

NITROGEN CONTAINING COMPOUNDS (Amines)

A

SINGLE CORRECT CHOICE TYPE

Each of these questions has 4 choices (a), (b), (c) and (d) for its answer, out of which ONLY ONE is correct.

1. Nitrobenzene can be reduced to aniline by

$$H_2/Ni$$

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

2.
$$Cl_{3}N^{+}$$
 $N_{3}Cl_{3}$ $Z; Z is$



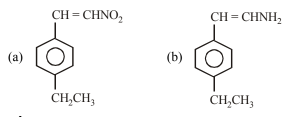
(b)
$$\left\langle \begin{array}{c} N \\ N \end{array} \right\rangle$$

(c)
$$N$$

(d)
$$\bigcap_{N}$$

3.
$$CH = CHNO_2$$

$$\xrightarrow{Zn/Hg} Product. Here the product is COCH3$$



4. Which statement is true regarding the following structure?

$$CH_3$$
 C_2H_5
 C_3H_7

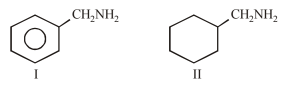
- (a) It is a chiral molecule
- (b) It exists in two resolvable optically active forms
- (c) Both a) and b)
- (d) Neither a) nor b)
- 5. Introduction of a methyl group in ammonia markedly increases the basic strength of ammonia in aq. solution, introduction of the second methyl group increases only marginally the basic strength of methyl amine in water. This is due to
 - (a) different type of hybridisation in the two amines
 - (b) protonated dimethyl amine is more solvated than methyl amine
 - (c) protonated dimethyl amine is more solvated than the protonated methyl amine
 - (d) protonated dimethyl amine is less stable than the protonated methyl amine
- **6.** The basic character of ethyl amine, diethyl amine and triethyl amine in chlorobenzene is
 - (a) $C_2H_5NH_2 < (C_2H_5)_2NH < (C_2H_5)_3N$
 - (b) $C_2H_5NH_2 < (C_2H_5)_3N < (C_2H_5)_2NH$
 - (c) $(C_2H_5)_3N < (C_2H_5)_2NH < C_2H_5NH_2$
 - (d) $(C_2H_5)_3N < C_2H_5NH_2 < (C_2H_5)_2NH$



Mark Your	1. abcd	2. abcd	3. abcd	4. abcd	5. abcd
Response	6. abcd				

- 7. The correct order of decreasing basic character of the three aliphatic primary amines is

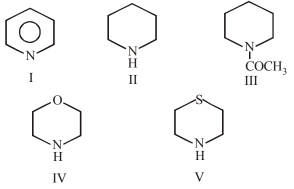
 - (a) I > II > III
- (b) III > II > I
- (c) $I > II \approx III$
- (d) $I = II \equiv III$
- Which of the statement is true regarding the basicity of 8. the following two primary amines?



- (a) Both are equally basic because both are 1° amines
- (b) I > II because it is an aromatic amine
- (c) II > I because it is an aliphatic amine
- (d) I < II because of difference in the nature of β -carbon
- 9. The correct order of decreasing basic character is

- (a) II > I > III > IV
- (b) IV > II > I > III
- (c) IV > III > II > I
- (d) IV > II > III > I
- 10. The correct order of increasing basicity is

- (a) II < III < I
- (b) I ≈ III < II
- (c) I < II < III
- (d) III < I < II
- 11. The relative order of basic character of the following compounds is



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

- 12. When aniline is treated with acetyl chloride in presence of anhydrous aluminium chloride, the main product is
 - (a) o aminoacetophenone (b) p-aminoacetophenone
 - (c) both (a) and (b)
- (d) *m*-aminoacetophenone
- $CH_3CH_2CH_2NH_2 \xrightarrow{NaNO_2, HCl} P$. P is 13.
 - (a) CH₃CH₂CH₂OH
- (b) (CH₃)₂CHCl
- (c) Both (a) and (b)
- (d) Reaction not possible
- 14. Benzenediazonium chloride when treated with phenols gives azo dyes, to get best result the pH of the medium should be
 - (a) around 4
- (b) around 8
- (c) around 10
- (d) 12
- 15. In the following reaction, the reagent X should be

$$RCOOH + [X] \xrightarrow{conc. H_2SO_4} RNH_2$$

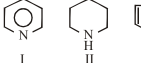
- (a) NH₃
- (b) HN₃
- (c) either of the two
- (d) None of the two
- 16. Which of the following can undergo electrophilic substitution when treated with nitrous acid at 0° C?
 - (a) $C_6H_5NH_2$
- (b) C₆H₅NHCH₃
- (c) $C_6H_5N(CH_3)_2$
- (d) None
- Which of the following does not reduce C₆H₅NO₂ to 17. aniline?
 - (a) Sn/HCl
- (b) SnCl₂/HCl
- (c) Zn/HCl
- (d) LiAlH₄
- 18. Which of the following amines can be resolved into two enantiomers?

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



M. By Voys	7. abcd	8. abcd	9. abcd	10. abcd	11. abcd
Mark Your Response	12. a b c d	13. a b c d	14. abcd	15. abcd	16. abcd
	17. abcd	18. abcd			

- 19. Which of the following statement is false?
 - (a) Dimethyl amine as well as trimethyl amine are soluble in water
 - Trimethyl amine forms hydrogen bond neither with itself nor with water.
 - (c) Trimethyl amine can act as hydrogen bond acceptor only, while dimethyl amine can serve as both a hydrogen bond donor and acceptor
 - (d) All the three statements are false
- 20. Ephedrine is a secondary amine. It is widely used in cold and allergy conditions in the form of its hydrochloride but not as such because
 - (a) the amine itself has unpleasant smell, while its salt is odourless
 - (b) the amine is insoluble in water, while the salt is soluble in water
 - (c) the amine is unstable and easily oxidised by air, while the salt is resistant to atmospheric oxidation
 - (d) of all the above facts
- 21. Benzamide and benzyl amine can be distinguished by
 - (a) cold dil. NaOH
 - (b) cold dil. HCl
 - (c) both a & b
 - (d) NaNO₂, HCl, 0°C, then β-naphthol
- The correct order for the basic character of the compounds 22. I to IV should be

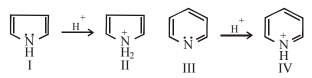


- (a) IV < III < I < II
- (b) IV < I < III < II

IV

- (c) IV < II < III < I
- (d) IV < III < II < I
- 23. Which of the following is true regarding basic character of pyridine and pyrrole?
 - (a) Pyrrole is more basic because its nonbonding electrons occupy sp³ orbital
 - (b) Pyridine is more basic because its nonbonding electrons is not a part of aromatic sextet.
 - (c) Both are equally basic
 - (d) Pyridine is less basic because it is 3° amine

Pyrrole and pyridine both are basic and form salts with 24.



Which of the following statement is true regarding the aromatic character of the four species?

- (a) All the four are aromatic
- (b) I, III and IV are aromatic
- (c) I, II and III are aromatic
- (d) I and III are aromatic
- 25. Ethylene can be prepared in good yield by

(a)
$$CH_3CH_2N^+(CH_3)_3I^- \xrightarrow{heat}$$

$$CH_2 = CH_2 + (CH_3)_3 N + HI$$

(b)
$$CH_3CH_2N^+(CH_3)_3OH^- \xrightarrow{heat}$$

$$CH_2 = CH_2 + (CH_3)_3 N + H_2O$$

- Both a and b
- $CH_3CH_2NH_2 \xrightarrow{heat} CH_2 = CH_2 + NH_3$
- 26. Which one of the following is not an oxidation product of a primary amine?
 - (a) A hydroxylamine
- (b) A nitroso compound
- (c) A nitro compound
- (d) None of these
- 27. Which of the following method is used for eliminating nitrogen of an amine present outside the ring?
 - (a) Hofmann elimination
- (b) Cope elimination
- (c) Both
- (d) Emde degradation
- 28. Which of the following does not react with nitrous acid?
 - (a) $C_6H_5NH_2$
- (b) $C_6H_5NHCH_3$
- (c) $C_6H_5N(CH_3)_2$
- (d) None of these
- 29. Which of the following leads to carbon-carbon double bond?
 - (a) 1° Amine + RCHO \rightarrow (b) 2° Amine + R₂CO \rightarrow
 - (c) 2° Amine + RCHO \rightarrow
- (d) both b & c
- 30. Electrophilic aromatic substitution of pyridine resembles with
 - (a) benzene
- (b) aniline
- (c) nitrobenzene
- (d) none of these



M. D. Voys	19.abcd	20. a b c d	21. a b c d	22. abcd	23. abcd
Mark Your Response	24. a b c d	25. a b c d	26. abcd	27. abcd	28. abcd
	29. a b c d	30. abcd			

- 31. Which of the following will react most readily with NaOH to form ethanol?
 - $(CH_3)_4 N^+ I^-$
- (b) $(CH_3)_4 S^+ I^-$
- (CH₃)₃CCl
- (d) CH₃OCH₃
- NOH Ш CCH₃ 32.

