# Cambridge (CIE) A Level Chemistry



## **Arenes**

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- \* Reactions of Arenes
- \* Electrophilic Substitution of Arenes
- \* Location of Halogenation on Arenes
- \* Directing Effects of Substituents on Arenes



### **Reactions of Arenes**



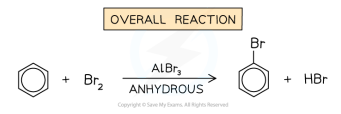
## **Reactions of Arenes**

- Arenes are very stable compounds due to the **delocalisation of**  $\pi$  **electrons** in the ring
  - This is because the negative charge is spread out over the molecule instead of being confined to a small area
- During chemical reactions such as substitution reactions, this delocalised ring is maintained
  - Addition reactions however, disrupt the aromatic stabilisation
- Benzene undergoes a wide range of reactions including combustion -(complete and incomplete) and the following reactions:
  - Halogenation
  - Nitration
  - Friedel-Craft's alkylation
  - Friedel-Craft's acylation
  - Complete Oxidation
  - Hydrogenation

## Halogenation

- Halogenation reactions are examples of electrophilic substitution reactions
- Arenes undergo **substitution** reactions with chlorine (Cl<sub>2</sub>) and bromine (Br<sub>2</sub>) in the presence of anhydrous AICI<sub>3</sub> or AIBr<sub>3</sub> catalyst respectively to form halogenoarenes (aryl halides)
  - The chlorine or bromine acts as an **electrophile** and replaces a hydrogen atom on the benzene ring
  - The catalyst is required for the reaction to take place, due to the stability of the benzene structure

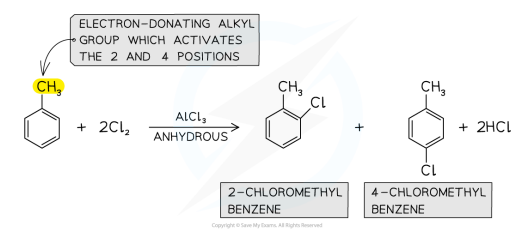
## Halogenation of benzene



Arenes undergo substitution reactions with halogens to form aryl halides

- Alkylarenes such as methylbenzene undergo halogenation on the 2 or 4 positions
- This is due to the **electron-donating** alkyl groups which activate these positions
  - Phenol ( $C_6H_5OH$ ) and phenylamine ( $C_6H_5NH_2$ ) are also activated in the 2 and 4 positions
- The halogenation of alkylarenes, therefore, results in the formation of **two products**

### Halogenation of alkylarenes



#### Alkylarenes are substituted on the 2 or 4 position

■ Multiple substitutions occur when excess halogen is used

### Halogenation of alkylarenes using an excess of halogen

In the presence of excess halogen, multiple substitutions occur

### **Nitration**

- Another example of a substitution reaction is the **nitration** of arenes
- In these reactions, a nitro (-NO<sub>2</sub>) group replaces a hydrogen atom on the arene
- The benzene is reacted with a mixture of concentrated nitric acid (HNO<sub>3</sub>) and concentrated sulfuric acid ( $H_2SO_4$ ) at a temperature between 25 and 60  $^{\circ}C$

### Nitration of benzene



Your notes





### During nitration, a hydrogen atom is replaced by an $NO_2$ group

• Again, due to the **electron-donating** alkyl groups in alkylarenes, nitration of methylbenzene will occur on the 2 and 4 position

### Nitration of alkylarenes

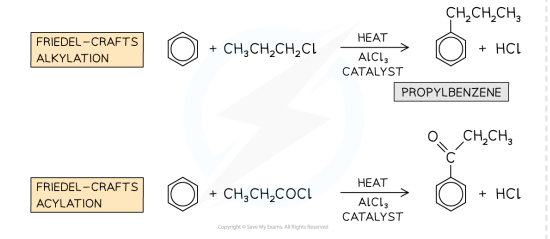
### Alkylarenes are nitrated on the 2 or 4 position

