ORGANIC CHEMISTRY - SOME BASIC PRINCIPLES AND TECHNIQUES

- 3-phenylpropenoic acid is IUPAC name of:
 - a) Mendaleic acid
- b) Pyruvic acid
- c) Succinic acid
- d) Cinnamic acid
- How many isomers are possible for the compound having molecular formula C₃H₅Br₃? 2.

b) 4

c) 6

d) 8

- 3. The strain in bonds of cyclopropane is:
 - a) 0°44'
- b) 24°44'
- c) 9°44'
- d) 5°16'

- Chlorine in vinyl chloride is less reactive because:
 - a) sp^2 -hybridized carbon has more acidic character than sp^3 -hybridized carbon
 - b) C-Cl bond develops partial double bond character
 - c) Of resonance
 - d) All of the above are correct
- The alkene that exhibits geometrical isomerism is
 - a) Propene
- b) 2-methyl propene
- c) 2-butene
- d) 2-methyl-2-butene
- 6. Pick out the alkane which differs from the other members of the group
 - a) 2,2-dimethyl propane b) Pentane
- c) 2-methyl butane
- d) 2, 2-dimethyl butane

7. The IUPAC name of CH=CH is:

- a) 1-amino prop-2-enal
- b) 3-amino prop-2-enal
- c) 1-amino-2-formylethene
- d) 3-amino-1-oxoprop-2-ene
- Detection of sulphur in sodium extract is done by
 - a) Lead acetate

b) Sodium nitroprusside

c) Both (a) and (b)

d) None of these

The IUPAC name for

$$\begin{array}{c} \operatorname{CH_3} \\ \operatorname{CH_3CHOHCH_2-C-OH} \\ \operatorname{CH_3} \\ \end{array}$$

- a) 1,1-dimethyl-1,2-butanediol
- b) 2-methyl-2,4-pentanediol
- c) 4-methyl-2,4-pentanediol
- d) 1,3,3-dimethyl-1,3-propanediol
- 10. In the following carbocations, the stability order is:



(I)
$$RCH_2 \overset{+}{C}H_3$$
 (II) $CH_3 \overset{-}{C}H_3$

- a) III > II > IV > I
- b) IV > I > II > III
- c) IV > III > II > I
- d) III > IV > II > I

- 11. The shape of the π electron cloud in acetylene is
 - a) Linear
- b) Planar
- c) Cylinder
- d) Doughtnut
- 12. Acidified sodium fusion extract on addition of ferric chloride solution gives blood red colouration which confirm the presence of
 - a) S and Cl
- b) N and S
- c) N

d) S

- 13. Conversion of chlorobenzene to phenol involves
 - a) Electrophilic substitution

b) Nucleophilic substitution

c) Free radical substitution

- d) Electrophilic addition
- 14. In sulphur detection of an organic compound, sodium nitroprusside solution is added to sodium extract. Formation of violet colour is due to
 - a) Na₃Fe(CN)₆
- b) Na₃[Fe(CN)₅NOS]
- c) Fe(CNS)₃
- d) None of these

- 15. The maximum bond energy is present
 - a) C H
- b) C C
- c) C N
- d)C 0
- 16. The number of secondary hydrogens in 2, 2-dimethyl butane is
 - a) 8

b) 6

c) 4

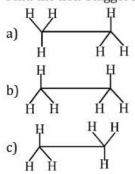
d) 2

17. The name of the compound,

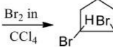
$$\mathrm{CH_3CH_2CH_2} \ \mathrm{CCH_3} \ \mathrm{is:}$$
 O

- a) 2-pentanone
- b) Pentanone-2
- c) Pentan-2-one
- d) All are correct

18. Find the non-staggered form(s) of ethane:



- d) None of these
- 19. With a change in hybridisation of the carbon bearing the charge, the stability of a carbanion increase in the
 - a) $sp < sp^2 < sp^3$
- b) $sp < sp^{3} < sp^{2}$
- c) $sp^3 < sp^2 < sp$ d) $sp^2 < sp < sp^3$
- 20. The addition reaction among the following is



- d) All of the above



21.
$$:\bar{C}H_2-C-CH_3$$
 and CH_2-C-CH_3 are $:\bar{C}H_3$ o $:\bar{C}H_3$ $:\bar{C}H_$

- a) Resonating structures
- b) Tautomers
- c) Geometrical isomers
- d) Optical isomers
- 22. The correct definition for organic chemistry is:
 - a) Chemistry of carbon compounds
 - b) Chemistry of compounds derived from living organisms
 - c) Chemistry of hydrocarbons and their derivatives
 - d) None of the above
- 23. Which of the organic compounds will give red colour in Lassaigne test?
 - b) || c) || NH₂ C NH₂ $NH_2 C NH_2$ a) NaCNS
- 24. The compound formed in the positive test for nitrogen with the Lassaigne solution of an organic compound is
 - a) $Fe_4[Fe(CN)_6]_3$
- b) $Na_3[Fe(CN)_6]$
- c) Fe(CN)₃
- d) Na₄[Fe(CN)₅NOS]

d) None of these

- - a) 1,2,3-trieyanopropane
 - b) Propane-1,2,3-tricarbonitrile
 - c) 1,2,3-cyanopropane
 - d) Propane tricarbylamine
- 26. Which of the following reactions proceeds via secondary free radical?

a)
$$CH_3 - CH = CH_2 \xrightarrow{HBr} CH_3 - CH - CH_3$$
|
| Br
| CH₃ - CH = CH₂ $\xrightarrow{HBr} CH_3 - CH_2 - CH_2$ Br
| Br
| C) $C_2H_2 \xrightarrow{Br_2/FeBr_3} C_2H_2$ Br
| C) $C_3H_4 \xrightarrow{Br_2/FeBr_3} C_4H_2$ Br

- c) $C_6H_6 \xrightarrow{Br_2/FeBr_3} C_6H_5Br$
- 27. The production of an optically active compound from a symmetric molecule without resolution is called:
 - a) Walden inversion
 - b) Asymmetric synthesis
 - c) Partial racemisation
 - d) None of these
- 28. Among the following, which one has more than one kind of hybridization?
 - (i) CH₃CH₂CH₂CH₃
 - (ii) $CH_3CH = CHCH_3$
 - (iii) $CH_2 = CH CH \equiv CH$
 - (iv) $CH \equiv CH$
 - a) (ii) and (iii)
- b) (ii) and (i)
- c) (iii) and (iv)
- d) (iv)

- 29. The IUPAC name of C₆H₅COCl is
 - a) Benzoyl chloride

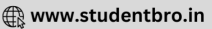
b) Benzene chloro ketone

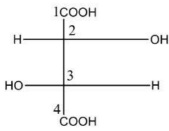
c) Benzene carbonyl chloride

d) Chloro phenyl ketone

30. In the compound,







Configuration at C_2 and C_3 atoms are

a) S, S

b) R, S

c) S, R

- d) R, R
- 31. The number of isomeric alkenes with molecular formula C₆H₁₂ are
 - a) 8

b) 10

c) 11

d) 13

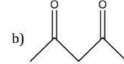
- 32. Which is wrong IUPAC name?
 - a) CH₃CH₂CH₂COOCH₂CH₃ (Ethyl butanoate)

b)
$$CH_3$$
— $CH \cdot CH_2CHO(3$ -methyl butanal) CH_3

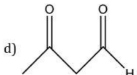
c)
$$\begin{array}{c} CH_3 \cdot CH \cdot CH \cdot CH_3 \text{(2-methyl butanal)} \\ I \\ OH \ CH_3 \end{array}$$

- 33. Which of the following statements is wrong?
 - a) In general organic compounds have low m.p. and b.p.
 - b) Isomerism is common in organic compounds
 - c) Organic compounds cannot be synthesized in the laboratory
 - d) The number of organic compound is very large
- 34. Nitroethane can exhibit one of the following kind of isomerism
 - a) Metamerism
- b) Optical activity
- c) Tautomerism
- d) Position isomerism
- 35. Which of the following would show configurational enantiomorphism?
 - a) NH₃
 - b) (CH₃)₃N
 - c) Methyl, ethyl, propylamine
 - d) Methyl, allyl, phenyl, benzyl ammonium iodide
- 36. Heterolysis of carbon-chlorine bond produces:
 - a) Two free radicals
 - b) Two carbonium ions
 - c) Two carbanions
 - d) One cation and one anion
- 37. Maximum enol content is in









- 38. Which of the following compounds will show metamerism?
 - a) $CH_3 CO C_2H_5$
- b) $C_2H_5 S C_2H_5$
- c) $CH_3 O CH_3$
- d) $CH_3 O C_2H_5$

39. The IUPAC name of the compound,

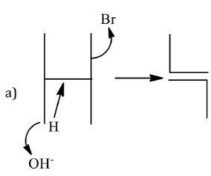
$$CH_2$$
= C - CH_2 - C = CH is:

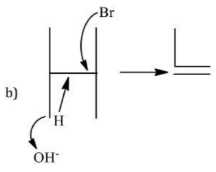
a) 2-methylpent-1-en-4-yne

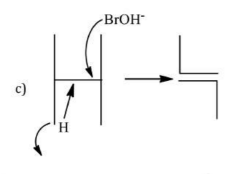


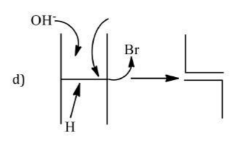
- b) 4-methylpent-4-en-1-yne
- c) 2-methylpent-2-en-4-yne
- d) 4-methylpent-1-en-4-yne
- 40. Which of the following is elimination reaction

- b) CH₃CH₂CH₂Cl + aq. KOH → CH₃CH₂CH₂OH +N(CH₃)₃
- c) $CH_3 C CH_2 CH_3 \xrightarrow{Alc.KOH} CH_3CH = CHCH_3$ CH₃ - CH - CH₂Br + Alc. KOH
- d) $\stackrel{\Delta}{\rightarrow}$ CH₃CH CH₂OH OH
- 41. Dehydrohalogenation in presence of OH is correctly represented by









42.

The IUPAC name of

- a) But-3-enoic acid
- b) But-1-enoic acid
- c) Pent-4-enoic acid
- d) Prop-2-enoic acid

- 43. On exciting Cl₂ molecules by UV light, we get

c) Cl-

- d) All of these
- 44. Mixture of sugar and common salt is separated by crystallisation by dissolving in
 - a) H₂O
- b) C₂H₅OH
- c) C_5O_6
- d) None of these

45. The structure,





shows:

- a) Geometrical isomerism
- b) Optical isomerism
- c) Geometrical and optical isomerism
- d) Tautomerism
- 46. The general formula for cycloalkanes is:
 - a) $C_n H_{2n+2}$
- b) $C_n H_{2n}$
- c) $C_n H_{2n-2}$
- d) C_nH_n

47. The IUPAC name of the compound

- a) 2(carboxymethyl)-pentane-l,5-dioic acid
- b) 3-carboxyhexane-l, 6-dioic acid
- c) Butane-l, 2, 4-tricarboxylic acid
- d) 4-carboxyhexane-l, 6-dioic acid
- 48. $Na_2S + Na_2[Fe(CN)_5NO] \rightarrow Purple colour.$ It is due to
 - a) Na₄[Fe(CN)₃NOS]
- b) Na₃[Fe(CN)₅NOS]
- c) Na₄[Fe(CN)₅NO]
- d) Na₄[Fe(CN)₅NOS]

- 49. The bond that undergoes heterolytic cleavage most easily is
 - a) C O
- b) C C
- c) C H
- d) 0 H
- 50. Increasing order of stability among the three main conformations (i.e., Eclipse, Anti, Gauche) of 2fluoroethanol is
 - a) Eclipse, Gauche, Anti b) Gauche, Eclipse, Anti c) Eclipse, Anti, Gauche d) Anti, Gauche, Eclipse
- 51. Phosphorus is estimated as
 - a) Na₃PO₄
- b) P2O5
- c) P_2O_3
- d) $Mg_2P_2O_7$
- 52. The number of asymmetric carbon atoms and the number of optical isomers in CH₃(CHOH)₂COOH are respectively:
 - a) 3 and 4
- b) 1 and 3
- c) 2 and 4
- d) 2 and 3
- 53. Species containing carbon with three bonds and an electron are called:
 - a) Carbenes
- b) Caarbanions
- c) Carbocation
- d) Free radicals

- 54. Which of the aldehyde is most reactive?
 - a) $C_6H_5 CHO$

b) CH₃CHO

c) HCHO

- d) All the equally reactive
- 55. Which of the following cannot show S_N1 reaction?









- 56. 3-methyl penta-1,3-diene is:
 - a) $CH_2 = CH(CH_2)_2CH_3$
 - b) $CH_2 = CHCH(CH_3)CH_2CH_3$
 - c) $CH_3CH = C(CH_3)CH = CH_2$
 - d) $CH_3 CH = CH(CH_3)_2$
- 57. Which of the following compounds is optically active?
 - a) 1 butanol
- b) Isopropyl alcohol
- c) Acetaldehyde
- d) 2-butanol
- 58. How many optically active forms are possible for a compound of the formula, CHO. CHOH. CHOH. CHOH. CH2OH?







		***	*** * **	(n. n//			
120	a) 2	b) 4	c) 3	d) 8			
59.	그 그 그렇게 하나 있는데 바다 하나 하나 하는데	iddendum adds on the carb	on atom joined to the least	number of hydrogen atoms."			
	This statement is called:						
	a) Markownikoff's rule						
	b) Peroxide effect						
	c) Baeyer's strain theory						
	d) Thiele's theory						
60.		eric carbocations possible					
	a) 3	b) 4	c) 2	d) 5			
61.	The correct order for homolytic bond dissociation energies. (ΔH in kcal/mol) for $CH_4(A)$, $C_2H_6(B)$ and						
		l experimental conditions					
	a) C>B>A	b) B>C>A	c) C>A>B	d) A>B>C			
62.	The sodium extract of an organic compound on treatment with FeSO ₄ solution, FeCl ₃ and HCl gives a red						
	solution. The organic compound contains						
	a) Both nitrogen and sulp	hur	b) Nitrogen only				
	c) Sulphur only		d) Halogen				
63.	d-tartaric acid and l-tarta	ric acid are :					
	a) Structureal isomers	b) Diastereoisomers	c) Tautomers	d) Enantiomers			
64.	Which of the following is	a pair of functional isomer:	s?				
	a) CH ₃ COCH ₃ , CH ₃ CHO		b) C ₂ H ₅ CO ₂ H, CH ₃ CO ₂ CH	3			
	c) C ₂ H ₅ CO ₂ H, CHCO ₂ C ₂ H	5	d) CH ₃ CO ₂ H, CH ₃ CHO				
65.	Which of the following is	an optically active compou	nd?				
	a) Lactic acid	b) Chloro acetic acid	c) Meso-tartaric acid	d) Acetic acid			
66.	Give the correct IUPAC na	ime for					
	CH ₃						
	Î						
	CH ₃ . CH ₂ OCH. CH ₂ . CH ₂ . CH ₂ Cl						
	a) 2-ethoxy-5-chloropent	ane	b) l-chloro-4-ethoxy-4-m	ethylbutane			
	c) 1-chloro-4-ethoxypentane		d) Ethyl-1-chloropentylether				
67.	The IUPAC name of the compound,						
	CH ₂ —CH—CH ₂ is:						
	он он он						
	a) 1,2,3-trihydrosypropa	ne					
	b) 3-hydroxypentane-1,5-diol						
	c) 1,2,3-hydroxypropane						
	d) Propane-1,2,3-triol						
68.	3. Bond energywith the increase in number of lone pairs on the bonded atoms.						
	a) Decreases	b) Increases	c) Does not change	d) None of these			
69.	A liquid decomposes at it	s normal boiling point. It ca	an be purified by				
	a) Sublimation		b) Steam distillation				
	c) Vacuum distillation		d) Fractional distillation				
70.	On monochlorination of 2	-methyl butane, the number	er of chiral compounds for	med are :			
	a) 2	b) 4	c) 6	d) 8			
71.	Stability of which interme	ediate is not governed by h	yperconjugation?				
	a) Carbon cation	b) Carbon anion	c) Carbon free radical	d) None of these			
72.	The ammonia evolved from the treatment of 0.30g of an organic compound for the estimation of nitrogen						
	was passed in 100mL of 0	as passed in 100mL of 0.1M sulphuric acid. The excess of acid required 20mL of 0.5 M sodium hydro					
	solution for complete neutralisation. The organic compound is						
	a) Acetamide	b) Benzamide	c) Urea	d) Thiourea			
73.	Conversion of CH ₄ to CH ₃	Cl is an example of which o	of the following reaction?				

a) Electrophilic substitution

b) Free radical addition

c) Nucleophilic substitution

- d) Free radical substitution
- 74. Number of possible isomers of glucose are:
 - a) 10

b) 14

c) 16

d) 20

75. The reaction

 $CH_3CH_2CHCH_3 \xrightarrow{NaNH_2} Butene-1$ and butane -2 (major) | Br

The correct statement (s) are

a) 2-butene is Saytzeff product

- b) 1-butene is Hofmann (s) product
- c) The elimination reaction follows Saytzeff rule
- d) All of the above
- 76. Consider the following carbanions

(II)
$$O_2N$$
— $\tilde{C}H_2$

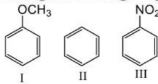
$$(III)$$
 $\langle \bigcirc \rangle$ $-\ddot{C}H_2$

Correct order of stability is

- a) I>II>III
- b) III>II>I
- c) II>III>I
- d) I>III>II
- 77. The stability of 2,3-dimethyl but-2-ene is more than 2-butene. This can be explained in terms of:
 - a) Resonance
- b) Hyperconjugation
- c) Electromeric effect
- d) Inductive effect

- 78. Protin solvent is
 - a) Diethyl ether
- b) n-hexane
- c) Acetone
- d) Ethanol

- 79. Addition of Br2 on trans-butene-2 gives:
 - a) A racemic mixture of 2,3-dibromobutane
 - b) Meso form of 2,3-dibromobutane
 - c) Dextro form of 2,3-dibromobutane
 - d) Laevo form of 2,3-dibromobutane
- 80. Among the following compounds (I-III) the correct order of reaction with electrophilic reagand is



- a) II>III>I
- b) III<I<II
- c) I>II>III
- d) I=II>III
- 81. During AgNO₃ test for detection of halogens, sodium extract is boiled with few drops of conc. HNO₃ to decompose
 - a) NaCN
- b) Na₂S
- c) Both (a) and (b)
- d) None of these

82. Which is true about following?



- a) Only III is a chiral compound
- b) Only II and IV are chiral compounds
- c) All four are chiral compounds
- d) Only I and II are chiral compounds
- 83. How many chiral compounds are possible on monochlorination of 2-methyl butane?

b) 4

- 84. How many isomers of C₅H₁₁OH will be primary alcohols?

b) 4

d) 2

- 85. The epoxide ring consists of which of the following?
 - a) Three membered ring with two carbon and one oxygen
 - Four membered ring with three carbon and one oxygen
 - c) Five membered ring with four carbon and one oxygen.
 - d) Six membered ring with five carbon and one oxygen.
- 86. The reaction which is not the example of nucleophilic substitution among the following is
 - a) $CH_3C Br + CH_3OH \rightarrow CH_3C OCH_3 + HBr$
- b) $CH_3C Cl + aq. KOH \rightarrow CH_3C OH + KCl$
- c) \sim CI+ alc. KOH \rightarrow + KCI+H₂O d) \rightarrow Br + aq. KOH \rightarrow OH + HO
- 87. Consider the following reaction

$$>$$
C=O + H₂NOH \rightarrow $>$ C=NOH +H₂O

Is an example of

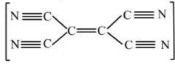
- a) Substitution
- b) Elimination
- c) Addition
- d) Addition elimination
- 88. An important chemical method to resolve a racemic mixture makes use of the formation of:
 - a) meso compound
- b) Enantiomer
- c) Racemers
- d) diastereoisomers
- 89. Red colour complex ion formed on adding FeCl₃ to sodium extract when N and S both are present in organic compound is
- a) $[Fe(CN)_6]^{4-}$
- c) [Fe(CNS)₂]⁺
- d) $[Fe(CN)_6]^{3-}$

- 90. (I) $CH_3CH_2Br \xrightarrow{LAH} C_2H_6$ and (II)
 - $(CH_3)_3CBr \xrightarrow{LAH}$ alkene, The reason for this is
 - a) (I)S_N2(II) E₁ mechanism

b) (I) S_N1, (II) E₂ mechanism

c) (I)S_N1, (II) E₁ mechanism

- d) (I) S_N2,(II) E₂ mechanism
- 91. How many σ and π -bonds are there in the molecule of tetracyanoethylene?



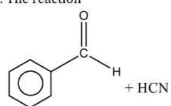
- a) 9σ and 9π
- b) 5σ and 9π
- c) 9σ and 7π
- d) 5σ and 8π
- 92. Which of the following complex formation indicates presence of sulphur in the organic compound when sodium nitroprusside is added to sodium extract of the compound?
 - a) $Fe_4[Fe(CN)_6]_3$
- b) $Na_2[Fe(NO)(CN)_5]$
- c) $Fe_4(CNS)_3$
- d) Na₄[Fe(CN)₅NOS]

93. Who pointed out the concept hyperconjugation?

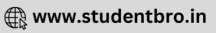




94.	a) Nathan and Baker Alkyl halide can be conver	ted into alkene by	c) Kekule	d) Kolbe		
	a) Nucleophilic substitution reaction b) Elimination reaction					
		itution and elimination rea	action			
	d) Rearrangement					
95.	The order of reactivities of the following alkyl halides for a $S_N 2$ reaction is : a) $RF > RCI > RBr > RI$					
	b) $RF > RCI > RI$					
	c) $RCI > RBr > RF > RI$					
06	d) RI > RBr > RCl > RF The entirally active alkane with lowest molecular weight is:					
90.	The optically active alkane with lowest molecular weight is :					
	а) СН₃СН₂С≡СН	b) CH ₃ CH ₂ —CH—CH ₃	c) $CH_3 - C$ C_2H_5	d) CH ₃ CH ₂ ·CH ₂ CH ₃		
97.	Which type of isomerism i	s most common among eth	ners?			
00	a) Metamerism	b) Functional	c) Chain	d) Position		
98.	With a change in hybridisation of the carbon bearing the charge, the stability of a carbanion increase in the order					
		b) $sp < sp^3 < sp^2$	c) $\operatorname{sp}^3 < \operatorname{sp}^2 < \operatorname{sp}$	d) $\operatorname{sp}^2 < \operatorname{sp} < \operatorname{sp}^3$		
99.	A molecule is R_3C —H. If H is replaced by $Z(R_3C$ —Z) and on doing so electron density on R_3 —C par					
	increases, then Z is: a) Electron attracting grounds	ın				
	b) Electron withdrawing g	(3)				
	c) Electron repelling grou	p				
100	d) Either of the above 100. Which of the following compounds are not arranged on order of decreasing reactivity towards electrophilic substitution?					
100.						
	a) Fluorobenzene > chlorobenzene > bromo benzene					
	b) Phenol> n-propyl benzene> benzoic acid>					
	c) Chlorotoluene >para-nitrotoluene>2-chloro-4-nitro toluene d) Benzoic acid> phenol>n-propyl benzene					
101.	101. A mixture of camphor and benzoic acid can be separated by					
	a) Sublimation		b) Extraction with a solve			
102	c) Chemical method		d) Fractional crystallisation ion of π -electrons. Each π -electron is attached with :			
102.	a) 4 carbon	b) 2 carbon	c) 3 carbon	d) 6 carbon		
103.	Grignard reagent adds to			u, o our oor		
	a) >c==0	b) $-C \equiv N$	c) >c—s	d) All of these		
104.	04. Resonance energy is more for					
	a) C ₆ H ₆		b) Cylohexene			
105	c) Cycloheptene The reaction		d) Cyclohexa -1,2,3-triene			
	O O					







is an example of a) Electrophilic addition b) Electrophilic substitution c) Nucleophilic substitution d) Nucleophilic addition 106. Which of the following is not chiral? a) 3-bromopentane b) 2-hydroxy propanoic acid c) 2-butanol d) 2,3-dibromopentane 107. The structures (CH₃)₃CBr and CH₃[CH₂]₃Br represent a) Chain isomerism b) Position isomerism c) Chain as well as position isomerism d) Functional isomerism 108. Detection of phosphorus in the compound can be done by its conversion into phosphate. Reagent to identify phosphate ion is a) Sodium nitroprusside b) Ammonium molybdate c) Potassium ferrocyanide d) Potassium ferricyanide 109. The hemolytic fission of a hydrocarbon results in the formation of: a) Carbonium ions b) Free radicals c) Carbanions d) Carbenes 110. Which does not have sp^2 -hybridised carbon atom? a) Acetamide b) Acetic acid c) Acetonitrile d) Acetone 111. Methoxy methane and ethanol are a) Position isomers b) Chain isomers c) Functional isomers d) Optical isomers 112. Which one of the following monoenes does not exhibit geometric isomerism? b) C₃ H₆ c) C₅H₁₀ a) C_4H_8 d) C₈H₁₆ 113. Addition of Br₂ on cis - butene - 2 gives : a) A racemic mixture of 2,3-dibromobutane b) Meso form of 2,3-dibromobutane c) Dextro form of 2,3-dibromobutane d) Laevo form of 2,3-dibromobutane 114. When two halogen atoms are attached to two adjacent carbon atoms, the dihaloalkane is called: a) Alkylidene dihalide b) Alkane dihalide c) Alkylene dihalide d) Alkyl halide 115. The electrophile involved in the sulphonation of benzene is a) SO₃⁺ b) SO_3^{2-} c) H₃O⁺ d) SO_3 116. The number of meso forms in the following compound is HOOC. CH(CH₃). CH(OH). $CH(CI).CH(OH)CH(CH_3).$ COOH a) 3 b) 4 c) 8 d) 16 117. In Kjeldahl's method, then nitrogen present in the organic compound is quantitatively converted into a) Gaseous ammonia b) Ammonium sulphate c) Ammonium phosphate d) Ammonia 118. The reaction; NH2 $NHCOCH_3$ COCH₃ is called: a) Substitution reaction b) Elimination reaction c) Rearrangement reaction

- d) None of the above
- 119. The compound which gives the most stable carbonium ion on dehydration is
 - a) CH₃CH(CH₃)CH₂OH
- b) $(CH_3)_3COH$
- c) $CH_2 = CHCH_2CH_2OH$ d) $CH_3CHOHCH_2 CH_3$
- 120. A similarity between optical and geometrical isomerism is that:
 - a) Each forms equal number of isomers for a given compound
 - b) If in a compound, one is present then so is the other
 - c) Both are included in stereoisomerism
 - d) They have no similarity
- 121. Reaction of phenol with chloroforms/sodium hydroxide to give o-hydroxy benzaldehyde involves the formation of
 - a) Dichloro carbene
- b) Trichloro carbene
- c) Chlorine atoms
- d) Chlorine molecules

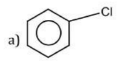
- 122. Which kind of fission is favoured by sunlight?
 - a) Heterolytic fission
- b) Homolytic fission
- c) Both (a) and (b)
- d) None of these
- 123. The stability of the free radicals allyl, benzyl, 3°, 2°, 1° and CH3 is in the order
 - a) Benzyl > allyl > 3° > 2° > 1° > CH_3
- b) Allyl $> 3^{\circ} > \text{benzyl} > 2^{\circ} > 1^{\circ} > \text{CH}_3$
- c) $3^{\circ} > 2^{\circ} > 1^{\circ} > CH_3 > \text{allyl} > \text{benzyl}$
- d) $3^{\circ} > 2^{\circ} > 1^{\circ} > CH_3 > allyl = benzyl$
- 124. Which class of compounds can exhibit geometrical isomerism?
 - a) $C_6H_5CH = NOH$
 - b) $CH_3CH = CHCH_3$
 - HOOCCH $-CH_2$ —CHCOOH

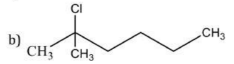
 - d) All of the above
- 125. The correct order of increasing basicity of the given conjugate bases ($R=CH_3$) is
 - a) $RCO\overline{O} < HC \equiv \overline{C} < \overline{R} < \overline{N}H_2$

b) $\overline{R} < HC \equiv \overline{C} < RCO\overline{O} < \overline{N}H_2$

c) $RCO\overline{O} < NH_2 < HC \equiv \overline{C} < \overline{R}$

- d) $RCO\overline{O} < HC \equiv \overline{C} < \overline{N}H_2 < \overline{R}$
- 126. Which of the following shows S_N1 reaction most readily?





- 127. Which of the following compounds is optically active?
 - a) (CH₃)₂CHCH₂OH
- b) CH₃CH₂OH
- c) CCl₂F₂
- d) CH₃CHOHC₂H₅
- 128. To which ring size cycloalkanes, Baeyer's strain theory is not valid?
- b) 4 carbon
- c) 5 carbon
- d) \geq 6 carbon
- 129. The S_N1 mechanism for substitution reaction by nucleophile is favoured by :
 - a) Low concentration of nucleophile
 - b) Weak nature of nucleophile
 - c) Polar solvent
 - d) All of the above
- 130. Which of the following orders is not correct regarding the -I effect of the substitutents?

a)
$$-I < -Cl < -Br < -F$$

b)
$$-\stackrel{+}{N} R_3 < -\stackrel{+}{O} R_2$$

c)
$$-{}^{+}_{N}R_{2} < -0R < -F$$

d)
$$-SR < -OR < -OR_2$$

131. Lactic acid shows optical activity in:



a) Solution state b) Liquid state c) Crystalline state d) In all states 132. In cyclopropane, cyclobutane and cyclohexane, the common group is a) -C b) CH2 d) - CHc) $-CH_3$ 133. Total number of isomeric aldehydes and ketones that can exist with the molecular formula $C_5H_{10}O$: a) 5 b) 8 c) 6 d) 7 134. Allyl isocyanide has: a) 9σ and 4π -bonds b) 8σ and 5π -bonds c) 9σ , 3π and 2 non-bonded electrons d) 8σ , 3π and 4 non-bonded electrons 135. +/effect is shown by a) $-CH_3$ b) -Br c) -Cl d) - NO₂136. LiAlH₄ is used as: b) Reducing agent d) A water softener a) Oxidizing agent c) A mordant 137. 0.765g of an acid gives 0.535g of CO2 and 0.138 g of H2O. Then, the ratio of the percentage of carbon and hydrogen is a) 19:2 b) 18:11 c) 20:17 d) 1:7 138. Which one of the following is the stable structure of cyclohexatriene? b) Boat form c) Half chair form d) Planar form a) Chair form 139. The IUPAC name of compound shown below is a) 2-bromo-6- chlorocyclohex-1-ene b) 6-bromo-2-chlorocyclohexene c) 3-bromo-1-chlorocyclohexene d) 1-bromo-3-chlorocyclohexene 140. Total number of rotational conformers of n-butane are : a) 2 b) 6 c) 5 d) 3 141. Sublimation is a process in which a solid a) Changes into vapour form b) Changes into another allotropic form c) Changes into liquid form d) None of the above 142. IUPAC name of the compound a) 5- methyl-4-isopropyl-6, 6'diethyloctane b) 3, 3-dimethyl, 3-ethyl-5- isopropyl octane c) 3, 3-diethyl-4-methyl-5-(1,1-dimethyl) octane d) 3, 3- diethyl-4-methyl-5-(1'-methylethyl) octane 143. The group named as benzal possessesnature. d) Tetravalent a) Monovalent b) Bivalent c) Trivalent 144. A secondary(2°) carbon is one that is joined to:

a) 1-alkyl group

b) 2-alkyl groups

c) 3-alkyl groups

d) None of these

145. Which type of strain is present in fully eclipsed conformation of butane?

a) Angle strain

b) Steric strain

c) Both (a) and (b)

d) Neither (a) nor (b)

146. 29.5 mg of organic compound containing nitrogen was digested according to Kjeldahl's method and the evolved ammonia was absorbed in 20mL of 0.1 M HCl solution. The excess of the acid required 15mL of 0.1M NaOH solution for complete neutralisation. The percentage of nitrogen in the compound is

a) 59.0

b) 47.4

c) 23.7

d) 29.5

147. The highest electrical conductivity of the following aqueous solutions is of

a) 0.1 M difluoroacetic acid

b) 0.1 M fluoroacetic acid

c) 0.1 M chloroacetic acid

d) 0.1 M acetic acid

148. Consider the following carbanions

Correct order of stability is

a) 1>2>3

b) 3>2>1

c) 2 > 3 > 1

d) 1>3>2

149. Which of the following compounds (s) has 'Z' configuration?

a) (i) only

b) (ii) only

c) (iii) only

d) (i) and (iii)

150. Nucleophiles are:

a) Electron loving

b) Electron hating

c) Nucleus loving

d) Nucleus hating

151. Ethyl acetoacetate exhibits:

- a) Optical isomerism
- b) Geometrical isomerism
- c) Tautomerism
- d) enantiomerism

152. The total number of cyclic isomers possible for a hydrocarbon with the molecular formula C₄H₆ is

153.

Cl is: The IUPAC name of

- a) 2-ethyl-3-methylbutanoyl chloride
- b) 2,3-dimethylpentanoyl chloride
- c) 3,4-dimethylpentanoyl chloride
- d) 1-chloro-l-oxo-2,3-dimethylpentane

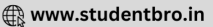
154. A molecule of urea can show

a) Chain isomerism

b) Position isomerism

c) Geometrical isomerism

d) Tautomerism



155. Cyclic hydrocarbon molecule (*A*) has all the C and H atoms in single plane. All the C–C bonds have same length, less than 1.54 Å but more than 1.34Å. The ∠ (angle) CCC is :

- a) 190°28′
- b) 100°
- c) 180°
- d) 120°

156. The number of π -electrons present in cyclobutadienyl ion, $(C_4H_3)^-$ is :

a) 8

b) 6

c) 4

d) 2

157. Geometrical isomerism is possible in case of

- a) Pentene-2
- b) Propane
- c) Pentane
- d) Ethene

158. The strongest best among the following is:

- a) NH₄⁺
- b) :NH₃
- c) : NH2
- d):ŌH

159. Anti-Markownikoff addition of HBr is not observed in:

- a) Propene
- b) Butene-1
- c) But-2-ene
- d) Pent-2-ene

160. The number of 1°, 2° and 3° carbon atoms present in isopentane are respectively:

- a) 3, 2, 1
- b) 2, 3, 1
- c) 3, 1, 1
- d) 2, 2,1

161. The restricted rotation about carbon-carbon double bond in 2-butene is due to:

- a) Overlap of two p-orbitals
- b) Overlap of one p and one sp^2 -hybridized orbitals
- c) Overlap of two sp^2 -hybridized orbitals
- d) Overlap of one s and one sp^2 -hybridized orbitals

162. Formation of acetylene from ethylene is an example of

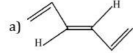
a) Addition reaction

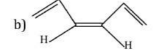
b) Substitution reaction

c) Elimination reaction

d) Condensation reaction

163. The structure of cis-bis (propenyl) ethane is:



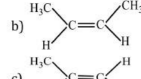


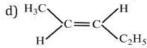




164. The compound which reacts with HBr obeying Markownikoff's rule is:

a) $CH_2 = CH_2$



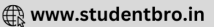


- 165. A molecule of benzene contains:
 - a) Twelve sigma-bonds and three pi-bonds
 - b) Eighteen sigma-bonds and three pi-bonds
 - c) Twelve pi-bonds and three sigma-bonds
 - d) Six hydrogen-bonds, six sigma-bonds and three pi-bonds

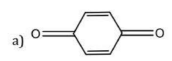
166. Zn—Cu couple used as reducing agent is:

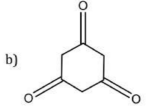
- a) Mixture of Zn and Cu powder
- b) Copper deposited on granulated zinc
- c) Zn deposited on copper fillings
- d) A rod half made of copper and half made of zinc

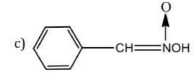


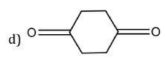


- 167. Considering the state of hybridization of carbon atoms, find out the molecule among the following which is linear?
 - a) CH₃-CH₂-CH₂-CH₃
 - b) $CH_3 CH = CH CH_3$
 - c) $CH_3 C \equiv C CH_3$
 - d) $CH_2 = CH CH_2 C \equiv CH$
- 168. Tautomerism is not exhibited by



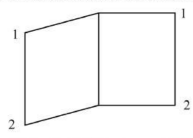






- 169. Which of the substance is purified by sublimation?
 - a) Naphthalene
- b) Benzoic acid
- c) Camphor
- d) All of these

170. The IUPAC name of the following compound is



- a) Bicyclo [2,2,0] octane b) Bicyclo [0,2,2] hexane c) Bicyclo [2,1,1] hexane d) Bicyclo [2,2,0] hexane
- 171. $\begin{array}{c} CH_3 \\ H_3C \end{array} \longrightarrow \begin{array}{c} CH_3 \\ OH \end{array} \xrightarrow{H^+} [F] \xrightarrow{Br_2,CCl_4} \underbrace{C_4H_8Br_2}_{\substack{5 \text{ such produc} \\ \text{are possible}}}$

How many structures of F are possible?

- a) 2
- b) 5
- c) 6
- d) 3
- 172. Example of chlorinolysis is:
 - a) $CH_2 = CH_2 \xrightarrow{Cl_2} C_2H_4Cl_2$
 - b) $CCl_4 + H_2O \rightarrow COCl_2 + 2HCl$
 - c) $CHCl_3 + 4NaOH \rightarrow HCOONa + 3NaCl + 2H_2O$
 - d) $C_3H_8 \stackrel{Cl_2}{\rightarrow} CCl_4 + C_2Cl_6 + 8HCl$
- 173. The number of optical enantiomorphs of tartaric acid:
 - a) 3

b) 2

c) 4

d) 1

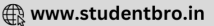
174. IUPAC name of $CH_3 - CH - CH_2 - CHO$ is

Cl

a) 3-chlorobutanol

b) 3-chlorobutanaldehyde





c) 3-chlorobutanal

d) 2-chlorobutanol

175. 4 g of hydrocarbon on complete combustion gave 12.571 g of CO2 and 5.143 g of water. What is the empirical formula of the hydrocarbon?

b) C₂H₃

c) CH₂

d) CH₃

176. The compound which contains all the four 1°, 2°, 3° and 4° carbon atoms is

a) 2, 3-dimethylpentane

b) 3-chloro-2, 3-dimethylpentane

c) 2, 3, 4-trimethylpentane

d) 3, 3-dimethylpentane

177. Which of the following is useful for making pure water from a solution of salt in water?

a) Filtration

b) Distillation

c) Chromotography

d) Steam distillation

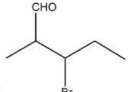
178. Which of the following does not contain chiral carbon atom?

a) Lactic acid

b) 2-chlorobutanoic acid c) Tartaric acid

d) Succinic acid

179.



The IUPAC name of

a) 2-methyl-3-bromohexanal

b) 3-bromo-2-methylbutanal

c) 2-bromo-3-bromobutanal

d) 3-bromo-2-methylpentanal

180. CH₃CH₂Cl undergoes homolytic fission, produces

181. Among the following orbital bonds, the angle is minimum between:

a)
$$sp^3 - sp^3$$
 bonds

b)
$$p_x$$
 and p_y -orbitals

d)
$$sp - sp$$
 bonds

182. Which of the following is the correct order of decreasing $S_N 2$ reactivity? ($X = \alpha$ halogen)

is

a)
$$RCH_2X > R_3CX > R_2CHX$$

b)
$$RCH_2X > R_2CHX > R_3CX$$

c)
$$R_3CX > R_2CHX > RCH_2X$$

d)
$$R_2CHX > R_3CX > R_2CH_2X$$

183. Write the IUPAC name of

$$\begin{array}{c} \text{OH} \\ \mid \\ \text{CH}_3 - \text{CH}_2 - \text{C} - \text{CH}_2 - \text{CH}_3 \\ \mid \\ \text{CH}_3 \end{array}$$

a) 3-methylpentane-3-ol

b) 3-hydroxyhexane

c) 3-hydroxy-3-methyl pentane

d) All of the above

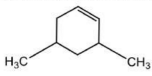
184. Polarization of electron in acrolein may be written as

a)
$$_{\text{CH}_2}^{-\delta}$$
=CH-CH=O

b)
$$_{\text{CH}_2}^{-\delta}$$
 = CH - CH = O

d)
$$^{+}_{\text{CH}_2}$$
=CH-CH=O

185. IUPAC name of the following compound is



a) 3, 5-dimethylcyclohexene

b) 3, 5-dimethly-1-cyclohexene

c) 1, 5-dimethly-5-cyclohexene

d) 1, 3-dimethyl-5-cyclohexene

186. In this reaction,

 $CH_3CHO + HCN \rightarrow CH_3CH(OH)CN$

 $\xrightarrow{H.OH}$ $CH_3CH(OH)COOH$

an asymmetric centre is generated. The acid obtained would be

- a) 50%D+50%L-isomer b) 20%D+80%L-isomer c) D-isomer
- d) L-isomer
- 187. Two crystalline forms of a substance, one being a mirror image of the other are called:
 - a) Pentane
- b) Chain isomers
- c) Stereoisomers
- d) Functional isomers
- 188. Which one of the following is an intermediate in the reaction of benzene with CH₃Cl in the presence of anhydrous AlCl₃?
 - a) Cl+

- b) CH₃
- c) CH₃+

- 189. The number of optical isomers of CH₃CH(OH)CH(OH)CHO is
- b) 2

- 190. The ratio of σ -to π -bonds in benzene is:
 - a) 2

b) 4

c) 6

d) 8

191. In a S_N2 substitution reaction of the type

$$R - Br + Cl^{-} \xrightarrow{DMF} R - Cl + Br^{+}$$

Which one of the following has the highest relative rate?

a)
$$\begin{array}{c} \text{CH}_3 - \text{CH} - \text{CH}_2 \text{Br} + \text{CI}^- \\ \text{CH}_3 \end{array}$$

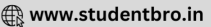
- c) CH₃CH₂Br
- d) CH₃-CH₂-CH₂Br
- 192. Hyperconjugation is
 - a) $\sigma \pi$ delocalisation
- b) No bond resonance
- c) $\sigma \pi$ odd electron
- d) All of these
- 193. Which one of the following reactions is a condensation reaction?
 - a) HCHO →para-formaldehyde
 - b) $CH_3CHO \rightarrow para$ -aldehyde
 - c) CH₃COCH₃→ mesityl oxide
 - d) $CH_2 = CH_2 \rightarrow polyethylene$
- 194. Which group has the maximum-Inductive effect?
 - a) $-NO_2$
- b) CN
- c) -COOH
- d) -F

195. The correct IUPAC name of the following compound is

 $CH_3CH_2CH - C = C - CHCH_2CH_2CH_2CH_3$

- a) 5, 6-dimethyl-8-methyl dec-6-ene
- b) 6-butyl-5-ethyl-3-methyl oct-4-ene
- c) 5, 6-diethyl-3-methyl dec-4-ene
- d) 2, 4, 5-triethyl non-3-ene
- 196. Which is incorrect about enantiomorphs?
 - a) They rotate the plane of polarized light in different directions
 - b) They have mostly identical physical properties
 - c) They have same configuration
 - d) They have different biological properties
- 197. Which one is the seniormost functional group in the nomenclature of an organic compound if it possesses more than one functional group?





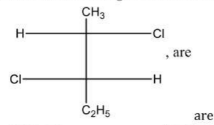
a) —CHO

b) —COOH

c) - OH

d) CO

198. The absolute configuration of the following



a) 2S, 3R

b) 2S, 3S

c) 2R, 3S

d) 2R, 3R

199. Which step is chain termination step in the following mechanism?

(i)
$$Cl_2 \xrightarrow{hv} Cl^{\bullet} + Cl^{\bullet}$$

(ii)
$$Cl^{\bullet} + CH_4 \longrightarrow \mathring{C}H_3 + HCl$$

$$(iii) \overset{\bullet}{\mathrm{CH}}_3 + \mathrm{Cl}_2 \longrightarrow \mathrm{CH}_3 \mathrm{Cl} + \mathrm{Cl}^{\bullet}$$

(iv)
$$Cl^{\bullet} + \mathring{C}H_3 \longrightarrow CH_3Cl$$

b) (ii)

c) (iii)

d) (iv)

200. The reaction intermediate produced, by homolytic cleavage of a bond is called

a) Carbene

b) Carbocation

c) Carbanion

d) Free redical

201. Fractional distillation is useful in distillation of

a) Petroleum

b) Coal-tar

c) Crude alcohol

d) All of these

202. Which of the following species is paramagnetic?

a) A carbocation

b) A free radical

c) A carbanion ion

d) All of these

203. Sulphur trioxide is:

a) An electrophile

b) A nucleophile

c) A homolytic reagent

d) A base

204. In Kjeldahl's method, ammonia from 5g of food neutralizes 30 cm³ of 0.1 N acid. The percentage of nitrogen in the food is

a) 0.84

b) 8.4

c) 16.8

d) 1.68

205. The number of isomeric alkanes having the molecular formula C₅H₁₂ is

a) Three

b) Five

c) Nine

d) Thirty two

206. Select the organic compound which was prepared for the first time in laboratory from its elements:

a) Urea

b) CH₃COOH

c) C₂H₅OH

d) None of these

207. Which of the following compounds can exist in optically active form?

a) 1-butanol

b) 2-butanol

c) 3-pentanol

c) CH₃CHO

d) 4-heptanol

208. The compound in which carbon uses only its sp^3 hybrid orbitals for bond formation is

a) (CH₃)₃COH

b) **НСООН**

d) $(H_2N)_2CO$

209. How many types of functional group can be present in an amine with the formula C₃H₉N?

b) 2

c) 3

210. Select the most reactive cycloalkane:

a) Cyclopropane

b) Cyclobutane

c) Cyclopentane

d) Cyclohexane

211. The -I effect is shown by :

a) -COOH

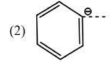
b) $-CH_3$

c) -CH₃CH₂

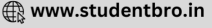
d) $-CHR_2$

212. The stability of carbanions in the following;

(1)
$$-RC = \overset{\Theta}{C}$$







(3) $R_2C = \overset{\Theta}{C}H$ (4) $R_3C - \overset{\Theta}{C}H_2$ is in the order of: a) (2) > (3) > (4) > (1)b) (4) > (2) > (3) > (1)c) (1) > (3) > (2) > (4)d) (1) > (2) > (3) > (4)213. Glyoxal is a) $CH_2OH - CH_2OH$ b) $CHO - CH_2OH$ c) COOH - CO - COOH d) CHO - CHO 214. IUPAC name of acraldehyde is b) Propenyl aldehyde a) But-3-en-1-al c) But-2-ene-1-al d) Prop-2-en-1-al 215. The IUPAC name of $CH_3 - C \equiv CH(CH_3)_2$ is a) 4-methyl-2-pentyne b) 4, 4-dimethyl-2-butyne d) 2-methyl-4-pentyne c) methyl isopropyl acetylene 216. What information is provided by reaction mechanism? a) The bonds broken and formed b) The reaction intermediates c) The relative rates of discrete steps, especially the slowest one d) All of the above 217. The enolic from of acetone contains a) 8σ bonds, 2π -bonds and 1 lone pair b) 9σ bonds, 1π -bonds and 2 lone pairs c) 9σ bonds, 2π -bonds and 1 lone pair d) 10σ bonds, 1π -bonds and 1 lone pair 218. Which of the following acids has the smallest dissociation constant? a) CH3CHFCOOH b) FCH2CH2COOH c) BrCH2CH2COOH d) CH₃CHBrCOOH 219. IUPAC name of, CH3CH(OH)CH2CH2COOH is: a) 4-hydroxypentanoic acid b) 1-carboxy-3-butanoic acid c) 1-carboxy-4-butanol d) 4-carboxy-2-butanol 220. The number of isomers for the compound with the molecular formula C2BrClFI is c) 5 221. Among the following the strongest nucleophile is a) C₂H₅SH b) CH₃COO⁻ c) CH₃NH₂ d) NCCH₂ 222. Which of the following statements is correct? a) +I group stabilises a carbocation b) +I group stabilises a carbanion c) -I group stabilises a carbocation d) -I group stabilises a free radical 223. Which of the following species does not exert a resonance effect? c) C₆H₅OH a) C₆H₅NH₂ d) C₆H₅Cl b) $C_6H_5NH_3$ 224. The number of different amines corresponding to the formula C₃H₉N is: b) 3 d) 5 c) 4 225. Lactic acid molecule has a) One chiral carbon atom b) Two chiral carbon atoms d) asymmetric molecule c) No chiral carbon atom 226. The arrangement of atoms that characterises a particular stereoisomer is called: a) Geometry of isomer b) Configuration c) Conformers d) None of these 227. An alkane forms isomers if the number of carbon atoms is:



a) ≥ 1

b) ≥ 2

c) ≥ 3

 $d) \ge 4$

228. Which of the following statements is correct?

- a) Desmotropism is another name for tautomerism
- b) Allyl carbocation is less stable than isopropyl carbocation
- c) -I effect is exhibited by-NH₃
- d) The formula CH2Cl2 is non-polar
- 229. The IUPAC name of CH₃COCH(CH₃)₂ is
 - a) Isopropylmethyl ketone

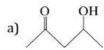
b) 2-methyl-3-butanone

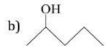
c) 4-methylisopropyl ketone

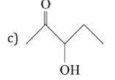
d) 3-methyl-2-butanone

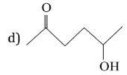
230. Qualitative test of halogens in an organic compound is made by

- a) Fleming's test
- b) Beilstein test
- c) Bayer's test
- d) Fehling's test
- 231. Which one of the following will most readily be dehydrated in acidic conditions?









