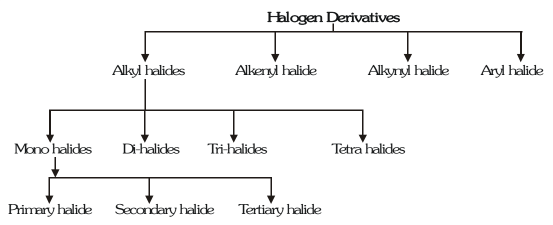
# ALKYL HALIDE AND ARYL HALIDE

Compounds derived from hydrocarbons by replacement of one or more H-atoms by corresponding no. of halogen atoms are known as halogen derivatives.

#### □ Classification:

On the basis of nature of hydrocarbon from which they are obtained, hydrocarbon derivatives can be classified as:



- (a) Alkyl halides: Halogen derivative of alkanes.
- (b) Alkenyl halides: Halogen derivative of alkenes.
- (c) Alkynyl halides: Halogen derivative of alkynes
- (d) Aryl halides: Halogen derivative of arenes (aromatic)
- ♦ Alkyl halides: They are further classified on the basis of halogen atoms introduced in the molecule. e.g.
  - (i) Mono halides: These involves replacement of one H-atom by halogen atom.

General formula  $C_n H_{2n+1} X$ 

CH<sub>3</sub>CH<sub>2</sub>Br Ethyl bromide (Bromo ethane)

(ii) Dihalides: Replacement of two H-atom by halogen atoms.

General formula  $C_nH_{2n}X_2$ 

Example: CH<sub>2</sub>X<sub>2</sub> Methylene dihalide

CH<sub>2</sub>X Ethylene dihalide or Vicinal dihalide

CH<sub>2</sub>X

CH<sub>3</sub> Ethylidene dihalide or geminal dihalide

CHX<sub>0</sub>

(iii) Trihalides: Replacement of three H-atoms by halogen atoms. General formula -  $C_nH_{2n-1}X_3$ . Example:  $CHX_3$  Trihalo methane or haloform

When all the H-atoms from an alkane are replaced by halogen atoms, then the compounds are called as perhalo compound - General formula  $C_nH_{2n-2}X_4$  (tetra halide).

 $CH_4 \longrightarrow CX_4$  (Per halo methane)

 $C_2H_6 \longrightarrow C_2X_6$  (Per halo ethane)

MONO HALIDES: These are classified on the basis of nature of C-atom carrying the halogen atom. **Primary halide or 1^0 alkyl halides:** Halogen atom attached with a primary or  $1^0$  C-atom. Halo methane or methylhalide Example: CH<sub>3</sub>—X  $CH_3$ — $CH_2$ —XHalo ethane or ethyl halide  $CH_3$ — $CH_2$ — $CH_2$ —X1-Halo propane or n-propyl halide **Secondary or 2<sup>0</sup> alkyl halides**: Halogen atom linked with 2<sup>0</sup> C-atom. Example: CH3-CH-CH3 2-halo propane or Iso propyl halide CH3-CH-CH2-CH3 2-halo butane or Sec. butyl halide (C) **Tertiary halide or 3<sup>o</sup> alkyl halide:** halogen atom linked with 3<sup>o</sup> C-atom. Example : R - C - XIsomerism: Alkyl halides shows position and chain isomerism -OH3CH2CH2CI

CH3CHCH3

Position isomers Example:  $C_3H_7Cl$ Example :  $C_4H_9Cl(a)$   $CH_3CH_2CH_2CH_2Cl$  1 - chloro butane

(b)  $CH_3$   $CH_2CH_2CH_2Cl$  1 - chloro-2-methyl propane

(c)  $CH_3$   $CH_2$   $CH_3$  2 - chloro butane

(d)  $CH_3$   $CH_3$  2 - chloro-2-methyl propane  $CH_3$   $CH_3$   $CH_3$  2 - chloro-2-methyl propane a, c  $\longrightarrow$  Chain isomers
a, d  $\longrightarrow$  Chain and position isomers
b, d  $\longrightarrow$  Position isomers
c, d  $\longrightarrow$  Chain isomers
vity order. The and find the content of the con Reactivity order: The order of reactivity of alkyl halides is - RI > RBr > RCl > RFBond energy values: C-I (57.4), C-Br (65.9), C-Cl (78.5) and C-F (105.4) K.cal/mole C-I bond is most reactive because lower energy is required to break the bond. On the basis of nature of alkyl group the reactivity order of alkyl halide is tert > Sec. > Primary

Since alkyl groups, are electron repelling or electron releasing, larger no. of alkyl groups on C-atom of C-X greater is the electron density on C-atom hence ease in release of X atom as  $X^-$  ion (+ I effect of alkyl group)

Alkyl halides are generally more reactive than the corresponding alkanes due to the presence of polar covalent  $(\overset{|_{\delta_+}}{C}\overset{\delta_-}{X})$  bond. So alkyl halides ( R–X) undergo nucleophilic substitution reaction.