### Friedel-Crafts reactions

- Friedel-Crafts reactions are also **electrophilic substitution** reactions
- Due to the aromatic stabilisation in arenes, they are often **unreactive**
- To use arenes as **starting materials** for the synthesis of other organic compounds, their structure, therefore, needs to be changed to turn them into more reactive compounds
- Friedel-Crafts reactions can be used to substitute a hydrogen atom in the benzene ring for an **alkyl group** (Friedel-Crafts alkylation) or an **acyl group** (Friedel-Crafts acylation)
- Like any other electrophilic substitution reaction, the Friedel-Crafts reactions consist of three steps:
  - 1. Generating the electrophile
  - 2. Electrophilic attack on the benzene ring
  - 3. Regenerating aromaticity of the benzene ring

### Examples of Friedel-Crafts alkylation and acylation reactions







During alkylation, an alkyl / R group is substituted on the benzene ring and during acylation, an acyl / RCO group is substituted on the benzene ring

## Friedel-Crafts alkylation

- In this type of Friedel-Crafts reaction, an **alkyl chain** is substituted into the benzene ring
- The benzene ring is reacted with a chloroalkane in the presence of an AICl<sub>3</sub> catalyst
- An example of an alkylation reaction is the reaction of benzene with chloropropane (CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CI) to form propylbenzene

### Example of a Friedel-Crafts alkylation reaction

### STEP 1: GENERATING THE ELECTROPHILE

CHLOROPROPANE

### STEP 2: ELECTROPHILIC ATTACK

### STEP 3: RESTORING AROMATICITY



Alkylation reactions of benzene follow the 3 steps of electrophile generation, electrophilic attack and regeneration of aromaticity

## Friedel-Crafts acylation

- In the Friedel-Crafts acylation reaction, an **acyl group** is substituted into the benzene ring
  - An acyl group is an alkyl group containing a carbonyl, C=O group
- The benzene ring is reacted with an acyl chloride in the presence of an AlCl<sub>3</sub> catalyst
- An example of an acylation reaction is the reaction of methylbenzene with propanoyl chloride to form an acyl benzene
  - Note that the acyl group substitutes on the 4 position due to the -CH<sub>3</sub> group on the benzene

### Example of a Friedel-Crafts acylation reaction

#### STEP 1: GENERATING THE ELECTROPHILE

PROPANOYL CHLORIDE

### STEP 2: ELECTROPHILIC ATTACK

$$CH_3$$
 $CH_3$ 
 $CH_3$ 
 $CH_3$ 
 $CH_3$ 
 $CH_2CH_3$ 
 $CH_3$ 
 $CH_3$ 
 $CH_3$ 

### STEP 3: RESTORING AROMATICITY



$$\begin{array}{c} CH_{3} \\ + [AlCl_{4}]^{-} \\ \end{array} \begin{array}{c} CH \\ + HCl + AlCl_{3} \\ C = O \\ CH_{2}CH_{3} \end{array}$$

Acylation reactions of benzene follow the same 3 steps of electrophile generation, electrophilic attack and regeneration of aromaticity

### Complete oxidation

- Normally, alkanes are not **oxidised** by **oxidising agents** such as potassium manganate(VII) (KMnO<sub>4</sub>)
- However, the presence of the benzene ring in alkyl arenes affects the properties of the alkyl side-chain
- The alkyl side-chains in alkyl arenes are **oxidised** to **carboxylic acids** when **refluxed** with alkaline potassium manganate(VII) and then acidified with dilute sulfuric acid ( $H_2SO_4$ )
  - For example, the complete oxidation of **ethylbenzene** forms **benzoic acid**

### Oxidation of alkylarenes



The complete oxidation of alkyl side chains in arenes gives a carboxylic acid

## **Hydrogenation**

- The hydrogenation of benzene is an addition reaction
- Benzene is **heated** with **hydrogen gas** and a **nickel** or **platinum catalyst** to form cyclohexane

