

Introduction:-

Nuclear Magnetic Resonance (NMR) is a spectroscopy technique which is based on the absorption of electromagnetic radiation in the **radio frequency region 4 to 900 MHz** by nuclei of the atoms.

Proton Nuclear magnetic resonance spectroscopy is one of the most powerful tools for elucidating the number of hydrogen or proton in the compound.

It is used to study a wide variety of nuclei:

- ^1H ^{15}N
- ^{19}F ^{19}F
- ^{13}C ^{31}P

Page 3

Theory of NMR:-

Spin quantum number (I) is related to the atomic and mass number of the nucleus.

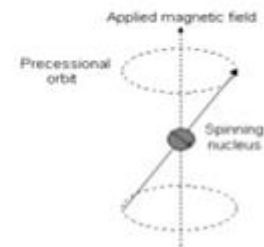
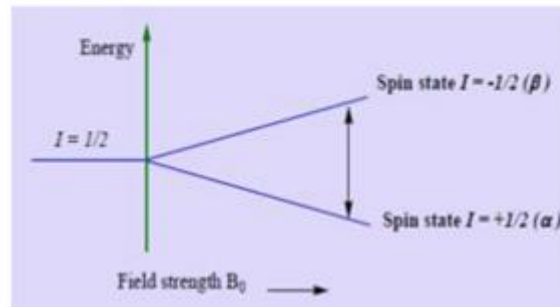
I	Atomic Mass	Atomic Number	Examples	
Half-integer	Odd	Odd	^1H (1/2)	NMR active
Half-integer	Odd	Even	^{13}C (1/2)	
Integer	Even	Odd	^2H (1)	
Zero	Even	Even	^{12}C (0)	Not NMR active

Elements with either **odd mass** or **odd atomic number** have the property of nuclear "spin".

Page 4

If an external magnetic field is applied, the number of possible orientations calculated by $(2I+1)$.

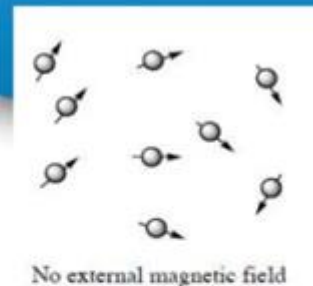
Hydrogen has spin quantum number $I=1/2$ and possible orientation is $(2*1/2+1=2)$ two $+1/2$ and $-1/2$.



Page 5

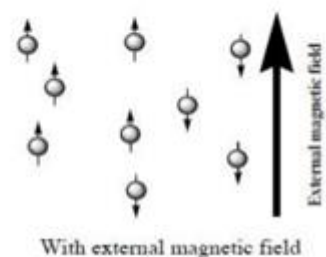
Principles of NMR

The theory behind NMR comes from the spin of a nucleus and it generates a magnetic field. Without an external applied magnetic field, the nuclear spins are random in directions.



No external magnetic field

But when an external magnetic field (B_0) is present the nuclei align themselves either with or against the field of the external magnet.

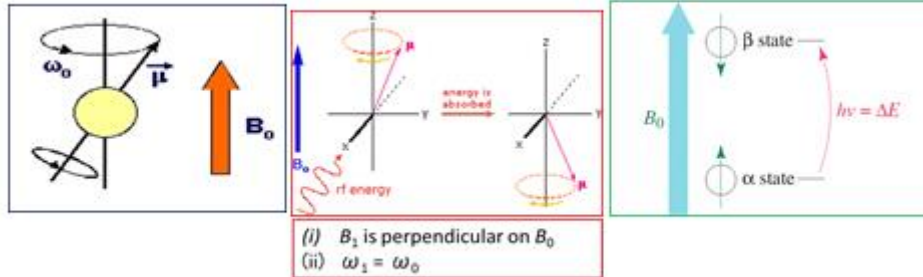
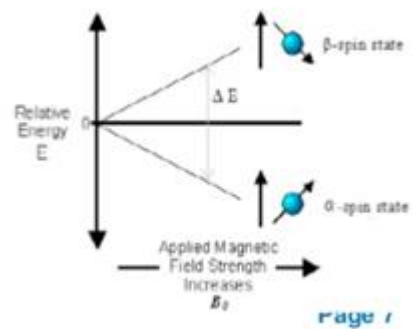
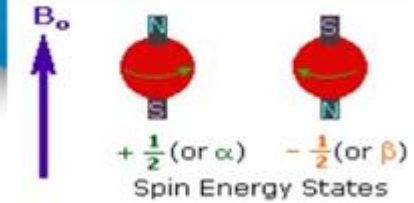


With external magnetic field

If an external magnetic field is applied, an energy transfer (ΔE) is possible between ground state to excited state.

when the spin returns to its ground state level, the absorbed radiofrequency energy is emitted at the same frequency level.

