# **NUCLEAR MAGNETIC RESONANCE SPECTROSCOPY**

Involves interaction of materials with the **low-energy radiowave region** of the electromagnetic spectrum

### **Origin of Spectra**

Theory All nuclei possess charge and mass. However, those with either an odd mass number or an odd atomic number also possess spin and have angular momentum.

> examples  ${}^{1}_{1}H$   ${}^{2}_{1}H$   ${}^{13}_{6}C$   ${}^{14}_{7}N$   ${}^{17}_{8}O$   ${}^{19}_{9}F$   ${}^{31}_{16}P$  possess spin but  ${}^{12}_{6}$ C  ${}^{16}_{8}$ O do not  $^{1}_{1}$ H  $^{2}_{1}$ H  $^{2}_{1}$ H  $^{13}_{6}$ C <sup>13</sup>C <sup>14</sup>N<br>6 7  $^{14}_{7}N$   $^{17}_{8}O$ <sup>17</sup>o <sup>19</sup>F <sup>9</sup>F <sup>31</sup><br>9<sup>F</sup> 15 15

A nucleus with spin can be detected by nuclear magnetic resonance (nmr) spectroscopy.

A spinning nucleus such as <sup>1</sup>H behaves as a spinning charge and generates a magnetic field. It can be likened to a bar magnet.



When placed in an externally applied field it can align with, or against, the field.

The energy difference between the two states (∆Ε) depends on the applied field.

The sample is placed in the field of a large electromagnet and a radio-frequency (RF) field is applied. The magnetic field is increased and the excitation or "flipping" of nuclei from one orientation to another is detected as an induced voltage resulting from the absorption of energy from the RF field.



**The basic arrangement of an nmr spectrometer**

An nmr spectrum is the plot of the induced voltage against the sweep of the field. The area under a peak is proportional to the number of nuclei "flipping"

# **Other**

uses NMR spectroscopy is the same technology as that used in **MRI (magnetic resonance imaging)** to obtain diagnostic information about internal structures in body scanners.

## **INTERPRETATION OF PROTON NMR SPECTRA**

**Introduction** Spectra provide information about the structure of organic molecules from the ...

- number of different signals in the spectrum
- position of the signals (chemical shift)
- splitting pattern of the signals
- intensity of the signals

# **Running**

- a liquid sample is placed in a long thin tube which spins in a magnetic field
	- solids are **dissolved in solvents which won't affect the spectrum CDCl<sup>3</sup>**
	- **tetramethylsilane, (CH3)4Si**, is added to provide a reference signal
	- the spectrum can be 'integrated' to find the relative peak heights

- **TMS** it produces a single intense peak
	- signal is just outside the range shown by most protons
	- it is inert, non-toxic and has a low boiling point it can be distilled off if required

$$
\begin{array}{c}\n\text{CH}_3\\ \n\text{I} \\
\text{CH}_3-\text{Si}-\text{CH}_3\\ \n\text{CH}_3\n\end{array}
$$

#### **TETRAMETHYLSILANE**

# **Chemical**

- each proton type is said to be **chemically shifted** relative to a standard
- the chemical shift is the difference between the field strength at which it absorbs and the field strength at which TMS protons absorb
- the delta (δ) scale is widely used as a means of reporting chemical shifts

$$
\delta = \frac{\text{Observed chemical shift (Hz) x 10}^6}{\text{Spectrometer frequency (Hz)}}
$$
 ppm (parts per million)

- the chemical shift of a proton is constant under the same conditions
- the **TMS** peak is assigned a value of ZERO ( $\delta$  = 0.00)
- all peaks in the spectrum are related to it and reported in parts per million
- H's near to an electronegative species are shifted "downfield" to higher δ values







# **Multiplicity** This occurs because the spin of one nucleus affects that of a **chemically different nucleus on an adjacent atom**.