The centre for attacking  $\stackrel{1}{Nu}$  is  $\stackrel{1}{-C}$  atom

## Genreal Method of Preparation of Monohalides:

1. By direct halogenation of alkanes:

2. By the addition of H-X on alkenes:

R—CH = CHR + HX 
$$\longrightarrow$$
 RCH<sub>2</sub>—CHXR  
CH<sub>2</sub>=CH<sub>2</sub> + HX  $\longrightarrow$  CH<sub>3</sub>—CH<sub>2</sub>X  
CH<sub>3</sub>-CH = CH<sub>2</sub> + HX  $\longrightarrow$  CH<sub>3</sub> — CH—CH<sub>3</sub>  
X

Isopropyl halide

## 3. By Alcohols:

(a) By the action of hydrogen halides :

$$\textbf{Example} \; : \; \text{R---CH}_2 - \text{OH} \xrightarrow{\quad H-X \\ \quad ZnCl_2} \rightarrow \text{RCH}_2 - X$$

Mechanism:

$$R-CH_{2}-\overset{\overset{}{\circ}H}{\overset{}{\circ}H}\overset{\overset{}{\longrightarrow}} R-CH_{2}\overset{\overset{}{\bigcirc}G}{\overset{}{\bigcirc}}-H\overset{-H_{2}O}{\overset{}{\longrightarrow}}R-\overset{\overset{}{\bigcirc}G}{\overset{}{\longrightarrow}}R-CH_{2}-X$$

$$(unstable) \qquad \qquad (Product)$$

In this reaction intermidiate carbocation is formed so rearrangement ( $H^-$  shifting or  $CH_3^-$  shifting) can take place.

 ${\rm ZnCl_2}$  act as dehydrating agent and absorbs  ${\rm H_2O}$  from the reaction so good yield of halide is obtained. Also it generates  ${\rm H^+}$  from HCl.

$$\mathsf{HCl} + \mathsf{ZnCl}_2^{\mathsf{1}} \quad \longrightarrow \quad \mathsf{ZnCl}_3^{\mathsf{1}} \, + \mathsf{H}^{\oplus}$$

Reactivity order for alcohol:

Reactivity 

stability of intermediate carbocation , so reactivity order: Tert. alc. > Sec. alc. > Pri. alc.

Reactivity order of H—X is: HI > HBr > HCl

HI is maximum reactive so it reacts readily with  $1^{\circ}$ ,  $2^{\circ}$  and  $3^{\circ}$  alcohols.

$$R$$
— $OH + HI$   $\longrightarrow$   $R$ — $I + H2O$ 

HCl and also  $1^{\circ}$  alcohol are less reactive so  $\text{ZnCl}_2$  or some amount of  $\text{H}_2\text{SO}_4$  is needed to increase the reactivity.

**Example**: 
$$CH_3$$
— $CH_2$ — $OH + HCl$   $\xrightarrow{ZnCl_2}$   $CH_3$ — $CH_2$ — $Cl$ 

At normal condition:

Note: HCl + ZnCl<sub>2</sub> is called as lucas reagent, alchol gives turbidity with lucas reagent.

Reactivity towards lucas reagent (difference in 1°, 2° and 3° alcohol).

	1° alcohol	2° alcohol	3° alcohol
Time to	in 30 min.	in 5 min.	in 1 min.
give turbidity			

(b) By the action of phosphorus halides ( $S_{N^1}$  mechanism) :

 $PBr_3$  and  $PI_3$  are less stable, thus for bromides and Iodides, (P +  $Br_2$ ) Or (P +  $I_2$ ) mixture is used.

(c) By reaction with thionyl chloride - (Darzen's procedure) (  $S_{N^i} and \, S_{N^2} \,$  mechanism) :

$$\label{eq:reconstruction} \text{R--OH} \quad + \quad \text{SOCl}_2 \qquad \xrightarrow{\text{Pyridine}} \quad \text{R--Cl} \ + \ \text{SO}_2 \ + \ \text{HCl}$$

One mole One mole

Because of less stability of  $SOBr_2$  and  $SOI_2$ , R—Br and RI does not obtained by this method.

4. Borodine - Hunsdicker's reaction:

R—COOAg + 
$$X_2 \xrightarrow{CCl_4} R$$
—X +  $CO_2$  + AgX  
Silver salt of  $(Cl_2 \text{ or } Br_2)$   
a fatty acid

5. By halide exchange:

R–Cl or R—Br + KI 
$$\xrightarrow{Acetone}$$
 R–I + KCl or KBr (Conant finkelstein reaction)   
  $2CH_3Cl + HgF_2 \longrightarrow 2CH_3$ –F +  $HgCl_2$  (Swart reaction)

R-I and R-F can be prepared by this method only.

6. By reaction of alkanes with sulphuryl chloride (SO<sub>2</sub>Cl<sub>2</sub>):

$$\text{R---H} \quad + \quad \text{SO}_2 \, \text{Cl}_2 \quad \xrightarrow{\quad \text{light} \quad \quad } \text{R----Cl} \quad + \quad \text{HCl} \quad + \quad \text{SO}_2$$

☐ Physical Properties :

- (a) The lower members  $CH_3F$ ,  $CH_3Cl$ ,  $CH_3Br$ ,  $C_2H_5Cl$  and  $C_2H_5F$  are gases at room temp.  $CH_3I$  and members upto  $C_{18}$  are colourless sweet smelling liquids.
- (b) Higher B.P. than parent alkanes.

