



Cambridge (CIE) A Level Chemistry



Halogenoalkanes

Contents

- * Producing Halogenoalkanes
- * Substitution Reactions of Halogenoalkanes
- * Elimination Reactions of Halogenoalkanes
- * S_N1 & S_N2
- * Reactivity of Halogenoalkanes



Production of Halogenoalkanes: Substitution & Addition Reactions

- **Halogenoalkanes** are alkanes that have one or more halogens
- They can be produced from:
 - **Free-radical substitution** of alkanes
 - **Electrophilic addition** of alkenes
 - **Substitution** of an alcohol

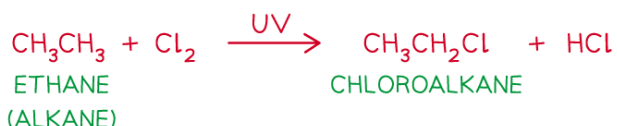
Free-radical substitution of alkanes

- A free-radical substitution reaction involves **three main steps**: **initiation**, **propagation**, and **termination**
- This reaction occurs between an alkane and a halogen, such as **chlorine (Cl_2)** or **bromine (Br_2)**, in the presence of **ultraviolet (UV) light**
 - **Initiation:**
 - UV light provides energy to break the Cl–Cl or Br–Br bond by **homolytic fission**, producing two identical **halogen free radicals** ($\text{Cl}\cdot$ or $\text{Br}\cdot$)
 - **Propagation:**
 - The halogen radicals react with alkane molecules in a **chain reaction**, producing new radicals and continuing the substitution of hydrogen atoms with halogen atoms
 - **Termination:**
 - The reaction stops when **two free radicals combine** to form a stable molecule, ending the chain process

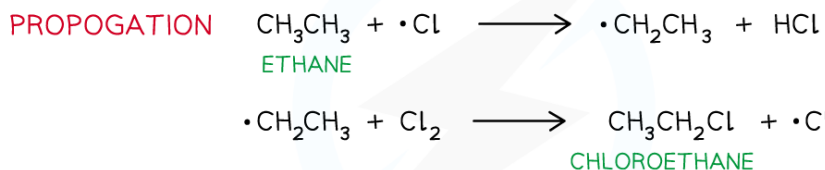
Free-radical substitution reactions of alkanes



Your notes



INITIATION $\text{Cl}-\text{Cl} \xrightarrow{\text{UV}} 2\cdot\text{Cl}$



$$\begin{array}{c} \cdot\text{CH}_2\text{CH}_3 + \cdot\text{CH}_2\text{CH}_3 \longrightarrow \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_3 \\ \text{BUTANE} \\ \cdot\text{Cl} + \cdot\text{Cl} \longrightarrow \text{Cl}_2 \\ \text{CHLORINE} \end{array}$$

Copyright © Save My Exams. All Rights Reserved

Free-radical substitution reactions of alkanes produce halogenoalkanes

Electrophilic addition

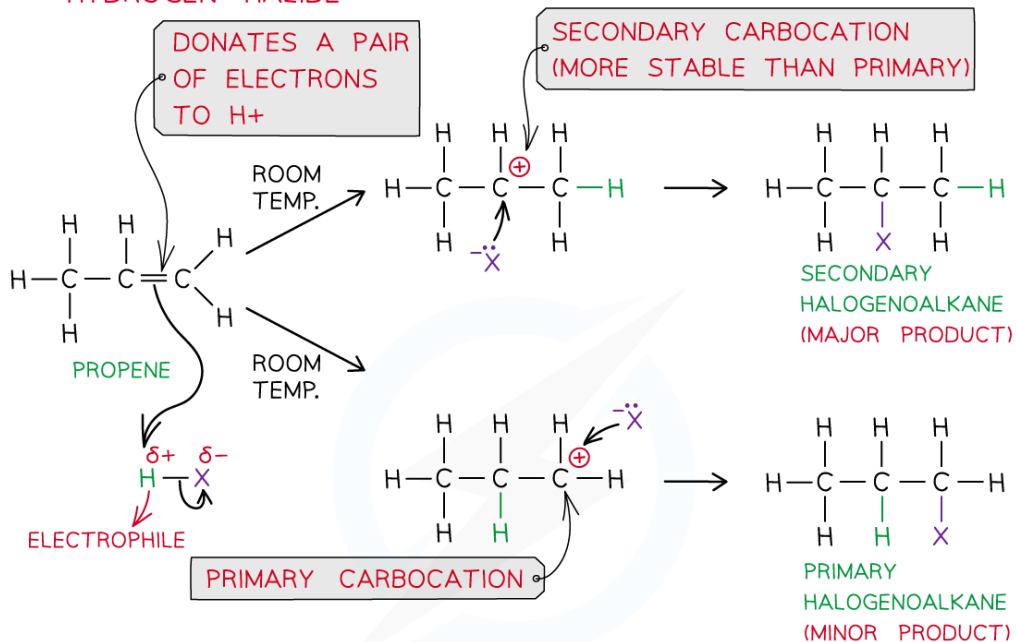
- Halogenoalkanes can also be produced from the addition of **hydrogen halides** (HX) or **halogens** (X_2) at **room temperature** to alkenes
 - In hydrogen halides, the hydrogen acts as the electrophile and accepts a pair of electrons from the C-C bond in the alkene
 - The major product is the one in which the halide is bonded to the most substituted carbon atom (**Markovnikov's rule**)
 - In the addition of halogens to alkenes, one of the halogen atoms acts as an electrophile and the other as a nucleophile