232. The family to which methoxyethene belongs, is:

- a) Hydrocarbon
- b) Ketone
- c) Unsaturated ether
- d) Ester

233. Electrophiles are:

- a) Electron loving species
- b) Electron hating species
- c) Nucleus loving reagents
- d) Nucleus hating reagents
- 234. Iso-propyl chloride undergoes hydrolysis by
 - a) S_N1 mechanism

b) S_N 2 mechanisms

c) S_N1and S_N2 mechanisms

- d) Neither S_N1 nor S_N2 mechanism
- 235. The IUPAC name of the compound,

$$CH_3$$
— CH_2 — CH — $CONH_2$ is:

- a) 2-ethylbutanamide
- b) 2-methylbutanamide
- c) 1-amino-2-methylpropane
- d) None of the above
- 236. Carbon and hydrogen are estimated in organic compounds by
 - a) Kjeldalhl's method
- b) Duma's method
- c) Leibig's method
- d) Carius method

237. The compound having highest dipole moment is:

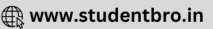
c)
$$H \subset O$$

238. A free radical is:

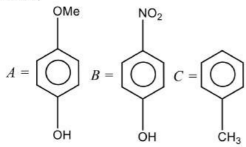
- a) Non-existing
- b) Short lived
- c) Diamagnetic
- d) Fairly stable

239. In 2-methyl-l-propanol, the hybrid carbons of sp^3 , sp^2 and sp are respectively:

- a) 3, 2, 1
- b) 4, 3, 0
- c) 4, 0, 0
- d) 1, 2, 3
- 240. In electrophilic aromatic substitution reaction, the nitro group is meta directing because it



- a) Decreases electron density at ortho and para positions
- b) Decreases electron density at meta position
- c) Increases electron density at meta postion
- d) Increases electron density at ortho and para positions
- 241. Given,



The decreasing order of the acidic character is

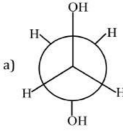
- a) A > B < C
- b) B>A>C
- c) B > C > A
- d) C>B>A

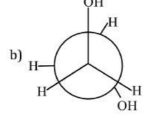
242. Give the IUPAC name for,

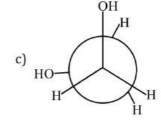
- a) Ethyl-4- oxoheptanoate
- b) Methyl-4- oxoheptanoate
- c) ethyl-4- oxohexanoate
- d) Methyl 4- oxohexanoate
- 243. The total number of acyclic isomers including the stereoisomers with the molecular formula C₄H₇Cl

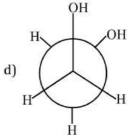
b) 12

- 244. Which of the following conformers for ethylene glycol is most stable?









- 245. Which of the following compounds is resistant to nucleophilic attack by hydroxy ion?
 - a) Methylacetate
- b) Acetonitrile
- c) Acetamide
- d) Diethyl ether

- 246. The stabilization due to resonance is maximum in:
 - a) Cyclohexane
- b) Cyclohexene
- c) 1,3-cyclohexadiene
- d) 1,3,5-cyclohexatriene
- 247. A mixture of camphor and benzoic acid can be easily separated by
 - a) Sublimation

b) Extraction with solvent

c) Fractional crystallisation

- d) Chemical method
- 248. Fractional crystallisations is carried out to separate a mixture of
 - a) Organic solids mixed with inorganic solids
 - b) Organic solids slightly soluble in water
 - c) Organic solids having small difference in their solubilities in suitable solvent
 - d) Organic solids having great difference in their solubilities in suitable solvent
- 249. The type of isomerism observed in urea molecule is:
 - a) Chain
- b) Position
- c) Geometrical
- d) Functional
- 250. Which of the following intermediate have the complete octet around the carbon atom?
 - a) Carbonium ion
- b) Carbanion
- c) Free radical
- d) Carbene

251. The name of, $(CH_3)_2$ HC— O— CH_2 — CH_2 — CH_3 is:





- a) Isopropyl propyl ether
- b) Dipropyl ether
- c) di-isopropyl ether
- d) Isopropyl propyl ketone
- 252. A neutral divalent carbon intermediate produced by the removal of two attached atoms is called:
 - a) Free radical
- b) Carbanion
- c) Carbocation ion
- d) Carbine

- 253. Which types of isomerism is shown by 2, 3-dichlorobutane?
 - a) Structural
- b) Geometric
- c) Optical
- d) Diastereo

254. The correct IUPAC name of the compound,

$$\frac{2}{3}$$
 is

- a) 3-(1-ethyl propyl) hex-1-ene
- b) 4-Ethyl-3-propyl hex-1-ene
- c) 3-Ethyl-4-ethenyl heptane
- d) 3-Ethyl-4-propyl hex-5-ene
- 255. IUPAC name of $(CH_3)_2N C_2H_5$ is:
 - a) Dimethyl ethyl amine
 - b) Dimethylaminomethane
 - c) Dimethylaminoethane
 - d) N, N-dimethylethanamine
- 256. Among the following compounds, the most acidic is
 - a) p-nitrophenol

b) p-hydroxybenzoic acid

c) o-hydroxybenzoic acid

d) p-toluic acid

- 257. Electrophiles are:
 - a) Lewis bases
- b) Lewis acids
- c) Amphoteric
- d) None of these
- 258. On monochlorination of n-pentane, the number of isomers formed is :
 - a) 4

b) 3

c) 2

d) 1

- 259. Cyclohexane is:
 - a) Aliphatic compound
 - b) Alicyclic compound
 - c) Aromatic compound
 - d) Heterocyclic compound
- 260. Which of the following is a primary halide?
 - a) Isopropyl iodide

b) Secondary butyl iodide

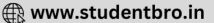
c) Tertiary butyl bromide

- d) Neo hexyl chloride
- 261. The percentage of s' character of the hybrid orbital of carbon in ethane, ethane and ethyne respectively are
 - a) 25, 33, 50
- b) 20, 50, 33
- c) 25, 50, 75
- d) 33, 66, 99

- 262. Which is a chiral molecule?
 - a) CH₃Cl
- b) CH₂Cl₂
- c) CHBr₃
- d) CHClBrI

- 263. The stability of a carbonium ion depends upon
 - a) The bond angle of the attached group
 - b) The substrate with which it reacts
 - c) The inductive effect and hyper-conjugative effect of the attached group
 - d) None of the above
- 264. The IUPAC name of the compound,





- a) 3,4-dimethyl-3-n-propylnonane
- b) 4-ethyl-4,5-dimethyldecane
- c) 6,7-dimethyl-7-n-propylnonane
- d) 6,7-dimethyl-7-ethyldecane
- 265. Bromination of alkanes involves
 - a) Carbanions
- b) Carbocations
- c) Carbenes
- d) Free radicals
- 266. The isomeric cis-2-butene and trans-2-butene can be distinguished on the basis of :
 - a) Their physical nature
 - b) Their reduction products
 - c) The products they give on ozonolysis
 - d) The products they give on addition to bromine
- 267. Lassaigne's test is not used for the detection of
 - a) Carbon
- b) Halogens
- c) Nitrogen
- d) Sulphur

268. Consider the following carbocations,

(I)
$$C_6H_5CH_2$$
 (II) $C_6H_5CH_2CH_2$

(III)
$$C_6H_5\overset{+}{C}HCH_3$$
 (IV) $C_6H_5\overset{+}{C}(CH_3)_2$

- b) II<III<IV
- c) III<I<II<IV
- d) IV<III<I<II
- 269. The simplest formula of a compound containing 50% of element X (at. wt 10) and 50% of element Y (at. wt. 20) is
 - a) XY

b) XY_2

c) X_2Y

- d) X_2Y_2
- 270. n-pentane, iso-pentane, and neo pentane are examples for isomers of the type
 - a) Geometrical
- b) Optical
- c) Chain
- d) Positional
- 271. Homolytic fission of C—C bond in ethane gives an intermediate in which carbon ishybridized.
 - a) sp^3

b) sp^2

c) sp

- d) sp^2d
- 272. Pick out the correct statement from the following and choose the correct answer from the codes given
 - I. Hexa-1, 5-diene is a conjugated diene
 - II. Prop-1, 2-diene is conjugated diene
 - III. Hexa-1, 3-diene is a conjugated diene
 - IV. Buta-1, 3-diene is an isolated diene
 - V. Prop-1, 2-diene is a cumulative diene
 - a) I,II

- b) II, III
- c) IV, V
- d) II, V

273. The IUPAC name of

$$\begin{array}{c|c} & \operatorname{CH_3} \\ | \\ \operatorname{CH_3} - \operatorname{CH} - \operatorname{CH_2} - \operatorname{C} - \operatorname{CH_3} \\ | & | \\ \operatorname{OH} & \operatorname{OH} \end{array}$$

a) 1, 1-dimethyl-1, 3-butanediol

b) 2-methyl-2, 4-pentanediol

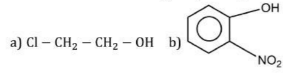
c) 4-methyl-2, 4-pentanediol

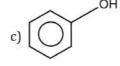
- d) l, 3, 3-trimethyl-1, 3-propane diol
- 274. Which among the following is the correct IUPAC name of isoamylene?
 - a) 1-pentene
- b) 2-methyl-2-butene
- c) 3-methyl-1-butene
- d) 2-mythyl-1-butene
- 275. Which of the following compounds exhibits geometrical isomerism?
 - a) C₂H₅Br
- b) $(CH)_2(COOH)_2$
- c) CH₃CHO
- d) $(CH_2)_2(COOH)_2$

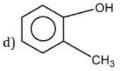




276. Which one of the following compounds, is most acidic?







- 277. An organic compound has carbon and hydrogen percentage in the ratio 6:1 and carbon and oxygen percentages in the ratio 3:4. The compound has the empirical formula
 - a) C₂H₆O
- b) CHO₂

- 278. Among the following the one which does not exhibit functional group isomerism is:
 - a) C_2H_6O
- b) C₃H₈O
- c) C₄H₁₀
- d) C₄H₁₀O

- 279. How many structural formulae are possible for C₅H₁₁Cl?
 - a) 6

b) 8

c) 10

d) 12

- 280. Which one of the following is a secondary alcohol?
 - a) 2-methyl-1-propanol b) 2-methyl-2-propanol c) 2-butanol
- d) l-butanol
- 281. Among the following anions (a) \overline{CH}_3 (b) \overline{NH}_2 , (c) OH^- , (d) F^- the order of basicity is:
 - a) a > b > c > d
- b) b > a > c > d
- c) c > b > a > d
- d) c > a > b > d

- 282. Electromeric effect is
 - a) Permanent effect
- b) Temporary effect
- c) Resonance effect
- d) Inductive effect
- 283. In the following groups, $-OAC OMe OSO_2Me OSO_2CF_3$ the order of leaving group ability is :
 - a) I > II > III > IV
- I < I < III < VI (d
- c) III > II > IV
- d) II > III > IV > I
- 284. The angle of rotation of plane of polarized light depends upon:
 - a) The nature of the light beam
 - b) The number of the molecules
 - c) The number of asymmetric carbon atoms in the molecule of the substance

b) *Tert* –butylchloride

- d) All of the above
- 285. Which of the following shows geometrical isomerism?
 - a) C_2H_5Br
- b) (CH₂)(COOH)₂
- c) $(CH)_2(COOH)_2$
- d) C_2H_6
- 286. Which of the following cannot undergo nucleophilic substitution under ordinary conditions?
 - a) Chlorobenzene
- c) Isopropyl chloride
- d) None of these
- 287. The C—C bond length of the following molecules is in the order.
 - a) $C_2H_6 > C_2H_4 > C_6H_6 > C_2H_2$
 - b) $C_2H_2 < C_2H_4 < C_6H_6 < C_2H_6$
 - c) $C_6H_6 > C_2H_2 > C_2H_6 > C_2H_4$
 - d) $C_2H_4 > C_2H_6 > C_2H_2 > C_6H_6$
- 288. Isomerism among compounds due to the migration of a proton is known as:
- a) Geometrical
- b) Optical
- c) Tautomerism
- d) Position

- 289. Removal of hydrogen atom is easier when it is attached to:
 - a) 1° carbon
- b) 2° carbon
- c) 3° carbon
- d) Same in all

- 290. The order of stability of carbanions is:
 - a) $CH_3^- > 1^\circ > 2^\circ > 3^\circ$
- b) $3^{\circ} > 2^{\circ} > 1^{\circ} > CH_3^-$
- c) $3^{\circ} > 1^{\circ} > 2^{\circ} > CH_{3}^{-}$
- d) $2^{\circ} > 3^{\circ} > 1^{\circ} > CH_3^-$

- 291. Glycerine contains
 - a) 1° carbon

b) 2° carbon

c) 3° carbon

- d) Both 1° and 2° carbon
- 292. Which of the following pairs of carbon skeletons in an example of isomerism?

a)
$$C - C - C - C$$
 and $C - C - C$



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c)
$$C - C - C$$
 and $C - C - C$

d)
$$C-C-C-C$$
 and $C-C-C$

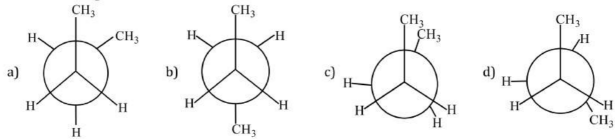
293. In cannizzaro reaction given below

2PhCHO
$$\xrightarrow{\text{OH}^{\Theta}}$$
 PhCH₂OH + PhCO₂ $\xrightarrow{\Theta}$

The slowest step is

- a) The attack of :: OH^{\odot} at the carboxyl group
- b) The transfer of hydride to the carbonyl group
- c) The abstraction of proton from the carboxylic group
- d) The deprotonation of PhCH2OH

294. In the following the most stable conformation of n-butane is :



- 295. $S_N 1$ mechanism for the reaction, $R X + KOH \rightarrow ROH + KX$ follow:
 - a) Carbocation mechanism
 - b) Carbanion mechanism
 - c) Free radical mechanism
 - d) Either of the above
- 296. An electrophilic reagent must have
 - a) A vacant orbital

- b) An orbital containing one electron
- c) An orbital containing two electrons
- d) All completely filled atomic orbitals
- 297. In which of the following structures the number of sigma bonds are equal to the number of π -bonds?
 - a) 1,2-propadiene
- b) 2,3-dicyanobut-2-ene c) Tetracyanoethylene
- d) None of these

298. Which one of the starred carbons is the asymmetric one?

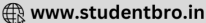
- b) CH3CH2CH2ČH2CH2OH
- c) CH₃ČH₂CH₂CH₂CH₂OH
- d) CH3CH2CH2CH2CH2CH
- 299. The chemical name of anisole is
 - a) Ethanoic acid
- b) Methoxy benzene
- c) Propanone
- d) Acetone
- 300. How many optically active stereomers are possible for butan-2, 3-diol?
 - a) 1

b) 2

c) 3

d) 4





- 301. Naphthalene molecule contains:
 - a) 10π -electrons
- b) 8π -electrons
- c) 12π -electrons
- d) 14π -electrons
- 302. The first organic compound urea was synthesized in the laboratory by:
 - a) Kekule
- b) Liebig
- c) Lavoisier
- d) Wöhler

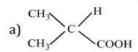
303. In the hydrocarbon

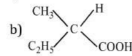
$$CH_3 - CH = CH - CH_2 - CE + CH_3 - CH_3 -$$

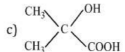
The state of hybridization of carbons 1, 3 and 5 are in the following sequence:

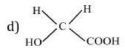
- a) sp, sp^3, sp^2
- b) sp, sp^2, sp^3
- c) sp^3 , sp^2 , sp
- d) sp^2 , sp, sp^3

- 304. Which of the following is not a nucleophile?
 - a) BF₃
- b) NH₃
- c) CN-
- d) OH-
- 305. Which of the following compounds can exhibit optical isomerism?









- 306. Which is the most stable carbocation?
 - a) iso-propyl cation

b) Triphenylmethyl cation

c) Ethyl cation

- d) n-propyl cation
- 307. The correct structure of 4-bromo-3-methyl-but-1-ene.
 - a) Br CH = C(CH₃)₂

b) $CH_2 = CH - CH(CH_3) - CH_2Br$

c) $CH_2 = C(CH_3)CH_2CH_2Br$

d) $CH_3 - C(CH_3) = CHCH_2 - Br$

- 308. IUPAC name of C6H5COCl is:
 - a) Benzoyl chloride
 - b) Benzenechloro ketone
 - c) Benzene carbonyl chloride
 - d) Chloro phenyl ketone
- 309. Stability order of ... is in order

$$C_6H_5$$
— $\overset{+}{\text{CH}}_2$, CH_2 = $\overset{+}{\text{CH}}$ — $\overset{+}{\text{CH}}_2$, $(CH_3)_3C^+$, CH_2 = $\overset{+}{\text{CH}}$ H (II) (IV)

- a) IV<III<II<I
- b) IV<II<I<III
- c) I<II<III<IV
- d) IV<I<III<II
- 310. Relative stabilities of the following carbocations will be in the order
 - \oplus
- Ð (
- CH₃ CH₂ CH₂ OCH₃
- \boldsymbol{A}

a) C>B>A

- B
- b) C<B<A
- c) B>C>A
- d) C>A>B

- 311. Which method is used to separate sugars?
 - a) Fractional crystallisation
- b) Sublimation
- c) Chromatography

- d) Benedict's reagent
- 312. Sublimation can't be used for purification of
 - a) Benzoic acid
- b) Camphor
- c) Urea
- d) Naphthalene

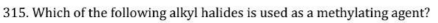
- 313. Which of the following is phenyl ethanoate?

b) COO2COOH

с) Соонз

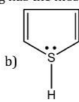
- 314. Zero inductive effect is shown by:
 - a) C_6H_5 —
- b) H
- c) CH3-
- d) Cl—

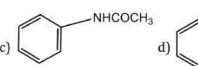


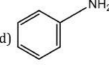


- b) C₂H₅Br
- d) CH₃I
- 316. Which one of the following has the most nucleophilic nitrogen?









317. Chlorobenzene is 0, p-directing in electrophilic substituting reaction. The directing influence is explained by

- a) +M of Ph
- b) +I of Cl
- c) +M of Cl
- d) -I of Ph

- a) 3°<2°<1°
- b) 3°>2°>1°
- c) 1°<2°>3°
- d) 3°>2°<1°

- a) Its planar structure
- b) Its regular tetrahedral nature
- c) Similar sizes of carbon and chlorine atoms
- d) Similar electron affinities of carbon and chlorine
- 320. IUPAC name of the compound

$$\begin{array}{c} \operatorname{CH_3} \\ \operatorname{CH_3} \\ \operatorname{CH_2-CH-CH_2-CH-CH_2-CH_2-CH_3} \\ \\ \operatorname{H_3C} \\ \end{array}$$

a) 4-isoprophyl, 6-methyl octane

b) 3-methyl, 5-(1-methylethyl) octane

c) 3-methyl, 5-isopropyl octane

- d) 6-methyl, 4-(1-methylethyl) octane
- 321. The isomers which are interconverted through rotation around a single bond are
 - a) Conformers
- b) Diastereomers
- c) Enantiomers
- d) Position isomers

- 322. The number of optical isomers of pent-3-en-2-ol is:

b) 4

c) 8

d) 16

323. Dehydrohalogenation of an alkyl halide is a/an

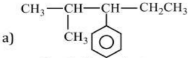
- a) Nucleophilic substitution reaction
- b) Elimination reaction
- c) Both nucleophilic substitution and elimination reaction
- d) Rearrangement
- 324. The minimum number of carbon atoms which a ketone may contain is :

b) 2

c) 3

d) 4

325. Which nomenclature is not according to IUPAC system?



2-methyl-3-phenylpentane

5-oxohexanoic acid

d)
$$CH_3$$
 CH_3 CH_3 CH_2 CH_2 CH_3 CH_3 CH_3 CH_3

- 326. What is the formula of tertiary butyl alcohol?
 - a) $CH_3 CH(CH_3) CH_2 OH$

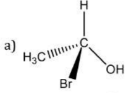
b) $CH_3 - (CH_2)_2OH$

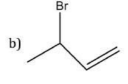
c) $CH_3 - CH(OH) - CH_2 - CH_3$

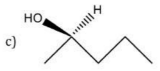
- d) $(CH_3)_3$. C OH
- 327. The IUPAC name of neopentane is
 - a) 2-methylbutane
- b) 2,2-dimethylpropane
- c) 2-methylpropane
- d) 2,2-dimethyl butane

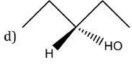
- 328. Select the strongest bond:
 - a) → C C ←
- b) >C=C<

329. Which of the following molecules is achiral?







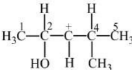


- 330. An enantiomerically pure acid is treated with racemic mixture of an alcohol having one chiral carbon. The ester formed will be
 - a) Optically active mixture

b) Pure enantiomer

c) Meso compound

- d) Racemic mixture
- 331. In the following carbocation, H/CH₃ that is most likely to migrate to the positively charged carbon is:



- a) CH₃ at C-4
- b) H at C-4
- c) CH₃ at C-2
- d) Hat C-2

- 332. The number of π -electrons in benzene molecule is :
 - a) 3×2
- b) 2^{3}

- c) 3×3
- 333. Which of the following statements is necessarily true in the case of isomeric organic compounds?
 - a) They are hydrocarbons
 - b) They are optically active
 - c) They yield the same products on complete combustion
 - d) They have same melting or boiling points
- 334. Cis trans, isomers generally
 - a) Contain an asymmetric carbon atom
- b) Rotate the plane of polarized light

c) Are enantiomorphs

- d) Contain a double bonded carbon atoms
- 335. Among the following compounds nitrobenzene, benzene, aniline and phenol, the strongest basic behaviour in acid medium is exhibited by:
 - a) Phenol
- b) Aniline
- c) Nitrobenzene
- d) Benzene

- 336. Geometrical isomerism is not shown by :
- Geometrica:

 CH₃

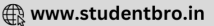
 a) CH₃CH₂C=CCH₂CH₃

 CH₃

 CH₃

 - c) $CH_2 = C(Cl)CH_3$
 - d) CH_3 — $CH = CH_2$ — $CH = CH_2$
- 337. The correct stability order for the following species as





a) II>IV>I>III

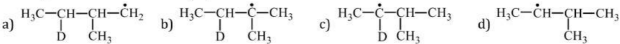
- b) I>II>III>IV
- c) II>I>IV>III
- d) I>III>IV

338. An optically active compound is:

- a) 1-bromobutane
- b) 2-bromobutane
- c) 1-bromo-2-methyl propane
- d) 2-bromo-2-methyl propane
- 339. Consider the following reaction,

$$\begin{array}{ccc} \mathbf{H_{3}C-CH-CH-CH_{3}+\mathring{B}r} & & \mathbf{\dot{B}r} & \mathbf{\dot{Y}'+HBr} \\ \mathbf{\dot{D}} & \mathbf{\dot{C}H_{3}} & & & \end{array}$$

Identify the structure of the major product X':



340. In Lassaigen's test, the organic compound is fused with a piece of sodium metal in order to

- a) Increase the ionization of the compound
- b) Decrease the melting point of the compound
- c) Increase the reactivity of the compound
- d) Convert the covalent compound into a mixture of ionic compounds

341. Which of the following sodium compound/compounds are formed when an organic compound containing both nitrogen and sulphur is fused with sodium?

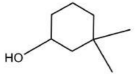
a) Cyanide and sulphide

b) Thiocyanate

c) Sulphite and cyanide

- d) Nitrate and sulphide
- 342. The IUPAC name of $CH_3 CH = CH C \equiv CH$ is
 - a) Pent-3-en-1-yne
- b) Pent-3-en-4-yne
- c) Pent-2-en-4-yne
- d) Pent-2-en-3-yne
- 343. The compound having molecular formula C₄H₁₀O can show:
 - a) Metamerism
- b) Functional isomerism c) Positional isomerism
- d) All of these

344. The IUPAC name of the compound



- a) 3, 3-dimethyl-1-hydroxy cyclohexane
- b) 1, 1-dimethyl-3- hydroxy cyclohexane

c) 3, 3- dimethy-1- cyclohexanol

- d) 1,1-dimethyl-3-cyclohexanol
- 345. In hyperconjugation, the atom involved is:
 - a) β-H atom
- b) α-H atom
- c) γ H atom
- d) All of these
- 346. Reactivity of hydrogen atoms attached to different atoms in alkanes has the order:
 - a) $3^{\circ} > 1^{\circ} > 2^{\circ}$
- b) $1^{\circ} > 2^{\circ} > 3^{\circ}$
- c) $3^{\circ} > 2^{\circ} > 1^{\circ}$
- d) None of these

- 347. Which has maximum percentage of chlorine?
 - a) Pyrene
- b) PVC
- c) Chloral
- d) Ethylidene chloride

- 348. $H_2C = 0$ behaves as:
 - a) Nucleophile
- b) Electrophile
- c) Both (a) and (b)
- d) None of these

- 349. The most stable carbocation is:
- b) CH₃CH₂
- c) (CH₃)₂CH
- d) (CH₃)₃C





The above reaction proceeds through

a) Free radicals substitution

b) Nucleophilic substitution

c) Electrophilic substitution

d) None of the above

351. Which reaction sequence would be best to prepare 3-chloro-aniline from benzene?

- a) Chlorination, nitration, reduction
- b) Nitration, chlorination, reduction
- c) Nitration, reduction, chlorination
- d) Nitration, reduction, acylation, chlorination, hydrolysis

352. Why is light necessary to bring in chlorination reactions of alkane?

- a) The dissociation of Cl2 gives Cl free radical
- b) The Cl₂ molecule absorbs light to show hemolytic bond fission
- c) The formation of Čl free radical propagate the chain reaction
- d) All of the above
- 353. IUPAC name of

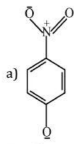
- a) 4-bromo-3-ethyl-1,4-pentadiene
- b) 2-bromo-3-ethyl-1,4pentadiene
- c) 2-bromo-3-ethyl-1-5-pentadiene
- d) None of the above
- 354. The product of reaction,

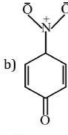
$$CH_3$$
 CH_3
 CH_2
 CH_2
 CH_3
 CH_3

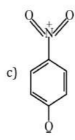
product is:

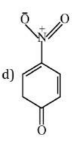
355. The most unlikely representation of resonance structures of p-nitrophenoxide ion is:











- 356. For all practical purposes, influence of inductive effect is neglected after:
 - a) 2nd carbon atom
- b) 1st carbon atom
- c) 3rd carbon atom
- d) None of these

357. CH₃CHCH₂CH₃Cl₂hv C₅H₁₁CI

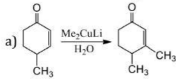
N (isomeric produces) $\xrightarrow{\text{Fractional}} M$ (isomeric product)

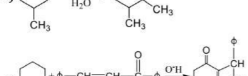
what are the no. of N and M?

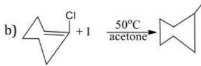
- a) 6, 6
- b) 6, 4
- c) 4, 4

d) 3, 3

- 358. Geometrical isomers differ in:
 - a) Position of functional groups
 - b) Position of atoms
 - c) Spatial arrangement of atoms
 - d) Length of carbon chain
- 359. Which of the following is an example of substitution reaction?







d) None of the above

360. The study of three dimensional structure of molecule is called:

- a) Stereochemistry
- b) Solid state chemistry
- c) Chirality
- d) None of these
- 361. Orbital interaction between the σ -bonds of a substituent group and a neighbouring π -orbital is known as
 - a) Hyperconjugation

b) Inductive effect

c) Steric effect

d) Electric quadrapole interactions

362. The shape of \overline{C}_{H_3} is:

- a) Linear
- b) Planar
- c) Pyramidal
- d) None of these

363. Which of the following contains only three pairs of electrons?

- a) Carbocation
- b) Carbanion
- c) Free radical
- d) None of these
- 364. 2-hexyne gives trans-2-hexene on treatment with
 - a) Li/NH₃
- b) Pd/BaSO₄
- c) LiAlH₄
- d) Pt/H₂

365. Lassaigne's test for the detection of nitrogen fails in

a) $H_2N - CO - NHNH_2$. HCl

- b) NH₂ NH₂. HCl

c) $NH_2 - CO - NH_2$

d) $C_6H_5 - NH - NH_2$. HCl

Which of the following compounds yields most stable carbanion after rupture $\begin{pmatrix} \text{C--C} \end{pmatrix}$ of bond?



a)
$$_{\text{CH}_3}$$
— $_{1}^{\text{C}}$ — $_{2}^{\text{C}}$ HI $_{2}$ b) $_{\text{CH}_3}$ — $_{1}^{\text{C}}$ — $_{2}^{\text{C}}$ — $_{1}^{\text{C}}$ — $_{2}^{\text{C}}$ H $_{2}$ I

d) None of these

367. The IUPAC name of $C_2H_5 - o - CH$

$$C_2H_5$$
—O—CH $\overset{\text{CH}_3}{\underset{\text{is}}{\smile}}$

- a) Ethoxy propane
- b) 1, 1-dimethyl ether
- c) 2-ethoxy iso-propane d) 2-ethoxy propane
- 368. The relative adsorption of each component of the mixture is expressed in terms of
 - a) adsorption factor

b) retention factor

c) co-factor

d) sorption factor

369. Following reaction,

$$(CH_3)_3CBr + H_2O \rightarrow (CH_3)_3COH + HBr$$

is an example of

a) Elimination reaction

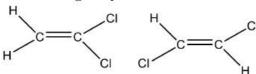
b) Free radical substitution

c) Nucleophilic substitution

- d) Electrophilic substitution
- 370. The most stable carbonium ion among the following is

- 371. t-butyl alcohol is
 - a) 2-methyl propane-2-ol
 - c) 3-methyl butan-1-ol

- b) 2-methyl propane-1-ol
- d) 3-methyl butan-2-ol
- 372. The following compound differ in



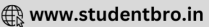
- a) Configuration
- b) Conformation
- c) Structure
- d) Chirality

- 373. A compound containing 80% C and 20% H is likely to be
 - a) C_6H_6
- b) C_2H_6
- c) C₂H₄
- d) C_2H_2
- 374. Overlap of which of the following atomic orbitals would be maximum to form the strongest covalent bond?
 - a) $1s 2s(\sigma)$
- b) $1s 2p(\sigma)$
- c) $2p 2p(\pi)$
- d) $2p 2p(\sigma)$

- 375. A strong base can abstract an α -hydrogen from :
 - a) Amine
- b) Ketone
- c) Alkane
- d) Alkene
- 376. During elimination reactions, the hybrid state of carbon atoms involved in change shows:
 - a) sp^3 to sp^2 nature
 - b) sp^2 to sp nature
 - c) No change in hybridized state
 - d) Either of the above
- 377. IUPAC name of C₆H₅CN is:
 - a) Phenyl nitrile
- b) Benzene nitrile
- c) Benzyl nitrile
- d) Phenyl cyanide
- 378. Who proposed the tetrahedral mirror image structures to a pair of enantiomers?
 - a) Kekule
- b) Wöhler
- c) van't Hoff
- d) None of these

- 379. The S_N1 reactivity of following halides will be in the order
 - (i) $(CH_3)_3CBr$
- $(ii)(C_6H_5)_2CHBr$
- $(iii)(C_6H_5)_2C(CH_3)Br(iv)(CH_3)_2CHBr$
- $(v) C_2 H_5 Br$
- a) (v)>(iv)>(i)>(ii)>(iii)

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



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

d) (iii)>(ii)>(iv)>(v)

380. Heterolysis of propane gives:

- a) Methyl and ethyl free radicals
- b) Methylium cation and ethyl anion
- c) Methyl anion and ethylium cation
- d) Methylium and ethylium cations
- 381. Delocalised electrons are present in
 - a) 1, 3- butadiene
- b) C_6H_6
- c) 1, 3, 5-hexatriene
- d) All of these

382. Compounds which rotate plane polarised light in clockwise direction are known as:

- a) Dextrorotatory
- b) Laevorotatory
- c) Optically inactive compounds
- d) Racemic

383. Carbanions initiate:

- a) Addition reactions
- b) Substitution reactions
- c) Both (a) and (b)
- d) None of these

384. Impure glycerine can be purified by

a) Steam distillation

b) Simple distillation

c) Vaccum distillation

d) Extraction with a solvent

385. IUPAC name of urea is:

- a) Diaminoketone
- b) 1-aminoethanamide
- c) 1-aminomethanamide
- d) aminoacetamide

386. Which of the following process is not used for the purification of solid impurities?

- a) Distillation
- b) Sublimation
- c) Crystallisation

387. When the hybridization state of a carbon atom changes from sp^3 to sp^2 and finally to sp, the angle between the hybridized orbitals:

- a) Is not affected
- b) Increases progressively
- c) Decreases considerably
- d) Decreases gradually

388. The chief reaction product of reaction in between n-butane and bromine at 130°C is:

389. Dehydration of alcohol is an example of which type of reaction?

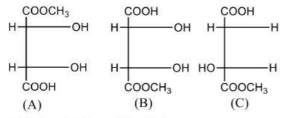
- a) Substitution
- b) Elimination
- c) Addition
- d) Rearrangement

390. The IUPAC name of

$$\begin{array}{ccc} \operatorname{CH_3--CH--CH=-C--CHO} \\ | & | \\ \operatorname{OH} & \operatorname{CH_3} \end{array}$$

- a) 4-hydroxy-1-methylpentanal
- b) 4-hydroxy-4-methylpent-2-en-1-al
- c) 2-hydroxy-4-methylpent-2-en-5-al
- d) 2-hydroxy-3-methylpent-2-en-5-al
- 391. The correct statement about the compounds (A), (B) and (C) is



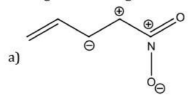


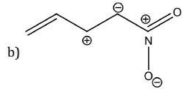
a) (A) and (B) are identical

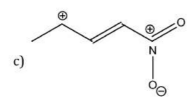
b) (A) and (B) are diastereomers

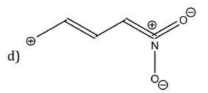
c) (A) and (C) are enantiomers

- d) (A) and (B) are enantiomers
- 392. Among the following the least stable resonance structure is







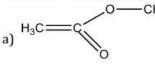


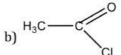
- 393. The organic liquid that mix freely with water is:
 - a) CHCl₃
- b) CCl₄
- c) CS₂

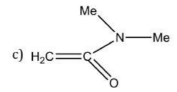
- d) C2H5OH
- 394. The increasing order of +ve I-effect shown by H, CH_3 , C_2H_5 and C_3H_7 is:
 - a) $H < CH_3 < C_2H_5 < C_3H_7$
 - b) $H > CH_3 < C_2H_5 > C_3H_7$
 - c) $H < C_2H_5 < CH_3 < C_3H_7$
 - d) None of the above
- 395. The best method for the separation of naphthalene and benzoic acid from their mixture is
 - a) Chromatography
- b) Crystallisation
- c) Distillation
- d) Sublimation

- 396. The reagent used in dehydrohalogenation process is:
 - a) Alcoholic KOH
- b) NaNH₂
- c) C₂H₅ONa
- d) All of these

397. The least active electrophile is

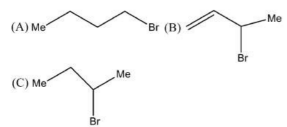






- H_3C C H_3C S
- 398. The isomerism which exists between CH3CHCl2 and CH2ClCH2Cl is:
 - a) Chain
- b) Functional
- c) Positional
- d) Metamerism

399. Consider the following bromides



The correct order is S_N1 reactivity is

- a) (B)>(C)>(A)
- b) (B)>(A)>(C)
- c) (C)>(B)>(A)
- d) (A)>(B)>(C)

400. The appropriate reagent for the following transformation,

- a) Zn(Hg), HCl
- b) NH2NH2, OH-
- c) H₂/Ni
- d) NaBH₄
- 401. Hydride shift from C-2 will give the most stable resonance stabilized carbocation as
 - a) CH_3 at C-4
- b) H at C 4
- c) CH_3 at C-2
- d) Hat C-2

- 402. Hyperconjugation involves overlap of the following orbitals

- b) $\sigma \rho$
- d) $\pi \pi$

- 403. Most stable carbonium ion is
 - a) C₂H₅
- b) (CH₃)₃C
- c) $(C_6H_5)_3 \overset{+}{C}$
- d) C₆H₅CH₂

- 404. During a nitration of benzene, the attacking electrophile is
- b) NO₂
- d) HNO₃
- 405. The (R) and (S) enantiomers of an optically active compound differ in
 - a) Their reactivity

b) Their optical rotation of plane polarised light

c) Their melting point

- d) Their solubility in achiral reagents
- 406. The number of chiral centres in (+) -glucose

c) 2

d) 1

- 407. Hydrogen cyanide and hydrogen isocyanide are:
 - a) Tautomers
 - b) Positional isomers
 - c) Metamers
 - d) Chain isomers
- 408. Which of the following hydrocarbons is most unsaturated?
 - a) C2H4
- b) C_2H_2
- c) C_2H_6
- d) $CH_3CH = CH_2$
- 409. Sometimes the behaviour of a compound is explained by assuming that it exists in a world between two or more different possible structures. This phenomenon is called:
 - a) Isomerism
- b) Resonance
- c) Mutarotation
- d) Allotropism
- 410. How many primary amines are possible with the formula C₄H₁₁N?

b) 2

c) 3

- d) 4
- 411. Which one of the following pairs represents stereoisomerism?
 - a) Geometrical isomerism, position isomerism
 - b) Geometrical isomerism, conformational isomerism
 - c) Optical isomerism, geometrical isomerism
 - d) Optical isomerism, metamerism





412. The large number of organic compounds is due to:			
a) Catenation property of carbon			
b) Covalent bond formation			
c) Isomerism			
d) polymerization			
413. The IUPAC name of $CH_3 - C - CH - CH_3$ is			
Ⅱ │ O CH ₃			
O CH ₃			
a) 2-methyl-3-butanone	b) 3-methyl-butan-2-one		
c) 3-methyl butanone	d) None of these		
414. Formic acid is a stronger acid than acetic acid. This	can be explained using		
a) $+M$ effect b) $-I$ effect	c) + <i>I</i> effect	d) – <i>M</i> effect	
415. The energy of C—C triple bond in acetylene in kcal			
a) 140 b) 192	c) 60	d) 100	
416. In which of the following molecules, the resonance	effect is not present?	80	
a) $\left(\begin{array}{c} \\ \\ \end{array}\right)$ \left			
a) NH ₂ b) NH ₃	c) ()—OH	d) 《	
417. Which of the following represents the given mode of	of hybridization $sn^2 - sn^2 -$	- sn - sn from left to right?	
		100 N	
a) $CH_2 = CH - C \equiv CH$ b) $HC \equiv C - C \equiv N$	c) $CH_2 = C - C = CH_2$	d) CH2	
418. The Lassaigne's extract is boiled with dil HNO_3 before	ore testing for halogens beca	ause	
a) AGCN is soluble in HNO ₃	b) Silver halides are solu	ble in HNO ₃	
c) Na ₂ S and NaCN are decomposed by HNO ₃	d) Ag ₂ S is soluble in HNC)3	
419. The tautomeric form which is less stable is called :			
a) Anion form b) Cation form	c) Labile form	d) All of these	
420. The effect involving the complete transfer of a sh	ared pair of electrons to o	ne of the atoms joined by a	
multiple bond at the requirement of attacking reage	ent is called :		
a) Inductive effect b) Mesomeric effect	c) Electromeric effect	d) None of these	
421. Which of the following acids does not exhibit optical	l isomerism?		
a) Lactic acid b) Tartaric acid	c) Maleic acid	d) α-amino acids	
422. Many organic compounds are prepared by using PC	l ₅ because :		
a) OH group of alcohol is easily replaced by chloring	e atom		
b) Chlorines are added to the unsaturated compour	nds		
c) It removes water from organic compounds			
d) Phosphorus atoms are entered in the alcohol			
423. Which of the following conformations of cyclohexar	ne is chiral?		
a) Twist boat b) Rigid	c) Chair	d) Boat	
424. Which type of isomerism is shown by propanal and	propanone?		
a) Functional group b) Metamerism	:#2		
dy runetional group	c) Tautomerism	d) Chain isomerism	
425. Identify the product in the given reaction:	. 경영	d) Chain isomerism	
	. 경영	d) Chain isomerism	
425. Identify the product in the given reaction:	. 경영	d) Chain isomerismd) CH₂(NO). CH₂. CH₂Cl	
425. Identify the product in the given reaction: $CH_3 - CH = CH_2 + NOCl \rightarrow Product$ a) $CH_3CHCl. CH_2. NO$ b) $CH_3CH(NO). CH_2Cl$ 426. A straight chain hydrocarbon has the molecular for	c) Tautomerism c) $CH_3CH_2CH(Cl)(NO)$ mula C_8H_{10} . The hybridizat	d) CH ₂ (NO). CH ₂ . CH ₂ Cl tion for the carbon atoms	
 425. Identify the product in the given reaction: CH₃ − CH = CH₂ + NOCl → Product a) CH₃CHCl. CH₂. NO b) CH₃CH(NO). CH₂Cl 426. A straight chain hydrocarbon has the molecular for from one end of the chain to the other are respective. 	c) Tautomerism c) $CH_3CH_2CH(Cl)(NO)$ mula C_8H_{10} . The hybridizat	d) CH ₂ (NO). CH ₂ . CH ₂ Cl tion for the carbon atoms	
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 425. Identify the product in the given reaction: CH₃ - CH = CH₂ + NOCl → Product a) CH₃CHCl. CH₂. NO b) CH₃CH(NO). CH₂Cl 426. A straight chain hydrocarbon has the molecular for from one end of the chain to the other are respective formula of the hydrocarbon would be: a) CH₃ - C ≡ C - CH₂ - CH = CH - CH = CH₂ 	c) Tautomerism c) $CH_3CH_2CH(Cl)(NO)$ mula C_8H_{10} . The hybridizat	d) CH ₂ (NO). CH ₂ . CH ₂ Cl tion for the carbon atoms	
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 425. Identify the product in the given reaction: CH₃ - CH = CH₂ + NOCl → Product a) CH₃CHCl. CH₂. NO b) CH₃CH(NO). CH₂Cl 426. A straight chain hydrocarbon has the molecular for from one end of the chain to the other are respective formula of the hydrocarbon would be: a) CH₃ - C ≡ C - CH₂ - CH = CH - CH = CH₂ 	c) Tautomerism c) $CH_3CH_2CH(Cl)(NO)$ mula C_8H_{10} . The hybridizat	d) CH ₂ (NO). CH ₂ . CH ₂ Cl tion for the carbon atoms	

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427. Reaction,

$$R \longrightarrow CO + HCN \longrightarrow R - C \longrightarrow OH \text{ is a } CN$$

- a) Electrophilic substitution
- b) Nucleophilic substitution
- c) Electrophilic addition
- d) Nucleophilic addition
- 428. The total number of acylic iosmers including the stereoisomers (geometrical and optical), with the molecular formula C4H7Cl is
 - a) 12

b) 11

c) 10

- d) 9
- 429. The best method to separate the mixture of ortho -and-para nitrophenol (1:1) is
 - a) Vaporisation
- b) Colour spectrum
- c) Distillation
- d) Crystallisation

- 430. Which of the following does not show electrometric effect?
 - a) Alkenes
- b) Ethers
- c) Aldehyde
- d) Ketones
- 431. Shifting of electrons of a multiple bond under the influence of a reagent is called:
 - a) I-effect
- b) E-effect
- c) M-effect
- d) T-effect
- 432. 0.4 g of a silver salt of a monobasic orgainc acid gave 0.26 g pure silver on ignition. the molecular weight of the acid is (atomic weight of silver=108)

b) 37

c) 89

- d) 105
- 433. The S_N 2 mechanism for, $R X + KOH(aq) \rightarrow R OH + KX$ follows with :
 - a) 100% inversion
- b) 50% inversion

is:

- c) 40% inversion
- d) 30% inversion

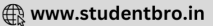
434.

- a) 4,5-dimethyl oct-4-ene
- b) 3,4-dimethyl oct-5-ene
- c) 4,5-dimethyl oct-5-ene
- d) None of the above
- 435. The reaction,

$$\begin{array}{ccc}
O & O & O \\
\parallel & \parallel & \parallel \\
R - C - X + Nu^{-} \rightarrow R - C - Nu + X
\end{array}$$

- is fastest when X is
- a) OCOR
- b) OC₂H₅
- c) NH₂
- d) Cl
- 436. Which pair represents chain isomers?
 - a) CH3CHCl2 and ClCH2CH2Cl
 - b) Propyl alcohol and isopropyl alcohol
 - c) 2-methylbutane and neopentane
 - d) Diethylether and dipropylether
- 437. The empirical formula of an acid is CH₂O₂, the probable molecular formula of the acid may be
- b) C₃H₆O₄
- c) $C_2H_2O_4$
- d) CH_2O_2
- 438. The number of valence electrons in the excited carbon atom is:
 - a) Two in s and two in p-orbitals
 - b) 4 single p-orbitals
 - c) One in s and three in p-orbitals
 - d) None of the above





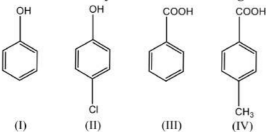
100 11 1	40.5	V. A. A. D. B.	140500
439. A hydrocarbon contains	10.5 g carbon and 1 g hydr	ogen. Its 2.4 g has 1 L volur	ne at 1 atm and127°C.
Hydrocarbon is	L) C II	-) C II	D.C. II
a) C ₆ H ₇	b) C ₆ H ₈	c) C ₅ H ₆	d) C ₆ H ₆
440. The number of stereoiso	mers possible for a compo	und of the molecular formu	$Ia CH_3 - CH = CH -$
CH(OH) — Me is			
a) 3	b) 2	c) 4	d) 6
441. The structural formula o	f methyl aminomethane is	*	
a) $(CH_3)_2CHNH_2$	b) $(CH_3)_3N$	c) $(CH_3)_2NH$	d) CH ₃ NH ₂
442. A mixture of oil and water	er is separated by		
a) Filtration		b) Fractional distillation	
c) Sublimation		d) Using separating funn	el
443. The stability of			
$CH_3 - CH = CH - CH_3, C$	$CH_3 - C = C - CH_3$		
	1 1		
	CH ₃ CH ₃		
(I)	(II)		
$CH_3 - C = CH_2 CH_3 - C$ $CH_3 CH_3 CH_4 CH_4 CH_5 CH_5 $	$C = CH - CH_3$		
CH ₃	CH ₃		
	(IV)		
In the increasing order is			
a) III <i<iv<ii< td=""><td>b) I<ii<iv< td=""><td>c) IV<iii<ii<i< td=""><td>d) II<iii<iv<i< td=""></iii<iv<i<></td></iii<ii<i<></td></ii<iv<></td></i<iv<ii<>	b) I <ii<iv< td=""><td>c) IV<iii<ii<i< td=""><td>d) II<iii<iv<i< td=""></iii<iv<i<></td></iii<ii<i<></td></ii<iv<>	c) IV <iii<ii<i< td=""><td>d) II<iii<iv<i< td=""></iii<iv<i<></td></iii<ii<i<>	d) II <iii<iv<i< td=""></iii<iv<i<>
444. The number of optically			2
a) 1	b) 3	c) 4	d) 2
445. The nodal plane in the π -		60. 4 0.0000	
a) The molecular plane			
b) A plane parallel to the	molecular plane		
		nich bisects the carbon-carb	oon sigma bond at right
angle	to the molecular plane w	nen bibeeto tire curbon curb	on organia bonia ac rigine
	r to the molecular plane wh	nich contains the carbon-ca	rhon sigma hond
446. Which of the following h	and the second s		. Don organia Donia
			D DE 0/E/)
a) CH ₃ OH	b) AlCl ₃	c) (CH ₃) ₄ N $\ddot{0}$ H	d) $BF_3O(Et)_2$
447. Which of the following is	heterocyclic aromatic spe	cies?	
	,,		B.L.
a) 🗸	b) [c) [d) \N
,O,	~	O,	l H
448. A mixture of o-nitrophen	ol and n-nitrophenol can b	ne congrated by	11
a) Fractional crystallisat	and and the contract of the co	b) Sublimation	
	1011	and the contract of the contra	
c) Chemical separation	:tt1 11	d) Steam distillation	
449. The total number of cycli	ic structural as well as ster	o isomers possible for a cor	npound with the molecular
formula C_5H_{10} is	1.3.4	2.6	J) 7
a) 2	b) 4	c) 6	d) 7
450. State the hybridization o	하는 사람들이 있었다. 그리고 아이를 잃었다면 하는 사람이 아니는 아이들이 아이들이 아니는 아이들이 아니다.		D.M. C.I
a) sp^3	b) <i>sp</i> ²	c) sp	d) None of these
451. Which of the following ca			***
a) Alkenes	b) Ketones	c) Aldehydes	d) Ethers
452. Hydride ion transfer take	es place in	\$250000 Te 800	
a) Frankland method		b) Wurts reaction	

c) Cannizzaro's reaction d) Wolff-Kishner reduction 453. An organic compound contains 29.27% carbon, 5.69 % hydrogen and 65.04% bromine. Its empirical formula is d) C₃H₇Br b) C₃H₃Br c) $C_2H_4Br_2$ a) C₃H₅Br 454. OH IUPAC name of is: OH a) but-2-ene-2,3-diol b) pent-2-ene-2,3-diol c) 2-methylbut-2-ene-2,3-diol d) Hex-2-ene-2,3-diol 455. The IUPAC name of $(CH_3)_3C - CH = CH_2is$ a) 1, 1, 1-trimethyl-2-propene b) 3, 3, 3-trimethyl-2-propene c) 2, 2-dimethyl-3-butene d) 3, 3-dimethyl-1-butene 456. The function of soda lime, a mixture of solid NaOH and solid CaO during decarboxylation of carboxylic acids is: a) To increase the rate of reaction b) To decrease the rate of reaction c) To change the rate of reaction d) None of the above is correct 457. t-butyl chloride reacts with OH^- by S_N1 mechanism and rate \propto [t-buty1 chloride]. One of the reasons for this is that a) Stereochemical inversion takes place b) t- buty1 carbocation is first formed which is more stable c) The product t-butyl alcohol is more stable d) The intermediate *t*-butyl carbocation is stabilized by solvation 458. Heterolysis of CH₃CH₂CH₃ result in formation of c) $\bar{C}H_3$ and C_2H_5 a) $\overset{\oplus}{\text{CH}_3}$ and $\overset{\bullet}{\text{C}_2}\text{H}_5$ b) $\overset{\bullet}{\text{CH}_3}$ and $\overset{\bullet}{\text{C}_2}\text{H}_5$ d) CH3and C2H5 459. Alkyl c yanide $R - C \equiv N$ and alkyl isocyanides $R - N \rightarrow C$ are: c) Functional isomers a) Tautomers b) Metamers d) Geometrical isomers 460. A racemic mixture is a mixture of : a) meso and its isomers b) *d*-and its *l*-isomers in equal proportions c) d-and its l-isomers in different proportions d) meso and d-isomers 461. A mixture of iodine and sodium chloride can be easily separated by a) Fractional distillation b) Steam distillation c) Chromatography d) Sublimation 462. The property by virtue of which a compound can turn the plane of polarization of light is known as: b) Phosphorescence a) Photolysis c) Optical activity d) polarization 463. Correct order of stability is a) $HC \equiv \overline{C} > CH_2 = \overline{C}H > CH_3 - \overline{C}H_2$ b) $CH_3 - \overline{C}H_2 > CH_2 = \overline{C}H > CH \equiv \overline{C}$ c) $CH_3 - \overline{C}H_2 > CH \equiv CH \cong CH_2 = \overline{C}H$ d) All are equally stable 464. In the estimation of sulphur in an organic compound, fuming nitric acid is used to convert sulphur into c) H_2SO_3 d) H₂SO₄ 465. The IUPAC name of compound

$$\begin{array}{c} O \\ \parallel \\ CH_2-C-OH \\ \mid OH \quad is: \\ COOH \\ CH_2-COOH \end{array}$$

- a) 1,2,3-tricarboxy-2,1-propane
- b) 3-carboxy-3-hydroxy-1,5-pentanedioic acid
- c) 3-hydroxy-3-carboxy-1,5-pentanedioic acid
- d) None of the above
- 466. Which of the following will be chiral?
 - a) CH₃CHCl₂
- b) CH₃CHBrCl
- c) CD₂Cl₂
- d) CH2ClBr
- 467. In the dehydration reaction $CH_3CONH_2 \xrightarrow{P_2O_5} CH_3C \equiv N$, the hybridization state of carbon change from
 - a) $lsp^3 to sp^2$
- b) lsp to sp
- c) $lsp^2 to sp$
- d) $lsp to sp^3$

468. The correct acidity order of the following is



- a) (III)>(IV)>(II)>(I)
- b) (IV)>(III)>(I)>(II)
- c) (III)>(II)>(IV)
- d) (II)>(III)>(IV)>(I)

- 469. Which of the following is not a nucleophile?
 - a) BF₃
- b) CN-
- c) OH-
- d) NH₃

- 470. Least stable conformer of cyclohexane is
 - a) Chair
- b) Boat
- c) Twist boat
- d) Planar hexagon

- 471. The compound isomeric with acetone is:
 - a) Propionaldehyde
- b) Propionic acid
- c) Ethoxy ethane
- d) None of these

- 472. Which of the chloride is less reactive towards hydrolysis?
 - a) Vinyl chloride
- b) Allyl chloride
- c) Ethyl chloride
- d) t —butyl chloride

- $473. \ Glycerol \ is an alcohol \ which \ can \ be \ classified \ as$
 - a) Trihydric
- b) Monohydric
- c) Dihydric
- d) Hexahydric
- 474. Addition of Br_2 on $CH_2 = CH_2$ in presence of NaCl(aq.) gives :
 - a) CH₂Br. CH₂Br
- b) CH₂Br. CH₂Cl
- c) CH₂Br. CH₂OH
- d) All of these

- 475. The electromeric effect in organic compounds is a:
 - a) Temporary effect
 - b) Permanent effect
 - c) Temporary-permanent effect
 - d) None of the above
- 476. The function of boiling the sodium extract with conc. HNO₃ before testing for halogen is
 - a) To make the solution acidic

b) To make the solution clear
 d) To destroy CN⁻ and S²⁻ ions

- c) To convert Fe2+ to Fe3+
- 477. Copper wire test of halogens is known as
- d) Beilstein's test

- a) Liebig's test b) Lassaigne's test 478. Which of the following is singlet carbine?
 - a) $(CH_3)_3C^+$
- b) C₂H₅ Č—H
- c) CH₃ C HCH₃

c) Fusion test

d) $CH_2 = CH - \dot{C}H_2$

479. Which of the following will be easily nitrated?

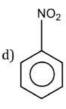








c) CH₃NO₂



480. Optical isomerism is shown by

- a) Propanol-2
- b) Butanol-2
- c) Ethanol

d) Methanol

481. Williamson's synthesis involves

- a) S_N1 mechanism
- c) S_N2 mechanism

- b) Nucleophilic addition
- d) S_E mechanism

482. Free radicals can undergo:

- a) Disproportionation to two species
- b) Rearrangement to a more stable free radical
- c) Decomposition to give another free radical
- d) All of the above are correct

483. During addition of bromine on ethene, the first species formed is

b) C₂H₄OH⁺

c) ⁺_{CH₂CH₂Br}

d) C₂H₅+

484. Metamers of ethyl propionate are

- a) C4H9COOH and HCOOC4H9
- c) CH₃COOCH₃ and CH₃COOC₃H₇
- b) C₄H₉COOH and CH₃COOC₃H₇
- d) CH3COOC3H7 and C3H7COOCH3

485. Which statement is correct about the hybridization of carbon atoms in,

$$H_{C}^{1} = {}^{2}_{C} - {}^{3}_{C} = {}^{4}_{CH}$$
?

