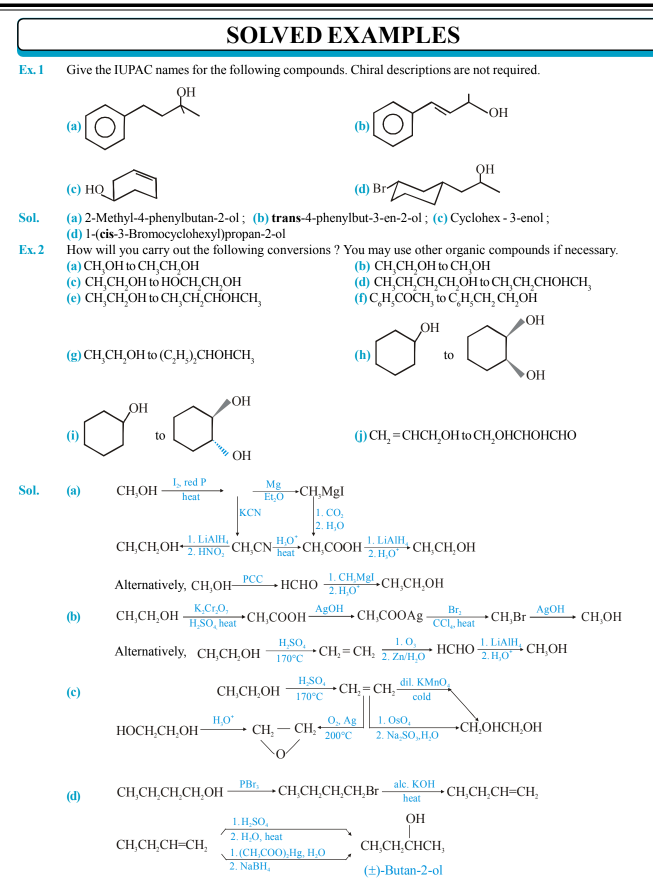
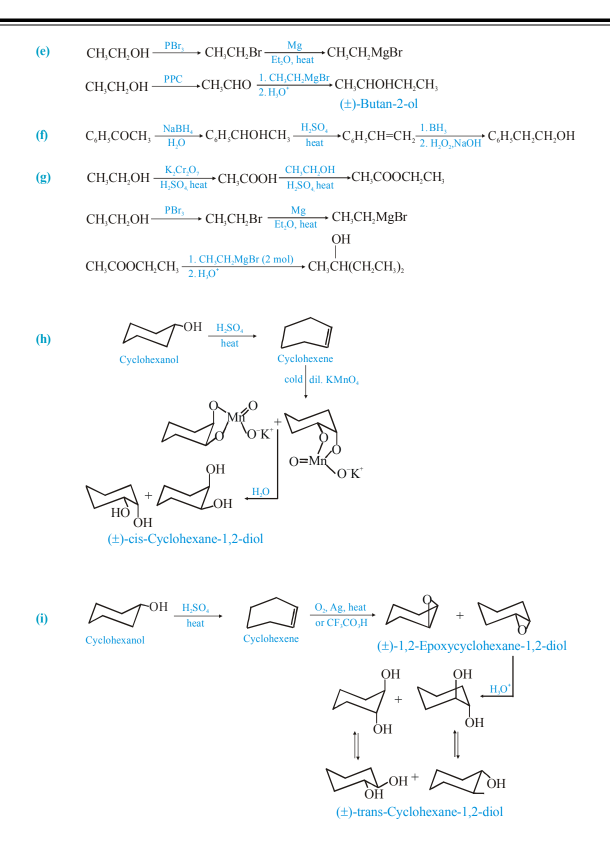
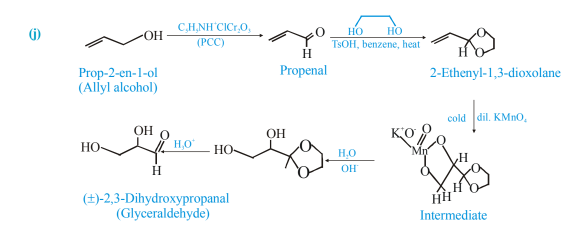


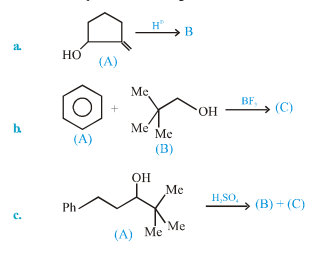
ALCOHOL PHENOL AND ETHER





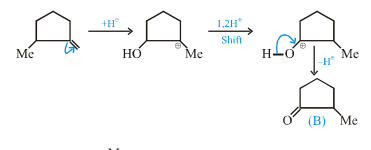


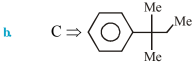
Ex.3 Complete the following reactions :



Sol.
$$B \Rightarrow \int_{Me}^{Me}$$

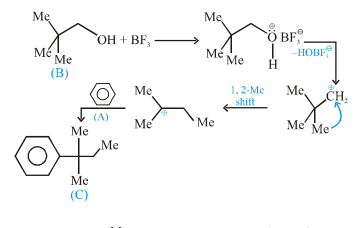
Mechanism :

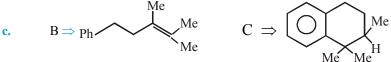




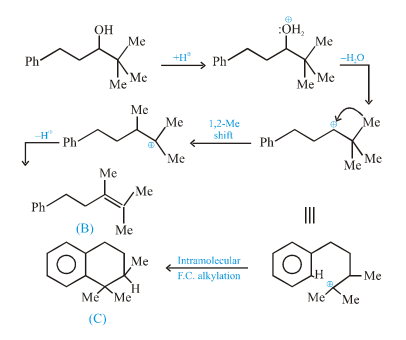
117

Mechanism :

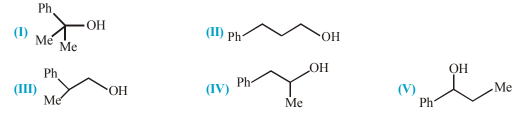


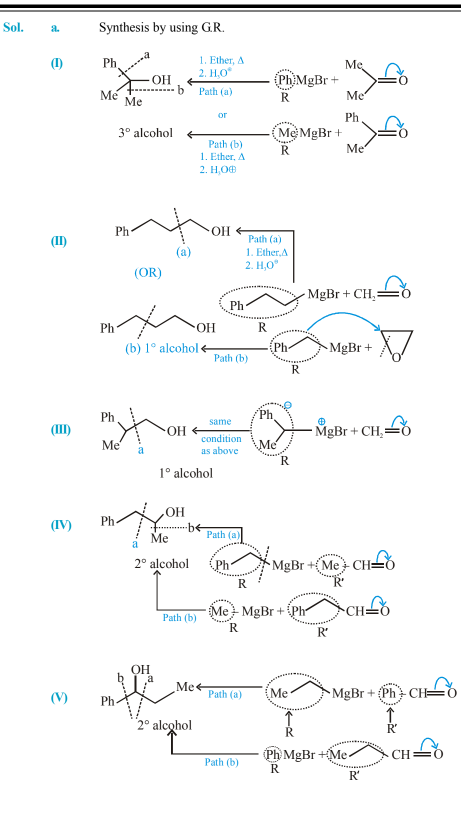


Mechanism :



Ex.4 Synthesise the following alcohols by using Grignard reagent (G.R.)





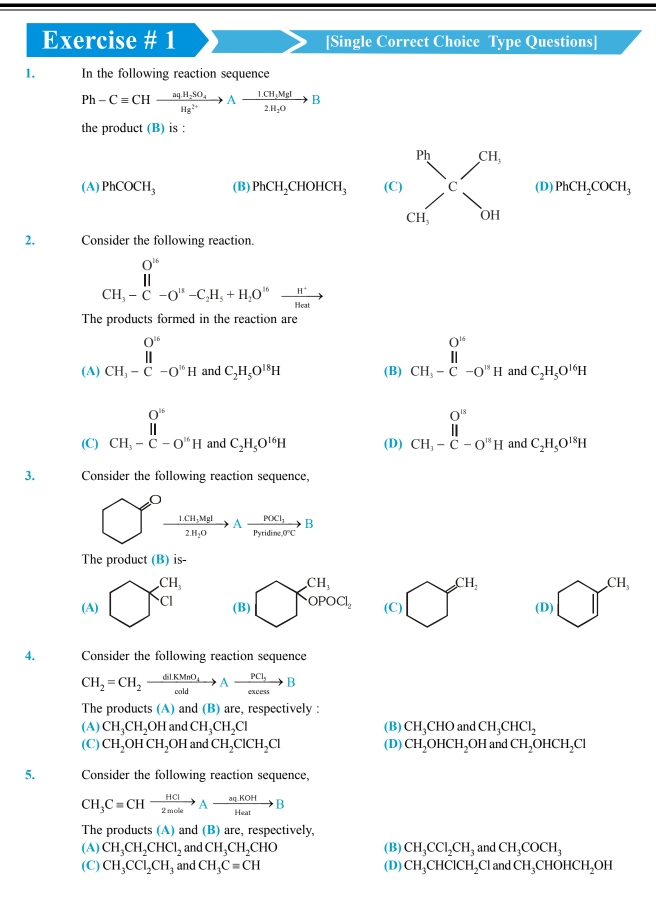
119

(a) $Cl_{2}/light (1eq.)$ (b) $Cl_{2} (excess) / hv$ (c) $Dil. H_{2}SO_{4} then H_{2}O$ (d) $Conc. H_{2}SO_{4} then heat$ (e) $(1) CH_{3}COCl$ $(2) AlCl_{3}$ (f) $(1) (CH_{3}CO)_{2}O$ $(2) AlCl_{3}$ (g) $CH_{3}-CH_{2}-O-CH_{2}-CH_{3} \xrightarrow{Cl_{4}/light} CH_{3}-CH-O-CH-CH_{3}$ Cl Cl Cl Cl Cl Cl(h) $CH_{3}-CH_{2}-O-CH_{2}-CH_{3} \xrightarrow{Cl_{4}(excess)} CCl_{3}-C-O-C+CCl_{3}$ (c) Cl Cl(c) $Cl_{3}-CH_{2}-O-CH_{2}-CH_{3} \xrightarrow{Cl_{4}(excess)} CCl_{3}-C-O-C+CCl_{3}$ (c) $Cl_{3}-CH_{2}-O-CH_{2}-CH_{3} \xrightarrow{dil. H_{3}SO_{4}} CH_{3}-CH_{2}-OH + CH_{3}-CH_{2}-HSO_{4} \xrightarrow{H_{4}O} CH_{3}-CH_{2}-OH$ (d) $CH_{3}-CH_{2}-O-CH_{2}-CH_{3} \xrightarrow{conc. H_{3}SO_{4}} CH_{3}-CH_{2}-OH + CH_{3}-CH_{2}-HSO_{4} \xrightarrow{A} CH_{2}=CH_{2}$

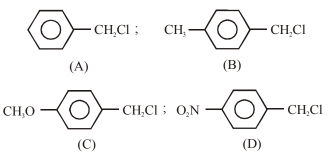
Complete the following reactions of dimethyl ether with the following reagents :

- (e) $CH_3-CH_2-O-CH_2-CH_3 \xrightarrow{(1) CH_3COCl} CH_3-CH_2-Cl+CH_3-COOCH_2-CH_3$
- (f) $CH_3-CH_2-O-CH_2-CH_3 \xrightarrow{(1) CH_3(CO)_2O} 2CH_3-COOCH_2-CH_3$

Ex.5



6. Consider the following chlorides :



The order of reactivity of A, B, C and D towards hydrolysis by S_N1 mechanism is : (A) A < B < C < D(B) D < C < B < A(C) D < A < B < C(D) C < B < A < D

7. Propene is allowed to react with m-chloroperoxobenzoic acid. The product (A) is then reduced with $LiAlH_4$ in dry ether to give (B).