Electrophilic addition to alkenes

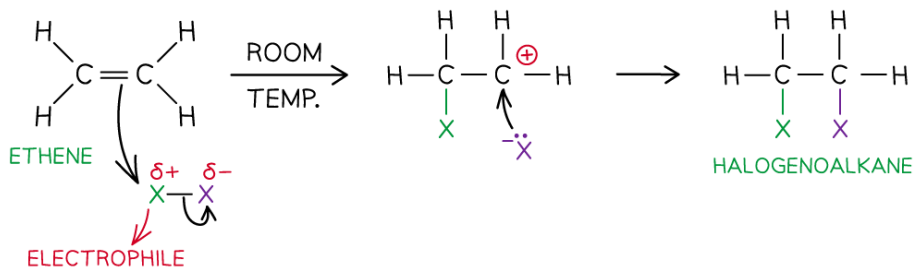


Your notes

• HYDROGEN HALIDE



• HALOGEN



Copyright © Save My Exams. All Rights Reserved

Electrophilic addition of hydrogen halides or hydrogen at room temperatures to alkenes results in the formation of haloalkanes

Substitution of alcohols

- In the **substitution of alcohols** an alcohol group is replaced by a halogen to form a haloalkane
- The substitution of the alcohol group for a halogen can be achieved by reacting the alcohol with:
 - HX (or KBr with H₂SO₄ or H₃PO₄ to make HX)
 - PCl₃ and heat
 - PCl₅ at room temperature
 - SOCl₂

Substitution of alcohols



Your notes

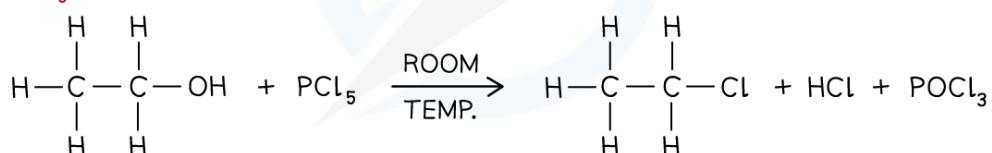
- HX (OR KBr WITH H_2SO_4 OR H_3PO_4)