(b)
$$O_2N$$
 NHCOCH₃ Br

(c)
$$O_2N$$
 CONHCH₃ Br

(d)
$$O_2N$$
 CCH_3

- 33. In Hofmann bromamide degradation, one of the important steps is the migration of
 - (a) an alkyl group without its electron pair to electron deficient N atom
 - (b) an alkyl group with its electron pair to electron deficient O atom
 - (c) an alkyl group with its electron pair to electron rich N atom
 - (d) an alkyl group with its electron pair to electron deficient N atom

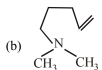
- $(CH_3)_3C \stackrel{||}{C} NH_2 \xrightarrow{(i) OD^-/Br_2} Product P is$ 34.
 - (a) $(CH_3)_3 CNH_2$
- (b) (CH₃)₃ CNHD
- (c) (CH₃)₃CND₂
- (d) no reaction
- 35. Reaction of ethyl amine with alkaline chloroform leads to the formation of carbylamine reaction. This reaction involves the attack of an electrophile on ethyl amine, the electrophile is
 - (a) H_2O^+
- (b) H⁺
- (c) RNH_3^+
- (d) : CCl₂
- 36. Predict the possible number of alkenes and the main alkene in the following reaction.

$$\begin{array}{c}
\text{NHCH}_2\text{CH}_3 & \xrightarrow{\text{(i) excess CH}_3\text{I}} & [] \xrightarrow{\text{heat}} & \text{Product}
\end{array}$$

- (a) 2 and /
- (b) 2 and /
- (c) 3 and $H_2 C = CH_2$ (d) 2 and $H_2 C = CH_2$

37.
$$(i) CH_3I \longrightarrow Product P is$$

$$CH_3 \xrightarrow{(ii) Ag_2O, heat} Product P is$$



(c)
$$CH_3$$
 CH_3

38.
$$(i) \xrightarrow{H_2O_2} Product P is$$

$$CH_3 N(CH_3)_2$$

(a)
$$V_{CH_3}$$
 CH_3









Mark Your	31.abcd	32. a b c d	33. abcd	34. abcd	35. abcd
RESPONSE	36. a b c d	37. a b c d	38. a b c d		



39.
$$R - C - CH - N = N$$
 Intermediate + N_2

What is the nature of its intermediate in this reaction?

- (a) Carbonium ion
- (b) Carbanion
- (c) Carbene
- (d) Free radial
- 40. The intermediate formed in the above reaction rearranges to

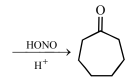
(a)
$$R - CH_2 - CH$$
 (b) $R - CH = C = O$

(b)
$$R - CH = C = O$$

(c)
$$R - C - CH_3$$

(d)
$$R - C - C = C$$

41.
$$\underbrace{\begin{array}{c} O \\ \\ CH_3NO_2 \\ \hline C_2H_5O^- \end{array}}_{C_2H_5O^-} [A] \longrightarrow [B]$$



Which reaction is/are involved in the above conversion?

- (a) Diazotisation and Beckmamn rearrangement
- (b) Diazotisation and Robinson annulation
- (c) Aldol condensation and diazotisation
- (d) Diazotisation and pinacol type of rearrangement
- Which of the reaction can not be used for introducing a 42. C = C linkage?
 - (a) Hofmann elimination
- (b) Aldol condensation
- (c) Wittig reaction
- (d) None of the three
- 43. Following reaction is an example of

$$\begin{array}{c}
O \\
R - C - OH + HN_3 \xrightarrow{H_2SO_4} R - NH_2
\end{array}$$

- (a) Hofmann reaction
- (b) Curtius reaction
- (c) Schmidt reaction
- (d) Lossen reaction

- 44. Which of the following is one of the intermediates in Hofmann, Curtius, Schmidt and Lossen reaction?
 - (a) RCO
- (b) RNCO
- (c) RCNO
- (d) RCONH
- 45. Which of the following involves migration to electrondeficient nitrogen?
 - (a) Benzilic acid rearrangement
 - (b) Wolf rearrangement
 - (c) Allylic rearrangement
 - (d) Beckmann rearrangement
- 46. Which of the following is not formed as an intermediate in the Hofmann rearrangement?

$$R-CONH_2 \xrightarrow{\quad Br_2 \ / \ NaOH \quad} RNH_2$$

$$\begin{array}{ccc}
 & O & H \\
 & \parallel & \parallel \\
 & (a) & R - C - N Br
\end{array}$$

(b)
$$R - C - N - Br$$

- (d) All the three are formed
- 47. Identify (C) and (D) in the following series of reactions

$$CH_3NH_2 \xrightarrow{excess \text{ of } \atop CH_3I} [A] \xrightarrow{AgOH} [B] \xrightarrow{heat} [C] + [D]$$

- (a) (CH₃)₃COH, CH₃NH₂
- (b) $(CH_3)_2C = CH_2, CH_3NH_2$
- (c) (CH₃)₃N, CH₃OH
- (d) $(CH_3)_2C=CH_2$, CH_3OH
- Identify (D) and (E) in the given reaction 48.

$$CH_3COOH \xrightarrow{(i) LiAlH_4} (A) \xrightarrow{(i) KCN} (B)$$

$$\xrightarrow{\text{CHCl}_3/\text{OH}^-} \text{(C)} \xrightarrow{\text{hydrolysis}} \text{(D)} + \text{(E)}$$

- (a) CH₃CH₂CH₂NHCH₃
- (b) (CH₃)₂CHNH₂, HCOOH
- (c) CH₃CH₂CH₂NH₂, HCOOH
- (d) CH₂CH₂CH₂COOH

Mark Your	39. a b c d	40. a b c d	41. a b c d	42. abcd	43. abcd
Response	44. a b c d	45. a b c d	46. abcd	47. abcd	48. abcd

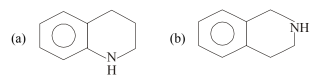
Z is

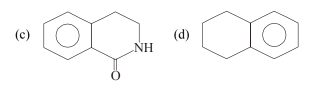
- (a) a single compound
- (b) a mixture of two compounds
- (c) a mixture of three compounds
- (d) a mixture of four compounds

51.
$$\frac{\text{(i) CHN}_2\text{COOC}_2\text{H}_5}{\text{(ii) Heat with Cu powder}} [Y] \xrightarrow{\text{oxidation}} [Z]$$

Compound Z should be

52. Which of the following undergoes bromination most easily?



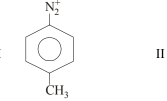


- 53. How many structural isomers of a Grignard reagent are possible for preparing *n*-butane by reaction with ethyl amine?
 - (a) 1

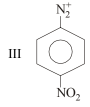
(b) 2

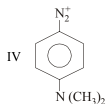
(c) 3

- (d) 4
- **54.** For the diazonium ions, the order of reactivity towards diazo-coupling with phenol in the presence of dil.NaOH is









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



Mark Your	49. a b c d	50. a b c d	51.abcd	52. abcd	53. abcd
RESPONSE	54. a b c d				



COMPREHENSION TYPE

This section contains groups of questions. Each group is followed by some multiple choice questions based on a paragraph. Each question has 4 choices (a), (b), (c) and (d) for its answer, out of which ONLY ONE is correct.

PASSAGE-1

When a quaternary ammonium halide is treated with moist silver oxide (a slurry of Ag₂O in water), a quaternary ammonium hydroxide is formed as the main product. Thermal decomposition of the quaternary ammonium hydroxide gives an alkene. This reaction is known as Hofmann elimination and has most of the characteristics of an E2 reaction.

$$\label{eq:ch3ch2} \begin{aligned} \mathsf{CH_3CH} &= \mathsf{CHCH_3} \, + \, \mathsf{CH_3CH_2CH} = \mathsf{CH_2} \\ &\quad \mathsf{X} &\quad \mathsf{Y} \end{aligned}$$

- The Hofmann eliminations are 1.
 - (i) concreted
- (ii) anti-stereoselective
- (iii) occur in separate steps (iv) syn-stereoselective
- (a) (i) and (iv)
- (b) (i) and (ii)
- (c) Only (i)
- (d) Only (iii)
- 2. In the above reaction, which of the following is correct regarding yield of X and Y?
 - (a) X = Y
- (b) $X \approx Y$
- (c) X > Y
- (d) X < Y

PASSAGE-2

The conversion of an amide to an amine with one carbon atom less by the action of alkaline hypohalite is known as Hofmann degradation.

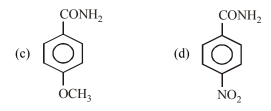
$$RCONH_2 + Br_2 + 4KOH \longrightarrow RNH_2$$

The most important feature of the reaction is the rearrangement of N-bromamide anion to isocyanate:

$$\begin{array}{ccc} O & O \\ || & || \\ R-C-N & \longrightarrow \begin{array}{c} C=N-R \\ II \end{array} & \xrightarrow{H_2O} \begin{array}{c} H_2O \\ \end{array} \\ \rightarrow H_2N-R+CO_2 \end{array}$$

Hofmann reaction is accelerated if the migrating group is more electron-releasing.

- 3. Which step is the driving force in the above reaction to proceed in right direction?
 - (a) conversion of I to II
 - (b) conversion of II to III
 - (c) conversion of III to RNH₂
 - (d) All
- Which of the following can undergo Hofmann reaction most easily?



- $-\text{CONH}_2 \xrightarrow{\text{OBr}^-}$ Product amine will be 5. C_6H_5
 - (+) α -Phenylpropionamide
 - (a) (+) –
- (b) (-) –
- (c) 50 : 50 racemic
- (d) non-50:50 racemic



6. Predict the product in the following reaction :

$$CONH_2$$
 $CONH_2$
 $CONH_2$
 $CONH_2$
 $CONH_2$
 $CONH_2$
 $CONH_2$
 $CONH_2$
 $CONH_2$

- (i) NH_2
- (ii) NH₂
- $\text{(iii)} \ \bigodot^{15}_{\text{NH}_2}$
- (iv) NH₂
- (a) (i) and (ii)
- (b) (ii) and (iii)
- (c) (i) and (iii)
- (d) All the four

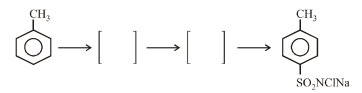
PASSAGE-3

Understand carefully the following two reactions and answer the questions mentioned below.

