

ال vibrational mode معناها كم حركة متوقعة لكل molecule في
عندي قانونين إذا كان المركب linear أو nonlinear إذا كان linear
القانون يكون $3N-5$ اذا كان nonlinear رح يكون $3N-6$ من وين جابو
رقم 3 رقم 3 معناه عدد الإحداثيات تبع الحركات وهو x,y,z ال N معناها
عدد ال atom الموجودة بال molecule معنى الرقم 6 بال nonlinear
هو ال 3rotation حركات وال حركتين وال 3transition حركات
بصير 6 بال linear في حركة وحدة بال rotation لانو حركة وحدة
خلتو active مثل ما قلنا عن CO_2 فبنقص من ال 6 واحد بصير 5..

Vibrational Modes

- Number of possible modes
 - Nonlinear molecule: $3N - 6$
 - Linear molecule: $3N - 5$
- 3 degrees of freedom – i.e., 3 coordinates in space
- 3 translations and 3 rotations account for 6 motions of molecule
- Rotation about center bond in linear molecule is indistinguishable



- Remaining degrees of motion represent vibrational motion (i.e., number of vibrations within the molecule)

Calculate Number of Vibrational Modes

The degrees of vibrational modes for linear molecules can be calculated using the formula:

$$3n - 5 \text{-----}(1)$$

The degrees of freedom for nonlinear molecules can be calculated using the formula:

$$3n - 6 \text{-----}(2)$$

n is equal to the number of atoms within the molecule of interest.

The following procedure should be followed when trying to calculate the number of vibrational modes:

- 1- Determine if the molecule is linear or nonlinear (i.e. Draw out molecule using VSEPR). If linear, use Equation 1. If nonlinear, use Equation 2
- 2- Calculate how many atoms are in your molecule. This is your n value. Plug in your n value and solve.

Example 1: CS₂

An example of a linear molecule would be CS₂. There are a total of 3 atoms in this molecule. Therefore, to calculate the number of vibrational modes, it would be:

$$3(3)-5 = 4 \text{ vibrational modes.}$$

Example 2: CCl₄

هاد المركب مش active على ال IR لكن اذا طلب ال vibrational mode إلو بعمل على القانون

CCl₄ is a nonlinear molecule. In this molecule, there are a total of 5 atoms.

Therefore, there are $3(5)-6 = 9$ vibrational modes.

Example 2: H₂O

H₂O is a nonlinear molecule. In this molecule, there are a total of 3 atoms.

Therefore, there are $3(3)-6 = 3$ vibrational modes.

How many vibrational modes?