The emitted radiofrequency signal that give the NMR spectrum of the concerned nucleus.



$$\Delta E = h\nu = \frac{\gamma B^0}{2\pi} \text{-----1}$$

$$\gamma = \frac{2\pi\mu}{hI} \text{-----2}$$

$$\omega^0 = \gamma B^0 \text{-----3}$$

Add equation 1 to 3 produced

$$\omega^0 = 2\pi\nu \text{-----4}$$

Classical Description

- *Angular velocity of this motion is given by:*

$$\omega_o = \gamma B_o$$

where the frequency of precession or **Larmor** frequency is:

$$\nu = \gamma B_o / 2\pi$$

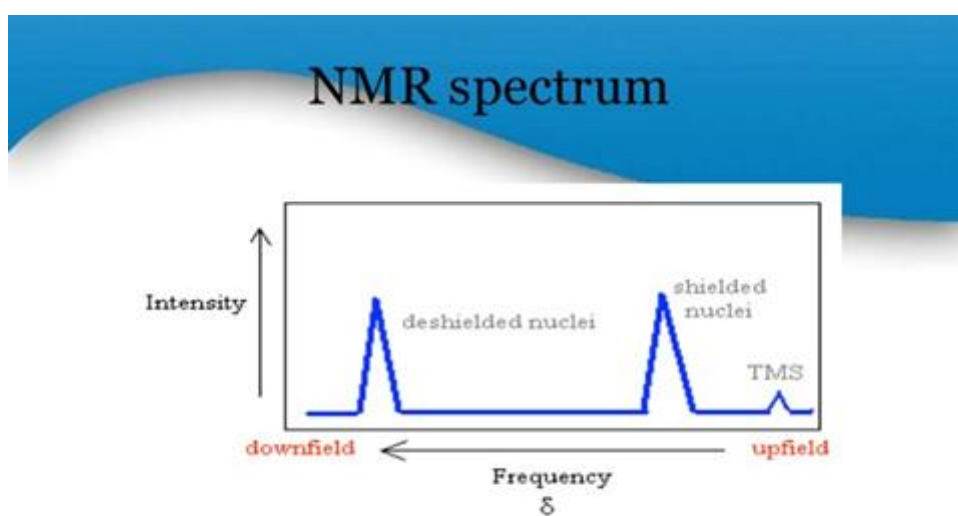
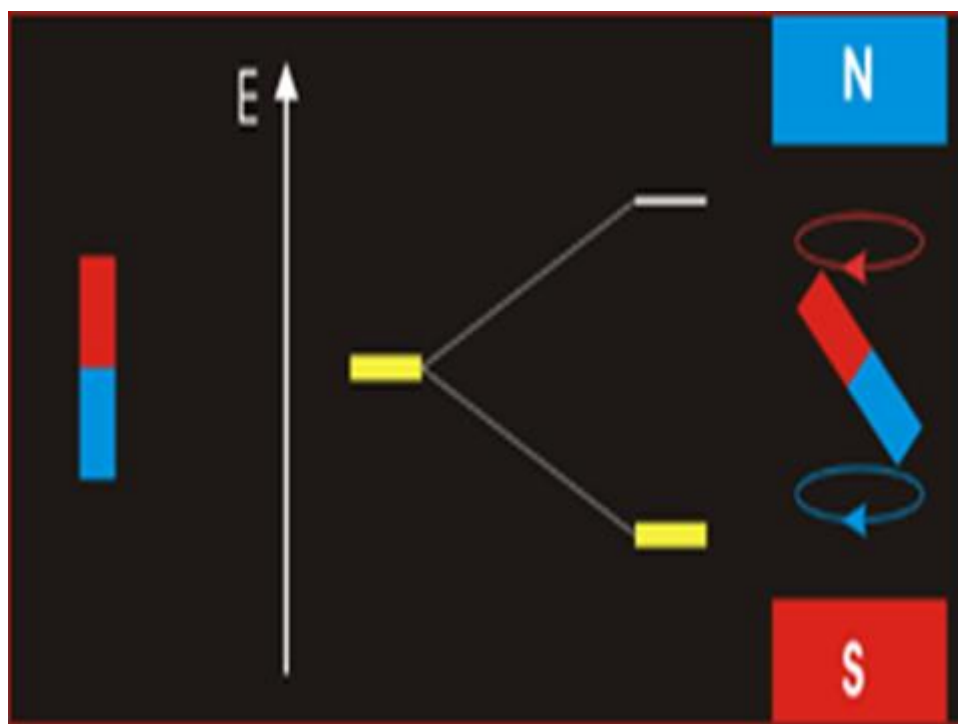
Same as quantum mechanical description

