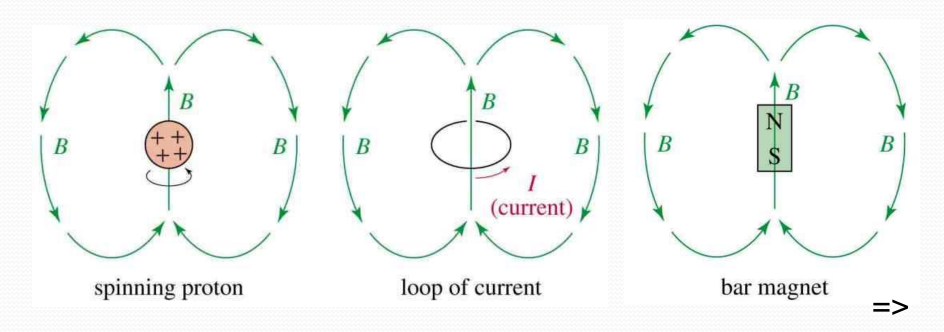
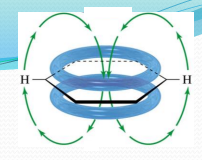
#### Introduction

- NMR is the most powerful tool available for organic structure determination.
- It is used to study a wide variety of nuclei:
  - <sup>1</sup>H
  - <sup>13</sup>C
  - <sup>15</sup>N
  - 19F
  - 31P

## Nuclear Spin

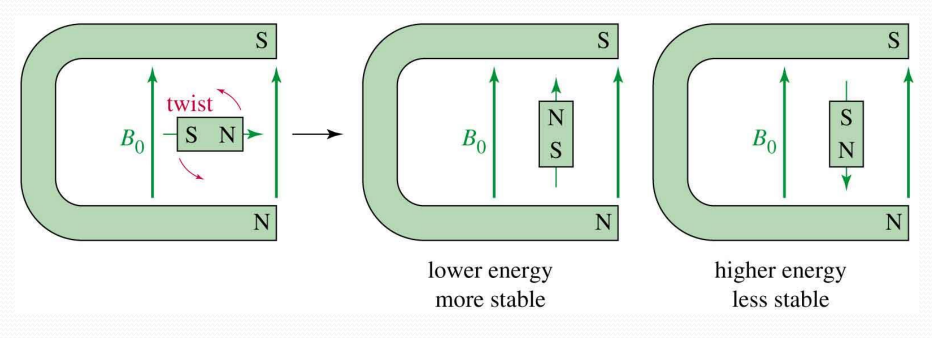
- A nucleus with an odd atomic number or an odd mass number has a nuclear spin.
- The spinning charged nucleus generates a magnetic field.

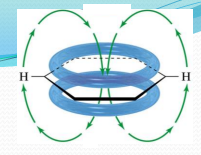




#### **External Magnetic Field**

# When placed in an external field, spinning protons act like bar magnets.

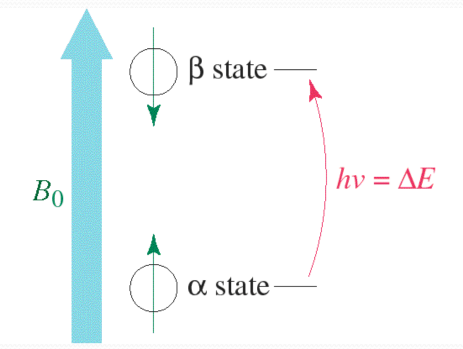




#### **Two Energy States**

The magnetic fields of the spinning nuclei will align either *with* the external field, or *against* the field.

A photon with the right amount of energy can be absorbed and cause the spinning proton to flip.



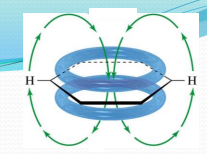
# $\Delta E$ and Magnet Strength

- Energy difference is proportional to the magnetic field strength.
- $\Delta E = h v = \gamma \underline{h} B_{o}$  $2\pi$
- Gyromagnetic ratio, γ, is a constant for each nucleus (26,753 s<sup>-1</sup>gauss<sup>-1</sup> for H).
- In a 14,092 gauss field, a 60 MHz photon is required to flip a proton.
- Low energy, radio frequency.

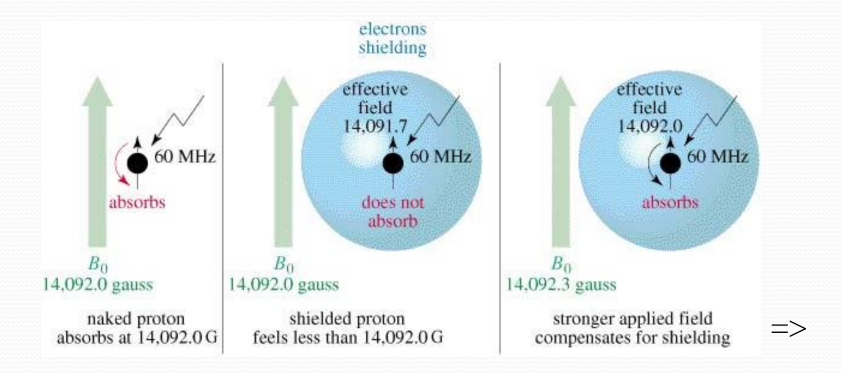
# Magnetic Shielding

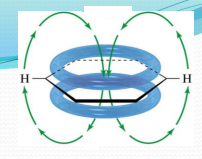
- If all protons absorbed the same amount of energy in a given magnetic field, not much information could be obtained.
- But protons are surrounded by electrons that shield them from the external field.
- Circulating electrons create an induced magnetic field that opposes the external magnetic field.

#### **Shielded Protons**



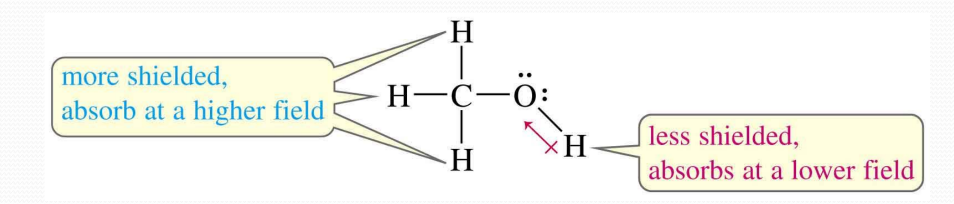
# Magnetic field strength must be increased for a shielded proton to flip at the same frequency.





