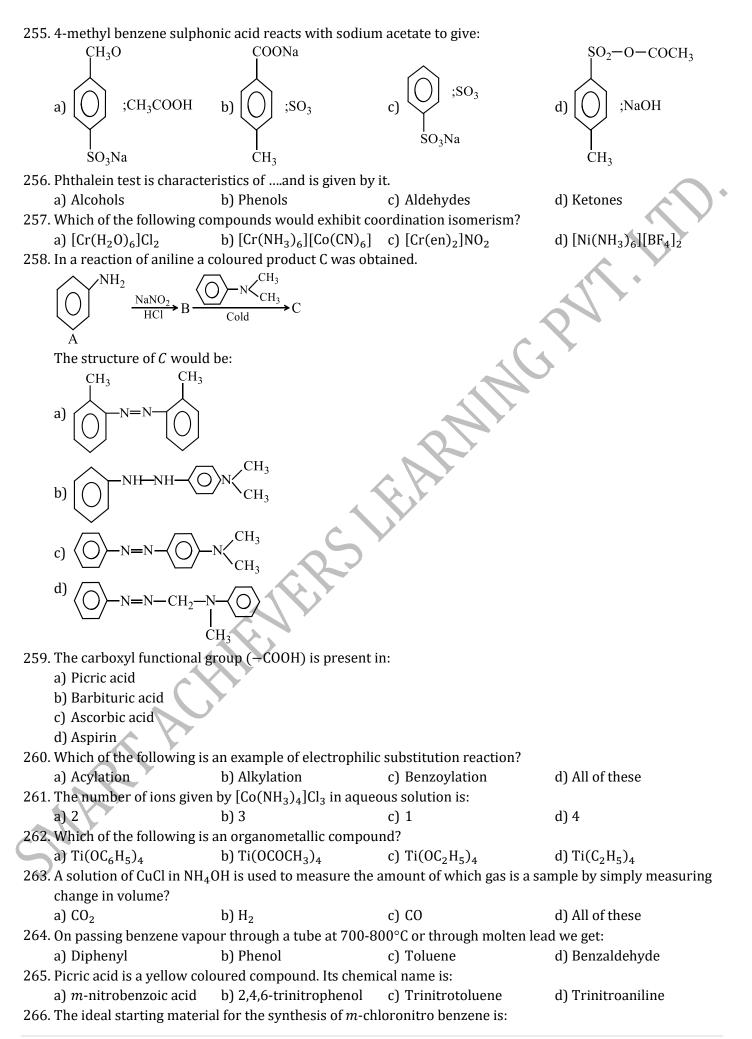


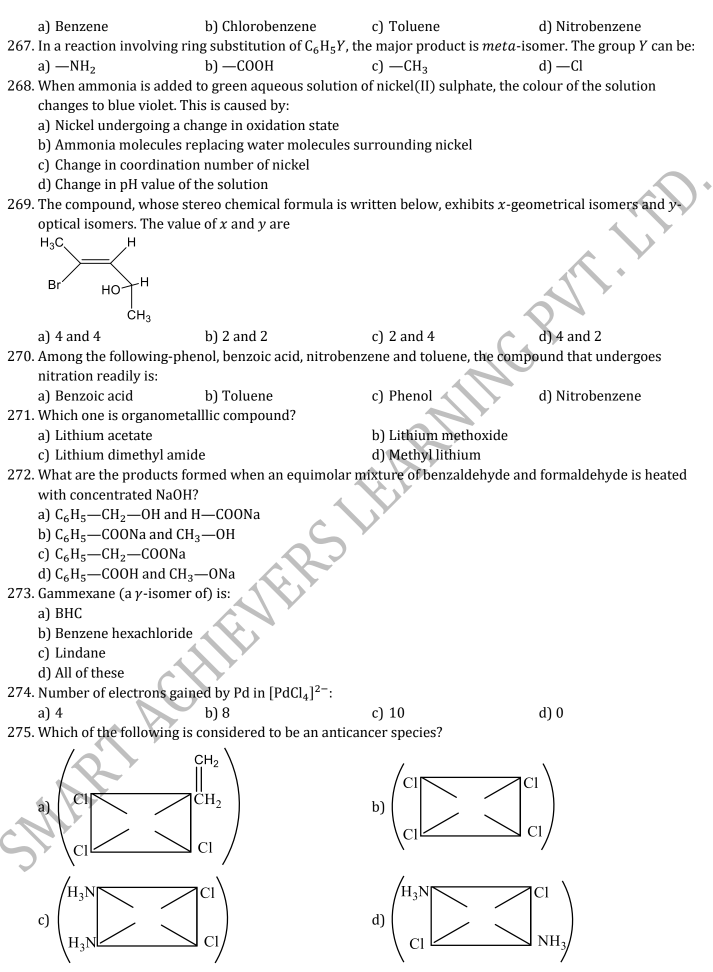


d) Me SPh SPh NO<sub>2</sub> 217. The oxidation number of cobalt in  $K[Co(CO)_4]$  is d) −3 a) –1 b) +3c) +1 218. Formaldehyde-phenol resin is: a) Orlon b) Nylon c) Teflon d) Bakelite 219. Among the ligands NH<sub>3</sub>, en, CN<sup>-</sup> and CO, the correct order of their increasing field strength, is a)  $CO < NH_3 < en < CN^$ b)  $NH_3 < en < CN^- < CO$ c)  $CN^{-} < NH_3 < CO < en$ d) en  $< CN^{-} < NH_3 < CO$ 220. Cyclopentadienyl anion is aromatic due to the presence of: d) 12 π-electrons a)  $6\pi$ -electrons b) 10  $\pi$ -electrons c) 4  $\pi$ -electrons 221. The IUPAC name of  $K_4[Fe(CN)_6]$  is b) Potassium hexa cyanoferrate (I) a) Potassium ferrocyanide d) Potassium hexa cyanoferrate (II) c) Tetra potassium hexa cyanoferrate (II) 222. Which xylene is most easily sulphonated? c) Meta a) Ortho b) Para d) All at the same rate 223. The IUPAC name of following polyfunctional compound is соон OHC н a) 2,4-dioxo cyclohexanoic acid b) 2,4-dioxo cycloheptanoic acid c) 4-formyl-2-oxo cyclohexane-1-carboxylic acid d) 2,4-dioxo cyclohexane-1-carboxylic acid 224. Alkyl groups are *o*- and *p*-directing because of: a) Resonance effect b) Inductive effect c) Resonance effect through hyperconjugation d) All of the above 225. Racemic modification can be resolved by a) The use of enzymes b) Fractional crystallisation c) Fractional distillation d) None of the above 226. Which of the following structure contain 1 primary and 7 secondary hydrogen atoms?  $CH_3 - CH - CH_2 - CH_2$ CH2  $CH - CH_3$  $CH_3$ CH<sub>2</sub> b)  $CH_3 - CH_2 - CH_2 - CH$  $CH_2 - CH_3$ CH<sub>3</sub> c)  $\mathrm{CH}_3-\mathrm{CH}_2-\mathrm{CH}-\mathrm{CH}_2-\mathrm{CH}_3$ d)  $CH_3 - CH_2 - CH_2 - CH_2$ 

 $CH_2 - CH_3$ 227. Which of the following compounds does not dissolve in conc. H<sub>2</sub>SO<sub>4</sub> even on warning? a) Ethylene b) Benzene c) Hexane d) Aniline 228. In the complex  $Fe(CO)_x$ , the value of x is and it is: c) 5, trigonal pyramidal a) 3, octahedral b) 4, tetrahedral d) 6, square pyramidal 229. The empirical formula of naphthalene is: b)  $C_5 H_4$ c)  $C_2H$ d)  $C_n H_{2n}$ a)  $CH_2$ 230. The chemical formula of diammine silver (I) chloride is b)  $[Ag(NH_3)_3]Cl$ c)  $[Ag(NH_4)_2]Cl$ d)  $[Ag(NH_3)_2]C$ a)  $[Ag(NH_3)Cl]$ 231. For the square planar complex [M(a)(b)(c)(d)] (where , *M*=central metal and *a*, *b*, *c*, and *d* are monodentate ligands), the number of possible geometrical isomers are a) 1 b) 2 c) 3 232. Which group is *meta* directing? a)  $-CCl_3$ b) -0H c)  $-NH_2$ 233. The IUPAC name of the compound  $[Cu(NH_3)_4(NO_3)_2]$  is: a) Cuprammonium nitrate b) Dinitratotetraamminecopper(II) c) Tetraamminecopper(II) dinitrite d) Tetraamminecopper(III) dinitrite 234. Coordination number of Fe in the complexes  $[Fe(CN)_6]^{4-}$ ,  $[Fe(CN)_6]^{3-}$  and  $[FeCl_4]^-$  would be respectively a) 6, 4, 6 b) 6, 6, 4 c) 6.3.3 d) 2.3.3 235. Which statement is true for cyclohexane? a) It has two possible isomers b) It has three conformations c) Boat conformation is most stable d) Chair and boat conformations differ in energy by 44 kJ/mol 236. Ligands in a complex salt are: a) Anions linked by coordinate bonds to a central metal atom or ion b) Cations linked by coordinate bonds to a central metal atom or ion c) Molecules linked by coordinate bonds to a central metal atom or ion d) Ions or molecules linked by coordinate bonds to a central metal atom or ion 237. The IUPAC name of CH<sub>3</sub>COCH<sub>2</sub>COOC<sub>2</sub>H<sub>5</sub> is a) Ethyl butanoate b) Ethyl-(3-oxo)butanoate c) Ethyl butan-1-oate-2-one d) Ethyl butan-4-oate-2-one 238. When benzene is treated with CO and HCl in presence of anhydrous aluminium chloride, benzaldehyde is formed. This reaction is known as: a) Friedel-Craft's reaction b) Rosenmund's reaction c) Stephen's reaction d) Gattermann-Koch's reaction 239.  $[Cr(NH_3)_6]^{3+}$  ion is: a) Paramagnetic b) Diamagnetic c) Square planar d) None of these 240. The following compound can exhibits a) Tautomerism b) Optical isomerism c) Geometrical isomerism d) Geometrical and optical isomerism 241. Which complex is diamagnetic?

a)  $[Fe(CN)_6]^{4-}$ b)  $[Cu(NH_3)_4]^{3+}$ c)  $[Ti(H_20)_6]^{3+}$ d) None of these 242. Meso-tartaric acid is optically inactive due to the presence of a) Molecular symmetry b) Molecular asymmetry c) External compensation d) Two asymmetric C-atoms 243. Complex forming tendency increases with: a) Increase in size of cation b) Decrease in size of cation c) Increase in size of anion d) None of the above 244. Ziegler-Natta catalyst is b)  $Al_2(C_2H_6)_6 + TiCl_4$ a) (Ph<sub>3</sub>P)<sub>3</sub>RhCl c)  $Fe(C_2H_5)_2$ d) K[PtCl<sub>3</sub>(C<sub>2</sub>H<sub>4</sub>)] 245. Among the following compounds the one that is most reactive towards electrophilic nitration is: d) Nitrobenzene a) Toluene b) Benzene c) Benzoic acid 246. Phenol on oxidation gives chloranil. The oxidant used is: b) KMnO<sub>4</sub> c)  $KClO_3 + HCl$ d) None of these a)  $K_2 S_2 O_8$ 247. The IUPAC name of the compound  $\mathsf{CH}_3 \underline{-\!\!\!\!-} \overset{}{\mathsf{CH}} \underline{-\!\!\!\!-} \mathsf{CH}_2 \underline{-\!\!\!\!-} \mathsf{CH}_3$  $-CH-CH-CH_2-CH_2-CH_3$ CH<sub>3</sub>-CH<sub>2</sub>is a) 3-sec-butyl-5-ethyl-3-methyloctane b) 4-sec-butyl-5-ethyl-3-methyloctane c) 5-sec-butyl-4-ethyl-3-methyloctane d) 4-sec-butyl-3-ethyl-5-methyloctane 248. All the common *m*-directing groups.....the benzene ring towards electrophilic substitution reactions. a) Deactivate b) Activate c) Both (a) and (b) d) None of these 249. Among the following, the coloured compound is : a) CuCl b)  $K_3C_4(CN)_4$ c)  $CuF_2$ d)  $[Cu(CH_3CN)_4]BF_3$ 250. The existence of two different coloured complexes with the composition of  $[Co(NH_3)_4Cl_2]^+$  is due to: a) Linkage isomerism b) Geometrical isomerism c) Coordination isomerism d) Ionisation isomersim 251. [Co(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub>] possesses: a) Square planar geometry b) Tetrahedral geometry c) Tetrahedral nature d) Octahedral geometry 252. Which one does not belong to ligand? a) PH<sub>3</sub> b) N0<sup>+</sup> c)  $BF_3$ d) Cl<sup>-</sup> 253. Product formed in the reaction;  $\xrightarrow{(CH_3)SO} \text{Product; is:}$ OH OН OH CH<sub>3</sub> ·CH<sub>2</sub>SCH<sub>3</sub> b) a) c) CH<sub>2</sub>SCH<sub>3</sub> 254. Which one of the following has square planar structure? c) [NiCl<sub>4</sub>]<sup>2-</sup> a)  $[Ni(CN)_4]^{2-}$ b)  $[Ni(CO)_4]$ d) All of these





276. For benzaldehyde which of the following is incorrect?

a) It is an aromatic aldehyde

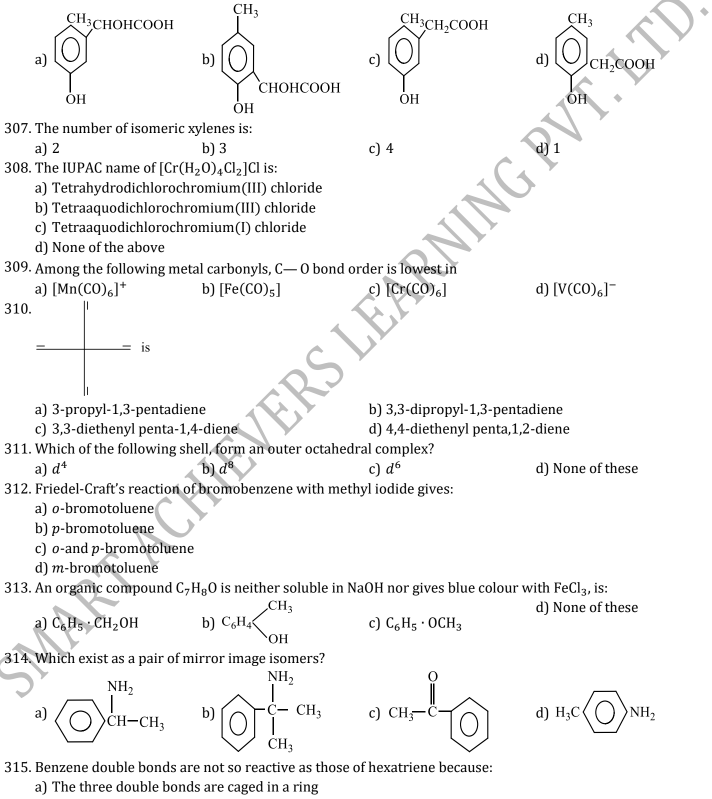
b) It is used in perfumery c) On oxidation it yields benzoic acid d) On reduction it yields phenol 277. The main source of aromatic compounds is: b) Petroleum d) Both (b) and (c) a) Wood c) Coal 278. Phenol on hydrogenation in presence of a nickel catalyst at 160°C gives: a) Benzene b) Cyclohexane c) Cyclohexanol d) *n*-hexanol 279. The IUPAC name of compound C≡N N is a) Hexane-1, 2, 5-tricarbonitrile b) Hexane-1, 3, 6-tricarbonitrile c) Butane-1, 2, 4-tricarbonitrile d) Butane-1, 3, 4-tircarbonitrile 280. Fac-mer isomerism is associated with which one of the following complexes? (M=central metal) d) [MABCD] a)  $[M(AA)_2]$ b)  $[MA_{3}B_{3}]$ c)  $[M(AA)_3]$ 281. Which of the following is the correct order of stability of the following four distinct conformation of *n* butane? a) Staggered > Gauche > Partially eclipsed > Fully eclipsed b) Gauche > Staggered > partially eclipsed > Fully eclipsed c) Staggered > Partially eclipsed > Gauche > Fully eclipsed d) Fully eclipsed > Staggered > Partially eclipsed > Gauche 282. *o*-nitrophenol can form hydrogen bonds within the molecule. It thus, has: b) Very high viscosity c) Low m.p. d) none of these a) Very high m.p. 283. The element which does not form mononuclear carbonyl is: d) W b) Mn c) Ni a) Fe 284. Which of the following is hexadentate ligand? b) Ethylene diamine tetra acetic acid a) Ethylene diamine c) 1,10-phenanthroline d) Acetyl acetonato 285. The molecular formula of a saturated compound is  $C_2H_4Cl_2$ . The formula permits the existence of two b) Position isomers c) Optical isomers a) Functional isomers d) cis – trans isomers 286. An octahedral complex is formed when hybrid orbitals of the following type are involved a)  $sp^3$ b)  $dsp^2$ c)  $d^2sp^3$ d)  $sp^2d^2$ 287. The IUPAC name of the given compound  $CH_3 - CH = CH - COOC_2H_5$  is a) Ethyl propenoate b) Ethyl-2-butenoate c) Ethyl-1-butenoate d) Propene ethyl methanoate 288. Which product is not obtained by heating wood or coal in the absence of air? c) Benzene a) Coal-tar b) Naphthalene d) Wax 289. Dry distillation of calcium benzoate with calcium formate gives: a) Acetaldehyde b) Benzoic acid c) Benzaldehyde d) Benzoic anhydride 290. Which will give Fe<sup>3+</sup> ions in solution? a)  $[Fe(CN)_6]^{3-1}$ b)  $Fe_2(SO_4)_3$ c)  $[Fe(CN)_6]^{4-}$ d)  $NH_4(SO_4)_2 \cdot FeSO_4 \cdot 6H_2O_4$ 291. Each metal possesses: a) Primary valencies satisfied by anions only b) Secondary valencies satisfied by donor molecules c) Coordination number d) All of the above 292. Aspirin is: a) Antibiotic b) Antipyretic c) Sedative d) Psychedelic

| 293. Hybridisation, shape and magnetic moment of $K_3$ [Co                             | $(CO_3)_3$ ] is                                     |                         |
|--|---|-------------------------|
| a) $d^2 s p^3$ , octahedral, 4.9 BM  | b) $sp^3d^2$ , octahedral, 4.9 E                    | 3M                      |
| c) <i>dsp</i> <sup>2</sup> , square planer, 4.9 BM                                     | d) $sp^3$ , tetrahedral, 4.9BM                      |                         |
| 294. Among the following complexes ( <i>K-P</i> ),                                     |   |                         |
| $K_3[Fe(CN)_6](K)$ , $[Co(NH_3)_6]Cl_3(L)$ ,   |   |                         |
| $Na_3[Co(ox)_3](M)$  |   |                         |
| $[Ni(H_2O)_6]Cl_2(N)$ , and  |   |                         |
| $[Zn(H_2O)_6](NO_3)_2(P)$ the diamagnetic complexes                                    |   |                         |
| a) <i>K, L, M, N</i> b) <i>K, M, O, P</i>  | c) <i>L, M, O, P</i>                                | d) <i>L, M, N, O</i>    |
| 295. Aniline when diazotised in cold and then treated wit                              | h dimethyl aniline gives a c                        | coloured product. It    |
| structure would be:  |   |                         |
| a) CH <sub>3</sub> NH-N=N-NHCH <sub>3</sub>  |   |                         |
|  |   |                         |
| b) $CH_3 \longrightarrow N = N \longrightarrow NH_2$                                   |   | 21                      |
| c) $(CH_3)_2N$ $N=N$   | 5   |                         |
| d)   |   |                         |
| $(CH_3)_2N$ $NH$   | Ar.   |                         |
| 296. Pyridine possesses:   |   |                         |
| a) Aromatic nature   |   |                         |
| b) Unsaturated aliphatic nature  |   |                         |
| c) Alicyclic nature  | X   |                         |
| d) Aliphatic nature  |   |                         |
| 297. A reagent used for identifying nickel ion is:                                     |   |                         |
| a) Potassium ferrocyanide  |   |                         |
| b) Phenolphthalein   |   |                         |
| c) Dimethyl glyoxime   |   |                         |
| d) EDTA  |   |                         |
| 298. Aniline was diazotised and subsequently reduced wi                                |   |                         |
| a) Phenyl aniline b) Phenyl hydrazine  | c) <i>p</i> -amino azobenzene                       | d) Diazoamino benzene   |
| 299. The reaction of toluene with $Cl_2$ in presence of FeCl <sub>3</sub>              | gives predominantly:                                |                         |
| a) <i>m</i> -chlorobenzene   |   |                         |
| b) Benzoylchloride   |   |                         |
| c) Benzyl chloride   |   |                         |
| d) <i>o</i> - and <i>p</i> -chlorobenzene  | ) 13+ complex?                                      |                         |
| 300. Which statement is not correct in the case of [Co(NH a) It is octahedral in shape | <sub>3</sub> ) <sub>6</sub> ] <sup>*</sup> complex? |                         |
| b) It involves $d^2sp^2$ -hybridization  |   |                         |
| c) It has diamagnetic nature   |   |                         |
| d) None of the above   |   |                         |
| 301. Pick out the complex compound in which the central                                | metal atom obevs FAN rul                            | e strictly              |
| a) $K_4[Fe(CN)_6]$ b) $K_3[Fe(CN)_6]$  | c) $[Cr(H_2O)_6]Cl_3$                               | d) $[Cu(NH_3)_4]SO_4$   |
| 302. Amongst the following, the compound that can be m                                 |   | ~, [0%(1113)4]004       |
| a) Benzene b) Methoxy benzene  | c) Toluene  | d) Chlorobenzene        |
| 303. <i>p</i> -chloroaniline and anilium hydrochloride can be di                       |   | . ,                     |
| a) $P_2O_5$ b) AgNO <sub>3</sub>   | c) Carbylamine test                                 | d) Sandmeyer's reaction |
| 304. Pyrogallol is trihydroxy benzene.   |   | - V                     |
|  |   |                         |

a) 1, 2, 4 b) 1, 2, 3 c) 1, 3, 5 d) None of these

305. Phenol is weakly acidic but does not react withNaHCO<sub>3</sub> like carboxylic acids hence:

- a) Phenol is weaker than carbonic acid
- b) Phenol is stronger than acid
- c) Phenol is stronger than carboxylic acid
- d) None of the above
- 306. *p*-cresol reacts with chloroform in alkaline medium to give compound (*A*) which adds hydrogen cyanide to form compound (*B*). The latter on acidic hydrolysis gives chiral carboxylic acid. The acid is:



- b) Benzene is aromatic and has six  $\pi$ -resonating electrons
- c) Benzene has no double bond

## d) Benzene is non-polar

| d) Benzene is non-polar  |   |                                       |
|--|---|---------------------------------------|
| 316. The most stable ion is  |   |                                       |
| a) $[Fe(OH)_5]^{3-}$ b) $[FeCl_6]^{3-}$  | c) $[Fe(CN)_6]^{3-1}$   | d) $[Fe(H_2O)_6]^{3+}$                |
| 317. Which of the following is/are threo isomers?  |   |                                       |
| 317. Which of the following is/are threo isomers?<br>$a \xrightarrow{b} d a \xrightarrow{b} d d \xrightarrow{b} a$<br>$a \xrightarrow{x} d d \xrightarrow{x} a d \xrightarrow{x} a$<br>(i) (ii) (iii)<br>a) Only (i) |   |                                       |
| a - d a - d d - a  |   |                                       |
| a - d d - a d - a  |   |                                       |
|  |   |                                       |
| (i) (ii) (iii)   |   |                                       |
| a) Only (i)  | b) Only (ii)  | $\sim$                                |
| c) Only (iii)  | d) All (i), (ii) and (iii)  |                                       |
| 318. In the coal-tar distillation of middle oil, the aroma   | tic compounds present ar  | e:                                    |
| a) Benzene, naphthalene, anthracene  |   |                                       |
| b) Naphthalene, pyridine, phenol   |   |                                       |
| c) Naphthalene, pyridine   |   |                                       |
| d) None of the above   |   |                                       |
| 319. The correct order of increasing reactivity of $C - X$   | bond towards nucleophili  | c in the following compound           |
| is:  | · · · · · · · · · · · · · · · · · · ·                                     |                                       |
| XX   |   |                                       |
| $\downarrow$ $\downarrow$ $_{NO_2}$  |   | *                                     |
| (CU) C V (CH) CH-V   |   |                                       |
| $(\bigcirc), (\bigcirc), (CH_3)_3C - X, (CH_3)_2CH - X$<br>III IV  |   |                                       |
|  |   |                                       |
| I I NO <sub>2</sub>  |   |                                       |
| ПŽ   |   |                                       |
| a) I <ii<iv<iii b)="" ii<iii<iv<="" td=""><td>c) IV<iii<i<ii< td=""><td>d) III<ii<iv< td=""></ii<iv<></td></iii<i<ii<></td></ii<iv<iii>  | c) IV <iii<i<ii< td=""><td>d) III<ii<iv< td=""></ii<iv<></td></iii<i<ii<> | d) III <ii<iv< td=""></ii<iv<>        |
| 320. Which of the following system is most stable for a  | chelate?  |                                       |
| a) Two fused cyclic system   | b) Three fused cyclic   | system                                |
| c) Four fused cyclic system  | d) Five fused cyclic sys  |                                       |
| 321. Which of the following reaction take place when a   | mixture of concentrated l   | $HNO_3$ and $H_2SO_4$ reacts on       |
| benzene at 300 K?  |   |                                       |
| a) Sulphonation b) Nitration   | c) Hydrogenation  | d) Dehydration                        |
| 322. Consider the following reaction:  |   |                                       |
| Phenol $\xrightarrow{Zn \text{ dust}} X \xrightarrow{CH_3Cl} X \xrightarrow{Anhydrous AlCl_3} Y \xrightarrow{Alkaline KMnO_4} Z$ , the product Z is  | 5:  |                                       |
|  |   |                                       |
| a) Benzene b) Toluene  | c) Benzaldehyde   | d) Benzoic acid                       |
| 323. The shortest C—O bond order exists in:  |   |                                       |
| a) $[Mn(CO)_6]^+$ b) $[Fe(CO)_5]$  | c) $[Cr(CO)_6]$   | d) [V(CO) <sub>6</sub> ] <sup>-</sup> |
| 324. Between <i>p</i> -nitrophenol and salicyladehyde, solub   | ility in base is:   |                                       |
| a) Almost nil in both cases  |   |                                       |
| b) Higher in <i>p</i> -nitrophenol   |   |                                       |
| c) Higher for salicyladehyde   |   |                                       |
| d) Equal in nature   |   |                                       |
| 325. (+) and (-) forms of optically active compounds a   | re different in   |                                       |
| a) Boiling points b) Melting points  | c) Specific gravity   | d) Specific rotation                  |
| 326. Benzene on treatment with dry HCN and HCl in pr   | resence of anhy. AlCl <sub>3</sub> follo                                  | owed by hydrolysis forms:             |
| a) Chlorobenzene b) Benzoic acid   | c) Benzaldehyde   | d) Cyanobenzene                       |
| 327. In which of the following compounds does the cer  |   |                                       |
| a) $K_3$ Fe(CN) <sub>6</sub> b) $K_4$ Fe(CN) <sub>6</sub>  | c) $Cu(NH_3)_4SO_4$   | d) All of these                       |
| 328. Pick the correct name of $[Co(NH_3)_5Cl]Cl_2$   |   | -                                     |
| a) Chloropentammine cobalt (III) chloride  | b) Chloropentammine   | cobalt (III)                          |

a) Chloropentammine cobalt (III) chlorideb) Chloropentammine cobalt (III)c) Chloropentammine cobalt (II) chlorided) Pentammine chloro cobalt(III) chloride

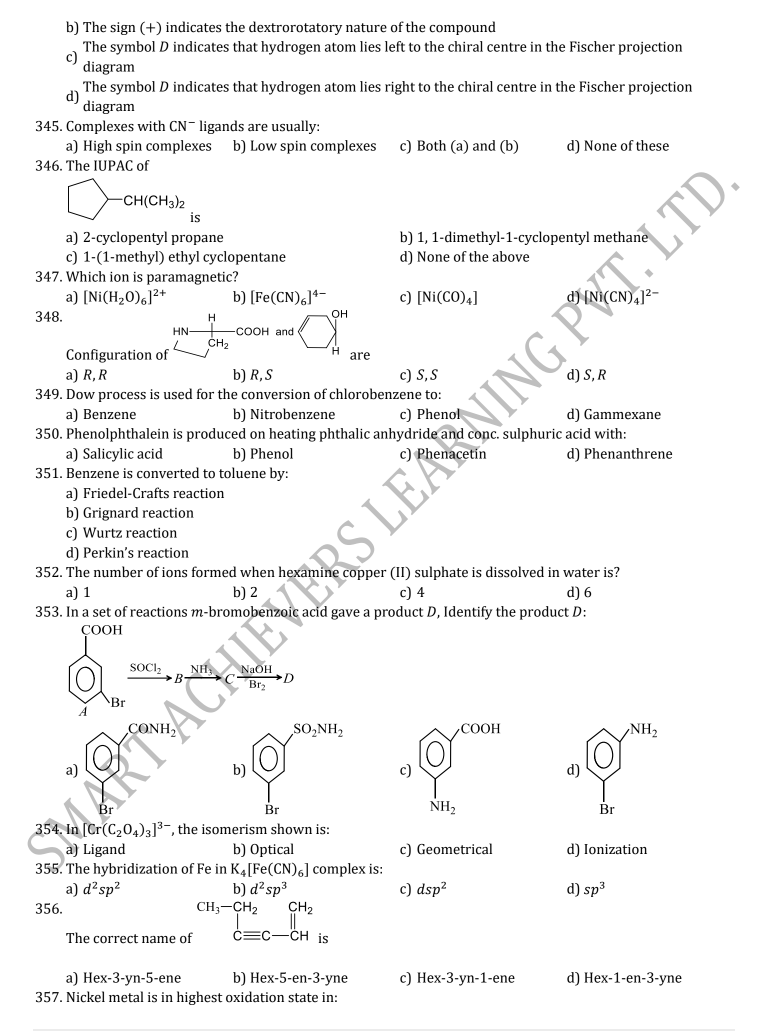
329. The geometry of  $Ni(CO)_4$  and  $Ni(PPh_3)_2Cl_2$  are

b) Both tetrahedral

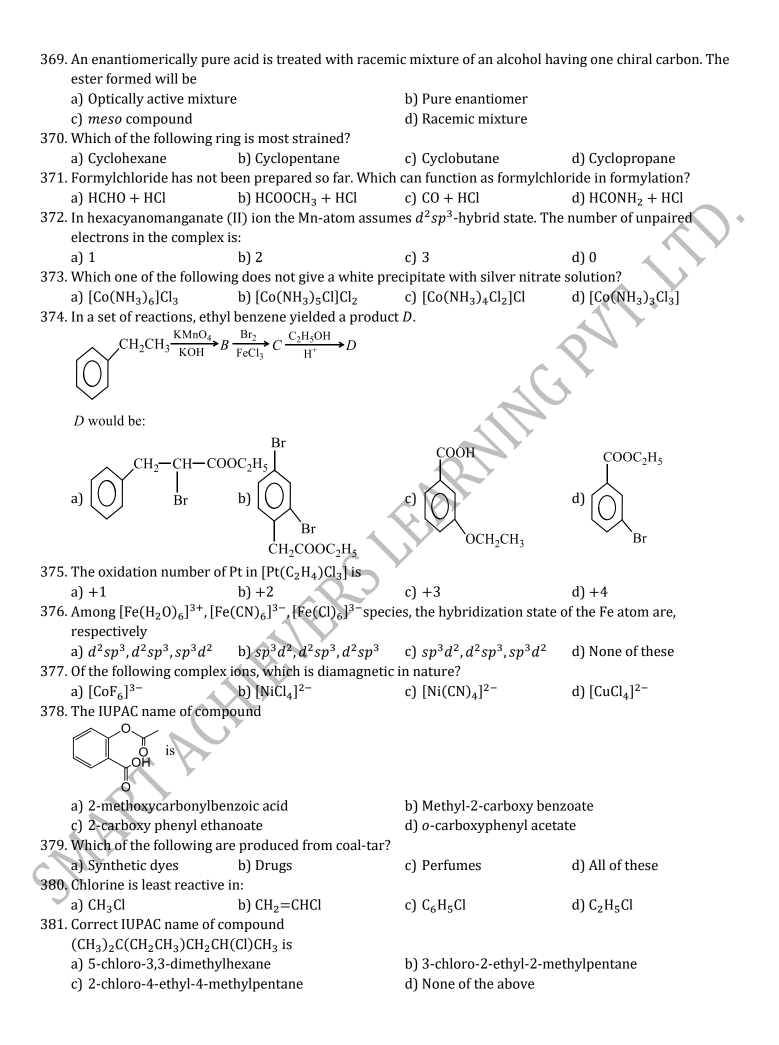
d) Both square planar

330. Select pair of chain isomers from the following

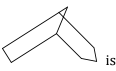
(I)(III) (III) b) II and III c) I and IV d) II and III a) I and II 331. Which ligand produces a high crystal field splitting (a strong ligand field)? c) CN<sup>-</sup> d) All are corre a) CO b)  $NO_{2}^{-}$ 332. Benzene reacts with *n*-propyl chloride in the presence of anhydrous  $AlCl_3$  to give predominantly: a) Isopropyl benzene b) No reaction c) *n*-propylbenzene d) 3-propyl-1-chlorobenzene 333. Which of the following coordination compounds would exhibit optical isomerism? a) Pentaamminenitrocobalt (III) iodide b) Diamminedinitroplatinum (II) d) Tris-(ethylenediamine) cobalt(III) bromide c) *trans*-dicyanobis (ethylenediamine) 334. What is the magnetic moment of  $K_3$ [FeF<sub>6</sub>]? a) 3.87 BM b) 4.89 BM d) 6.92 BM c) 5.91 BM 335. The EAN of Cr in  $[Cr(SCN)_6]^{3-}$  is: b) 33 a) 35 d) 37 336. Which has maximum paramagnetic character? a)  $[Fe(CN)_6]^{4-}$ b)  $[Cu(H_2O)_4]^{2+}$ c)  $[Cu(NH_3)_4]^{2+}$ d)  $[Mn(H_20)_6]^{2+}$ 337. Phenol, when it first reacts with concentrated sulphuric acid and then with concentrated nitric acid, gives: a) Nitrobenzene b) 2, 4, 6-trinitrobenzene c) o-nitrophenol d) p-nitrophenol 338. Activation of benzene ring by  $-NH_2$  in aniline can be reduced by treating with: b) Ethyl alcohol c) Acetic acid d) Acetyl chloride a) Dil. HCl 339. Sulphonation of benzoic acid produces mainly: a) o-sulphobenzoic acid b) *m*-sulphobenzoic acid c) p-sulphobenzoic acid d) o-p-disulphobenzoic acid 340. The IUPAC name for the complex  $[Co(NO_2)(NH_3)_5]Cl_2$  is a) Nitrito -N- pentamminecobalt (III) chloride b) Nitrito -N- pentamminecobalt (II) chloride c) Pentammine nitrito-N- cobalt (II) chloride d) Pentaammine nitrito-N- cobalt (III) chloride 341. The ionisation isomer of  $[Cr(H_2O)_4Cl(NO_2)C]$  is  $\sim$ a) [Cr(H<sub>2</sub>O)<sub>4</sub>(O<sub>2</sub>N)]Cl<sub>2</sub> b)  $[Cr(H_2O)_4Cl_2](NO_2)$ c)  $[Cr(H_2O)_4Cl(ONO)]$  Cl d)  $[Cr(H_2O)_4Cl_2(NO_2)]$ .  $H_2O$ 342. Salicylic acid, aspirin, nylon, plastics and picric acid have a common raw material, namely: a) Methane b) Formic acid c) Phenol d) Alcohol 343. Ulmann's reaction is used for the preparation of: b) Iodobenzene d) Naphthalene a) Diphenyl c) Toluene 344. Which of the following statements is/are incorrect for D - (+) –glyceradehyde? a) The symbol D not indicates the dextrorotatory nature of the compound



a) Ni(CO)<sub>4</sub> b)  $K_2 NiF_6$ c)  $[Ni(NH_3)_6](BF_4)_2$ d)  $K_4[Ni(CN)_6]$ 358. Which of the following complexes show six coordination number? d)  $[Cr(H_20)_6]^{3+}$ a)  $[Zn(CN)_4]^{2-}$ b)  $[Ni(NH_3)_4]^{2+}$ c)  $[Cu(CN)_4]^{2-}$ 359. Which of the following statements is wrong? a) The IUPAC name of alkenes ends with suffix-ene b) The IUPAC name of alkynes ends with suffix-yne c) The IUPAC name of acid amide is alkanamide d) The substituents get lower number in comparison to principal functional group 360. The possible number of isomers for the complex  $[MCl_2Br_2]SO_4$  is: a) 1 c) 4 d) 5 361.  $K_3[(Al)(C_2O_4)_3]$  is called b) Potassium alumino oxalate a) Potassium aliminium (III) oxalate c) Potassium trioxalato aluminate (VI) d) Potassium trioxalato aluminate (III) 362. In Fe(CO)<sub>5</sub>, the Fe — C bond possesses b) Both  $\sigma$  and  $\pi$  –characters a)  $\pi$  – Character only c) Ionic characters d)  $\sigma$  – Character only 363. The reaction,  $[Fe(CNS)_6]^{3-} \rightarrow [FeF_6]^{3-}$  taken place with a) Decrease in magnetic moment b) Increase in magnetic moment d) Increase in coordination number c) Decrease in coordination number 364. Which chloro derivative of benzene among the following would undergo hydrolysis most readily with aqueous NaOH to furnish the corresponding hydroxyl derivative? d)  $C_6H_5Cl$ 365. Some salts although containing two different metallic elements give test for only one of them in solution. Such salts are: b) Double salts a) Complex salts c) Normal salts d) None of these 366. Mixture X = 0.02 mole of  $[Co(NH_3)_5SO_4]Br$  and 0.02 mole of  $[Co(NH_3)_5Br]SO_4$  was prepared in 2 litre of solution. 1 litre of mixture X + excess AgNO<sub>3</sub>  $\rightarrow$  Y. 1 litre of mixture X + excess BaCl<sub>2</sub>  $\rightarrow$  Z. No. of moles of *Y* and *Z* are. a) 0.01, 0.01 b) 0.02, 0.01 d) 0.02, 0.02 c) 0.01, 0.02 367. The hybridization of central metal ion and shape of Wilkinson's catalyst is a)  $sp^3d$ , trigonal bipyramidal b) *sp*<sup>3</sup>,tetrahedral c)  $dsp^2$ , squre planar d)  $d^2sp^2$ , octahedral 368. The *d*-electron configurations of  $Cr^{2+}$ ,  $Mn^{2+}$ ,  $Fe^{2+}$  and  $Co^{2+}$  are  $d^4$ ,  $d^5$ ,  $d^6$  and  $d^7$  respectively. Which one of the following will exhibit minimum paramagnetic behaviour? a)  $[Cr(H_2O)_6]^{2+}$ b)  $[Mn(H_20)_6]^{2+}$ c)  $[Fe(H_2O)_6]^{2+}$ d)  $[Co(H_2O)_6]^{2+}$ (At. Nos. Cr = 24, Mn = 25, Fe = 26, Co = 27)



 $NO_2$ 382. Sn + HCl In the above reaction 'X' stands for: a)  $NH_2$ b) Cl c)  $SnCl_2$ d)  $^+_{NH_3Cl}$ 383. Which follows EAN rule? a)  $Fe(CO)_5$ b) Ni(CO)<sub>4</sub> c)  $K_4[Fe(CN)_6]$ d) All are correct 384. Which one is bidentate ligand? a)  $C_2 O_4^{2-}$ b)  $NH_2 \cdot CH_2 \cdot CH_2 \cdot NH_2$  c) Both (a) and (b) d) None of these 385. The reagent used for conversion of benzene diazonium chloride to benzene is: d) All of these a)  $H_3PO_2 + H_2O$ b)  $Na_2SnO_2 + NaOH$ c)  $C_2H_5OH$ 386. Which will not give the usual test for iron? a)  $K_2Fe_2(SO_4)_4 \cdot 24H_2O$ b)  $(NH_4)_2 Fe(SO_4)_2 \cdot 6H_2O$ c)  $K_3[Fe(CN)_6]$ d)  $Fe_2(SO_4)_3$ 387.  $[Co(NH_3)_5SO_4]Br$  and  $[Co(NH_3)_5 Br]SO_4$  are a pair of ..... isomers. a) Ionisation b) Ligand c) Coordination d) Hydrate 388. The first organic compound prepared in the laboratory was a) Acetic acid b) Acetylene c) Urea d) Methane 389. Aniline on heating with conc.  $H_2SO_4$  at 460 K gives: a) Aniline sulphate b) Benzene sulphonic acid c) Sulphanilic acid d) None of the above 390. Which of the following statements regarding phenols is not correct? a) Phenols are stronger acid than water and alcohols b) Phenols are weaker acids than carboxylic acids c) Phenols are soluble in both aqueous NaOH and aqueous NaHCO<sub>3</sub> d) Phenoxide ions are more stable than the corresponding phenols 391. Which would decolourise cold, aq. potassium permanganate solution? a) Benzoic acid b) Cinnamic acid c) p-toluic acid d) *m*-toluic acid 392. The magnetic moment of  $K_3$  [Fe(CN)<sub>6</sub>] is found to be 1.7 BM. How many unpaired electron (s) is/are present per molecule? a) 1 b) 2 c) 3 d) 4 393. The IUPAC name of the compound CH<sub>2</sub>-CH-CH<sub>2</sub>-CH<sub>2</sub> ĊΗ2 is ĊH<sub>2</sub>—ĊH—CH<sub>2</sub>—ĊH<sub>2</sub> a) Bicyclo [2,5,0] nonane b) Bicyclo [5,0,2] nonane c) Bicyclo [5,2,0] nonane d) Bicyclo [0,2,5] nonane 394. The IUPAC name of the compound COOH is a) 2-oxocyclohexane-1-carboxylic acid b) Cyclohexane-2-oxo-1-carboxylic acid c) 6-oxocyclohexane-1-carboxylic acid d) None of the above 395. The IUPAC name of



NH<sub>2</sub>

a) Spiro [3.2.1] octane b) Bicyclo [3.2.2] octane c) Bicyclo [3.2.1] octane d) None of these 396. Which of the following deactivates benzene substitution?

c) Chlorobenzene

a) -NHRb) - 0H c) - 0R d) -COOR

d) Phenol

397. Aniline, chloroform and alc. KOH on heating give:

a) Phenyl isocyanide b) Phenyl cyanide 398. In the chemical reactions,

$$\underbrace{\bigcirc}_{\text{HCl},278\text{K}}^{\text{NaNO}_2} A \xrightarrow{\text{HBF}_4} B$$

the compounds ''A'' and ''B'' respectively are :

a) Nitrobenzene and chlorobenzene

b) Nitrobenzene and fluorobenzene

c) Phenol and benzene

d) Benzenediazonium chloride and fluorobenzene

399. The incorrect statement for IUPAC system of nomenclature is

a) In an organic compound, the longest carbon chain is always selected for assigning the root word

b) There is no compound with the name 3-ethyl pentane

- c) Out of NH<sub>2</sub> and OH groups present in an organic compound, NH<sub>2</sub> is treated as substituent
- d) Different alkyl groups are written alphabetically while, writing the IUPAC name
- 400. When sodium benzene sulphonate is fused with sodium hydroxide (solid), followed by hydrolysis the product formed is:

b) Sod. phenoxide a) Benzene c) Benzene thiophenol d) Phenol 401. The correct order of stability of conformations of cyclohexane is

b) Twist boat > chair > boat

a) Chair > twist boat > boat

c) Boat > chair > twist boat 402. Phenol with dilute HNO<sub>3</sub> gives:

a) meta and para nitrophenol

b) ortho and para nitrophenol

c) Trinitrophenol

d) ortho and meta nitrophenol

403. The increasing order of boiling points of compounds given below is:

- (I) 1,2-dihydroxy benzene
- (II) 1,3-dihydroxy benzene

(III) 1,4-dihydroxy benzene

(IV) Hydroxyl benzene

```
a) I < II < III < IV
                                b) I < II < IV < III
                                                            c) IV < I < II < III
                                                                                        d) IV < II < I < III
404. The pair of the compounds in which both the metals are in the higher possible oxidation state is
```

b)  $[Co(CN)_6]^{3-}$ , MnO<sub>3</sub> a)  $CrO_2Cl_2$ ,  $MnO_4^-$ 

c)  $TiO_3$ ,  $MnO_2$ 

405. The number of ions given by  $K_2[PtCl_6]$  in aqueous solution is: a) 2 b) 3 c) 4

406. Which of the following are functional isomers? a) CH<sub>3</sub>CH<sub>2</sub>Cl and CH<sub>3</sub>CH<sub>2</sub>Br

c) C<sub>2</sub>H<sub>5</sub>OC<sub>2</sub>H<sub>5</sub> and CH<sub>3</sub>OC<sub>3</sub>H<sub>7</sub>

b)  $CH_3CHBr_2$  and  $CH_2Br_2 \cdot CH_2Br$  $CH_3CH_2CHO \text{ and } CH_3-d$ 

d) Zero

d)  $[Fe(CN)_6]^{3-}$ ,  $[Co(CN)_6]^{3-}$ 

d) Boat > twist boat > chair

407. Phenol is:

c) Strongly basic

d) Weakly basic

a) Strongly acidic b) Weakly acidic 408. The correct IUPAC name of  $KAl(SO_4)_2 \cdot 12H_2O$  is:

- a) Aluminium potassium sulphate-12-water
- b) Potassium aluminium(III) sulphate-12-water
- c) Potassium aluminate(III) sulphatehydrate
- d) Aluminium(III) potassium sulphate hydrate-12
- 409. A complex shown below can exhibit:

- a) Optical isomerism only
- b) Geometrical isomerism only
- c) Both optical and geometrical isomerism
- d) None of the above
- 410. The IUPAC name of the complex  $[Co(NH_3)_4Cl_2]Cl$  is
  - a) Dichloro tetraammine cobalt (III) chloride
  - c) Tetraammine dichloro cobalt (II) chloride
- b) Tetraammine dichloro cobalt(III) chloride
- d) Tetraammine dichloro cobalt (IV) chloride
- 411. The correct decreasing order of their reactivity towards hydrolysis is:
  - (i)  $C_6H_5COCl$

(iv) OHC

| (ii) $O_2N$ —COCl   |  |
|---------------------|--|
| (iii) $H_{2}C$ COCl |  |

a) (i)>(ii)>(iii)>(iv)

c) (ii)>(iv)>(i)>(iii) d) (ii)>(iv)>(iii)>(i)

d) All of these

412. Nitrobenzene is generally used for:a) Preparing shoe polish b) Preparing floor polish c) Preparing aniline

COCI

413. In the coordination compound,  $K_4[Ni(CN)_4]$ , the oxidation state of nickel is

b) (iv)>(ii)>(i)>(iii)

| COM                               | pound, 14[111(011)4], the t                                  | Shutter of mer   | CI 15  |   |  |
|-----------------------------------|--|--|--|---|--|
|                                   | b) 0   | c) +1  |  | d) +2   |  |
| npare                             | ed to benzoic acid:  |  |  |   |  |
|                                   | b) Has same acidity  | c) Has less acidity  | /  | d) None of these  |  |
| oected                            | d to be bidentate?   |  |  |   |  |
|                                   | b) $CH_3C \equiv N$  | c) Br <sup>-</sup>   |  | d) CH <sub>3</sub> NH <sub>2</sub>  |  |
| ollowi                            | ing is most reactive towar                                   | ds aqueous NaOH?   |  |   |  |
|                                   | b) C <sub>6</sub> H <sub>5</sub> CH <sub>2</sub> Cl          | c) C <sub>6</sub> H <sub>5</sub> Br  |  | d) BrC <sub>6</sub> H <sub>4</sub> Br   |  |
| mati                              | c compound?  |  |  |   |  |
|                                   | b) Naphthalene   | c) Xylene  |  | d) Cyclohexane  |  |
| ollowi                            | ing is wrongly matched?                                      |  |  |   |  |
| —                                 | Square planar  | b) [Ni(CO) <sub>4</sub> ]  | —  | Neutral ligand  |  |
| —                                 | $sp^3 d^2$   | d) [Co(en) <sub>3</sub> ] <sup>3+</sup>  | —  | Follows EAN rule  |  |
| 419. Stereoisomers have different |  |  |  |   |  |
| ıla                               |  | b) Structural form   | nula   |   |  |
|                                   | npare<br>oected<br>ollowi<br>omatio<br>ollowi<br><br>e diffe | b) 0<br>mpared to benzoic acid:<br>b) Has same acidity<br>bected to be bidentate?<br>b) $CH_3C \equiv N$<br>b) $C_6H_5CH_2Cl$<br>b) $C_6H_5CH_2Cl$<br>b) Naphthalene<br>b) Naphthalene<br>b) Naphthalene<br>c) Naphthalene<br>b) Naphthalene<br>c) Naphtha | b) 0 c) +1<br>mpared to benzoic acid:<br>b) Has same acidity c) Has less acidity<br>b) CH <sub>3</sub> C $\equiv$ N c) Br <sup>-</sup><br>b) CH <sub>3</sub> C $\equiv$ N c) Br <sup>-</sup><br>b) CH <sub>3</sub> C $\equiv$ N c) Br <sup>-</sup><br>b) C <sub>6</sub> H <sub>5</sub> CH <sub>2</sub> Cl c) C <sub>6</sub> H <sub>5</sub> Br<br>b) C <sub>6</sub> H <sub>5</sub> CH <sub>2</sub> Cl c) C <sub>6</sub> H <sub>5</sub> Br<br>matic compound?<br>b) Naphthalene c) Xylene<br>blowing is wrongly matched?<br>- Square planar b) [Ni(CO) <sub>4</sub> ]<br>- sp <sup>3</sup> d <sup>2</sup> d) [Co(en) <sub>3</sub> ] <sup>3+</sup><br>e different | inpared to benzoic acid:b) Has same acidityc) Has less acidityb) Has same acidityc) Has less acidityb) CH_3C ≡ Nc) Br <sup>-</sup> ollowing is most reactive towards aqueous NaOH?b) C_6H_5CH_2Clc) C_6H_5Bromatic compound?b) Naphthalenec) Xyleneollowing is wrongly matched?-Square planar-sp <sup>3</sup> d <sup>2</sup> -d) [Co(en)_3]^{3+} -e different |  |

d) Molecular mass

c) Configuration

420. Which of the following will show optical isomerism?

a)  $[Cu(NH_3)_4]^{2+}$ 

- b)  $[ZnCl_4]^{2-}$
- c)  $[Cr(C_2O_4)_3]^{3-1}$
- d)  $[Co(CN)_6]^{3-1}$

421. A complex of cobalt has five ammonia molecules, one nitro group and two chlorine atoms for each cobalt atom. One mole of this compound produces three mole ions in aqueous solution which on treating with excess of AgNO<sub>3</sub> give two mole of AgCl. The formula of the compound is: d)  $[Co(NH_3)_5][(NO_2)_2Cl_2]$ 

a)  $[Co(NH_3)_4NO_2Cl][(NH_3b) [Co(NH_3)_5Cl][ClNO_2] c) [Co(NH_3)_5NO_2]Cl_2$ 422. Which one group is trivalent in nature? a) Benzo b) Benzal

c) Benzyl

423. Benzene contains double bonds but does not give addition reactions because:

- a) Double bonds in benzene are strong
- b) Double bonds change their position rapidly
- c) Resonance lowers the energy of benzene molecule and leads to greater stabilization
- d) None of the above

424. Low spin complex of  $d^6$ -cation in an octahedral field will have the following energy:

a) 
$$\frac{-12}{5}\Delta_0 + P$$
  
b) 
$$\frac{-12}{5}\Delta_0 + 3P$$
  
c) 
$$\frac{-2}{5}\Delta_0 + 2P$$
  
d) 
$$\frac{-2}{5}\Delta_0 + P$$

 $(\Delta_0 = \text{Crystal field splitting energy in an octahedral field, } P = \text{Electron pairing energy})$ 425. C<sub>7</sub>H<sub>8</sub>O show how many isomers?

