

Interpreting ^1H (Proton) NMR Spectra

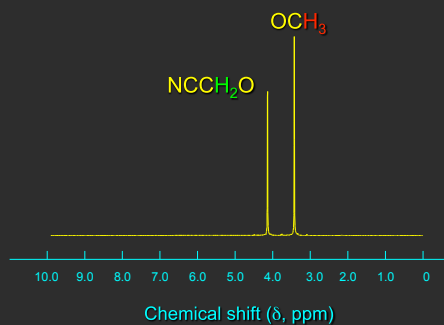
Information contained in an NMR spectrum includes:

1. number of signals
2. their intensity (as measured by area under peak)
3. splitting pattern (multiplicity)

Number of Signals

protons that have different chemical shifts are chemically nonequivalent
exist in different molecular environment

NMR $\text{N}\equiv\text{CCH}_2\text{OCH}_3$

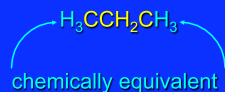


Chemically equivalent protons

are in identical environments

have same chemical shift

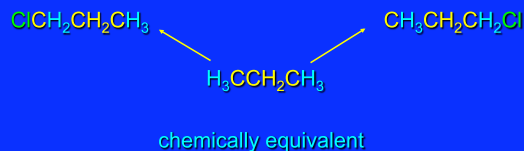
replacement test: replace Hs with an arbitrary "test group" → generates the same compound



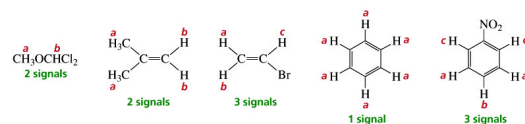
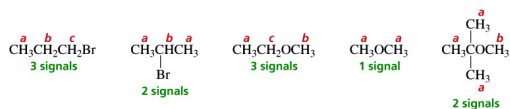
Chemically equivalent protons

Replacing protons at C-1 and C-3 gives same compound (1-chloropropane)

C-1 and C-3 protons are chemically equivalent and have the same chemical shift



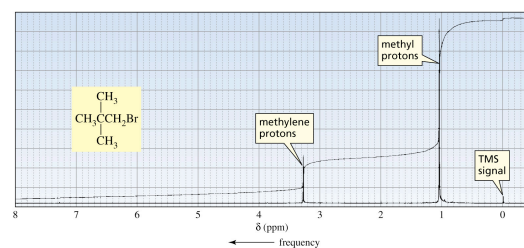
Chemical Shift: Chemically equivalent protons
 Each set of chemically equivalent protons in a compound gives rise to a signal in an ^1H NMR spectrum of that compound



Question

- How many chemically non-equivalent kinds of protons are there in 2,2-dimethylbutane?
- A) 2
- B) 3
- C) 4
- D) 5

^1H NMR spectrum of 1-bromo-2,2-dimethylpropane



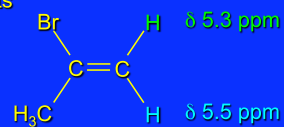
Question

- How many proton signals would you expect to find in the ^1H -NMR spectrum of 2-chloropentane?
- A) 2
- B) 3
- C) 4
- D) 5

Diastereotopic protons

replacement by some arbitrary test group generates diastereomers: non-superimposable, non-mirror images; multiple chiral carbon atoms

diastereotopic protons can have different chemical shifts



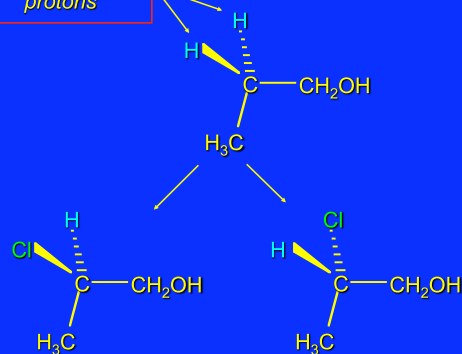
Enantiotopic protons

are in mirror-image environments

replacement by some arbitrary test group
generates enantiomers

enantiotopic protons have the same
chemical shift

Enantiotopic protons



http://chemconnections.org/organic/Movies%20Org%20Flash/nmr-chem-equivalence-Flash/W13_06v3.swf

© To determine whether similar appearing protons are equivalent, substitute another atom "Z" for each of the protons in question. Then determine their relationship and categorize the atoms as *same*, *diastereotopic*, or *enantiotopic*. Consider the green H's then click "Start" to begin.