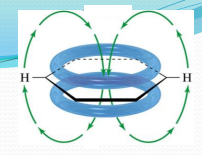
#### Protons in a Molecule

#### Depending on their chemical environment, protons in a molecule are shielded by different amounts.

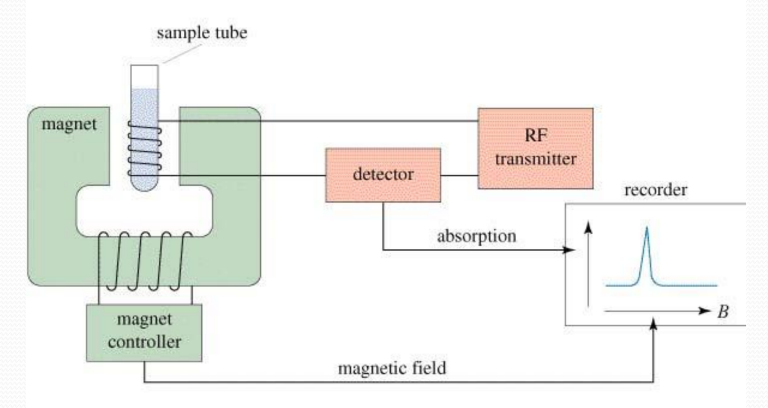


#### **NMR Signals**

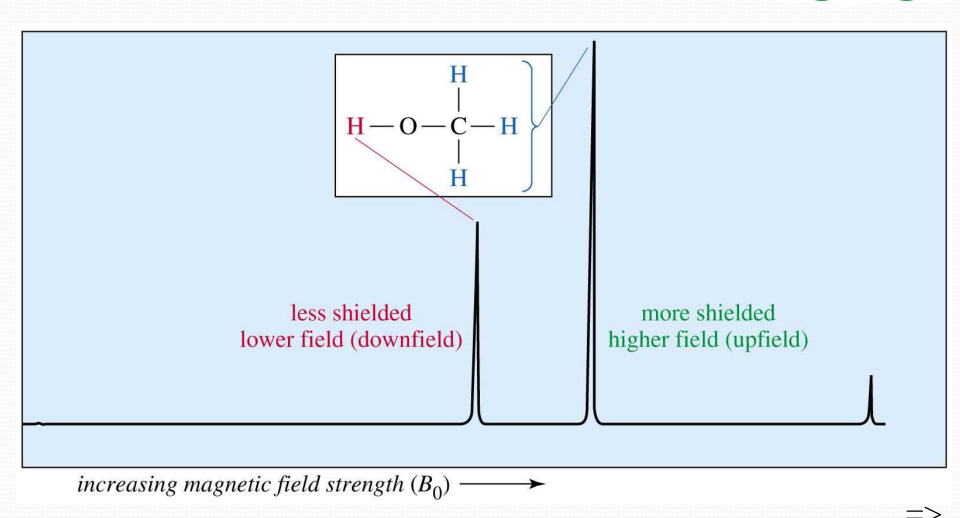
- The *number* of signals shows how many different kinds of protons are present.
- The *location* of the signals shows how shielded or deshielded the proton is.
- The *intensity* of the signal shows the number of protons of that type.
- Signal *splitting* shows the number of protons on adjacent atoms.



#### The NMR Spectrometer







#### СH<sub>3</sub> H<sub>3</sub>C\_Si-CH<sub>3</sub> Tetramethylsilane

- TMS is added to the sample.
- Since silicon is less electronegative than carbon, TMS protons are highly shielded. Signal defined as zero.
- Organic protons absorb downfield (to the left) of the TMS signal.



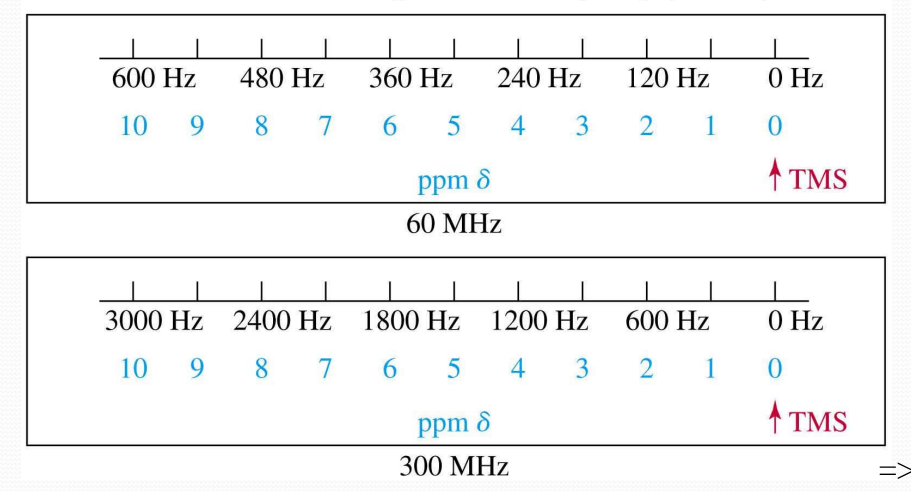
# **Chemical Shift**

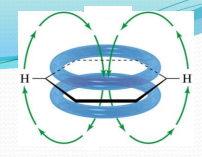
- Measured in parts per million.
- Ratio of shift downfield from TMS (Hz) to total spectrometer frequency (Hz).
- Same value for 60, 100, or 300 MHz machine.
- Called the delta scale.



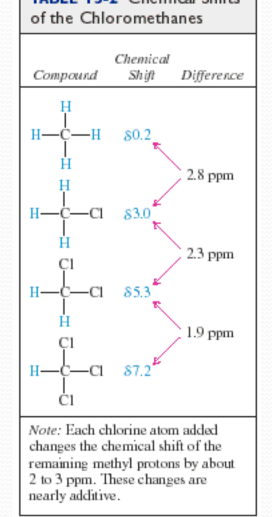
#### **Delta Scale**

chemical shift, ppm  $\delta = \frac{\text{shift downfield from TMS (in Hz)}}{\text{spectrometer frequency (in MHz)}}$ 





# Location of Signals



- More electronegative atoms deshield more and give larger shift values.
- Effect decreases with distance.
- Additional electronegative atoms cause increase in chemical shift.

