

Answer Key Version - Q (NEET FRESH All Batches)

Physics					Chemistry				
Sec.A	11. 1	22. 2	33. 1	43. 3	Sec. A	61. 3	72. 3	83. 1	93. 1
01. 1	12. 3	23. 3	34. 4	44. 1	51. 4	62. 1	73. 3	84. 1	94. 3
02. 4	13. 4	24. 1	35. 3	45. 1	52. 2	63. 1	74. 2	85. 4	95. 3
03. 3	14. 3	25. 2	Sec. B	46. 1	53. 3	64. 4	75. 1	Sec. B	96. 2
04. 4	15. 1	26. 2	36.502900	14 7 015 ber	fie54. 4	65. 3 _®	76. 1	86. 4	97. 4
05. 4	16. 3	27. 2	37. 1	48. 2	55. 3	66. 2	77. 1	87. 2	98. 1
06. 1	17. 2	28. 3	38. 2	49. 3	56. 2	67. 1	78. 2	88. 3	99. 4
07. 3	18. 2	29. 1	39. 3	50. 3	57. 1	68. 1	79. 4	89. 2	100. 2
08. 1	19. 2	30. 1	40. 4	KNE	58. 3	69. 4	80. 3	90. 2	
09. 1	20. 4	31. 1	41. 1		59. 4	70. 1	81. 3	91. 3	
10. 2	21. 3	32. 3	42. 3		60. 3	71. 4	82. 1	92. 2	
Biology									
Part-I	110. 1	121. 4	132. 2	142. 3	Part-II	160. 2	171. 1	182. 1	192. 1
Sec.A	111. 1	122. 2	133. 4	143. 4	Sec.A	161. 1	172. 4	183. 1	193. 4
101. 2	112. 3	123. 3	134. 3	144. 1	151. 4	162. 4	173. 4	184. 4	194. 4
102. 3	113. 1	124. 3	135. 3	145. 3	152. 1	163. 4	174. 2	185. 2	195. 2
103. 4	114. 1	125. 2	Sec.B	146. 3	153. 4	164. 1	175. 2	Sec. B	196. 3
104. 3	115. 1	126. 2	136. 1	147. 2	154. 2	165. 1	176. 2	186. 3	197. 2
105. 1	116. 4	127. 1	137. 4	148. 4	155. 2	166. 4	177. 1	187. 2	198. 3
106. 4	117. 4	128. 1	138. 2	149. 2	156. 2	167. 3	178. 1	188. 3	199. 3
107. 1	118. 1	129. 1	139. 2	150. 1	157. 4	168. 3	179. 2	189. 1	200. 2
108. 3	119. 1	130. 1	140. 3		158. 4	169. 4	180. 3	190. 3	
109. 1	120. 3	131. 1	141. 3		159. 1	170. 4	181. 2	191. 3	

SECTION - A (35 Questions)

01. (1) As the regions are of equipotentials, so Work done $W = q\Delta V$

 ΔV is same in all the cases hence work-done will also be same in all the cases.

02. **(4)** Electrostatic potential energy of the given system is given as,

$$U = \frac{1}{4\pi\varepsilon_0} \times \frac{1}{a} (Qq + Qq + q^2)$$
$$U = 0 \qquad 2Qq = -q^2 \quad \Rightarrow \quad Q = -q/2$$

03. **(3)** In uniform electric field dipole experience only torque, but no force.

04. (4)
$$\begin{array}{c} Q \\ \downarrow 2 \\ \downarrow 3 \\ \downarrow 4 \\ \downarrow 2 \\ \downarrow 2 \\ \downarrow 2 \\ \downarrow 3 \\ \downarrow 4 \\$$

 \Rightarrow value of q such that all three charges are in equilibrium.

equilibrium.
i.e. F...at B = zero

$$\hat{F}_{BA} + \hat{F}_{BC} = 0$$
 or $|\hat{F}_{BA}| = |\hat{F}_{BC}|$
 $\frac{1}{4\pi\epsilon_0} \frac{Q^2}{l^2} = \frac{1}{4\pi\epsilon_0} \frac{Qq}{l^2/4}$
 $\Rightarrow Q = q(4) \Rightarrow q = \left(\frac{Q}{4}\right)$
As, $\hat{F}_{BA} = -\hat{F}_{BC} \Rightarrow q = \left(-\frac{Q}{4}\right)$
(4)

The flux passing through the square of 1 m placed in xy-plane inside the electric field is zero because

$$\phi = EA\cos 90^\circ = 0$$

05.

