6: Organic Chemistry I

6A. Introduction to Organic Chemistry

Hydrocarbon is a compound consisting of hydrogen and carbon only

Saturated: Contain single carbon-carbon bonds only

Unsaturated : Contains a C=C double bond

to know

Basic definitions

Molecular formula: The formula which shows the actual number of each type of atom

Empirical formula: shows the simplest whole number ratio of atoms of each element in the compound

General formula: algebraic formula for a homologous series e.g. CnH2n

Structural formula shows the minimal detail that shows the arrangement of atoms in a molecule, eg for butane: $CH_3CH_2CH_2CH_3$ or $CH_3(CH_2)_2CH_3$,

Displayed formula: show all the covalent bonds present in a molecule

Drawing Displayed formulae



When drawing organic compounds add the hydrogen atoms so that each carbon has 4 bonds Remember that the shape around the carbon atom in saturated hydrocarbons is tetrahedral and the bond angle is 109.5°



Skeletal formula shows the simplified organic formula, shown by removing hydrogen atoms from alkyl chains, leaving just a carbon skeleton and associated functional Groups.



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2-methylbutane

But-2-ene

Butan-1-ol

cyclohexane

cyclohexene

Homologous series are families of organic compounds with the same functional group and same general formula.

- •They show a gradual change in physical properties (e.g. boiling point).
- Each member differs by CH₂ from the last.
- same chemical properties.

Functional group is an atom or group of atoms which when present in different molecules causes them to have similar chemical properties

homologous series	functional group	prefix / suffix (* = usual use)	example
Alkane	cc	-ane	CH ₃ CH ₂ CH ₂ CH ₃ Butane
Alkenes	 C==C 	suffix -ene	H H propene
Alcohols	сон	suffix* -ol prefix hydroxy-	н н н Propan-1-ol <u>H C C C C O H</u> <u>H H H</u> H H H
Halogenoalkanes	Chalogen	prefix chloro- bromo- iodo-	H H H H H - C - C - C - C - C - C - C -
Aldehydes	о —с—н	suffix -al prefix formyl-	но Ш н-с-с-н Н
Ketones	c	suffix* -o ne prefix oxo-	нон O Propanone нсн н н
carboxylic acids	о Ш с—он	suffix -oic acid	H O O Ethanoic acid O H O O H O O O O O O O O O O O O O O
Esters	oo	-yl –oate	H = C + C + C + C + C + C + C + C + C + C

When compounds contain more than one functional group, the order of precedence determines which groups are named with prefix or suffix forms. The highest precedence group takes the suffix (and the lowest number on the carbon chain), with all others taking the prefix form. However, double and triple C-C bonds only take suffix form. **Order of priority highest first:**

Carboxylic acids >aldehydes>ketones>alcohols>alkenes>halogenoalkanes



code	no of carbons
meth	1
eth	2
prop	3
but	4
pent	5
hex	6
hept	7
oct	8
non	9
dec	10

Basic rules for naming functional groups

The functional group is named by a prefix or suffix. e.g. bromoethane, ethanol, propene

•When using a suffix, add in the following way :

If the suffix starts with a vowel- remove the –e from the stem alkane name e.g. Propan-1-ol, butan-1-amine, ethanoic acid, ethanoylchloride, butanamide

If the suffix starts with a consonant or there are two or more of a functional group meaning di, or tri needs to be

used then **do not remove the the –e** from the stem alkane name

e.g. Propanenitrile, ethane-1,2-diol, propanedioic acid, propane-1,2,3-triol, Pentane-2,4-dione.

•The position of the functional group on the carbon chain is given by a number – counting from the end of the molecule that gives the functional group the lowest number. For aldehydes, carboxylic acids & nitriles, the functional group is always on carbon 1.

•We only include numbers, H = C = C = C = H methylpropane to avoid ambiguity. H = C = H

•The functional groups take precedence over branched chains in giving the lowest number

•Where there are two or more of the same groups, *di-, tri- , tetra-, penta- or hexa-* are used. Note the point made above about the addition of 'e' to the stem

•Words are separated by numbers with dashes

numbers are separated by commas

•If there is more than one functional group or side chain, the groups are listed in alphabetical order (ignoring any *di*, *tri*).



