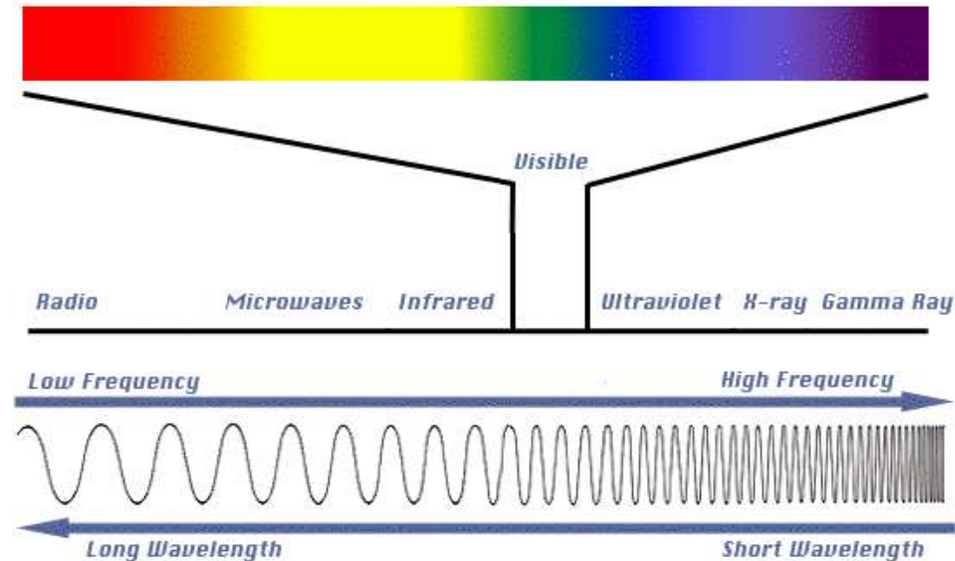


Nuclear Magnetic Resonance Spectroscopy



NMR Spectroscopy

NMR spectroscopy is a form of absorption spectrometry.



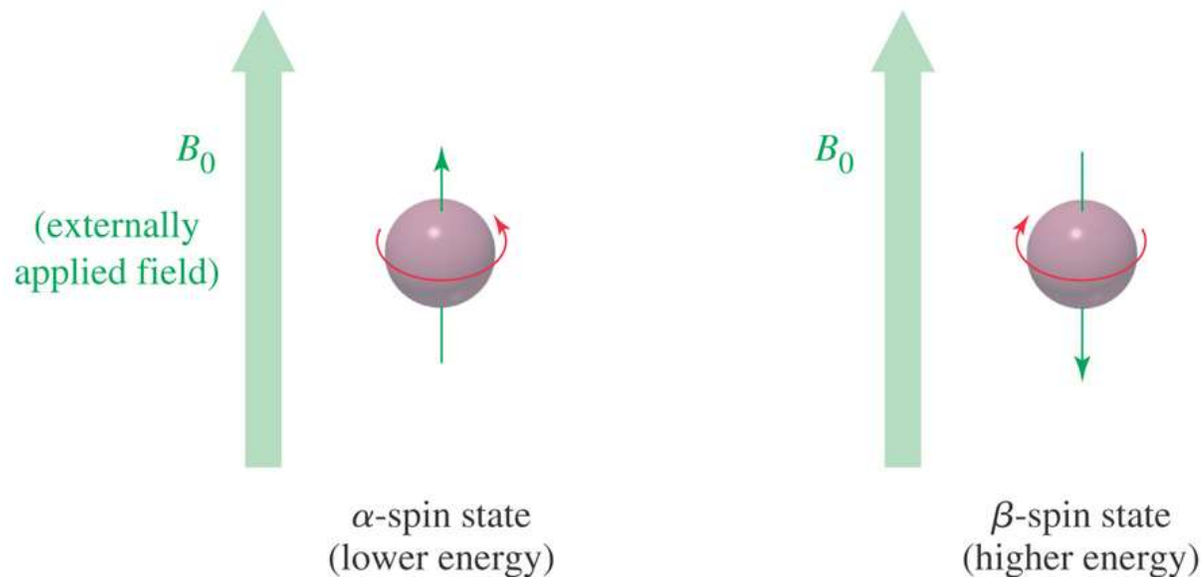
Most absorption techniques (e.g. – Ultraviolet-Visible and Infrared) involve the electrons... in the case of NMR, it is the nucleus of the atom which determines the response.

An applied (magnetic) field is necessary for the absorption to occur.

Nuclear Magnetic Resonance (NMR) Spectroscopy

direct observation of the H's and C's of a molecules

Nuclei are positively charged and spin on an axis; they create a tiny magnetic field



Not all nuclei are suitable for NMR.

^1H and ^{13}C are the most important NMR active nuclei in organic chemistry

Natural Abundance

^1H 99.9%

^{13}C 1.1%

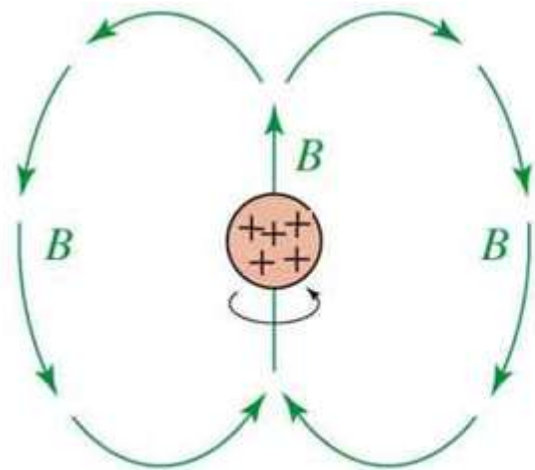
^{12}C 98.9% (not NMR active)

Spectral Properties, Application and Interactions of Electromagnetic Radiation

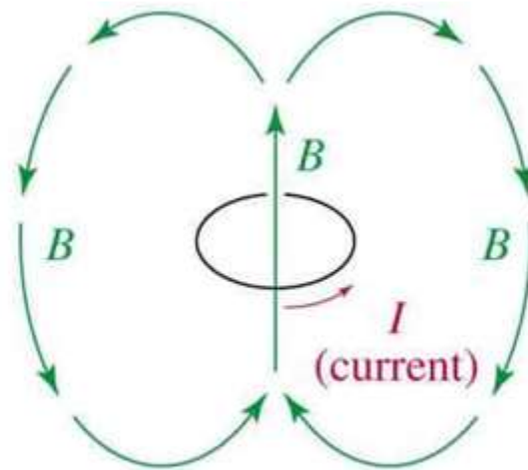
Energy	Wave Number ν	Wavelength λ	Frequency ν	Type Radiation	Type spectroscopy	Type Quantum Transition	
Kcal/mol	Electron volts, eV	cm^{-1}	cm	Hz			
9.4×10^7	4.9×10^6	3.3×10^{10}	3×10^{-11}	10^{21}	Gamma ray	Gamma ray emission	Nuclear
9.4×10^3	4.9×10^2	3.3×10^6	3×10^{-7}	10^{17}	X-ray	X-ray absorption, emission	Electronic (inner shell)
9.4×10^1	4.9×10^0	3.3×10^4	3×10^{-5}	10^{15}	Ultra violet	UV absorption	Electronic (outer shell)
9.4×10^{-1}	4.9×10^{-2}	3.3×10^2	3×10^{-3}	10^{13}	Infrared	IR absorption	Molecular vibration
9.4×10^{-3}	4.9×10^{-4}	3.3×10^0	3×10^{-1}	10^{11}	Micro-wave	Microwave absorption	Molecular rotation
9.4×10^{-7}	4.9×10^{-8}	3.3×10^{-4}	3×10^3	10^7	Radio	Nuclear magnetic resonance	Magnetically induced spin states

Nuclear Spin

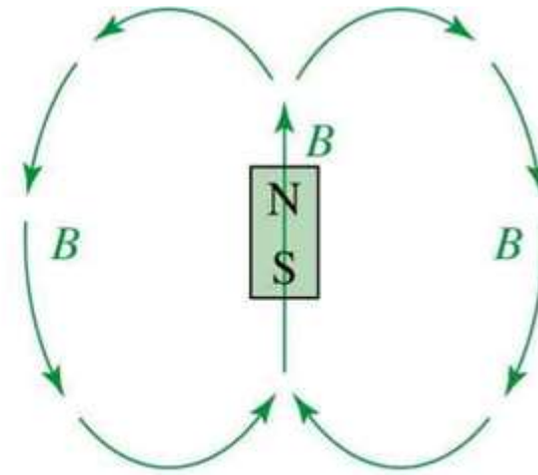
- A nucleus with an odd atomic number or an odd mass number has a nuclear spin.
- The spinning charged nucleus generates **a magnetic field**.



spinning proton



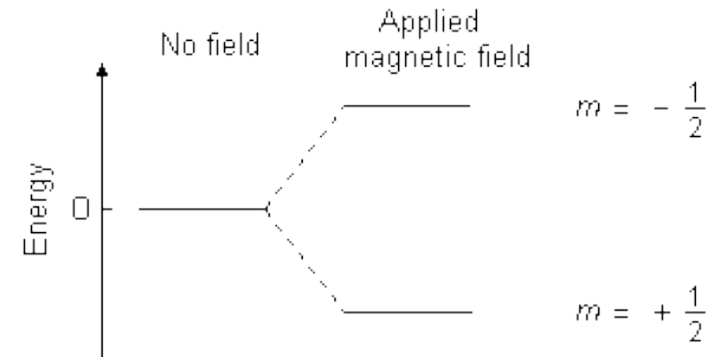
loop of current



bar magnet

Spin Quantum Numbers

Energy levels for a nucleus with spin quantum number $1/2$



The number of spin states is $2I + 1$,
where I is the spin quantum number.

- If the number of neutrons **and** the number of protons are both even, then the nucleus has **NO** spin.
- If the number of neutrons **plus** the number of protons is odd, then the nucleus has a half-integer spin (i.e. $1/2, 3/2, 5/2$)
- If the number of neutrons **and** the number of protons are both odd, then the nucleus has an integer spin (i.e. 1, 2, 3)

SPIN QUANTUM NUMBERS OF SOME COMMON NUCLEI

The most abundant isotopes of C and O do not have spin.

Element	¹ H <small>P=1, N=0</small>	² H <small>P=1, N=1</small>	¹² C <small>P=6, N=6</small>	¹³ C <small>P=6, N=7</small>	¹⁴ N <small>P=7, N=7</small>	¹⁶ O <small>P=8, N=8</small>	¹⁹ F <small>P=9, N=10</small>
Nuclear Spin Quantum No (I)	1/2	1	0	1/2	1	0	1/2
No. of Spin States	2	3	0	2	3	0	2

Elements with either odd number of protons or neutrons (odd mass number or odd atomic number) have the property of nuclear “spin”.

The number of spin states is $2I + 1$, where I is the spin quantum number.

Table. General rules for determination of nuclear spin quantum numbers

Mass Number	Number of Protons	Number of Neutrons	Spin (I)	Example
Even	Even	Even	0	^{16}O
	Odd	Odd	Integer (1,2,...)	^2H
Odd	Even	Odd	Half-Integer (1/2, 3/2,...)	^{13}C
	Odd	Even	Half-Integer (1/2, 3/2,...)	^{15}N

Atoms active in NMR

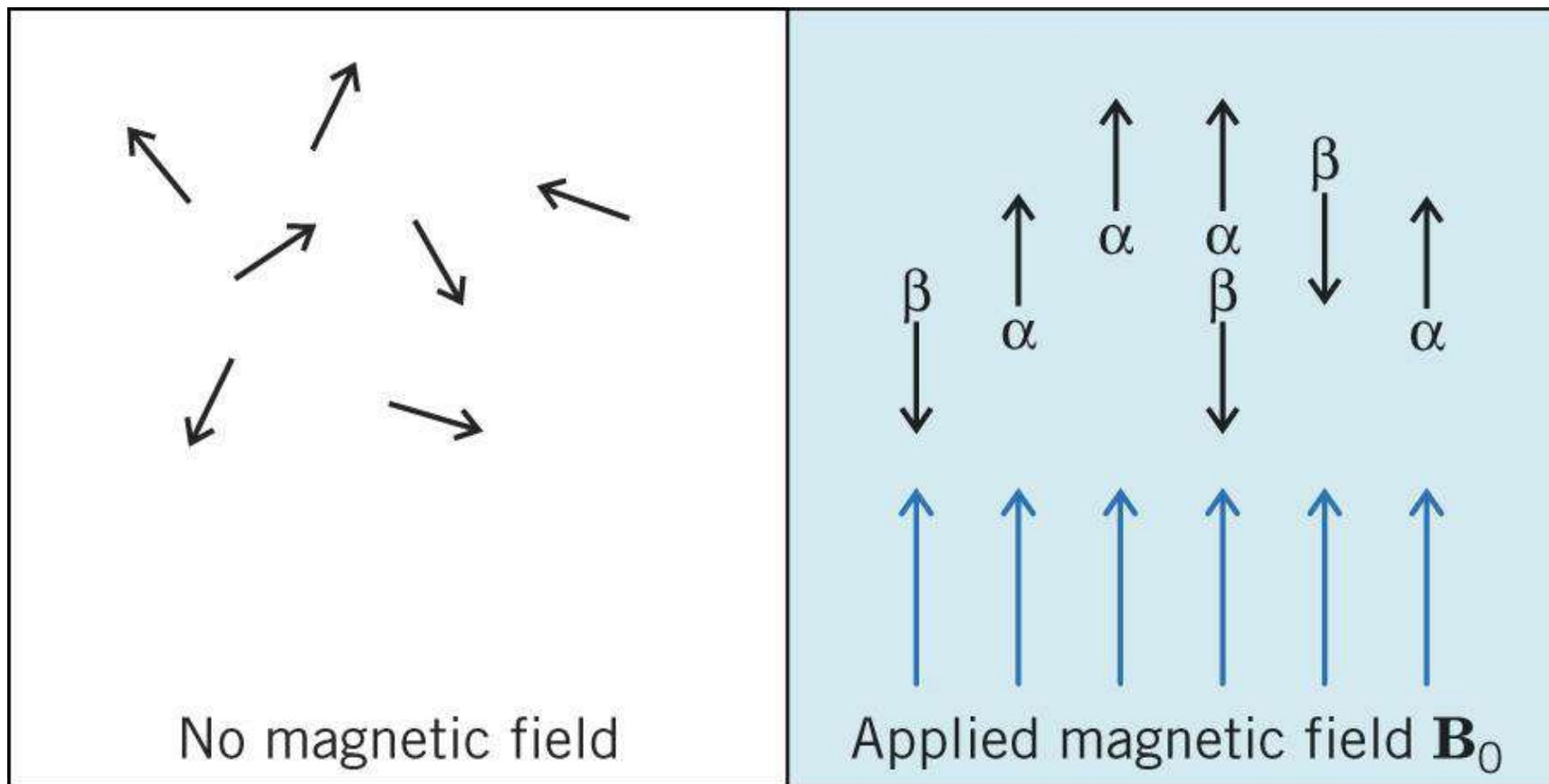
$I = 1/2$: ^1H , ^{13}C , ^{19}F , ^{31}P

