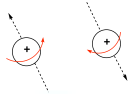


Nuclear Magnetic Resonance (NMR) Spectroscopy

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.

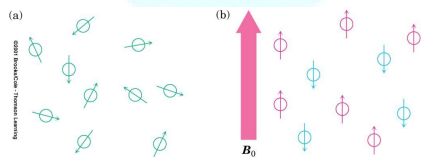
¹H and ¹³C are the most important NMR active nuclei in organic chemistry Natural Abundance

¹H 99.9%

¹³C 1.1%

¹²C 98.9% (not NMR active)

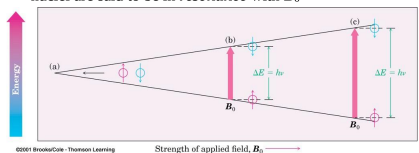
- (a) Normally the nuclear magnetic fields are randomly oriented
- (b) When placed in an external magnetic field $(\mathbf{B_0})$, the nuclear magnetic field can either be aligned with the external magnetic or oppose the external magnetic field



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



Applied EM radiation (radio waves) causes the spin to flip and the nuclei are said to be in *resonance* with $B_{\rm o}$



$$DE = h_n$$
 $DE = \frac{1}{2p}$

 B_o = external magnetic field strength g= gyromagnetic ratio 1H = 26,752 ^{13}C = 6.7

gB h

Note that $\frac{h}{}$ is a constant and is sometimes denoted as $\frac{h}{}$

NMR Active Nuclei: nuclear spin quantum number (I) atomic mass and atomic number

Number of spin states = 2I + 1 (number of possible energy levels)

Even mass nuclei that have even number of neutron have I=0 (NMR inactive)

Even mass nuclei that have odd number of neutrons have an integer spin quantum number ($I=1,\,2,\,3,\,etc$)

Odd mass nuclei have half-integer spin quantum number (I = 1/2, 3/2, 5/2, etc)

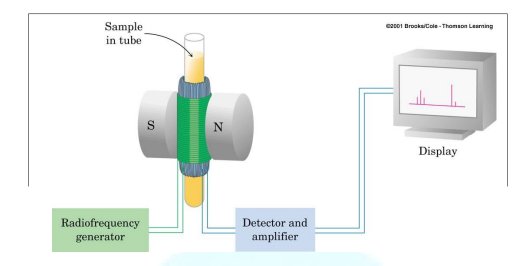
I= 1/2: ¹H, ¹³C, ¹⁹F, ³¹P

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

I=3/2: ¹⁵N

I=0: 12 C, 16 O



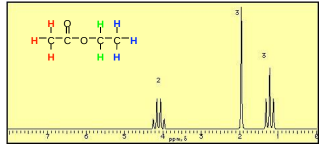


Continuous wave (CW) NMR Pulsed (FT) NMR

Different nuclei absorb EM radiation at different wavelength (energy required to bring about resonance)

Nuclei of a given type, will resonate at different energies depending on their chemical and electronic environment.

The position (*chemical shift*, d) and pattern (*splitting* or *multiplicity*) of the NMR signals gives important information about the chemical environment of the nuclei

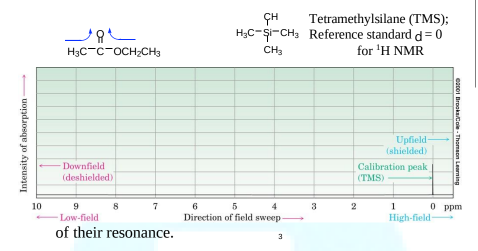


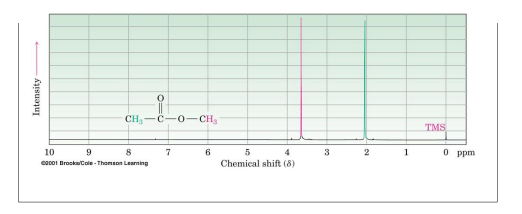
Chemical shift: the exact field strength (in ppm) of a nuclei comes into resonance relative to a reference standard (TMS



Electron clouds "shield" nuclei from the external magnetic field causing then to absorb at slightly higher energy

Shielding: influence of neighboring functional groups on the electronic structure around a nuclei and consequently the chemical shift







Vertical scale= intensity of the signal

Horizontal scale= chemical shift (d), dependent upon the field strength of the external magnetic field; for ¹H, d is usually from 1-10 ppm