(D) CH₃-

$$CH_{3}CH = CH_{2} \xrightarrow{MCPBA} A \xrightarrow{1.LiAlH_{4}} B$$

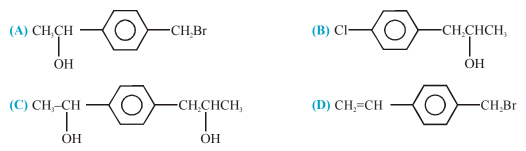
The structure of the product (B) is:
(A) CH_{3}CHOHCH_{2}OH (B) CH_{3}CH_{2}CH_{2}OH (C) CH_{3}CHOHCH_{3}

8. 2-Phenylethanol may be prepared by the reaction of phenylmagnesium bromide with :

(A) HCHO (B)
$$CH_3CHO$$
 (C) CH_3COCH_3 (D)

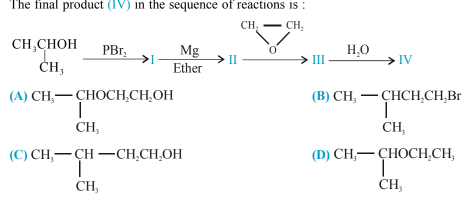
9. In the reaction

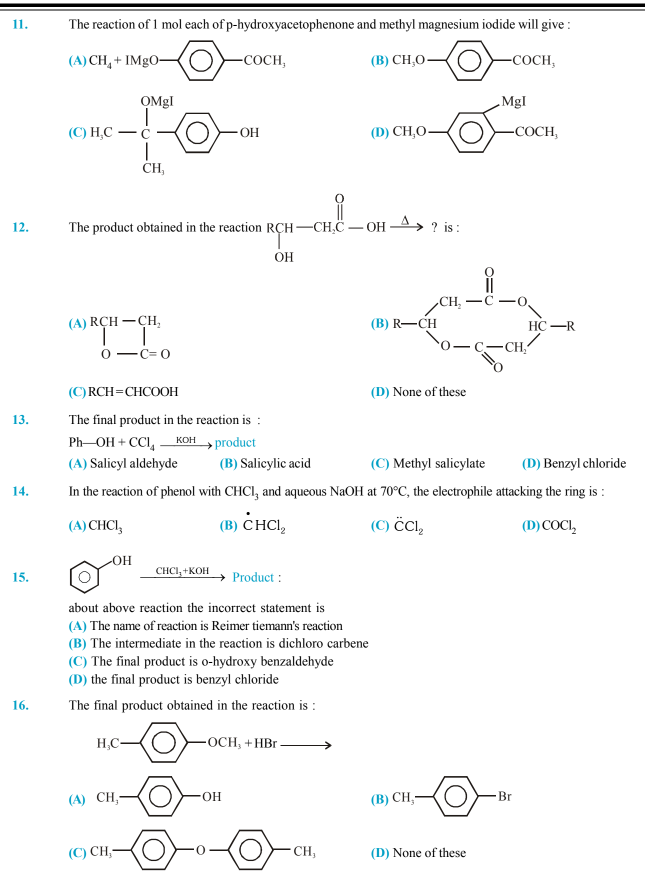
$$Cl \longrightarrow CH_2Br + CH_3CHO + Mg \xrightarrow{dry ether} A \xrightarrow{H_3O^+} B$$



10.

The final product (IV) in the sequence of reactions is :



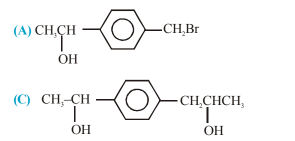


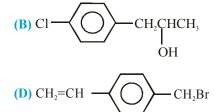
123

- 17.The acidity of the compounds RCOOH, H_2CO_3 , C_6H_5OH , ROH decreases in the order :(A) RCOOH > $H_2CO_3 > C_6H_5OH > ROH$ (B) $C_6H_5OH > RCOOH > H_2CO_3 > ROH$ (C) ROH > $C_6H_5OH > RCOOH > H_2CO_3$ (D) $H_2CO_3 > RCOOH > C_6H_5OH > ROH$
- 18. Arrange the following in order of decreasing acidic strength. p- nitrophenol (I), p-cresol (II), m-cresol (III), phenol (IV):
 (A) I>II>II>III>IV
 (B) IV>III>II>I
 (C) I>III>II>IV
 (D) III>II>I
 - (A) 12 II 2 III 2
- **19.** In the reaction

Cl — CH₃Br + CH₃CHO + Mg
$$\xrightarrow{\text{Dry ether}}$$
 A $\xrightarrow{\text{H}_3\text{O}^+}$

the product (B) is :



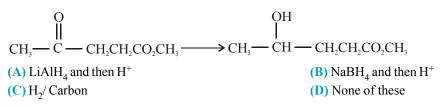


В

20. The products formed in the reaction are :

$$C_{6}H_{5} - C - OH + CH_{3}OH \xrightarrow{H_{2}SO_{4}}_{Heat} \rightarrow$$
(A) $C_{6}H_{5} - C - OCH_{3}$ and $H_{2}O$
(B) $C_{6}H_{5} - C - OCH_{3}$ and $H_{2}O$
(C) $C_{6}H_{5} - C - CH_{2}OH$ and $H_{2}O$
(D) $C_{6}H_{5}OCH_{3}$, CO and $H_{2}O$

21. The conversion



(B) CH₃CH₂OH

22. Identify the final product of the reaction :

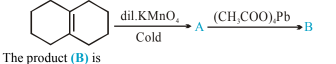
 $CH_3MgBr + CH_2CH_2 \longrightarrow H_3O^+$

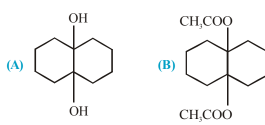
(A) CH₃OH

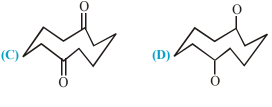


 $\textbf{(D)} CH_3 CH_2 CH_2 OH$

- For the cleavage of ethers by halogen acids, the order of reactivity of halogen acids is : 23. (A) HI > HBr > HCl(B) HBr>HI>HCl (C) HCl>HBr>HI (D) Ethers do not undergo cleavage
- 24. Consider the following reactions :

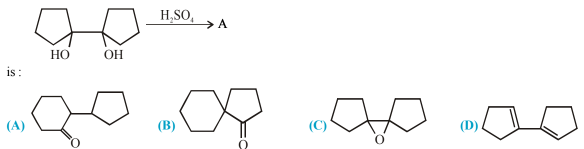






25.

The major product (A) formed in the reaction



26. Decreasing order of relative nucleophilicity of the following nucleophiles in protic solvent is -

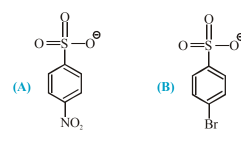
 $H_{2}O$

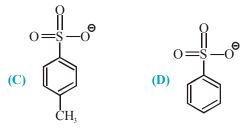
- $\overset{\Theta}{SH}, \quad \overset{\Theta}{AcO}, \quad \overset{\Theta}{PhO}, \quad \overset{\Theta}{OH},$ (A) $\overset{\Theta}{SH} > \overset{\Theta}{OH} > H_2O > A_{CO} \overset{\Theta}{O} > PhO$ (B) $\overset{\Theta}{SH} > \overset{\Theta}{OH} > PhO > A_{CO} \overset{\Theta}{O} > H_2O$ (C) $\overset{\otimes}{SH} > Ph\overset{\otimes}{O} > \overset{\otimes}{OH} > H_2O > Ac\overset{\otimes}{O}$ (D) $\overset{\otimes}{OH} > \overset{\otimes}{SH} > Ph\overset{\otimes}{O} > Ac\overset{\otimes}{O} > H_2O$
- 27. Electrophiles are (A) Electron deficient species (C) Electron rich species

- (B) having atleast one pair of electron (D) having vacant p or d-orbital
- Which of the following is an electrophilic reagent? 28. (A) H₂O (B) OH⁻

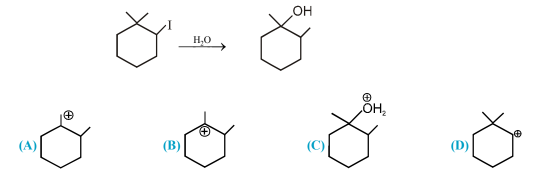
(C)
$$NO_2^+$$
 (D) None

29. The best leaving group is :

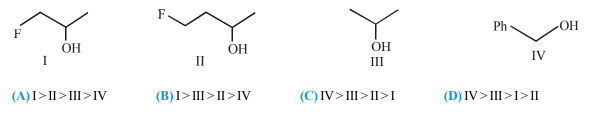




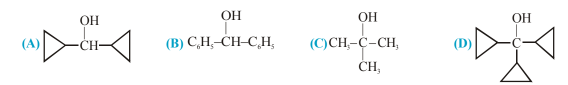
30. Which of the following is not expected to be intermediate of the following reaction ?



31. The correct order of reactivity of following alcohols towards conc.HCl/ZnCl₂ is :



32. Which one of the following will be most reactive for S_N^1 reaction?



33. Which describes the best stereochemical aspects of the following reaction ?

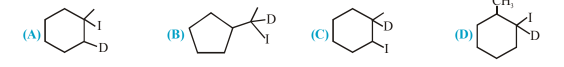
$$\begin{array}{c} CH_{3} \\ \hline \\ Ph \\ OH \end{array} \xrightarrow{H-Br} Product$$

(A) Inversion of configuration occurs at the carbon undergoing substitution.

- (B) Retention of configuration occurs at the carbon undergoing substitution.
- (C) Racemization occurs at the carbon undergoing substitution.
- (D) The carbon undergoing substitution is not stereogenic.