## **Typical Values**

| Type of Proton  | Approximate $\delta$ | Type of Proton    | Approximate $\delta$  |
|---|----------------------|-------------------|-----------------------|
| alkane ( $-CH_3$ )  | 0.9                  | >c=c              | 1.7                   |
| alkane (—CH <sub>2</sub> —)                                     | 1.3                  | CH <sub>3</sub>   |                       |
| alkane $\begin{pmatrix}\mathbf{C}\mathbf{H} \\ - \end{pmatrix}$ | 1.4                  | Ph—H              | 7.2                   |
|   |                      | $Ph-CH_3$         | 2.3                   |
| O<br>II   |                      | R—CHO             | 9–10                  |
|   | 2.1                  | R—COOH            | 10-12                 |
| $-C \equiv C - H$   | 2.5                  | R—OH              | variable, about 2–5   |
| $R-CH_2-X$  | 3-4                  | Ar—OH             | variable, about 4–7   |
| (X = halogen, O)  |                      | R—NH <sub>2</sub> | variable, about 1.5–4 |
| ≥c=c< <sub>H</sub>  | 5-6                  |                   |                       |

*Note:* These values are approximate, as all chemical shifts are affected by neighboring substituents. The numbers given here assume that alkyl groups are the only other substituents present. A more complete table of chemical shifts appears in Appendix 1.

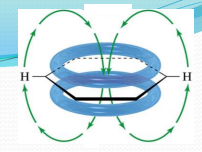
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#### Aromatic Protons, $\delta$ 7- $\delta$ 8

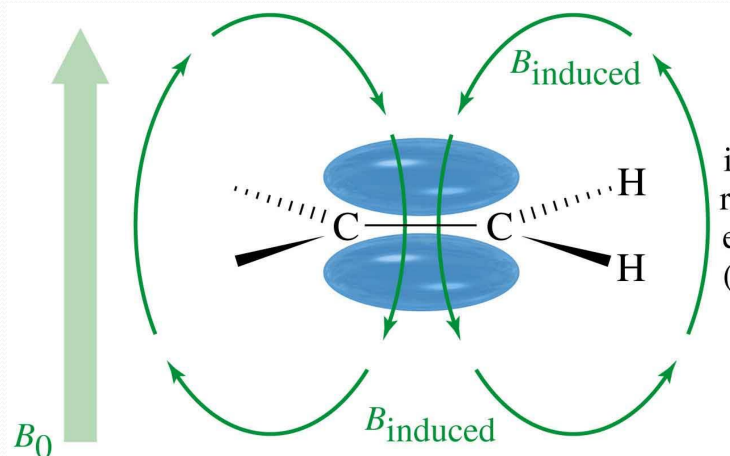
circulation of electrons (ring current) H Η *B*<sub>induced</sub> induced magnetic field

induced field reinforces the external field (deshielding)

 $B_0$ 

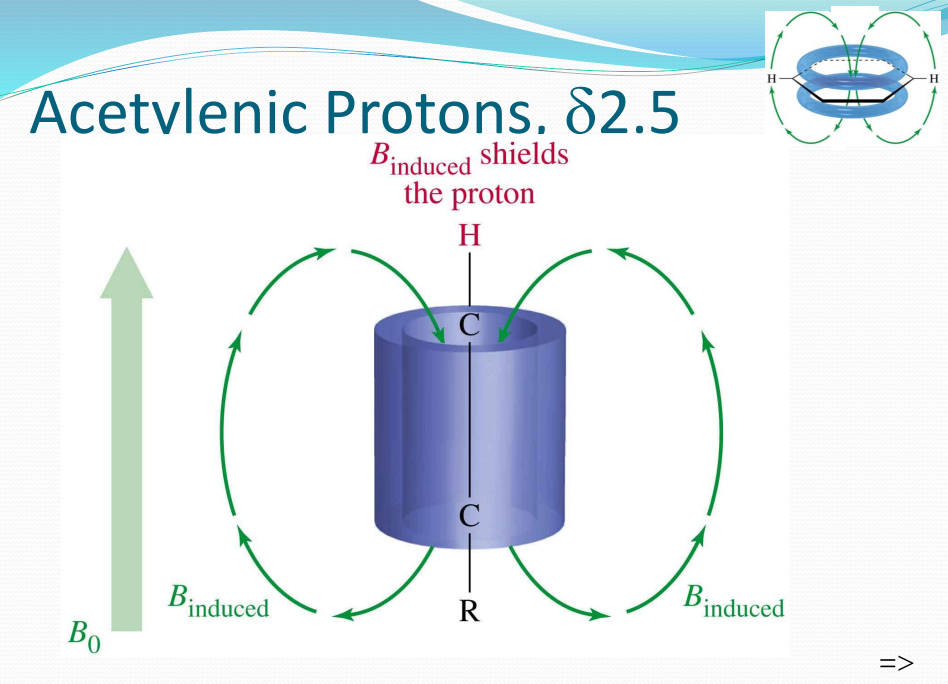


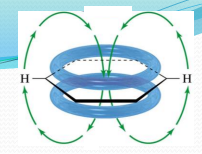
#### Vinyl Protons, $\delta 5-\delta 6$



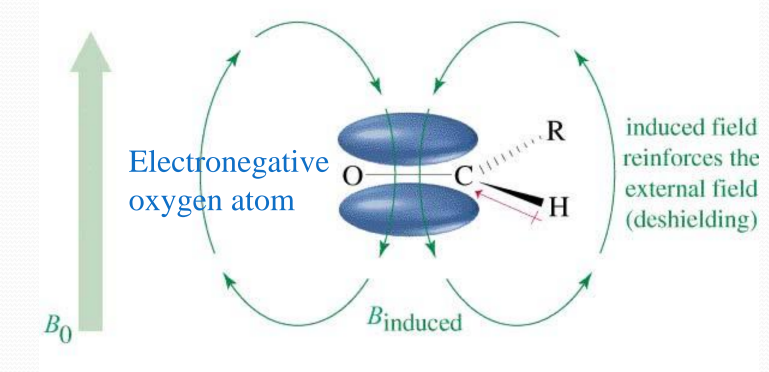
induced field reinforces the external field (deshielding)

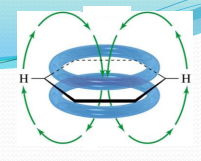
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#### Aldehyde Proton, $\delta 9-\delta 10$

