

# **WJEC Chemistry AS-Level**

## 2.8: Instrumental Analysis

**Detailed Notes** 

**English Specification** 

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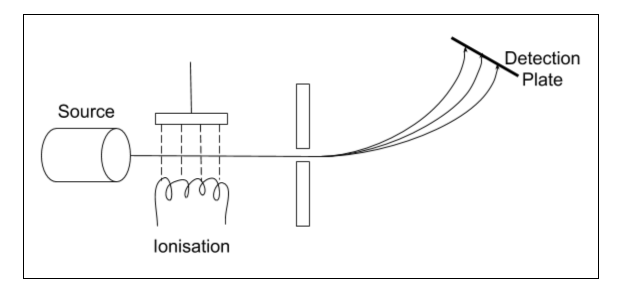
## **Mass Spectrometry**

This is an **analytical technique** used to identify different **isotopes** and find the overall relative **atomic mass** of an element.

## Time of Flight (TOF) Mass Spectrometry

This form of mass spectrometry records the **time** it takes for ions of each isotope to reach a detector. Using this, **spectra** can be produced showing each isotope present. The process of TOF mass spectrometry is as follows:

- Ionisation A sample of an element is vapourised and injected into the mass spectrometer where a high voltage is passed over the chamber. This causes electrons to be removed from the atoms (they are ionised) leaving +1 charged ions in the chamber.
- 2. **Acceleration** The positively charged ions are **accelerated** towards a negatively charged **plate**.
- 3. **Ion Drift** The ions are **deflected** by a **magnetic field** into a **curved path**. The radius of their path is dependent on the charge and mass of the ion.



- 4. **Detection** When the positive ions hit the detection plate, they gain an electron producing a flow of charge. The greater the abundance, the greater the current produced.
- 5. **Analysis** The **current** values are then used in combination with the **flight times** to produce a **spectra** print-out with the **relative abundance** of each isotope displayed.



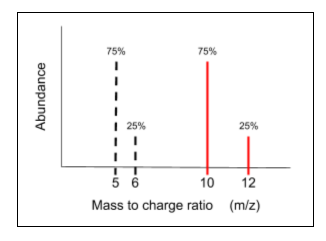






During the ionisation process, a **2+ charged ion** may be produced. This means it will be affected more by the magnetic field producing a curved path of **smaller radius**. As a result, its mass to charge ratio (m/z) is **halved** and this can be seen on spectra as a trace at half the expected m/z value.

Example:



Using this spectra, the relative atomic mass can be calculated:

$$Ar = \frac{m/z \times abundance}{Total \ abundance}$$

Example:

$$Ar = (10x75) + (12x25) = 10.5$$

$$(75 + 25)$$

## **High Resolution Mass Spectrometry**

This is a much more **sensitive** form of mass spectrometry which allows the Mr of a substance to be determined to **several decimal places**. Precise atomic masses are given which can then be used to calculate the molecular formula of the compound being tested, using the same method as above.

## **Infrared Spectroscopy**

This analytical technique uses **infrared** (**IR**) **radiation** to determine the **functional groups** present in organic compounds. The IR radiation is passed through a sample where the different types of bonds **absorb radiation** in different amounts. These varying amounts of absorbance are measured and recorded, allowing certain bonds, and therefore functional groups, to be





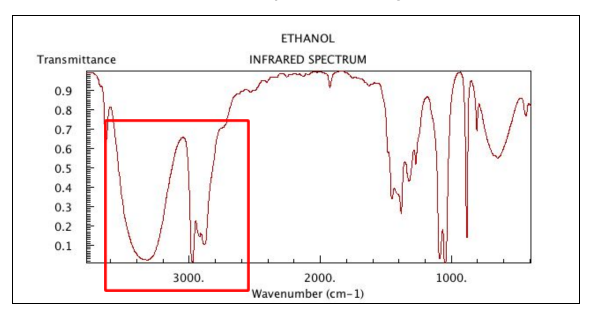






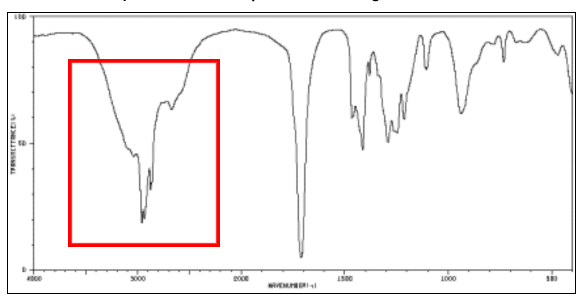
identified. A **spectrum** is produced from the measurements which has **characteristic curves** for the different functional groups:

#### -OH Alcohol Group - characteristic peak is in the range 3230 - 3550 cm<sup>-1</sup>



(https://commons.wikimedia.org/wiki/File:EtOH\_Spectra.png)
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## -OH Acid Group - characteristic peak is in the range 2500 - 3000 cm<sup>-1</sup>