MHz

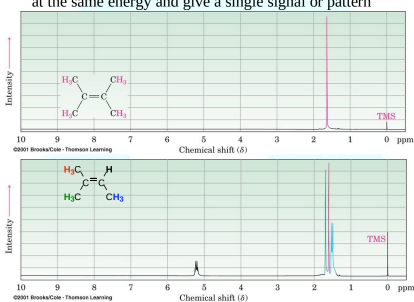
14,100 gauss: 60 MHz for ¹H (60 million hertz) ppm= 60 Hz

15 MHz for ¹³C

140,000 gauss: $600 \text{ MHz for }^{1}\text{H}$ ppm = 600 Hz

150 MHz for ¹³C

Equivalence: chemically and magnetically equivalent nuclei resonate at the same energy and give a single signal or pattern



Test of Equivalence:

- 1. Do a mental substitution of the nuclei you are testing with an arbitrary label (your book uses X)
- 2. Ask what is the relationship of the compounds with the arbitrary label

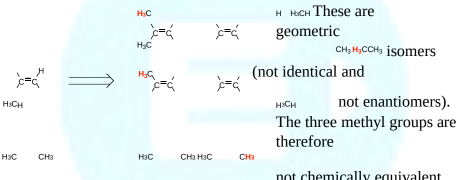


3. If the labeled compounds are identical (or enantiomers), then the original nuclei are chemically equivalent and will normally give rise to a single resonance in the NMR spectra

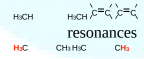
If the labeled compounds are not identical (and not enantiomers), then the original nuclei are not chemically equivalent and can give rise to different resonances in the NMR spectra

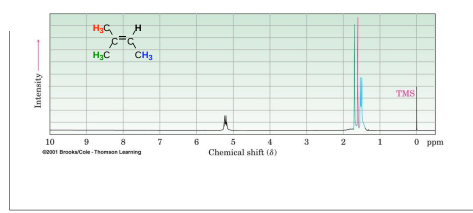
c_{H₃}Identical, so the protons are equivalent

Identical, so the methyl groups are equivalent



not chemically equivalent and *can* give rise to different







Homotopic: equivalent Enantiotopic: equivalent



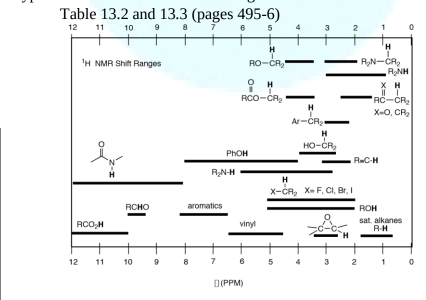
Diastereotopic: non-equivalent

Cyclohexane: two different types of protons, axial and equitorial

The chair-chair interconversion interchanges the axial and equatorial protons and is a fast process at room temperature. NMR is like a camera with a slow shutter speed and a blurred image of fast processes is observed.

At room temperature the cyclohexane protons are observed as a timeaverage and appear as a single resonance. At -90 °C the chairchair interconversion is sufficiently slow that axial and equatorial are observed as two separate resonances.

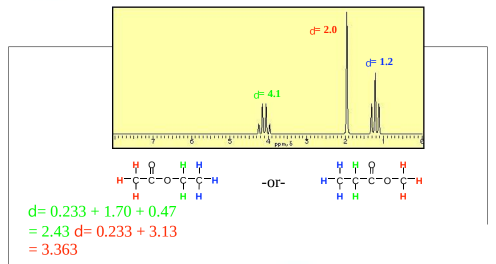
Typical ¹H NMR chemical shifts ranges also see





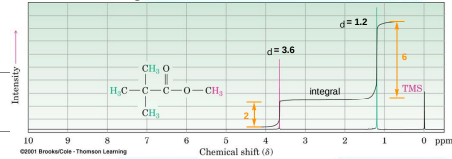
```
The influence of neighboring groups (deshielding) on <sup>1</sup>H chemical
         shifts is additive (to an extent)
Shoolery's additivity rules for predicting the chemical shift of protons
         of the type:y
                                          хсн
                                                                                   хсн
                      for protons on
                                                                 sp3 carbons only
                            d(ppm) = 0.233 + S S_i
 Funtional Group (X,Y)
                              S<sub>i</sub> (ppm)
        -Cl
                               2.53
                               2.33
        -Br
       -1
                               1.82
        -OH
                               2.56
        -OR
                               3.23
                               3.13
               OR
        -SR
                               1.64
        -NR<sub>2</sub>
                              1.57
 -CH<sub>3</sub>
                              0.47
                               2.53
                                                Funtional Group (X,Y)
                                                                             S<sub>i</sub> (ppm)
                                                                1.85
                                                                              1.44
                                                                              1.70
                                                                1.55
                                                                              1.59
                                                        NR<sub>2</sub>
                                                     -CF<sub>3</sub> 1.14
                                                                     -CN
                                                                             1.70
 d = 0.233 + 3.13 + 0.47 = 3.833 d =
 0.233 + 1.70 = 1.933
```