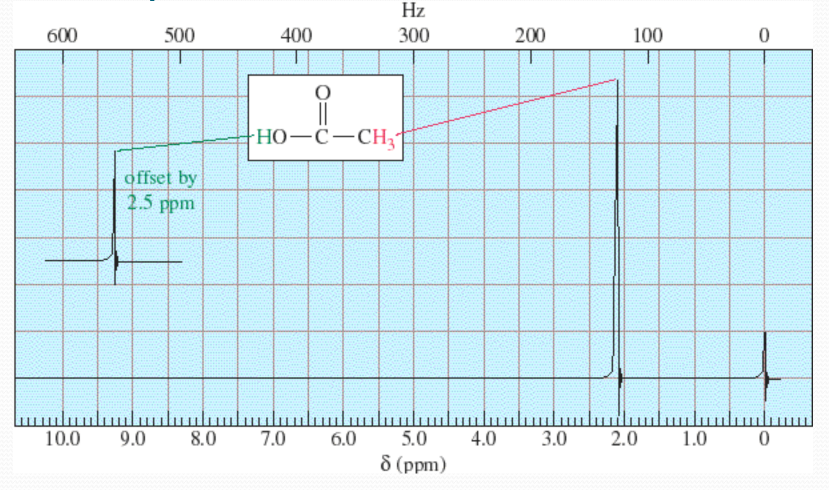




#### **O-H and N-H Signals**

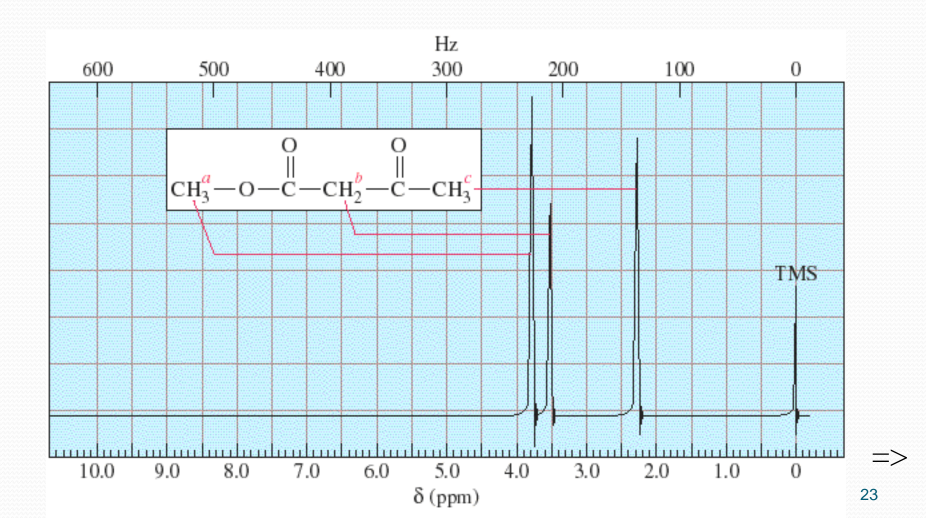
- Chemical shift depends on concentration.
- Hydrogen bonding in concentrated solutions deshield the protons, so signal is around δ3.5 for N-H and δ4.5 for O-H.
- Proton exchanges between the molecules broaden the peak.

# Carboxylic Acid Proton, $\delta$ 10+



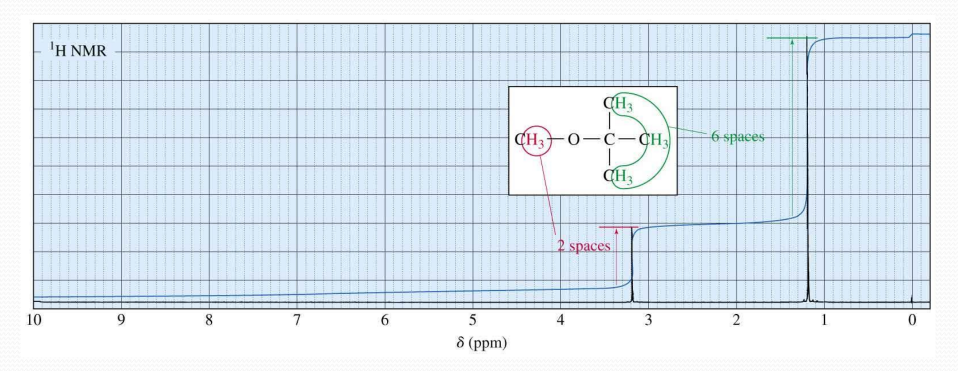
## Number of Signals

Equivalent hydrogens have the same chemical shift.



#### Intensity of Signals

- The area under each peak is proportional to the number of protons.
- Shown by integral trace.



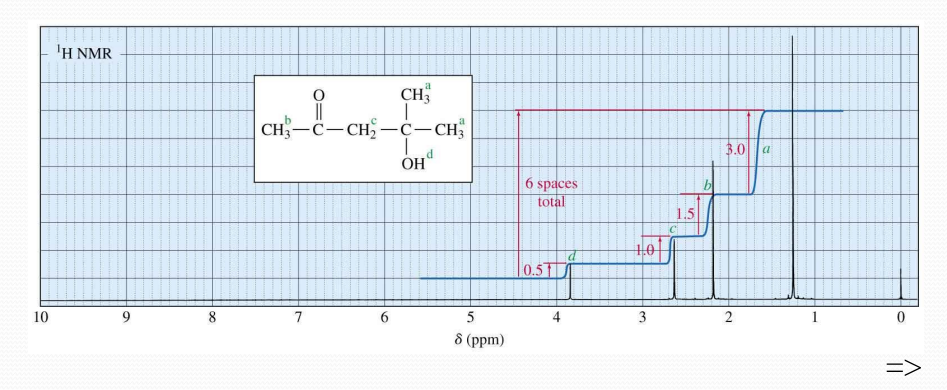
Chapter 13

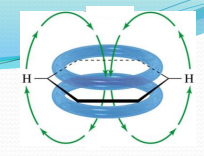
24

# H

#### How Many Hydrogens?

When the molecular formula is known, each integral rise can be assigned to a particular number of hydrogens.



