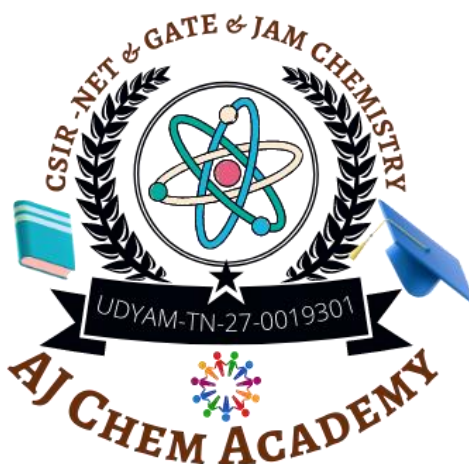


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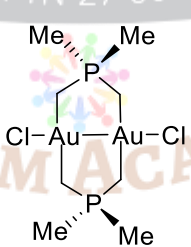
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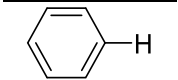
Q.21 – Q.70 MCQ, carry TWO marks each (for each wrong answer: -0.5).

You are required to Answer Maximum 35 Questions.

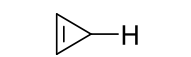
21. The HOMO (highest occupied molecular orbital) to LUMO (lowest unoccupied molecular orbital) electronic transition responsible for the observed colours of halogen molecules (gas) is
 (a) $\pi^* \rightarrow \sigma^*$ (b) $\pi \rightarrow \pi^*$ (c) $\sigma \rightarrow \sigma^*$ (d) $\pi \rightarrow \sigma^*$
22. In the hydrolysis of $\text{trans-}[\text{Co}(\text{en})_2\text{Cl}(\text{X})]^+$, if the leaving group is chloride, the formation of cis product is the least, when X is
 (a) NO_2^- (b) NCS^- (c) Cl^- (d) OH^-
23. The expected number of ^{19}F -NMR spectral lines, including satellites, for $[\text{XeF}_5]^-$ is [Abundance of ^{129}Xe ($I = \frac{1}{2}$) = 26%]
 (a) Two (b) Twenty one (c) Three (d) One
24. The expected H-H-H bond angle in $[\text{H}_3]^+$ is
 (a) 180° (b) 120° (c) 60° (d) 90°
25. The number of bridging ligand(s) and metal-metal bond(s) present in the complex $[\text{Ru}_2(\eta^5\text{-Cp})_2(\text{CO})_2(\text{Ph}_2\text{PCH}_2\text{PPh}_2)]$, respectively are ____ (obeys 18-electron rule)
 (a) 0 and 1 (b) 2 and 1 (c) 3 and 1 (d) 1 and 2
26. The oxidation state of gold in the following complex is
- 
- (a) 0 (b) 1 (c) 2 (d) 3
27. The rate of alkene coordination to $[\text{PtCl}_4]^{2-}$ is highest for
 (a) norbornene (b) ethylene (c) cyclohexene (d) 1-butene
28. The nephelauxetic parameter ' β ' is highest for
 (a) Br^- (b) Cl^- (c) CN^- (d) F^-
29. The ${}^2\text{E}_g \leftarrow {}^4\text{A}_{2g}$ transition in the electronic spectrum of $[\text{Cr}(\text{NH}_3)_6]^{3+}$ occurs nearly at
 (a) 650 nm (b) 450 nm (c) 350 nm (d) 200 nm
30. In the catalytic hydration of CO_2 by carbonic anhydrase, CO_2 first interacts with
 (a) OH group of the active site of the enzyme and then with zinc



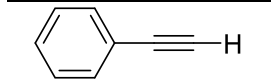
- (b) H_2O of the active site of the enzyme and then with zinc
 (c) Zinc of the active site of the enzyme and then with OH group
 (d) Zinc of the active site of the enzyme and then with H_2O
31. For the reaction; $\text{HX}_{(\text{aq})} + \text{H}_2\text{O}_{(\text{l})} \rightleftharpoons \text{H}_3\text{O}^+_{(\text{aq})} + \text{X}^-_{(\text{aq})}$, The highest value of $[\text{X}^-]_{(\text{aq})}$, when X^- is
 (a) OCl^- (b) F^- (c) Cl^- (d) NO_2^-
32. The correct statement for d.c. polarography is
 (a) $E_{1/2}$ is concentration dependent
 (b) Dropping mercury electrode is a macro electrode
 (c) Limiting current is equal to diffusion current
 (d) A large excess of supporting electrolyte eliminates migration current
33. Saturation factor in neutron activation analysis is,
 (A = induced radioactivity; ϕ = neutron flux; σ = effective nuclear cross section; N = no of target atoms; λ = decay constant)
 (a) $\frac{A}{\phi\sigma N}$ (b) $\frac{\phi\sigma NA}{\lambda}$ (c) $\frac{\lambda}{A\phi\sigma N}$ (d) $\frac{\phi\sigma N}{A}$
34. The primary analytical method is (not using a reference)
 (a) Inductively coupled plasma emission spectrometry
 (b) Energy dispersive X-ray fluorescence spectrometry
 (c) Anodic stripping voltammetry
 (d) Isotopic dilution mass spectrometry
35. The number of inorganic sulphur (or sulphide) atoms present in the metalloprotein active sites of rubredoxin, 2-iron ferredoxin and 4-iron ferredoxin, respectively, are
 (a) 0, 2 and 4 (b) 2, 4 and 3 (c) 0, 4 and 2 (d) 0, 2 and 3
36. The metal iodide with metallic lustre and high electrical conductivity is
 (a) NaI (b) CdI_2 (c) LaI_2 (d) BiI_3
37. The correct order of the bond dissociation energies for the indicated C–H bond in following compounds is
- X**



Y

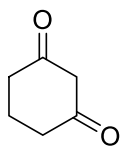


Z

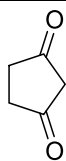

- (a) $Z > Y > X$ (b) $X > Y > Z$ (c) $X > Z > Y$ (d) $Z > X > Y$
38. The correct order of the acidity for the following compounds is



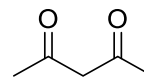
P



Q

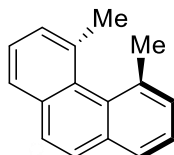


R



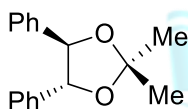
- (a) $Q > R > P$ (b) $R > Q > P$ (c) $Q > P > R$ (d) $R > P > Q$

39. The **correct statement** about the following compound is



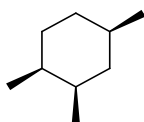
- (a) Compound is chiral and has P configuration
 (b) Compound is chiral and has M configuration
 (c) Compound is achiral as it possesses C_2 -axis of symmetry
 (d) Compound is achiral as it possesses plane of symmetry

40. **Methyl groups** in the following compound are



- (a) Homotopic (b) Diastereotopic
 (c) Enantiotopic (d) Constitutionally heterotopic

41. Among the structures given below, the **most stable conformation** for the given compound is



(a)



(b)



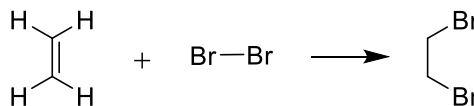
(c)



(d)



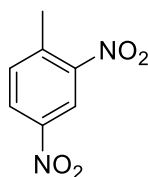
42. **Molecular orbital interactions** involved in the **first step** of the following reaction is



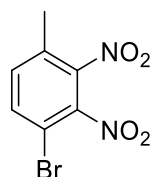
- (a) $\pi_{C=C} \rightarrow \sigma_{Br-Br}^*$ (b) $n_{Br} \rightarrow \sigma_{C-Br}^*$ (c) $\pi_{C=C} \rightarrow \sigma_{Br-Br}$ (d) $n_{Br} \rightarrow \pi_{C=C}$

43. The **major product** formed in the **dinitration** of **4-bromotoluene** is

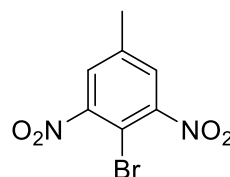
(a)



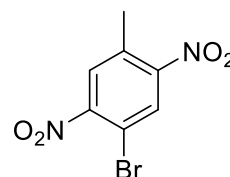
(b)



(c)

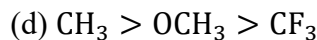
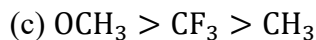
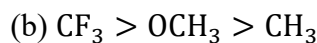
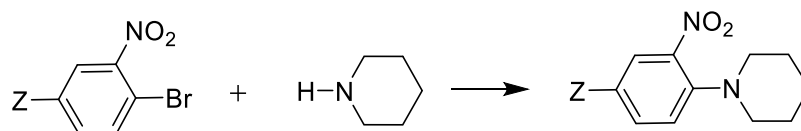


(d)

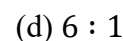
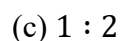
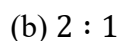
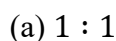


44. The **correct order** of the rate constants for the following series of reactions is,

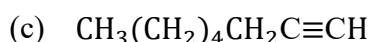
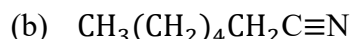




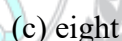
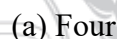
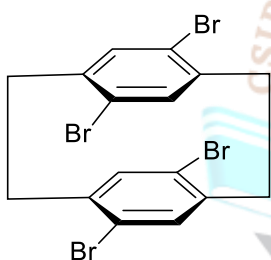
45. $^1\text{H-NMR}$ spectrum of a mixture of benzene and acetonitrile shows two singlets of equal integration. The molar ratio of benzene : acetonitrile is



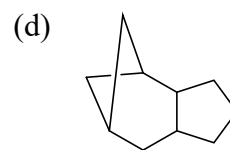
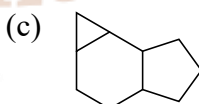
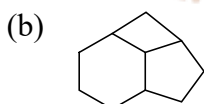
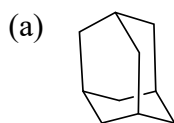
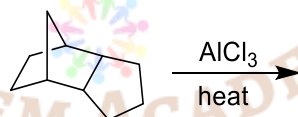
46. The compound which shows IR frequencies at both 3314 and 2126 cm^{-1} is,