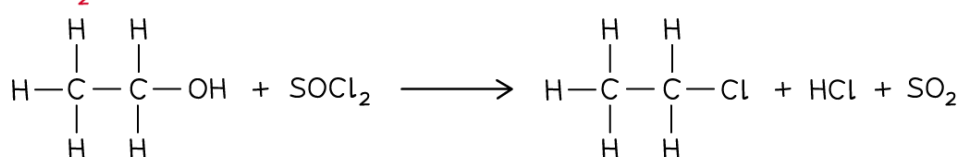
- PCl_3 + HEAT



- PCl_5



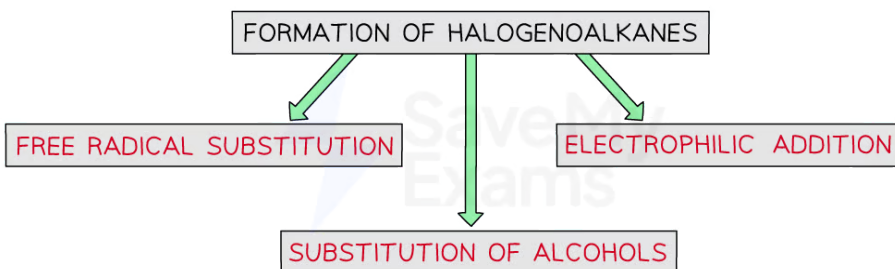
- SOCl_2

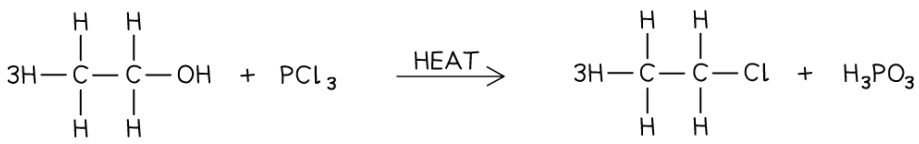


Copyright © Save My Exams. All Rights Reserved

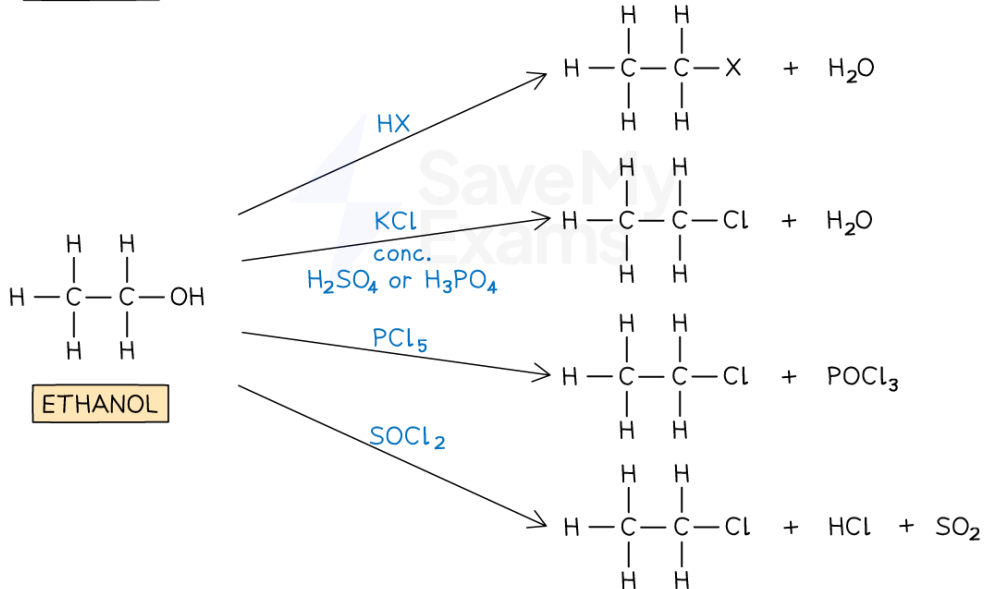
Substitution of alcohols produces halogenoalkanes

Different methods of forming halogenoalkanes





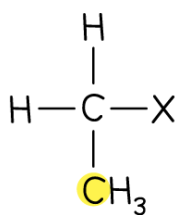
ETHANOL



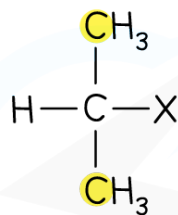
Classifying Halogenoalkanes

- Depending on the carbon atom the halogen is attached to, halogenoalkanes can be classified as **primary**, **secondary** and **tertiary**
 - A **primary halogenoalkane** is when a halogen is attached to a carbon that itself is attached to one other alkyl group
 - A **secondary halogenoalkane** is when a halogen is attached to a carbon that itself is attached to two other alkyl groups
 - A **tertiary halogenoalkane** is when a halogen is attached to a carbon that itself is attached to three other alkyl groups

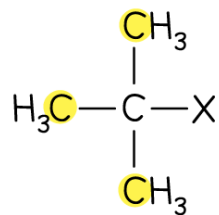
Primary, secondary and tertiary halogenoalkanes



