Spectroscopy and Chromatography

The effect of different types of radiation on molecules

i **infrared in analysis** – infra red energy causes bonds to vibrate. This can be used to identify the types of bond in a molecule

ii microwaves for heating- certain molecules absorb the microwaves causing them to rotate

iii **radio waves in nmr** – can cause the hydrogen nucleus to **change its spin state.** This can give us information about the arrangements of hydrogens in a molecule.

iv **ultraviolet in initiation of reactions** – UV energy can break bonds such as the CI-CI bond or C-CI bond

NMR spectroscopy

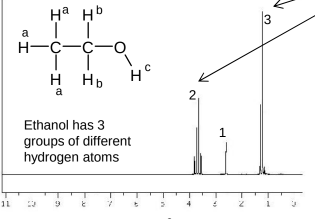
NMR spectroscopy involves interaction of materials with the lowenergy radiowave region of the electromagnetic spectrum

NMR spectroscopy is the same technology as that used in 'magnetic resonance imaging' (MRI) to obtain diagnostic information about internal structures in body scanners e.g. scanning for brain disorders

The radio waves used in proton nmr cause the hydrogen nucleus to **change** its spin state.

Equivalent Hydrogen atoms.

In an H NMR spectrum, there is one signal for each set of equivalent H atoms.



3 sets of equivalent H's: ratio 3:2:9

$$^{\mathrm{a}}_{\mathrm{H_3}}\mathrm{C--CH}=^{\mathrm{c}}_{\mathrm{CH_2}}$$

3 sets of equivalent H's: ratio 3:1:2

In addition the **intensity (integration value)** of each signal is proportional to the **number of equivalent H atoms** it represents.

$$H_3C$$
— C — CH_3 1 signal

3 sets of equivalent H's: ratio 3:2:3

4 sets of equivalent H's: ratio 3:1:2:3

Solvents

Samples are dissolved in solvents without any ¹H atoms, e.g. CCl₄, CDCl₃.

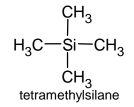
This means that in the H NMR the solvent will not give any peaks

Calibration and shift

A small amount of TMS (tetramethylsilane) is added to the sample to calibrate the spectrum

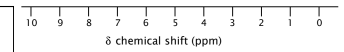
TMS is used because:

- •its signal is away from all the others
- •it only gives one signal
- •it is non-toxic
- •it is inert
- •it has a low boiling point and so can be removed from sample easily



The spectra are recorded on a scale known as the chemical shift (), which is how much the field has shifted away from the field for TMS..

The is a measure in parts per million (ppm) is a relative scale of how far the frequency of the proton signal has shifted away from that for TMS.

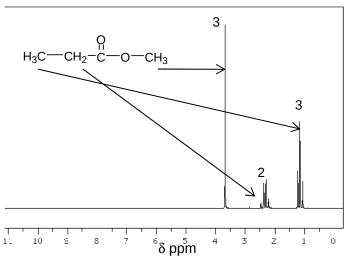


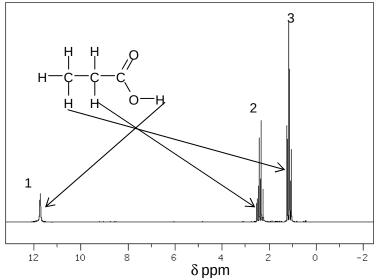
H NMR shift

¹H n.m.r. chemical shift data

Type of proton	δ/ ppm	
RO H	0.5-5.0	
RCH_3	0.7 - 1.2	
RNH_2	1.0 - 4.5	
R_2CH_2	1.2 - 1.4	
R ₃ CH	1.4 - 1.6	
R-C-C- 0 H	2.1-2.6	
R-O-C- H	3.1-3.9	
RCH ₂ Cl or Br	3.1-4.2	
R-C-O-C- 0 H	3.7-4.1	
C = C	4.5 – 6.0	
R-C H	9.0-10.	
R-C O-H	10.0-12.	

The depends on what other atoms/groups are near the H – more electronegative groups gives a greater shift.





Spin-Spin coupling in H NMR

In high resolution H NMR each signal in the spectrum can be split into further lines due to inequivalent H's on neighbouring C atoms.

Nuclei in identical chemical environments do not show coupling amongst themselves!