Decreasing order of B.P. is : 
$$R-I > R-Br > R-CI > R-F$$
  
among isomeric R—X decreasing order of B.P. is : Primary > Secondary > tertiary

R—Br and R—I 
$$\longrightarrow$$
 heavier than water

Decreasing order of density is : R-I > R-Br > R-CI > R-F

- (d) R—X are polar co-valent compounds but insoluble in water because they can not form H-bonds. They dissolve in organic solvents.
- (e) R—X burns with a green flame due to interaction of X with Cu wire.(Beilstein test)
- (f) The stability order is : R F > R Cl > R Br > R I

R—I is least stable and darken in light due to photodecomposition.

$$2R-I \xrightarrow{hv} R-R + I_2$$

☐ Chemical Properties :

A. Nucleophilic substitution reaction ( $S_N$ ): Due to electronegativity difference the -  $\stackrel{|}{C}$  X bond is highly

Thus the C-atom of the  $\overset{\delta_+}{C}-\overset{\delta_-}{X}$  bond becomes centre to attack by a nucleophile  $\overset{1}{(Nu)}$ .

 $X^1$  ion from R—X molecule is substituted by a  $\overset{1}{N}u$ . i.e.  $S_N$  reaction are the most common reactions in R—X. R—X +  $\overset{1}{N}u$  —  $\longrightarrow$  R—Nu +  $X^1$ 

These may be takes place by two ways -

(a) 
$$S_{N^1}$$
 mechanism

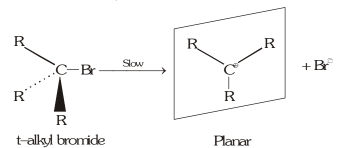
(b) 
$$S_{N^2}$$
 mechanism

Reactivity order is :

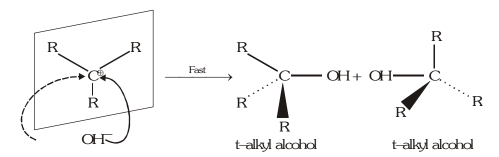
# $\square$ Mechenism of $S_{N^1}$ and $S_{N^2}$ :

 $\bullet$   $S_{N^1}$  Mechanism:  $SN^1$  stands for uni molecular nucleophilic substitution. The mechanism involves two steps. Consider the hydrolysis of tert, butyl bromide with aqueous NaOH.

**Step 1:** The alkyl halide ionises to give a planar corbonium ion. The corbonium ion is planar because the central positively charged carbon is  $sp^2$  hybridized.



Step-2: The nucleophile can attack the planar carbonium ion from either side to give the product.



- (i) Ionisation is the rate determining step because it is the slow step. In other words, the rate at which alcohol is formed should depend upon the concentration of tertiary alkyl halide alone.
- $\text{Rate} = K[R_3C Br]$  It is obvious that the reaction follows first order kinetics, therefore reaction is called  $S_{N^1}$ .
- (ii) The reactivity order for  $S_{N^1}$  reaction  $\infty$  stability of carbocations formed by halides.
- $\therefore$  reactivity order of halides ( $S_{N^1}$ ) varies as follows : Benzyl halide > Allylhalide > 3°halide > 2° halide > 1° halide > methyl halide.
- (iii) Remember that in case alkyl halide is optically active, SN<sup>1</sup> reactions lead to racemisation.

 $\bullet$   $S_{N^2}$  mechanism:  $S_{N^2}$  stands for bimolecular nucleophilic substitution. In this type of nucleophilic substitution reaction, there occurs bond making and bond breaking simultaneously.

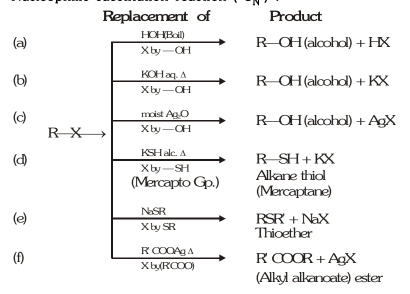
Transition state

- (i) Reactivities of alkyl halides in  $S_{N^2}$  substitution is governed by steric factors. The bulkier the group, that less reactive it will be.
- (ii) Reactivity order of alkyl halide varies as follows:  $CH_3X > 1$ °halide > 2°halide > 3° halide
- (iii) The order of reactivity among 1° alkyl halides is :  $CH_3X > C_2H_5X > C_3H_7X$  etc. Remember that in case alkyl halide is optically active,  $SN^2$  reactions lead to Walden inversion.
- (iv) Thus in short 3° alkyl halides react by  $S_{N^1}$ , 1° by  $S_{N^2}$  and 2° by either or both of them  $SN^1$  and  $SN^2$  depend upon the nature of the alkyl halide and the reagent.
- (v) For a given alkyl group the order of reactivity is (for  $S_{N^1}$  and  $S_{N^2}$  both) : RI > RBr > RCl > RF
- (vi) In addition to substitution reaction alkyl halide also undergo elimination reactions to form alkene with the removal of a molecule of hydrogen halide (dehydrohalogenation). In dehydrohalogenation, hydrogen and halogen atoms are eliminated from two adjacent carbon atoms, the reaction also known as  $\beta$ -elimination may proceed by  $E_1$  &  $E_2$  mechanism (analogous to  $S_{N^1}$  and  $S_{N^2}$  mechanism).

