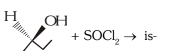
EXERCISE-01

CHECK YOUR GRASP

SELECT THE CORRECT ALTERNATIVE (ONLY ONE CORRECT ANSWER)

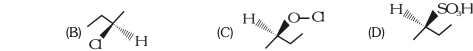
1.



The product formed in the reaction $^{\rm H}_{\rm SOCl_2} \rightarrow {\rm is}^{-}$ $^{\rm "CULTIVATING}$ EXCELLENCE IN EVERY STUDENT" $^{\rm 9814516618}$







2.

proceeds by the mechanism

- (A) S_{N^1}
- (B) S_{N^2}
- (C) $S_N i$

(D) S_{F^2}

- 3. 1, 3- Dibromopropane reacts with metallic zinc to form
 - (A) propene
- (B) cyclopropane
- (C) propane
- (D) hexane

4. Consider the following reaction sequence,

$$CH_3C \equiv CH \xrightarrow{\quad \text{aq.H}_2SO_4 \quad } A \xrightarrow{\quad PCl_5 \quad } B.$$

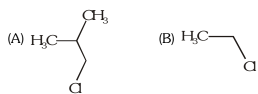
RAKESH SIR "CULTIVATING EXCELLENCE IN EVERY STUDENT"

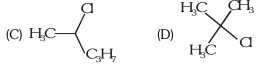
The products (A) and (B) are, respectively,

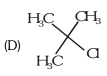
- (A) CH₃COCH₃ and CH₃CCl₂CH₃
- (B) CH₃CH₂CHO and CH₃CH₂CHCl₂
- (C) CH₃CHOHCH₃ and CH₃CHClCH₃
- (D) CH₃CH₂CH₂OH and CH₃CH₂CH₂Cl
- 5. Which of the following has highest dipole moment:
 - (A) CH₂Cl
- (B) CH₃F
- (C) CH₃Br
- (D) CH₃I

- 6. In S_{N^1} the first step involves the formation of
 - (A) free radical
- (B) carbanion
- (C) cabocation
- (D) final product

- 7. To form alkane isonitrile, alkyl halide is reacted with:
 - (A) KCN
- (B) AgCN
- (C) HCN
- (D) NH₄CN
- 8. Which one of the following compounds most readily undergoes substitution by S_N^2 mechanism?







- 9. Sec. Butyl chloride undergo alkaline hydrolysis in the polar solvent by
 - (A) S_{N^2}
- (B) S_{N^1}
- (C) S_{N^1} and S_{N^2}
- (D) none of these

10. Grignard reagent can be prepared by

(A)
$$CH_3$$
— CH_2 — $Cl + Mg \xrightarrow{dry} ether$

$$\begin{array}{c} CH_3 \\ \downarrow \\ (C) CH_3 - C - OH + Mg \xrightarrow{dry \\ ether} \\ CH_3 \end{array}$$

- (D) All of them
- 11. Most stable carbocation formed from $(CH_3)_3C$ -Br, $(C_6H_5)_3CBr$, $(C_6H_5)_2CHBr$ and $C_6H_5CH_2Br$ would be
 - (A) $C_6H_5\overset{\oplus}{C}H_9$

- (B) $(CH_{3/3} \overset{\oplus}{C})$ (C) $(C_{6}H_{5/3} \overset{\oplus}{C})$ (D) $(C_{6}H_{5/2} \overset{\oplus}{C})$
- 12. For the reaction

$$CH_3CH(X)CH_2CH_3 \xrightarrow{alc.\ KOH} CH_3-CH_3-CH_2-CH_3$$

- (A) CH₃—CH=CH—CH₃ predominates
- (B) $CH_9 = CH CH_9 CH_3$ predominates
- (C) Both are formed in equal amounts
- (D) The product ratio depends on the halogen
- 13. The products of reaction of alcoholic silver nitrite with ethyl bromide are
 - (A) Ethane
- (B) Ethene
- (C) Ethul alcohol
- (D) Nitro ethane
- The reaction $CH_0Br + OH^- \longrightarrow CH_0OH + Br^-$ obeys the mechanism 14.

- (B) S_{N^2}
- (C) E₁

- (D) E_2
- 15. Ethylidene chloride can be prepared by the reaction of HCl and
 - (A) Ethane
- (B) Ethylene
- (C) Acetylene
- (D) Ethylene glycol
- 16. 1-phenyl-2-chloropropane on treating with alc. KOH gives mainly
 - (A) 1-phenylpropene

(B) 2-phenylpropene

(C) 1-phenylpropane-2-ol

- (D) 1-phenylpropan-1-ol
- 17. Grignard reagent is obtained when magnesium is treated with
 - (A) Alkyl halide in presence of alcohol
- (B) Alkyl halide in presence of phenol
- (C) Alkyl halide in presence of dry ether
- (D) Alkyl halide in presence of alcoholated ether
- 18. Ethylene reacts with bromine to form -
 - (A) Chloroethane
- (B) Ethylene dibromide
- (C) Cyclohexane
- (D) 1-bromo propane

- $C_2H_4 \xrightarrow{Br_2} X \xrightarrow{KCN} Y ; Y is$ 19.
 - (A) CH₂CH₂CN

(B) NC—CH₂—CH₂—CN

(C) Br—CH₂—CH₂CN

- (D) Br—CH=CHCN
- 20. Reactivity order of halides for dehydrohalogenation is
 - (A) R F > R Cl > R Br > R I
- (B) R I > R Br > R Cl > R F
- (C) R I > R Cl > R Br > R F
- (D) R F > R I > R Br > R Cl
- 21. Which of the following is least reactive in a nucleophilic substitution reaction
 - (A) CH₂=CHCl
- (B) CH₂CH₂Cl
- (C) CH_o=CHCH_oCl
- (D) (CH₂)₂C-Cl

22. The correct reactivity order of alcohols towards H-X will be

$$(II)$$
 H_3 C \longrightarrow CH

(III)
$$CH_3$$
— CH_2 — OH

- VI < III < I > III > IV
- (B) IV > III > II > I
- (C) II > IV > I > III
- (D) II > IV > III > I
- Identify 'Z' in the following reaction series, CH_3 : $CH_2CH_2Br \xrightarrow{aq.NaOH} (X) \xrightarrow{Al_2O_3} (Y) \xrightarrow{HOCl} (Z)$: 23.