$I = 1$: ^2H , ^{14}N

$I = 3/2$: ^{15}N

Atoms active in NMR

$I = 0$: ^{12}C , ^{16}O



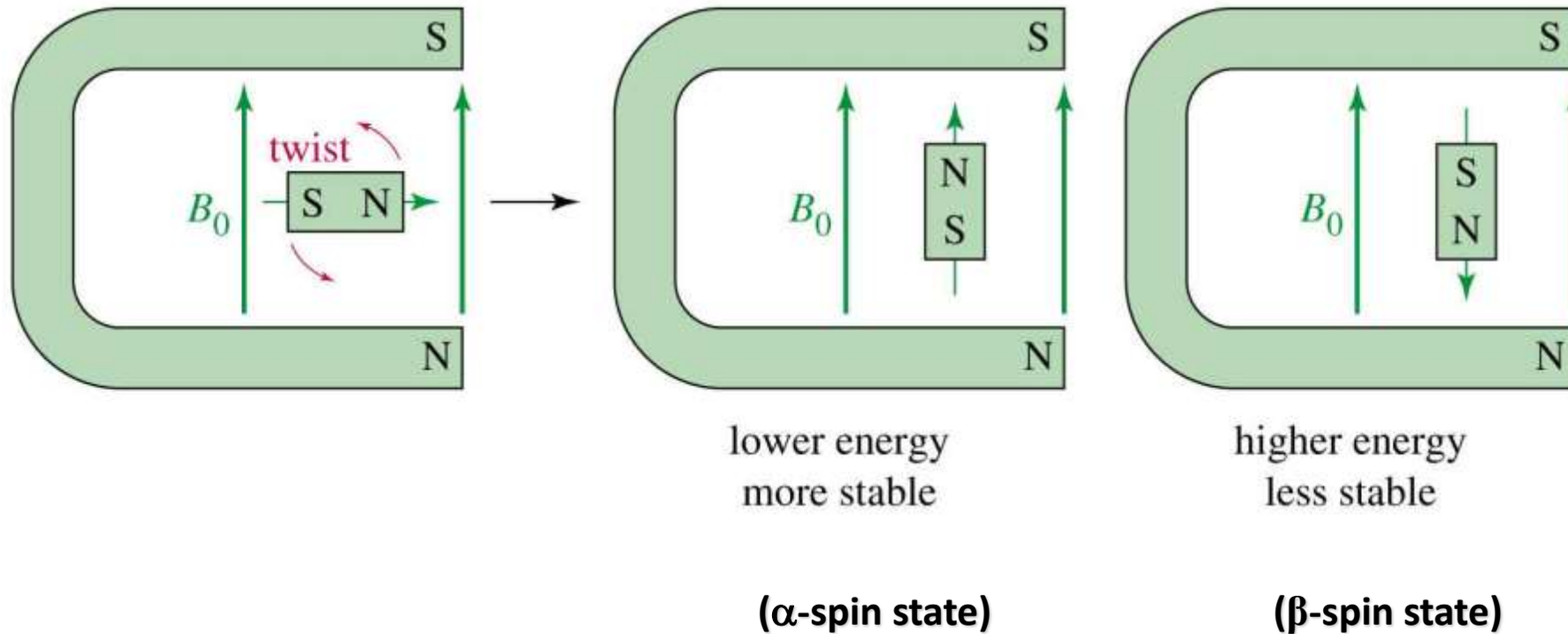
No magnetic field

Applied magnetic field \mathbf{B}_0

(a)

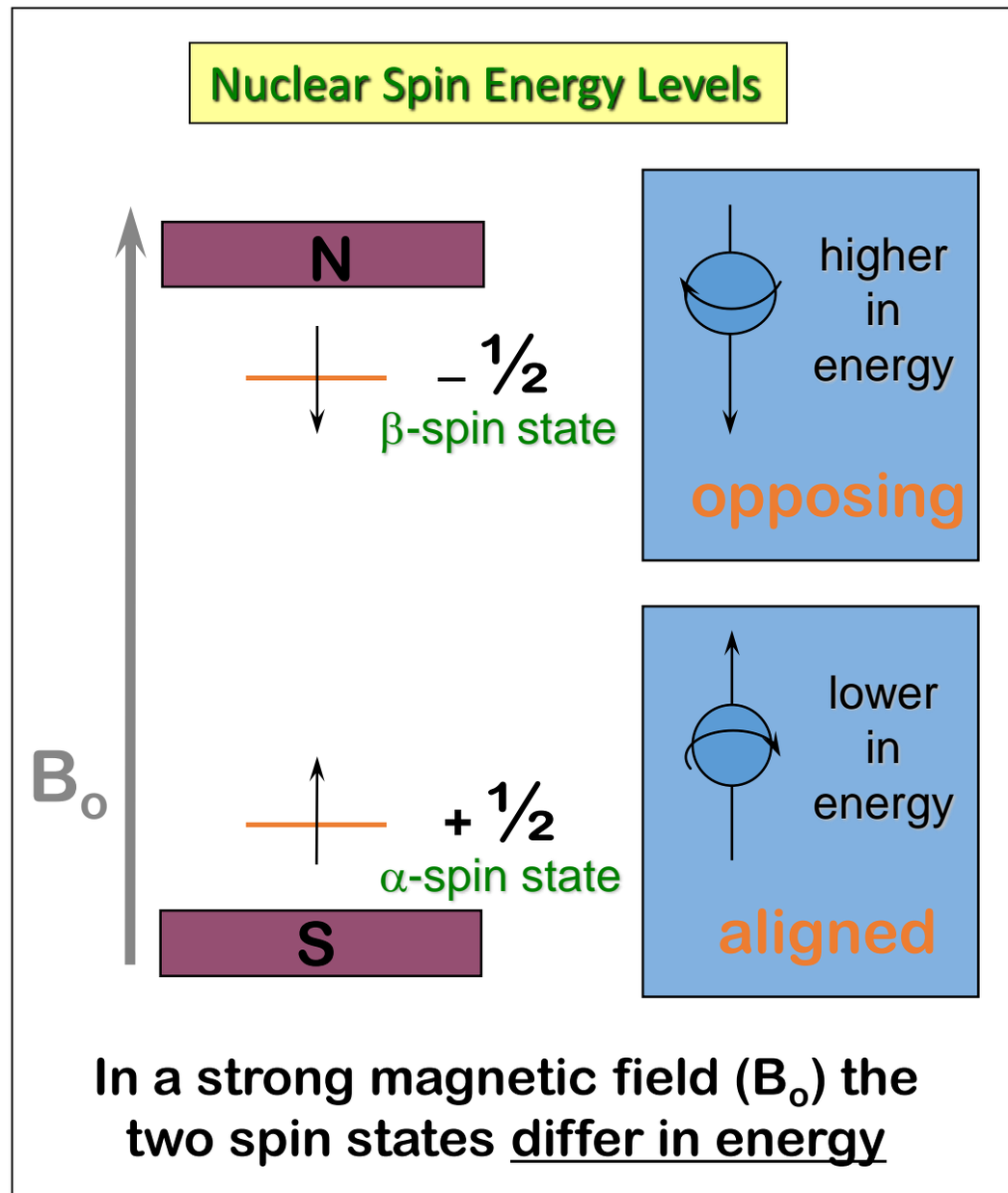
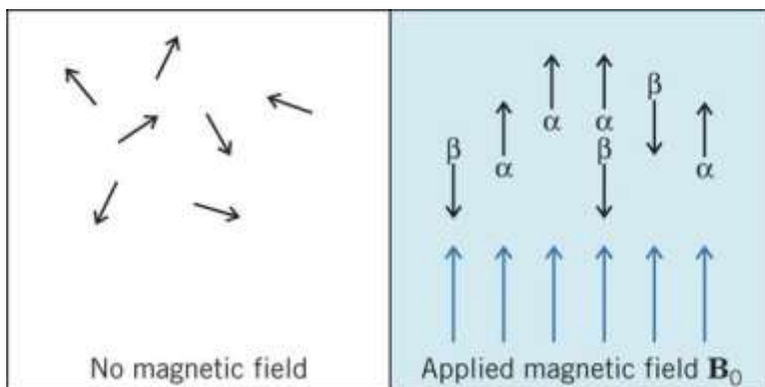
(b)

Behavior of spinning protons with external magnetic field



THE NUCLEUS IN A MAGNETIC FIELD

When nuclei are exposed to external magnetic field of strength B_0 , their spins line up parallel to the applied field, either spin aligned (α -spin state) or spin opposed (β -spin state) to the external field.



Energy of spinning nuclei

The energy of the nucleus in these two states (orientations) is given by:

$$E = -\frac{\gamma m h}{2\pi} B_0$$

**Energy of α -state
(spin aligned)**

$$E_{+1/2} = -\frac{\gamma(+\frac{1}{2})h}{2\pi} B_0$$

$$E_{+1/2} = -\frac{\gamma h}{4\pi} B_0$$

**Energy of β -state
(spin opposed)**

$$E_{-1/2} = -\frac{\gamma(-\frac{1}{2})h}{2\pi} B_0$$

$$E_{-1/2} = \frac{\gamma h}{4\pi} B_0$$

**Energy difference
between the two
state**

$$\Delta E = \frac{\gamma h}{4\pi} B_0 - \left(-\frac{\gamma h}{4\pi} B_0\right) = \frac{\gamma h}{2\pi} B_0$$

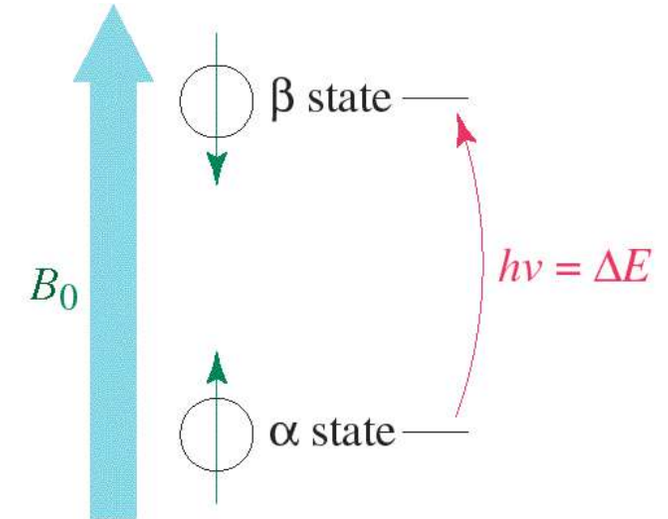
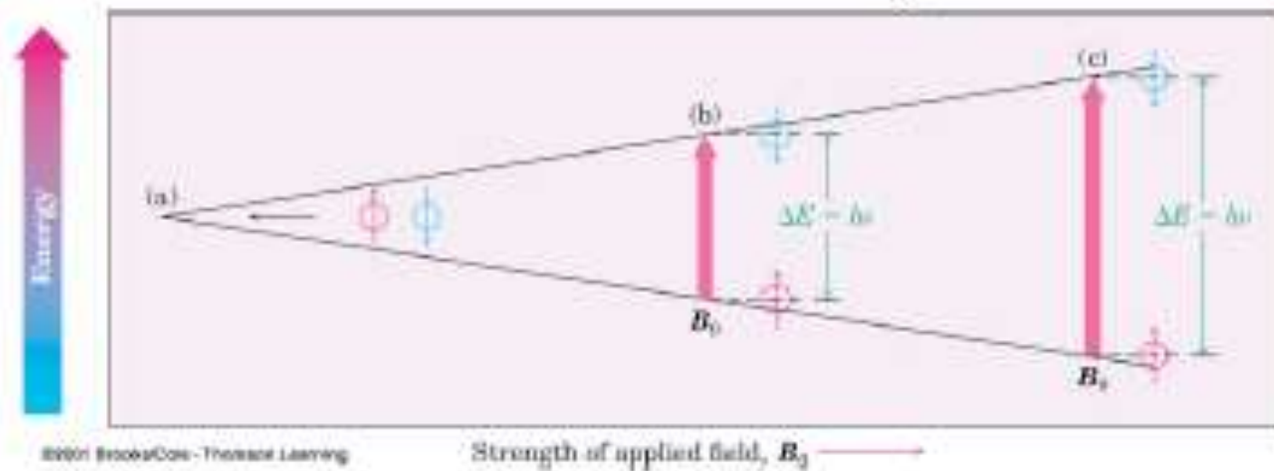
Absorption of electromagnetic radiation of frequency ν that correspond to in energy to ΔE

$$\Delta E = \cancel{h\nu} = \frac{\cancel{\gamma h}}{2\pi} B_0$$

ΔE and Magnet Strength

The energy difference between aligned and opposed to the external magnetic field (B_0) is generally small and is dependent upon B_0

Applied EM radiation (radio waves) causes the spin to flip and the nuclei are said to be in **resonance** with B_0



$$\Delta E = h\nu = \frac{\gamma}{2\pi} B_0$$

B_0 = external magnetic field strength

γ = gyromagnetic ratio

$^1\text{H} = 26,752 \text{ s}^{-1}\text{gauss}^{-1}$

$^{13}\text{C} = 6.7 \text{ s}^{-1}\text{gauss}^{-1}$

. In a 14,092 gauss field, a 60 MHz (60,000,000) photon is required to flip a proton.

. Low energy, radio frequency.

Magnetic Shielding

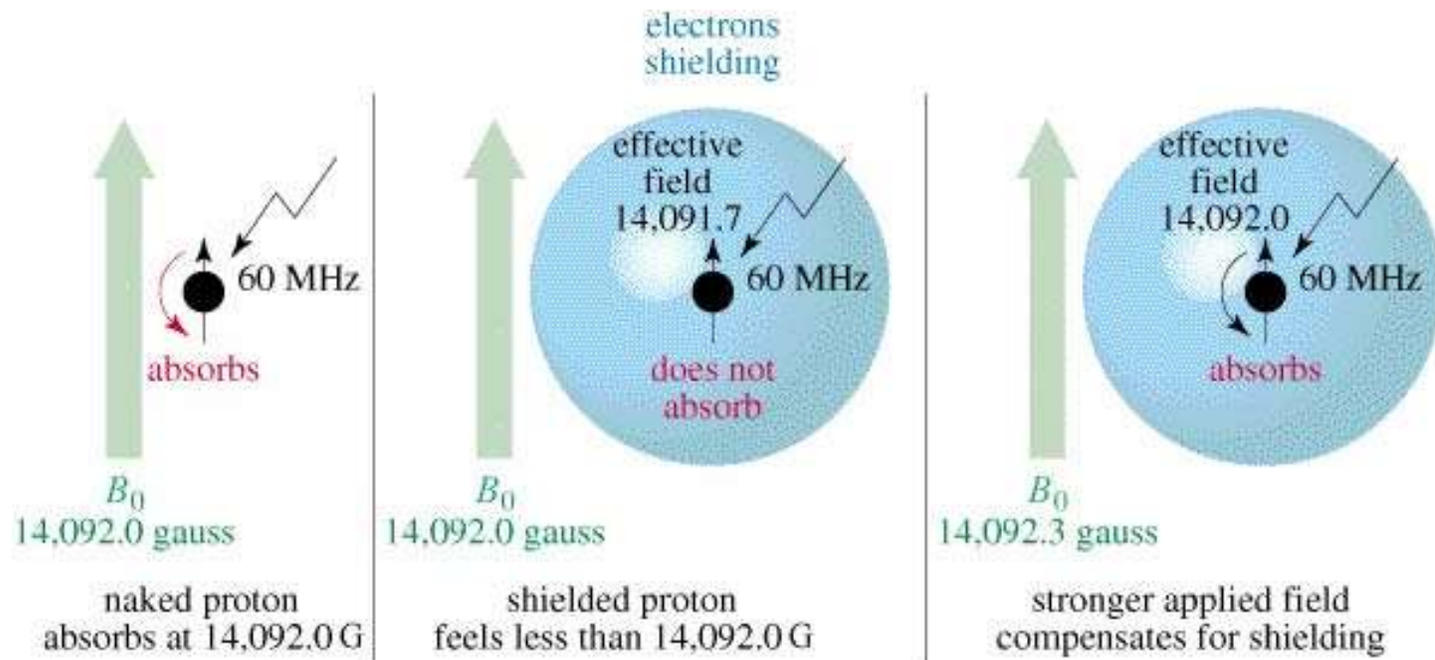
If all protons absorbed the same amount of energy in a given magnetic field, not much information could be obtained.

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.