Splitting of peak = number of inequivalent H's on neighbouring C atoms + 1

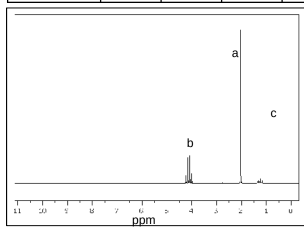
signal	singlet	doublet	triplet	quartet	quintet
appearance					ML
Split number of peaks	1	2	3	4	5
number of neighbouring inequivalent H atoms	0	1	2	3	4
relative size		1:1	1:2:1	1:3:3:1	1:4:6:4:1

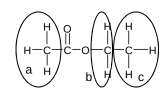
The peak due to group **a** will be a **triplet** as it is next to **b** (a carbon with 2 H's)

The peak due to group **b** will be a **quartet** as it is next to **a** (a carbon with 3H's)

The peak due to group **c** will be a **singlet** as it is next to a carbon with no H's)

For 6 split peaks use the term hextet or multiplet





The peak due to group **a** will be a **singlet** as it is next to a carbon with 0 H's Shift 2.1-2.6 Integration trace 3

The peak due to group **c** will be a **triplet** as it is next to a carbon with 2 H's Shift 0.7-1.2 Integration trace 3

The peak due to group **b** will be a **quartet** as it is next to a carbon with 3 H's Shift 3.7 -4.1 Integration trace 2

Mass spectrometry

Measuring the M_r of an organic molecule

If a molecule is put through a mass spectrometer it will often break up and give a series of peaks caused by the fragments. The peak with the largest m/z, however, will be due to the complete molecule and will be equal to the M, of the molecule. This peak is called the parent ion or

Fragmentation

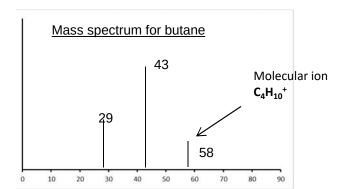
When organic molecules are passed through a mass spectrometer, it detects both the whole molecule and fragments of the molecule.

Several peaks in the mass spectrum occur due to fragmentation. The Molecular ion fragments due to covalent bonds breaking: [M]+-

Relatively stable ions such as carbocations R+ such as CH₂CH₂+ and acylium ions [R-C=O]+ are common. The more stable the ion, the greater the peak intensity.

The peak with the highest mass/charge ratio will be normally due to the original molecule that hasn't fragmented (called the molecular ion). As the charge of the ion is +1 the mass/ charge ratio is equal to Mr.

molecular ion

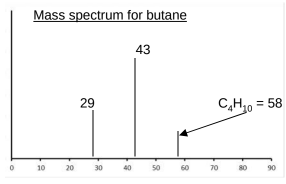


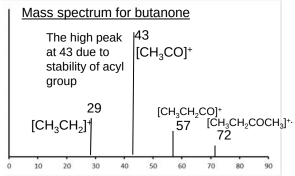
Spectra for C₄H₁₀

Molecular ion formed: [M]+·+ e-M

The molecule loses an electron and becomes both an ion and a free radical

> This process produces an ion and a free radical. The ion is responsible for the peak





Equation for formation molecular ion

$$C_4H_{10} \rightarrow [C_4H_{10}]^{+} + e^-$$
 m/z 58

Equations for formation of fragment ions from molecular ions

$$[C_4H_{10}]^{+\cdot} \rightarrow [CH_3CH_2CH_2]^+ + \cdot CH_3 \text{ m/z } 43$$

X+ + Y-

$$[C_4H_{10}]^{+} \rightarrow [CH_3CH_2]^{+} + \cdot CH_2CH_3 \quad m/z \ 29$$

Equation for formation molecular ion

$$CH_3CH_2COCH_3 \rightarrow [CH_3CH_2COCH_3]^{+} + e^- m/z 72$$

Equations for formation of fragment ions from molecular ions

$$[CH_3CH_2COCH_3]^{+} \rightarrow [CH_3CH_2CO]^+ + \cdot CH_3 \text{ m/z } 57$$

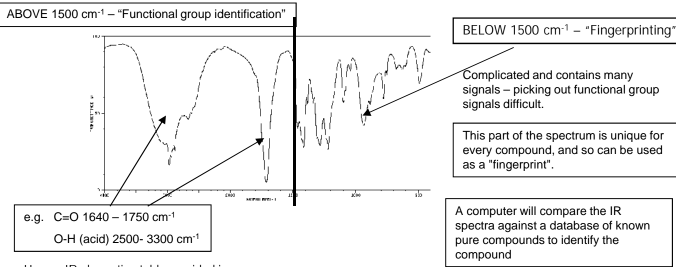
$$[CH_3CH_2COCH_3]^{+} \rightarrow [CH_3CO]^{+} + \cdot CH_2CH_3 \text{ m/z } 43$$