24. For the reaction,

 $C_0H_1OH + HX \xrightarrow{ZnX_2} C_0H_1X$, the order of reactivity is

- (A) HI > HCl > HBr

- (B) HI > HBr > HCI (C) HCI > HBr > HI (D) HBr > HI > HCI
- 25. Ethyl alcohol reacts at a faster rate with HI than with HCl in forming the corresponding ethyl halides under identical conditions mainly because -
 - (A) HI, being a stronger acid, protonates ethyl alcohol at oxygen much better and helps substitution
 - (B) the bond length in HI is much shorter than that in HCl
 - (C) I is a much better leaving group

RAKESH SIR

(D) I is a much better nucleophile than Cl-

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CHECK YOUR GRASP						A	ANSWER KEY				EXERCISE -1				
Que.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15
Ans.	В	В	В	Α	Α	С	В	В	С	А	С	D	D	В	С
Que.	16	17	18	19	20	21	22	23	24	25					
Ans.	Α	С	В	В	В	Α	D	В	В	D					

EXERCISE-02 BRAIN TEASERS

SELECT THE CORRECT ALTERNATIVES (ONE OR MORE THEN ONE CORRECT ANSWERS)

- 1. Which of the following does/do produce a white precipitate of AgCl on warming with alcoholic silver nitrate?
 - (A) Allyl chloride
- (B) t-Butyl chloride
- (C) Benzyl chloride
- (D) Vinyl chloride
- 2. What is the order of reactivity of the following compounds towards nucleophilic substitution?









- (A) I < II < III < IV
- (B) IV < III < II < I
- (C) IV < I < II < III
- (D) IV < II < I < III

- **3.** The order of decreasing nucleophilicity of the following is
 - (A) $H_2O > OH^- > CH_3COO^- > CH_3O^-$
- (B) $CH_{3}O^{-} > OH^{-} > CH_{3}COO^{-} > H_{9}O$
- (C) $CH_3COO^- > CH_3O^- > OH^- > H_2O$
- (D) $HO^{-} > CH_{3}O^{-} > CH_{3}COO^{-} > H_{9}O$
- **4.** The order of decreasing $S_N 1$ reactivities of the halides

CH₂CH₂CH₂Cl

CH₂= CHCHClCH₃

CH₃CH₂CHClCH₃

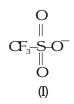
I (A) I > II > III

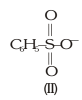
(B) II > I > III

(C) II > III > I

(D) III > II > I

5. Consider the following anions.





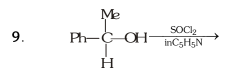




When attached to sp^{3} - hybridized carbon, their leaving group ability in nucleophilic substitution reactions decreases in the order

- (A) I > II > III > IV
- (B) I > II > IV > III
- (C) IV > I > II > III
- (D) IV > III > II > I
- **6.** The basicity of RO⁻,HO⁻, RCOO⁻, ROH, and H₂O are of the order -
 - (A) $HO^{-} > RO^{-} > H_{9}O > ROH > RCOO^{-}$
 - (B) $RO^{-} > HO^{-} > RCOO^{-} > ROH > H_{0}O$
 - $(C) H_{\circ}O > ROH > RCOO^{-} > HO^{-} > RO^{-}$
 - (D) ROH $> H_{\circ}O > HO^{-} > RCOO^{-} > RO^{-}$
- 7. Which of the following are aprotic solvents:
 - (A) DMSO
- (B) DMF
- (C) H₂O
- (D) CH₃COOH

- 8. Which is/are true statements (s):
 - (A) Protonation increases electrophilic nature of carbonyl group
 - (B) $CF_3SO_3^-$ is better leaving group than $CH_3SO_3^-$
 - (C) Benzyl carbonium ion is stabilised by resonance
 - (D) $CCl_3CH < OH$ is stable, due to H-Bonding



Which statement is true for the above reaction?

(A) Retention of configuration

(B) Inversion of configuration

- (C) Inversion and Retention both
- (D) None
- 10. Which of the following undergoes hydrolysis most easily



$$(C) \bigcirc NO_2$$

$$NO_2$$

$$(D) \bigcirc O$$

(C)
$$NO_2$$
 NO_2

- 11. A compound 'A' formula of C₃H₆Cl₂ on reaction with alkali can give 'B' of formula C₃H₆O or 'C' of formula C_3H_4 . 'B' on oxidation gave a compound of the formula $C_3H_6O_2$. 'C' with dilute H_2SO_4 containing Hg^{2+} ion gave 'D' of formula C_3H_6O , which with bromine and alkali gave the sodium salt of $C_2H_4O_2$. Then 'A' is
 - (A) CH₃CH₂CHCl₂
- (B) CH₃CCl₂CH₃
- (C) CH₂ClCH₂CH₂Cl (D) CH₃CHClCH₂Cl
- 12. Isobutyl magnesium bromide with dry ether and absolute alcohol gives
 - (A) CH₃—CH—CH₂OH and CH₃CH₂MgBr
- (C) CH_3 — CH_3 — CH_3 , CH_2 = CH_2 and Mg(OH)Br (D) CH_3 — CH_3 and CH_3CH_2OMgBr CH_3
- 13. Following reaction is

$$CH_{3}(CH_{2})_{5} \xrightarrow{\overset{H}{\equiv}} C-Br \xrightarrow{OH} HO - \overset{H}{C} \xrightarrow{\overset{H}{\equiv}} (CH_{2})_{5}CH_{3}$$