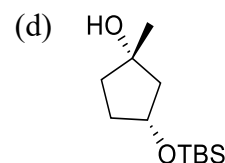
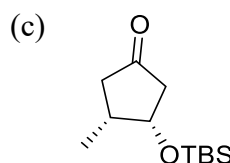
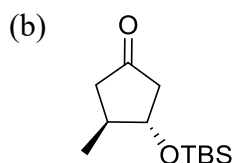
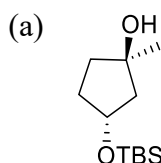
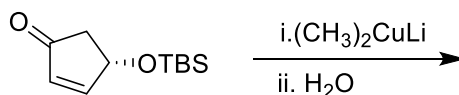
47. Number of signals present in the proton decoupled $^{13}\text{C-NMR}$ spectrum of the given compound is



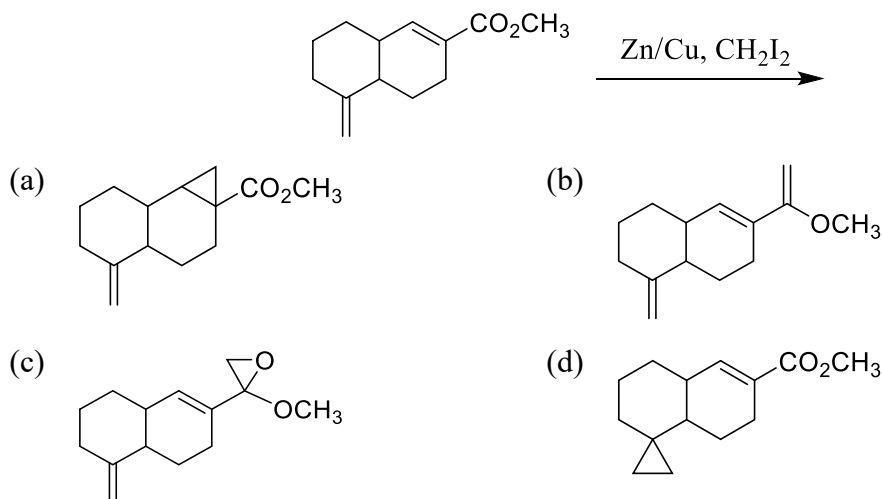
48. The most stable product formed in the following reaction is



49. The major product in the following reaction is



50. The major product formed in the following reaction is



51. Correct characteristics of the functional groups of **adenine** in **DNA base pair** are
- (a) N(3) is a hydrogen bond acceptor and C(6)NH₂ is a hydrogen bond donor
 (b) N(1) is a hydrogen bond acceptor and C(6)NH₂ is a hydrogen bond donor
 (c) Both N(3) and C(6)NH₂ are hydrogen bond acceptors
 (d) Both N(1) and C(6)NH₂ are hydrogen bond acceptors
52. ¹H-NMR spectrum of an organic compound recorded on a **500 MHz** spectrometer showed a **quartet** with a line positions at **1759, 1753, 1747, 1741 Hz**. Chemical shift (δ) and coupling constant (Hz) of the quartet are
- (a) 3.5 ppm, 6 Hz (b) 3.5 ppm, 12 Hz (c) 3.6 ppm, 6 Hz (d) 3.6 ppm, 12 Hz
53. The weight of the configuration with **two up and three down spins** in a system with **five spin $\frac{1}{2}$ particles** is
- (a) 120 (b) 60 (c) 20 (d) 10
54. For a reaction with a **activation energy** of **49.8 kJ mol⁻¹**, the **ratio of the rate constants** at **600 K** and **300 K**, (k_{600}/k_{300}), is approximately
- ($R = 8.3 \text{ J mol}^{-1} \text{ K}^{-1}$)
- (a) $\ln(10)$ (b) 10 (c) $10 + e$ (d) e^{10}
55. Covariance is defined by the relation **Cov(x,y) = $\langle xy \rangle - \langle x \rangle \langle y \rangle$** . Given the arbitrary constants A, B and C, Cov(x,y) will be **zero** only when
- (a) $y = Ax^2$ (b) $y = Ax^2 + B$ (c) $y = Ax + B$ (d) $y = Ax^2 + Bx + C$
56. Each void in a **two dimensional hexagonal close-packed layer** of circles is surrounded by
- (a) six circles (b) three circles (c) four circles (d) twelve circles
57. The **ionic mobilities** of **NH₄⁺** and **HCO₃⁻** are **$6 \times 10^{-4} \text{ V}^{-1} \text{ s}^{-1}$** and **$5 \times 10^{-4} \text{ V}^{-1} \text{ s}^{-1}$** ,



- respectively. The transport numbers of NH_4^+ and HCO_3^- are respectively.
- (a) 0.545 and 0.455 (b) 0.455 and 0.545 (c) 0.090 and 0.910 (d) 0.910 and 0.090
58. The ionic strength of a solution containing 0.008 M AlCl_3 and 0.005 M KCl is
- (a) 0.134 M (b) 0.053 M (c) 0.106 M (d) 0.086 M
59. The correct normalized wavefunction for one of the sp^2 hybrid orbitals is
- (a) $\frac{1}{3}\phi_{2s} + \frac{1}{3}\phi_{2p_x} + \frac{1}{3}\phi_{2p_y}$ (b) $\frac{1}{\sqrt{3}}\phi_{2s} + \frac{2}{\sqrt{3}}\phi_{2p_x} + \frac{1}{\sqrt{6}}\phi_{2p_y}$
 (c) $\frac{1}{\sqrt{3}}\phi_{2s} + \frac{1}{\sqrt{2}}\phi_{2p_x} + \frac{1}{\sqrt{6}}\phi_{2p_y}$ (d) $\frac{1}{\sqrt{3}}\phi_{2s} + \frac{1}{2\sqrt{3}}\phi_{2p_x} + \frac{1}{\sqrt{6}}\phi_{2p_y}$
60. The correct statement in the context of NMR spectroscopy is
- (a) Static magnetic field is used to induce transition between the spin states
 (b) Magnetization vector is perpendicular to the applied Static magnetic field
 (c) Static magnetic field is used to create population difference between the spin states
 (d) Static magnetic field induces spin-spin coupling
61. The parameter which always decreases during a spontaneous process at constant S and V, is
- (a) U (b) H (c) C_p (d) q
62. Triple point pressure of the substances P, Q, R and S are 0.2, 0.5, 0.8 and 1.2 bar, respectively. The substance which sublimates under standard conditions on increasing temperature is
- (a) P (b) Q (c) R (d) S
63. According to the transition state theory, the plot with slope equal to $\frac{-\Delta H^\ddagger}{R}$ is
- (a) $\ln k$ vs. T (b) $\ln\left(\frac{k}{T}\right)$ vs. T (c) $\ln\left(\frac{k}{T}\right)$ vs. $\frac{1}{T}$ (d) $\ln k$ vs. $\frac{1}{T}$
64. The transition that belongs to the Lyman series in the hydrogen-atom spectrum is
- (a) $1s \leftarrow 4s$ (b) $1s \leftarrow 4p$ (c) $2s \leftarrow 4s$ (d) $2s \leftarrow 4p$
65. The molecule that possesses S_4 symmetry element is
- (a) ethylene (b) allene (c) benzene (d) 1,3-butadiene
66. Vibrations of diatomic molecules are usually modelled by a harmonic potential. If the potential is given by x^2 , the correct statement is
- (a) Force is $2x$ and force constant is 2 (b) Force is $-2x$ and force constant is 2
 (c) Force is $2x$ and force constant is -1 (d) Force is $-2x$ and force constant is -1
67. When 1×10^{-5} g of a fatty acid ($M = 602.3$ g/mol) was placed on water as a surface film, a monomolecular layer of area 100 cm^2 was formed on compression.



- The cross-sectional area (in $\text{\AA}^2$) of the acid molecule is
 (a) 50 (b) 100 (c) 150 (d) 200
68. Mark-Houwink equation ($[\eta] = KM^a$) is used for the determination of
 (a) number-average molar mass (b) weight-average molar mass
 (c) viscosity-average molar mass (d) z-average molar mass
69. Many properties of nanoparticles are significantly different that the corresponding bulk material due to
 (a) Smaller band gap of nanoparticles compared to bulk
 (b) Higher heterogeneity of the nanoparticles solutions
 (c) Larger ratio of surface area to volume of the nanoparticles compared to the bulk
 (d) Smaller ratio of surface area to volume of the nanoparticles compared to the bulk
70. The correct match for the following is

Column-I		Column-II		
i)	camphor	P.	Structural protein	i
ii)	insulin	Q.	hormone	ii
iii)	keratin	R.	enzyme	iii
		S.	steroid	
		T.	Terpene	

- (a) P ; R ; T
 (b) T ; Q ; P
 (c) S ; R ; P
 (d) T ; Q ; S

Q.71 – Q.145 MCQ, carry FOUR marks each (for each wrong answer: -1).
You are required to Answer Maximum 25 Questions.