PRIMARY
HALOGENOALKANE



SECONDARY
HALOGENOALKANE



TERTIARY
HALOGENOALKANE

WHERE X IS A HALOGEN

Copyright © Save My Exams. All Rights Reserved



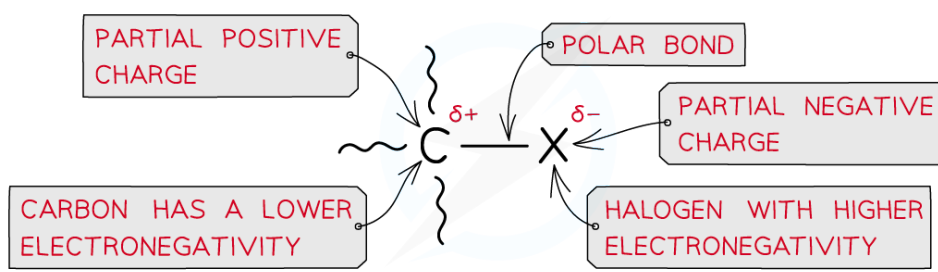
The number of alkyl groups attached determines if the halogenoalkane is primary, secondary or tertiary



Nucleophilic Substitution Reactions of Halogenoalkanes

- **Halogenoalkanes** are much more reactive than alkanes due to the presence of the **electronegative** halogens
 - The halogen-carbon bond is polar causing the carbon to carry a partial positive and the halogen a partial negative charge
- A **nucleophilic substitution** reaction is one in which a **nucleophile** attacks a carbon atom which carries a **partial positive charge**
- An atom that has a **partial negative charge** is replaced by the nucleophile

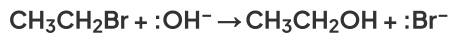
Explaining the polarity of a carbon-halogen bond



Due to large differences in electronegativity between the carbon and halogen atom, the C-X bond is polar

Reaction with NaOH

- The reaction of a halogenoalkane with **aqueous alkali** results in the formation of an alcohol
- The halogen is **replaced** by the OH^-
- The **aqueous hydroxide** (OH^- ion) behaves as a **nucleophile** by donating a pair of electrons to the carbon atom bonded to the halogen
- For example, bromoethane reacts with aqueous alkali when heated to form ethanol
 - Hence, this reaction is a **nucleophilic substitution**
 - The halogen is replaced by a nucleophile, $:\text{OH}^-$



Reaction with KCN

- The **nucleophile** in this reaction is the **cyanide**, CN^- ion

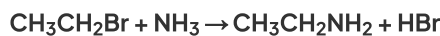


Your notes

- **Ethanolic solution of potassium cyanide** (KCN in ethanol) is **heated under reflux** with the halogenoalkane
- The product is a **nitrile**
 - For example, bromoethane reacts with ethanolic potassium cyanide when heated under reflux to form propanenitrile
 - The halogen is replaced by a nucleophile, :CN⁻
- The nucleophilic substitution of halogenoalkanes with KCN adds an **extra** carbon atom to the carbon chain
- This reaction can therefore be used by chemists to make a compound with one more carbon atom than the best available organic starting material

Reaction with NH₃

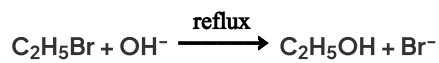
- The **nucleophile** in this reaction is the **ammonia**, NH₃ molecule
- An **ethanolic solution** of **excess ammonia** (NH₃ in ethanol) is **heated under pressure** with the halogenoalkane
- For example, bromoethane reacts with excess ethanolic ammonia when heated under pressure to form ethylamine
 - The product is a **primary amine**
 - The halogen is replaced by an amine group NH₂



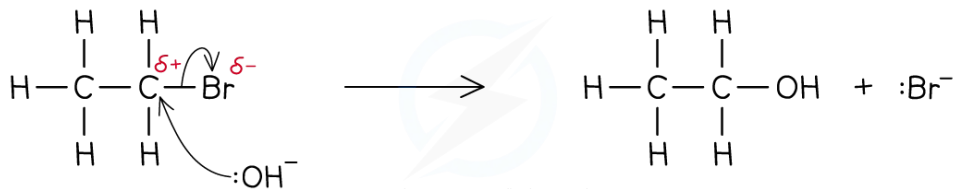
- It is very important that the ammonia is in excess as the product of the nucleophilic substitution reaction, the **ethylamine**, can act as a nucleophile and attack another bromoethane to form the **secondary amine, diethylamine**