426. CCl<sub>3</sub>

The above structural formula refers to:

H

a) 2

c) DDT

c) 4

d) RNA

d) 5

d) All of these

Ö Have its IUPAC name as a) Octa dec-9-enoic acid

b) Oleic acid c) Ethyl hexadic-9-enoic acid d) All of these 428. The type of isomerism present in nitropentaammine-chromium (III) chloride is :

Ja) Optical b) Linkage c) Ionization 429. Which complex compound possesses  $sp^3d^2$  hybridisation?

b) 3

b) DNA

d) polymerization

c)  $[Fe(CN)_6]^{3-}$ a)  $[Fe(NH_3)_6]^{3+}$ b)  $[Fe(CN)_6]^{4-}$ d)  $[Fe(Cl)_6]^{3-1}$ 430. Amongst the following carboxylic acids the strongest acid is:

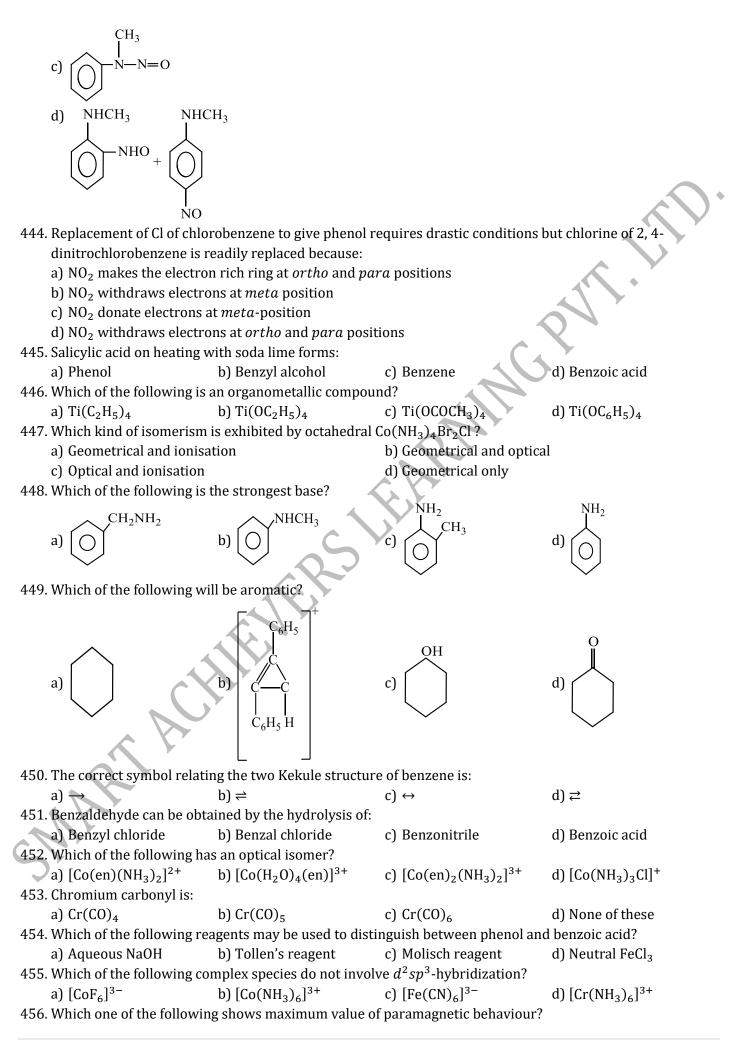
a) Benzoic acid

b) *o*-methoxybenzoic acid

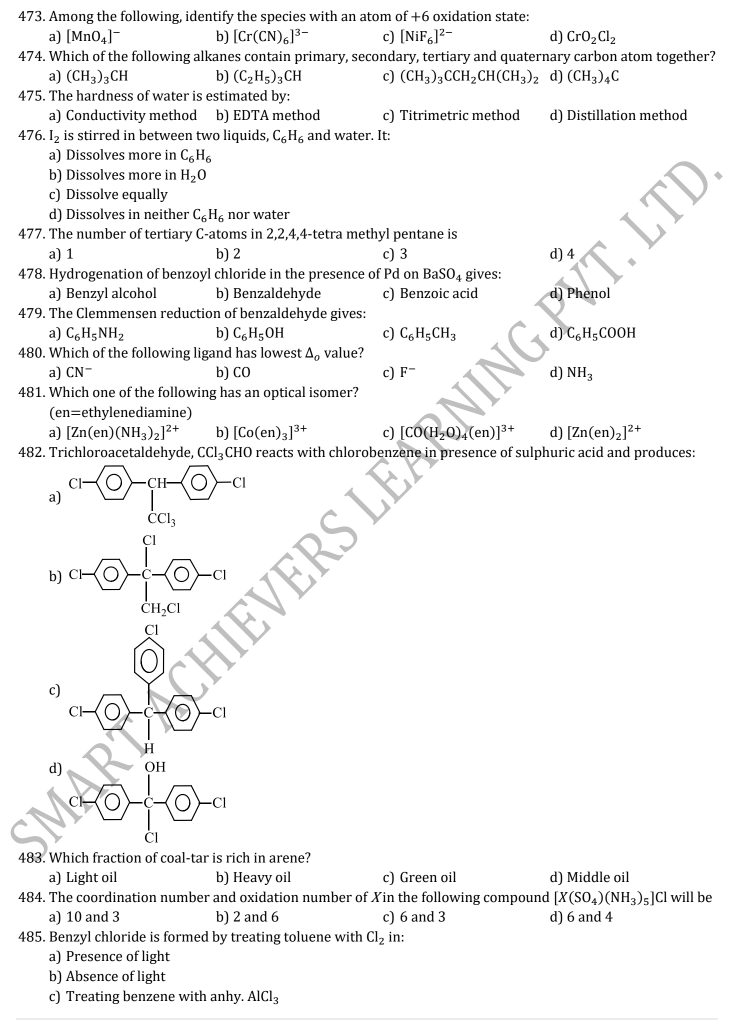
c) *m*-nitrobenzoic acid

d) p-nitrobenzoic acid

431. When EDTA solution is added to Mg<sup>2+</sup> ion solution, then which of the following statements is not true? Four coordinate sites of Mg<sup>2+</sup> are occupied by EDTA and remaining two sites are occupied by water a) molecules. b) All six coordinate sites of  $Mg^{2+}$  are occupied. c)  $P^{H}$  of the solution is decreased. d) Colourless  $[Mg - EDTA]^{2-}$  chelate is formed. 432. The energy difference between chair and the boat conformation of cyclohexane is a) 29.7 kJ b) 44 kJ c) 151 kJ d) 36 kJ 433. Compounds having the same molecular formula but different properties are called a) Isotopes b) Isobars c) Isomers d) Isomorphs 434. CH<sub>3</sub>OCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>OCH<sub>2</sub>CH<sub>3</sub> is a) Ethylmethylpropyl diether b) Ethylmethoxypropyl ether d) 1-ethoxy-3-methoxy propane c) 3-ethoxy-1-methoxy propane 435. The benzene molecule contains: a) Six *sp*<sup>2</sup>-hybridized carbons b) Three  $sp^2$ -hybridized carbons c) Six *sp*<sup>3</sup>-hybridized carbons d) Three  $sp^3$ -hybridized carbons 436. The correct order of stability of conformations of  $NH_2 - CH_2 - CH_2 - OH$  is a) Gauche > eclipsed > anti b) Gauche > anti > eclipsed c) Eclipsed > gauche > anti d) Anti > eclipsed > gauche 437. The solubility of AgCN increases by the addition of KCN because of: a) Complex formation b) Redox change c) Salt formation d) None of these 438. Alicyclic compounds are b) Aliphatic cyclic compounds a) Aromatic cyclic compounds c) Both (a) and (b) d) None of the above 439. Which of the following compounds reacts slower than benzene in electrophilic bromination? a)  $C_6H_5 - NO_2$ b)  $C_6H_5 - NH_2$ c)  $C_6H_5 - OH$ d)  $C_6H_6 - CH_3$ 440. The fraction of chlorine precipitated by  $AgNO_3$  solution from  $[Cu(NH_3)_5Cl]Cl_2$  is: a) 1/2 b) 2/3 c) 1/3 d) 1/4441. Number of possible optical isomers in  $[Co(en)_2Cl_2]^+$  is b) 3 a) 2 c) 4 d) 6 442. Dimethyl glyoxime gives a red precipitate with Ni<sup>2+</sup> which is used for its detection. To get this precipitate readily, the best pH range is a) < 1 b) 3 – 4 c) 9 – 11 d) 2 – 3 443. Predict the product:  $NHCH_3$ — $NaNO_2$ +HCl  $\rightarrow$  ProductOH CH<sub>3</sub> b)



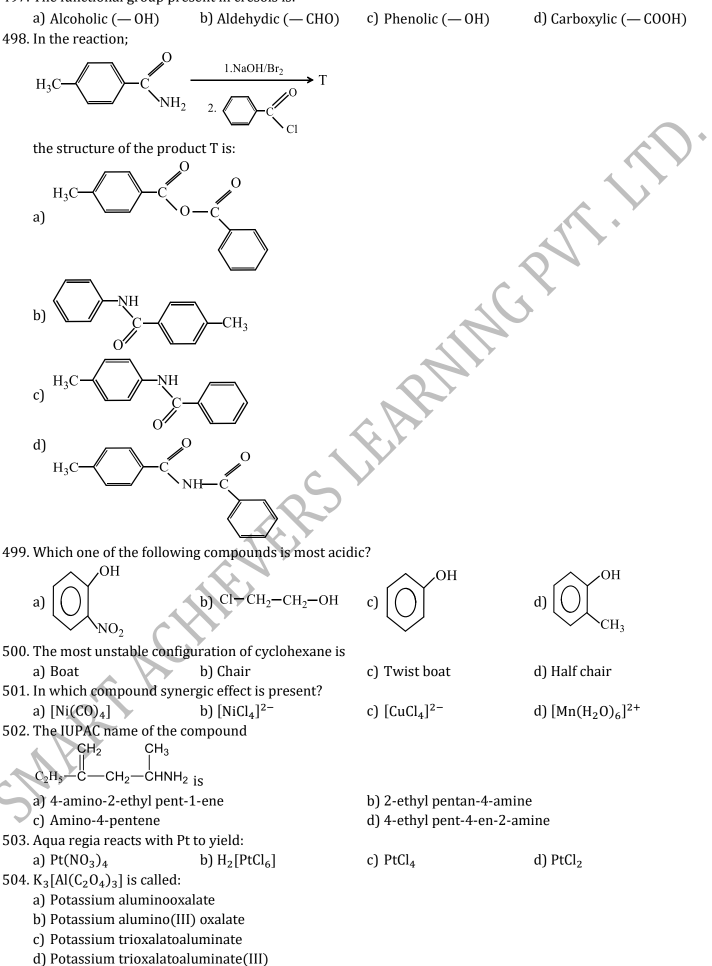
| a) [Sc(CN) <sub>6</sub> ] <sup>3-</sup>                                   | b) [Co(CN) <sub>6</sub> ] <sup>3–</sup>   | c) $[Ni(CN)_4]^{2-}$                        | d) $[Cr(CN)_6]^{3-}$                     |
|---|---|---|--|
| 457. The IUPAC name of<br>HOOC – CH <sub>2</sub> – CH <sub>2</sub> –      | $CH_2 - CH - CH_2 - COOH$   |   |  |
|   |   |   |  |
|   | CH <sub>2</sub> COOH  |   |  |
| is  |   |   |  |
| <ul><li>a) 3-(carboxymethyl)</li><li>b) 5-(carboxymethyl)</li></ul>       | •   |   |  |
|   | pentane dicarboxylic acid   |   | $\sim$                                   |
| d) 4-(carboxymethyl)  | pentane dicarboxylic acid   |   | $\langle \cdot \rangle$                  |
| 458. Which of the following   | g species will be diamagneti  |   |  |
| a) [Fe(CN) <sub>6</sub> ] <sup>4–</sup>                                   | b) [FeF <sub>6</sub> ] <sup>3–</sup>  | c) $[Co(C_2O_4)_3]^{3-1}$                   | d) [CoF <sub>6</sub> ] <sup>3–</sup>     |
|   | wing is an outer orbital com  |   |  |
| a) $[Cr(NH_3)_6]^{3+}$  | b) $[Co(NH_3)_6]^{3+}$  | c) $[Ni(NH_3)_6]^{2+}$                      | d) $[Zn(NH_3)_6]^{2+}$                   |
| 460. Moth balls contain:  |   |   |  |
| a) Camphor  | b) Benzoic acid   | c) Naphthalene                              | d) Cinnamic acid                         |
| a) Oxidation number   | tate ligands in the complex i   | b) Primary valency                          | <b>*</b>                                 |
| c) Coordination numb  | or  | d) EAN                                      |  |
|   | lle, the number of π-electror   |   |  |
| a) 12   | b) 14   | c) 10                                       | d) 20                                    |
| •   | ane which one of the follow   |   | -  |
| a) Eclipsed and chair o   |   | b) Staggered and chair                      |  |
| c) Staggered and boat   |   | d) Eclipsed and boat co                     |  |
| 464. Among the following v  | which is not $\pi$ -bonded organ  | ometallic compound?                         |  |
| a) K[PtCl <sub>3</sub> ( $\eta^2 - C_2H_4$ )                              | ] b) Fe(η <sup>5</sup> – C <sub>5</sub> H <sub>5</sub> ) <sub>2</sub>           | c) $Cr(\eta^6 - C_6H_6)_2$                  | d) $(CH_3)_4Sn$                          |
| 465. <i>o</i> , <i>p</i> -directing groups a                              | 9   | *   |  |
| a) Activating groups  | b) Deactivating groups  | c) Neutral groups                           | d) None of these                         |
| 466. Aryl halides are less re   | active towards nucleophilic   | substitution reaction as co                 | ompared to alkyl due halides             |
| to:   |   |   |  |
| b) Resonance stabiliza  | s stable carbonium ion  |   |  |
| c) Longer carbon-halo   |   |   |  |
| d) The inductive effect   |   |   |  |
| 467. Which would be least   |   |   |  |
| a) Nitrobenze   | b) Anisole  | c) Phenol                                   | d) Chlorobenzene                         |
| 468. Which has a smell of o   | ,   |   |  |
| a) Benzaldehyde   | b) Benzoic acid   | c) Ethyl salicylate                         | d) Methyl salicylate                     |
|   | ber of Pt in [Pt(NH <sub>3</sub> ) <sub>4</sub> Cl <sub>2</sub> ] <sup>2+</sup> |   |  |
| a) 2  | b) 4  | c) 6  | d) 8                                     |
| 470. $C_6H_5Cl$ on treating wit   | h NaOH at 300°C gives phen  | ol. However the yield is po                 | or because of side reaction              |
| producing:  |   |   |  |
| a) C <sub>6</sub> H <sub>5</sub> Na                                       | b) C <sub>6</sub> H <sub>5</sub> OCH <sub>3</sub>                               | c) $C_6H_5OC_6H_5$                          | d) None of these                         |
| 471. In $Cr(NH_3)_4Cl_2$ ]Cl the  |   |   |  |
| a) NH <sub>3</sub> only   | b) Cl <sup>-</sup> only   | c) Both NH <sub>3</sub> and Cl <sup>-</sup> | d) Cr, NH <sub>3</sub> , Cl <sup>-</sup> |
| 472. Which statement is no  |   |   |  |
| a) It is less basic than<br>b) It can be steam dist                       |   |   |  |
| <ul><li>b) It can be steam dist</li><li>c) It reacts with sodiu</li></ul> |   |   |  |
| d) It is soluble in wate  |   |   |  |
|   |   |   |  |



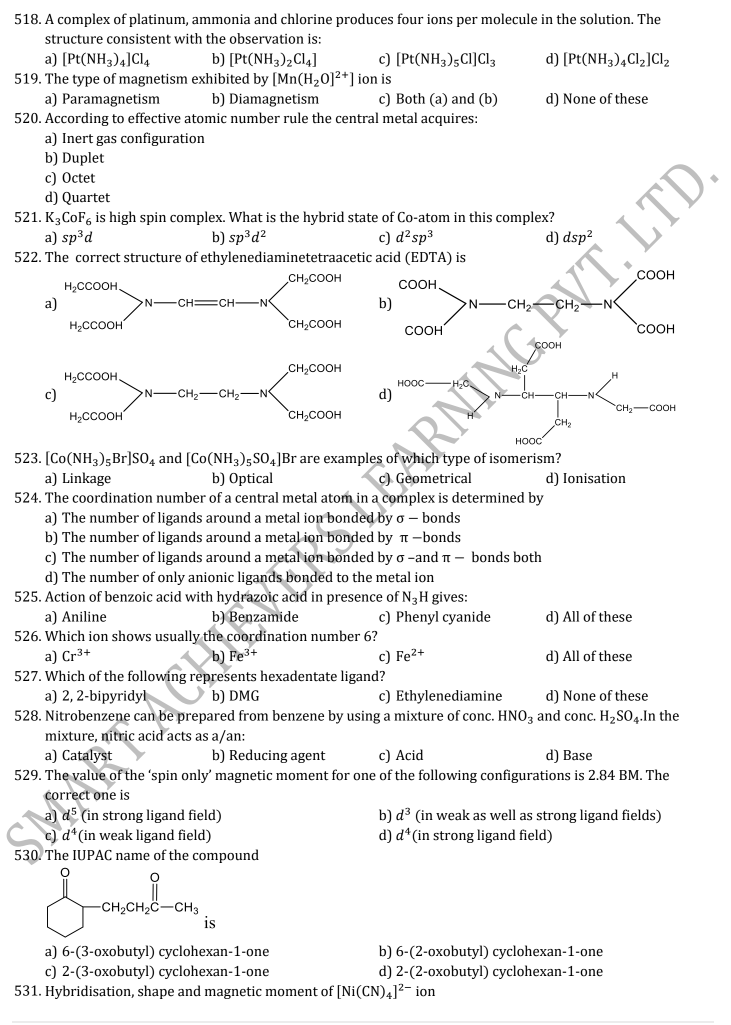
d) Treating benzene with As<sub>2</sub>S<sub>3</sub> 486. Which complex cannot ionize in solution? a)  $[CoCl_3(NH_3)_3]$ b)  $K_4(Fe(CN)_6]$ c)  $K_2[Pt(F_6)]$ d)  $[Pt(NH_3)_6]Cl_4$ 487. [Ni  $(CN)_4$ ]<sup>2–</sup>, [MnBr<sub>4</sub>]<sup>2–</sup> and [CoF<sub>6</sub>]<sup>3–</sup>, geometry, hybridisation and magnetic moment of the ions respectively, are Tetrahedral, square planar, octahedral : a) *sp*<sup>3</sup>, *dsp*<sup>2</sup>, *sp*<sup>3</sup>*d*<sup>2</sup>: 5.9, 0, 4.9 Tetrahedral, square planar, octahedral : b)  $dsp^2$ ,  $sp^3$ ,  $sp^3d^2$ : 0, 5.9, 4.9 Square planar, tetrahedral, octahedral : c)  $dsp^2$ ,  $sp^3$ ,  $d^2sp^3$ : 5.9,4.9,0 Square planar, tetrahedral, octahedral : d)  $dsp^2$ ,  $sp^3$ ,  $sp^3d^2$ : 0, 5.9, 4.9 488. Ozonolysis of benzene gives: a) 1 molecule of glyoxal b) 2 molecules of glyoxal c) 3 molecules of glyoxal d) None of these 489. In benzene, C—C bond length is 1.39 Å; the C—H bond length is: d) 1.46 b) 1.08 c) 1.54 a) 1.39 490. The IUPAC name of following compound is b) 3-N,N-dimethyl, 3-methyl pentanamine a) N,N-dimethyl, 3-methyl pentan-3-amine d) 3-methyl-3-N, N-dimethyl butane c) 3-methyl-3-N, N-dimethyl pentane 491. Which of the following may be used as food preservative? a) Benzene b) Ethylene c) Sodium benzoate d) Sodium metaaluminate 492. Which compound is formed when sodium phenoxide is heated with ethyl iodide? a) Phenetole b) Ethyl phenyl alcohol c) Phenol d) None of these 493. In metal carbonyl (organometallic) complexes, the M— C bond is a) Ionic b) Covalent with ionic character c) Covalent d) Coordinate covalent 494. Octahedral complex is a) *cis* b) *trans* c) mer d) fac 495. The correct order of magnetic moments (spin only values in BM) among the following is (Atomic no. Mn=25, Fe=26, Co=27) a)  $[MnCl_4]^{2-} > [CoCl_4]^{2-} > [Fe(CN)_6]^{4-}$ b)  $[MnCl_4]^{2-} > [Fe(CN)_6]^{4-} > [CoCl_4]^{2-}$ c)  $[Fe(CN)_6]^{4-} > [MnCl_4]^{2-} > [CoCl_4]^{2-}$ d)  $[Fe(CN)_6]^{4-} > [CoCl_4]^{2-} > [MnCl_4]^{2-}$ 496. Aniline and methyl amine can be differentiated by: a) Diazotisation followed by coupling with phenol b) Reaction with chloroform and aqueous solution of KOH c) Reaction with HNO<sub>2</sub>

d) None of the above

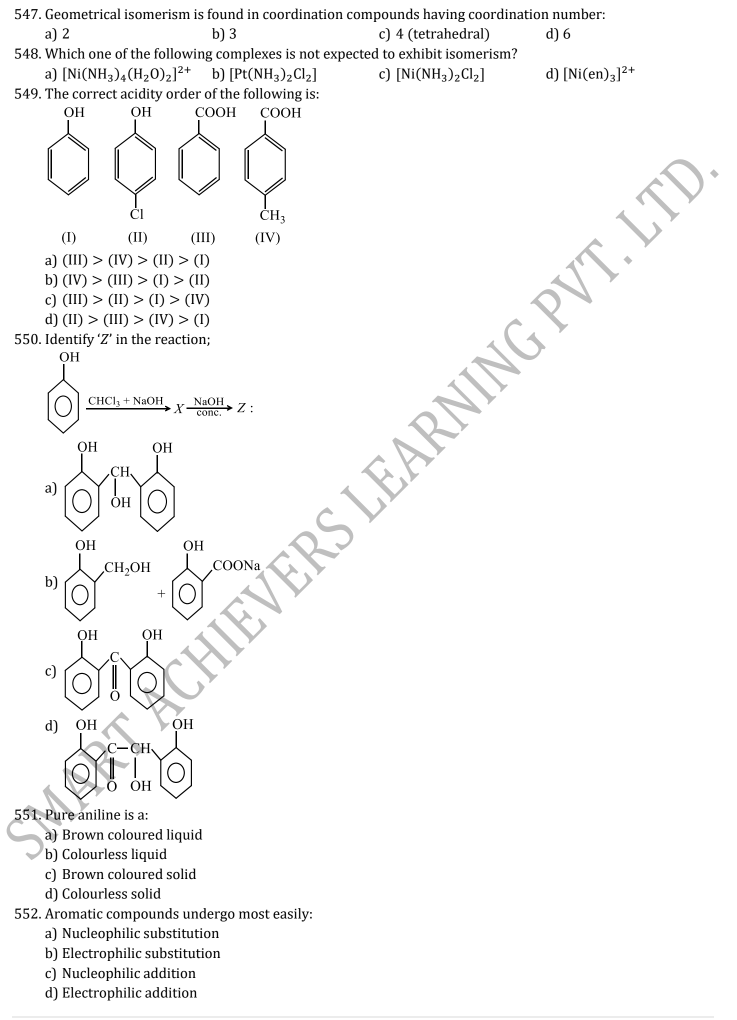
497. The functional group present in cresols is:

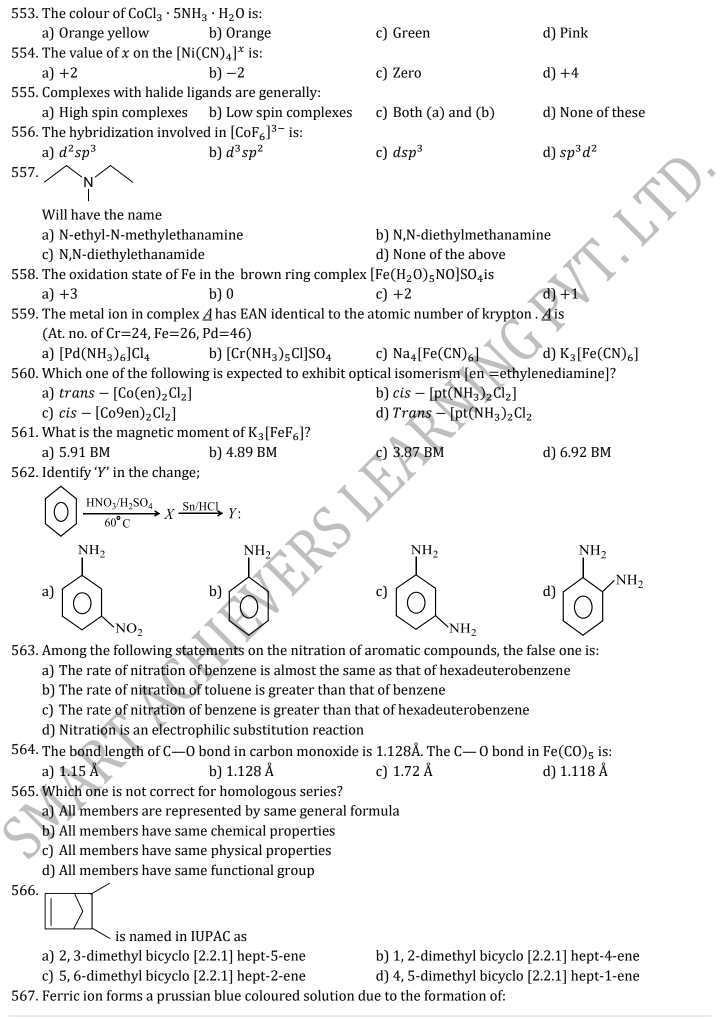


505. The IUPAC name of



| a) $dsp^2$ , square planar, zero   | b) <i>dsp</i> <sup>2</sup> , square planar, |                                |
|--|---|--------------------------------|
| c) $sp^2d^2$ , octahedral, zero  | d) $d^2 s p^3$ , octahedral, 1.             | 73                             |
| <sup>532.</sup> Choose the IUPAC name of $\Box$  |   |                                |
| a) Dicyclobutane   | b) Bicyclo [2.2.0] hexan                    | e                              |
| c) Bicyclo [2.2.1] hexane  | d) None of these                            |                                |
| 533. Which of the following is a heterocyclic compound   | ?   |                                |
| a) Phenanthrene b) Thiophene   | c) Phenol                                   | d) Aniline                     |
| 534. $[Sc(H_2O)_6]^{3+}$ ion is  |   |                                |
| a) Colourless and diamagnetic  | b) Coloured and octahe                      | dral                           |
| c) Colourless and paramagnetic   | d) Coloured and parama                      | agnetic                        |
| 535. Benzene reacts with $CH_3Cl$ in the presence of anhy  | drous AlCl <sub>3</sub> to form:            |                                |
| a) Xylene b) Toluene   | c) Chlorobenzene                            | d) Benzylchloride              |
| 536. The magnetic moment of $[Co(NH_3)_6]Cl_3$ is  |   | •                              |
| a) 1.73 b) 2.83  | c) 6.6                                      | d) Zero                        |
| 537. The correct order of reactivity towards electrophi  | lic substitution is:                        | $\sim$                         |
| a) Phenol >Benzene>Chlorobenzene>Benzoic ac  | id 🔨  | X                              |
| b) Benzoic acid>Chlorobenzene>Benzene>Phenc  | ol  | <b>A</b>                       |
| c) Phenol >Chlorobenzene>Benzene>Benzoic ac  | id  | <i>J</i>                       |
| d) Benzoic acid>Phenol>Benzene>Chlorobenzen  | e   |                                |
| 538. The product formed by the reaction of $C_6H_5CN$ and  | d CH <sub>2</sub> N <sub>2</sub> is:        |                                |
|  |   | d) None of these               |
| a) $(\bigcirc CH = CH_2N_2 \qquad N \qquad N$  | c) CH <sub>2</sub> CH <sub>2</sub>          |                                |
|  |   |                                |
| 539. Increasing order of expected enol content   |   |                                |
| a) $CH_3COCH_2CHO > CH_3COCH_3 > CH_3CHO > CH_3CHO = CH$ | СОСН-СОСН-                                  |                                |
| b) $CH_3COCH_2COCH_3 > CH_3COCH_2CHO > CH_3CO$ |   |                                |
| c) $CH_3CHO > CH_3COCH_3 > CH_3COCH_2CHO > CH_3COCH_2CHO > CH_3COCH_3 > CH_3COCH_2CHO > CH_3C$ |   |                                |
| d) $CH_3COCH_3 > CH_3COCH_2COCH_3 > CH_3CHO >$ |   |                                |
| 540. Out of the following the metal which forms polynu   |   |                                |
| a) Na b) Mg  | c) Mn                                       | d) All of these                |
| 541. Picric acid and benzoic acid can be distinguished b   | •   |                                |
| a) Aqueous NaHCO $_3$ b) Aqueous NaOH  | c) Aqueous FeCl <sub>3</sub>                | d) Aqueous $Na_2CO_3$          |
| 542. The compound having the lowest oxidation state of   |   | a) nquoouo nu <sub>2</sub> 003 |
| a) $K_4$ Fe(CN) <sub>6</sub> b) $K_2$ FeO <sub>4</sub>   | c) $Fe_2O_3$                                | d) Fe(CO) <sub>5</sub>         |
| 543. The name of $[Pt(NH_3)_4Cl_2]^{2+}$ , $[PtCl_4]^{2-}$ is  | 0) 20203                                    |                                |
| a) Tetramminedichloroplatinum(IV) tetrachlorop   | latinate(II)                                |                                |
| b) Dichloroplatinum (IV) tetrachloroplatinate  |   |                                |
| c) Tetrachloroplatinum (II) tetrammineplatinate  |   |                                |
| d) Tetrachloroplatinum (II) dichlorotetraammine  | platinate                                   |                                |
| 544. <i>m</i> -dihydroxybenzene is also called:  | 1   |                                |
| a) Catechol b) Resorcinol  | c) Quinol                                   | d) Pyrogallol                  |
| 545. The ion which exhibits green colour   |   |                                |
| a) $Cu^{2+}$ b) $Mn^{2+}$  | c) Co <sup>2+</sup>                         | d) Ni <sup>2+</sup>            |
| 546. $X \xrightarrow{\text{Cl}_2}$ Benzotrichloride $\xrightarrow{\text{Hydrolysis}} Y$  | -   | -                              |
|  |   |                                |
| X and Y respectively are:  |   |                                |
| a) Benzene, benzaldehyde   |   |                                |
| b) Toluene, benzaldehyde   |   |                                |
| c) Toluene, benzoic acid<br>d) Benzene, benzoic acid   |   |                                |
| uj Denzene, Denzoit atiu   |   |                                |

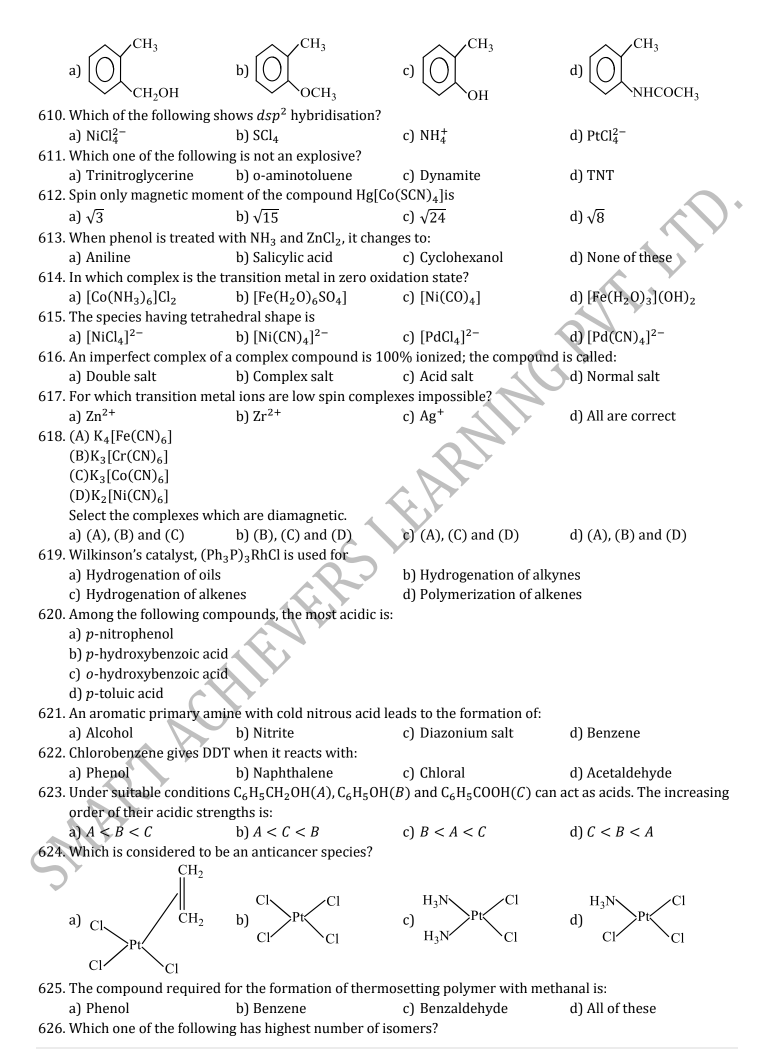




|      | a) $K_4[Fe(CN)_6]$                                | b) Fe(CNS) <sub>3</sub>                                | c) $Fe_4[Fe(CN)_6]_3$                | d) $K_3[Fe(CN)_6]$                                      |
|------|---|--|--------------------------------------|---|
| 568. | . What is the magnetic mor                        | ment of [FeF <sub>6</sub> ] <sup>3-</sup> ?            |                                      |   |
|      | a) 5.92   | b) 5.49  | c) 2.34                              | d) 4  |
| 569. | . Which of the following ca                       | n exhibit geometrical isom                             | erism?                               |   |
|      | a) $[MnBr_4]^{2-}$                                | b) [Pt(NH <sub>3</sub> ) <sub>3</sub> Cl] <sup>+</sup> | c) $[PtCl_2.P(C_2H_5)_3]_2$          | d) [Fe(H <sub>2</sub> O) <sub>5</sub> NO] <sup>2+</sup> |
| 570. | . A compound contains 2 d                         | lissimilar asymmetric C-ato                            | ms. The number of optical            | isomers are   |
|      | a) 2  | b) 3   | c) 4                                 | d) 5  |
| 571. | . Coordination number of                          | Ni in $[Ni(C_2O_4)_3]^{4-}$ is:                        |                                      |   |
|      | a) 3  | b) 6   | c) 4                                 | d) 5  |
| 572. | . Which compound exhibit                          | s optical isomerism?                                   | ,                                    |   |
|      | a) Pentaamminenitrocob                            | =  |                                      |   |
|      | b) Diamminedichloroplat                           | , ,  |                                      |   |
|      | · ·   | hylenediamine) chromium                                | (III) chloride                       |   |
|      | d) Tris-(ethylenediamine                          |  | () ••                                |   |
| 573  | . Ruthenium carbonyl is:                          | jeobale (III) bronnae                                  |                                      |   |
| 575  | a) $Ru(CO)_4$                                     | b) Ru(CO) <sub>5</sub>                                 | c) Ru(CO) <sub>8</sub>               | d) Ru(CO) <sub>6</sub>                                  |
| 574  | . Oxidation state of nitroge                      | , ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,                |                                      | u) (u(00) <sub>6</sub>                                  |
| 574  | -   | tion state   |                                      |   |
|      | -   | 0  | b) NH <sub>2</sub> OH –              | 1   |
|      | · · · · · ·                                       | -2   |                                      | 3   |
| 575  |   |  | )032                                 | 5   |
| 5/5. | -   | in participate in linkage ison                         |                                      |   |
|      | a) NH <sub>3</sub>                                | b) H <sub>2</sub> O                                    | c) $H_2NCH_2CH_2NH_2$                | d) $NO_2^-$   |
| 576. |   | s soluble in water than <i>p</i> -an                   | a <i>m</i> -nitrophenois because:    |   |
|      | , .   | ntramolecular H-bonding                                |                                      |   |
|      |   | ntermolecular H-bonding                                |                                      |   |
|      |   | ophenol is lower than those                            |                                      |   |
|      |   | volatile in steam than those                           | e of <i>m</i> -and <i>p</i> -isomers |   |
| 577. | . Among the following mos                         |  |                                      |   |
|      | a) Benzyl amine                                   | b) Aniline   | c) Acetanilide                       | d) <i>p</i> -nitro aniline                              |
| 578. | . The EAN of platinum in p                        | otassium hexachloroplatina                             | ate (IV) is:                         |   |
|      | a) 46   | b) 86  | c) 36                                | d) 84   |
| 579. | . The number of ions form                         | ed when copper ammoniun                                | n sulphate is dissolved in w         | ater is:  |
|      | a) 1  | b) 2   | c) 4                                 | d) Zero   |
| 580. |   | nnot show linkage isomeri                              | sm?                                  |   |
|      | a) $NO_2^-$                                       | 6) 1113  | c) CN <sup>-</sup>                   | d) SCN <sup>-</sup>                                     |
| 581. | . Xylenes on oxidation with                       | h acidic KMnO <sub>4</sub> gives:                      |                                      |   |
|      | a) Phthalic acid                                  | b) Isophthalic acid                                    | c) Terephthalic acid                 | d) All of these   |
| 582. | . The ratio of $\sigma$ -and $\pi$ -bond          | ds in benzene is:                                      |                                      |   |
|      | a) 2  | b) 4   | c) 6                                 | d) 8  |
| 583. | . The order of decreasing r                       | eactivity towards S <sub>E</sub> reacti                | on for the given compound            | is:   |
|      | (i)C <sub>6</sub> H <sub>6</sub>                  |  |                                      |   |
| c    | (ii)C <sub>6</sub> H <sub>5</sub> CH <sub>3</sub> |  |                                      |   |
|      | (iii)C <sub>6</sub> H <sub>5</sub> Cl             |  |                                      |   |
| ¥    | (iv)C <sub>6</sub> H <sub>5</sub> OH              |  |                                      |   |
|      | a) (ii)>(iv)>(i)>(iii)                            | b) (iv)>(iii)>(ii)>(i)                                 | c) (iv)>(ii)>(i)>(iii)               | d) (i)>(ii)>(iii)>(iv)                                  |
| 584  |   | ompounds is not optically a                            |                                      |   |
|      | Η CI  | ÇI Br  | Br H                                 | н ң   |
|      | Н   | н  |                                      |   |
|      | a) <sup>CI</sup> H                                | b) <sup>H</sup>  |                                      | d) Br/Br Di H   |
|      | H   | I<br>H   | H Br                                 | H Br  |
|      |   |  |                                      |   |

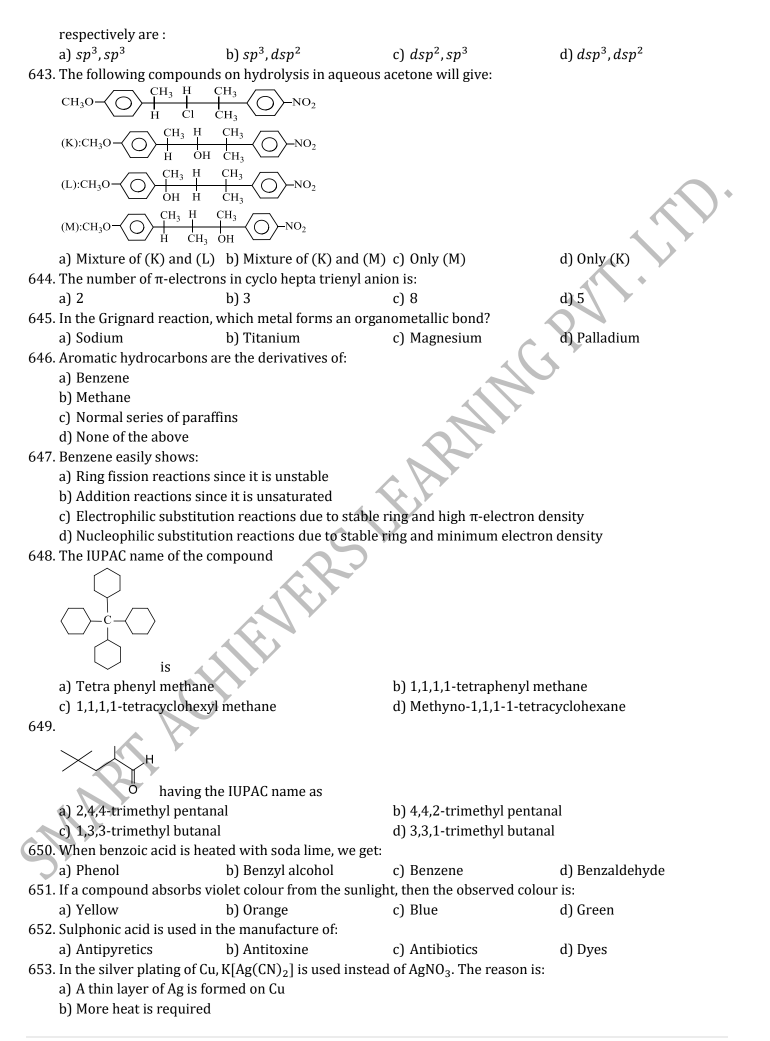
| 585. The number of geometric   | cal isomers of [Co(NH <sub>3</sub> ) <sub>3</sub> (N   | $(0_2)_3$ ] are:   |                                      |
|--|--|--|--------------------------------------|
| a) Zero  | b) 2   | c) 3   | d) 4                                 |
| 586. Phenol is less acidic than:   |  |  |                                      |
| a) Water   | b) <i>p</i> -methoxyphenol   | c) <i>p</i> -nitrophenol   | d) Ethanol                           |
| 587. In the reaction,  | )H Soda lime   |  |                                      |
| $C_6H_5CH_3 \xrightarrow{Oxidation} A \xrightarrow{NaO}$   | $B \xrightarrow{\text{Source}} C$  |  |                                      |
| a) C <sub>6</sub> H <sub>5</sub> OH  | b) C <sub>6</sub> H <sub>6</sub>   | c) C <sub>6</sub> H <sub>5</sub> COONa   | d) C <sub>6</sub> H <sub>5</sub> ONa |
| 588. Incorrect statement is  |  |  |                                      |
| -  | inite number of conformation of conformation of conformation of the second |  |                                      |
|  | e has considerable angle s<br>e is less stable then stagge   |  |                                      |
|  | on possess maximum energy  |  |                                      |
| 589. The complex $[Co(NH_3)_5B$  | -  |  |                                      |
| a) PbCl <sub>2</sub>   | b) AgNO <sub>3</sub>   | c) KI  | d) None of these                     |
| 590. Which of the following co   |  | est paramagnetic behaviour   | 2                                    |
| a) $[Fe(en)(bpy)(NH_3)_2]^2$   | +  | Ć.   |                                      |
| b) $[Co(OX)_2(OH)_2]^-$  |  |  |                                      |
| c) $[\text{Ti}(\text{NH}_3)_6]^{3+}$   | +  |  |                                      |
| $[V(gly)_2(OH)_2(NH_3)_2]$   | en = ethylenediamine and   | hny — hinyridylmoities   |                                      |
| (At. No. Ti= $22$ , V= $23$ , 1  | -  | bpy – bipyridyinoides  |                                      |
| 591. The coordination number   | •  | av increase to 8.  |                                      |
| a) Cobalt  | b) Osmium  | c) Nickel  | d) Iron                              |
| 592. Compound used for cover   | ring wounds caused by bit  | e of mad dog is:   |                                      |
| a) Benzoic acid  | b) Aniline   | c) Phenol  | d) Salicylic acid                    |
| 593. Cinnamic acid on decarbo  |  |  |                                      |
| a) Benzene   | b) Toluene   | c) Styrene   | d) Benzaldehyde                      |
| 594. In which of the following   |  | ow optical isomerism?<br>b) [PtCl(dien)]Cl, [NiCl <sub>2</sub> B   | r 12-                                |
| a) <i>Cis</i> -[Cr(C <sub>2</sub> O <sub>4</sub> ) <sub>2</sub> Cl <sub>2</sub> ] <sup>3-</sup> ;<br>c) [Co(NO <sub>3</sub> ) <sub>3</sub> (NH <sub>3</sub> ) <sub>3</sub> ], <i>cis</i> |  | d) $[Co(en)_3]Cl_3, cis-[Co(e)_3]Cl_3, cis-[Co(e)_$ |                                      |
| 595. The name of the ring stru   |  |  | -                                    |
|  | b) Chelate complex   | c) Polynuclear complex   | d) None of the above                 |
| 596. IUPAC name of   | $\frown$   |  |                                      |
| $Cl_2CH - CH - CH - CCl_3$   | is   |  |                                      |
| $Cl_2CH - CH - CH - CCl_3$ $ $ $C_2H_5$ $C_2H_5$   |  |  |                                      |
|  |  |  |                                      |
| a) 1,1,1,4,4-pentachloro-2<br>b) 3-(dichloromethyl)-4-   | (trichloromethyl)-hexane   |  |                                      |
|  | (dichloromethyl)-hexane  |  |                                      |
| d) 1,1,4,4,4-pentachloro-2   |  |  |                                      |
| 597. Which statement is wron   |  | yde and benzaldehyde?  |                                      |
| a) Both react with hydrox  | xylamine to form oximes  |  |                                      |
| b) Both react with HCN to  |  |  |                                      |
| c) Both react with NaOH  |  |  |                                      |
| d) Both react with hydraz  | _  | 0) 12+ -   |                                      |
| 598. The coordination number   |  |  | d) 1                                 |
| a) 4<br>599. Which reaction sequence   | b) 3<br>would be best to prepare   | c) 2<br>3-chloroaniline from henze   | d) 1<br>ne?                          |
| a) Chlorination, nitration   |  |  |                                      |
| b) Nitration, chlorination   |  |  |                                      |
|  |  |  |                                      |

| a) Nitration reduction chlorination  |                   |
|--|-------------------|
| c) Nitration, reduction, chlorination  |                   |
| d) Nitration, reduction, acetylation, chlorination, hydrolysis   |                   |
| 600. The complexes $(Co(NH_3)_6)$ [ $Cr(C_2O_4)_3$ ] and $[Cr(NH_3)_6]$ [ $Co(C_2O_4)_3$ ]             |                   |
| a) Geometrical isomerism<br>b) Ionization energy<br>b) Link and isomerical                             |                   |
| c) Coordination isomerism d) Linkage isomerism   |                   |
| 601. The reaction,   |                   |
| $C_6H_5NHCOCH_3 \xrightarrow{B_2/Fe} BrC_6H_4NHCOCH_3$   |                   |
| is an example of:  |                   |
| a) Substitution reaction   |                   |
| b) Addition reaction   |                   |
| c) Condensation reaction   |                   |
| d) Elimination reaction  |                   |
| 602. Given the molecular formula of the hexa coordinated complexes is                                  |                   |
| (A) CoCl <sub>3</sub> .6NH <sub>3</sub>  |                   |
| (B) $CoCl_3$ .5NH <sub>3</sub>   |                   |
| (C) $CoCl_3 .4NH_3$  |                   |
| If the number of coordinated NH <sub>3</sub> molecules in A, B and C respectively are 6, 5 and 4 the p | rimary valency    |
| in $(A)$ , $(B)$ and $(C)$ are   | initially valency |
| a) 6, 5, 4 b) 3, 2, 1 c) 0, 1, 2 d) 3, 3, 3  |                   |
|  |                   |
| $603. C_6 H_{14}$ has two tertiary carbons. The IUPAC name is  | mathulbutana      |
|  | methylbutane      |
| 604. The compound $[Co(NO_2)(NH_3)_5]Cl_2$ and $[Co(ONO)(NH_3)_5]Cl_2$ are examples of:                |                   |
|  | tion isomers      |
| 605. Which is not a $\pi$ -bonded complex?   |                   |
| a) Zeise salt b) Ferrocene c) Dibenzene chromium d) Tetrae   | -                 |
| 606. When phenol is treated with $PCl_5$ , the yield of chlorobenzene is generally poor because of     |                   |
|  | ry phosphate      |
| 607. Which will show tautomerism?  |                   |
|  |                   |
|  |                   |
| $a) \circ ( ) = ( ) = ( ) ( ) ( ) ( ) ( ) ( ) ( )$   |                   |
|  |                   |
| Ů  |                   |
|  |                   |
| o O  |                   |
|  |                   |
|  |                   |
| c) d)  |                   |
|  |                   |
| 0  |                   |
| 608. The IUPAC name of compound  |                   |
|  |                   |
|  |                   |
|  |                   |
| a) N-phenylaminoethanone b) N-phenylethanamide   |                   |
|  |                   |
| c) N-phenylmethanamide d) N-phenylaminomethane   |                   |
| 609. Which one of the following is most reactive towards electrophilic reagent?                        |                   |



Page | 50

a) [Co(NH<sub>3</sub>)<sub>5</sub>Cl]<sup>2+</sup> c)  $[Ru(NH_3)_4Cl^-]$ d)  $[In(PP_3)_2H(CO)]^{2+}$ b)  $[Co(en)_2Cl_2]^+$ 627. Which group is *o*- and *p*-directing? b)  $-SO_3H$ c) — COOH d)  $-NHCOCH_3$ a)  $-NO_2$ 628. When benzyl chloride is boiled with aqueous solution of lead nitrate in current of carbon dioxide, the main product is: b) Benzyl alcohol c) Benzaldehyde d) Nitrobenzene a) Benzoic acid 629. Ligands in complex compounds a) Donates electron pair b) Accept electron pair c) Neither accept electron pair nor donate d) All of the above 630. Aniline is separated by: a) Fractional crystallisation b) Fractional distillation c) Steam distillation d) Vacuum distillation 631. In which of the following octahedral complexes of Co (at. No. 27), will be magnitude of  $\Delta_0$  be the highest? a)  $[Co(CN)_6]^{3-1}$ b)  $[Co(C_2O_4)_3]^{3-1}$ c)  $[C_0(H_2O)_6]^{3+}$ d)  $[Co(NH_3)_6]^{3+}$ 632. The IUPAC name of K<sub>2</sub>[PtCl<sub>6</sub>] is b) Potassium hexachloroplatinate (IV) a) Hexachloroplatinate potassium d) Potassium hexachloroplatinum(IV) c) Potassium hexachloroplatinate 633. Aqueous solution of nickel sulphate on treating with pyridine and then adding a solution of sodium nitrite gives dark blue crystals of: c)  $[Ni(py)_4(NO_2)_2]$  d)  $[Ni(py)_3(NO_2)]_2SO_4$ a)  $[Ni(py)_4]SO_4$ b)  $[Ni(py)_2(NO_2)_2]$ 634. Benzyl alcohol is obtained from benzaldehyde by: b) Cannizzaro's reaction c) Kolbe's reaction a) Fittig's reaction d) Wurtz's reaction 635. The structure of the compound that gives a tribromo derivative on treatment with bromine water is: CH<sub>3</sub> CH<sub>2</sub>OH OH b) a) 636. The coordination number and the oxidation state of the element 'E' in the complex  $[E(en)_2(C_2O_4)]NO_2$ (where (en) is ethylene diamine) are, respectively : a) 6 and 3 b) 6 and 2 c) 4 and 2 d) 4 and 3 637. Benzaldehyde reacts with PCl<sub>5</sub> to give: a) Benzyl chloride b) Benzo trichloride d) Chlorobenzene c) Benzal chloride 638. Which one of the following complex ions has geometrical isomers? c)  $[Co(NH_3)_2(en)_2]^{3+}$ a)  $[Co(en)_3]^{3+}$ b)  $[Ni(NH_3)_5Br]^+$ d)  $[Cr(NH_3)_4(en)]^{3+}$ 639. The strongest acid among the following aromatic compounds is: b) para-chlorophenol a) Ortho-nitrophenol d) meta-nitrophenol c) *para*-nitrophenol 640. The isomers observed in alkanes is a) Metamerism b) Chain isomerism c) Position isomerism d) Geometrical isomerism 641. The two compounds pentaamminesulphatocobalt (III) bromide and pentaamminesulphatocobalt(III) chloride represent: a) Linkage isomerism b) Ionization isomerism c) Coordination isomerism d) No isomerism 642. Both  $[Ni(CO)_4]$  and  $[Ni(CN)_4]^{2-}$  are diamagnetic. The hybridisation of nickel in the compounds



c) Ag<sup>+</sup> ions are completely removed from solution d) Less availability of Ag<sup>+</sup> ion as Cu cannot displace Ag from Ag(CN)<sub>2</sub> 654. The strongest *o*-, *p*-directing group among the following is: a) — OH b) —Cl c)  $-C_{6}H_{5}$ d) —Br 655. Out of  $TiF_6^{2-}$ ,  $CoF_6^{3-}$ ,  $Cu_2Cl_2$  and  $NiCl_4^{2-}$  (Z of Ti = 22, Co = 27, Cu = 29, Ni = 28) the colourless species are: b) TiF<sub>6</sub><sup>2-</sup> and CoF<sub>6</sub><sup>3-</sup> c)  $Cu_2Cl_2$  and  $NiCl_4^{2-}$  d)  $TiF_6^{2-}$  and  $Cu_2Cl_2$ a)  $CoF_6^{3-}$  and  $NiCl_4^{2-}$ 656. Which is true in the case of  $[Fe(CN)_6]^{3-}$  complex? a)  $d^2sp^3$ -hybridization of Fe b) Paramagnetic c) One unpaired electron d) All of the above are correct 657. The IUPAC name of [Ni(PPh<sub>3</sub>)<sub>2</sub>Cl<sub>2</sub>]<sup>2+</sup> is b) Dichloro bis (triphenylphosphine)nickel(II) a) Bis-dichloro (triphenylphosphine)nickel(II) d) Triphenyl phosphine nickel (II) dichloride c) Dichloro triphenylphosphine nickel(II) 658. The complex [Co(NH<sub>3</sub>)<sub>3</sub>Cl<sub>3</sub>]is: a) Neutral b) Cationic c) Anionic d) None of these 659. From the stability constant (hypothetical values) given below, predict which is the strongest ligand? a)  $Cu^{2+} + 4NH_3 \rightleftharpoons [Cu(NH_3)_4]^{2+}$ ;  $(K = 4.5 \times 10^{11})$ b)  $Cu^2 + 4CN \rightleftharpoons [Cu(CN)_4]^{2-}$ ;  $(K = 2.0 \times 10^{27})$ c)  $Cu^{2+} + 2en \rightleftharpoons [Cu(en)_2]^{2+}$ ;  $(K = 3.0 \times 10^{15})$ d)  $Cu^{2+} + 4H_20 \rightleftharpoons [Cu(H_20)_4]^{2+}$ ;  $(K = 9.5 \times 10^8)$ 660. Which has highest m.p.? c) *p*-bromophenol a) o-bromophenol b) *m*-bromophenol d) m-chlorophenol 661. Hexafluorocobaltate(III) ion is found to be high spin complex, the probable hybrid state of cobalt in it is: b)  $sp^3$ d)  $sp^3d^2$ a)  $d^2sp^3$ c)  $sp^3d$ 662. Which isomeric dibromotoluene is most difficult to make from toluene? b) 2,4 c) 3,5 d) 2,6 a) 2,3 663. Which one of the following forms with an excess of CN<sup>-</sup>(cyanide) a complex? c)  $Ni^{2+}$ d)  $Fe^{2+}$ a) Cu<sup>+</sup> b) Ag 664. Nitration of salicylic acid gives: a) 2,4,6-trinitrosalicylic acid b) 2,4,6-trinitrophenol c) 2,4,6-trinitrobenzoic acid d) None of the above 665. The IUPAC name of the compound CH<sub>3</sub>-CH<sub>2</sub>-C--CH<sub>2</sub>-CH<sub>2</sub> OH a) N-hydroxy-3-amino pentane b) N-hydroxyamino pentane c) N-hydroxy-3-imino pentane d) None of the above 666. Which is not true of the coordination compound  $[Co(en)_2Cl_2]Cl_2$ a) Exhibits geometrical isomerism b) Exhibits optical isomerism c) Exhibits ionisation isomerism d) Is an octahedral complex 667. The IUPAC name of CH<sub>3</sub>-CH-CH-CH-ĊH<sub>2</sub>Br is a) 3-(bromomethyl)-2-methyl butanoyl chloride b) 3-(bromomethyl)-2-methyl propanoyl chloride c) 2-(bromomethyl)-3-methyl butanoyl chloride d) None of the above

668. Aniline is reacted with bromine water and the resulting product is treated with an aqueous solution of

sodium nitrite in the presence of dilute HCl. The compound so formed is treated with fluoroboric acid which is subsequently heated dry. The final product is:

a) *p*-bromofluorobenzene

b) p-bromoaniline

c) 2,4,6-tribromofluorobenzene

69. Which of the following is a common donor atom in ligands?  
a) Nitrogen b) Oxygen c) Arsenic d) Both (b) and (c) for the reaction of aniline with acetyl chloride in presence of NaOH gives:  
a) Acetanilide b) Aniline hydrochloride c) *p*-chloroaniline d) A red dye  
671. In the reaction, the compound "X" is:  

$$Me - O - CH = CHCOOH$$
  
a) CH<sub>3</sub>COOH

b) Br  $\cdot$  CH<sub>2</sub>COOH

c)  $(CH_3CO)_2O$ 

d) CHO · COOH

672. Which of the following will exhibit maximum ionic conductivity?

a) 
$$K_4[Fe(CN)_6]$$
 b)  $[Co(NH_3)_6]Cl_3$ 

c) Uric acid

d)  $[Ni(CO)_4]$ 

b) 3-carboxy-3-hydroxy-pentane-1, 5-dioic acid d) 3-carboxy-3-hydroxy-hexane-1, 6-dioic acid

d) None of these

a) Greater than (*X*) and (*Y*)

- b) Smaller than (*X*) and (*Y*)
- c) Greater than (*X*) but smaller than (*Y*)

d) Equal to zero

CH<sub>2</sub>

b) Ferrous ferricyanide a) Ferricyanide c) Ferrous cyanide d) Ferri ferrocyanide 676. The correct IUPAC name of OH

COOH. COOH COOH

a) 2-hydroxypropane

c) 2 carboxy-4 hydroxy-pentane-1, 5-dioic acid 677. The trivial name among the following is

a) Acetone b) Acetylene

678. The IUPAC name of  $[Pt(NH_3)_4(NO_2)Cl]SO_4$  is a) Chloronitro tetrammine platinum (IV) sulphate

b) Tetrammine chloronitro platinum (II) sulphate

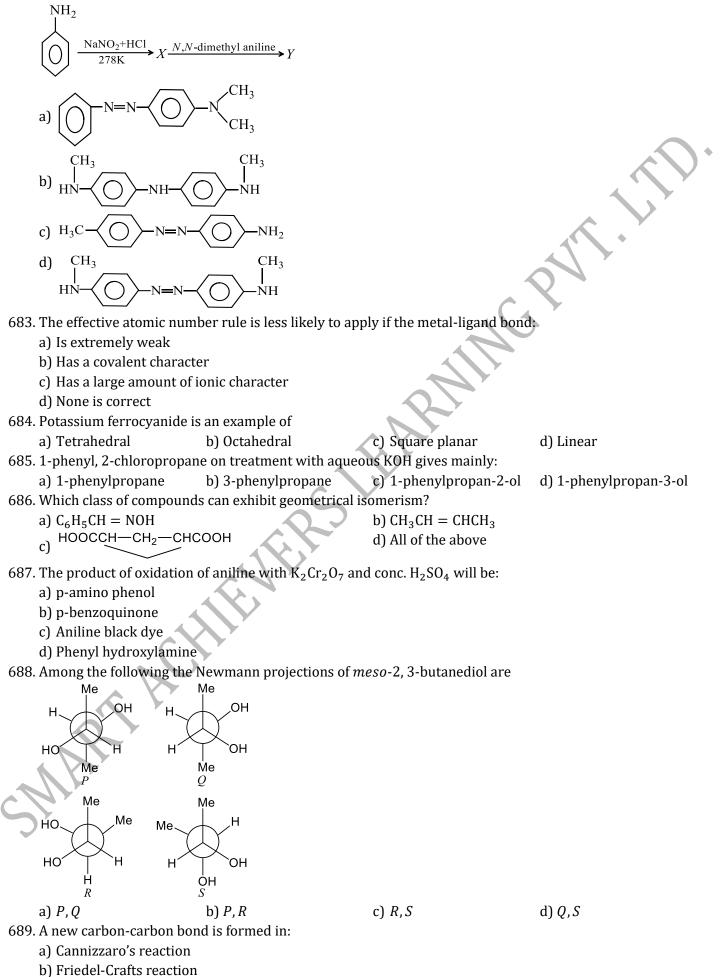
c) Tetrammine chloronitro platinum (IV) sulphate

d) Chlorotetrammine nitroplatinum (IV) sulphate

679. The overlapping in benzene is in carbon-carbon orbitals of the type:

a) 
$$p - p$$
 b)  $sp - sp$  c)  $sp^2 - sp^2$  d)  $sp^3 - sp^3$   
680. Change in composition of coordination sphere yields which type of isomer?  
a) Geometrical b) Ionization c) Optical d) None of these  
681. The IUPAC name of K<sub>2</sub>[Ni(CN)<sub>4</sub>] is  
a) Potassium tetracyanonickelate (II) b) Potassium tetracyanatonickelate (III)

c) Potassium tetracyanatonickel (II) d) Potassium tetracyanonickel (III) 682. Aniline in a set of the following reactions yielded a coloured compound *Y*:

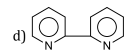


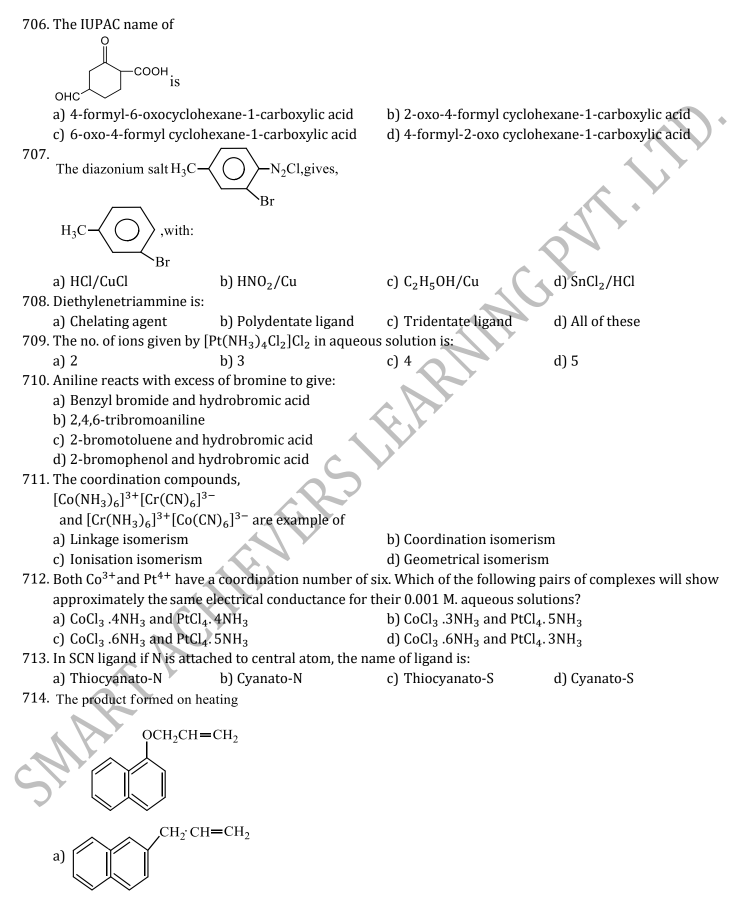
c) Clemmensen reduction

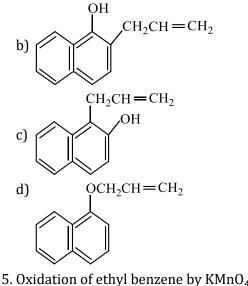
d) None of the above

690. Which of the following compounds can exhibit tautomerism?

| 690. Which of the following compounds can exhibit tautomerism? |  |  |                                |  |
|--|--|--|--------------------------------|--|
| СНО  |  | NO <sub>2</sub>                                    | $CH_3 - CH - CH_3$             |  |
| a) [O]   | b) [ ]   |  | d)                             |  |
|  | 5)   |  | NH <sub>2</sub>                |  |
|  |  |  |                                |  |
|  | npound among the following is  |  |                                |  |
| a) Benzylamine   | b) Aniline   | c) Acetanilide                                     | d) <i>p</i> -nitroaniline      |  |
|  | ving has least oxidation state o   |  |                                |  |
| a) $K_3[Fe(OH)_6]$   |  | b) $K_2[FeO_4]$                                    |                                |  |
| c) $FeSO_4(NH_4)_2SC$  |  | d) [Fe(CN) <sub>6</sub> ] <sup>3–</sup>            |                                |  |
|  | netic moment value (in Bohr n  |  |                                |  |
| a) 0   | b) 2.84  | c) 4.90  | d) 5.92                        |  |
| 694. Which is an excelle                                       | ent antiseptic?  |  |                                |  |
| a) Phenol  | b) Benzyl alcohol  | c) Benzaldehyde                                    | d) Acetic acid                 |  |
| 695. Scientist who expl  | ained the structures and isome   | erism in the complex comp                          | ound was:                      |  |
| a) Sidgwick  | b) Pauling   | c) Powell  | d) Werner                      |  |
| 696. The cation that do  | es not form an ammine comple   | ex with excess of ammonia                          |                                |  |
| a) Al <sup>3+</sup>  | b) Ag <sup>+</sup>   | c) Cu <sup>2+</sup>                                | d) Cd <sup>2+</sup>            |  |
| 697. The complex ion w   | hich has the highest magnetic  | moment among the follow                            | ving is                        |  |
| a) [CoF <sub>6</sub> ] <sup>3–</sup>                           | b) [Co(NH <sub>3</sub> ) <sub>6</sub> ] <sup>3+</sup>  | c) $[Ni(NH_3)_4]^{2+}$                             | d) $[Ni(CN)_4]^{2-}$           |  |
| 698. For square planar   | complex of platinum (II), [Pt(N  | $(H_3)(Br)(Cl)Py]^{2+}$ , how m                    | any isomeric forms are         |  |
| possible?  |  |  | -                              |  |
| a) Two   | b) Three   | c) Four  | d) Six                         |  |
| 699. Which of the follow                                       | ving has highest boiling point?  |  | 2                              |  |
| a) Benzene   | b) Phenol  | c) Toluene   | d) Ethyl benzene               |  |
| ,  | ing organic compound on heat   | ,  | 5 6                            |  |
| _  | ig vapours. The compound cou   | -  | y                              |  |
| a) Nitrobenzene  | b) Benzamide   | c) <i>N</i> , <i>N</i> -dimethyl amin              | ne d) Aniline                  |  |
| -  | -  |  | nimum electrical conductivity? |  |
|  | tinum (IV) chloride  |  | ne platinum (IV) chloride      |  |
| <i>,</i> .   | mine platinum (IV) chloride  | , ,  | ne platinum (IV) chloride      |  |
| 702. False statement is  |  | -,   |                                |  |
|  | s increase the enol content in t   | automerism   |                                |  |
|  | rom the normal bond angles in  |  | olecule                        |  |
|  | nave identical physical propert  | _  | olecule                        |  |
|  | an also be position isomers  | 1105   |                                |  |
|  | name of the compound is  |  |                                |  |
| $CH_3 - CH - CH -$   | -  |  |                                |  |
| $c_{13} - c_{11} - c_{11} - c_{11}$                            | $G_{11} = G_{112} = G_{113}$   |  |                                |  |
| Cl Br  | l<br>I   |  |                                |  |
| a) 4-bromo-5-chlo  | I  | h) 2 bromo 2 chloro                                | 4 jada hayana                  |  |
|  |  | b) 3-bromo-2-chloro-                               |                                |  |
| C) 3-bromo-4-iodo  |  | d) 2-bromo-3-bromo-                                |                                |  |
|  | $_{6}$ H <sub>5</sub> CH <sub>2</sub> Cl) can be prepared from the solution of the s |  |                                |  |
| a) $SO_2Cl_2$<br>705 The compound 2.2                          | b) SOCl <sub>2</sub>   | c) S <sub>2</sub> Cl <sub>2</sub>                  | d) NaOCl                       |  |
| $\sim$ $\sim$ $\sim$   | '-bipyridine has the structure   |  |                                |  |
|  |  | $m\left( \bigcirc \right) \left( \bigcirc \right)$ |                                |  |
| $a_{N} \sim N$   |  |  |                                |  |
|  |  |  |                                |  |







715. Oxidation of ethyl benzene by KMnO<sub>4</sub> gives:

a) Benzyl alcohol b) Benzophenone c) Acetophenone d) Benzoic acid 716. One of the following statements regarding Reimer-Tiemann reaction is false:

- a) Reaction of phenol with CHCl<sub>3</sub> and KOH
- b) CCl<sub>2</sub> acts as a nucleophile
- c) Reaction of phenol with CCl<sub>4</sub> and NaOH

d) Reaction of phenol with formaldehyde to form bakelite

- 717. The structure representing a heterocyclic compound is
  - $CH_2$ a)  $CH_2 - CH_2$

718. Phenol reacts with Br<sub>2</sub>in CCl<sub>4</sub> at low temperature to give:

a) m-bromophenol

- b) o-and p-bromophenol
- c) p-bromophenol
- d) 2,4,6-tribromophenol

719. The correct name of the compound  $[Cu(NH_3)_4](NO_3)_2$ , according to IUPAC system is

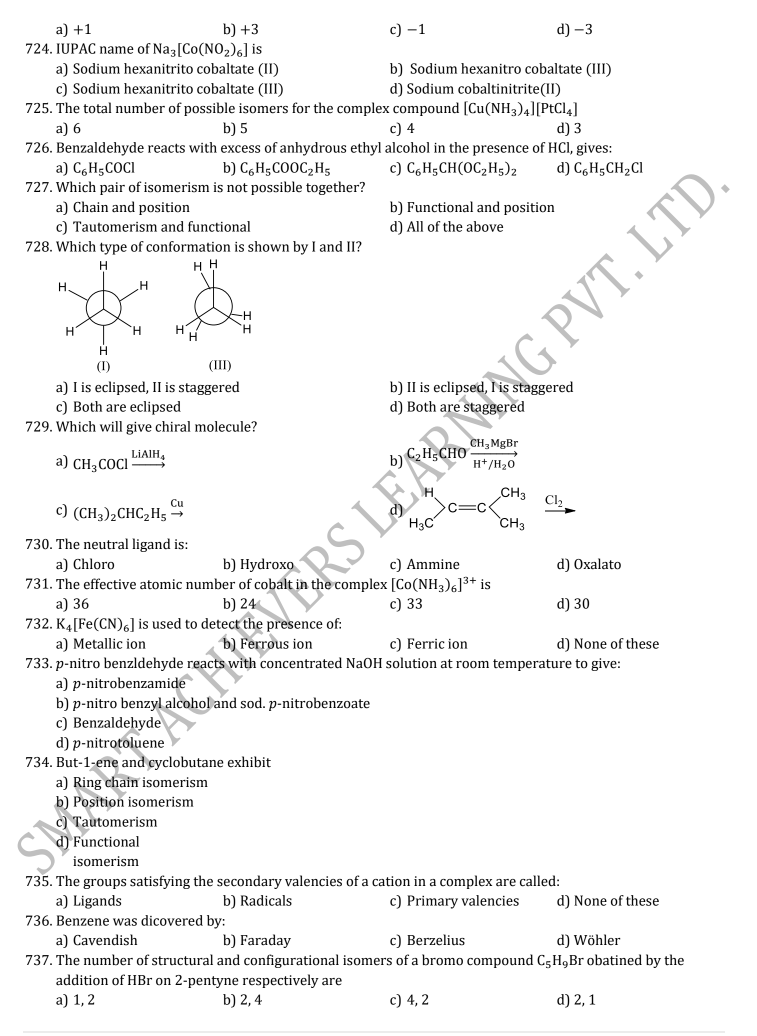
- a) Cuprammonium nitrate b) Tetrammine copper (II) dinitrate
- c) Tetrammine copper (II) nitrate d) Tetrammine copper (II) dinitrite

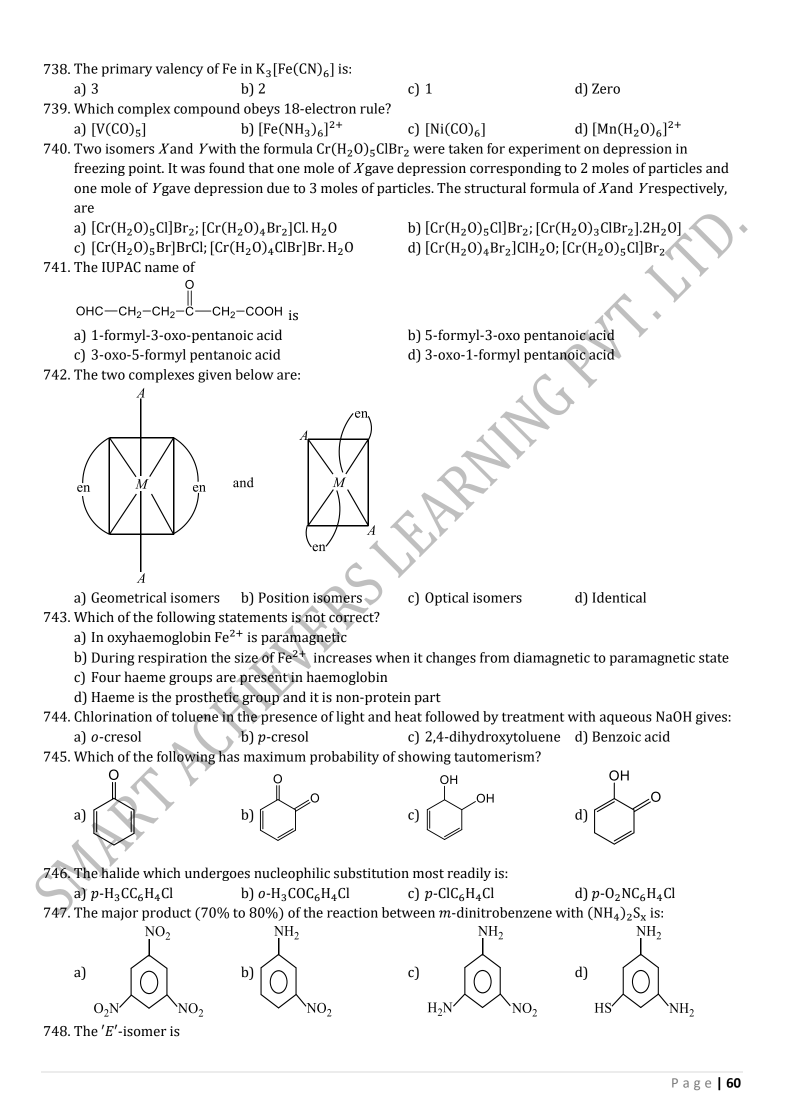
720. Nitroethane can exhibit one of the following kind of isomerism

a) Metamerism b) Optical activity c) Tautomerism d) Position isomerism 721. What would be the correct IUPAC name of

CH<sub>2</sub>

- a) 3,3-dimethyl-3-cyclopentyl propanal
- b) 3-methyl-3-cyclopentyl butan-1-al
- c) 1-(1-methyl-1-formyl) methylethyl cyclopropane
- d) None of above
- 722. The number of unpaired electrons in the square planar  $[Pt(CN)_4]^{2-}$  ion is
  - c) 0 d) 3 a) 2 b) 1
- 723. The oxidation number of cobalt in  $K[Co(CO)_4]$  is





>c=c< d) None of the above 749. The Baeyer angle strain is minimum in a) Cyclopropane b) Cyclobutane c) Cyclopentane d) Cyclohexane 750. Among the following ions, which one has the highest unpaired electrons? a)  $[Cr(H_2O)_6]^{3+}$ b)  $[Zn(H_20)_6]^{2+}$ c)  $[Fe(H_20)_6]^{2+}$ d)  $[Cr(H_2O)_6]^3$ 751. Which will give a white precipitate with AgNO<sub>3</sub> in aqueous solution? a)  $[Co(NH_3)_5Cl](NO_2)_2$ b)  $[Pt(NH_3)_6]Cl_4$ c)  $[Pt(en)Cl_2]$ d)  $[Cu(NH_3)_4]S$ 752. The organic product formed in the reaction;  $C_6H_5COOCH_3 \xrightarrow{(I)LiAlH_4}{(II)H_2O}$ : a) C<sub>6</sub>H<sub>5</sub>CH<sub>2</sub>OH and CH<sub>3</sub>OH b) C<sub>6</sub>H<sub>5</sub>COOH and CH<sub>4</sub> c) C<sub>6</sub>H<sub>5</sub>CH<sub>3</sub> and CH<sub>3</sub>OH d)  $C_6H_5CH_3$  and  $CH_4$ 753. Complexes with bidentate ligands are called: b) Chelates a) Ligands c) Complex d) None of these 754. Excited state configuration of Mn<sup>2+</sup> is b)  $t_{2g}^3 e_g^2$ d)  $t_{2a}^5 e_a^0$ a)  $t_{2g}^4$ 755. The IUPAC name of  $CH_3$ ,COOC₂H₅ is b) Ethyl methyl butenoate a) Ethyl acetylate d) Ethyl (3-methyl) but-2-enoate c) Ethyl acetoethanoate 756. The compound which result from the coordination of carbon monoxide are known as b) Electronic a) Carbon permono c) Carbonyls d) None of these 757. The correct IUPAC name of AlCl<sub>3</sub>(EtOH)<sub>4</sub> is: a) Aluminium(II) chloride-4-ethanol b) Aluminium(III) chloride-4-ethanol c) Aluminium(IV)chloride-4-hydroxy ethane d) Aluminium chloride-4-ethanol 758. The IUPAC name of  $[Co(NH_3)_6][Cr(C_2O_4)_3]$  is a) Hexaamine cobalt (III) tris (oxalato ) chromium b) Hexaamine cobalt (III) tris (oxalato ) chromate(III) c) Hexaamine cobalt tris (oxalato ) chromium(III) d) Hexaamine cobalt (III) chromium (III) oxalate 759. The insecticide, germicide gammexane is a formulation for: a) DDT b) Benzene hexachloride c) Hexachlorobenzene d) Chloral 760. Among  $[Ni(CO)_4]^{2-}$ ,  $[Ni(CN)_4]^{2-}$ ,  $[NiCl_4]^{2-}$  species, the hybridisation states of the Ni atom are, respectively (Atomic no. of Ni=28) c)  $sp^3$ ,  $sp^3$ ,  $dsp^2$ d)  $dsp^2$ ,  $sp^3$ , spa)  $sp^3$ ,  $dsp^2$ ,  $dsp^2$ b)  $sp^3$ ,  $dsp^2$ ,  $sp^3$ 761. Which of the following complex ions is expected to absorb visible light?

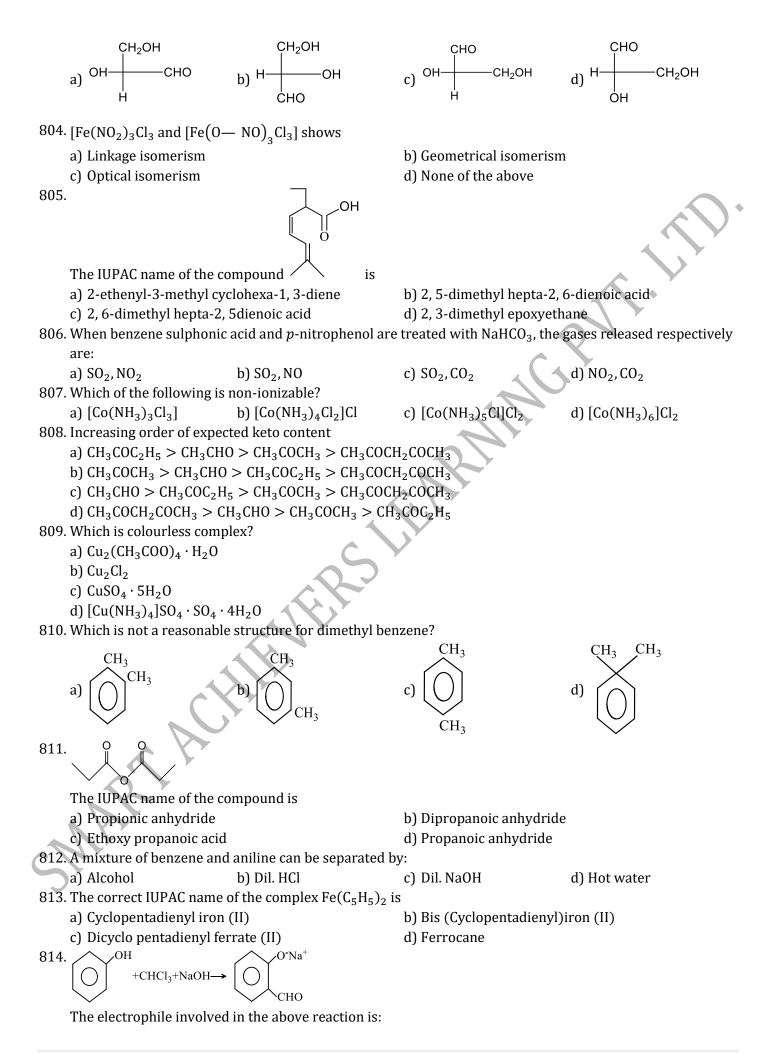
a)  $[Zn(NH_3)_6]^{2+}$ b)  $[Sc(H_2O)_3(NH_3)_3]^{3+}$ c)  $[Ti(en)_2(NH_3)_2]^{4+}$ d)  $[Cr(NH_3)_6]^{3+}$ [At. no. Zn = 30, Sc = 21, Ti = 22, Cr = 24] 762. Chain isomers of CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>OH is/are a) 2 b) 3 c) 4 d) 5 763. Although chlorobenzene does not give Ulmann's reaction. However, presence of ... group in chlorobenzene at o-, p-position enables it to give Ulmann's reaction. d)  $SO_3H$ a)  $NO_2$ b)  $NH_2$ c) OH 764. Which statement is true? a) A compound with *R* configuration is the (+) enantiomer b) If configuration changes from + to -, that essentially means inversion of configuration take place c) An achiral molecule reacts to give a chiral molecule, always racemic forms d) By breaking two bonds on the chiral centre configuration changes 765. Which can be used for carrying out electrophilic aromatic substitution? d) Hydride ion b) Liquid NH<sub>3</sub> c) Oleum a) Water 766. Which of the following can participate in linkage isomerism? b) H<sub>2</sub>NCH<sub>2</sub>CH<sub>2</sub>NH<sub>2</sub> c) H<sub>2</sub>O d) : NH<sub>3</sub> a)  $NO_2^-$ 767. Aniline in a set of reactions yielded a product *D*.  $\operatorname{NH}_2 \xrightarrow{\operatorname{NaNO}_2} A \xrightarrow{\operatorname{CuCN}} B \xrightarrow{\operatorname{H}_2} C \xrightarrow{\operatorname{HNO}_2} D$ The structure of the product *D* would be: a)  $C_6H_5NHCH_2CH_3$ b) C<sub>6</sub>H<sub>5</sub>CH<sub>2</sub>OH c)  $C_6H_5CH_2NH_2$ d) C<sub>6</sub>H<sub>5</sub>NHOH 768. The number of ions formed when cuprammonium sulphate is dissolved in water is a) Zero c) 2 b) 1 d) 4 769. Tautomerism is not exhibited by : a)  $C_6H_5 - CH = CH - OH$  b) 770. Benzaldehyde reacts with NH<sub>3</sub> to give: a) Aniline b) Benzamide d) Hydrobenzamide c) Phenylcyanide 771. In coal-tar fraction of heavy oil, the aromatic compound present is: a) Cresol b) Pyridine c) Benzene d) Anthracene 772. Optical isomerism is shown by octahedral complexes a) Having all monodentate ligands b) Having all the three bidentate ligands c) Having two *trans* bidentate ligands d) Having two trans monodentate ligands 773. Which can be hydrolysed most easily? a)  $(C_6H_5)_3CCl$ b)  $C_6H_5CH_2Cl$ c)  $(C_6H_5)_2$ CHCl d)  $C_6H_5Cl$ 774. The most stable configuration of *n* butane will be c) Gauche a) Skew boat b) Eclipsed d) Staggered-anti 775. Anhydrous aluminium chloride is used in Friedel-Craft's reaction because it is: a) Electron rich b) Soluble in ether c) Ionizable to chloride and aluminium ions

## d) Electron deficient molecule

776. The two isomers given below are

COOH (i) I H - C - OHHO - C - HТ COOH (ii) HOOC H - C - OHH - C - OHHOOC b) Diastereomers c) Measomers d) Position isomers a) Enantiomers 777. Which of the following has lowest boiling point? a) Phenol b) o-nitrophenol c) *m*-nitrophenol d) *p*-nitrophenol 778. The IUPAC name of [Ni(NH<sub>3</sub>)<sub>4</sub>][NiCl<sub>4</sub>] is a) Tetrachloro nickel (II) - tetraammine nickel (II) b) Tetraammine nickel (II) -tetrachloro nickel(II) c) Tetraammine nickel (II) -tetrachloro nickelate(II) d) Tetrachloro nickel (II) -tetraammine nickelate(0) 779. All ligands are: c) Neutral a) Lewis acid b) Lewis base d) None of these 780. Aspirin is known as: b) Acetyl salicylate c) Methyl salicylic acid a) Phenyl salicylate d) Acetyl salicylic acid 781. Which of the following has on optical isomer? a)  $[Co(NH_3)_3Cl]^+$ b)  $[Co(en)(NH_3)_2]^{2+}$ c)  $[Co(H_2O)_4(en)]^{3+}$ d)  $[Co(en)_2(NH_3)_2]^{3+}$ 782. The IUPAC name of the compound a) 1,1,1-trichloro-2,2-diphenyl ethane b) 2,4,5-trichloro hexanol c) 2,2,2-trichloro bicyclo [4.4.0] nenone d) 2,2,2-trichloro-1,1-diphenyl ethane 783. The property by virtue of which a compound can rotate the plane of polarised light is known as a) Polarisability b) Phosphorescence c) Optical activity d) Polarization 84. The molecules represented by the following two structures are CH<sub>3</sub> н он но Br Ŕr  $\dot{C}_2H_5$ a) Epimers b) Diastereomers c) Enantiomers d) Identical 785. The IUPAC name of the coordination compound  $K_3$  [Fe(CN)<sub>6</sub>] is a) Tripotassium hexacyanoiron (II) b) Potassium hexacyanoiron(II) c) Potassium hexacyanoferrate (III) d) Potassium hexacyanoferrate (II) 786. Which one of the following is an inner orbital complex as well as diamagnetic in nature?

| a) $[Cr(NH_3)_6]^{3+}$                    |   |                                    | d) $[Zn(NH_3)_6]^{2+}$                |  |  |
|---|---|------------------------------------|---------------------------------------|--|--|
| 787. How many unpaired elec               | -   | tral metal ion of $[CoCl_4]^{2-1}$ | ?                                     |  |  |
| a) 3                                      | b) 4  | c) 5                               | d) 2                                  |  |  |
| 788. Show the coordination n              | umber of the metal ion, its o                         | oxidation number, the num          | ber of electrons in <i>d</i> -        |  |  |
| orbitals and the number                   | of unpaired electrons <i>d</i> -orb                   | itals respectively in comple       | $ex [Co(H_2O)_4SO_3]Cl.$              |  |  |
| a) 6, 3, 6, 4                             | b) 6, 3, 6, 0   | c) 5, 3, 6, 4                      | d) 5, 3, 6, 0                         |  |  |
| 789. Benzene reacts withto                | o give acetophenone.                                  |                                    |                                       |  |  |
| a) Acetyl chloride                        |   |                                    |                                       |  |  |
| b) Acetyl chloride in pres                | sence of anhy. AlCl <sub>3</sub>                      |                                    |                                       |  |  |
| c) Anhy. AlCl <sub>3</sub>                |   |                                    | $\langle \nabla \rangle$              |  |  |
| d) None of the above                      |   |                                    |                                       |  |  |
| 790. Which group would you                | introduce into a drug or a d                          | ye to make it water soluble        | ?                                     |  |  |
| a) $-NO_2$                                | b) —Cl  | c) $-SO_3H$                        | d) —OH                                |  |  |
| 791. In the coordination com              | pound, K <sub>4</sub> [Ni(CN) <sub>4</sub> ],oxidatio | on state of nickel is              |                                       |  |  |
| a) —1                                     | b) +1   | c) 0                               | d) +2                                 |  |  |
| 792. The IUPAC name of [Cr(I              | ,   |                                    |                                       |  |  |
| a) Tetraaminodichlorocl                   |   | Ć                                  |                                       |  |  |
| b) Tetraaminodichlorocl                   |   |                                    |                                       |  |  |
| c) Dichlorotetraammine                    | , ,   |                                    |                                       |  |  |
| d) Tetraaminodichlorocl                   | , ,   |                                    |                                       |  |  |
| 793. Vanillin, used as a flavou           |   |                                    |                                       |  |  |
| a) An aliphatic alcohol                   | b) An aromatic aldehyde                               | c) A hydrocarbon                   | d) A carbohydrate                     |  |  |
| 794. Which of the following w             |   |                                    |                                       |  |  |
| a) $[Cr(en)(H_2O)_4]^{3+}$                | -   |                                    | d) $[Cr(NH_3)_6]^{3+}$                |  |  |
| 795. Which one is a mixed ket             |   |                                    |                                       |  |  |
| a) Benzophenone                           | b) Benzenone  | c) Acetophenone                    | d) Dibenzyl ketone                    |  |  |
| 796. Transition metals can for            | ,   | -)                                 |                                       |  |  |
| a) Zero oxidation state                   | b) Cation form  | c) Anion form                      | d) All of these                       |  |  |
| 797. Toluene on oxidation wi              |   | •                                  | - ,                                   |  |  |
| a) Phenol                                 | b) Benzoic acid                                       |                                    | d) Benzyl alcohol                     |  |  |
| 798. $[Pt((NH_3)_4]Cl_2$ is               |   |                                    |                                       |  |  |
| a) Pyramidal                              | b) Pentagonal   | c) Tetrahedral                     | d) Square planar                      |  |  |
| 799. In Fe(CO) <sub>5</sub> , the FE—C bo |   | .,                                 |                                       |  |  |
| a) $\pi$ -character only                  | r   |                                    |                                       |  |  |
| b) Both $\sigma$ and $\pi$ -character     | ers   |                                    |                                       |  |  |
| c) Ionic character                        | · ·   |                                    |                                       |  |  |
| d) $\sigma$ -character only               |   |                                    |                                       |  |  |
| 800. Which molecule has tetra             | ahedral geometry?                                     |                                    |                                       |  |  |
| a) $[Co(NH_3)_6]^{3+}$                    | e .   | c) Fe(CO) <sub>5</sub>             | d) [NiCl <sub>4</sub> ] <sup>2–</sup> |  |  |
| $801. [Co(NH_3)_5Br]SO_4$ and [C          |   |                                    | - 7 L - 43                            |  |  |
| a) Linkage isomerism                      |   | 1                                  |                                       |  |  |
| b) Geometrical isomeris                   | m   |                                    |                                       |  |  |
| c) Ionization isomerism                   |   |                                    |                                       |  |  |
| d) Optical isomerism                      |   |                                    |                                       |  |  |
|   | 802. The compounds $R - NO_2$ and $R - ONO$ are       |                                    |                                       |  |  |
| a) Geometrical isomers                    | -   | b) Functional isomers              |                                       |  |  |
| c) Metamers                               |   | d) Optical isomers                 |                                       |  |  |
| 803. Which of the following F             | ischer projection formula is                          |                                    | ?                                     |  |  |
| 8-  | . ,   | 0,,                                |                                       |  |  |



| a) dichloromethyl cati<br>b) Dichlorocarbene (:                   |   |   |   |
|---|---|---|---|
| ,   |   |   |   |
| c) Trichloromethyl ani  | < 5, <sup>-</sup>   |   |   |
| d) Formyl cation (CH  | 0)  |   |   |
| 815. Benzoyl Chloride is p  | repared from benzoic acid                                 | by:   |   |
| a) Cl <sub>2</sub> , <i>hv</i>                                    | b) SO <sub>2</sub> Cl <sub>2</sub>                        | c) SOCl <sub>2</sub>                                | d) Cl <sub>2</sub> , H <sub>2</sub> O   |
|   | g ions forms most stable co                               |   |   |
| a) Fe <sup>3+</sup>   | b) Mn <sup>2+</sup>                                       | c) Ni <sup>2+</sup>                                 | d) Cu <sup>2+</sup>                     |
|   |   | uld exhibit the lowest value                        | e of paramagnetic behaviour?            |
|   | n=25, Fe=26, Co=27)                                       | $\sum \left[ 1 - \left( \alpha \right) \right]^{2}$ |   |
|   | b) [Fe(CN) <sub>6</sub> ] <sup>3–</sup>                   |   | d) [Cr(CN) <sub>6</sub> ] <sup>3–</sup> |
|   | g statements is not correct                               |   |   |
|   |   | fer in the state of hybridisa                       |   |
| -   |   | fer in the magnetic propert                         | les.                                    |
| -   | $[Cl_4]^{2-}$ and $[Ni(CN)_4]^{2-}$ dif                   | fer in geometry.<br>fer in primary valencies of s   | nickel                                  |
|   |   |   | more stability is shown by:             |
| a) $[Fe(H_20)_6]^{3+}$  | b) $[Fe(CN)_6]^{3-}$                                      | c) $[Fe(C_2O_4)_3]^{3-1}$                           | d) $[FeCl_6]^{3-}$                      |
| 820. In the reaction,   | b) [i c(civ)6]  |   |   |
|   |   |   |   |
| $C_8H_6O_4 \xrightarrow{\text{NH}_3} X \xrightarrow{\text{NH}_3}$ | CONH <sub>2</sub>   |   |   |
|   | СООН  |   |   |
| the intermediate 'X' is   | 3:  |   |   |
| a) Phthalic anhydride   | b) Phthalic acid 👝  | c) <i>o</i> -xylene                                 | d) Benzoic acid                         |
| 821. Which of the followin  | g is $\pi$ complex?                                       | Y   |   |
| a) Trimethyl alumini  | ım b) Ferrocene   | c) Diethyl zinc                                     | d) Nickel carbonyl                      |
|   |   |   |   |
| •   |   | -   | ound formed is salicyladehyde           |
|   | ace of chloroform the prod                                |   |   |
| a) Salicyladehyde   | b) Phenolphthalein  | c) Salicylic acid                                   | d) Cyclohexanol                         |
|   |   | g (c) complexing, the set of                        | properties shown by CN <sup>-</sup> ion |
| towards metal specie  |   |   | d)                                      |
| a) B, c   | b) A, b, c  | c) C, a   | d) A, b                                 |
| a) —NO <sub>2</sub>   | g is most powerful <i>meta</i> d<br>b) —SO <sub>3</sub> H | c) —CHO   | d) —COOH                                |
|   | owing compounds will sho                                  | -   | u) —coon                                |
| a) $CH_3COC_3H_7$   | b) $CH_3OC_2H_5$  | c) $CH_3SC_2H_5$                                    | d) CH <sub>3</sub> OCH <sub>3</sub>     |
| 826. The hybridization of   | 5 2 5   | 0, 0113002115                                       |   |
| a) $d^2sp^3$  | b) $sp^2d^3$  | c) $sp^3d$  | d) $sp^3d^2$                            |
| 827. The correct name of [  | <i>, , ,</i>  |   |   |
|   | um (II) dichloro tetrammir                                | ne platinate  |   |
|   | ne platinum (IV) tetrachlo                                |   |   |
| c) Tetrammine dichlo  | oro platinum (IV) tetrachlo                               | ro platinate (II)                                   |   |
| d) Tetrachloro platin   | um (II) tetrammine platina                                | ite (IV)  |   |
| 828. The oxidation state of                                       | firon in K <sub>4</sub> [Fe(CN) <sub>6</sub> ]is          |   |   |
| a) 1  | b) 4  | c) 3  | d) 2                                    |
|   | compound can be detecte                                   |   |   |
| a) Change in colour   | b) Change in solubilit                                    | cy c) Change in pH                                  | d) All are correct                      |
| 330. The complex that viol  | ates the EAN:   |   |   |

| a) Potassium ferrocyanide<br>b) Potassium ferricyanide<br>c) Nickel carbonyl   |  |  |
|--|--|--|
| d) Cobalt(III) hexaammine chloride   |  |  |
| 831. Chlorobenzene on heating with aqueous $\rm NH_3$ und                      | ler pressure in presence of            | Cu <sub>2</sub> Cl <sub>2</sub> gives: |
| a) Aniline b) Benzamide  | c) o-dichlorobenzene                   | d) Chloroaminobenzene                  |
| 832. The complex, [Pt(Py)(NH <sub>3</sub> )BrCl] will have how m               | any geometrical isomers?               | -                                      |
| a) 2   |  |  |
| b) 3   |  |  |
| c) 4   |  | $\sim$                                 |
| d) 0   |  |  |
| 833. Which one doesn't have $\pi$ –bond?                                       |  |  |
| a) Grignard reagent  | b) Dibenzene chromiu                   | m                                      |
| c) Zeise's salt  | d) Ferrocene                           |  |
| 834. The IUPAC name of the compound $CH_2 - CH - CC$                           |  |  |
|  |  |  |
| NH <sub>2</sub> OH   | Ć                                      |  |
| a) 1-hydroxy-2-aminopropanoic acid   | b) 2-hydroxy-3-aminor                  | propanoic acid                         |
| c) 3-amino-2-hydroxypropanoic acid   | d) 2-hydroxy-1-aminoj                  | propanoic acid                         |
| 835. EDTA is aligand.  |  |  |
| a) Monodentate   |  |  |
| b) Hexadentate   |  |  |
| c) Bidentate   |  |  |
| d) Tridentate  |  |  |
| 836. Thymol, a phenol derivative is mainly used as: 🙏                          |  |  |
| a) Germicide   |  |  |
| b) Insecticide   |  |  |
| c) Antibiotic  |  |  |
| d) Fragrance compound and antiseptic   |  |  |
| 837. Which of the following complex has zero magneti                           | c moment (spin only)?                  |  |
| a) $[Ni(NH_3)_6]Cl_2$ b) $Na_3[FeF_6]$   | c) $[Cr(H_2O)_6]SO_4$                  | d) $K_4[Fe(CN)_6]$                     |
| 838. Which compound is zero valent metal complex?                              |  |  |
| a) $[Ni(CO)_4]$ b) $K_3[Fe(CN)_6]$   | c) $[Pt(NH_3)_2Cl_2]$                  | d) $[Cu(NH_3)_4SO_4]$                  |
| 839.   |  |  |
| The IUPAC name of is   |  |  |
| a) Bicyclo [5.5.0] nonane  | b) Biphenyl                            |  |
| c) Cyclopropyl cyclohexane   | d) Spiro [3.5] nonane                  |  |
| 840. The tetrahedral crystal field splitting is onlyof                         | the octahedral splitting.              |  |
| a) 1/9 b) 2/9  | c) 4/9                                 | d) 5/9                                 |
| 841. IUPAC name of [Co(ONO)(NH <sub>3</sub> ) <sub>5</sub> ]Cl <sub>2</sub> is |  |  |
| a) Pentammine nitrocobalt (II) chloride  | b) Pentammine nitrosc                  |  |
| c) Pentammine nitritocobalt (III) chloride                                     | d) Pentammine oxo-nit                  | trocobalt (III) chloride               |
| 842. Point out the central ion ligand in the complex $K_2$                     |  |  |
| a) $Cd^+$ , $CN^{1-}$ b) $Cd^{2+}$ , $CN^{1-}$                                 | c) Cd <sup>2+</sup> , CN <sup>4-</sup> | d) $Cd^{2+}$ , $CN^{2-}$               |
| 843. Number of chiral centres in   |  |  |
|  |  |  |
|  |  |  |
| [ ] is/are   |  |  |
|  |  |  |
| a) 1 b) 2  | c) 3                                   | d) 4                                   |

844. From the equation,  $3C_2H_2 \rightarrow C_6H_6$ , find the volume of acetylene (NTP) for the manufacture of 3 mole of benzene: b) 134.4 litre a) 67.2 litre c) 201.6 litre d) 33.8 litre 845. According to IUPAC nomenclature sodium nitroprusside is named as a) Sodium pentacyanonitrosyl ferrate(II) b) Sodium pentacyanonitrosyl ferrate(III) c) Sodium nitroferricyanide d) Sodium nitroferrocyanide 846. Among  $[Ni(CO)_4]$ ,  $[Ni(CN)_4]^2$  and  $[NiCl_4]^2$  species the hybridisation states of Ni atom are respectively: a)  $sp^3$ ,  $dsp^2$ ,  $dsp^2$ b)  $sp^3$ ,  $dsp^2$ ,  $sp^3$ c)  $sp^3$ ,  $sp^3$ ,  $dsp^2$ d)  $dsp^2$ ,  $sp^3$ ,  $sp^3$ 847. The chemical name of DDT is: a) Dichloro dinitro toluene b) Dichloro dimethyl toluene c) p, p'-dichloro diphenyl trichloroethane d) None of the above 848. The stability of complexes of  $Cu^{2+}$ ,  $Ni^{2+}$ ,  $Co^{2+}$  and  $Fe^{2+}$  varies in the order b)  $Cu^{2+} > Fe^{2+} > Ni^{2+}$ a)  $Cu^{2+} > Ni^{2+} > Co^{2+} > Fe^{2+}$ d)  $Cu^{2+} < Ni^{2+} < Co^{2+} <$ c)  $Ni^{2+} > Co^{2+} > Fe^{2+} > Cu^{2+}$ 849. The number of unpaired electrons in  $Ni(CO)_4$  is a) 0 c) 3 b) 1 850. In sodium tetrafluorooxochromate(....),  $Na_3[Cr(0)F_4]$  the left out place should be filled with which of the following roman numerals? d) None of these a) VI b) III 851. The IUPAC name of compound  $CN - CH_2 - CH - CH_2 - COOCH_3$  is OCH<sub>3</sub> a) 3-methoxy-4-cyano methyl butanoate b) Methyl-4-cyano-3-methoxy butanoate c) 4-cyano-3-methoxy methyl butanoate d) Methyl-3-methoxy-4-cyano butanoate 852. Cumene is: a) *o*-methyl phenol b) *p*-cresol c) Isopropyl benzene d) Phenyl *n*-propane 853. In Etard's reaction toluene is oxidised to benzaldehyde using: a)  $H_2O_2$ b)  $Cl_2$ c) Chromium trioxide or  $CrO_2Cl_2$ d) KMnO<sub>4</sub> 854. Which of the following will exhibit geometrical isomerism? a) Propene b) Butene-2 c) Butene-1 d) 1, 1-dichloro butane 855. Ferrocene is: c)  $Cr(\eta^5 - C_5H_5)_5$  d)  $Os(\eta^5 - C_5H_5)_2$ a)  $Fe(\eta^5 - C_5H_5)_2$ b) Fe( $\eta^2 - C_5 H_5$ )<sub>2</sub> 856. Which one is an outer orbital complex? a)  $[Ni(NH_3)_6]^{2+}$ b)  $[Mn(CN)_{6}]^{4-}$ c)  $[Co(NH_3)_6]^{3+}$ d)  $[Fe(CN)_6]^{4-}$ 857. The pair of  $[Co(SO_4)(NH_3)_5]Cl$  and  $[CoCl(NH_3)_5]SO_4$  constitutes ) Optical isomers b) Linkage isomers c) Coordination isomers d) Ionisation isomers 858. The IUPAC name of  $K_2[Cr(CN)_2O_2(O)_2(NH_3)]$  is a) Potassiumammine dicyanodioxoperoxochromate (VI) b) Potassiumammine cyanoperoxodioxochrometic (IV) c) Potassiumammine dicyanodioxoperoxochromium (IV) d) Potassiumammine dicyanodioxoperoxochromium (IV) 859. In spectrochemical series chlorine is above than water *i.e.*,  $Cl > H_2O$ , this is due to a) Good  $\pi$ -acceptor properties of Cl

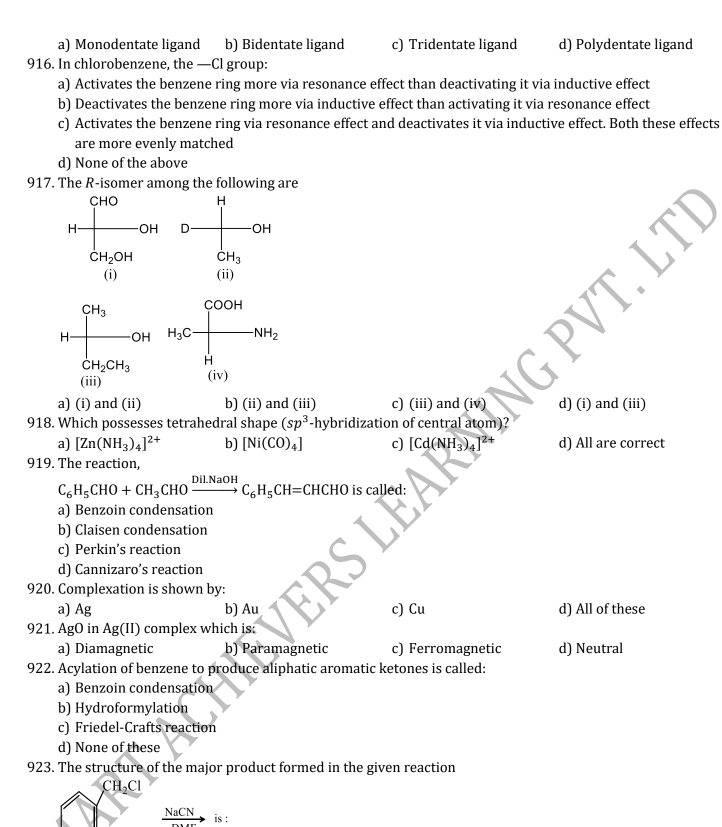
b) Strong  $\sigma$  –donor and good  $\pi$ -acceptor properties of Cl c) Good  $\pi$  –donor properties of Cl d) Larger size of Cl than  $H_2O$ 860. The type of isomerism shown by  $[Co(en)_2(NCS)_2]Cl$  and  $[Co(en)_2(NCS)Cl]NCS$  is: a) Coordination b) Ionization c) Linkage d) All of these 861. Which ion shows only the coordination number 4 in complexes? a) Pt<sup>2+</sup> b) Cr<sup>3+</sup> c) Fe<sup>3+</sup> d) Pt<sup>4+</sup> 862. The spin magnetic moment of cobalt in  $Hg[Co(SCN)_4]$  is : a) √3 b) √8 c)  $\sqrt{15}$ d)  $\sqrt{24}$ 863. Which of the following is not an isomer of but-1-yne? a) But-2-yne b) Buta-1-3-diene c) Methyl cyclopropene d) But-2-ene 864. How many unpaired electrons are present in the central metal ion of [CoCl<sub>4</sub>] b) 3 a) 2 c) 4 0.5 865. The brown ring complex compound is formulated as  $[Fe(H_2O)_5NO]SO_4$ . The oxidation state of Fe is: a) +1 b) +2c) +3 d) Zero 866. Correct IUPAC name of Cl is CCl<sub>3</sub>-CH b) Dichloro diphenyl trichloroethane a) Gammexane c) Diparachlorophenyl trichloroethane d) 1,1,1-tirchloro-2,2-bis (4-chlorophenyl) ethane 867. IUPAC name of is a) Cumene b) 2-phenyl propane c) Phenyl propane d) 1-(2-propyl) benzene 868. Which of the following gives violet colour with an alcoholic solution of FeCl<sub>3</sub>? b) Toluene c) Salicylic acid a) Benzoic acid d) Nitrobenzene 869. Which of the following is wrong statements? a) Ni(CO)<sub>4</sub>, has zero oxidation number for Ni b) Ni(CO)<sub>4</sub>, has oxidation number +4 for Ni c) Ni is metal d) CO is gas 870. Which of the following represents a chelating ligand? b) Cl<sup>-</sup> c) OH<sup>-</sup> d) DMG a)  $H_2O$ 871. The correct order of reactivity of PhMgBr with; O Ph-d -Ph CH<sub>3</sub>-C - CH<sub>2</sub> is: -H CH<sub>3</sub>· (II) (II) CIID a) I > II > IIIb) III > I > II c) II > III > I d) II > I > III872. Which of the following will give maximum number of isomers? b)  $[Ni(en)(NH_3)_4]^{2+}$ a)  $[Co(NH_3)_4Cl_2]$ c)  $[Ni(C_2O_4)(en)_2]$ d)  $[Cr(SCN)_2(NH_3)_4]^+$ 873. CuCl reacts with KCN solution forming a complex. Coordination number of copper in the complex is:

| a) 2 b) 3  | c) 4                          | d) 6   |  |  |
|--|-------------------------------|--|--|--|
| 874. The terms stereoisomers, enantionmers and diaster   | •                             | 2  |  |  |
| a) Only to configurational isomers including geome   | tric isomers                  |  |  |  |
| b) Only to configurational isomers   |                               |  |  |  |
| c) To both configurational as well as conformational isomers   |                               |  |  |  |
| d) To neither configuration nor conformational isor  |                               |  |  |  |
| 875. Aniline was acetylated. The product on nitration fol  |                               | sis gave:  |  |  |
| a) <i>o</i> -nitroacetanilide b) <i>o</i> -and <i>p</i> -nitroaniline  | c) <i>m</i> -nitroaniline     | d) Acetanilide   |  |  |
| 876. The IUPAC name of the compound $[CuCl_2(CH_3NH_2)]$   | •                             | $\sim$   |  |  |
| a) Dichloro bis (dimethyl amine) copper(II)  | b) Dichloro bis (methyl a     | amine) copper(II)  |  |  |
| c) Dimethyl amine copper (II) chloride   | d) Bis (dimethyl amine )      |  |  |  |
| 877. Which is the structure of compound 2-(1-cyclobute   |                               |  |  |  |
| (1 0)000000  | $\wedge$                      |  |  |  |
| $\langle \rangle$  | $\langle \rangle$             |  |  |  |
| a)   | b)                            |  |  |  |
|  |                               | $\langle \rangle$  |  |  |
|  |                               |  |  |  |
|  |                               | <b>&gt;</b>  |  |  |
| $\langle \rangle$  |                               |  |  |  |
| c)   | d)                            |  |  |  |
|  |                               |  |  |  |
|  |                               |  |  |  |
| 878. On explosion TNT gives:   |                               |  |  |  |
| a) $CO + N_2 + H_2 + CH_4 + CO_2$  |                               |  |  |  |
| b) $CO + N_2 + H_2$  | $\mathbf{X}$                  |  |  |  |
| c) $CO_2 + N_2 + H_2O$   | Y                             |  |  |  |
| d) $CO + N_2 + H_2O$   |                               |  |  |  |
| 879. Hexafluoroferrate(III) ion is an outer orbital complete   | ex. The number of unpaire     | d electrons present in it is:  |  |  |
| a) 1 b) 5  | c) 4                          | d) Unpredictable   |  |  |
| 880. The EAN of Fe in $K_3$ [Fe(CN) <sub>6</sub> ] is:   |                               |  |  |  |
| a) 36 b) 37  | c) 38                         | d) 35  |  |  |
| 881. The IUPAC name of the compound  |                               |  |  |  |
| OH   |                               |  |  |  |
|  |                               |  |  |  |
| CH <sub>3 is</sub>   |                               |  |  |  |
| a) 4-methyl cyclopent-1-en-3-ol  | b) 5-methyl cyclopent-2       | -en-1-ol   |  |  |
| c) 2-methyl cyclopent-4-en-1-ol  | d) 3-methyl cyclopent-1       | -en-2-ol   |  |  |
| 882. Which one amongst the following, exhibit geometries   | cal isomerism?                |  |  |  |
| a) [Co <sup>III</sup> (NH <sub>3</sub> ) <sub>5</sub> Br]SO <sub>4</sub> b) Co <sup>III</sup> [EDTA] <sup>1–</sup> | c) $[Cr^{III}(SCN)_{6}]^{3-}$ | d) [Pt <sup>II</sup> (NH <sub>3</sub> ) <sub>2</sub> Cl <sub>2</sub> ] |  |  |
| 883. Chiral molecules are those which are  |                               |  |  |  |
| a) Superimposable on their mirror images   | b) Non-superimposable         | on their mirror images   |  |  |
| c) Unstable molecules  | d) Capable of showing g       | eometrical isomerism   |  |  |
| 884. At room temperature the eclipsed and the staggered  | d forms of ethane cannot b    | e isolated because   |  |  |
| $\checkmark$ a) Both the conformers are equally stable   | b) They interconvent raj      | pidly  |  |  |
| There is a large energy barrier of rotation about  | d) The energy difference      | e between the  |  |  |
| c) the $\sigma$ -bond  | conformers is large           |  |  |  |
| 885. A group of atoms can function as a ligand only when   | n                             |  |  |  |
| a) It is a small molecule  | b) It has an unshared ele     | ectron pair  |  |  |
| c) It is a negatively charged ion  | d) It is a positively charg   | ged ion  |  |  |
| 886. The IUPAC name of $Ni(CO)_4$ is:  |                               |  |  |  |
| a) Tetracarbonyl nickelate(0)  |                               |  |  |  |
|  |                               |  |  |  |

|       | b) Tetracarbonyl nickelat                    |                             |                                |                             |
|-------|--|-----------------------------|--------------------------------|-----------------------------|
|       | c) Tetracarbonyl nickel(0                    |                             |                                |                             |
|       | d) Tetracarbonyl nickel(I                    | I)                          |                                |                             |
| 887   | 7. 2-methyl phenol is:                       |                             |                                |                             |
|       | a) <i>o</i> -cresol                          | b) Catechol                 | c) <i>p</i> -cresol            | d) <i>m</i> -cresol         |
| 888   | 3. $NH_2 \cdot NH_2$ serves as:              |                             |                                |                             |
|       | a) Monodentate ligand                        | b) Chelating ligand         | c) Bridging ligand             | d) Both (a) and (c)         |
| 886   | <ol> <li>For blasting purpose TNT</li> </ol> |                             |                                |                             |
|       | a) NH <sub>4</sub> Cl                        | b) $NH_4NO_3$               | c) $NH_4NO_2$                  | d) $(NH_4)_2SO_4$           |
| 890   | ). During the debromination                  |                             |                                |                             |
|       | a) <i>cis</i> -2-butene                      | b) 1-butene                 | c) <i>n</i> -butane            | d) <i>trans</i> -2-butene   |
| 891   | I. The IUPAC name of $K_2$ [Cr               |                             |                                |                             |
|       | -  | yano dioxoperoxochromate    | e b) Potassium ammine cya      |                             |
|       | (VI)   |                             | peroxodioxochromium            |                             |
|       | c) Potassium ammine cya                      |                             |                                | no peroxodioxochromatic     |
| 007   | peroxodioxochromium                          |                             | (IV)                           |                             |
| 892   | 2. Benzene on reaction with                  |                             | $SO_4$ followed by reaction of | $\Gamma Cl_2/FeCl_3$ gives: |
|       | a) 3-chloro-1-nitrobenzer                    |                             |                                | /                           |
|       | b) 2-chloro-1-nitrobenzer                    |                             |                                |                             |
|       | c) 4-chloro-1-nitrobenzer                    | and 4-chloro-1-nitrobenzer  |                                |                             |
| 803   | 8. The number of isomeric f                  |                             |                                |                             |
| 075   | a) 2   | b) 3                        | () 4                           | d) 1                        |
| 894   | l. Nitration of benzene is:                  | 5 5                         |                                |                             |
| 0,7,1 | a) Nucleophilic substituti                   | on                          | $\mathbf{X}$                   |                             |
|       | b) Electrophilic substituti                  |                             | ) Y                            |                             |
|       | c) Electrophilic addition                    |                             |                                |                             |
|       | d) Nucleophilic addition                     |                             |                                |                             |
| 895   | 5. Reimer-Tiemann reaction                   | n involves a:               |                                |                             |
|       | a) Carbonium ion interme                     |                             |                                |                             |
|       | b) Carbene intermediate                      |                             |                                |                             |
|       | c) Carbanion intermediat                     | e                           |                                |                             |
|       | d) Free radical intermedia                   | ate                         |                                |                             |
| 896   | 5. Which does not have a ca                  | rboxyl group?               |                                |                             |
|       | a) Picric acid                               | b) Ethanoic acid            | c) Aspirin                     | d) Benzoic acid             |
| 897   | 7. In Cannizaro's reaction gi                | ven below:                  |                                |                             |
|       | 2PhCHO <u></u> PhCH2OH                       | +PhCO <sup>°</sup>          |                                |                             |
|       | the slowest step is:                         |                             |                                |                             |
|       | a) The transfer of hydride                   | e to the carbonyl group     |                                |                             |
|       |  | ton from the carboxylic gro | แท                             |                             |
|       | c) The deprotonation of F                    |                             | up                             |                             |
| C     | d) The attack of $:$ $\mathring{O}H$ at the  | —                           |                                |                             |
|       |  |                             |                                |                             |
| 898   | 3. The oxidation state of Ag                 | -                           |                                |                             |
|       | a) Zero                                      | b) +1                       | c) +2                          | d) +1.5                     |
| 899   | ). Hybridization of Fe in $[K_3$             |                             |                                |                             |
|       | a) $sp^3$                                    | b) $d^2sp^3$                | c) $sp^3d^2$                   | d) $dsp^3$                  |
| 900   | ). Which of the following is                 | not isomeric with diethyl e |                                |                             |
|       | a) Methyl <i>n</i> -propyl ether             |                             | b) Butan-1-ol                  |                             |
|       | c) 2-methyl propan-2-ol                      |                             | d) Butan-2-one                 |                             |
|       |  |                             |                                |                             |

901. In the given conformation  $C_2$  is rotated about  $C_2 - C_3$  bond anticlockwise by an angle of 120° then the conformation obtained is

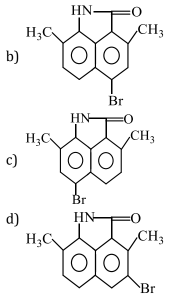
| comormation obtained is   |   |  |
|---|---|--|
| C <sub>4</sub><br>CH <sub>3</sub>   |   |  |
| H, H  |   |  |
| $C_2$   |   |  |
| H H   |   |  |
| ĊH <sub>3</sub>   |   |  |
| C <sub>1</sub>  |   | $\frown$   |
| a) Fully eclipsed conformation  | b) Partially eclipsed con                                 | formation  |
| c) Gauche conformation  | d) Staggered conformati                                   | on   |
| 902. Crystal field stabilization energy for high spin $d^4$ oc                                      | ctahedral complex is:                                     |  |
| a) $-1.8 \Delta_0$ b) $-1.6 \Delta_0 + P$   | c) −1.2 Δ <sub>0</sub>                                    | d) $-0.6 \Delta_0$                               |
| 903. Which kind of isomerism is exhibited by octahedral   | l [Co(NH <sub>3</sub> ) <sub>4</sub> Br <sub>2</sub> ]Cl? |  |
| a) Geometrical and ionisation   |   |  |
| b) Geometrical and optical  |   | $\langle \rangle$                                |
| c) Optical and ionisation   | Ċ   |  |
| d) Geometrical only   |   |  |
| 904. The IUPAC name of the following compound is  |   |  |
| H <sub>2</sub> C  |   |  |
|   |   |  |
| a) 5-cyclopropyl pent-2-en-1-oic acid   | b) 6-cyclopropyl pent-2-                                  | en-1-oic acid                                    |
| c) 5-cyclopropyl pent-1-en carboxylic acid  | d) 6-cyclopropyl pent-1-                                  |  |
| 905. Which of the following compounds will show a nega  |   | -  |
| a) Glucose b) Ethyl alcohol   | c) A cetaldehyde  | d) Benzophenone                                  |
| 906. Friedel-Craft's reaction is not possible in:   | ejneetalaenyae  | aj benzopnenone                                  |
| a) $C_6H_5OH$ b) $C_6H_5C_2H_5$   | c) C <sub>6</sub> H <sub>5</sub> NO <sub>2</sub>          | d) C <sub>6</sub> H <sub>5</sub> CH <sub>3</sub> |
| 907. The geometry of Ni(CO) <sub>4</sub> and Ni(PPh <sub>3</sub> ) <sub>2</sub> Cl <sub>2</sub> are | 0) 061151102  | 4) 661156113                                     |
| a) Both square planar   | b) Tetrahedral and squa                                   | re planar respectively                           |
| c) Both tetrahedral   | d) Square planar and tet                                  |  |
| 908. The number of isomers possible for square planar c   |   | · ·  |
| a) 2 b) 3   | c) 4  | d) 6   |
| 909. The correct order for the wavelength of absorption   | in the visible region is                                  |  |
| a) $[Ni(NO_2)_6]^{4-} < [Ni(NH_3)_6]^{2+} < [Ni(H_2O)_6]^{2+}$                                      |   | $[I_20]_6]^{2+} < [Ni(NO_2)_6]^{4-}$             |
| c) $[Ni(H_2O)_6]^{2+} < [Ni(NH_3)_6]^{2+} < [Ni(NO_2)_6]^{4-}$                                      |   | $[I_20)_6]^{2+} < [Ni(NH_3)_6]^{2+}$             |
| 910. The IUPAC name of CCl <sub>3</sub> CH <sub>2</sub> CHO is                                      |   |  |
| a) Chloral  | b) 1,1,1-trichloropropan                                  | ol   |
| c) 2,2,2-trichloropropanol  | d) 3,3,3-trichloropropan                                  | ol   |
| 911. The coordination number of Cu in $[Cu(H_2O)_4]^{2+}$ cou                                       | mplex is  |  |
| a) 2 b) 1   | c) 3  | d) 4   |
| 912. Among the following, the correct statement is  |   |  |
| a) Prefixes are written before the name of compour  | nd  |  |
| $igsirclevel{b}$ b) Suffixes are written after the name of compound                                 |   |  |
| c) The IUPAC name is always written as a single wo  | ord   |  |
| d) All of the above   |   |  |
| 913. In which of the following <i>p</i> -electrons of the halogen                                   |   |  |
| a) Chlorobenzene b) Bromobenzene  | c) Allyl chloride   | d) Vinyl chloride                                |
| 914. Which of the following does not have optical isomer  |   |  |
| a) $[Co(en)(NH_3)_2Cl_2]Cl$ b) $[Co(en)_2Cl_2]Cl$   | c) $[Co(NH_3)_3Cl]$                                       | d) $[Co(en)_3]Cl_3$                              |
| 915. Ethylene diamine is an example of  |   |  |
|   |   |  |