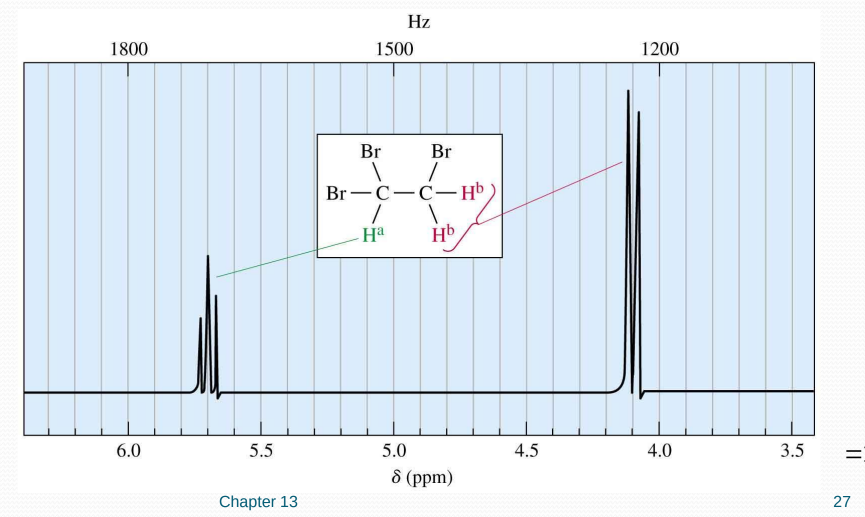


## **Spin-Spin Splitting**

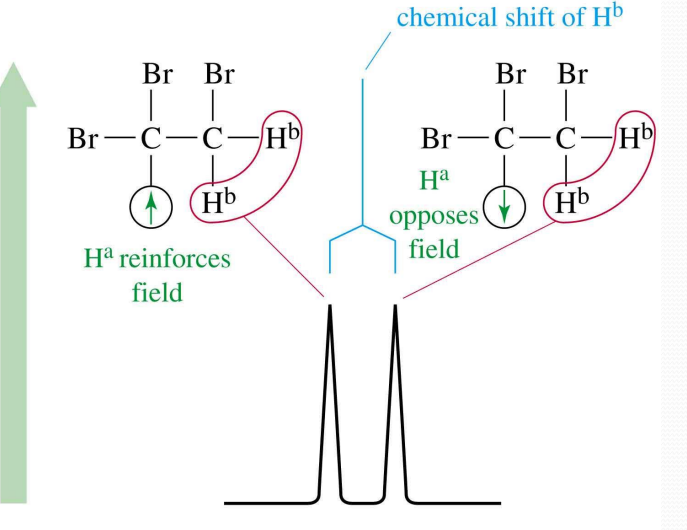
- Nonequivalent protons on adjacent carbons have magnetic fields that may align with or oppose the external field.
- This magnetic coupling causes the proton to absorb slightly downfield when the external field is reinforced and slightly upfield when the external field is opposed.
- All possibilities exist, so signal is split. =>

#### 1,1,2-Tribromoethane

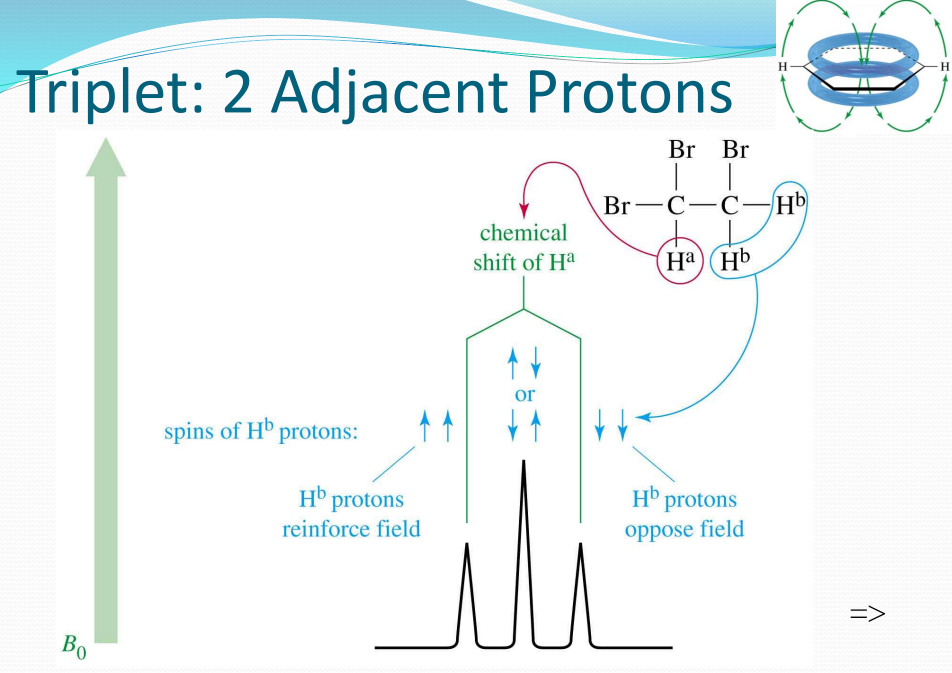
#### Nonequivalent protons on adjacent carbons.



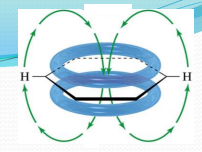
## Doublet: 1 Adjacent Proton



 $B_0$ 



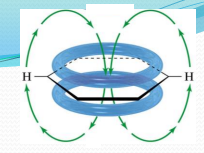
#### The N + 1 Rule



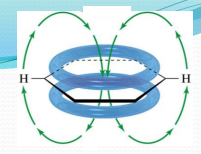
#### If a signal is split by N equivalent protons, it is split into N + 1 peaks.

| Number of Equivalent<br>Protons Causing Splitting | Number of Peaks<br>(multiplicity) | Area Ratios<br>(Pascal's triangle) |
|---|-----------------------------------|------------------------------------|
| 0   | 1 (singlet)                       | 1                                  |
| 1   | 2(doublet)                        | 1 1                                |
| 2   | 3 (triplet)                       | 1 2 1                              |
| 3   | 4 (quartet)                       | 1 3 3 1                            |
| 4   | 5 (quintet)                       | 14641                              |
| 5   | 6 (sextet)                        | 1 5 10 10 5 1                      |
| 6   | 7 (septet)                        | 1 6 15 20 15 6                     |