Definitions

Start

Question

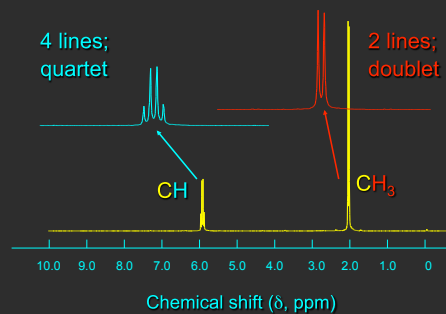
- How many proton signals would you expect to possibly find in the ^1H -NMR spectrum of 2-chloropentane?
- A) 6
- B) 7
- C) 8
- D) 9
- E) More than 9

Spin-Spin Splitting in NMR Spectroscopy

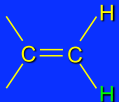
not all peaks are singlets
signals can be split by coupling of
nuclear spins

NMR

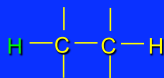
Cl_2CHCH_3



Two-bond and three-bond coupling

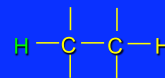
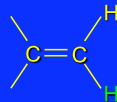


protons separated by
two bonds
(geminal relationship)



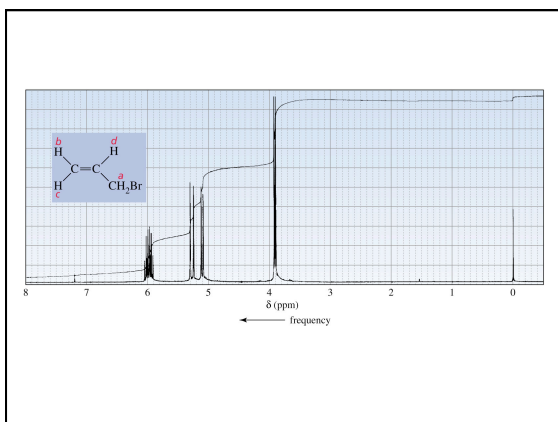
protons separated by
three bonds
(vicinal relationship)

Two-bond and three-bond coupling



in order to observe splitting, protons cannot
have same chemical shift

coupling constant (2J or 3J) is independent
of field strength

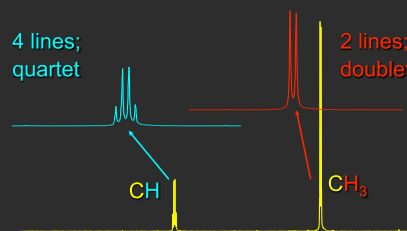


NMR

Cl_2CHCH_3

4 lines;
quartet

2 lines;
doublet

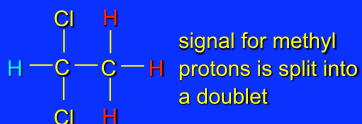


coupled protons are vicinal (three-bond coupling)

CH splits CH₃ into a doublet

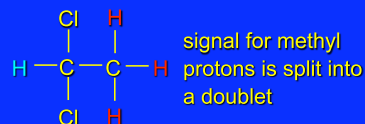
CH₃ splits CH into a quartet

Why do the methyl protons of
1,1-dichloroethane appear as a doublet?



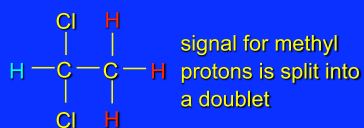
To explain the splitting of the protons at C-2,
we first focus on the two possible spin
orientations of the proton at C-1

Why do the methyl protons of
1,1-dichloroethane appear as a doublet?



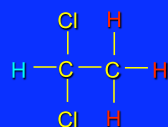
There are two orientations of the nuclear spin
for the proton at C-1. One orientation shields the
protons at C-2; the other deshields the
C-2 protons.

Why do the methyl protons of 1,1-dichloroethane appear as a doublet?



The protons at C-2 "feel" the effect of both the applied magnetic field and the local field resulting from the spin of the C-1 proton.