## Hydrogenation of benzene

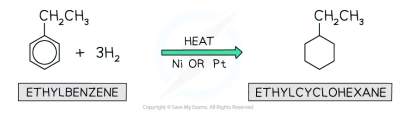




### Hydrogenation of benzene results in a loss of aromaticity

• The same reaction occurs when **ethylbenzene** undergoes hydrogenation to form cycloethylbenzene

### Hydrogenation of methylbenzene



Hydrogenation of alkylarenes also results in a loss of aromaticity

## **Summary of Reactions of Arenes Table**

Reaction	Conditions	Products
Halogenation	$\text{Cl}_2$ with an AICl $_3$ catalyst $\text{Br}_2$ with an AIBr $_3$ catalyst	Aryl halide
Nitration	A mixture of concentrated H <sub>2</sub> SO <sub>4</sub> and concentrated HNO <sub>3</sub> Temperature between 25 °C and 60 °C	Nitroarene
Friedel-Crafts alkylation	Halogenoalkane and anhydrous AlCl <sub>3</sub> catalyst	Alkylbenzene
Freidel-Crafts acylation	Acyl chloride and anhydrous AlCl <sub>3</sub> catalyst	Acylbenzene
Complete oxidation	Hot, alkaline KMnO $_4$ and then dilute acid	Benzoic acid
Hydrogenation	Heating with hydrogen and Pt / Ni catalyst	Cyclohexane

## **Electrophilic Substitution of Arenes**



## **Electrophilic Substitution of Arenes**

- The **electrophilic substitution** reaction in arenes consists of **three steps**:
  - 1. Generation of an electrophile
  - 2. Electrophilic attack
  - 3. Regenerating aromaticity

## Generation of an electrophile

- The **delocalised**  $\pi$  system is extremely stable and is a region of **high electron density**
- Consequently, the first step of an electrophilic substitution reaction involves the generation of an electrophile
  - The electrophile can be a positive ion or the positive end of a polar molecule
- There are numerous electrophiles which can react with benzene:

### Table of electrophiles commonly used with benzene

Reaction type	Electrophile*
halogenation	X <sup>+</sup> , e.g. CI <sup>+</sup>
nitration	NO <sub>2</sub> <sup>+</sup>
Friedel-Craft's alkylation	R <sup>+</sup>
Friedel-Craft's acylation	R-C=O <sup>+</sup>

- Typically electrophiles cannot simply be added to the reaction mixture
  - The electrophile is produced in situ, by adding appropriate reagents\* to the reaction mixture

## Electrophilic attack

- A pair of electrons from the benzene ring is donated to the electrophile to form a covalent bond
- This disrupts the aromaticity in the ring as there are now only four π electrons and there is a positive charge spread over the five carbon atoms

## Regenerating aromaticity



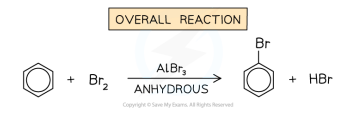
- In the **final step** of electrophilic substitution, the aromaticity of the benzene ring system is restored
- Your notes

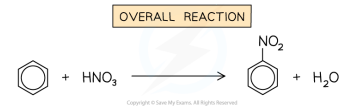
- This happens by **heterolytic cleavage** of the C-H bond
  - This means that the electrons in this bond go into the benzene  $\pi$  bonding system

## Electrophilic substitution mechanism

- The halogenation and nitration of arenes are both examples of electrophilic substitution reactions
  - A hydrogen atom is replaced by a halogen atom or a nitro (-NO<sub>2</sub>) group

### Bromination and nitration of benzene



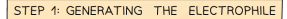


During bromination, a hydrogen atom is substituted by a bromine atom and during nitration, a hydrogen atom is substituted by a nitro group