71. Consider the following statements for KC_8 :
 [P] It is paramagnetic
 [Q] It has eclipsed layer structure
 [R] Its electrical conductivity is greater than that of graphite
 The correct answer is
 (a) P and Q (b) P and R (c) Q and R (d) P, Q and R
72. Among the following, choose the correct products that are formed in the reaction of S_2Cl_2 with ammonia is CCl_4 :

P	Q	R	S
NH_4Cl	S_4N_4	S_8	$\text{S}_3\text{N}_3\text{Cl}_3$

 (a) P, Q and R (b) P, Q and S (c) Q, R and S (d) P, R and S
73. For $[\text{Ce}(\text{NO}_3)_4(\text{OPPh}_3)_2]$, from the following
 [P] Its aqueous solution is yellow-orange in colour



[Q] Coordination number of Ce is ten

[R] It shows metal to ligand charge transfer

[S] It is diamagnetic in nature

The correct answer is

- (a) P and Q (b) P and R (c) P, Q and S (d) Q, R and S

74. Consider the following statements, I and II :

I : $[\text{Rh}(\text{CO})_2\text{I}_2]^-$ catalytically converts CH_3I and CO to CH_3COI

II : $[\text{Rh}(\text{CO})_2\text{I}_2]^-$ is diamagnetic in nature

The correct statement from the following is

- (a) I and II are correct and II is an explanation of I
 (b) I and II are correct and II is not an explanation of I
 (c) I is correct and II is incorrect
 (d) I and II are incorrect

75. In a direct isotopic dilution method for determination of phosphate, 2 mg of $^{32}\text{PO}_4^{3-}$ (specific activity 3100 disintegration $\text{s}^{-1} \text{mg}^{-1}$) was added to 1 g of a sample solution. The 30 mg of phosphate isolated from it has an overall activity of 3000 disintegration s^{-1} . The % mass of PO_4^{3-} in the sample is

- (a) 30 (b) 6 (c) 9 (d) 15

76. Consider the following statements for $[\text{FeO}_4]^{4-}$,

[P] It is paramagnetic

[Q] It has T_d symmetry

[R] Adopts distorted square planar geometry

[S] Shows approximately D_{2d} symmetry

The correct answer is

- (a) P, Q and R (b) P, R and S (c) P and S (d) P and Q

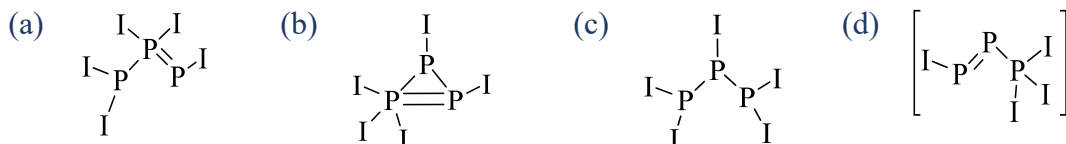
77. The geometry of $[\text{ReH}_9]^{2-}$ is

- (a) Monocapped square antiprism (b) Monocapped cube
 (c) tricapped trigonal prism (d) Heptagonal bipyramid

78. The reaction between PI_3 , PSCl_3 and zinc powder gives P_3I_5 as one of the products. The solution state ^{31}P -NMR spectrum of P_3I_5 shows a doublet ($\delta = 98$) and a triplet ($\delta = 102$). The correct structure of P_3I_5 is



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79. Some molecules and their **properties in liquid ammonia** are given in columns I and II respectively. Match column I with column II

	Column I	Column II						
(P)	Cl ₂	(i)	Weak acid					
(Q)	S ₈	(ii)	Strong acid	P	Q	R	S	
(R)	CH ₃ CO ₂ H	(iii)	Disproportionation	(a)	i	ii	iii	iv
(S)	Urea	(iv)	Solvolysis and	(b)	ii	iii	iv	i
			Disproportionation	(c)	iii	iv	i	ii
				(d)	iv	iii	ii	i

80. The **spectroscopic ground state term symbols** for the octahedral aqua complexes of **Mn(II), Cr(III) and Cu(II)**, respectively are

(a) ²H, ⁴F and ²D (b) ⁶S, ⁴F and ²D (c) ²H, ²H and ²D (d) ⁶S, ⁴F and ²P

81. From the following transformations

[P] Epoxidation of alkene

[Q] Diol dehydrase reaction

[R] Conversion of ribonucleotide-to-deoxyribonucleotide

[S] 1, 2-Carbon shift in organic substrates

Those promoted by **coenzyme B₁₂** are

(a) P and Q (b) Q, R and S (c) P, Q and S (d) P, Q and R

82. Match the items in column-I with the appropriate items in column-II

	Column-I	Column-II
(P)	Metallothioneins	(i) Cis-[Pd(NH ₃) ₂ Cl ₂]
(Q)	Plastocyanin	(ii) Cysteine rich protein
(R)	Ferritin	(iii) Electron transfer
(S)	Chemotherapy	(iv) Iron Transport
		(v) Iron Storage
		(vi) Carboplatin

The correct Answer is

P	Q	R	S
(a)	ii	iii	v ; iv



- (b) ii ; iii ; iv ; i
 (c) ii ; iii ; v ; vi
 (d) iii ; v ; vi ; ii

83. For OH^- catalysed $\text{S}_{\text{N}}1$ conjugate base mechanism of $[\text{Co}(\text{NH}_3)_5\text{Cl}]^{2+}$, the species obtained in the first step of the reaction is/are

- (a) $[\text{Co}(\text{NH}_3)_5\text{OH}]^{2+} + \text{Cl}^-$ (b) $[\text{Co}(\text{NH}_3)_4(\text{NH}_2)\text{Cl}]^+ + \text{H}_2\text{O}$
 (c) $[\text{Co}(\text{NH}_3)_4(\text{NH}_2)]^{2+} + \text{Cl}^-$ (d) $[\text{Co}(\text{NH}_3)_5\text{Cl}(\text{OH})]^+$ Only

84. Match the species in column-X with their properties in column-Y

Column-X		Column-Y	
[P]	Heme A	(i)	Oxo-bridged Mn_4 cluster
[Q]	Water splitting enzyme	(ii)	Tetragonal elongation
[R]	$[\text{Mn}(\text{H}_2\text{O})_6]^{2+}$	(iii)	Predominantly $\pi \rightarrow \pi^*$ electronic transitions
[S]	$[\text{Cr}(\text{H}_2\text{O})_6]^{2+}$	(iv)	d $\rightarrow$ d spin-forbidden transitions
		(v)	Tetragonal compression

The correct answer is

P	Q	R	S
(a)	iii ; i ; v ; ii		
(b)	iii ; i ; iv ; ii		
(c)	v ; iii ; iv ; ii		
(d)	iii ; i ; iv ; v		

85. According to isolobal analogy, the right set of fragments that might replace $\text{Co}(\text{CO})_3$ in $[\text{Co}_4(\text{CO})_{12}]$ is

- (a) CH, BH and $\text{Mn}(\text{CO})_5$ (b) P, CH and $\text{Ni}(\eta^5\text{-C}_5\text{H}_5)$
 (c) $\text{Fe}(\text{CO})_4$, CH_2 and SiCH_3 (d) BH, SiCH_3 and P

86. According to wade's rules, the correct structural types of $[\text{Co}(\eta^5\text{-C}_5\text{H}_5)\text{B}_4\text{H}_8]$ and $[\text{Mn}(\eta^2\text{-B}_3\text{H}_8)(\text{CO})_4]$ are

- (a) Closo and nido (b) Nido and arachno
 (c) Closo and arachno (d) nido and nido

87. The correct geometry of $[\text{Rh}_6\text{C}(\text{CO})_{15}]^{2-}$ is

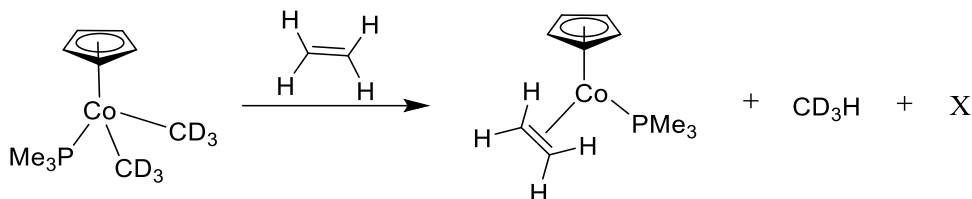
- (a) octahedron (b) Pentagonal pyramid
 (c) Trigonal prism (d) Monocapped square pyramid

88. The final product(s) of the reaction of arachno borane B_4H_{10} with NMe_3 is/are



- (a) $[\text{BH}_3 \cdot \text{NMe}_3]$ and $[\text{B}_3\text{H}_7 \cdot \text{NMe}_3]$ (b) $[\text{BH}_2(\text{NMe}_3)_2]^+ [\text{B}_3\text{H}_8]^-$
 (c) $[\text{B}_4\text{H}_{10} \cdot \text{NMe}_3]$ (d) $[\text{B}_4\text{H}_{10} \cdot \text{NMe}_3]$ and $[\text{BH}_2(\text{NMe}_3)_2]^+ [\text{B}_3\text{H}_8]^-$

89. **Product-X** in the following reaction is



- (a) $\text{CD}_2=\text{CD}_2$ (b) $\text{D}_3\text{C}-\text{CD}_3$ (c) $\text{CH}_2=\text{CD}_3$ (d) $\text{CH}_2=\text{CD}_2$

90. Treatment of $\text{Fe}(\text{CO})_5$ with 1,3-butadiene gives X that shows two signals in its ^1H -NMR spectrum. X on treatment with HCl yields Y which shows four signals in its ^1H -NMR spectrum. The compound Y is



91. In the following redox reaction with an equilibrium constant $(K) = 2.0 \times 10^8$,



The self-exchange rates for oxidant and reductant are $5.0 \text{ M}^{-1}\text{s}^{-1}$ and $4.0 \times 10^3 \text{ M}^{-1}\text{s}^{-1}$, respectively. The approximate rate constant ($\text{M}^{-1}\text{s}^{-1}$) for the reaction is

- (a) 3.16×10^6 (b) 2.0×10^6 (c) 6.32×10^6 (d) 3.16×10^4

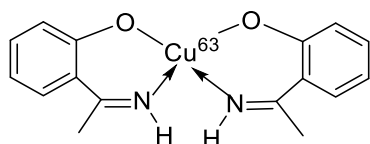
92. The correct statement for a Fischer carbene complex is

- (a) The carbene carbon is electrophilic in nature
 (b) Metal exists in high oxidation state
 (c) Metal fragment and carbene are in the triplet states
 (d) CO ligands destabilize the complex