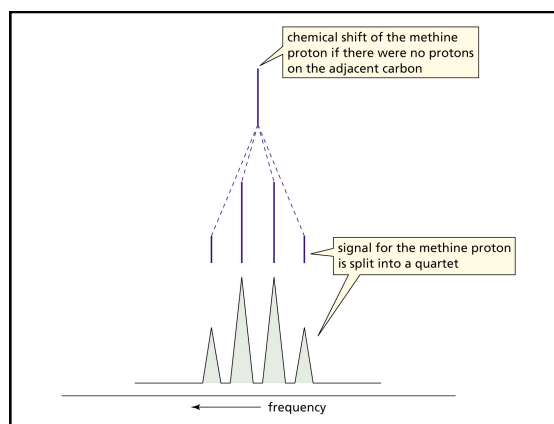
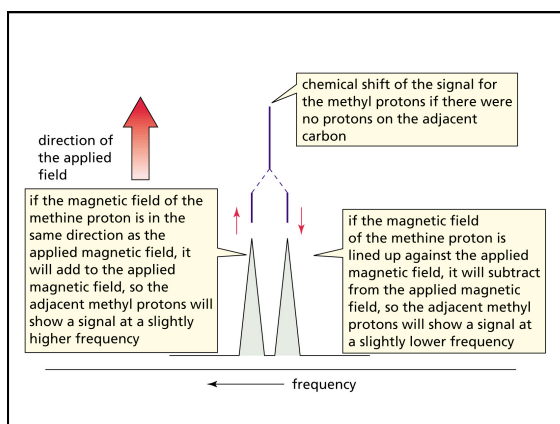
Why do the methyl protons of 1,1-dichloroethane appear as a doublet?



"true" chemical shift of methyl protons (no coupling)

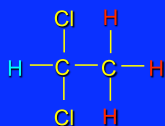
this line corresponds to molecules in which the nuclear spin of the proton at C-1 reinforces the applied field

this line corresponds to molecules in which the nuclear spin of the proton at C-1 opposes the applied field



Why does the methine proton of 1,1-dichloroethane appear as a quartet?

signal for methine proton is split into a quartet



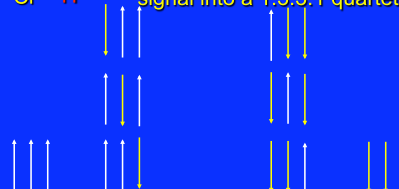
The proton at C-1 "feels" the effect of the applied magnetic field and the local fields resulting from the spin states of the three methyl protons. The possible combinations are shown on the next slide.

Why does the methine proton of 1,1-dichloroethane appear as a quartet?



There are eight combinations of nuclear spins for the three methyl protons.

These 8 combinations split the signal into a 1:3:3:1 quartet.



The splitting rule for ^1H NMR

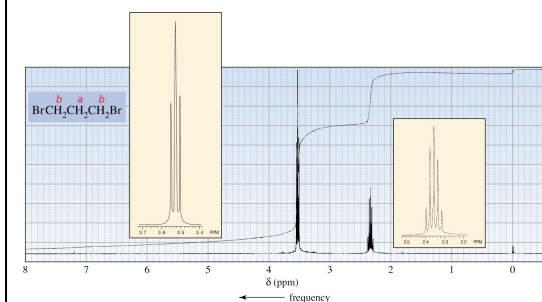
For simple cases, the multiplicity of a signal for a particular proton is equal to the number of chemically equivalent vicinal protons + 1.

Table

Splitting Patterns of Common Multiplets

| Number of equivalent protons to which H is coupled | Appearance of multiplet | Intensities of lines in multiplet |
|--|-------------------------|-----------------------------------|
| 1 | Doublet | 1:1 |
| 2 | Triplet | 1:2:1 |
| 3 | Quartet | 1:3:3:1 |
| 4 | Pentet | 1:4:6:4:1 |
| 5 | Sextet | 1:5:10:10:5:1 |
| 6 | Septet | 1:6:15:20:15:6:1 |

