

Chemistry 2

Lecture 8

IR Spectroscopy of Polyatomic Molecules



Assumed knowledge

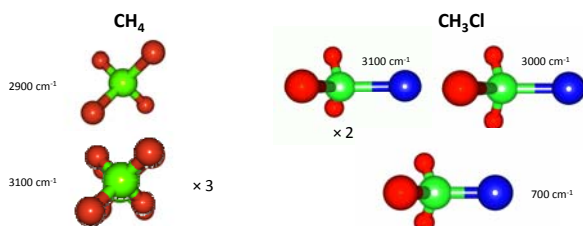
There are $3N-6$ vibrations in a non-linear molecule and $3N-5$ vibrations in a linear molecule. Only modes that lead to a change in the dipole moment are IR active. For each mode, bands with $\Delta v = \pm 1, 2, 3$ are possible but the overtones ($\Delta v > 1$) are increasingly weak. Combination bands where more than one mode is simultaneously excited are also possible.

Learning outcomes

- Be able to pick the vibrational modes in a molecule which are likely to be coupled and those which are likely to be local modes
- Be able to explain the appearance of IR spectra in terms of (strong) fundamental transitions and weaker overtones and combinations
- Be able to use Jablonski diagrams to assign spectra

Local Modes

- Although all vibrations in a molecule can couple, for a number of normal modes the bulk of the vibrational amplitude can be found on a small number of atoms.
- This frequently occurs when the natural frequency of an individual vibration is does not closely match that of other vibrations in the molecule



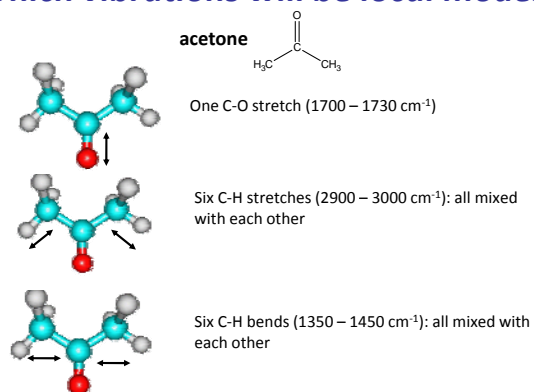
- All C-H equivalent: all C-H stretches are completely mixed
- All C-H equivalent: all C-H stretches are completely mixed
- C-Cl frequency is much lower: almost completely uncoupled

How can you pick local modes?

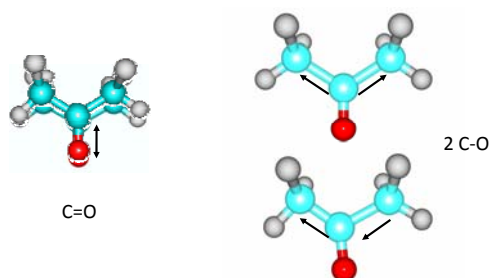
2 main things to look for:

1. Vibration involves atom(s) with only one bond: this means the atom vibrates into space, rather than against other atoms.
2. The vibrating atom(s) have a different natural vibrational frequency to the connected groups (atoms and bonds):- this means their vibration is OUT OF RESONANCE with their neighbors.

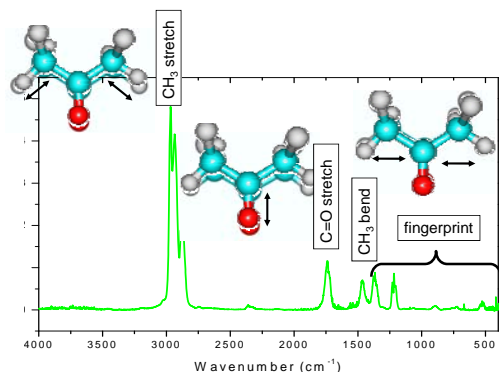
Which vibrations will be local modes?



Acetone: C-O and C-C stretches



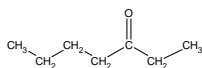
IR Spectrum of Acetone



Which vibrations will be local modes?



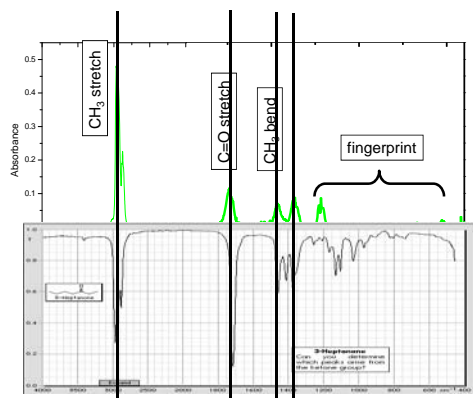
acetone



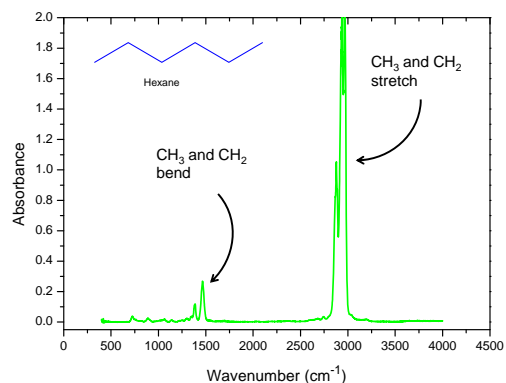
3-heptanone

- What type of vibrations will be local modes (with characteristic frequencies)?