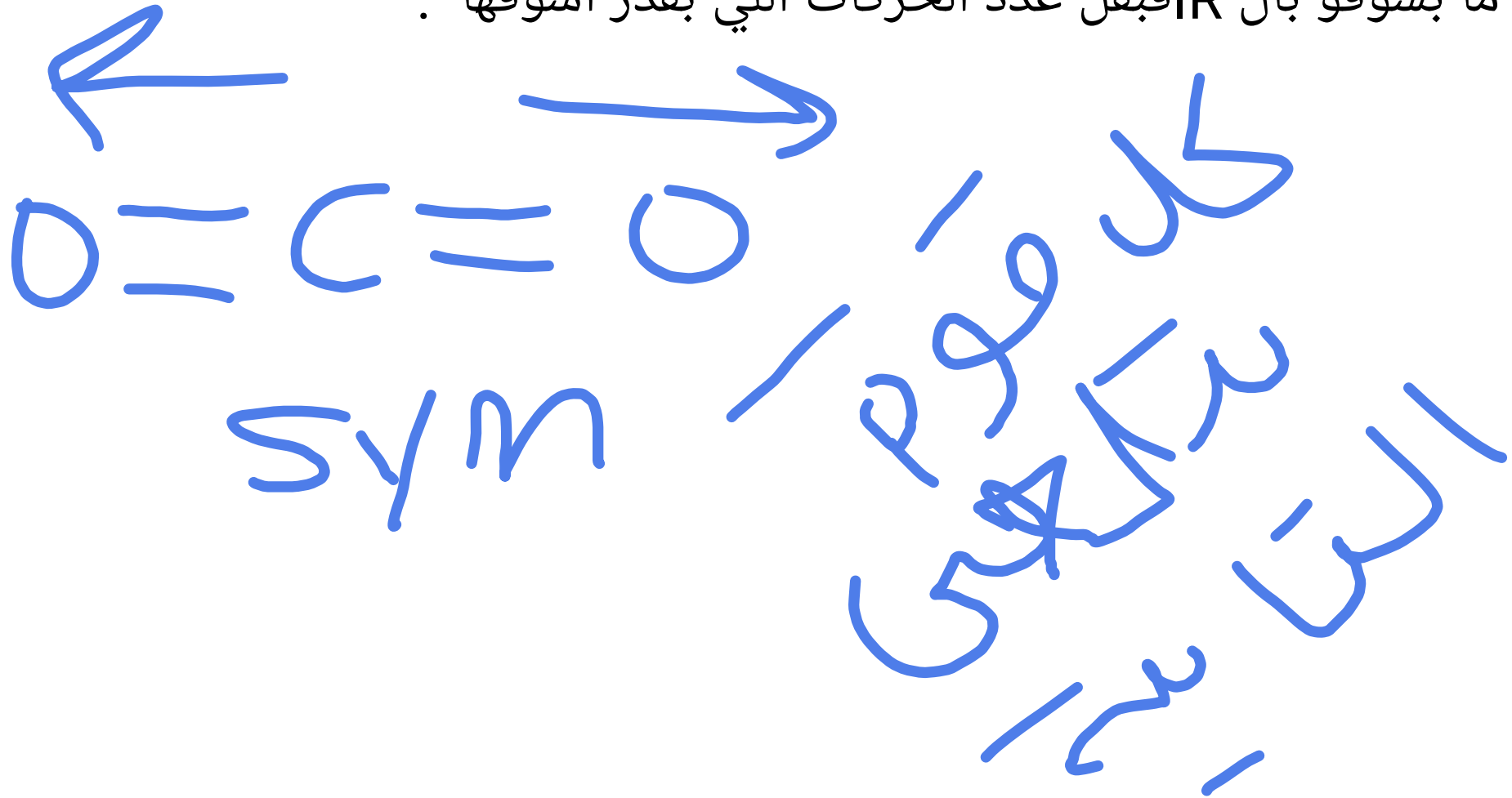
2 atoms (H_2) - 1 vibration (stretch ν)

3 atoms (H_2O) - 3 vibrations (ν_s , ν_{as} , σ)

3 atoms (CO_2) - 4 vibrations (ν_s , ν_{as} , σ , ω)

4 atoms (H_2CO) - 6 vibrations (ν_s , ν_{as} , σ , ω , $\rho(\text{CH}_2)$, $\nu(\text{C=O})$)

في عوامل بتزيد من أو بتقلل الحركات عن النظري أو ال peak بتكون أقل أو أكثر أول أشي رح نحكي عنو هو ال fewer peak والعوامل المؤثره عليها ال symmetric بمعنى آخر إنو ال symmetric ما بأثر على ال dipole بالتالي ما بشوفو بال IR فبقل عدد الحركات اللي بقدر أشوفها .



تاني عامل إنو صح بكون عندي ست حركات لكن هاي الحركات البعض منها طاقتها أقل من طاقة ال IR بالتالي ما بظهورو كلهم اول مثلا بكون ال compound في identical group بالتالي لما يصير في absorption رح تظهر زي كأنها واحد ..