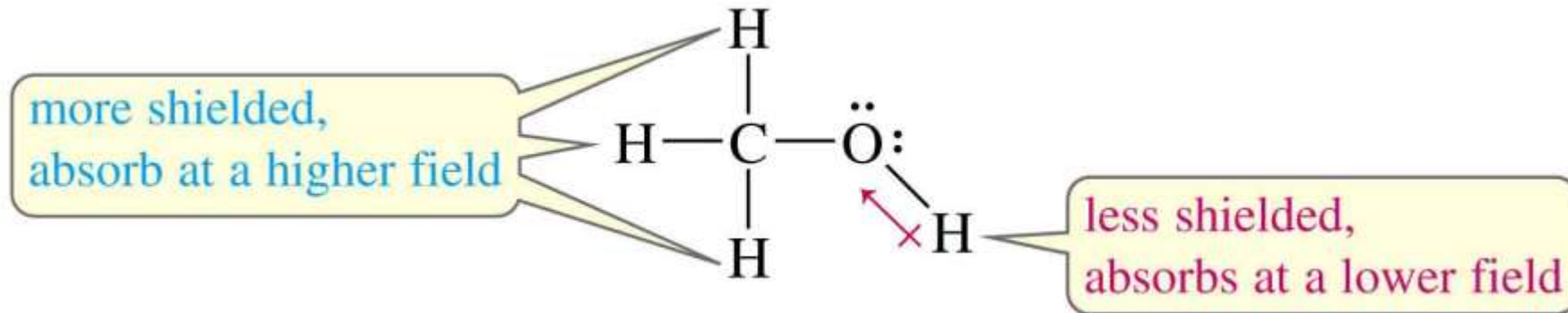
Shielded Protons

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



Protons in a Molecule

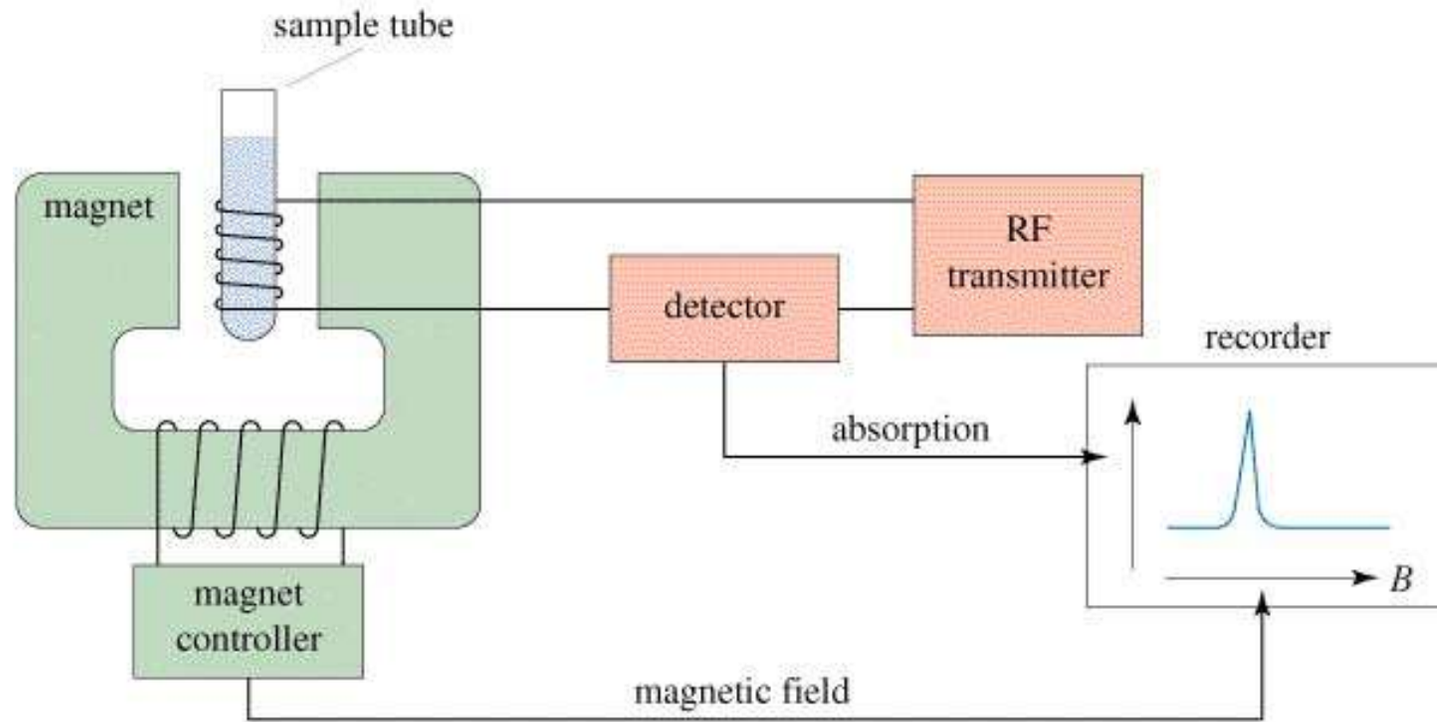
Depending on their **chemical environment**, protons in a molecule are **shielded by different amounts**.



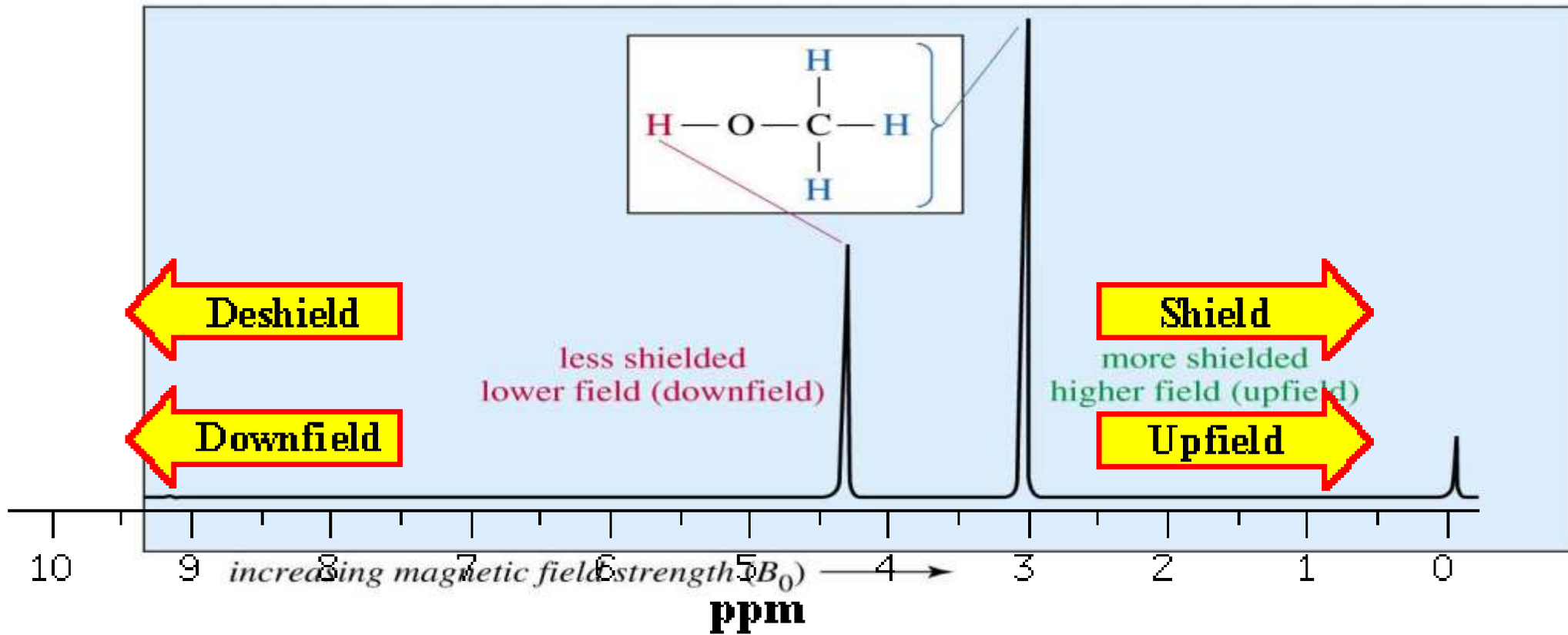
NMR Signals

- 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.

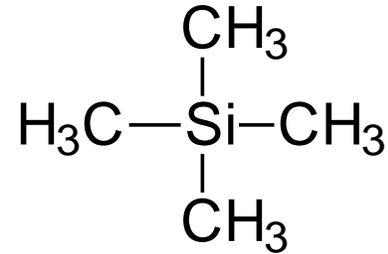
The NMR Spectrometer



The NMR Graph



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 (δ).

Chemical Shift calculation

$$\delta = \frac{V_H - V_{\text{TMS}}}{V_{\text{NMR}}} \times 10^6 \text{ ppm}$$

δ = Chemical shift (ppm)

V_H = Frequency of proton

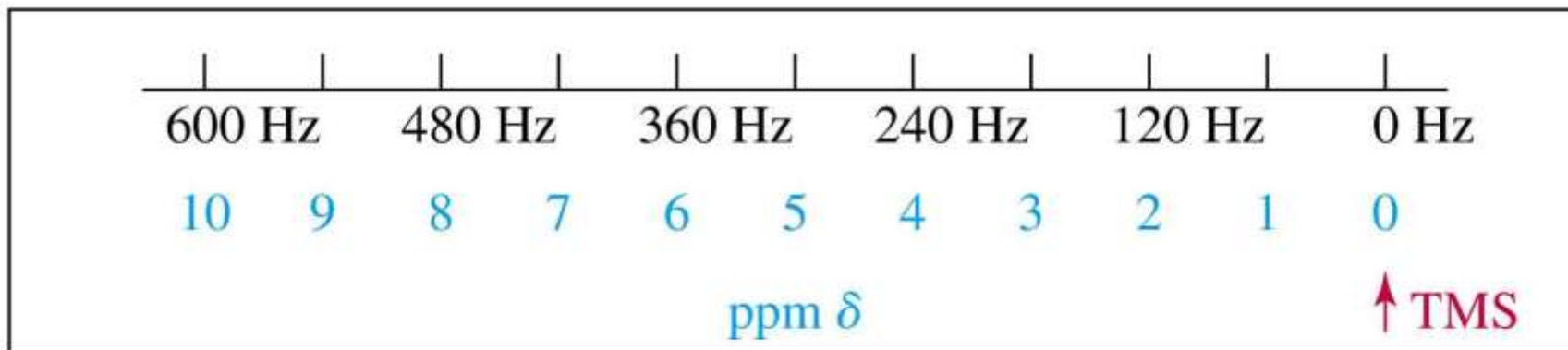
$V_{\text{TMS}} = 0$

$V_{\text{NMR}} = 60 \text{ MHz or } 100 \text{ MHz or } 300 \text{ MHz}$

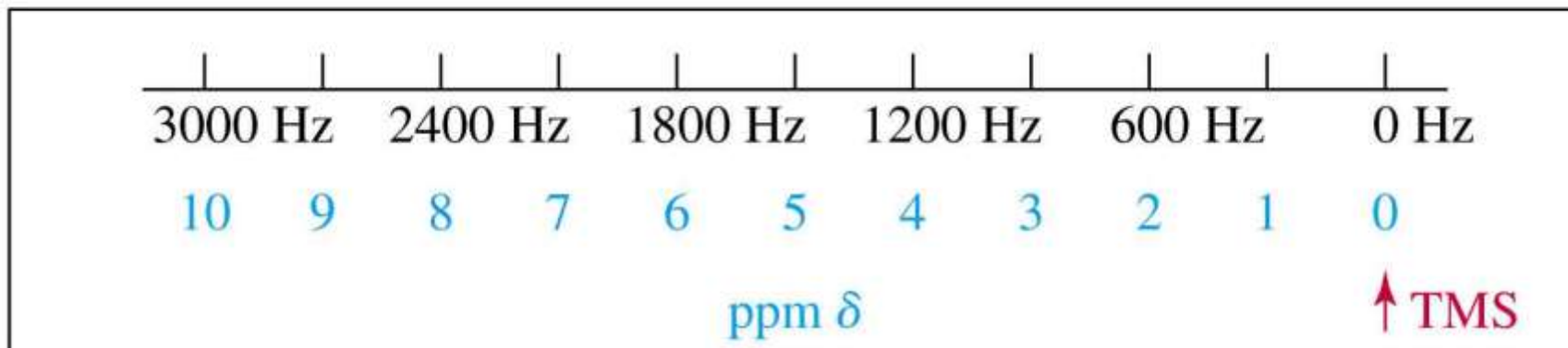
$1 \text{ MHz} = 10^6 \text{ Hz}$

Delta Scale

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

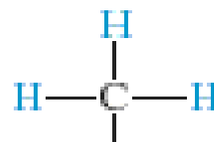
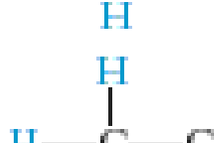
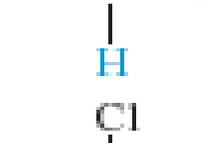
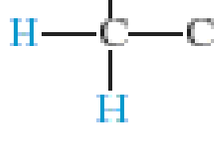


60 MHz



300 MHz

TABLE 13-2 Chemical Shifts of the Chloromethanes

<i>Compound</i>	<i>Chemical Shift</i>	<i>Difference</i>
	$\delta 0.2$	
	$\delta 3.0$	2.8 ppm
	$\delta 5.3$	2.3 ppm
	$\delta 7.2$	1.9 ppm

Note: Each chlorine atom added changes the chemical shift of the remaining methyl protons by about 2 to 3 ppm. These changes are nearly additive.

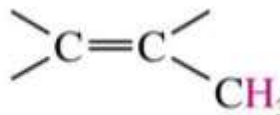
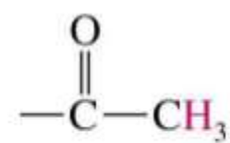
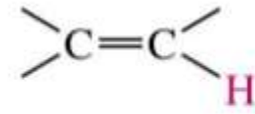
Location of Signals

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

CHCl_3	CH_2Cl_2	CH_3F	CH_3OH	CH_3Cl	CH_3Br	CH_3I	CH_4	$(\text{CH}_3)_4\text{Si}$
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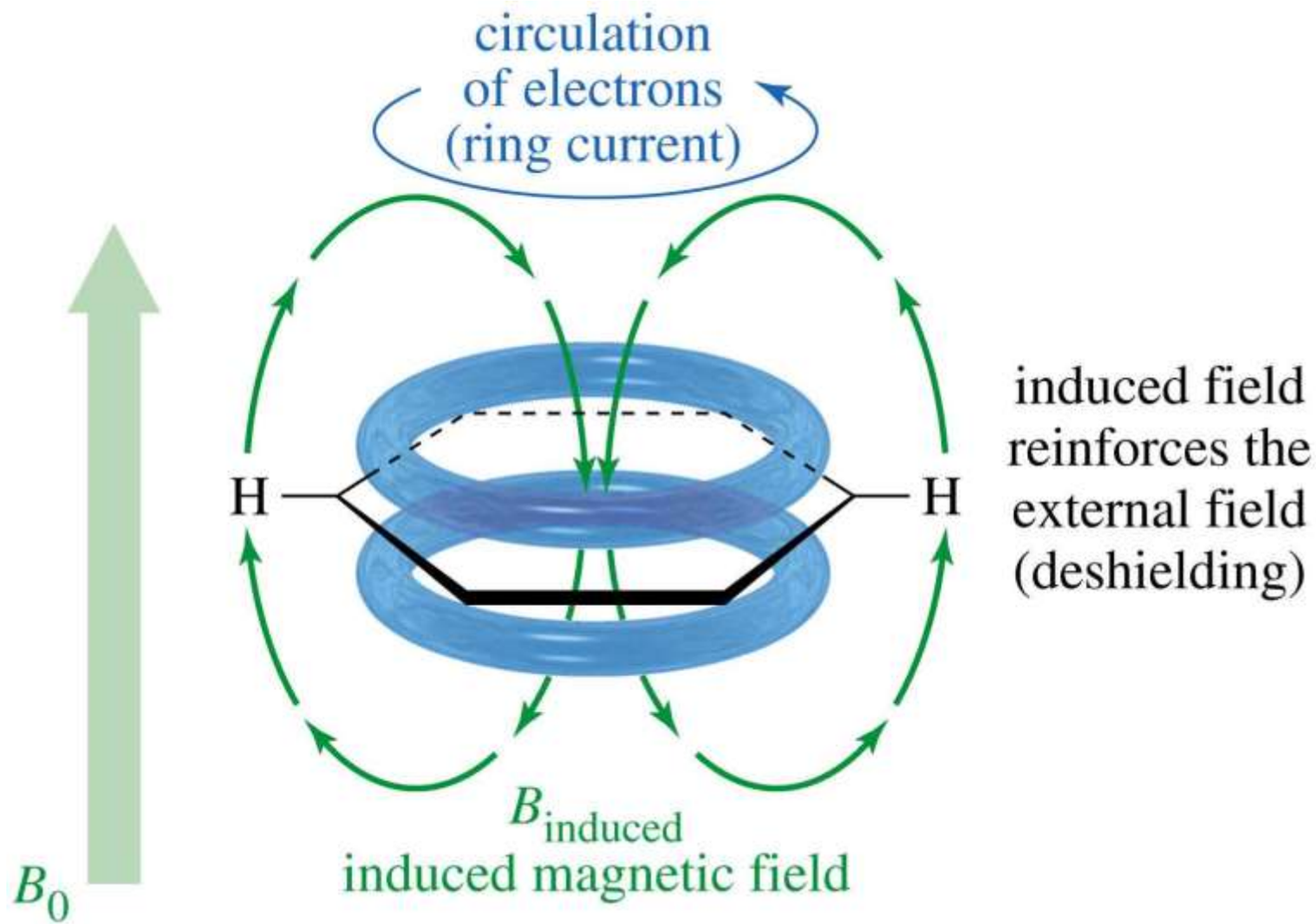
7.27	5.30	4.26	3.4	3.05	2.68	2.16	0.23	0
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Typical Values