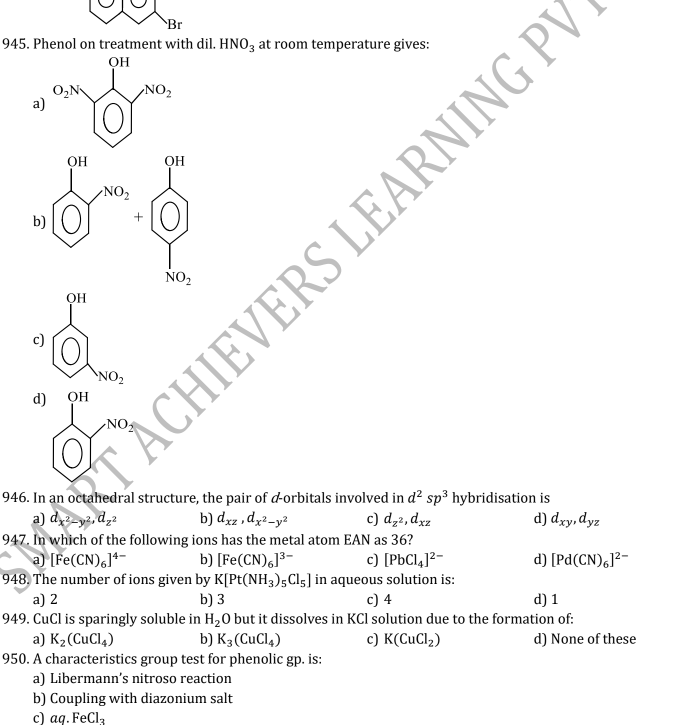
CH<sub>2</sub>CN

CH<sub>2</sub>Cl b)  $CH_2Cl$ c) CH<sub>2</sub>CN d) 924. Chlorobenzene is prepared commercially by: a) Grignard reaction b) Raschig process c) Wurtz - Fittig reaction d) Friedel-Crafts reaction 925. An aqueous solution of CoCl<sub>2</sub> on addition of excess of concentrated HCl turns blue to formation of a) [CoCl<sub>4</sub>]<sup>2–</sup> c)  $[Co(H_2O)_22Cl_4]^{2-}$ b)  $[Co(H_2O)_2Cl_4]^{2-}$ d)  $[Co(H_2O)_4Cl_2]$ 926. Which one of the following will not show geometrical isomerism? a)  $[Cr(NH_3)_4Cl_2]Cl$ b)  $[Co(en)_2Cl_2]Cl$ c)  $[Co(NH_3)_5NO_2]Cl_2$ d)  $[Pt(NH_3)_2Cl_2]$ 927. When ethyl benzoate is hydrolysed with aqueous alkali, the products present in the medium are: c)  $C_2H_5OH$ ,  $C_6H_5COOH$ a)  $C_6H_5COOH, C_2H_5O^$ b)  $C_6H_5COO^-, C_6H_5OH$ d)  $C_6H_5COO^-, C_2H_5O^-$ 928. The IUPAC name of CONH<sub>2</sub> .СНО a) 2-carbamovl hexanal b) 2-carbamoyl hex-3-en-1-al c) 6-keto-2-methylhexanamide d) 5-formyl-2-methylpent-3-en-1-amide 929. Which of the following is more basic than aniline? b) Benzylamine a) *p*-Nitroaniline c) Diphenylamine d) Triphenylamine 930. Name of some compounds are given below. Which one is not in IUPAC system?  $CH_3 - CH - CH - CH_3$ b)  $\frac{CH_3 - C \equiv C - CH(CH_3)_2}{4 \text{ methyl-2-pentyne}}$ a) OH CH-4-methyl-2-butanol CH<sub>3</sub> d)  $CH_3 - CH_2 - CH_2 - CH - CH - CH_2CH_3$ ethyl-3- methyl - but -1- ene CH<sub>2</sub>CH<sub>3</sub> 3-methyl-4-ethyl heptane 931. For which transition metal ions are low spin complexes possible? a) Rh<sup>3+</sup> b) Mn<sup>3+</sup> c) Ru<sup>2+</sup> d) All are correct 932. Which one is monodentate ligand? a) F<sup>-</sup> b)  $NO_{2}^{-}$ c)  $H_20$ d) All are correct 933. Cyclic hydrocarbon molecule A has all the carbons and hydrogens in a single plane. All the carbon-carbon bonds are of same length and less than 1.54 Å and more than 1.34 Å. The C—C—C bond angle will be: a) 120° b) 180° c) 100° d) 109°28' 934. Chlorine reacts with benzaldehyde to give:

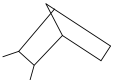
| a) Benzyl chloride  | b) Benzal chloride                     | c) Benzoyl chloride                      | d) Chlorobenzene      |
|---|--|--|-----------------------|
| 935. Phenol is:   | b) belizai cilioriue                   | cj benzoyi chioride                      | u) chiorobenzene      |
| a) A base weaker than N   | IH.                                    |  |                       |
| b) An acid stronger than  | -                                      |  |                       |
| c) An acid weaker than  |  |  |                       |
| d) Neutral  |  |  |                       |
| 936. Which one is example of  | f octahedral complex?                  |  |                       |
| a) $Cu(NH_3)_4^{2+}$  | b) $FeF_6^{3-}$                        | c) $Zn(NH_3)_4^{2+}$                     | d) Ni(CN) $_{4}^{2-}$ |
| 937. Which one of the follow  | -                                      | ·)(3)4                                   |                       |
|   |  | ling potassium ferrocyanid               | e solution.           |
|   |  | $\frac{1}{3}$ ions, we get a precipitate |                       |
|   | a violet vortex test in redu           |  |                       |
| d) From a mixed precipi   | tate of AgCl and AgI, amm              | onia solution dissolves only             | y AgCl                |
| 938. Which of the following f   | ractions obtained in fracti            | onal distillation of coal-tar            | contains benzene and  |
| toluene?  |  |  |                       |
| a) Light oil  |  |  | X                     |
| b) Heavy oil  |  | . (                                      | *                     |
| c) Middle oil   |  |  | <b>&gt;</b>           |
| d) Green oil  |  |  |                       |
| 939. The tetrahedral complex  |  | nber                                     |                       |
| a) 3  | b) 6                                   | c) 4                                     | d) 8                  |
| 940. The C—C bond length in   |  |  |                       |
| a) Less   | b) More                                | c) Equal                                 | d) None of these      |
| 941. Which are generally use  |  | e of aldehydes and ketones?              |                       |
| a) Hydroxylamine hydro  |  |  |                       |
| b) 2,4-dinitrophenylhyd   |  |  |                       |
| c) Phenylhydrazinehydr<br>d) All of the shows   | rochloride                             |  |                       |
| d) All of the above<br>942. In the reaction,  |  |  |                       |
|   |  |  |                       |
| Phenol $\xrightarrow{Zn}$ $(A) \xrightarrow{Conc. H_2SO_4}$<br>$\xrightarrow{Conc. HNO_3 at 60^\circ}$  | $\sim C^{\rightarrow}(B)$              |  |                       |
|   | $(C) \leftarrow \frac{Zn}{N_2 OH(an)}$ |  |                       |
| The compounds $(A)$ $(B)$   | ) and ( <i>C</i> ) are the following   | ינ                                       |                       |
| a) Benzene, nitrobenzer   |  | 2.                                       |                       |
| b) Benzene, dinitrobenz   |  |  |                       |
| c) Toluene, <i>m</i> -nitrobenz   |  |  |                       |
| d) Benzene, nitrobenzer   |  |  |                       |
| 943. En is an example of a:   |  |  |                       |
| a) Monodentate ligand   | b) Bidentate ligand                    | c) Tridentate ligand                     | d) Hexadentate ligand |
| 944. The major product obtain   | ined when Br <sub>2</sub> /Fe is treat | ed with                                  |                       |
| $HC$ $HN$ $O$ $CH_3$  |  |  |                       |
|   |  |  |                       |
| $\mathbf{O}$ $\mathbf{O}$ is:   |  |  |                       |
|   |  |  |                       |
| H <sub>2</sub> C <sub>2</sub> $\downarrow$ |  |  |                       |
| a) H <sub>3</sub> C   | - 3                                    |  |                       |
| Br  |  |  |                       |
| ы, 🔨 🔨  |  |  |                       |



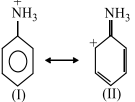
945. Phenol on treatment with dil. HNO<sub>3</sub> at room temperature gives:



#### d) All of the above 951. Write the IUPAC name of the compound



b) 2, 3-dimethyl bicyclo [2,2,1] heptane a) 5, 6-dimethyl bicyclo [2,2,1] heptane c) 2, 3-dimethyl bicyclo [1,2,2] heptane d) 3, 4-dimethyl bicyclo [2,1,2] heptane 952. Choose the correct statement from the ones given below for two anilium in:



a) II is not an acceptable canonical structure because carbonium ions are less stable than ammonium ions

- b) II is not an acceptable canonical structure because it is non-aromatic
- c) II is not an acceptable canonical structure because the nitrogen has 10 valence electrons
- d) II is an acceptable canonical structure
- 953. Which of the following statements is/are incorrect?
  - a) Metamerism belongs to the category of structural isomerism
  - b) Tautomeric structures are the resonating structures of a molecule
  - c) The violet colouration produce by a molecule with neutral ferric chloride solution indicates the presence of enolic group in the molecule
  - d) Geometrical isomerism is not shown by alkenes
- 954. Gives are (i) cyclohexanol; (ii) acetic acid; (iii) 2, 4, 6-trinitrophenol; and (iv) phenol. In these the order of decreasing acidic character will be:
- c) (ii) > (iii) > (iv) > (i)a) (iii) > (ii) > (iv) > (i)b) (ii)>(iii)>(i)>(iv) d) (iii)>(iv)>(ii)>(i) 955. Phenol and benzoic acid can be distinguished by:

956. The functional groups – OH, –COOH, –CHO, –OCH<sub>3</sub> attached to a chiral carbon is in the preference order a) OH > COOH > CHO > OCH<sub>3</sub> b

$$0 \text{ OCH}_3 > 0 \text{H} > C 0 \text{OH} > C \text{H} 0$$

$$OCH_3 > OH > CHO > COOH$$

d) Conc.  $H_2SO_4$ 

$$(J) UCH_3 > COUH > CHU > OH$$

c) Aqueous NaOH

957. The hypothetical complex chloro diaquatriammine cobalt(II) chloride can be represented as:

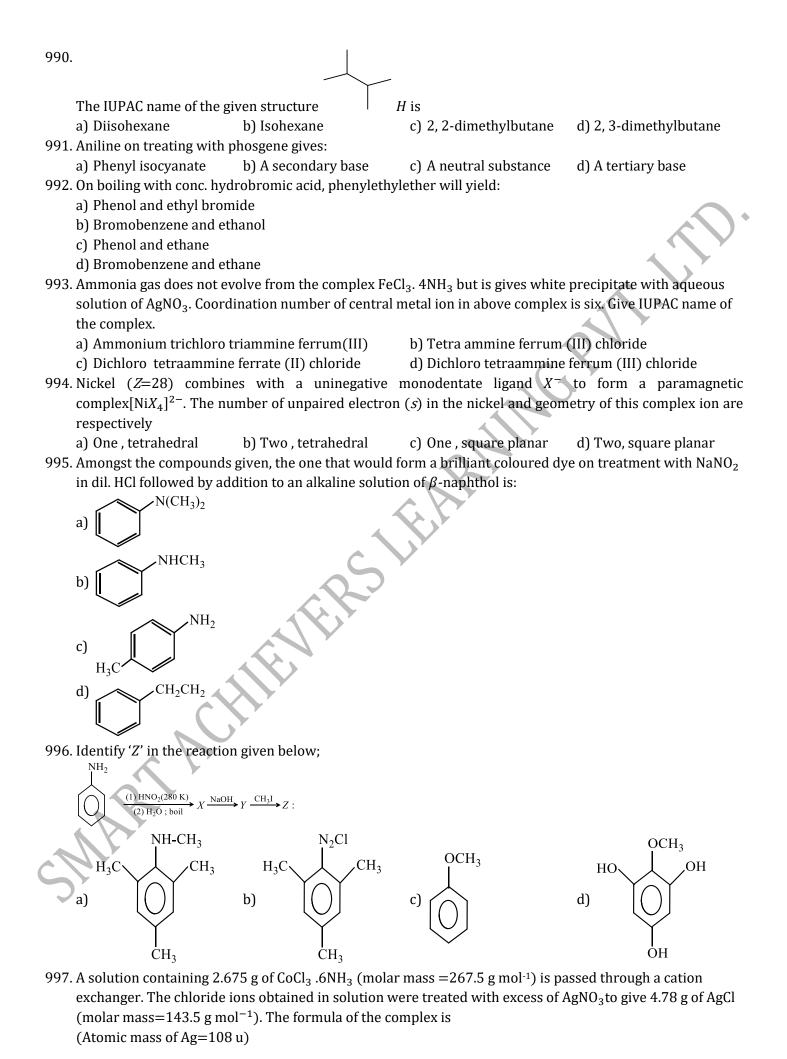
- a)  $[CoCl(NH_3)_3(H_2O)_2]Cl_2b) [Co(NH_3)_3(H_2O)Cl_3]$ c)  $[Co(NH_3)_3(H_2O)_2Cl]$ d)  $[Co(NH_3)_3(H_2O)_3]Cl_3$ 958. Which is expected to be paramagnetic?
- a)  $[Ni(H_20)_6]^{2+}$ c)  $[Zn(NH_3)_4]^{2+}$ d)  $[Co(NH_3)_6]^{3+}$ b)  $[Ni(CO_4)]$ 959. The molecular formula of diphenyl methane

$$\bigcirc$$
 CH<sub>2</sub>  $\bigcirc$  is C<sub>13</sub>H<sub>12</sub>

a) *A, B* b) *B, C* c) *C, A* d) A, B, C 961. the double bonds are In

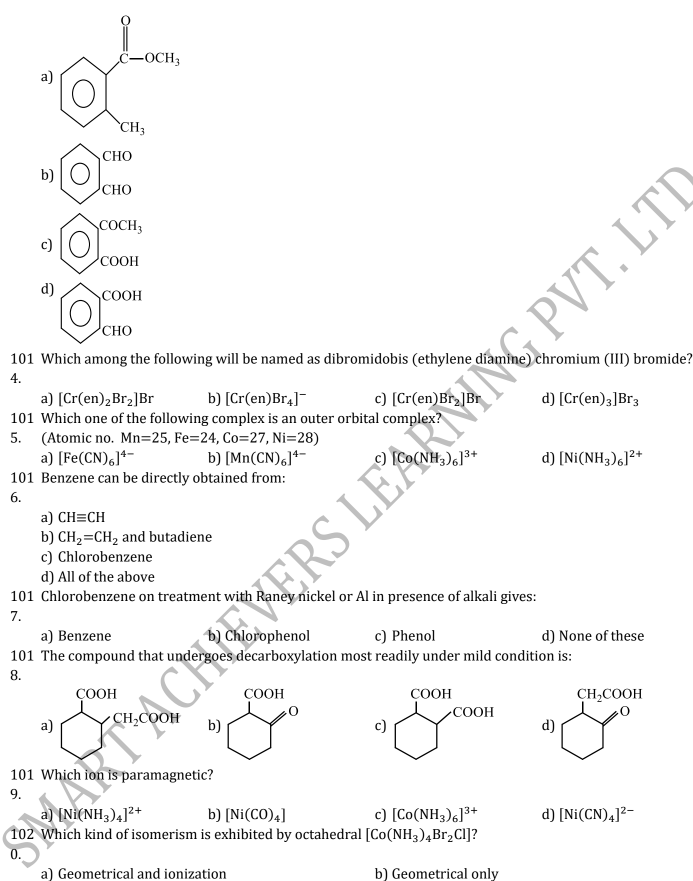
a) cis, cis b) cis, trans c) trans, cis d) trans, trans 962. The reaction of toluene with Cl<sub>2</sub> in presence of FeCl<sub>3</sub> gives 'X' and the reaction in presence of light gives '*Y*'.Thus, '*X*' and '*Y*' are: a) X=benzal chloride; Y = o-chlorotoluene b) X = m-chlorotoluene; Y = p-chlorotoluene c) X = o-and p-chlorotouene; Y=trichloro methyl benzene d) X = benzal chloride; Y = m-chlorotoluene 963. Among the following four compounds: a) Phenol b) Methyl phenol c) *meta*-nitrophenol d) para-nitrophenol 964. Which gives phthalic anhydride on reaction with hot, conc.  $H_2SO_4$  in presence of Hg? b) Phenol a) Naphthalene c) *p*-xylene d) *m*-xylene 965. *Cis-trans*-isomerism is found in square planar complexes of the molecular formula: (*a* and *b* are monodentate ligands) a) Ma<sub>4</sub> d) Mab b)  $Ma_3b$ c)  $Ma_2b_2$ 966. Which ion produces a small crystal field splitting (a weak ligand field)? a) I<sup>-</sup> b) Cl<sup>-</sup> d) All of these c) F<sup>-</sup> 967. Benzene undergoes substitution reaction more easily than addition because: a) It has a cyclic structure b) It has three double bonds c) It has six hydrogen atoms d) Of resonance 968. Isomers have essentially identical a) Structural formula b) Chemical properties c) Physical properties d) Molecular formula 969. Which of the following pair is not correctly matched? a) Absorption peak for  $[Cr^{III}(NH_3)_6]^{3+} = 21680 \text{ cm}^{-1}$ b) Effective atomic no. of Pt in  $[PtCl_6]^{2-} = 84$ c) Crystal field stabilization energy of  $d^2$  in weak ligand field =  $(-)0.8 \Delta_0$ d) Example of weak ligand field for  $d^5$  configuration =  $[Mn^{II}F_6]^{4-}$ 970. Aspirin (or acetyl salicylic acid) is obtained by action of CH<sub>3</sub>COCl with: a) Salicylic acid b) Phenol c) Benzaldehvde d) Aniline 971. CuCl dissolves in ammonia forming a complex. The coordination number of copper in the complex is: b) 2 c) 4 d) 6 a) 1 972. IUPAC name of the following cycloalkane is CH<sub>3</sub> a) 8-methyl bicyclo [4,3,0] nonane b) 1-methyl bicyclo [4,3,0] nonane c) 3-methyl bicyclo [4,3,0] nonane d) 4-methyl bicyclo [4,3,0] nonane 973. Schiff's bases are formed when aniline is condensed with: a) Phenols b) Aromatic aldehydes c) Aryl chlorides d) Aliphatic alcohols 974. Which of the following is not an organometallic compound? a) Zeise' s salt c) Sodium ethoxide d) Ferrocene b) TEL 975. Molecular formula C<sub>5</sub>H<sub>12</sub>O will show a) Position b) Optical isomerism c) Functional isomerism d) All of these 976. Both Co<sup>3+</sup> and Pt<sup>4+</sup>have a coordination number of six. Which of the following pairs of complexes will show approximately the same electrical conductance for their 0.001 M aqueous solutions? b) CoCl<sub>3</sub>. 3NH<sub>3</sub> and PtCl<sub>4</sub> .5NH<sub>3</sub> a) CoCl<sub>2</sub>. 4NH<sub>3</sub> and PtCl<sub>4</sub>. 4NH<sub>3</sub> c) CoCl<sub>3</sub>. 6NH<sub>3</sub> and PtCl<sub>4</sub> .5NH<sub>3</sub> d) CoCl<sub>3</sub>. 6NH<sub>3</sub> and PtCl<sub>4</sub> .3NH<sub>3</sub>

977. Which of the following is not an organometallic compound? a) Sodium ethoxide b) Trimethyl aluminium c) Tetraethyl lead d) Ethyl magnesium bromide 978. The number of water molecule(s) directly bonded to the metal centre in  $CuSO_4$ .  $SH_2O$  is b) 2 d) 4 a) 1 c) 3 979. The formula of sodium nitroprusside is: a)  $Na_4[Fe(CN)_5NO_2]$ b)  $Na_2[Fe(CN)_5NO]$ c) NaFe[Fe(CN)<sub>6</sub>] d)  $Na_2[Fe(CN)_6NO_2]$ 980. The IUPAC name of the compound  $H_2N - CH - CH_2OH$  is COOH a) 2-amino-2-carboxy pentanol b) 1-amino-2-hydroxy propanoic acid c) 1-hydroxy-2-amino-3-propanoic acid d) 2-amino-3-hydroxy propanoic acid 981. Which of the following complex species does not involve inner orbital hybridisation? a)  $[CoF_6]^{3-1}$ b)  $[Co(NH_3)_6]^{3+}$ c)  $[Fe(CN)_6]^{3-1}$ d)  $[Cr(NH_3)_6]^{3+}$ 982. The EAN of nickel in  $K_2[Ni(CN)_4]$  is: d) 38 a) 35 b) 34 c) 36 983. The type of isomerism shown by, 6,6'-disitrodiphenic acid is b) Optical c) Geometrical a) Conformational d) Functional 984. Which one of the following compounds forms benzoic acid on oxidation? c) Chlorobenzene a) Chlorophenol b) Benzylchloride d) Chlorotoluene 985. Glycinato ligand is: a) CH<sub>2</sub> COO b) Bidentate ligand c) Two donor sites N and O<sup>-</sup> d) All of the above 986. Which one is the most likely structure of  $CrCl_3 \cdot 6H_2O$ , if 1/3 of total chlorine of the compound is precipitated by adding AgNO<sub>3</sub> to its aqueous solution? a)  $CrCl_3 \cdot 6H_2O$ b)  $[Cr(H_2O)_3Cl_3] \cdot (H_2O)_3$ c)  $[CrCl_2(H_2O)_4] \cdot Cl \cdot 2H_2O$ d)  $[CrCl(H_2O)_5]Cl_2 \cdot H_2O$ 987. Carbon in benzene undergoes  $sp^2$ -hybridization and the bond angle is 120°. The shape of benzene molecule is: a) Linear b) Planar d) Planar hexagonal c) Pyramidal 988. The example of coordination isomerism is  $[Co(NH_3)_6]$   $[Cr(CN)_6]$  and b)  $[Co(NH_3)_5Br]SO_4$  and  $[Co(NH_3)_5SO_4]Br$  $[Cr(NH_3)_6] [Co(CN)_6]$ c)  $Co(NH_3)_5NO_3$  SO<sub>4</sub> and  $[Co(NH_3)_5SO_4]NO_3$ d)  $[Pt(NH_3)_4Cl_2]Br_2$  and  $[Pt(NH_3)_4Br_2]Cl_2$ 989. Coordination compounds have great importance in biological systems. In this context which of the following statement is incorrect? a) Haemoglobin is the red pigment of blood and contains iron b) Cyanocobalamin is B<sub>12</sub> and contains cobalt c) Chlorophylls are green pigments in plants and contains calcium d) Carbocypeptidase-A an enzyme and contains zinc



a)  $[Co(NH_3)_6]Cl_3$ b)  $[CoCl_2(NH_3)_4]Cl$ c)  $[CoCl_3(NH_3)_3]$ d)  $[CoCl(NH_3)_5]Cl_2$ 998.  $[Cr(H_2O)_6]^{3+}$  ion has *d*-electrons equal to: a) 2 b) 3 c) 4 d) 5 999. Enol form is more stable in a) CH<sub>3</sub>CHO b) CH<sub>3</sub>COCH<sub>3</sub> c) CH<sub>3</sub>COCH<sub>2</sub>COOC<sub>2</sub>H<sub>5</sub> d) Cyclohexanone 100 The coordination number of cobalt in  $[Co(en)_2Br_2]Cl_2$  is: 0. a) 2 b) 4 d) 8 c) 6 100 Which one readily accepts a proton? 1. a) Acetylene c) Aniline b) Nitrobenzene d) Phenol 100 Identify '*Z*' in the reaction; 2.  $CH_2 - OH$ RMMCR  $\frac{\text{Vigorous}}{\text{oxidation}} X \xrightarrow{\text{Dry}} Z$ CH<sub>2</sub>-OH СООН a) COOH СООН b) .CO c) d) COOH CH<sub>2</sub>OH 100 The number of  $\sigma$  and  $\pi$ -bonds in a molecule of benzene is: 3. a)  $6\sigma$  and  $9\pi$ b)  $9\sigma$  and  $3\pi$ c)  $12\sigma$  and  $3\pi$ d)  $6\sigma$  and  $6\pi$ 100 The phenomenon of optical activity will be shown by: 4. b) c) en d) R D À

| 100<br>5  | CH <sub>3</sub><br>H <sub>3</sub> C  |
|-----------|--|
| 5.        |  |
|           | The correct name of the compound $CH_3$ is   |
|           | The correct name of the compound CH <sub>3</sub> is<br>a) 1,3,4-trimethyldecaline b) 1,3,9-trimethyldecaline   |
|           | c) 1,8,10-trimethyldecaline d) 1,3,10-trimethyldecaline  |
|           | If NH <sub>4</sub> OH is added to the (PtCl <sub>4</sub> ) <sup>2–</sup> ion, the complex formed represents:   |
| 6.        |  |
| 100       | a) Zero dipoleb) Finite dipolec) Infinite dipoled) All of theseWhich one of the following will be able to show <i>cis-trans</i> -isomerism?d) All of these |
| 7.        |  |
|           | a) $M_{A_{3}B}$  |
|           | b) $M_{(AA')_2}$   |
|           | c) $M_{A_2BCD}$  |
|           | d) $\frac{M_{A_4}}{(AA' \text{ is unsymmetrical bidentate ligand, } ABCD \text{ are unidentate ligands.})}$  |
| 100       | The coordination number of a metal in coordination compound is   |
| 8.        |  |
|           | a) Same as primary valency b) Sum of primary and secondary valencies   |
| 100       | c) Same as secondary valency d) None of the above<br>The IUPAC name of $K_4[Ni(CN)_4]$ is  |
| 9.        | The for Ac hance of K4[M(Ch)4] is  |
|           | a) Tetrapotassium tetracyanonickelate (II) b) Potassium tetracyanonickel (II)  |
| 101       | c) Potassium tetracyanonickelate (0) d) Potassium tetracyanonickelate (II)   |
| 101<br>0. | Which of the following compounds shows optical isomerism?  |
|           | a) $[Co(CN)_6]^{3-}$ b) $[Cr(C_2O_4)_3]^{3-}$ c) $[ZnCl_4]^{2-}$ d) $[Cu(NH_3)_4]^{2+}$  |
| 101       | $[C_6H_5]_2Pd(SCN)_2]$ and $[(C_6H_5)_2Pd(NCS)_2]$ are:  |
| 1.        | a) Linkage isomers b) Coordination isomers c) Ionization isomers d) Geometrical isomers  |
| 101       | Mark the correct statement   |
| 2.        |  |
|           | a) Ethane has two conformations of which staggered conformation is more stable than the eclipsed   |
|           | conformation<br>b) Ethane has an infinite number of conformations of which eclipsed conformation is more stable than the                                   |
|           | staggered conformation   |
|           | c) Ethane has an infinite number of conformation of which staggered conformation has the maximum   |
|           | energy<br>d) Ethane has an infinite number of conformation of which the staggered conformation is possessed by   |
|           | majority of the molecules at room temperature  |
| 101       |  |
| 3.        |  |
|           | Lactone Can be obtained by which   |
|           |  |
|           | Of the following on heating with alkali followed with  |
|           | acid hydrolysis?   |

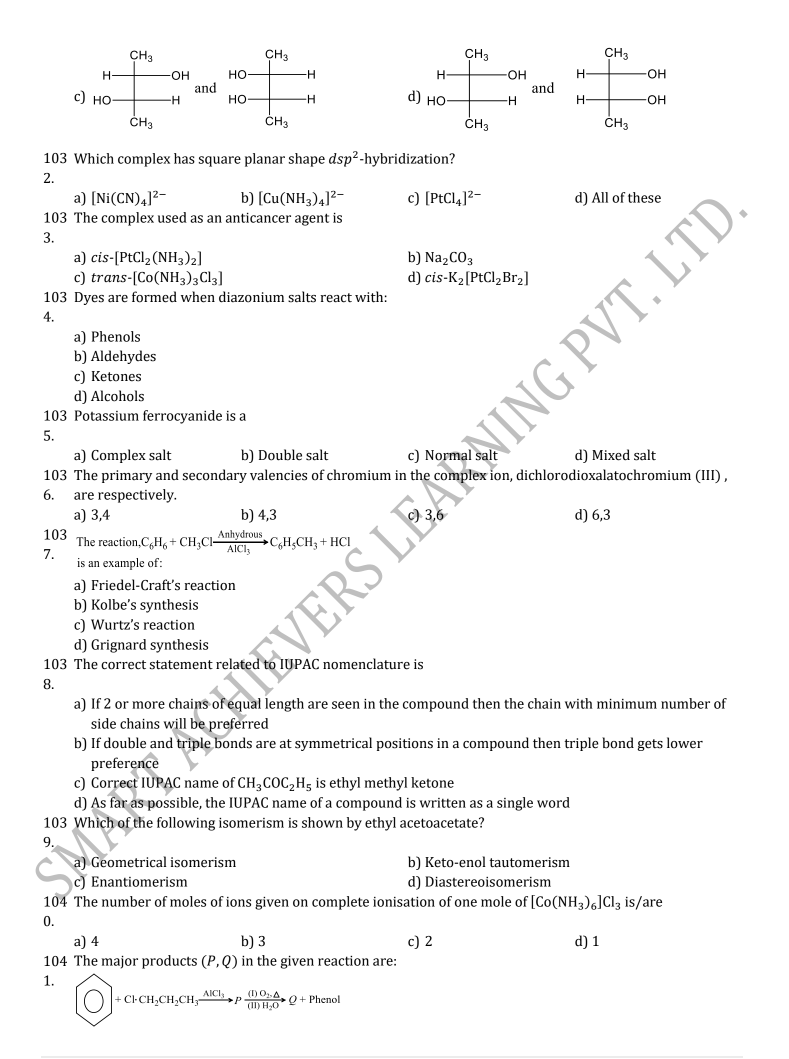


c) Geometrical and optical d) Optical and ionisation

102 Resorcinol and conc.  $H_2SO_4$  in presence of phthalic anhydride produce a compound which is: 1.

a) A dyeb) An antisepticc) An indicatord) A detergent102Which of the following compounds shows optical isomerism?2.

c)  $[Co(CN)_6]^{3-}$ a)  $[Cr(C_2O_4)_3]^{3-}$ b)  $[Cu(NH_3)_4]^{2+}$ d)  $[ZnCl_4]^{2-}$ 102 The IUPAC name of  $[Co(NH_3)_6]Cl_3$  is 3. a) Hexamine cobalt (II) chloride b) Triammine cobalt (III) trichloride c) Hexamine cobalt (III) chloride d) None of the above 102 In the following compounds, the order of acidity is: OH 4. OH OH OH (II) (III) (IV) (I) a) III > IV > I > IIb) I > IV > III > IIc) II > I > III > IV d) IV > III >102 Consider the following structure and choose the correct statements 5.  $NH_2$ Ш b) I and III have R-configuration a) I and II have R-configuration d) Both (a) and (c) are correct c) Only III has S-configuration 102 Benzaldehyde, when heated with concentrated KOH solution, gives: 6. a) C<sub>6</sub>H<sub>5</sub>CH<sub>2</sub>OH b) C<sub>6</sub>H<sub>5</sub>COOH c)  $C_6H_5COOK$ d) Mixture of C<sub>6</sub>H<sub>5</sub>COOK and C<sub>6</sub>H<sub>5</sub>CH<sub>2</sub>OH 102 Write the IUPAC name of the compound 0、 7. a) Bicyclo-[2.2.2] octane-2,6-dione b) Bicyclo-[2.2.2] octane-3,5-dione c) Bicyclo -[2.2] octane 2,6-dione d) Bicyclo [2,2] octane-3,5-dione 102 3-chloro-4-methyl benzene sulphonic acid on steam distillation gives: 8. *m*-chloro benzene a) Toluene c) p-methyl benzene sulphonic acid b) sulphonic acid d) o-chloro toluene 102 The oxidation number of platinum in  $[Pt(NH_3)_5Cl]Cl_3$  is 9. a) 2 b) 3 c) 4 d) 6 103 Which of the following is not an organometallic compound? 0. a)  $C_2H_5ONa$ b) CH<sub>3</sub>Mgl c) Tetraethyl tin d)  $KC_4H_9$ 103 Which of the following pairs of compounds are enantiomers? 1. CH<sub>3</sub> н — ОН НО — b) <sub>HO</sub> — H <sup>A</sup> H— H HO H H and HO H HO-CHa



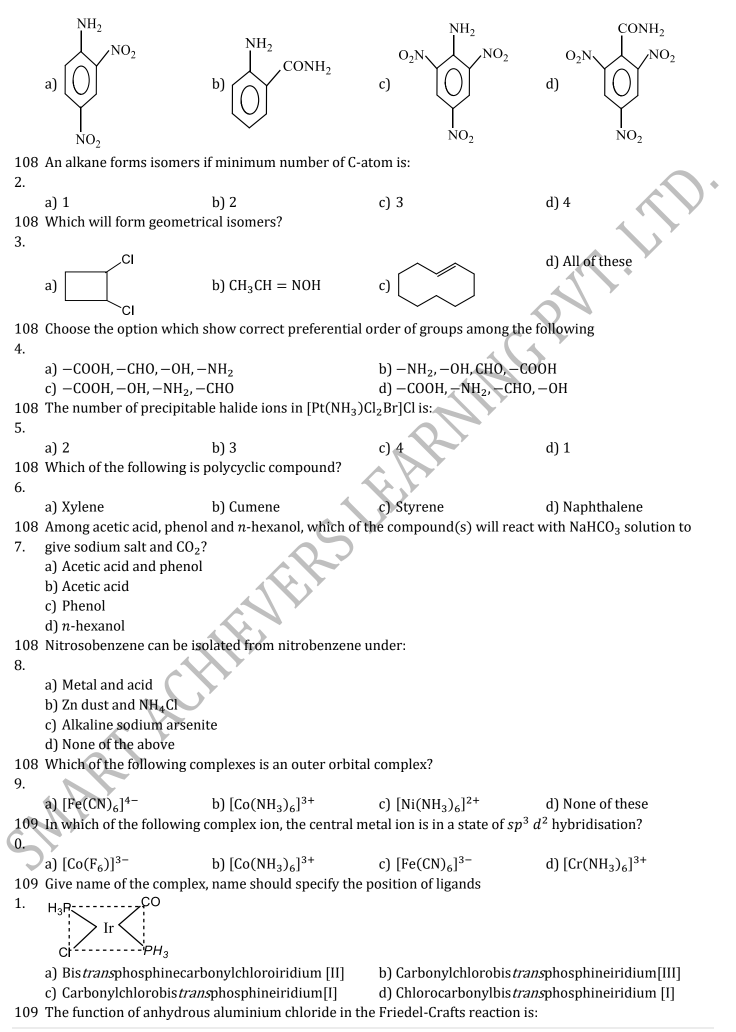
0. a) The complex is high spin complex b) Both Fe atoms are in the same oxidation state c) The coordination number of iron is 4 d) Both Fe atoms are in different oxidation state 105 The number of chiral carbon atoms present in the molecule 1. is c) 2 d) 1 a) 3 b) 4 105 The complex that doesn't give a precipitate with AgNO<sub>3</sub> solution 2. a)  $[Co(NH_3)_33Cl_3]$ b)  $[Co(NH_3)_6]Cl_3$ c)  $[Ag(NH_3)_2]Cl$ d)  $[Cr(NH_3)]$ 105 The IUPAC name of the given compound  $[Co(NH_3)_5Cl]Cl_2$  is 3. b) Cobalt penta ammine chloro chloride a) Penta amino cobalt chloride chlorate c) Pentamine chloro cobalt (III) chloride. d) Penta amino cobalt (III) chlorate 105 Amongst Ni(CO)<sub>4</sub>,  $[Ni(CN)_4]^{2-}$  and  $[NiCl_4]^{2-}$ 4. a) Ni(CO)<sub>4</sub> is diamagnetic,  $[NiCl_4]^{2-}$  and  $[Ni(CN)_4]^{2-}$  are paramagnetic b) Ni(CO)<sub>4</sub> and  $[NiCl_4]^{2-}$  are diamagnetic and  $[Ni(CN)_4]^{2-}$  is paramagnetic c) Ni(CO)<sub>4</sub> and  $[Ni(CN)_4]^{2-}$  are diamagnetic and  $[NiCl_4]^{2-}$  is paramagnetic d)  $[NiCl_4]^{2-}$  and  $[Ni(CN)_4]^{2-}$  are diamagnetic and  $Ni(CO)_4$  is paramagnetic 105 Which aromatic acid among the following is weaker than simple benzoic acid? 5. NO<sub>2</sub>  $CH_2$ SO<sub>3</sub>H a) b) COOH СООН CO<sub>2</sub>H 105 Which statement is incorrect? 6. a) Ni(CO)<sub>4</sub>-tetrahedral, paramagnetic b) [Ni(CN)<sub>4</sub>]<sup>2–</sup>-square planar, diamagnetic c) Ni(CO)<sub>4</sub>-tetrahedral, diamagnetic d)  $[NiCl_4]^{2-}$  -tetrahedral, paramagnetic 105 Which of the following has asymmetric C-atom? 7. H Cl H Cl Η D b) H - C - C - Clc) H − C − C − H d)  $CH_3 - C - C - CH_3$ - H Ι Br OH Η Η H H H H 105 The IUPAC name of CH2-CH2-CH2-OH 8. is a) 1-phenyl-3-propanol b) 3-phenyl-1-propanol c) 1-hydroxy-3-phenyl-propane d) None of the above 105 The complexes [Co(NH<sub>3</sub>)<sub>6</sub>][Cr(CN)<sub>6</sub>] and [Cr(NH<sub>3</sub>)<sub>6</sub>][Co(CN)<sub>6</sub>] are the examples of which type of

| 9.         | isomerism?   |   |  |   |
|------------|--|---|--|---|
|            | a) Geometrical isomeris  | m   |  |   |
|            | b) Linkage isomerism   |   |  |   |
|            | c) Ionization isomerism  |   |  |   |
|            | d) Coordination isomeri  | sm  |  |   |
| 106        | Racemic tartaric acid is   | optically inactive due to                   |  |   |
| 0.         |  |   |  |   |
|            | a) External compensation   | on  | b) Internal compensation                     |   |
|            | c) Presence of plane of s  |   | d) All of the above                          | $\frown$  |
| 106        | Nitration of aniline is do   |   | ·) · · · · · · · · · · · · · · · · · ·       |   |
| 1.         |  |   |  |   |
|            | a) Acidic medium   |   |  |   |
|            | b) Alkaline medium   |   |  |   |
|            | c) Neutral medium  |   |  |   |
|            | ,  | first converting it into aceta              | nilida before nitration                      |   |
| 106        | A bridging ligand posses   | —   |  | $\bigcirc$  |
| 2.         | A bi luging liganu posses  | 5565.                                       |  | ×.  |
| Ζ.         | a) Deludentate er mene   | dontato noturo                              |  |   |
|            | a) Polydentate or mono   |   |  | /   |
|            | b) Two or more donor c   |   |  |   |
|            |  | tself attached to two metal i               | ons  |   |
|            | d) All of the above  |   |  |   |
|            | What is the neutralization   | on equivalent of benzoic aci                | d?   |   |
| 3.         |  |   |  |   |
|            | a) 122   | b) 61                                       | c) 244                                       | d) 488  |
|            | <i>m</i> -chlorobenzaldehyde   | on reaction with conc. KOH                  | at room temperature gives:                   |   |
| 4.         |  |   |  |   |
|            | •  | enzoate and <i>m</i> -hydroxy bei           | -  |   |
|            |  | yde and <i>m</i> -chlorobenzylalc           |  |   |
|            |  | ol and <i>m</i> -hydroxy benzylalo          |  |   |
|            | 5  | enzoate and <i>m</i> -chlorobenzy           |  |   |
| 106        | The oxidation number o   | f Fe in brown ring [Fe(H <sub>2</sub> 0)    | <sub>5</sub> NO] <sup>2+</sup> is            |   |
| 5.         |  |   |  |   |
|            | a) 0   | b) +1                                       | c) +2  | d) +3   |
| 106        | $[Cr(H_2O)_6]Cl_3$ (at. No. o  | f Cr=24) has a magnetic mo                  | ment of 3.83 BM. The corre                   | ct distribution of 3 <i>d</i> -                   |
| 6.         | electrons in the chromiu   | ım of the complex:                          |  |   |
|            | a) $3d_{xy}^{1}$ , $3d_{yz}^{1}$ , $3d_{xz}^{1}$   | b) $3d_{xy}^1$ , $3d_{yz}^1$ , $3d_{z^2}^1$ | c) $(3d_{x^2-y^2}^1), 3d_{z^2}^1, 3d_{xz}^1$ | d) $3d_{xy}^1$ , $(3d_{y^2-y^2}^1)$ , $3d_{yz}^1$ |
| 106        |  | solution is added to 100 mL                 | -  | -   |
| 7.         |  | ver chloride obtained in gra                |  |   |
| <i>,</i> . | a) $287 \times 10^{-3}$  | b) 143.5 × 10 <sup>-3</sup>                 | c) 143.5 × $10^{-2}$                         | d) 287 × $10^{-2}$                                |
| 106        |  | sible structural isomers of t               | -  | ,   |
| 8.         | The total number of pos  | sible structural isolilers of t             |  | [rt Cl <sub>4</sub> ] ale.                        |
| 0.         | a) 3   | b) 5  | a) 4   | d) 6  |
| 106        | . F  | ,   | c) 4   | d) 6  |
|            | A similarity between op  | tical and geometrical isome                 |  |   |
| 9.         |  |   | J  |   |
|            |  | ber of isomers for a given c                | =  |   |
|            |  | is present then so is the oth               | er   |   |
|            | c) Both are included in s  |   |  |   |
| 4 6 -      | d) They have no similar  |   | 1 (11)                                       | 1   |
| 107        | In $ Ni(NH_3)_4 SO_4$ , the value va | alency and coordination nur                 | nber of Ni will be respective                | ely   |

107 In  $[Ni(NH_3)_4]SO_4$ , the valency and coordination number of Ni will be respectively

0.

a) 3 and 6 b) 4 and 4 c) 4 and 2 d) 2 and 4 107 C<sub>6</sub>H<sub>5</sub>CHO is different from aliphatic aldehyde in its reaction towards: 1. a) Tollen's reagent b) Schiff's reagent c) NaHSO<sub>3</sub> d) Fehling's solution 107 Oxidation of naphthalene by acidic KMnO<sub>4</sub> gives: 2. b) Benzaldehyde d) Benzoic acid a) Toluene c) Phthalic acid 107 The number of possible theoretical conformations of *n*-butane are 3. a) Two b) Three c) Four d) Infinite 107 Which is correct order for acidic nature of following acids? (I) PhCOOH (II)  $o - NO_2C_6H_4COOH$ 4. (III)  $p - NO_2C_6H_4COOH$ (IV) m- NO<sub>2</sub>C<sub>6</sub>H<sub>4</sub>COOH a) II > III > IV > I b) II > IV > III > I c) II > IV > I > III> III > IV d) [ 107 Salicylic acid when treated with zinc dust gives: 5. d) Benzoic acid b) Salicyladehyde c) Benzene a) Phenol 107 Action of PCl<sub>5</sub> on salicylic acid produces: 6. a) o-chlorobenzoyl b) o-hydroxybenzoyl chloride c) o-chlorobenzoic acid d) None of the above 107 Which of the following species is most stable? 7. a)  $p - O_2 N - C_6 H_4 - C_{H_2}^T$ b)  $_{C_6H_5}-_{CH_2}^+$ c)  $_{p-C1-C_{6}H_{4}-CH}$ d)  $p-CH_3O-C_6H_4-CH_2$ 107 Give the IUPAC name of the following 8.  $CH_3 CH_2 - CH_2 - CH_3$ CH<sub>3</sub>-CH<sub>2</sub>-CH<sub>2</sub>-CH<sub>3</sub> a) 5-ethyl-4, 4-dimethyloctane b) 4-ethyl-5, 5-dimetyloctane c) 3-ethyl-2-methyl-2-propyl hexane d) 4-ethyl-5-methyl, 5-propyl hexane 107 Which of the following reacts with KCN to form benzoin? 9. a) C<sub>6</sub>H<sub>5</sub>CHO b)  $C_6H_5Cl$ c)  $C_2H_5Cl$ d)  $C_6H_5CH_3$ 108 Which one is an organometallic compound in the following? 0. a) C<sub>2</sub>H<sub>5</sub>ONa b)  $C_2H_5 - S - C_2H_5$ d) Al( $C_6H_5S$ )<sub>3</sub> c)  $Al_2(CH_3)_6$ 108 The formula of picramide is: 1.



2. a) To absorb water b) To absorb hydrochloric acid c) To produce an electrophile d) To produce nucleophile 109 Coordination isomerism is caused by interchange of ligands between the 3. a) Complex cation and complex anion b) Inner sphere and outer sphere c) Low oxidation and higher oxidation states d) cis and trans structure 109 Which aldehyde is used in the manufacture of perfumes? 4. a) Cinnamaldehyde c) Benzaldehyde d) None of thes b) Salicyladehyde 109 Which of the following statements is not correct? 5. a) A meso compound has chiral centres but exhibits no optical activity b) A meso compound has no chiral centres and thus are optically inactive A meso compound has molecules in which one half of molecule is superimposable on the other even c) through chiral centre is present in them A meso compound is optically inactive because the rotation caused by one half of molecule is cancelled by the rotation produced by another half 109 The volume (in mL) of 0.1 M AgNO<sub>3</sub> required for complete precipitation of chloride ions present in 30 mL of 0.01 M solution of [Cr(H<sub>2</sub>O)<sub>5</sub>Cl]Cl<sub>2</sub> , as silver chloride is close to 6. a) 3 b) 4 c) 5 d) 6 109 Benzene is a resonance hybrid mainly of two Kekule structures. Hence: 7. a) Half of the molecules correspond to one structure, and half to the second structure b) At low temperatures benzene can be separated into two structures c) Two structures make equal contribution to resonance hybrid d) An individual benzene molecule changes back and forth between two structures 109 Keto form is more stable in 8. a)  $CH_3COCH_2COOC_2H_5$  b)  $CH_3COCH_2COCH_3$ c)  $CH_3COCH_3$ d) CH<sub>3</sub>COCH<sub>2</sub>COC<sub>2</sub>H<sub>5</sub> 109 The oxidation state and effective atomic number(EAN) of cobalt  $(CoF_6)^{2-}$  are respectively 9 c) 4 and 37 a) 3 and 36 b) 4 and 35 d) 2 and 35 110 Benzamide on reaction with POCl<sub>3</sub> gives: 0. a) Aniline b) Chlorobenzene c) Benzylamine d) Benzonitrile 110 Which pair of carbon skeleton is an example of isomerism? 1. С a) C - C - C - C and C - C - CС С С b) C - C - C and C - C - CС С

Page | 92

| 1.  |   |   |   |                       |
|-----|---|---|---|-----------------------|
|     | a) [CoCl <sub>4</sub> ] <sup>2–</sup>                   | b) [FeCl <sub>4</sub> ] <sup>2–</sup>                               | c) [NiCl <sub>4</sub> ] <sup>2–</sup>                           | d) $[PtCl_4]^{2-}$    |
| 111 | Which exhibits highest m                                |   |   |                       |
| 2.  |   |   |   |                       |
|     | a) $[Co(NH_3)_6]Cl_3$                                   | b) [Co(NH <sub>3</sub> ) <sub>5</sub> Cl]Cl <sub>2</sub>            | c) $[Co(NH_3)_4Cl_2]Cl$   | d) $[Co(NH_3)_3Cl_3]$ |
|     | =   |   | piological systems. In this co                                  | ontext which of the   |
| 3.  | following statement is inc                              |   |   |                       |
|     |   | gment in plants and contain   |   |                       |
|     |   | l pigment of blood and con<br>min B <sub>12</sub> and contains coba |   |                       |
|     |   | an enzyme and contains coba an enzyme and contains $zi$             |   |                       |
| 111 |   | e by the combination of [Co   |   |                       |
| 4.  | dompron buit buil be muue                               |   | (1113)501] 11101  |                       |
|     | a) Cl <sup>-</sup>                                      | b) 2Cl <sup>-</sup>   | c) PO <sub>4</sub> <sup>3-</sup>                                | d) 2K <sup>+</sup>    |
| 111 | Which of the following pa                               | irs represents linkage ison   | _   |                       |
| 5.  |   |   |   | N T                   |
|     |   | $[Pt (NH_3)_4][CuCl_4]$   | b) $[Pd(PPh_3)_2 (NCS)_2]$ and                                  |                       |
|     | c) $[Co(NH_3)_5]NO_3SO_4$ and                           |   | d) $[PtCl_2(NH_3)_4]Br_2$ and $[$                               | $PtBr_2(NH_3)_4]Cl_2$ |
| 111 | The reaction products of                                | $C_6H_5OCH_3 + HI \xrightarrow{\Delta} is:$                         |   |                       |
| 6.  |   |   |   |                       |
| 111 |   | b) $C_6H_5I + CH_3OH$   | c) $C_6H_5CH_3 + HOI$<br>otash and another compour              | d) $C_6H_6 + CH_3OI$  |
| 7.  |   |   | ed by reacting a compound                                       |                       |
|     | presence of slaked lime. C                              |   | ca by routing a compound  |                       |
|     | a) $C_6H_5NH_2$   | b) CH <sub>3</sub> OH   | c) CH <sub>3</sub> COCH <sub>3</sub>                            | d) CHCl <sub>3</sub>  |
| 111 | Chlorine is most reactive                               | in:   |   |                       |
| 8.  |   |   |   |                       |
|     | a) CH <sub>3</sub> Cl                                   | b) CH <sub>2</sub> =CHCl  | c) C <sub>6</sub> H <sub>5</sub> Cl                             | d) $C_6H_5CH_2Cl$     |
|     | The C—C bond order in b                                 | enzene is close to:   |   |                       |
| 9.  | a) 1.5  | b) 2.5  | c) 3.0  | d) 6.0                |
| 112 | 2   |   | 2 mole of [Co(NH <sub>3</sub> ) <sub>5</sub> Br]SO <sub>4</sub> | 2                     |
| 0.  | solution  | [00(1113)5004]21 010 010  |   |                       |
|     | 1 L of mixture <i>X</i> + excess                        | $AgNO_3 \rightarrow Y$  |   |                       |
|     | 1 L of mixture <i>X</i> + excess                        | $\operatorname{BaCl}_2 \longrightarrow Z$                           |   |                       |
|     | Number of moles of Y and                                |   |   |                       |
|     | a) 0.01, 0.01   | b) 0.01,0.02  | c) 0.02, 0.01   | d) 0.02, 0.02         |
|     | Phenol can be converted i                               | into salicylic acid by:   |   |                       |
| 1.  | a) Etard's reaction                                     |   |   |                       |
|     | a) Etard's reaction<br>b) Kolbe's reaction              |   |   |                       |
|     | c) Reimer-Tiemann react                                 | ion   |   |                       |
| 5   | d) Both (b) and (c)                                     |   |   |                       |
| 112 |   | Which of the following reas   | sons is correct?  |                       |
| 2.  |   |   |   |                       |
|     | a) Presence of one CO as l                              |   |   |                       |
|     | b) Presence of monodenta                                | -   |   |                       |
|     | c) Metal-metal (Fe-Fe) bo                               |   |   |                       |
| 117 | d) Resonance hybridization<br>The formula of dichlorobi |   |   |                       |
| 112 |   | a (urca) copper (11) 15.  |   |                       |
|     |   |   |   |                       |

3.

a)  $[CuO = C(NH_2)_2]Cl_2$ b)  $[CuCl_2 \{0 = C(NH_2)\}]$ 

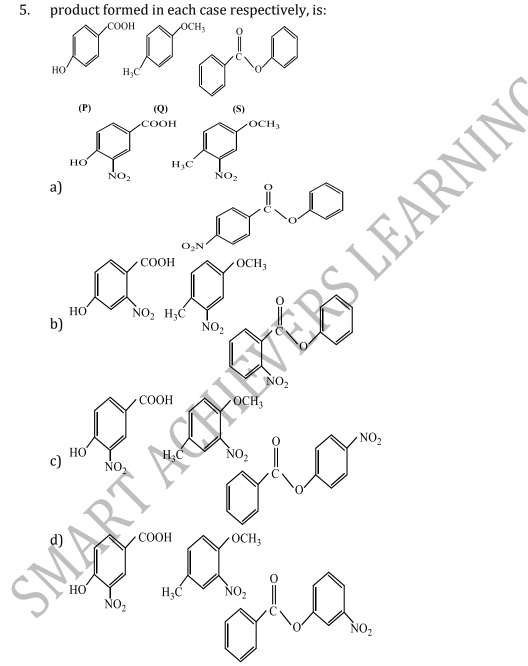
c)  $[Cu{0 = C(NH_2)_2}Cl]Cl$ 

d)  $[CuCl_2][0 = C(NH_2)_2]H_2$ 

112 Which of the following facts about the complex  $[Cr(NH_3)_6]Cl_3$  is wrong?

4.