Reaction (i)

$$\stackrel{\text{NHCOCH}_3}{\longrightarrow} \left[\quad \right] \longrightarrow \left[\quad \right] \longrightarrow \stackrel{\text{NH}_2}{\bigodot}$$

Reaction (ii)



- 7. Which of the steps is(are) common in the two reactions?
 - (a) 1st step
- (b) 2nd step
- (c) both
- (d) none

- **8.** Which of the final products are medicinally important?
 - (a) Product from reaction (i)
 - (b) Product from reaction (ii)
 - (c) Products from both reactions
 - (d) None of the two products
- **9.** Which one of the following reagent can be used for the introduction of Cl and Na in reaction (ii)?
 - (a) Cl_2 / Na_2CO_3
- (b) HCl/NaOH
- (c) NaCl/NaOH
- (d) NaOCl / NaOH
- 10. Saccharin, an important sweetening agent, of the following

structure
$$N^{-}K^{+}$$
 is prepared by one of the

products of the reaction (ii), that product should be

(a)
$$OCCH_3$$
 $OCCH_3$

(c)
$$OONH_2$$

 SO_2NH_2

PASSAGE-4

$$+ CH_3CH = CH_2 \xrightarrow{AlCl_3}$$

$$\xrightarrow{\text{COOH}} \xrightarrow{\text{(i) PCl}_5}$$

$$C_6H_5CON_3 \xrightarrow{\text{heat}} C_6H_5NCO \xrightarrow{C_2H_5OH} \text{Product}$$

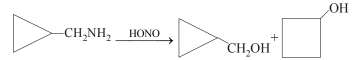
11. What will be the product(s) when isopropylbenzene is oxidised with oxygen and product is acidified

(d)
$$OH$$
 + $(CH_3)_2C = O$

- **12.** The final product in the above series of reactions is
 - (a) an amine
 - (b) a mono substituted urea
 - (c) a disubstituted urea
- (d) a urethane
- 13. Conversion of C₆H₅CON₃ to C₆H₅NCO is an example of rearrangement where an alkyl group migrates to
 - (a) electron deficient carbon atom
 - (b) electron deficient oxygen atom
 - (c) electron deficient nitrogen atom
 - (d) electron rich nitrogen atom

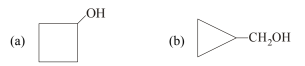
PASSAGE-5

Nitrous acid reacts with amines forming different products depending upon the type of amine. Aliphatic primary amines react with HONO forming alcohol as the major product, other products being alkene and alkyl halide. Certain cyclic primary amines can undergo ring expansion or ring contraction on treatment with HONO again forming alcohol as the major product.



Aromatic primary amines when treated with HONO undergo diazotization to form diazonium salt as the stable product.

- **14.** Which of the following statement is not true?
 - (a) All primary amines first form diazonium salt when treated with nitrous acid.
 - (b) All aliphatic primary amines when treated with nitrous acid form primary alcohols as the major product
 - (c) The three classes of amines give different products on treatment with nitrous acid.
 - (d) None of the three





16. $OH \xrightarrow{NaNO_2} Stable \text{ product } Z \text{ may be}$

(a)
$$CH_2OH$$
 (b) OH

17.
$$(+)$$
 - CH₃CHCH₂CH₃ $\xrightarrow{\text{HONO}}$ Product is

- (a) enantiomer of the original compound
- (b) diastereomer of the original compound
- (c) racemic mixture of the original compound
- (d) (+) isomer



Mark Your	11. abcd	12. a b c d	13. abcd	14. abcd	15. abcd
RESPONSE	16.abcd	17. abcd			

PASSAGE-6

Hofmann reaction is found to follow the following path.

$\begin{array}{ccc} & & & O & \\ R - \overset{\parallel}{C} - NH_2 & \xrightarrow{Br_2} & R - \overset{\parallel}{C} - NHBr \xrightarrow{OH^-} \end{array}$

$$R - \overset{O}{\overset{||}{C}} \overset{\bigcirc}{\overset{||}{\overset{||}{O}}} Br \xrightarrow{-Br^{-}} R - \overset{O}{\overset{||}{C}} \overset{||}{\overset{||}{\overset{||}{O}}} \longrightarrow$$

$$\begin{array}{c}
O \\
\parallel \\
C = \dot{N} - R \xrightarrow{H_2O} H_2NR + CO_2
\end{array}$$
IV

- Number of moles of NaOH used in the complete reaction 18.
 - (a) 1

(b) 2

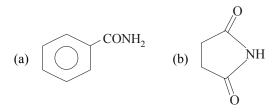
(c) 3

- (d) 4
- Hofmann reaction, when carried out in presence of 19. methanol containing sodium methoxide, instead of NaOH,

$$C_6H_5CONH_2 \xrightarrow{Br_2,CH_3ONa} Product is$$

20.
$$CH_3 - C - ND_2 \xrightarrow{NaOH} Amine, Z is$$

- (a) $CH_3 ND_2$
- (b) CH₃NH₂
- (c) mixture of (a) and (b) (d) Reaction not possible
- *21. Which of the following will not undergo Hofmann bromamide reaction?



- C₆H₅CONHCH₃
- (d) C₆H₅CONHBr

- 22. Which of the following species in the above mechanism is electron deficient?
 - (a) I

(b) II

(c) III

- (d) IV
- In presence of alkali, (b) undergoes hydrolysis to form CH2CONH2

which when undergoes Hofmann reaction. CH₂COO

PASSAGE-7

Beckmann rearrangement is used for determining the configuration of ketoxines since the two isomers form different N-substituted amides when heated with some acidic reagent. The different steps of the mechanism are as follows.

$$\begin{array}{c} R \\ R' \end{array} C = N \xrightarrow{OH} \xrightarrow{H^+} \begin{array}{c} R \\ R' \end{array} C = N \xrightarrow{OH_2} \xrightarrow{OH_2}$$

$$\begin{array}{c} R \\ R' \end{array} > C = \stackrel{}{N}^{+} \xrightarrow{-H^{+}} R - \stackrel{+}{C} = \stackrel{\cdot}{N}R' \xrightarrow{H_{2}O} \end{array}$$
II III

$$R \to R' \longrightarrow R' \longrightarrow R' \longrightarrow R'$$

- Pick up the species which has an electron deficient centre? 23.
 - (a) I

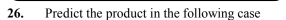
(b) II

(c) III

- (d) Two of the three
- 24. The mechanism involves migration of
 - R group with its bonding electron pair
 - (b) R' group with its bonding electron pair
 - R' group without its bonding electron pair
 - (d) OH group with its bonding electron pair
- 25. Which of the following reaction/step is involved in Beckmann rearrangement?
 - (a) Formation of conjugated acid
 - (b) Migration of an alkyl group to electron deficient N
 - Enlolisation
 - All the three



Mark Your	18. abcd	19. abcd	20. abcd	21. abcd	22. abcd
RESPONSE	23. a b c d	24. a b c d	25. a b c d		



$$\begin{array}{c} \text{CH}_{3} \\ \text{C}_{2}\text{H}_{5} \end{array} \text{C} = \text{N} \xrightarrow{\begin{array}{c} \text{(i) PCl}_{5} \\ \text{(ii) D}_{2}\text{O} \end{array}} \text{P}$$

- (a) $CH_3CON(D)C_2H_5$
- (b) $C_2H_5CON(D)CH_3$
- (c) CH₂CONHC₂H₅
- (d) C₂H₅CONHCH₂

27.
$$CH_3 C = N$$
 $C_2H_5 C = N$ OH $CH_2O^{18} \rightarrow Q$ is

- (a) C₂H₅NHCOCH₃
- (b) $C_2H_5NHCO^{18}CH_3$
- (c) $CH_3NHCO^{18}C_2H_5$ (d) $C_2H_5CONHCH_3$

PASSAGE-8

Like ammonia, an amine (1°, 2° or 3°) can react with an alkyl halide to form next higher class of amine. Here, again it is the presence of electron pair on nitrogen which makes amines to behave as nucleophile and alkyl halides thus undergo nucleophilic substitutions.

$$\begin{array}{c} \vdots \\ R\,N\,H_2 \\ \text{$^{1^\circ}$ Amine} \\ \text{(Nucleophile)} \end{array} \xrightarrow{1^\circ\text{Alkyl halide}} \begin{array}{c} H & X^- \\ | \\ \end{array} \\ H \end{array}$$

$$\longrightarrow R \overset{\dots}{N} - CH_2R' + HX$$

$$H$$
2° Amine

A second alkylation may follow, converting the secondary amine to a tertiary amine which may be alkylated to give a quaternary ammonium salt.