(Modified from https://chem.libretexts.org/Bookshelves/Physical and Theoretical Chemistry Textbook Maps
/Supplemental Modules (Physical and Theoretical Chemistry)/Spectroscopy/Vibrational Spectroscopy/Infrared Spectro
scopy/Interpreting Infrared Spectra/IR8. More Complicated IR Spectra)
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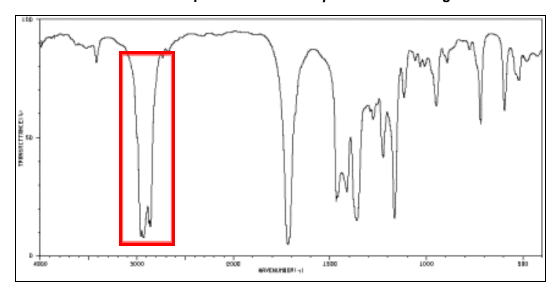








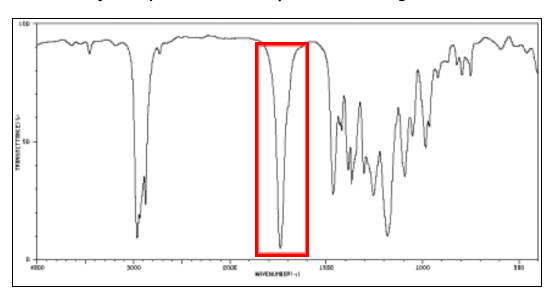
## C=C Unsaturated Group - characteristic peak is in the range 1620 - 1680 cm<sup>-1</sup>



(Modified from https://employees.csbsju.edu/cschaller/Principles%20Chem/structure%20determination/IRHydrocarbon.htm)

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## C=O Carbonyl Group - characteristic peak is in the range 1680 - 1750 cm<sup>-1</sup>



(Modified from https://employees.csbsju.edu/cschaller/Principles%20Chem/structure%20determination/IRHydrocarbon.htm)

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### **Fingerprint Region**

Each IR spectrum has a **fingerprint region** to the far right-hand side (between 500cm<sup>-1</sup> and 1500cm<sup>-1</sup>), which appears as an area of lots of peaks very close together. This region appears due to tiny differences in species which act as a molecules' 'fingerprint', allowing it to be **specifically identified**.

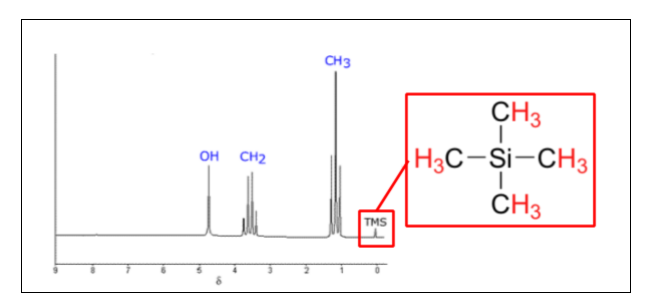
Fingerprint regions are **very difficult to interpret** without specialist knowledge. Therefore, at A Level, it is not a requirement to be able to identify compounds from them.

## **NMR Spectroscopy**

This is an **analytical technique** that allows the **structure** of a molecule to be determined by analysing the **energy of bond environments**. Different bond environments within a molecule absorb different amounts of energy, so they show as **different peaks** on a spectra print out.

The bond environment peaks are measured against a **standard** molecule, **tetramethylsilane**  $(Si(CH_3)_4)$ , known as TMS. This is a standard molecule as it contains four **identical** carbon and hydrogen environments. It is seen as a peak at  $\partial=0$  ppm on the x-axis. This makes it easy to **distinguish** from the other peaks.

Example:



(Modified from https://chemistry.stackexchange.com/questions/42757/why-only-one-peak-is-observed-in-nmr-spectrum-of-h2)

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Using the peaks from an NMR spectrum, the structure of a chemical compound can be determined. This is a method of particular use in fields such as **forensics**, as it allows for the analysis and **identification of unknown substances**.







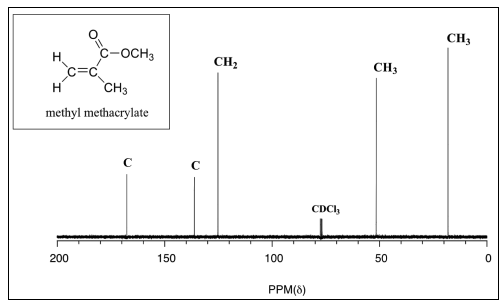




#### C<sup>13</sup> NMR

This form of NMR spectroscopy analyses how many different carbon environments are present in the molecule. The different environments are shown as peaks at different  $\partial$  values.

Carbon environments that are near to an **oxygen** have ∂ values that are **shifted** to the right. This is because oxygen is **very electronegative** and it acts to pull electrons away from the carbon atom.