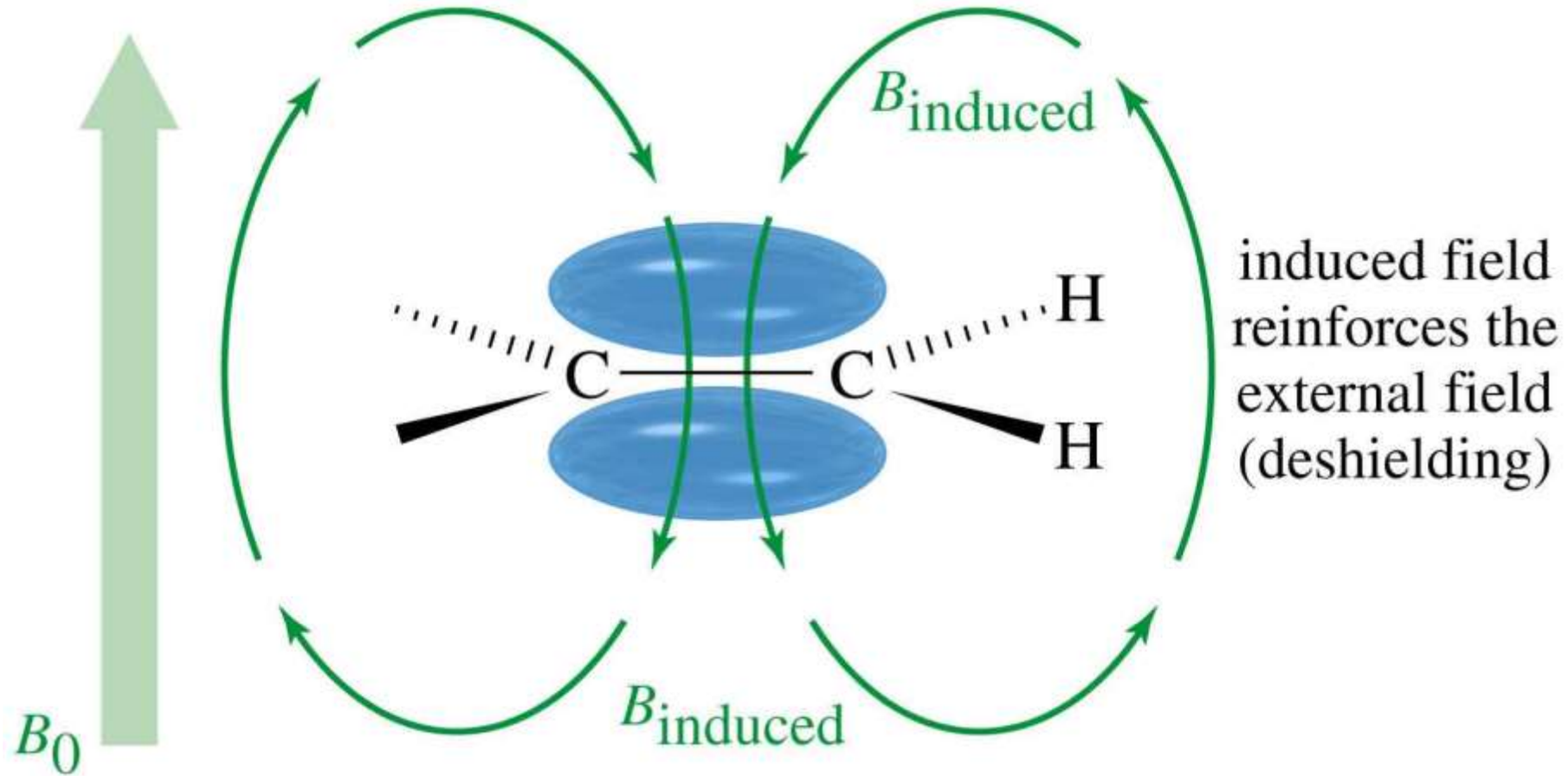
Type of Proton	Approximate δ	Type of Proton	Approximate δ
alkane ($-\text{CH}_3$)	0.9		1.7
alkane ($-\text{CH}_2-$)	1.3	Ph—H	7.2
alkane ($-\underset{ }{\text{CH}}-$)	1.4	Ph—CH ₃	2.3
	2.1	R—CHO	9–10
$-\text{C}\equiv\text{C}-\text{H}$	2.5	R—COOH	10–12
R—CH ₂ —X	3–4	R—OH	variable, about 2–5
(X = halogen, O)		Ar—OH	variable, about 4–7
	5–6	R—NH ₂	variable, about 1.5–4

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.

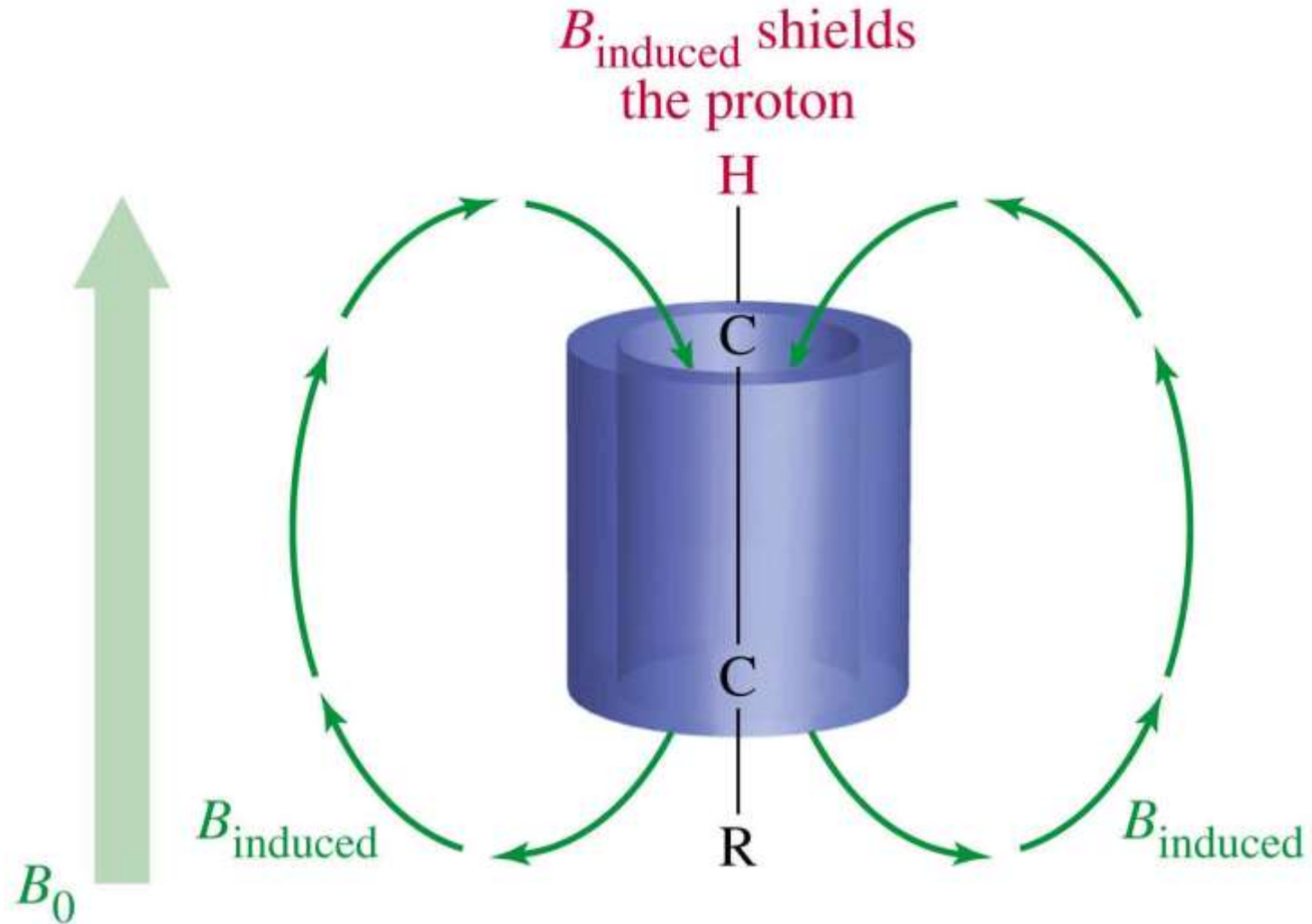
Aromatic Protons, $\delta 7\text{-}\delta 8$



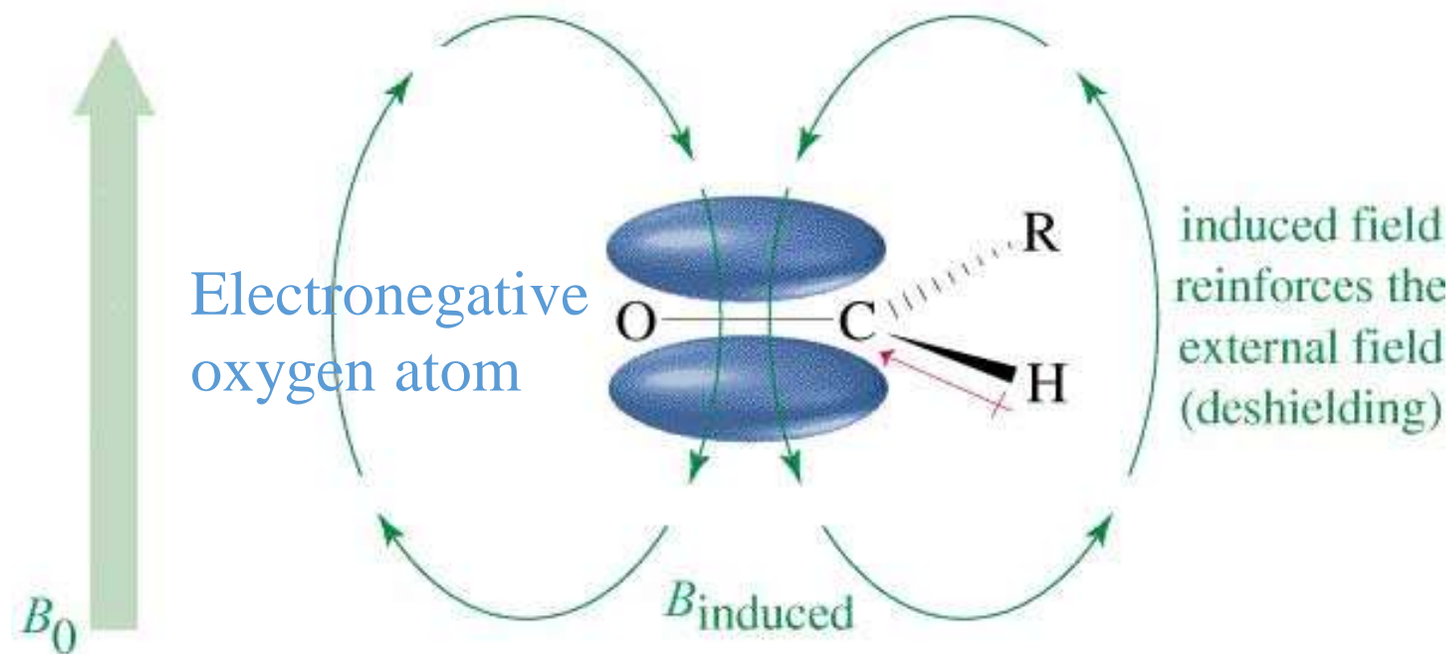
Vinyl Protons, $\delta 5\text{-}\delta 6$



Acetylenic Protons, $\delta 2.5$

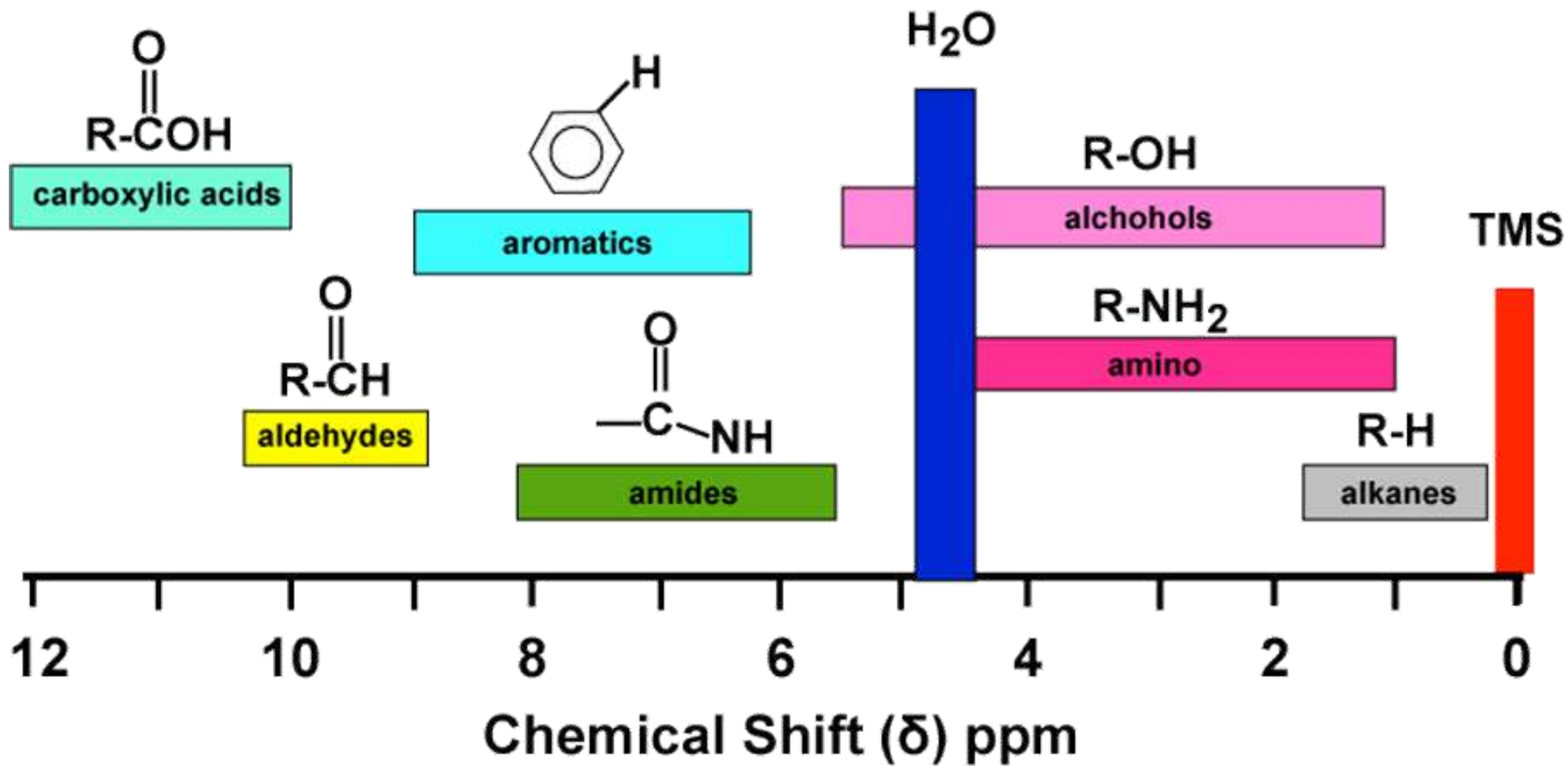


Aldehyde Proton, $\delta 9$ - $\delta 10$

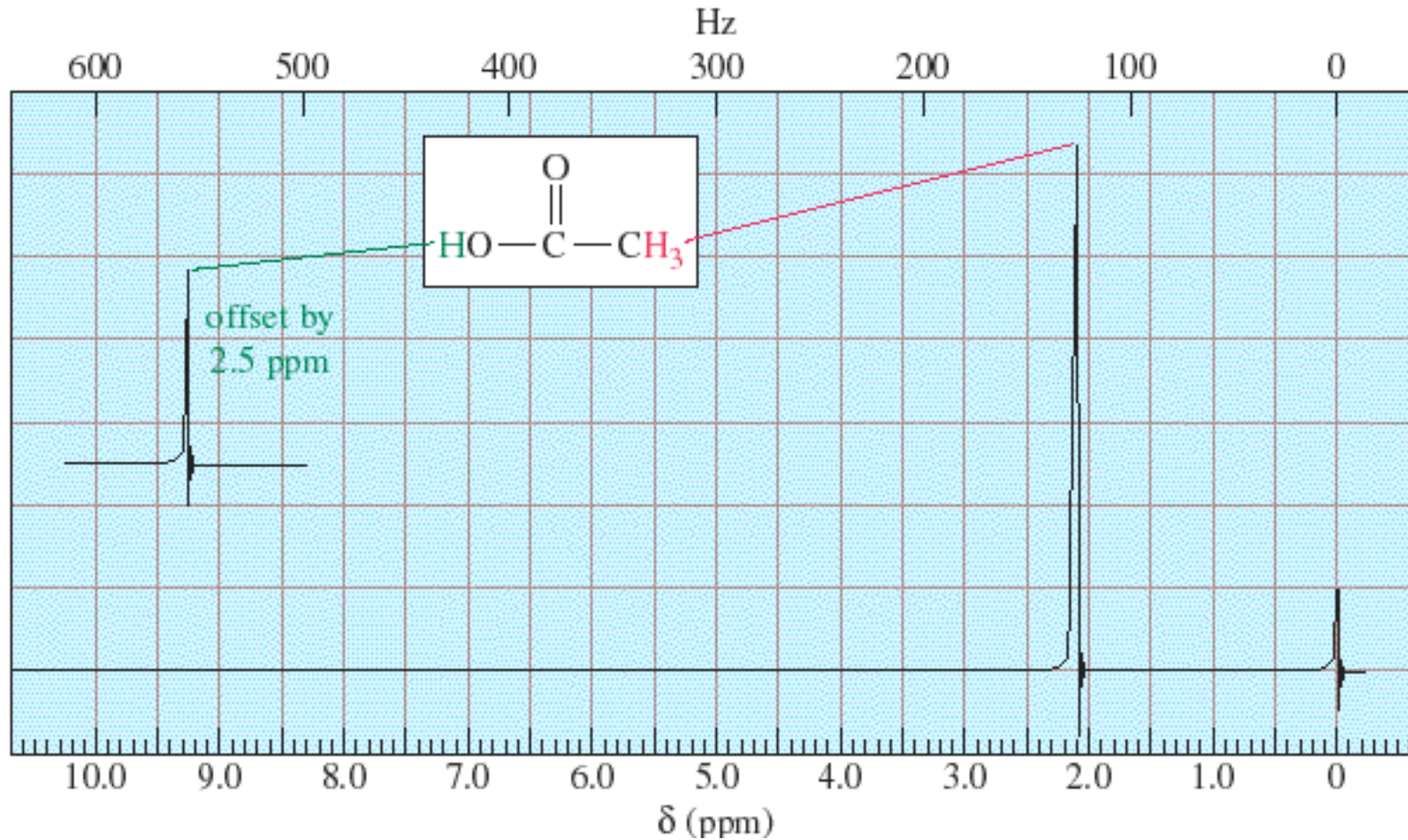


O-H and N-H Signals

- Chemical shift depends on concentration.
- Hydrogen bonding in concentrated solutions deshield the protons, so signal is around $\delta 3.5$ for N-H and $\delta 4.5$ for O-H.
- Proton exchanges between the molecules broaden the peak.