(A) E,

(C) E_o

- (D) S_{N^2}
- 14. On treatment with chlorine in presence of sunlight, toluene gives the product -
 - (A) o-chloro toluene

(B) 2, 5-dichloro toluene

(C) p-chloro toluene

- (D) Benzyl chloride
- 15. In S_{N^1} reaction an optically active substrates mainly gives :
 - (A) Retention in configuration

(B) Inversion in configuration

(C) Racemic product

- (D) No product
- 16. Alkyl iodides can be prepared by :-

 - (A) $RCH_2COOAg + I_2 \xrightarrow{CCl_4} RCH_2I$ (B) $RCH_2Cl + NaI \xrightarrow{acetone} RCH_2I + NaCl$
 - $(C)R OH + HI \longrightarrow RI + H_2O$
- (D) $CH_4 + I_2 \longrightarrow CH_3I$

- 17. Which of the following reagents can be used to prepare an alkyl halide:-
 - (A) NaCl
- (B) $HCl + ZnCl_2$
- (C) SOCl₂
- (D) PCl₅
- **18.** Which of the following reactions depict the nucleophilic substitution of C_2H_5Br :
 - (A) $C_2H_5Br + C_2H_5SNa \longrightarrow C_2H_5SC_2H_5 + NaBr$
 - (B) $C_2H_5Br \xrightarrow{Na+C_2H_5OH} C_2H_6 + HBr$
 - (C) $C_9H_5Br + AgCN \longrightarrow C_9H_5NC + AgBr$
 - (D) $C_2H_5Br + KOH \longrightarrow C_2H_5OH + KBr$
- 19. For an $\,S_{N^2}\,$ reaction, which of the following statements are true :-
 - (A) The rate of reaction is independent of the concentration of the nucleophile
 - (B) The nucleophile attacks the C-atom on the side of the molecule opposite to the group being displaced
 - (C) The reaction proceeds with simultaneous bond formation and rupture
 - (D) None of these
- **20.** Which of the following is an S_{N^2} reaction:
 - (A) $CH_3CH_2Br + KOH \longrightarrow CH_3CH_2OH + KBr$
 - (B) $CH_3CH_2Br + CH_3CH_2ONa \longrightarrow CH_3CH_2OCH_2CH_3 + NaBr$
 - (C) $(CH_3)_3CBr + KOH \longrightarrow (CH_3)_3COH + KBr$

BRAIN TEASERS						F	ANSWER KEY			EXERCISE -2					
Que.	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15
Ans.	A,B,C	С	В	С	В	В	A,B	A,B,C,D	В	D	Α	D	D	D	С
Que.	16	17	18	19	20										
Ans.	B.C	B,C,D	A,C,D	В,С	A,B										

TRUE OR FALSE:

1. Alkyl halides follow the reactivity sequence,

$$R-I > R-Br > R-Cl > R-F$$

- 2. Vinyl chloride reacts with dilute NaOH to form vinyl alcohol.
- 3. Allyl chloride is more reactive than vinyl chloride.
- **4.** Tertiary butyl bromide undergoes $S_N 1$ reactions.
- **5.** Both vic- and gem-dihalides on heating with zinc dust in presence of alcohol form same alkene.

FILL IN THE BLANKS:

- 1. The interaction of elemental sulphur with Grignard reagent gives
- 2. An alkyl halide may be converted into alcohol byreaction
- **4.** Allyl chloride is a compound while vinyl chloride is inert towards nucleophilic substitution.
- 6. Alkyl halides are formed when thionyl chloride and are refluxed in presence of pyridine.
- 7. Ethylene chloride on hydrolysis with ag. KOH forms..........

MATCH THE COLUMN

1. Match the column I with column II.

	Column-I (reaction)	Co	lumn-II (Mechanism)
(A)	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	(p)	$S_{_{\rm N}}1$
(B)	Phr (+) H +SOCI ₂ Pyridine H ₃ C H	(q)	S _N 2
(C)	$CH_{3}-CH-Br + \overset{\Theta}{OC_{2}}H_{5}/EtOH \longrightarrow CH_{3}$	(r)	$S_{_{ m N}}$ i
(D)	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	(s)	E_2

2. Match the column I with column II.

	Column-I (Substrate)		Column-II (Relative rate of solvolysis in 50% aqueous ethanol at 45°C)
(A)	≫_a	(p)	7700
(B)	\wedge a	(q)	1
(C)	H ₃ C O	(r)	91
(D)	Ph/\d	(s)	1,30,000

ASSERTION & REASON QUESTION:

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: Iodination of akanes is carried out by heat in presence of readucing agent.

Because

Statement-II: Iodination of alkanes takepalce explosively.

2. Statement-I: Chloropropane has higher boiling point than chloroethane.

Because

Statement-II: Haloalkanes are polar molecules.

3. Statement-I : Polar solvent slows down S_{N^2} reaction.

Because

Statement-II: CH_3 -Br is less reactive than CH_3Cl .

4. Statement-I: Primary benzylic halides are more reactive than primary alkyl halides towards S_{N^1} reaction.

Because

Statement-II: Reactivity depends upon the nature of the nucleophile and the solvent.

5. Statement-I: Vinylic halides are reactive towards nucleophilic substitution reaction.

Because

Statement-II: Reactivity is due to the polarity of carbon-halogen bond.

6. Statement-I: Aryl halides undergo electrophilic substitution less readily than benzene.

Because

Statement-II: Aryl halide gives only meta product w.r.t. electrophilic substitution.

7. Statement-I: Optically active 2-iodibutane on treatment with NaI in acetone undergoes racemisation.

Because

Statement-II: Repeated Walden inversions on the reactant and its product eventually gives a racemic mixture.

Statement-I: Free radical chlorination of n-butane gives 72% of 2-chlorobutane and 28% of 1-chlorobutane though it has six primary and four secondary hydrogens.