34.
$$(H_3)$$
 (H_3) (H_3)

Identify the major product :



 $CH_3(CH_2)_2 - CH_2OH \xrightarrow{\text{NaBr, } H_2SO_4} X,$ 35. 1-butanol Identify X and the mechanism of the reaction (A) $CH_3 - CH_2 - CH_2 - CH_2 - Br \& S_N 1$ **(B)** $CH_3 - CH_2 - CH_2 - CH_2 - Br \& S_N 2$ (D) $CH_3 - CH - CH_2 - CH_3 \& S_N^2$ | Br (C) $CH_3 - CH - CH_2 - CH_3 \& S_N 1$ Br $\xrightarrow{\text{SOCl}_2} \text{Product}$ 36. СН'ОН (X) Product of the above reaction is : **(B) (C) (D)** CH, $- \text{OH} \xrightarrow{\text{PCl}_5} X$; (X) is 37. Н- C_2H_5 (D-2-Butanol) (A) S-2-Chlorobutane (B) R - 2-Chlorobutane (C) Mixture of R and S 2-Chlorobutane (D) 1-Chlorobutane $\xrightarrow{\text{PBr}_3} (X), X \text{ is : } (X)$ 38. (B) **(C) (D)** $SOCl_2 \rightarrow Product$ 39. H₋C CH₂ Identify the product Cl (B) Cl $C_{6}H_{5}$ (C) HS $C_{6}H_{5}$ (D) H₅C₆ H_{5} (D) H₅C₆ (D) H₅ (D) H₅C₆ (D) H₅C₆ (D) H₅ (D) H₅C₆ (D) H₅ (D) H₅C₆ (D) H₅ (A) H₅C₆

OH

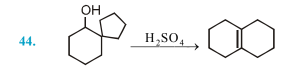
SH

Consider the following reaction. **40.** OH Cl $CH_3 - CH_2 - CH - CH_3 \xrightarrow{SOCl_2} CH_3 - CH_2 - CH - CH_3$ In the above reaction which phenomenon will take place : (B) Retention (A) Inversion (C) Racemisation **(D)** Isomerisation 41. In the given reaction : $CH_3 - CH_2 - CH_2 - O - CH_2 - CH_3 \xrightarrow{HCI/\Delta} [X] + [Y]$ [X] and [Y] will respectively be : (A) $CH_3 - CH_2 - CH_2OH \& CH_3 - CH_2 - CI$ (B) $CH_3 - CH_2 - CI \& CH_3 - CH_2 - OH$ (C) CH_3 - CH_2 - CH_2 - $CI \& CH_2$ = CH_2 (D) CH_3 -CH= $CH_2 \& CH_2$ = CH_2 $CH_{3} \xrightarrow{CH_{3}} CH_{2} \xrightarrow{EtO^{-}K^{-}/EtOH} X$ 42. The product X and Y are respectively : CH₃ CH₃ CH₃ CH₃ (A) $CH_3 - C - CH_2OCH_2CH_3$ & $CH_3 - C - CH_2OH$ (B) $CH_3 - C - CH_2OH$ & $CH_3 - C - CH_2OCH_2CH_3$ ЬĤ OCH₂CH₃ ÓН OCH₂CH₃ CH, CH, CH₃ CH. (C) $CH_3 - C - CH_2OH \& CH_3 - C - CH_2OCH_2CH_3$ (D) $CH_3 - C - CH_2OCH_2CH_3 \& CH_3 - C - CH_2OH_3$ ÓН ÓН ÓН ÔН 43. The relative rate of acid catalysed dehydration of following alcohols would be, CH₃ CH₃ $CH - CH - CH_3$ $\dot{C}H - CH_2 - CH_2 - OH$ OH Ι Π CH₃ CH₃ $\dot{C} - CH_2 - CH_3$ Ċ – CH₂OH ЬĤ CH₃

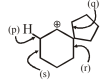
(A) III > I > IV > II (B) III > IV > I > II (C) I > III > IV > II (D) III > IV > I > II

Ш

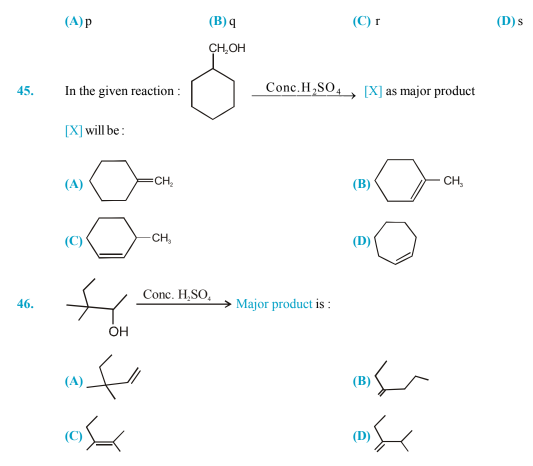
IV



In the above reaction the major product is shown, which is formed through the intermediate (carbocation) given below :

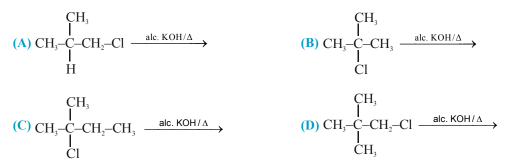


Which bond will migrate to form the above product ?





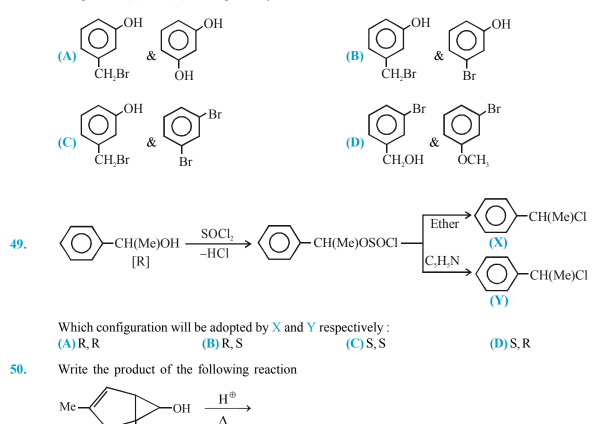
. In which of the following reaction, regioselectivity can be observed.

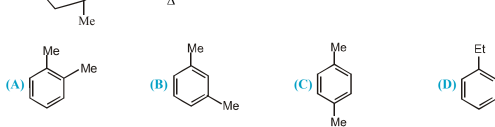




$$\bigcup_{\substack{\text{CH,OH}}} \stackrel{\text{OH}}{\longrightarrow} \stackrel{\text{HBr}}{\longrightarrow} (\mathbf{A}), \quad \bigcup_{\substack{\text{OCH}}} \stackrel{\text{OH}}{\longrightarrow} \stackrel{\text{HBr}}{\longrightarrow} (\mathbf{B})$$

The product (A) and (B) are respectively :





51. When (1R, 2R)-1, 2-Dibromo-1, 2-diphenyl ethane is treated with alcoholic solution of KOH, the most probable product would be :

(A) trans-1, 2-diphenyl ethene (C) cis-alkenyl bromide

(B) A mixture of cis, trans alkenyl bromide

(D) trans-alkenyl bromide

52.
$$CH_3 - CH_2 - CH_2 - CH_3 - CD_3 \xrightarrow{H_3PO_4} Total no. of alkenes (X) \xrightarrow{Fractional} No. of Fractions obtained (Y)$$

 $CH_3 OH$
X and Y are :
(A) 6,6 (B) 9,5 (C) 9,6 (D) 6,4

(D) H₃O[⊕]

53. Number of fractions on fractional distillation of solvolysis in ethanol of 1-chlorobut-2-ene is : **(A)**1 **(B)**2 **(C)**3 **(D)**4

54.
$$CH_3CH_2OH + HCl \xrightarrow{ZnCl_2} CH_3CH_2Cl + H_2O$$

In the above reaction, the leaving group is :
(A) HO^{Θ} (B) H_2O

(B) H₂O

- 55. Which of the following statement is not true?
 - (A) Nucleophiles possess unshared pairs of electron which are utilized in forming bonds with electrophilic substrate.
 - (B) The cyanide ion is an ambident nucleophile and causes nucleophilic substitution of alkyl halide by either of its carbon atom or nitrogen atom.
 - (C) The nitrite ion is an ambident nucleophile and causes nucleophilic substitution of alkyl halide by either of its oxygen atoms or nitrogen atom.
 - (D) Strength of Nucleophile generally decreases on going down a group in the periodic table.

56.
$$Cl-CH_2CH_2CH_2CH_2OH \xrightarrow{NaOH(aq.)} A$$
, the product A is :
(A) HO-CH_2CH_2CH_2CH_2OH
(B) $\bigcirc \bigcirc \bigcirc \bigcirc \bigcirc \bigcirc$
(C) Cl-CH_2-CH-CH_2-CH_2OH
(D) ClCH_2CH_2CH_2CH_2ONa

57. Which of the following reactions is the best choice for preparing methyl cyclohexyl ether?

(A)
$$(H_{3}I)$$
 (B) $(H_{3}ONa)$
(C) $(H_{3}I)$ (C) $(H_{3}I)$ (C) $(H_{3}OH)$ (C) $(H_{3}OH)$

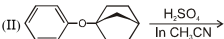
58.

$$R - OH \xrightarrow{X^{\odot}} R - X + \overleftarrow{O}H$$

(A) Reaction will not take place because OH is poor leaving group; X is weak base and OH is strong base
(B) Reaction will not take place because OH is poor leaving group; X is strong base and OH is weak base.
(C) Reaction will not take place because OH is good leaving group; X is strong base and OH is weak base.
(D) Reaction will not take place because OH is good leaving group; X is weak base and OH is strong base.

59.

$$(I) \longrightarrow - \circ \longleftarrow \xrightarrow{H_2SO_4} H_2SO_4 \longrightarrow$$



Which of the following statement is correct about these reaction.

- (A) Both the reactions take place at the same rate
- (B) The first reactions take place faster than second reaction.
- (C) The second reactions take place faster than first reaction.
- **(D)** Both the reactions take place by S_N^{1} mechanism

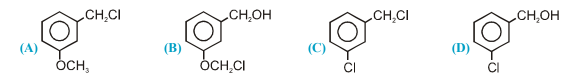
60.

DCH.

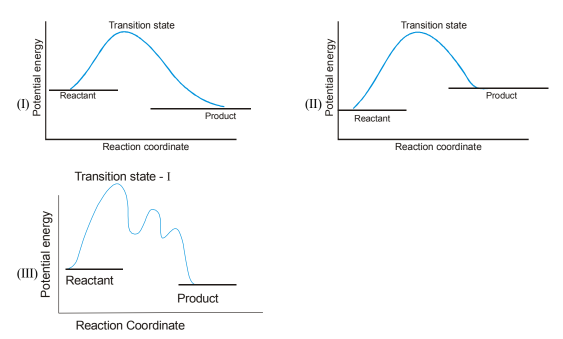
$$CH_2OH$$

$$\xrightarrow{SOCl_2} A + SO_2 + HCl$$
Pyridine

The product A in the above reaction is



61. Following are the curves for nucleophilic substitution reaction.

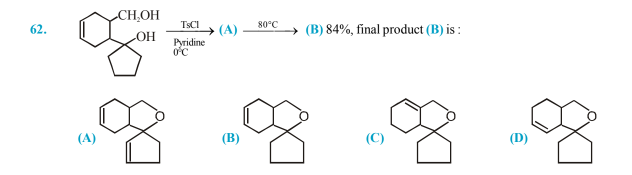


The correct statement (s) is (are)

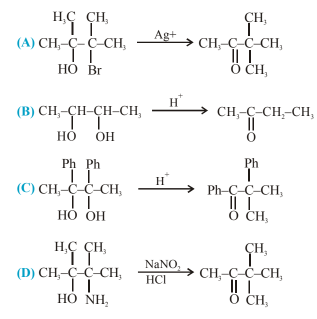
(A) 'I' is potential energy diagram for S_N^2 reaction that takes place with a negative potential energy change.