06. (1) As per Coulomb's law, $F = k \frac{q_1 q_2}{r^2}$

According to questions,

$$F = k \frac{(q_1/2)(q_1/2)}{(r/2)^2} = k \frac{q_1 q_2}{r^2} = F$$

This, force between them will remain same.

07. **(3)** Electric field is perpendicular to the equipotential surface and is zero every where inside the metal.

- 08. (1) By using QE = mg $\Rightarrow E = \frac{mg}{Q} = \frac{10^{-6} \times 10}{10^{-6}} = 10$ V/m upward because charge is positive.
- 09. (1) We know that, electric field at distance r from an infinitely long line charge is given by

$$E = \frac{\lambda}{2\pi\varepsilon_0 r} \Longrightarrow \lambda = 2\pi\varepsilon_0 \cdot rE$$

10. (2) Using, q = ne

$$\Rightarrow 2 = n \times 1.6 \times 10^{-19}$$

$$\Rightarrow n = \frac{2}{1.6 \times 10^{-19}} = 1.25 \times 10^{19} = 125 \times 10^{17}$$

electrons

11. **(1)**
$$E = \frac{\sigma}{2\varepsilon_0}$$

12. **(3)** At
$$O, E \neq 0, V = 0$$

13. **(4)** Conducting surface behaves as equipotential surface.

14. (3) By using
$$W = Q \cdot \Delta V \Rightarrow \Delta V = \frac{2}{20} = 0.1$$
 volt

15. (1) Electric field due to all hollow spherical conductor is governed by following equation E = 0, for r < R

and
$$E = \frac{Q}{4\pi\varepsilon_0 r^2}$$
 for $r \ge R$...(i)

i.e. inside the conductor field will be zero and outside the conductor will vary according to

$$E \propto \frac{1}{r^2}$$

16. **(3)** Change in kinetic energy = Force \times Displacement = qEy

17. **(2)**
$$KE - 0 = q(V_1 - V_2)$$

= 2 × 1.6 × 10⁻¹⁹ × (70 - 50) = 40 KeV

18. (2) $E_{\text{inside}} = \frac{\rho}{3\varepsilon_0} r \quad (r < R)$ $E_{\text{outside}} = \frac{\rho R^3}{3\varepsilon_0 r^2} \quad (r \ge R)$

i.e. inside the uniformly charged sphere field varies linearly $(E \propto r)$ with distance and outside varies

according to $E \propto \frac{1}{r^2}$

19. **(2)** Force on electron
$$F = QE = Q\left(\frac{V}{d}\right)$$

$$\Rightarrow F = (1.6 \times 10^{-19}) \left(\frac{1000}{2 \times 10^{-3}}\right) = 8 \times 10^{-14} \text{ N}$$

