## **BASIC ORGANIC SPECTROSCOPY**

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#### **Electromagnetic Radiation**

is propagated at the speed of light has properties of particles and waves the energy of a photon is proportional to its frequency

#### Figure 13.1: The Electromagnetic Spectrum

#### Shorter Wavelength $(\lambda)$

400 nm

Longer Wavelength  $(\lambda)$ 

<u>75</u>0 nm

Visible Light

Higher Frequency ( $\nu$ )

Lower Frequency (v)

Higher Energy (E)

Lower Energy (E)



Shorter Wavelength ( $\lambda$ ) Longer Wavelength ( $\lambda$ )

Higher Frequency (v)

Lower Frequency ( $\nu$ )

Higher Energy (E)

Lower Energy (E)

Energy

Cosinic rays W REALS <u>X-161/3</u> Ultraviolet light **Visiple light** Infrared radiation Microwaves Radio waxes

Figure 13.1: The Electromagnetic Spectrum

Principles of Molecular Spectroscopy: Quantized Energy States



Electromagnetic radiation is absorbed when the energy of photon corresponds to difference in energy between two states. What Kind of States?

electronicUV-Visvibrationalinfraredrotationalmicrowavenuclear spinradiofrequency

Introduction to <sup>1</sup>H NMR Spectroscopy The nuclei that are most useful to organic chemists are:

<sup>1</sup>H and <sup>13</sup>C

both have spin =  $\pm 1/2$ 

<sup>1</sup>H is 99% at natural abundance

<sup>13</sup>C is 1.1% at natural abundance



A spinning charge, such as the nucleus of <sup>1</sup>H or <sup>13</sup>C, generates a magnetic field. The magnetic field generated by a nucleus of spin +1/2 is opposite in direction from that generated by a nucleus of spin -1/2.

The distribution of nuclear spins is random in the absence of an external magnetic field.





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An external magnetic field causes nuclear 🦟 magnetic moments to align parallel and antiparallel to applied +field. + +

There is a slight excess of nuclear magnetic moments aligned parallel to the applied field.

+

+

+

+

#### Energy Differences Between Nuclear Spin States



no difference in absence of magnetic field proportional to strength of external magnetic field

The frequency of absorbed electromagnetic radiation is proportional to

the energy difference between two nuclear spin states which is proportional to

the applied magnetic field

#### Units

The frequency of absorbed electromagnetic radiation is proportional to

#### Ηz

the energy difference between two nuclear spin states which is proportional to

the applied magnetic field

kJ/mol (kcal/mol)

tesla (T)

The frequency of absorbed electromagnetic radiation is different for different elements, and for different isotopes of the same element.

For a field strength of 4.7 T: <sup>1</sup>H absorbs radiation having a frequency of 200 MHz (200 x 10<sup>6</sup> s<sup>-1</sup>) <sup>13</sup>C absorbs radiation having a frequency of 50.4 MHz (50.4 x 10<sup>6</sup> s<sup>-1</sup>)

The frequency of absorbed electromagnetic radiation for a particular nucleus (such as <sup>1</sup>H) depends on its molecular environment.

This is why NMR is such a useful tool for structure determination.

Nuclear Shielding and <sup>1</sup>H Chemical Shifts

What do we mean by "shielding?" What do we mean by "chemical shift?"

# Shielding

An external magnetic field affects the motion of the electrons in a molecule, inducing a magnetic field within the molecule.





## Shielding

An external magnetic field affects the motion of the electrons in a molecule, inducing a magnetic field within the molecule.

The direction of the induced magnetic field is opposite to that of the applied field.





The induced field shields the nuclei (in this case, C and H) from the applied field.

A stronger external field is needed in order for energy difference between spin states to match energy of rf radiation.



## Chemical Shift

Chemical shift is a measure of the degree to which a nucleus in a molecule is shielded.

Protons in different environments are shielded to greater or lesser degrees; they have different chemical shifts.



# DownfieldUpfieldDecreased shieldingIncreased shielding



Chemical shift ( $\delta$ , ppm) measured relative to TMS



Chemical shift ( $\delta$ , ppm)

### Effects of Molecular Structure on <sup>1</sup>H Chemical Shifts

Protons In Different Environments Experience Different Degrees Of Shielding And Have Different Chemical Shifts Electronegative substituents decrease the shielding of methyl groups

 $CH_3F$  $\delta$  4.3 ppm $CH_3OCH_3$  $\delta$  3.2 ppm $CH_3N(CH_3)_2$  $\delta$  2.2 ppm $CH_3CH_3$  $\delta$  0.9 ppm $CH_3Si(CH_3)_3$  $\delta$  0.0 ppm

Electronegative substituents decrease the shielding of methyl groups

 $CH_3F$  $\delta$  4.3 ppm*least shielded H* $CH_3OCH_3$  $\delta$  3.2 ppm $CH_3N(CH_3)_2$  $\delta$  2.2 ppm $CH_3CH_3$  $\delta$  0.9 ppm $CH_3Si(CH_3)_3$  $\delta$  0.0 ppm

Effect is cumulative

 $CHCI_3$  $CH_2CI_2$  $CH_3CI$ 

δ 7.3 ppm
δ 5.3 ppm
δ 3.1 ppm

Protons attached to sp<sup>2</sup> hybridized carbon are less shielded than those attached to sp<sup>3</sup> hybridized carbon



Table 13.1 (p 496)



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Type of proton Chemical shift ( $\delta$ ), ppm

H—NR 1-3

H—OR 0.5-5

H—OAr 6-8

О || HO—C— 10-13