The order of elimination reaction is :  $3^{\circ}$  halides  $> 2^{\circ}$  halides  $> 1^{\circ}$  halides

- (vii) In general 3° halides tend to react by elimination; 1° halides by substitution and 2° halides by either or both of the reactions.
- B. Elimination Reactions (ER): Alkyl halides also undergo ER in the presence of base as Nu (Loss of H—X and formation of = bond)

1. Nucleophilic substitution reaction  $(S_N)$ :



Reaction with  $KNO_2$  and  $AgNO_2$ (h)

$$\stackrel{\delta_{+}}{R} \stackrel{\delta_{-}}{-X} + \stackrel{+}{K} \stackrel{N}{N} O_{2} \stackrel{Alc.}{\longrightarrow} \stackrel{Alc.}{\longrightarrow} \stackrel{R}{-} O \stackrel{N}{\longrightarrow} = O + KX + RNO_{2}$$
 (Alkyl nitrites) major

(i) Reaction with NaOR' (Sodium alkoxide): (williamson synthesis reaction)

$$R$$
— $X$  +  $NaOR'$  — —  $R$ — $OR'$  +  $NaX$ 

(j) Reaction with  $NH_3$ :

Example: (i) 
$$R - X + NH_3 \xrightarrow{\Delta} R - NH_2 + H - X$$

(ii) 
$$R - X + NH_3 \xrightarrow{\Delta} R - NH_2 + H - X$$
  
(excess)

(iii) 
$$R-X+NH_3 \xrightarrow{\Delta} R-NH_2 \xrightarrow{R-X} R-NH-R \xrightarrow{R-X} R-N-R$$

$$\downarrow R \\ \downarrow R-X$$

$$\downarrow R$$

$$\downarrow$$

(iv) 
$$CH_3$$
— $CH_2$ — $Cl+ NH_3$ — $\Delta$ — $CH_3$ — $CH_2$ — $NH_2 + HCl$ 

(v) 
$$CH_3 - CH_3 + NH_3 \xrightarrow{\Delta} CH_3 - CH_3 + HCI$$
  
 $CI$ 
 $CI$ 

(v) 
$$CH_3 - CH_3 + NH_3 \xrightarrow{\Delta} CH_3 - CH_3 + HCI$$
 $CH_3$ 
 $CH_3$ 

(vi)  $CH_3 - CC_3 + NH_3 \xrightarrow{\Delta} CH_3 - CC_3 + NH_4CI$ 
 $CH_3$ 
 $CH_3$ 

(k) Reaction with CH  $\equiv \bar{C} \stackrel{+}{Na}$ :

$$R-X + CH \equiv \overline{C} \stackrel{+}{Na} \xrightarrow{\Delta} R-C \equiv CH + NaX$$

2. Elimination Reaction : (Dehydrohalogenation) Alkyl halides undergo  $\beta$  - elimination on treatment with KOH (alc.)

$$R - \overset{\beta}{C}H_{2} - \overset{\alpha}{C}H_{2} - X + KOH(alc.) \xrightarrow{\Delta} R - CH = CH_{2} + HX$$

$$CH_{3} - CH_{2} - CH_{2} - CH_{2} - Br + KOH(alc.) \xrightarrow{\Delta} CH_{3} - CH_{2} - CH = CH_{2} + HBr$$

$$\overset{\beta}{C}H_{3} - CH_{2} - CH_{2} - CH_{2} - HBr$$

$$CH_{3} - CH_{3} - CH_{3} - CH_{3} - CH_{3} - CH_{3} - CH_{2} - CH = CH_{2} + HBr$$

$$2 - \text{butene (80\%)} \qquad 1 - \text{butene (20\%)}$$

♦ Competition between substitution and elemination reactions :

Reactivity order of alkyl halides : 
$$E_1$$
 - Reaction :  $1^\circ < 2^\circ < 3^\circ$  
$$E_2$$
 - Reaction :  $1^\circ < 2^\circ < 3^\circ$  
$$S_{N^1}$$
 - Reaction :  $1^\circ < 2^\circ < 3^\circ$  
$$S_{N^2}$$
 - Reaction :  $1^\circ > 2^\circ > 3^\circ$ 

Alkyl halides	S <sub>N<sup>2</sup></sub> /E <sub>2</sub>	S <sub>N1</sub> /E <sub>2</sub>	
1° Alkyl halides	Mainly give substitution unless	Can not undergo SN <sup>1</sup> /E <sup>1</sup>	
	sterically hindered alkyl halide		
	or sterically hindered base in		
	which case elimination is favoured		
2° Alkyl halides	Both substitution and elemination,	Both substitution and elemination	
	stronger base/ bulkier base or high	high temp. greater of percentage	
	temp. greater percentage of elimination.	of elimination	
3° Alkyl halides	Mainly elimination	Both substitution and elimination, high	
		temp greater percentage of elimination.	

- (i)  $S_{N^2}/E_2$  is favoured by high conc. of good neucleophile or strong base. (CH $_3$ O $^1$ , HO $^1$ ) Rate of Reaction  $\propto$  (Substrate) (Reagent)
- (ii)  $S_{N^1}/E_1$  is favoured by low conc. of poor neucleophile or weak base (CH<sub>3</sub>OH, H<sub>2</sub>O)
- (iii) If an alkyl halide, undergoes  $S_{N^2}/S_{N^1}$  Reaction then  $S_{N^2}$  will be favoured by high conc. of good neucleophile (negetively charged) in presence of polor aprotic solvent . where as  $S_{N^1}$  reaction is favoured by low conc. of poor neucleophile (neutral) is presence of polar protic solvent.

Polar protic solvent : H<sub>2</sub>O, CH<sub>3</sub>OH, HCOOH.