Because

Statement-II: A secondary hydrogen is abstracted more easily than the primary hydrogen.

9. **Statement-I**: Nucleophilic substitution reaction on an optically active alkyl halide gives a mixture of enantiomers.

Because

Statement-II : The reaction occurs by $\textbf{S}_{_{N^{1}}}$ mechanism.

10. Statement-I: Boiling point of alkyl halide increases with increase in molecular weight.

Because

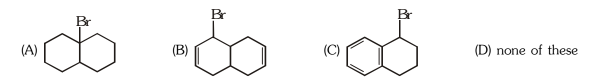
Statement-II: Boiling point of alkylhalides are in the order RI > RBr > RCI > RF.

COMPREHENSION BASED QUESTIONS:

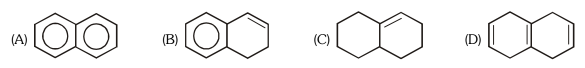
Comprehension # 1

An organic compound A has molecular formula $C_{10}H_{17}Br$ and it is non-resolvable. A does not decolourize brown colour of bromine water solution. A on treatment with $(CH_3)_2COK/(CH_3)_3COH$ yields B as major product. B on treatment with H_2/Pt yields $(C_{10}H_{16})$ which on treatment with Cl_2/hv yields three monochloro derivative. Also B on boiling with acidic permanganate solution yields $C(C_{10}H_{16}O_3)$. C on heating with sodalime yields D $(C_9H_{16}O)$. D on reducing with LiAlH $_4$ followed by heating the product with concentrated H_2SO_4 yields E (C_9H_{16}) as major product. E on treatment with ozone followed by work-up with Zn– H_2O yields 6-Ketononanal.

1. Compound A is:



2. Compound B is :



3. Compound C is:

4. Compound D is:

$$(A) \bigcirc O \qquad (B) \bigcirc O \qquad (C) \bigcirc O \qquad (D) \bigcirc O \qquad (D)$$

Comprehension # 2

Nucleophilic substitution reactions generally expressed as

$$Nu^- + R - L \longrightarrow R - Nu + L^-$$

Where $Nu^- \longrightarrow Nucleophile$; $R \longrightarrow L \longrightarrow substrate$; $L \longrightarrow leaving$ group

The best leaving groups are those that become the most stable ions after they depart. Since most leaving groups leave as a negative ion, the best leaving groups are those ions that stabilize a negative charge most effectively. A good leaving group should be

- (a) electron-withdrawing to polarize the carbon
- (b) stable once it has left (not a strong base)
- (c) polarisable- to maintain partial bonding with the carbon in the transition state (both $S_N 1$ and $S_N 2$). This bonding helps to stabilise the transition state and reduces the activation energy.

1. Among the following which is feasible?

(A)
$$X^- + CH_3 - CH_2 - H \longrightarrow CH_3 - CH_2 - X + H^-$$

(B)
$$X^- + CH_3 -OH \longrightarrow CH_3 -X + \overline{O}H$$

(C)
$$X + H_3C \longrightarrow CH_3 - X + H_2O$$

(D)
$$X^- + CH_3 - CH_3 - CH_3 - X + \overline{C}H_3$$

- 2. Among the following which is false statement?
 - (A) The weaker the base after the group departs, the better the leaving group
 - (B) A reactive leaving group would raise the energy of the product, driving the equilibrium towards the
 - (C) Relative leaving group ability may vary with change of solvent
 - (D) Better leaving group only increases S_N^2 rate, not S_N^2 1.

$$\textbf{3}. \hspace{1cm} \begin{array}{cccc} CH_3Br & CH_3F & CH_3OH & CH_3OSO_2CF_3 \\ \text{ $_{(I)}$} & \text{ $_{(II)}$} & \text{ $_{(II)}$} \end{array}$$

The correct order of decreasing reactivity of the above compounds towards CH_3O^- in an S_N2 reaction is:

(A)
$$I > IV > II > III$$

$$(B) IV > I > II > II$$

(C)
$$IV > I > III > II$$

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

The correct order of increasing leaving group capability of above anoins

(A)
$$III < IV < II < I$$

(B)
$$II < III < I < IV$$

(C)
$$II < IV < III < I$$

(D)
$$I < III < II < IV$$

Comprehension # 3

Nucleophilic aliphatic substitution reaction is mainly of two types : $S_N 1$ and $S_N 2$. The $S_N 1$ mechanism is a two step process. Reaction velocity of $S_{
m N}1$ reaction depends only on the concentration of the substrate. Since product formation takes place by the formation of carbocation, optically active substrate gives (+) and (-) forms of the product. In most of the cases the product usually consits of 5-20% inverted product and 80-95% racemised species. The more stable the carbocation, the greater is the proportion of racemisation. In solvolysis reaction, the more nucleophilic the solvent, the greater is the proportion of inversion.

1. Which one of the following compound will give $S_N 1$ reaction predominantly?

(A)
$$H_5C_6$$
 B_r (B) H_3C (C) H_3C B_r (D) All of these

2. Which of the following compounds will give $S_N 1$ and $S_N 2$ reactions with considerable rate ?

Select the correct answer from the codes given below

3. For the given reaction

$$\begin{array}{c|c} R_1 & R_1 \\ \hline R & R & R_1 \\ \hline R_2 & R_2 \end{array} OH$$

Which substrate will give maximum racemisation?