20. (4) Given, $E = (2\hat{k} + 3\hat{j} + \hat{k}) NC^{-1}$ and $S = 10\hat{i} m^{2}$ We know that, Electric flux, $\phi = \overline{E} \cdot \overline{S} = (2\hat{i} + 3\hat{j} + \hat{k}) \cdot (10\hat{i}) = 20 \text{ Nm}^2 \text{ C}^{-1}$ (3) $KE = QV \Longrightarrow KE = (2e)200 \text{ V} = 400 \text{ eV}$ 21. (2) Relation for electric field is given by $E = \frac{\lambda}{2\pi\epsilon_0 r}$ 22. (Given : $E = 7.182 \times 10^8 \,\text{N/C}$) $r = 2 \text{ cm} = 2 \times 10^{-2} \text{ m}$ $\frac{1}{4\pi\varepsilon_0} = 9 \times 10^{-9} \Longrightarrow \lambda = 2\pi\varepsilon_0 rE = \frac{2 \times 2\pi\varepsilon_0 rE}{2}$ $=\frac{2\times10^{-2}\times7.182\times10^8}{2\times9\times10^9}=7.98\times10^{-4}\,\mathrm{C/m}$ (3) Potential energy of the system 23. $\frac{-kqQ}{r} - \frac{kQq}{r} - \frac{kq^2}{2r} = 0$

$$\frac{-4kqQ + kq^2}{2x} = 0 \implies kq^2 = 4kQq$$
$$\implies \frac{q}{Q} = 4$$

24. (1) By Gauss' theorem,

$$|\phi_E| = \left| \frac{q}{\varepsilon_0} \right| = \left| \frac{-2e}{\varepsilon_0} \right| = \frac{2e}{\varepsilon_0}$$

25. (2)

26. (2) Charge flows from higher potential to low potential

$$\frac{KQ_1}{1} = \frac{KQ_2}{3} \implies Q_2 = 3Q_1$$
$$Q_1 + Q_2 = 4 \times 10^{-2} \implies 4Q_1 = 4 \times 10^{-2}$$
$$Q_1 = 10^{-2} \text{ C} \implies Q_2 = 3Q_1 = 3 \times 10^{-2}$$

Final charge on the bigger sphere

$$Q_1 = 3 \times 10^{-2} \text{ C}$$

(2) The electric field will be maximum at B, as the 27. di-rection of electric field is along decreasing potential $V_B > V_C > V_A$.

28. (3)



Hence time period

$$T = 2\pi \sqrt{\frac{l}{g'}} = 2\pi \sqrt{\frac{l}{\left(g + \frac{QE}{m}\right)}}$$
29. (1)
Equipotential
Surfaces

30. (1) From symmetry of the figure all corner have same electric potential. Therefore work done in moving the charge q from the corner to the diagonally opposite corner is zero.



31.

3

By using
$$E = 9 \times 10^9 \cdot \frac{2 pr}{(r^2 - l^2)^2}$$
; where
 $p = (500 \times 10^{-6}) \times (10 \times 10^{-2}) = 5 \times 10^{-5} \text{c} \times \text{m},$
 $r = 25 \text{ cm} = 0.25 \text{ m}, l = 5 \text{ cm} = 0.05 \text{ m}$
 $0 \times 10^9 \times 2 \times 5 \times 10^{-5} \times 0.25$

$$E = \frac{9 \times 10^9 \times 2 \times 5 \times 10^{-5} \times 0.25}{\{(0.25)^2 - (0.05)^2\}^2} = 6.25 \times 10^7 \text{ N/C}$$

32. (3) Electric potential inside a conductor is constant and it is equal to that on the surface of the conductor.

V m

33. (1) Here
$$v^2 - u^2 = 2as$$
 $v^2 = 0^2 + 2\left(\frac{qE}{m}\right)e$
 $v = \sqrt{\frac{2qE}{m}e} \implies v = \sqrt{\frac{2qEI}{m}}$

34. **(4)**
$$\vec{E} = -\frac{\partial V}{\partial x}\hat{i} - \frac{\partial V}{\partial y}\hat{j} - \frac{\partial V}{\partial z}\hat{k}$$



=

÷.

$$= -[6 - 8y]\hat{i} + (-8x - 8 + 6z)\hat{j} + (6y)\hat{k}]$$

At (1,1,1), $\vec{E} = 2\hat{i} + 10\hat{j} - 6\hat{k}$
 $\Rightarrow (\vec{E}) = \sqrt{2^2 + 10^2 + 6^2} = \sqrt{140} = 2\sqrt{35}$
 $\therefore F = q\vec{E} = 2 \times 2\sqrt{35} = 4\sqrt{35}$

35. (3) Option (1) shows lines of force starting from one positive charge and terminating at another. Option (2) has one line of force making closed loop.