Polar aprotic solvent : DMSO,  $CH_3CN$ ,  $C_2H_5$ -O- $C_2H_5$ 

$$\begin{array}{c} \textbf{Example}: \ \ CH_3CH_2CH_2 - Br + CH_3O^1 - \xrightarrow{CH_3OH} \ \ CH_3CH_2CH_2 - OCH_3 + CH_3CH = CH_2 \\ \qquad \qquad \qquad \qquad \qquad \text{major} \qquad \qquad \text{minor} \end{array}$$

$$S_{N^2} \qquad E_2$$
 
$$CH_3 \qquad CH_3 \qquad CH_3 \qquad CH_3$$
 
$$Example: CH_3-CH-CH_2-Br+CH_3O^2-\frac{CH_3OH}{2} \rightarrow CH_3-CH_2+CH_2+CH_3-C=CH_2$$
 
$$minor \qquad major \qquad S_{N^2} \qquad E_2$$

$$\begin{array}{c} CH_{_3}\\ |\\ CH_{_2}CH_{_2}CH_{_2}CH_{_2}Br \ +\ CH_{_3}-C-ONa \longrightarrow CH_{_3}-CH_{_2}CH_{_2}CH_{_2}Br \\ |\\ CH_{_3} \end{array}$$
 (E\_2) mech.

Example: 
$$CH_3-CH-CH_3$$

Br

$$CH_3CH_2OH$$

$$CH_3-CH-CH_2$$

$$(major)$$

$$CH_3CH_2OH$$

$$CH_3-CH-CH_3$$

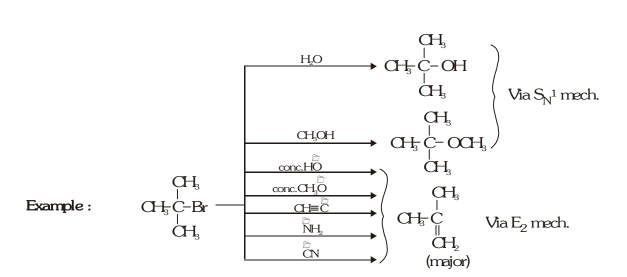
$$CH_3COH$$

$$CH_3-CH-CH_3$$

$$CH_3COH$$

$$CH_3-CH-CH_3$$

$$CH_3-CH-C$$



**Saytzeff rule:** According to this rule "The dehydrohalogenation of an alkyl halide results preferentially in the production of more alkylated alkene i.e. more stable alkene.

3. Wurtz Reaction: When a mixture of different alkyl halides,  $(R_1 - X)$  and  $(R_2 - X)$  is used a mixture of alkane is formed -

$$R_1$$
— $X + 2Na + X$ — $R_2$   $\xrightarrow{Ether \Delta}$   $R_1$ — $R_2 + R_1$ — $R_1 + R_2$ — $R_2 + NaX$ 

If 'Zn' is used in place of 'Na' the reaction is called Frankland's Reaction.

4. Halogenation of alkyl halides:

$$\text{CH}_{3}\text{--Cl} + \text{Cl}_{2} \xrightarrow{\quad \Delta \text{ or. U.V.} \\ \quad \text{light(-HCl)}} \text{CH}_{2}\text{Cl}_{2} \xrightarrow{\quad \text{Cl}_{2} \\ \quad \text{-HCl}} \text{CHCl}_{3} \xrightarrow{\quad \text{Cl}_{2} \\ \quad \text{-HCl}} \text{CCl}_{4}$$

Similarly 
$$C_2H_5Cl \xrightarrow{Cl_2} C_2H_4Cl_2 \xrightarrow{Cl_2} C_2H_3Cl_3 \xrightarrow{Cl_2} C_2Cl_6$$

**5. Reduction**: R - X are reduced to corresponding alkane.

$$R-X + 2H \xrightarrow{Reducing agent} R-H + HX$$

(i) 
$$CH_3CH_2Br + H_2 \xrightarrow{N_i} CH_3CH_3 + HBr$$

(ii) 
$$CH_3CH_2CH_2Br + 2H \xrightarrow{Zn/HCl} CH_3 - CH_2 - CH_3 + HBr$$

(iii) 
$$C_2H_5I + HI \xrightarrow{\text{Red.P}} C_2H_6 + I_2$$

6. Formation of Organometalic compounds :

(i) 
$$R-X + Mg \xrightarrow{dry ether} RMgX$$
 (Grignard reagent)

(ii) 
$$2C_2H_5Br + 2Zn \xrightarrow{dry \ ether} (C_2H_5)_2 Zn + ZnBr_2$$

(iii) 
$$2C_2H_5Br + Hg(Na) \longrightarrow (C_2H_5)_2Hg + 2NaBr$$
  
Sodium Amalgam

(iv) 
$$4C_2H_5Cl + 4Na / Pb \longrightarrow (C_2H_5)_4 Pb + 4NaCl + 3Pb$$
  
Sodium lead Alloy Tetra ethyl lead (used as antiknocking agent)

7. Friedel - Crafts reaction:

8. Action of heat:

(i) 
$$CH_3$$
— $CH_2$ — $CI$   $\xrightarrow{\text{At or below } 300^{\circ}C}$   $CH_3$ — $CH_$ 

(ii) 
$$CH_3-CH-CH_2 \xrightarrow{Above} CH_3CH=CH_2+H$$
 (Elimination) Propylene

- ♦ Uses :
  - (a) As alkylation agent (Wurtz reaction)
  - (b) As synthetic reagent
  - (c) Lower members used as anaesthetic agent, refrigerent or solvent.