### Step 1: Generating the Br+ and NO<sub>2</sub>+ electrophiles

- For the **halogenation** reaction:
  - This is achieved by reacting the halogen with a halogen carrier
  - The halogen molecules form a **dative bond** with the halogen carrier by donating a lone pair of electrons from one of its halogen atoms into an empty 3p orbital of the halogen carrier

Step 1 of the halogenation of arenes





$$\begin{array}{c} \text{DATIVE COVALENT} \\ \text{BOND} \\ \\ \text{Br} \xrightarrow{\delta^{+}} \text{Br} & \text{AlBr}_{3} \longrightarrow \text{Br}^{+} + [\text{AlBr}_{4}]^{-} \\ \\ & \xrightarrow{\text{ELECTROPHILE}} \end{array}$$

During bromination, an AlBr<sub>3</sub> halogen carrier catalyst is used and during chlorination an AICI<sub>3</sub> halogen carrier catalyst is used

- For the **nitration** reaction:
  - The electrophile NO<sub>2</sub>+ion is generated by reacting it with concentrated nitric acid  $(HNO_3)$  and concentrated sulfuric acid  $(H_2SO_4)$

Step 1 of the nitration of arenes

STEP 1: GENERATING THE ELECTROPHILE

HNO<sub>3</sub> + 
$$2H_2SO_4$$
  $\longrightarrow$   $NO_2^+$  +  $2HSO_4^-$  +  $H_3O^+$ 

ELECTROPHILE

During nitration, concentrated nitric acid and concentrated sulfuric acid react to form the NO<sub>2</sub><sup>+</sup> electrophile

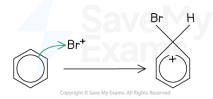
## Step 2: Electrophilic attack by the Br+ and NO<sub>2</sub>+ electrophiles

- Once the electrophile has been generated, it will carry out an electrophilic attack on the benzene ring
  - The nitrating mixture of  $HNO_3$  and  $H_2SO_4$  is **refluxed** with the arene at 25 60 °C
- A pair of electrons from the benzene ring is donated to the electrophile to form a covalent bond
  - This disrupts the aromaticity in the ring as there are now only four  $\pi$  electrons and there is a **positive charge** spread over the five carbon atoms

Step 2 of the halogenation of arenes

### STEP 2: ELECTROPHILIC ATTACK





A pair of electrons from the benzene ring is donated to the Br<sup>+</sup> electrophile to form a covalent bond causing a loss in aromaticity

Step 2 of the nitration of arenes





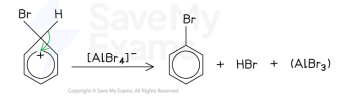
A pair of electrons from the benzene ring is donated to the NO<sub>2</sub><sup>+</sup> electrophile to form a covalent bond causing a loss in aromaticity

### Step 3: Regenerating / restoring aromaticity

- In the **final step** of the reaction, this aromaticity is restored by **heterolytic cleavage** of the C-H bond
  - This means that the bonding pair of electrons goes into the benzene  $\pi$  bonding system

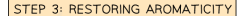
### Step 3 of the halogenation of arenes

### STEP 3: RESTORING AROMATICITY



Step 3 of the nitration of arenes









In both reactions, the C-H bond of the substituted carbon atom breaks and the electrons go back into the benzene  $\pi$  bonding system, restoring aromaticity

### Addition reactions of arenes

- The delocalisation of electrons (also called aromatic stabilisation) in arenes is the main reason why arenes predominantly undergo substitution reactions over addition reactions
- In substitution reactions, the **aromaticity** is restored by **heterolytic cleavage** of the C-H bond
- In addition reactions, on the other hand, the aromaticity is **not restored** and is in some cases completely lost
  - The **hydrogenation** of arenes is an example of an addition reaction during which the aromatic stabilisation of the arene is completely lost
  - The cyclohexane formed is **energetically less stable** than the benzene