Reaction with aqueous silver nitrate

- Halogenoalkanes can be broken down **under reflux** by water to form **alcohols**
- The breakdown of a substance by water is also called **hydrolysis**
- The water in aqueous silver nitrate will hydrolyse the halogenoalkane
 - The fastest nucleophilic substitution reactions take place with the iodoalkanes as the C-I bond is the **weakest (longest)**
 - The slowest nucleophilic substitution reactions take place with the fluoroalkanes as the bond is the **strongest (shortest)**
- For example, bromoethane reacts with aqueous silver nitrate solution to form ethanol and a Br⁻ ion
 - The Br⁻ ion will form a cream precipitate with Ag⁺
- This reaction is classified as a nucleophilic substitution reaction with **water molecules** in aqueous **silver nitrate** solution acting as nucleophiles, **replacing** the halogen in the



Nucleophilic substitution with OH⁻



In nucleophilic substitution with OH⁻, the bond that forms and the bond that breaks must both involve the carbon atom that is bonded to the leaving group

- This reaction is similar to the nucleophilic substitution reaction of halogenoalkanes with aqueous alkali, however, hydrolysis with water is much **slower** than with the OH⁻ ion in alkalis
 - The hydroxide ion is a better nucleophile than water as it carries a full formal negative charge
 - In water, the oxygen atom only carries a partial negative charge

Comparing water and the hydroxide ion as nucleophiles



A hydroxide ion is a better nucleophile as it has a full formal negative charge whereas the oxygen atom in water only carries a partial negative charge; this causes the nucleophilic substitution reaction with water to be much slower than with aqueous alkali

- The halogenoalkanes have different **rates of hydrolysis**
- This reaction can be used as a **test** to identify halogens in a halogenoalkane by measuring how long it takes for the test tubes containing the halogenoalkane and aqueous silver nitrate solutions to become **opaque**



Elimination Reactions of Halogenoalkanes

- An elimination reaction involves the loss of a small molecule from a larger organic molecule
- In halogenoalkanes, this small molecule is usually a **hydrogen halide** (e.g. HBr or HCl)
- The product is typically an **alkene**
 - The elimination reaction of bromoethane with ethanolic sodium hydroxide

Reaction Conditions for Elimination

- Halogenoalkanes are heated with **ethanolic sodium hydroxide** (NaOH dissolved in ethanol)
- Under these **anhydrous conditions**, elimination occurs:
 - The **C-X bond** (where X = halogen) breaks **heterolytically**
 - A **halide ion (X⁻)** is released
 - A **double bond** forms, producing an **alkene**
- E.g. Elimination of bromoethane:



- One **hydrogen atom** and the **bromine atom** are eliminated
- The carbon chain forms a **C=C double bond** to make **ethene**



Examiner Tips and Tricks

Reaction **conditions are crucial** in determining the product.

Elimination = hot, ethanolic NaOH

Substitution = warm, aqueous NaOH



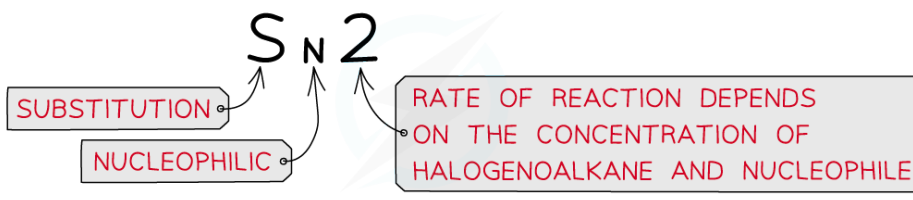
Halogenoalkanes: SN1 & SN2 Mechanisms

- In **nucleophilic substitution** reactions involving halogenoalkanes, the halogen atom is replaced by a nucleophile
- These reactions can occur in two different ways (known as **SN2** and **SN1** reactions) depending on the structure of the halogenoalkane involved

SN2 reactions

- In **primary halogenoalkanes**, the carbon that is attached to the halogen is bonded to **one alkyl group**
- These halogenoalkanes undergo nucleophilic substitution by an **SN2** mechanism
 - 'S' stands for 'substitution'
 - 'N' stands for 'nucleophilic'
 - '2' means that the rate of the reaction (which is determined by the slowest step of the reaction) depends on the concentration of both the halogenoalkane and the nucleophile ions