> Option (4) shows all lines making closed loops. All these are not correct. Only option (3) is correct.

Section - B (Attempt Any 10 Questions)

36. (2) $A \rightarrow (2); B \rightarrow (3); C \rightarrow (4); D \rightarrow (1)$

37. (1) Net field at origin

$$E = \frac{q}{4\pi\varepsilon_0} \left[\frac{1}{1^2} + \frac{1}{2^2} + \frac{1}{4^2} + \dots \infty \right]$$
$$= \frac{q}{4\pi\varepsilon_0} \left[1 + \frac{1}{4} + \frac{1}{16} + \dots \infty \right]$$
$$= \frac{q}{4\pi\varepsilon_0} \left[\frac{1}{1 - \frac{1}{4}} \right] = 12 \times 10^9 q \,\mathrm{N/c}$$

$$38. \quad \textbf{(2)} \ V = \frac{1}{4\pi\varepsilon_0} \frac{q}{r}$$

Here, the electric potential at point A

$$V = 2V_{+\mathrm{ve}} + 2V_{-\mathrm{ve}}$$

39. (3) When charge q is released in uniform electric field E, then its acceleration

$$a = \frac{qE}{m}$$
 (constant)

So, its motion will be uniformly accelerated motion and its velocity after t second is given by

$$v = at = \frac{qE}{m}t$$

The KE of charged particle,

$$KE = \frac{1}{2}mv^2 = \frac{1}{2}m\left(\frac{qE}{m}t\right)^2 = \frac{q^2E^2t^2}{2m}$$
$$V = \frac{1}{4\pi\varepsilon_0}\left[\frac{2q}{L} - \frac{2q}{L\sqrt{5}}\right] \quad V = \frac{2q}{4\pi\varepsilon_0L}\left(1 - \frac{1}{\sqrt{5}}\right)$$

(4) Work done is equal to charge in potential 40. energy.

In Ist case, when charge +Q is situated at C.

+q	+Q	q
Ă		B
H		>
₩	2L	>

Electric potential energy of system

$$U_1 = \frac{1}{4\pi\varepsilon_0} \frac{(q)(-q)}{2L} + \frac{1}{4\pi\varepsilon_0} \frac{(-q)Q}{L} + \frac{1}{4\pi\varepsilon_0} \cdot \frac{qQ}{L}$$

In IInd case, when charge +Q is moved from C to D.

Electric potential energy of system in that case

$$U_2 = \frac{1}{4\pi\varepsilon_0} \cdot \frac{(q)(-q)}{2L} + \frac{1}{4\pi\varepsilon_0} \cdot \frac{qQ}{3L} + \frac{1}{4\pi\varepsilon_0} \frac{(-q)(Q)}{L}$$

Work done =
$$\Delta U = U_2 - U_1$$

$$= \left(-\frac{1}{4\pi\varepsilon_0}\frac{q^2}{2L} + \frac{1}{4\pi\varepsilon_0}\frac{qQ}{3L} - \frac{1}{4\pi\varepsilon_0}\frac{qQ}{L}\right)$$
$$= \left(-\frac{1}{4\pi\varepsilon_0}\frac{q^2}{2L} - \frac{1}{4\pi\varepsilon_0}\cdot\frac{qQ}{L} + \frac{1}{4\pi\varepsilon_0}\cdot\frac{qQ}{L}\right)$$
$$= \frac{qQ}{4\pi\varepsilon_0}\cdot\left[\frac{1}{3L} - \frac{1}{L}\right] = \frac{qQ}{4\pi\varepsilon_0}\frac{(1-3)}{3L}$$
$$= \frac{-2qQ}{12\pi\varepsilon_0L} = -\frac{qQ}{6\pi\varepsilon_0L}$$