Integration of ¹H NMR resonances

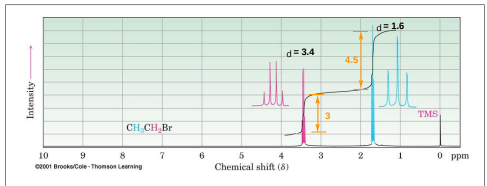
The area under an NMR resonance is proportional to the number of nuclei that give rise to that resonance.

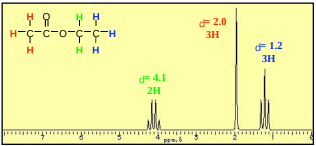


The relative area under the resonances at d= 3.6 and 1.2 is 1:3

The integral is superimposed over the spectrum as a "stair-step" line. The height of each "step" is proportional to the area under the resonance.

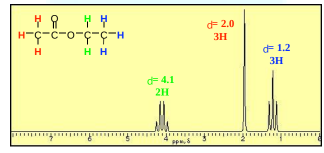






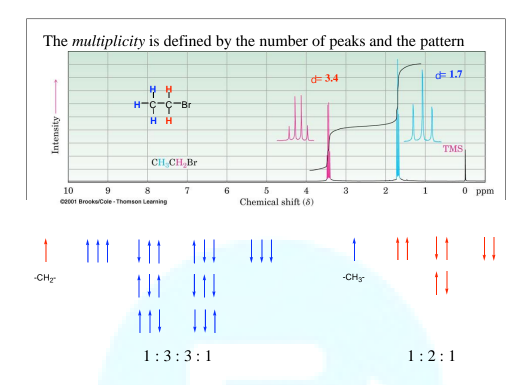
Spin-Spin Coupling (splitting) protons on adjacent carbons will interact and "split" each others resonances into multiple peaks (multiplets)

 $\mathbf{n} + \mathbf{1}$ rule: equivalent protons that have n equivalent protons on the adjacent carbon will be "split" into $\mathbf{n} + \mathbf{1}$ peaks.



Resonances always split each other. The resonance at d=4.1 splits the resonance at d=1.2, therefore the resonance at d=1.2 must split the resonance at d=4.2.



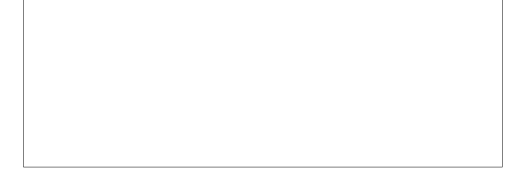


One proton on an adjacent carbon will split a proton into a **doublet** (d), two peaks of 1:1 relative intensity

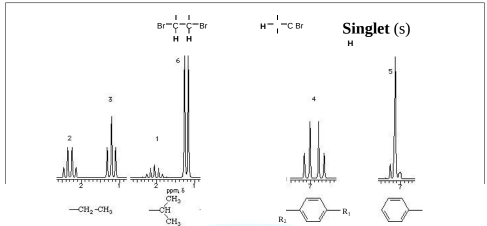
Two proton on an adjacent carbon will split a proton into a **triplet** (t), three peaks of 1:2:1 relative intensity

Three proton on an adjacent carbon will split a proton into a **quartet** (q), four peaks of 1:3:3:1 relative intensity

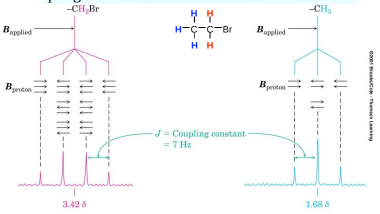
Equivalent protons do not show spin-spin coupling







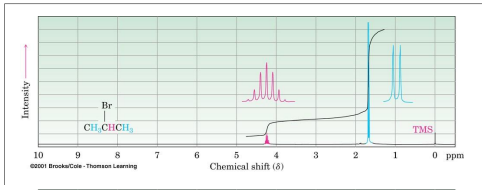
The resonance of a proton with n equivalent protons on the adjacent carbon will be "split" into n+1 peaks with a *coupling constant J*. Coupling constant: distance between peaks of a split pattern; expressed in Hz. Protons coupled to each other have the same coupling constant J.