(A)
$$H_5C_6$$
 $\xrightarrow{CH_3}$ B_r C_2H_5

$$(B) \begin{array}{c|c} H_2C & CH_3 \\ \hline & Br \\ C_2H_5 \end{array}$$

(C)
$$H_3C_6$$
 CH_3

Comprehension #3: 1. (A)

$$H_{5}C_{6} \xrightarrow{Br} NO_{2}$$

$$(D) \qquad \qquad \downarrow \\ NH_{3}$$

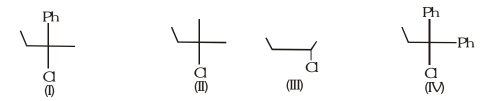
ISCELLANEOUS TYPI	E QUESTION	AN	SWER KE	Y		EXERCISE -3
True / False						
1 . T	2. F	3. T	4. T	5. F		
Fill in the Bla	nks					
 thioalcohol reactive glycol 	<u>-</u>	lic substitution es or alkylene		3. sodium6. alcoho	,	villiamson's synthesis
Match the Co.	<u>lumn</u>					
1. (A) \rightarrow r; B \rightarrow c	$l_1 : (C) \rightarrow s : (D)$	\rightarrow p	2. (A) –	\rightarrow q; (B) \rightarrow r;	$(C) \rightarrow s ; (D)$	\rightarrow p
Assertion - Re	ason Questic	ons				
1 . D	2. B	3. C	4. B	5 . D	6. C	7. A
8. A	9 . A	10 . B				
Comprehensio.	n Based Qu	estions				
Comprehension Comprehension			3 . (A) 3 . (B)	4. (C) 4. (B)		

3. (C)

2. (A)

1. Arrange the following compounds in order of :

Decreasing $S_{_{\!N}}1$ reaction rate :



- 2. Select the member of each pair that shows faster rate of S_N^2 reaction with KI in acetone.
 - (a) $\mathrm{CH_3-CH_2-CH_2-CH}$ and $\mathrm{CH_3-CH-CH_2-CI}$ and $\mathrm{CH_3-CH-CH_2-CI}$ $\mathrm{CH_3}$
 - (I) (II) (II) (b) $CH_3-CH_2-CH_2-CI$ and $CH_3-CH_2-CH_2-Br$ (I) (II)
 - (c) CH₃-CH-CH₂-CH₂-C and CH₃-C-CH₂CI
 CH₃
 (I)
 (II)

(d)
$$CH_3$$
 CH_2 CH_2 CH_3 CH_3 CH_3 CH_4 CH_5 $CH_$

- 3. Of the following statements which are true for $S_N 1$ reaction.
 - (a) Tertiary alkyl halides react faster than secondary.
 - (b) The absolute confuguration of the product is opposite to that of the reactant when an optical active substrate is used.
 - (c) The reaction shows first order kinetics.
 - (d) The rate of reaction depends markedly on the nucleophilicity of the nucleophile.
 - (e) The mechanism is two step.
 - (f) Carbocations are intermediate.
 - (g) Rate ∝ [Alkyl halides]
 - (h) The rate of the raction depends on the nature of the leaving group.
- **4.** Of the following statements, which are true for S_N^2 reaction.
 - (a) Tertiary alkyl halides reacts faster than secondary.
 - (b) The absolute configuration of product is opposite to that of the reactant when an optically active substrate is used.
 - (c) The reaction shows first order kinetics.
 - (d) The rate of the reaction depends markedly on the nucleophilicity of the attacking reagent.
 - (e) The mechanism is one step.

- (f) Carbocations are intermediate.
- (g) Rate ∞ [Alkyl halides]
- (h) The rate of the raction depends on the nature of the leaving group.
- 5. Arrange the isomers of molecular formula C_4H_9Cl in order of decreasing rate of reaction with sodium iodide in acetone.
- 6. There is an overall 29-fold difference in reactivity of 1-chlorohexane, 2-chlorohexane towards potassium iodide in acetone. Which one is the most reactive? why?
- 7. Identify the product when A reacts with $CH_2CH = CH_2$
 - (a) Br₂/Fe
- (b) Br₂/CCl₄
- (c) NBS
- (d) HBr
- $\textbf{8.} \qquad \text{Identify major product in the following}: \ \text{CICH}_2\text{CHCl}_2 \xrightarrow{\quad \text{OH}^- \quad} \text{CH}_2 = \text{CCl}_2$
 - (a) CH3CH2CHCH3 ETO-

- (b) CH₃CH₂CH₂CH₂Br EtO
- (c) CH2CH2CHCH3 EtO-Br
- (d) H₃CCCH₃CH₂CH₃CCK (d) H₃CCCH₃CCH (dH₃CC)H

- (e) EtO-EtOH
- 9. Which is faster in the following pairs of halogen compounds undergoing $S_{\scriptscriptstyle N}2$ reactions?
 - (a) \bigcap and \bigcap
- (b) \prod and \bigcap
- (c) \longrightarrow and \longrightarrow \bigcirc
- (d) Br and
- 10. R-Mg-Br (A) on reaction with H_2O forms a gas (B), which occupied 1.4 L/g at NTP. What is product when R-Br reacts with benzene in presence of $AlCl_3$?

- 1. IV > I > II > III
- 2. (a) I (b) II (c) I (d) I
- **3**. (a) T (b) F (c) T (d) F (e) T (f) T (g) T (h) T
- **4.** (a) F (b) T (c) F (d) T (e) T (f) F (g) F (h) T
- **5.** $1^{\circ} > 2^{\circ} > 3^{\circ}$ Anion of acetic acid is more stabilised by resonance than phenoxide ion.
- **6.** 1-chlorohexane

Because it follows Sn2 path.

7. (a) $CH_2CH = CH_2$ and para Ortho

$$(d) \begin{array}{|c|c|} \hline CHCH_2CH_3 \\ \hline Br \end{array}$$

(1, 2-H shift gives more stable benzylic carbocation)

8. (a) $CH_3CH = CHCH_3$ (b) $CH_3CH = CHCH_3$ (c) $CH_3CH_2CH_3$

- 9. Ease of backside attack (less steric hindrance) decides which undergoes S_N^2 faster (except in (b) in which iodide is better leaving group). In all cases first one is fater than the other for S_N^2 reaction.
- 10. Gas B is CH₄, hence A is CH₃MgBr. CH₃Br forms CH₃ on reaction with benzene.