28. Identify the product(s) obtained when Br(CH₂)₄Br is heated with 1 equivalent of ethyl amine

(b)
$$C_2H_5NH$$
 NHC₂H₅



29. Identify the product(s) obtained when Br(CH₂)₄Br is heated with excess of methyl amine

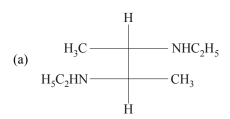
(c)
$$N$$
 CH_3

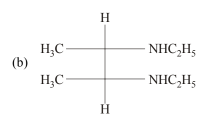
30.

$$CH_3$$

$$N + C_2H_5NH_2 \longrightarrow Z, \text{ here } Z \text{ is}$$

$$CH_3^{\text{MMV}}$$





- (c) Both of these
- (d) None of the two



PASSAGE-9

Summary of the four related name reactions for the preparation of primary amines are given below.

$$RCONH_2 \xrightarrow{Br_2, OH^-} RNH_2$$

Hofmann

$$RCON_3 \xrightarrow{H_2O} RNH_2$$

Curtius

$$RCONHOH \xrightarrow{OH^{-}\atop H_2O} RNH_2$$

Lossen

$$RCOOH \xrightarrow{HN_3} RNH_2$$

- **31.** Reactant of which reaction is not a derivative of carboxylic acid?
 - (a) Hofmann
- (b) Curtius
- (c) Lossen
- (d) None of the three

- 32. In which of the reactions, leaving group is carboxylate anion?
 - (a) Hofmann
- (b) Cutius
- (c) Lossen
- (d) Schmidt
- **33.** In which of the reactions, nitrogen constitutes the leaving group?
 - (a) Curtius
- (b) Schmidt
- (c) Both
- (d) Hofmann
- **34.** Which of the reaction does not require alkaline medium?
 - (a) Hofmann
- (b) Curtius
- (c) Schmidt
- (d) Lossen
- **35.** Which of the reaction is of least importance?
 - (a) Schmidt
- (b) Hofmann
- (c) Curtius
- (d) Lossen



MARK YOUR RESPONSE

31.@bcd

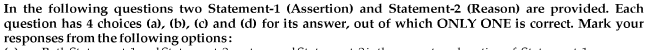
32. a b c d

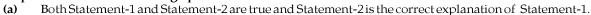
33. abcd

34. (a) b) c) d)

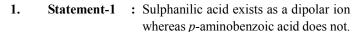
35. abcd







- (b) Both Statement-1 and Statement-2 are true and Statement-2 is not the correct explanation of Statement-1.
- (c) Statement-1 is true but Statement-2 is false.
- (d) Statement-1 is false but Statement-2 is true



Statement-2 : Carboxyl group, being more acidic than $-SO_3H$ group, can easily transfer a H^+ to the amino group.

Statement-1 : Nitration of aniline can be conveniently done by protecting the amino group by acetylation.

Statement-2: Acetylation increases the electrondensity in the benzene ring.

3. **Statement-1**: Reduction of *m*-dinitrobenzene with ammonium sulphide gives *m*-nitroaniline.

Statement-2 : *m*-Nitroaniline formed gets precipitated and hence further reduction is prevented.

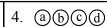
 Statement-1 : Benzonitrile is prepared by the reaction of chlorobenzene with potassium cyanide.

Statement-2: Cyanide (CN⁻) is a strong nucleophile.

5. Statement-1: 1° Amides react with Br₂ + NaOH to give 1° amines with one carbon atom less than the parent amide.

Statement-2: The reaction occurs through intermediate formation of acylnitrene.









6. Statement-1 : In strongly acidic solutions, aniline becomes more reactive towards electrophilic reagents.

Statement-2: The amino group being completely protonated in strongly acidic solution, the lone pair of electrons on the nitrogen is no longer available for resonance.

7. **Statement-1**: In order to convert R-Cl to pure R-NH₂, Gabriel-phthalimide synthesis can be used.

Statement-2: With proper choice of alkyl halides, phthalimide synthesis can be used to prepare 1°, 2° or 3° amines.

8. Statement-1 : Aniline does not undergo Friedel-Crafts reaction.

Statement-2 : -NH₂ group of aniline reacts with AlCl₃ (Lewis acid) to give acid-base reaction.

Statement-1 : Acetamide reacts with Br₂ in presence of methanolic CH₃ONa to form methyl N-methylcarbamate.

Statement-2: Methyl isocyanate is formed as an intermediate which reacts with methanol to form methyl N-methylcarbamate.

10. Statement-1 : 1°, 2° and 3° amines are hydrogenbonding donors, while 1° and 2° amines are hydrogen-bonding acceptors.

Statement-2: Low molecular weight amines are soluble in water.

11. Statement-1 : Pyridine is basic.

Statement-2: Lone pair of electrons on its nitrogen is not involved in aromatic sextet.

12. Statement-1 : Pyridine is more basic than pyrrole.

Statement-2 : Lone pair of electrons on N in pyridine and pyrrole are different in nature, these form a part of aromatic sextet in pyrrole, while not in pyridine

while not in pyridine.

13. Statement-1: All compounds containing an odd number of nitrogen atoms have odd masses and those containing even number of N atoms have even masses.

Statement-2: Nitrogen rule can be applied to both aliphatic and aromatic compounds.

14. Statement-1 : The main product of reaction of alcoholic silver nitrite and ethyl bromide is nitroethane.

Statement-2: Silver nitrite is predominantly covalent compound.

15. Statement-1 : HCN on hydrolysis gives formic acidStatement-2 : HCN is a weak monoprotic acid

16. Statement-1 : Hofmann elimination gives alkeneStatement-2 : Alkenes are less acidic than NH₃

17. Statement-1: Nitromethane can give aldol condensation.

Statement-2 : α -Hydrogen of nitromethane is acidic.



Mark	Your
RESP	ONSE

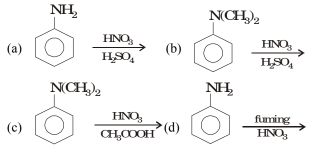
6. abcd	7. abcd	8. abcd	9. abcd	10. abcd
11. abcd	12. a b c d	13. a b c d	14. abcd	15. abcd
16. a b c d	17. a b c d			



MULTIPLE CORRECT CHOICE TYPE

Each of these questions has 4 choices (a), (b), (c) and (d) for its answer, out of which ONE OR MORE is/are correct.

1. Nitro group can't be introduced easily by which reaction?



- **2.** Which of the following compound can be methylated by diazomethane?
 - (a) C₂H₅COOH
- (b) $C_2H_5NH_2$
- (c) C_6H_5OH
- (d) CH₃COCH₂COOC₂H₅
- **3.** Which of the following amine can be prepared by Gabriel method?
 - (a) CH₃CH₂NH₂
- (b) $(CH_3)_2CHNH_2$
- (c) (CH₃)₃CNH₂
- (d) $C_6H_5NH_2$



MARK	Your
RESI	PONSE

- 1. **abcd**
- 2. (a)(b)(c)(d)
- 3. a

- **4.** Which of the following can't be used as an alkylating agent for an amine?
 - (a) CH₃CH₂Cl
- (b) $CH_2 = CHCl$
- (c) C₆H₅Cl
- (d) $(C\tilde{H_3})_3CCl$
- 5. $(CH_3)_3CNH_2$ can't be prepared by
 - (a) reductive amination
- b) oxime reduction
- (c) Hofmann degradation
- (d) Gabriel synthesis.
- **6.** Which of the following reaction can be used for preparing aniline?
 - (a) $C_6H_5COOH \xrightarrow{N_3H, conc. H_2SO_4}$
 - (b) $C_6H_5NC \xrightarrow{H_3O^+}$
 - (c) $C_6H_5NC \xrightarrow{LiAlH_4}$
 - (d) $C_6H_5CONH_2 \xrightarrow{Br_2/NaOH}$
- 7. Which of the following can exist as inner salt?
 - (a) p-Aminobenzenesulphonic acid
 - (b) p-Aminobenzoic acid
 - (c) Aminoacetic acid
 - (d) Alanine
- **8.** Which of the following pairs show coupling reaction?

(a)
$$N_2^+ Cl^-$$
 OCH₃ $+$ OCH_3

$$(b) \qquad \begin{array}{c} N_2^+ \text{Cl}^- \\ NO_2 \\ NO_2 \end{array} + OCH_3$$

(d) Diazotised sulphanilic acid + Dimethylaniline

9. Which of the following statement is true regarding reaction of *p*-aminophenol with arenediazonium chloride?

$$HO \longrightarrow \underbrace{\begin{array}{c} 3 & 2 \\ \\ 5 & 6 \end{array}} \longrightarrow NH_2 + ArN_2^+Cl^- \longrightarrow$$

- (a) Reaction takes place at position 2 in presence of HCl.
- (b) Reaction takes place at position 3 in presence of NaOH.
- (c) Only two positions (2 and 6) can be coupled in presence of OH^-
- (d) Four azo groups can be introduced in the molecule.
- **10.** Which of the following reacts with nitrous acid?
 - (a) Acetamide
 - (b) 2-Nitrobutane
 - (c) 2-Methyl-2-nitropropane
 - (d) Diethylamine
- **11.** *p*-Nitroaniline can be obtained by

(a)
$$\frac{\mathrm{SO_3H}}{\mathrm{HNO_3}}$$

$$(b) \quad \overbrace{ \qquad \qquad }^{\text{NH}_2} \xrightarrow{(i) \text{ HNO}_3}$$

(c)
$$(ii) \xrightarrow{\text{(i) } (\text{CH}_3\text{CO)}_2\text{O}} \xrightarrow{\text{(iii) } \text{HNO}_3/\text{H}_2\text{SO}_4,}$$

$$(d) \qquad \stackrel{\text{NH}_2}{\bigodot} \qquad \xrightarrow{\text{HNO}_3} \rightarrow$$

$$\text{SO}_3\text{H}$$

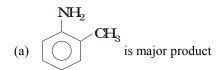


Mark Your	4. abcd	5. abcd	6. abcd	7. abcd	8. abcd
RESPONSE	9. abcd	10. abcd	11. abcd		

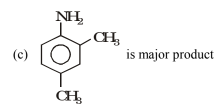


12. $\langle \bigcirc \rangle$ NH₂ + CH₃Cl $\xrightarrow{AlCl_3}$?