The emitted radio frequency is directly proportional to the strength of the applied field.

$$\nu = \frac{\gamma B_o}{2\pi}$$

B_o = External magnetic field experienced by proton

γ = Magnetogyric ratio (The ratio between the nuclear magnetic moment and angular moment)

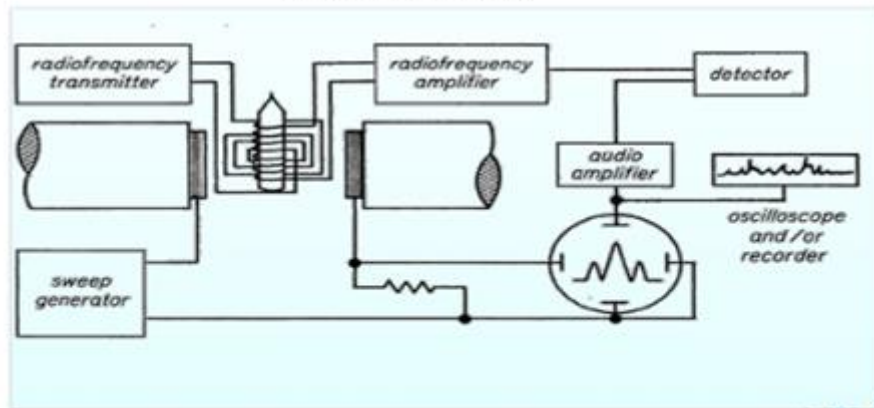


The NMR spectrum is a plot of intensity of NMR signals VS magnetic field (frequency) in reference to TMS

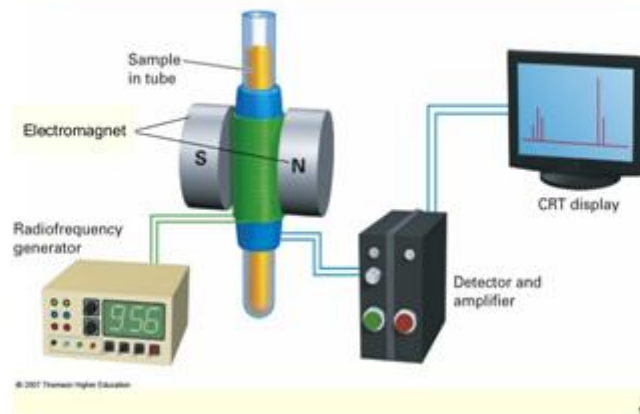
NMR instrumentation

1. Sample holder
2. Permanent magnet
3. Magnetic coils
4. Sweep generator
5. Radio frequency transmitter
6. Radio frequency receiver

7. Read out systems



Schematic diagram of NMR set-up



- 1. Sample holder** :- Glass tube with 8.5 cm long, 0.3 cm in diameter
- 2. Permanent magnet** :- It provides homogeneous magnetic field at 60-100 MHZ
- 3. Magnetic coils** :- These coils induce magnetic field when current flows through them.
- 4. Sweep generator** :- To produce the equal amount of magnetic field pass through the sample

5. Radio frequency transmitter :- A radio transmitter coil that produces a short powerful pulse of radio waves

6. Radiofrequency Receiver :- A radio receiver coil that detects radio frequencies emitted as nuclei relax to a lower energy level

7. Readout system :- A computer that analyses and record the data

Solvents used in NMR

The following solvents are normally used in which hydrogen replaced by deuterium.