(B) 'II' is potential energy diagram for S_N^2 reaction with a positive potential energy change

(C) 'III' shows potential energy diagram for S_N^1 reaction with large potential energy of activation for first (slowest) step (D) All of the above



63. Which of the following does not representing the correct product



64. What are the products of the following reaction ?

$$(A) p Br - C_6H_4 - OCH_2CH_2 - Br$$

$$(B) Ph - Br + Br - CH_2CH_2 - Br$$

$$(C) Ph - Br + Br - CH_2CH_2 - Br$$

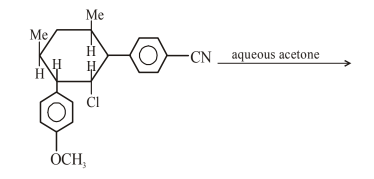
$$(D) Ph - OH + CH_3 - CHBr_2$$

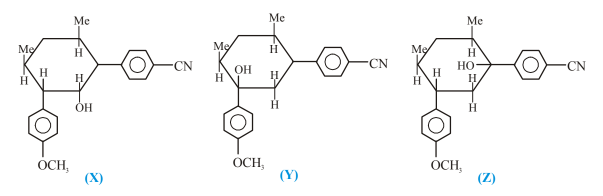


- 1. Which of these statements are correct about nucleophiles :
 - (A) Nucleophiles have an unshared electron pair and can make use of this to react with an electron deficient species.
 - (B) The nucleophilicity of an element (as electron donor) generally increases on going down a group in the periodic table.
 - (C) A nucleophile is electron-deficient species
 - (D) All good nucleophiles are good bases when we deal across the period.
- 2. The correct nucleophilicity order is/are :

(A) $(CH_3)_3 O^{\Theta} > CH_3^{\Theta}$ (B) $CH_3 S^{\Theta} > CH_3 SH$ (C) $CH_3 CH_2 CH_2 O^{\Theta} > (CH_3)_3 CO^{\Theta}$ (D) $(CH_3 CH_2)_3 N > (CH_3 CH_2)_3 P$

3. Which of the following are correct regarding the products of the given reaction

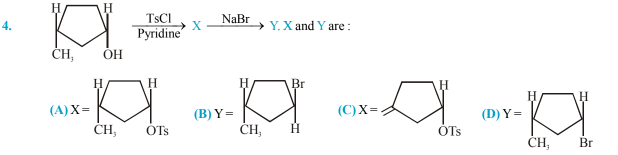




(A) only X and Z are formed (B) Y > Z (amount)

(C) X > Y (amount)

(D) Y > X (amount)



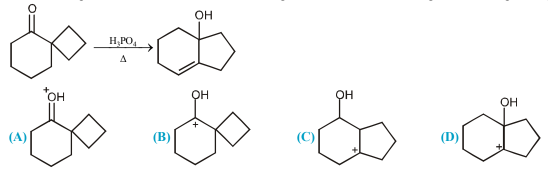
5.
$$Y \xleftarrow{CH_{3}OH}_{H^{\bigoplus}} CH_{3} \xrightarrow{C}_{O}CH_{2} \xrightarrow{H_{2}O^{18}}_{H^{\bigoplus}} X, \text{ Identify } X \& Y :$$

$$(A) X = CH_{3} \xrightarrow{C}_{O}C \xrightarrow{C}_{O}CH_{2} \xrightarrow{H_{2}O^{18}}_{H^{\bigoplus}} X, \text{ Identify } X \& Y :$$

$$(A) X = CH_{3} \xrightarrow{C}_{O}C \xrightarrow{C}_{O}CH_{2} \xrightarrow{C}_{O}CH_{2} \xrightarrow{C}_{O}CH_{2} \xrightarrow{C}_{O}CH_{3} \xrightarrow{$$

6.

What are the possible intermediates of the following reaction which form during this reaction path way ?



- 7. Which of the following order is/are correct for the rate of E2 reaction.
 - (A) 5-Bromocycloheptene > 4-Bromocycloheptene
 - (B) 2-Bromo-1-phenylbutane > 3-Bromo-1-phenylbutane
 - (C) 3-Bromocyclohexene > Bromocyclohexane
 - (D) 3-Bromo-2-methylpentane > 2-Bromo-4-methylpentane
- 8. Which of the following statement (s) is/are correct
 - (A) E2 is a concerted reaction in which bonds break and new bonds form at the same time in a single step.
 - (B) Order of reactivity of alkyl halides towards E2 dehydrohalogenation is found to be $3^{\circ} > 2^{\circ} > 1^{\circ}$
 - (C) In E2 reaction both β hydrogen and leaving group should be antiperiplanar.
 - (D) In E2 elimination different stereoisomer (diastereomer) converts into different stereo product.
- 9. Observe the following reaction and tick correct option (s)

$$CH_3 - CH - CH_3 + CH_3ONa \xrightarrow{r_1} CH_3 - CH - OCH_3 \xrightarrow{(I)} CH_3$$

$$CH_3 - I + CH_3 - CH - ONa \xrightarrow{r_2} CH_3 - CH - OCH_3 \dots (II)$$

 $CH_3 CH_3 CH_3 CH_3 CH_3$

(A) $r_1 < r_2$

I

- (B) Transition state in reaction (I) is less stable then transition state in reaction (II)
- (C) Substrate of both the reactions are equally reactive to a particular nucleophile
- (D) r_1 and r_2 both are directly proportional to the concentration of nucleophile CH₃ONa

10.

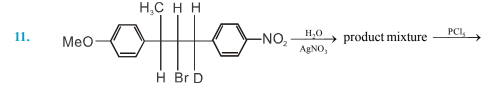
$$CH_{3} - CH - CH_{2} \xrightarrow{CH_{3}C} C^{\Theta} \times (X) \xrightarrow{CH_{3}I} (Y). \text{ Identify } X \& Y:$$

$$(A) X = CH_{3} - CH - CH_{2} - C \equiv CCH_{3} \qquad (B) Y = CH_{3} - CH_{2} - CH - C \equiv CCH_{3}$$

$$O - CH_{3} \qquad OCH_{3} \qquad (D) X = CH_{3} - CH - CH_{2} - C \equiv CCH_{3}$$

$$O - CH_{3} \qquad (D) X = CH_{3} - CH - CH_{2} - C \equiv CCH_{3}$$

$$O - CH_{3} \qquad (D) X = CH_{3} - CH - CH_{2} - C \equiv CCH_{3}$$



No of product 'm' (Total isomers) $\xrightarrow{\text{Fractional}}$ No. of Fractions 'n'.

- (A) Value of m & n are 4,2 respectively.
- (B) Product mixture, and all fractions are optically active.
- (C) Value of m & n are 4,4 respectively.
- (D) Reaction of product mixture with PCl_5 is S_N^2 mechanism.
- 12. Which of the following reaction are not feasible ?

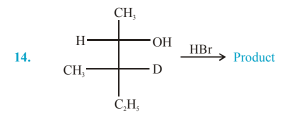
(A)
$$\bigcirc - \Theta \oplus ONa + CH_3Br \longrightarrow \bigcirc -OCH_3$$

(B) $CH_3 \longrightarrow CH_3 \oplus CH_3 \oplus OR$
(C) $CH_3 \longrightarrow CH_3 \oplus CH_3 \oplus CH_3 \oplus CH_3 \oplus OR$
(D) $\bigcirc -Br + RONa \oplus \odot \bigcirc OR$

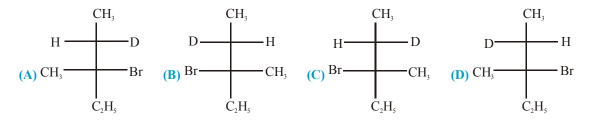
13. Which of the following reaction takes place by S_N^2 mechanism :

(A)
$$CH_3 - CH_2 - ONa + CH_3 - CH_2 - Cl \longrightarrow$$

(B) $Cl + NaOH \longrightarrow$
(C) $(CH_3)_3 - C - CH_2 - CH_2 - Br + excess NH_3 \longrightarrow$
(D) $CH_3 - CH_2 - OH + HBr \longrightarrow$



Which of the following structures are possible products.



 $C_{2}H_{5}C \equiv CH \xrightarrow{CH_{3}MgBr} A \xrightarrow{1.HCHO} B$

The product (B) is

(A) CH₂CH₂OH

(B) $C_2H_5C - CHCH_2OH$ CH₃

(C)
$$C_2H_5C \equiv C - CH - CH_3$$

OH
(D) $C_2H_5C \equiv C - CH_2OH$

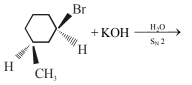
16. In the transformations

PhCH = CH₂
$$\xrightarrow{\text{ArCO}_3\text{H}}_{\text{CH}_2\text{Cl}_2}$$
 A $\xrightarrow{1.\text{LiAlH}_4, \text{Et}_2\text{O}}_{2.\text{H}_2\text{O}}$ B
the end product (B) is
(A) PhCH—CH₂ (B) PhCHOH CH₃ (C) PhCH₂CH₂COAr (D) PhCH₂CH₂OH

17. The reactivities of methanol (I), 1- propanol (II), 2- butanol (III) and 2- methyl-2- propanol (IV) towards sodium metal follow the order (D) IV > II > III > I

(A) I > II > III > IV**(B)** IV > III > II > I (\mathbf{C}) I > IV > II > III

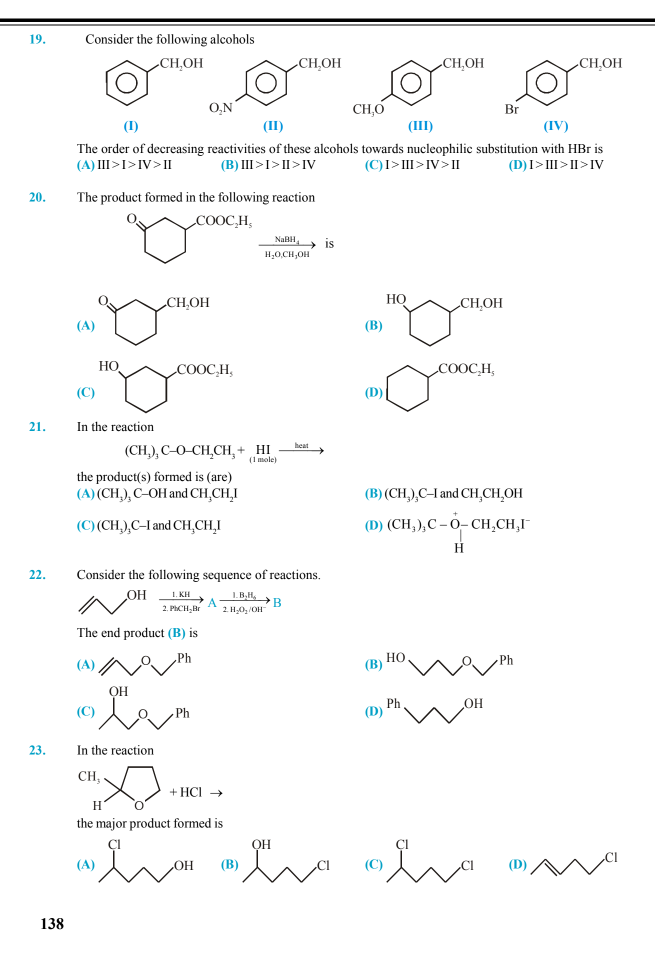
18. Consider the following reaction.