- 1. Explain the following observations:
 - (a) Azide ion (N_3^-) react with 2- bromopentane thousand times faster than with neopentyl bromide in a S_N^- 2 reaction though former is a secondary halide while latter is primary.
 - (b) What will happen to the stereochemistry of product of the following reaction:\

$$H \xrightarrow{Br} CH_3 + N_3^{-} \xrightarrow{S_1 2}$$

- (c) What will happen to the rate if the concentration of alkyl bromide in (b) is doubled?
- (d) What will happen to the rate if the concentration of azide ion in (b) is doubled?
- (e) How the sign of optical rotation of reactant and product are related in (b)
- (f) When allowed to stand in dilute H_2SO_4 , laevo-rotatory 2-butanol slowly loses optical activity.
- 2. Provide structure of major product in the following reaction indicating stereochemistry where appropriate:

3. Propose mechanism of the following reactions:

4. Which of the following alkyl halide could be successfully used to synthesize Grignard reagent and why other fail?

$$HO$$
 (I)
 O
 (III)
 BI
 (III)
 BI
 (III)
 BI
 (III)
 BI

- An alkyl bromide A has molecular formula $C_8H_{17}Br$ and four different structures can be drawn for it, all optically active. A on refluxing with ethanolic KOH solution yields only one elimination product $B(C_8H_{16})$ which is still enantiomeric. B on treatment with H_2/Pt yields $C(C_8H_{18})$ which does not rotate the plane polarized light, B on ozonolysis followed by work-up with H_2O_2 yields $D(C_7H_{14}O)$ as one product which is still resolvable. Deduce structures of A to D.
- **6.** Identify A to G in the following.

(a)
$$A \xrightarrow{Br_2CCl_4} A \xrightarrow{KCN} B \xrightarrow{H_3O^+} C$$

(b)
$$O \xrightarrow{Br_2COl_4} D \xrightarrow{KON} E \xrightarrow{H_3O^+} G$$

7. Br
$$\xrightarrow{(A)}$$
 C $\xrightarrow{Mg/ether}$ C $\xrightarrow{D_2O}$ C $\xrightarrow{Na/ether}$ D $\xrightarrow{Na/ether}$ D $\xrightarrow{Na/ether}$ C $\xrightarrow{Na/ether}$ D $\xrightarrow{Na/ether}$ D $\xrightarrow{Na/ether}$ D $\xrightarrow{Na/ether}$ D

Identify B to F

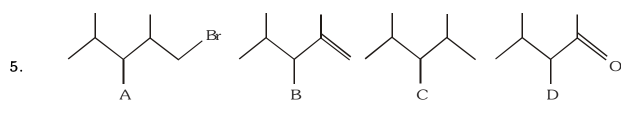
- 8. Vinyl chloride does not give S_N reaction but allyl chloride gives. Explain.
- 9. Arrange the following in the increasing order of their ability as a leaving group:
 - (a) CH_3S^- , CH_3O^- , CF_3^- and F^-
 - (b) $CF_3SO_3^-$, $CH_3SO_3^-$ and CH_3COO^-
- 10. RBr when treated with AgCN in a highly polar solvent gives RNC whereas when it is treated with NaCN it gives RCN. Explain.

1. (a) Though neopentylbromide is primary, bulky tertiary butyl group possess very large steric hindrance to the attack of bulky nucleophile N_3^- .

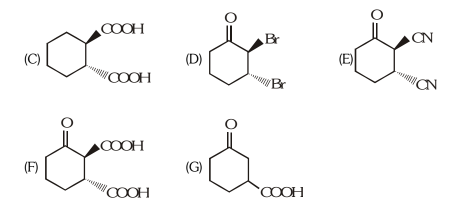
(b)
$$H \longrightarrow CH_3 + N_3^- \xrightarrow{S_1/2} CH_3 \longrightarrow H$$

- (c) Rate will double
- (d) Rate will double
- (e) not related
- (f) Recemization occur through carbocation intermediate

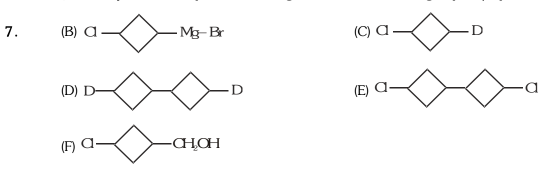
4. Only II can be used for successful synthesis of Grignard reagent, rest all contain acidic proton and will react with R^- (from Grignard reagent) forming alkane.



Product• H₂O



(decarboxylation takes place on heating when there is a keto group at β - position)



8. In Vinyl chloride, C - Cl bond is stable due to resonance (as in chlorobenzene)

Hence S_N reaction in which CI is replaced by nucleophile is not possible. In addition to this, sp^2 - hybridised carbon is more acidic than sp^3 - carbon, hence removal of proton (H⁺) is easier than removal of halide (Cl⁻) In allyl chloride, S_N reaction is easier since allyl carbocation formed after removal of Cl⁻ is stabilised by resonance

$$CH_2 = CHCH_2CI \longrightarrow CH_2 = CHCH_2+CI$$
Allyl carbocation

$$CH_2 = CH - CH_2 \longleftrightarrow CH_2 - CH = CH_2$$

- 9. (a) $CF_3^- < CH_3O^- < CH_3S^-$; (b) $CH_3COO^- < CH_3SO_3^- < CF_3SO_3^-$
- 10. As $[CN]^-$ is an ambident nuicleophile which abve two nucleophile which have two nucleophilic sites and can attack from either side. In a highly polar solvent, AgCN promotes the formation of carbocation R^+ , precipitation of AgBr.