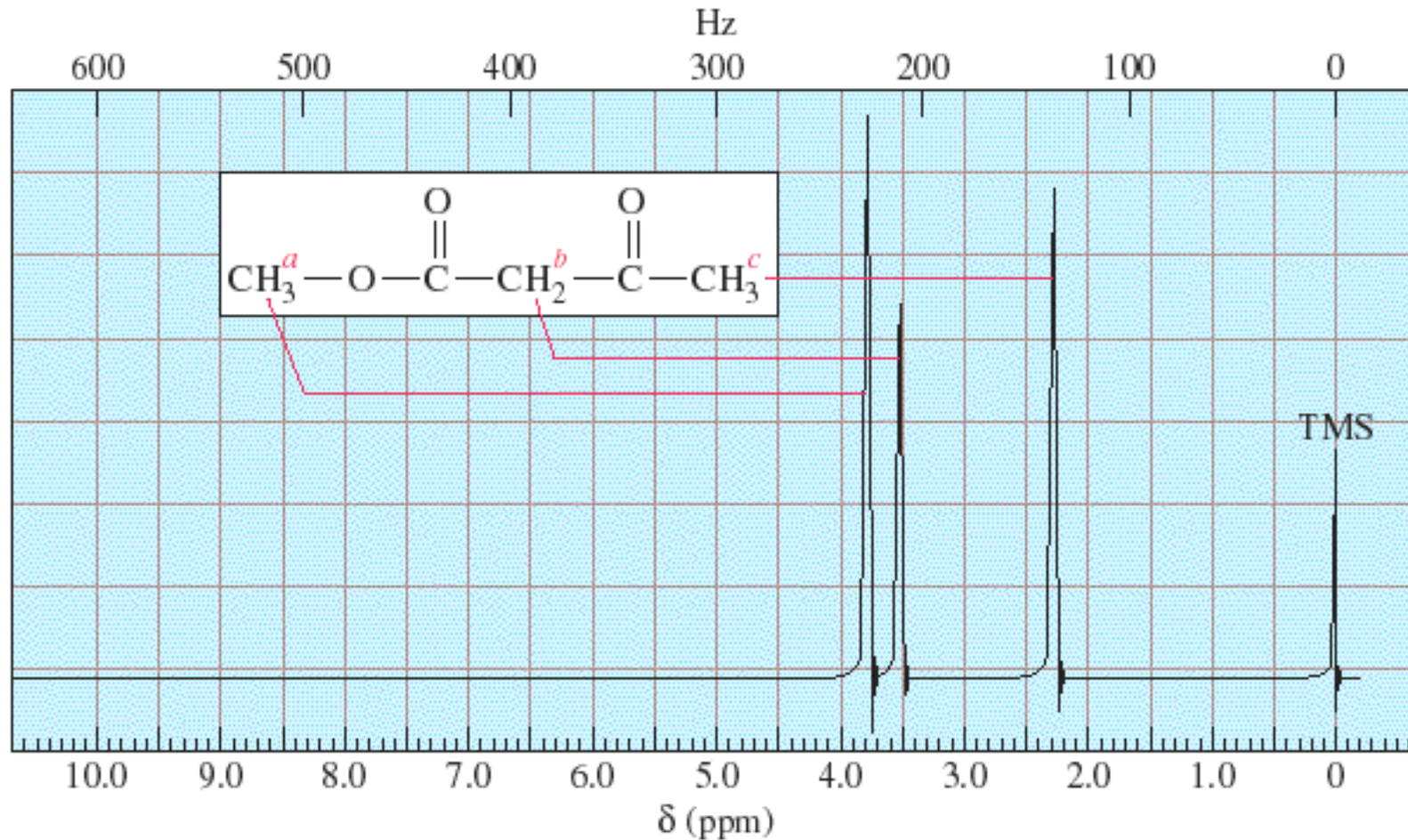


Carboxylic Acid Proton, $\delta 10+$



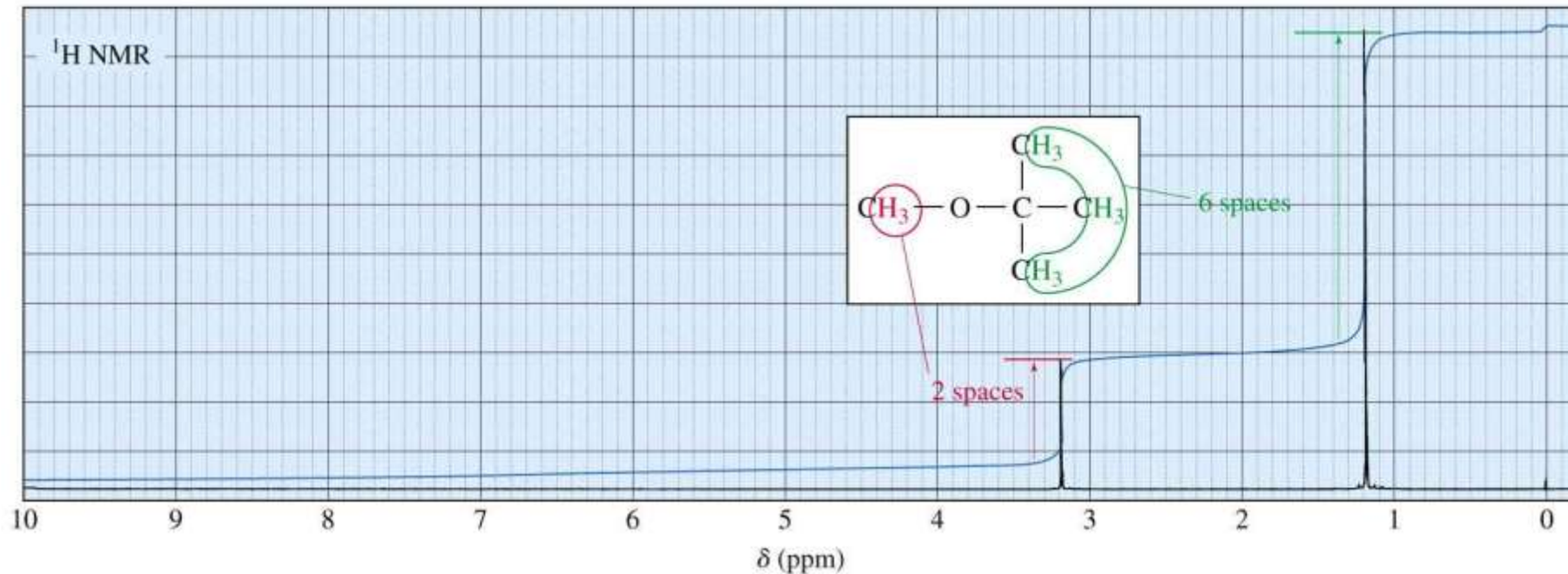
Number of Signals

Equivalent hydrogens have the same chemical shift.



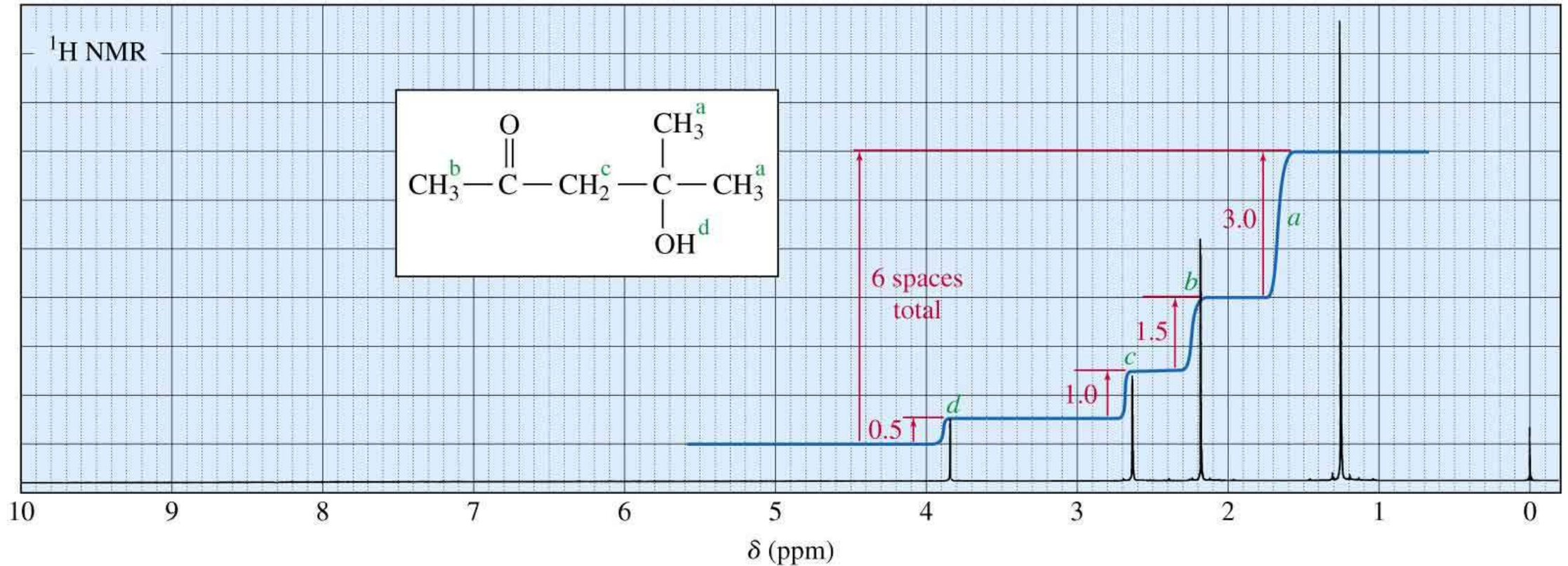
Intensity of Signals

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



How Many Hydrogens?

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



The $N + 1$ Rule

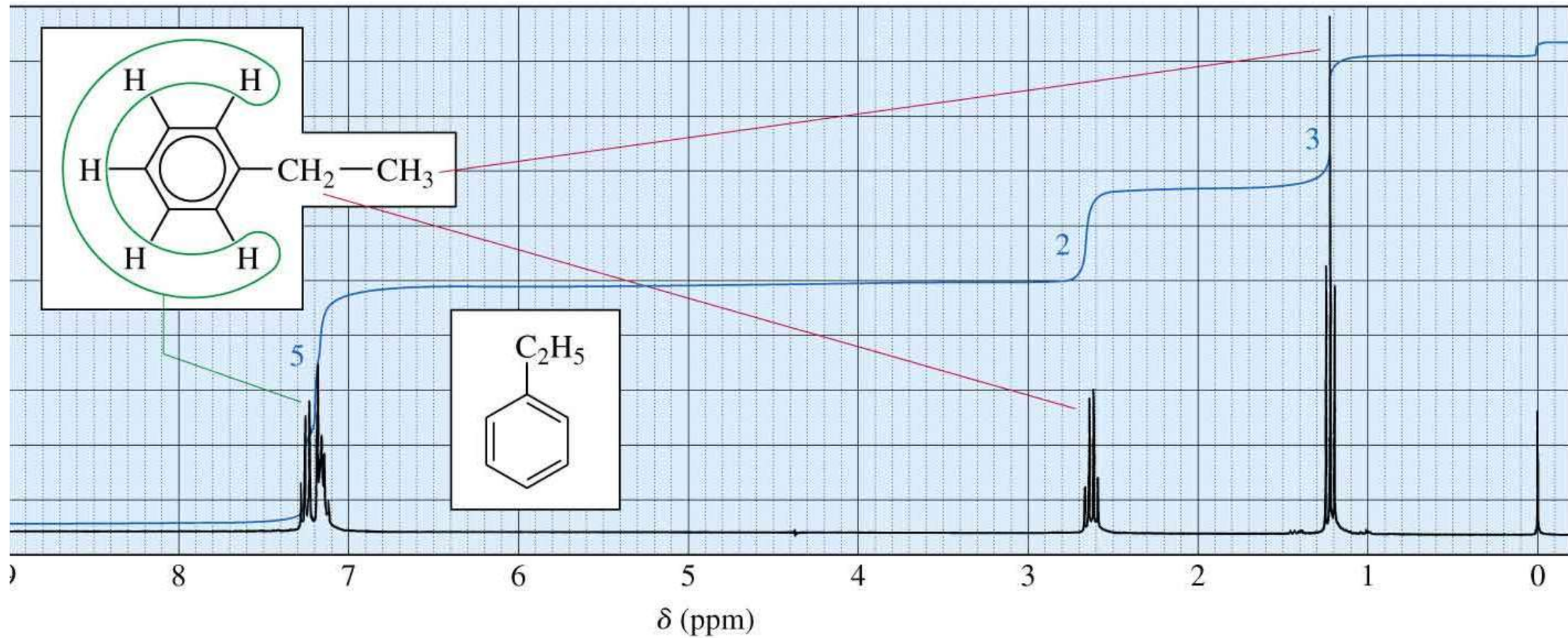
If a signal is split by N equivalent protons, it is split into $N + 1$ peaks.

Relative Peak Intensities of Symmetric Multiplets		
<i>Number of Equivalent Protons Causing Splitting</i>	<i>Number of Peaks (multiplicity)</i>	<i>Area Ratios (Pascal's triangle)</i>
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)	1 4 6 4 1
5	6 (sextet)	1 5 10 10 5 1
6	7 (septet)	1 6 15 20 15 6 1

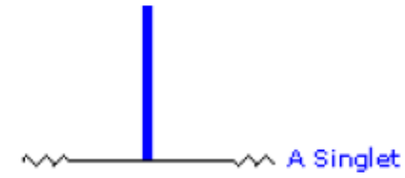
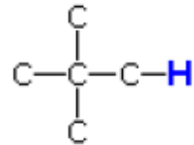
Range of Magnetic Coupling

- 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.