General formula  $C_nH_{2n}X_2$ . Two H - atom of alkanes, replaced by two halogen atoms to form dihalides.

Dihalides are classified as:

(a) Gem dihalide: The term Gem is derived from geminal means - same position.

Two similar halogen atoms are attached to same C - atom

**Example:**  $CH_3CHX_2$  ethylidene dihalide (1, 1 - dihalo ethane)

 $CH_3$  Isopropylidene dihalide (2, 2 - dihalo propane)

**(b) Vic dihalides :** Vic term from - Vicinal means adjacent C - atoms Two halogen atoms are attached on adjacent carbon atom.

Example :  $CH_2-CH_2$   $CH_2-CH_2$   $CH_2-CH_2$   $CH_2$   $CH_$ 

ethylene dihalide Propylene dihalide (1,2-dihaloethane) (1,2-dihalopropane)

(c)  $\alpha$ ,  $\omega$  dihalides: Halogen atoms are attached with terminal C - atom. They are separated by 3 or more C - atom. They are also known as polymethylene halides.

Ex. OH\_-CH\_-CH\_2-CH\_2 (1,4-dichloro butane)
X Tetramethylene dichloride

#### ☐ General Method of Preparation :

- (a) Gem dihalides:
- (i) By the reaction of  $PCl_5$  on carbonyl compound.

$$\begin{array}{c} O \\ \parallel \\ CH_3-C-H \\ +PCI_5 \\ \longrightarrow CH_3-C-H+PCCI_3 \\ \end{array}$$

Acetaldehyde

Ethylidene chloride

$$\begin{array}{cccc}
O & Br \\
& & | \\
CH_3-C-CH_3 & +PBr_5 \longrightarrow CH_3-C-CH_3 + POBr_3 \\
& & | \\
Br
\end{array}$$

Acetone

2, 2 - dibromo propane

(ii) By addition of halogen acids on alkynes:

CH
$$\equiv$$
CH + HBr $\longrightarrow$  CH<sub>2</sub> $=$ CHBr $\longrightarrow$  CH<sub>3</sub> CHBr<sub>2</sub>  
Vinyl bromide 1, 1 - dibromo ethane

## (b) Vic-dihalides:

(i) By the addition of halogens to alkenes:

$$\begin{array}{c} CH_2 \\ \parallel \\ CH_2 \end{array} + Br_2 \longrightarrow \begin{array}{c} CH_2Br \\ CH_2Br \\ 1, \ 2 \ \text{- dibromo ethane} \end{array}$$

$$CH_3-CH=CH_2+Br_2\longrightarrow CH_3-CH-CH_2$$
 $Br$   $Br$ 
 $1, 2$  - dibromo propane

(ii) By the action of  $PCl_5$  on glycols :

$$\begin{array}{ccc} \text{CH}_2\text{OH} & +2\text{PCI}_5 & \longrightarrow & \text{CH}_2\text{CI} \\ \text{CH}_2\text{OH} & +2\text{PCCI}_3 + 2\text{HCI} \\ \text{CH}_2\text{CI} & +2\text{CI}_3 & +2\text{CI}_3 \end{array}$$

(c)  $\alpha - \omega$  dihalides:

(i) 
$$CH_2$$
= $CH$ - $CH_2$ Br +  $HBr$   $\longrightarrow$   $Br$ - $CH_2$ - $CH_2$ - $CH_2$ Br   
1, 3 - dibromo propane

(iii) 
$$+ Cl_2 \xrightarrow{hv}$$
  $+ HCl$ 

Note: Here has less tendency to open the ring.

# ☐ Physical Properties :

- (i) Lower members are colourless, oily liquids with sweet smell. Higher members are solid.
- (ii) The reactivity of gemdihalides is lesser than vicinal or mono halides. [Reason in presence of one halogen atom (Strong attracting –I effect) the other halogen atom can not be so easily replaced.
- (iii) These are heavier than water.

## ☐ Chemical Properties :

(i) Action of KOH(alc.) : (Dehydrohalogenation)

$$\begin{array}{cccc} CH_2X & CH_3 & \text{ (i) alc. KOH} \\ | & \text{ or } & | & \text{ (ii) NaNH}_2 \end{array} \quad \begin{array}{c} CH \\ || \\ CH \end{array}$$

(ii) Action of KOH(aq.): (Hydrolysis) It is a distinction test for gem and vic dihalides.

(a) 
$$CH_2-CI + 2KOH(aq.) \longrightarrow CH_2-OH + 2KOI$$
  
 $CH_2-OH + 2KOI$   
 $CH_2-OH + 2KOI$   
 $CH_2-OH + 2KOI$ 

Vic - dihalide. 1, 2 - ethane diol.