In the absence of such promotion by Ag^+ , with $Na+[CN]^-$, the resulting S_N^2 reaction is found to proceed with preferential attack on the atom in the nucleophile which is more polarisable i.e. C.

$$NC^-\!\!+\!R-Br \longrightarrow [NC^{\delta^-}\!....R....Br^{\delta^-}] \longrightarrow N \equiv C-R+Br^-$$
 Transition State

1.	Chlorination of toluene in the presence of light and heat followed by treatment with aqueous NaOH gives											
	(A) o-cresol		(B) p-cresol	(B) p-cresol								
	(C) 2, 4 dihydroxytolu	ene	(D) Benzoic acid	(D) Benzoic acid								
2.	Aryl halides are less reactive towards nucleophilic substitution reaction as compared to alkyl halides d											
	(A) The formation of I	ess stable carbonium ion	(B) Resonance stabiliza	(B) Resonance stabilization								
	(C) The inductive effe	et	(D) sp^2 hybridised carl	(D) sp^2 hybridised carbon attached to the								
3.	1-Chlorobutane on re	action with alcoholic potas	sh gives -	ives -								
	(A) 1-butene	(B) 1-butanol	(C) 2-butene	(D) 2-butane	ol							
4.	In the addition of HB	r to propene in the absen	ce of peroxides, the first st	tep involves the	addition of -							
	(A) H ⁺	(B) Br ⁻	(C) H•	(D) Br•	[IIT-93]							
5.	Arrange the following	compounds in order of in	creasing dipole moment :		[IIT-96]							
	(I) Toluene ; (II) m-dich	(I) Toluene ; (II) m-dichlorobenzene ; (III) o-dichlorobenzene ; (IV) p-dichlorobenzene										
	(A) I < IV < II < III	(B) $IV < I < II < III$	(C) $I < IV < II < III$	(D) IV < II <	I < III							
6.	In the reaction of p-cl	nloro toluene with KNH_2 in	n liq. NH_3 , the major prod	luct is -	[IIT-97]							
	(A) o-toluidine	(B) m-toluidine	(C) m-chloroaniline	(D) p-chloro	aniline							
7.	(CH ₃) ₃ CMgCl reactio	n with D ₂ O produces :			IIT-97]							
	(A) (CH ₃) ₃ CD	(B) (CH ₃) ₃ OD	(C) (CD ₃) ₃ CD	(D) (CH ₃) ₃ OI	D							
8.	The intermediate duri	ng the addition of HCl to	propene in presence of pe	eroxide is -	[IIT-97]							
	(A) CH3CHCH2C1	(B) CH3CHCH3	(C) $CH_3\dot{C}HCH_2$	(D) CH3CH	ġĊH ₂							
9.	The number of possib	The number of possible enantiomeric pairs that can be produced during monochlorination of isopentane										
	(A) 2	(B) 3	(C) 4	(D) 1	[IIT-97]							
10.	During debromination is -	of meso-2,3-dibromobutar	ne, with Zn dust/CH ₃ COO	H the major com	npound formed [IIT-97]							
	(A) n-butane	(B) 1-butene	(C) cis-2-butene	(D) trans-2-b	outene							
11.	Benzyl chloride (C ₆ H	₅ CH ₂ Cl) can be prepared t	from toluene by chlorinatio	on with	[IIT-98]							
	(A) SO_2Cl_2	(B) SOCl ₂	(C) PCl ₅	(D) NaOCl								
12.	Toluene, when treated	d with Br ₂ /Fe, gives o and	l p-bromotoluene, because	the CH ₃ group) -							
	(A) is ortho and para	directing	(B) is meta directing	(B) is meta directing								
	(C) deactivates the ring by hyperconjugation (D) deactivates the ring											
13.	A solution of (+) 2 -ch $SbCl_5$ due to the form		uene racemises slowly in th	ne presence of sn	mall amount of [IIT-99]							
	(A) Carbanion	(B) Carbene	(C) Free-radical	(D) Carbocat	tion							
14.	The order of reactivity	y of the following alkyl hal	lides for a S_{N^2} reaction is	s for a ${}^{{}^{\circ}}\!\!\!\!S_{N^2}$ reaction is -								
	(A) $RF > RCl > R - B$	r > R-I	(B) $R-F > R -Br > R$	(B) $R-F > R -Br > R-Cl > R-I$								
	(C) $R-Cl > R -Br > R-F > R-I$ (D) $R-I > R -Br > R-Cl > R-F$											
15 .	Which of the following	g has the highest nucleoph	ilicity:		[IIT-2000]							
	(A) F ⁻	(B) OH ⁻	(C) CH ₃ ⁻	(D) $\mathrm{NH_2}^-$								
16.	An S_{N^2} reaction at an	An S_{N^2} reaction at an asymmetric carbon of a compound always gives :										
	(A) an enantiomer of	the substrate	(B) a product with opp	(B) a product with opposite optical rotation								
	(C) a mixture of diaste	ereomers	(D) a single stereoison	(D) a single stereoisomer								

- 17. The number of isomer for the compound with molecular formula C_2 BrCl FI is [IIT-01] (A) 3 (B) 4 (C) 5 (D) 6
- 18. In the presence of peroxide, hydrogen chloride and hydrogen iodide do not give anti-Markovnikov's addition to alkenes because : [IIT-01]
 - (A) both are highly ionic
 - (B) one is oxidising and the other is reducing
 - (C) one of the steps in endothermic in both the cases
 - (D) all the steps are exothermic in both the reactions
- 19. Identify the set of reagents/reaction conditions 'X' and 'Y' in the following set of transformations -[IIT-02]

$$CH_3-CH_2-CH_2Br \xrightarrow{X} Product \xrightarrow{Y} CH_3-CH-CH_3$$

- (A) X = dilute aqueous NaOH, 20°C, Y = HBr/acetic acid, 20°C
- (B) X = concentrated alcoholic NaOH, 80°C ; Y = HBr/acetic acid, 20°
- (C) X = dilute aqueous NaOH 20°, $Y = Br_2/CHCl_3$, 0°C
- (D) X = concentrated alcoholic NaOH, 80° C; Y = $Br_2/CHCl_3$, 0° C