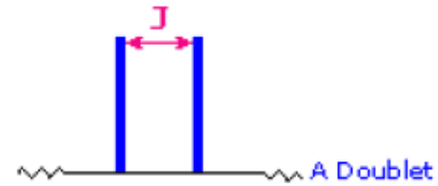
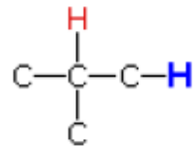
Splitting for Ethyl Groups



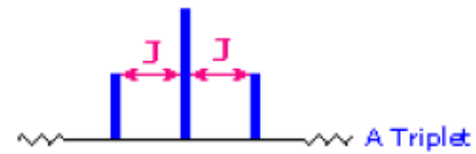
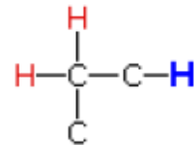
No Coupled
Hydrogens



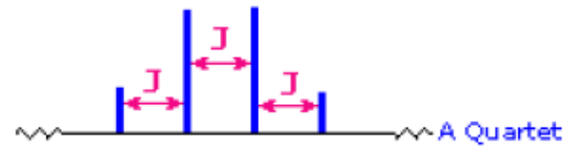
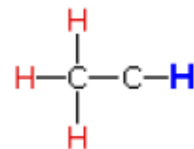
One Coupled
Hydrogen



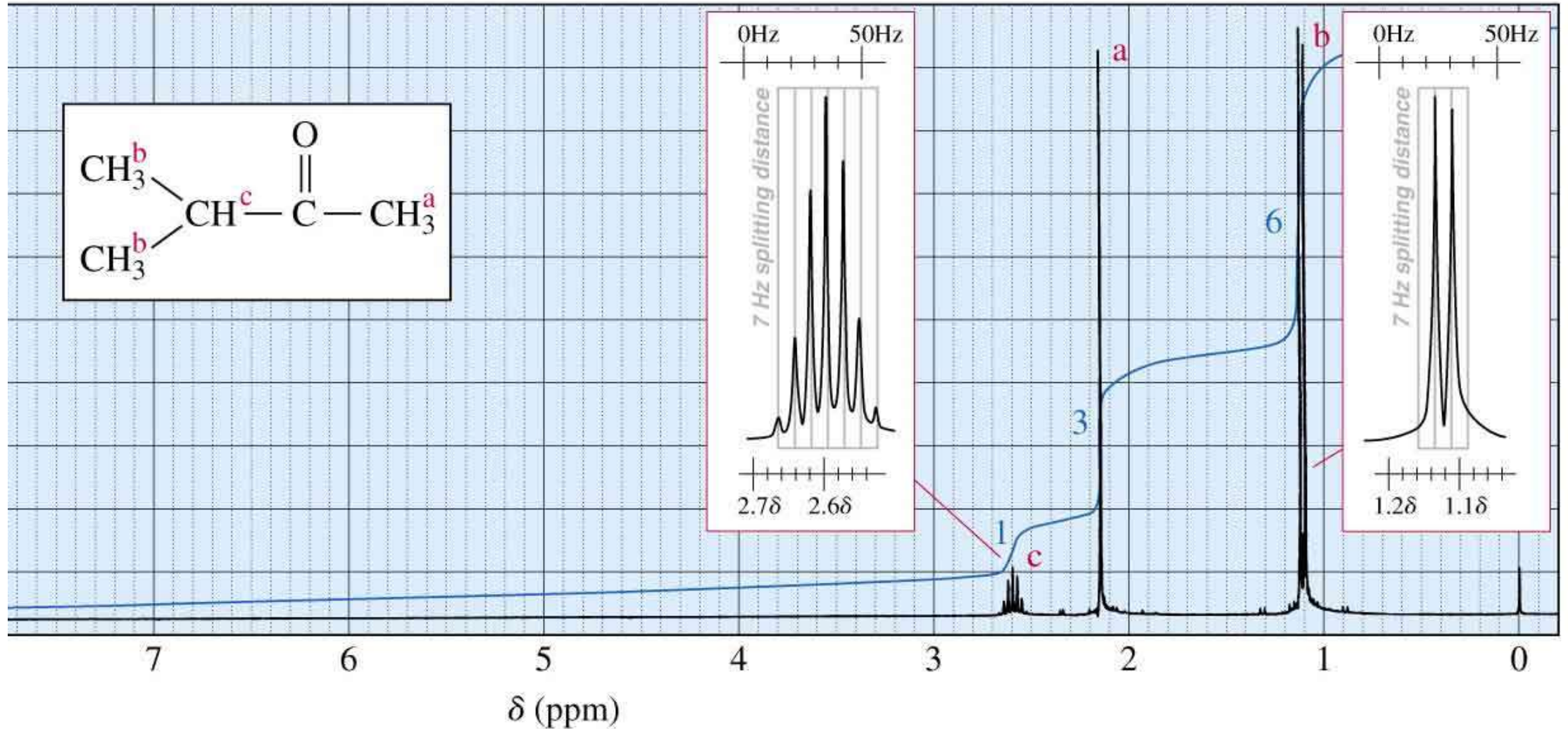
Two Coupled
Hydrogens



Three Coupled
Hydrogens



Splitting for Isopropyl Groups

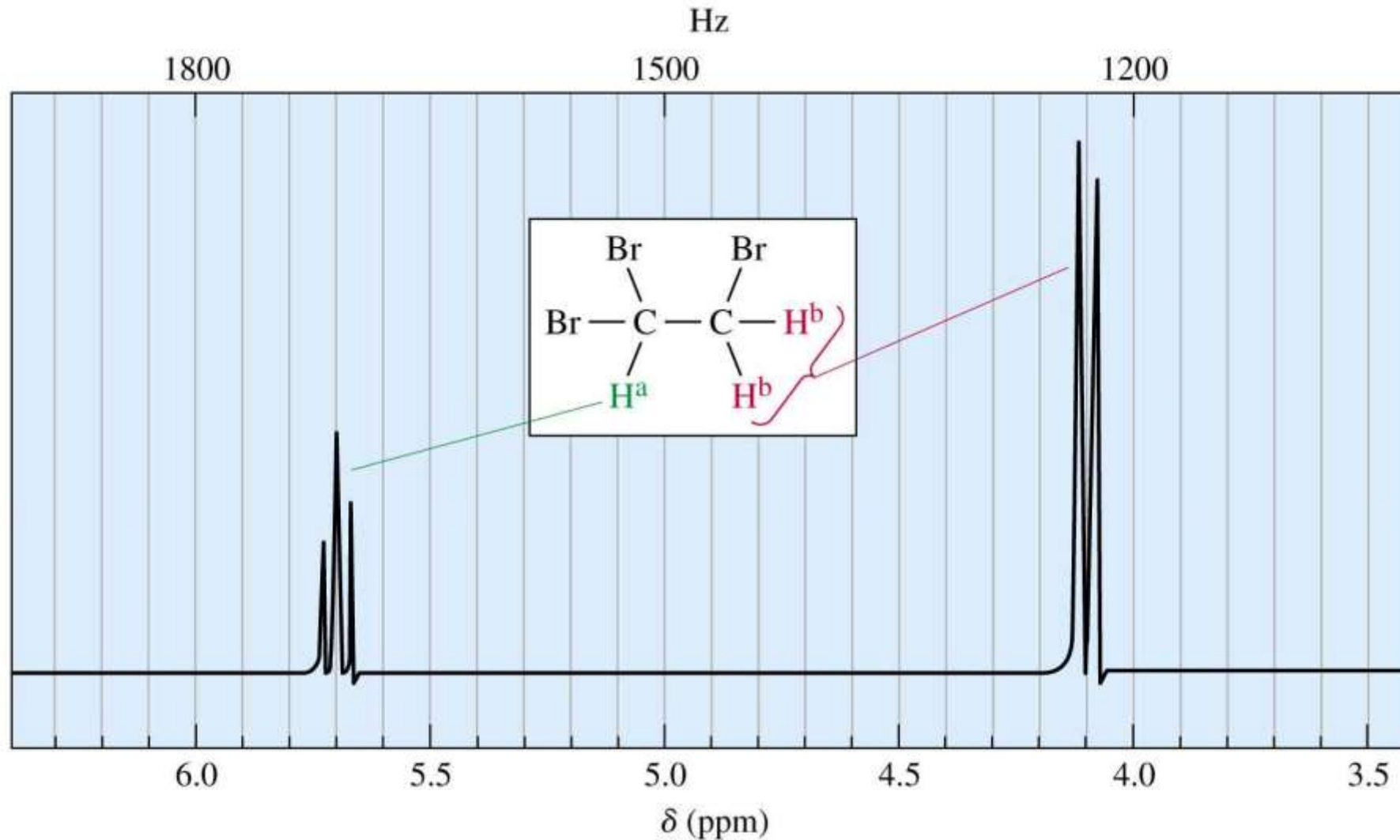


Spin-Spin Splitting

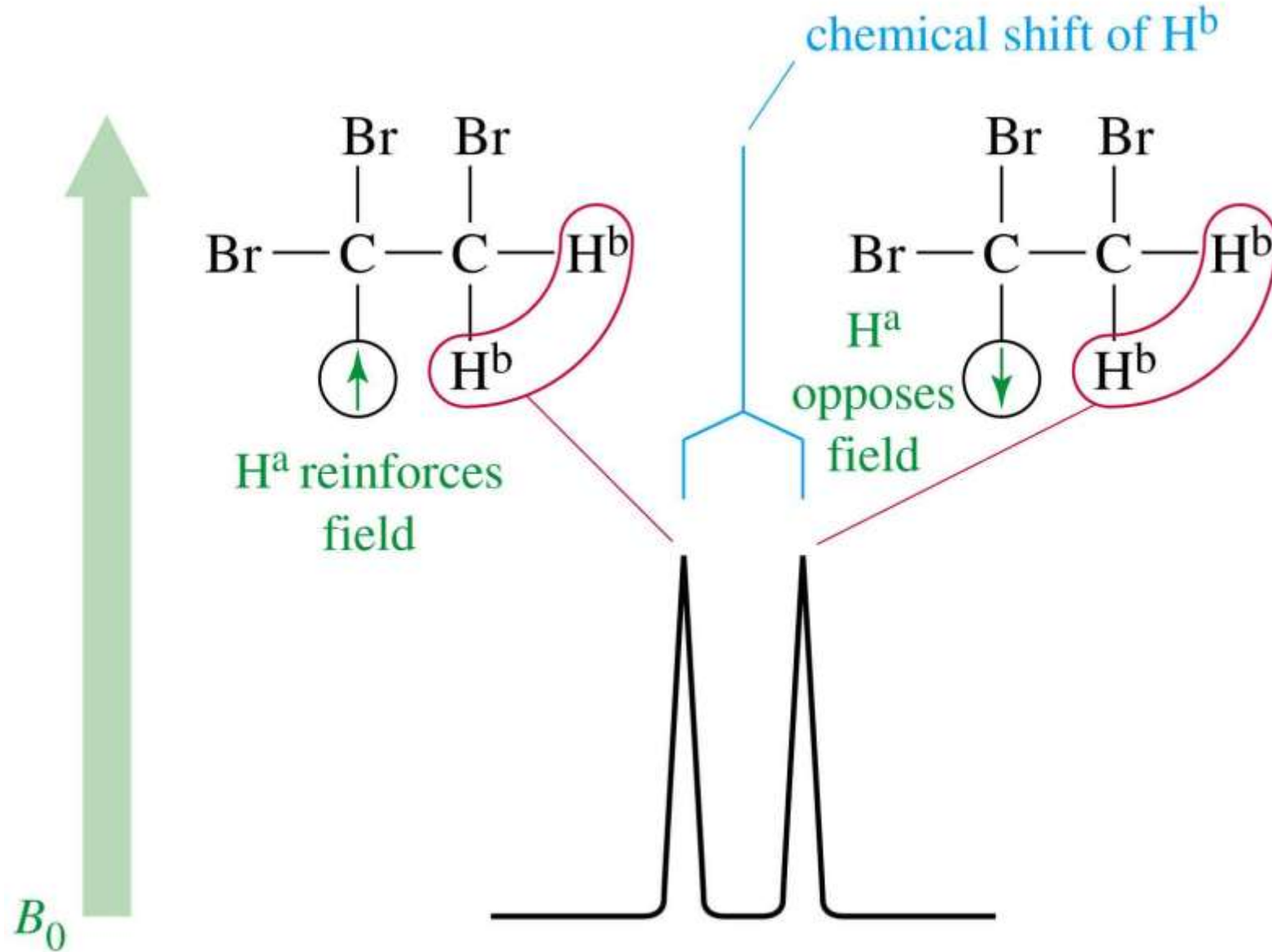
- Nonequivalent protons on **adjacent carbons** have magnetic fields that may align with or oppose the external field.
- This magnetic coupling causes the proton to absorb slightly downfield when the external field is reinforced and slightly upfield when the external field is opposed.
- All possibilities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

1,1,2-Tribromoethane

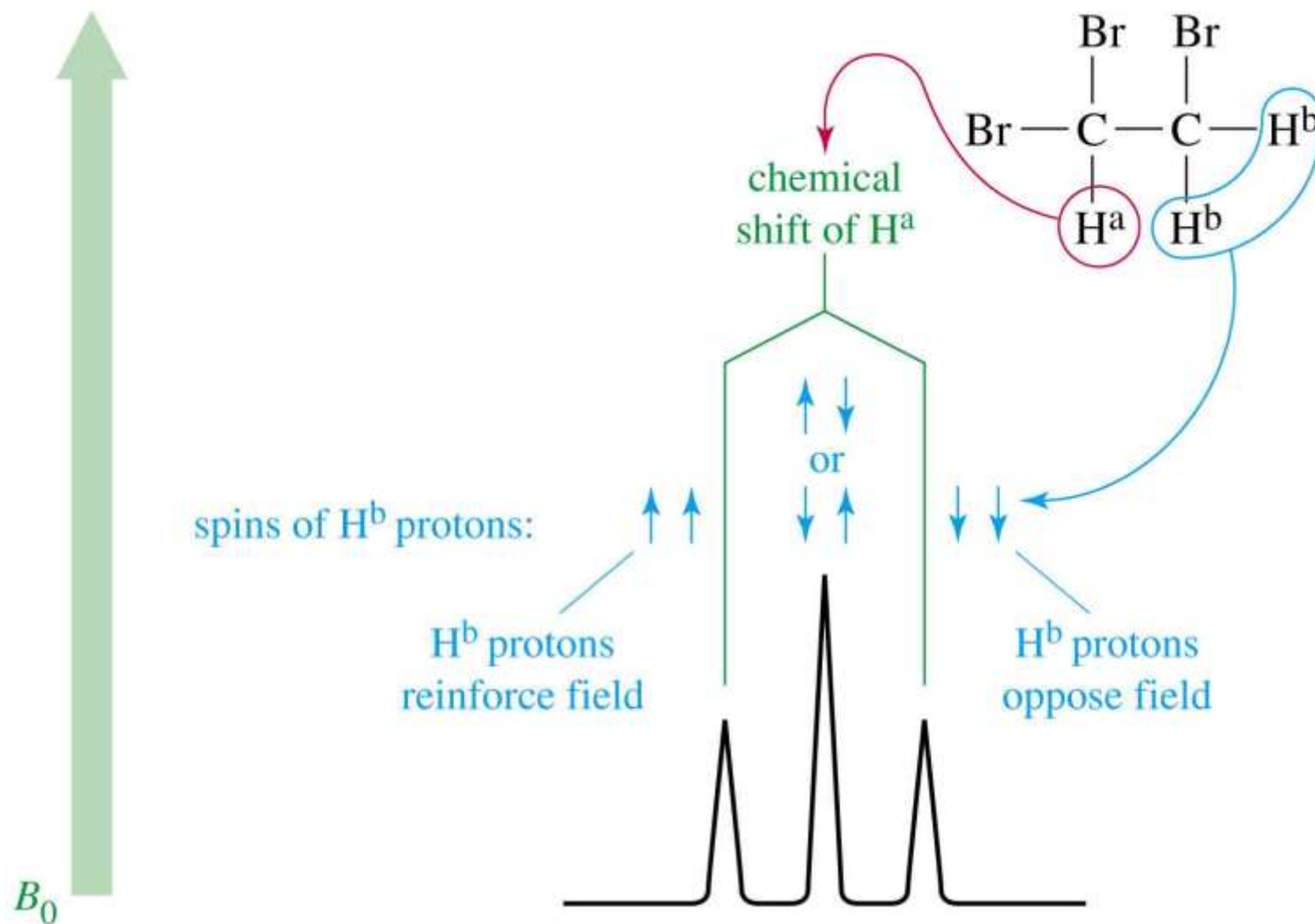
Nonequivalent protons on adjacent carbons.



Doublet: 1 Adjacent Proton



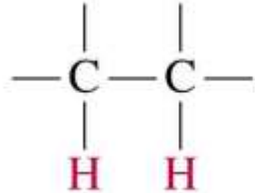
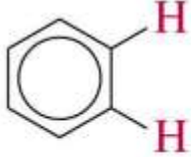
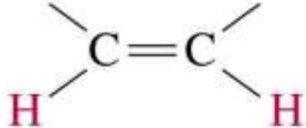
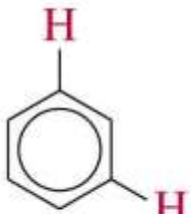
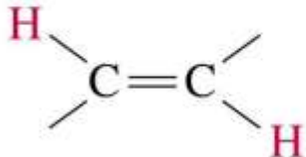
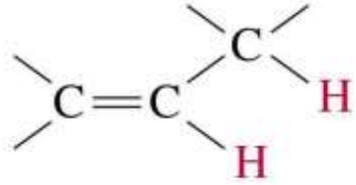
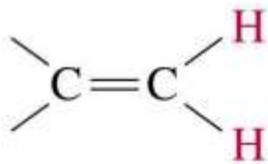
Triplet: 2 Adjacent Protons



Coupling Constants

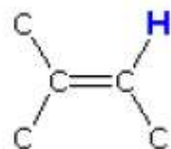
- Distance between the peaks of multiplet
- Measured in Hz
- Not dependent on strength of the external field
- Multiplets with the same coupling constants may come from adjacent groups of protons that split each other.