93. The acidic solution containing trimethylamine (X), dimethylamine (Y) and methylamine (Z) ($\text{P}K_a$ of cations 9.8, 10.8 and 10.6, respectively) was loaded on a cation exchange column. The order of their elution with a gradient of increasing $\text{pH} > 7$ is

- (a) $X < Z < Y$ (b) $Y < Z < X$ (c) $Y < X < Z$ (d) $Z < Y < X$

94. For complex-P, deuteration of NH proton does not alter the EPR spectrum. The number of hyperfine lines expected in the EPR [$I(^{63}\text{Cu}) = 3/2$] spectrum of P is,

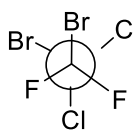
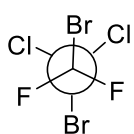


- (a) 20 (b) 12
(c) 60 (d) 36

95. The numbers of triangular faces in square antiprism, icosahedron and tricapped trigonal prism (Capped on square faces), respectively, are

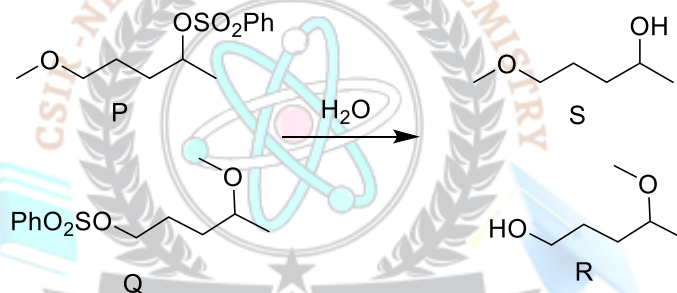
- (a) 8, 20 and 14 (b) 8, 20 and 12 (c) 10, 12 and 14 (d) 10, 12 and 12

96. Number of lines in the ^{19}F -NMR spectrum of $\text{F}_2\text{C}(\text{Br})-\text{C}(\text{Br})\text{Cl}_2$ at -120°C assuming it a mixture of static conformations given below, are



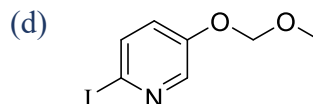
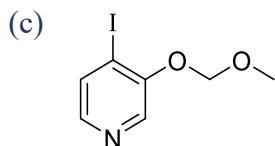
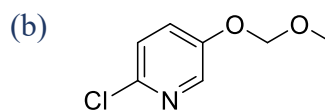
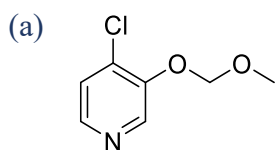
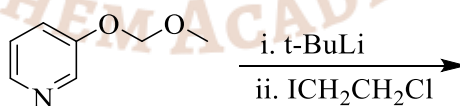
- (a) one (b) two
(c) four (d) five

97. The correct statement for the reactants P, Q to give products R, S is

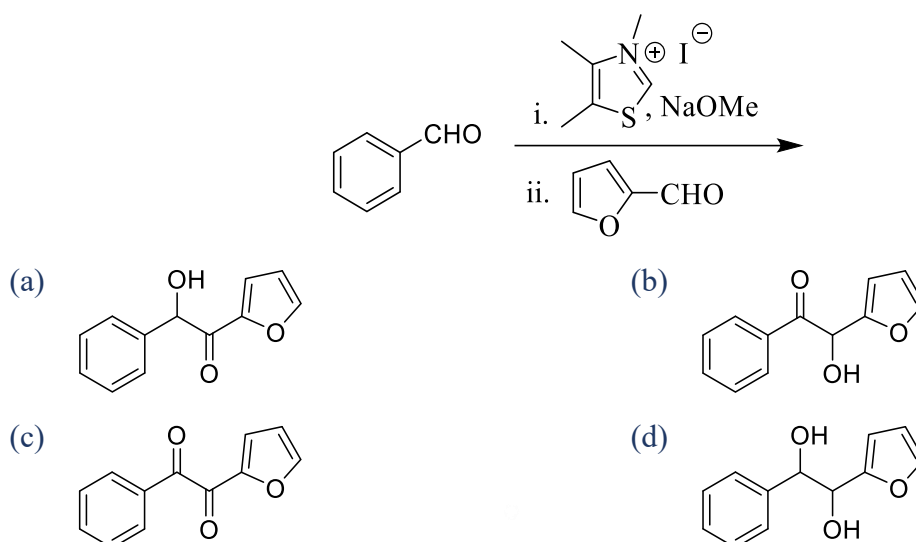


- (a) P gives R and Q gives S (b) P gives S and Q gives R
(c) P and Q give identical amounts of R and S (d) P and Q give S

98. The major product formed in the following reaction is

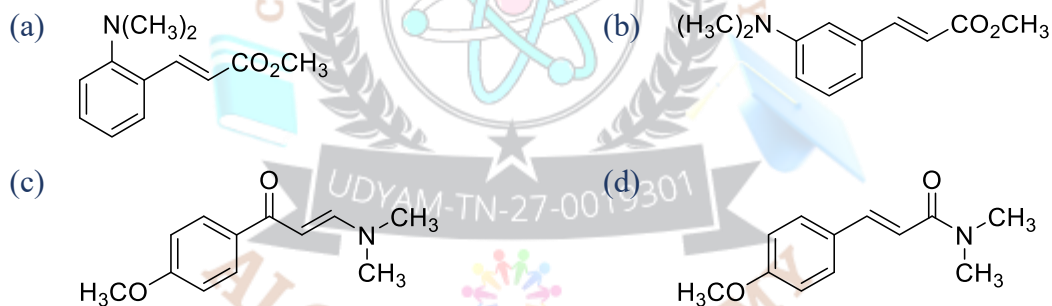


99. The major product formed in the following reaction is

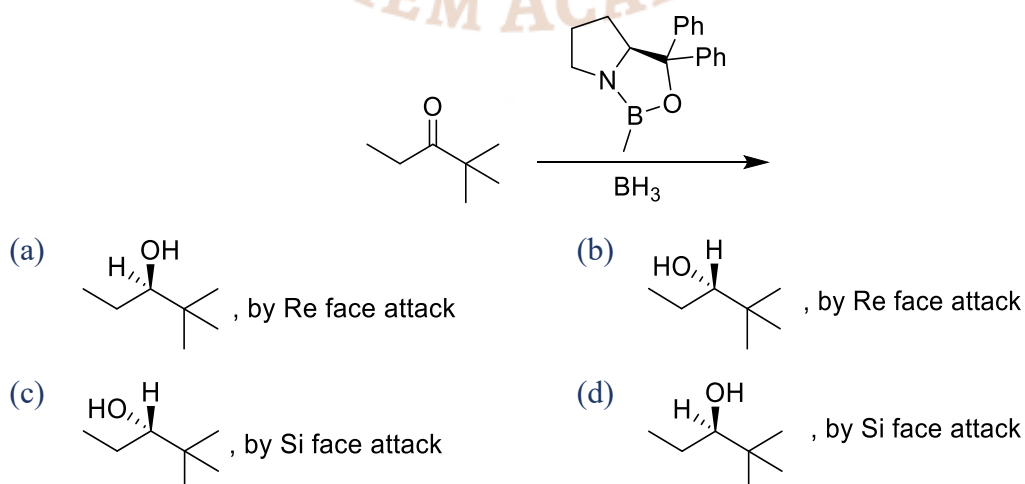


100. The compound that exhibits following spectral data is

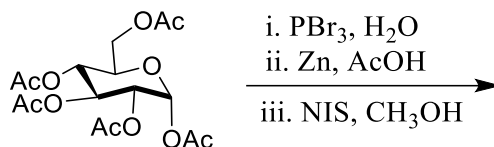
$^1\text{H-NMR}$: 8.0 (d, $J = 12.3$ Hz, 1H), 7.7 (d, $J = 8.0$ Hz, 2H), 3.8 (s, 3H),
 (δ in ppm) 6.8 (d, $J = 8.0$ Hz, 2H), 5.8 (d, $J = 12.3$ Hz, 1H), 3.0 (s, 6H)



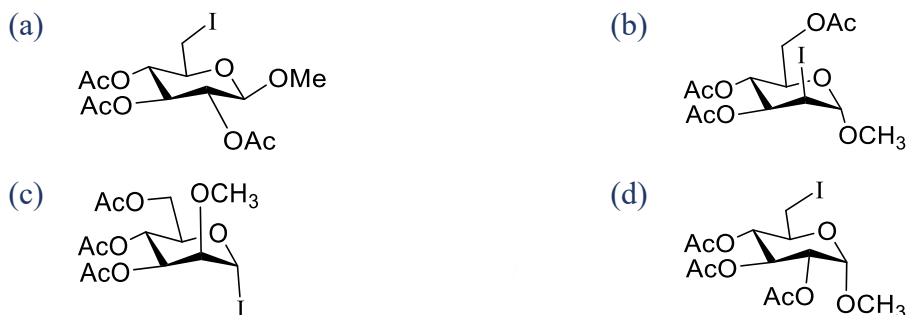
101. The major product formed in the following reaction is



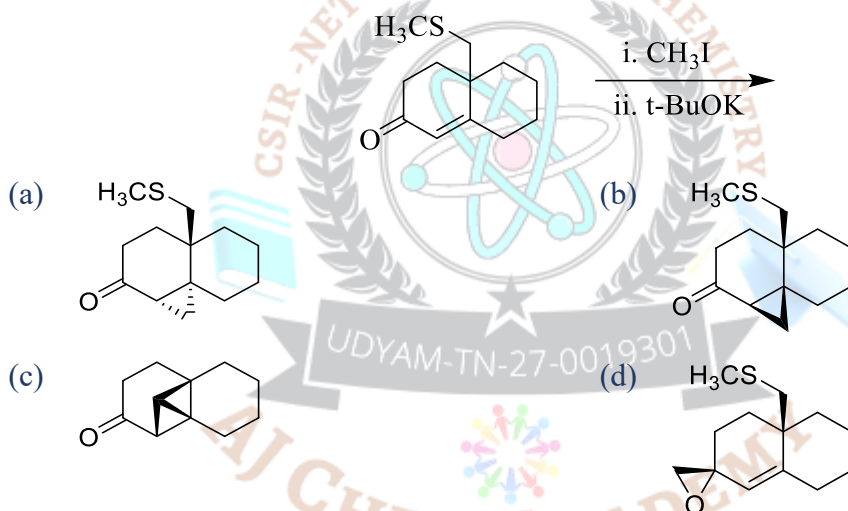
102. The major product formed in the following reaction is



