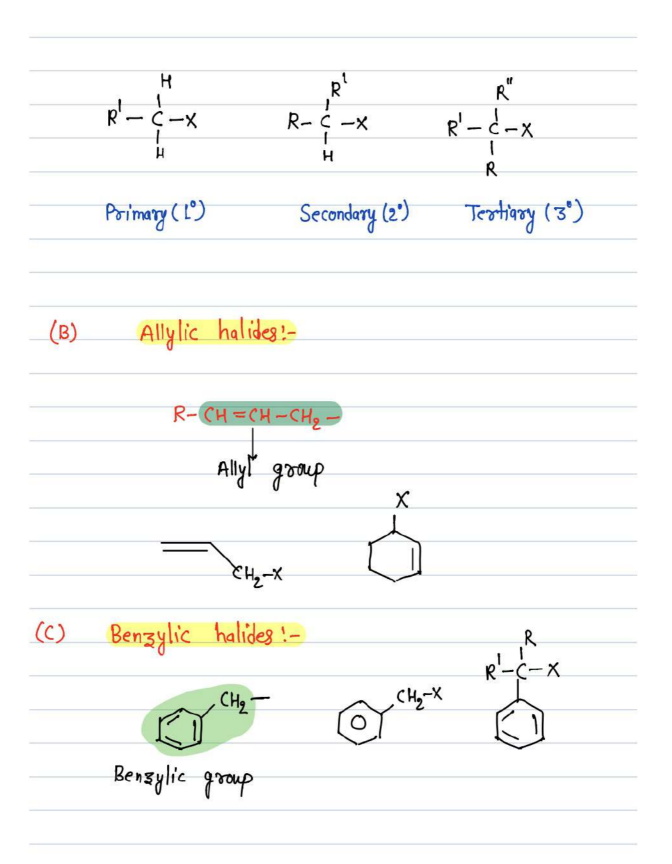
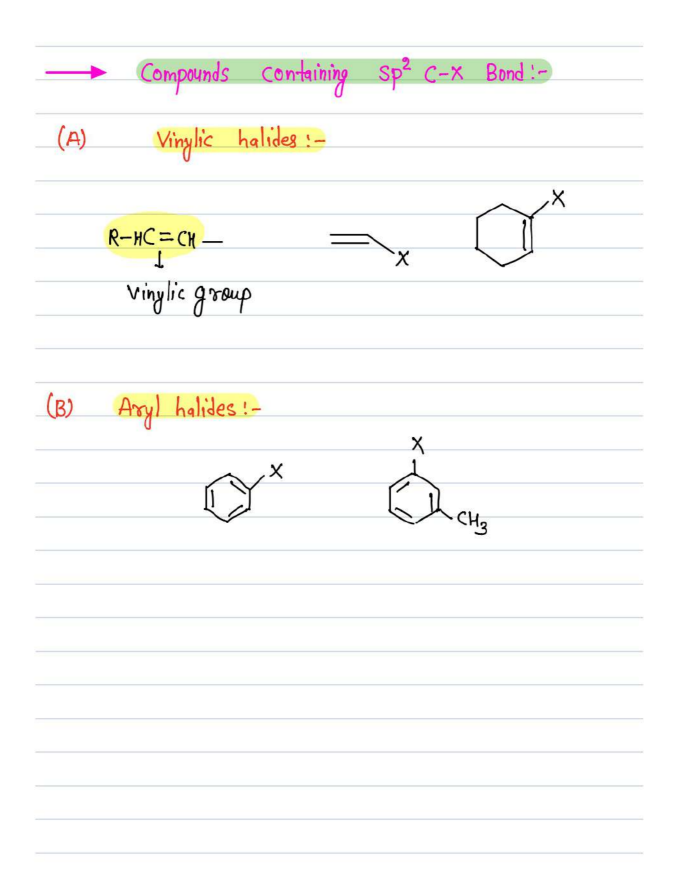
aliphatic or in the format aryl halide (haloo		
→ On the Basis	of Number of	Halogen Atoms:-
Monohaloalkane	Dihaloalkane	Tribalealkane
CH3-CH2-X	CH ₂ - X I CH ₂ -X	CH2-X CH-X CH2-X
Monohalogrenes	Dihalogrenes	Trihalogrenes
© X	o x	× ×
Compound	Confaining sp	(X=F,Cl,Br,I)
(A) Alkyl hali	ides!	