Values for Coupling Constants

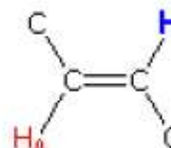
		<u>Approx. J</u>		<u>Approx. J</u>
	(free rotation)	7 Hz ^a		8 Hz
			(ortho)	
	(cis)	10 Hz		2 Hz
			(meta)	
	(trans)	15 Hz		6 Hz
			(allylic)	
	(geminal)	2 Hz		

^aThe 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.

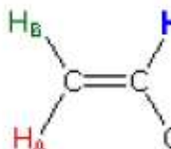
No Coupled
Hydrogens



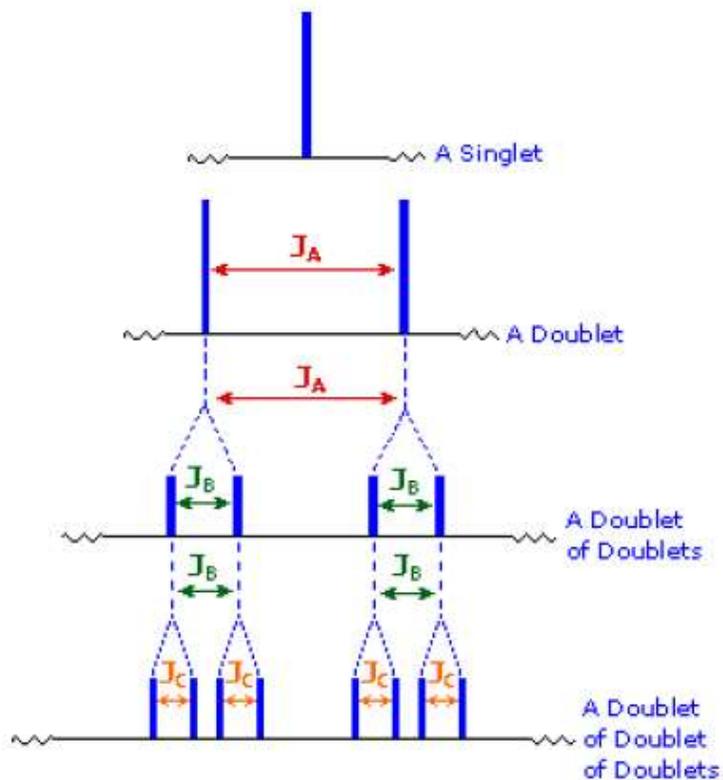
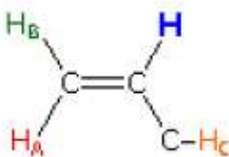
One Coupled
Hydrogen



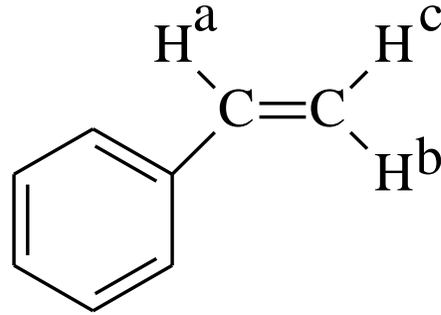
Two Coupled
Hydrogens



Three Coupled
Hydrogens

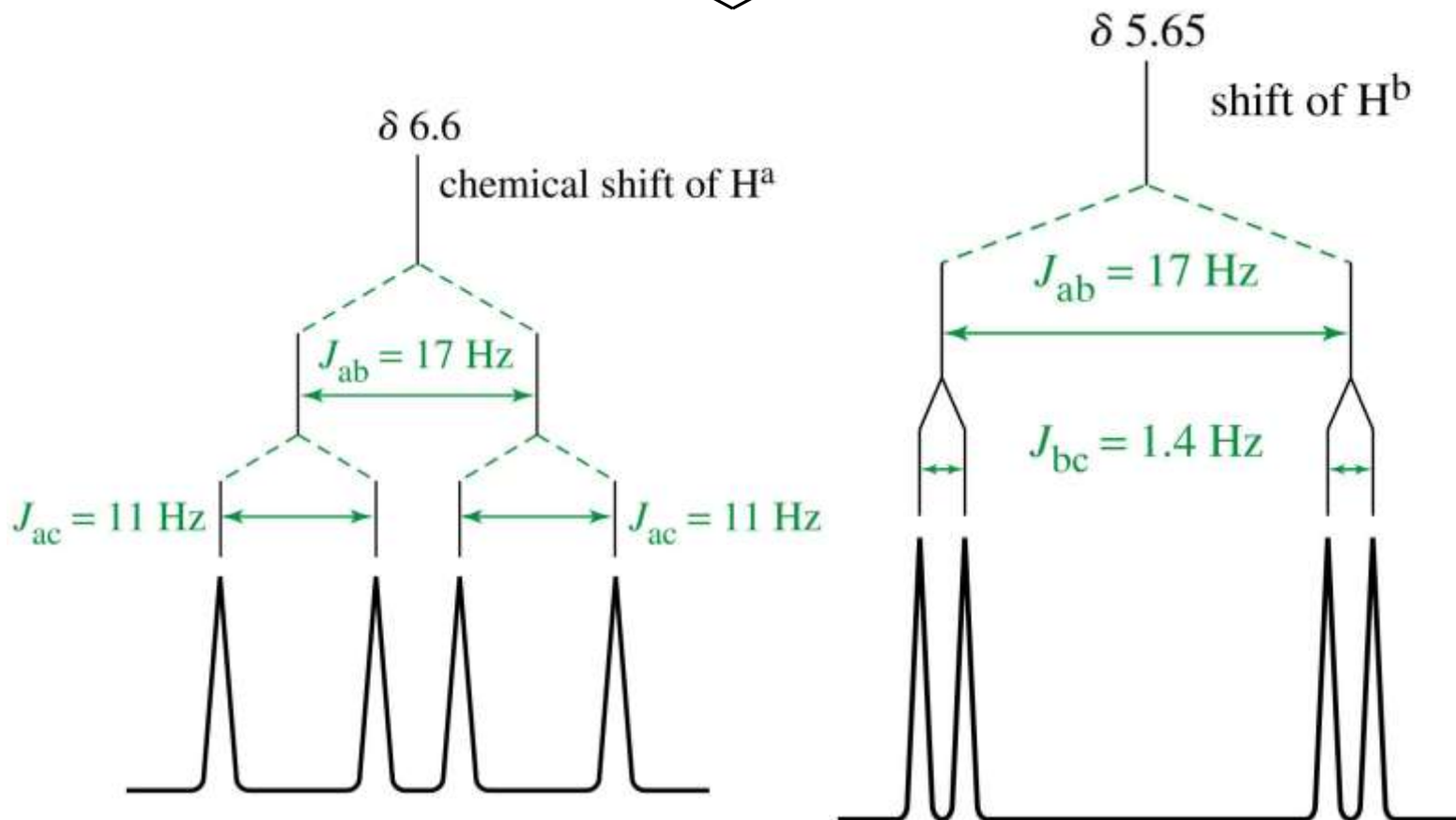
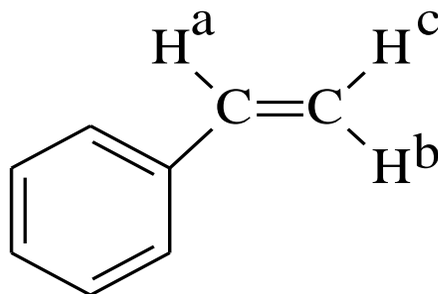


Complex Splitting

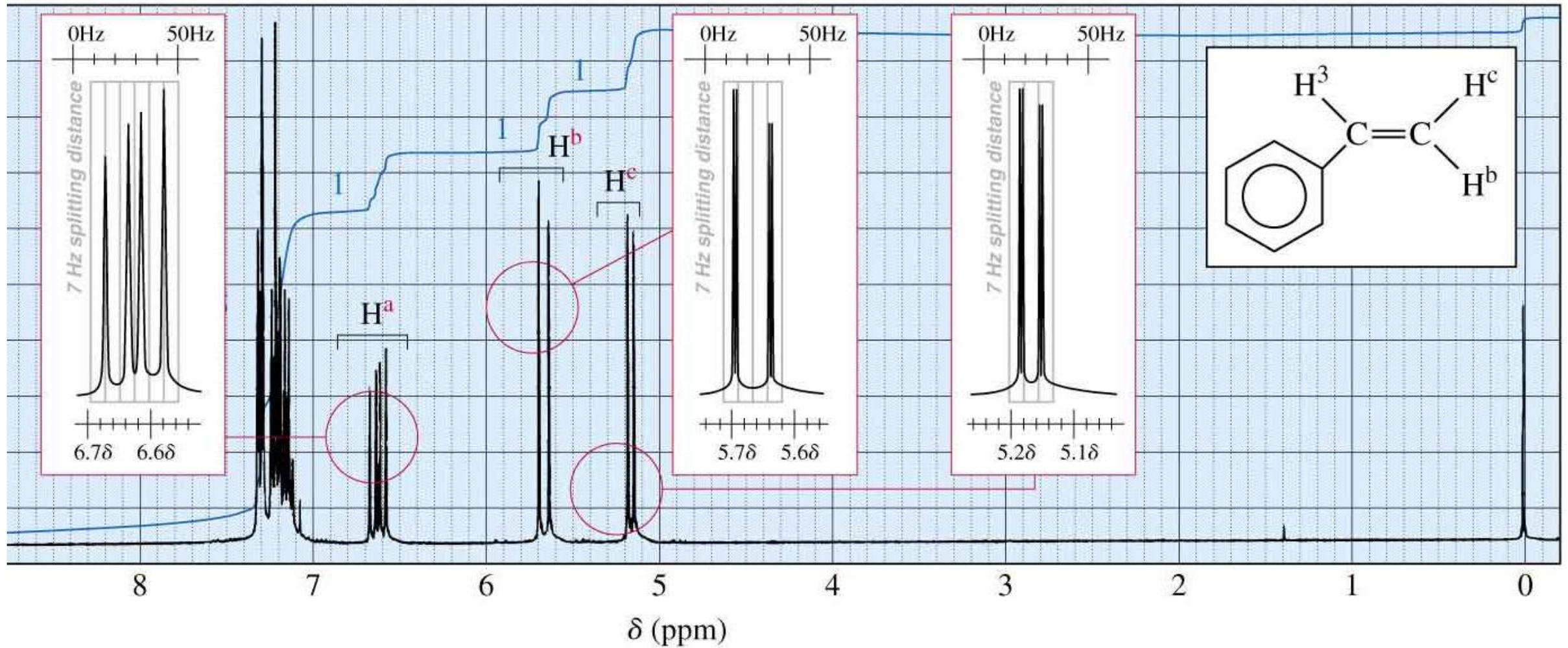


- Signals may be split by adjacent protons, different from each other, with different coupling constants.
- Example: H^a of styrene which is split by an adjacent H *trans* to it ($J = 17$ Hz) and an adjacent H *cis* to it ($J = 11$ Hz).

Splitting Tree



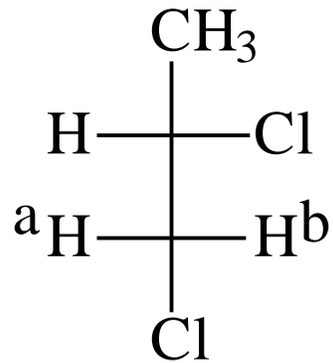
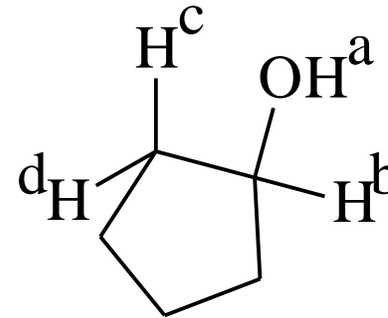
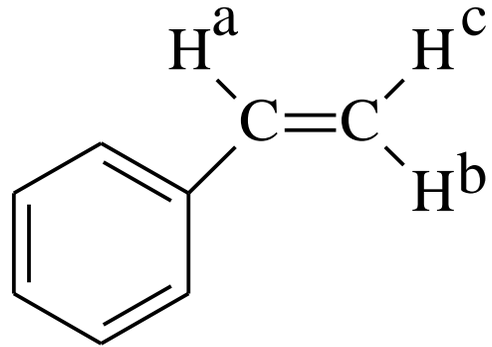
Spectrum for Styrene



Stereochemical Nonequivalence

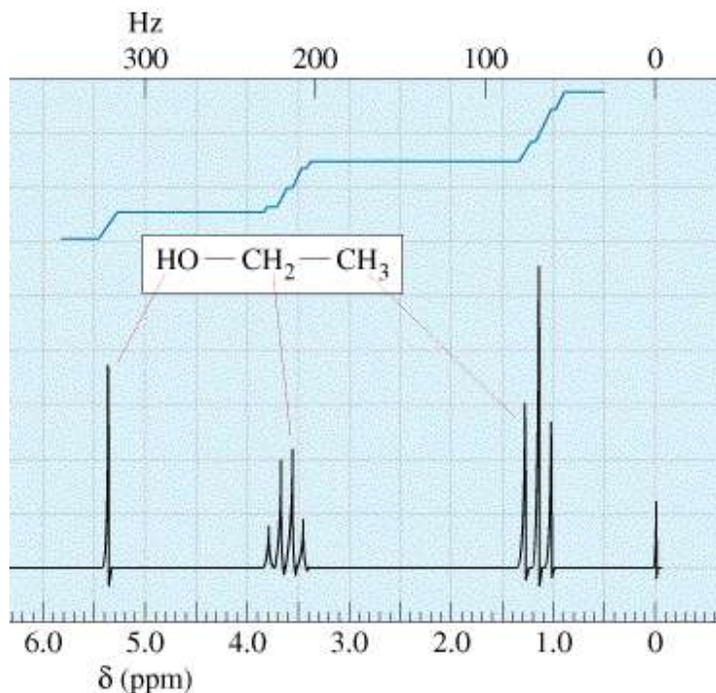
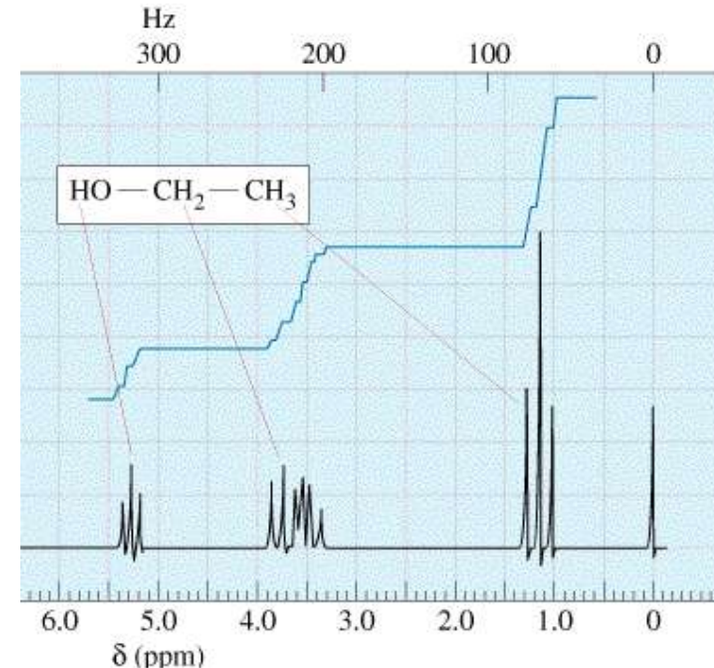
- Usually, two protons on the same C are equivalent and do not split each other.
- If the replacement of each of the protons of a -CH_2 group with an imaginary “Z” gives stereoisomers, then the protons are non-equivalent and will split each other.

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.