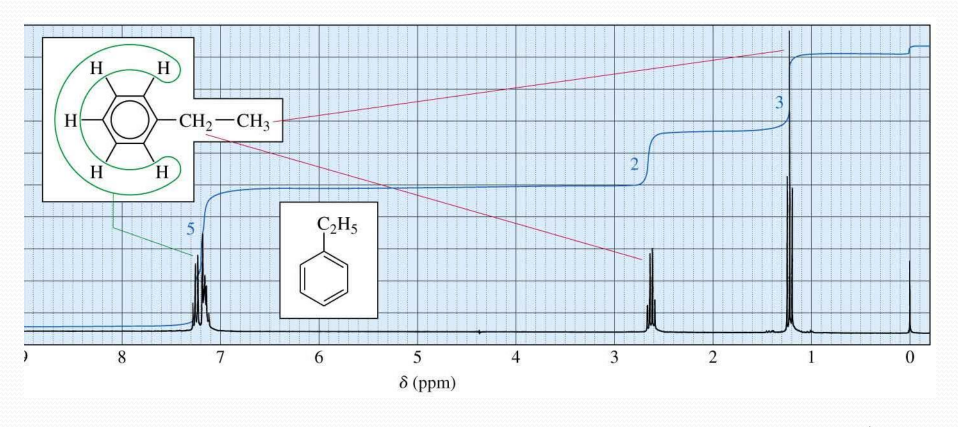
# Range of Magnetic Coupling



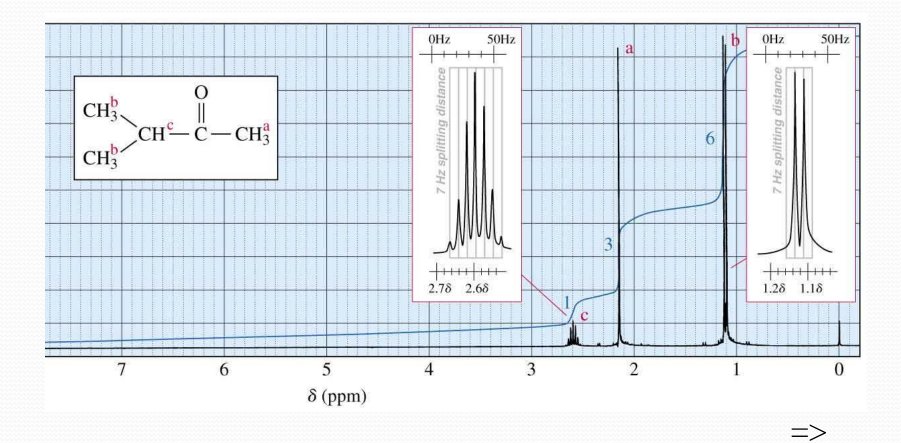
- Equivalent protons do not split each other.
- Protons bonded to the same carbon will split each other only if they are not equivalent.
- Protons on adjacent carbons normally will couple.
- Protons separated by four or more bonds will not couple.



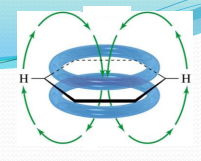
# Splitting for Ethyl Groups



# Splitting for Isopropyl Groups



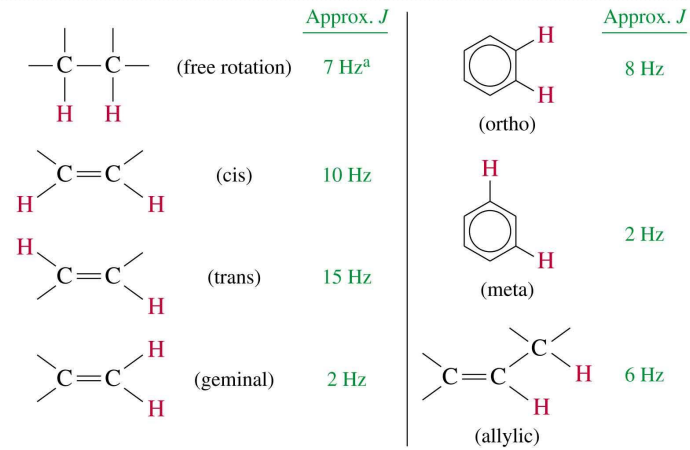
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#### **Coupling Constants**

- Distance between the peaks of multiplet
- Measured in Hz
- Not dependent on strength of the external field
- Multiplets with the same coupling constants may come from adjacent groups of protons that split each other.

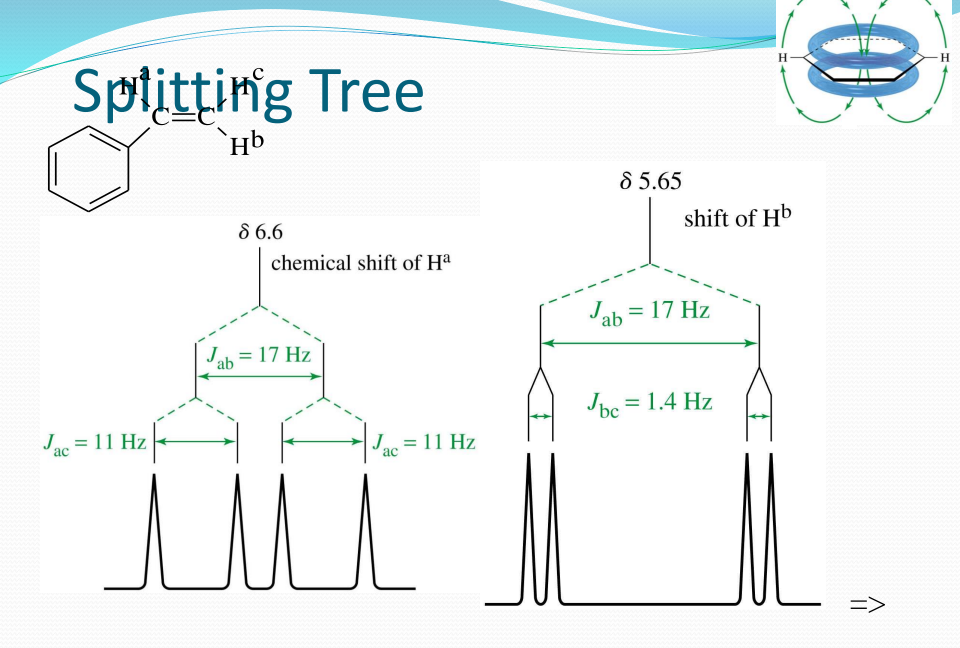
# Values for Coupling Constants

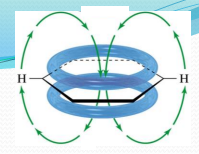


<sup>a</sup>The value of 7 Hz in an alkyl group is averaged for rapid rotation about the carbon–carbon bond. If rotation is hindered by a ring or bulky groups, other splitting constants may be observed.