(iii) Reaction with KCN: Gem and Vic dihalide gives different products

Vic.:

$$\begin{array}{c} CH_2CI \\ -I_2CI \\ CH_2CI \end{array} + 2KCIN \xrightarrow{\qquad -2KCI \qquad } CH_2-CIN \xrightarrow{\qquad -H_2O/H^+ \qquad } CH_2-COOH \xrightarrow{\qquad \Delta \qquad -H_2O} CH_2-COOH \\ & CH_2-COOH \xrightarrow{\qquad -H_2O} CH_2-COOH \end{array}$$

Gem.:

$$CH_{3}-CH \stackrel{C}{\underbrace{CN}} \xrightarrow{2KCN} CH_{3}-CH \stackrel{CN}{\underbrace{CN}} \xrightarrow{H_{2}O/H^{\dagger}} CH_{3}-CH \stackrel{COOH}{\underbrace{COOH}} \xrightarrow{\Delta} CH_{3}-CH_{2}COOH$$
Propionic acid

- (i) CN group on acid hydrolysis gives COOH
- (ii) Two COOH group on one C atom on heating always loose CO2 to form monocarboxylic acid.
- (iii) Two COOH group on vic. C atom on heating loose H<sub>2</sub>O to form anhydride.
- (iv) Dehalogenation:

# GRIGNARD REAGENT

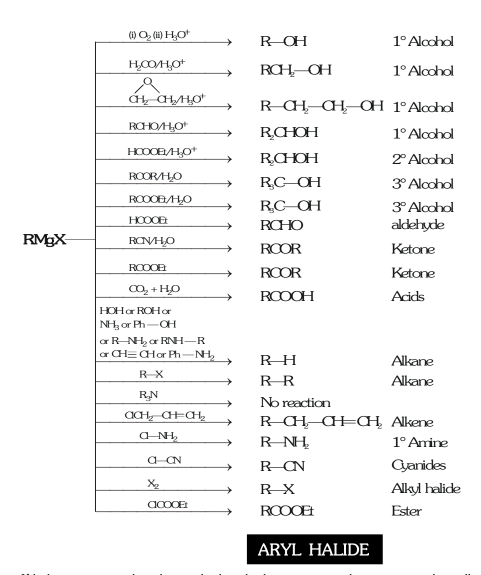
#### ☐ General Method of Preparation :

Grignard reagents are prepared in the laboratory by the action of alkyl halides on magnesium metal in the presence of dry ether.

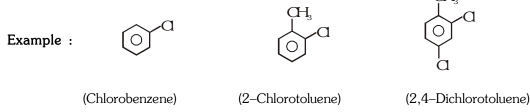
$$R-X+Mg \xrightarrow{dry} RMgX$$
 (Grignard reagent) (Alkyl magensium halide)

The ease of formation of Grignard reagent is in the order RI > RBr > RCIEther is used to dissolved the Grignard reagent by coordination.

#### ☐ Chemical Reactions :



If halogen atom is directly attached to the benzene ring, then compound is called as Haloarene.



- ☐ General Methods of Preparation :
- ♦ Halogenation of Benzene :

$$\bigcirc + C_2 \xrightarrow{ACC_3} \bigcirc + C$$

#### From Phenol :

$$3C_6H_5OH + POCl_3 \longrightarrow (C_6H_5)_3 PO_4 + 3HCl$$
(major)

# ♦ Sandmeyer's reaction :

$$\bigcirc \qquad N_2 \square \\ + CuBr \qquad \xrightarrow{HBr} \qquad \bigcirc \qquad Br \\ + N_2$$

# ♦ Reaction with KI:

#### ♦ Balz-schiemann's reaction :

$$\bigcirc \stackrel{N_2Cl}{\bigcirc} + HBF_4 \quad \longrightarrow \quad \bigcirc \stackrel{\oplus \ \square}{\stackrel{\square}{\nearrow}} HBF_4 \quad \longrightarrow \quad \stackrel{\Delta}{\bigcirc} + BF_3 + N_2$$

#### ♦ Hunsdicker reaction :

#### ♦ Raschig Process :

$$2 \bigcirc + 2H\Omega + O_2 \xrightarrow{GL_2G_2} \qquad 2 \bigcirc + 2H_2O$$

## ☐ Chemical Properties :

- (i) Chlorobenzene is essentially inert to aqueus sodium hydroxide at room temperature.
- (ii) Aryl halide are very less reactive than alkyl halides in nucleophilic substitution reactions.
- (iii) The carbon-halogen bonds of aryl halides are too strong & aryl cations are too high in energy to permit aryl halides to ionize readily  $S_{N^1}$ -type process.
- (iv) The optimal transition state geometry required for  $S_{N^2}$  process cannot be achieved.
- (v) Nucleophilic attack from the side opposite the carbon-halogen bond is blocked by the aromatic ring.

# The Elimination-Addition Mechanism of Nucleophilic Aromatic Substitution (Benzyne) :

(i) Very stong base such as sodium or potassium amide react with anyl halide, even those without electron withdrawing substituents to give products corresponding to nucleophilic substitution of halide by the base.

#### Mechanism

**Step-1**: Elimination stage; Amide ion is a very strong base and brings about the dehydrohalogenation of chlorobenzene by abstracting a proton from the carbon adjacent to the one that bears the leaving group. The product of this step is an unstable intermediate called benzyne.

**Step-2**: Beginning of addition phase; Amide ion acts as a nucleophile and adds to one of the carbons of the triple bond. The product of this step is a carbanion.

**Step-3**: Completion of addition phase; The aryl anion abstracts a proton from the ammonia used as the solvent in the reaction.

# **NOTE:** The $sp^2$ orbital in the plane of the ring in benzyne are not properly aligned for good overlap thus $\pi$ bonding is weaker than alkyne.

# Nucleophilic Aromatic Substitution by the addition Elimination Mechanism:

(i) The generally accepted mechanism for nucleophilic aromatic substitution in nitro-substituted aryl halides.