Which of the following statement is false regarding above reaction?



$$\begin{array}{c} NH_2 \\ \hline \\ CH_3 \end{array}$$



- (d) No substitution
- 13. The correct name for $CH_3 N \stackrel{\triangle}{=} C$ is
 - (a) methyl carbylamine
- (b) methyl isocyanide
- (c) methyl isonitrile
- (d) acetoisonitrile
- **14.** Urea on heating with ethyl alcohol gives
 - (a) urethane
- (b) semicarbazide
- (c) malonylurea
- (d) ethyl carbonate

15.
$$(i) O_2, V_2O_5, heat \atop (iii) NH_3, heat \atop (iii) NaOH / Br_2$$
 Product

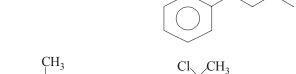
Which of the following statement (s) is (are) true regarding above series of reactions?

(a) Product formed is
$$\begin{array}{c} \text{COOH} \\ \text{CONH}_2 \end{array}$$

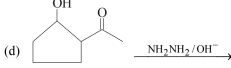
- (b) Product formed is $\begin{array}{c} NH_2 \\ COOH \end{array}$
- (c) Reaction involved is oxidation
- (d) Reaction involved is Hofmann degradation

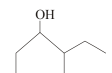
- **16.** Oximes can be converted into amides by
 - (a) SO₃
- (b) BF₃
- (c) PCl₅
- (d) $C_6H_5SO_2Cl$
- 17. Pick up the correct statement
 - (a) Toluene and ethane, both react at the same rate with chlorine in presence of light.

(b)
$$\xrightarrow{H_2/Pt}$$



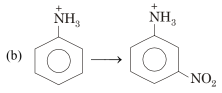
(c)
$$+ \text{Cl}_2 \xrightarrow{\text{light}}$$
 OH





18. Which of the following is/are correct regarding nitration of aniline with conc HNO₃ and conc. H₂SO₄?

(a)
$$\xrightarrow{NH_2} \xrightarrow{\text{conc. HNO}_3} \xrightarrow{NH_2} \xrightarrow{NH_2} \xrightarrow{NH_2} \xrightarrow{NH_2} \xrightarrow{NH_2} \xrightarrow{NO_2} \xrightarrow{NO_$$



- (c) The substitution can be explained on the basis of inductive effect (– I)
- (d) The substitution can be influenced by +M and +E effects.



Mark Your	12. a b c d	13. a b c d	14. abcd	15. abcd	16. abcd
RESPONSE	17. a b o d	18. abcd			

19. In which of the following reaction, Hofmann elimination is the major product?

(a)
$$Me_3COK$$
 Br

(b)
$$\xrightarrow{\text{conc.H}_2\text{SO}_4 \atop \text{heat}}$$

(c)
$$\stackrel{+}{N}Me_3OH^- \xrightarrow{heat}$$

(d)
$$C1 \xrightarrow{\text{alco. KOH}}$$

- Which of the following intermediates are likely to be 20. formed when 2-methylpropyl amine is treated with nitrous acid?
 - (a) $(CH_3)_2CHCH_2$ (b) $(CH_3)_3COH_2$
 - (c) (CH₃)₂CHCH₂OH₂ (d) (CH₃)₂CCH₃

- 21. Carbenes are the reactive intermediates in
 - (a) Reimer Tiemann reaction
 - (b) Wittig reaction
 - (c) Hofmann bromamide reaction
 - (d) Carbylamine reaction
- 22. Which of the following are intermediates in Sandmeyer reaction?
 - (a) $C_6H_5N^+ \equiv NC1^-$ (b) $C_6H_5N^+ \equiv N$
 - (c) \dot{C}_6H_5
- (d) C_6H_5Cl

23.
$$C_4H_{11}N + HONO \longrightarrow C_4H_{10}O$$
(X) (3° alcohol)

X should

- (a) give carbylamine reaction
- (b) undergo diazotisation
- (c) react with water
- (d) give Hofmann bromamide reaction



E

Mark Your

19 (a)(b)(c)(d)

20. (a) (b) (c) (d)

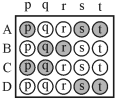
21. (a) (b) (c) (d)

22. (a) (b) (c) (d)

23. (a)(b)(c)(d)

MATRIX-MATCH TYPE

Each question contains statements given in two columns, which have to be matched. The statements in Column-I are labeled A, B, C and D, while the statements in Column-II are labelled p, q, r, s and t. Any given statement in Column -I can have correct matching with ONE OR MORE statement(s) in Column-II. The appropriate bubbles corresponding to the answers to these questions have to be darkened as illustrated in the following example: If the correct matches are A-p, s and t; B-q and r; C-p and q; and D-s then the correct darkening of bubbles will look like the given.





- (A) Hydrogen bonding
- Column-II Ethyl acetoacetate
- (B) β-Amino acids
- **Proteins**
- COOH r. (C) H₂N
 - Zwitterion

(D) Aniline

2.

3.

Diazotisation

Column-I

Column-II

NH

- (B) $H_2N \ddot{C} NH_2$
- Two types of N
- Aromatic
- (D) $C_6H_5N^+ \equiv NCl^-$
- Non-aromatic

Column-I

- (A) $CHCl_3 \xrightarrow{OH^-}$
- Column-II Hofmann product
- (B) $CHCl_2 CF_3$
- E2 mechanism

$$OC_2H_5^-$$

α-Elimination

$$\xrightarrow{\text{Me}_3\text{CO}^-}$$

(D)
$$CH_3 - CH - CH_2CH_3$$
 s. $N^+(CH_3)_3$

$$\xrightarrow{OH^-}$$

4. Column-I

- (A) ArS_{N_1}
- Carbocation

Column-II

- as intermediate
- (B) ArS_{N_2}
- Carbanion
- as intermediate
- (C) $OH^- > (CH_3)_3CO^-$
- Basic character
- (D) $OH^- < NH_2^-$
- Nucleophilic character

5. Column-I

- Column-II
- (A) Allylic rearrangement Carbanions
- (B) Hofmann rearrangement q. Electron deficient
 - species
- (C) Wittig reaction
- Carbocations
- (D) Carbylamine reaction
- Carbenes

Column-I 6.

- Column-II
- (A) Gabriel phthalimide reaction
- C₆H₅CH₂NH₂
- (B) Reduction with LiAlH₄
- $C_6H_5NH_2$
- (C) Reaction with alc. KOH + CHCl₃
- C₆H₅CN
- (D) 1° Amide with $Br_2 + KOH$
- s. CH₃CH₂NH₂

Ø1

Mark Your RESPONSE

- 1. BPQTS
 - C

- 2.
 - (P)(q)(r)(s)**PQTS**
- 5.

- 3.
 - (P)(q)(r)(s)(P)(I)
- 6.

7. Column I

Column II

Column I

(B) $CH_2 = CHC1$

9.

(A)

Column II

: CCl₂

CHCl₃ / alc. KOH

- (A) Pyrolysis of quaternary p. ammonium salts
- syn-Elimination
- (B) β-Dehydrohalogenation q.
 - anti-Elimination
- (C) γ-Dehydrohalogenation r.
- Saytzeff product
- (D) Dehydration of alcohol s.
- Hofmann product

8. Column I

Column II

- (A) Normal base with 1° alkyl fluoride
- E1 reaction
- (B) Normal base with 2°

(C) Bulkyl base with 1°

- E2 reaction
- alkyl chloride
- Saytzeff product
- alkyl halide
- (D) Dehydration of alcohol s.

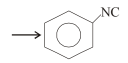
Hofmann product

- OH (C)
- NaNH₂

q.

 \rightarrow CH \equiv CH

- NH_2 (D)
- C_2H_5OH / KOH



10.

Column - I

Column - II

- (A) Hofmann degradation
- (B) Beckmann
- Amides Isocyanates
- rearrangement
- (C) Curtius rearrangement NaOH/Br₂
- RCON₃ (D) Lossen rearrangement



Mark Your RESPONSE

- pqrs 7. $\mathbb{P}(\mathbb{Q})$
- 10. **PQTS P9TS** \mathbf{C} **PQTS**
- p q r s 8. P(0)
- 9.



newarkey

SINGLE CORRECT CHOICE TYPE

1.	(b)	7.	(a)	13.	(c)	19.	(b)	25.	(b)	31.	(a)	37.	(b)	43.	(c)	49.	(d)
2.	(c)	8.	(d)	14.	(b)	20.	(d)	26.	(d)	32.	(b)	38.	(d)	44.	(b)	50.	(c)
3.	(b)	9.	(b)	15.	(b)	21.	(b)	27.	(c)	33.	(d)	39.	(c)	45.	(d)	51.	(c)
4.	(a)	10.	(d)	16.	(c)	22.	(a)	28.	(d)	34.	(c)	40.	(b)	46.	(d)	52.	(a)
5.	(d)	11.	(c)	17.	(d)	23.	(b)	29.	(d)	35.	(d)	41.	(c)	47.	(c)	53.	(b)
6.	(a)	12.	(d)	18.	(d)	24.	(b)	30.	(c)	36.	(c)	42.	(d)	48.	(c)	54.	(b)

$oxed{\mathbf{B}}$ $oxed{oxed}$ Comprehension Type $oxed{oxed}$

1	(b)	7	(c)	13	(c)	19	(b)	25	(d)	31	(d)
2	(d)	8	(c)	14	(b)	20	(b)	26	(c)	32	(c)
3	(b)	9	(d)	15	(a,b)	21	(c)	27	(c)	33	(c)
4	(c)	10	(a)	16	(d)	22	(c)	28	(a, d)	34	(b)
5	(a)	11	(d)	17	(c)	23	(d)	29	(c,d)	35	(d)
6	(c)	12	(d)	18	(d)	24	(b)	30	(a)		

REASONING TYPE **≡**

1	(c)	4	(d)	7	(c)	10	(b)	13	(b)	16	(b)
2	(c)	5	(a)	8	(a)	11	(a)	14	(a)	17	(a)
3	(c)	6	(d)	Q	(a)	12	(a)	15	(d)		