- The complex involves  $d^2sp^3$  hybridisation and isb) The complex is paramagnetic. octahedral in shape.
- a)
- c) The complex is an outer orbital complex.
- d) The complex gives white precipitate with silver nitrate solution.
- 112 The compounds P, Q and S were separately subjected to nitration using HNO<sub>3</sub>/H<sub>2</sub>SO<sub>4</sub> mixture. The major



- 112 Aromaticity of benzene is due to:
- 6.
- a) Ring
- b) Three double bonds
- c) Delocalisation of  $\pi$ -electrons

d) None of the above 112 7. The IUPAC name of is a) 2, 2, 4, 4-tetramethyl pentane b) 2, 2-dimethyl propane c) 4-ethyl-3-methyl hex-3-ene d) Ethyl isopropyl ethene 112 Phenol is heated with a solution of mixture of KBr and KBrO<sub>3</sub>. The major product obtained in the above 8. reaction is: d) 2,4,6-tribromophenol a) 2-bromophenol b) 3-bromophenol c) 4-bromophenol 112 The coordination number of a central metal atom in a complex is determined by 9. a) The number around a metal ion bonded by pi-bonds b) The number of only anionic ligands bonded to the metal ion c) The number of ligands around a metal ion bounded by sigma and pi-bonds both d) The number of ligands around a metal ion bonded by sigma bonds 113 The true statement about benzene is: 0. a) Because of Monosubstitution of There are two types of There is a cyclic unsaturation benzene benzene gives three b) C—C bonds in benzene c) delocalisation of  $\pi$ easily undergoes isomeric products molecule electrons in benzene additions 113 Which reagent can convert CO group to  $C(C_6H_5)OH$ ? 1. c)  $C_6H_5MgBr$ a) C<sub>6</sub>H<sub>5</sub>OH b) C<sub>6</sub>H<sub>5</sub>CH<sub>2</sub>OH d)  $C_6H_5Cl$ 113 Which has highest paramagnetism? 2. d)  $[Zn(H_20)_6]^{2+}$ a)  $[Cr(H_2O)_6]^{3+}$ b)  $[Fe(H_2O)_6]^2$ c)  $[Cu(H_2O)_6]^{2+}$ 113 Which is not true ligands metal complex 3. a) Larger the ligand, the more stable is the metal-ligand complex b) Highly charged ligand forms stronger bonds c) Larger the permanent dipole moment of ligand, the more stable is the bond d) Greater the ionization potential of central metal, the stronger the bond 113  $[Co(NH_3)_4Cl_2]NO_2$  and  $[Co(NH_3)_4Cl \cdot NO_2]Cl]$  are 4. a) Optical isomers b) Geometrical isomers c) Ionization isomers d) Linkage isomers 113 Acetophenone on oxidation by perbenzoic acid gives phenyl acetate. The reaction is named as: 5. a) Baeyer-Villiger oxidation b) Perkin's reaction c) Claisen condensation d) Reformatsky reaction 113 Friedel-Craft's reaction does not occur in case of: 6. a) Toluene b) Benzene c) Naphthalene d) pyridine 113 One mode of a complex compound  $Co(NH_3)_5Cl_3$  gives three moles of ions on dissolution in water. One of the same complex reacts with two moles of AgNO<sub>3</sub> solution to yield two moles of AgCl(s). The structure of 7. the complex is a)  $[Co(NH_3)_3Cl_3] \cdot 2NH_3$ b)  $[Co(NH_3)_4Cl_2] \cdot Cl \cdot NH_3$ c)  $[Co(NH_3)_4Cl]Cl_2 \cdot NH_3$ d)  $[Co(NH_3)_5Cl]Cl_2$ 

| 113<br>8. | $C_6H_6$ is a very good indus  | trial solvent for:  |   |   |
|-----------|--|---|---|---|
| 113       | a) Oil<br>Salol is used as:  | b) Fat  | c) Rubber   | d) All of these                           |
| 9.        | Saloi is used as.  |   |   |   |
| ).        | a) Antiseptic  | b) Antipyretic  | c) Both (a) and (b)   | d) None of these                          |
| 114       | Presence of nitro gp. in be  |   |   |   |
| 0.        | or or  | 0   |   |   |
|           | a) Deactivates the ring for  | $r S_E$ reaction  |   |   |
|           | b) Activates the ring for S  | $_E$ reactions  |   |   |
|           | c) Renders the ring basic  |   |   |   |
|           | d) Deactivates the ring for  |   |   |   |
|           | _  | mplexes will show geometr   | rical as well as optical isom   | erism? (en =ethylene                      |
| 1.        | diamine)   |   |   |   |
|           | a) $[Pt(NH_3)_2Cl_2]$  |   | c) $[Pt(en)_3]^{4+}$  | d) [Pt(en) <sub>2</sub> Cl <sub>2</sub> ] |
|           | The huge number of organ   | nic compounds is due to the   | e fact that   | X   |
| 2.        |  |   |   |   |
|           | a) Tetravalency of carbon  |   | b) Carbon possesses prop  | erty of catenation                        |
| 111       | c) Carbon compounds ex   |   | d) Both (b) and (c)   | formedia                                  |
| 114<br>3. |  | ated with Br <sub>2</sub> in presence of<br>ements which are related to |   | Iormea is m-                              |
| 5.        |  | ensity on <i>meta</i> carbon is m                                       |   | nara positions                            |
|           | -  | en Br <sup>+</sup> attacks at the <i>ortho</i>                          |   |   |
|           |  | ain aromaticity from the m  |   |   |
|           | d) None of the above   | ,   |   |   |
| 114       | -  | g compounds when dissolv  | red in water, gives a solutio   | on with pH more than 7?                   |
| 4.        |  |   |   | -   |
|           | a) C <sub>6</sub> H <sub>5</sub> NH <sub>2</sub>                           | b) C <sub>6</sub> H <sub>5</sub> OH                                     | c) C <sub>2</sub> H <sub>5</sub> OH                                       | d) CH <sub>3</sub> COCH <sub>3</sub>      |
| 114       | Formula of ferrocene is:   |   |   |   |
| 5.        |  |   |   |   |
|           | a) $[Fe(CN)_6]^{4-}$   | b) $[Fe(CN)_6]^{3+}$  | c) $[Fe(CO)_5]$   | d) $[(C_6H_5)_2Fe]$                       |
|           | What is the EAN of nickel  | in $Ni(CO)_4$ ?   |   |   |
| 6.        |  |   |   | N 00                                      |
| 111       | a) 38  | b) 30   | c) 36   | d) 32                                     |
|           |  |   |   | olution in water . One mole               |
| 7.        | of the complex is  | ts with two moles of AgNO <sub>3</sub>                                  | solution to yield two mole  | s of Ager(s). The structure               |
|           | a) $[Co(NH_3)_5Cl]Cl_2$  |   | b) [Co(NH <sub>3</sub> ) <sub>3</sub> Cl <sub>2</sub> ]. 2NH <sub>3</sub> |   |
|           | c) $[Co(NH_3)_5CI_2CI_2]$  |   | d) $[Co(NH_3)_4Cl]Cl_2.NH_3$  |   |
| 114       | , , , , <u>,</u> ,   | g has largest number of iso   | ) = ( ), I = <b>1</b> )   |   |
| 8,        | (R=alkyl group, en=ethyl)  | • •   |   |   |
|           | a) $[Ru(NH_3)_4Cl_2]^+$  | b) $[Co(NH_3)_5Cl]^{2+}$  | c) $[Ir(PR_3)_2H(CO)]^{2+}$   | d) $[CO(en)_2Cl_2]^+$                     |
|           | Which complex is likely to   |   |   | ) [ ] / 2 23                              |
| 9.        |  |   |   |   |
|           | a) Trans-[Co(NH <sub>3</sub> ) <sub>4</sub> Cl <sub>2</sub> ] <sup>+</sup> |   |   |   |
|           | b) $[Cr(H_2O)_6]^{3+}$   |   |   |   |
|           | c) Cis- $[Co(NH_3)_2(en)_2]^{3+1}$   | F   |   |   |
|           | d) Trans-[Co(NH <sub>3</sub> ) <sub>2</sub> (en) <sub>2</sub>              | 2] <sup>3+</sup>  |   |   |
|           | A square planar complex  | is formed by hybridization  | of which atomic orbitals?   |   |
| 0.        |  |   |   |   |

| a) $s, p_x, p_y, d_{yz}$<br>115 The IUPAC nam<br>1. $CH_2 - CH - CH$<br>  $ $ $ $ | the of the compound $I - CH_2COCl$ is  | c) $s, p_x, p_y, d_{z^2}$   | d) $s, p_x, p_y, d_{xy}$ |
|---|--|---|--------------------------|
| c) 1, 2, 4-butan<br>115 Nitrobenzene c  | anetetrachlorocarbonyl<br>etricarboxylic acid<br>ran be prepared from benzene by ເ | b) 1, 2, 3, 4-butanete<br>d) None of the above<br>using a mixture of conc. Hi |                          |
| a) Base<br>115 In the compour   | re HNO <sub>3</sub> acts as a:<br>b) Acid<br>nd lithium tetrahydroaluminate, th    | c) Reducing agent<br>ne ligand is   | d) Catalyst              |
| 3.<br>a) H  | Ъ) Н+  | c) H-   | d) None of these         |
| SMAR  | Actilities   |   |                          |

# **COORDINATION COMPOUNDS**

CHEMISTRY

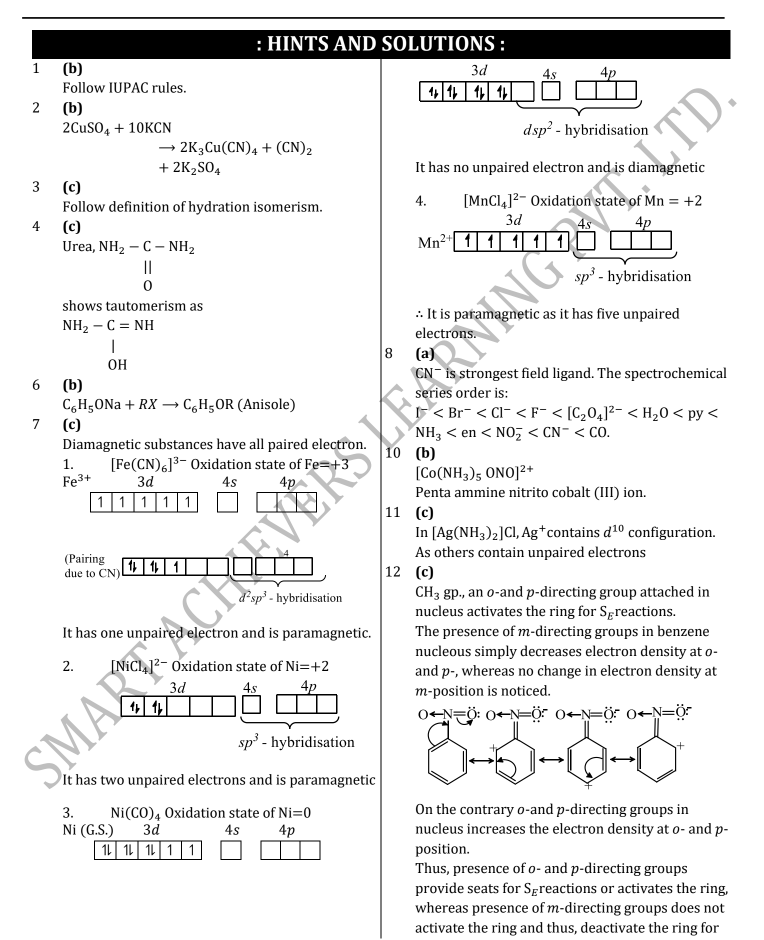
|      |   |      |   |      |   | : ANS | W | ER K | EY |      |   |      |   |      |
|------|---|------|---|------|---|-------|---|------|----|------|---|------|---|------|
| 1)   | b | 2)   | b | 3)   | С | 4)    | С | 177) | С  | 178) | b | 179) | a | 180) |
| 5)   | b | 6)   | b | 7)   | С | 8)    | а | 181) | С  | 182) | b | 183) | а | 184) |
| 9)   | С | 10)  | b | 11)  | С | 12)   | С | 185) | b  | 186) | d | 187) | С | 188) |
| 13)  | d | 14)  | d | 15)  | С | 16)   | С | 189) | d  | 190) | b | 191) | b | 192) |
| 17)  | b | 18)  | d | 19)  | b | 20)   | b | 193) | d  | 194) | d | 195) | a | 196) |
| 21)  | а | 22)  | С | 23)  | d | 24)   | d | 197) | b  | 198) | b | 199) | С | 200) |
| 25)  | d | 26)  | d | 27)  | d | 28)   | d | 201) | b  | 202) | b | 203) | d | 204) |
| 29)  | b | 30)  | а | 31)  | d | 32)   | d | 205) | b  | 206) | d | 207) | d | 208) |
| 33)  | d | 34)  | С | 35)  | а | 36)   | b | 209) | b  | 210) | С | 211) | С | 212) |
| 37)  | d | 38)  | С | 39)  | С | 40)   | d | 213) | С  | 214) | С | 215) | b | 216) |
| 41)  | а | 42)  | С | 43)  | С | 44)   | d | 217) | а  | 218) | d | 219) | b | 220) |
| 45)  | b | 46)  | d | 47)  | b | 48)   | d | 221) | d  | 222) | C | 223) | С | 224) |
| 49)  | b | 50)  | d | 51)  | d | 52)   | d | 225) | а  | 226) | b | 227) | С | 228) |
| 53)  | а | 54)  | b | 55)  | b | 56)   | С | 229) | b, | 230) | d | 231) | С | 232) |
| 57)  | а | 58)  | С | 59)  | а | 60)   | b | 233) | b  | 234) | b | 235) | d | 236) |
| 61)  | С | 62)  | С | 63)  | а | 64)   | d | 237) | b  | 238) | d | 239) | а | 240) |
| 65)  | С | 66)  | d | 67)  | С | 68)   | а | 241) | a  | 242) | a | 243) | b | 244) |
| 69)  | d | 70)  | d | 71)  | С | 72)   | a | 245) | а  | 246) | С | 247) | b | 248) |
| 73)  | а | 74)  | b | 75)  | d | 76)   | b | 249) | С  | 250) | d | 251) | d | 252) |
| 77)  | С | 78)  | b | 79)  | С | 80)   | d | 253) | С  | 254) | a | 255) | а | 256) |
| 81)  | d | 82)  | d | 83)  | С | 84)   | d | 257) | b  | 258) | С | 259) | d | 260) |
| 85)  | b | 86)  | b | 87)  | С | 88)   | С | 261) | d  | 262) | d | 263) | С | 264) |
| 89)  | d | 90)  | а | 91)  | С | 92)   | а | 265) | b  | 266) | d | 267) | b | 268) |
| 93)  | С | 94)  | b | 95)  | a | 96)   | С | 269) | b  | 270) | С | 271) | d | 272) |
| 97)  | b | 98)  | С | 99)  | b | 100)  | С | 273) | d  | 274) | b | 275) | С | 276) |
| 101) | d | 102) | С | 103) | d | 104)  | b | 277) | d  | 278) | С | 279) | С | 280) |
| 105) | d | 106) | а | 107) | b | 108)  | С | 281) | а  | 282) | С | 283) | b | 284) |
| 109) | а | 110) | a | 111) | b | 112)  | С | 285) | b  | 286) | С | 287) | b | 288) |
| 113) | b | 114) | a | 115) | d | 116)  |   | 289) | С  | 290) | b | 291) | d | 292) |
| 117) | С | 118) | a | 119) | b | 120)  | b | 293) | b  | 294) | С | 295) | С | 296) |
| 121) | а | 122) | d | 123) | d | 124)  |   | 297) | С  | 298) | b | 299) | d | 300) |
| 125) | d | 126) | b | 127) | а | 128)  | d | 301) | а  | 302) | b | 303) | b | 304) |
| 129) | b | 130) | С | 131) | d | 132)  |   | 305) | а  | 306) | b | 307) | b | 308) |
| 133) |   | 134) | d | 135) | d | 136)  |   | 309) | b  | 310) | С | 311) | b | 312) |
| 137) |   | 138) | d | 139) | b | 140)  |   | 313) | С  | 314) | а | 315) | a | 316) |
| 141) | b | 142) | а | 143) | а | 144)  |   | 317) | b  | 318) | b | 319) | d | 320) |
| 145) | а | 146) | d | 147) | d | 148)  |   | 321) | b  | 322) | d | 323) | d | 324) |
| 149) | а | 150) | С | 151) | а | 152)  |   | 325) | d  | 326) | С | 327) | b | 328) |
| 153) | b | 154) | С | 155) | а | 156)  |   | 329) | b  | 330) | d | 331) | d | 332) |
| 157) | d | 158) | b | 159) | b | 160)  |   | 333) | d  | 334) | С | 335) | b | 336) |
| 161) | b | 162) | С | 163) | d | 164)  |   | 337) | d  | 338) | d | 339) | b | 340) |
| 165) | а | 166) | а | 167) | b | 168)  |   | 341) | b  | 342) | С | 343) | а | 344) |
| 169) | b | 170) | b | 171) | С | 172)  |   | 345) | b  | 346) | С | 347) | а | 348) |
| 173) | С | 174) | а | 175) | С | 176)  | С | 349) | С  | 350) | b | 351) | а | 352) |

| 353)         | d      | 354)         | b      | 355)         | b      | 356) d           | 557)         | а | 558)         | d      | 559)         | С      | 560) c           |
|--------------|--------|--------------|--------|--------------|--------|------------------|--------------|---|--------------|--------|--------------|--------|------------------|
| 357)         | b      | 358)         | d      | 359)         | d      | 360) d           | 561)         | а | 562)         | b      | 563)         | С      | 564) a           |
| 361)         | d      | 362)         | b      | 363)         | b      |                  | 565)         |   | 566)         | С      | 567)         | С      | 568) a           |
| 365)         | а      | 366)         | а      | 367)         | С      |                  | 569)         |   | 570)         | С      | 571)         | b      | 572) d           |
| 369)         | а      | 370)         | d      | 371)         | С      | 372) a           | ,            |   | 574)         | С      | 575)         | d      | 576) a           |
| 373)         | d      | 374)         | d      | 375)         | С      | 376) c           | 577)         |   | 578)         | b      | 579)         | b      | 580) b           |
| 377)         | С      | 378)         | а      | 379)         | d      | 380) c           | 581)         |   | 582)         | b      | 583)         | С      | 584) d           |
| 381)         | d      | 382)         | а      | 383)         | d      | 384) c           | 585)         |   | 586)         | С      | 587)         | b      | 588) d           |
| 385)         | d      | 386)         | С      | 387)         | a      | 388) c           | 589)         |   | 590)         | С      | 591)         | b      | 592) c           |
| 389)         | С      | 390)         | С      | 391)         | b      | 392) a           | 593)         |   | 594)         | d      | 595)         | b      | 596) b           |
| 393)         | С      | 394)         | a      | 395)         | C      |                  | 597)         |   | 598)         | a      | 599)         | b      |                  |
| 397)         | а      | 398)         | d      | 399)         | b      |                  | 601)         |   | 602)         | b      | 603)         | d      | 604) b           |
| 401)         | a      | 402)         | b      | 403)         | C      |                  | 605)         |   | 606)         | d      | 607)         | d      | 608) b           |
| 405)         | b      | 406)         | d      | 407)         | b      | ,                | 609)         |   | 610)         | d      | 611)         | b      | 612) b           |
| 409)         | a      | 410)         | b      | 411)         | С      |                  | 613)         |   | 614)         | С      | 615)         | а      | 616) a           |
| 413)         | b      | 414)         | а      | 415)         | a      | -                | 617)         |   | 618)         | С      | 619)         | С      | 620) c           |
| 417)<br>421) | d      | 418)<br>422) | C      | 419)<br>422) | C      | 420) c           | ,            |   | 622)<br>626) | C      | 623)<br>627) | a<br>d | 624) c           |
| 421)<br>425) | c<br>d | 422)<br>426) | a<br>c | 423)<br>427) | с<br>а | 424) b<br>428) b | ,            |   | 626)<br>630) | b<br>C | 627)<br>631) | d<br>a | 628) c<br>632) b |
| 429)         | d      | 430)         | d      | 431)         | a<br>a | 432) a           | (00)         |   | 634)         | b      | 635)         | a<br>a | 636) a           |
| 433)         | u<br>C | 430)<br>434) | u<br>C | 435)         | a      | -                | 637)         |   | 638)         | C      | 639)         | a<br>C | 640) b           |
| 437)         | a      | 438)         | b      | 439)         | a      | -                | 641)         |   | 642)         | b      | 643)         | a      | 644) c           |
| 441)         | b      | 442)         | c      | 443)         | b      | ,                | 645)         |   | 646)         | a      | 647)         | c<br>c | 648) d           |
| 445)         | a      | 446)         | a      | 447)         | a      | -                | 649)         |   | 650)         | c      | 651)         | a      | 652) d           |
| 449)         | b      | 450)         | d      | 451)         | b      | 452) <b>(</b>    |              |   | 654)         | a      | 655)         | d      | 656) d           |
| 453)         | c      | 454)         | d      | 455)         | a      | -                | 657)         |   | 658)         | a      | 659)         | b      | 660) c           |
| 457)         | а      | 458)         | а      | 459)         | С      | 460) c           | 661)         |   | 662)         | С      | 663)         | b      | 664) b           |
| 461)         | С      | 462)         | b      | 463)         | b      |                  | 665)         |   | 666)         | С      | 667)         | С      | 668) c           |
| 465)         | а      | 466)         | b      | 467)         | a      |                  | 669)         | d | 670)         | а      | 671)         | С      | 672) a           |
| 469)         | С      | 470)         | С      | 471)         | с      | 472) d           | 673)         | а | 674)         | b      | 675)         | b      | 676) a           |
| 473)         | d      | 474)         | С      | 475)         | b      | 476) a           | 677)         | С | 678)         | С      | 679)         | С      | 680) b           |
| 477)         | b      | 478)         | b      | 479)         | С      | 480) c           | 681)         | а | 682)         | а      | 683)         | С      | 684) b           |
| 481)         | b      | 482)         | а      | 483)         | a      | 484) c           | 685)         | C | 686)         | d      | 687)         | b      | 688) b           |
| 485)         | а      | 486)         | a      | 487)         | d      | 488) c           | 689)         | b | 690)         | b      | 691)         | а      | 692) c           |
| 489)         | b      | 490)         | а      | 491)         | С      | 492) a           | 693)         | а | 694)         | а      | 695)         | d      | 696) a           |
| 493)         | d      | 494)         | d      | 495)         | a      | ,                | 697)         | а | 698)         | b      | 699)         | b      | 700) d           |
| 497)         | С      | 498)         | C      | 499)         | a      |                  | 701)         |   | 702)         | С      | 703)         | b      | 704) a           |
| 501)         | а      | 502)         | d      | 503)         | b      | ,                | 705)         |   | 706)         | d      | 707)         | С      | 708) d           |
| 505)         | b      | 506)         | d      | 507)         | a      |                  | 709)         |   | 710)         | b      | 711)         | b      | 712) c           |
| 509)         | b      | 510)         | d      | 511)         | b      | -                | 713)         |   | 714)         | b      | 715)         | d      | 716) d           |
| 513)         | d      | 514)         | С      | 515)         | b      | -                | 717)         |   | 718)         | b      | 719)         | С      | 720) c           |
| 517)         | a      | 518)         | С      | 519)         | a      | -                | 721)         |   | 722)         | С      | 723)         | С      | 724) b           |
| 521)         | b      | 522)         | С      | 523)         | d      | -                | 725)         |   | 726)         | С      | 727)         | С      | 728) b           |
| 525)<br>520) | a      | 526)<br>520) | d      | 527)<br>521) | d      | -                | 729)         |   | 730)<br>724) | C      | 731)         | a      | 732) c           |
| 529)<br>522) | d<br>h | 530)<br>524) | c      | 531)<br>525) | a<br>h | ,                | 733)         |   | 734)<br>729) | a      | 735)<br>720) | a<br>h | 736) b           |
| 533)<br>527) | b      | 534)<br>539) | a      | 535)<br>520) | b<br>h | -                | 737)         |   | 738)<br>742) | a<br>d | 739)<br>742) | b      | 740) d           |
| 537)<br>541) | a      | 538)<br>542) | a<br>d | 539)<br>542) | b      | ,                | 741)         |   | 742)<br>746) | d<br>d | 743)<br>747) | a<br>h | 744) d<br>749) c |
| 541)<br>545) | C<br>d | 542)<br>546) | d<br>c | 543)<br>547) | a<br>d | -                | 745)         |   | 746)<br>750) | d<br>C | 747)<br>751) | b<br>b | 748) c<br>752) a |
| 545)<br>549) | d      | 546)<br>550) | C<br>h | 547)<br>551) | d<br>b | -                | 749)         |   | 750)<br>754) | C<br>h | 751)<br>755) | b<br>d | 752) a<br>756) c |
| 549)<br>553) | a<br>d | 550)<br>554) | b<br>b | 551)<br>555) | b<br>a | -                | 753)<br>757) |   | 754)<br>758) | b<br>b | 755)<br>759) | d<br>b | 756) с<br>760) b |
| 555J         | d      | 534J         | U      | 555J         | a      | 550J U           | /3/]         | U | / 30]        | U      | /39]         | U      |                  |
|              |        |              |        |              |        |                  |              |   |              |        |              |        |                  |

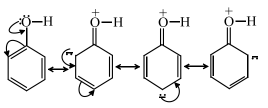
|      |   | -    |     | -    |   |      |   | 0.44  |   | 0 ( 0 ) |   |       |            |       |   |
|------|---|------|-----|------|---|------|---|-------|---|---------|---|-------|------------|-------|---|
| 761) | d | 762) | b   | 763) | a | 764) |   | 961)  | С | 962)    | C | 963)  | a          | 964)  | a |
| 765) | С | 766) | a   | 767) | b | 768) | C | 965)  | C | 966)    | d | 967)  | d          | 968)  | d |
| 769) | b | 770) | d   | 771) | a | 772) | b | 969)  | b | 970)    | а | 971)  | b          | 972)  | а |
| 773) | a | 774) | d   | 775) | d | 776) | b | 973)  | b | 974)    | С | 975)  | d          | 976)  | С |
| 777) | b | 778) | С   | 779) | b | 780) | d | 977)  | а | 978)    | d | 979)  | b          | 980)  | d |
| 781) | d | 782) | a   | 783) | С | 784) | d | 981)  | a | 982)    | b | 983)  | b          | 984)  | b |
| 785) | С | 786) | b   | 787) | а | 788) | С | 985)  | d | 986)    | С | 987)  | d          | 988)  | а |
| 789) | b | 790) | С   | 791) | С | 792) | С | 989)  | С | 990)    | d | 991)  | а          | 992)  | а |
| 793) | b | 794) | b   | 795) | С | 796) |   | 993)  | d | 994)    | b | 995)  | С          | 996)  | C |
| 797) | b | 798) | d   | 799) | b | 800) | d | 997)  | a | 998)    | b | 999)  | c          | 1000) |   |
| 801) | С | 802) | b   | 803) | С | 804) | а | ,     |   | 1002)   |   | 1003) |            | 1004) |   |
| 805) | С | 806) | d   | 807) | а | 808) | С | 1005) |   | 1006)   |   | 1007) |            | 1008) |   |
| 809) | b | 810) | d   | 811) | d | 812) | b | ,     |   | 1010)   |   | 1011) |            | 1012) |   |
| 813) | b | 814) | b   | 815) | С | 816) | b | 1013) | b | 1014)   |   | 1015) | - <b>F</b> | 1016) | d |
| 817) | а | 818) | d   | 819) | С | 820) | а | 1017) |   | 1018)   |   | 1019) |            | 1020) |   |
| 821) | b | 822) | С   | 823) | С | 824) | а | 1021) | а | 1022)   |   | 1023) |            | 1024) | d |
| 825) | а | 826) | d   | 827) | С | 828) | d | 1025) | а | 1026)   |   | 1027) |            | 1028) | d |
| 829) | d | 830) | b   | 831) | а | 832) | b |       |   | 1030)   |   | 1031) |            | 1032) | d |
| 833) | а | 834) | С   | 835) | b | 836) | d | 1033) | а | 1034)   |   | 1035) | а          | 1036) | С |
| 837) | d | 838) | а   | 839) | d | 840) | С | 1037) | a | 1038)   | d | 1039) | b          | 1040) | а |
| 841) | С | 842) | b   | 843) | b | 844) | С | 1041) | c | 1042)   |   | 1043) | d          | 1044) | а |
| 845) | b | 846) | b   | 847) | а | 848) | d |       |   | 1046)   | b | 1047) | d          | 1048) | С |
| 849) | а | 850) | b   | 851) | b | 852) | С | 1049) | d | 1050)   |   | 1051) | С          | 1052) | а |
| 853) | С | 854) | b   | 855) | а | 856) | a | 1053) | С | 1054)   | С | 1055) | b          | 1056) | а |
| 857) | d | 858) | а   | 859) | b | 860) | b | 1057) | d | 1058)   | b | 1059) | d          | 1060) | а |
| 861) | а | 862) | С   | 863) | d | 864) | b | 1061) | d | 1062)   | d | 1063) | а          | 1064) | d |
| 865) | а | 866) | d   | 867) | d | 868) | С | 1065) | b | 1066)   | a | 1067) | а          | 1068) | С |
| 869) | b | 870) | d   | 871) | c | 872) | d | 1069) | С | 1070)   | d | 1071) | d          | 1072) | С |
| 873) | С | 874) | а   | 875) | b | 876) | b | 1073) | d | 1074)   | а | 1075) | d          | 1076) | а |
| 877) | b | 878) | a   |      | b | 880) |   | 1077) |   | 1078)   | а | 1079) | а          | 1080) | С |
| 881) | а | 882) | d   | 883) | b | 884) | b | 1081) | С | 1082)   | d | 1083) | d          | 1084) | а |
| 885) | b | 886) | С   | 887) | а | 888) | d | 1085) | d | 1086)   | d | 1087) | b          | 1088) | d |
| 889) | b | 890) | d 🗸 | 891) | а | 892) |   | 1089) |   | 1090)   | а | 1091) | С          | 1092) | С |
| 893) | а | 894) | b   | 895) | b | 896) |   | 1093) |   | 1094)   | С | 1095) | b          | 1096) | d |
| 897) | а | 898) | b   | 899) | b | 900) | d | 1097) | С | 1098)   | С | 1099) | b          | 1100) | d |
| 901) | С | 902) | d   | 903) | а | 904) |   | 1101) |   | 1102)   | а | 1103) | d          | 1104) | С |
| 905) | b | 906) | C   | 907) | С | 908) |   | 1105) |   | 1106)   |   | 1107) |            | 1108) |   |
| 909) | a | 910) | d   | 911) | d | 912) |   | 1109) |   | 1110)   |   | 1111) |            | 1112) |   |
| 913) | C | 914) | С   | 915) | b | 916) |   | 1113) |   | 1114)   |   | 1115) | b          | 1116) | а |
| 917) | a | 918) | d   | 919) | b | 920) |   | 1117) |   | 1118)   | d | 1119) | а          | 1120) | а |
| 921) | a | 922) | С   | 923) | d | 924) |   | 1121) |   | 1122)   |   | 1123) | b          | 1124) |   |
| 925) | а | 926) | С   | 927) | b | 928) |   | 1125) |   | 1126)   |   | 1127) |            | 1128) |   |
| 929) | b | 930) | d   | 931) | d | 932) |   | 1129) |   | 1130)   |   | 1131) | С          | 1132) | b |
| 933) | а | 934) | С   | 935) | С | 936) |   | 1133) |   | 1134)   | С | 1135) | а          | 1136) | d |
| 937) | d | 938) | а   | 939) | С | 940) |   | 1137) |   | 1138)   |   | 1139) |            | 1140) |   |
| 941) | d | 942) | d   | 943) | b | 944) |   | 1141) |   | 1142)   | d | 1143) | a          | 1144) | а |
| 945) | b | 946) | а   | 947) | а | 948) |   | 1145) |   | 1146)   | С | 1147) | а          | 1148) | d |
| 949) | b | 950) | d   | 951) | b | 952) | С | 1149) | С | 1150)   | b | 1151) | a          | 1152) | а |
| 953) | b | 954) | а   | 955) | а | 956) | С | 1153) | С |         |   |       |            |       |   |
| 957) | а | 958) | а   | 959) | d | 960) | С |       |   |         |   |       |            |       |   |
|      |   |      |     |      |   |      |   |       |   |         |   |       |            |       |   |

# **COORDINATION COMPOUNDS**

#### CHEMISTRY



 $S_E$  reactions



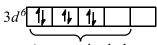
#### 13 **(d)**

In both  $[Co(NH_3)_6]^{3+}$  and  $[CoF_6]^{3+}$ , Co is present as  $Co^{3+}$ .

Thus, the electronic configuration of Co is  $_{27}\text{Co}=[\text{Ar}] 3d^7, 4s^2$  $_{27}\text{Co}^{3+}=[\text{Ar}]3d^6, 4s^0$ 

In case of  $[Co(NH_3)_6]^{3+}$ , NH<sub>3</sub> is a strong field ligand, so pairing of electrons in 3*d*-orbital takes place.

 $_{27}$ Co<sup>3+</sup>=[Ar]3d<sup>6</sup>, 4s<sup>0</sup>



(no unpaired electron)

In  $[CoF_6]^{3+}$ , F is a weak field ligand, thus doesn't cause pairing. Hence,

 $_{27}\text{Co}^{3+}=[\text{Ar}]3d^6, 4s^0$ 

| $3d^6$ | 1, | 1 | 1 | 1 | 1 |
|--------|----|---|---|---|---|
|        |    |   |   |   |   |

4 unpaired electron

#### 18 **(d)**

It is a test for primary amines. No doubt 2,4dimethylaniline is also primary amine but it does not give test due to steric hindrance.

# 19 **(b)**

 $CN^-$  is strong field ligand because it is an example of pseudohalide. Pseudohalide ions are stronger coordinating ligands and they have the ability to form  $\sigma$  bond and  $\pi$ -bond.

#### 20 **(b)**

Higher the charge and smaller the size of ligand, more stable is the complex formed

# 21 (a)

Trinitrobenzene is an explosive compound formed during nitration of  $C_6H_6$  with fuming HNO<sub>3</sub>.

# 22 (c)

A ligand is a species that is capable of donating an electron pair(s) to the central metal ion. The substances which are capable of donating an electron pair are called Lewis base, so a ligand is also a Lewis base.

#### 23 **(d)**

In  $Ni(CO)_4$ , Ni is in zero oxidation state. It has tetrahedral geometry but is diamagnetic. In

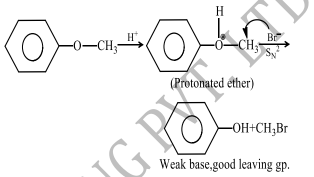
 $[Ni (CN)_4]^{2-}$ , Ni is in +2 oxidation state. It is  $dsp^2$  hybridised and have square planar shape. The compound is diamagnetic.

# 24 **(d)**

 $[Co(CN)_6]^{3-}$  has  $d^2sp^3$ -hybridisation and six *d*-electrons are paired due to strong field ligand. Thus no unpaired electron.

# 25 **(d)**

 $HBr \rightarrow H^+ + Br^-$ 

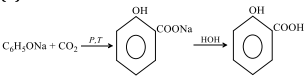


Ether reacts with acid to give protonated ether. The next step involves nucleophilic attack by halide ion with the displacement of weakly basic alcohol molecule.

#### 26 **(d)**

Octahedral complex should have six hybridized orbitals.





Kolbe-Schmidt's reaction.

# 29 **(b)**

The pair of electron present with nitrogen will not be available to be donated as H<sup>+</sup> will consume that one.

#### 30 **(a)**

It provides maximum number of ions (five) on ionization.

31 **(d)** 

Follow Vorlander's rule.

# 32 **(d)**

Organometallic compounds are those in which metal is linked directly with carbon.  $CH_3Li$ , methyl lithium due to the presence of metal-carbon bond, is an organometallic compound.

33 **(d)** 

The directive influence order is:

$$0^- > NR_2 > NHR > NH_2 > OH > OCH_3$$
  
 $\approx NHCOCH_3 > CH_3 > X$ 

34 **(c)** 

Hybridisation Shape

| dsp <sup>2</sup> | Square planar   |
|------------------|-----------------|
| sp <sup>3</sup>  | Tetrahedral     |
| $sp^2$           | Trigonal planar |

Hence, in tetrahedral complexes metal atom is  $sp^3$ hybridised.

#### 36 **(b)**

The number of ligands attached to the central metal ion is called the coordination number. So, coordination numbers of Fe in  $[{\rm Fe}~({\rm CN})_6]^{4-}, [{\rm Fe}({\rm CN})_6]^{3-}$  and  $[{\rm FeCl}_4]^-$  are 6, 6

and 4 respectively.

#### 37 (d)

Tautomers may or may not be metamers

#### 38 (c)

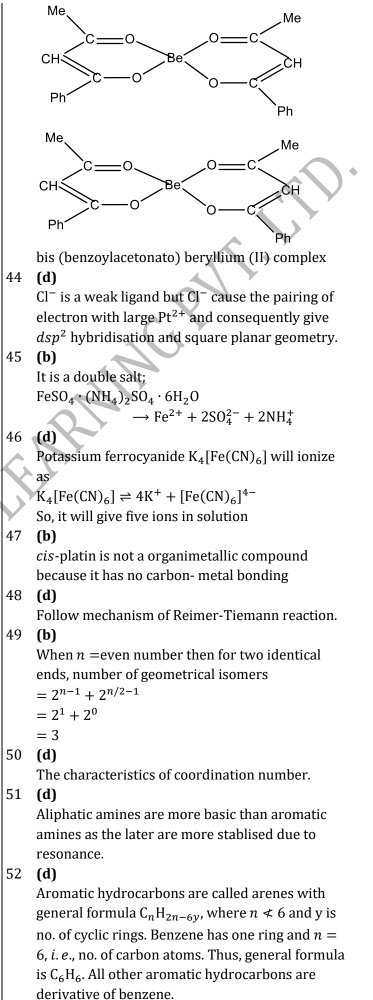
EAN=(Atomic number  $-0.S + 2 \times C.N.$ ) Hence, EAN of Ni in  $[Ni(CN)_4]^{2-} = (28 - 2 + 2 \times 2)^{2-}$ 4) = 34

#### 39 (c)

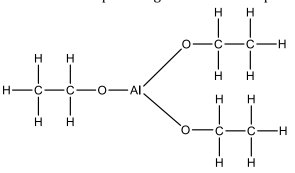
Electron repelling nature of methoxy gp. facilitate the protonation of alcohol.

#### 40 (d)

 $[Ni(Cl)_4]^{2-}$  oxidation state of Ni is +2 So, configuration of  $Ni^{2+} =$  $1s^2, 2s^22p^6, 3s^23p^63d^8$ In Ni<sup>2+</sup> 3d4*s* 11 11 11 1 In  $[NiCl_4]^2$  **1 1 1**  $sp^3$  - hybridisation Thus, due to  $sp^3$ -hybridisation of Ni<sup>2+</sup> in  $[NiCl_4]^{2-}$ , the shape of  $[NiCl_4]^{2-}$  is tetrahedral. 41 (a) This is Sandmeyer's reaction. 42 (c) *p*-nitrophenols are more acidic. 43 (c) Benzoylacetonato beryllium exhibit optical isomerism as follows

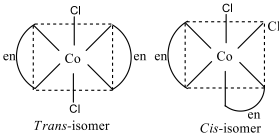


# Al $(OC_2H_5)_3$ doesn't have metal-carbon bond.(*i.e.*, it is not an example of organometallic compound). 60

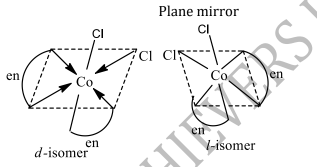


#### 54 **(b)**

In  $[Co(en)_2Cl_2]$ , four isomers are possible, two geometrical isomers and two optical isomers.



Now, *cis*-isomer also show optical isomerism. *Cis* isomer exists in enantiomeric form as it is unsymmetrical.



55 **(b)** 

A carbon atom which is attached by four different group is called an asymmetric carbon atom or chiral centre

HOOC(CHOH)<sub>2</sub>COOH has two asymmetric carbon atom

56 **(c** 

Each  $\pi$ -electron is delocalised on each C-atom. **57** (a)

An orange-red dye is formed with  $C_6H_5NH_2$ .

59 **(a)** 

Thiophene reacts more readily with  $H_2SO_4$  than  $C_6H_6$  giving thiophene sulphonic acid which is water soluble and thus, can be separated from  $C_6H_6$ . This can not be made by fractional

distillation because thiophene and  $C_6H_6$  both have nearly same b.p.

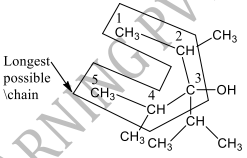
#### (b)

61

As cobalt is present as  $CO^{3+}$  and coordination number of cobalt is 6, the molecular formula of compound should be  $CoCl_3$ .  $yNH_3$ . Now, as it gives a total of three ions when dissolved in water, its structural formula must be  $[CoCl(NH_3)_5]Cl_2 \approx [CoCl(NH_3)_5]^{2+} + 2Cl^{-}$ 

Thus, only one Cl<sup>-</sup> ion is satisfying both primary and secondary valency of Co<sup>3+</sup> in this compound.

(c) The structure of alcohol is



2,4-dimethyl-3-(1-methyl) ethyl pentan-3-ol 62 (c)

The transition metal cations during complex formation show *d*-*d* transition to give coloured ions.

#### 63 **(a)**

 $-CH_3$  gp. Shows +ve inductive effect and -OH gp. shows resonance effect which increases the electron density on  $C_6H_6$  ring.

#### 64 **(d)**

It produces least number of ions in solution.

# 66 **(d)**

The process is known as aromatisation or cyclization.

67 **(c)** 

$$CH_3$$

$$|$$

$$CH_3 - C - CH_3$$

$$|$$

$$CH_3$$

neo-pentane

The structure shows that all the hydrogen atoms are attached to primary C-atoms hence these are primary hydrogens

68 **(a)** 

Follow IUPAC rules.

69 **(d)** 

H<sub>3</sub>C H<sub>3</sub>C NO<sub>2</sub>
H<sub>3</sub>C NO<sub>2</sub>
H<sub>3</sub>C NO<sub>2</sub>
H<sub>3</sub>C NO<sub>2</sub>
has no α-hydrogen. Hence, it will not show tautomerism
70 (d)
Both CN<sup>-</sup> and NO<sub>2</sub><sup>-</sup> are strong field ligands.
71 (c)
Prussian blue is Fe<sup>41</sup><sub>4</sub>[Fe<sup>II</sup>(CN)<sub>6</sub>]<sub>3</sub> or M<sup>1</sup>Fe<sup>III</sup>[Fe<sup>II</sup>(CN)<sub>6</sub>], where M<sup>I</sup> is Na, K, Rb, Li, Cs.
73 (a)
Co<sup>3+</sup>, Fe<sup>3+</sup> and Cr<sup>3+</sup> have 6*d*-electrons, 5*d*-electrons and 3*d*-electrons respectively. Mn<sup>7+</sup> has no *d*-electron.
74 (b)

All complexes of Co(III) have six ligands or coordination number of six and thus, are octahedral in shape.

#### 75 **(d)**

NH<sub>3</sub> is weak as well as strong field ligand.

#### 77 **(c)**

 $[Pt(NH_3)_3Br(NO_2)Cl]Cl$ 

Triamminebromochloronitro platinum (IV) chloride.

#### 78 **(b)**

Both the carbon attached to 0 are part of aromatic system.

# 79 **(c)**

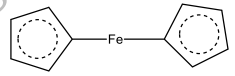
Phenol is weak acid.

#### 80 **(d)**

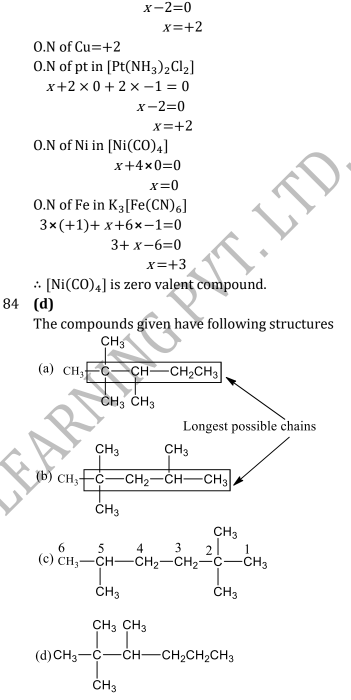
[EDTA]<sup>4–</sup> is a hexadentate ligand because it donates six pairs of electrons to central metal atom in a complex.

#### 82 **(d)**

Ferrocene of bi-(cyclopentadienyl) iron is an orange-crystalline solid. It is  $Fe(\eta^5 - C_5H_5)_2$ . The structure of ferrocene is regarded as sandwiche structure, in which the iron atom is sandwiched between two  $C_5H_5$  organic rings. The planes of the rings are parallel so that all the carbon atoms are at the same distance from the iron atom. It is a  $\pi$ -bonded complex. Its structure is as



83 (c)  $[Cu(NH_3)_4]SO_4$ Oxidation number of  $Cu \Rightarrow x+4 \times 0 - 2 = 0$ 



Out of these the (a) and (b) contain 5 C-atoms in their longest possible chains hence, these could not be the correct options for 2, 2, 3-trimethylhexane. Out of (c) and (d), the (c) is 2, 2, 5-trimethyl hexane and (d) is 2, 3, 3-trimethyl hexane

#### 85 **(b)**

Phenoxy benzene is diphenyl ether.

# 86 **(b)**

Ziegler-Natta catalyst is an organometallic compound containing titanium. It is  $TiCl_4$  and  $(C_2H_5)_3$  Al. It is used in the preparation of polyethylene.

$$nCH_2 = CH_2 \xrightarrow{330-350 \text{ K},1-2 \text{ atm}}_{\text{TiCl}_4 + (C_2H_5)_3\text{Al}} (-CH_2 - CH_2 -)_n$$

polyethylene

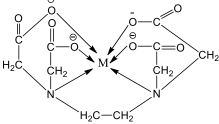
87 **(c)** 

Al<sub>2</sub>(C<sub>2</sub>H<sub>5</sub>)<sub>6</sub> + TiCl<sub>4</sub> is Zeigler Natta catalyst. 88 **(c)** 

Transition metals have empty or half filled *d*-orbitals to accept electron pairs.

89 **(d)** 

The number of atom of the ligand that are directly bound to the central metal atom or ion by coordinate bonds is known as the coordinate number of the metal or ion. It is actually the number of chemical bonds which the ligand form with the central metal atom or ion

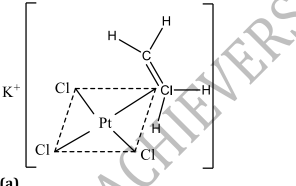


# 90 **(a)**

Acyl chlorides or acid amhydrides are used in acylation.

92 **(a)** 

Zeise's salt,  $K[PtCl_3(C_2H_4)]$  is a  $\pi$ -bonded organometallic compound. Its structure is as



95 **(a)** 

Follow IUPAC rules.

96 **(c)** 

Since the complexes

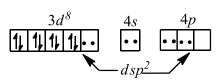
 $[PtCl_2(NH_3)_4]Br_2$  and  $[PtBr_2(NH_3)4Cl_2$  have the same molecular formula but on ionisation they give different ions, they exhibit ionisation isomerism.

 $P [PtCl_2(NH_3)_4]Br_2 \rightleftharpoons [PtCl_2(NH_3)_4]^{2+} + 2Br^{-}$  $[PtBr_2(NH_3)_4]Cl_2 \rightleftharpoons [PtBr_2(NH_3)_4]^{2+} + 2Cl^{-}$ 

97 **(b)** 

 $\mathrm{Ni}^{2+} + 4\mathrm{CN}^{-} \rightarrow [\mathrm{Ni}(\mathrm{CN})_4]^{2-}$ 

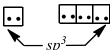
Here  $Ni^{2+}$  has  $d^8$ -configuration with  $CN^-$  as strong ligand.



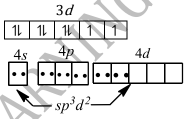
*d*<sup>8</sup>-configuration in strong ligand field gives *dsp*<sup>2</sup> hybridisation, hence square planar geometry.

 $\mathrm{Ni}^{2+} + 4\mathrm{Cl}^{-} \rightarrow [\mathrm{Ni}\mathrm{Cl}_4]^{2-}$ 

Here Ni<sup>2+</sup> has  $d^8$ -configuration with CN<sup>-</sup> as weak ligand.



 $d^{8}$ -configuration in weak ligand field gives  $sp^{3}$ hybridisation, hence tetrahedral geometry. Ni<sup>2+</sup> with H<sub>2</sub>O forms [Ni(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup> complex and H<sub>2</sub>O is a weak ligand.



Therefore, [Ni(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup>has octahedral geometry. 98 **(c)** 

Benzene ring is activates for  $S_E$  reaction by the +Ieffect as well as hyperconjugation of CH<sub>3</sub> group -Cl deactivates as -I effect predominates over +M effect. -NO<sub>2</sub> group deactivates ring by -Ieffect and -M effect.

100 **(c)** 

Alcohols are neutral.

101 **(d)** 

—OH is *o*-and *p*-directing gp.

102 **(c)** 

 $[{\rm Fe}(\eta^5-{\rm C_2H_5})_2]$  is the organometallic compound which has  $\sigma$  and  $\pi$  bonds present

# 103 **(d)**

BHC is  $C_6H_6Cl_6$  a saturated cyclic molecule.

#### 104 **(b)**

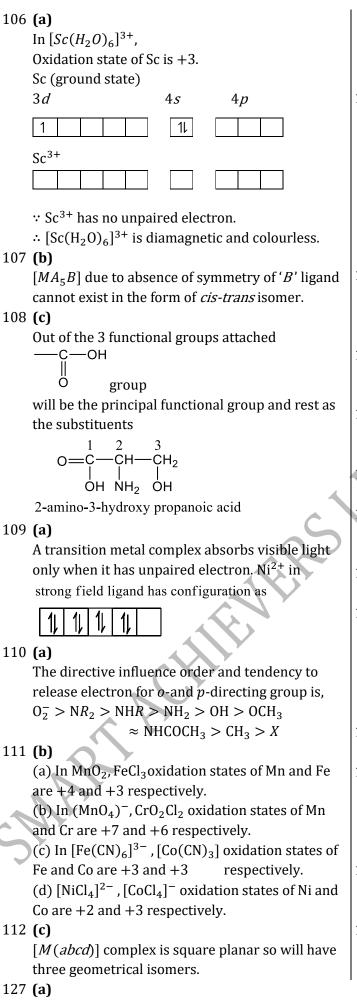
The complexes can be written as follows  $[Co(NH_3)_6]Cl_3, [Co(NH_3)_5Cl]Cl_2, [Co(NH_3)_4Cl_4]Cl$ Hence, no. of primary valencies are 3, 2 and 1 respectively

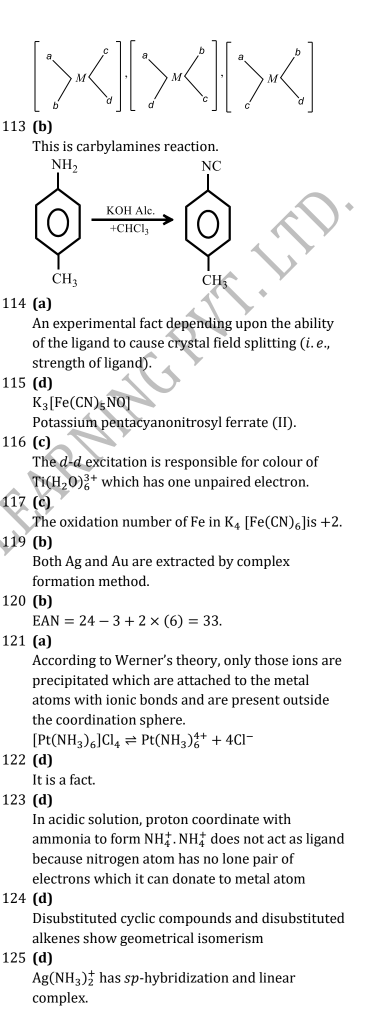
105 **(d)** 

 $[Cr(NH_3)_5 NO_2]Cl_2$  compound shows linkage isomerism because it has  $NO_2$  group which is ambidentate ligand.

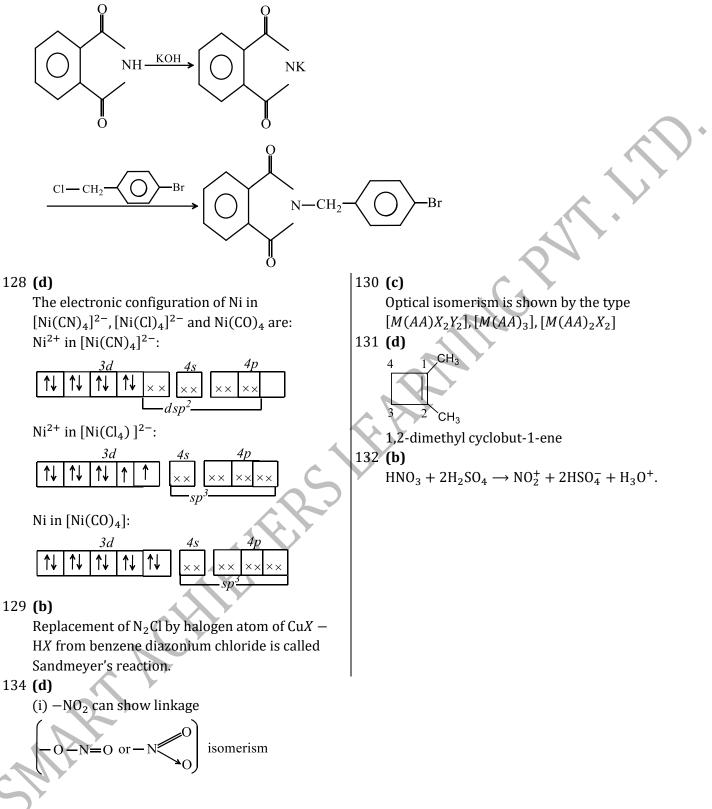
It can be linked *via* N atom  $(-NO_2)$  or *via* O atom (-ONO) to form two different isomers.

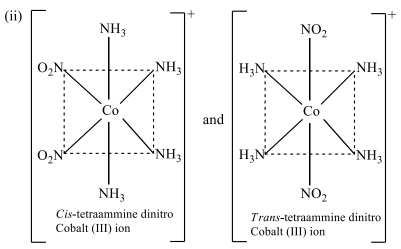






The replacement of Cl is due to the formation of stable benzyl carbocation. Alternatively Cl is present in side chain and thus replaced whereas Br is attached in benzene nucleus.





(iii) Also  $[Co(NH_3)_4(NO_2)_2]$ Cl has its ionisation isomer as  $[Co(NH_3)_4NO_2Cl]NO_2$ .

135 (d)

| • •                 |                  |
|---------------------|------------------|
| Complex             | Hybridization    |
| $[Ni(CO)_4]$        | sp <sup>3</sup>  |
| $[Ni(CO)_4]^{2-}$   | dsp <sup>2</sup> |
| $[CoF_6]^{3-}$      | $sp^3d^2$        |
| $[Fe(CN)_{6}]^{3-}$ | $d^2sp^3$        |

#### 136 **(d)**

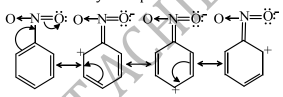
2, 4, 6-trinitrophenol is known as picric acid, an explosive.

#### 137 **(c)**

It is a fact.

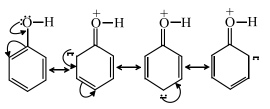
#### 139 **(b)**

OH gp., an *o*- and *p*-directing group activates ring for reactions. The presence of *m*-directing groups in benzene nucleous simply decreases electron density at *o*- and *p*-, whereas no change in electron density at *m*-position is noticed.



On the contrary *o*-and *p*-directing groups in nucleus increases the electron density at *o*- and *p*-position.

Thus, presence of o- and p-directing groups provide seats for  $S_E$  reactions or activates the ring, whereas presence of m-directing groups does not activate the ring and thus, deactivate the ring for  $S_E$  reactions



## 140 **(a)**

If magnetic moment is zero the species should not have unpaired electrons.

141 **(b)** 

Chlorophyll is a complex having Mg-atom.

## 142 **(a)**

Primary valency of metal is satisfied only by the anion. It is simply ionic valency. While secondary valency is satisfied by ligands (which can give a lone pair of electron). The ligands satisfying secondary valency, are always written in coordination sphere. This concept was given by Werner.

In  $K_3[Fe(CN)_6]$ , the CN<sup>-</sup> ions satisfy both the primary as well as secondary valency of Fe<sup>3+</sup> ion. (a)

# 143 **(a)**

The following isomers the alkene have

(i) 
$$CH_3 - CH_2 - CH = CH_2$$
  
(ii)  $CH_3 - CH = CH - CH_3$   
(iii)  $H_3C$   $C=C$   $H_3$   
 $H_3C$   $H_3C$   $H_3C$   $CH_3$   $CH_3$   $CH_3$ 

(iv) 
$$H_3C$$
  $C=C$   $H_3$   $CH_3$   $H_3C$   $CH_3$ 

(v) 
$$CH_3 - C = CH_2$$

145 **(a)** 

According to postulates of Werner's theory for coordination compounds, metal atoms exhibit two types of valencies *i.e.*, primary valency and secondary valency. The primary valency is ionisable whereas the secondary valency is nonionisable.

146 **(d)** 

 $[Fe(CN)_6]^{4-}$  is **diamagnetic** (Fe<sup>2+</sup> has  $3d^6$ configuration and the 6 electron pairs up in three *d*-orbitals followed by  $d^2sp^3$ -hybridisation).  $[Cr(NH_3)_6]^{3+}$  is paramagnetic  $(Cr^{3+}$  has  $3d^3$ configuration. Hybridisation is  $d^2sp^3$ . Due to 3 unpaired electrons it is paramagnetic)

 $[Cr(CO)_{6}: Cr(Z =$ 

25):  $[Ar]^{18}4s^1, 3d^5$ .

The one 4*s*-electron pairs up with five 3*d*electrons in three *d*-orbitals. This is followed by  $d^2sp^3$ -hybridisation to give octahedral complex. No unpaired electron and hence complex is diamagnetic.

$$Fe(CO)_5 : Fe(Z =$$

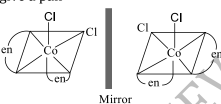
26):  $[Ar]^{18}4s^2, 3d^6$ .

The six electrons in *d*-subshell pairs up in three *d*orbitals. This is followed by  $d^2sp^3$ -hybridisation to give octahedral geometry with one vacant hybridised orbital. The resulting shape of the complex is square based pyramid. As there is no unpaired electron, the complex is **diamagnetic**.

## 147 (d)

A modified or extended Friedel-Crafts reaction. 148 (a)

cis[Co(en)<sub>2</sub>Cl<sub>2</sub>]Cl is optically active hence, it will give a pair



#### 149 (a)

 $\Delta_t$  is roughly 4/9 times to  $\Delta_0$ 

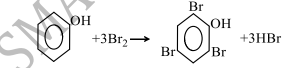
150 (c)

Follow IUPAC rules

151 (a)

Alkanes having less than four carbon atoms in basic chain will not show chain isomerism

152 **(b)** 



3 mole of Br<sub>2</sub> are needed.

153 (b)

Diamethyl glyoxime forms a colour complex with nickel

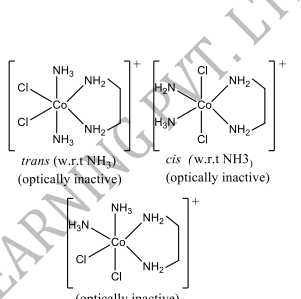
154 (c)

 $BF_4^-$  has  $sp^3$ -hybridisation and tetrahedral.

155 (a)

Oxidation state of iron in haemoglobin is +2.

- 156 (b)
  - 5. Geometrical isomers have same structural formula but differ in spatial arrangement of groups.
  - 6. Different arrangement of atoms or groups in three dimensional space results in two optical isomers which are image of each other.



(optically inactive)

Therefore, number of geometrical isomers, optical isomers and total number of isomers are 2, 2 and 3 respectively.

## 157 (d)

Hetero aromatics show aromatic nature due to  $4n + 2\pi$  electrons.

158 (b)

CN<sup>-</sup> ligand has strong ligand field because of higher value of  $\Delta$ .

159 (b)

% Enantiomeric excess

observed specific rotation

specific rotation of pure enantiomer × 100 Observed specific rotation  $=\frac{3/4}{100} \times (+16^{\circ}) \times 100$ 

 $= +12^{\circ}$ 

160 **(c)** 

Follow IUPAC rules.

161 (b)

 $[(C_6H_5)_3P)_3RhCl]$  or  $[(Ph_3P)_3RhCl]$  is a Wilkinson's catalyst, the most widely used of all catalysts for homogeneous hydrogenation.

162 (c)

Halogens attack double bond of C<sub>6</sub>H<sub>6</sub> in presence

of light. In absence of light as well as in presence of only AlCl<sub>3</sub>, S<sub>E</sub> reactions are noticed. 163 (d) 2 = 6. $[Pt(NH_3)_6]Cl_4$  complex gives five ions in the 177 (c) solution.  $[Pt(NH_3)_6]Cl_4 \rightleftharpoons [Pt(NH_3)_6]^{4+} + 4Cl^ [Ni(CO)_4]$ 164 (a) The EAN for Cu in  $[Cu(NH_3)_4]^{2+}$  is 35 and not 36,  $x + (0 \times 4) = 0$ the next inert gas at. No. x=0165 (a) 178 (b)  $1 \times 3 + a + 6 \times (-1) = 0$ ,  $\therefore a = +3$ 179 (a) 166 (a) In  $NaOC_2H_5$ , Na is attached to O-atom. 167 **(b)** In  $[Mn(H_2O)_6]^{2+}$ , Mn is present as  $Mn^{2+}$  or Mn 180 **(b)**  $3C_2H_2 \xrightarrow{\Delta} C_6H_6$ (II), so its electronic configuration 182 **(b)**  $=1s^2, 2s^22p^6, 3s^23p^63d^5$ 3*d* 4p183 (a) 4d186 (d) In  $[Mn(H_2O)_6]^{2+}$ , the coordination number of Mn 0 is six, but in presence of weak field ligand, there Ш will be no pairing of electrons in 3*d*. So, it will form high spin complex due to presence of five unpaired electron. OH  $\ln [Mn(H_20)_6]^{2+}$ unpaired electrons 187 (c) 170 (b) Due to aromatic nature;  $C_6H_5CH_2OH$  is exception nature. and does not burn with sooty flame. 189 (d) 171 (c) EDTA (Ethylenediaminetetraacetic acid) 191 **(b)** CH2COO H<sub>2</sub>CCOO CH<sub>2</sub>COO<sup>-</sup> It is H<sub>2</sub>CCOO 192 (a) hexadentate (6 electron pairs) that's why for octahedral complex only one EDTA 193 (d) is required. 173 (c) complex. It is Friedel-Crafts reaction. 194 (d) 174 (a) Tartaric acid is Resonance in phenoxide ion makes it more stable. More stable is ion less stable is phenol or more is acidic nature. 175 (c) Triethylenediamine cobalt(III) chloride is

 $[Co(NH_2CH_2CH_2NH_2)_3]Cl_3; NH_2CH_2CH_2NH_2 is$ bidentate ligand and thus, coordination no. =  $3 \times$ CO is a neutral ligand, so the oxidation state of metal in metal carbonyls is always zero.  $\operatorname{FeCl}_3 + \operatorname{Cl}_2 \longrightarrow \operatorname{FeCl}_4^- + \operatorname{Cl}^+$  $[Ni(CN)_4]^{2-}$  has  $dsp^2$ -hybridization while  $[Ni(Cl_4)^{2-}]$  and  $[Ni(CO_4)]$  have  $sp^3$ -hybridization.  $Cr^{2+}$ ,  $Mn^{2+}$ ,  $Fe^{2+}$  and  $Ni^{2+}$  have 4, 5, 4 and 2 unpaired electrons respectively. It is a reason for the fact.  $CH_3CH_2 - C - CH_2CH_3 \leftrightarrow$ (keto form)  $CH_3 - CH = C - CH_2CH_3$ (enol form) Non-polr part C<sub>6</sub>H<sub>5</sub> —shows more hydrophobic All involve  $d^2sp^3$ -hybridization. Aromatic amines are less basic than aliphatic amines. Also presence of electron attracting group decreases the basic character of aromatic amines. Follow IUPAC rules. All are weak field ligands and thus, give high spin

HO 
$$- C - C - COOH$$
  
HO  $- C - C - COOH$   
HO  $- C - C - COOH$   
H

2,3-dihydroxybutane-1,4-dioic acid

#### 195 (a)

$$\begin{split} \beta_4 \mbox{ for } [ML_4]^{2-} \mbox{can be written as} \\ \beta_4 = & \frac{[ML_4]^{2-}}{[M^{2+}][L^-]^4} = 2.5 \, \times \, 10^{13} \end{split}$$

The overall formation equilibrium constant can be written as

$$k = \frac{[ML_4]^{2^-}}{[M^{2^+}][L^{-1}]^4}$$
  
$$k = \beta_4 = 2.5 \times 10^{13}$$

## 196 (d)

 $[Cr(NH_3)_4Cl_2]^+$ Let oxidation state of Cr = x $NH_3=0$ Cl = -1Net charge =+1 $\therefore [Cr(NH_3)_4Cl_2]^+$ 

## 197 (b)

:.