41. (1) Electric field at the center of uniformly charged half ring is

$$E = \frac{2k\lambda}{R} = 2 \cdot \frac{1}{4\pi\varepsilon_0} \cdot \frac{q}{\pi R} \cdot \frac{1}{R} = \frac{q}{2\pi^2\varepsilon_0 R^2}$$

42. (3) Using $dV = -\vec{E} \cdot \vec{dr}$



$$\Rightarrow E = \frac{-(20 - 10)}{10 \times 10^{-2} \cos 120^{\circ}}$$

$$=\frac{-10}{10\times10^{-2}(-\sin 30^{\circ})}=\frac{-10^{2}}{-1/2}=200\,\mathrm{V/m}$$

43. (3) Suppose the field vanishes at a distance r, we have



$$(-q)$$
 $(0,0,0)$ $(0,0,0)$ $(0,0,0)$ $(0,0,0)$

$$\frac{Kq}{x^2} = \frac{Kq}{2(x-a)^2}$$
$$\Rightarrow 2(x-a)^2 = x^2 \Rightarrow \sqrt{2}(x-a) = x$$
$$\Rightarrow (\sqrt{2}-1)x = \sqrt{2}a \Rightarrow x = \frac{\sqrt{2}a}{\sqrt{2}-a}$$

44. (1) Electric field at a point is equal to the negative gradient of the electrostatic potential at that point.

Potential gradient relates with electric field
according to the following relation
$$E = \frac{-dV}{dr}$$

$$\vec{E} = -\frac{\partial V}{\partial x} = \left[-\frac{\partial V}{\partial x} \hat{i} - \frac{\partial V}{\partial y} \hat{j} - \frac{\partial V}{\partial z} \hat{k} \right]$$
$$= \left[\hat{i} (2xy + z^3) + \hat{j} x^2 + \hat{k} 3x z^2 \right]$$

45. (1) Intensity at 5m is same as at any point between B and C because the slope of BC is same throughout (*i.e.*, electric field between B and C is uniform). Therefore electric field at R = 5m is equal

to the slope of line *BC* hence by $E = \frac{-dV}{dr}$;

$$E = -\frac{(0-5)}{6-5} = 2.5\frac{V}{m}$$

46. (1) When dipole is given a small angular displacement θ about it's equilibrium position, the restoring torque will be

$$\tau = -pE\sin\theta = -pE\theta \quad (a\sin\theta = \theta)$$

or
$$I \frac{d^2 \theta}{dt^2} = -pE\theta$$
 (as $\tau = I\alpha = I \frac{d^2 \theta}{dt^2}$)
or $\frac{d^2 \theta}{dt^2} = w^2 \theta$ with $\omega^2 = \frac{pE}{I} \Longrightarrow \omega = \sqrt{\frac{pE}{I}}$

- 47. (1) If a body acquires positive charge, then it means that it lost few electrons. In this way its mass decreases.
- 48. (2)

$$\theta = \frac{\pi}{3} + \alpha \text{ where } \tan \alpha = \frac{1}{2} \tan \frac{\pi}{3}$$

$$\Rightarrow a = \tan^{-1} \sqrt{3}/2 \text{ so, } \theta = \frac{\pi}{3} + \tan^{-1} \sqrt{3}/2$$

- 49. **(3)** Since $V = \frac{W}{Q}$, more work will be done for a positive charge of two units as compared to positive charge of one unit, but the ratio $\frac{W}{Q}$ is same. Therefore potential difference is same.
- 50. (3) After following the guidelines mentioned above



0



CHEMISTRY

SECTION - A (35 Questions)

51. (4)

Lanthanons and Aluminium have no comparison, neither they are in same group nor in period.

52. **(2)**

Ferrous ammonium sulphate

53. **(3)**

As the number of ions if complex compounds when dissolved in water increases, molar conductivity also increases.