^1H NMR Splitting



Question

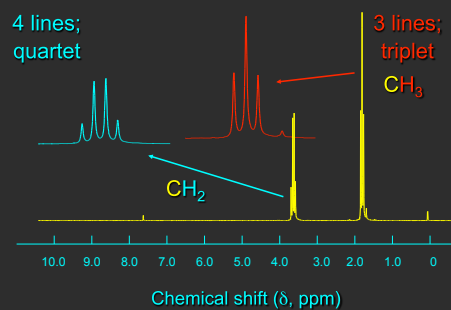
- What is the multiplicity (spin-spin splitting) of the protons of 1,2-dichloroethane?
- A) one singlet
- B) two singlets
- C) one doublet
- D) one triplet

Splitting Patterns: The Ethyl Group

$\text{CH}_3\text{CH}_2\text{X}$ is characterized by a triplet-quartet pattern (quartet at lower field than the triplet)

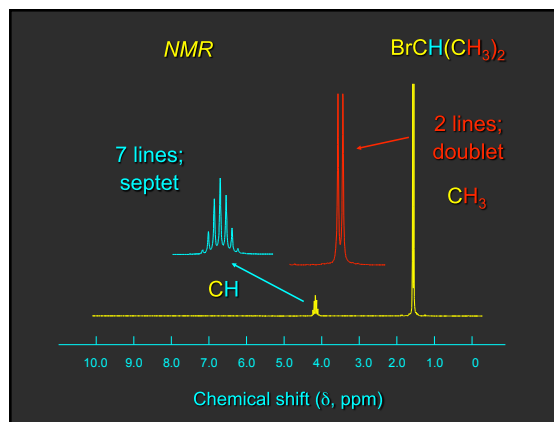
NMR

BrCH_2CH_3



Splitting Patterns: The Isopropyl Group

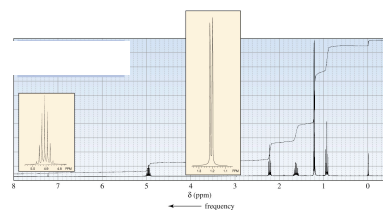
$(\text{CH}_3)_2\text{CHX}$ is characterized by a doublet-septet pattern (septet at lower field than the doublet)



Question

- Which isomer of $\text{C}_4\text{H}_9\text{Br}$ has only one peak in its ^1H -NMR spectrum having the chemical shift δ 1.8?
- A) 1-bromobutane
- B) 2-bromobutane
- C) 1-bromo-2-methylpropane
- D) 2-bromo-2-methylpropane

Question



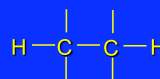
Which compound is the most reasonable for the NMR spectrum above?

- A) Ethyl pentanoate
- B) Propyl isobutyrate
- C) Butyl propanoate
- D) Isopropyl butyrate
- E) Isopentyl acetate

Splitting Patterns: Pairs of Doublets

Splitting patterns are not always symmetrical, but lean in one direction or the other when "coupled".

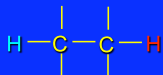
Pairs of Doublets



Consider coupling between two vicinal protons.

If the protons have different chemical shifts, each will split the signal of the other into a doublet.

Pairs of Doublets



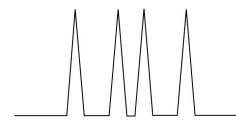
Let $\Delta\nu$ be the difference in chemical shift in Hz between the two hydrogens.

Let J be the coupling constant between them in Hz.

The Difference between a Quartet and a Doublet of Doublets

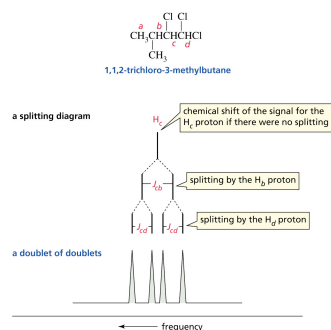


relative intensities: 1 : 3 : 3 : 1



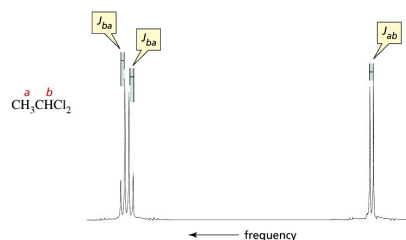
relative intensities: 1 : 1 : 1 : 1

A Splitting Diagram for a Doublet of Doublets



Coupling Constants

The coupling constant (J) is the distance between two adjacent peaks of a split NMR signal in hertz