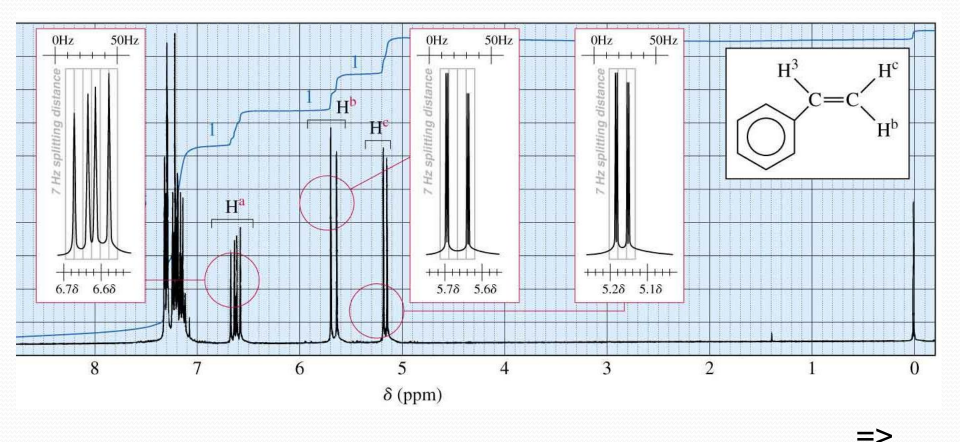


- Signals may be split by adjacent protons, different from each other, with different coupling constants.
- Example: H<sup>a</sup> of styrene which is split by an adjacent H *trans* to it (J = 17 Hz) and an adjacent H *cis* to it (J = 11 Hz).





# **Spectrum for Styrene**



#### Stereochemical

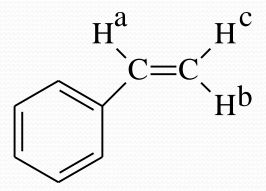
# H

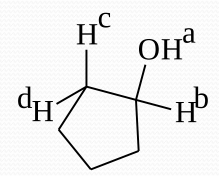
## Nonequivalence

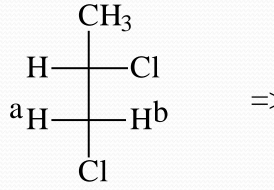
- Usually, two protons on the same C are equivalent and do not split each other.
- If the replacement of each of the protons of a -CH<sub>2</sub> group with an imaginary "Z" gives stereoisomers, then the protons are non-equivalent and will split each other.

# Some Nonequivalent

# Protons

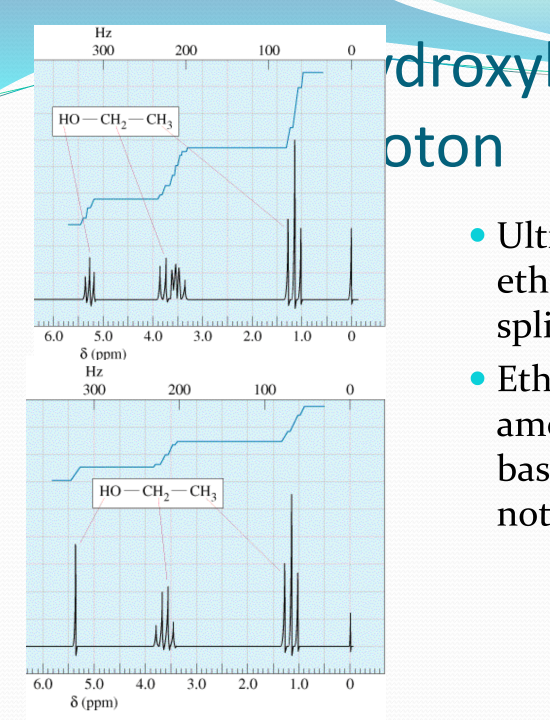






#### Time Dependence

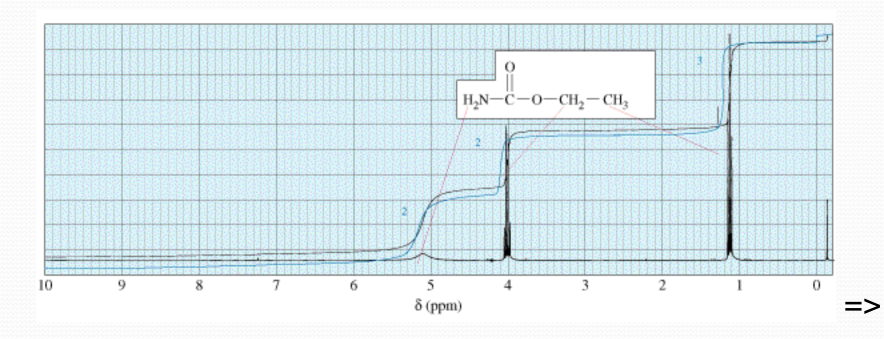
- Molecules are tumbling relative to the magnetic field, so NMR is an averaged spectrum of all the orientations.
- Axial and equatorial protons on cyclohexane interconvert so rapidly that they give a single signal.
- Proton transfers for OH and NH may occur so quickly that the proton is not split by adjacent protons in the molecule.



- Ultrapure samples of ethanol show splitting.
- Ethanol with a small amount of acidic or basic impurities will not show splitting.

#### **N-H Proton**

- Moderate rate of exchange.
- Peak may be broad.



# Identifying the O-H

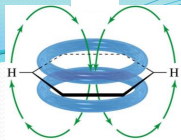
#### or N-H Peak

- Chemical shift will depend on concentration and solvent.
- To verify that a particular peak is due to O-H or N-H, shake the sample with D<sub>2</sub>O
- Deuterium will exchange with the O-H or N-H protons.
- On a second NMR spectrum the peak will be absent, or much less intense.

## Carbon-13

- <sup>12</sup>C has no magnetic spin.
- <sup>13</sup>C has a magnetic spin, but is only 1% of the carbon in a sample.
- The gyromagnetic ratio of <sup>13</sup>C is one-fourth of that of <sup>1</sup>H.

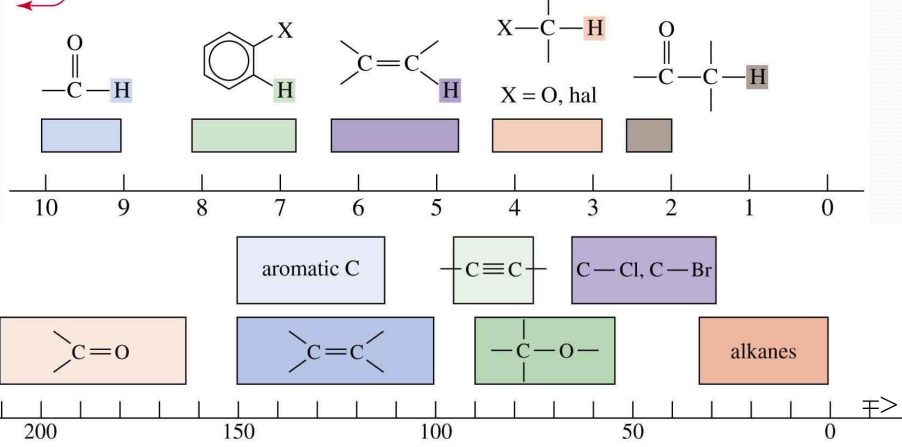
- Signals are weak, getting lost in noise.
- Hundreds of spectra are taken, averaged.