$$[CH_3CH_2COCH_3]^{+\cdot} \rightarrow [CH_3CH_2]^+ + \cdot COCH_3 \text{ m/z } 29$$

Infrared spectroscopy

Certain bonds in a molecule absorb infra-red radiation at characteristic frequencies causing the covalent bonds to vibrate

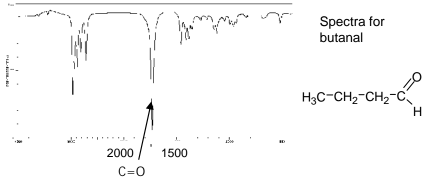
Complicated spectra can be obtained than provide information about the types of bonds present in a molecule



Use an IR absorption table provided in exam to deduce presence <u>or</u> absence of particular bonds or functional groups

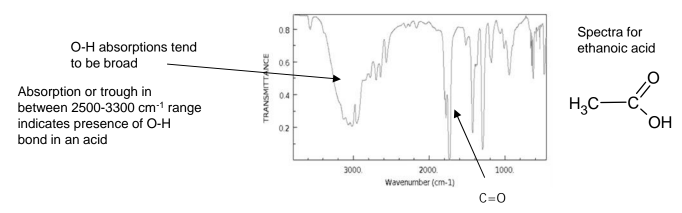
Bond Wavenumber C-O 1000-1300 C=O 1640-1750 C-H 2850 - 3100 2500-3300 O-H Carboxylic acids Very broad N-H 3200-3500 O-H 3200-3550 Acohols, phenols broad

use spectra to identify particular functional groups limited to data presented in wavenumber form e.g. an alcohol from an absorption peak of the O–H bond,



Absorption or trough in between 1640-1750 cm⁻¹ range indicates presence of C=O bond

Always quote the wave number range from the data sheet



Modern breathalysers measure ethanol in the breath by analysis using infrared spectroscopy

Chromatography

Chromatography is an analytical technique that separates components in a mixture between a mobile phase and a stationary phase.

Separation by column chromatography depends on the balance between solubility in the moving phase and retention in the stationary phase.

A solid stationary phase separates by adsorption, A liquid stationary phase separates by relative solubility

HPLC stands for high performance liquid chromatography.

HPLC: stationary phase is a solid silica

HPLC: mobile phase a liquid

The mobile phase may be a liquid or a gas. The stationary phase may be a solid (as in thin-layer chromatography, TLC) or either a liquid or solid on a solid support (as in gas chromatography, GC)

If the stationary phase was polar and the moving phase was non- polar e.g. Hexane. Then non-polar compounds would pass through the column more quickly than polar compounds as they would have a greater solubility in the non-polar moving phase.

(Think about intermolecular forces)

In gas-liquid chromatography GC the **mobile** phase is a inert **gas** such as nitrogen, helium, argon.

The **Stationary** phase is a **liquid** on an inert solid.

Gas-Liquid Chromatography

Gas-liquid chromatography can be used to separate mixtures of volatile liquids.

The time taken for a particular compound to travel from the injection of the sample to where it leaves the column to the detector is known as its **retention time.** This can be used to identify a substance.

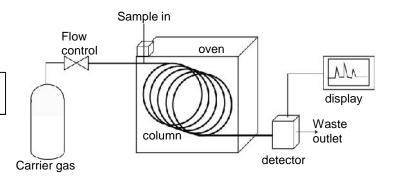
Some compounds have similar retention times so will not be distinguished.

Basic gas-liquid chromatography will tell us how many components there are in the mixture by the number of peaks. It will also tell us the abundance of each substance. The area under each peak will be proportional to the abundance of that component.