CHCl₂

3-methylbut-1-ene is correct and not 2-methylbut-3ene



trichloromethane

CH₂FCCl₂CH₂CH₃

2,2-dichloro-1-fluorobutane.

 $CH_2FCH_2CHBrCH_2CH_3$

3-bromo-1-fluoropentane

Alkenes

The double bond will be between two carbons. Use the lower number of the two to show the position of the double bond

The name for alkenes may include E or Z at start to show the type of stereoisomer

If more than one double bond is present then suffix ends **di**ene or **tri**ene. The stem ends in **a**

The suffix **-en** for alkenes can go in front of other suffixes. The alcohol and carboxylic acid groups have higher priority than the alkene group so take precedence with numbering



2-bromobutane

Halogenoalkanes

Class the halogen as a substituent on the C chain and use the prefix **-fluoro**, **-chloro**, **-bromo**, or **-iodo**. (Give the position number if necessary)



2,3-dichloro-1-fluoro-3-methylpentane

Multiple functional group and side chains are listed in alphabetical order (ignoring any *di*, *tri*).



н н

Br

Br

The alkene group has higher priority than the halogenoalkane group so it takes the lowest number on the carbon chain

н н

Alcohols

These have the ending **-ol** and if necessary the position number for the OH group is added between the name stem and the –ol

If there are two or more -OH groups then di, tri are used.

Add the **'e'** on to the stem name though.

The OH group has a higher priority than the halogenoalkane group and alkene so takes precedence in numbering. The OH is on carbon 1

If the compound has an –OH group in addition to another functional group with a higher priority. The priority group gets the suffix ending and the OH can be named with the prefix **hydroxy**-:





Isomers



Structural isomerism can arise from

- •Chain isomerism
- •Position isomerism
- •Functional group isomerism

EDEXCEL does not split structural isomers into the different categories. They are all classed as structural isomers.





Note: alkene and cyclo alkanes have the same general formula. Hexene and cyclohexane have the same molecular formula but have a different functional group

Stereoisomerism



Cis- but-2-ene

Can also be called

trans- but-2-ene

6A Alkanes

Alkanes are saturated hydrocarbons

General formula alkane C_nH_{2n+2}

Fuels from crude oil

Saturated: Contain single carbon-carbon bonds only

Hydrocarbon is a compound consisting of hydrogen and carbon only

Alkanes are used as fuels They are obtained from the crude oil in the order of fractional distillation, cracking and reformation of crude oil fuel gas (bottled) **Refining crude oil** 20° C 40° C **Fractional Distillation:** naptha (chemicals) Petroleum is a mixture consisting mainly <u>A</u>PP of alkane hydrocarbons Crude kerosene (jet fuel) 180° C oil ւրրը Petroleum fraction: mixture of hydrocarbons with 250° C a similar chain length and boiling point range ዋዋዋ 300° C JUL Furnace ubricating oils Oil is pre-heated

- then passed into column. .
- The fractions condense at different heights •
- The temperature of column decreases upwards
- The separation depends on boiling point.
- Boiling point depends on size of molecules.
- The larger the molecule the larger the London forces
- Similar molecules (size, bp, mass) condense together .
- Small molecules condense at the top at lower temperatures
- and big molecules condense at the bottom at higher temperatures.

Cracking

Cracking: conversion of large hydrocarbons to smaller molecules by breakage of C-C bonds

High Mr alkanes \rightarrow smaller Mr alkanes+ alkenes + (hydrogen)

Economic reasons for catalytic cracking

- The petroleum fractions with shorter C chains (e.g. petrol and • naphtha) are in more demand than larger fractions.
- To make use of excess larger hydrocarbons and to supply demand for shorter ones, longer hydrocarbons are cracked.
- The products of cracking are more useful and valuable than the • starting materials (e.g. ethene used to make poly(ethene) and ethane-1,2-diol, and ethanol) The smaller alkanes are used for motor fuels which burn more efficiently.