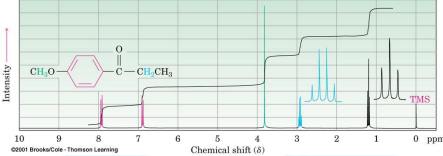


Quartet due to coupling with -CH₃

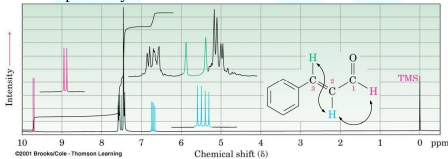
Triplet due to coupling with $-CH_2Br$





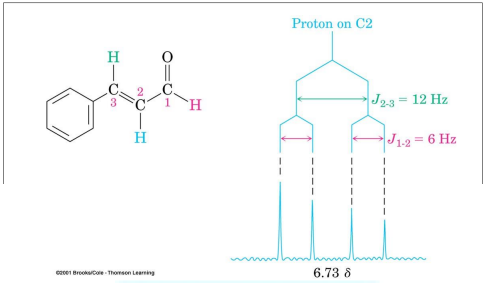


More complex spin-spin coupling: non equivalent protons will couple independently.



 H_2 splits H_3 into a doublet with coupling constant J_{2-3} H_2 splits H_1 into a doublet with coupling constant J_{1-2} H_1 splits H_2 into a doublet; H_3 splits H_2 into a doublet (doublet of doublets) with coupling constants J_{1-2} and J_{2-3} .





Summary of ¹H-¹H spin-spin coupling

- chemically equivalent protons do not exhibit spin-spin coupling to each other.
- the resonance of a proton that has n equivalent protons on the adjacent carbon is split into n+1 peaks (multiplicity) with a coupling constant *J*.
- protons that are coupled to each other have the same coupling constant
- non-equivalent protons will split a common proton independently.complex coupling.

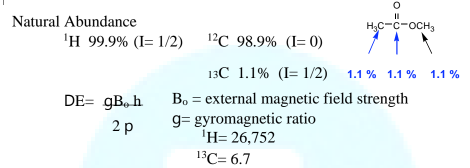
Spin-spin coupling is normally observed between nuclei that are one, two and three bonds away. Four-bond coupling can be observed in certain situations but is not common.

Summary of ¹H-NMR Spectroscopy



- the number of proton resonances equals the number of nonequivalent protons
- the chemical shift (d, ppm) of a proton is diagnostic of the chemical environment (shielding and deshilding)
- Integration: number of equivalent protons giving rise to a resonance
- spin-spin coupling is dependent upon the number of equivalent protons on the adjacent carbon

¹³C NMR Spectroscopy:

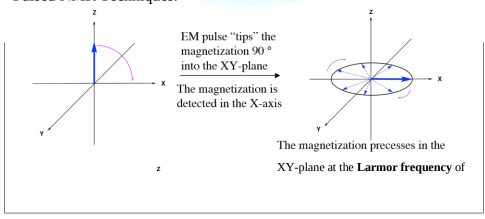


¹³C is a <u>much</u> less sensitive nuclei than ¹H for NMR spectroscopy

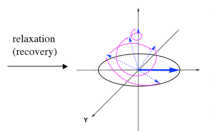
New techniques (hardware and software) has made ¹³C NMR routine

- Pulsed NMR techniques (FT or time domain NMR)
- Signal averaging (improved signal to noise)

Pulsed NMR Techniques:







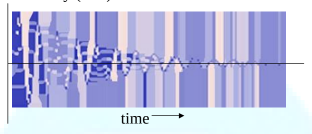
the nuclei, which is directly related to the chemical shift (d) of the nuclei

The magnetization will relax (recover) back to the Z-axis. As the magnetization precesses in XY-plane, it "spirals" back to the Z-axis.

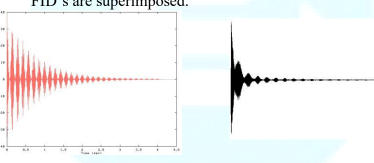
 ${\bf Animation:} \ \underline{{\bf http://www.nmr.ethz.ch/education/PCV/anim/puls_evol.html}}$

http://www.nmr.ucdavis.edu/BCM230_F2001/Flash/PresentationSecondWeek.swf

Free Induction Decay (FID)- time domain NMR



In pulse (FT) NMR, all nuclei are tipped at the same time and the FID's are superimposed.



Fourier Transform (FT) deconvolutes all of the FID's and gives an NMR spectra.

Signal averaging: pulsed NMR allows for many FID's (NMR spectra) to be accumulated over time. These FID's are added together and averaged. Signals (resonances) build up while the "noise" is random and cancels out during the averaging.