MULTIPLE CORRECT CHOICE TYPE

ANSWER KEY												
1.	(a,b,d)	5.	(a,b,d)	9.	(a,b,d)	13.	(a,b,d)	17.	(c,d)	21.	(a,b,d)	
2.	(a,c,d)	6.	(a,b,d)	10.	(a,b,d)	14.	(a,d)	18.	(b,c)	22.	(b,c)	
3.	(a,b)	7.	(a,c,d)	11.	(c,d)	15.	(b,c,d)	19.	(a,c)	23.	(a,b,c)	
4.	(b,c,d)	8.	(b,c,d)	12.	(a,b,c)	16.	(a,b,c)	20.	(b,d)			

- 1. A-p, q; B-r, s; C-r, s; D-s
- 3. A-r; B-s; C-p, q; D-p, q
- 5. A-q, r; B-q; C-p; D-q, s
- 7. A-s; B-q, r, s; C-p; D-r
- 9. A-r, s; B-r; C-p, q; D-p, q

- 2. A-q, r; B-p, s; C-p, r; D-p, r
- 4. A-p; B-q; C-s; D-r
- 6. A-p, s; B-p, s; C-p, q, s; D-p, q, s
- 8. A-q, s; B-q, r; C-q, s; D-p, r
- 10. A-p, q, r; B-p; C-q, s; D-q

A

SINGLE CORRECT CHOICE TYPE

- 1. **(b)** LiAlH₄ does not reduce $-NO_2$ group to $-NH_2$ group.
- 2. (c) $\xrightarrow[N \ H_3 \ H_3 \ Cl^-]{heat} \xrightarrow[N \ H]{heat} +NH_4Cl$
- 3. **(b)** Amalgamated -Zn and HCl reduces carbonyl group to methylene group without affecting the double bond. However, Zn and HCl reduces -NO₂ to -NH₂ group.
- **4.** (a) In amines, N is sp^3 hybridised and thus has pyramidal shape. In the given structure, since the three alkyl groups are different, and the fourth corner of the pyramid is occupied by lone pair of electrons, the molecule is chiral. However, the two enantiomers of the amine are not resolvable because of their rapid interconversion through a transition state having planar structure (sp^2 hybridised nitrogen)
- 5. (d) The basic character of an amine in water is determined by (i) electron availability on the N atom and (ii) the extent of stabilization of the cation (protonated amine) due to solvation by hydrogen bonding

- **6. (a)** In presence of chlorobenzene, hydrogen bonding is not possible between the protonated amine and the solvent and thus the stabilization factor (solvation effect) is absent. Hence basicity is explained on the basis of the number of electron releasing groups in an amine.
- 7. (a) Note the point of difference in the given compounds which here lies at β -carbon. In I, II, III, the β -carbon atoms are sp^3 , sp^2 and sp hybridised respectively which in turn cause the difference in their s character. We know that more is the s character of an atom, greater will be its electron-withdrawing nature. Thus sp (50% s character) hybridised carbon is most electron-withdrawing, while sp^3 (25% s character) is least electron-withdrawing. Further, we know that presence of an electron-withdrawing group decreases basicity of an amine. Thus

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8. (d) Here again the two amines differ in the nature of β -carbon atom

9. (b) Greater the delocalisation of electron pair on N, lesser is its availability for protonation leading to lesser basic character.



10. (d) II is most basic because delocalisation of electron pair leads to negative charge on N making it electron-rich and hence liable to be attacked by proton very easily. Moreover, the corresponding protonated species is very much stable because of equivalent contributing structures.

$$CH_{3} - C \xrightarrow{NH_{2}} CH_{3} - C \xrightarrow{NH_{2}} CH_{3}$$

Species III is least basic because of delocalisation of electron pair on N, making it less available for protonation. Species I lies in mid-way, thus

11. (c)
$$\sum_{N=1}^{N} S_{N} > \sum_{N=1}^{N} S_{N$$

- **12. (d)** In presence of AlCl₃ (a Lewis acid), aniline is converted into anilinium cation, which being *m*-directing gives *m*-aminoacetophenone.
- 13. (c) Although aliphatic 1° amines form diazonium salts, these are unstable and decompose to alkyl carbocation which can form variety of normal as well as rearranged products.

$$\text{CH}_{3}\text{CH}_{2}\text{CH}_{2}\text{NH}_{2} \xrightarrow{\text{HONO}} \left[\text{CH}_{3}\text{CH}_{2}\text{CH}_{2}\text{N}_{2}^{+}\text{Cl}^{-} \right] \rightarrow \text{CH}_{3}\text{CH}_{2}^{+}\text{CH}_{2} + \text{N}_{2} + \text{Cl}^{-}$$

$$\text{Unstable}$$

$$CH_{3}CH_{2}CH_{2}OH \xleftarrow{H_{2}O} CH_{3}CH_{2}\overset{+}{C}H_{2} \xrightarrow{CH_{3}\overset{+}{C}HCH_{3}} \xrightarrow{Cl^{-}} CH_{3}\overset{-}{C}HCH_{3}$$

$$\stackrel{Cl}{\longrightarrow} CH_{3}CHCH_{3}$$

$$\stackrel{Cl}{\longrightarrow} CH_{3}CHCH_{3}$$

$$\stackrel{Cl}{\longrightarrow} CH_{3}CHCH_{3}$$

- 14. (b) Couplings of arenediazonium cations and phenols take place most rapidly in slightly alkaline medium because
 - (i) acidic medium (pH < 7) will suppress the ionisation of C_6H_5OH to the more reactive $C_6H_5O^-$, while the alkaline medium (pH > 7) will enhance ionisation of phenol to phenoxide ion.
 - (ii) strongly alkaline medium (pH > 10) causes the arenediazonium salt to react with the OH⁻ ion to form a relatively unreactive diazohydroxide or diazoate ion.

$$Ar\overset{+}{N} \equiv \overset{OH^{-}}{\longrightarrow} Ar - \overset{\cdot}{N} = \overset{\cdot}{N} - \overset{OH}{OH} \stackrel{OH^{-}}{\longleftarrow} Ar - \overset{\cdot}{N} = \overset{\cdot}{N} - \overset{\cdot}{O} \stackrel{\cdot}{:}$$
Diazohydroxide

Diazoate ion

15. (b) Reaction is an example of **Schmidt reaction** in which carboxylic acids are heated with hydrazoic acid in presence of a mineral acid to form primary amines.

$$RCOOH + HN_3 \xrightarrow{conc. H_2SO_4} RNH_2 + CO_2$$

16. (c) Nitrosonium ion, NO from HONO is a weak electrophile, hence it can attack only on highly activated benzene nucleus, provided proper position, *p*- or *o*- is free.

$$\begin{array}{c} H_3C \\ \searrow \dots \swarrow CH_3 \\ & \stackrel{+}{\longrightarrow} \\ & \stackrel{N(CH_3)_2}{\longrightarrow} \\ & \stackrel{NO}{\longrightarrow} \\ & \stackrel{NO}{\longrightarrow} \\ \end{array}$$

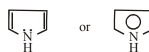
- 17. (d) LiAlH₄ does not reduce –NO₂ group.
- 18. (d) Most of the simple amines those having smaller alkyl (groups) can't be resolved, although they may contain three different alkyl groups.
- 19. (b) Hydrogen bonding is a strong intermolecular attraction between an electrophilic O–H or N–H hydrogen atom and a pair of non-bonding electrons. Thus a hydrogen bond requires both a **hydrogen bond donor** (a molecule having O–H or N–H group) and a **hydrogen bond acceptor** (a molecule having lone pair of electrons). Thus dimethyl amine can serve as hydrogen bond donor due to N–H group as well as hydrogen bond acceptor due to lone pair of electrons on N. Such molecules can form hydrogen bond with themselves and also with water (or other hydrogen bond donor). On the other hand, trimethyl amine can serve only as hydrogen bond acceptor since it has lone pair of electrons but not hydrogen on N. Hence it can form hydrogen bond with water but not with themselves.

- **20. (d)** Nearly all amines have unpleasant odour, while their salts are odourless, Amines having more than 6 carbon atoms are insoluble in water while their salts are soluble in water, hence the latter can be easily converted into syrup. Amines are susceptible to oxidation because of presence of lone pair of electrons on N.
- 21. (b) Cold dil. NaOH does not attack to either of the compound, while cold dil. HCl reacts only with benzyl amine $C_6H_5CH_2NH_2$.
- 22. (a) The four compounds differ in two respects: Compounds II has sp^3 hybridised N, compounds I and III have sp^2 while compound IV has sp hybridised N. Now we know that greater the s character of an orbital, more tightly its electrons are held and hence lesser will be their availability for protonation causing weak basic character. Thus the basic character of the three N's is $sp^3N > sp^2N > spN$. However, in pyrrole electrons on N are part of aromatic sextet, i.e., these are delocalised and hence lesser available for protonation and thus pyrrole is a weaker base than pyridine because in pyridine nonbonding electrons present in sp^2 orbital do not form a part of aromatic sextet. Thus the basic character should be in the following order.

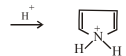
23. (b) Explained in the above question.



24.



I (Sextet present, hence aromatic) (e pair on N forms a part of aromatic sextet)



II (not aromatic) (Aromatic sextet is not present because *e* pair is involved in protonation)



or



 $\xrightarrow{\operatorname{H}^+}$



III (sextet present, hence aromatic) (e pair on N, not a part of sextet)

IV (sextet present, hence aromatic)

- **25. (b)** This is an example of Hofmann elimination which generally takes place by E2 mechanism and the latter requires a strong base (recall that OH⁻ is a strong base than I⁻). The NH₂⁻, being a strong base, can't be eliminiated easily.
- **26.** (d) All of the three are oxidation produces of a 1° amine.

