

ORGANIC STRUCTURE DETERMINATION

How do we know:

- · How atoms are connected together?
- Which bonds are single, double, or triple?
- What functional groups exist in the molecule?
- · If we have a specific stereoisomer?

The field of organic structure determination attempts to answer these questions.

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Infrared Spectroscopy

- Four common spectroscopic techniques used to determine structure:
 - Infrared Spectroscopy (IR)
 - Mass Spectrometry (MS or Mass Spec)
 - Nuclear Magnetic Resonance Spectroscopy (NMR)
 - Ultraviolet Spectroscopy

INSTRUMENTAL METHODS OF STRUCTURE DETERMINATION

- Nuclear Magnetic Resonance (NMR) Excitation of the nucleus of atoms through radiofrequency irradiation. Provides extensive information about molecular structure and atom connectivity.
- Infrared Spectroscopy (IR) Triggering molecular vibrations through irradiation with infrared light. Provides mostly information about the presence or absence of certain functional groups.
- Mass spectrometry Bombardment of the sample with electrons and detection of resulting molecular fragments. Provides information about molecular mass and atom connectivity.
- 4. Ultraviolet spectroscopy (UV) Promotion of electrons to higher energy levels through irradiation of the molecule with ultraviolet light. Provides mostly information about the presence of conjugated π systems and the presence of double and triple bonds.

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SPECTRUM INTERPRETATION PROCESS

- 1. Recognize a *pattern*.
- 2. Associate patterns with physical parameters.
- 3. Identify possible meanings, i.e. *propose explanations.*

Once a spectrum is obtained, the main challenge is to extract the information it contains in abstract, or hidden form. This requires the recognition of certain patterns, the association of these patterns with physical parameters, and the interpretation of these patterns in terms of meaningful and logical explanations.

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ELECTROMAGNETIC SPECTRUM

Most organic spectroscopy uses $\ensuremath{\textbf{electromagnetic}}$ energy, or radiation, as the physical stimulus.

Electromagnetic energy (such as visible light) has no detectable mass component. In other words, it can be referred to as "pure energy."

Other types of radiation such as alpha rays, which consist of helium nuclei, have a detectable mass component and therefore cannot be categorized as electromagnetic energy.

The important parameters associated with electromagnetic radiation are:

- Energy (E): Energy is directly proportional to frequency, and inversely proportional to wavelength, as indicated by the equation below.
- Frequency (μ)
 Wavelength (λ)

 $E = h\mu$





INFRARED SPECTROSCOPY

- Infrared radiation stimulates molecular vibrations.
- Infrared spectra are traditionally displayed as %T (percent transmittance) versus wave number (4000-400 cm-1).
- Useful in identifying presence or absence of functional groups.

Infrared radiation is largely thermal energy. It induces stronger molecular vibrations in covalent bonds, which can be viewed as springs holding together two masses, or atoms. Specific bonds respond to (absorb) specific frequencies equilibrium bond length www.nilmishra.name

















INFORMATION OBTAINED FROM IR SPECTRA

- IR is most useful in providing information about the presence or absence of specific functional groups.
- IR can provide a molecular fingerprint that can be used when comparing samples. If two pure samples display the same IR spectrum it can be argued that they are the same compound.
- IR does not provide detailed information or proof of molecular formula or structure. It provides information on molecular fragments, specifically functional groups.
- Therefore it is very limited in scope, and must be used in conjunction with other techniques to provide a more complete picture of the molecular structure.

















IR SPECTRUM OF ALKYNES

The most prominent band in alkynes corresponds to the **carbon-carbon triple bond**. It shows as a sharp, weak band at about **2100** cm⁻¹. The reason it's weak is because the triple bond is not very polar. In some cases, such as in highly symmetrical alkynes, it may not show at all due to the low polarity of the triple bond associated with those alkynes.

Terminal alkynes, that is to say those where the triple bond is at the end of a carbon chain, have C-H bonds involving the *sp* carbon (the carbon that forms part of the triple bond). Therefore they may also show a sharp, weak band at about **3300 cm**⁻¹ corresponding to the C-H stretch.

















- 1745 cm⁻¹

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