- a) C₁ and C₄ are sp²-hybridized
- b) C₂ and C₃ are sp²-hybridized
- c) All are sp-hybridized
- d) All are sp2-hybridized

486. Which one is not is IUPAC system?

a)
$$CH_3$$
— CH — CH — CH_3 (3-methyl-2-butanol) OH CH_3

b)
$$CH_3CH_2CH_2$$
— CH — CH — CH_2CH_3
 CH_2CH_3
 CH_2CH_3
 CH_2CH_3

.......

d) $CH_3 - C \equiv C - CH(CH_3)_2$ (4-methyl-2-pentyne)

- 487. The compound which exhibits optical isomerism is:
 - a) CH₃CHOHCH₃
- b) $(CH_3)_2CHCH_2CH_3$
- c) CH₃CHClCH₂CH₃
- d) CH3CCl2CH2CH3

488. Which of the following applies in the reaction $CH_3CHBrCH_2CH_3 \xrightarrow{Alco.KOH} ?$

- $(I)CH_3CH = CHCH_3(Major product)$
- $(II)CH_2 = CHCH_2CH_3(Minoe product)$
- a) Hofmann's rule
- b) Saytzeff's rule
- c) Kharasch effect
- d) Markownikoff's rule

489. Homologous compounds have:





- a) Same chemical properties
- b) Same molecular weight
- c) Same physical properties
- d) Same m.p. and b.p.
- 490. How many chiral compound are possible on mono chlorination of 2-methyl butane?

b) 4

d) 8

- 491. Which of the following may exist in enantiomorphs?
 - -ÇН—СООН
 - CH₃
 - b) $CH_3 = CHCH_2CH_2CH_3$
- 492. How many isomers are possible for the alkane C₄H₁₀?

b) 5

c) 2

d) 4

- 493. Which of the following IUPAC names is correct?
 - a) 2-methyl-3-ethylpentane
 - b) 2-ethyl-3-methylpentane
 - c) 3-ethyl-2-methylpentane
 - d) 3-methyl-2-ethylpentane
- 494. Ethyl acetoacetate shows, which type of isomerism?
 - a) Chain
- b) Optical
- c) Metamerism
- d) Tautomerism

- 495. Enol content is highest in
 - a) Acetone
- b) Acetophenone
- c) Acetic acid
- d) Acetyl acetone
- 496. The maximum number of stereoisomers possible for 3-hydroxy-2-methyl butanoic acid is:

b) 2

c) 3

d) 4

- 497. Which of the following will exhibit cis trans isomerism?
 - a) CH₂Br CH₂Br
- b) $CBr_3 CH_3$
- c) CHBr = CHBr
- d) $CBr_2 = CH_2$
- 498. Which of the following is most reactive towards electrophilic nitration?
 - a) Toluene
- b) Benzene
- c) Benzoic acid
- d) Nitrobenzene
- 499. The number of structural isomers possible for an organic compound with molecular formula C₅H₁₂ is:

b) 3

c) 4

- 500. Ethylene dichloride and ethylidine chloride are isomeric compounds. The false statement about these isomers is that they
 - a) React with alcoholic potash and give the same product
 - b) Are position isomers
 - c) Contain the same percentage of chlorine
 - d) Are both hydrolysed to the same product
- 501. The IUPAC name of

$$\begin{array}{c} \mathsf{CI} \\ \mathsf{CH}_3 - \mathsf{C} - \mathsf{CH}_2 - \mathsf{CH} = \mathsf{CH} - \mathsf{CH}_3 \\ \mathsf{H} \end{array}$$

- a) 5-chloro-hex-2-ene
- b) 2-chloro-hex-5-ene
- c) l-chloro-1-methyl-pent-3-ene
- d) 5-chloro-5-methyl-pent-2-ene
- 502. Which of the following compounds has incorrect IUPAC nomenclature?



a) $CH_3 CH_2 CH_2 COC_2 H_5$ ethylbutanoate O \parallel $CH_3 CHCCH_2 CH_3$ \mid CH_3

2-methyl-3-pentanone

503. The IUPAC name for tertiary butyl iodide is

- a) 4-iodo butane
- c) 1-iodo-3-methyl propane
- 504. Geometry of methyl free radical is
 - a) Pyramidal
- b) Planar
- 505. Dehydrogenation of ethanol to give ethanal is:
 - a) Addition reaction
 - b) α - α elimination reaction
 - c) α - β elimination reaction
 - d) α - γ elimination reaction
- $506. \ The \ arrangement \ of \ decreasing \ order \ of \ stability \ of$

ČH₃, Č₂H₅, (CH₃)₂ČH and (CH₃)₃ Č free radicals is :

- a) $\dot{C}H_3 > \dot{C}_2H_5 > (CH_3)_2\dot{C}H > (CH_3)_3\dot{C}$
- b) $(CH_3)_3\dot{C} > (CH_3)_2\dot{C}H > \dot{C}_2H_5 > \dot{C}H_3$
- c) $\dot{C}_2H_5 > \dot{C}H_3 > (CH_3)_2\dot{C}H > (CH_3)_3\dot{C}$
- d) $(CH_3)_3$ ° $\dot{C} > (CH_3)_2$ ° $\dot{C}H > \dot{C}H_3 > \dot{C}_2H_5$

507. Which one of the following compounds will be most readily dehydrated?

508. (1)
$$\phi - \overset{\circ}{C} - \phi$$
 (2) $\phi - \overset{\circ}{C} H_2$ (3) $CH_2 = CH - \overset{\circ}{C}H_2$
(4) Θ

CH₃CHCH₂CHO

- b) CH_3
 - 3-methyl butanal

CH₃CHCHCH₃

- d) H_3C OH
 - 2-methyl-3-butanol
- b) 2-iodo butane
- d) 2-iodo-2-methyl propane
- c) Tetrahedral
- d) Linear

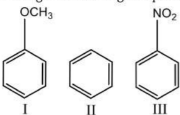


Correct order of stability is

- a) 1>4>2>3
- b) 1>2>3>4
- c) 1>2>4>3
- d) 1>3>4>2

509. The organic chloro compound, which shows complete stereochemical inversion during and $S_N 2$ reaction, is

- a) CH₃Cl
- b) $(C_2H_5)_2$ CHCl
- c) (CH₃)₃CCl
- d) (CH₃)₂CHCl
- 510. Among the following compounds (I-III) the correct order of reaction with the electrophile is



- a) II>III>I
- b) III<I<II
- c) I>II>III
- d) l≈II>III

511. Which of the following is an electrophile?

- a) :CCl₂
- b) CO2

- c) H₂O
- d) NH₃

512. In the following reactions,

(I)
$$CH_3$$
— CH — CH — CH_3
 $H^{\dagger}/Heat$
 A
 $Minor$
 OH
 $Minor$
 $Product$

(II)
$$A \xrightarrow{\text{HBr, dark}} C + D$$

$$(\text{Minor Product}) + (\text{Minor Product})$$

the major products (A) and (C) are respectively:

$$CH_3$$
 CH_3
 CH_3

$$CH_3$$
 CH_3
 CH_3
 CH_3
 CH_3
 CH_3
 CH_3
 CH_3

d)
$$CH_3$$
 CH_3 — C — CH — CH_3 and CH_3
 CH_3 — CH — CH — CH_3

- 513. An organic compound having carbon, hydrogen and sulphur contains 4% of sulphur. The minimum molecular weight of the compound is
 - a) 500
- b) 800

- c) 400
- d) 100



514. The structure of tertiary butyl carbonium ion is:

- a) Pyramidal
- b) Trigonal planar
- c) Tetrahedral
- d) Square planar

515. A carbonium ion contains:

- a) A + vely charged carbon centre
- b) A -vely charged carbon centre
- c) A carbon with odd electron on it
- d) None of the above

516. The formula of 3-chloro-2,2-dimethylbutane is:

- a) CH₃CH(CH₃)C(CH₃)₂Cl b) CH₃(CH₃)₂CH₂Cl
- c) CH₃C(CH₃)₂CH₂Cl
- d) CH₃CHClC(CH₃)₃

517. Which shows the easier electrophilic substitution in ring?

- a) N-acetyl aniline
- b) C₆H₅NH₃Cl
- c) Aniline
- d) Nitrobenzene

518. The number of isomeric pentyl alcohols possible is

- b) Four
- c) Six

d) Eight

519. Naphthalene can be easily purified by

- a) Sublimation
- b) Crystallisation
- c) Distillation
- d) Vaporisation

520. The total number of contributing structures showing hyperconjugation (involving - C - H bonds) for the following carbocation is

- a) Three
- b) Five
- c) Eight
- d) Six

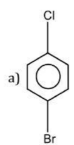
521. What is the structural formula of fumaric acid?

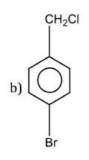
522. The correct structure of dimethylbutyne is:

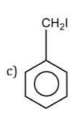
- a) $CH_3CH_2 C \equiv C CH_2CH_3$
- b) $(CH_3)_3C C C \equiv CH$
- c) $CH_3 C \equiv CCH(CH_3)_2$
- d) CH₃—Ç—CH₃ CH₃ CH₃

523. Which one of the following gives white precipitate with $AgNO_3$?









d) None of these

- 524. The reaction $(CH_3)_3CBr \xrightarrow{H_2O} (CH_3)_3$ C. OH is:
 - a) Elimination reaction
 - b) Free radical reaction
 - c) Substitution reaction
 - d) Displacement reaction
- 525. Which of the following is free radical?
 - a) Cl+

b) Cl⁻

c) Č1

- d) NO2
- 526. An organic compound X having molecular formula C₆H₇O₂N has 6 carbons in a ring system, two double bonds and also a nitro group as substitutent. The X is:
 - a) Homocyclic but not aromatic
 - b) Aromatic but not homocyclic
 - c) Homocyclic and aromatic
 - d) Heterocyclic
- 527. Quantitative measurement of nitrogen in an organic compound is done by the method
 - a) Berthelot method
- b) Belstein method
- c) Lassaigne test
- d) Kjheldahl method
- 528. During pyrolysis of an alkane, C—C bond breaks faster than the C—H bond because:
 - a) C—C bond is stronger
 - b) C— H bond is weaker
 - c) C— C bond involves π -bond in alkane
 - d) The bond energy of C-C bond is less than that of C-H bond
- 529. State of hybridization of carbon atom of carbene in the singlet state is:
 - a) sp^2

c) sp^3

d) None of these

- 530. IUPAC name of (CH₃)₃CClis
 - a) n-butyl chloride

b) 3-chloro butane

c) 2-chloro 2-methyl propane

- d) t -butyl chloride
- 531. Most stable carbocation is formed during the heating of which one of the following compound with conc.H2SO4?
 - a) (CH₃)₃COH
- b) C₆H₅CH₂OH
- c) (CH₃)₂CHOH
- d) CH3CH(OH)CH2CH3
- 532. The number of 1° and 2° carbon atoms in *n*-pentane are respectively:

- b) 3, 2
- d) 1, 3
- 533. In benzene, all the C C bonds are of equal length because of:
 - a) Isomerism
- b) Resonance
- c) Tautomerism
- d) Inductive effect

- 534. Nitration of benzene is
 - a) Electrophilic substitution

- b) Electrophilic addition
- c) Nucleophilic substitution
- d) Nucleophilic addition
- 535. The following reaction is described as
 - $\begin{array}{c} \text{CH}_3(\text{CH}_2)_5 \\ \text{H}_3\text{C} \\ \end{array} \begin{array}{c} \text{C} \text{Br} \xrightarrow{\text{OH}^-} \text{HO} \text{CH}_2 \\ \downarrow \\ \text{H} \end{array} \begin{array}{c} \text{(CH}_2)_5\text{CH}_3 \\ \text{CH}_3 \end{array}$
 - a) S_E 2
- c) S_N 1
- d) $S_N 0$
- 536. Nitrogen containing organic compound when fused with sodium forms
 - a) Sodium azide
- b) Sodium cyanide
- c) Sodamide
- d) Sodium cyanate





PARTYMORE CONTROL	englight good physiologists of exceptation of a population of these		
537. Which of the following is	1970) icu	n c u o
a) Br	b) OH	c) :CN	d) C ₂ H ₅ O:
538. In Kjeldahl's method for			1 4 wN
a) % of N = $\frac{1.4 \text{ W}}{N}$	b) % of N = $\frac{1.4 VN}{w}$	c) % of N = $\frac{VW}{1.8}$	d) % of N = $\frac{1.4 wN}{V}$
539. The most satisfactory me			
a) Fractional crystallisat		b) Chromatography	
c) Benedict's reagent		d) Carius method	
540. The IUPAC name of an u	nsymmetrical ether with th	e molecular formula C ₄ H ₁₀	0
a) Ethoxy propane	b) Methoxy ethane	c) Ethoxy ethane	d) Methoxy propane
541. S_N 1 reaction on optically		ives :	
a) Retention in configura			
b) Inversion in configura	ition		
c) Racemic product			
d) No product			
542. The structures that do not a) Tautomers	ot actually exist are called :		
b) Conformational isome	ore		
c) Canonical structures	.13		
d) Ontical isomers			
543.	$\hat{\Rightarrow}$		
543. The correct IUPAC name of,) is:		
	Y		
	\downarrow		
 a) Isopropyl benzene 	b) Cumene	c) Phenyl isopropane	d) 2-phenyl propane
544. When SCN ⁻ is added to a			
그 그 그 그 그 그 그 그 그 그 그 그 그 그 그 그 그 그 그	b) [Fe(OH ₂) ₅ (SCN)] ²⁺		d) [Fe(OH ₂)(SCN)] ⁶⁺
545. Which of the following is			
a) $^+_{ m C~H_3}$	b) _{RC H2}	c) _{R2} C H	d) R_3 ⁺ C
546. Which one of the followi	ng compound is most acidi	c?	ypyc Minnesot
	,OH	OH	OH
a) $Cl - CH_2 - CH_2 - OH$	b) ()	c) ()	a) (O)
.,	NO ₂	, ()	CH ₃
547. The number of carbon at	0.00	e are :	- 13
a) Four 1° carbon, one 4			
b) two 1° carbon, two 2°			
c) one 1° carbon, three 4			
d) None of the above is c	orrect		
548. The Cl—C—Cl angle in 1	,1,2,2-tetrachloroethene ar	id tetrachloromethane resp	ectively will be about :
a) 120° and 109.5°	b) 90° and 109.5°	c) 109.5° and 90°	d) 109.5° and 120°
549. $C_6H_5C \equiv N \text{ and } C_6H_5N$			
a) Position	b) Functional	c) Metamerism	d) Dextroisomerism
550. The number of possible	enantiomeric pairs that can	be produced during mono	chlorination of 2-methyl
butane is :	F) 2	-2.4	J) 1
a) 2	b) 3	c) 4	d) 1
551. (O)-ç-o-(O)			
In U			
Electrophilic substitution	n occurs at		
term und der som en			

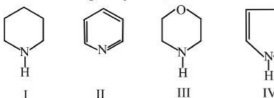
a) ortho/para at first ring

b) meta at first ring

c) ortho/para at second ring

- d) meta at second ring
- 552. In estimation of nitrogen by Duma's method 1.18 g of an orgain compound gave 224 mL of N₂ at NTP. The percentage of nitrogen in the compound is
- b) 11.8
- c) 47.7
- d) 23.7

553. In the following compounds,



the order of basicity is:

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

CH₃

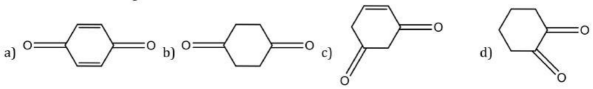
- c) II > I > III > IV
- d) I > III > IV

554.

In the structure,
$$H_3C$$
 — CH — CH_2 — C — CH_3 — CH_3

the number of carbons are:

- a) One primary, two secondary and one tertiary
- b) Four primary, two tertiary and one secondary
- c) One primary, one secondary, one tertiary and one quaternary
- d) Five primary, one secondary, one tertiary and one quaternary
- 555. Which of the following does not exhibit tautomerism?



556. Chromatography technique is used for the separation of

a) Small sample of mixture

b) Plant pigments

c) Dyestuff

d) All of the above

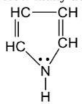
557. CH₃CH₂OH and CH₃OCH₃ are the example of

- a) Chain isomerism
- b) Functional isomerism c) Position isomerism
- d) Metamerism 558. The number of geometrical isomers in case of a compound with the structure, $CH_3 - CH = CH - CH =$ $CH-C_2H_5$ are.
 - a) Four
- b) Three
- c) Two
- d) five

559. The stability of the carbocation decreases in the order

- a) $R_2CH^+ > R_3C^+ > RCH_2^+ > CH_3^+$
- b) $R_3C^+ > R_2CH^+ > RCH_2^+ > CH_3^+$
- c) $CH_3^+ > R_2CH^+ > RCH_2^+ > R_3C^+$
- d) $CH_3^+ > RCH_2^+ > R_2CH^+ > R_3C^+$

560. How many delocalized π -electrons are there in the compounds



b) 2

c) 4

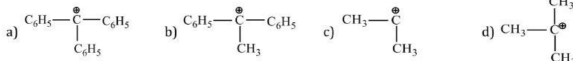
561. What will be the compound if two valencies of carbonyl group are satisfied by two alkyl groups?

- a) Aldehyde
- b) Ketone
- c) Acid
- d) Acidic anhydride





562. The most stable carbocation is:



b)
$$C_6H_5$$
— $\overset{\oplus}{C}$ — C_6H

563. Which of the following belongs to -I group?

- a) $-C_6H_5$
- b) $-CH_3$
- c) -CH₂CH₃
- d) $-C(CH_3)_3$

564. IUPAC name of following compound,

$$CH_3$$
 $-C$ $-CH_2$ $-CH_3$ is:

- a) 2-cyclohexylbutane
- b) 2-phenylbutane
- c) 3-cyclohexylbutane
- d) 3-phenylbutane

- 565. Which is most commonly used to dry organic liquids?
 - a) Lithium
- b) Sodium
- c) Potassium
- d) Rubidium
- 566. In Lassaigne's solution, pink/violet colouration is produced when sodium nitroprusside solution is added. It indicates the presence of
 - a) Sulphur
- b) Nitrogen
- c) Chlorine
- d) None of these
- 567. A carbonium ion is formed when a covalent bond between two atoms in an organic compound undergoes:
 - a) Homolysis
- b) Heterolysis
- c) Cracking
- d) Pyrolysis

- 568. Racemic mixture is formed by mixing two
 - a) Isomeric compounds

b) Chiral compounds

c) meso compounds

- d) Enantiomers with chiral carbon
- 569. In a solution, solvent can be separated from solute by one of the following process
 - a) Decantation
- b) Filtration
- c) Distillation
- d) Sedimentation

- 570. Buta-1,3-diene and But-2-yne are:
 - a) Position isomers
- b) Functional isomers
- c) Chain isomers
- d) Tautomers
- 571. CH₃ CHCl CH₂ CH₃ has a chiral centre. Which one of the following represent its *R*-configuration?

- 572. Which of the following statements (s) is (are) not true?
 - a) Carbanions and carbonium ions, usually exist in ion pairs or else solvated

Acidity increases and basicity decreases in going from left to right across a row of Periodic Table

b) $CH_4 < NH_3 < H_2O < HF(acidity)$

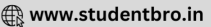
$$CH_3^- > NH_2^- > OH^- > F^-$$
(basicity)

- c) RCOOH like RCOR reacts with H₂NOH to give an oxime
- Decreasing order of ionizing power of solvents is
- $CF_3COOH > HCOOH > H_2O > CH_3COOH > CH_3OH > C_2H_5OH > (CH_3)_2SO > CH_3CN$
- 573. The intermediate during the addition of HCl to propene in the presence of peroxide is :
 - a) CH₃ CHCH₂Cl
- b) CH₃ ČHCH₃
- c) CH₃CH₂ CH₂
- d) CH₃CH₂ ČH₂

- 574. Which of the following represents neo -pentyl alcohol?
 - a) $CH_3CH(CH_3)CH_2CH_2OHb$) $(CH_3)_3C CH_2OH$
 - c) CH₃(CH₂)₃OH
- d) CH₃CH₂CH(OH)C₂H₅

575. 2-methyl-2-butene will be represented as:





- 576. The most abundant organic compound in the world is:

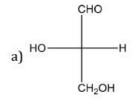
- b) Chlorophyll
- c) Alkaloids
- d) Cellulose
- 577. The chain initiating species in free radical chlorination of methane is:
 - a) Cl free radical
- b) HCl
- c) CH₃ radical
- d) Methylene radical

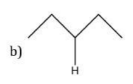
- 578. Which of the following belongs to +I group?
 - a) OH
- b) $-OCH_3$
- c) -COOH
- $d) CH_3$
- 579. Different structures generated due to rotation about, C C axis, of an organic molecule, are examples of
 - a) Geometrical isomerism

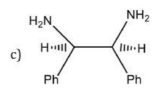
b) Conformational isomerism

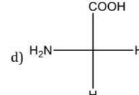
c) Optical isomerism

- d) Structural isomerism
- 580. Which of the following molecules is expected to rotate the plane of plane-polarised light?









- 581. Chromatography was discovered by
 - a) Kekule
- b) Pauling
- c) Rutherford
- d) Tswett
- 582. Sodium nitroprusside when added to an alkaline solution of sulphide ions produces a colouration

- b) brown
- c) Blue
- d) Purple

- 583. (CH₃)₄ N is neither an electrophile, nor a nucleophile because it:
 - a) Does not have electron pair for donation as well as cannot attract electron pair
 - b) Neither has electron pair available for donation nor can accommodate electron since all shells of N are fully occupied
 - c) Can act as Lewis acid and base
 - d) None of the above
- 584. Isopentane can form four isomeric mono bromo derivatives. How many of them are optically active?

b) 2

c) 3

d) None of these

- 585. Which one of the following does not show resonance?
 - a) Carbon dioxide
- b) Benzene
- c) Nitromethane

d) Propane

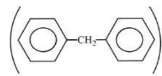
- 586. Select the organic compounds aliphatic in nature but burn with smoky flame:
 - a) CCl₄
- b) CHCl₃
- c) C₆H₅CH₂OH
- d) Both (a) and (b)

- 587. Which of the following is an electrophilic reagent?

b) BF₃

- c) NH₃
- d) RÖH

588. The molecular formula of diphenyl methane is $C_{13}H_{12}$.



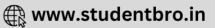
How many structural isomers are possible when one of the hydrogen is replaced by a chorine atom?





a) 6	b) 4	c) 8	d) 7
589. A mixture of iron filling	s and sulphur cannot be se	parated by	
a) Heating		b) Magnet	
c) Shaking with CS ₂		d) Washing in a curre	nt of water
590. Isomers of propionic ac	id are		
a) HCOOC ₂ H ₅ and CH ₃ C	COOCH ₃	b) HCOOC ₂ H ₅ and C ₃ H	H ₇ COOH
c) CH ₃ COOCH ₃ and C ₃ H	I ₇ OH	d) C ₃ H ₇ OH and CH ₃ CO	
591. Reactions involving het	erolytic fission are said to p	proceed via :	
a) Ionic mechanism	b) Polar mechanism	c) Both (a) and (b)	d) None of these
592. Which of the following	orders is true regarding the	e acidic nature of COOH?	
a) Formic acid > acetic	acid> propanoic acid	b) Formic acid > aceti	c acid< propanoic acid
c) Formic acid < acetic	acid> propanoic acid	d) Formic acid > aceti	c acid< propanoic acid
593. Which behaves both as		electrophile?	it ti
a) CH ₃ OH	b) CH ₃ NH ₂	c) CH ₃ CN	d) CH ₃ Cl
594. Alkaline hydrolysis of a		70, 50 cm months and a contract of the contrac	
CH ₃ −C−O−CH−CH ₂ C	H ₂ OH Alcohol + salt		
CII.	3		
The second state to the second state of the se	hand the man than to		
The correct statement a			
	ion about chiral carbon ato		
	ion about chiral carbon ato	m is inverted	
c) Alcohol loses optical	550		
d) All statement are inc		<u></u>	
595. In which case the carbo	크리아스 (CHO) 이 경기 등록 전에 가장하고 있다. 이 교육이 (CHO) 이 이 (CHO) 등 이 이 등로 하는데 되었다.		922199
a) 2-butene	b) Benzene	c) 1-butene	d) 1-propyne
596. Incorrect statement is			
 a) Aniline can be purified 	73		
b) Beilstein test is not g			
	used for estimation of sulp		
(1997) 12 12 12 12 12 12 12 12 12 12 12 12 12	ed in the qualitative detect		compounds
597. The increasing order of	stability of the following fr	ee radicals are	
, (CH ₂) ₂ CH < (CH ₂) ₂ C	Stability of the following in $C < (C_6H_5)_2 CH$ $< (C_6H_5)_3 C$		
a) 3/2 3/3	< (C ₆ H ₅) ₃ C		
•	• •		
b) $(C_6H_5)_3 \overset{\bullet}{C} < (C_6H_5)_2 \overset{\bullet}{C}$	$CH < (CH_3)_3 C$		
٠,	< (CH ₃) ₂ CH		
(C.H.), CH < (C.H.)	, C < (CH ₂), C		
c) (C6113)2 C11 (C6113)	3 C < (CH ₃) ₃ C < (CH ₃) ₂ CH		
35	• (6113)2 611		
d) (CH ₃) ₂ CH < (CH ₃) ₃ C	$C < (C_6H_5)_3 C$		
	< (C ₆ H ₅) ₂ CH		
598. Which one of the follow	ing explain, why propene ι	ındergo electrophilic add	ition with HBr, but not with
HCN?	8478 NT NO 05000 150	1780 M	
a) Br ⁻ is better nucleop	hile than CN ⁻		
(1.1 m), 400 1 (10 f) 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	rce of proton as it is strong	er acid than HCN	





c) HCN attacks preferentially via lone pair of nitrogen

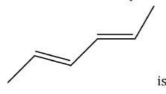
599. The structural formula of 2,2,3-trimethyl hexane is :

d) The C-Br bond being stronger is formed easily as compared to C-CN bond

d)
$$CH_3 - CH - CH_2 - CH_2 - CH_3$$
 CH_3 CH_3 CH_3

600. The IUPAC name of the compound,

- a) 2-iodo-3-chloro-4-pentanoic acid
- b) 4-oxo-3-chloro-2-iodo pentanoic acid
- c) 4-carboxy-4,3-chloro-2-butanone
- d) 3-chloro-2-iodo-4-oxo-pentanoic acid
- 601. Select the correct statement about the detection of sulphur in organic compounds
 - a) Sulphur present in organic compound on fusion with sodium is converts to Na₂S₂O₃
 - b) FeCl₃ gives purple colour when added to the sodium fusion extract
 - c) Sodium nitroprusside is used to detect the presence of sulphur
 - d) All of the above
- 602. The name of the compound



a) (2Z,4Z)-2, 4-hexadiene

b) (2Z-, 4E)-2, 4-hexadiene

c) (2E, 4Z)-2, 4-hexadiene

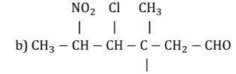
- d) (4E, 4Z)-2, 4-hexadiene
- 603. Rotation of plane polarized light can be measured by :
 - a) Manometer
- b) Calorimeter
- c) Polarimeter
- d) Viscometer
- 604. A molecule having three different chiral carbon atoms, how many stereoisomers it will have?
 - a) 8

b) 3

c) 9

- d) 6
- 605. Correct the structural formula of compound-5-nitro-3-methoxy-3-methylhexanoyl chloride is

NO₂ OCH₃ O | | | | | | | | | | | a) $CH_3 - CH - CH_2 - C - CH_2 - C - CI$ | CH_3 CH_3



 $\begin{array}{ccccc} \mathrm{CH_3} - \mathrm{CH_2} - \mathrm{CH} - \mathrm{C} - \mathrm{CH} - \mathrm{CH_3} \\ \mathrm{d)} & | & | & | \\ & \mathrm{Cl} & \mathrm{O} & \mathrm{OCH_3} \end{array}$

OCH₃



OCH₃

- 606. Which of the following is the strongest base?
 - a) Acetamide
- b) Aniline
- c) Methylamine
- d) Dimethylamine
- 607. The isomeric monosubstitution products theoretically possible for the structure,

$$\mathrm{CH_2} = \mathrm{HC} - \mathrm{CH_2} - \mathrm{CH_2} - \mathrm{CH} = \mathrm{CH_2}$$
 are :

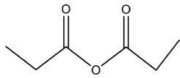
c) 4

- d) 6
- 608. Intermediate product formed in the acid catalysed dehydration of n-propyl alcohol is :
 - a) $CH_3 CH_2 CH_3$
- b) $CH_3 CH = CH_2$ c) $CH_3 CH_3 CH_3$
- d) CH₃—CH₂—CH₂

609. Product [A] in the reaction:

$$\begin{array}{c|c}
\text{OH} & \text{OH} \\
-\text{CH}_3 & \xrightarrow{\text{H}_2\text{SO}_4} [A], \text{ is :}
\end{array}$$

610. The IUPAC name of the following compound is



a) Propionic anhydride

b) Dipropanoic anhydride

c) Ethoxy propanoic acid

- d) Propanoic anhydride
- 611. Which of the following has the highest nucleophilicity?
 - a) F-

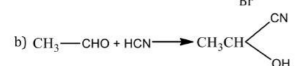
- b) OH-
- c) CH₃
- d) NH₂

- 612. Anthracene is purified by
 - a) Filtration
- b) Distillation
- c) Crystallisation
- d) Sublimation

613. Which of the following requires radical intermediate?

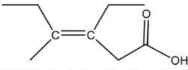
$$CH_3 - CH = CH_2 + HBr \rightarrow CH_3 - CH - CH_3$$

a)



- c) $CH_3 CH = CH_2 + HBr \rightarrow CH_3 CH_2 CH_2 Br$
- d) $CH_3CHO + NH_2OH \xrightarrow{H^+} CH_3 CH = N OH$
- 614. Chiral molecules are those which:
 - a) Are not superimposable on their mirror images
 - b) Are superimposable on their mirror images
 - c) Show geometrical isomerism
 - d) Are unstable molecules
- 615. Who is called the 'Father of Chemistry'?
 - a) Faraday
- b) Priestley
- c) Rutherford
- d) Lavoisier
- 616. With a change in hybridization of the carbon bearing the charge, the stability of a carbanion increase in the
 - a) $sp < sp^3 < sp^2$
- b) $sp < sp^2 < sp^3$ c) $sp^2 < sp < sp^3$ d) $sp^3 < sp^2 < sp$

617. The correct IUPAC name of the acid



- a) Z-3-ethyl-4-methyl hex-3-en-1-oic acid
- b) Z-3-ethyl-4-methyl hexanoic acid
- c) Z-3, 4-diethylpent-3-en-1-oic acid
- d) E-3-ethyl-4-methylhex-4-en-1-oic acid

618. IUPAC name of,

$$\begin{array}{c} \text{CH}_3 \\ \text{I} \\ \text{CH}_3 - \text{C} = \text{C} - \text{CH} - \text{CH} = \text{CH}_2 \text{ is:} \\ \text{CH}_2 \\ \text{CH}_3 \\ \end{array}$$

- a) 2-ethyl-3-methyl-hexa-l-en-4-yne
- b) 5-ethyl-4-methyl-hexa-2-yn-5-ene
- c) 3-methylene-4-methylhepta-5-yne
- d) 5-methylene-5-ethyl-4-methylhepta-2-yne
- 619. The total number of alkenes possible by dehydromination of 3-bromo-3-cyclopentylhexane using alcoholic KOH is
 - a) 1

b) 3

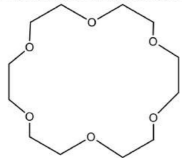
c) 5

d) 7

- 620. The Cl—C—Cl bond angle in dichloro methane will be :
 - a) > 109°28′
- b) < 109°28′
- c) 109°28'
- d) 120°
- 621. In the following reaction sequence, the chain initiation steps is :
 - a) C1—C1→Č1+Č1
 - b) $CH_4 + \mathring{C}l \longrightarrow \mathring{C}H_3 + HCl$
 - c) $\overset{\bullet}{C}H_3 + Cl_2 \longrightarrow CH_3Cl + \overset{\bullet}{Cl}$
 - d) $\overset{\bullet}{C}H_3 + \overset{\bullet}{C}l \longrightarrow CH_3Cl$
- 622. Amongst the following which of the above are true for S_N 2 reaction?
 - (i) The rate of reaction is independent of the concentration of the nucleophile.
 - (ii) The nucleophile attacks the carbon atom on the side of the molecule opposite to the group being displaced.
- (iii) the reaction proceeds with simultaneous bond formation and bond rupture.
 - a) (i), (ii)
- b) (i), (iii)
- c) (i), (ii), (iii)
- d) (ii), (iii)

- 623. Predict the nature of principal product in the reaction,
 - $BrCH_2CH_2CH_2CH_2Br + KOH (alc.) \rightarrow Product :$
 - a) 1,3-butadiene
- b) Cyclobutane
- c) $BrCH_2CH_2CH = CH_2$ d) None of these

- 624. The most stable carbanion is
 - a) CH₃
- b) RCH₂
- c) R₃C^Θ
- d) CH2CHO
- 625. Crown ethers are named as X-crown-Y. In the following crown ether, X and Y are respectively



- a) 6 and 12
- b) 18 and 6
- c) 24 and 6
- d) 6 and 24





626. Which one of the nitrogen atoms in

0

 $H_2N - NH - C - NH_2$ is the most nucleophilic?

Ш

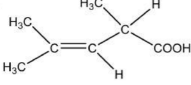
a) III

b) I

c) II

d) All three N atoms

627.



Compound can exhibit

a) Geometrical isomerism

b) Tautomerism

c) Optical isomerism

- d) Geometrical and optical isomerism
- 628. The order of stability of the alkenes

$$R_2C = CR_2, R_2C = CHR, R_2C = CH_2,$$

RCH = CHR,

II

IV

 $RCH = CH_2$

- a) I>II>IV>III>V
- b) I=II>III>IV>V
- c) II>I>IV>III>V
- d) V>IV>III>I>II
- 629. If a compound on analysis was found to contain C = 18.5%, H = 1.55%, Cl = 55.04% and O = 24.81% then its empirical formula is
 - a) CH2OCl
- b) CH2ClO2
- c) ClCH₂O
- d) CHClO
- 630. 2-pentanone and 3-methyl-2-butanone are a pair of isomers.
 - a) Functional
- b) Chain
- c) Positional
- d) Stereo
- 631. The number of isomeric ethers with molecular formula $C_4H_{10}O$ is/ are. b) Two
 - a) One

- c) Three
- d) Four

- 632. Liebig's test is used to estimate
 - a) H

b) C

- c) C and H Both
- d) N

- 633. Number of isomers possible for C_4H_8O is

c) 5

d) 6

- 634. Most stable carbonium ion is
 - a) $\overset{\tau}{C}_2H_5$
- b) (CH₃)₃ C
- c) (C₆H₅)₃ C

635.



In a compound electrophilic substitution has occurred. The substitute-*E* are methyl - CH2Cl, -CCl3 and -CHCl2. The correct increasing order towards electrophilic substitution is

- a) $-CH_3 < -CH_2Cl < -CHCl_2 < -CCl_3$
- b) $-CH_3 < -CHCl_2 < -CH_2Cl < -CCl_3$
- c) $-CCl_3 < -CH_2Cl < -CHCl_2 < -CH_3$
- d) $-CCl_3 < -CHCl_2 < -CH_2Cl < -CH_3$
- 636. In fructose, the possible optical isomers are

b) 8

c) 16

- d) 4
- 637. Which structure can be explained by taking ground state configuration of atom?
 - a) BeH₂
- b) BF₃
- c) CH₄
- d) H₂O

- 638. Which one of the following carbanions is the least stable?
 - a) CH₃CH₂
- b) $HC \equiv C^-$
- c) $(C_6H_5)_3C^-$
- d) $(CH_3)_3C^-$
- 639. Which one of the following is the most energetic conformation of cyclohexane?

- b) Twisted boat
- c) Chair
- d) Half chair
- 640. The energy difference between the chair and boat form of cyclohexane is:





a) 44 kJ mol⁻¹

b) 24 kJ mol⁻¹

c) 34 kJ mol⁻¹

d) 68 kJ mol⁻¹

641.

The electrophile involved in the above reaction is

- a) Dichloromethyl cation (CHCl₂)
- b) Dichlorocarbene (:CCl₂)

c) Trichloromethyl anion (CCl₃)

- d) Formyl cation (CHO)
- 642. Addition of HI on double bond of propene yields isopropyl iodide as major product. It is because the addition proceeds through:
 - a) More stable carbocation
 - b) more stable carbanion
 - c) More stable free radical
 - d) Homolysis
- 643. For which of the following parameters the structural isomers C_2H_5OH and CH_3OCH_3 would be expected to have the same values? (Assume ideal behaviour)
 - a) Heat of vaporisation
 - b) Vapour pressure at the same temperature
 - c) Boiling points
 - d) Gaseous densities at the same temperature and pressure

644.

The IUPAC name of

- a) Bicyclo (2,1,0) pentane
- b) 1,2-cyclopropyl cyclobutane
- c) Cyclopentane (4,3)annulene
- d) 1,2-methylene cyclobutane
- 645. The IUPAC name of neopentane is:
 - a) 2,2-dimethylpropane b) 2-methylpropane
- c) 2,2-dimethylbutane
- d) 2-methylbutane
- 646. In Carius method of 0.099 g orgainc compound gave 0.287 g AgCl. The percentage of chlorine in the compound will be
 - a) 28.6
- b) 71.7
- c) 35.4
- d) 64.2
- 647. The most common type of reaction in aromatic compounds is
 - a) Elimination reaction

- b) Addition reaction
- c) Electrophilic substitution reaction
- d) Rearragement reaction
- 648. Select the correct order of basic nature:
 - a) $CH_3CH_2^- > CH_2 = CH^- > CH \equiv C^- \rightarrow OH^-$
 - b) $CH_3CH_2^- > CH \equiv C^- > CH_2 = CH^- > OH^-$
 - c) $CH_3CH_2^- > OH^- > CH \equiv C^- > CH_2 = CH^-$
 - d) $OH^- > CH \equiv C^- > CH_2 = CH^- > CH_3 CH_2^-$
- 649. Arrange the following carbocations in order of stability

benzyl allyl methyl vinyl

- II a) IV>III>II>I
- IV III
- c) II>IV>III>I
- d) III>II>IV

- 650. The prefix name of -SH group in IUPAC system is:
 - a) Mercapto
- b) Thiol

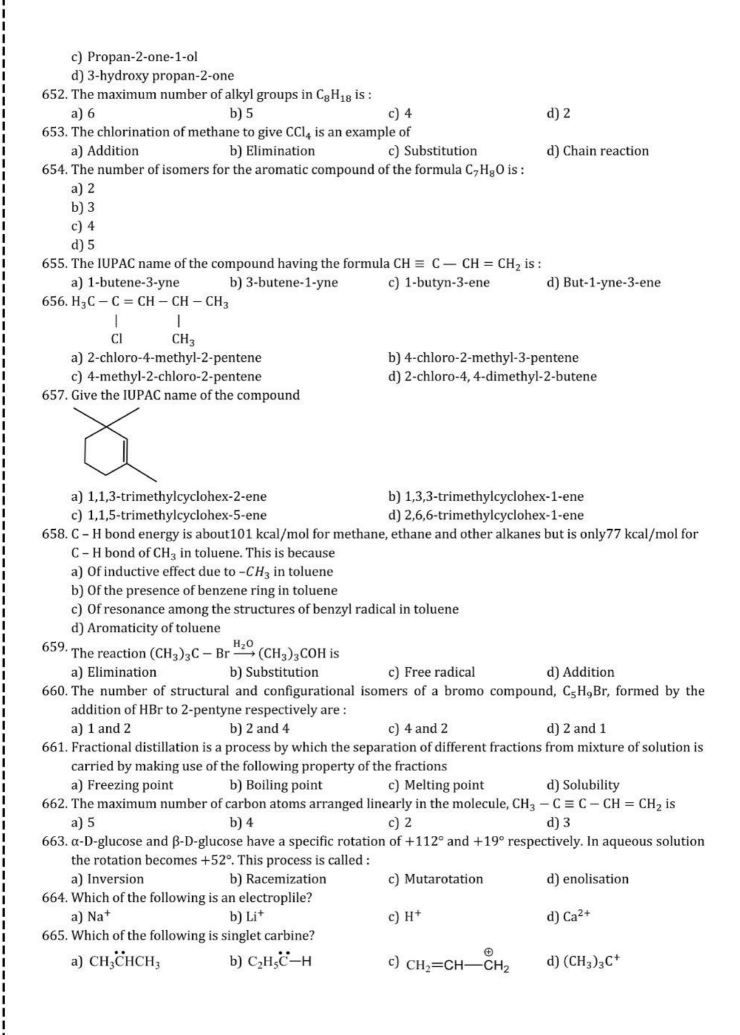
b) I>II>III>IV

- c) Sulphide
- d) None of these

- 651. The correct name for CH₃COCH₂OH is:
 - a) 2-keto propanol
 - b) 1-hydroxy propan-2-one







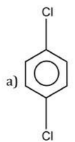
666. Which of the following is the correct order of priority of groups in D-glyceraldehyde?

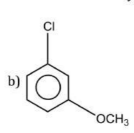
- a) OH(1), CHO(2), CH₂OH(3) and H(4)
- b) OH (1), CH₂OH(2), CHO(3) and H(4)
- c) CH₂OH(1), CHO(2), OH(3) and H(4)
- d) CHO(1), OH(2), CH2OH(3) and H(4)

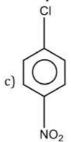
667. o-hydroxytoluene and benzyl alcohol are:

- a) Position isomers
- b) Keto-enol tautomers
- c) Chain isomers
- d) None of these

668. Which of the following would react most readily with nucleophiles?







669. Which group has the highest + Inductive effect?

- b) CH₃CH₂ -
- c) $(CH_3)_2CH -$
- d) $(CH_3)_3 C -$

670. The Prussian blue colour obtained during the test of nitrogen by Lassaigne;s test is due to the formation of

- a) Fe[Fe(CN)₆]₃
- b) $Na_4[Fe(CN)_6]$
- c) Fe₃[Fe(CN)₆]₄
- d) Fe₂[Fe(CN)₆]

671. How many π -electrons are there in the following structure?



b) 4

c) 6

d) 8

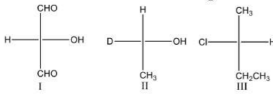
672. Which of the following statements is correct?

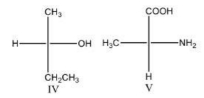
- a) Allyl carbonium ion (CH_2 =CH— CH_2) is more stable then propyl carbonium ion
- b) Propyl carbonium ion is more stable than the allyl carbonium ion
- c) Both are equally stable
- d) None of the above

673. Which of the following possesses an *sp*-hybridized carbon in its structure?

- a) $CCl_2 = CCl_2$
- b) $CH_2 = C = CH_2$
- c) $CH_2 = CH CH = CH_2$
- d) $CH_2 = CCI CH = CH_2$

674. Select R-isomers from the following





- a) I and III
- b) II, IV and V
- c) I, II and III
- d) II and III

675. Di-chloroacetic acid is a stronger acid than acetic acid.

This due to occurrence of



a) Mesomeric effect

b) Hyperconjugation

c) Inductive effect

d) Steric effect

676.

The IUPAC name of,

- a) 3-methyl cyclo-1-buten-2-ol
- b) 4-methyl cyclo-2-buten-1-ol
- c) 4-methyl cyclo-1-buten-3-ol
- d) 2-methyl cyclo-3-buten-1-ol
- 677. The sodium extract of an organic compound on boiling with HNO3 and addition of ammonium molybdate of solution gives a yellow precipitate. The compound contains
 - a) Nitrogen
- b) Phosphorus
- c) Sulphur
- d) Chlorine

678. Unpaired electron in CH3 occupies:

- a) sp-hybrid orbital
- b) sp^3 -hybrid orbital
- c) p-orbital
- d) sp2-hybrid orbital
- 679. In which of the following ways does the hydride ion tend to function?
 - a) An electrophile
- b) A nucleophile
- c) A free radical
- d) An acid

680. The number of isomers of the compound with molecular formula C2H2Br2is

d) 2

681. The treatment of CH_3MgX with $CH_3C \equiv C$ — H produces:

- a) CH₄
- b) $CH_3 CH = CH_2$ c) $CH_3C \equiv C CH_3$
- d) CH₃-C=C-CH₃
- 682. The correct decreasing order of priority for the functional groups of organic compounds in the IUPAC system of nomenclature is
 - a) -COOH, $-SO_3H$, $-CONH_2$, -CHO
- b) $-SO_3H$, -COOH, $-CONH_2$, -CHO
- c) -CHO, -COOH, SO₃H, -CONH₂
- d) -CONH2, -CHO, -SO3H, -COOH
- 683. Which of the following is most likely to show optical isomerism?

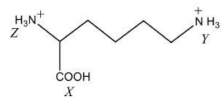
d)
$$CI$$

 $HC \equiv C - C = CH_2$

- 684. Which of the following statements is incorrect?
 - a) The rate of reaction increases with increase in water concentration in the hydrolysis of tertiary butyl bromide in methanol and water
 - The relative nucleophilicity in protic solvent is
 - $CN^- > I^- > \overline{O}H > Br^- > CI^- > F^- > H_2O$
 - In S_N 2 reactions, the order of reactivity of alkyl halides is in the order c) methyl>primary>secondary> tertiary
 - d) S_N2 reaction involves carbonium ions
- 685. Arrange in order of increasing acidic strength.







a) X>Z>Y

b) Z<X>Y

c) X>Y>Z

d) Z>X>Y

686. For the purification, isolation and separation of organic compounds, the latest technique followed is

a) Chromatography

b) Steam distillation

c) Fractional crystallisation

d) Sublimation

687. Arrange *p*-toluidine (I) N,N-dimethyl-*p*- toluidine (II) *p*- nitroaniline (III) and aniline (IV) in order of decreasing basicity

a) I>IV>III>II

b) I>II>III>IV

c) II>I>IV>III

d) III>I>II>IV

688. The reagent showing addition on alkene against the Markownikoff's rule of:

a) Br₂

b) H₂S

c) HF

d) HBr

689. Carbocation can undergo:

a) Loss of a proton

b) Addition to multiple bond

c) Combination with anions

d) All of the above

690. Lactic acid is:

a) Propionic acid

b) β-hydroxypropanoic acid

c) a-hydroxypropanoic acid

d) None of the above

691. Of the following compounds which will have a zero dipole moment?

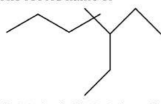
a) 1,1-dichloroethylene

b) trans-1,2-dichloroethylene

c) cis-1,2-dichloroethylene

d) None of the above

692. The IUPAC name of



a) 1,1-diethyl1-2, 2-dimethylpentane

b) 4,4-dimethyl-5,5-diethylpentane

c) 5, 5-diethyl-4,4-dimethylpentane

d) 3-ethyl-4,4-dimethylheptane

693.