It is also possible for gas-liquid chromatography machine to be connected to a mass spectrometer, IR or NMR machine, enabling all the components in a mixture to be identified.

GC-MS is used in analysis, in forensics, environmental analysis, airport security and space probes.

In gas-liquid chromatography, the mobile phase is a gas such as helium and the stationary phase is a high boiling point liquid absorbed onto a solid.



Most commonly a mass spectrometer is combined with GC to generate a mass spectra which can be analysed or compared with a spectral database by computer for positive identification of each component in the mixture.

TLC Chromatography (thin-layer chromatography)

A mixture can be separated by chromatography and identified from the amount they have moved. (Can be used with mixtures of amino acids)

Method: Thin-layer chromatography

- a) **Wearing gloves**, draw a **pencil line** 1 cm above the bottom of a TLC plate and mark spots for each sample, equally spaced along line.
- b) Use a capillary tube to add a **tiny drop** of each solution to a different spot and allow the plate to air dry.
- c) Add solvent to a chamber or large beaker with a lid so that is no more than **1cm in depth**
- d) Place the TLC plate into the chamber, making sure that the level of the solvent is below the pencil line. Replace the lid to get a tight seal.
- e) When the level of the solvent **reaches about 1 cm from the top of the plate**, remove the plate and mark the solvent level with a pencil. Allow the plate to **dry in the fume cupboard**.
- f) Place the plate under a **UV lamp** in order to see the spots. Draw around them lightly in pencil.
- g) Calculate the Rf values of the observed spots.

Wear plastic gloves to prevent contamination from the hands to the plate

pencil line -will not dissolve in the solvent

tiny drop – too big a drop will cause different spots to merge

Depth of solvent– if the solvent is too deep it will dissolve the sample spots from the plate

lid– to prevent evaporation of toxic solvent

Will get more accurate results if the solvent is allowed to rise to near the top of the plate but the Rf value can be calculated if the solvent front does not reach the top of the plate

dry in a fume cupboard as the solvent is toxic

UV lamp used if the spots are colourless and not visible

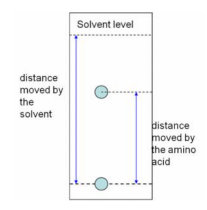
If using amino acids then ninhydrin spray can be used instead of UV lamp to locate the spots

 R_f value = distance moved by amino acid distance moved by the solvent

Each substance has its own R_f value

Measure how far each spot travels relative to the solvent front and calculate the *R*f value.

Compare Rf values to those for known substances.



Some substances won't separate because similar compounds have similar *R*f values. So some spots may contain more than one compound

Bringing it all together

1. Work out empirical formula

Elemental analysis C 66.63% H 11.18% O 22.19%

C H O 66.63/12 11.18/1 22.19/16 =5.5525 =11.18 =1.386875 =4 =8 =1

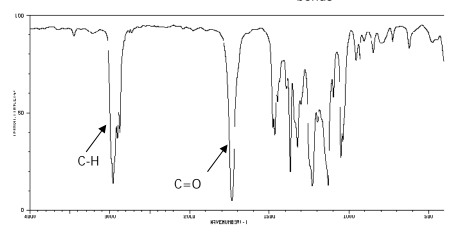
2. Using molecular ion peak m/z value from mass spectrum calculate Molecular formula

molecular ion peak m/z value= 144

Mr empirical formula $C_4H_8O = 72$ If Mr molecular formula 144 then compound is $C_8H_{16}O_2$

3. Use IR spectra to identify main bonds/functional group

 $\rm C_8H_{16}O_2$ could be an ester, carboxylic acid or combination of alcohol and carbonyl. Look for IR spectra for C=O and O-H bonds



There is a C=O but no O-H absorptions, so must be an ester.

4. Use NMR spectra to give details of carbon chain

4 peaks – only 4 different environments.

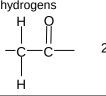
Peak at δ 4 shows H-C-O

Area 2 suggests CH₂ Quartet means next to a CH₃ H

5

-0-C-H Peak at δ 2.2 shows H–C=O

Area 2 suggests CH₂ Singlet means adjacent to C with no hydrogens



Peak at δ 1.2 shows R-CH₃ Area 3 means CH₃ Triplet means next to a CH₂

2

--СH₃

Put all together to give final structure

2

4

3

 δppm