Phenols are acidic; alcohols are neutral.

x = +3

#### 198 **(b)**

 $2 \times a + 4 \times (-2) + 2 \times 0 + 2 \times 0 = -$ = +3

 $x+4 \times 0+2(-1)=+1$ 

## 199 (c)

CH<sub>3</sub>MgI (Grignard reagent) is an organometallic compound due to C— Mg bond.

#### 200 (c)

Effective atomic number =electrons in Cr<sup>3+</sup> +electrons form 6NH<sub>3</sub> ligands.

 $=21+6\times2=33$ 

## 203 (d)

Hückel rule for aromaticity suggests that an aromatic compound must possess  $(4n + 2)\pi$ electrons, where n = 0, 1, 2..., etc., as well as  $\pi$ electrons cloud should embrace all the carbon atoms of the cyclic systems.

# 204 (a)

 $\bigcirc C_6H_5OH + Zn dust \rightarrow C_6H_6.$ 

#### 205 (b)

CH<sub>3</sub>-CO-N

N-bromo-N-chloro ethanamide

#### 206 (d)

It is condensation reaction.

$$\bigcirc -C - CH_3 \xrightarrow{C_2H_5O'}_{-C_2H_5OH} \bigcirc -C - \ddot{C}H_2 \xrightarrow{CH_3}_{-C_2H_5OH} \bigcirc -C - \ddot{C}H_2 \xrightarrow{CH_3}_{-C_2H_5OH} \bigcirc -C - CH_2 \xrightarrow{CH_3}_{-C_2H_5OH} \frown -C - CH_2 \xrightarrow{CH_5OH} \frown -C - CH_2 \xrightarrow{$$

#### 207 (d)

Due to more canonical forms.

208 (c)

$$2C_6H_5SO_2.0H \xrightarrow{P_2O_5} (C_6H_5SO_2)_2O$$

209 (b)

Isomeric substances that differ only in the arrangement of carbon atoms forming the base chain are known as chain isomers

CH<sub>3</sub>CH<sub>2</sub>CHCH<sub>2</sub>CH<sub>3</sub> 

Cl

$$Cl$$

$$CH_3 - CH_2 - C - CH_3$$

$$CH_2$$

If the compound with the same molecular formula differ in the position of the same functional group on the identical base chain the compounds are called position isomers

## 210 (c)

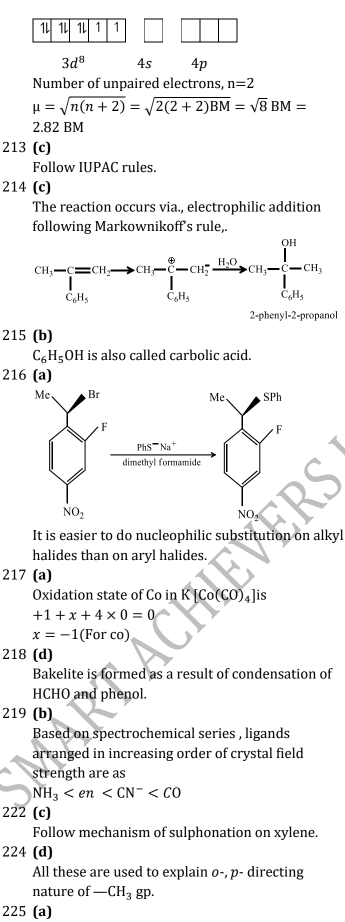
In  $K_3$ [Fe(CN)<sub>6</sub>], the ligands are negative which is present in coordination spheres shows a dual behavior. It may satisfied both primary and secondary valencies while, neutral ligand satisfied only secondary valencies

#### 211 (c)

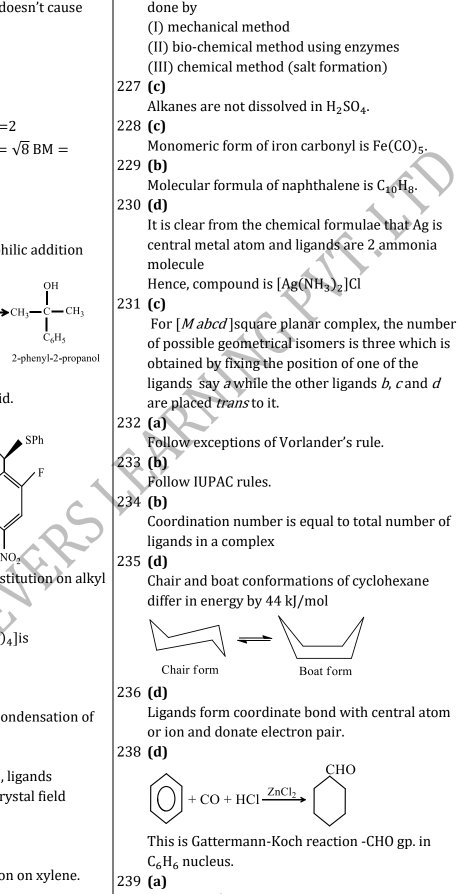
A number of transition metals form polymetallic carbonyls.

 $Ni: 3d^84s^2 \quad Ni^{2+}: 3d^8$ 

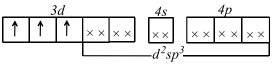
Since, Cl is a weak field ligand, it doesn't cause paring of electron.



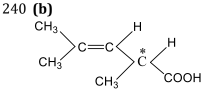
The separation of racemic mixture back into d and l isomers is known as resolution. It can be



 $[Cr(NH_3)_6]^{3+}$  has three unpaired electrons. Electronic configuration of  $Cr^{3+}$  in  $Cr(NH_3)_6$  is:



 $\times \times$  Electron pair donated by NH<sub>3</sub>.



The above compound has chiral centre Hence, it can exhibit optical isomerism while geometrical isomerism is not possible due to presence of identical groups on double bonded carbon atom

#### 241 (a)

It has no unpaired electron.

#### 242 (a)

COOH | H - C - OH | plane of symmetry H - C - OH | COOH

*Meso* tartaric acid is optically inactive due to the presence of molecular symmetry. It I optically inactive due to internal compensation, *ie*, the effect of one half of the molecule is neutralized by other

#### 243 **(b)**

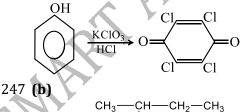
Smaller is cation, more is effective nuclear charge, more is the tendency to attract electron pair from ligands.

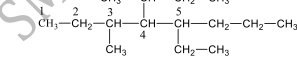
#### 245 (a)

Presence of *o*-, *p*-directing gp. facilitates the SE reactions.

#### 246 **(c)**

 $K_2S_2O_8$  gives quinol; KMnO<sub>4</sub> gives mesotartaric acid.



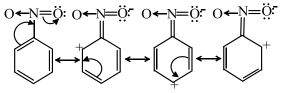


The compound is substituted octane, it has branches at carbon-3, carbon-4, carbon-5. The name is

4-sec-butyl-5-ethyl-3-metyl octane 248 (a)

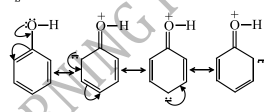
The presence of *m*-directing groups in benzene

nucleous simply decreases electron density at *o*and *p*-, whereas no change in electron density at *m*-position is noticed.



On the contrary *o*-and *p*-directing groups in nucleus increases the electron density at *o*- and *p*-position.

Thus, presence of o- and p-directing groups provide seats for  $S_E$  reactions or activates the ring, whereas presence of m-directing groups does not activate the ring and thus, deactivate the ring for  $S_E$  reactions



249 **(c)** 

 $CuF_2$  is blue coloured crystalline solid. 250 (d)

- Different ioniz
- Different ionization gives different colour.

251 (d)

The complex has coordination number of six which is found in octahedral complex.

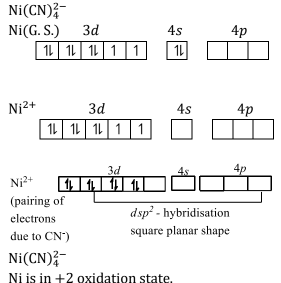
## 252 **(c)**

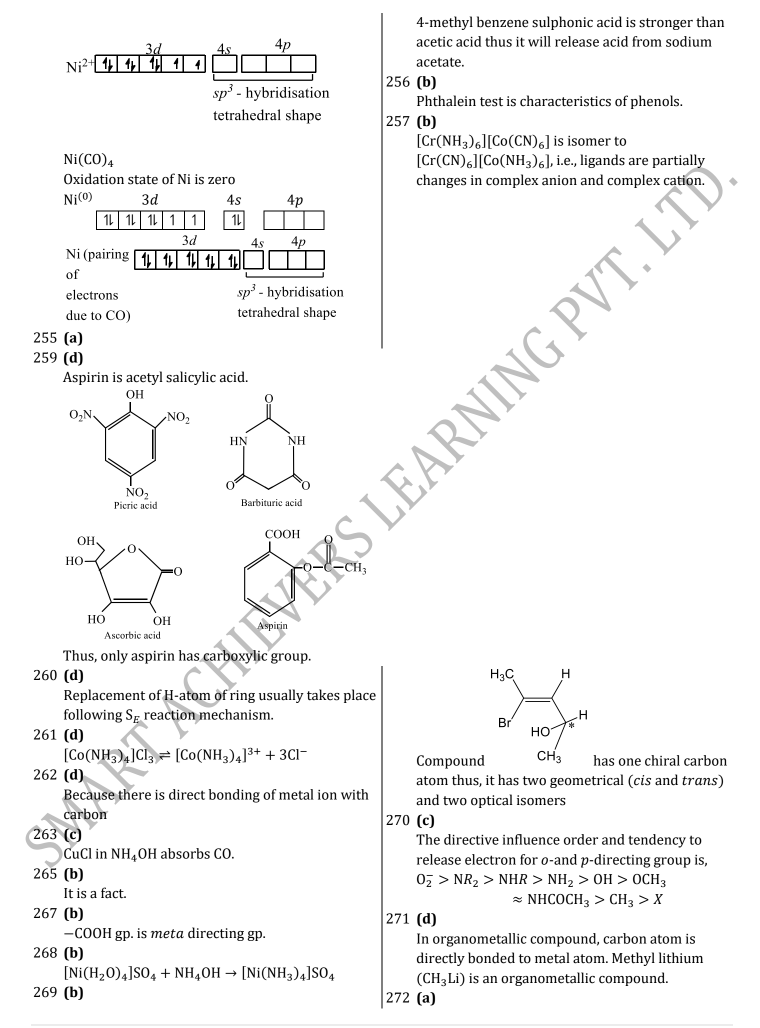
 $\mathrm{BF}_3$  has incomplete octet and is Lewis acid; it cannot donate electron pair.

## 253 **(c)**

Methyl thiomethyl group is inserted at *ortho* position by heating phenol with dimethyl sulphoxide and pyridine  $-SO_3/(CH_3CO)_2O$ .

## 254 **(a)**





This is crossed Cannizzaro's reaction in which HCHO is oxidized.

$$C_6H_5CHO + HCHO \xrightarrow{NaOH} C_6H_5CH_2OH + HCOONa$$
.

273 **(d)** 

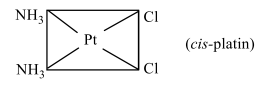
Gammexane is  $C_6H_6Cl_6$ .

274 **(b)** 

Each ligand donates one electron pair.

275 **(c)** 

7. is isomer of  $[Pt(NH_3)_2Cl_2]$  which is used as an anticancer drug for treating several types of malignant tumours.



## 276 **(d)**

$$C_6H_5CHO \xrightarrow{\text{Reduction}} C_6H_5CH_2OH.$$
  
Benzyl alcohol.

277 **(d)** 

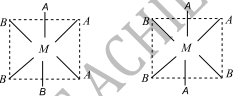
Petroleum and coal are main sources of aromatic compounds.

## 279 **(c)**

Butane-1,2,4-tricarbonitrile

280 **(b)** 

*Fac-mer* isomerism is associated with  $[MA_3B_3]$  type complexes.



Cis isoment (-Fac isomer) trans isoment (Mer- isomer)

282 **(c)** 

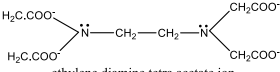
Intramolecular H-bonding gives rises to lower m.p.

## 283 **(b)**

Mn does not form mononuclear carbonyl.

284 **(b)** 

Ethylenediamine tetraacetic acid is a hexadentate ligand because it has six donor centres.



ethylene diamine tetra acetate ion

285 **(b)** 

H H | | (i) - H - C - C - Cl 1,1-dichloro ethane | | H Cl (ii) H H | | H - C - C - H 1,2-dichloro ethane

Both are position isomers

Cl Cl

286 <u>(</u>c)

| Hybridisatio                   | Geometry of   |  |  |
|--------------------------------|---------------|--|--|
| n                              | complex       |  |  |
| sp <sup>3</sup>                | Tetrahedral   |  |  |
| $dsp^2$                        | Square planar |  |  |
| $d^2sp^3$                      | Octahedral    |  |  |
| sp <sup>2</sup> d <sup>2</sup> | Not possible  |  |  |

287 (b)

```
CH_3 - CH = CH - COOC_2H_5 is
Ethyl-2-butenoate
```

288 (d)

Waxes are not obtained obtained by destructive distillation of wood or coal.

289 **(c)** 

Formaldehyde and benzophenone are also obtained.

290 **(b)** 

It ionizes to  $Fe^{3+}$  and  $SO_4^{2-}$ .

## 291 **(d)**

These are the concepts of Werner's theory.

## 293 **(b)**

In K<sub>3</sub>[Co(CO<sub>3</sub>)<sub>3</sub>], cobalt shows the +3 oxidation state *i.e.*,  $(d^6)$  ion. Hence, Co (+3) has four unpaired electrons so, it is paramagnetic. The magnetic moment of Co(+3)

In K<sub>3</sub>[Co(CO<sub>3</sub>)<sub>3</sub>] = 
$$\sqrt{n(n+2)}$$
 BM  
=  $\sqrt{4(4+2)}$  BM=4.9 BM

Where, n=number of unpaired electrons  $CO_3^{2^-}$  is a weak field bidentate ligand, so  $3CO_3^{2^-}$ ligands occupy six orbitals, thus it shows  $sp^3 d^2$ hybridisation and octahedral in shape.

## 296 **(a)**

Some heterocyclic compounds (hetero aromatics) possess aromatic nature. Follow Hückel rule.

## 297 **(c)**

Nickel reacts with dimethylglyoxime to give red ppt. of nickel-dimethyl glyoxime complex.

$$\begin{aligned} 3d^{\theta}. \\ 3d$$

 $Cr^0 = 3d^4$ ,  $4s^2$ . Effective configuration = 10.

Page | 117

spin octahedral complex.

erythro-isomer has identical

Chief

constituents

Temp.

range

| 1. | Light oil or  | 80-   | Benzene,          |
|----|---------------|-------|-------------------|
|    | crude         | 170°C | toluene, xylenes, |
| 2. | naphtha       |       | etc.              |
|    | Middle oil or | 170-  | Phenol,           |
| 3. | carbolic acid | 230°C | naphthalene,      |
|    | Heavy oil or  |       | pyridine, etc.    |
| 4. | creosote oil  | 230-  | Cresols,          |
|    | Green oil or  | 270°C | naphthalene       |
| 5. | anthracene    |       | quinolone, etc.   |
|    | oil           | 270-  | Anthracene,       |
|    | Pitch         | 360°C | phenanthrene,     |
|    |               |       | etc.              |
|    |               | Resid | 90-94% of         |
|    |               | ue    | carbon            |

## 319 (d)

The order of reactivity depends on the stability of intermediate carbocation formed due to heterolytic cleavage of C-X bond.

#### 320 (d)

Greater is the number of chelate rings, greater is stability of the chelate. Hence, five fused cyclic system is most stable for a chelate.

#### 321 **(b)**

 $HNO_3 + H_2SO_4$  acts as nitration mixture.

#### 323 (d)

The negative charge density on V-atom favours easy electron pair donation.

#### 324 **(b)**

Due to acidic nature.

#### 325 **(d)**

If a substance rotates the plane polarised light in clockwise direction it is dextrorotatory (+). If it rotates the plane polarised light in anticlockwise direction then it is laevorotatory (-)

#### 326 **(c)**

 $C_6H_6 \xrightarrow{\text{HCN+HCl}} C_6H_5\text{CH} = \text{NH} \xrightarrow{\text{HOH}} C_6H_5\text{CHO}$ 

EAN of Fe in  $K_4$ Fe(CN)<sub>6</sub> = 26 - 2 + 2 × 6 = 36; the at. no. of next inert gas.

#### 329 **(b)**

Both Ni (CO)<sub>4</sub> and Ni(PPh<sub>3</sub>)<sub>2</sub>Cl<sub>2</sub> have  $sp^3$ -hybridisation

#### 331 (d)

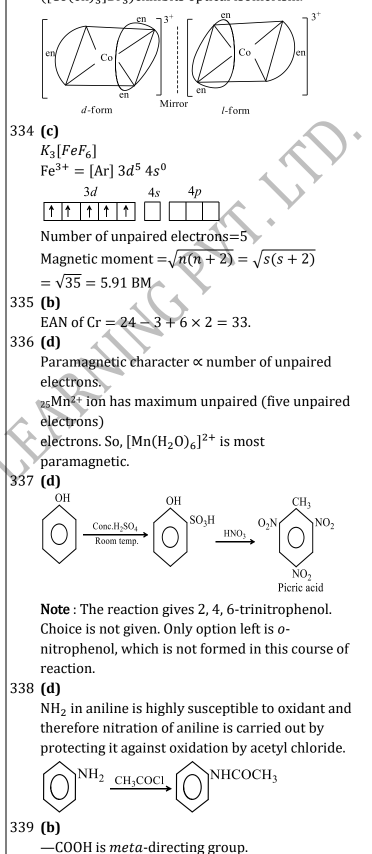
All are examples of strong ligand field, because all have greater value of  $\Delta$ ;  $\Delta$  represents the strength of ligand field.

#### 332 (a)

333 (d)

Due to rearrangement because 2° carbon is more stable than 1° carbon.

Tris –(ethylenediamine) cobalt(III) bromide ([Co(en)<sub>3</sub>]Br<sub>3</sub>)exhibits optical isomerism.



340 (d)

 $[Co(NO_2)(NH_3)_5]Cl_2$ 

341 **(b)** 

The ionisation isomer of

 $[Cr(H_2O)_4Cl(NO_2)]Cl$  is  $[Cr(H_2O)_4Cl_2](NO_2)$ because of exchanging of ligand and counter ions. 342 (c)

# All can be prepared from phenol.

## 343 (a)

Only iodobenzene gives Ulmann's reaction,

$$2C_6H_5I \xrightarrow{Cu} C_6H_5 - C_6H_5$$

## 344 **(c)**

Fischer projection can be manipulated by rotating a group of any three ligands in clockwise (D) or anticlockwise (L) direction, the fourth ligand does not change its position

## 345 **(b)**

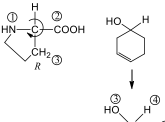
A strong filed ligand produces low spin complexes.

## 347 **(a)**

Ni in  $[Ni(H_2O)_6]^{2+}$  has two unpaired electrons in it.

## 348 **(b)**

Compounds in which a chiral centre is part of a ring are handed in a analogous fashion



 $C_6H_6 + CH_3Cl \xrightarrow{AlCl_3 anhy.} C_6H_5CH_3$ 

352 **(b)** 

The formula of hexamine copper (II) sulphate  $is[Cu(NH_3)_6]SO_4$ . It dissolve in water as

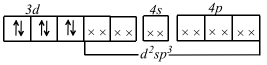
$$[Cu(NH_3)_6]SO_4 \underbrace{[Cu(NH_3)_6]^{2+} + SO_4^2}_{2 \text{ ions}}$$

## 354 **(b**)

Complex of type [M(AA)<sub>3</sub>] show optical isomerism.

## 355 **(b)**

Electronic configuration of  $Fe^{2+}$  in  $K_4Fe(CN)_6$  is:



 $\times$  × Electron pair donated by CN<sup>-</sup>.

356 **(d)** 

$$\begin{array}{c} \operatorname{CH}_3 - \operatorname{CH}_2 - \operatorname{C} \equiv \operatorname{C} - \operatorname{CH} = \operatorname{CH}_2\\ 6 & 5 & 4 & 3 & 2 & 1 \end{array}$$

357 **(b)** 

Oxidation state of Ni in  $K_2NiF_6$  is +4; the highest among all.

## 359 **(d)**

Substituents always get higher number than the principal functional group while, numbering the longest possible chain

## 360 **(d)**

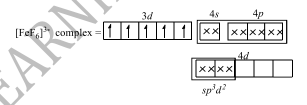
Draw different isomers.

## 362 **(b)**

Metal carbonyl organometallic compounds possess both  $\sigma$ -and  $\pi$ -characters.

## 363 **(b)**

 $\rm F^-$  is a weak field ligand and  $\rm [FeF_6]^{3-}$  is an outer-orbital complex



 $[FeF_6]^{3-}$  shows  $sp^3 d^2$  hybridisation and  $Fe^{3+}$  has five unpaired electrons.

In  $[Fe(CNS)_6]^{3-}$ ,  $CNS^-$  is a strong field ligand and is inner orbital complex.

 $[Fe(CNS)_{6}]^{3-1}$ 

 $sp^3d^2$  - hybridisation

 $[Fe(CNS)_6]^{3-}$  shows  $d^2 sp^3$  hybridisation and has one unpaired electron.

Hence, the reaction.

 $[Fe(CNS)_6]^{3-} \rightarrow [FeF_6]^{3-}$ 

takes place with increase in magnetic moment. 364 **(a)** 

Presence of three  $-NO_2$ gp. in chlorobenzene activates Cl atom to show  $S_N$  reactions.

365 **(a)** 

Complex compounds or complex salts containing two different metallic elements give tests for only one element. For example, potassium hexacyanoferrate (II),  $K_4$ [Fe(CN)<sub>6</sub>] gives tests

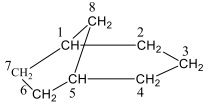
only for K<sup>+</sup> ions and not for Fe<sup>2+</sup> ions. 379 (d)  $K_4[Fe(CN)_6] \rightleftharpoons 4K^+ + [Fe(CN)_6]^{4-}$ Coal-tar is source of all these. 366 (a) 380 (c) Only primary valencies are ionized. Halogen attached to benzene nucleus is stabilized  $[Co(NH_3)_5Br]SO_4 \xrightarrow{BaCl_2} [Co(NH_3)_5Br]^{2+} + BaSO_4$ due to resonance. 382 (a)  $[Co(NH_3)_5SO_4]Br \xrightarrow{AgNO_3} [Co(NH_3)_5SO_4] + AgBr$  $-NO_2$  group is reduced to  $-NH_2$  by Sn/HCl. 0.01 mole of each by 0.01 mole of reactants. 383 (d) 367 (c) Each central atom attains the EAN equal to at. No. Wilkinson's catalyst, (Ph<sub>3</sub>P)<sub>3</sub>RhCl of next inert gas Kr, i.e., 36.  $RH^{+} = [Kr] 4d^{8}s^{0}$ 384 (c) *ie*,  $dsp^2$  hybridisation A bidentate ligand has two donor sites available Rh atom in Wilkinson's catalyst is  $dsp^2$  hybridised for coordination, e.g., giving a square planar shape to the molecule COO 368 (d) ; NH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NH<sub>2</sub>  $H_2O$  is weak field ligand, thus  $Co^{2+}$  has only 3 COO unpaired electrons. 385 (d) 369 (a)  $-N_2$ Cl is reduced to H by either of these reducing If an enantiomerically pure acid is treated with agents. racemic mixture of an alcohol having a chiral 386 **(c)** carbon, the product formed will be optically active Fe is present in the form of complex ion, *i.e.*, mixture  $[Fe(CN)_6]^{3-}$  which is not ionized to  $Fe^{3+}$  and  $CN^-$ . 370 (d) 387 (a) Cyclopropane is most strained since it has a  $[Co(NH_3)_5SO_4]Br \rightleftharpoons [Co(NH_3)_5SO_4]^+ + Br^$ maximum angle strain of  $24^{\circ} - 44'$  $[Co(NH_3)_5Br]SO_4 \rightleftharpoons [Co(NH_3)_5Br]^{2+} + SO_4^{2-}$ 371 (c) The molecular formula of both of the above CHO compounds is same but on ionisation they give different ions in solution, so they are called ZnCl<sub>2</sub> + CO + HCIionization isomers. 390 (c) Phenols are weak acids and do not react with this is Gattermann-Koch reaction to introduce  $NaHCO_3$  (a weak base). -CHO gp. in  $C_6H_6$  nucleus. 391 (b) 372 (a)  $C_6H_5CH = CHCOOH$  is cinnamic acid; it has Mn in  $Mn(CN)_6^{4-}$  has configuration: unsaturation. 392 (a) Magnetic moment of  $K_{3}[Fe(CN)_{6}] = 1.7 BM$  $d^2 s n^3$ Magnetic moment= $\sqrt{n(n+2)}$ 373 (d) *n* =number of unpaired electrons present in It does not ionize to give Cl<sup>-</sup> ions and thus, white molecule ppt. of AgCl will not be obtained.  $1.7 = \sqrt{n(n+2)}$ 375 (c)  $-n^2 + 2n - 2.89 = 0$  then n = 0.97 or 1  $\bigvee$  [Pt(C<sub>2</sub>H<sub>4</sub>)Cl<sub>3</sub>]  $x + 0 + (-1) \times 3 = 0$ 393 (c) x + (-3) = 0x = +3377 (c) –ĊН—СН<sub>2</sub>—ĊН<sub>2</sub>  $Ni^{2+}$  has two unpaired (3 $d^8$ ) electrons.  $CN^-$  is This compound contains 9 carbon atoms and strong field ligand and thus all the eight electrons are paired giving  $dsp^2$ -hybridisation.

corresponding alkane is nonane. Three bridges

contain 5, 2 and 0 carbon atoms. Therefore, the name of the compound is bicyclo [5.2.0] nonane

#### 395 (c)

The compound have structure as written below



bicyclo [3.2.1] octane

396 (d)

It is *m*-directing gp.

397 (a)

Carbylamines reaction.

 $\begin{array}{l} C_6H_5SO_3Na + NaOH \longrightarrow C_6H_5ONa + NaHSO_3\\ C_6H_5ONa \xrightarrow{HOH}{H^+} C_6H_5OH^+NaOH \end{array}$ 

#### 401 (a)

The stability order of conformations of cyclohexane is Chair > twist boat > boat > half chair

#### 402 **(b)**

Phenolic group is susceptible for oxidation and thus, to obtain *o*- and *p*-nitrophenol dil.  $HNO_3$  is used in place of conc.  $HNO_3 + H_2SO_4$ .

#### 403 (c)

Lower is mol. Wt. lower is b.p., also 1, 2-dihydroxy benzene show chelation and thus have lower b.p. than 1,3 and 1,4-derivatives.

enzene

<110°C

<

Phenol 1,2-dihydroxyb 1,3-dihyroxyb 1,4-dihydroxyb

enzene

< 105

enzene

m.p. 43°C

- <170°C b. p.
- <

405 **(b)** 

 $K_2[PtCl_6] \rightleftharpoons 2K^+ + [PtCl_6]^-$ 

407 **(b)** 

Phenols are weakly acidic due to resonance. 408 **(b)** 

Follow IUPAC rules.

#### 409 (a)

Compounds having coordination number six and following the general formula show geometrical and optical isomerism.

 $M_{A_4B_2}, M_{A_4BC}, M_{A_3B_3}$  and  $M_{(AA)_2B_2}$  show geometrical isomerism and

 $M_{A_2X_2Y_2}, M_{A_2X_2YZ}, M_{A_2XYZL}$  $M_{ABXYZL}, M_{(AA)_3}, M_{(AA)_2}X_2$  show optical isomerism. (AA) is bidentate ligand.

#### 410 **(b)**

 $[Co(NH_3)_4Cl_2]Cl$ , Its IUPAC name is tetraammine dichloro cobalt III chloride.

#### 411 **(c)**

The ease of hydrolysis depends upon the magnitude of the +ve charge on the carbonyl group.

## 412 **(d)**

All are the common uses of nitrobenzene.

#### 413 **(b)**

$$K_4[Ni(CN)_4] \rightarrow 4K^+ + [Ni(CN)_4]^{4-}$$
  
x+(4×-1)=-4  
x-4--4

414 **(a)** 

Presence of -OH gp. in  $C_6H_6$  nucleus increases acidic nature.

# 416 **(b)**

Halogen attached on side chain behaves as in aliphatic molecule.

## 417 **(d)**

Cyclohexane is an aliphatic cyclic compound.

12. In 
$$[Cu(NH_3)_4]^{2+}$$
, Cu is present as  $Cu^{2+}$ 

$$\mathrm{Cu}^{2+} = [\mathrm{Ar}]3d^94s^0$$

$$[Cu(NH_3)_4]^{2+} = [Ar]$$

 $(NH_3 being a strong field ligand shifts one electron from 3$ *d*-orbital to 4*p*-orbital.)

- 13. In  $[Ni(CO)_4]$ , CO is a neutral ligand
- 14. In  $[Fe(CN)_6]^{3-}$ , Fe is present as  $Fe^{3+}$ .

$$\mathrm{Fe}^{3+} = [\mathrm{Ar}]3d^54s^0$$

$$[Fe(CN)_6]^{3-} = [Ar]$$

$$\overbrace{1411}^{3d} \times \times \xrightarrow{4s} \xrightarrow{4p}_{x \times x}$$

Thus, its hybridization is  $d^2sp^3$  not  $sp^3d^2$ , i.e., it is

an inner orbital complex.

15.  $[Co(en)_3]^{3+}$  contains total 36 electrons, *i. e.* follows EAN rule.

#### 420 (c)

Optical isomerism is very common in octahedral complexes having general formula

$$\begin{bmatrix} M_{A_2B_2C_2} \end{bmatrix}^{n-}, \begin{bmatrix} M_{ABCDE} \end{bmatrix}^{n-}, \begin{bmatrix} M_{(AA)_3} \end{bmatrix}^{n-}, \\ \begin{bmatrix} M_{(AA)_3B_2} \end{bmatrix}^{n-} \begin{bmatrix} M_{(AA)_3BC} \end{bmatrix}^{n-} \text{ and } \begin{bmatrix} M_{(AB)_3} \end{bmatrix}^{n-} \\ \text{where } AA \text{ is symmetrical bidentate ligand like } \\ \text{COO}^- \\ \text{and } AB \\ \text{COO}^- \end{bmatrix}$$

is unsymmetrical bidentate ligand.

#### 421 (c)

 $[Co(NH_3)_5NO_2]Cl_2 \rightleftharpoons [Co(NH_3)_5NO_2]^{2+} + 2Cl^{-}$ 2Cl<sup>-</sup> + Ag<sup>+</sup>(excess)  $\rightarrow$  2AgCl  $\downarrow$ 

422 **(a)** 

 $C_6H_5C$  is benzo gp.

#### 423 (c)

It is the reason for given fact.

#### 424 **(b)**

 $d^6$ -cation with low spin has electronic

configuration  $t_{2g}^6 e_g^0$ .

Total energy=  $(-0.4 \Delta_0 \text{ per } e^- \times 6) + (e^- \text{ pairing energy of 3 pairs})$ 

 $= -2.4 \Delta_0 + 3P$  $= -\frac{12}{5}\Delta_0 + 3P$ 

#### 426 **(c)**

It is DDT, *i*. *e*., p, p' -dichloro diphenyl trichloroethane.

#### 428 **(b)**

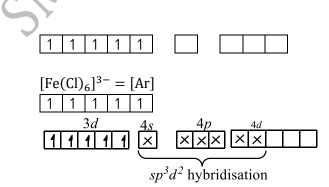
 $[Cr(NO_2)(NH_3)_5]$  Cl show linkage isomerism.

$$NO_2^-$$
 can link  $O-N=O$  or  $N = O$ 

#### 429 (d)

Cl, being a weak field ligand, does not cause pairing of *d*-electrons of the metal atom and thus, forms outer orbital complex as.

In  $[Fe(Cl)_6]^{3-}$ , Fe is present as  $Fe^{3+}$  $Fe^{3+} = [Ar]3d^5 4s^0 4p^0$ 



#### 430 (d)

<sub>p</sub>*K*<sub>a</sub> for (a), (b), (c) and (d) are 4.17, 4.09, 3.49 and 3.43 respectively.

## 431 **(a)**

 $[\mathrm{H_2EDTA}]^{2-} + \mathrm{Mg}^{2+} \rightarrow [\mathrm{MgEDTA}]^{2+} + 2\mathrm{H^+}$ 

- 16. In this complex, four donor sites are occupied by oxygen and two donor sites are occupied by nitrogen.
- 17. This complex is six coordinated.
- 18. Complex  $[MgEDTA]^{2-}$  is colourless.
- 19. Increase in [H<sup>+</sup>]decreases pH of the solution.

433 (c)

Different compounds having the same molecular formula but different properties are called isomers

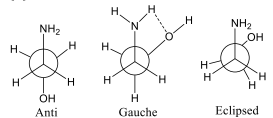
#### 434 **(c)**

Numbering will be done from this end because both are side chains and  $- \text{OCH}_3$  is smaller than  $- \text{OC}_2\text{H}_5$ 

3-ethoxy-1-methoxy-propane

435 **(a)** 

Each carbon in benzene is  $sp^2$ -hybridized. 436 **(b)** 



So, gauche form stabilized by intermolecular hydrogen bonding hence, gauche is more stable than anti

437 **(a)** 

 $AgCN + KCN \rightarrow K[Ag(CN)_2].$ 

438 **(b)** 

Carbocyclic compounds which resemble aliphatic compounds in their properties are called alicylic compounds

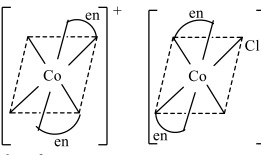
## $-NO_2$ gp. is deactivating gp.

#### 440 **(b)**

The molecule contains three chlorine atoms out of which only two are ionized.

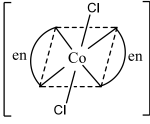
#### 441 **(b)**

 $[Co(en)_2Cl_2]^+$  have three optical isomers which are given below.



*d*-*cis* form





trans-meso form

#### 442 **(c)**

The reaction carried out in alkaline pH, *ie*, 9 – 11 444 **(d)** 

 $NO_2$  gp. withdraws electrons from o –and p-position and thus, deactivates the ring. This deactivation stabilises the negatively charged intermediates formed during reaction and thus, replacement of—Cl becomes easier.

#### 446 **(a)**

 $Ti(C_2H_5)_4$  is an organometallic compound because there is direct bonding of metal ion with carbon.

#### 447 (a)

Octahedral  $Co(NH_3)_4Br_2Cl$  shows ionisation and geometrical isomerism.

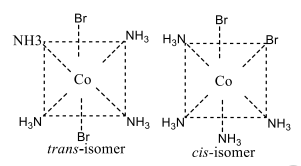
In ionisation isomerism ligands show different coordination sphere and the anions present outside the coordination sphere.

These are exchanged with each other as follows  

$$Co(NH_3)_4Br_2Cl \approx [Co(NH_3)_4Br_2]^+ + Cl^-$$

$$[Co(NH_3)_4BrCl]Br \rightleftharpoons [Co(NH_3)_4BrCl]^+ + Br^-$$
  
II

In geometrical isomerism, coordination number of central atom (cobalt) is six and shape is octahedral, so it shows following geometrical isomers.



## 448 **(a)**

Aliphatic amines are stronger base than aromatic amines.

449 **(b)** 

Follow Hückel rule.

$$C_6H_5CHCl_2 \xrightarrow{HOH} C_6H_5CHO$$

453 **(c)** 

Cr has coordination no. 6 in its carbonyl and other complexes.

## 454 **(d)**

Neutral FeCl<sub>3</sub> (aq.) gives violet coloured complex with phenol.

455 **(a)** 

 $[CoF_6]^{3-}$  is an outer complex having  $sp^3d^2$ -hybridization.

456 **(d)** 

$$[Sc(CN)_6]^{3-}$$
  
Sc=21=1s<sup>2</sup>, 2s<sup>2</sup>, 2p<sup>6</sup>, 3s<sup>2</sup>, 3p<sup>6</sup>, 4s<sup>2</sup>, 3d<sup>1</sup>

3d

 $Ni(CN)_4^{2-}$ 

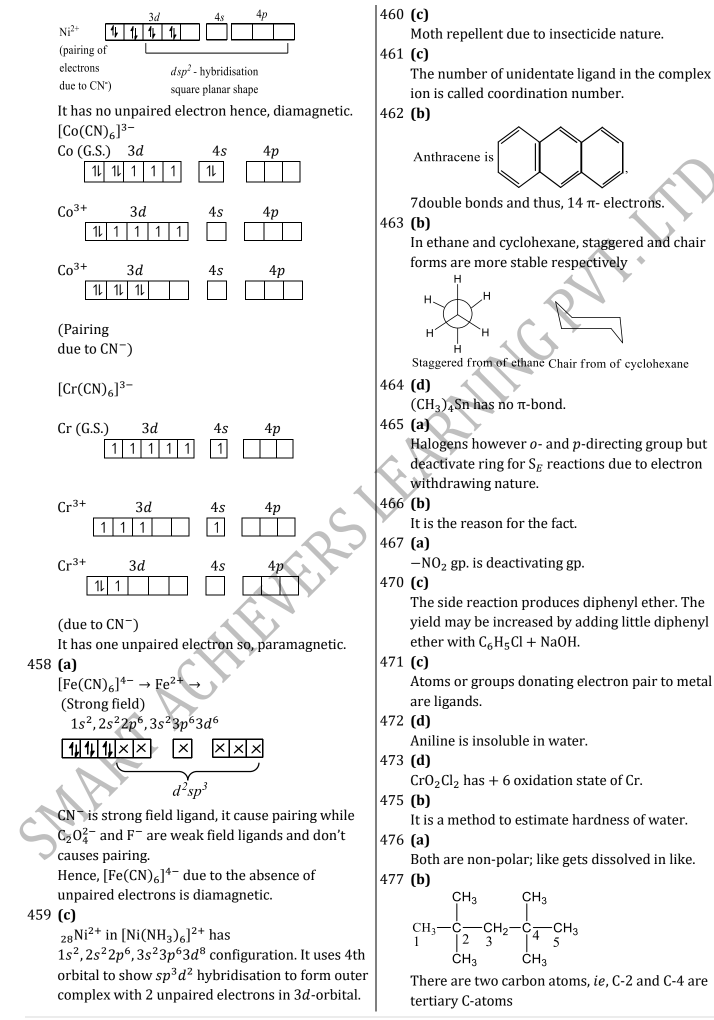
 $Sc^{3+} =$ 

Ni(G. S.)
 
$$3d$$
 $4s$ 
 $4p$ 

 11
 11
 1
 11

Ni<sup>2+</sup> 3d 4s 4p   

$$1 1 1 1 1 1$$



478 **(b)** 

Rosenmund's reaction.

479 **(c)** 

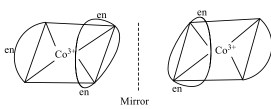
 $C_6H_5CHO \xrightarrow{Zn-Hg/HCl} C_6H_5CH_3$ 

480 **(c)** 

 $F^-$  has lowest  $\Delta_o$  value depending upon the splitting power of *d*-orbitals

481 **(b)** 

Complex  $[Co(en)_3]^{3+}$  has no plane of symmetry and centre of symmetry that's why it is optically active.



482 **(a)** 

It is preparation of DDT.

483 **(a)** 

Light oil mainly contains  $C_6H_6$ ,  $C_7H_8$ ,  $C_8H_{10}$ , etc. 484 **(c)** 

Coordination number is the number of ligand  $\sigma$ bonded to metal-atom. Hence, coordination number of X in  $[X(SO_4)(NH_3)_5]$ Cl is 6. Let oxidation state of X in the complex be 'y' then y+(-2)+5(0)+(-1)=0

y - 2 - 1 = 0y - 2 - 1 = 0v = +3

485 **(a)** 

 $C_6H_5CH_3 \xrightarrow{Cl_2} C_6H_5CH_2Cl$ 

In presence of light substitution occurs is side chain.

## 486 **(a)**

 $[CoCl_3(NH_3)_3]$  cannot ionize in solution because three chloride ions satisfy primary and secondary valencies. It will not be precipitated by the addition ofAgNO<sub>3</sub>.

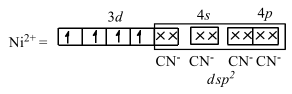
487 (d)

For  $[Ni(CN)_4]^{2-}$ , oxidation state of Ni is +2.

CN<sup>-</sup>=strong field ligand Ni<sup>2+</sup>(ground state)=

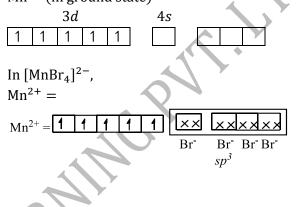
$$\begin{array}{c|c} 3d^8 & 4s^0 \\ \hline 1 & 1 & 1 & 1 \\ \hline \end{array}$$

 $In [Ni(CN)_4]^{2-}$  $Ni^{2+} =$ 

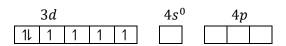


*dsp*<sup>2</sup> hybridisation, *i.e.*, square planar geometry, zero unpaired electron, *i.e.*, zero magnetic moment

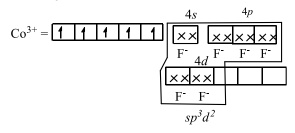
For  $[MnBr_4]^{2-}$ , oxidation state of Mn is +2. Br<sup>-</sup>= weak field ligand Mn<sup>2+</sup> (in ground state)



 $sp^3$  hybrisation, *i.e.*, tetrahedral geometry, five unpaired electrons, *i.e.*, magnetic moment=5.9 Co<sup>3+</sup> in ground state=



For  $[CoF_6]^{3-}$ , oxidation state of Co is +3. F<sup>-</sup>=weak field ligand In  $[CoF_6]^{3-}$ 



sp<sup>3</sup> d<sup>2</sup> hybridisation *i.e.*, octahedral geometry four unpaired electrons *i.e.*, magnetic moment is 4.91 BM.

489 **(b)** 

An experimental value.

- 491 **(c)** A commonly used food preservative.
- 492 **(a)**

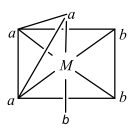
$$C_6H_5ONa + C_2H_5I \rightarrow C_6H_5OC_2H_5 + NaI$$
  
Phenetole

#### 493 (d)

Metal carbonyl are regarded as the coordination compounds formed by the donation of lone pair of electron of CO into the suitable empty orbital of zero valent transition metals such as Ni, Fe etc. Therefore, the M— C bond is coordinate covalent.

#### 494 (d)

When the three ligands (with same donor atoms) are on the same triangular face of the octahedron, the isomer is called *facial* or *fac* isomer. The octahedral complex is *facial* or *fac* isomer.



In this complex, the three ligands are on the same triangular face of the octahedron.

#### 495 (a)

Number of unpaired electrons in  $[Fe(CN)_6]^{4-}$  is zero.

Thus, magnetic moment

 $=\sqrt{n(n+2)}=0$  BM

(*n*=unpaired electrons)

$$n \text{ in } [\text{MnCl}_4]^{2-}=5, \sqrt{35} \text{ BM}$$
  
 $n \text{ in } [\text{CoCl}_4]^{2-}=3, \sqrt{15} \text{ BM}$ 

496 **(a)** 

Orange-red dye is formed with aniline.

499 (a)

It is a fact. Follow ortho effect.

#### 500 (d)

Half chair is transition state conformation between the chair and boat conformation. The energy difference between the chair and half chair conformation being 44 kJ mol<sup>-1</sup>. Hence it is most unstable

## 501 (a)

```
The M—C \pi-bond in metal carbonyl which is
formed by the donation of an electron pair from a
filled d-orbital of metal into the vacant
antibonding \pi-orbital of CO, strengthens the M—
C \sigma— bond. This is called synergic effect and is
usually observed in metal carbonyls. Thus
[Ni(CO)<sub>4</sub>] exhibits synergic effect.
```

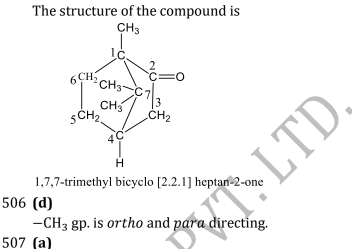
503 **(b)** 

 $Pt \xrightarrow{Aqua \text{ regia}} H_2[PtCl_6] \xrightarrow{\Delta} PtCl_4 + 2HCl$ 

504 (d)

Follow IUPAC rules.

505 **(b)** 



Estimation of calcium and magnesium is done by EDTA.

508 (d)

$$\begin{array}{c} \mathrm{CH}_{3}-\mathsf{CH}-\mathsf{CH}-\mathsf{CH}_{3}+\mathsf{Cl}_{2} \longrightarrow \mathrm{CH}_{3}-\mathsf{CH}-\mathsf{CH}_{2}\mathsf{CH}_{2}\mathsf{CH}_{2}\mathsf{CH}_{2}\mathsf{CH}_{3}\\ \mathsf{CH}_{3} \quad \mathsf{CH}_{3} \quad \mathsf{CH}_{3}\\ \mathsf{2},\mathsf{3}\text{-dimethyl butane} \\ \end{array}$$

Due to the presence of chiral centre it shows the optical activity and its mirror image are non superimposable hence it shows one enantiomer pair

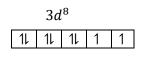
$$\begin{array}{c} \mathrm{CH}_3-\mathrm{CH}-\mathrm{CH}-\mathrm{CH}_2\mathrm{CI}\\ \mathrm{I}\\\mathrm{CH}_3\\\mathrm{CH}_3\\\mathrm{CH}_3\end{array} \qquad \begin{array}{c} \mathrm{CICH}_2-\mathrm{CH}-\mathrm{CH}-\mathrm{CH}_3\\ \mathrm{I}\\\mathrm{CH}_3\\\mathrm{CH}_3\\\mathrm{CH}_3\end{array}$$

509 **(b)** 

 $C_6H_6$  and other aromatic compounds show characteristics  $S_E$  reactions.

510 (d)  

$$CH_2 = CH - CH = CH - CH = CH - CH_3$$
  
 $1 \quad 2 \quad 3 \quad 4 \quad 5 \quad 6 \quad 7$   
Hepta-1,3,5-triene  
511 (b)  
 $1 \quad 2 \quad 3$   
 $CH_2 - CH - CH_2$   
 $| \quad | \quad |$   
 $CHO \quad CHO \quad CHO$   
 $Propane-1, 2, 3$ -tricarbaldehyde  
512 (a)  
The attacking species in sulphonation is SO<sub>3</sub>  
 $H_2SO_4 \rightarrow H_3O^+ + SO_3 + HSO_4^-$   
513 (d)  
 $CHCl_3$  has no reaction with  $Br_2$ .  
514 (c)  
 $[NiCl_4]^{2-}$ ; oxidation number of Ni,  
 $x-4=-2$   
 $\therefore x=+2$   
 $Ni_{(28)} = [Ar]3d^8, 4s^2$ 





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

 $\begin{array}{c|c} \times & \times & \times \\ \hline & sp^3 & sp^3 & sp^3 & sp^3 \\ \end{array}$ 

11

## $sp^3$ -hydrid orbitals, tetrahedral

Cl<sup>-</sup> is a weak ligand and thus unpaired electrons are not paired. Lone pairs from 4Cl<sup>-</sup>are accommodated in four  $sp^3$  hybrid orbitals. *N*=unpaired electron=2, paramagnetic Magnetic moment (spin only)

 $=\sqrt{N(N+2)}$  BM  $=\sqrt{8}$  =

2.828 BM

## 515 **(b)**

Pyridine shows S<sub>E</sub>reactions at position-3 preferentially and at 2,4-positions under specific conditions.

## 516 **(a)**

 $K[Ag(CN)_2] \rightleftharpoons K^+ + [Ag(CN)_2]^1$ 

## 517 **(a)**

Its coordination number will be 6 because it is bonded with three bidentate ligands

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

## 518 **(c)**

The four ions on ionisation are possible only when three Cl<sup>-</sup> are outside the coordination sphere.

#### $[Pt(NH_3)_5Cl]Cl_3 \rightleftharpoons [Pt(NH_3)_5Cl]^{3+} + 3Cl^{-}$ 519 (a)

In presence of  $H_2O$  which is a weak ligand no pairing occurs which results in unpaired electrons left in the compound, due to which it shows paramagnetism

4s

 $\frac{3d}{\mathrm{Mn}^{2+}} \uparrow \uparrow \uparrow \uparrow \uparrow \uparrow$ 

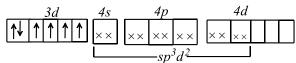
520 **(a)** 

This is a fact.

521 **(b)** 

It is outer complex having  $sp^3d^2[CoF_6]^{3-}$  ion.

Electronic configuration of  $Co^{3+}$  in  $[CoF_6]^{3-}$  is:



 $\times$  × Electron pair donated by F<sup>-</sup>

## 523 **(d)**

The two given compound have same composition but in solution both will give different ions. The isomerism is known as ionisation isomerism

#### 524 **(a)**

Coordination number is the maximum covalency shown by a metal or metal ion. It is the maximum number of ligands attached to metal by sigma bonds or coordinate bonds.

## 525 (a)

 $C_6H_5COOH+N_3H\longrightarrow C_6H_5NH_2+CO_2+N_2$  ; this is Schmidt's reaction.

## 526 **(d)**

Each possess the tendency to have coordination number equal to six.

## 527 (d)

Hexadentate ligand donates six pair of electrons to central atom.

(a) 2, 2-dipyridyl-bidentate ligand

(b) DMG-bidentate ligand  $CH_3 - C = N - O \rightarrow$ 

$$CH_3 - C = N \rightarrow$$

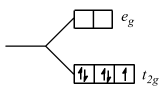
(c) Ethylenediamine-pentadentate ligand

∴ None of the given ligand is hexadentate ligand.

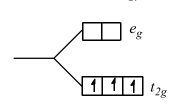
$$\frac{\text{HNO}_3}{\text{Base}} + \frac{\text{H}_2\text{SO}_4}{\text{Acid}} \rightarrow \text{NO}_2^+ + \text{H}_3\text{O}^+ + \text{HSO}_4^-$$

20.  $d^5$  in strong field

*n* =unpaired electron=1



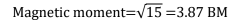
Magnetic moment= $\sqrt{n(n+2)BM}$ 



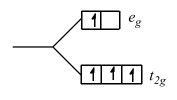
21.

*n* = 3

 $d^3$  in strong/weak field



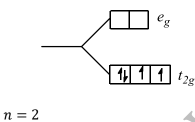
22.  $d^4$  in weak field



n = 4

Magnetic moment= $\sqrt{24}$  =4.90 BM

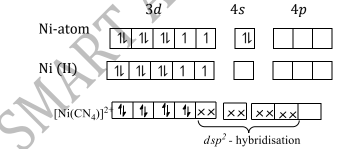
23.  $d^4$  in strong field



Magnetic moment= $\sqrt{8}$  =2.83 BM

## 531 **(a)**

 $[Ni(CN)_4]^{2-}$ ion has  $dsp^2$  hybridisation, zero magnetic moment and square planar structure.



It has no unpaired electrons hence, its magnetic moment is zero.

## 534 **(a)**

 $_{21}$ Sc=[Ar] $3d^{1}4s^{2}$ 

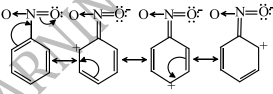
 $Sc^{3+} = [Ar]3d^04s^0$ no unpaired electrons in *d*-subshell, so it is diamagnetic and colourless.

```
536 (d)
```

In  $[Co(NH_3)_6]^{3+}$  the oxidation state of Co is +3.

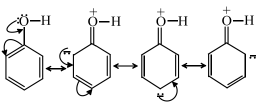
 $[Co(NH_3)_6]^{3+}$  does not contain unpaired electron hence, its magnetic moment is zero.

Presence of o-, p-directing groups in benzene nucleus activates ring for  $S_E$  reaction. Presence of m-directing deactivates ring for  $S_E$  reaction. Also halogens deactivating gp. Due to -IE inspite of oand p-directing nature. The presence of mdirecting groups in benzene nucleous simply decreases electron density at o- and p-, whereas no change in electron density at m-position is noticed.



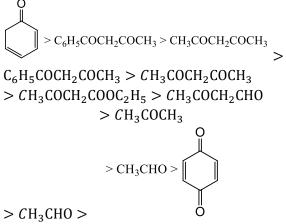
On the contrary *o*-and *p*-directing groups in nucleus increases the electron density at *o*- and *p*-position.

Thus, presence of o- and p-directing groups provide seats for  $S_E$  reactions or activates the ring, whereas presence of m-directing groups does not activate the ring and thus, deactivate the ring for  $S_E$  reactions



539 **(b)** 

The percentage of enolic contents of some common compounds in decreasing order will be



| 540          |   |     | Geometrical isomerism is found in compounds   |
|--------------|---|-----|---|
| <b>F</b> 4 1 | Mn forms $Mn_2(CO)_{10}$ carbonyl.  |     | having coordination no. 4 (square planar and not  |
| 541          | Picric acid has phenolic gp.  |     | tetrahedral shape) as well as coordination no. 6.<br>Coordination no. 4 (square planar) |
| 542          |   |     | $M_{A_2BC}, M_{A_2B_2}$   |
| 572          | (i) The sum of oxidation states of all atoms in a   |     |   |
|              | compound is zero.   |     | Showing geometrical isomerism $M_{ABCD}$ Coordination no. 6 (octahedral)                |
|              |   |     | $M_{A_4B_2}, M_{A_4BC}$   |
|              | (ii) Oxidation state of metal in carbonyl is zero.  |     | Showing geometrical isomerism.  |
|              |   |     | $M_{A_3B_3}, M_{(AA)_2B_2}$   |
|              | (a) $K_4 Fe(CN)_6$  | 548 |   |
|              | Let, oxidation state of Fe in $K_4$ Fe(CN) <sub>6</sub> = x   | 510 | Follow text.  |
|              |   | 549 |   |
|              | $\therefore \qquad +4+x+(-1\times 6)=0$   |     | Presence of electron repelling gp. decreases the  |
|              | $\therefore$ $x=+2$   |     | acidic strength.  |
|              |   |     | $C_6H_5COOH$ <i>p</i> -methyl benzoic <i>p</i> -chloro                                  |
|              | (b) $K_2 FeO_4$   |     | phenol phenol   |
|              | Let, oxidation state of Fe in $K_2$ FeO <sub>4</sub> = x  |     | acid  |
|              | Let, 0xitiation state of re in $K_2 re0_4 - x$  |     | <i>Ka</i> $6.76 \times 10^{-5}$ $1.26 \times 10^{-5}$ $4.16 \times$                     |
|              | $\therefore \qquad +1 \times 2 + x + (-2 \times 4) = 0$   |     | $10^{10}$ $1.05 \times 10^{-10}$  |
|              |   | 551 | (b)   |
|              | $\therefore$ $x=+6$   |     | It becomes brown (due to oxidation) on standing   |
|              | (c) Fe <sub>2</sub> 20 <sub>3</sub>   |     | in air.   |
|              |   | 552 |   |
|              | Let, oxidation state of Fe in $Fe_2O_3 = x$   |     | It is characteristics of aromatic compounds.  |
|              | $\therefore \qquad 2x + (-2 \times 3) = 0$  | 553 |   |
|              |   |     | The colour of the complex $COCl_3 \cdot 5NH_3 \cdot H_2O$ is                            |
|              | or 2 <i>x</i> =6  | 554 | pink.   |
|              | 6   | 554 | $2 + 4 \times (-1) = x, \qquad \therefore x = -2$                                       |
|              | $\therefore \qquad \qquad X = \frac{6}{2} = +3$   | 555 |   |
|              | (d) Fe(CO) <sub>5</sub>   |     | Halide ligands have low values of $\Delta$ .  |
|              |   | 556 | _   |
|              | Oxidation state of Fe in $Fe(CO)_5 = 0$   |     | Electronic configuration of $Co^{3+}$ in $[CoF_6]^{3-}$ is:                             |
|              | $\therefore$ Oxidation state of Fe is least in Fe(CO) <sub>5</sub> .  |     | 3d $4s$ $4p$ $4d$   |
|              | $\cdots$ Oxidation state of reast in re(CO) <sub>5</sub> .  |     |   |
| 543          | (a)   |     | $rac{1}{2}$   |
|              | The name of $[Pt(NH_3)_4Cl_2]^{2+}$ , $[PtCl_4]^{2-}$ is  |     | ×× Electron pair donated by $F^-$   |
|              | tetraamminedichloroplatinum (IV)  | 558 |   |
|              | tetrachloroplatinate (II). Since, positive ion is   |     | Let the oxidation state of Fe in  |
|              | written first and negative ion later.   |     | $[Fe(H_2O)_5NO]SO_4$ is x.  |
| 544          |   |     | $[Fe(H_2O)_5NO]^{2+}$   |
|              | Resorcinol is <i>meta</i> hydroxyphenol.  |     | $\Rightarrow \qquad x+0+1=2$  |
| 545          |   |     | $\therefore$ x=+1   |
|              | The compounds of nickel are green coloured due<br>to $d_{i}$ d transition in presence of ligand in Ni <sup>2+</sup> |     | Here, NO exists as nitrosyl ion $(NO^+)$ .  |
|              | to $d - d$ transition in presence of ligand in Ni <sup>2+</sup> cations.  | 559 |   |
| 546          |   |     | EAN = Z - (ON) + 2 (C.N.)   |
| 540          |   |     | where, O.N.=oxidation number  |
|              | $C_6H_5CH_3 \xrightarrow{Cl_2} C_6H_5CCl_3 \xrightarrow{HOH} C_6H_5COOH$  |     | C.N.=coordination number  |
| 547          | (a)   |     | Z = atomic number   |

$$[Fe(CN)_6]^{4-}$$
:  
EAN=26-(2)+2(6)  
=26-2+12=36

## 561 **(a)**

In the complex  $K_3$  [FeF<sub>6</sub>], Fe is present in +3 oxidation state.