Reforming

Turns straight chain alkanes into branched and cyclic alkanes and Aromatic hydrocarbons

Branched and cyclic hydrocarbons burn more cleanly and are used to give fuels a higher octane number.

Used for making motor fuels

fuel oil

bitumen

This is a chemical process

requires high temperatures.

involving the splitting of strong covalent bonds so

This is a physical process

involving the splitting of

weak London forces

between molecules

Alkanes as Fuels

Fuel : releases heat energy when burnt

Complete Combustion

In excess oxygen alkanes will burn with complete combustion

The products of <u>complete</u> combustion are CO_2 and H_2O .

 $C_8H_{18}(g)$ + 12.5 $O_2(g) \rightarrow 8CO_2(g)$ + 9 $H_2O(I)$

Incomplete Combustion

If there is a **limited amount of oxygen** then <u>incomplete</u> combustion occurs, producing CO (which is very toxic) and/or C (producing a sooty flame)

 $\begin{array}{rll} {\rm CH}_4(g) \ + \ {}^3\!\!/_2 \, {\rm O}_2(g) \ \to \ {\rm CO}(g) \ + \ 2 \ {\rm H}_2 {\rm O}(l) \\ {\rm CH}_4(g) \ + \ {\rm O}_2(g) \ \to \ {\rm C}(s) \ + \ 2 \ {\rm H}_2 {\rm O}(l) \end{array}$

Carbon monoxide is an highly toxic but odourless gas. It can cause death if it builds up in a enclosed space due to faulty heating appliances.

Pollution from Combustion

Sulfur containing impurities are found in petroleum fractions which produce SO_2 when they are burned.

$$S+O_2 \rightarrow SO_2$$
 $CH_3SH+3O_2 \rightarrow SO_2+CO_2+2H_2O$

SO₂ is acidic and will dissolve in atmospheric water and can produce acid rain.

Nitrogen Oxides NO_x

Nitrogen oxides form from the reaction between N_2 and O_2 inside the car engine. The **high temperature** and **spark** in the engine provides sufficient energy to break strong N_2 bond

$$N_2 + O_2 \rightarrow 2NO$$
 $N_2 + 2O_2 \rightarrow 2NO_2$

Pollutant	Environmental consequence
Nitrogen oxides (formed when N_2 in the air reacts at the high temperatures and spark in the engine)	NO is toxic and can form smog NO_2 is toxic and acidic and forms acid rain
Carbon monoxide	toxic
Carbon dioxide	Contributes towards global warming
Unburnt hydrocarbons (not all fuel burns in the engine)	Contributes towards formation of smog
Soot/particulates	Global dimming and respiratory problems

Catalytic converters

These remove CO, NO_x and unburned hydrocarbons (e.g. octane, C_8H_{18}) from the exhaust gases, turning them into 'harmless' CO₂, N₂ and H₂O.

 $\begin{array}{l} 2 \; \text{CO} \; + \; 2 \; \text{NO} \; \rightarrow \; 2 \; \text{CO}_2 \; + \; \text{N}_2 \\ \\ \text{C}_8 \text{H}_{18} \; + \; 25 \; \text{NO} \; \rightarrow \; 8 \; \text{CO}_2 \; + \; 12 \frac{1}{2} \; \text{N}_2 \; + \; 9 \; \text{H}_2 \text{O} \end{array}$

Converters have a ceramic honeycomb coated with a thin layer of catalyst metals p**latinum, palladium, rhodium** – to give a large surface area.

Incomplete combustion produces less energy per mole than complete combustion.

Carbon (soot)/particulates can cause global dimming- reflection of the sun's light

CO is toxic to humans as CO can from a strong bond with haemoglobin in red blood cells. This is a stronger bond than that made with oxygen and so it prevents the oxygen attaching to the haemoglobin.

Coal is high in sulfur content, and

large amounts of sulfur oxides are emitted from power stations.