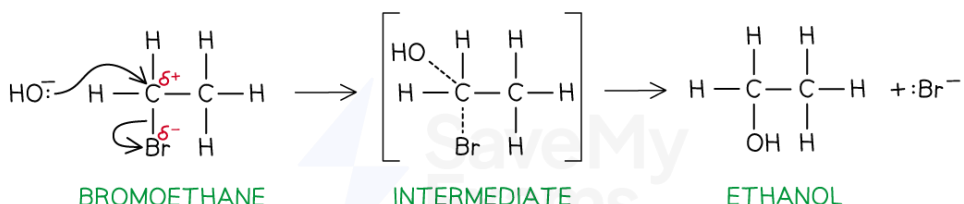
Defining an SN2 mechanism



Each term in the SN2 expression has a specific meaning

- The **SN2** mechanism is a **one-step** reaction
 - The nucleophile donates a pair of electrons to the $\delta+$ carbon atom to form a new bond
 - At the same time, the C-X bond is breaking and the halogen (X) takes both electrons in the bond (**heterolytic fission**)
 - The halogen leaves the halogenoalkane as an X^- ion
- For example, the nucleophilic substitution of bromoethane by hydroxide ions to form ethanol

The nucleophilic substitution of bromoethane by hydroxide ions



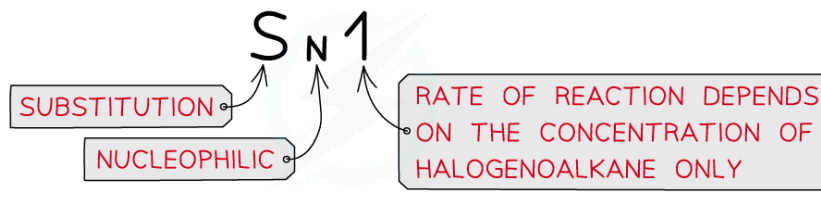
 Your notes

In this mechanism, the bromoethane is a primary halogenoalkane

S_N1 reactions

- In **tertiary halogenoalkanes**, the carbon that is attached to the halogen is bonded to **three alkyl groups**
 - These halogenoalkanes undergo nucleophilic substitution by an **S_N1** mechanism
 - ‘S’ stands for ‘substitution’
 - ‘N’ stands for ‘nucleophilic’
 - ‘1’ means that the rate of the reaction (which is determined by the slowest step of the reaction) depends on the concentration of only one reagent, the **halogenoalkane**

Defining an S_N1 mechanism



Each term in the SN1 expression has a specific meaning

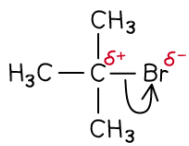
- The S_N1 mechanism is a two-step reaction
 - In the first step, the C-X bond breaks **heterolytically** and the halogen leaves the halogenoalkane as an X⁻ ion (this is the **slow** and **rate-determining step**)
 - This forms a **tertiary carbocation** (**which is a tertiary carbon atom with a positive charge**)
 - In the second step, the tertiary carbocation is attacked by the **nucleophile**
 - For example, the nucleophilic substitution of 2-bromo-2-methylpropane by hydroxide ions to form 2-methyl-2-propanol

The nucleophilic substitution of 2-bromo-2-methylpropane by hydroxide ions



Your notes

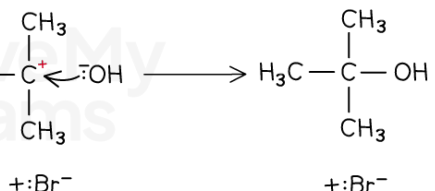
TERTIARY HALOGENOALKANE



2-BROMO-2-METHYL PROPANE

TERTIARY CARBOCATION

SLOW
(RATE DETERMINING STEP)



2-METHYL-2-PROPANOL

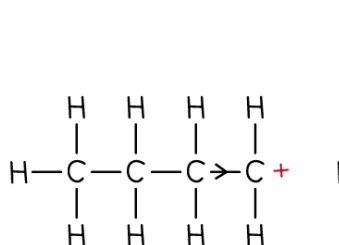
Copyright © Save My Exams. All Rights Reserved

In this mechanism, the 2-bromo-2-methylpropane is a tertiary halogenoalkane

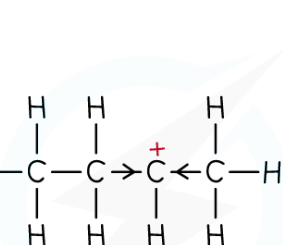
Carbocations

- In the S_N1 mechanism, a **tertiary carbocation** is formed
- This is not the case for S_N2 mechanisms as a **primary carbocation** would have been formed which is much **less** stable than tertiary carbocations
- This has to do with the **positive inductive effect** of the alkyl groups attached to the carbon which is bonded to the halogen atom
 - The alkyl groups push electron density towards the positively charged carbon, reducing the **charge density**
 - In tertiary carbocations, there are three alkyl groups stabilising the carbocation whereas in primary carbocations there is only one alkyl group
 - This is why tertiary carbocations are much more stable than primary ones