Factors Influence the Normal Modes

Four factors tend to produce **fewer experimental peaks** than would be expected from the **theoretical** number of normal modes. (1) the symmetry of the molecules is such that no change in dipole results from a particular vibration; (2) the energies of two or more vibrations are identical or nearly identical; (3) the absorption intensity is so low as to be undetectable by ordinary means; or the vibrational energy is in a wavelength region **beyond** the range of the instrument

العوامل التي بتعطيني more peak هم اول وحدة ال over tone
معناه إنو ال molecule صار ال excitation لل vibrational level
بس ما نتقل من v_0 ل v_1 لا انتقل ل v_2 او v_3 هسا اذا رجع واعطاني
طاقة للإلكترون اللي جنبو وبدأ يصعد فيهم بنسميه coupling mode

Fewer and more experimental peaks than calculated

- Fewer peaks

- (1) the symmetry of the molecules is such that no change in dipole results from a particular vibration
- (2) the energies of two or more vibrations are identical or nearly identical
- (3) the absorption intensity is so low as to be undetectable by ordinary means

- More peaks

- (1) Overtone
- (2) Combination bands

Fundamental Peaks and Overtones

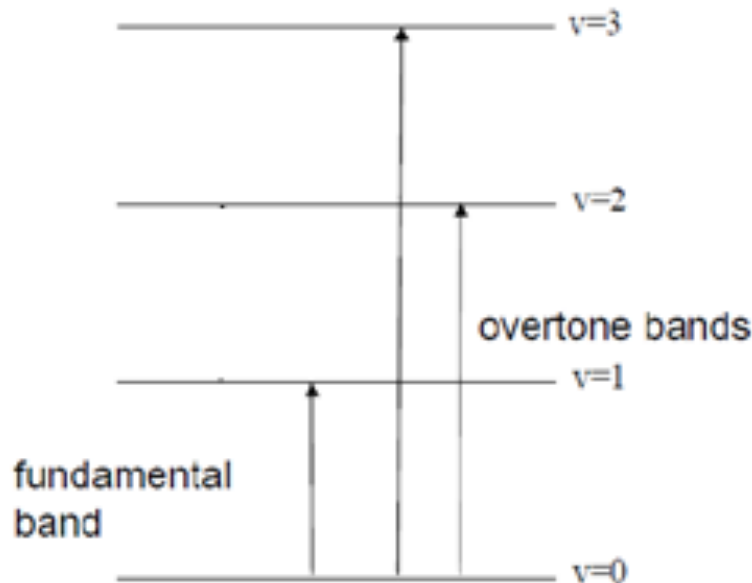
Fundamental transition:

The excitation from the ground state V_0 to the first excited state V_1 is called the fundamental transition. It is the most likely transition to occur.

Fundamental absorption bands are strong compared with other bands that may appear in the spectrum due to overtone, combination, and difference bands.

Overtone bands:

Result from excitation from the ground state to higher energy states V_2 , V_3 , and so on.



- If the fundamental absorption occurs at frequency ν , the overtones will appear at about 2ν , 3ν , and so on.
- Overtones are weak bands and may not be observed under real experimental conditions.

Coupling modes:

- Vibrating atoms may interact with each other. Two vibrational frequencies may couple to produce a new frequency $\nu_3 = \nu_1 + \nu_2$. The band at ν_3 is called a combination band.
- If two frequencies couple such that $\nu_3 = \nu_1 - \nu_2$, the band is called a difference band.
- Not all possible combinations and differences occur.

Factors that affect the frequency of light absorbed

1. Bonds which have one lighter and one heavier atom vibrate faster than bonds which have two heavier atoms.

العوامل اللي بتأثر على ال frequency

هون ثبتنا ال single bond وغيرنا ال atom اذا كانو مختلفات الطاقة بتزيد

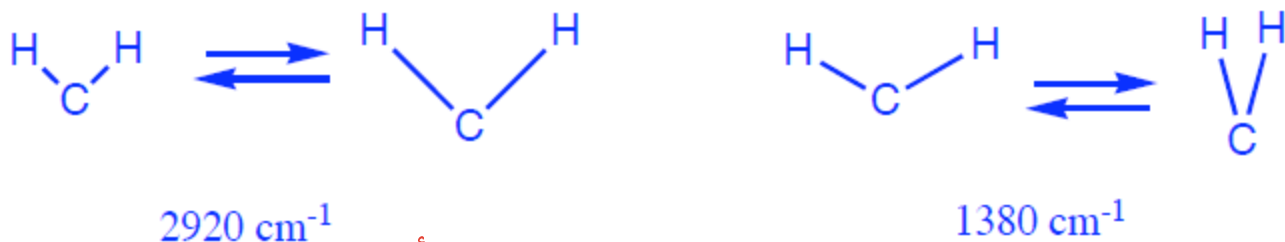
$\text{C}-\text{H}$	$\text{C}-\text{C}$
3000 cm^{-1}	1200 cm^{-1}