NOMENCLATURE

1-Chloro-2-methylbenzene

NATURE OF C-X BOND :-

$$\frac{1}{6}$$
 $\frac{1}{6}$ $\frac{1}{6}$

The carbon-halogen bond of alkyl halide is polar since electronegativity difference is non zero between Carbon and halogen atom.

(Vuestion!	Explain dipole moment	of CHz-F, CHz-Cl, CHz-Br and CHz-I.
Solution!	W	Dipole moment (Debye)
	CH3-CI	1.860
	CH3-F	1.847
	CH3-Br	1.830
	CH3-I	1.636
•	\$500 cm	and d, dipole moment of CHz-Cl is max
÷	Y	and d, dipole moment of CHz-Cl is max
	Result of both 9	and d, dipole moment of CHz-Cl is max. Compounds.
	Result of both 2 amony the above oreign 2015, All India 2010 Grive reason -	and d, dipole moment of CHz-Cl is maxi
	Result of both 2 amony the above oreign 2015, All India 2010 Grive reason -	and d, dipole moment of CHz-Cl is max Compounds. O, Delhi 2011, 2013 of chlorobenzene is lower than that

As sp2-hybridised carbon has more s-character so, it is more electronegative, thus the density of electrons of C-C
bond near the CI-atom is less in chlorobenzene that in cyclohexyl chloride. Hence, the C-CI bond of cycloher chloride is more polar, 1:e. it has higher dipole moment.
METHOD OF PREPRATION:—
1. FROM ALCOHOLS:—
R-OH + HCI ZnClg → R-C1 + H2O

$$\begin{array}{c} CI \\ P-CI+0 \\ \hline \\ CI \\ \end{array} \longrightarrow \begin{array}{c} R-CI+POCI_3+HCI \\ \hline \end{array}$$

$$\begin{array}{ccc} R-OH & \xrightarrow{\text{red P}/X_2} & R-X \\ \hline X_2 = Br_2, I_2 & \end{array}$$

2.	FROM	HYDRO	CARBON	S:-
----	------	-------	--------	-----

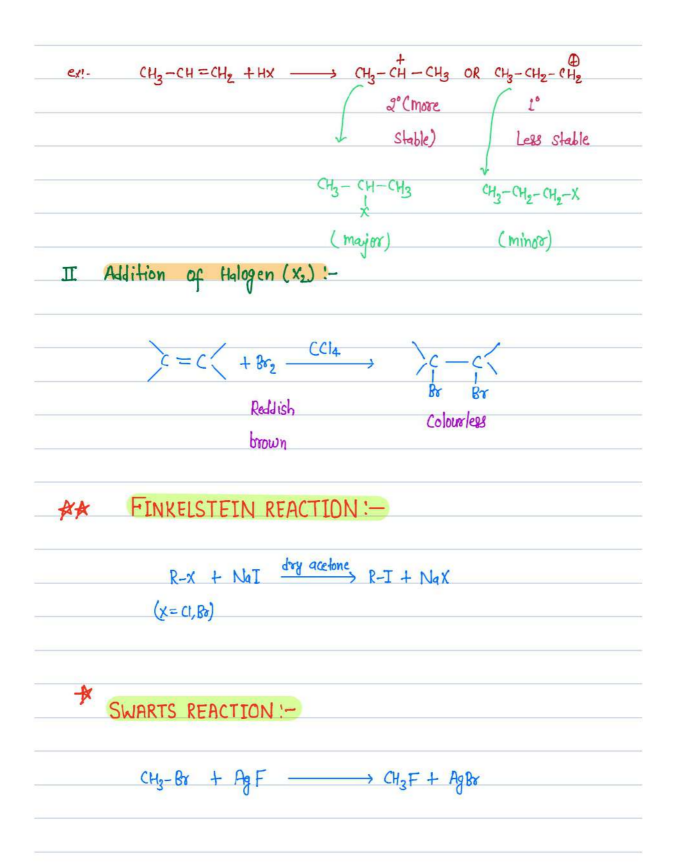
$$c = c < + Hx \longrightarrow c - c <$$

$$CH_3 - CH = CH_2 + HI \longrightarrow CH_3 - CH - CH_3 + CH_3 CH_2 CH_2 T$$

Major Minor (Markovnikov's rule)

SHORT Trick For Major product: Add halogen to the more stable.

Carbocation and H to less stable.