1 mole of [Pt(NH₃)₂Cl₂] gives one mole of ions

1 mole of $[Co(NH_3)_4Cl_2]$ Cl gives two mole of ions

1 mole of $K_4[Fe(CN)_6]$ gives five moles of ions

1 mole of $[Fe(CO)_5]$ gives zero moles of ions

[because it is a neutral molecule]

Hence $K_4[Fe(CN)_6]$ has the highest molar conductivity.

54. **(4)**

 V^{2+}, VO^{2+}

55. **(3)**

 $Fe^{2+} \xrightarrow{\text{oxidises}} Fe^{3+}$

56. **(2)**

By applying the EAN rule, the stability of metal carbonyls can be predicted. All the given metal carbonyls expect $Mn(CO)_5$ follow the EAN rule and thus, it exists as dimer $Mn_2(CO)_{10}$

57. (1)

Octahedral complexes of the type $[Ma_4B_2]$, $[Ma_2B_4]$, $[MA_3B_3]$ exhibit geometrical isomerism.

58. **(3)**

Due to d⁵ configuration, metallic bonds are weak. d⁵ orbital is half filled as a result 3d electrons are more tightly held by the nucleus and this reduces the de-localization of electrons resulting in weaker metallic bonding.

59. (4)

The actinoids exhibit more number of oxidation states in general than the lanthanoids. hence, (4) is correct.

- 60. **(3)**
- 61. **(3)**

3

 $[Ni(NH_3)_6]Cl_2$ has two unpaired electron.

62. **(1)**

The solubility of silver bromide in hypo solution due to the formation of $[Ag(S_2O_3)_2]^{3-}$.

63. **(1)**

It is one third of its molecular weight in alkaline medium

64. **(4)**

Fe(CO)₅ has dsp³-hybridisation so it show trigonal bipyramidal geometry.

65. **(3)**

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

O.N. of Co = +3



Due to paired e- it is diamagnetic

66. **(2)**

Decreases from +6 to +3.

$$\frac{K_2 C r_2 O_7 + 4 H_2 S O_4 \rightarrow K_2 S O_4 + C r_2 (S O_4)_3 + 4 H_2 O + 3 O_4}{[H_2 S + [O] \rightarrow S + H_2 O] \times 3}$$

$$\frac{K_2 C r_2 O_7 + 4 H_2 S O_4 + 3 H_2 S \rightarrow K_2 S O_4 + C r_1 (S O_4)_3 + 7 H_2 O + 3 S_4 + C r_2 (S O_4)_3 + 7 H_2 O + 3 F_2 + C r_2 (S O_4)_3 + 7 H_2 O + 3 F_2 + C r_2 + C$$

67. **(1)**

$$2KMnO_4 + KBr + H_2O$$

$$\rightarrow 2KOH + 2MnO_2 + KBrO_3$$

68. **(1)**

$$-0.6\Delta_{0}$$

$$[Co(en)_3]^{3+}, [Co(NH_3)_6]^{3+}, [Co(H_2O)_6]^{3+}$$

70. **(1)**

$$3M_{nO_{4}^{-}} + 5(Fe^{2+} + C_{2}O_{4}^{2-}) + 24H^{+} \rightarrow$$

$$3Mn^{2+} + 5Fe^{3+} + 10CO_{2} + 12H_{2}O$$

Thus 5M of FeC_2O_4 is oxidized by 3M of $KMnO_4$ then 1M of FeC_2O_4 is oxidized by 3/5 mole of $KMnO_4$.

71. (4)



There are six equivalent Cr–O bonds and one Cr–O–Cr bond.

72. **(3)**

 $Ni(CO)_4$ and $[Ni(CN)_4]^{2-}$ are diamagnetic and $NiCl_4^{2-}$ is paramagnetic.

73. **(3)**

Here Co is present as Co²⁺ ion which has 3 unpaired electrons. So the spin magnetic moment will be $\sqrt{3(3+2)}$, i.e., $\sqrt{15}$ B.M.