- (ii) An ortho nitro group exert a comparable rate-enhancing effect, m-chloronitrobenzyne while much more reactive than chlorobenzyne itself, is thousand of times less reactive than either o-or p-chloronitrobenzene.
- (iii) The effect of o- & p-nitro substituents is cummulative, as the rate data for substitution with methoxide ion in a series of nitro-substituted chlorobenzene derivative demostrate increasing rate of reaction as :

Chlorobenzene 1-Chloro-4-nitrobenzene Relative rate: 1.0 7 410<sup>10</sup>

1-Chloro-2, 4-dinitrobenzene 2.4 410<sup>15</sup>

2-Chloro-1, 3, 5-trinitrobenzene (too fast to measure)

(iv) In contrast to nucleophilic substitution in alkyl halides, where alkyl fluorides are exceedingly unreactive, aryl fluorides undergo nucleophilic substitution readily when the ring bears an o-or a p-nitro group.

- (v) Indeed, the order of leaving group reactivity in nucleophilic aromatic substitution is the opposite of that seen in alphatic substitution.
- (vi) Fluoride is the best reactive leaving group in nucleophilic aromatic substitution, iodide the least reactive.

Relative reactivity towards sodium methoxide in methanol (50°C) 
$$X = F \qquad Cl \qquad Br \qquad I \\ 312 \qquad 1.0 \qquad 0.8 \qquad 0.4$$
 NO,

(vii) Kinetic studies of many of the reactions described in teh section have demostrated that they follow a second-order rate law.

Rate = k[aryl halide] [nucleophile]

(viii) Second order kinetics is usually interpreated in terms of a bimolecular rate determining step.

p-Fluoronitrobenzene Sodium methoxide

p-Nitroanisole Sodium Fluoride

# Mechanism:

**Step -1 -** Addition stage. The nucleophile, in this case methoxide ion, adds to the carbon atom that bears the leaving group to give a cyclohexadienyl anion intermediate.

Methoxide ion

**Step -2 -** Elimination stage. Loss of halide form the cyclohexadienyl intermediate restores the aromaticity of the ring and gives the product of nucleophilic aromatic substitution.

Cyclohexadienyl anion intermediate

#### (a) Dow process:

p-Fluoronitrobenzene

Presence of deactivating group in ortho and para position makes the nucleophilic substitution easier.

Reactivity Order: (Towards nucleophilic substatitution)

(c) 
$$\begin{array}{c} O \\ \downarrow \\ O \\ + CuCN \end{array} \qquad \begin{array}{c} ON \\ \downarrow \\ \text{high pressure} \end{array} \qquad \begin{array}{c} ON \\ \downarrow \\ O \\ \end{array} + CuCN$$

♦ Fitting reaction :

♦ Wurtz fitting reaction :

♦ Formation of aryl megnisium halide :

$$\langle \bigcirc \rangle$$
  $\Box$  +  $Mg$   $\longrightarrow$   $\langle \bigcirc \rangle$   $Mg$   $\Box$ 

♦ Reduction:

$$\begin{array}{c} X \\ \downarrow \\ \bigcirc \end{array} \qquad \begin{array}{c} N_{i} - AI/N_{i}OH \\ \end{array} \qquad \begin{array}{c} \bigcirc \end{array}$$

♦ With Chloral:

DDT (insecticide) p,p-Dichloro diphenyl trichloroethane

♦ Electrophilic Substitution Reaction :

Cl is o- and p- directing group.

(ii) 
$$\bigcirc + HNO_3 \longrightarrow \bigoplus_{\Lambda} H_2SO_4 \longrightarrow \bigoplus_{\Lambda} NO_2 + \bigoplus_{\Lambda} NO$$

(iii) 
$$\begin{array}{c} O \\ O \\ O \\ O \end{array} + H_2SO_4 \quad \xrightarrow{\Delta} \quad \begin{array}{c} O \\ O \\ O \\ \end{array} + \quad \begin{array}{c} O \\ O \\ SO_3H \\ \end{array}$$

(iv) Friedal Craft Reaction:

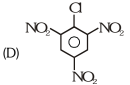
# SOLVED EXAMPLES

Ex.1 Which of the following undergoes Hydrolysis most easily:









Ans. (D)

- **Sol.** If there is more m-directing group then there will be more nuclephilic substitution reaction.
- **Ex.2** The product in the following reaction is:

$$\mathsf{Ph}-\mathsf{Cl}+\mathsf{Fe}/\mathsf{\;Br}_2{\longrightarrow}\mathsf{Product}$$

(A) o- bromo-chloro benzene

(B) p- bromo-chloro benzene

(C) (A) and (B) both

(D) 2, 4, 6-tribromo chloro benzene

Ans. (C)

- **Sol.** Since Cl group is deactivating and o/p directing group so only o– and p– products are formed.
- **Ex.3** The most reactive towards  $SN^1$  is :
  - (A) PhCH<sub>2</sub>Cl
- (B) Ph-Cl
- (C) CH<sub>3</sub>CHCl(CH<sub>3</sub>)
- (D) p-NO<sub>2</sub>--Ph--CH<sub>2</sub>--Cl

Ans. (A)

**Sol.**  $S_{N^1}$  the intermediate carbocation is formed.

$$C_6H_5$$
— $CH_2Cl$  —  $\to$   $C_6H_5CH_2$  is maximum stable due to resonance.

- Ex.4 Which of the following is used as insecticide:
  - (1) D.D.T.
- (2) Chloritone
- (3) Chloropicrin
- (4) (A) and (C) both **Ans**. (D)