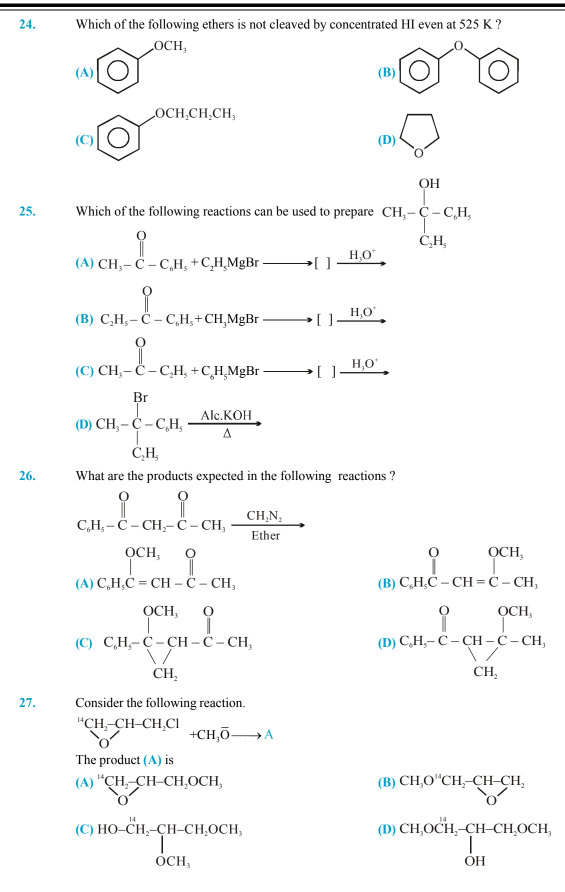


(1R, 3S)-cis-1 -Bromo -3-methylcyclohexane The product formed in the reaction is (A) (1R, 3S)-cis-3- methylcyclohexanol (C) (1S, 3S)-trans-3-methylcyclohexanol

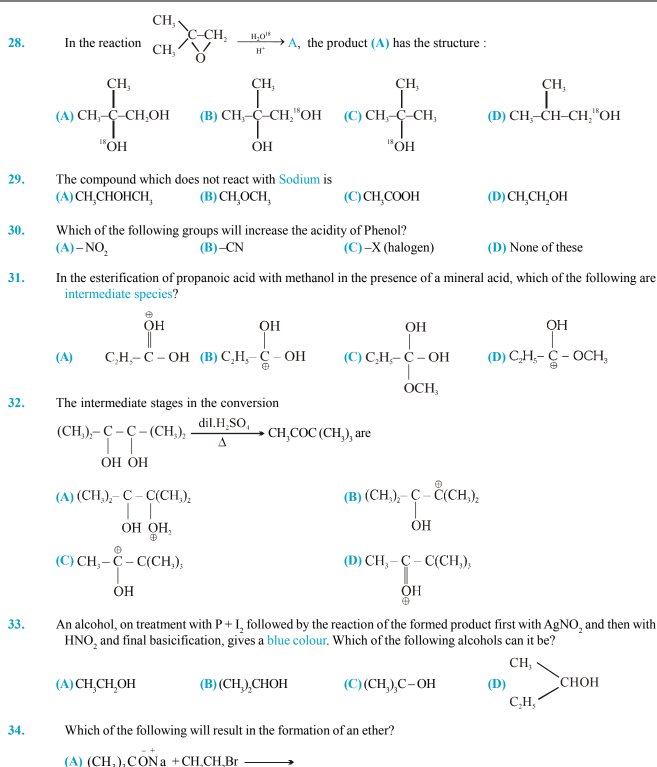
(B) (1S,3R)-cis-3-methylcyclohexanol (D) (1R,3R)-trans-3-methylcyclohexanol

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(A)
$$(CH_3)_3 CON a + CH_3 CH_2 Br$$

(B) $(CH_3)_3 CBr + C_2 H_5 ON a$
(C) $C_6 H_5 ON a + CH_3 Br$
(D) $C_6 H_5 Br + CH_3 ON a$

	Part # II	>>	[Assertion & Reason Type Questions]	
	 These questions contains, Statement-I (assertion) and Statement-II (reason). (A) Statement-I is True, Statement-II is True ; Statement-II is a correct explanation for Statement-I (B) Statement-I is True, Statement-II is True ; Statement-II is NOT a correct explanation for Statement-I (C) Statement-I is True, Statement-II is False. (D) Statement-I is False, Statement-II is True. 			
1.	Statement-I: Tert-Butoxide is a stronger base than OH^- or $C_2H_5O^-$ ion but is a much poorer nucleophile. Statement-II: A negatively charged ions is always more powerful nucleophile than its conjugate acid.			
2.	 Statement-I With Br₂-H₂O, phenol gives 2,4,6- tribromophenol but with Br-CS₂, it gives 4-bromophenol as the major product. Statement-II : In water ionization of phenol is enhanced but in CS₂, it is greatly suppressed. 			
3.		I : The relative	n n-alcohols in water decreases with increase in molecular we proportion of the hydrocarbon part in alcohols increases with ir its enhanced hydrogen bond with water.	•
4.	Statement-I: p- Nitrophenol is a stronger acid than o-nitrophenol. Statement-II: Intramolecular H-bond makes o-isomer weaker than p-isomer.			
5.			ore reactive than benzene towards of phenol, the intermediate carbocation is more resonance s	tabilized.
6.	Statement-	 Alcohol give strong acid 	es substitution reaction with HI but elimination reaction with l	H_2SO_4 . Although both are
	Statement-2		l nucleophile so it gives substitution reaction while $\mathrm{HSO_4}^{\Theta}$ ation reaction.	is weak nucleophile so it
7.	 Statement-1 : The anti-periplanar transition state is the most commonly seen in E² reactions. Statement-2 : The transition state for the anti-periplanar arrangement is a staggered conformation, with the base fa away from the leaving group. In most cases, this transition state is higher in energy than that for the syn-periplanar elimination. 			
8.	Statement-	1 : Tertiary-but methoxide.	yl methyl ether is not prepared by the reaction of tertiary-buty	l bromide with sodium
	Statement-	2 : Tertiary alky	yl halide gives elimination with strong base.	
9.			nnot be converted into alkyl bromides by reaction with NaBr. ery weak base cannot displace strong base OH ⁻ .	
10.	Statement-1 : S_N^2 reaction is carried out in the presence of polar aprotic solvents. Statement-2 : Polar aprotic solvents do not contain acidic hydrogen.			
11.	Statement-	1 : Iodide (I ⁻) is	s the strongest nucleophile (in protic solvent) among the hali	de anions

Part # I

Exercise # 3

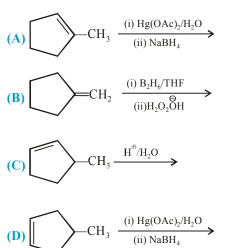
1. Match the column I with column II. **Column-I** (substrate + RMgX) (A) HCHO

(II) Here
$$O$$

(B) CH₃—C—CH₃
(C) CH₃—CH—CH—CH—CH₃

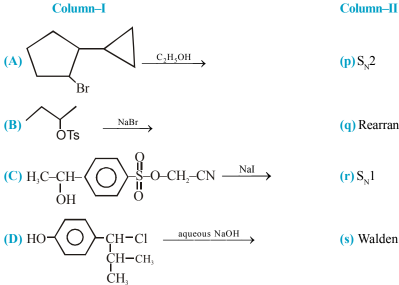
(D) Ester

2. Match the column I with column II. **Column-I (Reaction)**



Match the characteristics mentioned in Column -II with the reactions given in Column -I

3.



(p) S_{N}^{2}

(q) Rearrangement

 $(\mathbf{r}) \mathbf{S}_{N} \mathbf{1}$

(s) Walden Inversion

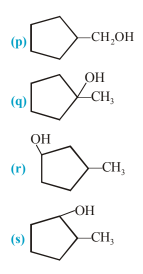
- **Column-II**
- (Product) (p) Tertiary alcohol

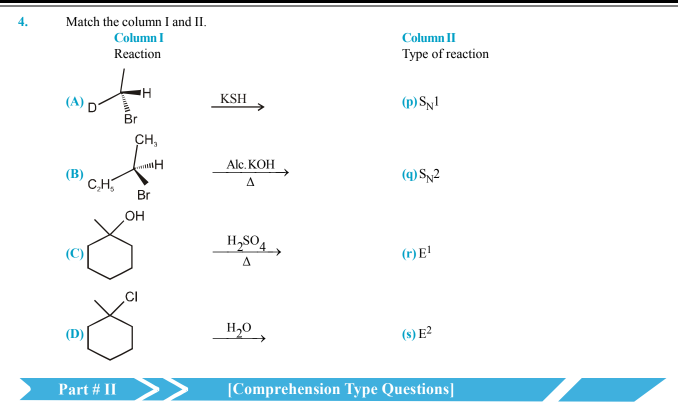
[Matrix Match Type Questions]

- (q) First ketone then 3° alcohol
- (r) Secondary alcohol

(s) Primary alcohol

Column-II (Possible products)





Comprehension #1

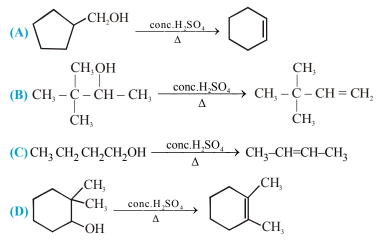
Alcohols undergo acid catalysed elimination reactions to produce alkenes. Because water is lost in the elimination, this reaction is called dehydration reaction. Secondary and tertiary alcohols always give E^1 reaction in dehydration. Primary alcohols whose β -carbon is branched also give E^1 reaction. The reactivity of alcohol for elimination reaction is tertiary alcohol > Primary alcohol.

1. In the given reaction :
$$CH_3 - CH - CH - CH_3 \xrightarrow[]{conc.H_2SO_4}{\Delta}$$
 Alkenes
 CD_3

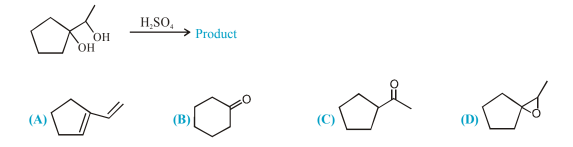
OII

Total number of alkenes (Including stereo isomers) formed will be(A) Two(B) Six(C) Four(D) Five

2. Which of the following dehydration product (major) is incorrect?

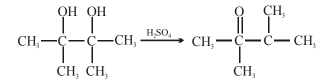


3. Identify the product in the given reaction :



Comprehension #2

When pinacol is treated with dilute H_2SO_4 , a re-arrangement reaction takes place which leads to the formation of a ketone.

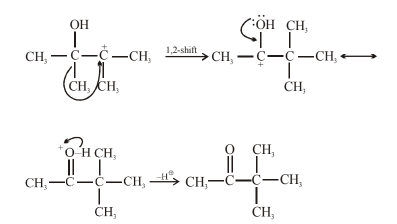


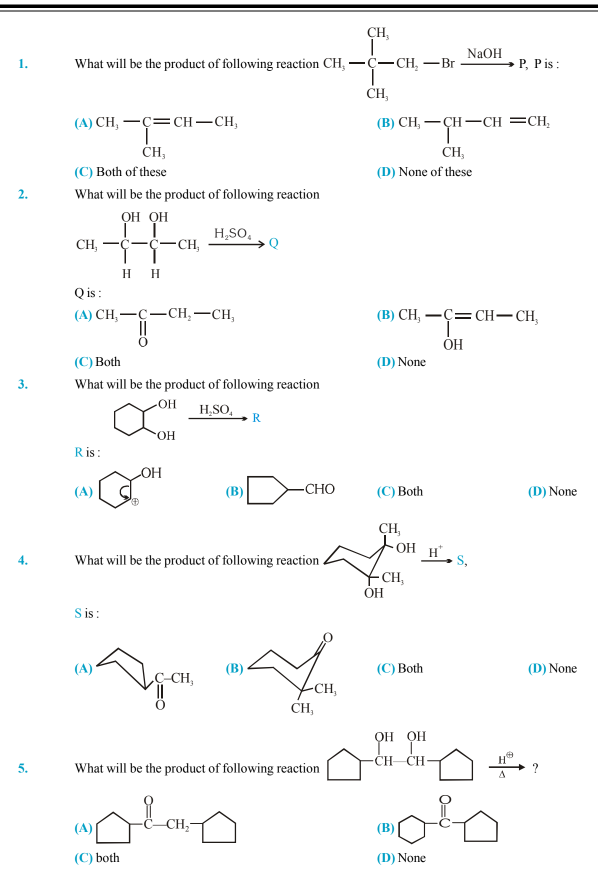
This reaction involves re-arrangement of carbocation.

