ALKENES

Structure

- form a homologous series of general formula C_nH_{2n} non cyclic alkenes only
- contain a carbon-carbon double bond somewhere in their structure
- unsaturated hydrocarbons can still have atoms added to their formula

Naming

- select the longest chain of C atoms containing the double bond; end in ENE
- number the chain starting from the end nearer the double bond;
 use a number to indicate the FIRST carbon in the double bond.
- prefix with substituents. Side chain positions are based on the number allocated to the first carbon in the double bond.
- if cis-trans isomerism exists, prefix with cis or trans

e.g.
$$CH_3 - CH = CH - CH_2 - CH(CH_3) - CH_3$$
 is called 5-methylhex-2-ene

ISOMERISM Two types of isomerism are found in alkenes.

Structural

due to • different positions for the double bond

but-1-ene and but-2-ene

· branching.

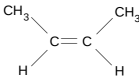
but-1-ene and 2-methylpropene

E/Z

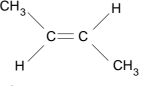
- due to restricted rotation of the carbon-carbon double bond
- single bonds can rotate so the groups around them are not 'frozen' in position
- double bonds can't rotate freely, thus "freezing" groups on either side of C=C
- not all alkenes exhibit this type of isomerism
- if there are two H's and two non-hydrogen groups attached to each carbon it is called *cis-trans* isomerism

but-1-ene

- Z higher priority groups on the same side of the C=C bond.
- E higher priority groups on opposite sides.



(Z) cis but-2-ene



(E) trans but-2-ene

$$CH_3$$
 $C=C$

2-methylpropene

Priority of atoms / groups

$$C_3H_7 > C_2H_5 > CH_3 > H$$

STRUCTURAL ISOMERS E/Z or CIS-TRANS ISOMERS but-1-ene, but-2-ene and 2-methylpropene cis (**Z**) and trans (**E**) but-2-ene

Q.1

Draw the structures of all the structural isomers of alkenes of formula C_6H_{12} . Name them and indicate those which exhibit E/Z (cis-trans) isomerism.

PHYSICAL PROPERTIES

Boiling point

- increases with molecular mass/size increased induced dipole-dipole attractions
- for isomers the greater the degree of branching, the lower the boiling point
- the lower members of the series are gases at rtp cyclohexene is a liquid.

Solubility

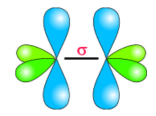
• non polar so they are insoluble in water but soluble in organic solvents.

CHEMICAL PROPERTIES

Theory

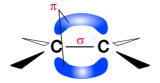
- spacial arrangement around the C=C is planar
- the bond angles are 120°

Three of the four electrons of carbon sp² hybridised and the remaining electron exists in a 2p orbital at right-angles to the three sp² orbitals.



[For more information, see notes on hybridisation]

The 2p orbitals overlap to form a new pi (π) orbital; it exists above and below the plane of the carbon-carbon bond.



- the main reaction of alkenes is **ELECTROPHILIC ADDITION**
- electrophiles are electron pair acceptors
- electrophiles are attracted to the electron rich carbon-carbon double bond.

AQA

ELECTROPHILIC ADDITION REACTIONS OF ALKENES

Addition of HBr

Reagent Hydrogen bromide; it is electrophilic because the H is slightly positive

Condition Room temperature.

Equation $C_2H_4(g)$ + HBr(g) -----> $C_2H_5Br(l)$ bromoethane

Mechanism

- As the HBr nears the alkene, one of the carbon-carbon bonds breaks and the pair of electrons attaches to the slightly positive H end of H-Br. This is an example of **HETEROLYTIC FISSION.** The HBr bond breaks to form a bromide ion. A carbocation (positively charged carbon species) is formed.
- **Step 2** The bromide ion behaves as a nucleophile and attacks the carbocation.

Overall there has been addition of HBr across the double bond.

Addition of H₂SO₄

SO₄ Reagent Concentrated sulphuric acid (85%).

Conditions 0°C

Equation $C_2H_4(g) + H_2SO_4(conc) \longrightarrow C_2H_5OSO_2OH(aq)$

ethyl hydrogensulphate

Mechanism

Importance Hydrolysis - the product is made into ethanol by boiling with water.

 $C_2H_5OSO_2OH(aq) + H_2O(l) \longrightarrow H_2SO_4(aq) + C_2H_5OH(g)$ ethanol

Industrially Phosphoric acid (H₃PO₄) and steam are used Ethanol can also be made by FERMENTATION

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Addition of Br₂

Reagent Bromine. (Neat liquid or dissolved in tetrachloromethane, CCI₄)

Condition Room temperature. No catalyst or UV light required!

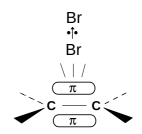
Equation $C_2H_4(g) + Br_2(g) \longrightarrow CH_2BrCH_2Br(l)$ 1,2 - dibromoethane

Mechanism

It is surprising that bromine should act as an electrophile as it is non-polar.