Biofuels

Most fossil fuels come from crude oil, which is a nonrenewable resource. Fossil fuel reserves will eventually run out

Alternative fuels have been developed from renewable resources. Alcohols and biodiesel, which can both be made from plants, are two examples of renewable plantbased fuels

Advantages of using Biofuels

Reduction of use of fossil fuels which are finite resources biofuels are renewable

Use of biodiesel is more carbon-neutral

Allows fossil fuels to be used as a feedstock for organic compounds

No risk of large scale pollution from exploitation of fossil fuels

Alcohols such as ethanol can be produced from the fermentation of sugars from plants.

Biodiesel is produced by reacting vegetable oils with a mixture of alkali and methanol

Disadvantages of Biofuels

Less food crops may be grown Land not used to grow food crops Rain forests have to be cut down to provide land Shortage of fertile soils

Introduction to Mechanisms

To understand how the reaction proceeds we must first understand how bonds are broken in organic mechanisms There are two ways to break a covalent bond:

1.HOMOLYTIC FISSION:

each atom gets one electron from the covalent bond

When a bond breaks by homolytic fission it forms two **Free Radicals**. **Free Radicals** do not have a charge and are represented by a

DEFINITION

A Free Radical is a reactive species which possess an **unpaired electron**

2. HETEROLYTIC FISSION: (one atom gets both electrons)

Heterolytic fission produces IONS

Most organic reactions occur via heterolytic fission, producing ions

The Mechanism:

To understand a reaction fully we must look in detail at how it proceeds step by step. This is called its **mechanism**

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

We use curly arrows in mechanisms to show the movement of an electron pair showing either breaking or formation of a covalent bond;

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

Free Radical Substitution Reactions of Alkanes

substitution

Reaction of alkanes with bromine / chlorine in UV light

In the presence of **UV light** alkanes react with chlorine to form a mixture of products with the halogens substituting hydrogen atoms.

In general, alkanes do not react with many reagents. This is because the C-C bond and the C-H bond are relatively strong.

It proceeds via a series of steps: Step one: Initiation Step two: Propagation Step three: Termination

 $CH_3CH_2CH_3 + Br^* \rightarrow HBr + CH_3CH^* CH_3$ $CH_3CH^*CH_3 + Br_2 \rightarrow CH_3CH_2BrCH_3 + Br^*$

If the question asks for the halogen to be substituted onto a middle carbon in the chain, it is important to put the free radical '**dot**' on the correct carbon in the propagation stages.

Further substitution

Excess Cl₂ present will promote further substitution and could produce CH₂Cl₂, CHCl₃ and CCl₄

 $\begin{array}{l} \mathsf{CH}_{3}\mathsf{CI} + \mathsf{CI}_{2} \rightarrow \mathsf{CH}_{2}\mathsf{CI}_{2} + \mathsf{HCI} \\ \mathsf{CH}_{2}\mathsf{CI}_{2} + \mathsf{CI}_{2} \rightarrow \mathsf{CHCI}_{3} + \mathsf{HCI} \\ \mathsf{CHCI}_{3} + \mathsf{CI}_{2} \rightarrow \mathsf{CCI}_{4} + \mathsf{HCI} \end{array}$

These reactions could occur

Example propagation steps that would lead to further substitution $CH_3CI + CI \rightarrow HCI + CH_2CI$ $CH_2CI + CI_2 \rightarrow CH_2CI_2 + CI$

Overall reaction equations

You should be able to write overall reaction equations for various reactions

Example 1. Write the overall reaction equation for the formation of CCI_4 from $CH_4 + CI_2$

 $CH_4 + 4 Cl_2 \rightarrow CCl_4 + 4 HCl$

Example 2. Write the overall reaction equation for the formation of $CFCI_3$ from $CH_3F + CI_2$

$$CH_3F + 3 Cl_2 \rightarrow CFCl_3 + 3 HCl$$

Note HCl is always the side product – never H_2

6C Alkenes

Addition Reactions of Alkenes

Addition reaction: a reaction where two molecules react together to produce one

1. Reaction of Alkenes with Hydrogen

Change in functional group: alkene → alkane Reagent: hydrogen Conditions: Nickel Catalyst Type of reaction: Addition/Reduction

Electrophilic Addition: Reactions of Alkenes

A bond is weaker than a bond so less energy is needed to break bond The bonds in alkenes are areas with high electron density. This is more accessible to electrophilic attack by electrophiles. Alkenes undergo addition reactions.