- 27. (c) Cope elimination is used for removing a nitrogen present outside the ring, while Hofmann elimination is used for removing nitrogen present inside as well as outside the ring.
- 28. (d) All the three react with HONO forming following respective products.

$$\begin{array}{ccc} C_6H_5 \stackrel{+}{N} \equiv NCl^- \\ \text{Diazonium chloride} \end{array} \qquad \begin{array}{cccc} C_6H_5 \stackrel{\cdots}{N} - N = O \\ & & \\ CH_3 \\ & & \\ 2^{\circ} \text{ N-Nirosoamine} \end{array}$$

p- Nitrosodimethylamine

29. (d)
$$RNH_2 + C = O \longrightarrow C \xrightarrow{OH} \xrightarrow{-H_2O} C = NR$$

1° Amine Aldehyde/ketone Carbinolamine Imine (Schiff's base (Schiff's base))

- 30. (c) Pyridine resembles a strongly deactivated benzene, here the deactivation is due to electron-withdrawing effect of the electrongative N atom from 2-and 4 positions making 3-positions to be more vulnerable for the attack of an electrophile. Remember that the nonbonding electrons on N are perpendicular to the π-system (they are present in sp² orbital), these are not involved in delocalisation, i.e., they can't stabilize the positively charged intermediate.
- **31. (a)** Due to greater electronegativity of N over S, positive charge on N will make the methyl groups more electron-deficient than that on positively charged S. Therefore, $(CH_3)_4 N^+I^-$ will undergo nucleophilic substitution more readily than the sulphur analogue.

$$HO \xrightarrow{} + CH_3 \rightarrow N^+(CH_3)_3 \longrightarrow CH_3OH + (CH_3)_3 N$$

32. (b) It is an example of Beckmann rearrangment; where the group *anti* to -OH migrates

33. (d) For this one should remember that Hofmann's degradation involves migration to electron deficient nitrogen atom, hence the alkyl group will migrate with its bonding pair of electrons.

34. (c) Remember that in Hofmann rearrangement, the two original H atoms of the –CONH₂ group are removed by base (OH⁻) and new H's are derived from H₂O.

$$(CH_3)_3C - \overset{O}{C} - NH_2 \xrightarrow{(i)OD^-/Br_2} (CH_3)_3C - ND_2$$

35. **(d)**
$$HCCl_3 \xrightarrow{OH^-} : \bar{C}Cl_3 \xrightarrow{-Cl^-} : CCl_2$$
Dichlorocarbene
(a neutral electron deficient species)

36. (c) The given amine has three β -hydrogens, hence it can form three alkenes of which least substituted (H₂C=CH₂) will be the major product (Hofmann product or Hofmann elimination).

$$\beta_{2} \qquad \beta_{1} \qquad (i) \text{ CH}_{3} \\ \beta_{3} \qquad (ii) \text{ Ag}_{2} \\ \text{NHCH}_{2} \text{CH}_{3} \qquad (iii) \text{ heat} \qquad (from \beta_{1}) \qquad + \text{ (from } \beta_{2}) \qquad (from \beta_{3})$$

In Hofmann elimination H atom (in the form of proton) is eliminated from that β carbon atom which has maximum number of H atom(s). This is due to the fact that Hofmann elimination (an example of E2 reaction) requires anti-coplanar arrangement of the β H atom and the leaving group. Further, higher the chances for this arrangement to remain in staggered conformation more will be the ease of elimination. Thus greater the number of H atom at β position, higher will be the chances for Me₃N⁺– and H atom to remain in this particular conformation (anti-coplanar arragnement of Me₃N⁺ – and H atom in staggered conformation).

37. **(b)**
$$CH_3$$
 CH_3 CH_3

38. (d) The reaction involves Cope eliminiation (heating of a 3° amine oxide to form an alkene with the elimination of a 2° hydroxylamine).

$$(CH_3)_2 \xrightarrow{H_2O_2} \beta \xrightarrow{\text{heat}} + \text{HON } (CH_3)_2$$

$$+(CH_3)_2 \text{NOH}$$

$$3^{\circ} \text{ Amine oxide (acts its own base)}$$

$$+(CH_3)_2 \text{NOH}$$

$$+(CH_3)_2 \text{NOH}$$

39,40. (c), (b)
$$R - C - CH = N$$

$$\begin{array}{c} O \\ \parallel \\ N \end{array} = N \\ \hline \begin{array}{c} -N_2 \\ -N_2 \end{array} \longrightarrow \begin{array}{c} O \\ \parallel \\ C - CH \end{array} \longrightarrow \begin{array}{c} O \\ \parallel \\ C = CHR \end{array}$$
Carbene (neutral)

Ketene

41. (c)
$$(CH_2NO_2)$$
 (CH_2NO_2) $(CH_2NO_$

42. (d) All the three reactions lead to the formation of C = C bonds under usual conditions.



43. (c) (a)
$$R - C - NH_2 \xrightarrow{Br_2/NaOH} R - NH_2$$
 (Hofmann reaction)

(b)
$$R - C - N_3 \xrightarrow{heat} R - NH_2$$
 (Curtius reaction)

(c)
$$R - C - OH + HN_3 \xrightarrow{H_2SO_4} R - NH_2$$
 (Schmidt reaction)

O
$$\parallel$$
(d) $R - C - NHOH \xrightarrow{OH^-} R - NH_2$ (Lossen reaction)

Hydroxamic acid

- **44. (b)** R N = C = O (Alkyl isocyanate) is the common intermediate in the four reactions.
- **45. (d)** In other three, the group migrates to electron deficient carbon.
- **46. (d)** All the three are formed

47. (c)
$$CH_3NH_2 \xrightarrow{CH_3I} (CH_3)_4N^+I^- \xrightarrow{AgOH} (CH_3)_4N^+OH^- \xrightarrow{heat} (CH_3)N + CH_3OH$$
(C) $(CH_3)H_2 \xrightarrow{CH_3I} (CH_3)_4N^+I^- \xrightarrow{AgOH} (CH_3)_4N^+OH^- \xrightarrow{heat} (CH_3)N + CH_3OH$
(C) $(CH_3)H_3 \xrightarrow{CH_3I} (CH_3)_4N^+I^- \xrightarrow{AgOH} (CH_3)_4N^+OH^- \xrightarrow{heat} (CH_3)N + CH_3OH$

48. (c)
$$CH_3COOH \xrightarrow{(i) LiAlH_4} CH_3CH_2Br \xrightarrow{(i) KCN} CH_3CH_2CH_2NH_2 \xrightarrow{CHCl_3,OH^-}$$

$$\text{CH}_{3}\text{CH}_{2}\text{CH}_{2}\text{NC} \xrightarrow[(+2\text{H}_{2}\text{O})]{\text{hydrolysis}} \text{CH}_{3}\text{CH}_{2}\text{CH}_{2}\text{NH}_{2} + \text{HCOOH}$$

49. (d)
$$\stackrel{\text{OH}}{\longrightarrow}$$
 $\stackrel{\text{NH}_2\text{OH}}{\longrightarrow}$ $\stackrel{\text{heat}}{\longleftarrow}$ $\stackrel{\text{conc. H}_2\text{SO}_4}{\longleftarrow}$ $\stackrel{\text{(Beckmann rearrangement)}}$

syn and anti-ketoximes

50. (c) The reaction is an example of hydrazinolysis.

$$\begin{array}{c|c}
O \\
\parallel \\
C \\
NH \\
C \\
NH
\end{array} + H_2NR$$

B E COMPREHENSION TYPE

- 1. **(b)** E2 eliminations are concreted and *anti*-stereoselective.
- 2. (d) Due to the bulk of the $-N^+Me_3$ group, the OH⁻ preferentially approaches and removes the least hindered β -hydrogen and thus form the least substituted alkene as the major product (difference from Saytzeff product).
- **3. (b)** Species II (acylnitrene) is electron-deficient (N has only 6 electrons), hence it has a tendency to get its octet completed by the migration of alkyl group.
- **4. (c)** -OCH₃ is more electron-releasing hence when the migrating aryl group has -OCH₃ in the para postion, its migration is accelerated.
- 5. (a) The migrating group (the group attached to C of the -CONH₂ part) never becomes free, so optical activity of the reactant is maintained.
- 6. (c) Since phenyl group (migrating group) does not become free, no cross product [i.e., (ii) and (iv)] is formed.

7-8-9.(c-c-d) Reaction (i)
$$\stackrel{\text{NHCOCH}_3}{\longleftrightarrow} \stackrel{\text{NHCOCH}_3}{\longleftrightarrow} \stackrel{\text{NHCOCH}_3}{\longleftrightarrow} \stackrel{\text{NHCOCH}_3}{\longleftrightarrow} \stackrel{\text{NH}_2}{\longleftrightarrow} \stackrel{\text{NH}_2}{\longleftrightarrow} \stackrel{\text{NH}_2}{\longleftrightarrow} \stackrel{\text{NH}_2}{\longleftrightarrow} \stackrel{\text{NH}_3}{\longleftrightarrow} \stackrel{\text{NH}_3}{\longleftrightarrow}$$

10. (a) Going backward, we can obtain answer

11. (d)
$$O_2$$
 O_2 O_3 O_4 O_3 O_4 O_4 O_4 O_5 O_7 O_8 O_8 O_9 $O_$

This step involves migration of phenyl group to electron-deficient oxygen.