 $_{26}$ Fe=[Ar] $3d^{6}4s^{2}$ 

 $\mathrm{Fe}^{3+} = [\mathrm{Ar}]3d^5$ 

Hence, number of unpaired electrons is five as F is weak ligand.

Magnetic moment=
$$\sqrt{n(n+2)}$$
  
=  $\sqrt{5(5+2)}$   
=5.91 BM

## 563 **(c)**

No doubt C— D bond cleavage is slower than C— H bond due to isotopic effect but rate of overall substitution is determined by the slow attachment of electrophile to carbocation (Morrison-Boyd 15.14P-532).

## 564 **(a)**

Due to synergic bond formation, bond order decreases and bond length increases a little.

## 566 **(c)**

It is a bicyclic compound having two common carbon atoms and three bridges. So, the IUPAC name is



5, 6-dimethyl bicyclo [2.2.1] hept-2-ene

#### 567 **(c)** Fe<sup>3+</sup>

$$(+ [Fe(CN)_6]^{4-} \rightarrow Fe_4[Fe(CN)_6]_3$$

568 **(a)** 

The effective magnetic moment of a paramagnetic substance is given by the relation

$$u = \sqrt{n(n+2)}$$
 BM.

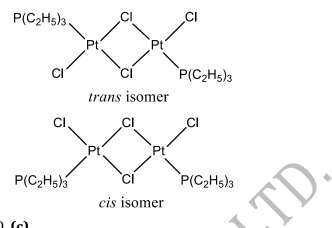
where, *n*=number of unpaired electrons.

In 
$$[FeF_6]^{3-}$$
,  $Fe^{3+}$  has five unpaired electrons

$$\mu \text{ of } [\text{FeF}_6]^{3-} = \sqrt{n(n+2)}$$
$$= \sqrt{5(5+2)}$$
$$= \sqrt{35} = 5.92 \text{ BM}.$$

569 **(c)** 

 $[PtCl_2, P(C_2H_5)_3]_2$  can exhibit geometrical isomerism, the geometrical isomers are



570 **(c)** 

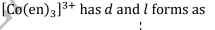
The minimum possible isomers of compound will be

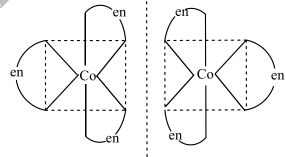
$$\begin{bmatrix} z \\ a - C - b \\ c - C - b \\ z' \end{bmatrix} \begin{bmatrix} z & z \\ b - C - a & b - C - a \\ d - C - c & c - C - d \\ z' & z' \end{bmatrix} \begin{bmatrix} z \\ a - C - b \\ d - C - c \\ z' \end{bmatrix}$$

571 **(b)** 

 $C_2O_4^{2-}$  is a bidentate group. As the complex contains three bidentate groups, the central metal ion has a coordination number of 6.

572 (d)





573 **(b)** 

Ru forms two carbonyls with zero oxidation number. Mononuclear  $Ru(CO)_5$  and trinuclear  $Ru_3(CO)_{12}$ .

574 (c)

Oxidation state of nitrogen in  $(N_2H_5)_2SO_4$ is 4x + 10 - 2 = 0

$$x = -2$$

575 **(d)** 

Linkage isomerism is exhibited by ambidentate ligands (ligands having two coordination sites). *e.g.*,  $NO_2^-$ .

If the bonding is through N, the ligand is named as nitro and if it is through O, it is named as nitrito.  $NO_2^- \rightarrow nitro - N$ 

$$0N0^- \rightarrow nitrito -0$$

Due to resonance of electron pair in aniline, 593 (c)  $C_6H_5CH = CHCOOH \xrightarrow{NaOH+CaO} C_6H_5CH = CH_2$ nitroaniline and acetanilide, these are weaker than C<sub>6</sub>H<sub>5</sub>CH<sub>2</sub> which does not involve lone pair of 594 (d) N in resonance. The basic order is: Benzyl amine  $[Co(en)_3Cl_3 ie, [Co(en)_3]^{3+}$ >Aniline > Acetanilide > Nitroaniline. 578 (b) Effective atomic no. (EAN) = at. No. of central atom -oxidation state  $+2 \times (no. of ligands) =$ en  $28 - 0 + 2 \times 4 = 36$  $EAN = 78 - 4 + 2 \times 6 = 86.$ 579 (b) Mirror  $[Cu(NH_3)_4]SO_3 \rightleftharpoons [Cu(NH_3)_4]^{2+} + SO_4^{2-}$  $cis[Co(en)_2Cl_2]Cl ie, cis [Co(en)_2Cl_2]$ 580 (b) Ammonia is not an ambident legand so it can donate electron only by N-atom 582 (b)  $C \alpha$  $12\sigma$  and  $3\pi$ . С 583 (c) -OH gp. is activating whereas Cl— is deactivating. Mirror  $-CH_3$  gp. is less activating than OH. 595 (b) 584 (d) The compound in which ligands form ring with the metal are called chelate complex. 597 (c) Benzaldehyde undergoes Cannizzaro's reaction. Br is symmetrical with The compound H 600 **(c)** respect to centre of the molecule Coordination isomerism is possible when both 585 (b) positive and negative ions of a salt are complex Two cis and trans forms. ions and the two isomers differ in distribution of 586 (c) ligands in the cation and the anion *p*-nitrophenol is more stronger acid than phenol. 601 (a) 587 (b) This is bromination of acetanilide, a  $S_E$  reaction.  $C_6H_5CH_3 \xrightarrow{[0]} C_6H_5COOH \xrightarrow{NaOH}$ 602 (b) H<sub>5</sub>COONa CaO+NaOH The primary valency is ionizable valency. It  $C_6H_6$ corresponds to oxidation state of metal. The 588 (d) primary valency is always satisfied by anion. Staggered conformation is most stable due to its  $[Co(NH_3)_6]Cl_3 \rightarrow [Co(NH_3)_6] + 3Cl^$ minimum energy (A)589 (a) Number of primary valency is 3  $[Co(NH_3)_5Br]SO_4 \rightleftharpoons [Co(NH_3)_5Br]^{2+} + SO_4^{2-}$ Pb<sup>2+</sup> + SO<sub>4</sub><sup>2-</sup>  $\rightarrow$  PbSO<sub>4</sub>  $\downarrow$ White insoluble :.  $[Co(NH_3)_5Cl]Cl2 \rightarrow [Co(NH_3)_5Cl] +$  $2Cl^{-}$ 590 (c) (B)Fe<sup>2+</sup>, Co<sup>5+</sup>, Ti<sup>3+</sup>, and V<sup>3+</sup> have 4, 4, 1, 2 unpaired Number of primary valency is 2 :. electron respectively. The pairing leads Fe<sup>2+</sup> with  $[Co(NH_3)_4Cl_2]Cl \rightarrow [Co(NH_3)_4] + Cl^$ no unpaired electron. Number of primary valency is 1. :. 591 (b) 603 (d) Os (Z=76) : [Xe] 4f<sup>14</sup>, 5d<sup>6</sup>, 6s<sup>2</sup> The carbon atom which is attached to three Hence, the coordination number in an osmium carbon atoms is called tertiary carbon atom. complex may increase to 8. C<sub>6</sub>H<sub>14</sub> has two tertiary carbons hence, its 592 (c) structure is as Phenol has antiseptic property.

2,3-dimethyl butane

### 604 **(b)**

The ligand  $NO_2$  has two types of linkage with central atom. In  $NO_2$ , it is the N-atom which is donor and in O—NO it is the O atom which donates electron pair.

## 605 **(d)**

Tetraethyl lead is organometallic compound.

606 **(d)** 

 $C_6H_5OH + PCl_5 \rightarrow (C_6H_5)_3PO_4$  is main product. 609 (c)

Directive influence order

 $-OH > -OCH_3 > -CH_3 > -NHCOCH_3 > -$ CH<sub>2</sub>OH of *o*-, *p*- gps. This is due to effect of +*R* directing influence of gp.

## 610 **(d)**

 $[PtCl_4]^{2-}$  shows  $dsp^2$  hybridization because internal *d*-orbitals participate in its hybridization.

## 611 **(b)**

Dynamite, TNT, TNB, trinitroglycerine are explosive.

## 612 **(b)**

In Hg[Co(SCN)<sub>4</sub>], Co is present as Co<sup>2+</sup>. The configuration of Co<sup>2+</sup> is given as following [Ar] $3d^7 4s^0$ 

$$\mathrm{Co}^{2+} = \boxed{1 \hspace{0.1cm} 1 \hspace{0$$

unpaired electrons (n)  $\therefore$  Magnetic moment ( $\mu$ )= $\sqrt{n(n+2)}$ 

$$\sqrt{3(3+2)} = \sqrt{15}$$
 BM

614 (c)

In metal carbonyls CO has ox. no. equal to zero. 615 **(a)** 

[NiCl<sub>4</sub>]<sup>2–</sup> has tetrahedral shape. In this complex, Ni is in the +2 oxidation state and Ni<sup>2+</sup> ion always forms tetrahedral complexes

616 **(a)** 

It is a differentiating point in between complex and double salt.

617 **(d)** 

All possess lesser number of unpaired electrons. 618 **(c)** 

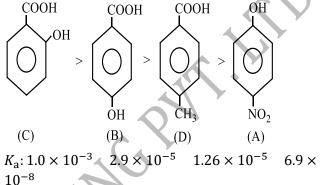
Structures  $K_4[Fe(CN)_6]$ ,  $K_3[Co(CN)_6]$ ,  $K_2[Ni(CN)_4]$  are diamagnetic.

## 619 **(c)**

Wilkinson's catalyst is used for hydrogenation of alkenes

#### 620 (c)

Due to *ortho* effect; *ortho* benzoic acid is most acidic because its anion is highly stabilized due to strong intramolecular H-bonding.



## 623 **(a)**

C<sub>6</sub>H<sub>5</sub>COOH is acid; phenol also as acid.

624 **(c)** 

*Cis*-isomer of  $[Pt(NH_3)_2Cl_2]$  is used as anticancer drug.

# 626 **(b)**

[Co(en)<sub>2</sub>Cl<sub>2</sub>]<sup>+</sup> shows geometrical as well as optical isomerism

628 **(c)** 

Lab method for preparation of benzaldehyde.

## 629 **(a)**

The ligand at least consist one donor atom having a lone pair of electrons which it can donate to metal atom or ion

## 630 **(c)**

Aniline is steam volatile.

## 631 **(a)**

CFSE (crystal field splitting energy) for octahedral complex,  $\Delta_0$  depends on the strength of negative ligand. Spectrochemically it has been found that the strength of splitting is as follows

 $\begin{array}{l} {\rm CO} > \ \underline{{\rm CN}^-} > NO_2^- > en > \ \underline{{\rm NH}_3} > py > \\ {\rm NCS}^- > {\rm H}_2 {\rm O} > {\rm O}^{2-} > O{\rm X}^{2-} > O{\rm H}^- > {\rm F}^- > \end{array}$ 

$$Cl^- > SCN^- > S^{2-} > Br^- > I^-$$

Therefore, magnitude of  $\Delta_0$  will be highest in case of  $[Co(CN)_6]^{3-}$ .

## 632 **(b)**

# K<sub>2</sub>[PtCl<sub>6</sub>]

Potassium hexachloroplatinate (IV).

633 **(c)** 

The complex formed by the reaction of  $NiSO_4$ , pyridine and  $NaNO_2$  gives  $[Ni(py)_4](NO_2)_2$  a

blue-coloured salt.

## 634 **(b)**

 $C_6H_5CHO \xrightarrow{KOH(aq.)} C_6H_5COOH + C_6H_5CH_2OH$ 635 (a)

Only m-cresols give tribromo derivatives on treatment with  $Br_2$  water.

#### 636 (a)

 $[E(en)_2C_2O_4]NO_2$   $\therefore \text{ Coordination number of } E = 6$  $\therefore \text{ Oxidation number of } E = 3[E^{3+} + 0 + (-2) + (-1) = 0]$ 

 $C_6H_5CHO \xrightarrow{PCl_5} C_6H_5CHCl_2$ 

638

|     | Complex                 | Isomerism shown |
|-----|-------------------------|-----------------|
| (a) | $[Co(en)]^{3+}$         | Optical only    |
| (b) | $[Ni(NH_3)_5Br]^+$      | No geometrical  |
|     |                         | isomer          |
| (C) | $[Co(NH_3)_2(en)_2]^3$  |                 |
| (d) | $[Cr(NH_3)_4(en)]^{3+}$ | No geometrical  |
|     |                         | isomer          |
| (c) |                         |                 |

# 639 (c)

Presence of  $-NO_2$  at *p*-position increases acidic character.

### 640 **(b)**

Alkanes are saturated hydrocarbons without any functional group, hence can show chain isomerism only

## 641 **(d)**

Both have different molecular formulae.

## 642 **(b)**

Ni in 
$$[Ni(CO)_4]$$
: ..... $3s^2, 3p^6, 3d^8, 4s^2$   
 $1 \downarrow 1 \downarrow 1 \downarrow 1$  1 1  $e^{4s}$   $e^{4p}$   
 $3d$   
Ni in  $[Ni(CN)_4]^{2-}$ : ..... $3s^2, 3p^6, 3d^8, 4s^2$ 

643 **(a)** 

The product (*K*) is formed through simple nucleophilic substitution while major product (*L*) is formed through  $\sim$ H<sup>-</sup> shift *via* S<sub>N</sub> 1 reaction and methoxy group stabilizes the carbocation intermediate of product(*L*).

## 645 **(c)**

In the Grignard reaction magnesium metal forms an organometallic bond

$$RX + Mg \xrightarrow{Dry \text{ ether}} R - Mg - X$$

## Grignard reagent

## 646 **(a)**

Aromatic hydrocarbons are called arenes with general formula  $C_nH_{2n-6y}$ , where  $n \not< 6$  and y is no. of cyclic rings. Benzene has one ring and n = 6, *i. e.*, no. of carbon atoms. Thus, general formula is  $C_6H_6$ . All other aromatic hydrocarbons are derivative of benzene.

## 647 **(c)**

It is a fact.

## 648 **(d)**

Tri and tetravalent bridges derived from methane are given the prefix methyno and methyno respectively

649 **(a)** 

The structure of the compound is

$$\begin{array}{c|ccccc}
CH_3 & CH_3 \\
5 & |4 & 3 & |2 & 1 \\
CH_3 - C - CH_2 - CH - C - H \\
| & || \\
CH_3 & O
\end{array}$$

2,4,4-trimethyl pentanal.

650 **(c)** 

$$C_6H_5COONa \xrightarrow{NaOH+CaO} C_6H_6.$$

651 **(a)** 

Complementary colours of absorbed light are seen.

652 **(d)** 

Presence of  $-SO_3H$  gp. increases solubility of drug or dyes.

653 **(d)** 

```
It is a fact.
```

654 **(a)** 

The directive influence order is:

$$0^- > NR_2 > NHR > NH_2 > OH > OCH_3$$

$$\approx$$
 NHCOCH<sub>3</sub> > CH<sub>3</sub> > X

655 **(d)** 

 $Ti^{4+}$ :  $3d^0$  and  $Cu^+$ :  $3d^{10}$  can not show *d*-*d*-transition and thus colourless.

656 **(d)** 

↑↓ | ↑↓

 $Fe^{3+}$  in  $[Fe(CN)_6]^{3-}$  is:

657 (b)  
The UPAC name of [N(PPh<sub>3</sub>)<sub>2</sub>Cl<sub>2</sub>]<sup>2+</sup> is dichlorobis (triphenylphosphine) nickel (I).  
658 (a)  
It is neutral complex as it does not ionize in solution state.  
659 (b)  
Higher is the stability constant of ligand, lesser is its dissociation, more is its stability.  
660 (c)  
R1s a fact.  
661 (d)  
Electronic configuration of Co<sup>3+</sup> in [CoF<sub>4</sub>]<sup>2-</sup> is:  

$$\frac{3d}{\sqrt{1+|1+|1+|1+||}} \frac{4\pi}{||x||} \frac{4\pi}{\sqrt{|x||}} \frac{4\pi}{\sqrt{$$

 $K_4[Fe(CN)_6]4K^+ + [Fe(CN)_6]^{4-}$ The oxidation number of Fe in  $[Fe(CN)_6]^{4-}$  is +2.  $Fe^{2+}: 1s^2, 2s^22p^6, 3s^23p^63d^6, 4s^04p$  $3d^6$ **4**s 11 11 1L × Х CN CN CN CN CN ... Since, CN<sup>-</sup>is a strong field ligand, pairing occurs and the hybridisation of  $[Fe(CN)_6]^{4-}$  is  $d^2sp^3$  and structure is octahedral. 685 (c) :.  $CH_3CHClCH_2C_6H_5 \xrightarrow{KOH(aq.)} CH_3CHOHCH_2C_6H_5$ :. 686 (d) All the compounds in which there should be 693 (a) restricted rotation about a bond in the molecule, show geometrical isomerism. Oximes of the type  $CH_3 - CH = N - OH, C_6H_5 - CH = N - OH,$  $C_6H_5 - C = N - OH$  and cyclic 694 (a) CH<sub>3</sub> 695 (d) HOOCCH--CH2-CHCOOH Compound like show geometrical isomerism 696 (a) 687 **(b)** The oxidation of aniline by  $K_2Cr_2O_7 + H_2SO_2$ (conc.) gives *p*-benzoquinone. 697 **(a)** 688 (b) In Rotate through 180° но HO Eclipsed Me Me 698 (b) HO Rotate through 60<sup>c</sup> но Ме Eclipsed meso Here, P and R represent meso-compound 689 **(b)** Friedel-Crafts reaction involves new C— C bond. 691 (a) Rest all show less tendency to donate electron pair due to resonance. 692 (c) 699 **(b)** (a)  $K_3[Fe(OH)_6]$ Let oxidation state of Fe in 700 (d)  $K_3[Fe(OH)_6] = x$  $(+1\times3)+x+(-1\times6)=0$ x = +3(b)  $K_2[FeO_4]$ 

Let oxidation state of Fe in  $K_2[FeO_4] = x$  $(+1\times2)+x+(4\times-2)=0$ x = +6FeSO<sub>4</sub>. (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>.6H<sub>2</sub>O Let oxidation state of Fe in  $FeSO_4$ .  $(NH_4)_2SO_4$ .  $6H_2O=x$ x+(-2)+2+(-2)=0x = +2(d)  $[Fe(CN)_6]^{3-}$ Let oxidation state of Fe in  $[Fe(CN)_6]^{3-}=x$  $x + (6 \times -1) =$  $FeSO_4$ . (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> has Fe in lowest oxidation state. CO is a strong ligand, all the six electrons of the

valence shell of Cr is paired and spin only magnetic moment=0

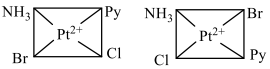
Phenol is used in carbolic soaps.

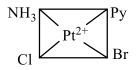
Werner proposed theory for complex compounds to explain the structure and isomerism in them.

It is p-block element and thus, has no tendency to form complex.

 $[CoF_6]^{3-}$  complex ion  $Co^{3+}$  is  $sp^3 d^2$  hybridized. F<sup>-</sup> is weak ligand and cannot pair up the *d*-electrons so, complex is high spin. Due to four unpaired electrons it is highly paramagnetic.

[*Mabcd*] type complexes exist in three isomeric forms.





Due to H-bonding.

A characteristics reaction of primary amine. This is carbylamines reaction.

701 (d)

The formula of given complex are as follows: (a) Hexammineplatinum (IV) chloride  $[Pt(NH_3)_6]Cl_4$ (b) Chloropentammine platinum (IV) chloride  $[Pt(NH_3)_5Cl]Cl_3$ (c) Dichhlorotetrammine platinum (IV) chloride  $[Pt(NH_3)_4Cl_2]Cl_2$ (d) Trichlorotriammine platinum (IV) chloride  $[Pt(NH_3)_3Cl_3]Cl$ In aqueous solution the complex ionise is  $[Pt(NH_3)_3 Cl_3]Cl \rightleftharpoons [Pt(NH_3Cl_3]^+ + Cl_3]$ Trichlorotriammine platinum (IV) 2 ions chloride gives the minimum number of ions in the solution. Hence, it has the minimum electrical conductivity. 702 (c) Diasteromers have different physical properties such as m. pt, b. pt solubilities 703 **(b)** The decreasing order of priority of prefix in numbering the carbon chain of an organic compound is Bromo > Chloro > Iodo  $\begin{array}{c} 1 \\ CH_3 \\ \hline CH_3 \\ \hline CH_3 \\ \hline CH_3 \\ \hline CH_2 \\ \hline CH_3 \\ \hline C$ 3-bromo-2-chloro-4-iodo hexane 705 (d) The structure of the compound 2, 2'-bipyridine is 706 (d) COO OHC 4-formyl-2-oxo-cyclohexane-1-carboxylic acid Note : If a compound contains two or more substituents then numbering is done in such a way that the sum of the locants is the lowest 707 (c)  $-N_2$ Cl gp. Is reduced to -H by reducing agent  $C_2H_5OH/Cu$ . 708 (d) All are the required facts for diethyl triamine. 709 (b)  $[\operatorname{Pt}(\operatorname{NH}_3)_4\operatorname{Cl}_2]\operatorname{Cl}_2 \rightleftharpoons [\operatorname{Pt}(\operatorname{NH}_3)_4\operatorname{Cl}_2]^{2+} + 2\operatorname{Cl}^-.$ 711 (b) When ligands are exchanged between metal atoms, coordination isomerism results. Hence,  $[Co(NH_3)_6][Cr(CN)_6]$  and  $[Cr(NH_3)_6][Co(CN)_6]$  re

presents coordination isomerism. 712 (c)  $Co^{3+}$  and  $Pt^{4+} = 6$  coordination number CoCl<sub>3</sub>.6NH<sub>3</sub> and PtCl<sub>4</sub>.5NH<sub>3</sub>  $[Co(NH_3)_6]Cl_3 \xrightarrow{In \text{ solution}} [Co(NH_3)_6]^{3+} + 3Cl^{-}$  $[PtCl(NH_3)_5]Cl_3 \xrightarrow{In \text{ solution}} [PtCl(NH_3)_5]^{3+} + 3Cl^{-}$ Number of ionic species are same in the solution of both complexes, therefore their equimolar solutions will show same conductance. 713 (a) Thiocyanato-N is the name when ligand SCN has electron pair donated by N-atom to metal. 715 (d) Any side chain is oxidised to -COOH. 716 (d) This is another reaction. 717 (c) CH=CH CH=CF (Furan) is heterocyclic compound 719 (c)  $[Cu(NH_3)_4](NO_3)_2$ tetrammine copper (II) nitrate. 720 (c) Nitorethane exhibits tautomerism  $CH_3CH_2-N$ nitro form aci form 722 (c) The electronic configuration Pt=[Xe]  $4 f^{14}, 5d^9, 6s^1$ :  $Pt^{2+} = [Xe]4f^{14}, 5d^8, 6s^0$  $[Pt(CN)_4]^{2-} = [Xe]4f^{14}$ 5d6s 6p×  $dsp^2$  - hybridisation : No unpaired electron is present in  $[Pt(CN)_4]^{2-1}$ ion. 723 (c) Let the oxidation number of cobalt is *x* in K  $[Co(CO)_4].$ 1 + x + 0 = 0x = -1724 (b) The IUPAC name of  $Na_3[Co(NO_2)_6]$  is sodium

hexanitrocobaltate (III).

725 (a)

- 1. $[Cu(NH_3)_4][PtCl_4]$ 2. $[Cu(NH_3)_3Cl] \cdot [PtCl_3(NH_3)]$ 3. $[Cu(NH_3)_2] \cdot [PtCl_2(NH_3)_2] cis$ 4. $[Cu(NH_3)_2Cl_2][PtCl_2(NH_3)_2] trans$ 5. $[Cu(NH_3)Cl_3] \cdot [Pt(Cl)(NH_3)_3]$ 6. $[Pt(NH_3)_4] \cdot [CuCl_4]$
- 727 **(c)**

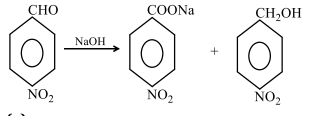
Tautomerism and functional isomerism is not possible together

732 (c)

$$Fe^{3+} + K_4Fe(CN)_6 \rightarrow KFe[Fe(CN)_6] + 3K^+$$
  
Prussian blue

733 **(b)** 

This is Cannizzaro's reaction.



 $\begin{array}{ll} \operatorname{CH}_3-\operatorname{CH}_2-\operatorname{CH}=\operatorname{CH}_3 \ \text{and} \\ \operatorname{CH}_2-\operatorname{CH}_2 \\ | & | \\ \operatorname{CH}_2-\operatorname{CH}_2 \\ \text{exhibit ring chain isomerism} \end{array}$ 

735 **(a)** 

Follow Werner's theory.

736 **(b)** 

Faraday for the first time isolated  $C_6H_6$  from coaltar.

737 **(b)** 

 $CH_3CH_2-C \equiv C - CH_3$ 2-pentyne

CH<sub>3</sub>CH<sub>2</sub>-C=C-CH<sub>3</sub> CH<sub>3</sub>-CH<sub>2</sub>-C=C-CH<sub>3</sub>  
Br H H Br  
3-bromo pent-2-ene 2-bromo pent-2-ene  

$$E, Z$$
  $E, Z$ 

Structural isomers (position) = 2 Stereo isomers = 4

738 (a)

According to Werner's theory, the primary valency of a metal is equal to the no. of charge on complex ion, *i. e.*, 3 on  $[Fe(CN)_6]^{3-}$ 

## 739 **(b)**

The complex which contains 18 valence electrons, follows 18-electron rule.

(a) In  $[V(CO)_5]$ 

The number of valence electrons

 $= 5 + (2 \times 5)$ 

 $= 15 e^{-1}$ (b)  $\ln[Fe(NH_3)_6]^{2+}$ , The number of valence electrons  $=6+(6\times2)=6+12=18 e^{-1}$ (c) In  $[Ni(CO)_6]$ , The number of valence electrons =  $10 + (2 \times 6) = 22$ e<sup>-</sup> (d) In  $[Mn(H_20)_6]^{2+}$ , The number of valence electrons= $5+(6\times 2)=17$ e<sup>-</sup> Thus, only  $[Fe(NH_3)_6]^{2+}$  follows 18-electron rule. 740 (d) One mole of X gave depression corresponding to 2 moles of particles, *i.e.*, on ionisation X gives 2 moles of ions, thus it contains only 1 ion outside the coordination sphere and its structural formula is  $[Cr(H_2O)_4Br_2]$ Cl. H<sub>2</sub>O while Y gives 3 moles of ions, thus it contains two ions outside the coordination sphere and its structural formula is  $[Cr(H_20)_5 Cl]Br_22$ 742 (d) Both represent only one molecule and no isomerism. 743 (a) Haemoglobin is porphyrin complex of ferrous iron being coordinated to four nitrogen atoms and additionally coordinated to a water reversible by a molecule. The water molecule appears to be replaceable reversible by a molecule of oxygen to give oxyhaemoglobin. Fe<sup>2+</sup> is diamagnetic due to strong field ligands. 746 (d) C1OH KOH. ; presence of  $-NO_2$ ,  $\dot{N}O_2$  $NO_2$ —CN and —COOH gp. at p-position facilitate replacement of Cl gp. by S<sub>N</sub> reactions to show normal S<sub>N</sub> reactions. 747 (b)  $(NH_4)_2S_x$  brings in selective reduction of one of

the two  $-NO_2$  group at *m*-position.

748 **(c)** 

If the highest priority groups on two carbon atoms of the double bond are on the opposite side, the configuration is *E*. (Entgegen)

#### 749 (c)

Cyclopentane possess  $0^{\circ}44'$  angle strain which is minimum

#### 750 (c)

 $[Fe(H_2O)_6]^{2+}$  has four unpaired electrons

751 **(b)**  $[Pt(NH_3)_6]Cl_4 \rightleftharpoons Pt(NH_3)_6 + 4Cl^ Ag^+ + Cl^- \rightarrow AgCl \downarrow$ White ppt.

#### 752 (a)

 $CH_{3}(CH_{2})_{3} \cdot NH_{2} \xrightarrow{KOH \text{ alc.}} CH_{3}(CH_{2})_{3}NC + 3KCl + 3H_{2}O$   $CH_{3} \longrightarrow C \equiv CH + Amm \cdot AgNO_{3} \longrightarrow CH_{3}C \equiv C \cdot Ag + HNO_{3}$ 

$$CH_3 \cdot CH_2COOCH_3 + NaOH \rightarrow CH_3CH_2COONa$$
  
+  $CH_3OH$   
OH

$$CH_3 - CH + anhy. ZnCl_2 + HCl$$
  
 $CH_3$   
 $CH_3$ 

 $\rightarrow$  Cloudiness appears within 5 minute.

## 754 **(b)**

Configuration of Mn<sup>2+</sup> is

CH2

 $[Ar]3d^{5}$ 

According to CFSE (crystal field stabilisation energy), in excited state of  $Mn^{2+}$  ion, three electrons go in  $t_{2g}$  level  $(d_{xy}, d_{yz} \text{ and } d_{zx})$  and two electrons go in  $e_g$  level  $(d_{z^2} \text{ and } D_{x^2-y^2})$ .

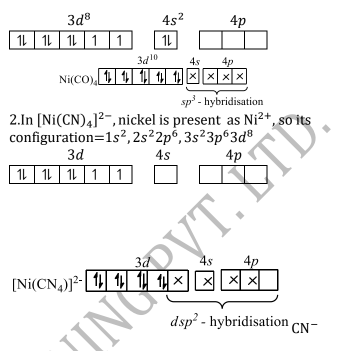
## 755 **(d)**

$$| CH_2 = C - CH_2 - COOC_2H_5$$

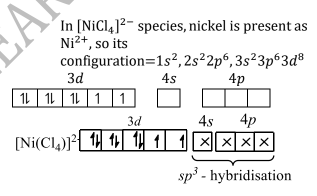
$$4 \quad 3 \quad 2 \quad 1$$
ethyl (3-methyl) but-2-enoate
$$756 \quad (c)$$
eg, Fe(CO)<sub>5</sub>, Ni(CO)<sub>4</sub>, etc.,
$$757 \quad (b)$$
Follow IUPAC rules.
$$758 \quad (b)$$

$$[Co(NH_3)_6][Cr(C_2O_4)]_3 \text{ its IUPAC name is hexa}$$
amine cobalt (III) tris (oxalato) chromate (III).
$$759 \quad (b)$$
Gammexane is C<sub>6</sub>H<sub>6</sub>Cl<sub>6</sub>.
$$760 \quad (b)$$
1.In Ni(CO)<sub>4</sub>, nickel is *sp*<sup>3</sup>-hybridised because in

it oxidation state of NI is zero. So, configuration of  ${}_{28}\text{Ni}=1s^22s^22p^6, 3s^23p^63d^8, 4s^2$ 



is strong field ligand, hence it makes Ni<sup>2+</sup> electrons to be paired up.



## Cl-

is weak field ligand, hence  $Ni^{2+}$  electrons are not paired.

761 **(d)** 

Ti<sup>4+</sup> :  $3d^0$ Cr<sup>3+</sup> :  $3d^3$  Completely filled or empty *d*orbitals are colourless. Zn<sup>2+</sup> :  $3d^{10}$ Sc<sup>3+</sup> :  $3d^0$ 

762 **(b)** 

Possible isomers are as follows :

$$\begin{array}{c} CH_{3} \\ | \\ CH_{3}CH_{2}CH_{2}CH_{2}OH \ CH_{3}CHCH_{2}OH, \\ (I) \qquad (II) \end{array}$$

CH<sub>3</sub>

$$CH_3 - C - OH$$
  
|  
 $CH_3$   
(III)

$$\begin{array}{c} CH_3 - CH - CH_2 - CH_3 \\ | \\ OH \\ (IV) \end{array}$$

Here, only (I), (II) and (III) are chain isomers 763 **(a)** 

2,2-dinitrodiphenyl or 4,4-dinitrodiphenyl is formed.

## 764 **(d)**

We have that by breaking two bond on the chiral centre configuration changes

#### 765 (c)

Use of oleum ( $H_2SO_4 + SO_3$ ) produces inclusion of  $-SO_3H$  gp. in  $C_6H_6$  ring.

## 766 **(a)**

 $NO_2^-$  can participate in linkage isomerism because it may be bonded to metal through nitrogen or through oxygen.

[(NH<sub>3</sub>)<sub>5</sub>CoNO<sub>2</sub>]Cl<sub>2</sub> and [(NH<sub>3</sub>)<sub>5</sub>CoONO]Cl<sub>2</sub> Pentaamminenitro pentaamminenitro cobalt (III) chloride cobalt (III) chloride

#### 768 **(c)**

Cuprammonium salt,  $[Cu(NH_3)_4]SO_4$  $[Cu(NH_3)_4]SO_4 \rightleftharpoons [Cu(NH_3)_4]^{2+} + SO_4^{2-}$ So, it will give two ions in water

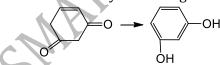
#### 769 **(b)**

(a) Shows tautomerism since aldehydes are more stable than vinyl alcohols

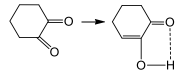
$$C_6H_5 - CH = CHOH \leftrightarrow C_6H_5CH_2 - CH_2 - CH_2$$
  
= 0

(b) Does not show tautomerism because it does not have hydrogens at  $\alpha$ -positions

( c ) Shows tautomerism because enol form is stabilized by H-bonding



(d) Shows tautomerism because enol form is stabilized by aromatic character



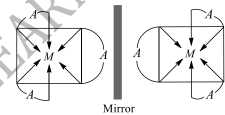
771 (a)

Main fractions of coal-tar and the compounds present there in are:

|    | Main fraction | Temp. | Chief             |
|----|---------------|-------|-------------------|
|    |               | range | constituents      |
| 1. | Light oil or  | 80-   | Benzene,          |
|    | crude         | 170°C | toluene, xylenes, |
| 2. | naphtha       |       | etc.              |
|    | Middle oil or | 170-  | Phenol,           |
| 3. | carbolic acid | 230°C | naphthalene,      |
|    | Heavy oil or  |       | pyridine, etc.    |
| 4. | creosote oil  | 230-  | Cresols,          |
|    | Green oil or  | 270°C | naphthalene       |
| 5. | anthracene    |       | quinolone, etc.   |
|    | oil           | 270-  | Anthracene,       |
|    | Pitch         | 360°C | phenanthrene,     |
|    |               |       | etc.              |
|    |               | Resid | 90-94% of         |
|    |               | ue    | carbon            |

#### 772 **(b)**

Octahedral complexes containing three bidentate ligands shows optical isomerism If A is a bidentate ligand then complex of type  $MA_3$  show optical isomerism



#### 773 **(a)**

Cl atom attached in side chain behaves as aliphatic in nature.

#### 775 **(d)**

Due to electron deficient molecule it accepts lone pair of electron to produce electrophile.

$$AlCl_3 + Cl_2 \rightarrow AlCl_4^- + Cl^+$$

#### 777 **(b)**

o –nitrophenol has intramolecular H-bonding.

778 **(c)** 

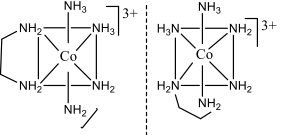
IUPAC name is tetraammine nickel (II) – tetrachloronickelate (II).

#### 779 **(b)**

Ligands are electron pair donor.

#### 781 **(d)**

*Cis*- form of  $[Co(en)_2(NH_3)_2]^{3+}$  is optically active.



784 (d) After two interchanges at each of the two chiral carbon atoms in second structure in such a way that  $CH_3$  group is held vertically upward and  $C_2H_5$ group vertically downward, we get first structure HO-Make two interchanges Thus, the two structures are identical 785 (c)  $K_3[Fe(CN)_6]$ cation anion Oxidation state of Fe in anion =+3Thus, it is potassium hexacyanoferrate (III). 786 (b) In  $[Zn(NH_3)_6]^{3+}$ , Zn exists as  $Zn^{2+}$ <sub>30</sub>Zn: 3d<sup>10</sup>, 4s<sup>2</sup> Zn<sup>2+</sup>: 3d<sup>10</sup>; Thus, no unpaired electron but it is outer orbital complex. In  $[Co(NH_3)_6]^{3+}$ , Co exists as  $Co^{3+}$ <sub>27</sub>Co: 3d<sup>7</sup>, 4s<sup>2</sup>  $Co^{3+}: 3d^6$ ; It is  $d^2sp^3$  inner orbital complex with 3 electron paired in 3d. 787 (a) In  $[CoCl_4]^{2-}$  ion, central metal atom i.e., cobalt is in +2 oxidation state. Hence,  $_{27}$ Co=[Ar] $3d^7 4s^2$  $\therefore {}_{27}\text{Co}^{2+}=[\text{Ar}]3d^7$ 11 11, Hence, number of unpaired electrons is three as Cl is weak ligand. 788 (c) Coordination number is the number of ligands in the coordination sphere. Hence, the coordination number of cobalt ion in  $[Co(H_2O)_4SO_3]Cl$  is 5. Let the oxidation number of Co is *x*. x+4(0)+(-2)+(-1)=0x + 0 - 2 - 1 = 0x=3Number of unpaired electrons in *d*-orbital are 4 because  $H_2O$  is a weak ligand and therefore, pairing of *d*-electrons is not possible. 789 (b)  $C_6H_6 + CH_3COCI \xrightarrow{AlCl_3} C_6H_5COOCH_3$ 790 (c)  $-SO_3H$  is water soluble. 791 (c)  $+1 \times 4 + x - 1 \times 4 = 0$  $4 + x - 4 = 0 \Rightarrow x = 0$  for Ni 792 (c)

Follow IUPAC nomenclature.

Due to bitter almond smell. It is  $CH_3O(OH)C_6H_3CHO$ .

# 795 **(c)**

 $C_6H_5COCH_3$  acetophenone is a mixed ketone having one alkyl and other phenyl gp. attached on -C=O gp.

# 796 **(d)**

These are the facts about transition metal atoms to act as central atom.

## 798 **(d)**

Since, hybridization is  $dsp^2$  so, it is square planar 799 **(b)** 

Metal-carbon bond in metal carbonyls has  $\sigma$  as well as  $\pi$  characters.

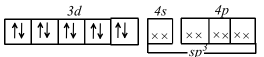
800 **(d)** 

The electronic configuration of Ni in  $[Ni(CN)_4]^{2-}$ ,  $[Ni(Cl)_4]^{2-}$  and  $Ni(CO)_4$  are:  $Ni^{2+}$  in  $[Ni(CN)_4]^{2-}$ :

$$\begin{array}{c|c} 3d \\ \uparrow \downarrow \uparrow \downarrow \uparrow \downarrow \uparrow \downarrow \downarrow \times \times \end{array} \xrightarrow{4s} 4s \\ \times \times \times \times \end{array} \xrightarrow{4p} \\ \times \times \times \times \end{array}$$

 $Ni^{2+}$  in  $[Ni(Cl_4)]^{2-}$ :

Ni in  $[Ni(CO)_4]$ :

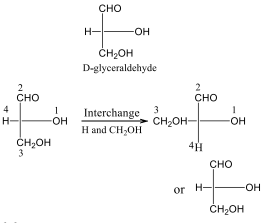


801 (c)

Both produce different ions in solution state:  $[Co(NH_3)_5Br]SO_4 \rightleftharpoons [Co(NH_3)_5Br]^{2+} + SO_4^{2-}$  $[Co(NH_3)_5SO_4]Br \rightleftharpoons [Co(NH_3)_5SO_4]^{1+} + Br^{-}$ 

803 **(c)** 

The configuration in which – OH group is on right side, H-atom is on left side, –CHO group is on upper side and  $CH_2OH$  is on lower side found in Fischer projection known as D-configuration



#### 804 (a)

Linkage isomerism is shown by those complexes which have an ambidentate ligand such as  $NO_2^-$ ,  $CN^-$  and  $SC\overline{N}$  etc. In [Fe  $(NO_2)_3Cl_3$ ], N is dono donor atom.

#### 805 (c)



2,6-dimethyl hepta-2, 5-dienoic acid

#### 806 (d)

NaHCO<sub>3</sub> reacts with acids to give  $CO_2$  from HCO<sub>3</sub> ion.

 $H^+ + HCO_3^- \rightarrow H_2O + CO_2 \uparrow$ 

#### 807 (a)

The species within the coordination sphere does not ionize.

#### 808 (c)

The reverse of enolic contents of compound is ketonic contents. Thus, the correct order of ketonic contents are  $CH_3CHO > CH_3COC_2H_5 > CH_3COCH_3$  $> CH_3COCH_2COCH_3$ 

#### 809 **(b)**

The anhydrous complexes of  $Cu_2^{2+}$  do not involve d-d transition and are thus, colourless.

#### 810 (d)

The valence of C-atom of ring is 5 at two methyl gp. attachment.

#### 812 **(b)**

Aniline is basic and thus, reacts with acid.

#### 814 **(b)**

The Riemer-Tiemann reaction is followed by dichloro carbene mechanism.

815 **(c)** 

 $C_6H_5COOH \xrightarrow{SOCl_2} C_6H_5COCl + SO_2 + HCl; -OH$ group is replaced by PCl<sub>5</sub>, PCl<sub>3</sub> or SOCl<sub>2</sub>. 816 **(b)** 

Mn<sup>2+</sup>will have half filled more stable *d*<sup>5</sup> configuration and without distributing it an outer orbital complex can be formed

817 **(a)** 

|                      | Hybridi<br>zation              | Unpair<br>ed    | Magneti        |
|----------------------|--------------------------------|-----------------|----------------|
|                      | zation                         | eu<br>electro   | c<br>momen     |
|                      |                                | ns              | t              |
| $1.[Co(CN)_6]^{3-}$  | d <sup>2</sup> sp <sup>3</sup> | 0               | 0              |
| $2.[Fe(CN)_6]^{3-1}$ | d <sup>2</sup> sp <sup>3</sup> | 1               | $\sqrt{3}$ BM  |
| 3.                   | $d^2sp^3$                      | 2               | $\sqrt{8}$ BM  |
| $[Mn(CN)_{6}]^{3-}$  |                                | $\land \bullet$ |                |
| $4.[Cr(CN)_6]^{3-1}$ | d <sup>2</sup> sp <sup>3</sup> | 3               | $\sqrt{15}$ BM |

Thus, least paramagnetism is in (a).

## 818 **(d)**

The primary valencies of Ni, in the complexes  $[Ni(Cl_4)]^{2-}$  and  $[Ni(CN)_4]^{2-}$  is same *i.e.*, (+II). Primary valencies are those valencies which a metal exhibits in the formation of its simple salt, these are non-directional. It is also referred as oxidation state of central metal atom.

## 819 (c)

The central metal ion is  $Fe^{3+}$  and  $C_2O_4^{2-}$  is negative bi-dentate ligand which forms more stable complex than neutral or mono-dentate ligand.

#### 821 **(b)**

In  $\pi$ -complex, organic ligands use their  $\pi$  system to bond with metal, *e*.g., ferrocene.

#### 823 **(c)**

CN<sup>-</sup> ions act both as reducing agent as well as good complexing agent

#### 824 (a)

The order of meta directing, gp. is:  $Me_3N^+ > NO_2 > CN > SO_3H > CHO > COCH_3$ > COOH

# 825 **(a)**

 $\begin{array}{cccc} CH_{3}COC_{3}H_{7} \mbox{ can exhibit metamerism} \\ 0 & 0 & CH_{3} \\ & & || & & || & | \\ CH_{3}-C-CH_{2}CH_{2}CH_{3}, \ CH_{3}-C-CH-CH_{3} \\ and & 0 \\ & & || \\ CH_{3}CH_{2}-C-CH_{2}CH_{3} \\ 826 \ \textbf{(d)} \end{array}$ 

It has coordination no. six and thus, octahedral or  $sp^{3}d^{2}$  -hybridization.

828 **(d)** 

Let the oxidation state of iron in  $K_4[Fe(CN)_6]$  is x.  $CH_3 CH_3$ 4(+1)+x+6(-1)=04 + x - 6 = 0In compound two chiral carbon x = +2atoms are present 829 (d) 844 (c) These are the methods to test complex formation. 9 mole or 9 × 22.4 litre of  $C_2H_2$  are needed. 830 (b) 845 (b) EAN of Fe in  $K_3$ Fe(CN)<sub>6</sub> is:  $26 - 3 + 2 \times 6 = 35$ , IUPAC name of sodium nitroprusside *i.e.*, not 36 the next inert gas. Na<sub>2</sub>[Fe(CN)<sub>5</sub>NO] is sodium pentacyanonitrosyl 833 (a) ferrate (III) because in it NO is neutral ligand and Grignard reagent is a  $\sigma$ -bonded organometallic the oxidation number of Fe is III, which is compound because all the bonds present in the calculated as reagent are single bonds.  $2 \times ON$  of Na+ ON of Fe +5 $\times ON$  of CN 835 (b)  $+1 \times ON \text{ of } NO=0$  $\ddot{N}(CH_2COO^-)_2$  $2 \times (+1) + 0$  N of Fe  $+5 \times (-1) + 1 \times 0 = 0$ EDTA is C<sub>2</sub>H<sub>4</sub>; ; it has six positions (*i.e.*, ON of Fe =5-2=+3N(CH<sub>2</sub>COO)<sub>2</sub> 846 (b) six electron pairs) available for attachment at The electronic configuration of Ni in central metal atom.  $[Ni(CN)_4]^{2-}, [Ni(Cl)_4]^{2-}$  and  $Ni(CO)_4$  are : 836 (d)  $Ni^{2+}$  in  $[Ni(CN)_4]^{2-}$ : Thymol is 3-hydroxy-l-isopropyl-4-methyl 3dbenzene  $(C_{10}H_{14}O)$ , a white crystalline phenol derivative, has smell of thyme, occurs in many essential oils used as fragrant material as well as Ni<sup>2+</sup> in [Ni(Cl)<sub>4</sub>]<sup>2-</sup> : mild antiseptic. 837 (d) 3d Ferrocyanide ion  $[Fe(CN)_6]^{4-}$  is diamagnetic in nature hence  $K_4[Fe(CN)_6]$  complex has zero magnetic moment.  $Ni^{2+}$  in  $[Ni(CO)_4]$  : Fe-atom 3d4s11 1 1 11 1 1 Fe(II) 11 1 1 1 1 848 (d)  $d^2 sp^3$  hybridisation When cations have same charge but number of *d*electrons are different then the stability (or CFSE) decreases with increase in the number of *d*-1L × × × [Fe(CN)<sub>6</sub>] electrons. Therefore, the correct order is  $Fe^{2+} > Co^{2+} > Ni^{2+} > Cu^{2+}$ CN CN CN CN CN CN 849 (a) 838 (a) Oxidation state of Ni in  $Ni(CO)_4$  is zero. CO ligand has zero oxidation state, that is why 3d4*s* 4p $[Ni(CO)_4]$  is a zero valent metal complex 11 11 840 (c) 11 11 1 1 A characteristic; follow ligand field theory. 842 **(b)** 3*d* Ni**| 1; | 1; | 1; |** 11 11 Central ion is  $Cd^{2+}$  and ligand is  $CN^{-}$ . 843 (b)  $sp^3$  - hybridisation CO is a strong ligand. It causes pairing of

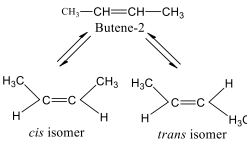
electrons. Hence, there is no unpaired electrons in Ni(CO)<sub>4</sub>.

850 **(b)** Ox. no. of Cr is calculated as:  $3 \times 1 + a + 1 \times (-2) + 4 \times (-1) = 0;$   $\therefore a = +3$ 853 **(c)** 

 $CrO_3$  or  $CrO_2Cl_2$  and a mixture of  $K_2Cr_2O_7$  +  $H_2SO_4$  + NaCl can also be used.

854 (b)

Butane-2 exhibit geometrical (*cis*, *trans*) isomerism



## 855 **(a)**

Ferrocene is a  $\pi$  complex Fe $(\eta^5 - C_5H_5)_2$ .

856 **(a)** 

Ni in  $[Ni(NH_3)_6]^{2+}$  has  $sp^3d^2$  (outer complex) having octahedral geometry.

## 857 (d)

These examples are ionisation isomers because of chloride and sulphate ions.

#### 859 **(b)**

When ligands are arranged in ascending order of crystal field splitting energy,  $\Delta$ , they produce a spectrochemical series.

In comparison to  $H_2O$ , Cl is strong  $\sigma$ -donor and good  $\pi$ -acceptor, therefore it is a strong ligand than  $H_2O$ . Hence, in the spectrochemical series Cl is above than water.

## 860 **(b)**

On ionization different species are formed to show ionization isomerism:

 $[Co(en)_2(NCS)_2]Cl \Rightarrow [Co(en)_2(NCS)_2] + Cl^ [Co(en)_2(NCS)Cl]NCS$ 

$$\rightleftharpoons$$
 [Co(en)<sub>2</sub>(NCS)Cl] + NCS<sup>-</sup>

# 861 **(a)**

Pt<sup>2+</sup> has square planar complexes with coordination number four.

# 862 (c)

 $\sqrt{15}$ 

In Hg[Co(SCN)<sub>4</sub>], Co exists as Co<sup>2+</sup>  $_{27}$ Co :  $3d^7$ ,  $4s^2$  $_{27}$ Co<sup>2+</sup> :  $3d^7$ 

SCN<sup>-</sup> a strong field ligand provides four electron to pair to show  $sp^3$ -hybridisation in  $[Co(SCN)_4]^{2-}$  and thus three unpaired electrons exists on  $CO^{2+}$ .

: Magnetic moment =  $\sqrt{n(n+2)} = \sqrt{3(3+2)} =$ 

863 **(d)** 

But-2-ene and but-1-yne do not have same molecular formula, thus are not isomers  $CH_3 - CH = CH - CH_3$   $CH_3 - CH_2 - C \equiv CH$ But-2-ene but-1-yne  $(C_4H_8)$   $(C_4H_6)$ 864 **(b)**  ${}_{27}Co \rightarrow [Ar]3d^74s^2$  $Co^{2+} \rightarrow 3d^74s^0$  $\uparrow\downarrow\uparrow\uparrow\downarrow\uparrow\uparrow\uparrow$ Number of unpaired electrons = 3 865 **(a)** NO in iron complexes has ox. no. equal to one.  $a + 5 \times 0 + 1 - 2 = 0$ ,  $\therefore a = +1$ 

868 (c)

Due to the presence of phenolic gp.

869 **(b)** 

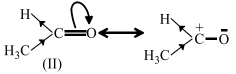
Ni(CO)<sub>4</sub> has a O.N. zero for Ni

870 (d)

Because it is a polydentate ligand which binds the central atom nickel forming a ring structure

## 871 **(c)**

Carbonyl compounds react with Grignard reagent following nucleophilic addition. More +ve is charge on C<sup>+</sup> centre of carbonyl gp., easier is nucleophile attack.



(Positive charge on C<sup>+</sup> is dispersed due to + I. E. of CH<sub>3</sub> gp.)

$$\underset{H_{3}C}{\overset{H_{3}C}{\underset{(I)}{\leftarrow}}} C \xrightarrow{H_{3}C} \overset{H_{3}C}{\underset{H_{3}C}{\leftarrow}} C \xrightarrow{+} C \xrightarrow{-} \overline{O}$$

(Positive charge on  $C^+$  is dispersed more due to +I.E. of two  $CH_3 gp.$ )

$$C_{6}H_{5} \xrightarrow{C_{6}H_{5}} C \xrightarrow{H_{5}C_{6}} \xrightarrow{H_{5}C_{6}} \xrightarrow{+} C \overline{O}$$

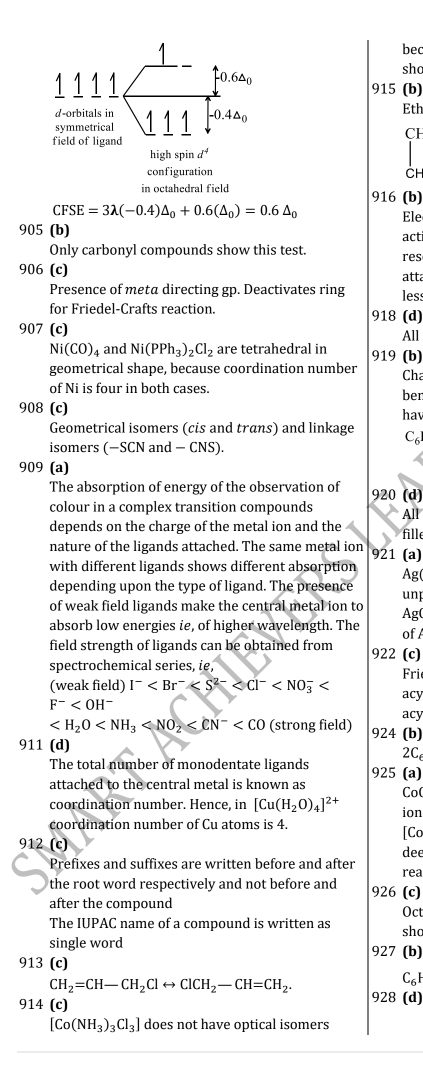
(Positive charge on C<sup>+</sup> is intensified due to -I.E. of C<sub>6</sub>H<sub>5</sub> gps.)

But in (III) conjugation of  $\sum$ CO gp. with  $\pi$  system

of benzene nucleus following resonance deactivates C<sup>+</sup> centre to attack by nucleophile. Resonance effect overpowers over -I.E. of  $C_6H_5$  gp.

 $[Cr(SCN)_2(NH_3)_4]^+$  shows geometrical (or *cis*-During debromination, meso-dibromobutane trans) and linkage isomerism. form *tran*-2-butene 873 (c)  $CuCl + 4KCN \rightarrow K_3[Cu(CN)_4] + KCl.$ Thus, coordination no. of Cu is four.  $CH_3$ Anti 876 **(b)** elimination The IUPAC name of the compound  $[CuCl_2(CH_3NH_2)_2]$  is dichloro bis-(methyl amine) trans-2-butene meso-2,3-dibromobutane copper (II). 891 (a) 877 (b) The IUPAC name of compound is  $K_2[Cr(CN)_2O_2(O)_2(NH_3)]$ is Potassium ammine dicyano dioxoperoxo chromate (VI) 2-(1-cyclobutenyl)-1-hexane 893 (a) It can show ionization isomerism:  $[Co(NH_3)_4Cl_2]^+$ 879 (b) and  $[Co(NH_3)_4Cl]^{2+}Cl$ . Electronic configuration of  $[FeF_6]^{3-}$  is: 894 (b) Replacement of H-atom of ring usually takes place following  $S_E$  reaction mechanism. 895 (b) ×× Electron pair donated by F<sup>-</sup> Follow mechanism of Reimer-Tiemann reaction. 880 (d) 896 (a) Effective atomic no. (EAN) = at. No. of central It is 2,4,6-trinitrophenol. atom -oxidation state  $+2 \times (no. of ligands) =$ 897 (a)  $28 - 0 + 2 \times 4 = 36$ Follow mechanism of cannizzaro's reaction.  $EAN = 26 - 3 + 2 \times (6) = 35$ 898 (b) 882 (d) Ag in Tollens' reagent exists as  $Ag_2O$ [Pt<sup>II</sup>(NH<sub>3</sub>)<sub>2</sub>Cl<sub>2</sub>] shows geometrical isomerism.  $2 \times a + 1 \times (-2) = 0$ NH<sub>2</sub> 7NH<sub>2</sub>  $\therefore a = +1$ 899 (b) NH.  $H_{3}$  $K_3[Fe(CN)_6]$ (orange yellow) (pale yellow) Electronic configuration of Fe =  $[Ar]3d^64s^2$ 884 (b) Electronic configuration of  $Fe^{3+} = [Ar]3d^5$ Staggered and eclipsed conformers cannot be Number of ligand (Coordination number)=6 physically separated because the energy Nature of ligand in strong field difference between them is so small that they 3d4smost readily interconvent at room temperature 885 (b) 3dA species or group of atoms can act as ligand only |↑↓|↑↓|↑|↑| when it carries an unshared pair, *i.e.*, lone pair of CN CN CN CN CN CN CN electrons. Hybridisation of Fe is  $d^2sp^3$ 886 (c) 900 (d) Follow IUPAC rules. Butan-2-one O 888 (d)  $NH_2 \cdot NH_2$  serves as monodentate as well as  $CH_3 - C - CH_2 - CH_3$  is not isomeric with bridging ligand because a 3-membered ring will diethyl ether  $CH_3CH_2 - 0 - CH_2 - CH_3$ . Because be too strained to be stable. both are differing in molecular formula 889 (b) 902 (d) TNT mixed with NH<sub>4</sub>NO<sub>3</sub> gives explosive material.

890 (d)



because it is of formula  $MA_3B_3$  which does not show optical isomerism

915 (b)

Ethylenediamine is a bidentate ligand.

$$\begin{array}{c} \text{CH}_2 & \overrightarrow{\text{NH}}_2 \\ | \\ \text{CH}_2 & \overrightarrow{\text{NH}}_2 \end{array}$$

916 (b)

Electron withdrawing nature or –*IE* increases the activation of ring more effectively, however resonance opposes inductive effect for attachment at o-and p-position and hence, makes less deactivation for *o*-and *p*-positions.

#### 918 (d)

All involves  $sp^3$ -hybridization.

#### 919 (b)

Chaisen condensation involves condensation of benzaldehyde with aliphatic aldehydes or ketones having two α-H-atoms, e.g.,

$$C_6H_5CHO + CH_3CH_2CHO \longrightarrow C_6H_5CH = C - CHO$$

## 920 (d)

All are transition elements with *d*-orbitals not filled to capacity.

Ag(II) has a  $d^9$ -configuration and must contain unpaired electron but AgOis diamagnetic because AgO does not contain Ag(II) but is a mixed oxide of Ag<sup>I</sup>Ag<sup>III</sup>O<sub>2</sub>.

#### 922 (c)

Friedel-Crafts reaction involves alkylation or acylation in benzene nucleus using alkylating or acylation reagents in presence of anhy. AlCl<sub>3</sub>.