20.
$$F \longrightarrow NO_2 \xrightarrow{(CH_3)_2NH} (A) \xrightarrow{(i) NaNO_2 + HO 0^\circ -5^\circ C} (B)$$
 [IIT-03]

$$(D) \begin{array}{c} H_3C \\ H_3C \end{array} N - \begin{array}{c} \\ \\ \end{array} N - \begin{array}{c} \\ \\ \end{array} N O_2 \end{array}$$

21. MeO NO
$$_2$$
 on hydrolysis in presence of acetone : [IIT-05]

- (A) K & L
- (B) only L
- (C) M only
- (D) K & M

22. The reagent (s) for the following conversion:

$$\frac{Br}{Br} + HC \equiv CH \text{ is /are -}$$

(A) alcoholic KOH

(B) Alcoholic KOH followed by NaNH₂

(C) aquesous KOH followed by NaNH₂

(D) Zn/CH₃OH

23. In the following groups: [IIT-97]

[IIT-07]

(I) -OAc; (II) -OMe; (III) $-OSO_2Me$; (IV) OSO_2CF_3

the order of leaving group ability is -

(A) I > II > III > IV

(B) IV > III > I > II

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

CH₂CH₂CHCl₂ Aq. KOH ? 24.

[IIT 1991]

25. Draw the stereochemical structure of the products in the following reaction.

[IIT 1994]

$$Br \xrightarrow{C_2H_5} H \xrightarrow{\stackrel{\circ}{\text{OH/DMSO}}} S_{N^2},$$

26. Give reasons: [IIT 2005]

(a) (i)
$$CH_3$$
 $C_2H_5OH(aq.)$ acidic solution ; (ii) CH_3 $C_2H_5OH(aq.)$ neutral CH_3

(b) (i)
$$O_2N$$
 O_3 O_2N O_3 O_2N O_3 O_3 O_4 O_4 O_5 O_5 O_5 O_5 O_5 O_6 O_7 O_8 O_8

- 27. An alkyl halide (A) of formula $C_6H_{11}Cl$ on treatment with potassium tertiary butoxide gives two isomeric alkenes (B) and (C) C_6H_{10} .Both alkene on hydrogenation give methycyclopentane. Predict the structure of (A), (B) and (C).
- 28. Which would be the major product in each of the following reactions?

(a)
$$H_3C$$
 $\xrightarrow{CH_3}$ $\xrightarrow{C_2H_5OH}$ $\xrightarrow{\Delta}$ \xrightarrow{C}

(b)
$$NO_2$$
 $NaOCH_3$ A

(c)
$$\xrightarrow{\text{base}}$$

(K) MeO
$$\longrightarrow$$
 H \longrightarrow NO₂ \longrightarrow NO₂

(L) MeO
$$\longrightarrow$$
 CH₃ H CH₃ NO.

It mainly gives

(A) K and L

(B) Only K

(C)L and M

(D) Only M

[IIT 2006]

30. Match the following:

Column I

- (A) ${\rm CH_3-CHBr-CD_3}$ on treatment with alc. KOH gives ${\rm CH_2=CH-CD_3}$ as a major product.
- (B) $Ph-CHBr-CH_3$ reacts faster than $Ph-CHBr-CD_3$.
- (C) Ph-CD $_2$ -CH $_2$ Br on treatment with C $_2$ H $_5$ OD/C $_2$ H $_5$ O gives Ph-CD=CH $_2$ as the major product.
- (D) $\rm PhCH_2CH_2Br$ and $\rm PhCD_2CH_2Br$ react with same rate.
- **31.** The major product of the following reaction is

Column II

- (P) E1 reaction
- (Q) E2 reaction
- (R) E1cb reaction
- (S) First order reaction

[IIT 2008]

$$(A) \qquad \begin{array}{c} H_3C \\ \hline \\ NO_2 \end{array}$$

$$(D) \qquad \begin{array}{c} H_3C \\ SPh \\ SPh \\ NO_2 \end{array}$$

- 33. The total number of alkenes possible by dehydrobromination of 3-bromo-3-cyclopentylhexane using alcoholic KOH is [IIT-2011]
- 34. KI in acetone, undergoes S_N^2 reaction with each of P, Q, R and S. The rates of the reaction vary as [IIT 2013]

(A)
$$P > Q > R > S$$

(B)
$$S > P > R > Q$$

(C)
$$P > R > Q > S$$

(D)
$$R > P > S > Q$$

35. Match the chemical conversions in List-I with the appropritate reagents in List-II and select the correct answer using the code given below this lists - [IIT 2013]

List-I

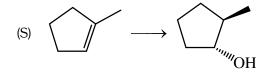
List-II

$$(P) \longrightarrow Cl \longrightarrow$$

- (1) (i) Hg(OAc)₂ (ii) NaBH₄
- $(Q) \longrightarrow ONa \longrightarrow OE$
- (2) NaOEt

$$(R) \qquad \longrightarrow \qquad \bigcirc OH$$

(3) Et-Br



(4) (i) BH_3 (ii) H_2O_2 /NaOH

Codes:

- (A) 2 3 1 4
- (B) 3 2 1 4
- (C) 2 3 4 1
- (D) 3 2 4 1

26. (a) (i) Products are
$$C_2H_3 + C_2H_5 + C_3H_5 + C$$

- (ii) No reaction takes place, as aryl halide is inert to nucleophilic substitution.
- (b) (i) $-NO_2$ group facilitates the nucleophilic attack because it stabilised the anion formed when attack of $\overline{O}H$ takes place.
 - (ii) Does not show reaction.

28. (a)
$$H_2C-C=CH-CH_3$$
 (b) CCH_3 (c) CCH_3

29. A,C **30**. D **31**. C **32**. D **33**. A **34**. B **35**. A