Definition Electrophile: an **electron pair acceptor**

2. Reaction of Alkenes with bromine/chlorine

ELECTROPHILIC (Br *).

3. Reaction of Hydrogen Bromide with alkenes

HBr is a polar molecule because Br is more electronegative than H. The H⁺ is attracted to the electron-rich pi bond.

If the alkene is unsymmetrical, addition of hydrogen bromide can lead to two isomeric products.

But-1-ene will form a mixture of 1-bromobutane and 2-bromobutane on reaction with hydrogen bromide

stable and a 50/50 split will be achieved)

4. Reaction of Potassium Manganate(VII) with Alkenes

Change in functional group: alkene → diol
Reagent: KMnO₄ in an acidified solution
Conditions: Room temperature
Type of reaction: Oxidation
Observation: purple colour of MnO₄⁻ ion will decolourise to colourless

This reaction with its colour change can be used as a test for the alkene functional group. It would not change colour with alkanes

5. Reaction of Bromine Water with Alkenes

Reagent: Bromine dissolved in water
Conditions: Room temperature
Type of reaction: Addition
Observation: Orange colour of bromine water will decolourise to colourless

This reaction with its colour change is used as a test for the alkene functional group.

Hydration of alkenes to form alcohols

Industrially alkenes are converted to alcohols in one step rather than the two in the above sulfuric acid reaction. They are reacted with water in the presence of an acid catalyst.

This reaction can be called **hydration:** a reaction where water is **added** to a molecule

$$\mathsf{CH}_2^{=}\mathsf{CH}_{2\ (g)}+\mathsf{H}_2\mathsf{O}_{\ (g)}\ \textbf{\rightarrow}\mathsf{CH}_3\mathsf{CH}_2\mathsf{OH}_{\ (I)}$$

Essential Conditions

High temperature 300 to 600°C

High pressure 70 atm

Catalyst of concentrated H₃PO₄

The high pressures needed mean this cannot be done in the laboratory. It is preferred industrially, however, as there are no waste products and so has a high atom economy. It would also mean separation of products is easier (and cheaper) to carry out. See equilibrium chapter for more on the industrial conditions for this reaction.

Addition Polymers

Addition polymers are formed from alkenes

This is called addition polymerisation

<u>Poly(ethene):</u> is used to make plastics bags, buckets, bottles. It is a flexible, easily moulded, waterproof, chemical proof, and low density plastic. Poly(alkenes) like alkanes are unreactive due to the strong C-C and C-H bonds.

Add the **n's** if writing an equation showing the reaction where 'n' monomers become 'n' repeating units

<u>Poly(propene)</u> is a stiffer polymer, used in utensils and containers and fibres in rope and carpets.

4

Methods of disposal of waste polymers

Incineration

Rubbish is burnt and energy produced is used to generate electricity.

Some toxins can be released on incineration. (e.g. Combustion of halogenated plastics (ie PVC) can lead to the formation of toxic, acidic waste products such as HCl.) Modern incinerators can burn more efficiently and most toxins and pollutants can be removed. Greenhouse gases will still be emitted though.

Volume of rubbish is greatly reduced.

Recycling

Saves raw materials- nearly all polymers are formed from compounds sourced/produced from crude oil. Saves precious resources.

Polymers need collecting/ sorting- expensive process in terms of energy and manpower.