Explanation ...

 as a bromine molecule approaches an alkene, electrons in the pi bond of the alkene repel the electron pair in the bromine-bromine bond thus inducing a dipole.



Importance

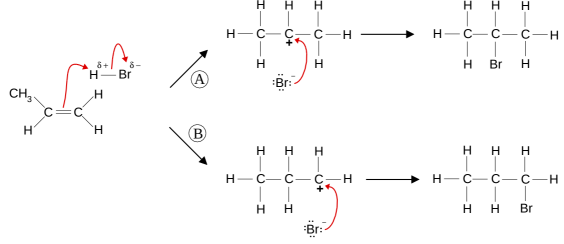
The addition of bromine dissolved in tetrachloromethane (CCl₄) or water (known as **bromine water**) is used as a **test for unsaturation**. If the reddish-brown colour is removed from the bromine solution, the substance possesses a double C=C bond.

Electrophilic addition to propene

Problem

- addition of HBr to propene gives two isomeric brominated compounds
- HBr is unsymmetrical and can add in two ways
- products are not formed to the same extent
- problem doesn't arise in ethene because it is symmetrical.

Mechanism



Markownikoff's Rule

Background The Russian scientist, Markownikoff, investigated the products of the addition of hydrogen halides to alkenes. He found that, when two products were formed, one was formed in a larger quantity. His original rule was based only on this reaction. The up-to-date version of Markownikoff's Rule uses carbocation stability as a criterion for predicting the products.

> It states that "In electrophilic addition to alkenes the major product is formed via the more stable carbocation (carbonium ion)."

> In the above reaction, path A involves a 2° carbocation, path B a 1° carbocation. As the 2° ion is more stable, the major product (i.e. 2-bromopropane) is formed via that route.

Carbocations / Carbonium ions

least stable methyl < most stable

Theory

Build up of charge in one place leads to instability. However, if it can be spread around or neutralised in some way, the stability is increased. Alkyl groups are electron releasing and can "push" electrons towards the carbocations thus reducing the charge density.

Q.2Draw the structures of the product(s) formed when HBr reacts with each of the isomers of C_4H_8 . If two products are formed, state which is the major product.

Outline the mechanism for the reaction between propene and conc. H_2SO_4 . Which alcohols are produced by hydrolysing the products?

Bond enthalpies (kJ mol-1)

H-Br 366

O-H 463

OTHER ADDITION REACTIONS

Direct Hydration

Reagent steam (330°C)

Conditions High Pressure (6MPa)

Catalyst - phosphoric acid

Product alcohol

Equation $C_2H_4(g) + H_2O(g) \longrightarrow C_2H_5OH(g)$ ethanol

Use ethanol manufacture

Note

O-H bonds are more polar than H-Br bonds yet the addition of H_2O requires a catalyst. An explanation is that O-H bonds are stronger so more energy is needed to break them.

Hydrogenation

Reagent hydrogen

Conditions nickel catalyst - finely divided

Product alkanes

Equation $C_2H_4(g) + H_2(g) \longrightarrow C_2H_6(g)$ ethane

Use margarine manufacture

POLYMERISATION

Process

- during polymerisation, an alkene undergoes an addition reaction with itself
- all the atoms in the original alkenes are used to form the polymer
- long hydrocarbon chains are formed

• the equation shows the original monomer and the repeating unit in the polymer

POLYMERS

Preparation Many are prepared by a free radical process involving high pressure, high temperature and a catalyst. The catalyst is usually a substance (an organic peroxide) which readily breaks up to form radicals which initiate a chain reaction.

> Another famous type of catalyst is a Ziegler-Natta catalyst (named after the scientists who developed it). Such catalysts are based on the compound TiCl₄.

Properties

Physical Can be varied by changing the reaction conditions (pressure, temperature etc).

Chemical Chemical properties are based on the functional groups within their structure.

> poly(ethene) is typical; it is fairly inert as it is basically a very large alkane. e.g. This means it is resistant to chemical attack and non-biodegradable.

Problems

Although polymers derived from alkenes are invaluable to modern society, their disposal creates widespread problems.

- they are unreactive to most chemicals and bacteria (non-biodegradable)
- if they are just discarded they add to the landfill problem
- recycling
- high cost of collection and re-processing
 - the different types of polymer have to be separated

burn waste • saves on landfill sites and produces **energy**

 toxic fumes (HCI) can be removed by 'scrubbing' from burning chlorinated polymers such as poly(chloroethene)

feedstock

- use the waste for the production of useful organic compounds
 - new technology can convert waste into hydrocarbons
 - hydrocarbons can then be turned back into polymers

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 ${\it Q.4}$ Complete the details showing the formation and use of different polymers.

Polymer Formula of monomer Formula of polymer Use(s)

poly(ethene)

n CH₂=CH₂

---->

 $-(CH_2-CH_2)_n-$

poly(propene)

poly(chloroethene)

poly(phenylethene)

poly(tetrafluoroethene)

poly(ethenyl ethanoate)

Q.5 Identify the monomer units in the following addition polymers.