3.3 Halogenoalkanes

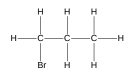
Naming Halogenoalkanes

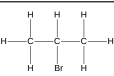
Based on original alkane, with a *prefix* indicating halogen atom: Fluoro for F; Chloro for Cl; Bromo for Br; lodo for I.

Substituents are listed alphabetically

Classifying halogenoalkanes

Halogenoalkanes can be classified as primary, secondary or tertiary depending on the number of carbon atoms attached to the C-X functional group.

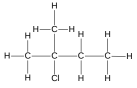




Primary halogenoalkane One carbon attached to the carbon atom adjoining the halogen Secondary halogenoalkane Two carbons attached to the carbon atom adjoining the halogen

Reactions of Halogenoalkanes

Halogenoalkanes undergo either substitution or elimination reactions



Tertiary halogenoalkane Three carbons attached to the carbon atom adjoining the halogen

Organic reactions are classified by their mechanisms

1. Nucleophilic substitution reactions

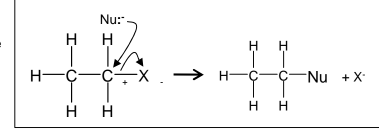
Substitution: swapping a halogen atom for another atom or groups of atoms

Nucleophile: electron pair donator e.g. :OH⁻, :NH₃, CN⁻

The Mechanism: We draw (or outline) mechanisms to show in detail how a reaction proceeds

:Nu represents any nucleophile – they always have a **lone pair** and act as **electron pair donators**

The nucleophiles attack the positive carbon atom



The carbon has a small positive charge because of the electronegativity difference between the carbon and the halogen

We use curly arrows in mechanisms (with two line heads) to show the movement of two electrons

A curly arrow will always **start** from a **lone pair** of electrons or the **centre of a bond**

The rate of these substitution reactions depends on the strength
of the C-X bond
The weaker the bond, the easier it is to break and the faster the reaction.

The iodoalkanes are the fastest to substitute and the fluoroalkanes are the slowest. The strength of the C-F bond is such that fluoroalkanes are very unreactive

	Bond enthalpy / kJmol ⁻¹	
C-I	238	
C-Br	276	
C-Cl	338	
C-F	484	

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I-bromopropane

2-chloro-2-methylbutane

Comparing the rate of hydrolysis reactions

Hydrolysis is defined as the splitting of a molecule (in this case a halogenoalkane) by a reaction with water

$CH_{3}CH_{2}X + H_{2}O \rightarrow CH_{3}CH_{2}OH + X^{-} + H^{+}$

Aqueous silver nitrate is added to a halogenoalkane. The halide leaving group combines with a silver ion to form a silver halide precipitate.

The precipitate only forms when the halide ion has left the halogenoalkane and so the rate of formation of the precipitate can be used to compare the reactivity of the different halogenoalkanes.

The **quicker** the **precipitate** is formed, the **faster** the **substitution** reaction and the **more reactive** the halogenoalkane.

The rate of these substitution reactions depends on the strength of the C-X bond . The weaker the bond, the easier it is to break and the faster the reaction.

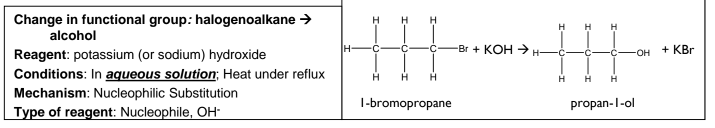
Water is a **poor nucleophile** but it can react **slowly** with halogenoalkanes in a substitution reaction

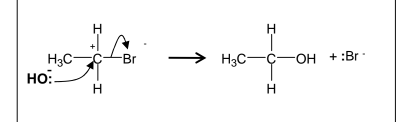
 $CH_3CH_2I + H_2O \rightarrow CH_3CH_2OH + I^- + H^+$ Ag⁺_(aq) + I⁻_(aq) → AgI _(s) - yellow precipitate

The iodoalkane forms a precipitate with the silver nitrate first as the C-I bond is weakest and so it hydrolyses the quickest

Agl $_{(s)}$ - yellow precipitate Ag**Br** $_{(s)}$ - cream precipitate Ag**Cl** $_{(s)}$ - white precipitate

Nucleophilic substitution with <u>aqueous</u> hydroxide ions

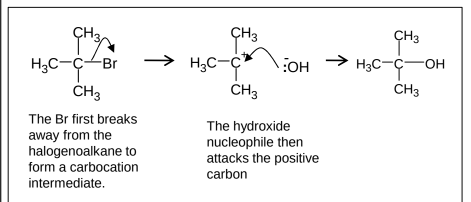




The **aqueous** conditions needed is an important point. If the solvent is changed to **ethanol** an elimination reaction occurs.