Step 1:

Step 2 :

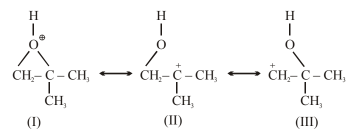
Carbocation rearrange by hydride, alkyl or aryl shift to get as stable as they can. Stability is the driving force for rearrangement migration of bond may also occur. Where by ring expansion and ring contraction takes place. The relief of strain can provide a powerful driving force for re-arrangement.





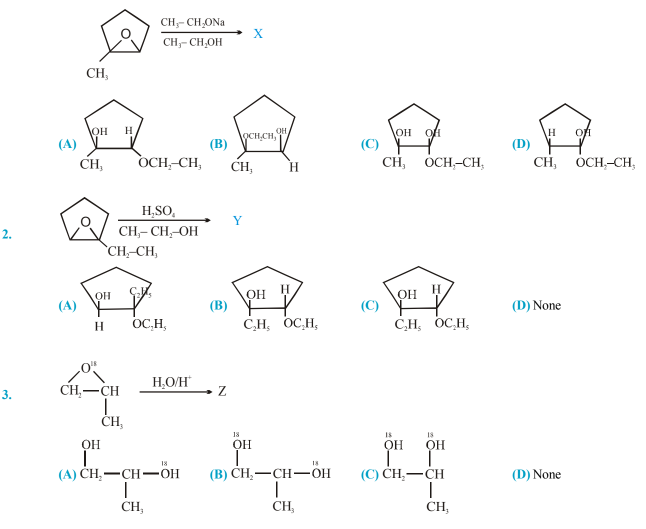
Comprehension #3

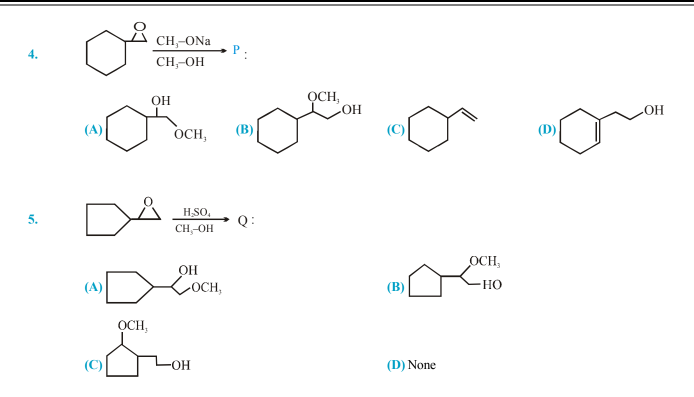
Symmetrically substituted epoxides give the same products in both the acid catalysed and base catalyzed ring opening. An unsymmetrical epoxide gives different products under acid catalysed and base catalysed conditions. Under basic conditions, the alkoxide ion simply attacks the less hindered carbon atom in an S_N^2 displacement. Under acidic conditions, the alcohol, attacks the protonated epoxide.



Structure II and III show that the oxirane carbon share part of positive charge. The tertiary carbon bear a larger part of positive charge and it is more strongly electrophilic. The bond between tertiary carbon and oxygen is weaker implying a lower transition state energy for attack at the tertiary carbon. Attack by the weak nucleophilic is sensitive to the strength of electrophile. Centre attack takes place at more electrophilic carbon which is usually the more substituted carbon because it can better support the positive charge.

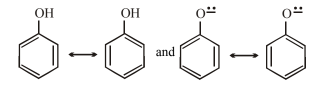
1. What will be the products in following reactions



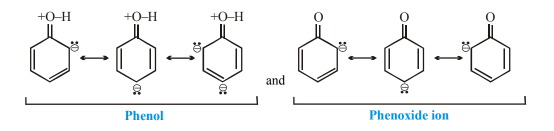


Comprehension #4

Phenols are converted into their salts by aqueous NaOH but not by aqueous bicarbonates. The salts are converted into the free phenols by aqueous mineral acids, carboxylic acid or carbonic acids. Most phenols have K_a value of about 10^{-10} , and are tremendously more acidic than alcohols. The difference in acidity are due to difference in stabilities of reactants and products. Phenol and phenoxide ions contain benzene ring and therefore must be hybrid of Kekuley structures



Being basic, oxygen can share more than a pair of electron with the ring.



Since energy must be supplied to separate opposite charge, the structure of phenols should contain more energy. The net effect of resonance is therefore to stabilise the phenoxide ion to a greater extent than the phenol and thus to shift the equilibrium towards ionization and make K_a larger than for an alcohol.

OH

 (\mathbf{B})

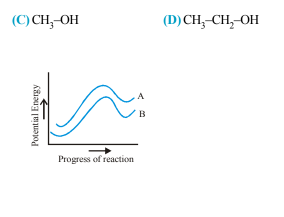
- 1.
- Which of the following is strongest acid?

2. Consider the following curves :

(A) Curve A represents the ionisation of alcohol
(B) Curve B represent the ionisation of Phenols
(C) Curve A represents the ionisation of phenol
(D) None

3. Which of the following is more stable:

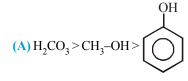


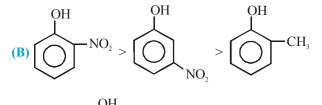


(C) Both

(D) Unpredictable

4. Correct order of acidity is





(C)
$$H_2CO_3 > \bigcirc > CH_3 - OH$$

(D) Both B and C

- Choose the correct statement:
 - (A) Phenol gives effervescence with NaHCO₃
 - (B) Picric acid is weaker acid than carbonic acid
 - (C) Picric acid gives effervescence with NaHCO₃

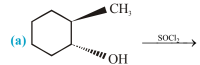
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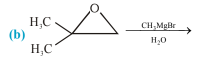
(D) $R-O^{\Theta}$ is more stable than

5

Exercise # 4 [Subjective Type Questions]

- 1. Explain why ArOR ethers are cleaved to give RI and ArOH rather than ArI and ROH.
- 2. Complete the following reactions :





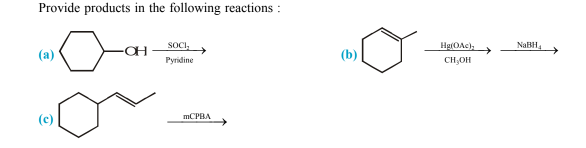
3. Complete the follofing reactions :



4. $C_2H_5OH \xrightarrow{PCl_5} (A) \xrightarrow{KCN} (B) \xrightarrow{H_3O^+} (C) \xrightarrow{NH_3} (D) \xrightarrow{heat} (E)$

5.
$$CH_3CH_2CH_2OH \xrightarrow{PBr_5} (A) \xrightarrow{KOH(Alc.)} (B) \xrightarrow{HBr} (C) \xrightarrow{NH_3} (D)$$

6. t-butyl alcohol reacts less rapidly with metallic sodium than the primary alcohol. Explain why

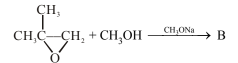


8.

7.

Identify the products A and B giving proper explanation :

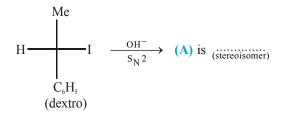
$$CH_{3}CH_{2} + H_{2}O^{18} \xrightarrow{H^{+}} A,$$



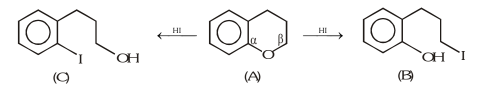
9. Indicate bonds which are cleaved I : in basic conditions II : in acidic conditions



10. In the following S_N^2 reaction



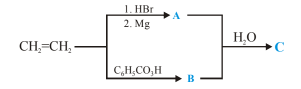
- 11. Hydration of 3-phenyl-1-butene in dilute H_2SO_4 is not a satisfactory method for preparing 3-phenyl-2-butanol, because 2-phenyl-2-butanol is obtained instead. Explain.
- 12. When A (given below) reacts with HI products is B and not C. Explain.



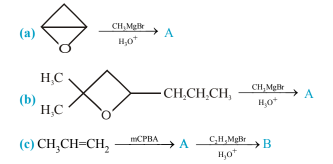
13. Isotopic carbon-14 in (A) appears at new position (as in B) when (A) reacts with CH₃ONa. Explain.

$$(A) \qquad (B) \qquad (B)$$

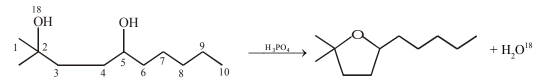
- 14. A compound (X) reacts with thionyl chloride to give a compound (Y). (Y) reacts with Mg to form a Grignard reagent, which is treated with acetone and the product is hydrolysed to give 2-methyl-2- butanol. What are structural formulae of (X) and (Y)?
- 15. Compound (A) $C_4H_{10}O$ reacts rapidly with metallic sodium, but undergoes almost no reaction with Lucas reagent. When (A) is treated with hot concentrated sulphuric acid, a new compound (B) C_4H_8 is formed. If C_4H_8 is hydrated with sulphuric acid a new compound (C) C_4H_9OH is formed, which is almost inert to metallic sodium but reacts rapidly with Lucas reagent. What are (A), (B) and (C) ?
- 16. Give the product of the reaction of Ph_2CHCH_2OH with HBr and explain its formation.
- 17. Complete the following reaction



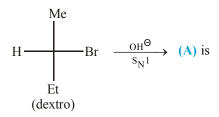
18. Complete the following reactions

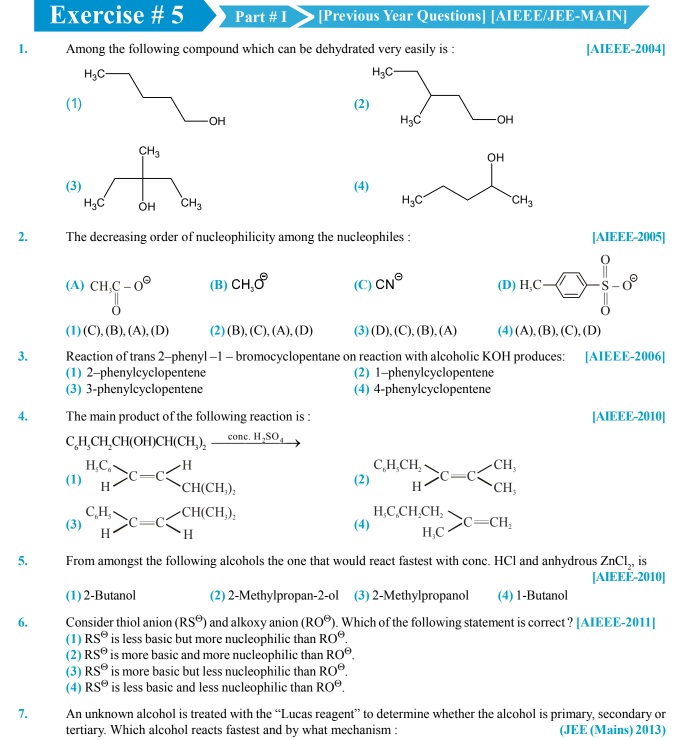


19. In the following dehydration of diol with H_3PO_4 , following product is formed such that isotopic ¹⁸O goes with H_2O . Explain.