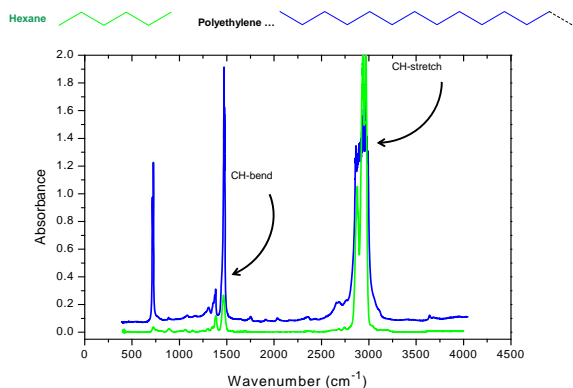
IR Spectrum of acetone and 3-heptanone



Local vs normal modes in hexane



Hexane vs polyethylene



Jablonski diagrams

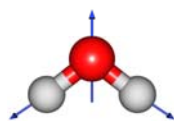
At least for simple molecules, a full analysis of the vibrational spectrum is often possible:

- Work out the number (3N-6) and IR activity of the vibrations
- The strongest bands are the *fundamentals*: single excitations of these modes ($\Delta v = 1$)
- First, second, third... overtones: $\Delta v = 2, 3, 4...$ with rapidly decreasing intensity
- Combinations: simultaneous excitation of different vibrations: $\Delta v = 1, 2, 3, 4$ for each mode with rapidly decreasing intensity

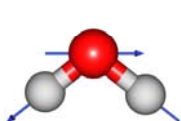
Normal Modes of H_2O

Polyatomic molecules can vibrate in many different ways

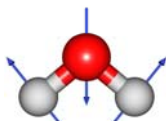
- H_2O contains 3 atoms so has $3N - 6$ ($3 \times 3 - 6$) = 3 vibrations
- Each is IR active



symmetric stretch
 $\nu_1 = 3657 \text{ cm}^{-1}$



asymmetric stretch
 $\nu_3 = 3756 \text{ cm}^{-1}$

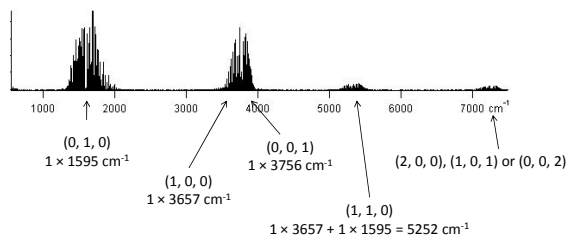


bend
 $\nu_2 = 1595 \text{ cm}^{-1}$

Vibrational Spectrum of H_2O

Polyatomic molecules can vibrate in many different ways

- $\nu_1 = 3657 \text{ cm}^{-1}$, $\nu_2 = 1595 \text{ cm}^{-1}$, $\nu_3 = 3756 \text{ cm}^{-1}$,
- Label excitations as (ν_1, ν_2, ν_3)
- As first approximation, neglect effect of anharmonicity on energies

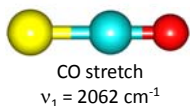


- Overtones carry on into visible with decreasing absorption

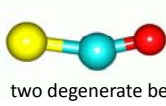
Vibrational Spectrum of OCS

Polyatomic molecules can vibrate in many different ways

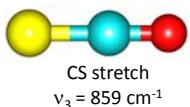
- Linear so $3N - 5 = 4$ vibrations
- Unlike CO_2 , the $C=O$ and $C=S$ natural frequencies are different leading to incomplete mixing:



CO stretch
 $\nu_1 = 2062 \text{ cm}^{-1}$

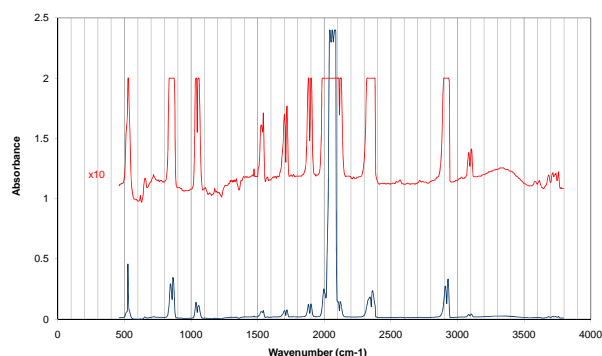


two degenerate bends
 $\nu_2 = 520 \text{ cm}^{-1}$



CS stretch
 $\nu_3 = 859 \text{ cm}^{-1}$

Vibrational Spectrum of OCS



Next lecture

- Raman spectroscopy

Week 12 tutorial and homework

- Spectroscopy worksheet in the tutorials
- Complete the practice questions from the lectures

Practice Questions

1. Assign the spectrum of OCS in as much detail as you can.
2. Two infrared spectra of hexane (C_6H_{14}) are shown overleaf. The top spectrum is a conventional mid-IR spectrum with assignment of the two main transitions indicated. The bottom spectrum extends from the mid-IR to the visible and shows a number of CH-stretch overtone transitions with ever decreasing intensity (note the scale factor for the intensity).
 - a) Propose an assignment for the unidentified peak at about 4400 cm^{-1} in the bottom spectrum
 - b) The wavenumber of the fundamental CH-stretch transition from the top spectrum is $\nu = 2962\text{ cm}^{-1}$. The wavenumber of the fourth CH-stretch overtone (i.e. the $\nu = 0 \rightarrow 5$ transition marked "5*"), from the bottom spectrum, is $\nu = 13401\text{ cm}^{-1}$. Calculate the harmonic constant, ω_e , and the anharmonicity constant, $\omega_e x_e$ for the CH bond.
 - c) Estimate the bond dissociation energy, D_0 , of the CH bond in hexane.
 - d) The tabulated CH bond energy is 411 kJ mol^{-1} . Comment on the agreement or disagreement with your answer to (c)