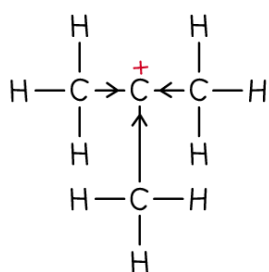
Stability of primary, secondary and tertiary carbocations



PRIMARY CARBOCATION (LEAST STABLE)



SECONDARY CARBOCATION



TERTIARY CARBOCATION (MOST STABLE)

Copyright © Save My Exams. All Rights Reserved

The carbocations become more stable moving from primary to secondary to tertiary

- **Secondary halogenoalkanes** undergo a **mixture** of both S_N1 and S_N2 reactions depending on their structure



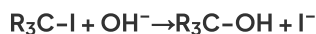
Reactivity of Halogenoalkanes

- The halogenoalkanes have different **rates** of **substitution reactions**
- Since substitution reactions involve **breaking** the **carbon-halogen** bond the **bond energies** can be used to explain their different reactivities

Halogenoalkane bond energy table

Bond	Bond Energy / kJ mol ⁻¹
C-F	467 (strongest bond)
C-Cl	346
C-Br	290
C-I	228 (weakest bond)

- The table above shows that the C-I bond requires the least energy to break, and is therefore the **weakest** carbon-halogen bond
- During substitution reactions the C-I bond will therefore **heterolytically** break as follows:



halogenoalkane → alcohol

- The C-F bond, on the other hand, requires the most energy to break and is, therefore, the **strongest** carbon-halogen bond
- Fluoroalkanes will therefore be less likely to undergo substitution reactions

Aqueous silver nitrate

- Reacting halogenoalkanes with **aqueous silver nitrate solution** will result in the formation of a **precipitate**
- The **rate of formation** of these precipitates can also be used to determine the reactivity of the halogenoalkanes

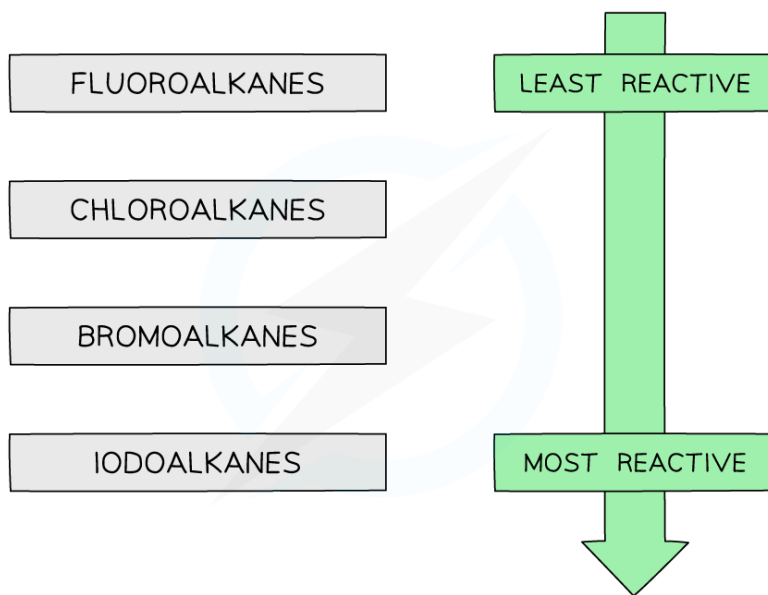
Halogenoalkane precipitates

- Chlorides
 - White (silver chloride) precipitate
- Bromides

- Cream (silver bromide)
- Iodides
- Yellow (silver iodide)
- The formation of the pale yellow silver iodide is the fastest (fastest **nucleophilic substitution** reaction)
- The formation of the silver fluoride is the slowest (slowest **nucleophilic substitution** reaction)
- This confirms that fluoroalkanes are the least reactive and iodoalkanes are the most reactive halogenoalkanes



The trend in reactivity of halogenoalkanes



Copyright © Save My Exams. All Rights Reserved

The halogenoalkanes become more reactive as you move down the group