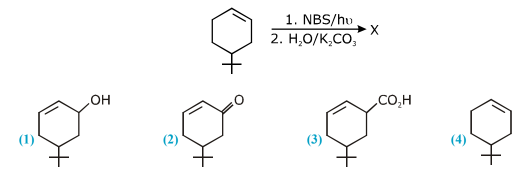
20. In the following S_N^1 reaction :





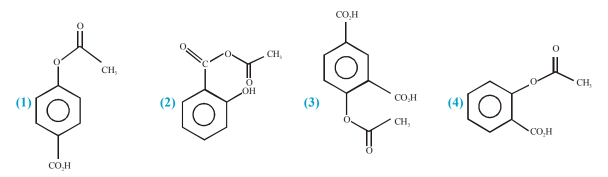
- (1) secondary alcohol by $S_N 1$ (2) tertiary alcohol by $S_N 1$
- (3) secondary alcohol by S_N^2 (4) tertiary alcohol by S_N^2

8. The product of the reaction given below is :

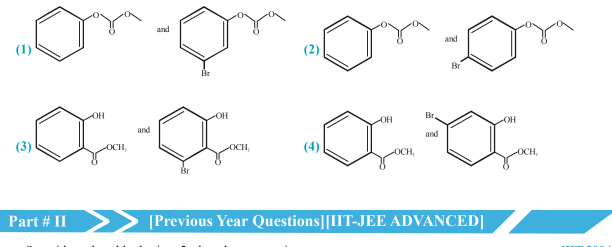


9. Phenol on treatment with CO_2 in the presence of NaOH followed by acidification produces compound X as the major product. X on treatment with $(CH_3CO)_2$ O in the presence of catalytic amount of H_2SO_4 produces :

(**JEE** (Mains) 2018)



Phenol reacts with methyl chloroformate in the presence of NaOH to form product A. A reacts with Br₂ to form product B. A and B are respectively : (JEE (Mains) 2018)



- 1.
 On acid catalysed hydration, 2-phenyl propene gives :
 [IIT-2004]

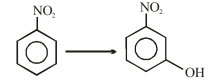
 (A) 3-phenyl-2-propanol
 (B) 2-phenyl-1-propanol
 (D) 2-phenyl-2-propanol

 (C) 1-phenyl-3-propanol
 (D) 2-phenyl-2-propanol
 [IIT-2005]

 2.
 Conversion of cyclohexanol into cyclohexene is most effective in :
 [IIT-2005]
- Conversion of cyclohexanol into cyclohexene is most effective in :

 (A) concentrated H₃PO₄
 (B) concentrated HCl
 (C) concentrated HCl/ZnCl₂
 (D) concentrated HBr

3. Carry out following conversion in four or less steps. Also mention all the reagents used and reaction conditions.



4. An organic compound $P(C_{3}H_{10}O)$ reacts 10^{15} times faster then ethylene with dilute $H_{2}SO_{4}$ to give two products Q and R. Both Q and R give positive iodoform test. Identify P, Q and R and also give reason for very high reactivity of P.

NaOH

CII

[IIT-2004]

[IIT-2004]

[**JEE - 2005**]

CU

Identify (X) and (Y).

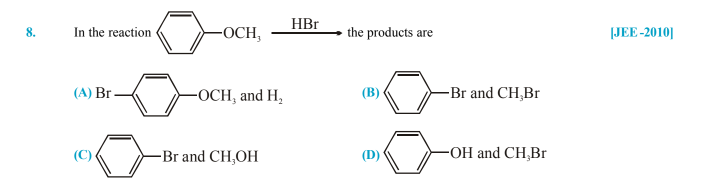
6. Phenyl magnesium bromide reacting with t-Butyl alcohol gives

 $\xrightarrow{H^{+}/\Delta} (X) \xrightarrow{1. O_{3}} (Y)$

(A)
$$Ph-OH$$
 (B) $Ph-H$ (C) $Ph-O-C-CH_3$ (D) $Ph-C-CH_3$ (D) $Ph-C-CH_3$ (H) CH_3

- 7.
 Statement-1: p-Hydroxybenzoic acid has a lower boiling point than o-hydroxybenzoic acid.
 [JEE 2007]

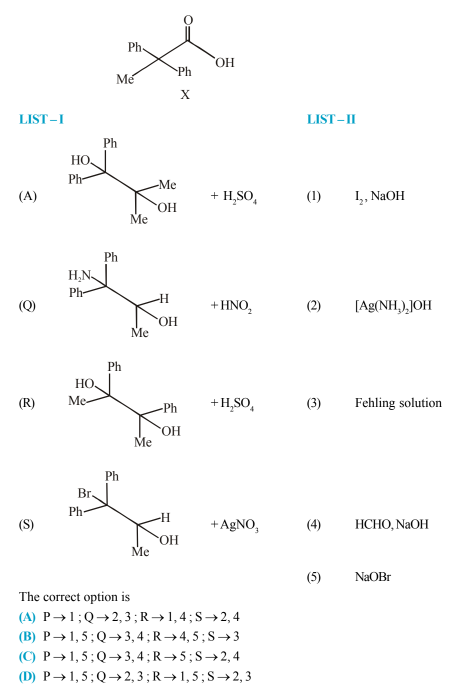
 Statement-2: o-Hydroxybenzoic acid has intramolecular hydrogen bonding.
 [JEE 2007]
 - (A) Statement-1 is True, Statement-2 is True; Statement-2 is a correct explanation for Statement-1.
 - (B) Statement-1 is True, Statement-2 is True; Statement-2 is NOT a correct explanation for Statement-1.
 - (C) Statement-1 is True, Statement-2 is False.
 - (D) Statement-1 is False, Statement-2 is True.

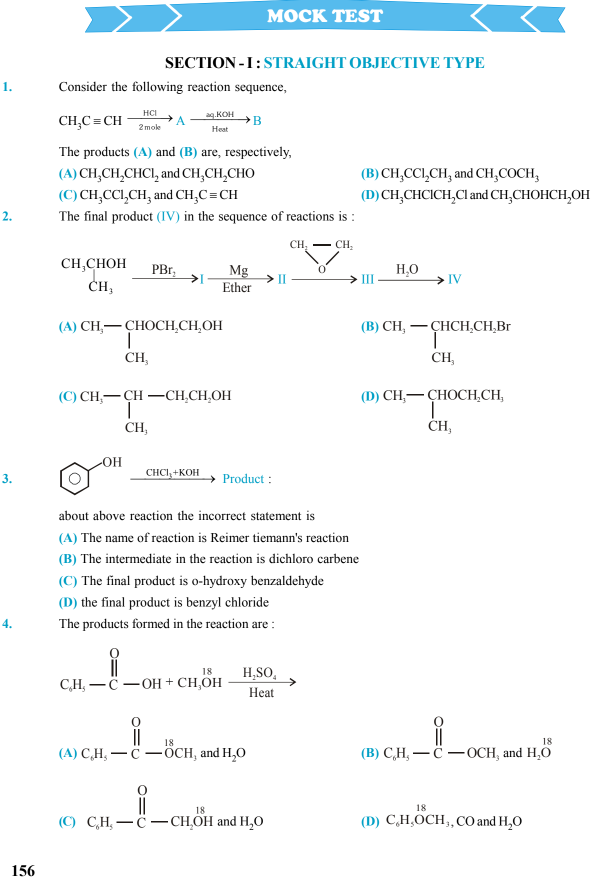


9. The desired product X can be prepared by reacting the major product of the reaction in LIST-I with one or more appropriate reagents in LIST - II.

(given, order of migratory aptitude : aryl > alkyl > hydrogen)

[JEE Advanced 2018]

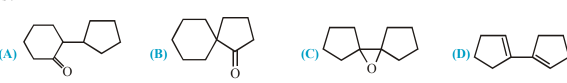




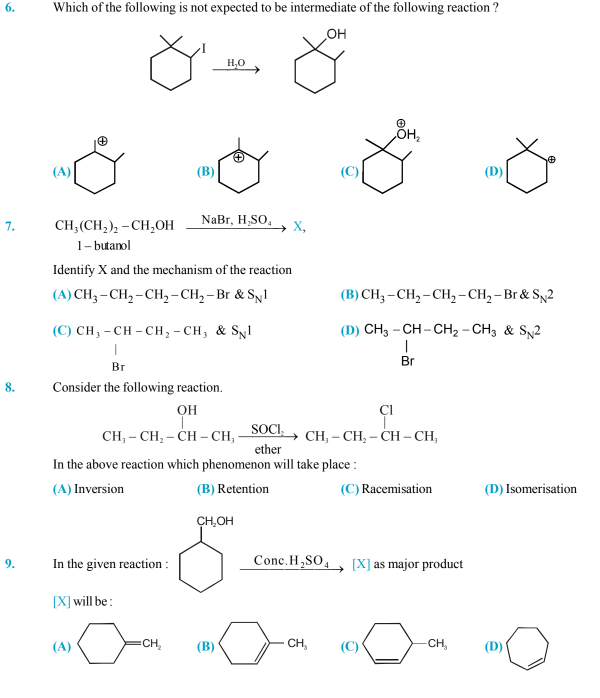
5. The major product (A) formed in the reaction

$$\underbrace{\bigcup_{HO}}_{HO} \underbrace{-H_2SO_4}_{OH} \xrightarrow{H_2SO_4} A$$

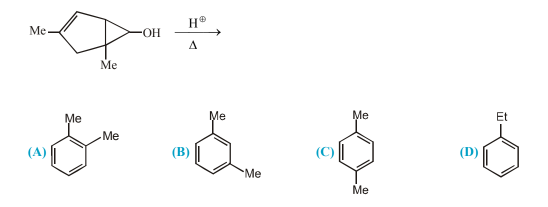
is :



6.



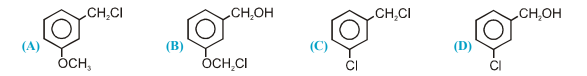
10. Write the product of the following reaction



- **11.** Which of the following statement is not true ?
 - (A) Nucleophiles possess unshared pairs of electron which are utilized in forming bonds with electrophilic substrate.
 - (B) The cyanide ion is an ambident nucleophile and causes nucleophilic substitution of alkyl halide by either of its carbon atom or nitrogen atom.
 - (C) The nitrite ion is an ambident nucleophile and causes nucleophilic substitution of alkyl halide by either of its oxygen atoms or nitrogen atom.
 - (D) Strength of Nucleophile generally decreases on going down a group in the periodic table.

12.
$$(\bigcirc CH_2OH) \xrightarrow{SOCl_2} A + SO_2 + HCl$$

The product A in the above reaction is



13. What are the products of the following reaction ?

$$(A) p Br - C_6H_4 - OCH_2CH_2 - Br$$

$$(B) Ph - Br + Br - CH_2CH_2 - Br$$

$$(C) Ph - Br + Br - CH_2CH_2 - Br$$

$$(D) Ph - OH + CH_3 - CHBr_2$$

SECTION - II : MULTIPLE CORRECT ANSWER TYPE

- 14. Which of these statements are correct about nucleophiles :
 - (A) Nucleophiles have an unshared electron pair and can make use of this to react with an electron deficient species.
 - (B) The nucleophilicity of an element (as electron donor) generally increases on going down a group in the periodic table.
 - (C) A nucleophile is electron-deficient species
 - (D) All good nucleophiles are good bases when we deal across the period.