2. Stronger bonds vibrate faster than weaker bonds.

هون ثبتنا ال atom وزدنا عدد ال bond كل ما زاد العدد كل ما زادت الطاقة

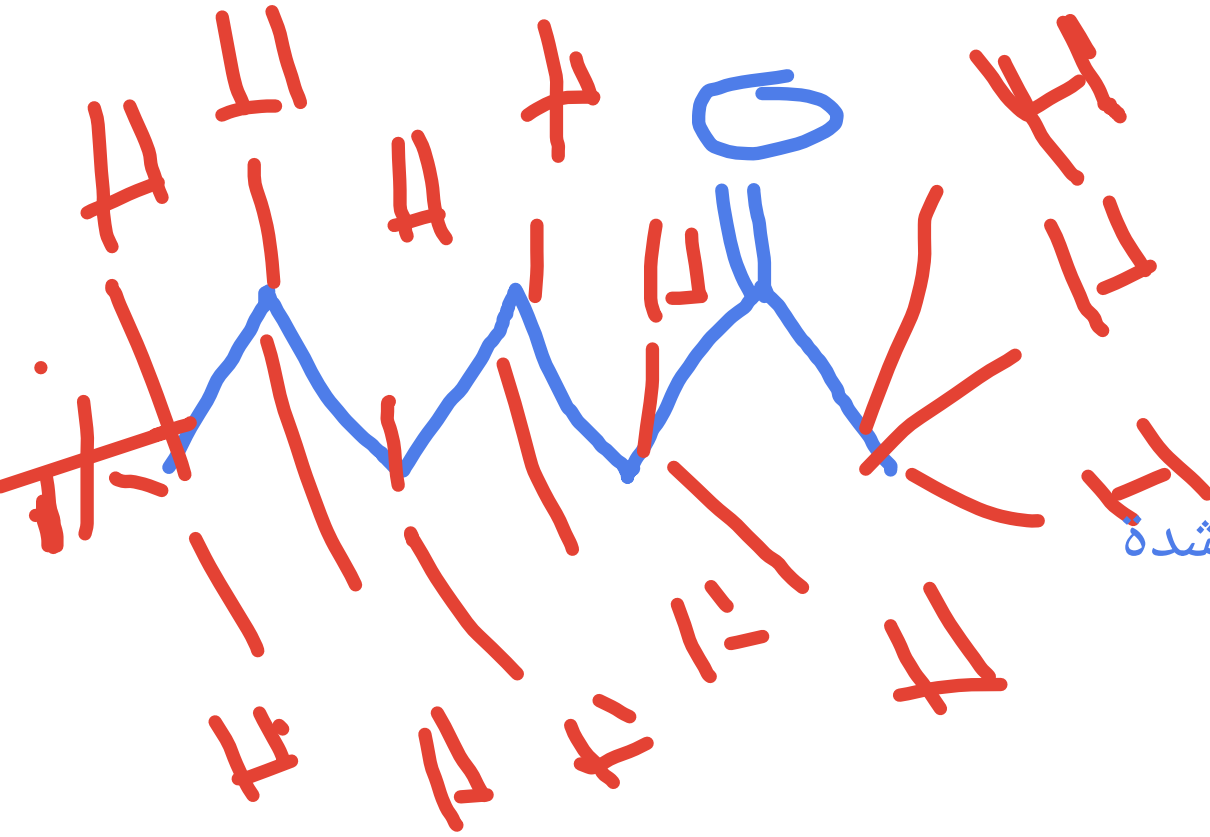
$\text{C}\equiv\text{C}$	$\text{C}=\text{C}$	$\text{C}-\text{C}$
2200 cm^{-1}	1660 cm^{-1}	1200 cm^{-1}