- also known as coupling or spin-spin splitting
- low resolution nmr gives 1 peak for each chemically different group of protons
- high resolution nmr gives more complex signals doublets, triplets, quartets...
- the signal produced indicates the number of protons on adjacent carbon atoms

# **No. of peaks = number of H's on adjacent chemically different atoms + 1**



- Theory Splitting patterns are worked out by considering the effect **adjacent, chemically different protons** have on another signal in a given environment. The spin of the proton producing the signal is affected by each of the two forms of the adjacent proton. One orientation augments/enhances its field and the other opposes/reduces it. Splitting patterns can be worked out by calculating the various **possible combinations** of alignment of adjacent protons.
	- 1 adjacent H can be aligned either with ( $\alpha$ ) or against (β) the field Fig. 1 only two equally probable possibilities the signal is split into **2 peaks** of **equal intensity**
	- 2 adjacent H's more possible combinations signal is more complex Fig. 2 get **3 peaks** in the ratio **1 : 2 : 1**
	- 3 adjacent H's even more possible combinations Tig. 3 get **4 peaks** in the ratio **1 : 3 : 3 : 1**



*Explain the splitting pattern when there are four adjacent protons.* 



expect a triplet signal (3 peaks) for the H in O-H but there is only one.

# **Integration** • the area under a signal is proportional to the number of hydrogen atoms present

- is achieved using an integration device which scans the peaks
- lines on the spectrum show the **relative abundance** of each hydrogen type

By measuring the distances between the integration lines (dotted line on the spectrum) one can work out a simple ratio between the various types of hydrogen.

## **The nmr spectrum of ethanol**



*Q.3 Measure the ratio of the heights of the integration lines in the ethanol spectrum. Does it correspond to the actual ratio of protons in the structure ?*

 **NOTE** Computers now analyse the area under each peak as the spectrum is run. The relative values are printed on the spectrum rather than showing the integration lines.

*Q.4 How many different proton environments are there in the following molecules? State the ratio of the number of protons in each environment. no. of H environments ratio of H's a*)  $CH_3CH_2CH_2CHBrCH_3$ *b*)  $CH_3CH_2CHBrCH_2CH_3$ 

*c) C6H6 (benzene)*

**Spectra of compounds containing OH groups**

Introduction Observation of spectra containing OH groups will reveal that **OH** protons ...

- only produce a single peak (singlet)
- are not counted as 'neighbouring protons' when working out splitting patterns



Explanation The proton (H) on any OH group can exchange rapidly with another H atom on any trace water. Because of this the H isn't around long enough to register a splitting signal for itself or any neighbouring protons.

### **D2O shake This is used to determine which signal is due to an OH proton**

- after a spectrum is run, the sample is shaken with a small amount of  $D_2O$
- any signal due to an OH proton disappears.



nmr spectrum of ethanol **before** a D<sub>2</sub>O shake



nmr spectrum of ethanol **after** a  $D_2O$  shake

Explanation **It is possible to exchange the H for deuterium, <sup>2</sup>H or D. The usual source is deuterium oxide, D2O, a form of water.**

**ROH + D2O ROD + HOD**

Deuterium doesn't exhibit nuclear magnetic resonance under the conditions used for proton nmr so the signal is removed to another part of the spectrum.

It also works with H's on -NH- or -NH<sub>2</sub>





# SOME TYPICAL PROTON CHEMICAL SHIFTS

F324

 $\overline{7}$ 

**INTERPRETING THE NMR SPECTRUM OF 1-BROMOPROPANE**



#### **Analysis**



Peaks There are three different signals so there are three chemically different protons.

Position The further the signals are shifted from TMS signal, the nearer they are to the Br.



The signals due to the protons attached to carbon ...



Integration The integration lines show that the ratio of protons is 2:2:3

**Summary** An nmr spectrum provides several types of information :-



In many cases this information is sufficient to deduce the structure of a molecule.

*Q.5 What would you expect the spectrum of 2-bromopropane to be like ? Explain your answer in terms of a) the chemical shift b) the splitting pattern c) integration*



Theory After hydrogen, the most useful atom providing information to organic chemists is carbon-13. Natural carbon contains about 1% of this isotope so the instruments for its detection need to be sensitive and spectra will take longer to record.

> Only the chemical shift is important as **each spectrum gives only single lines for each chemically equivalent carbon**. Carbon-13 nmr has wide applications in the study of natural products, biological molecules and polymers.



**Interpretation** for  $C_3H_7Br$ 









**The Carbon-13 spectrum of 2-methylbutane C5H<sup>12</sup>**



There are four chemically different carbon atoms in the molecule so there are four peaks in the <sup>13</sup>C nmr spectrum.



Other isomers of  $C_5H_{12}$  pentane  $CH_3CH_2CH_2CH_2CH_3$  3 peaks 2,3-dimethylpropane  $(CH_3)_4C$  2 peaks

## **SUPPLEMENTARY QUESTIONS**