Polymers can only be recycled into the same type – so careful separation needs to be done. Thermoplastic polymers can be melted down and reshaped.

feedstock for cracking

Polymers can be cracked into small molecules which can be used to make other chemicals and new polymers- Saves raw materials-

Chemists have designed ways to remove toxic waste products like HCl before they are emitted into the atmosphere. The waste gases from the incinerator are scrubbed/reacted with a base or carbonate. The base reacts with the acidic HCl gas, neutralising it (eg CaO + 2HCl \rightarrow CaCl₂ + H₂O)

Chemists have also develop biodegradable and compostable polymers. Biodegradable polymers can be made from substances such as maize and starch

6D Halogenoalkanes

Naming Halogenoalkanes

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

I-bromopropane

2-chloro-2-methylbutane

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

halogen

Reactions of Halogenoalkanes

Halogenoalkanes undergo either substitution or elimination reactions

Tertiary halogenoalkane Three carbons attached to the carbon atom adjoining the

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 bre	ak 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

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 and 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 haloalkane

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 OH^- is a stronger nucleophile than water as it has a full negative charge and so is more strongly attracted to the C ⁺

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

SN₂ nucleophilic substitution mechanism for halogenoalkanes

This mechanism occurs with primary halogenoalkanes

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

Primary halogenoalkanes do not do the SN₁ mechanism because they would only form an unstable primary carbocation.

Nucleophilic substitution with ammonia

formed leading to a lower yield of the amine. Using excess ammonia helps minimise this.

Elimination reaction of halogenoalkanes

Elimination with alcoholic hydroxide ions

Change in functional group: halogenoalkane \rightarrow
alkeneReagents:Potassium (or sodium) hydroxideConditions:In <u>ethanol</u>; HeatMechanism:EliminationRole of reagent:Base, OH⁻

Note the importance of the solvent to the type of reaction here.

Aqueous: substitution

Alcoholic: elimination

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

Halogenoalkanes have been used as refrigerants, fire retardants, pesticides and aerosol propellants.

Chloroalkanes and chlorofluoroalkanes can be used as solvents.

 CH_3CCl_3 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 ozone layer.

Some halogenoalkanes have low flammability.

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

+ KBr + H₂O

6E Alcohols

General formula alcohols C_nH_{2n+1}OH

Bond angles in alcohols

All the H-C-H bonds and C-C-O are 109.5° (tetrahedral shape), because there are 4 bond pairs of electrons repelling to a position of minimum repulsion.

The H-O- C bond is 104.5° (bent line shape), because there are 2 bond pairs of electrons and 2 lone pairs repelling to a position of minimum repulsion. Lone pairs repel more than bond pairs so the bond angle is reduced.

Different types of alcohols

Primary alcohols are alcohols where 1 carbon is attached to the carbon adjoining the oxygen

Secondary alcohols are alcohols where 2 carbon are attached to the carbon adjoining the oxygen

Tertiary alcohols are alcohols where 3 carbon are attached to the carbon adjoining the oxygen

1. Combustion of Alcohols

 $CH_3CH_2OH + 3O_2 \rightarrow 2CO_2 + 3H_2O$

Alcohols combust with a clean flame

2. Reaction of Alcohols with Sodium

Sodium reacts with alcohols

$$2CH_3CH_2OH + 2Na \rightarrow 2CH_3CH_2O^-Na^+ + H_2$$

This reaction can be used as a test for alcohols

3. Substitution reactions of Alcohols to form Halogenoalkanes

Various halogenating compounds can be used to substitute the –OH group for a halogen

Reaction with phosphorous (V) halide

Observations:

- effervescence,
- the mixture gets hot,
- sodium dissolves,
- a white solid is produced.

