## **Organic: Basic Concepts**

Hydrocarbon is a compound consisting of hydrogen and carbon only

Basic definitions to know

Saturated: Contain single carbon-carbon bonds only

Unsaturated: Contains a C=C double bond

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

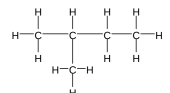
Empirical formula: shows the simplest whole number ratio of atoms of each element in the compound

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:  $CH_3CH_2CH_3$  or  $CH_3(CH_2)_2CH_3$ ,

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

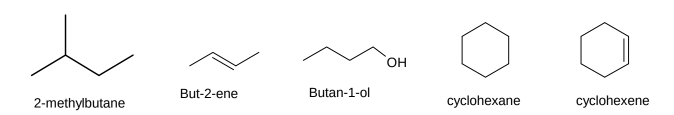
## **Drawing Displayed formulae**



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.



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

# **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 CH<sub>2</sub> from the last.
- same chemical properties.

homologous series	functional group	prefix / suffix (* = usual use)	example
Alkane		-ane	CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> CH <sub>3</sub> Butane
Alkenes		Suffix* -ene	C=C H propene
Alcohols	сон	suffix* -ol prefix hydroxy-	н н н н Propan-1-ol н—с—с—с—о—н ОН
Halogenoalkanes	C—halogen	Prefix* chloro- bromo- iodo-	H H H H H 1-chloropropane 1-chloropropane CI
Aldehydes	о    	suffix -al prefix formyl-	H O H O H O O ethanal
Ketones	c	suffix* -one prefix <b>oxo-</b>	H O H O Propanone
carboxylic acids	о    сон	suffix* -oic acid	H O O Ethanoic acid
Esters	co	-yl –oate	H O H O methylethanoate

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

## General rules for naming carbon chains

- •Count the longest carbon chain and name appropriately
- •Find any branched chains and count how many carbons they contain
- Add the appropriate prefix for each branch chain

Eg 
$$-CH_3$$
 methyl or  $-C_2H_5$  ethyl  $-C_3H_7$  propyl

3,5-dimethylheptane

carbons
1
2
3
4
5
6
7
8
9
10

no of

## 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.

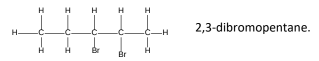
•We only include numbers, however, if they are needed to avoid ambiguity.

CHCl<sub>3</sub> trichloromethane

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

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

•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

CH<sub>2</sub>FCH<sub>2</sub>CHBrCH<sub>2</sub>CH<sub>3</sub>

3-bromo-1-fluoropentane

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

## 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



Double bond between C<sup>2</sup> and C<sup>3</sup> so But-2-ene

Z-but-2-ene

Penta-1,3-diene

CH<sub>2</sub>OHCHBrCH=CH<sub>2</sub>

2-bromobut-3-**en**-1-ol



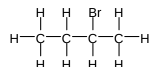
E-but-2-enoic acid

## Halogenoalkanes

Class the halogen as a substituent on the C chain and use the suffix **-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*).



2-bromobutane

5,5-dibromo-4-iodo-3-methylpent-1-ene

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

## 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  $\bf \'e'$  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**-:

Butan-2-ol

HO-CH<sub>2</sub>CH<sub>2</sub>-OH

Ethane-1,2-diol

$$\begin{array}{c|cccc} \mathsf{H}_2\mathsf{C} & \mathsf{CH} - \mathsf{CH}_2 \\ \mathsf{OH} & \mathsf{OH} & \mathsf{OH} \end{array}$$

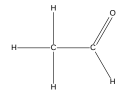
propan**e**-1,2,3-triol

E-3,6-dichlorohex-4-en-1-ol

2-hydroxypropanoic acid

## **Aldehydes**

An aldehyde's name ends in **–al**It always has the C=O bond on the first carbon of the chain so it does not need an extra number. It is by default number one on the chain.



Ethanal

If two aldehyde groups then **di** is put before **–al** and an **e** is added to the stem.

pentanedial

Aldehydes have a higher priority than alcohol so the –OH group uses the hydoxy prefix.

4-hydroxybutanal

#### **Ketones**

Ketones end in -one

When ketones have 5C's or more in a chain then it needs a number to show the position of the double bond. E.g. pentan-2-one

If two ketone groups then **di** is put before – **one** and an **e** is added to the stem.

Propan**one** 

Pentan**e**-2,4-dione

## Carboxylic acids

These have the ending **-oic acid** but no number is necessary for the acid group as it must always be at the end of the chain. The numbering always starts from the carboxylic acid end.

If there are carboxylic acid groups on both ends of the chain then it is called a - dioic acid

 ${\sf Ethan} \textbf{edioic acid}$ 

Note the **e** in this name

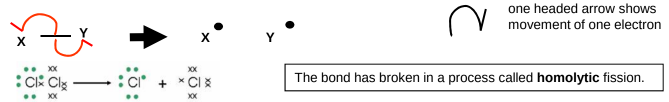
5-hydroxy-4-methylpentanoic acid

## **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



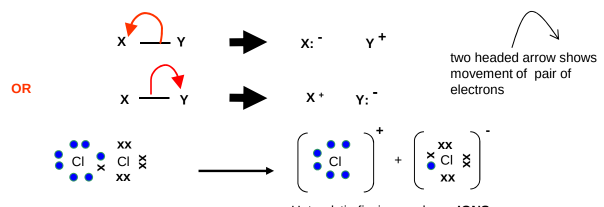
When a bond breaks by homolytic fission it forms two Free Radicals.