15.
$$Y \xleftarrow{CH_3OH}_{H^{\bigoplus}} CH_3 \xrightarrow{CH_3}_{O} CH_2 \xrightarrow{H_2O^{18}}_{H^{\bigoplus}} X$$
, Identify X & Y :

(A)
$$X = CH_3 - C - CH_2$$

B) $Y = CH_3 - C - CH_2$
 $H_3 - C - C$

(C)
$$X = CH_3 - CH_2$$
,
OH
(D) $Y = CH_3 - CH_2$,
OH
(D) $Y = CH_3 - CH_2$

16.
$$CH_3 - CH - CH_2 \xrightarrow{CH_3C} CH_3C \equiv C^{\Theta}$$
 (X) $CH_3I \rightarrow$ (Y). Identify X & Y :

(A)
$$X = CH_3 - CH - CH_2 - C \equiv CCH_3$$

|
 $O - CH_3$

(B)
$$Y = CH_3 - CH_2 - CH - C \equiv CCH_3$$

|
OCH₃

(C)
$$Y = CH_3 - CH - CH_2 - C \equiv CCH_3$$

 $\downarrow O - CH_3$

$$(\mathbf{D}) \mathbf{X} = \mathbf{CH}_3 - \mathbf{CH} - \mathbf{CH}_2 - \mathbf{C} \equiv \mathbf{CCH}_3$$

17. Consider the following sequence of reactions.

$$C_2H_5C \equiv CH \xrightarrow{CH_3MgBr} A \xrightarrow{1.HCHO} B$$

The product **(B)** is

(A)
$$CH_3CH_2OH$$

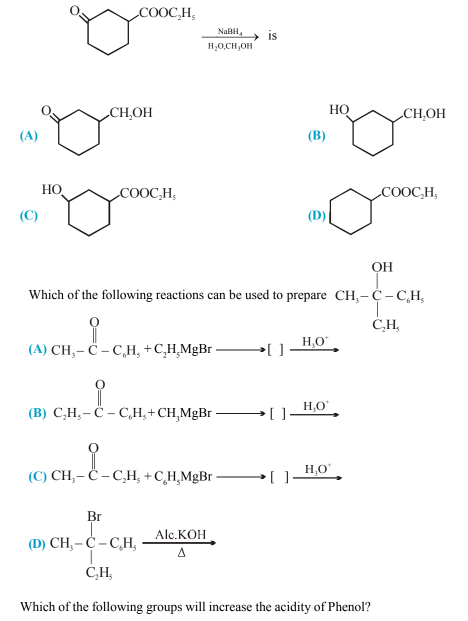
(B) $C_2H_5C - CHCH_2OH$
 CH_3

(C)
$$C_2H_5C \equiv C - CH - CH_3$$

|
OH
(D) $C_2H_5C \equiv C - CH_2OH$

CHEMISTRY FOR JEE MAIN & ADVANCED

18. The product formed in the following reaction



20. Which of the following groups will increase the acidity of Phenol?
 (A) - NO₂
 (B) -CN
 (C) -X (halogen)
 (D) None of these

21. Which of the following will result in the formation of an ether?

(A) $(CH_3)_3 CON a + CH_3 CH_2 Br \longrightarrow$ (B) $(CH_3)_3 CBr + C_2 H_5 ON a \longrightarrow$ (C) $C_6 H_5 ON a + CH_3 Br \longrightarrow$ (D) $C_6 H_5 Br + CH_3 ON a \longrightarrow$

19.

SECTION - III : ASSERTION AND REASON TYPE

- 22. Statement-1: The anti-periplanar transition state is the most commonly seen in E² reactions.
 - Statement-2: The transition state for the anti-periplanar arrangement is a staggered conformation, with the base far away from the leaving group. In most cases, this transition state is higher in energy than that for the syn-periplanar elimination.
- 23. Statement-1: Tertiary-butyl methyl ether is not prepared by the reaction of tertiary-butyl bromide with sodium methoxide.

Statement-2: Tertiary alkyl halide gives elimination with strong base.

- Statement-1: Alcohols cannot be converted into alkyl bromides by reaction with NaBr.
 Statement-2: Br⁻ being very weak base cannot displace strong base OH⁻.
- 25. Statement-1 : S_N2 reaction is carried out in the presence of polar aprotic solvents.

Statement-2: Polar aprotic solvents do not contain acidic hydrogen.

Statement-1 : Iodide (I⁻) is the strongest nucleophile (in protic solvent) among the halide anions
 Statement-2 : Iodide is the largest halide anion and is the most weakly solvated in a protic solvent

SECTION - IV : COMPREHENSION TYPE

Read the following comprehensions carefully and answer the questions.

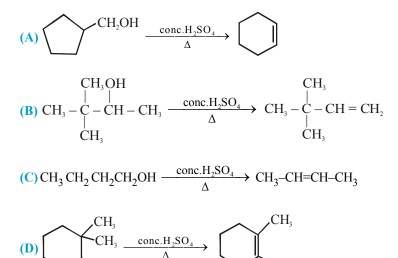
Alcohols undergo acid catalysed elimination reactions to produce alkenes. Because water is lost in the elimination, this reaction is called dehydration reaction. Secondary and tertiary alcohols always give E^1 reaction in dehydration. Primary alcohols whose β -carbon is branched also give E^1 reaction. The reactivity of alcohol for elimination reaction is tertiary alcohol > Secondary alcohol > Primary alcohol.

27. In the given reaction :
$$CH_3 - CH - CH - CH_3 \xrightarrow{\text{conc.}H_2SO_4} Alkenes$$

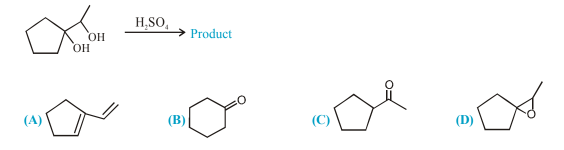
 CD_3

28. Which of the following dehydration product (major) is incorrect ?

OH



29. Identify the product in the given reaction :



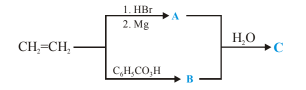
SECTION - V : MATRIX - MATCH TYPE

30. Match the column I and II.

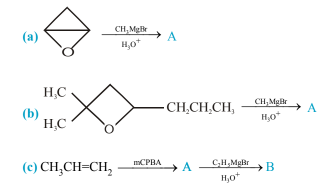
Column I Column II Reaction Type of reaction $\xrightarrow{\text{KSH}}$ (A) D (**p**) S_N^{1} Ŕr CH₃ Hum $\xrightarrow{\text{Alc. KOH}}$ **(B)** (q) S_N^2 Br OH $\frac{H_2SO_4}{\Delta}$ (r) E¹ **(C)** C H₂O (s) E² **(D**)

SECTION - VI : SUBJECTIVE TYPE

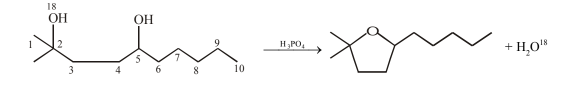
31. Complete the following reaction



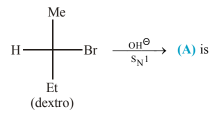
32. Complete the following reactions



33. In the following dehydration of diol with H_3PO_4 , following product is formed such that isotopic ¹⁸O goes with H_2O . Explain.



34. In the following S_N^1 reaction :



ANSWER KEY

EXERCISE - 1

۲

 1. C
 2. A
 3. C
 4. C
 5. B
 6. C
 7. C
 8. D
 9. B
 10. C
 11. A
 12. C
 13. B

 14. C
 15. D
 16. A
 17. A
 18. A
 19. B
 20. A
 21. B
 22. D
 23. A
 24. C
 25. A
 26. B

 27. D
 28. C
 29. A
 30. A
 31. C
 32. D
 33. C
 34. A
 35. B
 36. A
 37. B
 38. A
 39. A

 40. B
 41. A
 42. A
 43. A
 44. B
 45. B
 46. C
 47. C
 48. A
 49. B
 50. B
 51. D
 52. C

 53. C
 54. C
 55. D
 56. B
 57. A
 58. A
 59. B
 60. A
 61. D
 62. B
 63. C
 64. D

EXERCISE - 2 : PART # I

 1. A, B, D
 2. C, D
 3. B, D
 4. A, B
 5. A, D
 6. A, B, D
 7. B, C, D
 8. A, B, C, D
 9. A, B

 10. C, D
 11. B, C, D
 12. B, D
 13. A, B, C, D
 14. A, B, C, D
 15. D
 16. B
 17. A
 18. C
 19. A

 20. C
 21. B
 22. B
 23. B
 24. B
 25. A, B, C
 26. A, B
 27. B
 28. A
 29. B
 30. A, B, C

 31. A, B, C, D
 32. A, B, C, D
 33. B, D
 34. A, C

PART # II

1. B 2. A 3. C 4. A 5. A 6. A 7. C 8. A 9. A 10. B 11. A

EXERCISE - 3 : PART # I

1. $A \rightarrow s, B \rightarrow p, C \rightarrow r, D \rightarrow q$			2.	$A \rightarrow q, B \rightarrow p, C \rightarrow r, D \rightarrow q$		
3. $A \rightarrow q, r, B \rightarrow p, s, C \rightarrow p, D \rightarrow r$			4.	$A \rightarrow q, B \rightarrow s, C \rightarrow r, D \rightarrow p$		
					PART # II	
Comprehension #1:		1. B	2. B	3. C		
Comprehension #2:		1. A	2. A	3. B	4. A	5. B
Comprehension #3:		1. A	2. A	3. B	4. A	5. B
Comprehension #4:		1. A	2. B	3. B	4. D	5. C

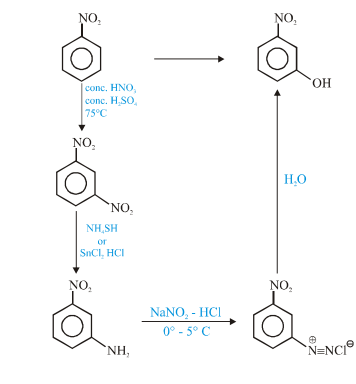
EXERCISE - 5 : PART # I

1. 3 **2.** 2 **3.** 3 **4.** 1 **5.** 2 **6.** 1 **7.** 2 **8.** 1 **9.** 4 **10.** 2

PART # II



3.



4. (P)
$$CH_3 - CH_2 - O - C - CH_3$$
; (Q) CH_3CH_2OH ; (R) $CH_3 - C - CH_3$

When ethylene reacts with dil. $H_2SO_4 CH_3 CH_2^{\oplus}$ is produced during rate determining step, whereas P gives resonance stabilized intermediate.

$$CH_{3} - CH_{2} - O - \bigcup_{\bigoplus}^{C} - CH_{3} \quad \longleftrightarrow \quad CH_{3} CH_{2} - \bigcup_{\bigoplus}^{C} = \bigcup_{i=1}^{C} - CH_{3}$$

Due to extra stability of intermediate the rate of reaction is very fast.



MOCK-TEST

 1. B
 2. C
 3. D
 4. A
 5. A
 6. A
 7. B
 8. B
 9. B
 10. B
 11. D
 12. A
 13. D

 14. A, B, D
 15. A, D
 16. C, D
 17. D
 18. C
 19. A, B, C
 20. A, B, C
 21. A, C
 22. C
 23. A

 24. A
 25. B
 26. A
 27. B
 28. B
 29. C
 30. A→q; B→s; C→r; D→p

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