### **Location of Halogenation on Arenes**



## **Halogenation in Arenes**

- Arenes will undergo substitution reactions with halogens to form aryl halides
  - This reaction is also called a **halogenation** reaction
- Depending on the reaction conditions, halogenation can occur:
  - In the aromatic ring
  - In the side chain

## Halogenation in the aromatic ring

• Halogenation of alkylarenes in the aromatic ring will occur when a halogen and anhydrous halogen carrier catalyst (such as AIBr<sub>3</sub> or AICl<sub>3</sub>) is used

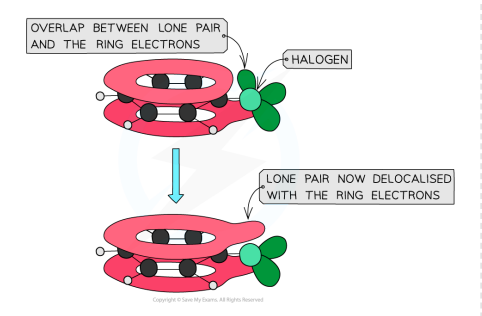
### Halogenation of alkylarenes in the aromatic ring

### A halogen carrier catalyst is used to generate the electrophile for the halogenation of alkylarenes

- Aryl halides are less reactive than halogenoalkanes as the carbon-halogen bond in aryl halides is stronger
- This is due to the partial overlap of the lone pairs on the halogen atom with the  $\pi$  system in the benzene ring
- The carbon-halogen bond, therefore, has a partial double bond character

## The lack of reactivity in alkylarenes / aryl halides





Aryl halides are unreactive due to the partial double bond character of the carbonhalogen bond

## Halogenation in the side chain

- Halogenation of alkylarenes in the side chain will occur when the halogen is passed into boiling alkylarene in the presence of ultraviolet (UV) light
  - This is a **free-radical substitution** reaction

## Halogenation of an alkylarene side chain

### Halogenation of alkylarenes in the side chain is an example of a free-radical substitution reaction

• If excess halogen is used, all hydrogen atoms on the alkyl side-chain will be substituted by the halogen atoms

## Halogenation of an alkylarene side chain using excess halogen





### In excess halogen, all hydrogen atoms on the alkyl side-chain will be replaced

• Note that no substitution into the benzene ring occurs under these conditions

### **Directing Effects of Substituents on Arenes**



## The Directing Effects of Substituents on **Arenes**

- Arenes readily undergo **electrophilic substitution** of one of their hydrogen atoms with another species
- Substituents that are already present on the arenes can affect where the substitution of the hydrogen atom on the arene takes place
  - These groups are said to **direct** substitution reactions to different **ring positions**

## Electron-withdrawing & electron-donating groups

- The substituents on the arenes can either be **electron-withdrawing** or **electron**donating groups
- Electron-withdrawing substituents remove electron density from the π system in the benzene ring making it less reactive
  - These groups **deactivate** attack by electrophiles and **direct** the incoming electrophile to attack the 3 and/or 5 positions
  - For example, the nitro group in nitrobenzene is an electron-withdrawing group
    - Upon bromination of nitrobenzene, the bromine electrophile will be directed to the 3 and/or 5 position
    - The products are 3-bromonitrobenzene and 5-bromonitrobenzene
- Electron-donating substituents **donate** electron density **into** the π system of the benzene ring making it more reactive
  - These groups activate attack by electrophiles and direct the incoming electrophile to attack the 2, 4 and/or 6 positions
  - For example, the methyl group in methylbenzene is an electron-donating group
    - Upon bromination of methylbenzene, the bromine electrophile will be directed to the 2 and/or 4 position
    - The products are 2-bromomethylbenzene and 4-bromomethylbenzene

### Electron-withdrawing & electron-donating substituents table

	Substituents	Activated positions
Electron withdrawing	-NO <sub>2</sub>	3 and / or 5
	-COOH	



	-COR	
Electron donating	-R	2, 4 and / or 6
	-OH	
	-NH <sub>2</sub>	