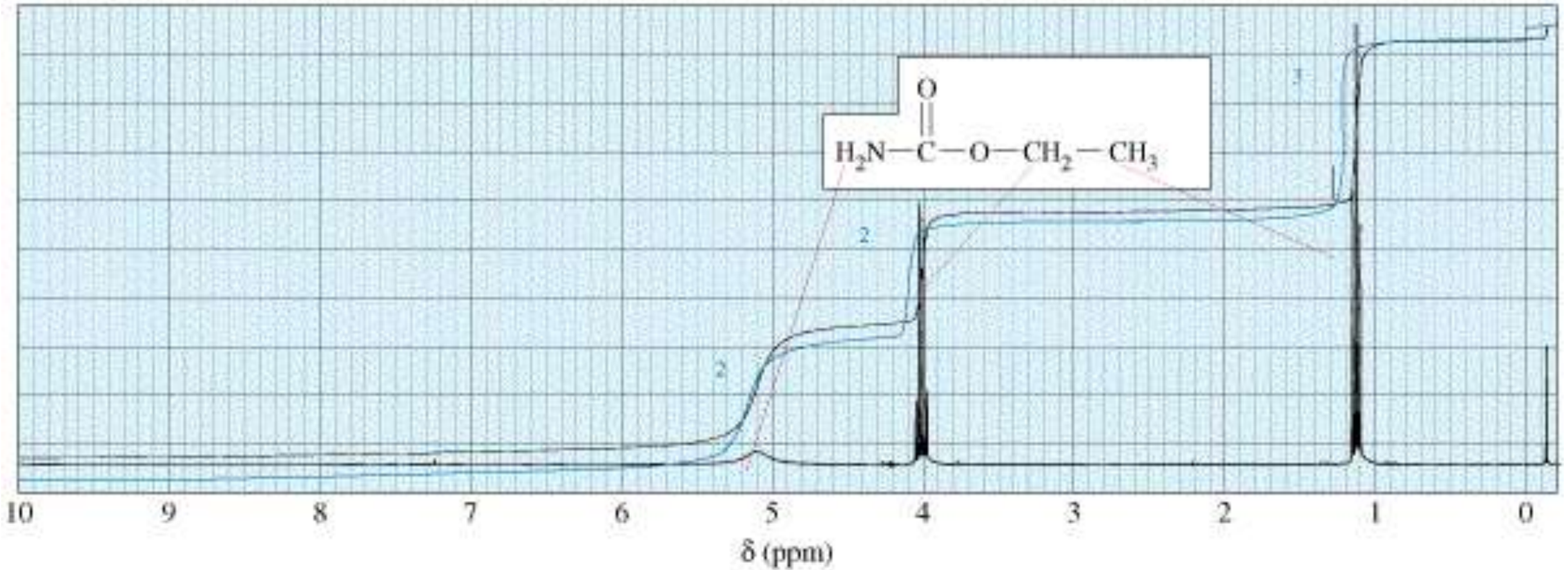


Hydroxyl Proton

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

N-H Proton

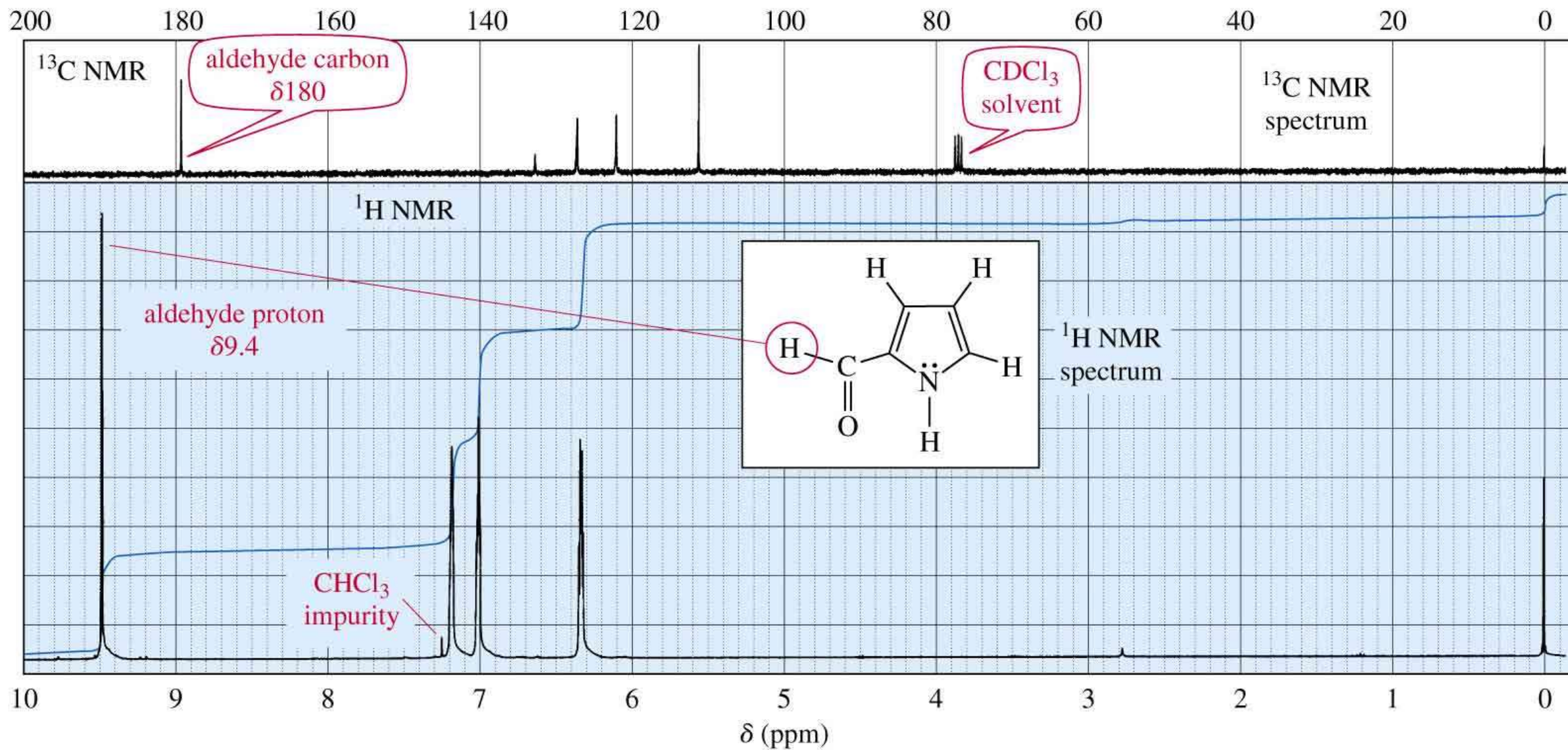
- Moderate rate of exchange.
- Peak may be broad.



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 D₂O
- 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.

Combined ^{13}C and ^1H Spectra



Differences in ^{13}C Technique

- Resonance frequency is \sim 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 ^{13}C would be adjacent to another ^{13}C , so splitting by carbon is negligible.
- ^{13}C will 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

- ^{13}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 ^{13}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.

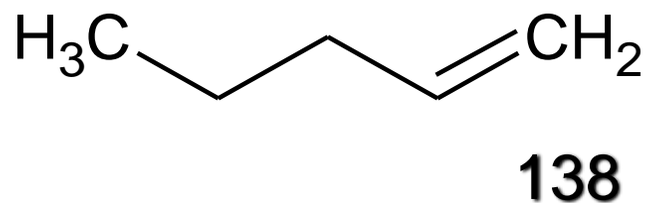
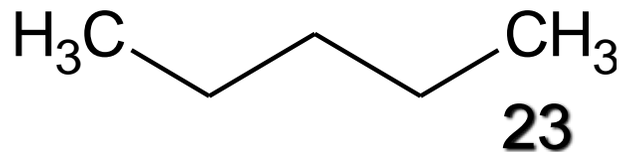
^{13}C Chemical Shifts

are measured in ppm (δ) from the carbons of TMS

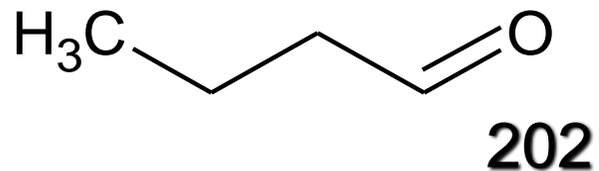
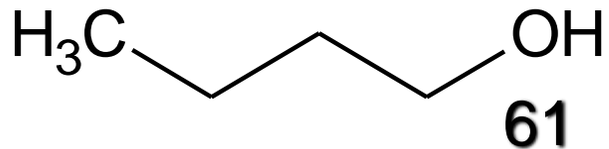
^{13}C Chemical shifts are most affected by:

- hybridization state of carbon
- electronegativity of groups attached to carbon

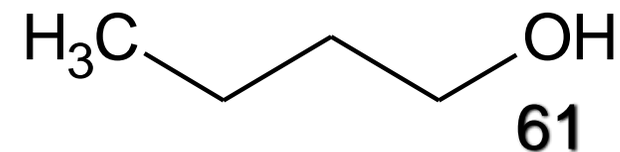
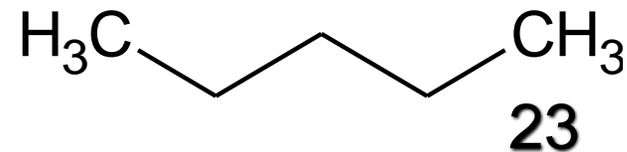
Examples (chemical shifts in ppm from TMS)



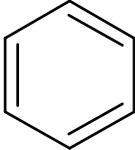
sp^3 hybridized carbon is more shielded than sp^2



sp^3 hybridized carbon is more shielded than sp^2



an electronegative atom deshields the carbon to which it is attached

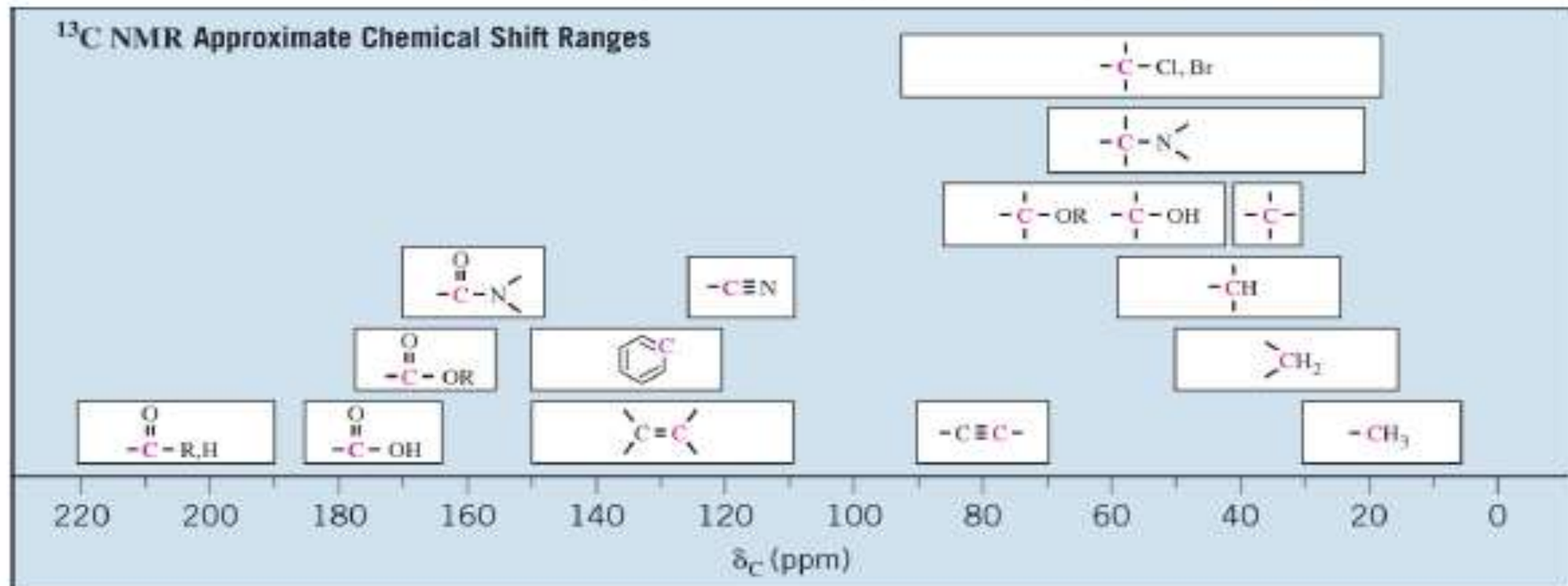
Type of carbon	Chemical shift (δ) ppm	Type of carbon	Chemical shift (δ) ppm
RCH_3	0-35	$RC\equiv CR$	65-90
R_2CH_2	15-40	$R_2C=CR_2$	100-150
R_3CH	25-50		110-175
R_4C	30-40		

Type of carbon	Chemical shift (δ) ppm	Type of carbon	Chemical shift (δ) ppm
RCH_2Br	20-40	$\begin{array}{c} \text{O} \\ \\ \text{RCOR} \end{array}$	160-185
RCH_2Cl	25-50	$\begin{array}{c} \text{O} \\ \\ \text{RCR} \end{array}$	190-220
RCH_2NH_2	35-50		
RCH_2OH	50-65		
RCH_2OR	50-65		

● ^{13}C Chemical Shifts

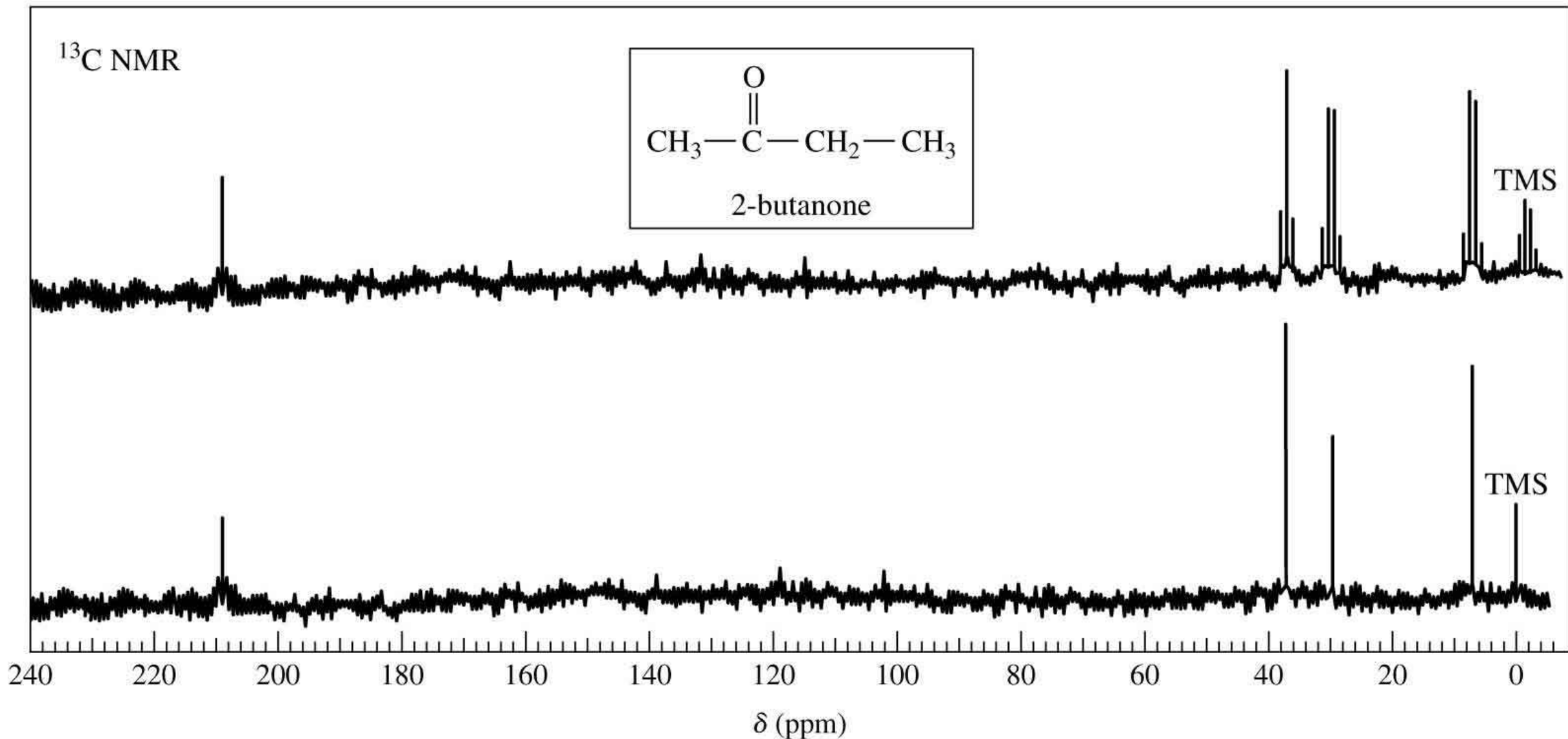
→ Just as in ^1H NMR spectroscopy, chemical shifts in ^{13}C NMR depend on the electron density around the carbon nucleus

- ⌘ Decreased electron density causes the signal to move downfield (*desheilding*)
- ⌘ Increased electron density causes the signal to move upfield (*sheilding*)



- Because of the wide range of chemical shifts, it is rare to have two ^{13}C peaks coincidentally overlap
- A group of 3 peaks at δ 77 comes from the common NMR solvent deuteriochloroform and can be ignored

Two ^{13}C NMR Spectra



Hydrogen and Carbon Chemical Shifts