74. **(2)**

Colour of transition metal ion salt is due to d - d transition of unpaired electrons of d-orbital. Metal ion salt having similar number of unpaired electron in d-orbitals shows similar colour in aqueous medium



Number of unpaired electrons = 1

75. **(1)**

 $Cr^{2+} + Mn^{3+} \rightleftharpoons Cr^{3+} + Mn^{2+}$

 Cr^{3+} is a reducing agent Mn^{3+} is an oxidising agent both Cr^{2+} & Mn^{3+} exhibit d^4 electronic configuration.

76. **(1)**

 6.08×10^{-6}

0

[Co(NH₃)₃Cl₃]

78. **(2)**

The complex is of the type [Mabcd]

M = metal





79. (4)

The complexes can be written as follows

$$[\mathrm{Co}(\mathrm{NH}_3)_6]\mathrm{Cl}_3[\mathrm{Co}(\mathrm{NH}_3)_5\mathrm{Cl}]\mathrm{Cl}_2[\mathrm{Co}(\mathrm{NH}_3)_4\mathrm{Cl}_2]\mathrm{Cl}$$

80. **(3)**

According to the relation

$$\Delta_{\rm t} = \frac{4}{9} \Delta_0$$

81. **(3)**

Maximum MP in 3d series is shown by Cr because of maximum no. of unpaired e^- and hence maximum extent of d-d overelapping.

82. (1)

La(OH)₃ is less basic than Lu(OH)₃

83. (1)

Cis-platin

84. (1)

 $I^- < Cl^- < F^- < H_2O < CN^-$

85. **(4)**

[Xe]4 $f^7 6s^2$, [Xe]4 $f^7 5d^1 6s^2$ and [Xe]4 $f^9 6s^2$



SECTION - B (Attempt Any 10 Questions)

86. **(4)**

- Co(Z = 27)
- 87. **(2)**

 $[CuCl_2 \{O = C(NH_2)_2\}_2]$

88. **(3)**

The assertion is correct but the reason is false. Actually transition metal show variable valency due to very small difference between the ns^2 and (n-1)d electrons, Therefore, assertion is correct but reason is false.

 $2MnO_{4}^{-} + 16H^{+} + C_{2}O_{4}^{--}$ $\rightarrow 2Mn^{+2} + 2CO_{2} + 8H_{2}O_{4}$

90. **(2)**

Fact

91. **(3)**

As +ve charge on the central metal atom increases, the less readily the metal can donate electron density into the π^* orbitals of CO ligand to weaken the C – O bond. Hence the C – O bond would be strongest in Mn(CO)₆⁺.

92. **(2)**

There is no regular trend in the $E^0(M^{2+}/M)$ values because of irregular variation of ionization energies and sublimation energies of the atoms of the members of the transition series.

93. (1)

$$VO_2^+ < Cr_2O_7^{2-} < MnO_4^-$$

94. **(3)**



Both points (1) and (2) are correct

96. **(2)**

Moles of CoCl₃.6NH₃ = $\frac{2.675}{267.5} = \frac{1}{100}$ mole = 0.01 mole

Moles of AgCl =
$$\frac{4.78}{143.5} = \frac{3}{100}$$
 mole = 0.03 mole

0.01 mole of compound gives moles of AgCl = 0.03 mole

So, 1 mole of compound gives moles of $AgCl = \frac{0.03}{0.01} = 3$ mole. So, structural formula of compound having $3Cl^{-1}$ ions out side of coordination sphere, so formula is $[Co(NH_3)_6]Cl_3$

97. **(4)**

98. (1)

In K_3 [Fe(CN)₆] the ligand are negative which is present in coordination spheres shows a dual behaviour. It may satisfy both primary & secondary valencies while neutral ligand satisfies only secondary valencies.

99. (4)

Co(III) is more stable in octahedral complexes and Osmium shows (VIII) oxidation state.

100. (2)

The basic character of any element changes with the oxidation state, low oxidation states are more basic and high oxidation state are more acidic. For example, MnO and Mn_2O_3 are basic while Mn_2O_7 is acidic in nature.