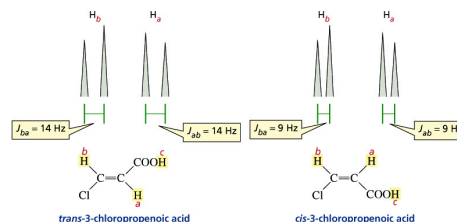


Coupled protons have the same coupling constant

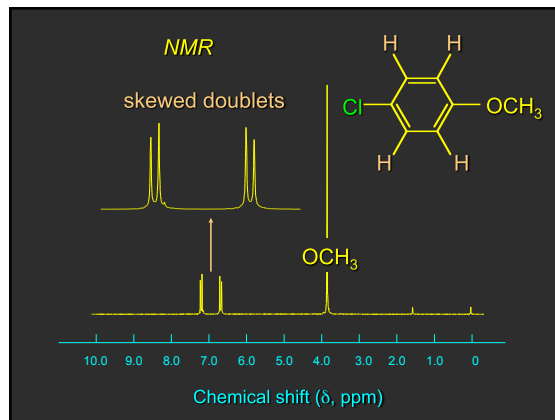
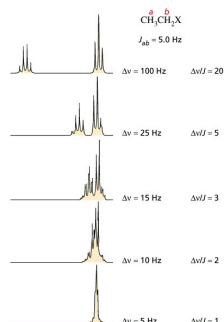
Approximate Values of Coupling Constants

| Approximate value of J_{ab} (Hz) | Approximate value of J_{ab} (Hz) |
|---|---|
| $\text{H}_a-\text{C}-\text{C}-\text{H}_b$ 7 | $\text{H}_a=\text{C}=\text{C}-\text{H}_b$ 15 (trans) |
| $\text{H}_a-\text{C}-\text{C}-\text{H}_b$ 0 | $\text{H}_a=\text{C}=\text{C}-\text{H}_b$ 10 (cis) |
| $\text{H}_a=\text{C}=\text{C}-\text{H}_b$ 2 (geminal coupling) | $\text{H}_a-\text{C}=\text{C}-\text{C}-\text{H}_b$ 1 (long-range coupling) |

A trans coupling constant is greater than a cis coupling constant

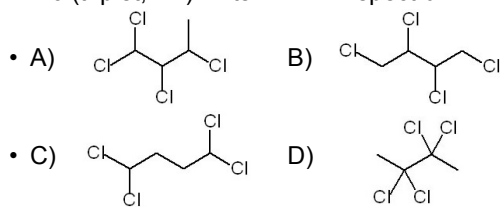


To observe well-defined splitting patterns, the difference in the chemical shifts (in Hz) must be 10 times the coupling constant values



Question

- Which isomer of formula C₄H₆Cl₄ has two signals at δ 3.9 (doublet, 4H) and δ 4.6 (triplet, 2H) in its ¹H-NMR spectrum?

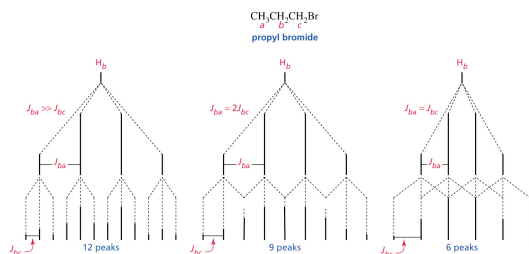


Complex Splitting Patterns

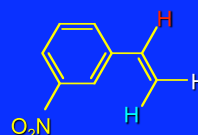
Multiplets of multiplets
aka "MESS-plets" *