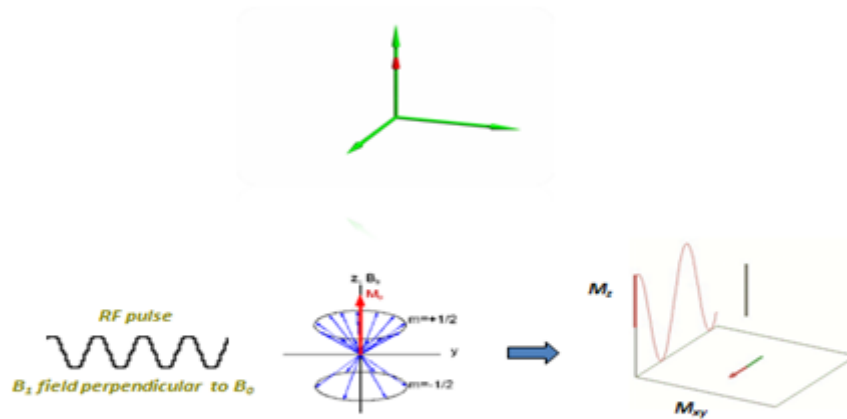
CCl_4 - carbon tetrachloride

CS_2 - carbon disulfide

CDCl_3 - Deuteriochloroform

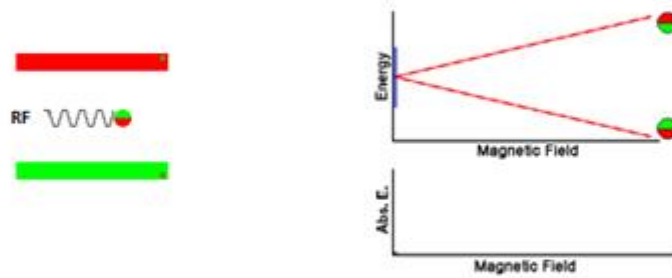
C_6D_6 - Hexa deuteriobenzene

D_2O - Deuterium oxide



Spins Orientation in a Magnetic Field (Energy Levels)

- Transition from the low energy to high energy spin state occurs through an absorption of a photon of radio-frequency (RF) energy



Frequency of absorption:

$$\nu = \gamma B_0 / 2\pi$$

Chemical shift

A **chemical shift** is defined as the difference in parts per million (ppm) between the resonance frequency of the observed proton and tetramethylsilane (TMS) hydrogens.

TMS is the most common reference compound in NMR, it is set at $\delta=0$ ppm

$$\text{Chemical shift, } \delta = \frac{\text{frequency of signal} - \text{frequency of reference}}{\text{spectrometer frequency}} \times 10^6$$

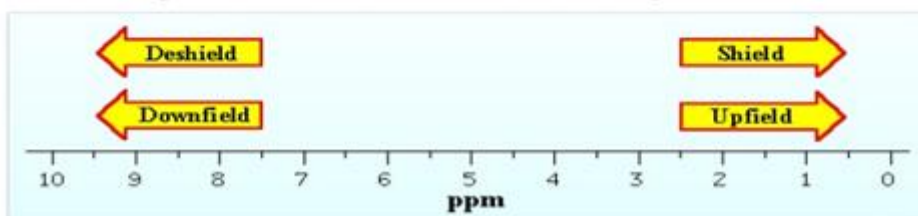
Page 14

Shielding of protons:-

High electron density around a nucleus **shields** the nucleus from the external magnetic field and the signals are **upfield** in the NMR spectrum

Deshielding of protons:-

Lower electron density around a nucleus **deshields** the nucleus from the external magnetic field and the signals are **downfield** in the NMR spectrum



Factors affecting chemical shift:-

- Electronegative groups
- Magnetic anisotropy of π -systems
- Hydrogen bonding

Electronegative groups:-

Electronegative groups attached to the C-H system decrease the electron density around the protons, and there is less shielding (*i.e.* deshielding) and chemical shift increases

Compound	Chemical shift
CH ₃ I	2.16
CH ₃ Br	2.65
CH ₃ Cl	3.10
CH ₃ F	4.26

Page 16

Magnetic anisotropy of π -systems:-

- The word "anisotropic" means "non-uniform". So magnetic anisotropy means that there is a "non-uniform magnetic field".
- Electrons in π systems (e.g. aromatics, alkenes, alkynes, carbonyls etc.) interact with the applied field which induces a magnetic field that causes the anisotropy.
- It causes both shielding and deshielding of protons.
- Example:-Benzene

Hydrogen bonding:-

- Protons that are involved in hydrogen bonding are typically change the chemical shift values.
- The more hydrogen bonding, the more proton is deshielded and chemical shift value is higher.

Page 17





















Proton NMR

The most common form of NMR is based on the hydrogen-1 (^1H), nucleus or proton.

It can give information about the structure of any molecule containing hydrogen atoms.

