Mass Spectra and IR

Mass Spectrometry

Fragmentation

fragments of the molecule.

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_r of the molecule. This peak is called the parent ion or **molecular ion**

When organic molecules are passed through a mass

spectrometer, it detects both the whole molecule and

Several peaks in the mass spectrum occur due to fragmentation.

Spectra for C₄H₁₀



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

acylium ions [R-C=O]⁺ are common. The more stable the ion, the greater the peak intensity.

Relatively stable ions such as carbocations R⁺ such as CH₃CH₂⁺ and

The Molecular ion fragments due to covalent bonds breaking: $[M]^+ \rightarrow X^+ + Y^-$

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.



Equation for formation molecular ion

 $CH_3CH_2CHO \rightarrow [CH_3CH_2CHO]^{+.} + e^{-} m/z 58$

Equations for formation of fragment ions from molecular ions

 $[\mathrm{CH_3CH_2CHO}]^{\text{+}} \rightarrow \ [\mathrm{CHO}]^{\text{+}} + \ \cdot \mathrm{CH_2CH_3} \quad \ \mathrm{m/z} \ \mathrm{29}$

 $[CH_3CH_2CHO]^{+} \rightarrow [CH_3CH_2]^+ + CHO m/z 29$

Equation for formation molecular ion

 $\mathrm{CH_3COCH_3} \ \rightarrow \ [\mathrm{CH_3COCH_3}]^{\text{+}} \ + \ \mathrm{e^{-}} \quad \mathrm{m/z} \ 58$

Equations for formation of fragment ions from molecular ions

 $[\text{CH}_3\text{COCH}_3]^{\text{+}\text{-}} \rightarrow [\text{CH}_3\text{CO}]^{\text{+}} + \ \cdot \text{CH}_3 \text{ m/z 43}$

It is not possible for propanone to fragment to give a peak at 29 so the fragmentation patterns can distinguish between the structural isomers of propanone and propanal

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-0	1000-1300
C=O	1640-1750
C-H	2850 -3100
O-H Carboxylic acids	2500-3300 Very broad
N-H	3200-3500
O-H Acohols, phenols	3200- 3550 broad

Use spectra to identify particular functional groups e.g. an alcohol from an absorption peak of the O–H bond, or C=O stretching absorption in aldehydes and ketones



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

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

Molecules which change their polarity as they vibrate can absorb infrared radiation. E.g. C-H, C=O, O-H

Molecules such as H_2 , O_2 and N_2 cannot change their polarity as they vibrate so can absorb infrared radiation and don't register on an infra red spectrum

The absorption of infra-red radiation by bonds in this type of spectroscopy is the same absorption that bonds in CO_2 , methane and water vapour in the atmosphere do that cause them to be greenhouse gases.

 H_2O , CO_2 , CH_4 and NO molecules absorb IR radiation and are greenhouse gases, whilst O_2 and N_2 are not.