The correct IUPAC name

a) 1- cyclopropyl cyclobutane

b) 1, 1-dicyclobutane

c) 1- cyclobutane-1- cyclopropane

d) None of the above

694. Reactivity towards nucleophilic addition reaction of

I. HCHO II.CH3CHO

III. CH₃COCH₃ is

a) II>III>I

b) III>II>I

c) I>II>III

d) I>III>II

695. Arrange the following compounds in order of their decreasing reactivity with an electrophile, E^{\oplus} .

(A) Chlorobenzene,

(B) 2,4-dinitrochlorobenzene,

(C) p-nitrochlorobenzene





a) C>B>A

b) B>C>A

c) A>C>B

d) A>B>C

696. Isomerism exhibited by acetic acid and methyl formate is:

a) Functional

b) Chain

c) Geometrical

d) Central

697. $C_3H_5Cl + aq. NaOH \rightarrow C_2H_5OH + NaCl;$

this reaction is

a) Electrophilic substitution of I order

b) Electrophilic substitution of II order

c) Nucleophilic substitution of I order

d) Nucleophilic substitution of II order

698. In IUPAC suffix name of —COX is:

a) Oyl halide

b) Halo carbonyl

c) Carbamoyl

d) None of these

699. IUPAC name of the compound,

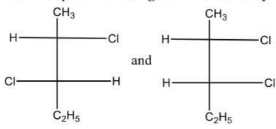
OH is: a) but-2-en-l-ol

b) l-hydroxy but-l-ene

c) 4-hydroxy butene-3

d) But-l-en-l-ol

700. The two optical isomers given below, namely



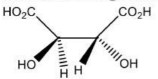
a) Enantiomers

b) Geometrical isomers

c) Diastereomers

d) Structural isomers

701. The absolute configuration of



a) S, S

b) R, R

c) R, S

d) S, R

702. Which one of the following compounds is most reactive towards nucleopilic addition?

a) CH3CHO

b) PhCOCH₃

c) PhCOPh

d) CH₃COCH₃

703. Among the following four structures I to IV

$$\begin{array}{c}\mathsf{CH}_3\\ \\ |\\ \mathsf{C}_2\mathsf{H}_5 & \mathsf{CH} & \mathsf{C}_3\mathsf{H}_7\\ (\mathrm{I})\end{array}$$

$$\begin{array}{c|ccccc} CH_{3} & CH_{3} & \\ & & & \\ & & & \\ CH_{3} & C & CH & C_{2}H_{5} \\ & & & \\ & & & \\ & & & \\ H & & & \\ & &$$

it is true that

a) All four are chiral compounds

b) Only I and II are chiral compounds

c) Only III is a chiral compound

d) Only II and IV are chiral compounds

704. Which of the following is the most stable cation?



a)
$$F_3C$$
— CH_2^{\oplus}

705. Write the IUPAC name of

$$\begin{array}{c} \mathrm{CH_2} - \mathrm{O} - \mathrm{CH} - \mathrm{CH_2} - \mathrm{CH_3} \\ | \\ \mathrm{CH_3} \end{array}$$

a) 3-methoxy butane

b) 2-methoxy butane

c) 3-methyl-3-methoxy propane

d) Butoxy methane

706. Which of the following species is not electrophilic in nature?

707. List the following alkoxide nucleophile in decreasing order of their S_N 2 reactivity

- a) 2>3>5>4>1
- b) 5>3>2>1>4
- c) 1>5>2>3>4
- d) 3>5>1>2>3

708. The Beilstein test for organic compound is used to detect

- a) Nitrogen
- b) Sulphur
- c) Carbon
- d) Halogens

709. Which of the following statements is not characteristic of free radical chain reaction?

- a) It gives major product derived from most stable free radical
- b) It is usually sensitive to change in solvent polarity
- c) It proceeds in three main steps like initiation, propagation and termination
- d) It may be initiated by UV light

710. The presence of carbon in an organic compound is detected by heating it with

- a) Sodium metal to convert it into NaCN
- b) CaO to convert it into CO which burns with a blue flame
- c) CuO to convert it into CO2 which turns lime water milky
- d) Cu wire to give a bluish green flame

711. IUPAC name of the compound, CH2-CH2CH2Cl is:



- a) 1-choloro-2,3-epoxypropane
- b) 3-chloro-1,2-epoxypropane
- c) 1-chloroethoxymethane
- d) None of the above

712. What is the correct IUPAC name of

- a) 4-methoxy-2-nitrobenzaldehyde
- b) 4-formyl-3-nitro anisole
- c) 4-methoxy-6-nitrobenzaldehyde
- d) 2-formyl-5-methoxy nitrobenzene
- 713. Which one is an elimination reaction?
 - a) $CH_3CH_3 + Cl_2 \rightarrow CH_3CH_2Cl + HCl$
 - b) $CH_3Cl + KOH(aq.) \rightarrow CH_3OH + KCl$
 - c) $CH_2 = CH_2 + Br \rightarrow CH_2BrCH_2Br$
 - d) $C_2H_5Br + KOH(alc.) \rightarrow C_2H_4 + KBr + H_2O$
- 714. Identify the compound that exhibits tautomerism





a) 2-butene	b) Lactic acid	c) 2-pentanone	d) Phenol
715. Which of the following is		a) NU	d) ROR
a) H ₂ O 716. The formula of ethaneni	b) SO ₃	c) NH ₃	u) KOK
a) C ₂ H ₅ NC	arie is .		
b) C ₂ H ₅ CN			
c) CH ₃ CN			
d) None of these			
717. Which of the following a	rids shows stereoisomeris	sm?	
a) Oxalic acid	b) Tartaric acid	c) Acetic acid	d) Formic acid
718. Among the following cor			a) I of fine dela
CH ₃	inpounds which can be def	lydrated very easily is	
a) CH ₂ – CH ₂ – C – CH ₂	- CH ₂		
 a) CH ₃ — CH ₂ — C — CH ₂ 			
OH			
$CH_3 - CH_2 - CH_2 - CH_3 - $	H - CH ₃		
b)	3		
ć	Н		
c) $CH_3 - CH_2 - CH_2 - C$			
$CH_3 - CH_2 - CH - CH$			
d)			
CH ₃			
719. Mark the incorrect state	ment in nitrogen Kjeldahl'	s method of estimation	
 a) Nitrogen gas is collect 	ed over caustic potash sol	lution	
b) Potassium sulphate is	used as boiling point elev	rator of H ₂ SO ₄	
c) Copper sulphate or m	ercury acts as a catalyst		
d) Nitrogen is quantitati	vely decomposed to give a	mmonium sulphate	
720. Which of the following o			
		c) $-NR_2 > -OR < -F$	$d) - NH_2 < -OR > -I$
721. The ease of dehydrohalo			
aneron vision in the		c) 3°<2°>1°	d) 3°>2°<1°
722. Lactic acid in which a me			.5 250
		omerism due to the molecul	ar geometry at the :
a) Carbon atom of the m			
b) Carbon atom of the ca	rboxylic acid group		
c) Central carbon atom	1		
d) Oxygen of the hydrox			
723. Which of the following p	rocess is suitable for the p		
a) Simple distillation		b) Fractional distillation	
c) Fractional crystallisat		d) Steam distillation	
724. Maleic and fumaric acids		a) Chain in anns	D. F et a al l'a anno anno
a) Tautomers		c) Chain isomers	
725. $CH_3Br + Nu^- \rightarrow CH_3 -$		g order of the rate of the abo	ove reaction with
nucleophiles $(Nu^-) A$ to			
11.75,	$c0^-, (C)H0^-, (D)CH_30^-$	0) 1 > D > C > D	d) P > D > C > 4
a) $D > C > A > B$		c) $A > B > C > D$	d) $B > D > C > A$
726. Which one is least reacti a) CH ₃ CH ₂ Cl	b) $CH_2 = CHCH_2CI$	c) $CH_2 = CHCl$	d) (CH ₃) ₃ CCl
aj GligGligGl	b) GH ₂ — GHGH ₂ G	c) GH ₂ – GHG	u) (6113)3661

	727. In methanol solution, bromine reacts with ethylene to yield BrCH ₂ CH ₂ OCH ₃ in addition to 1,2-			
	dibromoethane because :			
	a) The intermediate carbocation may react with Br ⁻ or CH ₃ OH			
	b) The methyl alcohol sol			
	c) The reaction follows M			
	d) This is a free radical m		achalia.	
	728. Number of tertiary carbo			4) 4
	a) 1	b) 2	c) Zero	d) 4
	729. Which step is chain propa		g mechanism:	
	(i) $Cl_2 \xrightarrow{hv} Cl^{\bullet} + Cl^{\bullet}$			
	(ii) $Cl^{\bullet} + CH_4 \longrightarrow CH_3$	+HCl		
	$(iii) Cl^{\bullet} + Cl^{\bullet} \longrightarrow Cl_2$			
	(iv) $\overset{\bullet}{C}H_3 + Cl \overset{\bullet}{\longrightarrow} CH_3Cl$			
	a) (i)	b) (ii)	c) (iii)	d) (iv)
	730. The IUPAC name of the c	compound $CH_3 - N \equiv C$ is:		
	a) Ethane nitrile	b) Methane isonitrile	c) Ethane isonitrile	d) None of these
	731. IUPAC name of			
	$CH_3CH_2C(Br) = CH - C$	lis		
	a) 2-bromo-1-chloro but	ene-1	b) 1-chloro-2-bromo but	ene-1
	c) 3-chloro-2-bromo but	ene-2	d) None of the above	
	732. Which of the following un	ndergoes nucleophilic subs	titution exclusively S _N 1 me	chanism?
	 a) Benzyl chloride 	b) Isopropyl chloride	c) Chlorobenzene	d) Ethyl chloride
	733. The sigma bond energy o			
	a) 99 kcal	b) 140 kcal	c) 200 kcal	d) 60 kcal
	734. The general formula C _n H	[25일] [16][[16] [16] [16] [16] [16] [16] [16		
	a) Diketones	b) Carboxylic acids	c) Diols	d) Dialdehydes
	735. The correct sequence of steps involved in the mechanism of Cannizzaro's reaction is a) Nucleophilic attack, transfer of H ⁻ and transfer of H ⁺			
		er of H ⁺ and nucleophilic a philic attack and transfer of		
	10 To	OH ⁻ , transfer of H ⁺ and tr		
	. 이 - 40 1			
736. Examine the following statements regarding S_N 2 reaction (1) The rate of reaction is independents of concentration of nucleophile				
	(2) The nucleophile attacks the carbon atom on the side of molecule opposite to the group being displaced			
(3) The reaction proceeds with simultaneous bond formation and rupture				
	Which of the above writt		ormation and rapture	
	a) 1, 2	b) 1, 3	c) 1, 2, 3	d) 2, 3
	737. Propanol and propanone	17-91-8-00-11 (AT-0-9-3-9		32.76
	a) Functional isomers	b) Enantiomers	c) Chain isomers	d) Structural isomers
	738. Diastereomers can be sep	parated by :		
	a) Fractional distillation) (15)	c) Electrophoresis	d) All of these
	739. Angle strain in cycloprop	ane is		A.S.
	a) 24°44′	b) 9°44'	c) 44'	d) -5°16′
	740. The function of AlCl ₃ in F	riedel-Craft's reaction is		
	a) To absorb HCl	b) To absorb water	c) To produce nucleophil	ed) To produce
				electrophile
	741. In Kjeldahl's method of e		₄ acts as	
	 a) Oxidising agent 	b) Reducing agent	c) Catalytic agent	d) Hydrolysis agent

742. A mixture of acetone and methanol can be separated by

a) Steam distillation

b) Vaccum distillation

c) Fractional distillation

d) None of these

743. The IUPAC name of,

- a) 4-hydroxy-1-methylpentanal
- b) 4-hydroxy-2-methylpentanal
- c) 3-hydroxy-2-methylpentanal
- d) 3-hydroxy-3-methylpentanal

744. The oxygen atom in phenol

- a) Exhibits only inductive effect
- b) Exhibits only resonance effect
- c) Has more dominating resonance effect than inductive effect
- d) Has more dominating inductive effect than the resonance effect

745. 2-methylpent-3-enoic acid shows:

- a) Optical isomerism
- b) Geometrical isomerism
- c) Both (a) and (b)
- d) None of these

746. In the reaction,

$$ROH - R'COOH \longrightarrow R' - C - OR + H_2O$$

water is formed by the combination of:

- a) Hydroxyl of acid with alcoholic hydroxyl hydrogen
- b) Hydroxyl of alcohol with carboxylic hydrogen
- c) Both the above changes
- d) None of the above

747. Pyridine is:

- a) An aromatic compound and a primary base
- b) A heterocyclic amino compound and a tertiary base
- c) An aromatic amino compound and forms salts
- d) A cyano derivative of benzene and secondary base

748. The reason for the loss of optical activity of lactic acid when - OH group is changed by H is that

- a) Chiral centre of the molecule is destroyed
- b) Molecules acquires asymmetry
- c) Due to change in configuration

d) Structural changes occurs

749. The correct order of nucleophilicity among the following is:

750. Which of the following compounds exhibits rotamers?

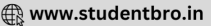
- a) 2-butene
- b) Maleic acid
- c) Butane
- d) Fumeric acid

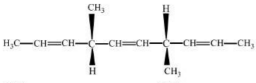
751. Ammonia molecule is:

- a) A nucleophile
- b) An electrophile
- c) A homolytic
- d) An acid

752. The number of optically active products obtained from the complete ozonolysis of the given compound is:







a) 0

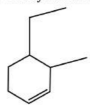
b) 1

c) 2

d) 4

753. The structures,

- a) Chain isomers
- b) Position isomers
- c) Stereo isomers
- d) mesomers
- 754. The systematic (IUPAC) name of the compound with the following structural formula shall be



a) 1-ethyl-2-methyl cyclohexene

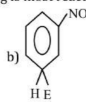
b) 2-methyl-1-ethyl cyclohexene

c) 3-ethyl-2-methyl cyclohexene

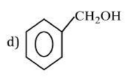
- d) 4-ethyl-3-methyl cyclohexene
- 755. 0.5 g of hydrocarbon gave 0.9 g water on combustion. The percentage of carbon hydrocarbon is

- b) 28.8
- c) 80.0
- d) 68.6
- 756. Which one of the following is most reactive towards electrophilic attack?









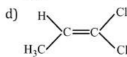
- 757. Identify, which of the below does not possess any element of symmetry?
 - a) (+)(-) tartaric acid
- b) Carbon tetrachloride
- c) Methane
- d) Meso-tartaric acid

758. Geometrical isomerism is shown by:

a)
$$H \subset C \subset C$$

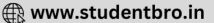
b)
$$H_{3}C$$
 $= C \begin{pmatrix} 1 \\ B_{1} \end{pmatrix}$

c)
$$H_3C$$
 $C = C$ Br



- 759. When thiourea is heated with metallic sodium, the compound which can't be formed is
 - a) NaCNS
- b) NaCN
- c) Na₂SO₄
- d) Na2S
- 760. An unknown compound A has a molecular formula C₄H₆. When A is treated with excess of Br₂ a new substance *B* with formula C₄H₆Br₄ is formed. *A* forms a white ppt. with ammoniacal silver nitrate solution. A may be:
 - a) But-1-yne
- b) But-2-yne
- c) But-1-ene
- d) But-2-ene
- 761. The racemisation of optically active compounds is driven by:
 - a) Entropy
- b) Enthalpy
- c) Entropy and enthalpy d) Element of symmetry





762. A cyclic stereoisomer having the molecular formula	a C ₄ H ₇ Cl are cla	assified and tal	oulated. Find o	out the
correct set of numbers.				

Geometrical

Optical

- a) 6 c) 6
- 2 0

- b) 4 d) 4
- 2 0

763. The correct name for the following hydrocarbon is



a) Tricycle [4.1.0]heptane

b) Bicyclo [5.2.1] heptane

c) Bicyclo [4.1.0] heptane

- d) Bicyclo [4.1.0] hexane
- 764. Which of the following is the most stable radical?
 - a) CH₃
- b) RCH₂
- c) R₂CH[•]
- d) R_3C
- 765. The number of 4° carbon atoms in 2,2,4,4-tetramethylpentane:
 - a) 1

b) 2

c) 3

d) 4

- 766. Inductive effect involves
 - a) Delocalisation of σ -electrons

- b) Displacement of σ -electrons
- c) Delocalisation of π –electrons

- d) Displacement of π -electrons
- 767. Compounds whose molecules are superimposable on their mirror images even though they contain asymmetric carbon atoms or chiral centres are known as:
 - a) Enantiomers
- b) Racemers
- c) Mesomers
- d) Conformers

- 768. Percentage of hydrogen is maximum in.
 - a) C2H4
- b) CH₄
- c) C_2H_2
- d) C_6H_6

- 769. Which of the following has most acidic hydrogen?
 - a) 3-hexanone
- b) 2, 4-hexanedione
- c) 2, 4-hexanedione
- d) 2, 3-hexanedione

770. $_{IUPAC\ name\ of\ CH_3}\cdot N\cdot CH_3$



- a) N, N- dimethylethanamine
- b) N-methyl, N-ethylmethanamine
- c) Dimethyl-ethylamine
- d) None of the above
- 771. Ease of abstraction of hydrogen is greater when attached to:
 - a) 1° carbon
- b) 2° carbon
- c) 3° carbon
- d) neo-carbon

- 772. neo-Heptyl alcohol is correctly represented as:
 - ·ÇH CH2CH3 CH₃ OH
 - -CH2CH2CH2CH3
 - C—CH₂CH₂CH₂OH



d)
$$C_2H_5$$
 C_2H_5 C_2H_5 C_2H_5

773. The IUPAC name for CH₃COCH(CH₃)₂ is:

- a) 4-methyl isopropyl ketone
- b) 3-methyl-2-butanone
- c) Isopropyl methyl ketone
- d) 2-methyl-3-butanone

774. Steam distillation is based on the fact that vaporisation of organic liquid takes place at

- a) Lower temperature than its boiling point
- b) Higher temperature than its boiling point
- c) Its boiling point
- d) Water and organic liquid both undergo distillation

775. IUPAC name of $CH_2 = CH - CH(CH_3)_2$ is:

- a) 1,1-dimethyl-2-propene
- b) 3-methyl-1-butene
- c) 2-vinylpropane
- d) 1-isopropyl ethylene

776. The hybridization of central carbon atom in 1,2- propadiene (allene) is

b) sp^2

d) None of these

777. The fairly neutral character of CH₃OH is changed to which of the following by adding sodium metal?

- a) Acidic
- b) Neutral
- c) An electrophile
- d) A nucleophile

778. The kind of delocalisation involving sigma bond is called

a) Inductive effect

b) Hyperconjugation effect

c) Electromeric effect

d) Mesomeric effect

779. In the case of homologous series of alkanes, which one of the following statements is incorrect?

- a) The members of the series have the general formula C_nH_{2n+2} , where n is an integer
- b) The difference between any two successive members of the series corresponds to 14 unit of relative atomic mass
- c) The members of the series are isomers of each other
- d) The members of the series have similar chemical properties

780. Which of the following reagents will be fruitful for separating a mixture of nitrobenzene and aniline?

- a) Aq. NaHCO₃
- b) H₂O
- c) Aq. HCl
- d) Aq. NaOH

781. The name formic acid was given for HCOOH because it was prepared from:

- a) Acetum
- b) Ant

- c) Wood
- d) Oxalis plant

782. 2, 3-dimethyl hexane contains tertiary secondary andprimary carbon atmos, respectively

- a) 2, 2, 1
- b) 2, 4, 3
- c) 4, 3, 2
- d) 3, 2, 4

783. Which one of the following is the correct formula for dichlorodiphenyltrichloro ethane?

$$CI \longrightarrow CI \longrightarrow CI \longrightarrow CI$$

$$CI \longrightarrow CI$$

$$CI \longrightarrow CI$$

$$CI \longrightarrow CI$$

$$CI \longrightarrow CI$$



- 784. How many sigma and pi bonds are there in the molecule of di cyano ethane (CN CH = CH CN)?
 - a) 3 sigma and 3 pi
- b) 5 sigma and 2 pi
- c) 7 sigma and 5 pi
- d) 2 sigma and 3 pi
- 785. Out of the following, the alkene that exhibits optical isomerism is
 - a) 3-methyl-2-pentene
- b) 4-methyl-1-pentene
- c) 3-methyl-1-pentene
- d) 2-methyl-2-pentene

- 786. The species which use sp^2 -hybrid orbitals in its bonding:
 - a) PH₃
- b) NH₃
- c) CH₃⁺
- d) CH₄

- 787. Carbanion can undergo:
 - a) Rearrangement
 - b) Combination with cation
 - c) Addition to a carbonyl group
 - d) All of the above are correct
- 788. An organic compound $C_5H_{11}X$ on dehydrohalogenation gives pentene-2 only. What is halide?
 - a) CH₃CH₂CHXCH₂CH₃
- b) $(CH_3)_2CHCHXCH_3$
- c) CH₃CH₂CH₂CHXCH₃
- d) CH₃CH₂CH₂CH₂CH₂X
- 789. Percentage composition of an organic compound is as follows

C=10.06, H=0.84, Cl=89.10

Which of the following corresponds to its molecular formula if the vapour density is 60.0?

- a) CH₃Cl
- b) CHCl₃
- c) CH₂Cl₂
- d) None of these
- 790. Which of the following is most reactive towards nucleophilic substitution reaction?
 - a) $CH_2 = CH Cl$
- b) C_6H_5Cl
- c) C₆H₅CH₂Cl
- d) $CICH_2 CH = CH_2$
- 791. Which among the following statements is correct with respect to the optical isomers?
 - a) Enantimoers are non-superimposable mirror images.
 - b) Diastereomers are superimposable mirror images.
 - c) Enantimoers are superimposable mirror image.
 - d) Meso forms have no plane of symmetry.
- 792. Consider thiol anion (RS^{Θ}) and alkoxy anion (RS^{Θ}) . Which of the following statement is correct?
 - a) RS^{Θ} is less basic and less nucleophilic than RO^{Θ}
 - b) RS^{Θ} is less basic but more nucleophilic than RO^{Θ}
 - c) RS^{Θ} is less basic and more nucleophilic than RO^{Θ}
 - d) RS^{Θ} is more basic but less nucleophilic than RO^{Θ}
- 793. The maximum number of alkene isomers for an alkene with molecular formula C4H8 is:

b) 3

- 794. The IUPAC name of the compound $(CH_3)_2CH CH = CH CHOH CH_3$ is
 - a) 5-methyl-hex-3-en-2-ol

b) 2-methyl-hex-3-en-5-ol





c) 2-hydroxy-5-nethyl-3-hexene

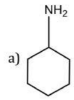
- d) 5-hydroxy-2-methyl-3-hexene
- 795. The number of isomers in C₄H₁₀O are
 - a) 7

b) 8

c) 6

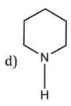
- d) 5
- 796. Which alkyl halide is preferentially hydrolysed by S_N1 mechanism?
 - a) (CH₃)₃C.Cl
- b) CH₃CH₂CH₂Cl
- c) CH₃CH₂Cl
- d) CH₃Cl

797. Which of the following is most basic?









798. The given compound in IUPAC may be called,

- a) Diacetone
- b) Acetoneamine
- c) Diacetoneamine
- d) 4-amino-4-methylpentan-2-one
- 799. The IUPAC name of the compound,

$$CH_2 - CH - COOH$$

OH NH₂

- a) 2-amino-3-hydroxy propanoic acid
- b) 1-hydroxy-2-amino propan-3-oic acid
- c) 1-amino-2-hydroxypropanoic acid
- d) 3-hydroxy-2-amino propanoic acid
- 800. Which of the following compounds is not chiral?
 - a) 1-chloro-2-methyl pentane

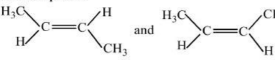
b) 2-chloropentane

c) 1-chloropentane

- d) 3-chloro-2-methyl pentane
- 801. If X is halogen the correct order for $S_N 2$ reactivity is:

a)
$$R_2CHX > R_3CX > RCH_2X$$

- b) $RCH_2X > R_3CX > RCH_2X$
- c) $RCH_2X > R_2CHX > R_3X$
- d) $R_3CX > R_2CHX > RCH_2X$
- 802. The compound



can be distinguished by their:

- a) Chlorinated products
- b) Products formed by addition of bromine
- c) Reaction with H₂/Ni
- d) None of the above
- 803. How many stereoisomers does this molecule have?

$$CH_3CH = CHCH_2CHBrCH_3$$

a) t

b) 8

c) 4

d) 2

- 804. What is the number of possible optical isomers in glucose?
 - a) 3

b) 4

c) 12

- d) 16
- 805. In which reaction addition takes place according to Markownikoff's rule?
 - a) $CH_3CH = CHCH_3 + Br \rightarrow$

b) $CH_2 = CH_2 + HBr \rightarrow$

			75 — 100 M
c) $CH_3CH = CH_2$		d) $CH_3CH = CH_2$	$+ Br_2 \rightarrow$
	gen in organic compounds (THE CONTROL OF THE PROPERTY O	
a) Leibig's test	b) Duma's test	c) Kjeldahl test	d) Beilstein's test
	for catenation next to carb		
a) N	b) S	c) Si	d) P
17. 17. 17. 17. 17. 17. 17. 17. 17. 17.	f alkyl halides by aqueous N	NaOH is best termed as :	
	ubstitution reaction		
b) Electrophilic a			
c) Nucleophilic a			
	ubstitution reaction		
	owing compounds exhibit s	stereoisomerism?	
a) 3-methyl buty		b) 2-methyl bute	
c) 2-methyl buta		d) 3-methyl buta	noic acid
810. The $+$ I.E.(induct	ive effect) is shown by :		
a) CH ₃	b) —OH	c) F	d) — C_6H_5
811. In paper chromat	tography		
1170	s liquid and stability phase		
b) Mobile phase i	s solid and stationary phas	se is liquid	
c) Both phases ar	re liquids		
d) Both phases ar	re solids		
812. Which one of the	following is not found in a	lkenes?	
a) Chain isomeris	sm		
b) Geometrical is	omerism		
c) Metamerism			
d) Position isome	erism		
813. Select the correct	t statement :		
a) The prefixes a	re written before the name	of compound	
b) The suffixes ar	e written after the name o	f compound	
c) The IUPAC nar	ne of a compound is alway	s written as one word	
d) All of the abov	e		
814. A compound cont	tains 2 dissimilar asymmet	ric carbon atoms. The num	ber of optically active isomers is :
a) 2	b) 3	c) 4	d) 5
815. The inductive effe			
	om's ability to cause bond p	oolarization	
b) Increases with increase of distance			
c) Implies the transfer of lone pair of electrons from more electronegative atom to the lesser			
	e atom in a molecule		
		rom lesser electronegative	atom to the more electronegative
atom in a mole			
	ne compound, CICH ₂ CH ₂ CO	OH is:	
a) 3-chloropropanoic acid			
b) 2-chloropropa			
c) 2-chloroethanoic acid			
d) Chlorosuccinio	acid		
817. Which one is a nucleophilic substitution reaction among the following?			
1.00 PM	$CN \rightarrow CH_3CH(OH)CN$		
b) CH3-CH=CH	$H_2 + H_2O \xrightarrow{H^+} CH_3 \xrightarrow{CH} CH$	H_3	
U)	OH		

c)
$$RCHO + R'MgX \longrightarrow R \longrightarrow CH \longrightarrow R$$

OH

d)
$$CH_3$$
 CH_3 CH_3 — CH_2 — CH_2 CH_3 CH_3 — CH_2 — CH_3 CH_3 — CH_2 — CH_2 CH_3

- 818. If there is no rotation of plane polarised light by a compound in a specific solvent, though to be chiral, it means that:
 - a) It is certainly meso
 - b) It is racemic mixture
 - c) It is certainly not chiral
 - d) No such compound
- 819. Formation of ethylene from acetylene is an example of
 - a) Elimination reaction

b) Substitutions reaction

c) Condensation reaction

- d) Addition reaction
- 820. Which of the following is nucleophilic addition reaction?
 - a) Hydrolysis of ethyl chloride by NaOH
- b) Purification of acetaldehyde by NaHSO₃ d) Decarboxylation of acetic acid

- c) Alkylation of anisol
- 821. The reagent used in dehalogenation process is:
 - a) KOH alc.
- b) Zn dust + alc.
- c) Na

d) KOH(aq)

- 822. Benzaldoxime exists in how many forms?
 - a) 1

b) 2

c) 3

d) 4

- 823. Resonance arises due to the:
 - a) Migration of atoms
 - b) Migration of proton
 - c) Delocalisation of σ -electron
 - d) Delocalisation of π -electron
- 824. In the given structure, which carbon atom is most electronegative?

$$CH_3$$
- CH_2 - CH = CH
 (I) (II) (III) (IV)

- a) (I)
- (II) (III) (IV)
- c) (III)
- d) (IV)

825. The following reactions is an example of Reaction.

$$C_2H_4Br_2 \xrightarrow{Alc.KOH} C_2H_2$$

a) Addition

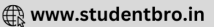
b) Dehydrobromination

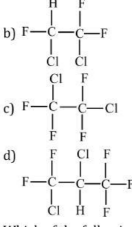
c) Substitution

- d) Debromination
- 826. Which one of the following pair represents stereoisomerism?
 - a) Structural and geometrical isomerism
 - b) Linkage and geometrical isomerism
 - c) Chain and rotational isomerism
 - d) Optical and geometrical isomerism
- 827. Freon-114 is an organic compound. It is chemically called 1,2-dichlorotetrafluoroethane. Its correct structural formula is:









- 828. Which of the following compounds is expected to be optically active?
 - a) $(CH_3)_2CHCHO$
- b) CH₃CH₂CH₂CHO
- c) CH₃CH₂CHBrCHO
- d) CH₃CH₂CBr₂CHO
- 829. Which of the following is least reactive in a nucleophilic substitution reaction?
 - a) $(CH_3)_3C CI$
- b) $CH_2 = CHCl$
- c) CH₃CH₂Cl
- d) $CH_2 = CHCH_2Cl$
- 830. During debromination of meso-dibromo-butane, the major compound formed is:
 - a) n-butane
- b) l-butene
- c) cis-2-butene
- d) trans 2 butene
- 831. What is the empirical formula of a compound having 40% carbon, 6.66% hydrogen and 53.34% oxygen?
 - a) C_2H_2O
- b) C₂H₄O
- c) CH₂O
- d) CHO

- 832. Which of the following can act as an nucleophile?
 - a) BF3
- b) FeCl₃
- c) ZnCl₂
- d) C2H5MgBr
- 833. The hybrid orbitals at carbon 2 and 3 in the compound CH₃CH = CHCH₃ are:
 - a) sp^3 , sp
- b) sp^2 , sp^2
- c) sp, sp
- d) sp^2 , sp

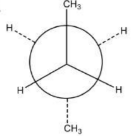
- 834. The alkyl halide that undergoes S_N1 reaction more radily is
 - a) Ethyl bromide
- b) Isopropyl bromide
- c) Vinyl bromide
- d) n-propyl bromide

835. The IUPAC name of,

$$\begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \end{array} \\ \text{CH} - \text{CH}_{2} - \text{C} - \text{CH}_{2} - \text{CH}_{2} \\ \text{CH}_{3} \end{array} \text{ is : }$$

- a) 2,4-dimethylhexanone-3
- b) 2,6-dimethylheptanone-4
- c) 2,6-dimethylhexanone-4
- d) 2,6-dimethylheptanone-5
- 836. In Lassaigne's test, a blue colour is obtained if the organic compound contains nitrogen. The blue colour is due to
 - a) $K_4[Fe(CN)_6]$
- b) $Fe_4[Fe(CN)_6]_3$
- c) Na₃[Fe(CN)₆]
- d) $Cu_2[Fe(CN)_6]$
- 837. According to Gahn-Ingold-Prelog sequence rules, the correct order of priority for the given group is
 - a) $-COOH > -CH_2OH > -OH > -CHO$
- b) $-COOH > -CHO > -CH_2OH > -OH$
- c) $-OH > -CH_2OH > -CHO > -COOH$
- d) $-OH > -COOH > -CHO > -CH_2OH$

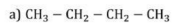
838.

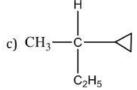


C₂ is rotated anticlockwise 102°C about C₂ - C₃ bond. The resulting conformer is



- a) Partially eclipsed
- b) Eclipsed
- c) gauche
- d) Staggered
- 839. Amongst the following compounds, the optically active alkane having lowest molecular mass is





- d) $CH_3 CH_2 C \equiv CH$
- 840. How many chiral isomers can be drawn from 2-bromo, 3-chloro butane?

b) 3

- d) 5
- 841. Glycerol can be separated from spent-lye in soap industry by
 - a) Steam distillation

- b) Fractional distillation
- c) Distillation under reduced pressure
- d) Ordinary distillation

842. The IUPAC name of

- a) 4-propoxy pentane
- b) Pentyl-propyl ether
- c) 2-propoxy pentane
- d) 2-pentoxy propane
- 843. Correct gradation of basic charactor
 - a) $NH_3CH_3NH_2 > NF_3$

b) $CH_3NH_2 > NH_3 > NF_3$

c) $NF_3 > CH_3NH_2 > NH_3$

- d) $CH_3NH_2 > NF_3 > NH_3$
- 844. An organic compound contains 49.3% carbon, 6.84% hydrogen and its vapour density is 73. Molecular formula of compound is
 - a) $C_6H_9O_3$
- b) $C_4H_{10}O_2$
- c) $C_3H_5O_2$
- d) $C_3H_{10}O_2$
- 845. Vital force theory of the origin of organic compounds was discarded by :
 - a) Kolbe's synthesis
- b) Haber's synthesis
- c) Wöhler's synthesis
- d) Berthelot's synthesis
- 846. In the presence of peroxide, hydrogen chloride and hydrogen iodide do not give anti-Markownikoff addition to alkenes because:
 - a) Both are highly ionic
 - b) One is oxidizing and the other is reducing
 - c) One of the steps are exothermic in both the cases
 - d) All the steps are exothermic in both the cases
- 847. Which of the following does not have a resonance structure?
 - a) Benzene
- b) Benzaldehyde
- c) Acetaldehyde
- d) Benzylamine
- 848. Which of the following is the correct order of stability of different conformations of butane?
 - a) Staggered > Gauche > Partially eclipsed > Fully eclipsed
 - b) Gauche > Staggered > Partially eclipsed > Fully eclipsed
 - c) Staggered > Fully eclipsed > Partially eclipsed > Gauche
 - d) None of the above
- 849. Glucose and fructose are:
 - a) Chain isomers
- b) Position isomers
- c) Functional isomers
- d) Optical isomers

- 850. The enol form of acetone after treatment with D20 gives:
 - a) CH₃-C=CH₂



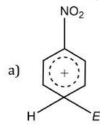
- 851. Eelipsed and staggered forms of *n*-butane are called a pair of :
 - a) Diastereomers
- b) Conformers
- c) Isomers
- d) Enantiomers
- 852. Arrange the following in order of increasing dipole moment (I) Toluene (II) m-dichlorobenzene (III) odichlorobenzene (IV) p-dichlorobenzene:
 - a) I < IV < II < III
- b) IV < I < II < III
- c) IV < I < III < II
- d) IV < II < I < III
- 853. In butane, which of the following forms has the lowest energy?
 - a) Gauche form
- b) Eclipsed form
- c) Staggered form
- d) None of these

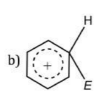
- 854. Molecular mass of a volatile substances may be obtained by
 - a) Beilstein method

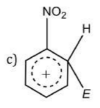
b) Lassaigne method

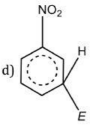
c) Victor Mayer's method

- d) Leibig's method
- 855. The electrophile, E^{\oplus} attacks the benzene ring to generate the intermediate σ –complex. Of the following, which σ –complex is of lowest energy?









- 856. Which is not deflected by a non-uniform electrostatic field?
- b) Chloroform
- c) Nitrobenzene
- d) Hexane

- 857. The reaction $C_2H_5l + KOH \rightarrow C_2H_5OH + KI$ is called
 - a) Hydroxylation substitution

b) Electrophilic substitution

c) Nucleophilic substitution

- d) dehydroiodination
- 858. Correct order of nucleophilicity is
 - a) $I^- > Br^- > Cl^- > F^-$

b) $F^- > Cl^- > Br^- > I^-$

c) $Cl^- > F^- > Br^- > I^-$

- d) $I^- > Cl^- > Br^- > F^-$
- 859. Due to the presence of an unpaired electron free radicals are
 - a) Cations
- b) Anions
- c) Chemically inactive
- d) Chemically reactive

- 860. Which of the following will have meso isomers also?
 - a) 2-hydroxy propanoic acid
 - b) 2,3-dichlorobutane
 - c) 2,3-dichloropentane
 - d) 2-chlorobutane
- 861. The addition of HBr on butene-2 in presence of peroxide follow the:
 - a) Electrophilic addition
 - b) Free radical addition
 - c) Nucleophilic addition
 - d) None of these
- IUPAC name of YIV
 - a) 5-methylhexanol
- b) 2-methylhexanol
- c) 2-methylhex-3-enol
- d) 4-methylpent-2-enol

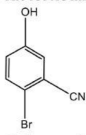




- 863. In which of the compounds given below there is more than one kind of hybridization (sp, sp^2, sp^3) for carbon?
 - (I)CH₃CH₂CH₂CH₃
- $(II)CH_3CH = CH CH_3$
- $(III)CH_2 = CH CH = CH_2$ $(IV)H C \equiv C H$ a) (II) and (IV)
 - b) (I) and (IV)
- c) (II) and (III)
- d) (II)

864. Which represents nucleophilic aromatic substitution reaction?

- a) Reaction of benzene with Cl2 in sunlight
- b) Benzyl bromide hydrolysis
- c) Reaction of NaOH with dinitrofluorobenzene
- d) Sulphonation of benzene
- 865. The IUPAC name of the following compound, is

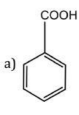


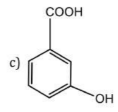
a) 4-bromo-3-cynophenoal

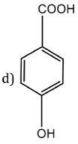
- b) 2-bromo-5-hydroxybenzonitrile
- c) 2-cyano-4-hydroxybromobenzene
- d) 6-bromo-3-hydroxybenzonitrile

866. Ethoxy ethane and methoxy propane are:

- a) Geometrical isomers
- b) Optical isomers
- c) Functional group isomers
- d) Metamers
- 867. Which of the following aromatic acid is most acidic?







868. The hybridization of carbon in diamond, graphite and acetylene is in the order:

- a) sp^3 , sp^2 , sp
- b) sp^2 , sp^3 , sp
- c) sp, sp^2, sp^3
- d) sp^2 , sp, sp^3

869. Which is optically active?

- a) Isobutyric acid
- b) β-chloropropionic acid
- c) Propionic acid
- d) α-chloropropionic acid

870. Which of the following statement is wrong?

- a) Using Lassaigne's test nitrogen and sulphur present in organic compound can be tested
- b) Using Beilstein's test the presence of halogen in a compound can be tested
- c) In Lassaigne's filtrate the nitrogen present in a organic compound is converted into NaCN
- d) In the estimation of carbon, an organic compound is heated with CaO in a combustion tube

871. The reaction, $CH_2 = CHCHO \xrightarrow{HX}$ gives :

- a) CH₃CHXCHO
- b) CH₂XCHCHO
- c) $CH_2 = CHCHX_2$
- d) None of these

872. What kind of isomerism is possible for 1-chloro-2-nitroethene?

a) Functional group isomerism

b) Position isomerism

c) E/Z isomerism

d) Optical isomerism



873. Acetonitrile is

- a) CH₃CN
- b) CH₃COCN
- c) C₂H₅CN
- d) C₆H₅CN

874. Formation of cyanohydrin from a ketone is an example of

a) Electrophilic addition

b) Nucleophilic addition

c) Electrophilic substitution

d) Nucleophilic substitution

875. An organic compound which produces a bluish green coloured flame on heating in presence of copper is

- a) Chlorobenzene
- b) Benzaldehyde
- c) Aniline
- d) Benzoic acid

876. The compound abd C-C abd will exist in:

- a) 3 forms
- b) 4 forms
- c) 5 forms
- d) 2 forms

877. Which of the following compounds has the maximum number of π -bonds?

a) $HC \equiv C - CH = CH_2$

b) $CH_2 = CH - CH = CH_2$

c) CH3CH2COCH3

d) $C_6H_5 - COOH$

878. The C - H bond distance is longest in

- a) C_2H_2
- b) C2H4
- c) C2H6
- d) C₂H₂Br₂

879. The yield in organic reactions is generally poor because the reactions are:

- a) Very fast
- b) Non-ionic
- c) Between covalent compounds
- d) Accompanied by side reactions

880. Which of the following resonating structures of 1-methoxy-1, 3-butadiene is least stable?

- $\stackrel{\Theta}{\text{Cl}} \stackrel{\oplus}{\text{CH}} \stackrel{\oplus}{\text{CH}} \stackrel{\ominus}{\text{CH}} \stackrel{\Box$

881. A student named the compound as 1,4-butadiene:

- a) The name is correct
- b) He committed an error in the selection of carbon chain
- c) He committed an error in position of double bond
- d) Unpredictable

882. The correct IUPAC name of (C2H5)4 C is:

- a) Tetraethyl methane
- b) 2-ethylpentane
- c) 3,3-diethylpentane
- d) None of these

883. The number of different substitution products possible when ethane is allowed to react with bromine is sunlight are:

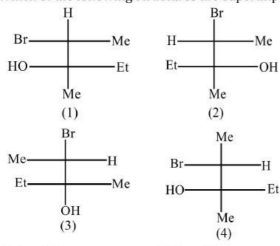
a) 9

b) 6

c) 8

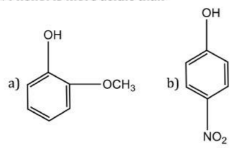
d) 5

884. Which of the following structures are superimposable?



- a) 1 and 2
- b) 2 and 3
- c) 1 and 4
- d) 1 and 3

885. Phenol is more acidic than



d) Both (a) and (c)

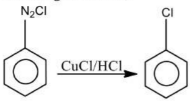
c) C₂H₂

886. During the fusion of an organic compound with sodium metal, nitrogen of the compound is converted into

- b) NaNH₂
- c) NaCN
- d) NaNC

887. The structure representing a heterocyclic compound is:

888. Following reaction is,



d) EI-CB

889. Which of the following reactions is an example of nucleophilic substitution reaction?

- a) $RX + Mg \rightarrow RMgX$
- b) $RX + KOH \rightarrow ROH + KX$
- c) $2RX + 2Na \rightarrow R R + 2NaX$
- d) $RX + H_2 \rightarrow RH + HX$

890. How many structural isomers are possible for C₄H₉Cl?

b) 4

d) 10

891. In which of the following species the central carbon atom is negatively charged?

- a) Carbonium ion
- b) Carbanion
- c) Carbocation
- d) Free radicals

892. Select the molecule having only one π -bond :

- a) CH≡ CH
- b) $CH_2 = CH CHO$
- c) $CH_3 CH = CH_2$
- d) $CH_3 CH = CHCOOH$

893. Optically active compound among the following is:

- a) 2-ethylbutanol-1
- b) *n*-butanol
- c) 2,2-dimethylbutanol
- d) 2-methylbutanol-1

894. Which of the following compounds will be most reactive towards nucleophilic addition reaction?

- a) CH3COCH2CH2CH2CH3
- b) CH₃CH₂COCH₂CH₂CH₃
- c) CH3CH2CH2CH2CHO
- d) $CH_3 CH_2 CO CH CH_3$

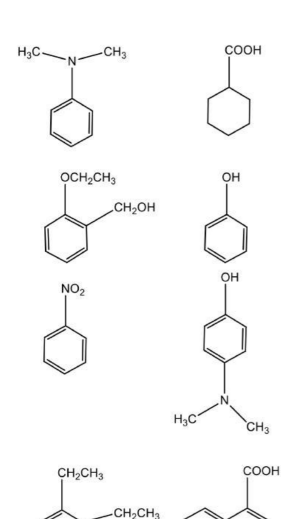




 CH_3 895. Lactic acid, CH₃CH(OH)COOH molecule shows: a) Geometrical isomerism b) Metamerism c) Optical isomerism d) Tautomerism 896. n-pentane and neopentane are: a) Functional isomers b) Geometrical isomers c) Chain isomers d) Position isomers 897. The IUPAC name of acryldehyde is a) Prop-2-en-1-al b) Propenylaldehyde c) But-2-en-1-al d) Propenal 898. Due to presence of an unpaired electron, free radicals are b) Anions c) Chemically inactive d) Chemically reactive a) Cations 899. 2-methylpent-3-ene is a chiral because it has: a) A centre of symmetry b) A plane of symmetry c) Symmetry at C2 carbon d) Both centre and a plane of symmetry 900. Which of the following molecules contain asymmetric carbon atom? a) CH₃CHClCOOH b) CH₃CH₂COOH c) CICH3. CH2COOH d) Cl₂CHCOOH 901. Cyclobutane and butene-1 are: a) Chain isomers b) Position isomers c) Ring-chain isomers d) Metamers 902. Which of the following is not true for carbanions? a) The carbon carrying the charge has eight valence electrons b) They are formed by heterolytic fission c) They are paramagnetic d) The carbon carrying the charge is sp^3 hydridised 903. Which of the following structures permits *cis-trans* isomerism? a) $X_2C = CY_2$ b) $XYC = CZ_2$ c) $X_2C = CXY$ d) XYC = CXY904. Which one of the following compound will show optical iosmerism? a) $(CH_3)_2 - CH - CH_2 - CH_3$ b) $CH_3 - CHOH - CH_3$ c) $CH_3 - CHCl - CH_2 - CH_3$ d) $CH_3 - CCl_2 - CH_2 - CH_3$ 905. The Kolbe's electrolysis proceeds via a) Nucleophilic substitution mechanism b) Electrophilic addition mechanism c) Free radical mechanism d) Electrophilic substitution reaction 906. Which of the following statements is not correct? a) Primary carbocation are more stable than secondary ones b) Secondary free radicals are more stable than primary free radicals c) Tertiary free radicals are more stable than secondary ones d) Tertiary carbonium ions are more stable than primary ones 907. Adsorbent is made of ... in TLC a) Silica gel b) Alumina c) Both (a) and (b) d) None of these

908. Amongst the following, the total number of compounds soluble in aqueous NaOH is





909. The ammonia evolved from the treatment of 0.30 g of an organic compound for the estimation of nitrogen was passed in 100 mL of 0.1 M sulphuric acid. The excess of acid required 20 mL of 0.5 M sodium hydroxide solution for complete neutralization. the organic compound is

a) acetamide

a) 1

b) benzamide

c) urea

c) 3

d) thiourea

d) 4

910. The structure remaining after one H is removed from hydrocarbon is:

b) 2

a) Alkyl group

b) Alkenyl group

c) Alkynyl group

d) All of these

911. C₆H₁₂ on addition of HBr in presence and in absence of peroxide gives some product. It is:

- a) Hexene-3
- b) 2,3-dimethyl butane-2
- c) Symmetrical alkene
- d) All of these

the IUPAC name is

a) 2-ethoxy pentane

b) 4-ethoxy pentane

c) Pentyl-ethyl ether

d) 2-pentoxy ethane

913. A solution of D(+)-2-chloro-2-phenylethane in toluene racemises slowly in the presence of small amount of SbCl5 due to the formation of

a) Carbanion

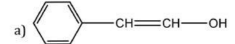
b) Carbene

c) Free radical

d) Carbocation

914. Tautomerism is exhibited by





915. The decreasing order of nucleophilicity among the nucleophilies

- (B) CH₃O
- (C) CN-

$$H_3C$$

- (D)
- a) (C), (B), (A), (D)
- b) (B), (C), (A), (D)
- c) (D), (C), (B), (A)
- d)(A),(B),(C),(D)
- 916. Which of the following statements is incorrect?
 - a) S_N 2 reaction proceeds with inversion
 - b) $S_N 1$ reaction proceeds with racemisation
 - c) S_N 2 reaction involves transition state
 - d) In transition state, one end carries δ^+ and another end carries δ^- charge
- 917. The hybridization of carbon atoms in C C single bond of $HC \equiv C CH = CH_2$ is

a)
$$sp^3 - sp$$

b)
$$sp^{3} - sp^{3}$$

c)
$$sp^2 - sp^3$$

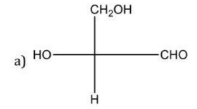
d)
$$sp - sp^2$$

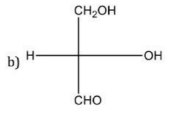
918. One of the stable resonating forms of methyl vinyl ketone is

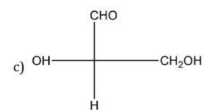
- 919. 5.6 g of an organic compound on burning with excess of oxygen gave 17.6g of CO_2 and 7.2 g of H_2O . The organic compound is
 - a) C_6H_6
- b) C₄H₈
- c) C_3H_8
- d) CH₃COOH

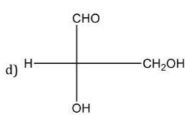
- 920. Base strength of,
 - 9 1) H₃CCH₂
 - 2) H₂C=CH and
 - 3) H-C= $\stackrel{\Theta}{=}$ is in the order of :
 - a) (3) > (2) > (1)
- b) (1) > (3) > (2)
- c) (1) > (2) > (3)
- d) (2) > (1) > (3)
- 921. Which of the following Fischer's projection formula is identical to D-glyceraldehyde?











- 922. 1.2g of organic compound of Kjeldahlization liberates ammonia which consumes
 - 30 cm^3 of 1N HCl. The percentage of nitrogen in the organic compound is
 - a) 30

b) 35

- c) 46.67
- d) 20.8

- 923. Among the following the dissociation constant is highest for
- b) C₆H₅CH₂OH
- c) $CH_3 C \equiv CH$
- d) CH₃NH₃+Cl-
- 924. How much of sulphur is present in an organic compound, if 0.53g of the compound gave 1.158g of BaSO₄on analysis?
 - a) 10%
- b) 15%
- c) 20%
- d) 30%

- 925. Which of the following is a dynamic isomerism?
 - a) Metamerism

b) Geometrical isomerism

c) Tautomerism

- d) Coordinate isomerism
- 926. Which among the following statements is correct with respect to the optical isomers?
 - a) Enantiomers are non-superimposable mirror images
 - b) Diastereomers are superimposable mirror images
 - c) Enantiomers are superimposable mirror images
 - d) Meso forms have no plane of symmetry
- 927. The stability order for carbocations given below is:

(I)
$$R \stackrel{+}{CH}_2$$
 (II) $R - \stackrel{+}{C} = CH_2$ (III) $R - \stackrel{+}{CH} - CH_3$

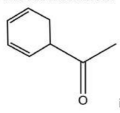
- a) I < II < III
- b) II > II < I
- c) III < I < II
- d) II < I < III
- 928. Duma's method involves the determination of nitrogen content in the organic compound in the form of
 - a) NH₃

b) N₂

c) NaCN

d) $(NH_4)_2SO_4$

929. The IUPAC name of



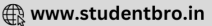
- a) 1- cyclohexa-2,4-dienylethanone
- b) 3- cyclohexa-2,4-dienylethanone
- c) 1- cyclohexa-3,5-dienylethanone
- d) 3- cyclohexa-3,5-dienylethanone
- 930. In the nucleophilic substitution reactions $(S_N 2 \text{ or } S_N 1)$, the reactivity of alkyl halides follows the sequence

a) R-I>R-Br>R-Cl>R-F

b) R-Cl>R-F>R-Br>R-I

c) R-F>R-Cl>R-Br>R-I

- d) R-I>R-F>R-Cl>R-Br
- 931. $\stackrel{\oplus}{R_3}$ N——CH——CH₂ $\stackrel{\text{HBr}}{\longrightarrow}$ product.