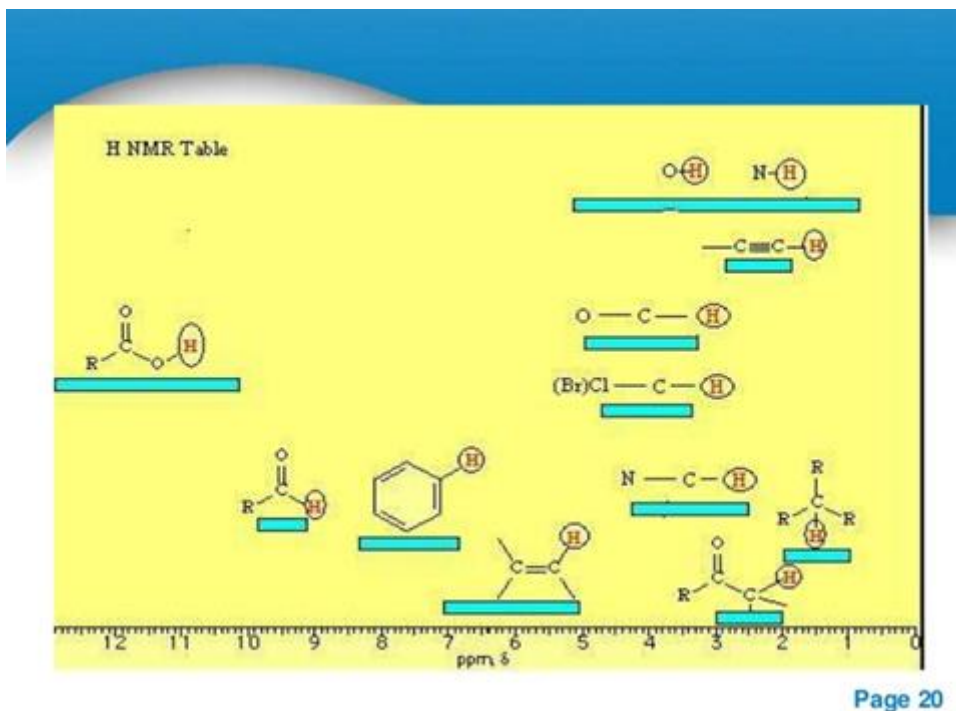
Page 18

^1H NMR chemical shift

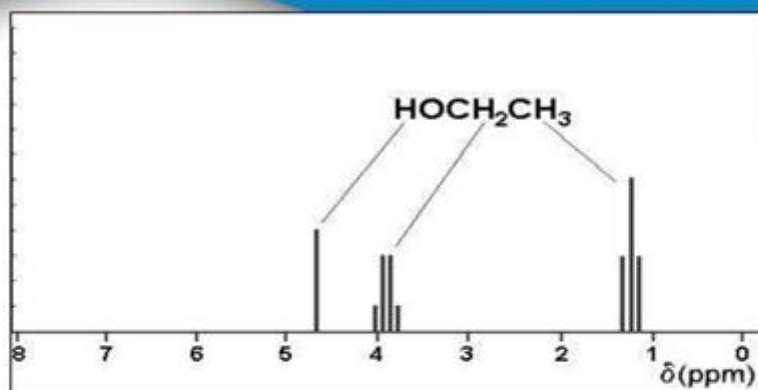
Type of proton	Approximate chemical shift (ppm)	Type of proton	Approximate chemical shift (ppm)	Type of carbon	Approximate chemical shift (ppm)	Type of carbon	Approximate chemical shift (ppm)
$(\text{CH}_3)_4\text{Si}$	0		6.5-8	$(\text{CH}_3)_4\text{Si}$	0	$\text{C}-\text{I}$	0-40
$-\text{CH}_3$	0.9		9.0-10	$\text{R}-\text{CH}_3$	9-35	$\text{C}-\text{Br}$	25-65
$-\text{CH}_2-$	1.3		2.5-4	$\text{R}-\text{CH}_2-\text{R}$	15-50	$\text{C}-\text{Cl}$	35-80
$-\text{CH}-$	1.4		2.5-4		20-60	$\text{C}-\text{N}$	40-60
$-\text{C}-\text{C}-\text{CH}_3$	1.7		3-4		30-40	$\text{C}-\text{O}$	50-80
	2.1		4-4.5	$\text{R}-\text{C}(\text{R})-\text{R}$	30-40		165-175
	2.3		4-5	$\text{R}-\text{C}(\text{R})-\text{R}$	30-40		165-175
$-\text{C}\equiv\text{C}-\text{H}$	2.4	RNH_2	Variable, 1.5-4	$\text{C}=\text{C}$	65-85		175-185
$\text{R}-\text{O}-\text{CH}_3$	3.3	ROH	Variable, 2-5	$\text{C}=\text{C}$	100-150		190-200
$\text{R}-\text{C}=\text{CH}_2$	4.7	ArOH	Variable, 4-7		110-170		205-220
$\text{R}-\text{C}(\text{R})=\text{C}-\text{H}$	5.3		Variable, 10-12				
			Variable, 5-8				

