Cambridge (CIE) A Level Chemistry



Halogen Compounds

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Producing Halogenoarenes



Production of Halogenoarenes

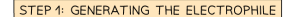
- Halogenoarenes are arenes which are bonded to halogen atoms
- They can be prepared from **substitution reactions** of arenes with chlorine or bromine in the presence of an anhydrous catalyst

Substitution of benzene to form halogenoarenes

- Chlorine gas is bubbled into benzene at room temperature and in the presence of an anhydrous AICl₃ catalyst to form **chlorobenzene**
- The AlCl₃ catalyst is also called a **halogen carrier** and is required to generate the electrophile (CI+)
- This electrophile attacks the electron-rich benzene ring in the first stage of the reaction which **disrupts** the delocalised π system in the ring
- To restore the aromatic stabilization, a hydrogen atom is removed in the second stage of the electrophilic substitution reaction to form chlorobenzene
- When this happens, the delocalised π system of the ring is **restored**
- The same reaction occurs with benzene and bromine in the presence of an AlBr₃ catalyst to form bromobenzene

Formation of chlorobenzene







$$\stackrel{\delta^{+}}{\text{Cl}} - \stackrel{\delta^{-}}{\text{Cl}} \longrightarrow \text{Cl}^{+} + [\text{AlCl}_{4}]^{-}$$

STEP 2: ELECTROPHILIC ATTACK

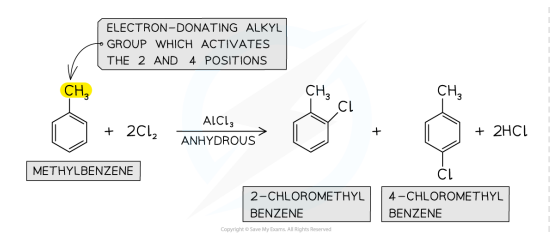
STEP 3: RESTORING AROMATICITY

Halogenoarenes can be formed from the electrophilic substitution reaction of arenes with halogens

Substitution of methylbenzene to form halogenoarenes

- The **electrophilic substitution** of **methylbenzene** with halogens results in the formation of multiple halogenoarenes as products
- This is because the methyl group (which is an alkyl group) in methylbenzene is electrondonating and pushes electron density into the benzene ring
- This makes the benzene ring more reactive towards electrophilic substitution reactions
- The methyl group is said to be 2,4-directing and as a result, the 2 and 4 positions are activated
- Electrophilic substitution of methylbenzene with chlorine and anhydrous AlCl₃ catalyst, therefore, gives 2-chloromethylbenzene and 4-chloromethylbenzene
- The reaction mechanism is **the same** as the substitution mechanism of benzene

Chlorination of methylbenzene





The methyl group on methylbenzene directs the incoming halogen on the 2 and 4 position

• In the presence of excess chlorine, substitution on the 6 position will also occur



Reactivity of Halogenoarenes



Difference in Reactivity of Halogenoalkanes & Halogenoarenes

- Halogenoarenes are very unreactive compared to halogenoalkanes
- The difference in reactivity between the two compounds is because of the carbonhalogen bond strengths

Halogenoalkanes

- The halogenoalkane chloroethane can take part in nucleophilic substitution reactions
- A nucleophile, such as a hydroxide (OH⁻) ion, will attack the slightly positive carbon
- A covalent bond is formed between that carbon atom and the nucleophile which causes the carbon-halogen bond to break
- Overall, the halogen is replaced by the nucleophile

Nucleophilic substitution of halogenoalkanes

Halogenoalkanes readily undergo nucleophilic substitution reactions due to the difference in electronegativity between the carbon and halogen

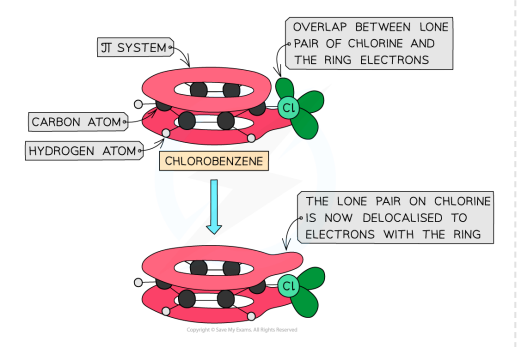
Halogenoarenes

- Halogenoarenes, such as chlorobenzene, **do not** readily undergo **nucleophilic** substitution reactions
 - Only under extremely **harsh** conditions, such as temperatures of 200 °C and a pressure of 200 atmospheres, will the chlorine in chlorobenzene get replaced by a nucleophile such as a hydroxide (OH-) ion
- This is because the carbon-chlorine bond is **very strong** and cannot be easily broken
 - One of the lone pairs of electrons on the chlorine will **interact** with the π system of the ring
 - This causes the carbon-chlorine bond to have a partial double-bond character, which strengthens the bond



How the carbon-chlorine bond affects the π system





The carbon-chlorine bond is very strong, as it has partial double-bond character

- Therefore, the **unreactivity** of halogenoarenes can be explained by the **delocalisation** of a lone pair on the halogen over the benzene
- This causes additional **stabilisation** of the system and **strengthens** the carbon-halogen bond, which affects the reactions that halogenoarenes will undergo
- It gets **harder** to break the carbon-halogen bond in halogenoarenes, which decreases reactivity