Predominant product is

a)
$$R_3$$
N —— CH —— CH₂

b)
$$R_3N - CH_2 - CH_2 - Br$$

c) CH2==CH-

- d) No reaction
- 932. n- pentane and 2-methyl butane are a pair of
 - a) Enantiomers

b) Stereoisomers

c) Diastereomers

d) Constitutional isomers

933.

IUPAC name of compound,

- a) 3-ethyl-4,4-dimethyl heptane
- b) 1,1-diethyl-2,2-dimethyl pentane
- c) 4,4-dimethyl-5,5-diethyl pentane
- d) 5,5-diethyl-4, 4-dimethyl pentane
- 934. IUPAC name of, (C₂H₅)₂CHCH₂OH is:
 - a) 2-ethylbutanol-1 b) 2-methylpentanol-1
- c) 2-ethylpentanol-1
- d) 3-ethylbutanol-1
- 935. Amongst the following the compound that can most readily get sulphonated is
 - a) Benzene
- b) Toluene
- c) Nitrobenzene
- d) Chlorobenzene
- 936. In E2 elimination, some compounds follow Hofmann's rule which means:
 - a) The double bond goes to the most substituted carbon
 - b) The compound is resistant to elimination
 - c) No double bond is formed
 - d) The double bond goes mainly towards the least substituted carbon
- 937. How many asymmetric carbon atoms are present in
 - (i) 1, 2-dimethylcyclohexane
 - (ii) 3-methylcyclopentane and
 - (iii) 3-methylcyclohexene?
 - a) Two, one, one b) One, one, one
- c) Two, none, two
- d) Two, none One

- 938. Which of the following is a chiral compound?
 - a) Hexane
 - b) n-butane
 - c) Methane
 - d) 2,3,4,trimethyl hexane
- 939. How many structures can compound with molecular formula C₇H₉N have?

b) 4

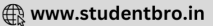
- d) 2
- 940. The maximum number of possible optical isomers in 1-bromo-2-methyl cyclobutane is

b) 2

c) 8

- 941. Which of the following types of reaction occurs when a substituent has got a double bond with evently distributed π electron cloud?
 - a) Electrophilic addition
 - b) Nucleophilic addition
 - c) Any of the (a) and (b)
 - d) None of the above
- 942. Vinyl alcohol and acetaldehyde are:
 - a) Geometrical isomers
 - b) Keto-enol tautomers c) Chain isomers
- d) None of these





943. 0.25 g of an organic compound on Kjeldahl's analysis gave enough ammonia to just neutralise 10cm3 of 0.5M H₂SO₄. The percentage of nitrogen in the compound is a) 28 b) 56 d) 112 944. Stereoisomers (geometrical or opticals) which are neither superimposable nor mirror image to each other are called: a) Enantiomers b) Mesomers c) Tautomers d) Diastereomers 945. Which one of the following will show optical isomerism? Η CH_3 CH_3 c) $H_3C - C - CO_2H$ a) $HO - C - CO_2H$ b) $H_3C - C - CO_2H$ d) $H_3C - C - CO_2H$ Cl OH H 946. The ion formed by the reaction of HNO_2 and H_2SO_4 is a) Nitronium ion b) Nitrosonium ion c) Nitrite ion d) Nitrate ion 947. Chloroacetic acid is a stronger acid than acetic acid. This can be explained using a) -M effect b) -l effect c) +Meffect d) +I effect 948. The basicity of aniline is less than that of cyclohexylamine. This is due to a) +R effect of - NH2 group b) -I effect of - NH2 group c) -R effect of -NH2 group d) Hyperconjugation effect 949. The compound is an example of: a) Aromatic compound b) Heterocyclic compound c) Annulene d) Xanthates 950. Dehydration of alcohol usually goes by a) E1 mechanism b) E2 mechanism c) E1 cb mechanism d) S_N2 mechanism 951. Geometrical isomerism is possible in b) Isobutene c) Acetophenone-oxime d) Benzophenone-oxime a) Acetone-oxime 952. Ethers are isomeric with a) Aldehydes b) Ketones c) Both aldehydes and ketones d) Alcohols 953. S_N1 reaction is fastest in CH_3 $CH_3 - CH - Cl$ $CH_3 - C - CI$ a) CH₃CH₂Cl d) CH_3 CH_2 CH_3 954. Vinyl chloride undergoes a) Only addition reactions b) Only elimination reactions c) Both (a) and (b) d) Substitution reactions 955. Fischer projection indicates: a) Horizontal substituents above the plane b) Vertical substituents above the plane

c) Both horizontal and vertical substituents below the planed) Both horizontal and vertical substituents above the plane

956. The reaction,



Is an example of

- a) Nucleophilic substitution
- b) Electrophilic addition
- c) Elimination reaction
- d) Nucleophilic addition
- 957. Acetone and propen-2-ol are:
 - a) Positional isomers b) Leto-enol tautomers c) Geometrical isomers d) Chain isomers
- 958. The number of stereoisomers obtained by bromination of trans 2 -butene is?
 - a) I

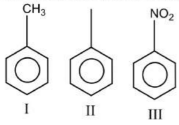
b) 2

c) 3

- d) 4
- 959. The compound which forms one monochloro product when treated with chlorine is:
 - a) n-pentane
- b) Isopentane
- c) neo-pentane
- d) None of these

- 960. Reactivity towards nucleophilic addition reaction of
 - (I)HCHO (II)CH3CHO (III)CH3COCH3is
 - a) II>III>I
- b) III>II>I
- c) I>II>III
- d) I>II<III

- 961. Maleic acid and fumaric acid are
 - a) Position isomers
- b) Geometric isomers
- c) Enantimoers
- d) Functional isomers
- 962. The ease of nitration of the following three hydrocarbons follows the order



- a) II=III≈I
- b) II>III>I
- c) III>II>I
- d) I=III>II

- 963. Which represents the condensed formula for pentanes?
 - a) CH₃(CH₂)₃CH₃
- b) $(CH_3)_3CCH_3$
- c) (CH₃)₂CHCH₂CH₃
- d) All of these

- 964. Which of the substance is purified by sublimation?
 - a) Benzoic acid
- b) Camphor
- c) Naphthalene
- d) All of these
- 965. The halogen compound which most readily undergoes nucleophilic substitutions is
 - a) $CH_2 = CHCl$

b) $CH_3CH = CHCl$

c) $CH_2 = CHC(Cl) = CH_2$

- d) $CH_2 = CHCH_2Cl$
- 966. Which of the following order is correct regarding the acidity of carboxylic acids?
 - a) Cl₃CCOOH > Cl₂CHCOOH > ClCH₂COOH
- b) Cl₃CCOOH > Cl₂CHCOOH < ClCH₂COOH
- c) Cl₃CCOOH < Cl₂CHCOOH > ClCH₂COOH
- d) Cl₃CCOOH < Cl₂CHCOOH < ClCH₂COOH
- 967. An $S_N 2$ reaction at an asymmetric carbon of a compound always gives
 - a) A mixture of diastereomers

- b) A single stereoisomer
- c) An enantiomer of the substrate
- d) A product with opposite optical rotation
- 968. The IUPAC name of the compound, $CH_3CH = CHC \equiv CH$ is:
 - a) Pent-4-yn-2-ene
- b) Pent-3-en-1-yne
- c) Pent-2-en-4-yne
- d) Pent-1-yn-3-ene
- 969. Reaction of methyl bromide with aqueous sodium hydroxide involves
 - a) Racemisation

b) S_N1 mechanism

c) Retention of configuration

- d) $S_N 2$ mechanism
- 970. An organic compound X(mol. formula $C_6H_5O_2N$) has six carbons in a ring system, three double bonds and also a nitro group as substituent. X is :
 - a) Homocyclic but not aromatic





- b) Aromatic but not homocyclic c) Homocyclic and aromatic d) heterocyclic
- 971. The compounds CH₃NH₂ and CH₃CH₂. NH₂ are:
 - a) Isomers
- b) Isobars
- c) Homologous
- d) Allotropes
- 972. The following compound will undergo electrophilic substitution more readily than benzene
 - a) Nitrobenzene
- b) Benzoic acid
- c) Benzaldehyde
- d) Phenol
- 973. Which of the following elements can't be detected by direct tests?
 - a) N

c) S

d) Br

974. IUPAC name of,

- a) 2-chloromethyl-4-methyl-hexanal
- b) 1-chloro-4-ethyl-2-pentanal
- c) 1-chloro-4-methyl-2-hexanal
- d) 1-chloro-2-aldo-4-methyl hexane
- 975. Position isomerism is shown by:
 - a) o-nitrolhenol and p-nitrophenol
 - b) Dimethyl ether and ethanol
 - c) Pentan-2-one and pentan-3-one
 - d) Acetaldehyde and acetone
- 976. Formulae of phenyl carbinol and chloral are respectively:
 - a) C₆H₅. CH₂CH₂OH and CHCl₂CHO
 - b) C₆H₅CH₂OH and CCl₃CHO
 - c) C₆H₅OH and CH₂Cl. CHO
 - d) C₆H₅CHO and CHCl₂CHO
- 977. How many primary carbon atoms are there in the compound,

a) 6

c) 4

d) 3

978. IUPAC name of,

- a) 4-butyl-2,5-hexadien-l-al
- b) 5-vinyloct-3-en-l-al
- c) 5-vinyloct-5-en-8-al
- d) 3-butyl-1,4-hexadien-6-al
- 979. The molecular formula of a saturated compound is C₂H₄Br₂.

This formula permits the existence of:

- a) Functional isomers
- b) Optical isomers
- c) Positional isomers
- d) cis trans isomers

- 980. Which of the following solvents are aprotic?
 - $(A)NH_3$
- (B)SO₂

(D)CH3CO2H

- (C)CH₃CN a) A,B,C
- b) A,C,D
- c) B,C
- d) A,C
- 981. The reaction of sodium ethoxide with iodoethane to from diethyl is termed as
 - a) Electrophilic substitution

b) Nuclephilic substitution

c) Electrophilic addition

d) Radical substitution



	k precipitate. The organic cor		d and addition of lead acetate
	[201] : 20 min 및 [발생시장으로 발표 - 12 min (2016) (2016) (2016) (2016)	3 4 TO 100 100 ME ON SECURE OF THE SECURE OF	d) Phoophomic
a) Nitrogen	b) Halogen	c) Sulphur	d) Phosphorus
	the compound CH ₃ CONHBr is		L DN GI
a) 1-bromoacetamid	그 그 그 그 그 그 그 그 그 그 그 그 그 그 그 그 그 그 그	c) N-bromoethanamic	-
	onobasic acid on ignition gav	370	75 A
a) 37	b) 57	c) 73	d) 88
985. The IUPAC name of o	[
$C_2H_5 - C - CH_2OH$	is		
CH ₂			
a) 2-ethylprop-2-en-		b) 2-hydroxymethylbi	
c) 2-methylenebutar		d) 2-ethyl-3-hydroxyp	prop-1-ene
	active stereoisomers are pos		
a) 0	b) 1	c) 2	d) 3
	e ion from a methane molecu		
 a) Methyl radical 	b) Carbonium ion	c) Carbanion	d) Methyl group
988. Which of the followi	ng will have a <i>meso-</i> isomer a	lso?	
a) 2-chlorobutane		b) 2, 3-dichlorobutane	e
c) 2, 3-dichloropenta	ane	d) 2-hydroxypropano	ic acid
989. Which chlorine atom	n is more electronegative in th	ne following?	
		CH ₃	CH ₃
		1	1
a) CH ₃ - Cl	b) $CH_3 - CH_2 - Cl$	c) $H - C - Cl$	d) $CH_3 - CH_2 - C - Cl$
a) CH ₃ — Cl	b) $CH_3 - CH_2 - CI$	c) H — C — Cl 	d) CH ₃ – CH ₂ – C – Cl
a) CH ₃ — Cl	b) $CH_3 - CH_2 - CI$	c) H – C – Cl CH ₃	d) $CH_3 - CH_2 - C - Cl$ CH_3
990. The resonating struc	ctures :	1	d) CH ₃ – CH ₂ – C – Cl CH ₃
990. The resonating struct	ctures : arrangement of electrons	CH ₃	d) CH ₃ – CH ₂ – C – Cl CH ₃
990. The resonating struction a) Differ only in the b) Differ in number of	ctures : arrangement of electrons of paired and unpaired electr	CH ₃	d) CH ₃ - CH ₂ - C - Cl CH ₃
990. The resonating struct a) Differ only in the b) Differ in number c c) Differ largely in the	ctures : arrangement of electrons of paired and unpaired electr neir energy contents	CH ₃	d) CH ₃ – CH ₂ – C – Cl CH ₃
990. The resonating struction a) Differ only in the b) Differ in number of	ctures : arrangement of electrons of paired and unpaired electr neir energy contents	CH ₃	d) CH ₃ - CH ₂ - C - Cl CH ₃
990. The resonating struction a) Differ only in the b) Differ in number c) Differ largely in the d) Do not lie in the s	ctures : arrangement of electrons of paired and unpaired electr neir energy contents	CH ₃	d) CH ₃ – CH ₂ – C – Cl CH ₃
990. The resonating struct a) Differ only in the second by Differ in number of c) Differ largely in the d) Do not lie in the second conformer	ctures : arrangement of electrons of paired and unpaired electroneir energy contents ame plane which are not enantiomers, a b) Diastereomer	CH ₃	d) $CH_3 - CH_2 - C - Cl$ CH_3
990. The resonating struct a) Differ only in the second biffer in number of c) Differ largely in the d) Do not lie in the second biffer largely in the second bif	ctures : arrangement of electrons of paired and unpaired electroneir energy contents ame plane which are not enantiomers, a b) Diastereomer	CH ₃	CH ₃
990. The resonating struct a) Differ only in the second by Differ in number of c) Differ largely in the d) Do not lie in the second conformer	ctures : arrangement of electrons of paired and unpaired electroneir energy contents ame plane which are not enantiomers, a b) Diastereomer	CH ₃	CH ₃
 990. The resonating structure a) Differ only in the second point of the second point of	ctures : arrangement of electrons of paired and unpaired electroneir energy contents ame plane which are not enantiomers, a b) Diastereomer	CH ₃	CH ₃
 990. The resonating struct a) Differ only in the struct b) Differ in number of c) Differ largely in the d) Do not lie in the structure 991. The optical isomers, a) Conformer 992. α-D-(+)-glucose and a) Enantiomers 	ctures : arrangement of electrons of paired and unpaired electroneir energy contents ame plane which are not enantiomers, a b) Diastereomer	CH ₃	CH ₃
 990. The resonating structure a) Differ only in the set of Differ largely in the description b) Differ largely in the set of Do not lie in the	ctures : arrangement of electrons of paired and unpaired electroneir energy contents ame plane which are not enantiomers, a b) Diastereomer	CH ₃	CH ₃
990. The resonating struct a) Differ only in the b) Differ in number c c) Differ largely in the d) Do not lie in the second isomers, a) Conformer 992. α-D-(+)-glucose and a) Enantiomers b) Conformers c) Epimers d) Anomers	ctures : arrangement of electrons of paired and unpaired electroneir energy contents ame plane which are not enantiomers, a b) Diastereomer	CH ₃ ons are called c) Mirror images	CH ₃
 990. The resonating structure a) Differ only in the set of Differ largely in the d) Do not lie in the set of Differ largely in the set of Diff	etures : arrangement of electrons of paired and unpaired electroneir energy contents ame plane which are not enantiomers, a b) Diastereomer I β-D-(+)-glucose are :	CH ₃ ons are called c) Mirror images	CH ₃
 990. The resonating structure a) Differ only in the structure b) Differ in number of c) Differ largely in the d) Do not lie in the structure 991. The optical isomers, a) Conformer 992. α-D-(+)-glucose and a) Enantiomers b) Conformers c) Epimers d) Anomers 993. Which of the following a) CH₃CH₂CH(Cl)CO 	etures : arrangement of electrons of paired and unpaired electroneir energy contents ame plane which are not enantiomers, a b) Diastereomer I β-D-(+)-glucose are :	CH ₃ ons ons ore called c) Mirror images g the acidity of carboxylic g	CH ₃
 990. The resonating struct a) Differ only in the struct b) Differ in number of c) Differ largely in the d) Do not lie in the struct 991. The optical isomers, a) Conformer 992. α-D-(+)-glucose and a) Enantiomers b) Conformers c) Epimers d) Anomers 993. Which of the following a) CH₃CH₂CH(Cl)CO b) CH₃CH₂CH(Cl)CO 	etures: arrangement of electrons of paired and unpaired electrons neir energy contents ame plane which are not enantiomers, a b) Diastereomer I β-D-(+)-glucose are:	CH ₃ ons ons one called c) Mirror images g the acidity of carboxylic g CICH ₂ CH ₂ CH ₂ COOH	CH ₃
 990. The resonating struct a) Differ only in the struct b) Differ in number of c) Differ largely in the struct d) Do not lie in the struct 991. The optical isomers, a) Conformer 992. α-D-(+)-glucose and a) Enantiomers b) Conformers c) Epimers d) Anomers 993. Which of the following a) CH₃CH₂CH(Cl)CO b) CH₃CH₂CH(Cl)CO c) CH₃CH₂CH(Cl)CO 	etures: arrangement of electrons of paired and unpaired electrons neir energy contents ame plane which are not enantiomers, a b) Diastereomer I β-D-(+)-glucose are: ong orders is correct regarding OH > CH ₃ CH(Cl)CH ₂ COOH > OH < CH ₃ CH(Cl)CH ₂ COOH <	CH ₃ ons are called c) Mirror images g the acidity of carboxylic g CICH ₂ CH ₂ CH ₂ COOH CICH ₂ CH ₂ CH ₂ COOH	CH ₃
 990. The resonating struct a) Differ only in the struct b) Differ in number of c) Differ largely in the struct d) Do not lie in the struct 991. The optical isomers, a) Conformer 992. α-D-(+)-glucose and a) Enantiomers b) Conformers c) Epimers d) Anomers 993. Which of the following a) CH₃CH₂CH(Cl)CO b) CH₃CH₂CH(Cl)CO c) CH₃CH₂CH(Cl)CO 	etures: arrangement of electrons of paired and unpaired electrons neir energy contents ame plane which are not enantiomers, a b) Diastereomer I β-D-(+)-glucose are: ong orders is correct regarding OH > CH ₃ CH(Cl)CH ₂ COOH > OH < CH ₃ CH(Cl)CH ₂ COOH < OH > CH ₃ CH(Cl)CH ₂ COOH <	CH ₃ ons are called c) Mirror images g the acidity of carboxylic g CICH ₂ CH ₂ CH ₂ COOH CICH ₂ CH ₂ CH ₂ COOH	CH ₃
 990. The resonating struct a) Differ only in the struct b) Differ in number of c) Differ largely in the d) Do not lie in the struct 991. The optical isomers, a) Conformer 992. α-D-(+)-glucose and a) Enantiomers b) Conformers c) Epimers d) Anomers 993. Which of the following a) CH₃CH₂CH(Cl)CO b) CH₃CH₂CH(Cl)CO c) CH₃CH₂CH(Cl)CO d) CH₃CH₂CH(Cl)CO 	etures: arrangement of electrons of paired and unpaired electrons neir energy contents ame plane which are not enantiomers, a b) Diastereomer I β-D-(+)-glucose are: ong orders is correct regarding OH > CH ₃ CH(Cl)CH ₂ COOH > OH < CH ₃ CH(Cl)CH ₂ COOH < OH > CH ₃ CH(Cl)CH ₂ COOH <	CH ₃ ons are called c) Mirror images g the acidity of carboxylic g CICH ₂ CH ₂ CH ₂ COOH CICH ₂ CH ₂ CH ₂ COOH	CH ₃



$$R$$
 $CH_2Br + N$ $Br^ CH_2 - N$ Br^-

is influenced by the hyper conjugation effect of group R. If R sequentially is

$$I.CH_3 -$$

II.
$$CH_3 - CH_2 - H_3C - CH - CH_3$$

the increasing order of speed of the above reaction is

- a) IV, III, II, I
- b) I, II, III, IV
- c) I, IV, III, II
- d) III, II, I, IV
- 995. The compound, whose stereo-chemical formula is written below, exhibits x geometrical isomers and y optical isomers

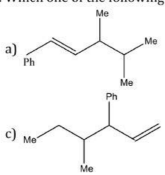
The values of x and y are

- a) 4 and 4
- b) 2 and 2
- c) 2 and 4
- d) 4 and 2

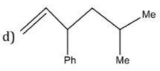
996. Geometrical isomerism is shown by

- a) -C-C-
- b) >c=<
- c) $-C \equiv C -$
- d) None of these

997. Which one of the following is s-butyl phynylvinyl methane?



b)



998. Arrange the carbanions,

(CH₃)₃C̄, CCl₃, (CH₃)₂CH, C₆H₅CH₂, in order of their decreasing stability

- a) $C_6H_5\bar{C}H_2 > \bar{C}CI_3 > (CH_3)_2\bar{C} > (CH_3)_2\bar{C}H$
- c) $\bar{C}Cl_3 > C_6H_5\bar{C}H_2 > (CH_3)_2\bar{C}H > (CH_3)_3\bar{C}$
- b) $(CH_3)_2\bar{C}H > \bar{C}Cl_3 > C_6H_5CH_2 > (CH_3)_3\bar{C}$
- d) $(CH_3)_3\overline{C} > (CH_3)_2\overline{C}H > \overline{C}H_2 > \overline{C}Cl_3$
- 999. RX + $I^- \rightarrow R I + X^-$ is an example of ... reaction.
 - a) Nucleophilic addition
 - c) Electrophilic addition

- b) Nucleophilic substitution
- d) Elimination







100 Bicyclo (1,1,0) butane is







100 The basic strength of

1.
$$CH \equiv \overline{C}, CH_2 = \overline{C}H, CH_3\overline{C}H_2$$

 $I \qquad II \qquad III$

Will be in order

- I II
- III a) I < II < III
 - b) II<III<I
- c) III<II<I
- d) III<I<II

100 Which of the following is most reactive towards elimination reaction?

- a) RCOO-
- c) NO₃
- d) RO⁻

100 IUPAC name of CH_3 — CH_2 —CH— NH_2 is :

- a) 1-methyl-1-aminopropane b) 2-aminobutane
- c) 2-methyl-3-aminopropane
- d) None of the above

100 The number of isomeric hexanes is

4.

a) 5

b) 2

c) 3

d) 4

100 The substitution reaction among the following is

$$_{b)}$$
 >C=O + NaHSO $_3$ --> $_{\rm C}$ $_{\rm SO_3Na}$

c)
$$CH_3$$
 $C-OH + \frac{Dry \ HCI/Anhy. \ ZnCl_2}{Lucas \ reagent} CH_3$ CH_3 $C-C$

d)
$$(CH_3)_2C = CH_2 + BrCl \rightarrow (CH_3)_2C - CH_2$$

100 The most stable carbocation is

6.









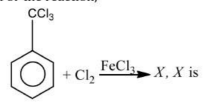
100 Among the following alkenes (I) 1-butene, (II) cis-2-butene, (III) trans-2-butene the decreasing order of

stability is:

- a) III > II > I
- b) II > I > II
- c) I > II > III
- d) II > I > III

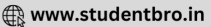
100 For the reaction,

8.



- a) Chloro benzene and carbon tetrachloride
- b) meta chloro benzotrichloride





c) ortho, para chloro benzotrichloride d) None of the above 100 Which of the following statements is not correct? a) A \searrow C == C \swarrow group is made up of 4σ -bond and 2π -bonds b) A σ -bond is stronger than π -bond c) A σ -bond can exist independently of π -bond d) A double bond is stronger than a single bond 101 The number of sp^3 - hybrid carbons in 2-butyne is: a) 4 b) 3 c) 2 d) 1 101 How many π -electrons are there in following? HC - CHHC CH d) 8 a) 2 c) 6 101 2. IUPAC name of a) Dimethyl amine b) 2-amino propane c) Isopropylamine d) 2-propanamine 101 An organic compound having molecular mass 60 is found to contain C=20%, H=6.67% and N=46.67% while rest is oxygen. On heating it gives NH₃ along with a solid residue. The solid residue gives violet colour with alkaline copper sulphate solution. The compound is a) CH3CH2CONH2 b) (NH₂)₂CO c) CH₃CONH₂ d) CH₃NCO 101 How many chiral carbon atoms are present in 2, 3, 4- trichloropentane? c) 2 d) 3 a) 4 b) 1 101 Which one of the following compounds is most polar? 5. b) CH₂F₂ c) CH₂Cl₂ d) CH₂Br₂ a) CH₂I₂ 101 Geometrical isomerism is not shown by 6. a) 1, 1-dichloro-1-pentene b) 1,2-dichloro-1-pentene c) 1, 3-dichloro-2-pentene d) 1, 4-dichloro-2-pentene 101 The change in optical rotation with time of freshly prepared solution of sugar is known as: a) Specific rotation b) Inversion c) Rotatory motion d) Mutarotation 101 Which of the following does not show stereo isomerism?



a)
$$CH_3$$
 CO CH_3 CH_3 CO CH_3 CH_3 CO CH_3

101 One of the following compounds exhibit geometrical iosmerism

9.

b)
$$CH_3 - HC(CH_3) - H(C)CH_3 - CH_3$$

c)
$$CH_3 - HC(CH_3) - CH_3$$

d)
$$CH_3CH = CH - CH_3$$

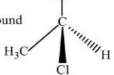
102 Which one of the following shows functional isomerism?

U.

b)
$$C_3H_6$$

102

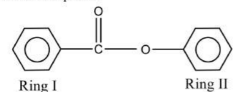
1. The chirality of the compound



a) R

102 In the compound

2.



electrophilic substitution occurs at

a) ortho/para position at ring I

b) meta position at ring I

c) ortho/para position at ring II

d) meta position at ring II

102 In the reaction of phenol with chloroform and aqueous solution of NaOH at 70°C, the electrophile attacking

3. the ring is

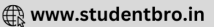
- a) CHCl₃
- b) CHCl₂
- $c): CCl_2$
- d) COCl2

102 The production of an optically active compound from a symmetric molecule without resolution in termed

4. as

- a) Walden inversion
- b) Partial racemisation
- c) Asymmetric synthesis
- d) Partial resolution





102 5.	An organic compound on	heating with CuO produce	es CO ₂ but no water. The or	ganic compound may be
Ů,	a) Carbon tetrachloride	b) Chloroform	c) Methane	d) Ethyl iodide
102	Which of the following st	atement is not applicable t	to Beilstein test?	
6.				
	a) Green or bluish green	flame is due to the formati	on of volatile cupric halides	i
	b) It does not tell us to w	hich halogen is present in	the organic compound	
	c) It is very sensitive test	can be easily performed		
	d) It is a sure test for the	presence of halogen		
102	Essential oils can be isola	ited by		
7.				
	a) Crystallization	b) Steam distillation	c) Sublimation	d) Distillation
102	Mesomeric effect involve	s delocalisation of		
8.				
	a) Pi-electrons	b) Sigma electrons	c) Protons	d) None of these
	The IUPAC name of the c	ompound,		
9.	HO			
	is:			
	a) 1,2-dimethyl-2-buteno	ol		
	b) 3-methylpent-3-en-2-			
	c) 3,4-dimethyl-2-buten-			
	d) 2,3-dimethyl-3-penter	nol		
103	Which of the following sp	oecies is paramagnetic in n	ature?	
0.				
	a) Carbonium ion	b) Free radical	c) Carbene	d) Nitrene
103	Isobutyl chloride is:			
1.				
	a) CH ₃ CH ₂ CH ₂ CH ₂ Cl	b) (CH ₃) ₂ CHCH ₂ Cl	c) CH ₃ CH ₂ CHClCH ₃	d) $(CH_3)_3 C - Cl$
	How many isomers will (C ₃ H ₆ have?		
2.				
	a) 1	b) 2	c) Zero	d) 4
	Which one of the following	ng compounds is capable o	f existing in a <i>meso</i> form?	
3.	- No. o. dul		A.N.A. Norway and property and and a second	
	a) 3, 3-dibromopentane		b) 4-bromo-2-pentanol	
100	c) 3-bromo-2-pentanol	,	d) 2, 3-dibromopentane	
	Geometrical isomerism is	s caused :		
4.	a) Drawatulated ustation	anaund C — Chand		
	a) By restricted rotation	around C = C bond e asymmetric carbon atom		
	3) (B) (B)	s attached to the same fun	ctional group	
	15)	atom between two polyva		
103	Lassaigne's test is used for		ient atoms	
5.	Lassaighe's test is used it	of the detection of		
٥.	a) Carbon only		b) Hydrogen only	
	c) Oxygen only		d) Nitrogen, sulphur and	halogens
103		arranged according to the		
6.		goa according to the		
38	a) Electrophile			
	b) Electrophile			



103 7.	d) Electrophile – Br^+ , N_3^-	N ₃ . Nucleophile – NO ₂ +, Br ⁺ , Nucleophile – <i>CH₃OH</i> , lkynes with molecular forn											
50.50	a) 3	b) 4	c) 5	d) 6									
103 8.	Example of geometrical is												
	a) 2-butanol	b) 2-butene	c) Butanal	d) 2-butyne									
	Which of the following is	an example of elimination i	reaction?										
9.													
	a) Chlorination of methane												
	b) Dehydration of ethanol												
	c) Nitration of benzene												
104	d) Hydroxylation of ethylene O4 The order of stability of the following carbanion is												
	1	ne following carballion is											
U.	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$												
	I II III	IV											
	a) I>II>III>IV	b) I>III>IV	c) IV>III>II>I	d) III>IV>I>II									
104	Which nomenclature in IU	JPAC is not correct?											
1.													
	a) Pentyne-3	b) Pentyne-2	c) Hexyne-3	d) None of these									
	The C—C bond angle in cy												
2.	0.7-200	82 25 25 25 25 25 25 25 25 25 25 25 25 25											
170 17	a) 60°	b) 120°	c) 109°28′	d) 180°									
	Absolute alcohol cannot b	e obtained by simple fracti	onal distillation because										
3.) D												
	a) Pure C ₂ H ₅ OH is unstab												
	b) C ₂ H ₅ OH forms hydroge		nton										
	101 2 TO 10 10 10 10 10 10 10 10 10 10 10 10 10	H is very close to that of wa ropic mixture is formed wit											
104	Racemic compound has	ropic illixture is formed wit	iii watei										
4.	Racellic compound has												
1.	a) Equimolar mixture of e	enantiomers	b) 1:1 mixture of enantion	ner and diastereomer									
	c) 1:1 mixture of diastere		d) 1:2 mixture of enantion										
104	[[[]][[]][[]][[][][][][[][][][][][][][from the preceding or the s										
5.	0		ı	0									
	a) a CH ₂ group	b) a CH ₃ group	c) Two hydrogen atoms	d) Four hydrogen atoms									
104	Vaccum distillation is use	d to purify liquids which	10 E 15%										
6.													
	a) Are highly volatile		b) Are explosive in nature	2									
	c) Decompose below their		d) Have high boiling point	t .									
	The number of isomeric s	tructures for C ₂ H ₇ N would	be:										
7.													
	a) 4	b) 3	c) 2	d) 1									
	The IUPAC name of CH ₃ -	- CH ₂ – CHO											
8.	-) D14	b) 2	-) Dut 1-1	J) Dt 1 4									
	a) Propanal-1	b) 2-methylbutanal	c) Butanal-1	d) Pentanal-1									

104 Which of the following is the weakest base?

a) Ethyl amine

b) Ammonia

c) Dimethyl amine

d) Methyl amine

105 The structure which has positive charge on the oxygen atom:

н**—**Ö—н а) | Н

b) CH₃—Ö—CH₃ c) CH₃—Ö:

d) н—Ö—Ö—н



ORGANIC CHEMISTRY - SOME BASIC PRINCIPLES AND TECHNIQUES

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

313)	c	314)	b	315)	d	316) a	[513]	b	514)	b	515)	a	516)	d
317)	c	318)	b	319)	b	320) ł	517)	c	518)	d	519)	a	520)	d
321)	a	322)	a	323)	b	324) 0	521)	d	522)	b	523)	b	524)	c
325)	c	326)	d	327)	b	328)	525)	С	526)	a	527)	d	528)	d
329)	d	330)	a	331)	d		529)	a	530)	С	531)	b	532)	a
333)	c	334)	d	335)	b	336) 0		b	534)	a	535)	b	536)	b
337)	d	338)	b	339)	b		537)	d	538)	b	539)	a	540)	d
341)	b	342)	c	343)	d		541)	С	542)	c	543)	d	544)	b
345)	b	346)	С	347)	a		545)	d	546)	b	547)	a	548)	a
349)	d	350)	a	351)	b		549)	b	550)	a	551)	С	552)	d
353)	b	354)	b	355)	c		553)	d	554)	d	555)	a	556)	d
357)	b	358)	c	359)	a		557)	b	558)	a	559)	b	560)	d
361)	a	362)	c	363)	a	364) a	1	b	562)	a	563)	a	564)	b
365)	b	366)	b	367)	d		565)	b	566)	a	567)	b	568)	b
369)	С	370)	c	371)	a	372) d		С	570)	b	571)	b	572)	c
373)	b	374)	b	375)	b		573)	b	574)	b	575)	d	576)	d
377)	b	378)	c	379)	d	380) (l	a	578)	d	579)	b	580)	a
381)	d	382)	a	383)	c	384)	1	d	582)	d	583)	b	584)	b
385)	c	386)	a	387)	b		585)	d	586)	d	587)	b	588)	b
389)	b	390)	b	391)	d	392) a	l	a	590)	a	591)	С	592)	a
393)	d	394)	a	395)	b	D. C.	593)	С	594)	a	595)	b	596)	c
397)	c	398)	c	399)	a		597)	a	598)	b	599)	С	600)	d
401)	d	402)	b	403)	c	404)	1	С	602)	a	603)	С	604)	a
405)	b	406)	a	407)	a	408) h	1	a	606)	d	607)	a	608)	c
409)	b	410)	d	411)	С		609)	b	610)	d	611)	С	612)	d
413)	b	414)	b	415)	b		613)	c	614)	a	615)	d	616)	d
417)	a	418)	c	419)	c	420)		С	618)	a	619)	c	620)	c
421)	С	422)	a	423)	С		621)	a	622)	d	623)	a	624)	d
425)	a	426)	d	427)	d	428) a	1	b	626)	b	627)	С	628)	a
429)	c	430)	b	431)	b	432) a		d	630)	b	631)	С	632)	c
433)	a	434)	a	435)	d	436)		d	634)	c	635)	d	636)	b
437)	d	438)	c	439)	a		637)	d	638)	d	639)	d	640)	a
441)	c	442)	d	443)	a		641)	b	642)	a	643)	d	644)	a
445)	c	446)	d	447)	c	-	645)	a	646)	b	647)	С	648)	a
449)	c	450)	c	451)	d		649)	b	650)	a	651)	b	652)	a
453)	d	454)	b	455)	d		653)	d	654)	d	655)	a	656)	a
457)	b	458)	c	459)	С		657)	b	658)	С	659)	b	660)	b
461)	d	462)	С	463)	a	WORKS ***	661)	b	662)	С	663)	c	664)	c
465)	b	466)	b	467)	c		665)	b	666)	a	667)	b	668)	c
469)	a	470)	d	471)	a		669)	d	670)	a	671)	С	672)	a
473)	a	474)	d	475)	a		673)	b	674)	c	675)	c	676)	b
477)	d	478)	b	479)	a		677)	b	678)	b	679)	b	680)	b
481)	С	482)	d	483)	a	여러하다라 ""	681)	а	682)	a	683)	b	684)	d
485)	С	486)	b	487)	c		685)	a	686)	a	687)	C	688)	d
489)	a	490)	a	491)	d		689)	d	690)	c	691)	b	692)	d
493)	c	494)	d	495)	d		693)	a	694)	c	695)	c	696)	a
497)	С	498)	a	499)	b	Strangerson	697)	d	698)	a	699)	d	700)	c
501)	a	502)	d	503)	d		701)	b	702)	a	703)	b	704)	b
505)	b	506)	b	507)	c		705)	b	706)	c	707)	a	708)	d
509)	a	510)	c	511)	a		709)	b	710)	c	711)	b	712)	a
						107	1 ,							

713)	d	714)	c	715)	b	716) c	885)	d	886)	c	887)	c	888)	a
717)	b	718)	a	719)	a	720) b	889)	b	890)	b	891)	b	892)	C
721)	b	722)	c	723)	d	724) b	893)	d	894)	c	895)	c	896)	c
725)	b	726)	C	727)	a	728) a	897)	a	898)	d	899)	С	900)	a
729)	b	730)	b	731)	a	732) a	901)	c	902)	c	903)	d	904)	c
733)	a	734)	b	735)	a	736) d	905)	c	906)	a	907)	c	908)	d
737)	a	738)	a	739)	a	740) d	909)	c	910)	d	911)	d	912)	a
741)	c	742)	c	743)	b	744) c	913)	d	914)	a	915)	b	916)	d
745)	c	746)	a	747)	b	748) a	917)	d	918)	b	919)	b	920)	c
749)	c	750)	a	751)	d	752) a	921)	b	922)	b	923)	c	924)	d
753)	b	754)	d	755)	C	756) b	925)	c	926)	a	927)	a	928)	b
757)	a	758)	b	759)	c	760) a	929)	a	930)	a	931)	b	932)	d
761)	a	762)	a	763)	c	764) d	933)	a	934)	a	935)	b	936)	d
765)	b	766)	b	767)	c	768) b	937)	a	938)	d	939)	a	940)	a
769)	b	770)	a	771)	C	772) c	941)	a	942)	b	943)	b	944)	d
773)	b	774)	a	775)	b	776) c	945)	b	946)	b	947)	b	948)	a
777)	d	778)	a	779)	C	780) c	949)	c	950)	a	951)	c	952)	d
781)	b	782)	a	783)	a	784) c	953)	c	954)	C	955)	a	956)	b
785)	C	786)	C	787)	d	788) a	957)	b	958)	a	959)	c	960)	C
789)	b	790)	d	791)	a	792) b	961)	b	962)	b	963)	d	964)	d
793)	c	794)	a	795)	a	796) a	965)	d	966)	a	967)	b	968)	b
797)	d	798)	d	799)	a	800) c	969)	d	970)	C	971)	c	972)	d
801)	c	802)	b	803)	d	804) d	973)	b	974)	a	975)	a	976)	b
805)	c	806)	d	807)	c	808) d	977)	a	978)	a	979)	c	980)	a
809)	c	810)	a	811)	c	812) c	981)	b	982)	c	983)	c	984)	c
813)	d	814)	C	815)	a	816) a	985)	a	986)	d	987)	b	988)	b
817)	d	818)	a	819)	d	820) b	989)	d	990)	a	991)	b	992)	d
821)	b	822)	b	823)	d	824) d	993)	a	994)	a	995)	b	996)	b
825)	b	826)	d	827)	C	828) c	997)	C	998)	C	999)	b	1000)	C
829)	b	830)	d	831)	C	832) d	1001)	a	1002)	d	1003)	b	1004)	a
833)	b	834)	C	835)	b		1005)		1006)		1007)		1008)	
837)		838)			C	840) c			1010)		1011)		1012)	
841)	C	842)	C	843)	b		1013)		1014)		1015)		1016)	
845)	C	846)	c	847)	С		1017)		1018)		1019)		1020)	
849)	C	15	a	851)	b		1021)		1022)		1023)		1024)	
853)	C	854)	С	855)	b		1025)		1026)		1027)		1028)	
857)	C	858)	a	859)	d		1029)		1030)		1031)		1032)	
861)	b	862)	d	863)	d	5	1033)		1034)		1035)		1036)	
865)	b	866)	d	867)	b		1037)		1038)		1039)		1040)	
869)	d	870)	d	871)	b		1041)		1042)		1043)		1044)	
873)	a	874)	b	875)	d		1045)		1046)		1047)	С	1048)	a
877)	d	878)	С	879)	d		1049)	b	1050)	a				
881)	C	882)	C	883)	a	884) d								
							ı							



ORGANIC CHEMISTRY - SOME BASIC PRINCIPLES AND TECHNIQUES

: HINTS AND SOLUTIONS :

- C_6H_5 CH = CHCOOH is cinnamic acid.
- 2 Draw position and chain isomers.
- 3 Strain = $\frac{1}{2}$ [Normal valence angel-valence angel]

 $= \frac{1}{2} \left[109^{\circ}28' - 60^{\circ} \right] = 24^{\circ}44'.$

- 6 (d) 2, 2-dimethyl butane is 6-carbon hydrocarbon (C_6H_{14})
 - CH_3 $CH_3 - C - CH_2 - CH_3$

Rest all are the chain isomers of pentane (C_5H_{12}).

 $H_3C - C - CH_3$ CH_3

2,2-dimethyl propane

(neo-pentane)

 CH_3 $H_3C - CH - CH_2 - CH_2$ 2- methyl butane (iso -pentane)

CH3CH2CH2CH2CH3 n-pentane

- (b) Follow IUPAC rules.
- 8 Detection of sulphur in sodium extract is done by lead acetate and sodium nitroprusside $Na_2S + (CH_3COO)_2Pb \rightarrow PbS + 2CH_3COONa$ lead acetate black ppt. $Na_2S + Na_2[Fe(CN)_5NO] \rightarrow Na_4[Fe(CN)_5NOS]$

Sodium nitroprusside sodium thio nitroprusside

(purple

colour)

- (b) -do -
- 10 (a) Cyclopropyl methyl carbocations are more stable than benzyl carbocations due to conjugation between bent orbitals of cyclopropyl group.
- 11 (c) The shape of π electron cloud in acetylene in cylindrical
- 12 (c) Acidified sodium fusion extract on addition of ferric chloride solution gives blood red colouration, which confirms the presence of N and

 $3NaCNS + aq. FeCl_3 \rightarrow Fe(CNS)_3 + 3NaCl$ (ferric thiocyanide)

13 **(b)** NaOH chlorobenzene phenol

> In this process one group is replaced by other, hence, it is a substitution process and both the leaving and attacking groups are nucleophilic, therefore it is an example of nucleophilic substitution reaction.

14 (b) (i) Fe(CNS)₃ is red in colour and is formed when both N and S are present in organic compound (ii) $Na_2S + Na_2[Fe(CN)_5NO] \rightarrow$ $[NaFe(C)_5NOS]$ from organic sodium violet colour



compound nitroprusside

15 (a)

Bond

$$C-H$$
 $C-C$ $C-$

N C - O

Bond energy (in kcal mol⁻¹) 97 83 73 86

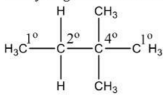
16 (d)

1° carbon is attached to one carbon atom.

2° carbon is attached to two carbon atms.

3° carbon is attached to three carbon atoms.

The hydrogen attached to 2° carbon atom are 2°.



 \div It has one 2° carbon atom and two 2°hydrogen atoms.

17 (d)

18 **(b)**

Non-staggered means eclipsed form.

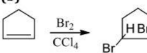
19 (c)

Carbanion is electron rich species. Stability of carbanion increases with increase in *s* —character of hybrid orbitals of carbon bearing the charge.

$$: sp^3 < sp^2 < sp$$

(25%s-character) (33%s-character) (50%s-character)

20 (b)



It is example of addition reaction

21 (a)

It is a fact.

22 (c)

It is the latest modified definition of organic chemistry.

23 **(b)**

In the lassaigne test, if organic compound consists of both N and S, then a red colour is obtained on adding aqueous

3

 ${\rm FeCl_3}$ To sodium extract. ${\rm NH_2-C-NH_2}$ contains both N and S hence, it will give red colour in Lassaigne test

NaCNS + FeCl₃ → [Fe(SCN)]Cl₂ + NaCl Blood red colour 24 (a)

If nitrogen is present in organic compound then sodium extract contains NaCN.

$$Na + C + N \xrightarrow{Fuse} NaCN$$

 $FeSO_4 + 6NaCN \rightarrow Na_4[Fe(CN)_6] + Na_2SO_4$
(A)

A changes to Prussian blue $Fe_4[Fe(CN)_6]_3$ on reaction with $FeCl_3$.

$$4\text{FeCl}_3 + 3\text{Na}_4[\text{Fe}(\text{CN})_6]$$

$$\rightarrow$$
 Fe₄[Fe(CN)₆]₃ + 12NaCl

25 **(b**)

Follow IUPAC rules.

27 **(b)**

It is the definition of asymmetric synthesis.

28 (a

(i) has sp^3 ; (ii) has sp^2 , sp^3 ; (iii) has sp^2 , sp; (iv)

31 (d

The number of isomeric alkenes with molecular formula C_6H_{12} are 13.

(1)
$$CH_2 = CH - CH_2 - CH_2 - CH_2 - CH_3$$

(2)
$$CH_3 - CH = CH - CH_2 - CH_2 - CH_3$$

(cis and trans)

(3)
$$CH_3 - CH_2 - CH = CH - CH_2 - CH_3$$

(cis and trans)

 CH_3

$$(4)CH_2 = CH - CH - CH_2 - CH_3$$

 CH_3

J

$$(5)CH_2 = C - CH_2 - CH_2 - CH_3$$

 CH_3

$$(6)CH_2 = CH - CH_2 - CH - CH_3$$

 CH_3

Ш

$$(7)CH_3 - CH = CH - CH - CH_3$$

(cis and trans)





$$CH_{3}$$

$$|$$

$$(8) CH_{3} - C = CH - CH_{2} - CH_{3}$$

$$CH_{3}$$

$$|$$

$$(9)CH_{3} - CH = C - CH_{2} - CH_{3}$$

$$(cis and trans)$$

32 **(c)**

It is 3-methyl butan-2-ol.

33 (c)

This statement is not true now.

34 (c)

Nitroalkanes exhibit tautomerism. In it, α -H-atom is labile and form nitrolic acid.

$$H_3C$$
— CH_2 — N
 O
 O
 H_3C — CH
 O
 O

36 (d)

37 **(b)**

Keto and enol forms are inter convertable. The enol content will be maximum when enol form is stabilised by hydrogen bonding.

In acetyl acetone, the enol form is stabilised by H-bonding, hence it has more enol content than other.

38 **(b)**

Compounds having bivalent functional group (like C=0, -0-, -S – etc) with atleast 4 carbon atoms (in case of ether and thioether) or atleast 5 carbon atoms (in case of ketones) exhibit metamerism. Hence, $C_2H_5 - S - C_2H_5$ will show metamerism.

39 (a)

Follow IUPAC rules.

40 (c)

$$^{+}N(CH_3)_3$$
 $|$
 $CH_3 - C - CH_2CH_3 \xrightarrow{Alc.KOH} CH_3CH = CHCH_3$
 $|$
 $|$
 $|$

is an example of elimination reaction

41 **(b)**

The dehydrohalogenation in presence of OH⁻is correctly represented by

$$\begin{array}{c|c} B & & \\ & H & \\ \hline OH^- & \end{array}$$

In this mechanism the base OH^- removes a proton from the β carbon.

43 (a)

Covalent bonds are cleaved in homolytic way in presence of UV light. It results in formation of free radical.

45 **(b)**

One asymmetric carbon atom is present.

46 **(b)**

Both alkene and cyclo alkane have general formula C_nH_{2n} .

47 **(b)**

IUPAC name of the above compound is 3-carboxyhexane-l, 6-dioic acid.

48 (d

 $Na_2S + Na_2[Fe(CN)_5NO] \rightarrow Na_4[Fe(CN)_5NOS]$ Sodium nitroprusside sodium thio nitro Solution prusside (purple colure)

49 (d)



Greater the difference in electronegativity of bonded atoms easier will be heterolytic cleavage

50 (c)

$$HO - CH_2 - CH_2 - F$$
 H
 $O - H$
 $S + CH$

Gauche conformation is comparatively more stable due to hydrogen linkage in between F and H (at O-atom), hence order is *Eclipse*, *Anti* (staggered), *Gauche*.

51 (d)

Phosphorous is estimated as $Mg_2P_2O_7$ P $\xrightarrow{\text{HNO}_3}$ H_3PO_4

$$H_3PO_4 + Mg^{2+} + NH_4OH \rightarrow MgNH_4PO_4$$

 $2MgNH_4PO_4 \rightarrow Mg_2P_2O_7 + H_2O + 2NH_3$
% of $P = \frac{62 \times wt.of Mg_2P_2O_7 \times 100}{222 \times w}$

52 **(c)**

$$n = 2$$
 and $a = 2^n = 2^2 = 4$.

54 **(c)**

Among carbonyl compounds, reactivity decrease with increase in alkyl groups as alkyl groups (+I effects) decrease positive character on C-atom. Thus, the correct order of reactivity is $HCHO > CH_3CHO > C_6H_5CHO$

55 (d)

Primary and secondary alkyl halides gives $S_N 2$ reaction

56 **(c)**

Follow IUPAC rules.

57 **(d)**

2-butanol is optically active as it contain as it contain chiral carbon atom.

$$\begin{array}{c} \operatorname{CH}_2 \\ | \\ \operatorname{CH}_3 - \operatorname{CH}_2 - \operatorname{C} - \operatorname{OH} \\ | \\ \operatorname{H} \end{array}$$

58 **(d)**

$$a = 2^n$$
; $n = 3$.

59 **(a**)

It is Markownikoff's rule.

61 (b)

The order of homolytic bond dissociation energies of CH_4 , C_2H_6 and CH_3Br is as

$$CH_4 > C_2H_6 > CH_3Br$$

63 (d)

Each d form has its l form and the pair is known as enantiomer.

64 **(b)**

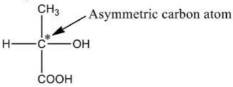
Carboxylic acid and esters show functional group isomerism. When two compounds have same molecular formula but different functional groups, then functional isomerism arises.

e.g.,

C2H5COOH and CH3COOCH3

65 (a)

One asymmetric carbon atom is present in a lactic acid molecule. Hence, it is an optically active compound.



66 (a)

According to IUPAC system, ether are named as alkoxy alkanes. The larger alkyl group froms the parent chain while lower alkyl group is taken ethereal oxygen and forms a part of alkoxy group.

$$CH_3$$
— CH_2 — O — CH — CH_2 — CH_2 — CH_2 — CH_2 CI 2-etoxy-5-chloropentane

67 (d)

Follow IUPAC rules.

68 (c)

It is a fact.

69 (c)

A liquid, which decomposes at its normal boiling point can be purified by vacuum distillation.