PHYSICAL PROPERTIES

Alkyl halides, though polar, are immiscible with water why?

Solution: Alkyl halides are polar molecules and so held together by dipole-dipole interactions. Similarly, strong H-bonds exist between the water molecules. The new force of attraction between the alkyl halides and water molecules is weaker than the forces of attraction already existing between alkyl halide-alkyl halide and water-water molecules. Hence, alkyl halides (though polar) are immiscible (or insoluble in) with water.

Boiling point :- R-I>R-B>R-CI>R-F

Reason: - Increase in size and mass of halogen atom, the magnitude of van der Waal forces increases.

For Isomeric haloalkanes:

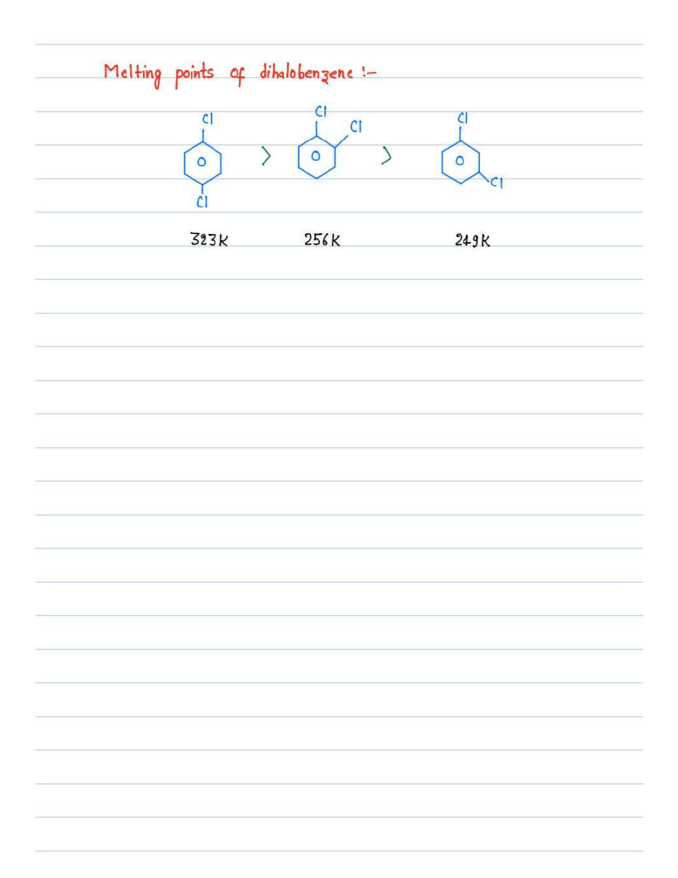
Boiling point \(\frac{1}{Branching} \)

CH3-CH2-CH2-CH2-Br \(\rangle \text{CH}_3-CH_2-CH-Br \)

CH3 \(\text{CH}_3 \)

CH3 \(\text{CH}_3 \)

CH3 \(\text{CH}_3 \)



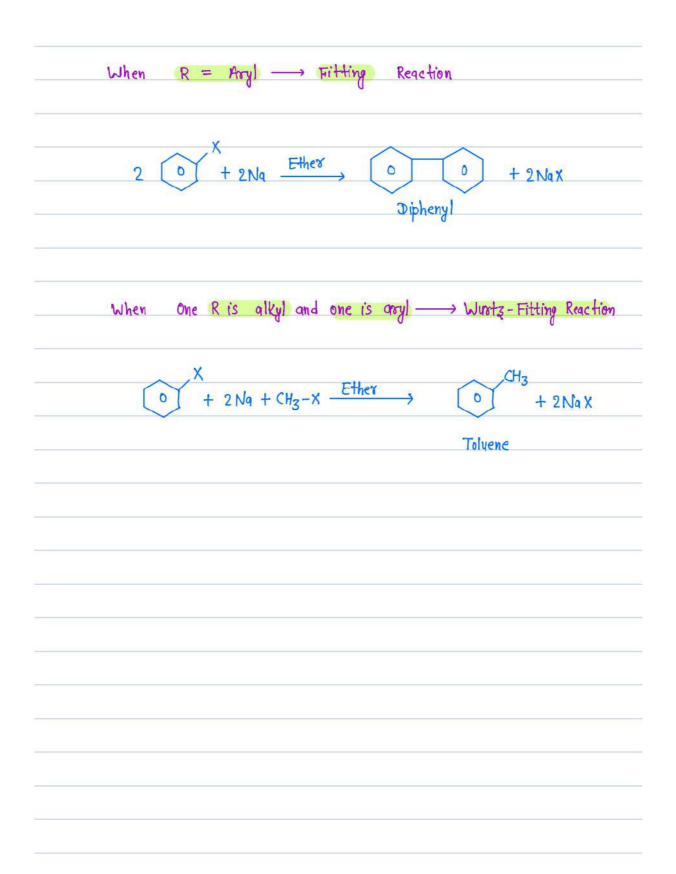
CHEMICAL PROPERTIES Haloalkanes Shows three type of reactions -Nucleophilic substitution Elimination reactions > Reaction with metals Nucleophilic Substitution: -> SNL See detail in ppt Haloalkanes react with KCN to foom alkyl cyanides as main product while Agen forms isocyanides as the chief (main) product. Explain KCN is predominantly ionic and provides cyanide ions Solytion! in solution. Although both carbon and nitrogen atoms are in a position to donate electron poios, the attack takes place mainly through carbon atom and not through nitrogen atom since C-C bond is more