*The values are approximate because they are affected by neighboring protons.

Page 19



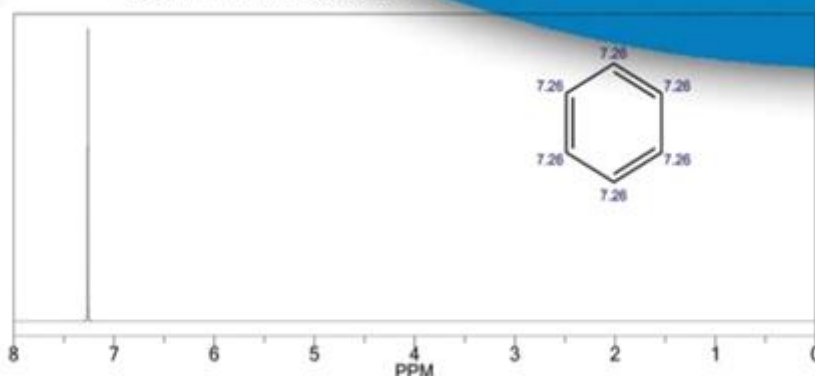
Proton NMR spectra of Ethanol:-



¹H spectrum of Ethanol:-
3 types of proton
CH₃, CH₂, OH

^1H NMR spectra of Benzene

ChemNMR ^1H Estimation



- Aromatic Hydrogen shows peak in the chemical shift scale 6.5-8.0 ppm.
- From the above spectrum Benzene has same type of protons and it shows single peak at 7.26

Page 22

Interpretation of ^1H NMR spectra:-

Number of signals	- Indicates how many "different kinds" of protons are present.
Position of signals	- Indicates something about (chemicalshift) magnetic (electronic) environment of protons
Relative intensity of signals	- Proportional to number of protons present
Splitting of signals (spin spin coupling)	- Indicates the number of nearby nuclei usually protons

Page 23

n+1 rule:-

- ❖ The multiplicity of signal is calculated by using n+1 rule.
- ❖ This is one of the rule to predict the splitting of proton signals. This is considered by the nearby hydrogen nuclei.

Therefore, n = Number of protons in nearby nuclei

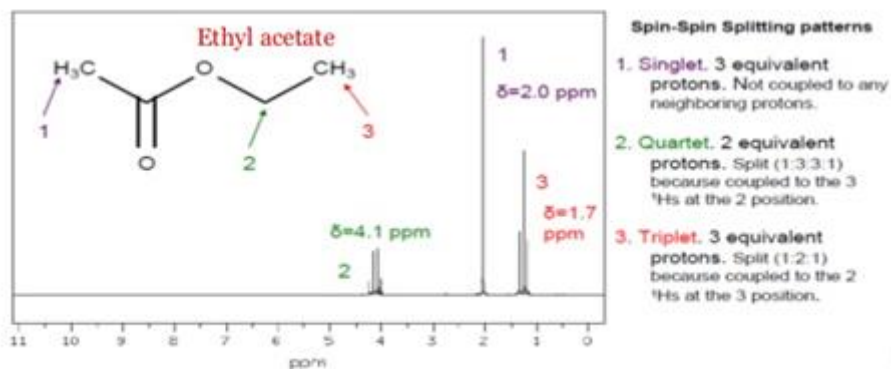
- Zero H atom as neighbour $n+1=0+1=1$ (singlet)
- One H atom as neighbour $n+1=1+1 = 2$ (doublet)
- Two H atom as neighbour $n+1=2+1 = 3$ (triplet)

Page 24

Spin-spin coupling (splitting)

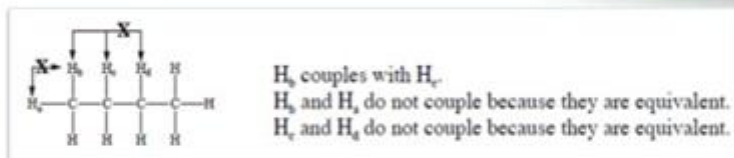
The interaction between the spins of neighbouring nuclei in a molecule may cause the splitting of NMR spectrum. This is known as spin-spin coupling or splitting.