71 (b)

Stability of carbanion is not governed by hyperconjugation. Its stability depends on the +I or -I group

72 (c)

Let unreacted $0.1M(=0.2N)H_2SO_4 = V'mL$ $\therefore 20 \text{ mL of } 0.5M \text{ NaOH}$

$$\therefore 20 \times 0.5 = V' \times 0.2$$

$$V' = 50 \text{mL}$$



Used
$$H_2SO_4 = 100 - 50 = 50mL$$

$$\%Nitrogen = \frac{1.4 \text{ NV}}{w}$$

where, N=normality of H2SO4

V=volume of H_2SO_4 used

∴ % nitrogen =
$$\frac{1.4 \times 0.2 \times 50}{0.30}$$

= 46.67%

% of nitrogen in

(a)
$$CH_3CONH_2 = \frac{14 \times 100}{59} = 23.73\%$$

(a)
$$CH_3CONH_2 = \frac{14 \times 100}{59} = 23.73\%$$

(b) $C_6H_5CONH_2 = \frac{14 \times 100}{122} = 11.48\%$
(c) $NH_2CONH_2 = \frac{28 \times 100}{60} = 46.67\%$

(c)NH₂CONH₂ =
$$\frac{28 \times 100}{60}$$
 = 46.67%

(d)
$$NH_2CSNH_2 = \frac{28 \times 100}{76} = 36.84\%$$

Therefore, the organic compound is urea.

73 (d)

$$CH_4 + Cl_2 \xrightarrow{hv} CH_3Cl$$

This is an example of free radical substitution

$$CI \longrightarrow CI^{\bullet} + CI^{\bullet}$$

$$CH_4 + CI^{\bullet} \longrightarrow CH_3^{\bullet} + HCI$$

$$CH_4 + CI^{\bullet} \longrightarrow CH_3^{\bullet} + HCI$$

$$CH_3^{\bullet} + Cl-Cl \longrightarrow CH_3Cl + Cl^{\bullet}$$

$$Cl^{\bullet}+ Cl^{\bullet} \longrightarrow Cl_2$$

Glucose has four dissimilar asymmetric carbon 82 atoms; $a = 2^4$.

75 (d)

The elimination takes place according to Saytzeff rule. The most substituted alkene (butane-2) is called Saytzeff product whereas less substituted alkene (butane-1) is called Hofmann product

76 (c)

-NO₂ group shows-M effect while CH₃O − group shows +M effect. (-Meffect stabilizes an anion) Hence, the order of stability is

$$\begin{array}{c|ccc}
\bar{C}H_2 & \bar{C}H_2 & \bar{C}H_2 \\
\hline
O & > & O \\
NO_2 & OCH_3 \\
\hline
(II) & (III) & (I)
\end{array}$$

77 **(b)**

The former possesses 12α-H atom whereas, later possesses six α -H atom. More is the no. of α -H atom, more is delocalisation and more is stability.

78 (d)

H2O, ROH, R - COOH etc are protic solvents because they are polar in nature and contain a hydrogen directly bonded to oxygen while other are aprotic solvents as they do not have a hydrogen bonded directly to oxygen. They are especially favourable for $S_N 1$ reactions. While aprotic solvents cannot have hydrogen bond to the nucleophile because they does not have hydrogen bonded to nitrogen or oxygen. They are favourable for S_N2 reactions.

79

Follow mechanism of addition reactions.

80 (c)

Methoxy group is electron releasing group it increases electron density of benzene nucleus - NO₂ group is electron withdrawing group, it decreases the electron density of benzene nucleus. Thus, the order of reaction with electrophilic regent is

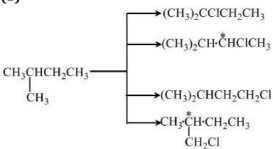
81

$$NaCN + HNO_3 \xrightarrow{\Delta} HCN \uparrow + NaNO_2$$

 $Na_2S + HNO_3 \xrightarrow{\Delta} H_2S \uparrow + 2NaNO_3$

The central carbon in I and II is asymmetric.

83 (b)



Thus, out of four isomers only two have chiral carbon. Each have two isomers.

85

The structural formula of epoxide is CH It consists three membered ring with two carbon

and one oxygen. $CH_3 - CH_2 - CH_2 - Cl + alc. KOH \rightarrow CH_3 - CH$

It is an example of elimination reaction





87 (d)

In the reaction

$$>$$
C=O + H₂NOH \rightarrow C=NOH + H₂O

Both addition and elimination takes place simultaneously. Thus, the reaction is addition elimination

88 (d)

Resolution of racemic mixture involves the 98 formation of distereoisomers.

89 (b)

When N and S both are present in the organic compound, then a red colour complex ion of $[Fe(CNS)]^{2+}$ is formed on adding $FeCl_3$ to sodium extract

NaCNS + FeCl₃ → [Fe(SCN)]Cl₂ + NaCl blood red colour

90 (a)

$$CH_3CH_2Br \xrightarrow{H-H} CH_3 - CH_3 + HBr$$

$$S_{N^2}$$

$$CH_{3}$$

$$CH_{3} - C - Br \xrightarrow{LAH} CH_{3} - C = CH_{2} + HBr$$

$$| (E_{2}) |$$

$$CH_{3} \qquad CH_{3}$$

91 (a)

Count σ and π -bonds.

92 (d)

The formation of sodium thionitroprusside (blue) shows the presence of sulphur.

$$Na_2S + Na_2[Fe(NO)(CN)_5]$$

Sodium sodium nitroprusside sodium thionitroprusside purple extract

93 **(a)**

It is a fact.

94 **(b)**

$$R - CH_2 - CH_2X + KOH(alc.) \rightarrow R - CH$$

= $CH_2 + KX + H_2O$

Alkyl halid undergo β -elimination to form alkene.

95 **(d)**

The rate of reaction follows the order : RI > RBr > RCl > RF; whether it obeys S_N1 or S_N2 mechanism due to steric hindrance of alkyl group.

96 (c)

Only this is optically active due to central carbon being asymmetric.

97 (a)

Ethers show metamerism. Metamerism arises when a polyvalent functional group

to different alkyl groups but the molecular formula remains same e.g.,

$$C_2H_5 - O - C_2H_5$$
 and $CH_3 - O - C_3H_7$

98 (6

Carbanion is electron rich species. Stability of carbanion increases with increase in *s* —character of hybrid orbitals of carbon bearing the charge.

 $sp^3 < sp^2 < sp$

(25%s-character) (33%s-character) (50%s-character)

99 (c)

The Z repells electrons and thus, electron density increases on R_3C part.

100 (d)

∵ −COOH group is a deactivating group

: Benzoic acid is less reactive towards electrophilic substitution.

So, benzoic acid> phenol> *n*-propyl benzene is not arranged correctly.

101 (c)

Aqueous solution of NaHCO₃ can be used to separate benzoic acid from its mixture with camphor. Benzoic acid form water soluble sodium benzoate with NaHCO₃.

102 (d)

Each π -electron is delocalised over six carbon atoms in ring.

103 (d)

Grignard reagent reacts with >C=0, $-C \equiv N$, >C=S as follows

104 (a)

Benzene is the most stable and we know that resonance energy is a direct measure of the stability of a molecule

105 (d)



It is an example of nucleophilic addition reaction.

107 (c)

 ${
m H_3C-C-Br}$ and ${
m CH_3CH_2CH_2Br}$ are chain as well as position isomers.

CH₃

108 (b)

Detection of phosphorus in the organic compound can be done by its conversion into phosphate. The ammonium molybdate is used to identify phosphate ion

$$Na_3PO_4 + 3HNO_3 \rightarrow H_3PO_4 3NaNO_3$$

$$H_3PO_4 + 12(NH_4)_2MoO_4 + 21HNO_3 \xrightarrow{\Delta}$$
 ammonium modybdate

$$(NH_4)_3PO_4.12MoO_3 + 21NH_4NO_3 + 12H_2O$$

yellow ppt.

109 (b)

Homolytic bond fission is one in which each entity involved in bond formation retains its electron involved in shared pair of electron to form free radicals.

110 (c)

 CH_3CN has sp^3 and sp-hybridised carbon atom.

111 (c)

CH₃OCH₃, C₂H₅OH

Methoxy methane Ethanol (ether)
(alcohol) functional group

 (C_2H_6O) (C_2H_6O) molecular formula

In methoxy methane and ethanol both molecular formula is same but functional groups are different, so they are functional isomers.

112 (b)

The main conditions for exhibiting geometrical isomerism are

- (i) Presence of double bond.
- (ii) Presence of different groups on same double bonded carbon.
- (iii) Presence of at least one similar group on adjacent double bonded carbon atoms.

$$C_3H_6(H_3C \xrightarrow{\hspace{1cm}} CH \xrightarrow{\hspace{1cm}} CH_2)_{\hspace{1cm}}$$
 does not exhibit geometric isomerism due to presence of same group on double bonded carbon atom (C_1) .

113 (a)

Follow mechanism of addition reaction.

114 (c)

Vicinal or alkylene dihalides.

115 (d)

The electrophile involved in the sulphonation of benzene is SO_3 .

$$2H_2SO_4 \rightarrow SO_3 + H_3O^+ + HSO_4^-$$

116 (b)

Number of meso structures in compound having odd number of chiral carbon atoms and

symmetrical molecule
$$=2^{\left(\frac{n}{2}-\frac{1}{2}\right)}$$

Given,
$$n = 5$$

$$\therefore \text{ Number of } meso \text{ forms} = 2^{\frac{5}{2} - \frac{1}{2}} = 2^2 = 4$$

117 (d)

In Kjeldahl's method, the nitrogen is quantitatively converted into ammonia by heating with NaOH

$$C + H + N \xrightarrow{\Delta} (NH_4)SO_4 + CO_2 + H_2O$$

(from organic compound)

 $(NH_4)_2SO_4 + 2NaOH \rightarrow Na_2SO_4 + 2NH_3 + 2H_2O$ In Dumas method nitrogen present in organic compound is quantitatively converted into N₂.

118 (c)

It is a fact.

119 (b)

$$(CH_3)_3 C$$
 OH $\xrightarrow{+H^+}$ $(CH_3)_3 \overset{+}{C}$ 3^o carbocation (more stable)

$$CH_2$$
— CH_2 — CH_2 OH $\xrightarrow{+H^+}$
 $-H_2$ O

 CH_2 — CH_2 —

Increasing order of stability of carbocation. 1°carbocation <2° carbocation<3°carbocation

120 (c)

Both geometrical and optical isomerism are included in stereoisomerism.

121 (a)

Phenol reacts with chloform and NaOH to give ohydroxy benzaldehyde or salicylaldehyde. In this reaction dichlorocarbene (: CCl₂) electrophile is



generated. This reaction is called Reimer-Tiemann reaction.

122 (b)

Homolytic fission is favoured by sunlight. In it, each bonded atom takes away its shared electrons and thus free radicals are produced.

$$CI \longrightarrow CI^{\bullet} + CI^{\bullet}$$

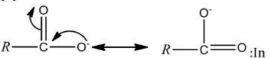
124 (d)

All show geometrical isomerism:

$$C_6H_5$$
 C —H and C_6H_5 C —H N —OH HO—N

$$CH_3$$
 C
 CCC
 CH_3
 CH_3
 CC
 $COOH$
 $COOH$

125 (d)



carboxylate ion, the negative charge is present on oxygen, a most electronegative element here, thus it is resonance stabilised.

 $HC \equiv C^-$: Carbon is sp-hybridised so its electronegativity is increased higher relative to nitrogen.

 $\overline{N}H_2$: Nitrogen is more electronegative than sp^3 -hybridised C-atom. From the above discussion, it is clear that the order of the stability of conjugated bases is as

$$RCOO^->HC\equiv C^->\overline{N}H_2>R^-$$
 and higher is the stability of conjugated bases, lower will be basic character. Hence, the order of basic character is as

 $RCOO^- < HC \equiv C^- < \overline{N}H_2 < R^-$

126 (b)

 $S_N {\bf 1}$ Reaction is most favourable for tertiary substance.

$$H_3C$$
 CH_3
 S_N1
 CH_3
 CH_3

127 (d)

 $CH_3CHOHC_2H_5$ is optically active because it has chiral C^* -atom

(most sable)

$$H_3C$$
 C_2H_5 Chiral carbon atom

128 (d)

The closed ring cycloalkanes beyond five carbon atoms has puckered ring structure maintaining tetrahedral nature or stainless rings, *e.g.*, cyclo hexane has chair and boat form.

129 (d)

These are characteristics of S_N 1 mechanism.

130 (c)

-I power of groups in decreasing order with respect to the reference H

$$NO_2 > CHO > COOR > F > Cl > Br > I > OH$$

 $> OR > NH_2$

131 (d)

Asymmetry is present in all states.

132 **(b)**

The structure of cyclopropane, cyclobutane and cyclo hexane are as

Hence, the common group in cyclopropane, cyclobutane and cyclohexane is $>CH_2$ group.

133 (d)

Draw all possible isomers.

135 (a

+I effect is shown by - CH₃ while -I effect is shown by -Br, -Cl and -NO₂.

136 (b)

It is a strong reducing agent.

% of C=
$$\frac{12}{44} \times \frac{0.535}{0.765} \times 100 = 19.07$$

% of H = $\frac{2}{18} \times \frac{0.138}{0.765} \times 100 = 2.004$

Ratio of % of C:H=19:2 (approx.)

138 (d)

Benzene has planar structure.

139 (c)

$$\begin{array}{c}
CI \\
6 \\
5
\end{array}$$
Br

Unsaturation (double bond) is given priority over halogen, then lowest set of locants. So, the correct IUPAC name is 3-bromo-1-chlorocyclohexene.

140 (c)

Follow conformation; The conformers for nbutane are two gauche, two eclipsed and one anti.

142 (d)

Select longest possible carbon atom chain, number it and name compound according to IUPAC, rules.

3, 3-diethyl-4-methyl-5-(1'-methyl ethyl)-octane

143 (b)

Benzal is C₆H₅CH group.

144 (b)

A 2° carbon is one of which two valencies are 150 (c) attached to carbon atom.

Eclipsed conformation of butane contain angle and steric strain both. Follow conformation.

146 (c)

HCl

Weight of organic compound =29.5mg

$$\begin{array}{ccc} & NH_3 + HCl \longrightarrow Na_4Cl \\ HCl & + & NaOH & \longrightarrow NaCl + H_2O \\ \text{(remaining)} & 15 \times 0.1 \ M \end{array}$$

=1.5 mmol

Total millimole of HCl=2

Millimole used by $NH_3 = 2 - 1.5 = 0.5$

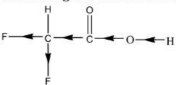
Weight of $NH_3=0.5 \times 17$ mg=8.5mg

Weight of nitrogen = $\frac{14}{17} \times 8.5mg = 7mg$

% Nitrogen =
$$\frac{7}{29.5} \times 1100 = 23.7\%$$

147 (a)

Fluoro group causes negative inductive effect increasing ionisation, thus 0.1M difluoroacetic acid has highest electrical conductivity.



148 (c)

-NO₂ group shows - M effect white CH₃O -group shows +M effect (-M effect stabilises an anion)

149 (d)

When the groups with higher priority (i.e., with high atomic number) are present on same side of double bond, then the configuration is Z but when present on opposite side of double bond, the configuration is E.

(i)
$$\overset{\text{CI}}{\underset{(Z)}{\longrightarrow}} c = c \overset{\text{Br}}{\underset{(Z)}{\longleftarrow}} c$$

(Priority : Cl > H and Br > F)

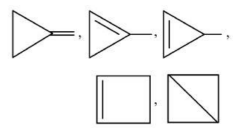
$$(ii)$$
 $\stackrel{CI}{\underset{(E)}{\longrightarrow}} c = c \stackrel{F}{\underset{(E)}{\longleftarrow}} c$

(Priority: C1 > H and Br > F)

Nucleophiles are electron rich species and can donate lone pair of electron to carbocation or any +ve centre.

152 (c)

 C_4H_6 can have five cyclic isomers.



153 (b)



154 (d)

Urea shows tautomerism as

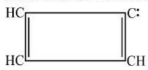
$$\begin{array}{ccc} & & & \text{OH} \\ & & & & \parallel \\ & & \text{NH}_2 - \text{C} - \text{NH}_2 \leftrightarrows \text{NH} = \text{C} - \text{NH}_2 \\ & \text{keto form} & \text{enolic form} \end{array}$$

155 (d)

The conditions given are for C₆H₆.

156 (c)

Count π -bonds. Delocalisation is not possible.



157 (a)

Pentene-2 exhibits cis and trans-isomerism.

159 (c)

The rule is valid for unsymmetrical alkene.

160 (c)

(CH₃)₂CHCH₂CH₃ is isopentane.

161 (a)

 π -bond in molecule give rise to hindered rotation.

162 (c)

$$CH_2 = CH_2 \xrightarrow{-H_2} CH \equiv CH$$

Conversion of ethylene into acetylene is a example of elimination reaction.

163 (d)

The two propenyl group attached to 1,2-position of carbon in *cis*-form.

$$CH_3$$
 CH_3 CH_3 CH_4 CH_5 CH_6 CH_7 CH_8 CH_8

164 (d)

Markownikoff's rule is obeyed during addition of unsymmetrical addendum on unsymmetrical alkene.

165 (a)

Benzene has 6 C—C and 6 C—H σ -bonds and 3 C = C π -bonds.

166 (b)

It is a fact.

167 (c)

Alkynes are linear due to sp-sp hybridized carbon.

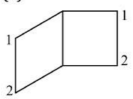
168 **(a**

For keto-enol isomerism a compound should have at least one α -hydrogen atom with respect to ketone group or in other words for tautomerism presence of α —hydrogen atom is essential.

does not exhibited.

tautomerism due to absence of α -hydroegn atom.

170 (d)



Hence, correct IUPAC name is bicyclo [2,2,0] hexane.

172 (d)

Chlorinolysis involves substitution reactions by chlorine.

173 (b)

 $a = 2^{n-1}$; where n is no. of asymmetric carbon; when molecule possesses symmetry.

174 (c)

$$\begin{array}{c} 4 \\ \text{CH}_{3} \longrightarrow \begin{array}{c} 3 \\ \text{CH}_{2} \longrightarrow \begin{array}{c} 1 \\ \text{CHO} \end{array} \\ \\ \text{CI} \\ \text{3-chlorobutanal} \end{array}$$

 \therefore The order is priority is -CHO > -Cl.

175 (c)

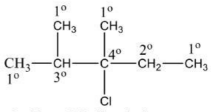
% C =
$$\frac{12}{44} \times \frac{12.517}{4.0} \times 100 = 85.7$$

% H =
$$\frac{2}{18} \times \frac{5.143}{4.0} \times 100 = 14.3$$

The mole ratio of C to H is $\frac{85.7}{12}$: $\frac{14.3}{1}$ = 7.14: 14.3 = 1: 2 = CH₂

176 (b)

3-chloro-2, 3-dimethyl pentane contains all the four $1^{\circ}, 2^{\circ}, 3^{\circ}$ and 4° carbon atoms.



3-chloro-2-3-dimethyl pentane

178 (d)

The carbon, four valencies of which are satisfied by four different groups, is termed as chiral carbon atom. The structures of the given compounds are as



(where,
$$\stackrel{*}{C} =$$
 =chiral carbon atom)

Hence, succinic acid does not contain any chiral carbon atom.

180 (a)

In homolysis, the covalent bond is broken in such a way that each resulting species known as free

$$CH_3CH_2$$
— $CI \xrightarrow{Homolytic} CH_3CH_2 + CI$

181 (b)

p-orbitals are at 90° to each other.

S_N2 reactions are greatly controlled by steric

$$R - CH_2 - X R_2CH - X R_3C - X$$
 1°
 2°
 3°

S_N2 reactivity decreases as bulkyness of alkyl group increases.

183 (a)

3-methyl pentane 3-ol

Hydroxy is used when - OH group is written in prefix. So, choice (b) and (c) are wrong.

184 (d)

Due to -R effect of – CHO group, oxygen carries $-\delta$ charge while the terminal carbon carries $+\delta$, ie,

185 (a)

Its IUPAC name is 3,5-dimethylcyclohexene.

186 (a)

Lactic acid obtained in the given reaction is an optically active compound due to the presence of chiral C-atom. It exists as d and l forms whose ratio 1:1.

187 (c)

The mirror-image isomerism is a class of stereoisomerism and are included in optical isomerism.

188 (c)

CH₃⁺ acts as an intermediate in the given reaction (Friedel Craft's alkylation). It is an example of electrophilic aromatic substitution. In this reaction CH_3^+ is electrophile.

189 (d)

Possible number of optical

isomers =
$$2^n$$

$$= 2^2 = 4$$

190 (b)

 C_6H_6 has 12σ and 3π -bonds.

191 (c)

For $S_N 2$ mechanism, there should be least steric hinderance.

193 (c)

Rest all are polymerisation.

The increasing order of inductive effect is: -F < $-COOH < -CN < -NO_2$.





197 (b)

-COOH is on top in preference table.

200 (d)

In homolytic cleavage, covalent bond is cleaved in such a way that each atom takes its shared electrons with itself and free radicals are formed.

$$A \cap \bigcap_{BH2} \frac{\text{Homolytic fission}}{\text{free radicals}} A^{\bullet} + B^{\bullet}$$

201 (d)

Fractional distillation is used for the separation of crude petroleum into various fractions like coaltar, crude alcohol and petroleum

202 **(b)**

Free radical has unpaired electron.

203 (a)

 SO_3 can accept lone pair of electron in d-subshell.

204 (a)

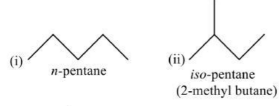
From Kjeldahl's method,

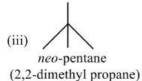
Percentage of nitrogen

$$= \frac{1.4 \times N \times V}{W} = \frac{1.4 \times 0.1 \times 30}{5}$$
$$= 0.84\%$$

205 (a)

The isomers alkanes having the molecular formula C_5H_{12} are as





206 (b)

Just after few years when Wöhler prepared urea from KCNO and $(NH_4)_2SO_4$, Kolbe prepared acetic acid in laboratory from its element and gave final blow to Vital force theory.

207 (b)

Due to the presence of asymmetric carbon atom, e.g.,

208 (a)

$$\begin{array}{c} sp^3\\ \mathsf{CH_3}\\ \mathsf{CH_3} & |\\ \mathsf{CH_3} & \mathsf{C}\\ |sp^3\\ \mathsf{CH_3}\\ sp^3\\ \end{array}$$

All bonds are σ - bonds hence C uses only its sp^3 -hybrid orbitals. In all other compounds there is one C = 0 double bond, therefore, this carbon is sp^2 - hybridized

210 (a)

Follow Baeyer's strain theory for stability of cycloalkane.

211 (a)

— COOH is electron withdrawing group.

213 **(d)**

CHO

is called glyoxal

214 (d)

IUPAC name of acraldehyde

$$(CH_2 = CH. CHO)$$
 is:

216 (d)

These are characteristics known from mechanism of reaction.

217 **(b)**

$$\begin{array}{c|c} H & \vdots \stackrel{\sigma}{\circ} \stackrel{H}{\circ} H \\ H & \stackrel{\sigma}{\circ} \stackrel{\sigma}{\circ} C \stackrel{\sigma}{\circ} C \stackrel{\sigma}{\circ} H \\ H & H \end{array}$$

enolic form of acetone

 9σ bonds, 1π bond, 2 lone pairs

218 (c)

BrCH₂CH₂COOH is the weakest acid and have lowest dissociation constant because. I.E. of Br is lesser than F and is far away from – COOH group.

219 (a)

Follow IUPAC rules.

220 (d)

Molecular formula C_2Br ClFl six isomers are possible.



$$C = C$$
 $C = C$
 C

$$c = c$$

221 (a)

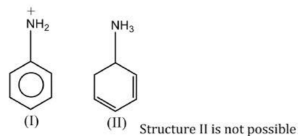
Nucleophilies are those substances which can donate a pair of electrons. They can be neutral or negatively charged. The nucleophilic power depends on the tendency of species to donate the electrons. Due to the presence of +I effect it increases. Hence, higher the +I effect, higher the nucleophilic power. The +I effect of ethyl is greater than +I effect of methyl group

$$C_2H_5 \longrightarrow S-H, CH_3 \longrightarrow C-O^-,$$

 $+I$
 $CH_3 \longrightarrow NH_2, NC \longrightarrow CH_2^-$

223 (b)

Among the given species $C_6H_5NH_3$ does not exert a resonance effect.



because in it nitrogen contains 10 valence electrons.

224 (c)

Draw all structures.

225 (a)

Chiral carbon atom has all four different groups attached to it.

 \div It has one asymmetric or chiral carbon atom.

226 (b)

Different spatial arrangement of atoms leads to its configuration.

227 (d)

Butane and isobutane and all higher alkanes show isomerism.

228 (a)

Desmo (bond), tropism (turn). Thus, desmotropism, *i.e.*, isomerism arised due to turning of bond was the name given to tautomerism.

229 (d)

$$\begin{array}{c|c}
CH_{3} & C & CH & CH_{3} \\
1 & 2 & 3 & 4 \\
CH_{3} & CH_{3}
\end{array}$$

3-methyl butan-2-one or 3-methyl 2-butanone

0

Keto (-C-) functional group is given priority.

230 (b)

Halogens can be identified in organic compounds by Beilstein-test.

231 (a)

It undergoes dehydration easily as the product obtained is conjugated and more stable.

232 (c)

Methoxy ethene is $CH_3O-CH = CH_2$; are unsaturated ether.

233 (a)

Electrophiles are electron deficient species which can share lone pair of electron with carboanion and are thus, called Lewis acids.

234 (c)

 Iso —propyl chloride is a 2° halide and 2° halides can undergo hydrolysis either by S_N1 or S_N2 mechanism depending upon the nature of solvent used.

235 (b)

Follow IUPAC rules.

236 (c)

Carbon and hydrogen are estimated in organic compounds by Liebig's method

$$C + 2CuO \xrightarrow{\Delta} 2Cu + CO_2$$

$$2H + CuO \xrightarrow{\Delta} Cu + H_2O$$

Percentage of carbon and hydrogen is calculated from the weight of CO_2 and H_2 produced

237 (c)



 μ is more for (c) then (d).

238 (b)

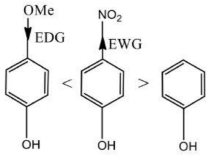
Free radical state is a transient state and thus, has short life.

240 (c)

When nitro group is present in the benzene nucleus, it withdraws electrons from o and ppositions. Thus, the electron density at the o and p -positions decreases. m-positions become positions of comparatively higher electron density and therefore, electrophilic attack occurs at mpositions.

241 (c)

According to Lewis, electron acceptor compounds are called acids. Therefore, compounds having tendency to accept electrons will be more acidic. The correct order of acidic character is as follows:



243 (b)

C4H7Cl is a monochloro derivative of C4H8 which itself exists in three isomeric forms.

(i) $CH_3 - CH_2 - CH = CH_2$: Its possible monochloro derivatives are

$$CH_3 - CH_2 - CH = CH - CI$$

2 isomers cis and trans forms

Optically active (exists in two forms)

$$ClCH_2 - CH_2 - CH = CH_2$$
 (one form)

$$H_3C - CH_2 - C = CH_2$$
 (one form)

(ii) $CH_3 - CH = CH - CH_3$: Its possible

monochloro derivatives are

$$CH_3 - CH = C - CH_3$$

Exists in two geometrical forms

$$CH_3 - CH = CH - CH_2CI$$

Exists in two geometrical forms

(iii) $CH_3 - C = CH_2$: Its possible monochloro derivatives are

$$CH_3 - C = CH - Cl$$
, (Only one form)

 CH_3

$$CICH_2 - C = CH_2(only one form)$$

 CH_3

Thus, the total acylic isomeric forms of C4H7Cl are

244 (d)

It shows intramolecular H-bonding.

245 (d)

Diethyl ether is resistant to nucleophilic attack by hydroxyl ion.

246 (d)

C₆H₆ has more canonical forms.

247 (d)

Chemical methods are based upon the distinguishing chemical properties of one class of organic compounds from the other. for example camphor and benzoic acid

248 (c)

Fractional crystalliation is used to purify organic solids which dissolve in a particular solvent. But their rate of solubility is different

249 (d)

NH4CNO is functional isomer of urea.

250 **(b)**

Carbanion (CH₃)

Here, the carbon atom carries a negative charge with lone pair of electrons and has eight electrons in outermost orbit and complete its octet.



Reactions in which carbanions are

formed as intermediate are said to proceed by a "Carbanion mechanism".

Carbanion is sp^3 hybridised, three sp^3 hybrid orbitals form covalent bonds with three atoms while the fourth sp^3 hybrid orbital has a nonbonding pair of electrons. It is pyramidal in shape as similar to NH3.

251 (a)

Ether group(-0-)has propyl and isopropyl group on its two sides.





252 (d)

Follow carbenes.

253 (c)

There are two chiral C-atoms (*) Thus, optical isomerism is possible.

254 (b)

Follow IUPAC rules.

255 (d)

-do-

256 (c)

A monosubstituted benzoic acid is stronger than a monosubstituted phenol as former being a carboxylic acid. Among the given substituted benzoic acid, *ortho* — hydroxy acid is strongest acid although — OH causes electron donation by resonance effect which tends to decreases acid strength. It is due to very high stabilisation of conjugate base by intramolecular H-bond which outweight the electron donating resonance effect of — OH.

The overall order of acid-strength of given four acids is *ortho*-hydroxybenzoic acid (pK_a = 2.98) > Toluic acid pk_a = 4.37) > p -hydroxybenzoic acid (pk_a = 4.58) > p -nitrophenol (pk_a = 7.15).

257 (b)

Electrophiles are electron pair acceptors.

258 (b

 $CH_3CH_2CHCICH_2CH_3$; $CH_3CHCICH_2CH_2CH_3$; $CH_2CICH_2CH_2CH_3$.

259 (b)

Benzene and all its derivatives along with heteroaromatics possess aromatic nature. Rest all possess aliphatic nature.

260 (d)

Neohexyl chloride is a primary halide as in it, Clatom is attached to a primary carbon.

$$\begin{array}{c} \operatorname{CH_3} \\ | \\ \operatorname{CH_3} - \operatorname{C} - \operatorname{CH_2} - \operatorname{CH_2} \operatorname{CI} \\ | \end{array}$$

 CH_3

261 (a)

Ethane, ethene and ethyne have sp^3 , sp^2 and sp-hybridization respectively.

262 (d)

Carbon is asymmetric as all its valencies are attached to different groups.

264 (b)

Follow IUPAC rules.

265 (d)

Bromination of alkanes in the presence of sunlight involves the formation of free radical, e.g.,

$$CH_4 \xrightarrow{Br_2} CH_3Br$$

Mechanism

Initiation

$$Br \longrightarrow Br \xrightarrow{hv} Br + Br$$

Propagation

$$CH_4 + Br - CH_3 + HBr$$

$$CH_3^{\bullet} + Br \longrightarrow CH_3Br + Br$$

Termination

$$Br + Br \longrightarrow Br_2$$

266 (d)

The two butene give different products on addition of $\mathrm{Br}_2\text{-}cis$ butene gives racemic mixture whereas trans butene gives meso form of 2,3,3-dibromo butene.

267 (a)

Lassaigne's test is used for the detection of halogens, nitrogen and sulphur.

268 (a)

Resonance and inductive effect decide stability of carbocations.

∴Correct order of stability is

269 (c)

Atom At mass (a) % (b)
$$\frac{b}{a}$$
 Ratio X 10 50 $\frac{50}{10} = 5$ 2 Y 20 50 $\frac{50}{20} = 2.5$ 1 Hence, empirical formula = X_2Y



270 (c)

The compounds which differ in the nature of carbon chain are called chain isomers, e.g.,

$$CH_3 - CH_2 - CH_2 - CH_2 - CH_3$$

n-pentane
 CH_3

$$CH_3 - CH - CH_2 - CH_3$$
iso-pentane

$$CH_3$$

$$H_3C-C-CH_3$$
 CH_3

neo-pentane

272 (c)

In conjugated diene alternate single and double bonds are present while in cumulative diene, double bonds are present at adjacent positions.

I.
$$H_2C = CHCH_2 CH_2 CH = CH_2$$
hexa-1, 5-diene

⇒ It is an isolated diene.

II.
$$H_2C = C = CH_2$$

prop-1, 2-diene

⇒ It is a cumulative diene.

hexa-1, 3-diene

⇒ It is a conjugated diene.

IV.
$$H_2C = CH = CH = CH_2$$

buta-1, 3-diene

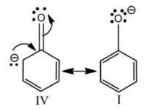
⇒ It is a conjugated diene.

V. Prop -1, 2-diene is a cumulative diene. Hence, statement 3 and 5 are correct.

276 (b)

Phenols are much more acidic than alcohol due to the stabilisation of phenoxide ion resonance.

Phenoxide ion is stabilised due to following resonating structures.



While, in alcohols

$$R \longrightarrow O$$
 $\longrightarrow H \longrightarrow R \longrightarrow O$ $\longrightarrow H^+$ alcohol (Not stabilised due to absence of resonance)

ortho nitrophenol is most acidic because in – NO_2 electron attracting group is attached to ortho position which helps in stabilising the negative charge on the oxygen of phenoxide ion. Hence, due to this reason acidic character of phenol is increased, while on attachment of – CH_3 group (electron donating group) acidic strength of phenol is decreased in cresol due to the destabilisation of phenoxide ion.

277 (d)

C: H: 0 = 6: 1: 8
=
$$\frac{6}{15} \times 100 : \frac{1}{15} \times 100 : \frac{8}{15} \times 100$$

40: 6.67: 53.3
= $\frac{40}{12} : \frac{6.67}{1} : \frac{53.3}{16}$
= 1: 2: 1 ie. CH₂O

278 (c)

Alkanes never show functional isomerism, metamerism, tautomerism and geometrical isomerism.

279 (b)

Draw all possible chain and position isomers.

280 (c)

2-butanol has following structure

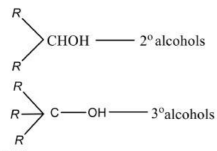
OH

|

CH₃ - CH - CH₂ - CH₃

2° carbon

R - CH₂OH - 1° alcohol



281 (a)

Stronger is acid, weaker is its conjugate base or weaker is nucleophilicity.

The acidic order $HF > H_2O > NH_3 > CH_4$.

282 (b)

Electromeric effect occurs only in the presence of attacking reagent. It operates in the molecules having multiple bonds. Since, it exists only on the demand of attacking reagent, it is a temporary effect. *e.g.*,

$$-c = N \xrightarrow{\text{Attacking reagent}} -c = N$$

283 (b)

Leaving group ability depends upon basicity of group.

284 (d)

Read optical activity.

285 (c)

The compounds must fulfill two conditions to show geometrical isomerism.

- (i) The compound should have at least one C=C.
- (ii) The two groups attached to same carbon must be different.

Out of given choices only (c) fulfill both conditions and shows geometrical isomerism.

$$H - C - COOH$$

H - C - COOH

(c)

2-butene-1,4-dioic acid

286 (a)

C — Cl bond is aryl chloride is stable due to delocalisation of electron by resonance. Also C —

Cl bond possess a double bond character like vinyl chloride, hence S_N reactions are not possible in chlorobenzene under ordinary conditions.

287 (b)

C-C, C=C and $C\equiv C$ bond length are 1.54 Å, 1.34Å and 1.20Å respectively. In benzene C=C is 1.40Å.

288 (c)

Definition of tautomerism.

289 (c)

The reactivity order is $3^{\circ}H > 2^{\circ}H > 1^{\circ}H$.

290 (a)

The +ve inductive effect of CH₃ group on carbanions intensifies negative charge on C⁻ centre and thus, 3° carbanion is more reactive.

291 (d)

Glycerine contains

$$^{\alpha}\text{CH}_2 - \text{OH} \leftarrow 1^{\circ} \text{ alcohol}$$

$$^{\beta}$$
CH − OH ← 2°alcohol

$$^{\alpha}$$
CH₂ − OH ← 1° alcohol

292 (a)

In rest all carbon chain is same.

293 (b)

In Cannizzaro reaction the transfer of H^- to another carbonyl group is difficult and slowest step. (rate determining step or key step)

$$Ph \longrightarrow C \longrightarrow H \xrightarrow{pH \longrightarrow CH \longrightarrow O} Ph \longrightarrow C \longrightarrow O$$
slowest step (rds)

$$+ \operatorname{PhCH_2OH} \longleftarrow \operatorname{Ph} \longrightarrow \operatorname{C} + \operatorname{PhCH_2O}$$

294 (b)

CLICK HERE



Anti conformation is the most stable form of n- 308 (c) butane (Bulky groups far apart).

295 (a)

$$R - X \longrightarrow R^+ + X^- \xrightarrow{OH^-} R - OH.$$

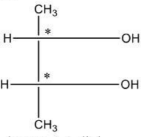
It has altogether different groups.

299 (b)



It is an ether and the name of ether is given as alkoxy alkane. So, its name is methoxy benzene.

300 (b)



butane-2-3-diol

Where C^* =asymmetric C atom

It is a symmetrical molecule, so the number of optically active stereomers= 2^{n-1}

(n=number of asymmetric C atom)

$$= 2^{2-1}$$

 $= 2^{-1} = 2$

302 (d)

Wöhler prepared urea from KCNO and (NH₄)₂SO₄

304 (a)

Electron donors having lone pair of electrons are nucleophile.

- (i) BF3 is not nucleophile because it does not have lone pair of electrons. It is infact Lewis acid because it accepts pair of electron
- (ii)NH₃, CN⁻ and OH⁻all have lone pair of electrons, so they are nucleophiles.

305 (b)

Due to asymmetric carbon atom.

306 **(b)**

Aryl carbonium ions are more stable than alkyl carbonium ions. The order of stability of carbocation is

Triphenyl methyl > Diphenyl methyl > Benzyl>Allyl>3°>2°>1°> methyl carbocation.

Follow IUPAC rules.

310 (a)

The dispersal of the charge stabilises the carbocation. More the number of alkyl groups; the greater the dispersal of positive charge and therefore, more the stability of carbocation, thus $C_2H_5^+ > CH_3^+$, $O - CH_3$ is also an electron donating group, thus it will increase the stability of carbocation, hence, the correct order of stability is C>B>A.

311 (c)

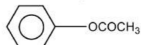
Chromatography method is used to separate sugars.

312 (c)

Only urea does not sublime while naphthalene, camphor and benzoic acid do

313 (c)

Esters are named by prefixing the name of the alkyl or aryl group (of OR' part) before the name of the parent acid and changing the suffix ic acid to ate. Hence, the structure of phenyl ehtanoate is



314 (b)

Inductive effect of groups is measured with respect

315 (d)

Methyl halides are methylating agents.

Nucleophiles are the species which have excess of electrons. Among the given species, the lone pair of nitrogen of pyrrole is involved in delocalisation of the ring, thus, are not available for donation. In aniline, the lone pair is involved in conjugation with the π -electrons of the ring while in pyridine, these are relatively free for donation. Thus, nitrogen of pyridine is most nucleophilic.



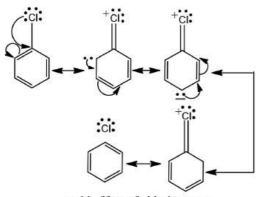
(phenyl and - COCH3 both are electron withdrawing groups, thus decreases the nuleophilicity of nitrogen).

317 (c)

Chlorobenzene is o, p directing in electrophilic substitution reaction. The directing influence is explained by +M of CI atom







+ M effect of chlorine atom

318 (b)

Free radicals are electrons deficient compounds. Alkyl groups are electron donor groups and they increase the stability of free radicals.

- : More the number of alkyl groups, more will be stability of free radicals.
- :: 3°>2°>1° is the correct order of stability of free radicals.

319 (b)

This give rise to net resultant of four C - Cl vectors equal to zero.

320 (b)

IUPAC name of compound.

Straight chain which contains large number of side chains taken as parent chain and counting starts from that side where the side chain is nearest.

3-methyl-5(1-methyl ethyl) octane.

321 (a)

The isomerism which arises due to rotation about a C-C is called conformational isomerism and the isomers are called conformational isomers or rotational isomers or conformers.

CH₃CH = CH₂CHOHCH₃ has one asymmetric carbon.

323 (b)

- (i) In nucleophilic substitution reaction more powerful nucleophile replaces weaker nucleophile.
- (ii) In rearrangement reaction atoms replace their position within molecule.

(iii) In elimination reaction small molecules (e.g., H2O, NH3) are lost.

$$R - CH2CH2Cl + KOH (alc.) \xrightarrow{\Delta} RCH$$
$$= CH2 + KCl + H2O$$

- : KCl and H2Omolecules are lost during reaction.
- : It is an elimination reaction.

324 (c)

CH₃COCH₃ is simplest ketone.

326 (d)

$$\begin{array}{c} \operatorname{CH_3} \\ | \\ \operatorname{CH_3} - \operatorname{C} - \operatorname{CH_3} \\ | \\ \operatorname{OH} \end{array}$$

is the formula of tertiary butyl alcohol as in it - OH group is attached to tertiary carbon.

327 (b)

The neopentane:

$$CH_3 \\ | \\ CH_3 - C - CH_3 \\ | \\ CH_3$$

IUPAC name:2,2-dimethyl propane

328 (d)

Triple bond possesses maximum bond energy.

329 (d)

If molecule having asymmetric carbon atom and is not superimoposable on its mirror image then it is chiral while if it is superimposable on its mirror image, it is achiral.

330 (a)

- (a) When optically active acid reacts with racemic mixture of an alcohol, it forms two types of isomeric esters. In each, the configuration of the chiral centre of acid will remain the same. So, the mixture will be optically active.
- 331 (d)

Due to H^- shift from C_2 to C_3 . Driving force is conjugation from oxygen. Also bulky gps hinders in hydride shift.

332 (a)

 C_6H_6 has six delocalized π -electrons.

333 (c)

Due to same molecular formula.

334 (d)

Cis - trans isomers generally contain double bonded carbon atoms.

335 (b)

Due to the presence of lone pair on N atom.

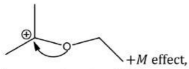


336 (c)

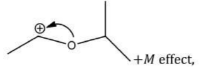
A molecule having doubly bonded carbon atoms shows geometrical isomerism only if both the doubly bonded carbon have altogether different group, i. e., $_{ba}C \equiv C_{ab}$ or $_{ab}C = C_{ac}$ or $_{dc}C = C_{ab}$.

337 (d)

I>III>IV

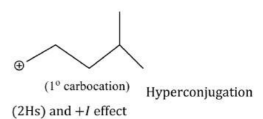


hyperconjugation (6Hs) and +I effect (2 Me-groups)



hyperconjugation (3Hs) and +I effect (1-Me group)

(2° carbocation) Hyperconjugation (5Hs) and +I effect



338 (b)

2-bromobutane has asymmetric carbon atom.

339 (b)

Br is less reactive and more selective and thus, formation of 3° free radical will be the major product.

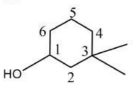
341 (b)

When organic compound containing both nitrogen and sulphur is fused with sodium, sodium thiocyanate is formed

343 (d)

Alcohols show position isomerism; Ethers show 353 (b) metamerism; Alcohols and ethers show functional isomerism.

344 (c)



Carbon with - OH group is given C1 thus it is 3, 3dimethyl-1-cyclohexanol.

345 **(b)**

-do-

346 (c)

The reactivity order for H atom is $3^{\circ}H > 2^{\circ}H >$ 1°H.

347 (a)

Pyrene is CCl₄.

349 (d)

3° carbonium ions are more stable as the +ve inductive effect disperses +ve charge on carbon atom.

352 (d)

Follow mechanism of free radical substitution.

The IUPAC name of this molecule is 2-bromo-3ethyl-1, 4-pentadiene.

355 (c)

N is pentavalent which is not possible.

356 (a)

Follow inductive effect.

357 (b)





M: (I), (II), (III), ; (I) and (II)

Cannot be separated by fractional distillation.

358 (c)

It is a fact.

359 (a)

Replacement of an atom or group by other atom or group is known as substitution reaction

$$\begin{array}{c}
O \\
\longrightarrow \\
CH_3
\end{array}$$

$$\begin{array}{c}
Me_2CuLi \\
\longrightarrow \\
H_2O
\end{array}$$

$$\begin{array}{c}
O \\
\longrightarrow \\
CH_3
\end{array}$$

$$\begin{array}{c}
CH_3
\end{array}$$

360 (b)

It is a fact.

361 (a)

Orbital interaction between the σ - bonds of a substituent group and a neighbouring π -orbital is known as hyperconjugation.

$$H \longrightarrow C \longrightarrow CH \longrightarrow CH_2 \longrightarrow H \longrightarrow C \longrightarrow CH_2$$

$$H \longrightarrow C \longrightarrow CH \longrightarrow CH_2 \longrightarrow H \longrightarrow CH_2$$

$$H \longrightarrow C \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow H^+ \longrightarrow CH_2$$

$$H \longrightarrow C \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2 \longrightarrow CH_2$$

$$H \longrightarrow CH \longrightarrow CH_2$$

$$H \longrightarrow CH$$

362 (c)

 sp^3 -hybridization with one position occupied by lone pair like NH_3 .

364 (a)

2-hexyne gives *trans*-2-hexene on treatment Li/NH₂

$$CH_3 - CH_2 - CH_2 - C \equiv C - CH_3 + H_2 \xrightarrow{Li/NH_3}$$

$$H$$

$$CH_3 - CH_2 - CH_2 - C = C - CH_3$$

H trans-2-hexene

365 (b)

Lassaigne's test is given by those nitrogenous compounds in which carbon is also present along with nitrogen.

In NH₂. NH₂. HCl, carbon is absent, so it does not given Lassaigne's test.

366 (b)

$$CH_3CH_2-\overset{\textstyle O}{\underset{\textstyle C}{\vdash}} -\overset{\textstyle O}{\underset{\textstyle C}{\vdash}} 1 \xrightarrow{\text{Heterolytic fission}} CH_3CH_2-\overset{\textstyle O}{\underset{\textstyle C}{\vdash}} + \check{C} \ I_3$$

 ${\overset{\circ}{C}}\ {\rm I}_3$ is most stable carbanion because of -I effect of I which disperses negative charge on ${\overset{\circ}{C}}$. Center



367 (d)

$$C_2H_5$$
—O—CH $\begin{pmatrix} 1 \\ CH_3 \\ CH_3 \\ 3 \end{pmatrix}$

2-ethoxy propane

The above compound is an ether and its name is written as alkoxy alkane. Oxy is attached with the lower group. Hence, the IUPAC name of above compound is 2-ethoxy propane.

368 (b)

In thin layer chromatography the relative adsorption of each component of the mixture is expressed in terms of retention factor (R_f)

distance moved by the spot centre from

$$R_f = \frac{\text{base line}}{\text{distance moved by the solvent from}}$$
the base line

369 (c)

 $(CH_3)_3CBr + H_2O \rightarrow (CH_3)_3C - OH + HBr$ Br is substituted by $-OH^-$ (nucleophile) S_N1 (unimolecular nuclerophilic substitution reaction)

370 (c)

 $C_6H_5\overset{ op}{C}HC_6H_5$ is the most stable since the +ve charge can be delocalized on both phenyl rings

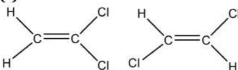
371 (a)

2-methyl propane-2-ol is tert-butyl alcohol. ${\it CH}_3$





372 (c)



Since, in the above structures, position of Cl is different, these are position isomers, which is a type of structural isomerism.

374 (b)

More directionally concentrated orbitals show more overlapping. Also more closer are shells to the nucleus more is overlapping.

375 (b)

Removal of H from ketone gives resonance stabilized carbanion.

376 (d)

380 (c)

$$\text{CH}_3\text{CH}_2\text{CH}_3 \xrightarrow{\text{Hetrolysis}} \overline{\text{CH}}_3 + \text{CH}_3 \xrightarrow{\text{CH}}_2, \text{CH}_3 \xrightarrow{\text{CH}}_2 \text{ is more stable than CH}_3 \xrightarrow{\text{CH}}_2 \overline{\text{CH}}_2$$

due to dispersal of +ve charge on ethylium ion on account of +ve inductive effect. Thus,

propane will not give $\overset{+}{\text{CH}}_3$ and CH_3 $\overline{\text{C}}\text{H}_2$.

381 (d)

Delocalised electrons are present in benzene, 1, 3-butadiene and 1,3,5-hexatriene

$$\bigcirc \longrightarrow \bigcirc _{(I)}$$

382 (a)

A characteristic of dextrorotatory.

383 **(c)**

It is a fact.

384 **(c)**

If a liquid decomposes at or below its boiling point, it is purified by vacuum distillation, impure glycerine is purified by this method

386 (a)

Distillation process is not used for purification of solid impurities. It is used for the purification of liquids which boils without decomposition and contains non-volatile impurities.

387 (b)

$$CH_3CH_2X \rightarrow CH_2 = CH_2(sp^3 \text{ to } sp^2);$$

 $CH_2 = CHX \rightarrow CH \equiv CH(sp^2 \text{ to } sp);$
 $CH_2XCH_2CH_2X \rightarrow \Delta \text{ (No change)}.$

377 **(b**)

Follow IUPAC rules.

378 (c)

It is a fact.

379 (d)

 $S_N 1$ (Unimolecular nucleophilic substitution reactions)

Rate∝ (substrate)

i.e., iii>ii>iv>v

Rate determining step in the formation of carbocation depends on the stability of carbocation formed. The stability of carbocations follow the order

$$(C_6H_5)_2 C^+(CH_3) > (C_6H_5)_2^+CH > (CH_3)_2^+CH > CH_3CH_2^+$$

 \therefore Order of $S_N 1$ reacticity is
 $(C_6H_5)_2C(CH_3)Br > (C_6H_5)_2CHBr > (CH_3)_3CBr$
 $> (CH_3)_2CHBr > C_2H_5Br$

 sp^3 , sp^2 and sp– orbitals are at 109°28′, 120° and 180°.

388 (b)

2°H is more reactive than 1°.

389 (b)

Dehydration of alcohol involves the loss of two atoms or groups from the adjacent carbon atoms, hence it is an example of β -elimination reaction.

$$\begin{array}{ccc} \mathbf{\beta} & \mathbf{\alpha} & & \\ \mathrm{CH_{3}CH_{2}OH} & \xrightarrow{+\mathrm{H}^{+}} & \mathbf{\beta} & & + \\ & \mathrm{CH_{3}CH_{2}OH_{2}} & & \\ & \mathrm{ethanol} & & & \end{array}$$

$$\beta$$
 α
 $CH_3CH_2^+$
 \rightarrow
 CH_2
 \rightarrow
 CH_2
 \rightarrow
 CH_2
 \rightarrow
 CH_2

391 (d)

(i) Eantiomers are pair of optical isomers which are related as non-superimposable mirror images of each other.