	in nature and	nitrogen is	free to donat	e elech
	pair forming	isocyanide a	8 the main p	moduct.
IMPORTAN	IT DEFINITION	IC ODAUT	APTICAL TOAM	1 C O C C 1
	1) ACLIVITION	12 MROU ! (DI LICHE 1301	IEKT 21.
CHIRAL (AS	YMETRIC) CARBO	H ATOM:-	The carbon a	tom Whi
CHIRAL (AS		N ATOM:—	The carbon a	tom Whi
is attac	YMETRIC) CARBOI	N ATOM:— ifferent atom	The carbon a	tom Who
is attac asymmetric	YMETRIC) CARBO hed to four di or chiral car	N ATOM:— ifferent atom atom ers are t	The carbon a	tom Wh
is attac asymmetric ENANTIO	ymetric) carbonal distribution of chiral car MERS:— Enantione age of each othe Substances which	N ATOM:— ifferent atom whom atom ers are the	The carbon and some sistem of the carbon and some supersing the plane polarion	tom who called pasable
is attac asymmetric ENANTIO	ymetric) carbon hed to four di or chiral car mers:— Enantione hage of each othe	N ATOM:— ifferent atom whom atom ers are the	The carbon and some sistem of the carbon and some supersing the plane polarion	tom who called pasable
is attac asymmetric ENANTIO misror in	ymetric) carbonal distribution of chiral car MERS:— Enantione age of each othe Substances which	N ATOM:— ifferent atom whom atom exs are the inch rotate the ialled dextoor	The carbon and some six he non-supersimple the plane polarion of the plane polarion of the plane polarion of the plane (+) and	tom who called pasable those u

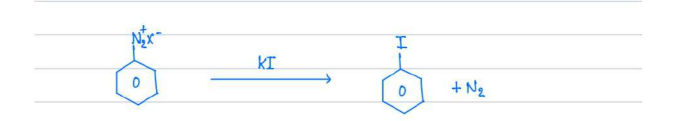
configura to	this process	IS (Alled	· s e tention	of
INVERSION	OF CONFI	GURATION	:- If the	beec
formed in	a reaction h	as a ster	eochemistry o	ppe8
to that	of reactant,	it leads to	inversion	ot
configuration	η			
RACEMIC M	IXTURE : - It	is q m	ixture containi	ng 🖪
	n equal propon			V
optically inc	active and ha	ve Zero	optical rotation	9m ·
3	If in a	reaction,	50% inversion	n o
50% retent	tion occurs,	the process	is called	
racemisation	1			

ELIMINATION REACTION: (B- Elimination OR Dehydrohalogenation) B = alc-KOH, NONH2 etc. Major product will be decided by saytzeff rule. AK Alkene which has the greater number of alkyl groups attached to the doubly bonded corbon atoms, will be major product. AIC-KOH CH3-CH2-CH= CH-CH3 Pent-2-ene (81%) Ex!-All India 2015 AIC-KOH CH3-CH2-CH2-CH=CH2 Pent-1-ene (19%) CBSF 2018 AIC-KOH Ex! -1- Bromo - 1- methylcyclohexane Major Minor