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

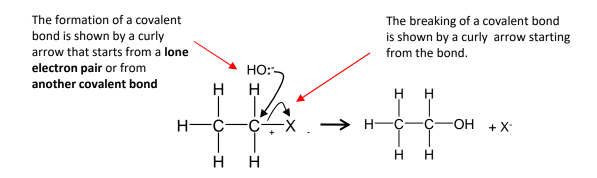
## The Mechanism:

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

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

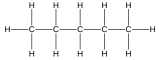


Structural isomers: same molecular formula different structures (or structural formulae)

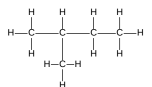
Structural isomerism can arise from

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

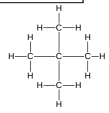
**Chain isomers**: Compounds with the same molecular formula but different structures of the carbon skeleton



pentane

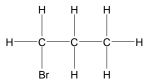


2-methylbutane

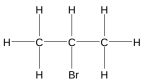


2,2-dimethylpropane

**position isomers**: Compounds with the same molecular formula but different structures due to different positions of the same functional group on the same carbon skeleton



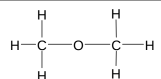
1-bromopropane H



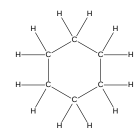
2-bromopropane

**Functional group isomers**: Compounds with the same molecular formula but with atoms arranged to give different functional groups

ethanol: an alcohol



Methoxymethane: an ether



Cyclohexane- cyclo alkane

CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CH=CH<sub>2</sub>

hexene- alkene

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

Stereoisomers have the same structural formulae but have a different spatial arrangement of atoms.

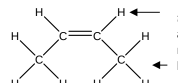
Alkenes can exhibit a type of isomerism called **E-Z stereoisomerism** 

E-Z isomers exist due to restricted rotation about the C=C bond

Single carbon-carbon covalent bonds can easily rotate

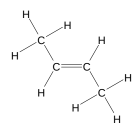
E-Z stereoisomers arise when:

- (a) There is **restricted rotation** around the C=C double bond.
- (b) There are two different groups/atoms attached both ends of the double bond.



two different groups attached either end of the restricted double bondleads to EZ isomers

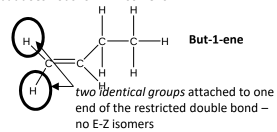
Z- but-2-ene

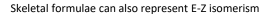


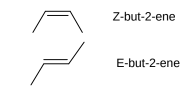
These are two isomers as the lack of rotation around the double bonds means one cannot be switched to the other.

E-but-2-ene

But-1-ene is a structural isomer of But-2-ene but does not show E-Z isomerism.

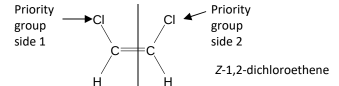






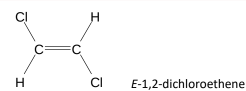
#### Naming E-Z stereoisomers

First determine the priority groups on both sides of the double bond



If the priority atom is on the same side of the double bond it is labelled Z from the german zusammen (The Zame Zide!)

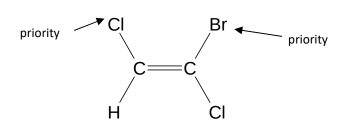
**Priority Group**: The atom with the bigger atomic number is classed as the priority atom



If the priority atom is on the opposite side of the double bond it is labelled E from the german entgegen (The Epposite side!)

## Cahn-Ingold-Prelog (CIP) priority rules.

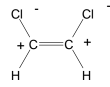
- 1. Compare the atomic number of the atoms directly attached to each side of the double bond; the atom of higher atomic number is given priority.
- 2. If the atoms are the same, consider the atoms at distance 2 from the double bond. Make a list of each atom bonded to the one directly attached to the double bond. Arrange list in order of decreasing atomic number. Compare the lists atom by atom; at the earliest difference, the group containing the atom of higher atomic number is given priority



$$C = C$$
 $C + C$ 
 $C = C$ 
 $C + C$ 
 $C +$ 

## The effect of EZ stereoisomerism on physical properties

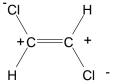
E-Z stereoisomers can have differing melting and boiling points.



*Z*-1,2-dichloroethene Boiling point =60°C

This molecule is **polar**. The polar C-Cl bonds are on the same side of the molecule. One side of the molecule is slightly negative.

The intermolecular forces are both van der waals and permanent dipole-dipole attractions.



E-1,2-dichloroethene Boiling point =48°C

This molecule is non- **polar**. The polar C-Cl bonds are on opposite sides of the molecule. The dipoles cancel out.

The intermolecular forces are is only van der waals so lower boiling point.