- (ii) Diasteriomers are pair of optical isomers which cannot be related as non-superimposable mirror images of each other.
- \therefore The only correct statement about given structures is that (*A*) and (*B*) are enantiomers.

392 (a)

Two positive charges present at the adjacent place, elevates the energy, thus lowers the stability most.

393 (d)

Due to H-bonding.

394 (a)

Follow inductive effect.

395 (b)

Naphthalene and benzoic acid cannot be separated by the sublimation method because the naphthalene and benzoic acid both are sublimes on heating. They are separated by hot water in which benzoic acid dissolves but naphthalene does not

396 (d)

All are used as dehydrohalogenating agent.

$$C_2H_5X$$
 $\xrightarrow{\text{(i) KOH alc.}}$ C_2H_5ONa $\xrightarrow{\text{or (iii) } C_2H_5ONa}$ $C_2H_4 + HX$.

397 (c)

In the given electrophile

$$\left[CH_3 - C \left< \begin{matrix} O \\ X \end{matrix} \right], \left[CH_3 C - O \right]$$

Group is the same. So, only X affects their activity, i.e., we have to discuss activity due to

(a)
$$- OCH_3$$
 (b) $-CI$

$$---$$
N $\stackrel{\mathsf{Me}}{\stackrel{}{\stackrel{}}{\stackrel{}}}$ (d) - S - CH₃

Since, amines are less actives, therefore, electrophile (c) will be least active.

398 (c)

The positions of Cl are different.

399 (a)

Higher the stability of carbocation, faster is the reaction because $S_N \mathbf{1}$ reactions involve the formation of carbocation intermediate.

Me
$$(2^{\circ} \text{ allylic}) \qquad (6 \text{ } \alpha \text{Hs}) \qquad (C)$$

$$(B) \qquad (C) \qquad \qquad \downarrow^{+} \text{CH}_{3}$$

$$(2 \text{ } \alpha \text{Hs}) \qquad (A)$$

400 **(b)**

Both Wolff-Kishner and Clemmensen reduction are used to convert CO to CH_2

The later is not suitable as it will also attack – OH group of ring.

401 (d)

In the following carbocation; H/CH_3 that is most likely to migrate to the positively charged carbon is

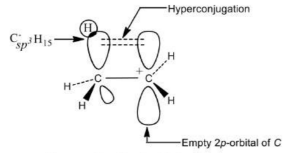
(A resonance stabilised carbocation)

402 (b)

Hyperconjugation arises due to the partial overlap of a $sp^3 - s$ (a C-H bond) with the empty p —orbital of an adjacent positively charged carbon atom.







Hyperconjugation in ethyl cation

403 (c)

In the triphenyl methyl carbonium ion the π electrons of all the three benzene rings are delocalised with the vacant p-orbital of central carbon atom. So, it is resonance stabilised. It is the most stable of all the carbonium ions given

$$CH_3$$
 CH_3
 CH_3
 CH_3
 CH_3
 CH_3 is stabilised by hyperconjugattion, a second order resonance.

404 (c)

During nitration of benzene the attacking electrophile is NO_2^+ . It is formed as follows by reaction between HNO3 and H2SO4.

(I)
$$HNO_3 + H_2SO_4 \longrightarrow H \longrightarrow O \longrightarrow NO_2 + HSO_4$$

(II)
$$H \longrightarrow O \longrightarrow NO_2 \longrightarrow H_2O + NO_2^+$$
 nitronium

405 (b)

The R and S enantiomers of an optically active compound differ in their optical rotation of plane polarised light.

406 (a)

A carbon atom which is attached by four different group is called chiral centre of asymmetric carbon atom. (+)-glucose has four chiral centres.

408 **(b)**

 C_2H_2 has two π -bonds.

409 (b)

The various structure derived for a molecule but none of them truly represents all the properties of that molecule are said to be canonical forms and the molecule is said to show resonance.

410 (d)

The possible primary amine with the formula C₄H₁₁N are

$$\begin{array}{c} \text{(i) } \text{CH}_{3}\text{CH}_{2}\text{CH}_{2}\text{CH}_{2}\text{NH}_{2} \\ \text{CH}_{3} \\ \text{|} \\ \text{(ii) } \text{CH}_{3} - \text{CH} - \text{CH}_{2} - \text{NH}_{2} \\ \text{CH}_{3} \end{array}$$

(iv)
$$\mathrm{CH_3} - \mathrm{CH_2} - \mathrm{CH} - \mathrm{NH_2}$$

411 (c)

Stereoisomerism is of two types-optical and geometrical.

412 (a)

Catenation is the tendency to unite atoms of an element to form a long carbon chain or ring.

414 (b)

Electron withdrawing group has – I effect while electron donating group has +I effect. In CH_3COOH , the alkyl group $(-CH_3)$ due to its greater +I effect increases the electron density on oxygen atom of the O-H bond. Due to this the release of H+ ion in acetic acid will be more difficult as compared to formic acid.

CLICK HERE



It is a fact.

416 (b)

If positive charge is present on nitrogen then positive charge will not be in conjugation to the ring because in this case nitrogen will become pentavalent

418 (c)

HNO3 is added to decompose Na2S and NaCN if present

$$Na_2S + 2HNO_3 \rightarrow 2NaNO_3 + H_2S \uparrow$$

 $NaCN + HNO_3 \rightarrow NaNO_3 + HCN \uparrow$

419 (c)

It is a fact.

420 (c)

The definition of electromeric effect.

421 (c)

It has no asymmetric carbon, however it shows geometrical isomerism.

422 (a)

Sbstituion of -OH group from a substrate can be easily made by PCl₅.

423 (c)



Chair form is unsymmetrical due to absence of any element of symmetry.

424 (a)

When two compounds have similar molecular formula but differ in the functional group then the isomerism is called functional group isomerism i.e.,

CH₃CH₂CHO and CH₃. CCH₃

426 (d)

Count σ -and π -bonds on each carbon and report hybridization.

427 (d)

Carbonyl compounds show nucleophilic addition.

428 (a)

Twelve in all

(a) Six geometrical isomers

$$(i)$$
 $C \longrightarrow C$
 $C \longrightarrow C$

$$(ii)$$
 H_3CH_2C $C = C$

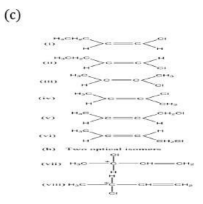
$$(iii) \xrightarrow{H_3C} C = C < CH_3$$

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

$$(v) \qquad \begin{array}{c} H_3C \\ \\ \\ H \end{array} \qquad C \longrightarrow C \\ \begin{array}{c} CH_2C \\ \\ \\ H \end{array}$$

(vi)
$$H_3C$$
 $C = C$ CH_2C

(b) Two optical isomers



429 (c)



Ortho and para- nitrohenol is separated by distillation because p —nitrophenol has higher boiling point than o-nitrophenol due to Hbonding.

430 **(b)**

Electromeric effect implies complete transfer of π electrons in presence of a reagent. Since, simple ethers do not contain a multiple bond, therefore, they do not show electromeric effect

431 (b)

It is definition of electromeric effect.

432 (a)

Mass of silver salt=0.4 g

Mass of silver =0.26 g

Eq. mass of silver salt/Eq. mass of Ag= wt.of silver salt

wt.of silver

Eq. mass of silver salt = $\frac{108 \times 0.4}{0.26}$ = 166

Eq. mass of acid = 166 - 108 = 58

433 (a)

The S_N 2 mechanism always involves 100% inversion since nucleophile attacks from back side of leaving group.

$$R - X \xrightarrow{OH} H\overline{O} - R - X$$
 $\downarrow HO - R + X \xrightarrow{\bullet}$

434 (a)

Follow IUPAC rules.

435 (d)

Best leaving group (poorest nucleophile) is Cl^{\oplus} , thus fastest reaction is with Cl.

436 (c)

Note that propyl (propan-l-ol) and isopropyl alcohol (propan-2-ol) are position isomers.

Empirical formula of acid = CH₂O₂

We know that molecular formula = n (empirical

If n = 1 molecular formula = $(CH_2O_2)_1 = CH_2O_2$

If n = 2 molecular formula = $(CH_2O_2)_2 = C_2H_4O_4$

If n = 3 molecular formula = $(CH_2O_2)_3 = C_3H_6O_6$

Thus, the probable molecular formula = CH_2O_2

Excited state of carbon is $2s^12p^3$.

$$C = 10.5 \text{ g} = \frac{10.5}{12} \text{ mol} = 0.87 \text{ mol}$$

$$H = 1 g = \frac{1}{1} mol = 1 mol$$

$$pV = nRT = \frac{w}{m}RT$$

$$1 \times 1 = \frac{2.4}{m} \times 0.082 \times 400$$

$$m = 79$$

Hence, the hydrocarbon is C₆H₇

440 (c)

There are four stereoisomers

$$cis - R$$
 $cis - S$

$$trans - R$$
 $trans - S$

441 (c)

IUPAC name is N-methyl methanamine.

442 (d)

Oil and water are immiscible liquids thus are separated by using separating funnel

443 (a)

Can be solved on the basis of hyperconjugative structures

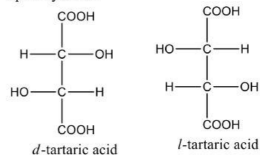
444 (d)

Total number of optical isomers= $(2)^n$

(where n=number of asymmetric carbon atom). $=(2)^2=4$

Out of these four optical isomers two are meso structures which are optically in active.

 \therefore Only two structures d and l- tartaric acid are optically active.



446 (d)

Three coordinate bond on O atom.

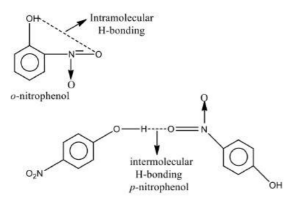
447 (c)

Furan is heterocyclic and aromatic due to Huckel's rule of aromaticity, i. e., 6π -electrons.

448 (d)

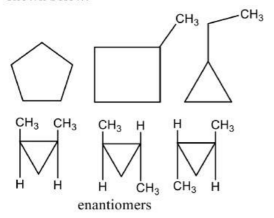
In mixture of o-nitrophenol, p-nitrophenol, onitrophenol is steam volatile due to intra molecular hydrogen bonding whereas pnitrophenol is less volatile due to inter molecular hydrogen bonding.





449 (c)

The total number of cyclic isomers are six as shown below.



451 (d)

Electromeric effect involves complete transfer of π -electron pair to more electronegative atom on the need of attacking reagent.

452 (c)

Cannizzaro reaction involves oxidation as well as reduction of aldehydes having lack of $\alpha-H$ atom. The mechanism of this reaction is as

(I) Attack of OH- on carbonyl carbon

$$R - C = O + OH - R - C = O$$

(ii) Transfer of hydride ion

(II) Transfer of hydride ion

453 (d)

C: H: Br =
$$\frac{2.27}{12}$$
: $\frac{5.69}{1}$: $\frac{65.04}{80}$
= 2.43: 5.69: 0.813
= 3: 7: 1
or empirical formula = C_3H_7Br

454 (b)

The terminal 'e' of ene is retained if suffix name starts with consonant.

455 (d)

IUPAC name=3, 3-dimethyl-1-butene.

456 (b)

CaO is added to NaOH to retard activity of NaOH, otherwise decarboxylation of acids will occur more violently.

 $RCOONa \xrightarrow{NaOH + CaO} R - H + Na_2CO_3$

457 (b)

$$\begin{array}{cccc} \operatorname{CH}_3 & \operatorname{CH}_3 \\ & | & | \\ \operatorname{CH}_3 - \operatorname{C} - \operatorname{Cl} + \operatorname{OH}^- & \rightarrow \operatorname{HO} - \operatorname{C} - \operatorname{CH}_3 + \operatorname{Cl}^- \\ & | & | \\ \operatorname{CH}_3 & \operatorname{CH}_3 \end{array}$$

Rate \propto [t-butyl chloride]

Tertiary butyl carbocation is first formed which is more stable

458 (c)

In heterolysis, the covalent bond is broken in such a way that one species (less electronegative) is deprived if its own electron, while the other species gain both the electrons

$$CH_3CH_2CH_3 \longrightarrow CH_3 + C_2H_5$$

459 (c)

Both have different functional groups, i.e., -CN and -NC.

460 (b)

A mixture of 50-50% of d and its l form is called racemic mixture.

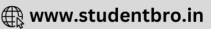
461 (d)

Sublimation is the process employed for those solids which convert directly into vapours on heating without converting into liquid phase

462 (c)

It is the definition of optical activity.

463 (a)



Stability of alkyl carbanion ∝

 $\xrightarrow[\text{magnitude of negative charge}]{\text{magnitude of negative charge}}$ and magnitude of negative charge $\propto +I$ power of the group. Hence, acetylenic carbanion is more stable than vinylic carbanion which is more stable than alkyl carbanion

465 (b)

Follow IUPAC rules.

466 (b)

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

468 (a)

A carboxylic acid is stronger acid than phenol, hence both III and IV are stronger acids than both I and II. Also IV has a methyl group that gives electrons donating inductive effect and decreases the acid strength. Therefore, III is stronger acid than IV. Between I and II, the dominate electron withdrawing inductive effect of chlorine increases acid strength of phenol slightly, hence II is stronger of phenol slightly, hence, II is stronger acid than I.

Thus, the overall order is: (a) III>IV>II>I.

469 (a)

All neutral covalent compound in which central atom has incomplete octet are electrophile. For example BeCl₂, BH₃, ZnCl₂, AlCl₃

470 (d)

Planar hexagon conformer has considerable angle strain due to the fact that its bonds are not 109.5°. It also has torsional strain. Due to presence of these strains planar hexagon conformer of cyclohexane is least stable.

471 (a)

Both have same molecular formula.

472 (a)

Due to resonance partial double bond character is created on vinyl chloride. So, chlorine atom is not replaced easily

473 (a)

Glycerol is a trihydric alcohol. It is $CH_2OH - CHOH - CH_2OH$. It contains three hydroxyl group. It is present in nature in oils and fats as its carboxylic esters.

474 (d)

Once the carbocation is formed as an intermediate, the nucleophile Cl^- and OH^- present in solution also attach it in addition of Br^- .

$$CH_{2} \xrightarrow{+\delta} CH_{2} + Br \xrightarrow{-\delta} Br \xrightarrow{+\delta} CH_{2} \xrightarrow{-CH_{2}} OH \xrightarrow{-CH_{2}} Products$$

$$Br$$

475 (a)

It is arised on the need of attacking reagent, e.g.,

$$C = C \left\langle \xrightarrow{A.R} \right\rangle \stackrel{+}{C} - \overline{C} \left\langle \right\rangle$$

478 **(b)**

An organic reaction intermediate, neutral species having divalent carbon atom with six valence electron out of which two are present in same orbital with opposite spin is called singlet carbene.

479 **(**a)

Nitration or aromatic compounds takes place by an electrophile. The electrophile will be more attracted towards electron rich position in benzene ring. Hence, electron donating groups will be easily nitrated.

Toluene will be most easily nitrated among these compounds due to presence of electron donating group (i.e., CH_3).

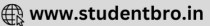
Nitrobenzene will be most slowly nitrated due to the presence of electron withdrawing group $(i.e., NO_2)$.

 ${\rm CH_3NO_2}$ will be formed by free radical substitution of CH_4

480 (b)

Optical isomerism is shown by an asymmetric carbon atom which has a carbon atom attached to four different atoms or groups.





So, butanol-2 is optically active.

$$\begin{array}{c} & \text{H} \\ \mid \\ \text{H}_3\text{C} - \text{CH}_2 - \text{C} - \text{CH}_3 \\ \mid \\ \text{OH} \\ \text{butanol-2} \end{array}$$

481 (c)

When sodium or potassium alkoxide is heated with an alkyl halide to give ether, this reaction is known as Williamson's synthesis.

$$RONa + R'X \rightarrow R - O - R' + NaX$$

This is an example of nucleophilic substitution and follow $S_N 2$ mechanism.

482 (d)

These are characteristics of free radicals.

483 (a)

Addition of \mbox{Br}_2 on ethane follow electrophilic addition

$$CH_2$$
= CH_2 + Br_2 - CH_2 - CH_2

Intermediate is cyclic bromonium ion

484 (d)

Metamers of ethyl propionate are as $CH_3COOC_3H_7, C_3H_7COOCH_3$

485 (c)

Each carbon has two σ -and two π -bonds.

486 (b)

It should be 4-ethyl-3-methyl heptane.

487 (c)

Due to asymmetric carbon atom in it.

488 (b)

This reaction is governed by Saytzeff's rule. According to this rule the elimination of β -hydrogen atom take place from the carbon having the lesser number of H-atoms or in other words a stable alkene is formed. (More substituted alkene is more stable)

489 (a)

Due to same functional group.

490 (a)

496 (d)

CH₃-CH-CH-COOH has two asymmetric carbon atoms and OH CH₂

molecule has no symmetry. Thus, number of optical isomers $= 2^n = 2^2 = 4$.

On chlorination of 2-methyl butane

$$\begin{array}{c|c}
\hline
Cl \\
\hline
Cl_2/h\nu
\end{array}
+
\begin{array}{c|c}
Cl \\
*
\end{array}$$

$$\begin{array}{c|c}
Cl \\
*
\end{array}$$

2-chiral compound are formed.

491 (d)

Due to the presence of asymmetric carbon atom.

492 (c)

Two isomers

$$\begin{array}{c} \operatorname{CH}_3 - \operatorname{CH} - \operatorname{CH}_3 \\ | \\ \operatorname{CH}_3 \\ \textit{Iso}\text{-butane} \\ \operatorname{CH}_3 - \operatorname{CH}_2 - \operatorname{CH}_2 - \operatorname{CH}_3 \end{array}$$

n-butane

494 (d)

Ethyl acetoacetate shows tautomerism.

0

||
$$CH_3 - C - CH_2 \cdot COOC_2H_5$$

Keto form

OH

|
 $CH_3 - C = CH \cdot COOC_2H_5$

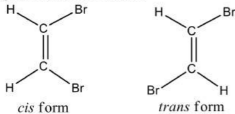
enol form

495 (d)

The amount of enolic form is highest in acetyl due to the stabilisation of enolic form by hydrogen bonding

497 (c)

Due to restricted rotation about double bond, the alkene shows geometrical isomerism because the relative position of atoms or groups attached to the carbon atoms of the double bond get fixed. If same groups or atoms attached with double bond bearing carbon, then alkene doesn't show geometrical isomerism.



498 (a)

Due to +I effect of CH_3 in toulene, it is more reactive than bezene. Due to electron withdrawing nature of - COOH group in benzoic acid and - NO2 group in nitrobenzene, both benzoic acid and nitrobenzene are less reactive than benzene.

499 (b)

These are isopentane, neopentane and n-pentane.

(a) Ethylene dichloride and ethylidine chloride both react with alc. KOH to produce ethyne.

$$H_3C - CHCl_2 + 2KOH(alc.) \rightarrow$$

$$HC \equiv CH + 2KCl + 2H_2O$$
 504 **(b)**

$$ClCH_2 - CH_2Cl + 2KOH alc. \rightarrow$$

$$HC \equiv CH + 2KCl + 2H_2O$$

(b) They are position isomers because they have same molecular formula but different position of chlorine atom.

ethylidene dichloride ethylene dichloride

- (c) : Their molecular formula is same
- : They have same percentage of chlorine.
- (d) They give different product on hydrolysis $CH_3CHCl_2 + 2KOH(aq) \rightarrow$ ethyledene chloride

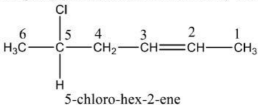
$$\begin{array}{c} & & & & \\ & \parallel \\ \text{CH}_3 - \text{CH(OH)}_2 \xrightarrow{-\text{H}_2\text{O}} \text{CH}_3 - \text{CH} \\ \text{Unstable} & \text{acetaldehyde} \\ \text{CH}_2\text{Cl} - \text{CH}_2\text{Cl} + 2\text{KOH (aq)} \rightarrow \text{CH}_2 - \text{CH}_2 \end{array}$$

ethylene dichloride

OH OH ethylene glycol

501 (a)

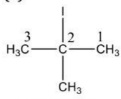
First the longest continuous chain of carbon atoms is selected. Now numbered the chain from the side containing senior functional group (i.e., the group placed above in the seniority table).



$$CH_3 - CH - CH - CH_3$$
 $CH_3 - CH - CH_3$
 $CH_3 - CH - CH_3$

2-methyl butan-2-ol

503 (d)



t- butyl iodide

Its IUPAC name is 2-iodo-2-methyl propane.

Number of hybrid orbitals = number of σ bonds+number of lps

∴Number of hybrid orbitals =3+0=3 Hence, hybridisation is sp^2 and geometry is planar.

506 (b)

Follow the concept of hyperconjugation.

507 (c)

On the basis of stability of carbocation formed.

508 (a)

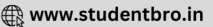
 $\boldsymbol{\varphi}$ is phenyl group, the question can be solved on the basis of number of conjugative structures

509 (a)

 $S_N 2$ order: methyl $> 1^{\circ} > 2^{\circ} > 3^{\circ}$.

510 (c)





Activating groups like – OCH_3 , –OH etc activates the benzene ring towards electrophilic substitution while deactivating groups like– NO_2 , –COOH etc. deactivates the benzene ring towards electrophilic substitution. Thus, order of reaction towards electrophile (of the given compounds) is as

I>II>III.

511 (a)

Electron deficient species or electron acceptor is electrophile. For example

 CH_3 , CH_2 , CX_2

513 (b)

As the min mol wt. must have at least one S-atom so

$$S\% = \frac{\text{wt. of one S} - \text{atom}}{\text{min. mol. wt}} \times 100$$
$$4 = \frac{32}{\text{min mol. wt}} \times 100$$
$$\text{Min mol wt.} \frac{32 \times 100}{4} = 800$$

514 (b)

A carbonium ion is sp^2 -hybridized.

515 (a)

It is a fact.

517 (c)

 $Or tho \ and \ para \ directing \ groups \ facilitate \ the \\ ring \ for \ electrophilic \ substitution \ reaction. - NH_2 \\ group \ increase \ electron \ density \ in \ ring, \ hece \\ activite \ it \ is \ to \ electrophilic \ substitutipon.$

518 (d)

n-pentanol, 2-pentanol, 3-pentanol, 2-methylbutanol, 2-methylbutan-2-ol, 3-methylbutanol, 2, 2-dimethypropanol, and 3-methylbutan-2-ol (8 isomers)

519 (a)

Substances which sublimes on heating is usually purified by sublimation. Hence, naphthalene is purified by sublimation.

520 (d)

There are total $6\alpha - H$ to sp^2 carbon and they all can participate in hyperconjugation.

521 (d)

The structural formula of fumaric acid is

523 (b)

A chloride linked with alkyl group is replaced with $AgNO_3$ and give white precipitate of AgCl.

$$\begin{array}{c|c} CH_2CI & CH_2NO_3 \\ \hline \\ + AgNO_3 & \\ \hline \\ Br & \\ \end{array} + AgCI \underset{precipitate}{\downarrow} \\ precipitate \\ \end{array}$$

524 (c)

Br is replaced by - OH.

525 (c)

Free radicals are represented by putting dot on entity.

527 (d)

Kjeldahl and Duma's methods are used for the quantitative estimation of nitrogen in an organic compound. In the Kjeldahl method, the nitrogen element of organic compound is changed to the ammonia.

528 (d)

Lesser is bond energy of 2p-2p overlapping in C—C, more is its reactivity than C—H bond showing 2p-1s overlapping.

529 (a)

Carbon atom in singlet carbone is sp^2 -hybridized.

531 (b)

C₆H₅CH₂⁺ is stabilized by conjugation while intermediates of rest of the compounds given are stabilized by hyperconjugation

532 (a)



CH₃CH₂CH₂CH₂CH₃ is n-pentane.

Resonance in benzene gives rise to identical C-C bond lengths.

534 (a)

During nitration benzene ring is attacked by NO₂⁺ and hydrogen of benzene ring is replaced by NO2

: Nitration of benzene is electrophilic substitution because NO₂⁺ is an electrophile.

535 (b)

$$\begin{array}{c|c} CH_3(CH_2)_5 \\ \hline \\ H_3C \\ \hline \\ H \end{array} \begin{array}{c} C-Br \\ \hline \\ optical \\ inversion \\ \hline \\ H \end{array} \begin{array}{c} OH^- \\ \hline \\ CH_3 \\ \hline \\ CH_3 \\ \hline \end{array}$$

In this reaction inversion takes place. Hence, it is an example of S_N2 reaction. In this mechanism the attack of OH⁻ ions take place from the back side while the Br ion leaves from the front side

536 (b)

Organic compound containing nitrogen is fused with a small piece of sodium metal to form NaCN. $Na + C + N \rightarrow NaCN$

> from organic compound

537 (d)

The order of nucleophilicity depends upon the nature of alkyl group 'R' on which a nucleophile to 548 (a) attack as well as on nature of solvent. However, if these are same, then weaker is acid, stronger is base, i.e., stronger is nucleophilicity. This acidic character is.

 $HI > HBr > HCl > HCN > H_2O > EtOH$

540 (d)

Only two unsymmetrical ether is possible from the formula $C_4H_{10}O$ viz

 $CH_3CH_2CH_2 - O - CH_3$ methoxy propane

541 (c)

S_N1 mechanism gives rise to 50% inversion as it involves front seat as well as back seat substitution. This leads to racemic products.

542 (c)

Canonical structures proposed in resonance are not the real structure of compound. The compound 554 (d) showing resonance has a definite structure which can however be not drawn on paper.

543 (d)

All the names are correct. Options (d) is IUPAC

544 (b)

On adding SCN⁻ to an aqueous solution of Fe(NO₃)₃, a blood red colour, due to formation of $[Fe(H_2O)_5(SCN)]^{2+}$ complex is obtained. This test is used for the detection of Fe^{3+} ion.

$$SCN^{-} + Fe(NO_3)_3 + 5H_2O \rightarrow [Fe(OH_2)_5(SCN)]^{2+} + 3NO_3^{-}$$

Blood red colour

545 (d)

In case of alkyl carbocations as the number of R group decreases stability decreases. Thus, the correct order of stability of carbocation is

$$R_3C^+ > R_2CH^+ > RCH_2 > CH_3$$

546 (b)

Ortho nitrophenol is the most acidic because electron withdraeing group increases acidic character due to - I effect of NO2

o - nitrophenol

 $CCl_2 = CCl_2$ has ethene like structure (i.e., sp^2 -CCl₄ has CH₄ hybridization); like structure, i. e., sp^3 -hybridization.

549 (b)

 $C_6H_5C \equiv N$ and $C_6H_5N \equiv C$ are functional isomers.

551 (c)

Second ring is in conjugation with lone pair of oxygen

552 (d)

% of N
$$\frac{28}{22400} \times \frac{\text{volume of N}_2 \text{at NTP}}{\text{wt. of compound}} \times 100$$

= $\frac{28}{22400} \times \frac{224}{1.18} \times 100$
= $\frac{28}{1.18} = 23.728$

553 (d)

In (II) and (IV) lone pair is involved in resonance.

A primary carbon is one which is joined to 1 carbon atom. A secondary carbon atom is joined to two



carbon atoms and the tertiary carbon is attached to three carbon atoms. A quaternary carbon has all its four valencies attached to carbon atoms.

555 (a)

The compounds which contain active methylene group at the adjacent position of carboxyl group show tautormerism.



This compound does not contain active methylene group, hence does not exhibit tautomerism.

Moreover, this compound is highly stable due to extensive cross-conjugation.

557 (b)

CH3CH2OH and CH3OCH3 have different functional groups.

(ie., -OH in a alcohol and -O-in ether), hence they are the example of functional isomerism.

558 (a)

Two pairs of *cis* and *trans* forms.

559 (b)

Stability of alkyl carbocations can be explained by inductive effect and hyperconjugation. According to these two effect the stability order is

$$R \stackrel{\oplus}{---} R > R \stackrel{\oplus}{---} CH - R > R \stackrel{\oplus}{---} CH_2 > CH_3$$

560 (d)

In the given compound four π - electrons of double bond and 1 lone pair on N atom leads to delocalization of six electrons

562 (a)

Due to resonance, extra stability in 3° carbocation. | 580 (a)

564 **(b)**

Follow IUPAC rules.

565 (b)

Na reacts with water less violently than K and Rb.

566 (a)

 $Na_2S + Na_2[Fe(CN)_5NO] \rightarrow Na_4[Fe(CN)_5NOS]$ Sod. extract pink/violet colour

567 **(b)**

Heterolytic bond fission produces +ve and -ve

568 (b)

Racemic mixture is formed by mixing two chiral compounds.

569 (c)

Distillation is applied if organic liquid is stable at its boiling point and contains a non-volatile impurity

570 **(b)**

Two double bonds are treated as different functional group with a triple bond.

571 (b)

Follow priority rule.

572 (c)

Aldehydes and ketones combine with a variety of compounds of the $Z - NH_2$ to form oxime

$$R_2$$
CO + NH₂OH \longrightarrow $\left[R_2$ C $\stackrel{\bigcirc}{\sim}$ NHOH $\right]$

$$\frac{\text{-H}_2\text{O}}{\text{-}} R_2\text{C=NOH}$$
 oxime

573 (b)

Addition of HCl is not peroxide effect and it occurs via electrophilic addition.

574 (b)

 $(CH_3)_3C - CH_2OH$ is neo -pentyl alcohol.

575 (d)

-do-

576 (d)

A lot of plant kingdom is made up of cellulose.

579 (b)

The different arrangement of atoms in space that results from the carbon-carbon single bond free rotation by 360° are called conformations or conformational isomers and this phenomenon is called conformational isomerism.

The molecule, which is optically active, has chiral centre, is expected to rotate the plane of polarised light.

One chiral center⇒optically active

Two chiral centres, but plane of symmetry within molecule⇒ optically inactive



583 (b)

Note the fact.

585 (d)

Alkanes do not show resonance.

586 (d)

Both CHCl3 and CCl4 burn with smoky flame although both are aliphatic. C₆H₅CH₂OH, aromatic but burns with non smoky flame. These are exceptions.

587 (b)

BF₃ is an electron deficient compound.

590 (a)

Isomers of propionic acid are as

$$\begin{array}{ccc} 0 & 0 \\ \parallel & \parallel \\ CH_3-C-OCH_3 & H-C-OC_2H_5 \\ \text{methyl ethanoate} & \text{ethyl formate} \end{array}$$

Heterolytic bond fission give rise to formation of ions.

592 (a)

This can be judged by comparing the stabilities of carboxylate ions formed. The most stable carboxylate ion is formed by strongest acid

593 (c)

CH₃NH₂ and CH₃OH are nucleophiles, CH₃ - Cl is

an electrophile. But $CH_3 - \overset{\delta_+}{C} = \overset{\delta_-}{=} \overset{\delta_-}{N}$ is a nucleophile due to the presence of a lone pair of electrons on N and is an electrophile due to the presence of a partial positive charge on C

594 (a)

No bond around chiral carbon is broken and so configuration will be retained.

595 (b)

Due to resonance in benzene.

Kjeldahl's method is used for estimation of nitrogen

597 (a)

Free radicals stability

Free radicals stability

$$C_6H_5$$
— \dot{c} — C_6H_5 > C_6H_5 — \dot{c} H C_6H_5

Highly stable by delocalisation

$$>$$
H₃C $-$ CH₃ $>$ H₃C $\stackrel{\bullet}{C}$ H₃ C C

9-hyperconjugative hydrogens and +I effect

598 (b)

HBr being better source of proton. It gives a H+ and a Br ion

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

Thus, H⁺attack the π bond of propene to form carbonium ion as

$$CH_3$$
 $\stackrel{+}{CH}$ + $Br^ \longrightarrow$ CH_3 $\stackrel{-}{CH}$ \longrightarrow CH_3 CH_3 CH_3 CH_3

599 (c)

Follow IUPAC rules.

600 (d)

Report prefixes in alphabetic order.

601 (c)

Sulphur is present in the sodium extract in the form of sodium sulphide (Na2S)

 $FeCl_3$ gives blood red colour with sodium extract contain N and S

602 (a)

The name of the compound (2Z,4Z)-2, 4-hexadiene.

603 (c)

It is a fact.

604 (a)

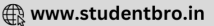
The number of stereoisomers=2' (Here, n =chiral carbon atom)

Thus, number of stereoisomers= 2^3 =8

605 (a)

The structural formula of the compound 5-nitro-3-methoxy-3-methyl hexanoyl chloride is as





606 (d)

Presence of methyl group on NH₃ molecule 608 (c) increases the tendency of N atom to lose electron

pair. However, tertiary, amines are less basic due to steric hindrance.

607 (a)

$$\begin{array}{lll} \text{CHCl} &=& \text{CHCH}_2\text{CH}_2\text{CH} &=& \text{CH}_2;\\ \text{CH}_2 &=& \text{CClCH}_2\text{CH}_2\text{CH} &=& \text{CH}_2;\\ \text{CH}_2 &=& \text{CHCHCH}_2\text{CH} &=& \text{CH}_2.\\ && & & & & & & \\ && & & & & & \\ && & & & & & \\ && & & & & & \\ && & & & & & \\ && & & & & & \\ && & & & & & \\ && & & & & & \\ \end{array}$$

2° carbocation is more stable.

609 (b)

In such cases where migrating group is cycloalkyl group, ring expansion may occur.

$$\begin{array}{c|c}
OH & OH \\
\hline
CH_3 & \xrightarrow{+H^+} \\
\hline
CH_3 & \xrightarrow{-H_2O}
\end{array}$$

$$\xrightarrow{\text{ring expansion}} \begin{array}{c} \overset{\text{CH}_3}{\longleftarrow} \overset{\text{CH}_3}{\longleftarrow} \\ \overset{\text{CH}_3}{\bigcirc} \end{array}$$

610 (d)

propanoic anhydride

In IUPAC system, anhydrides are named as alkanoic anhydride.

611 (c)

Stronger is an acid, weaker is its conjugate base or weaker is nucleophile. The acidic character order is:

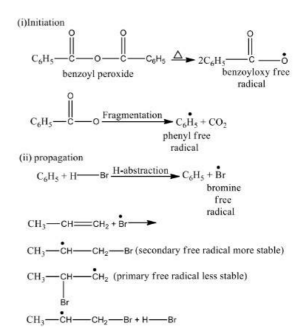
$$HF > H_2O > NH_3 > CH_4$$
.

613 (c)

$$CH_3 - CH = CH_2 + HBr \xrightarrow{Organic} CH_3 - CH_2 - CH_2 - Br$$

requires radical intermediate.

Mechanism



(iii) Termination

$$Br + Br \longrightarrow Br_2$$

614 (a)

Chiral molecules should not contain any kind of symmetry.

H-abstraction CH3-CH2-CH2-Br + Br

n - propyl bromide bromine

free radical

615 (d)

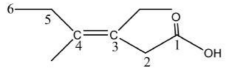
It is honour to Lavoisier.

616 (d)



Stability of carbanions increase with increasing in s-character of hybrid orbitals of carbon bearing charge therefore, the order is $sp^3 < sp^2 < sp$

617 (c)



E-3-ethyl-4-methyl hex-3-en-1-oic acid.

[The configuration of this compound is *E* because bulkier groups are present at opposite of the double bond.]

618 **(a)**

-do -

619 (c)

The substrate has three different types of B-H, therefore, first, three structural isomers of alkenes are expected as

The last two alkenes II and III are also a capable of showing geometrical isomerism hence two geometrical isomers for each of them will be counted giving a total of five isomers.

620 (c)

 $\mathrm{CH_2Cl_2}$ has sp^3 -hybridization and tetrahedral nature.

621 (a)

Chain initiation step involves formation of free radicals only.

622 (d)

Follow characteristics of S_N 2 mechanism.

623 (a)

Follow elimination of HBr from two ends.

624 (d)

 $^{\ominus}\mathrm{CH_{2}CHO}$ is the most stable carbanion since it is stabilize by resonance

0 || 0-

 $^-\mathrm{CH_2} - \mathrm{C} - \mathrm{H} \leftrightarrow \mathrm{CH_2} = \mathrm{C} - \mathrm{H}$

625 **(b)**

X-crown-Y, 18-crown-6

First number X is the total number of 'C' and 'O' atoms in the ring and second number Y is the number of oxygen atom in ring.

626 (b)

When the nucleophilic site is the same, nucleophilicity parallels basicity. It means more basic the nucleophile, stronger is the nucleophile.

 $H_{2}\overset{\bullet}{N}\left(I\right)$ is the most nucleophilic

Furthermore the NH_2 group away from the -C —group is not involved in resonance. Hence, its lone pair is reading available.

627 (c)

A compound could be optically active only when it contains, at least one asymmetric carbon atom or a chiral centre.

629 (d)

C: H: Cl:
$$0 = \frac{18.5}{12} : \frac{1.55}{1} : \frac{55.04}{35.5} : \frac{24.81}{16} = 1: 1: 1: 1$$

630 (b)

2-pentanone and 3-methyl-2-butanone are chain isomers because they differ in carbon skeleton.

CH₃CH₂CH₂COCH₃ 2-pentanone CH₃CHCOCH₃

CH₃ 3-methyl-2-butanone

631 (c)

Isomers of C4H10O are as follows

$$\begin{array}{c} \text{(i)} \ \ \text{H}_{3}\text{C} - \text{CH}_{2} - \text{CH}_{2} - \text{CH}_{2} - \text{OH} \\ \text{butanol-1} \end{array}$$

(ii)
$$H_3C - CH - CH_2 - CH_3$$

OH

butanol-2

 CH_3

(iii) CH₃ - CH - CH₂ - OH





2-methyl propan -2-ol

(v)
$$H_3C - CH_2 - O - CH_2 - CH_3$$

Diethyl ether

Hence, three isomeric ethers are possible.

632 (c)

Liebig's method is used to estimate carbon and hydrogen.

C and H
$$\xrightarrow{[O]}$$
 $CO_2 + H_2O$
%C = $\frac{12}{44} \times \frac{\text{weight of CO}_2}{\text{weight of compound}} \times 100$
%H = $\frac{2}{18} \times \frac{\text{weight of H}_2O}{\text{weight of compound}} \times 100$

633 (d)

There are six isomers possible for the compounds having molecular formula C_4H_8O , which are as follows

$$CH_{3}CH_{2}C \longrightarrow CH_{3}, CH_{3}CH_{2}CH_{2}CH_{0},$$

$$(i) \qquad (ii)$$

$$H_{3}C \longrightarrow CH.CHO$$

$$H_{3}C \longrightarrow CH_{2}OH$$

$$(cis) \qquad (iv)$$

$$H_{3}C \longrightarrow CH_{2}OH$$

$$(trans) \qquad (v)$$

$$CH_{2} \longrightarrow CH_{2}CH_{2}OH$$

634 (c)

$$C_2H_5$$
 and CH_3 C_{\oplus} C_{\oplus} C_{\oplus} C_{\oplus} C_{\oplus} are stabilised

(vi)

hyperconjugation. In ${\rm ^{C_6H_5CH_2}}$ benzyl group is resonance stabilised. In the triphenyl methyl carbonium ion, the $\pi\text{-electrons}$ of all the three benzene rings are delocalised with the vacant p- orbital of central carbon atom. So, it is resonance stabilised. Therefore, it is most stable of the given carbonium ions.

More the number of resonatic structures more will be the stability.

635 (d

Chlorine atoms are strongly electro negative (show negative inductive effect i.e., -I effect). They deactivate the ring towards electrophilic reaction.

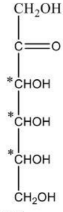
The increasing order of substituent-E towards electrophilic substitution is

$$-\mathsf{CCl}_3 < -\mathit{CHCl}_2 < -\mathit{CH}_2\mathsf{Cl} < -\mathit{CH}_3$$

636 (b)

Fructose has three chiral carbon atoms, hence the number of optical isomerism = $2^3 = 8$





637 (d)

O has two unpaired electrons in ground state.

638 (d)

An organic ion with a pair of available electrons and a negative charge on the central carbon atom is called a carbanion.

Electron attracting group (-CN, C=0) increases stability and electron releasing group $(-CH_3etc)$ decreases the stability of carbanion. In $(CH_3)_3C^-$, three $-CH_3$ groups (electron releasing group) are present, so it least stable.

639 (d)

The stability order of conformation of cyclohexane is chair>twist boat>boat>half chair. Hence, half chair is less stable due to torsional and angle strain.

640 (a)

It is a fact.

642 (a)

Formation of 2° carbocation, i.e.,

$$CH_3CH \xrightarrow{\Gamma} CH_2 \xrightarrow{H-H} CH_3 \overset{-\delta}{C}HCH_3$$

643 (d)

In CH₃CH₂OH, there is intermolecular H-bonding, while it is absent in isomeric ether CH₃OCH₃

- Larger heat is required to vaporise CH₃CH₂OH as compared to CH₃, OCH₃, thus (a) is incorrect.
- CH₃CH₂OH is less volatile than CH₃OCH₃, thus vapour pressures are different, thus
 (b) is incorrect.
- Boiling point of CH₃CH₂OH > CH₃OCH₃, thus (c) is incorrect.

Density = $\frac{\text{mass}}{\text{volume}}$, due to ideal behaviour at a given temperature and pressure volume and molar mass are same.

Hence, they have same vapour density.

644 (a)

Follow IUPAC rules.

646 (b)

% of chlorine =
$$\frac{35.5}{143.5} \times \frac{\text{mass of AgCl}}{\text{mass of the compound}} \times 100$$

= $\frac{35.5}{143.5} \times \frac{0.287}{0.099} \times 100$
= 71.71 %

647 (c)

Due to presence of delocalised π -electrons in the aromatic compounds, the electron density is maximum inside the ring. Therefore, aromatic compounds undergo electrophilic substitution reaction and resistance to addition reactions.

648 (a)

The acidic nature is $H_2O > C_2H_2 > C_2H_4 > C_2H_6$.

649 **(b)**

methyl CH_3 ,vinyl CH_2 =CH given options can be solved on the basis of conjugative and hyperconjugative structures

650 (a)

Follow IUPAC rules.

652 (a)

 $(CH_3)_3C - C(CH_3)_3$ has maximum number of alkyl groups in it.

653 (d)

Halogenation of methane is chain reaction, propagate through free radical

655 (a)

Follow IUPAC nomenclature.

658 (c)

 \because C – H bond in toluene has partial double bond character due to resonance.

∴ C – H bond in toluene has less energy as compared to others.

659 (b)

The reaction $(CH_3)_3C - Br \xrightarrow{H_2O} (CH_3)_3COH$ is an example of substitution reaction

660 (b)



Draw the isomers.

661 (b)

Distillation is used to purify liquids and based on difference in their boiling points. When the boiling points of liquids are very closed to each other, then fractional distillation is used.

662 (c)

The C-atoms attached to the triple bond lie in a straight line while the carbon of the ${\rm CH_2}$ group is inclined at an angle of 120°. Therefore, only 2 carbon atoms are linearly arranged

663 (c)

The conversion of α -glucose to β -glucose is called mutarotation.

664 (c)

Positively charged species in which central atom has incomplete octet is called electrophile, H^+, X^+, R^+ are electrophile

665 (b)

An organic reaction intermediate, neutral species having divalent carbon atom with six valence electron out of which two are present in same orbital with opposite spin is called singlet carbene

666 (a)

The structure of D-glyceraldehyde is as



The priority of groups is decided by the following rules

- (i) Atom having higher atomic number gets higher priority
- (ii) If the priority cannot be decided by rule 1 then the next atoms are considered for priority assignment.
- (ii) Where there is a = bond or=bond both atoms are considered to be duplicated or triplicated

$$\left(\begin{array}{c} H \\ - C \end{array}\right)$$
 has higher priority than – CH_2OH). Hence, the correct order of priority of groups in D-glyceraldehyde is as:

668 (c)

Nucleophile always attacks on electron deficient site. Presence of electron withdrawing groups such as NO₂, CHO etc decreases the electron density on benzene nucleus, hence such groups activate the ring towards nucleophilic attack. While presence of electron releasing groups such as R or OR increases the electron density, thus deactivates the nucleus towards nucleophilic attack.

 ${
m NO_2}$ group activates the ring more than Cl towards nucleophilic attack,

Hence reacts readily with nucleophile.



669 (d)

The increasing order is:

$$-CH_3 < CH_3 - CH_2 - < (CH_3)_2CH - < (CH_3)_3C -$$

670 (a)

The treatment with ${\rm FeCl_3}\,$ yield ferric ferrocyanide which has bulish green or prussian blue colouration

$$3Na_4[Fe(CN)_6] + 4FeCl_3 \rightarrow Fe_4[Fe(CN)_6]_3$$

671 (c)

Each π -bond contributes two π electrons and the two electrons of the lone pair are present in a p-orbital. Therefore, total number of π electrons is six

672 (a)

Allyl carbocations are more stable than the alkyl carbocations due to the resonance stabilization

673 (b)

The middle carbon has 2σ -and 2π -bonds.

675 (c)

Di-chloro acetic acid due to presence of two electron with drawing chloro groups (-I showing group) is more acidic than acetic acid $(+I \text{ showing} - \text{CH}_3\text{group})$.

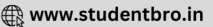
677 (b)

 $Na_3PO_4 + 3HNO_3 \rightarrow H_3PO_4 + 3NaNO_3$ sod. extract

$$H_3PO_4 + 12(NH_4)_2MoO_4 + 21HNO_3$$

 $\rightarrow (NH_4)_3PO_4.12MoO_3$
 $+ 21NH_4NO_3 + 12H_2O$





yellow

ppt.

678 **(b)**

1,2,3; 1,2,4 and 1,3,5-trimethyl benzene.

679 (b)

Hydride ion is formed when hydrogen accept a proton, so it has a tendency to donate electron. Since, hydride ion (H⁻) has a tendency to donate electron, it functions as nucleophile.

680 (b)

The structure of isomers from $C_2H_2Br_2$ are $CH_2 = CBr_2$;

$$CH_2 = CBr$$
;

$$c = c < Br$$

trans-isomer III

681 (a)

$$CH_3MgX + CH_3C$$

 $\equiv C - H \longrightarrow CH_4 + CH_3C$
 $\equiv C.MgX.$

682 (a)

The correct decreasing order of priority for the functional groups of organic compounds in the IUPAC system is

$$-COOH > SO_3H > -COOR > COCl > -CONH_2$$

> $-CN > -CH = O$

683 (b)

Optical isomerism is shown by compounds which have one or more chiral carbon atoms.

: It has asymmetric or chiral carbon atom.

: It shows optical isomerism.

684 (d)

 $S_N 2$ reaction does not involve ion formation, these infact involve formation of transition state

685 (a)

 pK_a value of carboxylic group is less than pK_a of $\stackrel{+}{\mathrm{NH_3}}$ in amino acid and $\stackrel{+}{\mathrm{---}\mathrm{NH_3}}$ (Z) will have

comparatively less pK_a than $\stackrel{---}{---} NH_3$ (Y) due to -I effect of carboxylic group. We know that acidic strength in inversely proportional to pK_a . Hence, correct order of acidic strength is

686 (a)

Chromatography is a modern technique used for the separation of mixtures into its components, purification of compounds and also to test the purity of compounds.

688 (d)

Kharasch effect involves addition of HBr.

689 (d)

These are characteristics of carbocations.

691 (b)

Symmetrical *trans*-form has non-polar nature.

694 (c)

The nucleophilic addition reaction is the characteristic addition of carbonyl compounds. Reactivity order of carbonyl compounds is in the order.

$$H C = 0 > H_3C C = 0 > H_3C C = 0$$

This is due to increase in the intensity of charge on carbon of carbonyl group due to +I effect of alkyl groups.

695 (c)

Chlorobenzene has only one deactivating group, .e., -Cl. In 2, 4-dinitrochlorobenzene three deactivating group, i.e., two $-\text{NO}_2$ and one -Cl are present and p —nitrochlorobenzene two deactivation groups, i.e., one NO_2 and one Cl is present. So, the order of reactivity is A > C > B.

696 (a

Acid has —COOH group whereas, ester has — COOR group.

697 (d)

The given reaction can be represented as $NaOH \rightarrow Na^+ + OH^-$

NaOH
$$\longrightarrow$$
 Na⁺ + OH⁻

OH

 $\begin{array}{c} CH_2CI \\ CH_3 \end{array}$
 $\begin{array}{c} \delta_- \\ CH_3 \end{array}$
 $\begin{array}{c} CH_3 \\ CH_3 \end{array}$

trasition state

Since in this reaction, a nucleophile replaces the other group, it is a example of nucleophilic substitution reaction.

The mechanism shows that the rate depends on the concentration of both alkyl halide and nucleophile. So, it is an example of S_N2(nucleophilic substitution of II order) reaction.

698 (a)

Follow IUPAC rules.

699 (d)

Follow IUPAC rules.

702 (a)

Carbonyl compounds undergoes nucleophilic addition reaction.

$$X - C - [X-shows negative inductive effect]$$

If group or atom attached with carbonyl carbon shows negative inductive effect, then it decreases electron density on carbonyl carbon and facilitate the attack of nucleophile, hence reactivity of carbonyl compound increases. The aromatic aldehydes and ketones are less reactive than their aliphatic analogues due to +R effect of benzene ring. The increasing order of the nucleophilic addition reaction in the following compounds will

 $CH_3CHO > CH_3COCH_3 > PhCOCH_3 > PhCOPh$

703 (b)

Chiral compounds which have one chiral centre. All four atoms or groups attached to carbon are different.

704 (b)

Due to the presence of methyl group positive inductive effect increases and the stability of carbocation also increases. The stability order of carbocation is

Tertiary > Secondary > Primary

705 (b)

According to IUPAC system ethers are named as alkoxy alkanes. The larger alkyl group forms the parent chain while lower alkyl group is taken with the ethereal oxygen and forms a part of alkoxy group.

$$CH_3$$
— O — CH_3 — CH_2 — CH_3
 CH_3

2-methoxybutane

706 (c)

H₃+O cannot accept electron pair.

707 (a)

Epoxide is ambident substrate for nucleophilic substitution reactions. In protonated epoxide carbon-2 and carbon-3 both aquire some positive charge due to the highly electronegative atom

708 (d)

Beilstein test is used to detect halogens in organic compounds.

709 (b)

Free radical chain reaction is initiated by UV light. It proceeds in three main steps like initiation, propagation and termination. It gives major products derived from most stable free radical

711 (b)

Follow IUPAC rules.

713 (d)

Elimination reactions involves removal of a molecule (HBr here) from a substrate.

714 (c)

ketoform

enol form

715 (b)

The species which are electron deficient and accept a pair of electron are called electrophile. Hence, SO3 is a electrophile as it contains an





electron deficient centre. While H_2O , NH_3 and R-O-R are nucleophiles.

718 (a)

$$\begin{array}{c|c} CH_3 & CH_3 \\ \hline CH_3-CH_2-C - CH_2-CH_3 & \stackrel{H^+}{\longrightarrow} & C_2H_5-\stackrel{C}{\oplus} - C_2H_5 \\ \hline OH & \end{array}$$

In the above reaction more stable carbocation is generated hence, the compound dehydrated very easily

719 (a)

In case of kjeldahl's method the percentage of N2 is then calculated from the amount of NH3

721 (b)

724 (b)

are geometrical isomers.