REACTION WITH METALS: - Compounds containing carbon metal bonds are known as organo-metallic compounds. Ex: - RMgX — Grignard Reagent R-x + mg dry ether RMgx Grignard reagent Cirignard reagents are highly reactive. Cirignood reagent react with water, alcohols, amines and formed hydrocarbon. RMgX + H2O - RH + Mg(OH)X WURTZ REACTION !- When alkyl halide react with Sodium in dry ether, reaction known as wortz reaction. 2RX + 2Na dry ether R-R + 2NaX



HALOARENES / ARYL HALIDES PREPRATION !-ELECTROPHILIC SUBSTITUTION :-CH3 0 0-Halotolyene P-Halotolyene SANDMEYER'S REACTION :-Benzene diazoniam halide + N2 (x= C1, Br) Aryl Halide Sandmeyer reaction



REACTIONS OF HALOARENES

Question! Anyl halides are extremely less reactive towards nucleophilic substitution. Why?

Ary) holides and vinyl holides are less reactive towards

SNI and SN2 reaction. Why?

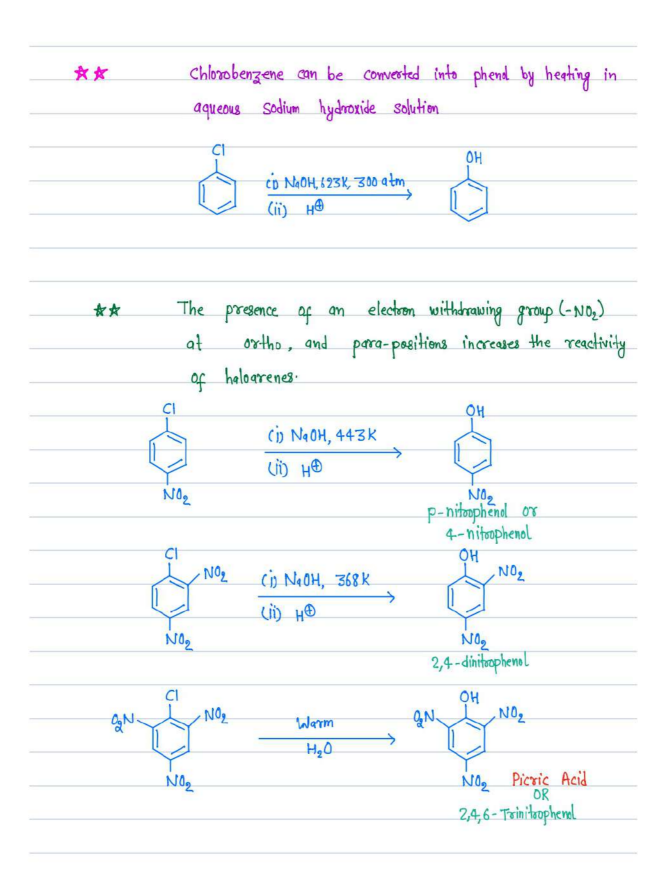
OR

Out of chlosobenzene and benzyl chloride, which one gets easily hydrolysed by agreeus NaOH and why?

Solution: 1. Resonance effect — In halo avenes or vinythalides, the electron pairs on halogen atom are in conjugation with T-electrons of the ring or alkyl group.

2. Difference in hybridisation of carbon atom in C-X bond:

The Sp2 hybridised carbon with a greater s-character is more electronegative and can hold the electron pair of C-X bond more tightly than Sp3-hybridised carbon in haloal kane with less s-character. Since it is difficult to break a shorter bond than a longer bond, therefore, halogrenes or vinythalides are less reactive than haloalkanes towards nucleophilic substitution reaction. Aryl halide 3. Instability of phenyl cation or vinyl cation - No resonance Stablization CH₂ = CH Vinyl Cation Phenyl Cation

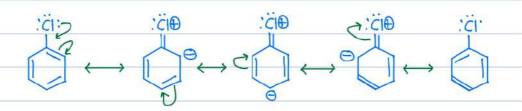


ELECTROPHILIC SUBSTITUTION REACTION:-

Ques:- Although chlorine is an electron withdrawing group, yet it is ortho-, para-directing in electrophilic arromatic substitution reactions. Why?

Solution: - Chlorine withdraws electrons through inductive effect and releases electrons through resonance.

In chlorobenzene, one of the lone pair on chlorine atom Conjugates with the ring.



Since, electron density is maximum at 0- and p-positions due to tR-effect, therefore an electrophile can easily attack at such positions.