The splitting pattern is related to the number of equivalent H-atom at the nearby nuclei.

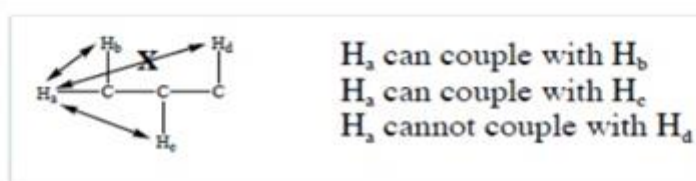


Rules for spin-spin coupling:-

- * Chemically equivalent protons do not show spin-spin coupling.
- * Only nonequivalent protons couple.



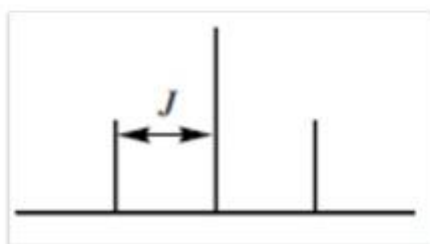
- * Protons on adjacent carbons normally will couple.
- * Protons separated by four or more bonds will not couple.



Page 26

Coupling Constant

- The distance between the peaks in a given multiplet is a measure of the splitting effect known as coupling constant.
- It is denoted by symbol J , expressed in Hz.
- Coupling constants are a measure of the effectiveness of spin-spin coupling and very useful in ^1H NMR of complex structures.



Page 27

Routine 2-D NMR Experiments:-

Correlation Spectroscopy (COSY) – Scalar Coupling

- » Identifies all coupled spins systems.

Nuclear Overhauser Effect Spectroscopy (NOESY) – Dipolar Coupling

- » Identifies neighboring spin systems ($\leq 5 \text{ \AA}$)
- » Identifies chemical exchange.

Heteronuclear Multiple/Single Quantum Correlation (HMQC/HSQC) – Scalar Coupling

- » Identifies coupling between heteronuclei (C-H, N-H)

Page 33

COSY: Homonuclear correlated spectroscopy. Correlation between protons that are coupled to each other

TOCSY: Total Correlation Spectroscopy- Uses a spin-lock for coherence transfer. During the spin-lock all protons of a coupled system become "strongly coupled," leading to cross peaks between all resonances of a coupled system.

HETCOR: Heteronuclear correlation, usually between ^1H and ^{13}C resonances mediated by J_{C-H}

Page 34

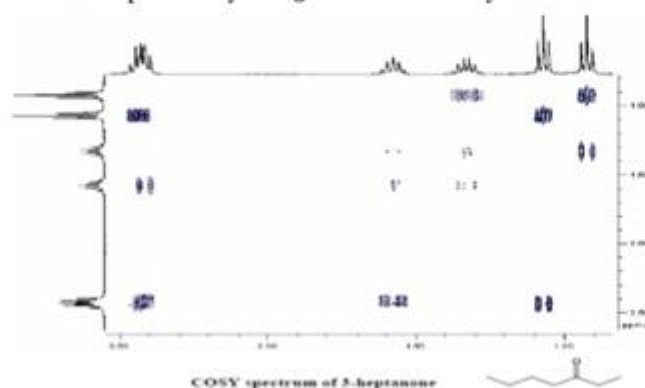
NOESY, ROESY: Proton-proton correlation mediated by dipolar coupling (NOE effect). Correlation between protons that are close in space. This is the single most powerful NMR technique for determining the 3-dimensional structure of molecules from conformations of small molecules to the 3-dimensional structure of small proteins.

HOESY: Heteronuclear Overhauser Effect Spectroscopy. Correlation between protons and heteronuclei that are close in space.