725 (b)

C₆H₅O⁻ possess less nucleophilicity due to stabilized nature of phenoxide ion. CH3OH is weaker acid than CH₃COOH and thus CH₃O⁻ is 735 (a) stronger base.

Acidic order: $CH_3COOH > H_2O > CH_3OH$

Vinyl chloride is least reactive for S_N reaction due to resonance

$$CH_2$$
=CH CH_2 -CH C

729 (b)

The chain propagation step involves the use of free radical and regeneration of another free radical.

730 (b)

CH3NC is methaneisonitrile.

732 (a)

Benzyl carbonium is more stable due to resonance and thus, benzyl chloride is more reactive.

733 (a)

It is a fact.

734 (b)

 $C_n H_{2n} O_2$ is general formula for open chain acid and ester.

$$n = 3 \quad C_3H_6O_2$$

$$0$$

$$\parallel$$

$$Acid \quad CH_3CH_2 - C - O - H$$

Such dehydrohalogenation follow E2 mechanism. The driving force of such reaction is the stability of alkene produced. Since, tertiary alkyl halide can give more substituted alkene, it reacts fastest followed by secondary and primary i. e., 3° > $2^{\circ} > 1^{\circ}$.

722 (c)

Central carbon atom is chiral carbon.

723 (d)

Those organic compounds, which are volatile in steam are purified by steam distillation. Since, aniline is a steam volatile compound, hence it is purified by steam distillation.

$$\begin{matrix} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\$$

Fumaric acid

The Cannizzaro reaction is as

$$\begin{array}{c} \text{HCHO} + \text{HCHO} \xrightarrow{\text{KOH(conc.)}} \text{CH}_3\text{OH} & + \text{HCOOK}^+ \\ & \text{methyl alcohol } \text{ acetic acid} \end{array}$$

The mechanism of Cannizzaro reaction is as Step I Attack of nucleophile OH- to the carbonyl carbon

Step II The transfer of hydride ion from anion (1) to second molecule of aldehyde and finally rapid transfer of proton takes place.



$$\begin{array}{c|c} O\\ \\ \\ \\ C \\ \\ acid \end{array} OK$$

737 (a)

Propanal and propanone are functional isomers

738 (a)

It is a fact.

739 (a)

Angle strain, $\alpha = \frac{1}{2} [109^{\circ}28' - \theta]$

In case of cyclopropane,

$$\theta = 60^{\circ}$$

$$\therefore \alpha = \frac{1}{2}(109^{\circ}28' - 60^{\circ}) = 24^{\circ}44'$$

740 (d)

The function of AlCl₃, in Friedel-Craft reaction, is to produce electrophile, which later add to benzene nucleus

745 (c)

The acid exist in cis and trans forms:

$$CH_3$$
 $CHCOOH$
 CH_3
 $CHCOOH$
 CH_3
 $CHCOOH$
 CH_3
 CH_3
 $CHCOOH$
 CH_3
 CH_3

746 (a)

747 (b)

Pyridine is a heterocyclic compound having six

$$CH_{3}-CH_{2}-CH_{2}CI+AlCl_{3}\longrightarrow$$

$$CH_{3}-CH_{2}-CH_{2}^{+}+AlCl_{4}^{-}$$

$$CH_{3}-CH-CH_{3}$$

$$(more stable)$$

$$CH_{3}-CH-CH_{3}$$

$$CH_{3}-CH-CH_{3}$$

741 (c)

Kejldahl's method is used for the estimation of nitrogen. The organic compound is heated with conc. H_2SO_4 in presence of K_2SO_4 (used to elevate boiling point of H_2SO_4) and $CuSO_4$ (used as catalyst) to convert all the nitrogen into $(NH_4)_2SO_4$.

742 (c)

Acetone and methanol have nearly equal boiling point. thus, they are separated by fractional distillation

743 **(b)**

Follow IUPAC rules.

744 (c)

The oxygen atom in phenol has more dominating resonance effect than inductive effect. Increase in charge separation decreases the stability of a resonating structure

Stability of resonating structure in decreasing order will be

$$III < VI \equiv II < I$$

Follow the mechanism of esterification.



membered ring formed with C and N-atoms.

748 (a)

When – OH group of lactic acid is replaced by H, then chiral carbon is lost.

OH H
$$CH_3 - C - COOH \rightarrow CH_3 - C - COOH$$

$$H$$

$$H$$

752 (a)

Ozonolysis of the compound may be given as:

$$H_{3}C-CH=CH-\underbrace{C}_{C}-CH=CH-\underbrace{C}_{C}-CH=CH-CH_{3}$$

$$\downarrow Ozonolysis (O_{3}, Zn/H_{2}O)$$

$$CH_{3}$$

$$\downarrow Ozonolysis (O_{3}, Zn/H_{2}O)$$

$$CH_{3}$$

$$\downarrow H_{3}C-CH=O+O=HC-\underbrace{C}_{C}-CH=O+H_{3}C$$

$$(achiral)$$

$$(achiral)$$

$$(achiral)$$

753 (b)

2-aminopentane and 3-aminopentane; Position is different.

755 (c)

% of H =
$$\frac{2}{18} \times \frac{\text{weight of H}_2\text{O}}{\text{weight of organic compound}} \times 100$$

$$= \frac{2}{18} \times \frac{0.9}{0.5} \times 100 = 20\%$$

 \therefore The percentage of carbon = 100 - 20 = 80 %

756 (b)

o- and p-directing groups facilitate S_E reactions whereas m-directing groups deactivate benzene ring for S_E reactions.

757 (a)

(+) and (-) tartaric acid does not possess any element of symmetry.

758 **(b)**

A molecule having doubly bonded carbon atoms shows geometrical isomerism only if both the doubly bonded carbon have altogether different group, i.e., $_{ba}C \equiv C_{ab}$ or $_{ab}C = C_{ac}$ or $_{dc}C = C_{ab}$.

759 (c)

The chemical formula of thiourea is $\rm NH_2CSNH_2$ so here $\rm Na_2S$, NaCN and NaCNS will be formed but not $\rm Na_2SO_4$

760 (a)

lactic acid

:Its optical activity is lost.

750 (a)

2-butene exhibit rotamers. Rotamers are the isomers formed by restricted rotation.

751 (d)

It contains lone pair electron on N atom.

A white precipitate with am. $AgNO_3$ confirms the presence of terminal alkyne.

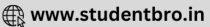
761 (a)

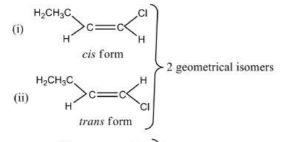
Racemisation involves change in entropy, *i.e.*, change in arrangement of groups position leading to a change in entropy of disorderness.

762 (a)

The acyclic stereoisomers of C₄H₇Cl are







(i)
$$CH_2CI$$
 $C = CC$
 CH_3
 CH_2CI
 CH_3
 CH_2CI
 CH_3
 CH_2CI
 CH_3
 CH_3

(i)
$$CI > C = C < CH_3$$
trans form

(ii) $CI > C = C < CH_3$
 $CI > C < CH_3$
 CI

Number of optical isomers $=2^n=2^1=2$ Hence, total number of geometrical isomers=6 Total number of optical isomers =2.

763 (c)

$$\bigcirc$$

bicyclo (4, 1, 0) heptane

This compound contains 7 carbon atoms, so the corresponding alkane is heptane. Two bridges contain 4 and 1 carbon atom respectively and one bridge does not contain any carbon atom. So the name of the compound is bicyclo (4,1,0) heptane.

764 (d)

Stability of alkyl free radicals can be explained by hyperconjugation and number of resonating structure due to the hyperconjugation. The decreasing order of stability of alkyl free radical is as follows

3° free radical > 2° free radical > 1° free radical > CH₃

766 (b)

Inductive effect involves only displacement (and not delocalisation) of σ –electrons.

767 (c)

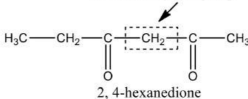
Meso forms are optically inactive as they are superimposable to their mirror images.

768 (b)

CH4 has highest ratio of H to C

769 (b)

active methylene group



When methylene group $(-CH_2)$ is attached with two electron withdrawing groups (like, -CHO, > C = O, -COOH, -CN, -X, etc), its acidity will increase due to -I effect of the electron withdrawing groups.

770 (a)

Follow IUPAC rules.

771 (c)

The reactivity order for H atom is $3^{\circ} > 2^{\circ} > 1^{\circ}$; Neocarbon does not have H atom.

773 **(b)**

-do-

774 (a)

Organic compound which are volatile in steam can be purified by steam distillation. It is based on the fact that vaporisation of organic liquid takes place at lower temperature than its boiling point

775 (b)

Follow IUPAC rules.

777 (d)

CH₃Ō is nucleophile;

$$CH_3OH + Na \longrightarrow CH_3O \overset{+}{N}a + (1/2)H_2$$

778 (a)

Inductive effect is the permanent effect on σ —electrons. It involve the electron displacement along the chain of saturated carbon atoms due to the presence of a polar covalent bond at one end of the chain.

779 (c)

Homologous differ by a group — CH₂ and cannot be isomer.

780 (c)

The reagent selected should be such that only one of components to be separated, reacts with it.

Aniline+aq. HCl→salt, which is water soluble

Nitrobenzene +aq. HCl→no reaction



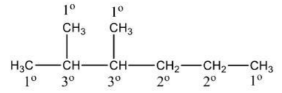
: aq. HCl is used to separate aniline and nitrobenzene.

781 (b)

Formic acid was obtained from ant (fromica in greek). This is trivial name for HCOOH.

782 (a)

The structure of 2, 3-dimethyl hexane is



So, the number of tertiary carbon atoms=2 The number of secondary carbon atoms=2 The number of primary carbon atoms=4

783 (a)

Follow IUPAC rules.

786 (c)

CH₃⁺ has planar structure.

787 (d)

These are characteristics of carbanion.

Follow Saytzeff rule for elimination. 3-halopentane will give only pentene-2.

789 (b)

Atom	Atomic	Percentage	$\frac{b}{a} = x$	
Ratio				
	Mass (a)	(\boldsymbol{b})		
C	12	10.06	$\frac{10.06}{12}$	1
Н	1	0.84	0.84	1
Cl	35.5	89.10	89.10 35.5	3

Empirical formula = CHCl₃

Empirical formula mass = 12 + 1 + 106.5 =

 $119.5 \approx 120$

Molecular mass = $2 \times V$. D = $2 \times 60 = 120$

$$n = \frac{\text{molar mass}}{\text{empirical formula mass}}$$
$$= \frac{120}{120} = 1$$

 $Molecular formula = (CHCl_3)_1 = CHCl_3$

790 (d)

During nucleophilic substitution weaker nucleophile is replaced by stronger nucleophile. The compound having C-Cl bond which can be most easily broken will be most reactive towards nuclophilic substitution reaction.

In vinyl chloride $CH_2 = CH - Cl$ and chlorobenzene C₆H₅Cl the C − Cl bond has partial double bond character due to resonance.

: They do not give nucleophilic substitution reaction easily

$$CH_2 \xrightarrow{\frown} CH \xrightarrow{\frown} CH_2 \xrightarrow$$

Benzyl chloride, give nucleophilic substitution easily because they carbocation formed is stabilised due to resonance.

CH₂=CH—CH₂CI
$$\xrightarrow{\text{-Cl}^-}$$
 CH₂ $\xrightarrow{\text{-Cl}^-}$ CH₂ allyl chloride

$$\xrightarrow{\text{-CH}_2}$$
 CH=CH₂ CH=CH₂ carbocation

$$\xrightarrow{\text{OH}^-}$$
 HOCH₂—CH=CH₂

791 (a)

Enantiomers are non-superimposable mirror images, e.g, lactic acid

Diastereomers are non-superimposable and are not the mirror images of each other. Moreover, meso form has plane of symmetry.

792 (b)

Nucleophilic strength increases down a column of the Periodic Table (in solvents that can have hydrogen bonds, such as water, alcohols, thio alcohols).

Nucleophilic strength $RO^- < RS^ RO^{\Theta} > RS^{-}$ Base strength

Thus, RO^{Θ} is more nucleophilic but less basic than RO-

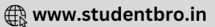
795 (a)

We know that there are seven isomers in $C_4H_{10}O$. Out of these seven isomers, four are of alcohol and three are of ether.

796 (a)

Tertiary halide always favours S_N1 mechanism (as they give comparatively stabler carbocation) white primary halide favours S_N2 mechanism.





Electron donors are bases. Since, electron density is highest at



(Piperidine), hence, it is most basic.

798 (d)

Follow IUPAC rules.

800 (c)

To be optically active, compound or structure should posses a chiral or asymmetric carbon atom. 1-chloropentane is not chiral.

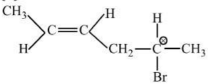
801 (c)

Stearic hinderance in tertiary halides give rise to less reactivity for S_N2.

802 (b)

Addition of Br2 gives altogether different products units cis and trans butene-2.

803 (d)



It has one chiral centre (two enantiomer) and two geometrical isomers

cis-d, trans-d, cis- and trans-l.

804 (d)

Glucose contains four chiral carbon atoms hence number of possible optical isomers are $2^4 = 16$.

805 (c)

Markownikoff's rule is for addition of unsymmetrical additive on unsymmetrical alkene.

806 (d)

Presence of halogen in organic compound can be detected by Beilstein's test.

The bond energy of catenation order is C > Si > S > P.

808 (d)

$$R - X \xrightarrow{\text{NaOH}} R - \text{OH} + \text{Na}X$$

 $R - X \xrightarrow{\text{OH}^-} R - \text{OH} + X^-$. This is nucleophilic substitution.

809 (c)

2-methyl butanoic acid exhibits stereo isomerism.

$$CH_3$$
 $|$ $CH_3 - CH_2 - CH - COOH$

It shows optical isomerism because it contains asymmetric carbon atom.

810 (a)

— CH₃ is electron repelling group.

812 (c)

Metamerism is found in molecules having polyvalent functional group.

813 (d)

There are IUPAC rules.

814 (c)

 $a = 2^n$; where n is no. of dissimilar asymmetric carbon atoms and a is no. of optically active

816 (a)

Follow IUPAC rules.

817 (d)

Nucleophile (-NH₃) replaces other nucleophile (-Br) in the reaction.

818 (a)

Meso form is optically inactive.

819 (d)

Formation of ethylene from acetylene is an example of addition reaction

$$\begin{array}{ccc} \text{CH} & \text{CH}_2 \\ ||| & +\text{H}_2 \xrightarrow{\text{Ni}} & || \\ \text{CH} & \text{CH}_2 \\ \text{Ethyne} & \text{ethene} \end{array}$$

820 (b)

Sodium hydrogen sulphite adds to aldehydes and ketones to form crystalline bisulphite addition products. The product is water soluble and can be converted back to the original carbonyl compound by treating it with dilute mineral acid or alkali. Therefore, these are useful for separation and purification of aldehydes like acetaldehydes.

821 (b)

Zn dust is used for dehalogenation,

$$CH_2X.CH_2X \xrightarrow{Zn \text{ dust}} CH_2 = CH_2.$$

823 (d)

Resonance in a molecule is arised due to delocalisation of π -electrons.

824 (d)

$$CH_3$$
- CH_2 - CH = CH
 sp^3
 sp^3
 sp^2
 sp

Electronegativity of different hybrid and unhybrid orbitals in decreasing order is as follows $s > sp > sp^2 > sp^3 > p$



825 (b)

$$\begin{array}{ccc} \text{CH}_2\text{Br} & \text{CH} \\ | & + & 2\text{KOH} \xrightarrow{\Delta} & ||| & + 2\text{KBr} + 2\text{H}_2\text{O} \\ \text{CH}_2\text{Br} & \text{CH} \end{array}$$

ethylene dibromide acetylene

This is a dehydrohalogenation reaction.

826 (d)

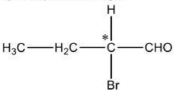
Stereoisomerism is of two types, geometrical and

827 (c)

Follow IUPAC rules.

828 (c)

Compounds having asymmetric C-atom is optically active, e.g.,



The C-atom whose four valencies are satisfied by four different groups is asymmetric C-atom.

829 (b)

Chlorine of vinyl chloride (CH₂ = CHCl) is nonreactive (less reactive) towards nucleophile in nucleophilic substitution reaction because it shows the following resonating structure due to +M effect of - Cl atom.

In structure II, Cl-atom have positive charge and partial double bond character with C of vinyl group, so it is more tightly attracted towards the nucleus and it does not get replaced by nucleophile in S_N - reaction.

830 (d)

Follow mechanism of debromination.

831 (c)

Atom	Atomic	Percentage	$\frac{b}{a} = x$
Ratio			
	Mass (a)	(\boldsymbol{b})	
C	12	40	$\frac{40}{12} = 3.33$
1			
H	1	6.66	$\frac{6.66}{1} = 6.66$
2			
0	16	53.34	$\frac{53.34}{16} = 3.33$
1			9778

832 (d)

Grignard reagents can act as electrophile and nucleophile.

833 (b)

Both these carbon atoms have 3σ -and 1π -bond. Recall hybridized orbitals never from π -bonds.

834 (c)

 S_N 1 mechanism involves the formation of carbocation intermediate. Hence, the species which gives the most stable carbocation readily undergoes S_N1 mechanism. t-butyl bromide gives the most stable carbocation, i. e., 3° carbocation, so it readily undergoes S_N1 reaction.

835 (b)

Follow IUPAC rules.

836 (b)

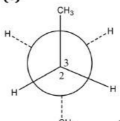
In the Lassaigne's test, a blue colour is obtained if the organic compound contains nitrogen. The blue colour is due to ferri-ferro cyanide i.e., $Fe_4[Fe(CN)_6]_3$.

837 (d)

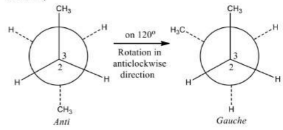
According to Cahn-Ingold-Prelog sequence rules, the priority of groups is decided by the atomic number of their atoms. When the atom (which is directly attached to the asymmetric carbon atom) of a group has higher atomic number, then the group gets higher priority. Groups which atoms of comparable atomic number having double or triple bond, have high priority than those have single bond.

Hence, the order of priority of group is $-OH > -COOH > -CHO > -CH_2OH$

838 (c)



Here, when C_2 is rotated anticlockwise 120° about $C_2 - C_3$ bond the resulting conformer is Gauche conformer. Hence,





CLICK HERE

Hence, empirical formula = CH_2O

carbon, thus optically active.

840 (c)

2-bromo 3-chloro butane

- ∴ Number of asymmetric carbon atoms=2
- \therefore Number of chiral isomers = $2^n = 2^2 = 4$

841 (c)

Glycerol can be separated from spent lye in soap industry by the distillation under reduced pressure because it decomposes near its boiling point

843 (b)

In gas phase tertiary amines are more basic than secondary amines which are more basic than ammonia

−*l* group present on central atom decreases electron density, hence decreases basicity $CH_3NH_2 > NH_3 > NF_3$

844 (a)

Atom	At mass (a)	% (b)	$\frac{b}{a}$
Ratio			u
C	12	49.3	$\frac{49.3}{12} = 4.10$
2			
H	1	6.84	$\frac{6.84}{1} = 6.84$
3			
0	16	43.86	$\frac{43.86}{16} = 2.74$
1			

Hence, empirical formula = (C_2H_3O) Molecular mass = $2 \times VD = 2 \times 73$

$$n = \frac{\text{molar mass}}{\text{empirical formula mass}} = \frac{146}{43} \approx 3$$

So, formula = $(C_2H_3O)_3 \approx C_6H_9O_3$

845 (c)

Wöhler prepared urea from inorganic compounds 861 (b) and rejected the vital force theory that organic compounds can only be synthesised from living organisms.

846 (c)

Follow mechanism of addition of HCl and HI in presence of peroxide. One of the chain propagation step is endothermic in both cases.

847 (c)

All aromatic compounds are resonance hybrid.

848 (a)

It is the stability order for various conformers.

849 (c)

Glucose has aldehyde group and fructose keto group. The general formula for both is C₆H₁₂O₆.

851 (b)

Follow conformation.

852 (b)

In o-, m-, p- derivatives vectors are at 60° , 120° and 180°. Thus, para has zero dipole moment. Also ortho form has more dipole moment than meta form.

853 (c)

The staggered form has lower energy than eclipsed form because of repulsive interaction between the H-atoms attached to two carbon atoms are minimum due to maximum distance between them.

854 (c)

Victor Mayer's method is applicable only for the determination of molecular mass of volatile substance

856 (d)

Hexane is non-polar molecule.

857 (c)

Nucleophilies may be neutral or negatively charged, whereas substrate undergoing nucleophilic substitution may be neutral or positively charged

 $C_2H_5 - I + OH^- \rightarrow C_2H_5OH + I^-$

858 (a)

Nucleophilicity increases on going down in the group of the Periodic Table

$$I^{\Theta} > Br^{\Theta} > Cl^{\Theta} > F^{\Theta}$$

859 (d)

Free radicals have unpaired electrons, but are neutrals and are reactive.

$$\dot{\text{CH}}_3 + \dot{\text{CH}}_3 \longrightarrow \text{CH}_3 \longrightarrow \text{CH}_3$$

Follow mechanism of Kharasch effect.

862 (d)

-do-

864 (c)



Reaction of NaOH with dinitrofluorobenzene represents nucleophilic aromatic substitution reaction because - NO₂ group is deactivating group. They make benzene nucleus electron deficient and facilitate the nucleophile to attack the ring.

$$O_2N$$
 O_2N
 O_2N
 O_2N
 O_3N
 O_4N
 O_4N

865 (b)

Cyano group has the highest priority therefore, parent name must be benzonitrile. Br occurs at 2position, and hydroxyl at 3-position, hence the IUPAC name is 2-bromo-5-hydroxy benzonitrile.

866 (d) Ethers show metamerism.

867 (b)

Due to resonance; the carbonyl group of benzoic acid is coplanar with the ring. If the electron withdrawing substituent (i.e., -I showing) is present at ortho position, it prevents the coplanarity and thus, the resonance. Hence, makes the acid more stronger.

Thus, among the given acids, ortho hydroxy benzene acid is the most acidic.

868 (a)

Diamond (sp^3) , Graphite (sp^2) , Acetylene (sp).

869 (d)

CH3CHClCOOH contains asymmetric carbon atom.

870 (d)

Statement (c) is wrong.

871 **(b)**

The -ve inductive effect of -CHO group play role to give anti Markownikoff's addition.

$$CH_2$$
 CH \rightarrow CHO $\rightarrow CHO$ $\rightarrow CHO$.

872 (c)

The structure of 1-chloro-2-nitroethene is as

$$C = C$$

In this compound E-Zisomerism is possible because it is highly substituted alkene. The E-Zsystem of nomenclature is developed by Cahn, Inglod and Prelog.

873 (a)

 $CH_3C = N$ is known as acetonitrile or methyl cyanide.

874 (b)

Ketone undergoes nucleophilic addition reaction because nucleophilic end of reagent attack first followed by electrophilic end of reagent.

$$X \longrightarrow C \longrightarrow R + CN^{-} \xrightarrow{\text{Slow}} R \longrightarrow C \longrightarrow R \xrightarrow{\text{Fast}} R \longrightarrow C \longrightarrow R$$

$$CN$$

$$CN$$

$$Extone$$

$$Cyanohydrin$$

875 (d)

Halogen containing compounds (C₆H₅Cl) When placed in a flame, the presence of halogen is revealed by a green to blue flame.

876 (a)

Two similar asymmetric carbon atoms; $\alpha =$ 2^{n-1} . Also meso form $=2^{\frac{n}{2}-1}$. Total = a + m.

877 (d)

In C_6H_5 ring there are three π -bonds and one π -

OH bond is present in group. Therefore, in all there are four π -bonds in C_6H_5COOH . In $CH_3CH_2COCH_3$ there is only one π bond in C = O group, in $CH_2 = CH - CH =$ CH_2 there are two π -bonds while in $HC \equiv C CH = CH_2$ there are three π - bonds

878 (c)

Order of bond length σ bond $(sp^3) > \sigma$ bond $(sp^2) > \sigma$ bond (sp)

879 (d)

It is a reason for the given fact.

880 (c)

The octet of all atoms are complete in structures a and b. The molecule in which all the atoms have completed octet is more stable than atom which have incomplete octet. Larger the number of resonating structures, larger will be the stability, thus structures a and b are stable.

In structure (d), the electron deficient of positive charged carbon is duly compensated by one pair electrons of adjacent oxygen atoms while such neighbour group support is not available in



structure (c). Hence, structure (c) is least stable in comparison to structure (d).

883 (a)

CH3CH2Cl; CH3CHCl2; CH2ClCH2Cl; CH3CCl3; CH2 CHCl2CCl3; CCl3CCl3

884 (d)

(1) and (3) are enantiomeric forms to each other.

885 (d)

Methoxy group, due to +I effect, increase electron density on OH- group, thus making it less acidic. Thus, o-methoxy phenol and acetylene are less than phenol.

p-nitrophenol is more acidic than phenol.

886 (c)

When organic compound is fused with sodium metal, nitrogen of the compound is converted into sodium cyanide as

887 (c)

It is structure of furan, a heterocyclic compound.

Diazonium salts are highly reactive. In Sandmeyer 900 (a) reaction diazo group is replaced by chlorine or bromine in presence of CuCl or CuBr.(Substitution 901 (c)

$${\rm C_6H_5N_2Cl} \stackrel{\Theta}{\longrightarrow} {\rm CuCl} \blacktriangleright {\rm C_6H_5Cl} + {\rm N_2}$$

X⁻is replaced by OH⁻.

There are four structural isomers are possible for C₄H₉Cl

(a)CH3CH2CH2CH2CI

$$(b)CH_3 - CH_2 - CH - CH_3$$

Cl

(c)
$$CH_3 - CH - CH_2CI$$

|

 CH_3

|

 CI

CH₃

$$(d)CH_3 - C - CH_3$$

891 (b)

A carbanion or carboanion has -ve charge on it.

894 (c)

The case with which a nucleophile attacks the carbonyl groups depends upon the electrondeficiency, i.e., magnitude of the positive charge on the carbonyl carbon. Since, an alkyl groups has electron-donating inductive effect.

(+I effect), therefore, greater the number of alkyl groups attached to the carbonyl groups greater is the electron-density on the carbonyl carbon and hence, lower is its reactivity towards nucleophilic addition reactions.

$$R > C = O > R > C = O$$

896 (c)

n-pentane and isopentane or 2-methylbutane are chain isomers since both have different hydrocarbon chain.

897 (a)

$$CH_2 = CH - CHO$$

Prop -2-en-1-al

898 (d)

Free radicals have unpaired electrons but are neutrals and are reactive.

The second carbon is asymmetric.

Both have different mode of linkage, i. e., chain and ring.

902 (c)

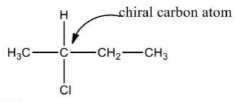
Carbanions contain even number of valence electrons and thus, show diamagnetic behaviour.

903 (d)

Molecules with two similar groups attached on either of the doubly bonded carbon do not show geometrical isomerism.

904 (c)

Compound $CH_3 - CHCl - CH_2 - CH_3$ shows optical isomerism due to the presence of chiral carbon atom.



905 (c)

The Kolbe's electrolysis proceeds via free radical mechanism. For example, when sodium propionate is electrolysed, n —butane, ethane, ethylene are obtained. The propionate ion discharge at the anode to form free radicals. $C_2H_5COO^- \rightarrow C_2H_5COO^{\bullet} + e^ C_2H_5COO^{\bullet} \rightarrow C_2H_5^{\bullet} + CO_2$



$$\begin{split} 2C_2H_5^{\scriptscriptstyle\bullet} &\to C_4H_{10} \\ C_2H_5^{\scriptscriptstyle\bullet} &+ C_2H_5^{\scriptscriptstyle\bullet} &\to C_2H_4 + C_2H_6 \end{split}$$

In TLC, adsorbent is made of silica gel or alumina

908 (d)

are soluble in aq. NaOH. Benzylic alcohol is less acidic than water so not soluble in aq. NaOH,

909 (c)

Lead unreached 0.1 M (=0.2 N) $H_2SO_4 = V \text{ mL}$ 20 Ml of 0.5 NaOH=V mL of 0.2 N H₂SO₄

$$20 \times 0.5 = V \times 0.2$$

$$V = \frac{20 \times 0.5}{0.2} = 50 \text{ mL}$$

Used H₂SO₄ = 100 - 50 = 50 mL
% of N =
$$\frac{1.4 \text{ NV}}{w}$$
 = $\frac{1.4 \times 0.2 \times 50}{0.30}$

= 46.67 %

% of nitrogen in

(a)
$$CH_3CONH_2 = \frac{14 \times 100}{59} = 23.73 \%$$

(a)
$$CH_3CONH_2 = \frac{14 \times 100}{59} = 23.73 \%$$

(b) $C_6H_5CONH_2 = \frac{14 \times 100}{121} = 11.57 \%$
(c) $NH_2CONH_2 = \frac{28 \times 100}{60} = 46.67 \%$

(c)
$$NH_2CONH_2 = \frac{28 \times 100}{60} = 46.67 \%$$

(d)
$$NH_2CSNH_2 = \frac{60}{76} = 36.84 \%$$

910 (d)

Removal of H from alkane, alkene and alkyne gives alkyl, alkenyl, alkynyl groups respectively.

There are two symmetrical hexenes as given in (a) and (b).

913 (d)

The solution of D(+)-2-chloro-2-phenyl ethane in toluene racemises slowly in the presence of SbCl₅ due to the formation in carbocation

914 (a)

Tautomerism it is functional isomerism in which the isomers are readily interchangeable and maintain a dynamic equilibrium with each other.

915 (b)

If acid is weak, its conjugate base (nucleophile) is strong and vice versa.

O
$$\parallel$$
 (A)CH₃ – C – O $^-$ is conjugated base of CH₃COH

- (B) CH₃O⁻ is a conjugate base of CH₃OH (II)
- (C) CN⁻ is a conjugate base of HCN (III)

$$H_3$$
C $SO_3^ SO_3^-$ is a conjugate base

Acidic nature of IV>I>III>II and nucleophilicity of B>C>A>D.

916 (d)

S_N2 reaction proceeds with inversion and a transition state is formed which does not carry any charge

918 (b)

The most stable one is that in which the positive and negative charges reside on the most electropositive and most electronegative atoms of the species respectively. Like

919 (b)

Organic compound
$$\xrightarrow{[o]} CO_2 + H_2O$$
17.6 g 7.2g

% of
$$C = \frac{12}{44} \times \frac{17.6}{5.6} \times 100 = 85.7\%$$

% of $H = \frac{2}{18} \times \frac{7.2}{5.6} \times 100 = 14.28\%$

Ele	Percen	Relative no.	Simplest ratio
me	tage	of atoms	F
nt			



С	85.7	85.7/12=7.1 4	7.14/7.14=1
Н	14.28	14.28/1=14. 28	14.28/7.14= 2

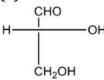
Hence, empirical formula of compound of = CH_2 \therefore Molecular formula of compound = C_4H_8

920 (c)

Stronger is acid, weaker is its conjugate base. Acidic nature : CH \equiv CH > CH $_2$ = CH $_2$ > CH $_3$ — CH $_3$

Conjugate basic : CH \equiv C^- < CH $_2$ = CH^- < CH $_3$ — CH $_2^-$ nature.

921 (b)



D-glyceraldehyde has the above formula. So, the Fischer's projection formula which is identical to it is

922 (b)

Percentage of Nin an organic

Compound =
$$\frac{1.4 \times N \times V}{W}$$
$$= \frac{1.4 \times 1 \times 30}{1.2} = 35$$

923 (c)

Dissociation of proton from ddd is very difficult due to -I effect of Cl^- and N^+ while in C_6H_5OH due to the reasonance stabilization of phenoxide ion proton eliminates easily. Similarly due to H-bonding in $C_6H_5CH_2OH$ it can be eliminate easily and in $CH_3C \equiv CH$ the proton is acidic in nature hence, it can be dissociated

924 (d)

% of S =
$$\frac{32}{233} \times \frac{\text{wt. of BaSO}_4}{\text{wt. of organic compound}} \times 100$$

= $\frac{32}{233} \times \frac{1.158}{0.53} \times 100$
= 30%

925 (c)

Tautomerism is a dynamic isomerism because two forms (keto and enol) of substance cannot be separated they are in dynamic equilibrium with each other.

926 (a)

Enantiomers are non-superimposable mirror images. e.g., lactic acid

927 (a)

Vinyl carbocations are more stable than primary carbocation but less stable than secondary carbocation.

928 (b)

Duma's method involve the determination of nitrogen content in the organic compound in the form of N_2

$$\begin{split} N_2O + Cu &\rightarrow N_2 + Cu \\ d\% \text{ of } N = \frac{28}{22400} \times \frac{\text{volume of N}_2 \text{at NTP}}{\text{weight of compound}} \times 100 \end{split}$$

930 (a)

The correct order of reactivity is

It is due to fact, that weaker the base, better it will be the leaving group.

Hence, I⁻ is the best leaving group.

931 (b)

$$R_3$$
N — CH — CH₂ HBr

Due to R_3^{Θ} (e^- withdrawing tendency) carbocation will appear farther to that (terminal). Hence, product is $R_3N - CH_2 - CH_2Br$.

932 (d)

n-pentane and 2-methyl butane are constitutional isomers or chain isomers or skeletal isomers.

934 (a)

Follow IUPAC rules.

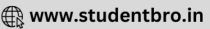
935 (b)

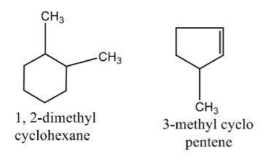
In toluene $(-CH_3)$ group is present which has +I effect and increases electron density on ortho and para position. While in sulphonation $-SO_3$ acts as electrophile. Therefore, it $(-SO_3H)$ attacks on ortho and para position readily.

936 (d)

Follow elimination rules.

937 (a)





contain two, one, one asymmetric carbon atoms respectively.

939 (a)

: Total of five isomers are possible by formula C7H9N.

941 (a)

Such a condition is seen when π bond is formed between similar atoms ie,



943 (b)

From kjeldahl's meuros,
Percent of nitrogen = $\frac{1.4 \times N \times V}{W}$ = $\frac{1.4 \times 0.5 \times 2 \times 10}{0.25}$ = 56%

944 (d)

It is the definition of distereoisomers.

945 (b)

The central carbon is attached to four different substituents, hence it is chiral, therefore optically

946 (b)

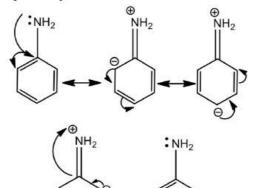
$$HNO_2 + H_2SO_4 \rightarrow NO^+ + HSO_4^- + H_2O$$
Nitrosonium ion

947 (b)

−Cl is an electron withdrawing (i.e., −I showing) group. It withdraws electrons when attached to the carboxylic acid and decreases the electron density on the oxygen atom. This will facilitate the release of H+ by making O - H bond more polar and thus - Cl increases the acidity of acetic acid when attached at, α position because of – *I* effect.

948 (a)

-NH2 has +R effect, it donates electrons to the benzene ring. As a result, the lone pair of electron on the N-atom gets delocalized over the benzene ring. As a result, the lone pair of electron on the Natom gets delocalized over the benzene ring and thus it is less readily available for protonation. Hence, aniline is a weaker base than cyclohexylamine.



Resonance in aniline

949 (c)

This is annulene.

950 (a)

Alcohols undergo dehydration usually by E1 mechanism. This is because elimination is preferred in case of tertiary alcohols, e.g.,

$$\xrightarrow[\text{-H}_2\text{O}]{\text{H}_3\text{C}} \text{H}_3\text{C}$$



Acetophenone oxime can show geometrical isomerism.

952 (d)

Ethers show functional isomerism with molecular formula $C_nH_{2n+2}O$. For example, C_2H_6O CH₃OCH₃, CH₃CH₂OH dimethyl ether, ethyl alcohol

953 (c)

The alkyl halide which produce stable carbonium ion undergo $S_N 1$ reaction. Tertiary alkyl halide is most stable. Thus, decreasing order of tendency of alkyl halides undergoes $S_N 1$ mechanism. t-alkyl halide>sec-alkyl halide>primary-alkyl halide

954 (c)

Vinyl chloride(CH₂ = CHCl) undergoes addition and elimination reactions. Substitutions reaction is shown by compounds having single bonds only.

955 (a)

Fisher projections are for illustration of optical isomers.

956 (b)

A hydrogen halide contain a highly polar H − ⊕

X bond can easily lose $^{\rm H}$ to the pi bond of an alkene. The result of the attack of ${\rm H}^{\oplus}$ is an intermediate carbocation, which quickly undergoes reaction with the negative halide ion (X^-) to yield an alkyl halide

960 (c)

Aldehydes and ketones readily undergo nucleophilic addition reaction. The order of reactivity, is as the +I effect of alkyl group increases

$$H > C = O > CH_3 > C = O > CH_3 > C = O$$

961 (b)

The structures of maleic and fumaric acids are given below

The structures of fumaric and maleic acid suggest that they are geometrical isomers because they have same molecular formula but different spatial arrangement of atoms around a double bond.

962 (b)

Stability order of arenium ion

963 (d)

In condensed structure formula, paranthesis is used for identical group of atoms.

964 (d)

Compound which sublime on heating can be purified by sublimation method.

Benzoic acid, camphor and naphthalene sublime on heating hence, they are purified by sublimation method.

965 (d)

CH₂ = CH. CH₂Cl compound undergoes nucleophilic substitution most readily.

966 **(a**

As the – I group increases at the α -carbon, acidity increases

967 (b)

In $S_N 2$ reactions, the nucleophile attaches itself from the direction opposite to that of the nucleophile already present in the second step, the previous nucleophile is removed and a single stereoisomer is obtained

968 (b)

Follow IUPAC rules.

$${}^{5}_{\text{CH}_{3}} \cdot {}^{4}_{\text{CH}} = {}^{3}_{\text{CH}} \cdot {}^{2}_{\text{CH}} = {}^{1}_{\text{CH}}$$

969 (d)

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

Since, the reaction rate depends upon the concentration of both reactant and nucleophile, it



is a $S_N 2$ reaction. It involves inversion of configuration.

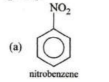
970 (c)

The given compound is nitrobenzene.

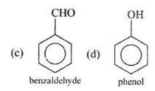
Both differs by a $-CH_2$ group.

972 (d)

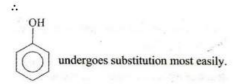
During electrophilic substitution electrophile attacks the double bond of benzene ring. The aromatic compounds having electron donating groups undergo electrophilic substitution more easily due to favouable effect of electron donating group.







- (i) NO₂, COOH and CHO groups are electron withdrawing groups so, they decrease the reactivity of organic compounds.
- (ii) OHgroup is electron donating group, so it increases the electron density in benzene ring ad increases the rate of reaction



973 (b)

Oxygen cannot be detected by direct test because oxygen is present in atmosphere and all tests are carried in atmosphere of oxygen

974 (a)

Follow IUPAC rules.

975 (a)

o-, m-, p — isomers are position isomers.

976 **(b)**

Carbinol is trivial name for HCH2OH. Thus, C₆H₅CH₂OH is phenyl carbinol and chloral is CCl₃CHO.

977 (a)

A primary carbon is one which is attached with one carbon atom.

978 (a)

Follow IUPAC rules.

979 (c)

1,1-dibromoethane and 1,2-dibromoethane.

980 (a)

A solvent molecule lacking a polar X-H bond is called aprotic solvent. NH3, SO2 and CH3CN are aprotic solvent while CH3COOH is protic solvent.

981 (b)

When sodium ethoxide reacts with iodoethane, diethyl ether is obtained (Williamson's synthesis) The mechanism of this reaction is as follows $C_2H_5ONa \rightleftharpoons C_2H_5O^- + Na^+$

$$C_2H_5O^- + C_2H_5 - I \frac{Slow}{C_2H_5 - O - C_2H_5 - I}$$
 transition state

Fast
$$C_2H_5OC_2H_5 + \Gamma$$

Since, the reaction involves substitution of a group by a nucleophile, it is an example of nucleophilic substitution reaction.

982 (c)

The organic compounds is fused with Na metal, Na2S is formed which is tested as It reacts with lead acetate and forms black ppt. of

$$Na_2S + (CH_3COO)_2Pb \rightarrow PbS + 2CH_3COONa$$

black ppt

983 (c)

984 (c)

$$\frac{E}{108} = \frac{100}{60}$$

Eq. wt of the silver salt $E = 108 \times \frac{100}{60} = 180$: Eq. wt. of the acid = E - 108 + 1 = 73

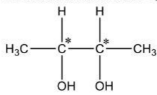
985 (a)

$$\begin{array}{c} 2 & 1 \\ {\rm C_2H_5} - {\rm C} - {\rm CH_2OH} \\ 3 \parallel \\ {\rm CH_2} \end{array}$$

2-ethylprop-2-en-1-ol

986 (d)

The structure of butane-2, 3-idol is as





: Optical isomers in compounds have similar asymmetric carbon atom, which are even in number = $2^n - 1$

$$n=2$$

: Total number of optically active stereoisomers $= 2^2 - 1 = 3$

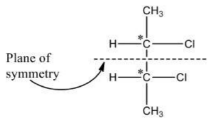
987 (b)

$$CH_4 \rightarrow CH_3^+ + H^-;$$

CH₃⁺ is methyl carbonium.

One asymmetric carbon atom, forms d, and l -optical and isomers.

(b) Two asymmetric carbon atoms, forms d, l and meso forms



Meso due to internal compensation

Two asymmetric carbon atoms but does not have symmetry. Hence, meso form is not formed.

One asymmetric carbon atom, meso form is not formed.

989 (d)

More the number of hyperconjugated structures, more will be electronegative chlorine atom.

 $CH_3 - Cl$ (a)

3-hyperconjugated structures

(b) $CH_3 - CH_2 - Cl$

2-hyperconjugated structures

$$\begin{array}{ccc} \text{(c)} & \text{CH}_3 \\ & \text{|} \\ \text{H} - \text{C} - \text{Cl} \\ & \text{|} \\ \text{CH}_3 \end{array}$$

6- hyperconjugated structures

(d)
$$CH_3 - CH_2 - C - CI$$

 CH_3

8-hyperconjugated structures

- :: 8-hyperconjugated structures are possible for
- :: Chlorine in this is most electronegative.

990 (a)

Follow resonance characteristics.

991 (b)

Diastereoisomers are a pair of optical isomers which cannot be related as non-superimposable mirror images of each other.

994 (a)

The rate of reaction is influenced by the hyperconjugation effect of group R. it depends on the electron donating power of alkyl group (R). The electron releasing power of R group depends on the number of hydrogen present on α carbon. The increasing order of speed with R group in the reaction is

$$H_3C$$
 CH_3 CH_3

i.e., IV<III<II<I

995 (b)

The given compound form two geometrical isomers and two optical isomers.

996 **(b)**

Geometrical isomerism is shown by >C=C< only when identical groups are not present on the double bonded carbon atoms.

999 (b)

$$RX + I^- \rightarrow R - I + X^-$$

This reaction is an example of nucleophilic substitution.

100 (c)

0



Its IUPAC name is bicyclo [1,1,0] butane.

100 (a)

Weak base → strong conjugate acid $CH \equiv C^- \rightarrow CH \equiv CH$ weakest base (strongest acid among the given) $CH_3 - CH_2^- \rightarrow CH_3CH_3$ (strongest base) (weakest acid)



100 (d)

With the increasing basicity of the added base, the 2 rates of the elimination reactions have been found to increase. Thus, RO is most reactive

100 (b)

Follow IUPAC rules. 3

100 (a)

Number of isomers in hexane are five as follows

100 (c)

(a) it is Diels Alder's reaction (cyclo addition)

$$>$$
C=O + NaHSO₃ \rightarrow $>$ C $<$ OH $>$ SO₃Na

It is nucleophilic addition reaction

$$CH_3$$
 CH_3
 $C-OH + \frac{Dry HCl/Anhy. ZnCl_2}{Lucas reagent} CH_3$
 CH_3
 CH_3

It is nucleophilic substitution reaction

$$(CH_3)_2C = CH_2 + BrCl \longrightarrow (CH_3)_2-C \longrightarrow CH_2$$
(d)

It is electrophilic addition reaction

100 (d)

(d) is with maximum conjugative structure among them

100 (a)

Based on heat of hydrogenation.

101 (c)

Butyne-2 is $CH_3 - C \equiv C - CH_3$; Two carbon of 101 (c) corner are sp3-hybridized.

101 (c)

Four π -electrons of double bond and 1 lone pair on N atom leads to delocalisation of six electrons.

101 (d)

$$\begin{array}{c} \mathrm{NH_2} \\ | \\ \mathrm{CH_3} - \mathrm{CH} - \mathrm{CH_3} \\ \mathrm{2-propanamine} \end{array}$$

101 (h)

Elem ent	Perce ntage	Percentage at. wt.	Simple ratio
С	20.0	$\frac{20.0}{20.0} = 1.66$	$\frac{1.66}{1.66} = 1$
Н	6.67	$\frac{12}{6.67} = 6.67$	$\frac{1.66}{6.67} = 4$
N	46.67	$\frac{1}{46.67} = 3.33$	$\frac{1.66}{3.33} = 2$
О	26.66	$\frac{26.66}{16} = 1.66$	$\frac{1.66}{1.66} = 1$

Empirical formula = CH_4N_2O

Empirical formula weight

$$= 12 + (4 \times 1) + (2 \times 14) + 16 = 60$$

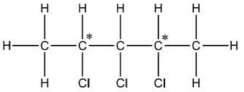
$$n = \frac{\text{mol. formula weight}}{\text{emp. formula weight}}$$
$$= \frac{60}{60} = 1$$

∴ Molecular formula=CH₄N₂O

Given compound gives biuret test. Thus, given compound is urea $(NH_2)_2CO$.

101 (c)

The structure of 2, 3, 4-trichloro pentane is



Hence, two chiral carbon atoms are present in 2, 3, 4-trichloropentane.

The electronegativity of F is maximum and thus, C—F bond is more polar.

101 (a)

1, 1-dichloro-1-pentene does not exhibit geometrical isomerism.

101 (b)

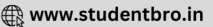
The hydrolysis of sugar solution (dextrorotatory) leads to formation of laevorotatory mixture due to formation of glucose (dextrorotatory) and fructose (laevorotatory).

First two and fourth will show stereoisomerism. 8

101 (d)

9 Geometrical isomers of $CH_3CH = CH - CH_3$ are





$$H_{3C}$$
 $C = C CH_3 H_{3C}$ $C = C CH_3$

6 Ethyl alcohol shows functional isomerism with dimetyl ether.

 $\begin{array}{ll} C_2H_5OH & CH_3-O-CH_3\\ alcohol & ether \end{array}$

102 (a)

- Highest to lowest priority (Br > Cl > CH₃) is clockwise than R.
- 102 (c)

2

Ring I

Ring II

Electrophilic substitution reaction takes place in compounds in which π —electrons are highly delocalised. The electrophile attacks the region of high electron density, therefore, electrophilic substitution occurs at ortho/para position at ring II.

102 (c)

When phenol reacts with chlrtoform and aqueous NaOH solution, it give salicyladehyde.

 $CHCl_3 + OH^- \leftrightharpoons H_2O + CCl_3^-$

$$\begin{array}{c} \overset{\circ}{\text{Ci}} \\ \overset{\circ}{\text{Ci$$

102 (c)

4 It is definition of asymmetric synthesis.

102 (a)

Since, the compound on heating with CuO produced CO_2 , it contains carbon. Again, it does not produce water, hence it does not contain hydrogen. So, the organic compound is carbon tetrachloride (CCl_4).

102 **(d)**

6

If organic compound gives blue or green colour at the tip of red hot copper wire, this indicates the presence of halogens in the compound. However, compounds like urea, thiourea etc also give blue or green colour in this test even in the absence of halogens, thus, it is not a sure test for halogens

102 (a)

8 Mesomeric effect involves complete transfer of π or lone pair of electrons to the adjacent atom or covalent bond. Hence, it involves delocalisation of pi (π) electrons.

103 (b)

0	Species	Valence electrons	Magnetic
	behavior		
	Carbonium ion	6	Diamagnetic
	Free radical	7	
	Paramagnetic		
	Carbene	6	Diamagnetic
	Nitrane	6	Diamagnetic

103 **(b)**

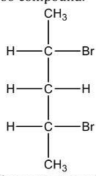
1 It should contain (CH₃)₂ CH – group to be named as iso.

103 (b)

2 One propene and one cyclopropane.

103 (c)

Those compounds which contain two or more asymmetric carbon atoms but are optically inactive due to presence of plane of symmetry, are called *meso* compounds. Meso compounds are optically inactive due to internal compensation. Out of the given compounds only 2, 4-dibromopentane have a plane of symmetry, so it is a meso compound.



2, 4-dibromopentane (meso compound)

103 (a)

4 It is a fact.

103 (a)

6 Electrophiles are the species having a tendency to accept a pair of electron, e. g., NO₂⁺, Br⁺ etc.
Nucleophiles are the species having a tendency to donate a pair of electron. e. g., CH₃OH. N₃⁻



103 (a)

 C_5H_8 has three possible alkynes. These are $CH_3CH_2 - CH_2C \equiv CH, CH_3CH_2C \equiv C - CH_3$ pent-1-yne pent-2-yne CH_3 $CH_3 - CH - C \equiv CH$ 3-methyl but-1-yne

104 (d)

I can have maximum 3 hyperconjugative structures. II has maximum 5 hyperconjugative structure, III has 2 conjugative structure while IV has 1 conjugative structure

104 (a)

 $CH_3C \equiv CCH_2CH_3$; It is always pentyne-2 and 104 (c) never pentyne-3.

104 (d)

The components of an azeotropic mixture are separated by special method, ie, fractional distillation. The simple fraction which distils at 337.8 K is a ternary azeotrope consisting of all water. Some alcohol and benzene

104 (a)

Hence, an equimolar mixture of the enantiomers (dextro or laevo forms) is called racemic mixture. It is represented as dl-form or \pm form and is optically inactive due to external compensation. Separation of racemic mixture into d- and l-forms is called resolution.

104 (a)

5 Two successive homologous differ in their formula by CH2 or have a difference of 14 units in their molecular weight.

104 (c)

6 If a liquid decomposes at or below its boiling point, it is purified by vaccum distillation

CH₃CH₂NH₂ and CH₃NHCH₃.

104 (a)

8 IUPAC name of CH₃CH₂CHO is propan-1-al.

104 (b)

Alkyl group (an electron releasing (+1 group)increases electron density at N-atom, hence, basic nature is increased. In ammonia, no alkyl group is present, so it is least basic.