* Dr. L. Burns

A Splitting Diagram for a Quartet of Triplets



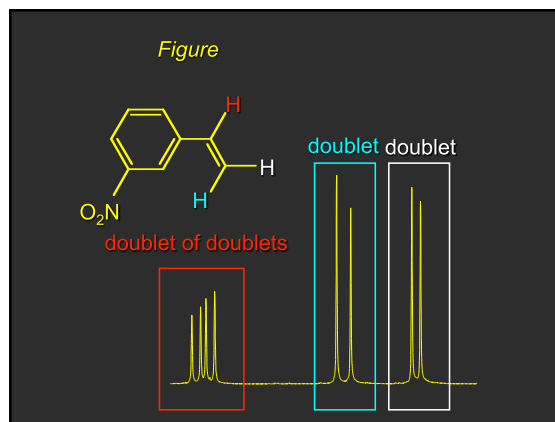
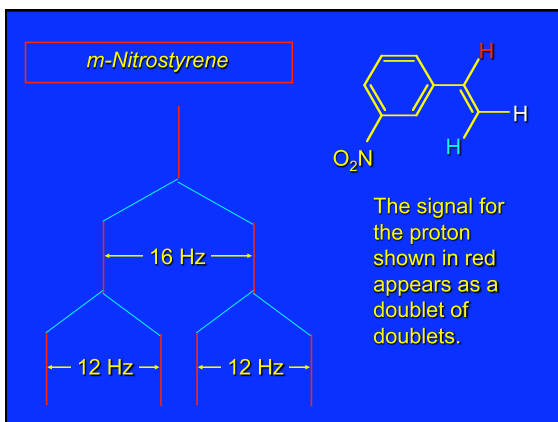
m-Nitrostyrene



Consider the proton shown in red.

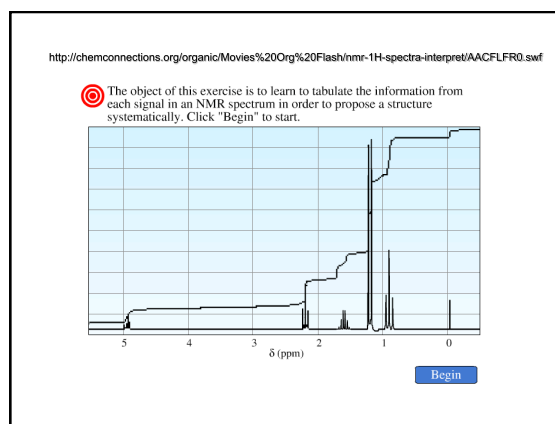
It is unequally coupled to the protons shown in blue and white.

J_{cis} = 12 Hz; J_{trans} = 16 Hz



<http://chemconnections.org/organic/Movies%20Org%20Flash/nmr-splitting.swf>

Click on the splitting pattern in an NMR spectrum that you would observe for the signal of the highlighted region(s).



¹H NMR Spectra of Alcohols & Amines

What about H bonded to O and N?

O—H

H—C—O—H

The chemical shift for O—H is variable (δ 0.5-5 ppm) and depends on temperature and concentration.

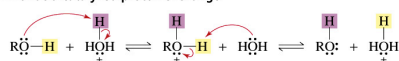
Splitting of the O—H proton is sometimes observed, but **often is not**. It usually appears as a broad peak.

Adding D₂O converts O—H to O—D. The O—H peak disappears.

Protons Bonded to Oxygen and Nitrogen

The greater the extent of the hydrogen bond, the greater the chemical shift

mechanism for acid-catalyzed proton exchange



These protons can undergo proton exchange with deuterium.

They appear as broad signals.

Question

Which of the following statements is false?

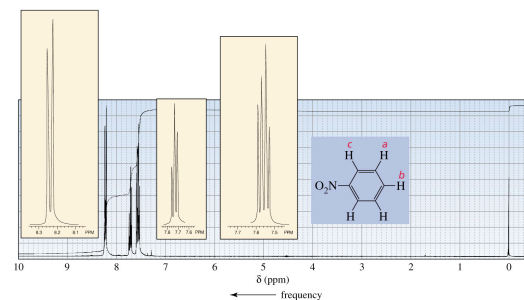
- A) Splitting of the hydroxyl proton of an alcohol is not usually observed.
- B) Alcohol protons shift to lower fields in more concentrated solutions.
- C) Addition of D_2O to alcohol will result in an increased intensity of the hydroxyl proton signal.
- D) The chemical shift of the hydroxyl proton depends on solvent, temperature, and concentration of the solution.

<http://chemconnections.org/organic/Movies%20Org%20Flash/nmr-peak-assignments.swf>

The object of this exercise is to assign each proton in a known chemical structure to its corresponding NMR signal. This procedure is necessary for any products you may isolate in lab. Click "Start" to begin.

Start

The signals for the H_a , H_b , and H_c protons do not overlap



The signals for the H_c , H_d , and H_e protons overlap