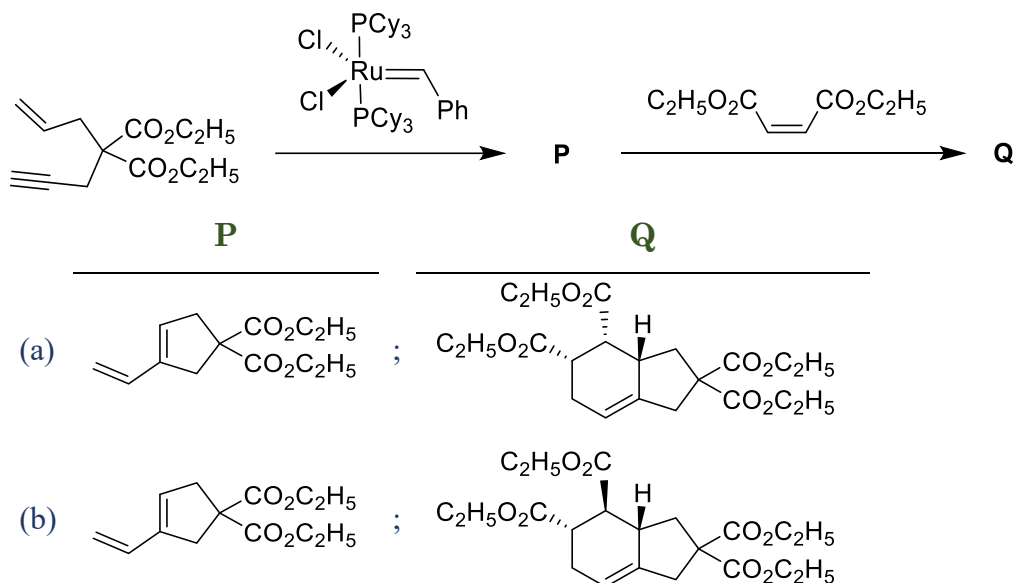
NIS = N-Iodosuccinimide

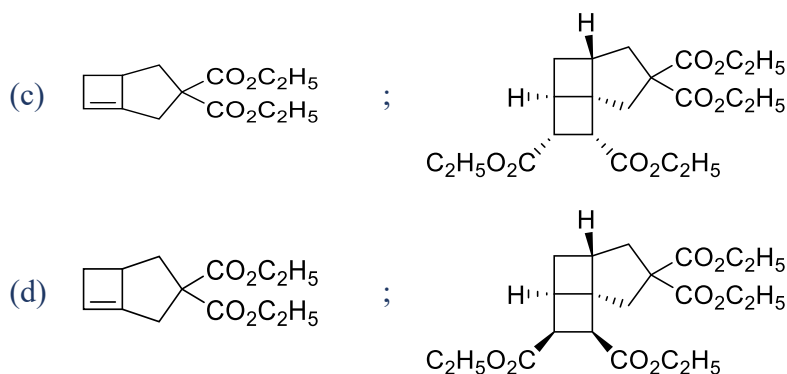


103. The major product formed in the following reaction is

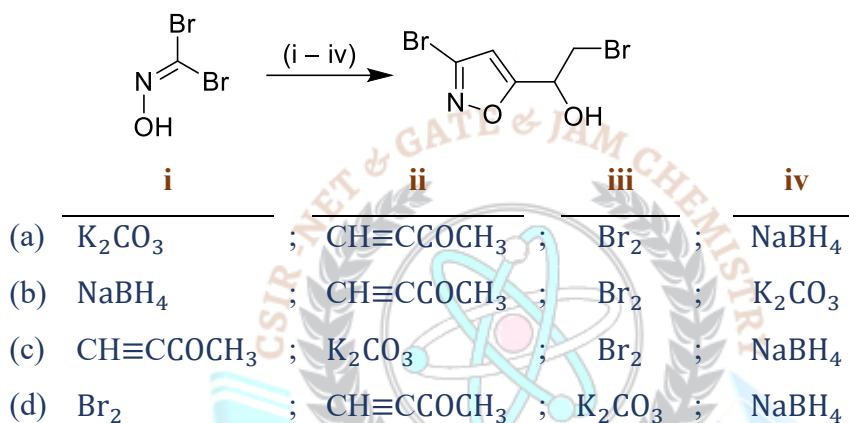


104. The major product formed in the following reaction is

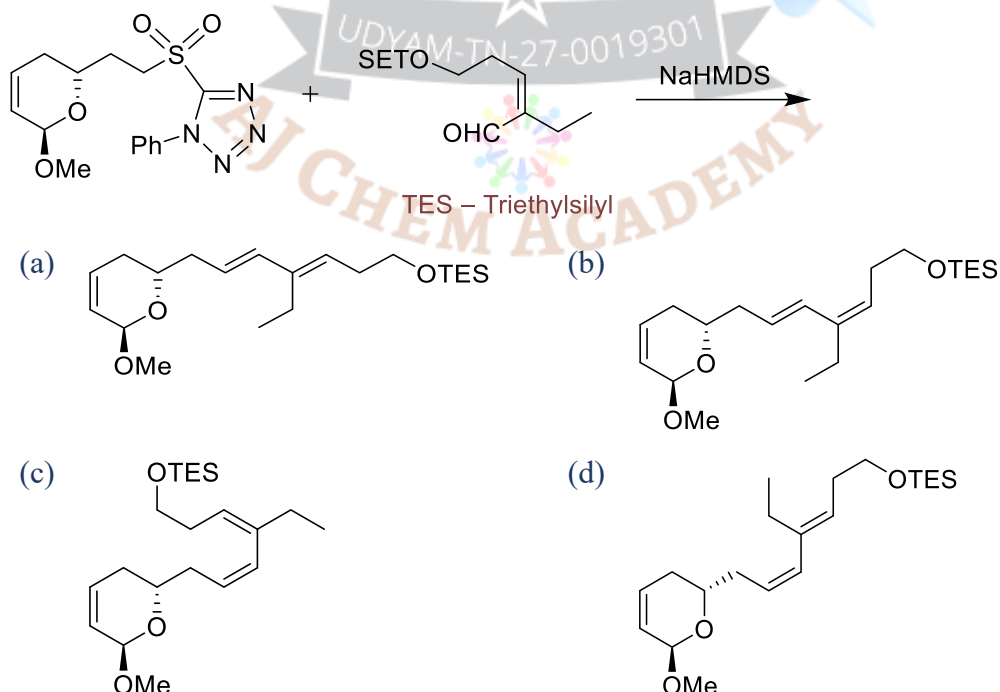




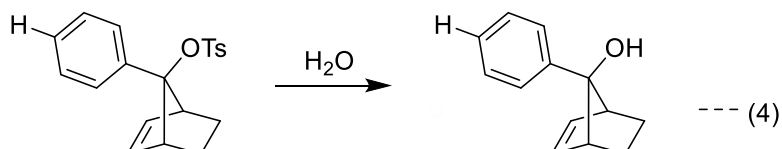
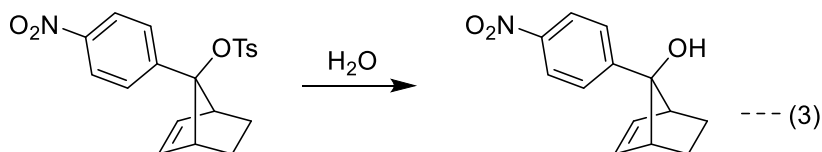
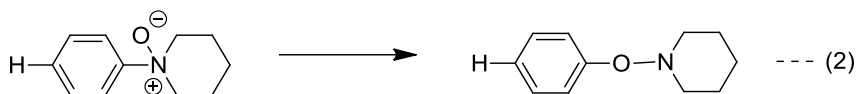
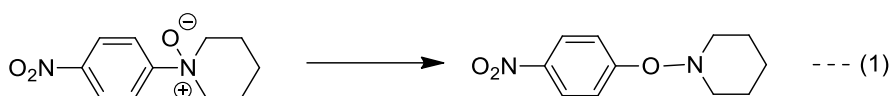
105. Correct sequence of reagents for the following conversion is



106. The major product formed in the following reaction is



107. For the four reactions given below, the rates of the reactions will vary as



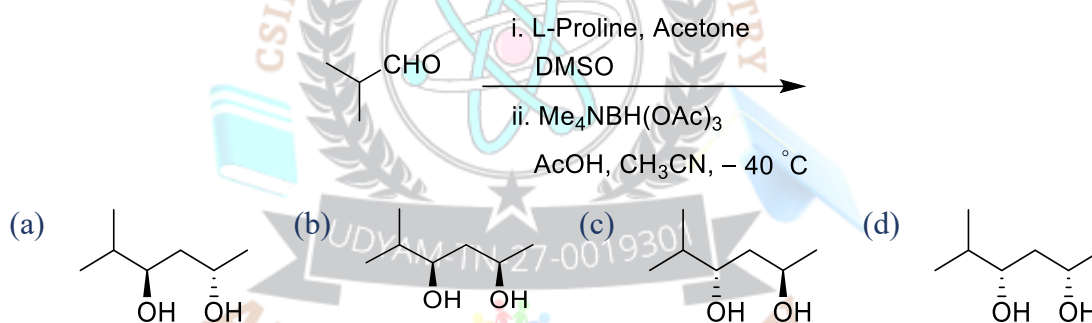
(a) 1 > 2 and 3 > 4

(b) 2 > 1 and 3 > 4

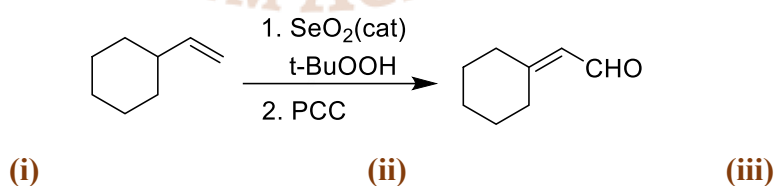
(c) 2 > 1 and 4 > 3

(d) 1 > 2 and 4 > 3

108. The major product formed in the following reaction is

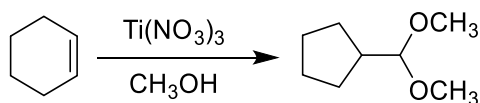


109. The correct sequence of pericyclic reactions involved in the following transformation is



(a)	ene reaction ; [2,3]-sigmatropic shift ; [3,3]-sigmatropic shift
(b)	ene reaction ; [3,3]-sigmatropic shift ; [1,3]-sigmatropic shift
(c)	[2,3]-sigmatropic shift ; ene reaction ; [1,3]-sigmatropic shift
(d)	[1,3]-sigmatropic shift ; [2,3]-sigmatropic shift ; [3,3]-sigmatropic shift