PCI₅ / PCI₃ / conc HCI / SOCI₂ / mixture of NaCI +

 H_2SO_4 can all be uses for substituting a CI

Full Oxidation of Primary Alcohols

Distinguishing between Aldehydes and Ketones

The fact that aldehydes can be further oxidised to carboxylic acids whereas ketones cannot be further oxidised is the chemical basis for tests that are commonly used to distinguish between aldehydes and ketones

Fehling's (Benedict's) solution

Reagent: Fehling's solution containing blue Cu ²⁺ ions.
 Conditions: heat gently
 Reaction: aldehydes only are oxidised by Fehling's solution into a carboxylic acid and the copper (II) ions are reduced to copper(I) oxide
 Observation: Aldehydes :Blue Cu ²⁺ ions in solution change to a red precipitate of Cu₂O. Ketones do not react

 $CH_3CHO + 2Cu^{2+} + 2H_2O \rightarrow CH_3COOH + Cu_2O + 4H^+$

The presence of a carboxylic acid can be tested by addition of sodium carbonate. It will fizz and produce carbon dioxide

Reaction of Alcohols with Dehydrating Agents

Producing alkenes from alcohols provides a possible route to polymers without using monomers derived from oil

Organic techniques

Distillation

In general used as separation technique to separate an organic product from its reacting mixture. Need to collect the distillate of the approximate boiling point range of the desired liquid.

Classic AS reaction using distillation Reaction: primary alcohol \rightarrow aldehyde Reagent: potassium dichromate (VI) solution and dilute sulfuric acid. Conditions: use a limited amount of dichromate and warm gently and distil out the aldehyde as it forms [This prevents further oxidation to the carboxylic acid] CH₃CH₂CH₂OH + [O] \rightarrow CH₃CH₂CHO + H₂O Observation Orange dichromate solution changes to green colour of Cr³⁺ ions

Reflux

Reflux is used when heating organic reaction mixtures for long periods. The condenser prevents organic vapours from escaping by condensing them back to liquids.

Never seal the end of the condenser as the build up of gas pressure could cause the apparatus to explode. This is true of any apparatus where volatile liquids are heated

Classic AS reaction using reflux Reaction: primary alcohol → carboxylic acid Reagent: potassium dichromate(VI) solution and dilute sulfuric acid Conditions: use an excess of dichromate, and heat under reflux: (distil off product after the reaction has finished using distillation set up) CH₃CH₂CH₂OH + 2[O] → CH₃CH₂CO₂H + H₂O Observation Orange dichromate solution changes to green colour of Cr³⁺ ions

Anti-bumping granules are added to the flask in both distillation and reflux to prevent vigorous, uneven boiling by **making small bubbles** form instead of large bubbles

It's important to be able to draw and label this apparatus accurately.

- Don't draw lines between flask and condenser.
- Don't have top of condenser sealed
- Condenser must have outer tube for water that is sealed at top and bottom
- Condenser must have two openings for water in and out that are open

Electric heaters are often used to heat organic chemicals. This is because organic chemicals are normally highly flammable and could set on fire with a naked flame.

Purifying an organic liquid

• Put the distillate of impure product into a separating funnel

- wash product by adding either
 - sodium hydrogencarbonate solution, shaking and releasing the pressure from CO₂ produced.
 - Saturated sodium chloride solution

•Allow the layers to separate in the funnel, and then run and discard the aqueous layer.

•Run the organic layer into a clean, dry conical flask and add three spatula loads of drying agent (anhydrous sodium sulfate) to dry the organic liquid.

• Carefully decant the liquid into the distillation flask

Mix organic solvent and oil-water mixture in a separating funnel then separate the oil layer. Distil to separate oil from organic solvent Add anhydrous CaCl₂ to clove oil to dry oil

•Distil to collect pure product

Solvent extraction

Sodium hydrogencarbonate will neutralise any remaining reactant acid.

Sodium chloride will help separate the organic layer from the aqueous layer

The drying agent should •be insoluble in the organic liquid • not react with the organic liquid

Separating funnel

Measuring boiling point

Decant to remove CaCl₂

Purity of liquid can be determined by measuring a boiling point. This can be done in a distillation set up or by simply boiling a tube of the sample in an heating oil bath.

Pressure should be noted as changing pressure can change the boiling point of a liquid

Measuring boiling point is not the most accurate method of identifying a substance as several substances may have the same boiling point.

To get a correct measure of boiling point the thermometer should be above the level of the surface of the boiling liquid and be measuring the temperature of the saturated vapour.