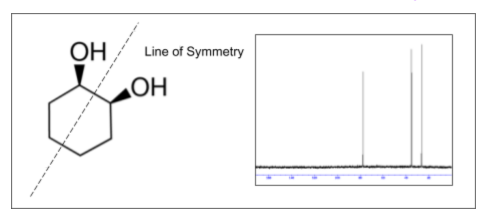


(https://chem.libretexts.org/Under Construction/Purgatory/Book%3A Organic Chemistry with a Biological

Emphasis (Soderberg)/Chapter 05%3A Structure Determination II/5.6%3A 13C-NMR spectroscopy)

Tim Soderberg / CC BY-SA 3.0

Molecules that have **symmetry**, such as 1,2-cyclohexanediol, may display **fewer** ∂ **peaks** than the number of carbon atoms in the molecule. Therefore in these cases, it is important to look at the given **molecular formula** of the compound in order to decipher its **displayed structure**:



(Modified from https://chem.libretexts.org/Bookshelves/Physical\_and\_Theoretical\_Chemistry\_Textbook\_Maps/
Supplemental\_Modules\_(Physical\_and\_Theoretical\_Chemistry)/Spectroscopy/Magnetic\_Resonance\_Spectroscopies/Nucle

ar\_Magnetic\_Resonance/NMR%3A\_Structural\_Assignment/NMR3.\_Symmetry\_in\_NMR)

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All  $C^{13}$  NMR  $\partial$  shift values can be found in most Chemistry data books and will be provided in the exam.

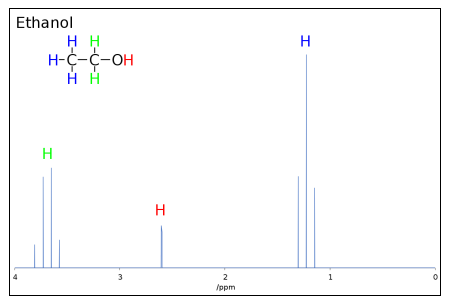
#### H<sup>1</sup> NMR (Proton NMR)

In this form of NMR, the different **hydrogen environments** in a molecule are analysed and displayed as peaks on a spectra. These peaks are also measured against the **TMS standard**.

The samples being analysed must be dissolved in a **non-hydrogen-containing solvent** so that it doesn't produce any peaks on the spectrum. **CCI**<sub>4</sub> is therefore a common solvent used along with **deuterated solvents** containing deuterium, an isotope of hydrogen.

H¹ NMR spectra are **more complex** than C¹³ spectra as the **heights** of the peaks show the relative intensity of each chemical shift value. These relative intensities correspond to the **number of hydrogens** in that certain environment within a molecule, shown as a number above the peak.

#### Example:



(https://nl.wikipedia.org/wiki/Ethanol)
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The peaks of a H¹ NMR spectra also inform where each environment is positioned within the molecule. Peaks are split into a small cluster with smaller peaks indicating how many hydrogens are on the adjacent carbon atom within the molecule. These smaller peaks are a splitting pattern and follow an 'n+1' rule, where n is the number of hydrogen on the adjacent carbon.



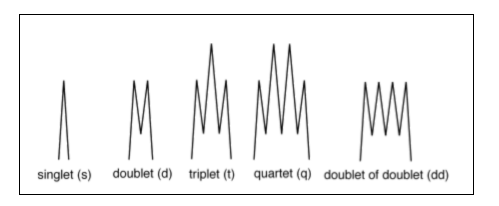






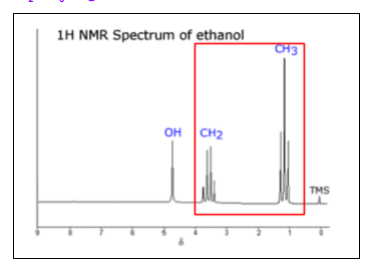


Singlet = no H on adjacent carbon
Doublet = 1 H on adjacent carbon
Triplet = 2 H on adjacent atom
Quartet = 3 H on adjacent carbon



(Modified fromhttps://chem.libretexts.org/Ancillary Materials/Reference
/Organic Chemistry Glossary/Splitting Pattern)
Gamini Gunawardena / CC BY-SA 3.0

There are some common combinations of peaks and splitting patterns that make deciphering the structure of the molecule easier. A triplet-quartet splitting pattern is a common combination as it represents a -CH<sub>2</sub>-CH<sub>3</sub> fragment.



#### The boxed peaks are produced by the -CH<sub>2</sub>-CH<sub>3</sub> fragment.

(Modified from <a href="https://chemistry.stackexchange.com/questions/42757/why-only-one-peak-is-observed-in-nmr-spectrum-of-h2">https://chemistry.stackexchange.com/questions/42757/why-only-one-peak-is-observed-in-nmr-spectrum-of-h2</a>)

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Multiple fragments can be worked out and pieced together to determine the **full molecular structure**.