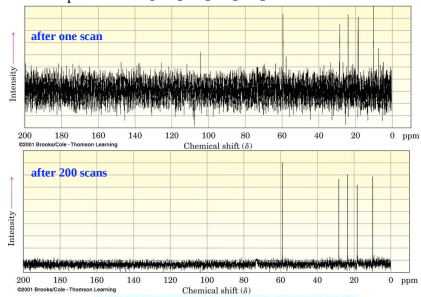


Enhanced signal to noise ratio and allows for NMR spectra to be collected on insensitive nuclei such as ¹C and small samples.

Chemical shifts give an idea of the chemical and electronic environment of the ¹³C nuclei due to shielding and deshielding effects range: 0 - 220 ppm from TMS

Signal-Averaging

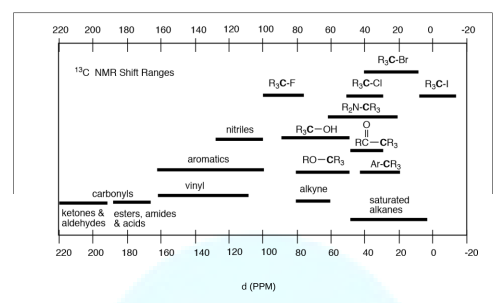


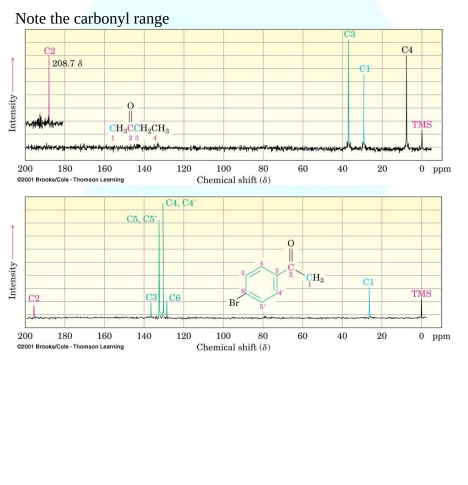


Chemical Shift Range of ${}^{13}\mathrm{C}$

¹ C NMR spectra will give a map of the carbon framework. The number of resonances equales the number of non-equivalent carbons.









¹H-¹³C spin-spin coupling: spin-spin coupling tells how many protons are attached to the ¹³C nuclei. (i.e., primary, secondary tertiary or quaternary carbon)

¹³C spectra are usually collected with the ¹H-¹³C coupling "turned off" (broad band decoupled). In this mode all ¹³C resonances appear as singlets.

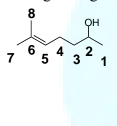
DEPT spectra (**D**istortionless **E**nhancement by **P**olarization **T**ransfer) a modern ¹³C NMR spectra that allows you to determine the number of attached hydrogens.

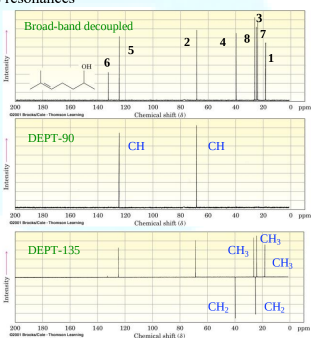
Run: broad-band decoupled spectra

DEPT-90: only CH's show up

DEPT-135: CH's and CH₃'s give positive resonances

CH₂'s give negative resonances





Solving Combined Spectra Problems:



Mass Spectra: Molecular

Formula

Nitrogen Rule Æ # of nitrogen atoms in the molecule

M+1 peak Æ # of carbons

Degrees of Unsaturation: # of rings and/or p-

bonds Infrared Spectra: Functional Groups

C=O

C=C

C≡C

¹H NMR:

Chemical Shift (d) Æ chemical environment of the H's

Integration Æ # of H's giving rise to the resonance

Spin-Spin Coupling (multiplicity) Æ # of non-equivalent H's on the adjacent carbons (vicinal coupling).

Shoolery's Rules: finial check on the structure assignment by ¹H NMR ¹³C NMR:

of resonances Æ symmetry of carbon framework
Type of Carbonyl

Each piece of evidence gives a fragment (puzzle piece) of the structure. Piece the puzzle together to give a proposed structure. The proposed structure should be consistent with all the evidence.

Magnetic Resonance Imaging (MRI): uses the principles of nuclear magnetic resonance to image tissue



MRI normally used the magnetic resonance of protons on water and very sophisticated computer methods to obtain images. Other nuclei within the tissue can also be used (³¹P) or a imaging (contrast) agent can be administered