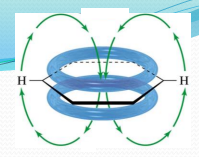
# Fourier Transform NMR

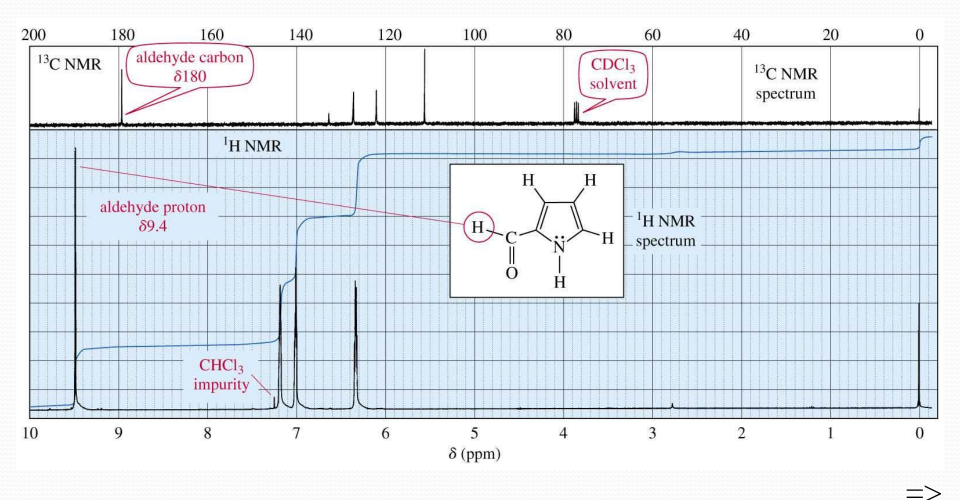
- Nuclei in a magnetic field are given a radiofrequency pulse close to their resonance frequency.
- The nuclei absorb energy and precess (spin) like little tops.
- A complex signal is produced, then decays as the nuclei lose energy.
- Free induction decay is converted to spectrum.

#### Hydrogen and Carbon Chemical Shifts -cooH all-al2



# Combined <sup>13</sup>C and <sup>1</sup>H Spectra





# Differences in

# <sup>13</sup>C Technique

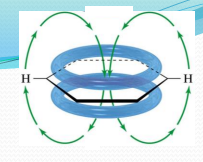
- Resonance frequency is ~ one-fourth, 15.1 MHz instead of 60 MHz.
- Peak areas are not proportional to number of carbons.
- Carbon atoms with more hydrogens absorb more strongly.

# Spin-Spin Splitting

- It is unlikely that a <sup>13</sup>C would be adjacent to another <sup>13</sup>C, so splitting by carbon is negligible.
- <sup>13</sup>C <u>will</u> magnetically couple with attached protons and adjacent protons.

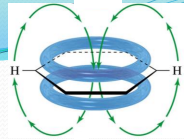
=>

• These complex splitting patterns are difficult to interpret.



#### **Proton Spin Decoupling**

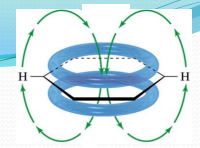
- To simplify the spectrum, protons are continuously irradiated with "noise," so they are rapidly flipping.
- The carbon nuclei see an average of all the possible proton spin states.
- Thus, each different kind of carbon gives a single, unsplit peak.



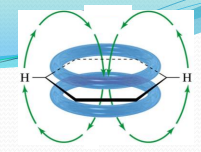
## **Off-Resonance Decoupling**

- <sup>13</sup>C nuclei are split only by the protons attached directly to them.
- The N + 1 rule applies: a carbon with N number of protons gives a signal with N + 1 peaks.

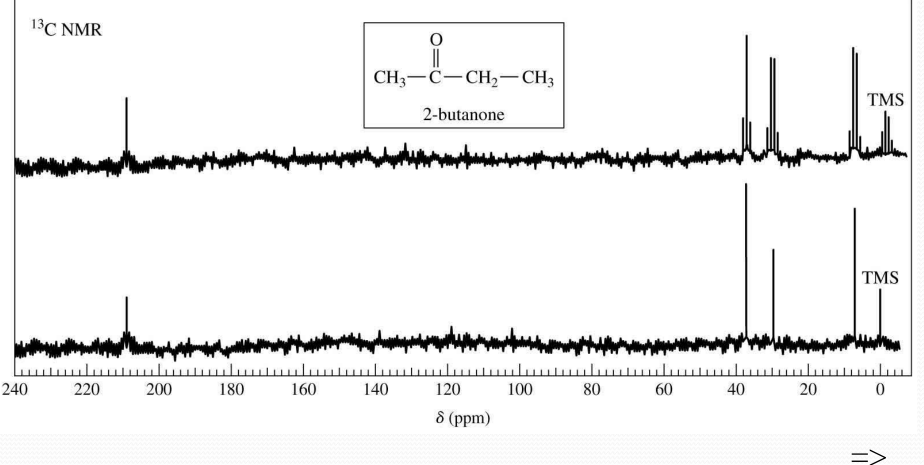
# Interpreting <sup>13</sup>C NMR



- The number of different signals indicates the number of different kinds of carbon.
- The location (chemical shift) indicates the type of functional group.
- The peak area indicates the numbers of carbons (if integrated).
- The splitting pattern of off-resonance decoupled spectrum indicates the number of protons attached to the carbon. =>

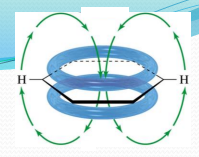


#### Two <sup>13</sup>C NMR Spectra



# MRI

- Magnetic resonance imaging, noninvasive
- "Nuclear" is omitted because of public's fear that it would be radioactive.
- Only protons in one plane can be in resonance at one time.
- Computer puts together "slices" to get 3D.
- Tumors readily detected.



# End of Chapter 13