#### 924 (b)

 $2C_6H_6 + 2HCl + O_2 \rightarrow 2C_6H_5Cl + H_2O$ 925 (a)

> CoCl<sub>2</sub> is a weak Lewis acid, reacting with chloride ions to produce salt containing the terrahedral  $[CoCl_4]^{2-}$ ion. CoCl<sub>2</sub> is blue when anhydrous, and a deep magenta colour when hydrated, for this reason it is widely used as an indicator for water

#### 926 (c)

Octahedral complexes of the type  $MA_5B$  do not show geometrical isomerism.

#### 927 (b)

 $C_6H_5COOC_2H_5 \xrightarrow{NaOH} C_6H_5COONa + C_2H_5OH$ 928 (d)

| —_C,—_NH <sub>2</sub>  | oil Anthracene,   |  |  |  |  |  |  |
|--|---|--|--|--|--|--|--|
| In this compound O is principal  | Pitch 270- phenanthrene,  |  |  |  |  |  |  |
| functional group   | 360°C etc.  |  |  |  |  |  |  |
| While – CHO is substituent group, hence  | 90-94% of   |  |  |  |  |  |  |
|  | Resid carbon  |  |  |  |  |  |  |
| CONH <sub>2</sub>  | ue  |  |  |  |  |  |  |
| 2 4 CHO  |   |  |  |  |  |  |  |
|  |   |  |  |  |  |  |  |
| 5-formyl-2-methyl pent-3-en-1-amide  | 939 <b>(c)</b>  |  |  |  |  |  |  |
| 929 <b>(b)</b>   | The coordination number (C.N.) of a metal atom  |  |  |  |  |  |  |
| $C_6H_5CH_2NH_2$ has least negative inductive effect   | in a complex is the total number of bonds formed  |  |  |  |  |  |  |
| and thus shows more basic nature.  | by metal with ligands.  |  |  |  |  |  |  |
| 930 <b>(d)</b>   | In case of tetrahedral complexes the number of  |  |  |  |  |  |  |
| CH <sub>3</sub>  | bonds formed between metal and ligand is four.  |  |  |  |  |  |  |
| CH <sub>3</sub> —CH <sub>2</sub> −CH <sub>2</sub> −CH−CH−CH <sub>2</sub> −CH <sub>3</sub>  | So, coordination number is also four.   |  |  |  |  |  |  |
| $\begin{array}{c} CH_3 - CH_2 - CH_2 - CH_2 - CH_2 - CH_3 \\ 7  6  5  \begin{vmatrix} 4 & 3 & 2 & 1 \\ 6 & -CH_2 - CH_3 \\ \hline \end{array}$ | 940 (b)   |  |  |  |  |  |  |
| 4-ethyl-3-methyl heptane   | Due to resonance bond length become identical and is 1.40 Å. Whereas in alkane C—C bond is                                  |  |  |  |  |  |  |
| <b>Note :</b> The prefix in a compound should be   | 1.54 Å and in alkene it is 1.34 Å.  |  |  |  |  |  |  |
| arranged in alphabetical order   | 941 (d)   |  |  |  |  |  |  |
| 931 (d)  | $C=0 + H_2 NNHC_6 H_5 \rightarrow C=NNHC_6 H_5$   |  |  |  |  |  |  |
| All possess lesser number of unpaired electrons.   | $\begin{array}{c} C = 0 + n_2 \text{NNRC}_{6}n_5 \implies C = \text{NNRC}_{6}n_5 \\ \text{All reagents do so.} \end{array}$ |  |  |  |  |  |  |
| 932 (d)  | 944 (c)   |  |  |  |  |  |  |
| A monodentate ligand has one donor site  | Electrophilic substitution occurs at electron rich  |  |  |  |  |  |  |
| available for coordination.  | centres usually at <i>o</i> - and <i>p</i> -positions. The ring   |  |  |  |  |  |  |
| 933 (a)  | attached with –NH will develop more electron  |  |  |  |  |  |  |
| It is benzene.   | density at <i>o</i> - and <i>p</i> -positions. Since <i>o</i> -position is  |  |  |  |  |  |  |
| 934 <b>(c)</b>   | blocked, thus electrophile will attach at <i>p</i> -  |  |  |  |  |  |  |
| $C_6H_5CHO \xrightarrow{Cl_2} C_6H_5COCl$  | position.   |  |  |  |  |  |  |
| 935 (c)  | 945 <b>(b)</b>  |  |  |  |  |  |  |
| Phenol is weak acid.   | Follow text.  |  |  |  |  |  |  |
| 936 <b>(b)</b>   | 946 <b>(a)</b>  |  |  |  |  |  |  |
| The coordination number in $[FeF_6]^{3-}$ is 6, hence it   | In the formation of $d^2sp^3$ hybrid orbitals, two  |  |  |  |  |  |  |
| is a octahedral complex  | $(n-1)d$ orbitals of $e_g$ set <i>i.e.</i> , $(n-1) d_{z^2}$ and $(n-1) d_{z^2}$  |  |  |  |  |  |  |
| 937 <b>(d)</b>   | 1) $d_{x^2-y^2}$ orbitals, one  |  |  |  |  |  |  |
| $AgCl + 2NH_3 \rightarrow [Ag(NH_3)_2]Cl$  | $ns$ and three $np(np_x, np_y$ and $np_z)$ orbitals   |  |  |  |  |  |  |
| 938 (a) Soluble  | combine together and form six $d^2sp^3$ hybrid  |  |  |  |  |  |  |
| Main fractions of coal-tar and the compounds   | orbitals.   |  |  |  |  |  |  |
| present there in are:  | 947 <b>(a)</b>  |  |  |  |  |  |  |
| Main fraction Temp. Chief  | EAN of Fe = $26 - 2 + 6 \times 2 = 36$ .  |  |  |  |  |  |  |
| range constituents   | 948 (a)   |  |  |  |  |  |  |
| 1. Light oil or 80- Benzene,   | $K[Pt(NH_3)_5Cl_5] \rightleftharpoons K^+ + [Pt(NH_3)_5Cl_5]^-$   |  |  |  |  |  |  |
| crude 170°C toluene, xylenes,  | 949 <b>(b)</b>  |  |  |  |  |  |  |
| 2. naphtha etc.  | $3\text{KCl} + \text{CuCl} \rightarrow \text{K}_3[\text{CuCl}_4]$ ; this is soluble in water                                |  |  |  |  |  |  |
| Middle oil or 170- Phenol,   | 950 (d)   |  |  |  |  |  |  |
| 3. carbolic acid 230°C naphthalene,  | All are characteristics tastes for phenol.  |  |  |  |  |  |  |
| Heavy oil or pyridine, etc.  | 952 (c)   |  |  |  |  |  |  |
| 4. creosote oil 230- Cresols,  | N cannot have more than 8 elements in its valence shell.  |  |  |  |  |  |  |
| Green oil or 270°C naphthalene   | 953 <b>(b)</b>  |  |  |  |  |  |  |
| 5. anthracene quinolone, etc.  |   |  |  |  |  |  |  |

Tautomeric structures of a molecule are not the resonating structures of the molecule

#### 954 (a)

2, 4, 6-trinitrophenol is strong acid than acetic acid but phenol is less acidic than acetic acid.

#### 955 (a)

Phenol being weak acid does not react with aq. NaHCO<sub>3</sub>.

#### 956 (c)

If two atoms directly attached to the double bond have the same atomic number, then the elative priority of the groups is determined by a similar comparison of the atomic numbers of the next elements in the groups. Thus, the preference order of given group is  $OCH_3 > OH > COOH > CHO$ 

#### 957 (a)

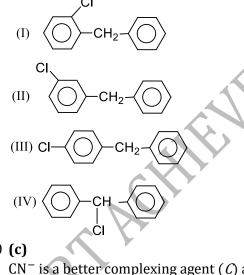
Follow IUPAC name.

958 (a)

It possesses  $d^8$  configuration of Ni<sup>2+</sup>.

959 (d)

There are four structural isomers possible for diphenyl methane when one H-atom is replaced by a Cl-atom



960 (c)

 $CN^{-}$  is a better complexing agent (*C*) as well as a reducing agent(A)

Thus, properties (*A*) and (*C*) are shown.

Property (C):Ni<sup>2+</sup> + 4CN<sup>-</sup>  $\rightarrow$  [Ni(CN)<sub>4</sub>]<sup>2-</sup> Property(*A*):

Π I  $CuCl2 + 5KCN \rightarrow K_3[Cu(CN)_4] + \frac{1}{2}(CN)_2 + 2KCl$  $(CN^{-} reduces Cu^{2+} to Cu^{+})$ 

#### 961 (c)

In the double bonds are trans and cis. The first and third bonds are identical

#### 963 (a)

*p* –nitrophenol is most acidic (among these) as it has electron withdrawing  $-NO_2$  gp.

#### 965 (c)

Geometrical isomerism is found in compounds having coordination no. 4 (square planar and not tetrahedral shape) as well as coordination no. 6. Coordination no. 4 (square planar)

#### $M_{A_2BC}, M_{A_2B_2}$

Showing geometrical isomerism Coordination no. 6 (octahedral)

$$M_{A_4B_2}$$
,  $M_{A_4BC}$ 

Showing geometrical isomerism.

 $M_{A_3B_3}, M_{(AA)_2B_2}$ 

#### 966 (d)

Follow crystal field theory.

#### 967 (d)

Due to resonance  $C_6H_6$  is stabilized and normal addition reactions (except addition of H<sub>2</sub>, Cl<sub>2</sub> and ozonolysis) are not observed in  $C_6H_6$ .

#### 969 (b)

EAN of Pt in  $[PtCl_6]^{2-} = 86$ 

 $CuCl + NH_3 \rightarrow [Cu(NH_3)_2]Cl.$  The coordination no. = No. of ligands attached.

#### 972 (a)

Longest system : 1,2,3,4,5,6 Next longest system : 6,7,8,1 Shortest system : 1,6 The IUPAC name of compound is 8-methyl bicyclo [4,3,0] nonane

973 (b)

Aniline on condensation with aromatic aldehyde gives Schiff's base.

#### 974 (c)

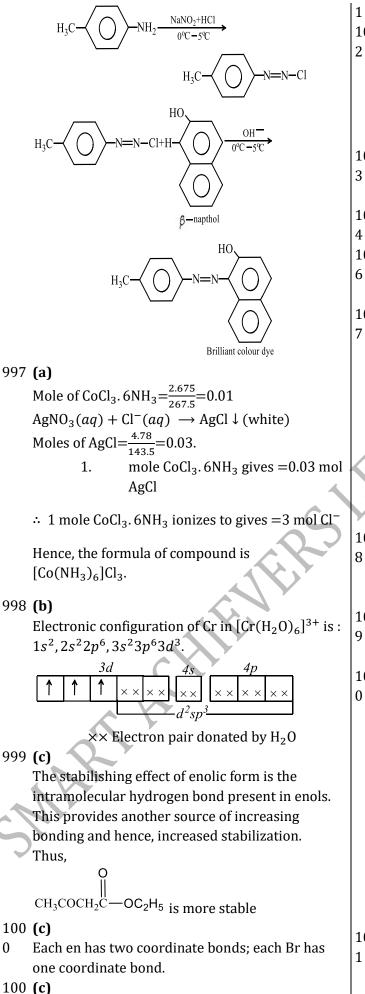
Organometallic compounds are those compounds in which metal is directly attached to the carbon atom. In sodium ethoxide, sodium attached to oxygen atom, hence it is not an organometallic compound.

975 (d)

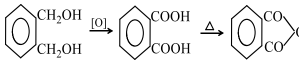
(i) CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>OH and  $CH_3CH_2CH_2 - CH - CH_3$ 

are Position isomers Due to restricted rotation about the carboncarbon single bond joining the two phenyl groups, Asymmetric carbon atom the molecule as a whole is chiral and thus shows (ii) CH3 optical isomerism 984 (b) Due the presence of asymmetry, optical Any side chain is oxidised to COOH gp. isomerism is possible 985 (d) (iii) CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>OH and These are facts about glycinato ligand. CH<sub>3</sub>CH<sub>2</sub>OCH<sub>2</sub>CH<sub>2</sub>CH<sub>3</sub> are functional isomerism 986 (c) 976 (c) According to Werner's theory, only those ions are Co<sup>3+</sup> and Pt<sup>4+</sup> have 6 coordination number. precipitated which are attached to the metal CoCl<sub>3</sub>. 6NH<sub>3</sub> and PtCl<sub>4</sub>. 5NH<sub>3</sub> atoms with ionic bonds and are present outside  $[Co(NH_3)_6Cl_3 \xrightarrow{\text{In solution}} [Co(NH_3)_6]^{3+} + 3Cl^{-}$   $[PtCl(NH_3)_5Cl_3 \xrightarrow{\text{In solution}} [PtCl(NH_3)_5]^{3+} + 3Cl^{-}$ the coordination sphere. 987 (d) sp<sup>2</sup>-hybridization leads to planar hexagonal Number of ionic species are same in the solution shape. of both complexes, therefore their equimolar 988 (a) solutions will show same conductance. Coordination isomerism is caused by interchange 977 (a) of ligands with the metal atoms. Organometallic compounds are those compounds 989 (c) in which carbon atom is directly linked to metal. Chlorophyll are green pigment in plant and But in sodium ethoxide as oxygen is attached to contain magnesium instead of caleium attached to sodium metal so, it is not a 991 (a) organometallic compound  $C_6H_5NH_2 + Cl_2OC \rightarrow C_6H_5N=C=O + 2HCl$ 978 (d) 992 (a)  $[Cu(H_2O_4)]SO_4$ . H<sub>2</sub>O coordination number of Cu is  $C_6H_5OC_2H_5 \xrightarrow{HBr} C_6H_5OH + C_2H_5Br$ 4. 993 **(d)** 979 (b)  $[Fe(NH_3)_4 . Cl_2]Cl$ Sodium nitroprusside is Na<sub>2</sub>[Fe(CN)<sub>5</sub>NO Tetraammine dichloro ferrum III chloride. 980 (d) 994 (b) 3  $NH_2 - CH - CH_2OH$  $_{28}$ Ni=[Ar]  $3s^2 3p^6 4s^2 3d^8$  $Ni^{2+} = [Ar] 3s^2 3p^6 3d^8$ COOH 2-amino-3-hydroxy propanoic acid 981 (a)  $sp^3$  $[CoF_6]^{3-}$  is an outer orbital complex ion. It Nickel has two unpaired electrons and geometry involves outer orbital hybridisation. It has  $sp^3d^2$ is tetrahedral due to  $sp^3$  hybridisation. hybridisation because  $F^-$  is a weak ligand. 995 (c) It is a test for  $-NH_2$  gp attached on benzene CoF<sub>6</sub>]<sup>3-</sup> **1**, **1**, **1 1** nucleus following diazotisation and coupling reaction. 982 (b) Effective atomic no. (EAN) = at. No. of central atom -oxidation state  $+2 \times (no. of ligands) =$  $28 - 0 + 2 \times 4 = 36$  $EAN = 28 - 2 + 2 \times 4 = 34.$ 

983 (b)



- 1 Aniline is base.
- 100 **(c)**
- 2 Two COOH on vicinal carbon atom lose  $H_2O$  on heating.



100 (c)

3 Benzene has 6 C—C, 6 C—H  $\sigma$ -bonds and 3C=C $\pi$ -bonds.

## 100 **(b)**

- 4 Rest all have plane of symmetry.
- 100 **(b)**
- 6 The number of unpaired electrons in complex  $[Pt(NH_3)_2]Cl_2$  are two.

100 **(c)** 

Geometrical isomerism is found in compounds having coordination no. 4 (square planar and not tetrahedral shape) as well as coordination no. 6. Coordination no. 4 (square planar)

#### $M_{A_2BC}, M_{A_2B_2}$

Showing geometrical isomerism  $M_{ABCD}$ 

```
Coordination no. 6 (octahedral)
```

 $M_{A_4B_2}, M_{A_4BC}$ 

Showing geometrical isomerism.

# $M_{A_3B_3}, M_{(AA)_2B_2}$

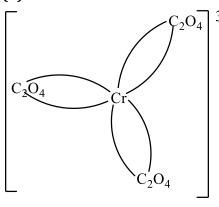
100 **(c)** 

According to the modern view primary valency of complex compound it its oxidation number while secondary valency is the coordination number

## 100 **(c)**

9 The IUPAC name of  $K_4[Ni(CN)_4]$  is potassium tetracyanonickelate (0).

101 **(b)** 



Mirror image

is not superimposable hence, optical isomerism is possible.

- 101 **(a)** 
  - S-atom is donor in SCN and N-atom is donor in NCS. The linkage isomerism arises when ligand has two possibilities to attach on central atom.,

#### 101 (d)

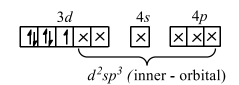
- 2 Ethane has an infinite number of conformation but staggered and eclipsed are preferred. Ethane molecule would exist in the staggered conformation due to its minimum energy and maximum stability
- 101 **(a)**
- 4 Follow IUPAC rules

#### 101 **(d)**

5 The complex in which nd orbitals are used in hybridisation, are called outer orbital complex.
26. [Fe(CN)<sub>6</sub>]<sup>4-</sup> =

27.

 $[Mn(CN)_6]^{4-} =$ 



28.

 $[Co(NH_3)_6]^{3+} =$ 

12+

29. 
$$\begin{array}{c|c} 3d & 4s & 4p \\ \hline 1 & 1 & 1 \\ d^2 s p^3 \ (\text{inner - orbital}) \end{array}$$

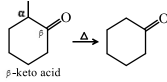
$$d^2sp^3$$
 (inner - orbital)

- 101 (d)
- 6 In each case aromatisation leads to formation of  $C_6H_6$ .

101 (a)  
7 
$$C_6H_5Cl \xrightarrow{H_2-Ni \text{ or } Al/NaOH} C_6H_6$$
;  
101 (b)

8  $\beta$ -keto acids undergo decarboxylation most easily on heating.





#### 101 **(a)**

9 Ni in  $[Ni(H_2O)_6]^{2+}$  has two unpaired electrons in it.

#### 102 **(d)**

4 An electron attracting group  $(-NO_2)$  disperses the negative charge on phenoxide ion and thus, makes it more stable or increases the acidic character of phenol. The substitution is more effective at *p*-position than in the *m*-position as the former involves a resonating structure bearing negative charge on the carbon attached to the electron withdrawing group. Also presence of electron repelling gp.  $(-CH_3)$  intensifies the negative charge on phenoxide ion and thus, makes phenol less acidic.

102 **(d)** 

6

$$2C_6H_5CHO \xrightarrow{NaOH} C_6H_5COOH + C_6H_5CH_2OH$$
  
Oxidised Reduced

102 **(c)** 9 Let

Let the ON of Pt in[Pt(NH<sub>3</sub>)<sub>5</sub>Cl]Cl<sub>3</sub> is x.  

$$x+5\times(0)-1-3=0$$
  
 $x-4=0$   
 $x=+4$ 

103 (a)

0

- Organometallic compounds have carbon-metal bond, hence  $CH_3Mgl$ , tetraethyl tin and  $KC_4H_9$  are organometallic compounds while  $C_2H_5ONa$  is not an organometallic compound due to absence of carbon-metal bond.
- 103 **(b)**

1

Optical isomers of a compound which are nonsuperimposable but related to each other as an object and its mirror are called enantiomers

$$\begin{array}{c|c} CH_3 \\ H \longrightarrow OH \\ HO \longrightarrow H \\ CH_3 \end{array} + \begin{array}{c} CH_3 \\ HO \longrightarrow H \\ HO \longrightarrow OH \\ CH_3 \end{array} + \begin{array}{c} CH_3 \\ HO \longrightarrow H \\ HO \longrightarrow OH \\ CH_3 \end{array}$$

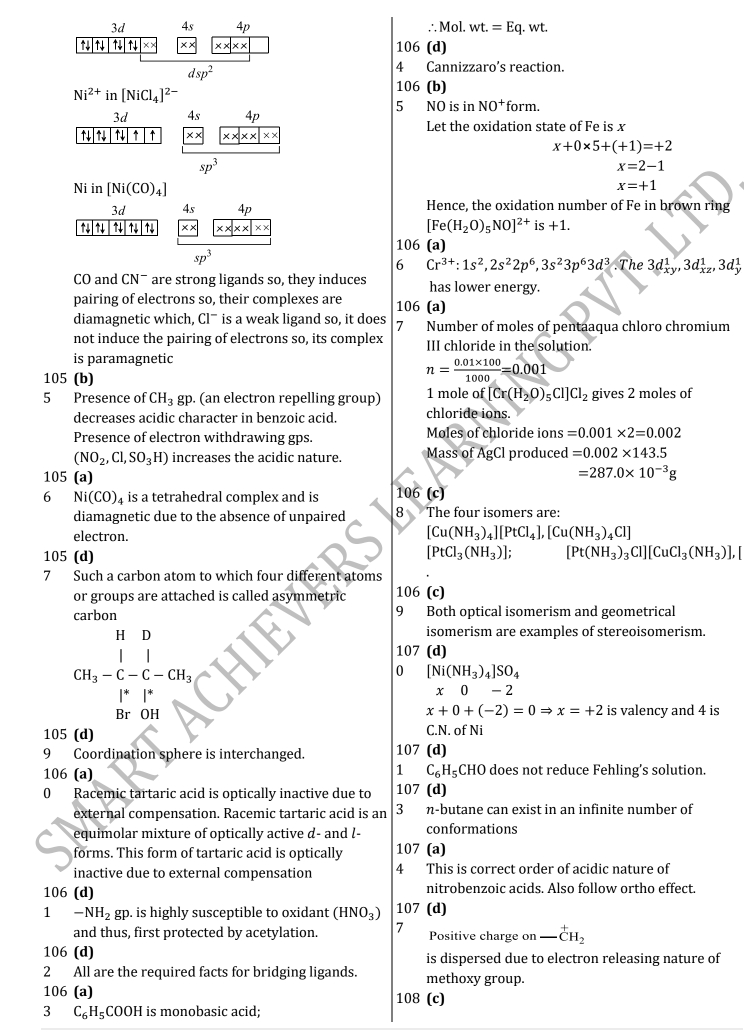
103 (d)

- 2 All involve  $dsp^2$  -hybridization.
- 103 (a)
- 3 Cis- isomer of [Pt(NH<sub>3</sub>)<sub>2</sub>Cl<sub>2</sub>] is used as an anticancer drugs for treating several type of malignant tumours when it is inject into the blood stream the more reaction Cl groups are lost so, the Pt atom bonds to a N-atom in guanosine (a part of DNA). This molecule can bond to two different guanosine units and by bridging between them it upsets the normal reproduction of DNA

4 
$$C_{H_1N_2C1} + \bigoplus_{P_2 \to d} \bigoplus_{p_2 \to d}$$

to zero.

CH



0 An organometallic compound is considered as a substance contains a carbon-metal bond e.g.,  $Al_2(CH_3)_6$ .

108 (c)

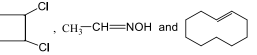
Picramide is 2, 4, 6-trinitroaniline. 1

108 (d)

Except alkynes, chain isomerism is observed 2 when the number of carbon atoms is four or more than four

#### 108 (d)

The isomerism which arises due to restricted 3 rotation about a bond in a molecule is known as geometrical isomerism



All of these form geometrical isomers

- 108 (a)
- 4 The choice of principal functional group is made on the basis of the following order Carboxylic acid > sulphonic acid > anhydride > esters > acid halide > acid amide > nitrile > aldehyde > ketone > alcohol > amine

#### 108 (d)

5  $[Pt(NH_3)Cl_2Br]Cl \rightleftharpoons [Pt(NH_3)Cl_2Br]^+ + Cl^ Cl^- + Ag^+ \rightarrow AgCl$ 

#### 108 **(b)**

-COOH gp. reacts with NaHCO<sub>3</sub> to give 7 effervescence.

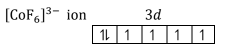
#### 108 (d)

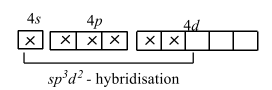
- $C_6H_5NO_2 \xrightarrow{Fe/H_2O_2(v)} C_6H_5NO_2$ 8
- 108 (c)

| Complex ion         | Hybridisation of central atom                                 |
|---------------------|---|
| $[Fe(CN)_{6}]^{4-}$ | $d^2 s p^3$ (inner)   |
| $[Mn(CN)_{6}]^{4-}$ | $d^2sp^3$ (inner)   |
| $[Co(NH_3)_6]^{3+}$ | $d^2 s p^3$ (inner)   |
| $[Ni(NH_3)_6]^{2+}$ | $sp^3d^2$ (outer)   |
|                     | $[Fe(CN)_6]^{4-}$<br>$[Mn(CN)_6]^{4-}$<br>$[Co(NH_3)_6]^{3+}$ |

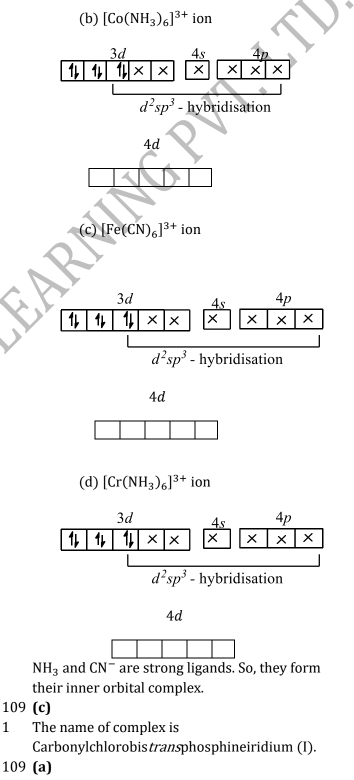
- 109 (a)
- Electronic configuration of  $Co^{2+}$  ion (a)







F<sup>-</sup> is a weak ligand. It cannot pair up electrons with *d*-subshell and forms outer orbital octahedral complex.



3 Coordination isomerism is caused by the interchange of ligands between complex cation and complex anion

109 (c)

Due to bitter almond smell. 4

#### 109 **(b)**

5 Compounds which do not show optical activity inspite of the presence of chiral carbon atoms are called meso-compounds

#### 109 (d)

6 mmol of complex  $=30 \times 0.01 = 0.3$  Also, 1 mole of complex [Cr(H<sub>2</sub>O)<sub>5</sub>Cl]Cl<sub>2</sub> gives only two moles of chloride ion when dissolved in solution.  $[Cr(H_2O)_5Cl]Cl_2 \rightarrow [Cr(H_2O)_5Cl]^{2+} + 2Cl^{-}$  $\Rightarrow$  mmol of Cl<sup>-</sup> ion produced from its 0.3 mmol =0.6

Hence, 0.6 mmol of Ag<sup>+</sup> would be required for precipitation.

- $\Rightarrow$  0.60 mmol of Ag<sup>+</sup>=0.1 M × V (in mL)
- $\Rightarrow$  V=6 mL

#### 109 (c)

This is Kekule's view for C<sub>6</sub>H<sub>6</sub> structure. 7

#### 109 (c)

C = 0 double bond of a carbonyl group is a 8 stronger bond (> C = 0,364 kJ/mol) than the C = C bond strength 250 kJ/mol) of the enol. Thus, CH<sub>3</sub>COCH<sub>3</sub> is more stable

#### 109 **(b)**

9 Effective atomic number (EAN) = Atomic no. of metal

-Oxidation no.+ Coordination no.×2  $For[CoF_6]^{2-}$ , the oxidation state of cobalt is 4.

 $EAN = (27-4) + 6 \times 2$ =23+12=35

#### 110 (d)

Metal atom or cation acts as Lewis acid or 3 electron pair acceptor.

110 (c)

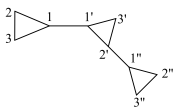
The nitration of  $C_6H_6$  does not occur at room 4 temperature. The solution becomes dark red due to absorption of NO<sub>2</sub> given out by HNO<sub>3</sub>.

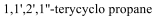
110 (c)

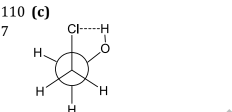
[Co(en)<sub>2</sub>NO<sub>2</sub>Cl]Br exhibits linkage isomerism 5 because the NO<sub>2</sub> group can exist as nitrito (--ONO) and nitro  $(-NO_2)$  group. The linkage isomers of [Co(en)<sub>2</sub>NO<sub>2</sub>Cl]Br are as  $[Co(en)_2NO_2Cl]Br$  and  $[Co(en)_2ONOCl]Br$ .

110 (c)

The compound will be numbered as and can be 6 named as unbranched assembles containing 3 or more identical cycles







The gauch conformation is most stable due to presence of H-bonding between H atom of OH and Cl

110 (c)

7

The electronic configuration of Ni in 8  $[Ni(CN)_4]^{2-}$ ,  $[Ni(Cl)_4]^{2-}$  and  $Ni(CO)_4$  are :  $Ni^{2+}$  in  $[Ni(CN)_4]^{2-}$ :

$$\begin{array}{c|c} 3d & 4s & 4p \\ \hline \uparrow \downarrow \uparrow \downarrow \uparrow \downarrow \uparrow \downarrow \times \times & \times \\ \hline dsp^2 \end{array}$$

Ni<sup>2+</sup> in [Ni(Cl)<sub>4</sub>]<sup>2-</sup> :  

$$3d \qquad 4s \qquad 4p$$

 $Ni^{2+}$  in  $[Ni(CO)_4]$  :

- 110 **(a)**
- 9 Follow IUPAC rules.
- 111 **(b)**
- Cd has no unpaired electron in  $[CdCl_4]^{2-}$  ion. 0
- 111 (d)

1

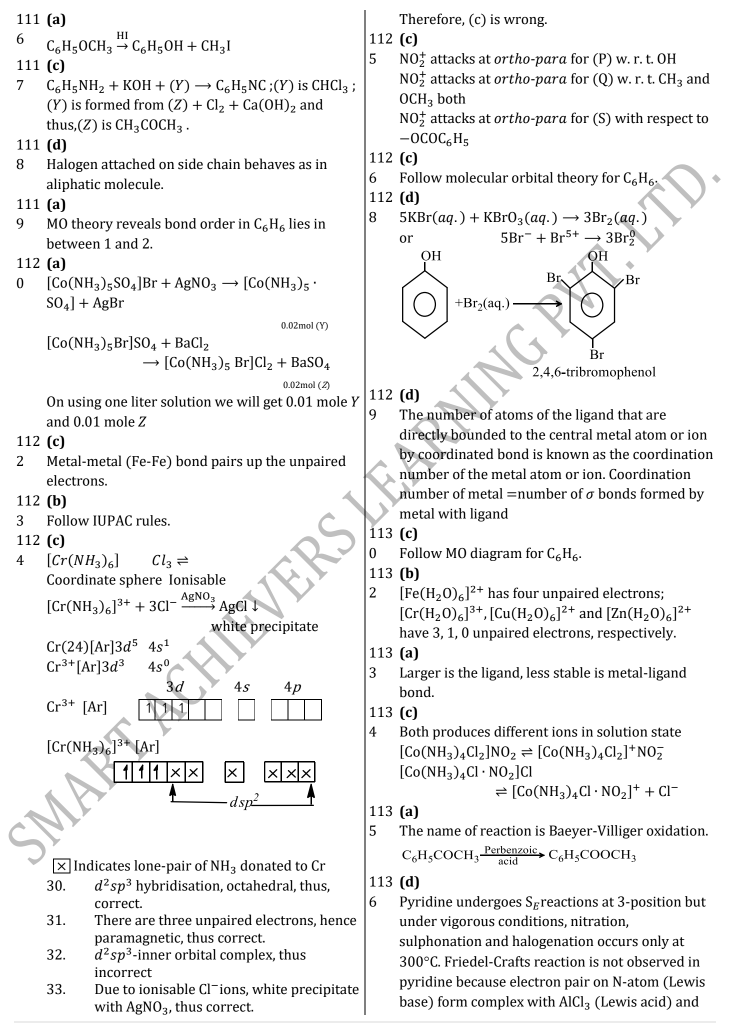
- It is a fact.
- 111 (a)
- 2 On ionization it gives maximum number of (four) ions.
- 111 (a)
- 3 Chlorophyll contains Mg, hence (a) is incorrect statement.
- 111 **(b)**

 $[Co(NH_3)_5Cl]^{2+} + 2Cl^- \rightarrow [Co(NH_3)_5Cl]Cl_2$ 4

111 (b)

5 Linkage isomerism is caused due to presence of ambidentate ligands.

 $[Pd(PPh_3)_2(NCS)_2]$  and  $[Pd(PPh_3)_2(SCN)_2]$  are linkage isomers due to SCN, ambidentate ligand.



a+ve charge on N-atom so produced decreases the activity of pyridine for  $S_E$  reaction.

113 (d)

7

- 2Cl<sup>-</sup> ions are inisable ∴ [Co(NH<sub>3</sub>)<sub>5</sub>Cl]Cl<sub>2</sub>  $\rightleftharpoons \underbrace{[Co(NH_3)_5Cl]^{2+} + 2Cl^{-}}_{3 \text{ ions}}$ 2Cl<sup>-</sup> + 2AgNO<sub>3</sub>  $\rightarrow$  2AgCl + 2NO<sub>3</sub><sup>-</sup>
- 113 (d)
- 8 Benzene is very good solvent.
- 113 **(a)**
- 9 Salol is phenyl salicylate and is used as antiseptic. 114 **(a)**
- 0  $-NO_2$  gp. is deactivating gp. for S<sub>E</sub> reaction.

#### 114 **(d)**

1 Geometrical isomerism is found in compounds having coordination no. 4 (square planar and not tetrahedral shape) as well as coordination no.6. Coordination no. 4 (square planar)

 $M_{A_2BC}, M_{A_2B_2}$ .,

Showing geometrical isomerism $M_{ABCD}$ Coordination no. 6 (octahedral)

 $M_{A_4B_2}, M_{A_4BC},$ 

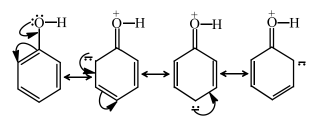
Showing geometrical isomerism.

 $M_{A_{3}B_{3}}, M_{(AA)_{2}B_{2}}$ 

#### 114 **(a)**

3 The presence of *m*-directing groups in benzene nucleus simply decreases electron density at *o*-and *p*- whereas no change in electron density at *m*-position is noticed.

On the contrary *o*- and *p*-directing groups in nucleus increase the electron density at *o*- and *p*-position.



Thus, presence of o- and p-directing groups provide seats for  $S_E$  reactions or activates the ring, whereas presence of m-directing groups does not activate the ring and thus, deactivate the ring for  $S_E$  reactions.

- 114 **(a)**
- 4 Aniline is basic.
- 114 **(d)**
- 5 Ferrocene is diphenyl iron complex.
- 114 **(c)**
- 6 Effective atomic number (EAN)

= Atomic no. 
$$-0.S. + 2 \times C.N.$$
  
= 28-0+2×4  
= 28+8

- 114 **(d)**
- 8  $[CO(en)_2Cl_2]$  forms optical and geometrical isomers.
- 114 **(c)**
- 9 Only *cis*-octahedral compounds show optical activity.
- 115 **(b)**
- 0 A square planar complex results from  $dsp^2$ hybridisation involving  $(n-1) d_{x^2-y^2}$ , ns,  $np_x$  and  $np_y$  atomic orbitals.
- 115 **(a)**
- 2 HNO<sub>3</sub> accepts a proton from H<sub>2</sub>SO<sub>4</sub>. H<sub>2</sub>SO<sub>4</sub>  $\rightleftharpoons$  H<sup>+</sup> + HSO<sub>4</sub><sup>-</sup> HNO<sub>3</sub> + H<sup>+</sup>  $\rightarrow$  H<sub>2</sub>O + NO<sub>2</sub><sup>+</sup>

115 **(c)** 

3

Lithium tetrahydroaluminate is Li[Al(H)<sub>4</sub>]

# **COORDINATION COMPOUNDS**

#### CHEMISTRY

#### Assertion - Reasoning Type

This section contain(s) 0 questions numbered 1 to 0. Each question contains STATEMENT 1(Assertion) and STATEMENT 2(Reason). Each question has the 4 choices (a), (b), (c) and (d) out of which **ONLY ONE** is correct.

- a) Statement 1 is True, Statement 2 is True; Statement 2 is correct explanation for Statement 1
- b) Statement 1 is True, Statement 2 is True; Statement 2 is not correct explanation for Statement 1
- c) Statement 1 is True, Statement 2 is False

d) Statement 1 is False, Statement 2 is True

| Statement 1: | The IUPAC name of the $CH_3 - CH = CH - CH_3 - CH = CH - CH_3 - CH = CH - CH_3 - CH - CH_3 -$ | $C \equiv CH$ is pent-3-en-1-yne and not pent-2-ene- |
|--------------|--|--|
|              | 4-yne  |  |

**Statement 2:** Lowest locant rule for multiple bond is preferred

#### 2

|   | Statement 1: | CH <sub>3</sub> O   |
|---|--------------|---|
|   |              | $CH_3 - \dot{C}H - CH_2 - \ddot{C} - OH_{is}$ 3-methyl butanoic acid                              |
|   | Statement 2: | In poly functional group, the substituent should be given lower number than the principal         |
| - |              | functional group  |
| 3 |              |   |
|   | Statement 1: | Highly charged cations are expected to form most strongly acidic hydrated cations                 |
|   | Statement 2: | The acidity of a hydrated metal ion depends on the strength of the bond between cation and oxygen |
| 4 |              |   |
|   | Statement 1: | CH <sub>3</sub>   |
|   |              | is 3-methyl cyclopentene  |
|   | Statement 2: | In the numbering, double bonded carbon atom gets preference to the alkyl group in                 |
| 5 | Nr.          | cycloalkenes  |
| 5 | Statement 1: | Hydrazine has two N as donor atoms and behaves as a chelating ligand                              |
|   | Statement 2: | Hydrazine is a neutral ligand   |
| 6 |              |   |
|   | Statement 1: | EDTA forms complexes with a large number of metal ions  |

Statement 2: It coordinates with 6 points of attachement to the metal, 4 0-atoms and two N atoms

|    | Statement 1: | A compound whose molecule has D configuration will always be dextrorotatory   |
|----|--------------|---|
|    | Statement 2: | Compounds having D configuration may be dextrorotatory or levorotatory  |
| 8  |              |   |
| U  | Cha.h        |   |
|    | Statement 1: | The IUPAC name for the compound, NCCH <sub>2</sub> CH <sub>2</sub> COOH is 3-cyano propanoic acid   |
|    | Statement 2: | -COOH is considered as substituent group while - CN is considered as the principal  |
| 9  |              | functional group  |
|    | Statement 1: | F- ion is weak ligand and forms outer orbital complex.  |
|    |              |   |
|    | Statement 2: | F- ion can not force the electrons of d $_{z^2}$ and d $_{x^2-y^2}$ orbitals to occupy d <sub>xy</sub> d <sub>yz</sub> and d <sub>zx</sub> orbitals of the same shell.          |
| 10 |              |   |
|    | Statement 1: | CH <sub>3</sub> – CH(Cl)OH is a optically active compound   |
|    | Statement 2: | Dissymetry arises in a chemical compound owing to the presence of chiral centre   |
| 11 |              |   |
| 11 |              |   |
|    | Statement 1: | Ions with more than five 3 <i>d</i> electrons have usually slightly larger magnetic moment than   |
|    | Statement 2: | calculated on the basis of $\mu = \sqrt{n(n+2)}$<br>The maximum number of unpaired <i>d</i> -electrons is five as in Mn <sup>2+</sup> and Fe <sup>3+</sup> and so, $\mu = 5.92$ |
|    |              | for 5 unpaired electrons  |
| 12 |              |   |
|    | Statement 1: | Geometrical isomerism is also called C is- trans isomerism.   |
|    | Statement 2: | Tetrahedral complexes shows geometrical isomerism.  |
| 13 |              |   |
|    | Statement 1: | [Cr(NH <sub>3</sub> ) <sub>6</sub> ] <sup>3+</sup> is paramagnetic.   |
|    |              |   |
|    | Statement 2: | $[Cr(NH_3)_6]^{3+}$ shows $d^2sp^3$ hybridisation.  |
| 14 | Ċ            |   |
|    | Statement 1: | A hydroxy group directly attached to a carbonyl group constitutes a carboxyl group  |
|    | Statement 2: | Ester is a family of carboxylic acid derivatives in which – OH group is altered by other  |
| 15 |              | group   |
| 15 |              |   |
|    | Statement 1: | In keto-enol tautomerism of dicarbonyl compounds, the enol form is preferred in contrast to the keto-form   |
|    | Statement 2: | The enol form is more stable due to resonance   |
| 16 |              |   |
|    | Statement 1: | $[Ti(H_2O)_6]^{3+}$ is coloured while $[Sc(H_2O)_6]^{3+}$ is colourless.  |
|    | Statement L  |   |

|    | Statement 2: | <i>d-d</i> transition is not possible in $[Sc(H_2O)_6]^{3+}$ .   |
|----|--------------|--|
| 17 |              |  |
|    | Statement 1: | The conversion of an optically active compound into its enantiomer is called Walden inversion                        |
|    | Statement 2: | A racemic mixture is optically inactive due to internal compensation   |
| 18 |              |  |
|    | Statement 1: | <i>d-d</i> transition is not possible in $[Sc(H_2O)_6]^{3+}$   |
|    | Statement 2: | $[Ti(H_20)_6]^{4+}$ is coloured while $[Sc(H_20)_6]^{3+}$ is colourless  |
| 19 |              |  |
|    | Statement 1: | Staggered form is less stable than the eclipsed form   |
|    | Statement 2: | The conformation in which the bond pairs of two central atoms are very far from one another is called staggered form |

# COORDINATION COMPOUNDS

|    |                |     |   |       | C   | HEM | ISTRY | ľ |     |   |     |   |     |   |  |
|----|----------------|-----|---|-------|-----|-----|-------|---|-----|---|-----|---|-----|---|--|
|    | : ANSWER KEY : |     |   |       |     |     |       |   |     |   |     |   |     |   |  |
| 1) | а              | 2)  | С | 3) b  | 4)  | a   | 13)   | b | 14) | b | 15) | а | 16) | а |  |
| 5) | d              | 6)  | а | 7) d  | 8)  | С   | 17)   | С | 18) | С | 19) | d |     |   |  |
| 9) | a              | 10) | a | 11) b | 12) | C   |       |   |     |   |     |   |     |   |  |

# **COORDINATION COMPOUNDS**

#### CHEMISTRY

## : HINTS AND SOLUTIONS :

8

9

#### 1 (a)

If however, there is a choice in numbering, the double bond is always given preference over the triple bond

#### 2 **(c)**

The functional group is – COOH, the numbering is done from RHS to give minimum number to carbon atom bearing the functional group. The given compound is a derivative of butane. The substituent is the methyl group. So, the above compound is 3-methyl butanoic acid

#### 3 **(b)**

Trivalent cations, as a group are more acidic than divalent cations and most monovalent cations have negligible acidic character

When cation-oxygen bond is strong, the bond between oxygen and hydrogen is weakened and the proton can be donated to a base

#### 4 **(a)**

In naming cycloalkanes, number the ring to give the double bonded carbons 1 and 2 choose the direction of numbering. So, that the substituents get the lowest numbers. The position of the double bond is not indicated because it is known to bond between C-1 and C-2

#### 5 **(d)**

 $\rm NH_2 - \rm NH_2$  is neutral ligand. It does not act as bidentate because when it acts as bidentate, a three membered ring will be formed, that will highly strained

## 6 **(a)**

EDTA binds to both Ca<sup>2+</sup>and Mg<sup>2+</sup>,*ie*, cations with noble gas configuration as well as to transition metal ions. It is so efficient in binding metal ions that it is used to remove traces of metal from distilled water

#### 7 **(d)**

The configuration in a compound is independent of its physical properties like optical activity

#### (c)

-COOH is the principal group while - CN is the substituent group

#### (a)

F<sup>-</sup> ion is weak field ligand and it forms outer orbital complex as it cannot force the electrons to pair up.

10 **(a)** 

In chiral centre, chiral axis and chiral plane, if any one is present, the molecule will be optically active

#### 11 **(b)**

The slightly larger value of  $\mu$  than expected from the formula  $\mu = \sqrt{n(n+2)}$  is due to a small contribution from the orbital anguler momentum of the electrons to the magnetic moment

12 (c)

Square planar complexes having  $dsp^2$  hybridisation shows geometrical isomerism.

#### 13 **(b)**

The  $Cr^{3+}$  ions has three unpaired electrons. It has  $d^2 sp^3$  hybridisation and it is paramagnetic.

## 14 **(b)**

If the acidic hydrogen of carboxylic acid is replaced by an aryl or alkyl group, the resulting structure is a carboxylate ester

(H)
$$RC \bigcirc O$$
 or (H) $RCO_2R'$ 

## 15 **(a)**

Resonance stabilisation of enol form can be shown as

$$\begin{array}{c} CH_3 - C = CH - C - CH_3 \\ | \\ O - H - \cdots - O \end{array} \xrightarrow{CH_3 - C - CH_3 CH_3 - C - - C - CH_$$

#### 16 **(a)**

 $[Sc[H_2O_6]^{3+}]$  has no unpaired electrons in its *d*-subshell and thus *d*-*d* transition is not possible

whereas  $[Ti(H_2O_6)]^{3+}$  has one unpaired electron in its d-subshell which gives rise to *d*-*d* transition to impart colour.

#### 17 **(c)**

Racemic mixture is optically inactive because the two enantiomers rotate the plane polarised light equally in opposite directions and cancel each others rotation. This phenomenon is called external compensation

#### 18 **(c)**

Both  $[Ti(H_2O)_6]^{4+}$  and  $[Sc(H_2O)_6]^{3+}$  are colourless due to absence of free electrons in 3d subshell

#### 19 **(d)**

The staggered form is more stable than the eclipsed form because the potential energy of staggered form in which the bond pairs of two carbons are far away from each other is minimum

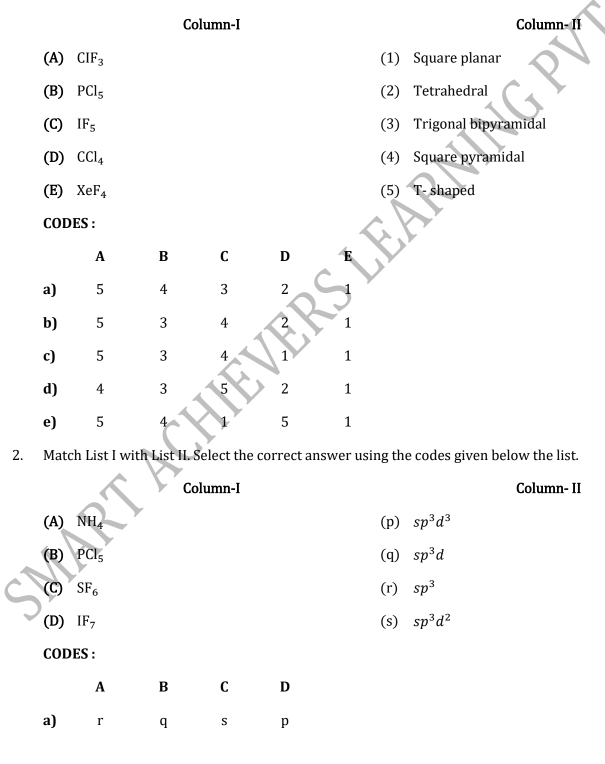
# **CHEMICAL BONDING AND MOLECULAR STRUCTURE**

#### CHEMISTRY

#### Matrix-Match Type

This section contain(s) 0 question(s). Each question contains Statements given in 2 columns which have to be matched. Statements (A, B, C, D) in **columns I** have to be matched with Statements (p, q, r, s) in **columns II**.

1. Match List I and List II and pick out correct matching codes from the given choices.



|    | b)         | р                             | q           | r            | S         |          |         |                              |
|----|------------|-------------------------------|-------------|--------------|-----------|----------|---------|------------------------------|
|    | c)         | q                             | r           | р            | S         |          |         |                              |
|    | d)         | S                             | р           | q            | r         |          |         |                              |
| 3. | Mate       | ch List I wit                 | h List II a | and choose   | the corre | ct match | ning co | odes from the choices given. |
|    |            |                               | C           | olumn-I      |           |          |         | Column- II                   |
|    | (A)        | PCl <sub>5</sub>              |             |              |           |          | (1)     | Linear                       |
|    | <b>(B)</b> | IF <sub>7</sub>               |             |              |           |          | (2)     | Pyramidal                    |
|    | (C)        | $H_{3}0^{+}$                  |             |              |           |          | (3)     | Trigonal bipyramidal         |
|    | (D)        | ClO <sub>2</sub>              |             |              |           |          | (4)     | Tetrahedral                  |
|    | <b>(E)</b> | $\rm NH_4^+$                  |             |              |           |          | (5)     | Pentagonal bipyramidal       |
|    |            |                               |             |              |           |          | (6)     | Angular                      |
|    | COD        | ES:                           |             |              |           |          |         |                              |
|    |            | Α                             | В           | С            | D         | Е        |         | 20                           |
|    | a)         | 3                             | 5           | 2            | 1         | 4        | 7~      |                              |
|    | b)         | 3                             | 5           | 4            | 1         | 4        | Ś       |                              |
|    | c)         | 3                             | 5           | 6            | 1         | 4        |         |                              |
|    | d)         | 3                             | 5           | 2            | 6         | 4        |         |                              |
|    | e)         | 3                             | 5           | 2            | 4         | 4        |         |                              |
| 4. | Mate       | ch List I wit                 | h List II a | and select t | he correc | t answer | r:      |                              |
|    |            |                               | C           | olumn-I      |           |          |         | Column- II                   |
|    | (A)        | ICI <sup>-</sup>              |             |              |           |          | (1)     | Linear                       |
|    | <b>(B)</b> | BrF <sub>2</sub> <sup>+</sup> |             |              |           |          | (2)     | Pyramidal                    |
|    | (C)        | ClF <sub>4</sub>              | *           |              |           |          | (3)     | Tetrahedral                  |
|    | (D)        | AlCl <sub>4</sub>             |             |              |           |          | (4)     | Square planar                |
|    | S          |                               |             |              |           |          | (5)     | Angular                      |
| 5  | COD        | ES :                          |             |              |           |          |         |                              |
|    |            | Α                             | В           | С            | D         |          |         |                              |

a)

b)

c)

| d) | 5 | 1 | 3 | 4 |
|----|---|---|---|---|
|----|---|---|---|---|

5. Match the compounds in the list I with that in List II.

| 5. | Mate                | ch the comp                   | ounds in t    | the list I w | rith that in | List II. |       |                               |
|----|---------------------|-------------------------------|---------------|--------------|--------------|----------|-------|-------------------------------|
|    |                     |                               | Col           | umn-I        |              |          |       | Column- II                    |
|    | (A)                 | XeO <sub>3</sub>              |               |              |              |          | (p)   | Planar triangular             |
|    | <b>(</b> B <b>)</b> | XeOF <sub>4</sub>             |               |              |              |          | (q)   | T- shape                      |
|    | (C)                 | BO <sub>3</sub> <sup>3-</sup> |               |              |              |          | (r)   | Trigonal pyramid              |
|    | (D)                 | CIF <sub>3</sub>              |               |              |              |          | (s)   | Square pyramid                |
|    | <b>(E)</b>          | $I_3^-(aq)$                   |               |              |              |          | (t)   | Linear                        |
|    |                     |                               |               |              |              |          | (u)   | Bent                          |
|    | COD                 | ES :                          |               |              |              |          |       |                               |
|    |                     | Α                             | В             | С            | D            | Ε        |       |                               |
|    | a)                  | р                             | S             | r            | q            | t        |       |                               |
|    | b)                  | q                             | S             | р            | r            | t        |       |                               |
|    | c)                  | r                             | S             | р            | q            | t        | 1.    |                               |
|    | d)                  | S                             | S             | р            | q            | t        | Ś     |                               |
| 6. | Mate                | ch List I with                | n List II. Se | elect the c  | orrect ans   | wer usi  | ng th | e codes given below the list. |
|    |                     |                               | Col           | umn-I        |              | 2        |       | Column- II                    |
|    | (A)                 | $\rm NH_4$                    |               |              | K)           |          | (p)   | $sp^3d^3$                     |
|    | <b>(</b> B <b>)</b> | PCl <sub>5</sub>              |               |              |              |          | (q)   | $sp^{3}d$                     |
|    | (C)                 | SF <sub>6</sub>               |               | $\sum$       |              |          | (r)   | sp <sup>3</sup>               |
|    | (D)                 | IF <sub>7</sub>               |               |              |              |          | (s)   | $sp^3d^2$                     |
|    | COD                 | ES:                           |               |              |              |          |       |                               |
|    |                     | A                             | В             | С            | D            |          |       |                               |
|    | a)                  | r                             | q             | S            | р            |          |       |                               |
|    | b)                  | р                             | q             | r            | S            |          |       |                               |
| 5  | c)                  | q                             | r             | р            | S            |          |       |                               |
|    | d)                  | S                             | р             | q            | r            |          |       |                               |
|    |                     |                               |               |              |              |          |       |                               |

# **CHEMICAL BONDING AND MOLECULAR STRUCTURE**

CHEMISTRY

| 1)       | b  | 2) | а    | 3) | d | : AN<br>4) | SWER<br>c | KEY : |   |   |   |                   |            |
|----------|----|----|------|----|---|------------|-----------|-------|---|---|---|-------------------|------------|
| 1)<br>5) | d  | 6) | a    | 55 | u | чj         | C         |       |   |   |   |                   | $\bigcirc$ |
|          |    |    |      |    |   |            |           |       |   |   |   | $\langle \rangle$ |            |
|          |    |    |      |    |   |            |           |       |   |   | Ś |                   |            |
|          |    |    |      |    |   |            |           |       |   | Q | 7 |                   |            |
|          |    |    |      |    |   |            |           |       |   |   |   |                   |            |
|          |    |    |      |    |   |            |           |       | 2 |   |   |                   |            |
|          |    |    |      |    |   |            |           | N     |   |   |   |                   |            |
|          |    |    |      |    |   |            | $\sim$    |       |   |   |   |                   |            |
|          |    |    |      |    |   | ~~~        | 2         |       |   |   |   |                   |            |
|          |    |    |      |    | 5 |            |           |       |   |   |   |                   |            |
|          |    |    |      | E. | Ŷ |            |           |       |   |   |   |                   |            |
|          |    |    | 5    |    |   |            |           |       |   |   |   |                   |            |
|          |    | 25 | , V, |    |   |            |           |       |   |   |   |                   |            |
| (        | 4, |    |      |    |   |            |           |       |   |   |   |                   |            |
| S        |    |    |      |    |   |            |           |       |   |   |   |                   |            |
|          |    |    |      |    |   |            |           |       |   |   |   |                   |            |

# **CHEMICAL BONDING AND MOLECULAR STRUCTURE**

CHEMISTRY

|        |   |  | : HINTS   | SAND    | 20 | LUTIONS :   |
|--------|---|--|---|---------|----|---|
| -      | <b>b)</b>   |  |   |         |    | $AlCl_4^- \Rightarrow 4bp + 0lp$ (thus $sp^3$ hybridisation) =  |
|        |   |  | (Structure)   |         |    | tetrahedral geometry  |
|        | (Compound   |  |   |         | 5  | (d)   |
|        | $\begin{array}{rrrr} 1. & CIF_{3} \\ 2. & PCI_{5} \\ 3. & IF_{5} \\ 4. & CCI_{4} \\ 5. & XeF_{4} \end{array}$ | 2.<br>3.   | <ol> <li>Trigonal<br/>bipyramidal</li> <li>Square<br/>pyramidal</li> <li>Tetrahedral</li> <li>Square</li> </ol> |         |    | XeO <sub>3</sub> ( $sp^3$ hybridisation)<br>= contain a loan pai<br>= trigonal pyram<br>shape<br>XeOF <sub>4</sub> ( $sp^3 d$ hybridisation) = one lone pair<br>= pyramidal shape<br>PO <sup>3-</sup> ( $ap^2$ hybridisation) = planar triangular |
| L      |   |  | planar  |         |    | $BO_3^{3-}(sp^2 \text{ hybridisation}) = \text{planar triangular}$<br>$ClF_3(sp^3d^2 \text{ hybridisation})$<br>= two lone pair<br>= T  shaped  |
| (<br>Г | (a)   |  | 1   |         |    |   |
|        | Molecule<br>/ion  | Type of<br>hybridis<br>ation   |   |         | 6  | $I_3^-$ ( <i>sp</i> <sup>3</sup> <i>d</i> hybridisation) = 3 loan pair=linear<br>(a)  |
|        | A. NH <sub>4</sub> <sup>+</sup>   | r. <i>sp</i> <sup>3</sup>  |   |         | C. | Ethane $CH_3 - CH_3 2sp^2$ hybrid carbon  |
|        | B. PCl <sub>5</sub><br>C. SF <sub>6</sub><br>D. IF <sub>7</sub>   | q. <i>sp</i> <sup>3</sup> <i>d</i><br>s. <i>sp</i> <sup>3</sup> <i>d</i> <sup>2</sup><br>p. <i>sp</i> <sup>3</sup> <i>d</i> <sup>3</sup> |   | 5       |    | Ethylene $CH_2 = CH_2 2sp^2$ hybrid carbon<br>Acetylene $CH \equiv CH 2sp$ hybrid carbon  |
| L      |   | <u> </u>   |   |         |    |   |
| (      | d)  |  |   |         |    |   |
|        | $PCl_5 = sp^3d$ (Trigonal pyramidal)  |  |   |         |    |   |
|        | $IF_7 = sp^3d^3$ (Pentagonal bipyramidal)   |  |   |         |    |   |
|        | $H_3O^+ = sp^3$ (Pyramidal)   |  |   | 1.      |    |   |
|        | $ClO_2 = sp^2$ (Angular) bond length are shorted than single bond due to resonance.                           |  |   | shorter |    |   |
|        | $NH_{4}^{+} = sp^{3}$ (Tetrahedral)   |  |   |         |    |   |
|        |   |  |   |         |    |   |
| Ι      | c)<br>$Cl_2^- \Rightarrow 2bp +$<br>hybridization   |  |   |         |    |   |
|        | $3rF_2^+ \Rightarrow 2bp$   |  | ometi y   |         |    |   |
|        |   |  | on)=angular geo   | metry   |    |   |
|        | $\mathrm{ClF}_4^- \Rightarrow 4bp$ -  | + 2 <i>lp</i> (thus, :   | $sp^3d^2$ hybridisat  | ion) =  |    |   |
| × .    | auare nlana   | r geometry   |   |         | 1  |   |