Page 35

COSY experiment

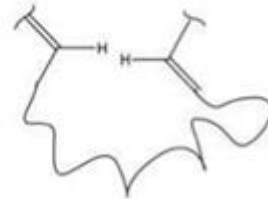
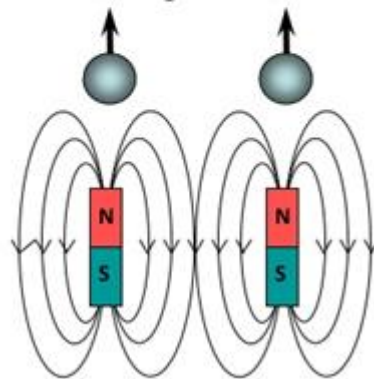
- Correlation Spectroscopy
- Commonly used for regiochemical assignment
- Cross-peaks appear if spin coupling is present
- Protons that are separated by 2 or 3 bonds are usually detected



Page 36

Nuclear Overhauser Effect

- NOE Caused by dipolar coupling between nuclei.
- The local field at one nucleus is affected by the presence of another nucleus. The result is a mutual modulation of resonance frequencies.



Page 37

Nuclear Overhauser Effect

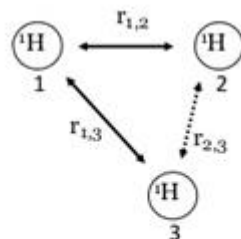
The intensity of the interaction is a function of the distance between the nuclei according to the following equation.

$$I = A (1/r^6)$$

I - intensity

A - scaling constant

r - internuclear distance



Arrows denote cross relaxation pathways

$r_{1,2}$ - distance between protons 1 and 2

$r_{2,3}$ - distance between protons 2 and 3

The NOE provides a link between an experimentally measurable quantity, I, and internuclear distance.

NOE is only observed up to 5 Å.

Page 38

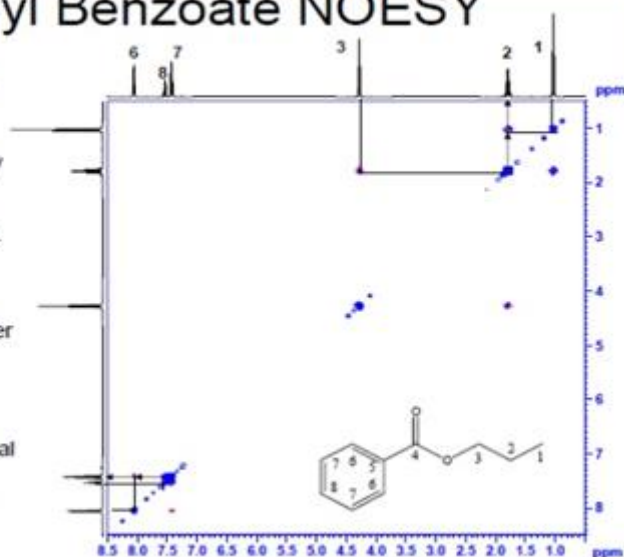
Propyl Benzoate NOESY

Nuclear Overhauser Effect Spectroscopy

Shows correlations to protons that are nearby in space to about 4 angstroms (most J coupling is filtered out).

Interpret by Noting which protons are close to one another (very useful for stereochemical analysis).

Protons that are geminal and vicinal will still appear because they are close.



Page 39

Contd...

- ✓ NMR is used in pharmaceutical science to study Pharmaceuticals and Drug metabolism.
- ✓ NMR is used in chemistry to ;
 - Determine the Enantiomeric purity.
 - Elucidate Chemical structure of organic and inorganic compounds.
 - Macromolecules –ligand interaction.

Page 41

¹H-NMR SPECTROSCOPY

applications

¹H widely used for structure elucidation.

Inorganic solids- Inorganic compounds are investigated by solid state ¹H-NMR. eg $\text{CaSO}_4 \cdot \text{H}_2\text{O}$.

Organic solids- Solid-state ¹H NMR constitutes a powerful approach to investigate the hydrogen-bonding and ionization states of small organic compounds.

- Direct correlation with hydrogen-bonding lengths could be demonstrated, e.g. for amino acid carboxyl groups.

Page 42

Polymers and rubbers- Examine hydrogen bonding and acidity.

Peptides and proteins

Clinical and scientific research

In vivo NMR studies-

- concerned with ¹H NMR spectroscopy of human brain.
- Many studies are concerned with altered levels of metabolites in various brain diseases.
- To determine the spatial distribution of any given metabolite detected spectroscopically IS (image selected *in vivo spectroscopy*).

Page 43

Application of NMR in medicine

MRI is specialist application of multi dimensional Fourier transformation NMR



- ✓ Anatomical imaging.
- ✓ Measuring physiological functions
- ✓ Flow measurements and angiography.
- ✓ Tissue perfusion studies.
- ✓ Tumors