12. (d)
$$C_6H_5 - N = C = O \xrightarrow{C_2H_5OH} C_6H_5 - NHCOOC_2H_5$$
Urethane

13. (c)
$$C_6H_5 - C - N = N = N : \xrightarrow{-N_2} C_6H_5 \xrightarrow{O} N : \xrightarrow{Nitrene} C = N - C_6H_5$$

C REASONING TYPE

- 1. (c) The correct reason is: -SO₃H group, being more acidic than -CO₂H group, can easily transfer a proton to the amino group.
- 2. (c) The correct reason is : Acetylation decreases the electron-density in the benzene ring thereby preventing oxidation.



- 3. (c) The correct reason is: The overall electron deficiency in m-nitroaniline is much less (due to -R-effect of NO₂ group and +R-effect of NH₂ group) than in m-dinitrobenzene (-R-effect of the two NO₂ groups) and hence does not accept additional electrons from a weak reducing agent such as (NH₄)₂S and thus further reduction is prevented.
- **4. (d)** Aryl halides (chlorobenzene) do not undergo nucleophilic substitution with KCN because of the low reactivity of the Cl atom, which is because of resonance in chlorobenzene. So assertion is wrong. Reason is correct.
- 5. (a) R is the correct explanation of A.
- 6. (d) In strongly acidic conditions, aniline becomes protonated with the result lone pair of electrons is not available to produce
 - +E and +M effects. On the other hand, the $-\stackrel{\oplus}{NH_3}$ group exerts strong –I effect causing deactivation of the ring.
- 7. (c) Correct R: Only primary aliphatic amines can be prepared by Gabriel phthalimide reaction.
- 8. (a) R is the correct explanation of A.
- 9. (a) R is the correct explanation of A.

D ■ MULTIPLE CORRECT CHOICE TYPE ■

- (a,b,d) Tertiary amino group is a strong activator, nitration can be carried out under milder conditions (HNO₃ + CH₃COOH).
 In Tertiary amines, electron pair on N is shielded by two bulky methyl groups, so these are not easily oxidised by mild oxidising agents.
- (a,c,d) Diazomethane is used for methylating acidic groups; compound IV has enolic —OH group, hence it can also be methylated by CH₂N₂.
- **3. (a,b)** For the preparation of Me₃CNH₂, the required alkyl halide is Me₃CX which will react with potassium phthalimide, a strong base, to form alkene rather than substituted product. For preparing C₆H₅NH₂, C₆H₅Cl will be the starting halide in which Cl is non-reactive.
- **4. (b,c,d)** Vinyl and aryl halides do not undergo S_{N^2} reactions, unless activated by electron withdrawing group (in the *o* and *p*-position in case of aryl halides). 3° Halides form alkenes on elimination.
- 5. (a,b,d) In reductive amination method, products can have only CH_3 , a 1°C, or 2°C bonded to N as in CH_3NH_2 , RCH_2NH_2 or R_2CHNH_2 respectively. In the reduction of oxime, amine formed can have only 1° or 2°C bonded to N. In Gabriel phthalimide method, the required Me_3CCl would undergo elimination (E2) rather than substitution (S_{N_2}).
- **6.** (a,b,d) Isonitriles (C₆H₅NC) on reduction give 2° amines (C₆H₅NHCH₃). All other three methods give aniline.
- 7. (a,c,d) In *p*-NH₂C₆H₄COOH, —COOH group is very weak so it can't transfer H⁺ to the weakly basic amino group. All other three form zwitterions.
- **8. (b,c,d)** In C₆H₅OCH₃, —OCH₃ does not sufficiently increase electron density on the ring. Recall that C₆H₅OH undergoes coupling in weakly alkaline medium which converts C₆H₅OH to the more reactive C₆H₅O⁻. In options (*b*) and (*c*), presence of electron-withdrawing —NO₂ groups increases electrophilic character to such an extent that these diazonium cations can couple even with the compounds having weak electron-releasing groups. Option (d) undergoes coupling reaction easily because –NMe₂ is sufficiently electron-releasing.
- **9. (a,b,d)** In *p*-aminophenol all the four positions (2, 3, 5 and 6) can be coupled (positions 2 and 6 in presence of H⁺ and positions 3 and 5 in presence of OH⁻).
- 10. (a,b,d) *tert*-Nitro compounds (Me₃CNO₂) do not react with HONO because they do not have any α -H. The three others react with HONO as usual.

$$\begin{array}{c} \text{NO}_2 \\ \text{CH}_3 - \text{CHCH}_2\text{CH}_3 \\ \text{2-Nitrobutane} \end{array} \xrightarrow{\begin{array}{c} \text{HONO} \\ \text{HONO} \end{array}} \begin{array}{c} \text{NO}_2 \\ \text{CH}_3 - \text{C} - \text{CH}_2\text{CH}_3 \\ \text{NO} \\ \text{Pseudonitrol (blue)} \end{array}$$

11. (c,d) — SO_3H group present in o- and p-positions are easily replaced.

12. (a,b,c)
$$C_6H_5NH_2 + AlCl_3 \longrightarrow C_6H_5N^+H_2 \xrightarrow{CH_3Cl} No reaction$$
Substituent $(-NH_2AlCl_3)$
a meta director

However, Friedel-Crafts reactions are the most sluggish electrophilic aromatic substitutions (halogenation occurs most easily), hence if a meta director is present on the ring, the compound does not undergo Friedel-Crafts alkylation/acylation.

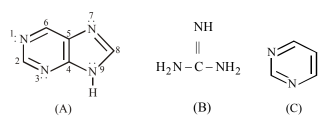
E

MATRIX-MATCH TYPE

1. A-p, q; B-r, s; C-r, s; D-s

- (A) Hydrogen bodning is possible in *ethyl acetoacetate* (due to the presence of enolic group) and in *proteins* (due to the presence of –CO–NH– grouping).
- (B) β -Amino acids, like α -amino acids, form *zwitterions* and also undergo *diazotisation* although the diazo salts, so formed, are unstable.
- (C) p-Amino benzoic acid, like β and α -amino acids forms *zwitterions* and being 1° aromatic amine also forms, well-known, *diazonium salt*.
- (D) Aniline undergoes diazotisation easily.

2. A-q, r; B-p, s; C-p, r; D-p, r



- (A) A is aromatic due to 10π electrons, N_1 , N_3 and N_7 are sp^2 hybridised their ℓp present in sp^2 orbital are localised hence undergo protonation. The N_9 is also sp^2 hybridised but its ℓp is in p-orbital hence involved in delocalisation, and thus not basic.
- (B) Due to resonance, all the three nitrogens of guanidine are identical. It is not cyclic, hence non-aromatic.
- (C-D) As mentioned above in (A), here the two nitrogens are identical, and the compound is aromatic due to the presence of aromatic sextet.

3. A-r; B-s; C-p, q; D-p, q

(A) $CHCl_3 \xrightarrow{OH^-} : CCl_2 + HCl$ (α -elimination)

(B) $CHCl_2 - CF_3 \xrightarrow{OC_2H_5^- \text{ fast}} Cl_2 \xrightarrow{C} - CF_3 \xrightarrow{Slow}$

 $Cl_2C = CF_2$

This reaction, although involves two steps; the rate determining step involves only one molecule hence the reaction is labelled as E1. Further, it is the carbanion (conjugate base of CHCl₂CF₃) that determines the rate of reaction, hence it is labelled as E1CB.

(C) When the proton to be removed is on the sterically hindered carbon, and the base also has sterically hindered carbon, less substituted alkene is the main product (*Hofmann elimination*)

$$CH_{3} \xrightarrow{CH_{3}} CH_{3} \xrightarrow{He_{3}CO^{-}} CH_{2} = C - CH_{2}CH_{3}$$

$$Rr \xrightarrow{Br} Major$$

(D) Elimination from quaternary ammonium ions usually gives the least substituted alkene (*Hofmann elimination*).

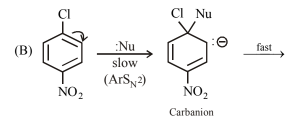
$$\begin{array}{c}
N^{+}Me_{3} \\
| CH_{3} - CHCH_{2}CH_{3} \xrightarrow{OH^{-}} CH_{2} = CHCH_{2}CH_{3} + Me_{3}N + H_{2}O
\end{array}$$

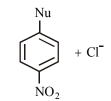
4. A-p; B-q; C-s; D-r

(A) ArS_{N1} reactions are rare, and involve carbocation as intermediate. The inportant example is substitution in aromatic diazonium salts.

$$ArN^{+} \equiv N \xrightarrow{slow} Ar^{+} + N_{2}$$
Highly stable Highly stable

$$Ar^+ + : Nu \xrightarrow{fast} Ar - Nu$$





(C) Higher the bulk of the group present on the nucleophilic centre, lesser is its nucleophilic character.

$$^{-}$$
O $-$ H $>$ $^{-}$ O $-$ C $($ CH $_{3}$) $_{3}$ bulkyl group

(D) Stronger the conjugated acid, weaker will be the base.

$$H_2O > NH_3$$
 stronger acid weaker acid

5. A-q, r; B-q; C-p; D-q, s

- (A) Allylic rearrangements involve the migration of the double bond (and the functional group, if present) from one position to other. These reactions proceed through SN¹, SN², SE¹ and SE² mechanism; SN¹ involves carbocations (electron-deficient species), while SE¹ involves carbanion.
- (B) Hofmann rearrangement involves nitrene
 O
 || ...

(R-C-N:) intermediate in which nitrogen is electron deficient.

- (C) Aldol condensation involves the attack of base on aldehyde or ketone, the reaction will involve the formation of carbanion.
- (D) Carbylamine reaction involves the formation of dichlorocarbene (:CCl₂) as intermediate which is electron deficient.