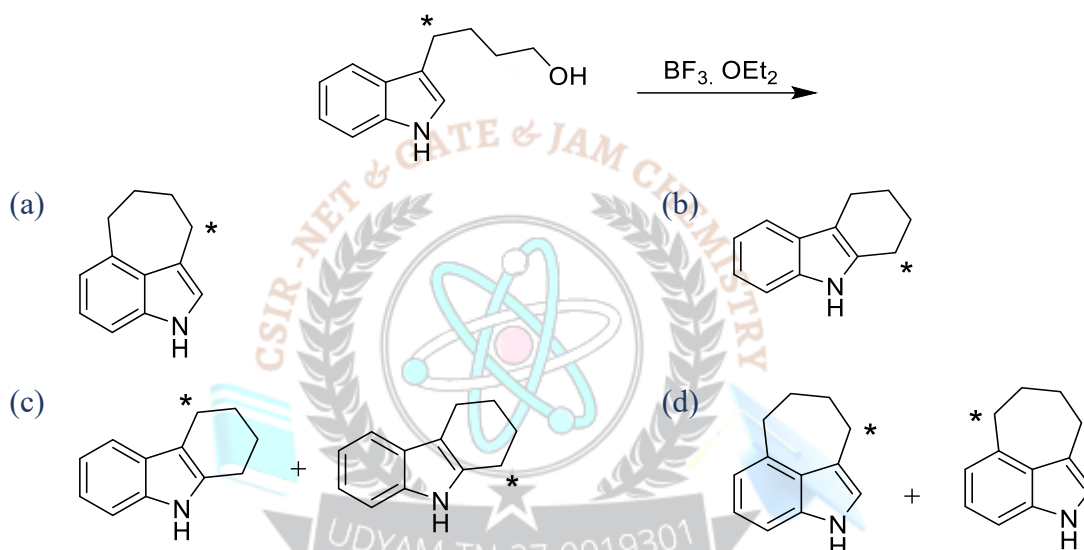
110. The intermediate that leads to the product in the following transformation is



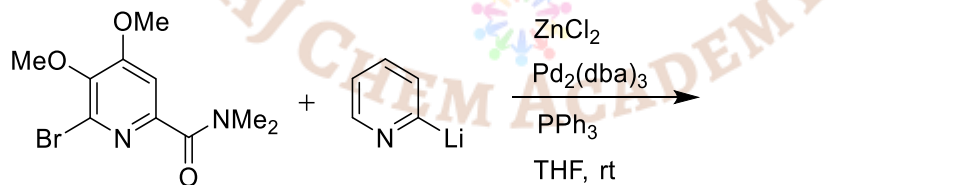


111. Product(s) of the following reaction is (are)

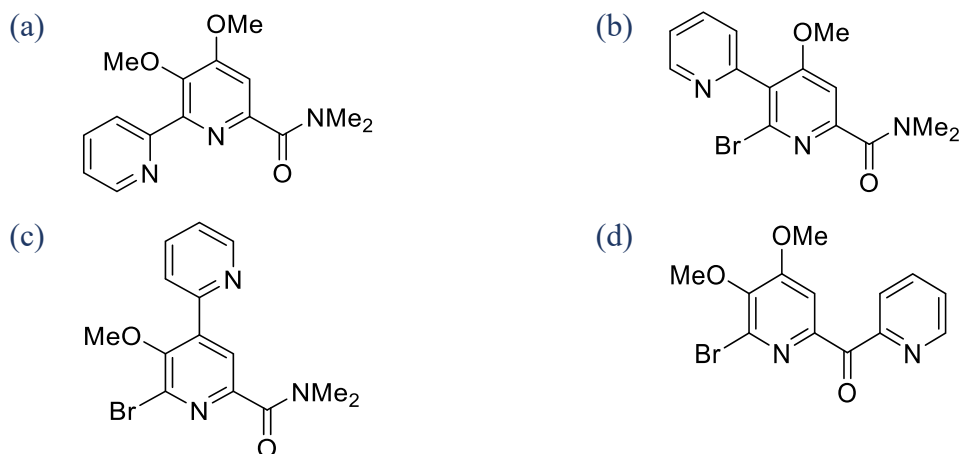
[*-indicates isotopically labelled carbon]



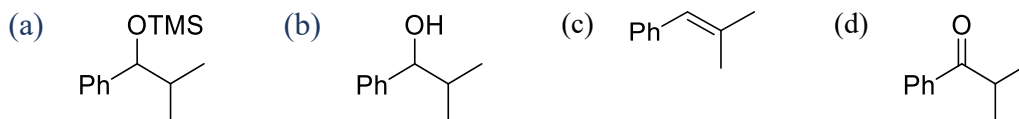
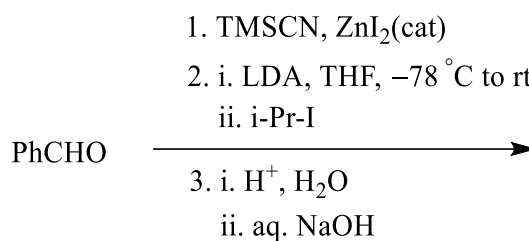
112. The major product formed in the following reaction is



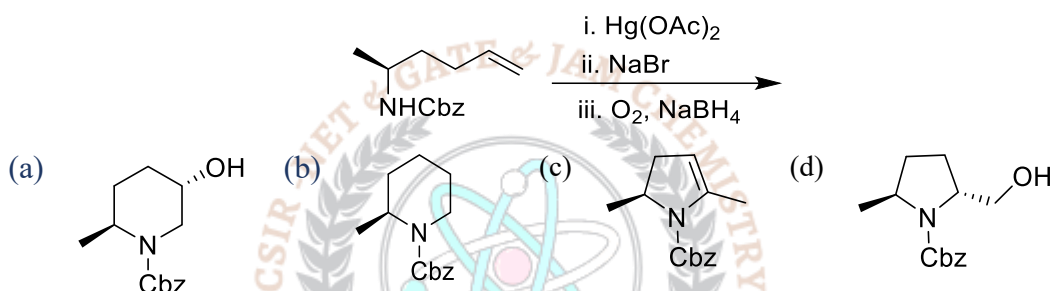
dba = Dibenzylidene acetone



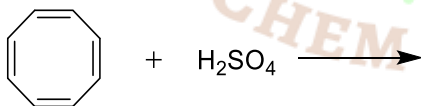
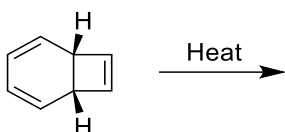
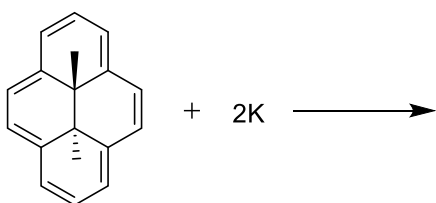
113. The **major product** formed in the following reaction is



114. The **major product** formed in the following reaction is



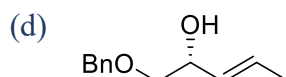
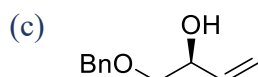
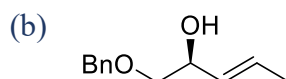
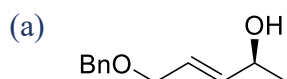
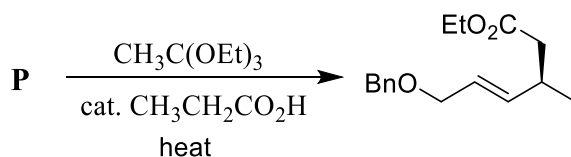
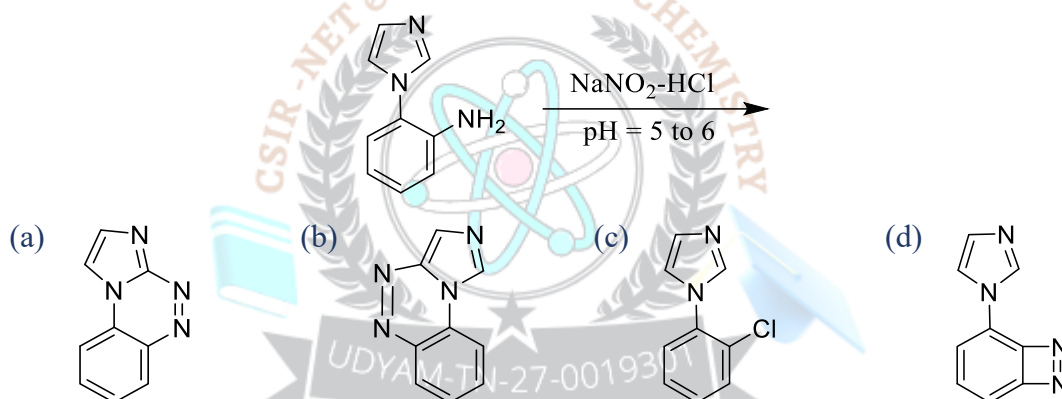
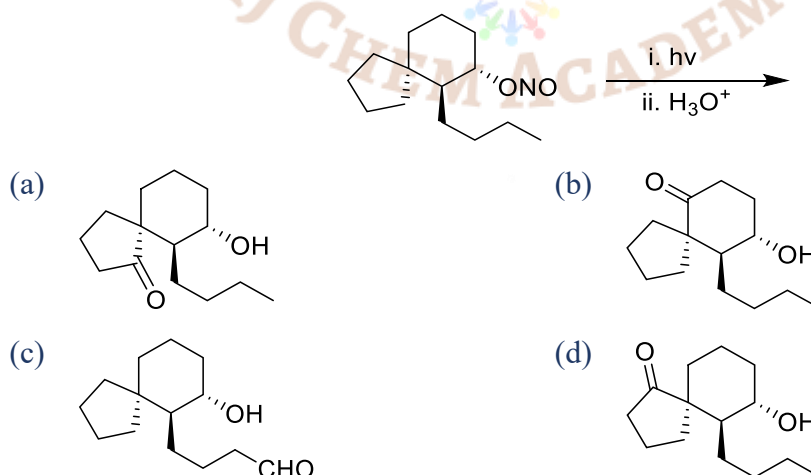
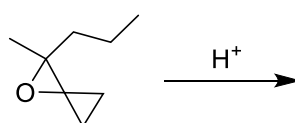
115. Correct match for the products of the reactions in column-I with the properties in column-II is