Alternative mechanism for tertiary halogenoalkanes

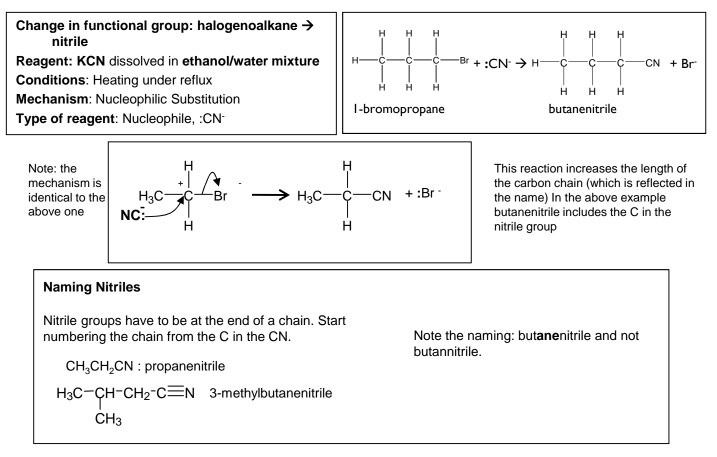
Tertiary halogenoalkanes undergo nucleophilic substitution in a different way



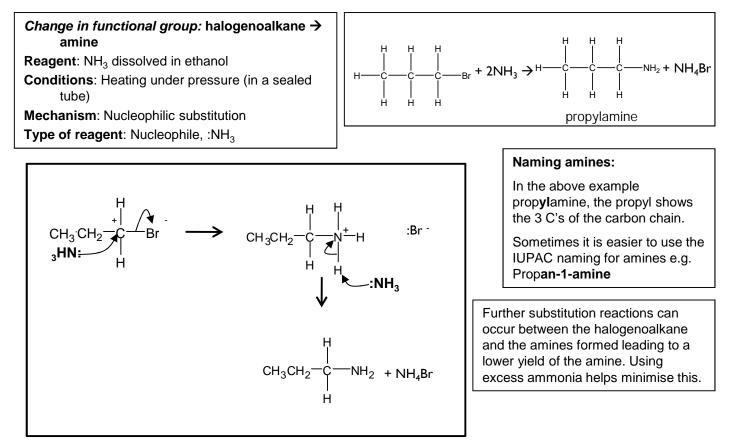
You don't need to learn this but there have been application of understanding questions on this

Tertiary halogenoalkanes undergo this mechanism as the tertiary carbocation is stabilised by the electron releasing methyl groups around it. (See alkenes topic for another example of this). Also the bulky methyl groups prevent the hydroxide ion from attacking the halogenoalkane in the same way as the mechanism above.

Nucleophilic substitution with cyanide ions



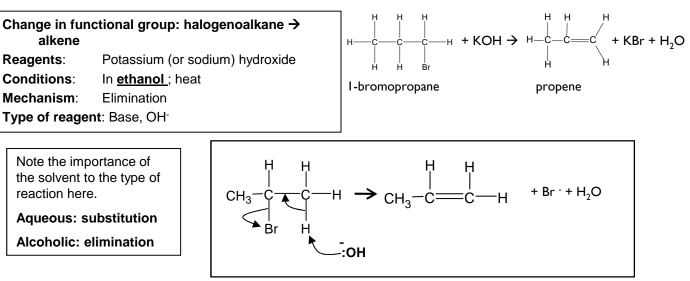
Nucleophilic substitution with ammonia



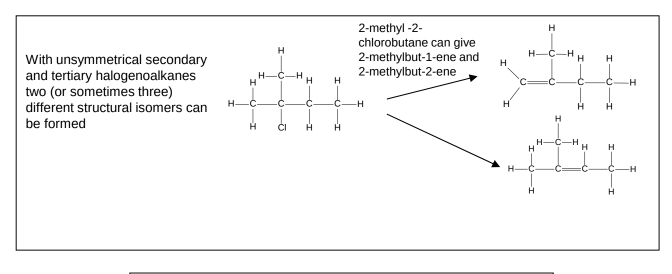
2. Elimination reaction of halogenoalkanes

Elimination with alcoholic hydroxide ions

Elimination: removal of small molecule (often water) from the organic molecule



Often a mixture of products from both elimination and substitution occurs



The structure of the halogenoalkane also has an effect on the degree to which substitution or elimination occurs in this reaction.

Primary tends towards substitution

Tertiary tends towards elimination

Uses of Halogenoalkanes

Chloroalkanes and chlorofluoroalkanes can be used as solvents.

CH₃CCl₃ was used as the solvent in dry cleaning.

Many of these uses have now been stopped due to the toxicity of halogenoalkanes and also their detrimental effect on the atmosphere.

Ozone Chemistry

The naturally occurring ozone (O_3) layer in the upper atmosphere is beneficial as it filters out much of the sun's harmful UV radiation.

Halogenoalkanes have also been used as refrigerants, pesticides and aerosol propellants

Ozone in the lower atmosphere is a pollutant and contributes towards the formation of smog.

Man-made chlorofluorocarbons (CFC's) caused a hole to form in the ozone layer.

Chlorine radicals are formed in the upper atmosphere when energy from ultra-violet radiation causes C–Cl bonds in chlorofluorocarbons (CFCs) to $CF_2Cl_2 \rightarrow CF_2Cl + Cl$ break.

The chlorine free radical atoms **catalyse** the decomposition of ozone, due to these reactions, because they are regenerated. (They provide an alternative route with a lower activation energy)

These reactions contributed to the formation of a hole in the ozone layer.

 $CI^{\bullet} + O_{3} \rightarrow CIO^{\bullet} + O_{2}$ $\rightarrow CIO^{\bullet} + O_{3} \rightarrow 2O_{2} + CI^{\bullet}$ Overall equation $2 O_{3} \rightarrow 3 O_{2}$

The regenerated Cl radical means that one Cl radical could destroy many thousands of ozone molecules.

Legislation to ban the use of CFCs was supported by chemists and that they have now developed alternative chlorine-free compounds. HFCs (Hydro fluoro carbons) e.g. CH_2FCF_3 are now used for refrigerators and air-conditioners. These are safer as they do not contain the C-Cl bond.

The C-F bond is stronger than the C-Cl bond and is not affected by UV.