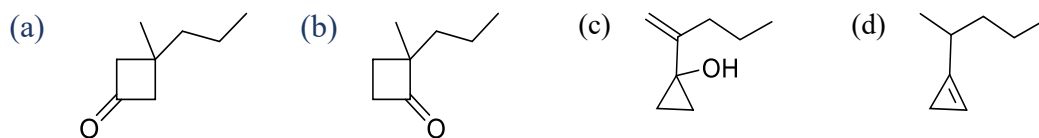
	Column-I	Column-II
i)		P) Aromatic
ii)		Q) Antiaromatic
iii)		R) Non-aromatic
iv)		S) Homoaromatic

	i	ii	iii	iv
(a)	P	S	R	Q
(b)	P	R	Q	S

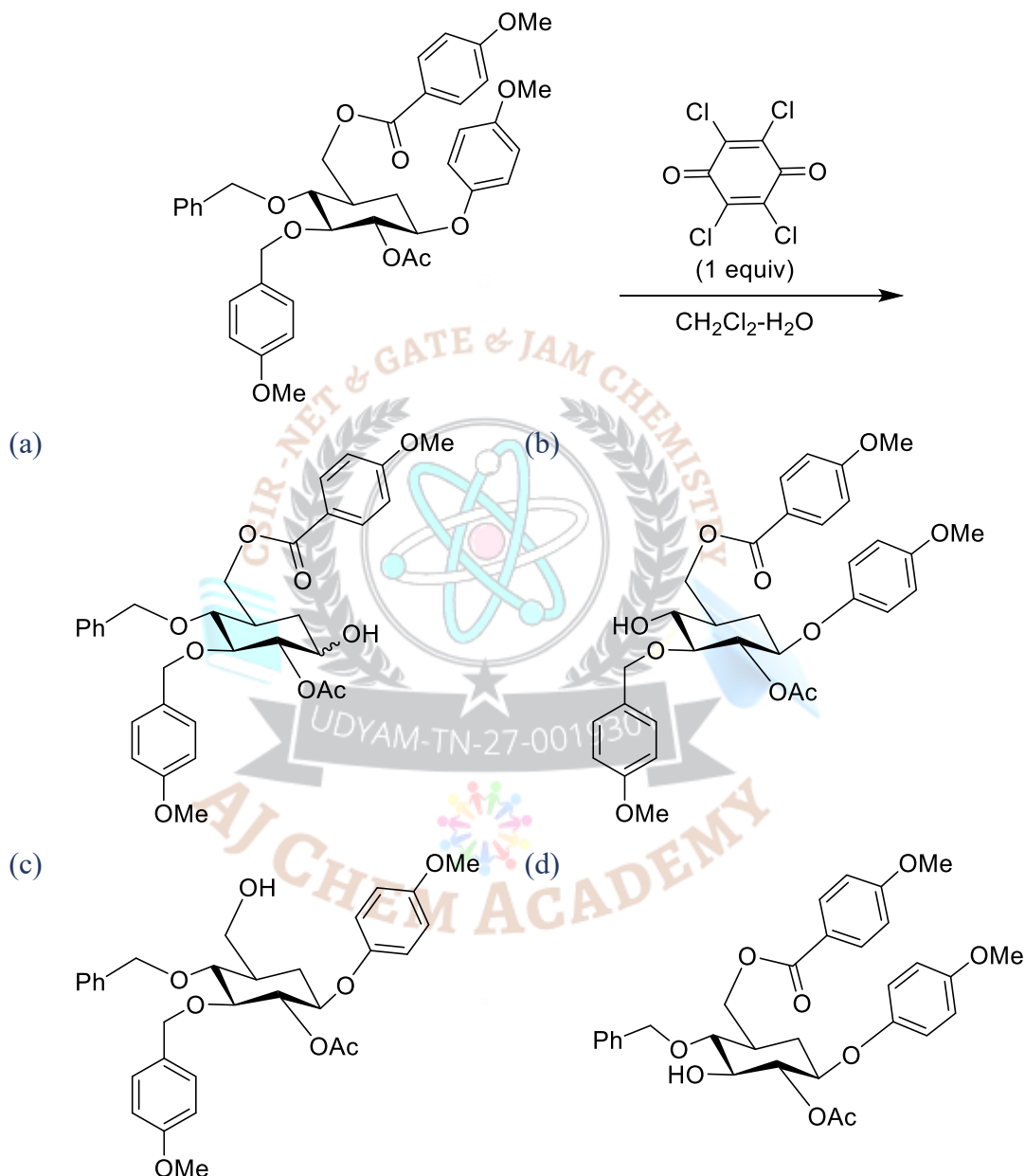
(c) Q ; R ; S ; P

(d) S ; Q ; R ; P

116. The correct **starting compound-P** in the following reaction is117. The **major product** formed in the following reaction is118. The **major product** formed in the following reaction is119. The **major product** formed in the following reaction is



120. The **major product** formed in the following reaction is



121. A constant of motion of **hydrogen atom** in the presence of **spin-orbit coupling** is

- (a) ℓ (b) s (c) $\ell + s$ (d) $\ell - s$

122. The **orbital degeneracy** of the level of a **one-electron atomic system** with **$Z = 5$** and energy $\approx -13.6\text{eV}$, is

- (a) 1 (b) 5 (c) 25 (d) 36

123. If we write a **normalized wavefunction** ψ as $\psi = \hat{A}\phi$, then ϕ is also normalized



When

(a) $\hat{A}$ is hermitian (b) $\hat{A}$ is anti-hermitian (c) $\hat{A}$ is unitary (d) $\hat{A}$ is any linear operator

124. The ground state of a certain system with energy E_0 is subjected to a perturbation V , yielding a first order correction E_1 . If E_0 is the true ground-state energy of the perturbed system, the inequality that always holds is

(a) $E_1 \geq 0$ (b) $E_1 \geq E_0$ (c) $E_0 + E_1 \leq E_0$ (d) $E_0 + E_1 \geq E_0$

125. The spatial part of an excited state $^3\Sigma_u^+$ of hydrogen molecule is proportional to $[1\sigma_g(1)1\sigma_u(2) - 1\sigma_g(2)1\sigma_u(1)]$. Using LCAO-MO expansion of $1\sigma_g$ and $1\sigma_u$ in terms of 1s-atomic orbitals, one can infer that this wavefunction has

(a) only ionic parts (b) only covalent parts
(c) both ionic and covalent parts (d) neither ionic nor covalent parts

126. The highest molecular orbitals for an excited electronic configuration of the oxygen molecule are $[1\pi_g]^1[3\sigma_u]^1$. A possible molecular term symbol for oxygen with this electronic configuration is

(a) $^1\pi$ (b) $^3\Sigma$ (c) $^1\Delta$ (d) $^1\Sigma$

127. For H_2O molecule, the electronic transition from the ground state to an excited state of B_1 symmetry is

C_{2v}	E	C_2	σ_v	σ'_v		
A_1	1	1	1	1	z, z^2, x^2, y^2	(a) not allowed
A_2	1	1	-1	-1	xy	(b) allowed with x polarisation
B_1	1	-1	1	-1	x, xz	(c) allowed with y polarisation
B_2	1	-1	-1	1	y, yz	(d) allowed with z polarisation

128. The pair of symmetry point groups that are associated with Only polar molecules is

(a) $C_{2v}, D_{\infty h}$ (b) C_{3v}, C_{2h} (c) D_{2h}, T_d (d) $C_{2v}, C_{\infty v}$

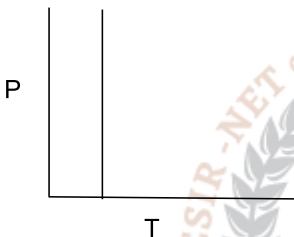
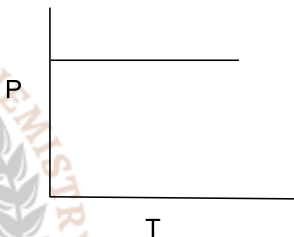
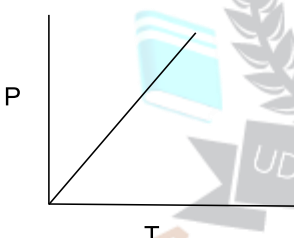
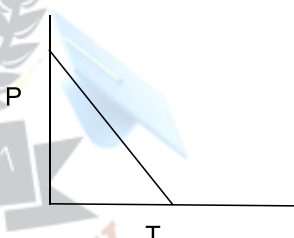
129. The rotational constant and the fundamental vibrational frequency of HBr are, respectively, 10 cm^{-1} and 2000 cm^{-1} . The corresponding values for DBr approximately are

(a) 20 cm^{-1} and 2000 cm^{-1} (b) 10 cm^{-1} and 1410 cm^{-1}
(c) 5 cm^{-1} and 2000 cm^{-1} (d) 5 cm^{-1} and 1410 cm^{-1}

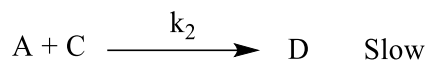
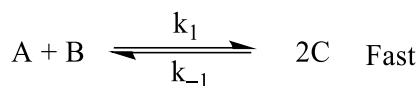
130. Among the following, both microwave and rotational Raman active molecule is

(a) CH_4 (b) N_2O (c) C_2H_4 (d) CO_2

131. In a 200 MHz NMR spectrometer, a molecule shows two doublets separated by

- 2 ppm.** The observed coupling constant is **10 Hz**. The separation between these two signals and the coupling constant in a **600 MHz** spectrometer will be, respectively
- (a) 600 Hz and 30 Hz (b) 1200 Hz and 30 Hz
(c) 600 Hz and 10 Hz (d) 1200 Hz and 10 Hz
132. The equation of state for one mole of a gas is given by $P(V - b) = RT$, where **b** and **R** are constants. The value of $\left(\frac{\partial H}{\partial p}\right)_T$ is
- (a) $V - b$ (b) b (c) 0 (d) $\frac{RT}{P} + b$
133. The volume change in a phase transition is **zero**. From this, we may infer that the **phase boundary** is represented by
- (a)  (b) 
(c)  (d) 
134. The partial derivative $\left(\frac{\partial T}{\partial V}\right)_P$ is equal to,
- (a) $-\left(\frac{\partial P}{\partial S}\right)_T$ (b) $-\left(\frac{\partial P}{\partial S}\right)_V$ (c) $-\left(\frac{\partial P}{\partial S}\right)_n$ (d) $-\left(\frac{\partial P}{\partial S}\right)_H$
135. If the energies of a bare proton aligned along and against an external static magnetic field (B_z) are $-\hbar\gamma B_z/2$ and $+\hbar\gamma B_z/2$. Respectively, then the **ratio of Probabilities of finding the proton along and against the magnetic field** is
- (a) $e^{-\hbar\gamma B_z/4k_B T}$ (b) $e^{-\hbar\gamma B_z/2k_B T}$ (c) $e^{\hbar\gamma B_z/2k_B T}$ (d) $e^{\hbar\gamma B_z/k_B T}$
136. **Partition function of a one-dimensional oscillator** having equispaced energy levels with energy spacing equal to $k_B T$ zero ground state energy is
- (a) e (b) $1/(e - 1)$ (c) $e/(e - 1)$ (d) $1/(e + 1)$
137. The reaction goes through the following elementary steps





Assuming that **steady state approximation** can be applied to C, on doubling the concentration of A, the rate of production of D will increase by

(assume $k_2[A] \ll k_{-1}[C]$)

- (a) 2 times (b) 4 times (c) 8 times (d) $2\sqrt{2}$ times

138. The rate of an **acid-catalysed reaction** in aqueous solution follows the rate equation $r = k [X^+][Y^{2-}][H^+]$, If k_{16} and k_4 are rate constants for the reaction at ionic strength of 16 mol L^{-1} and 4 mol L^{-1} respectively, $\ln \frac{k_4}{k_{16}}$ in terms of **Debye-Huckel constant** ($B = 0.51$), is

- (a) 4B (b) 8B (c) 10B (d) 12B

139. For two reactions



According to the **collision theory**, the ratio of squares of **pre-exponential factors** of reactions $2(A_2)$ and $1(A_1)$ at the same temperature $\left(\frac{A_2}{A_1}\right)^2$ is

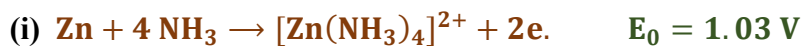
Species	Mass (g/mol)	Diameter (nm)	
X	5	0.3	(a) 4/5
Y	20	0.5	(b) 5/5
M	10	0.4	(c) 5/3
N	10	0.4	(d) 3/5

140. If the **specific conductances** of a sparingly soluble (1 : 1) salt ($MW = 200 \text{ g mol}^{-1}$) in its saturated aqueous solution at 25°C and that of water are $1.5 \times 10^{-3} \text{ ohm}^{-1} \text{ dm}^{-1}$ and $1.5 \times 10^{-5} \text{ ohm}^{-1} \text{ dm}^{-1}$, respectively, and the **ionic conductances** for its cation and anion at **infinite dilution** are 0.485 and $1.0 \text{ ohm}^{-1} \text{ dm}^{-1} \text{ mol}^{-1}$, respectively, the **solubility (in g L^{-1})** of the salt in water at 25°C is

- (a) 1×10^{-6} (b) 1×10^{-3} (c) 2×10^{-1} (d) 2×10^{-4}

141. Given,



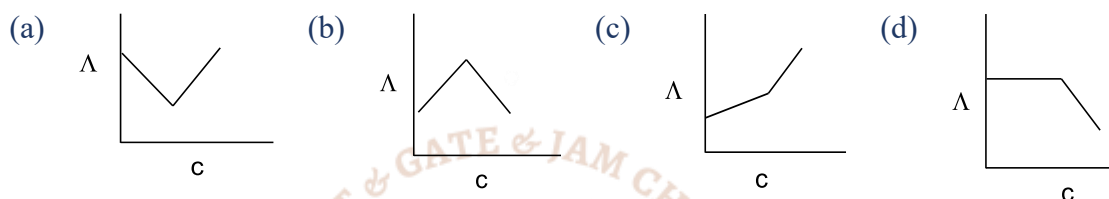


The formation constant of the complex $[\text{Zn}(\text{NH}_3)_4]^{2+}$ is approximately

$$\left(\frac{2.303RT}{F} = 0.0591 \right)$$

- (a) 1×10^5 (b) 1×10^7 (c) 1×10^9 (d) 1×10^{12}

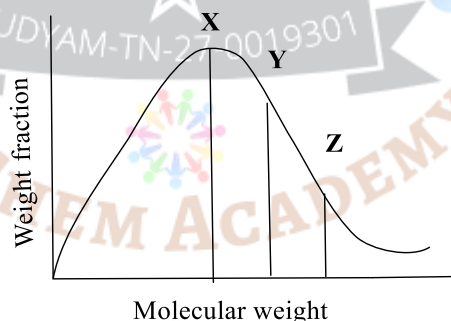
142. The molar conductivity (Λ) vs. Concentration (c) plot of sodium dodecylsulfate in water is expected to look like



143. The $\sin^2\theta$ values obtained from X-ray powder diffraction pattern of a solid are $2x$, $4x$, $6x$, $8x$ where x is equal to 0.06 . The wavelength of X-ray used to obtain this pattern is $1.54 \text{ \AA}$. The unit cell and the unit cell length, respectively, are

- (a) BCC, $3.146 \text{ \AA}$ (b) FCC, $3.146 \text{ \AA}$ (c) SCC, $6.281 \text{ \AA}$ (d) BCC, $1.544 \text{ \AA}$

144. Distribution of molar masses in a typical polymer sample shown below



The X, Y and Z represent

- (a) $\bar{M}_w$, $\bar{M}_v$, and $\bar{M}_n$, respectively (b) $\bar{M}_n$, $\bar{M}_v$, and $\bar{M}_w$, respectively
(c) $\bar{M}_v$, $\bar{M}_w$, and $\bar{M}_n$, respectively (d) $\bar{M}_n$, $\bar{M}_w$, and $\bar{M}_v$, respectively

145. Two bound stationary states, 1 and 2, of a one-electron atom, with $E_2 > E_1$ (E is the total energy) obey the following statement about their kinetic energy (T) and potential energy (V)

- (a) $T_2 > T_1$; $V_2 > V_1$ (b) $T_2 > T_1$; $V_2 < V_1$
(c) $T_2 < T_1$; $V_2 > V_1$ (d) $T_2 = T_1$; $V_2 > V_1$



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Answer Key

PART - B

Q.No	Ans
21.	a
22.	a
23.	c
24.	c
25.	c
26.	c
27.	a
28.	d
29.	a
30.	a
31.	c
32.	d
33.	a
34.	d
35.	a

Q.No	Ans
36.	c
37.	d
38.	c
39.	a
40.	a
41.	c
42.	a
43.	b
44.	a
45.	c
46.	c
47.	a
48.	a
49.	b
50.	d

Q.No	Ans
51.	b
52.	a
53.	d
54.	d
55.	c
56.	b
57.	a
58.	b
59.	c
60.	c

Q.No	Ans
61.	a
62.	d
63.	c
64.	b
65.	b
66.	b
67.	b
68.	c
69.	c
70.	b

PART - C

Q.No	Ans
71.	d
72.	a
73.	c
74.	b
75.	b
76.	c
77.	c
78.	c

Q.No	Ans
91.	b
92.	a
93.	a
94.	a
95.	a
96.	d
97.	c
98.	c

Q.No	Ans
111.	c
112.	a
113.	d
114.	d
115.	a
116.	b
117.	b
118.	a

Q.No	Ans
131.	d
132.	b
133.	a
134.	a
135.	d
136.	c
137.	d
138.	d



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79.	d
80.	b
81.	b
82.	c
83.	b
84.	b
85.	b
86.	b
87.	c
88.	a
89.	c
90.	b

99.	b
100.	c
101.	d
102.	b
103.	b
104.	a
105.	a
106.	b
107.	d
108.	a
109.	a
110.	b

119.	b
120.	d
121.	c
122.	c
123.	c
124.	d
125.	b
126.	a
127.	b
128.	d
129.	d
130.	b

139.	a
140.	c
141.	c
142.	d
143.	a
144.	b
145.	c

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