3. Stretching vibrations are faster than bending vibrations

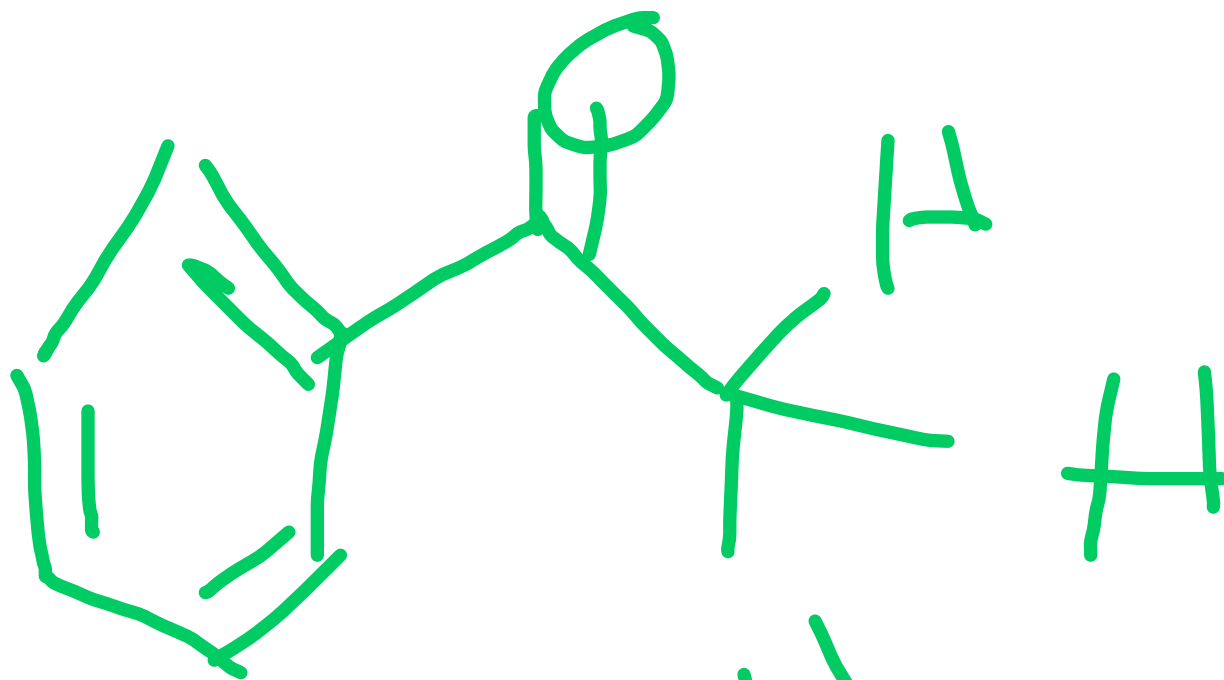


هون غيرنا ما بين ال stretch وال bend ال stretch أعلى من ال bend

كل ما زاد عدد ال C-H كل ما زادت الشدة تبعت ال absorption..



عدد ال C-H كبير رح تزيد شدة
الامتصاص ..



36 C-H bonds
↓
Ads



polar

↑ Abs

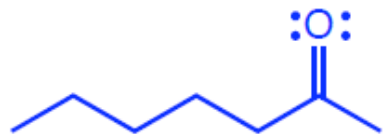


non
polar

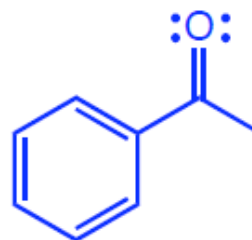
↓ abs

Factors that affect the amount of light that is absorbed

1. If there are many C-H bonds in a molecule absorbing light at the same frequency, the band will be much larger than if there are only a few.

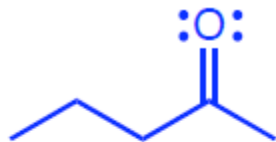


lots of C-H bonds
large absorption



fewer C-H bonds
smaller absorption

2. Strength of the dipole - bonds which have a strong dipole moment will absorb light more strongly than those which have a weaker dipole moment.



C=O very polar, strong dipole
large absorption



C=C nonpolar, small dipole
small absorption

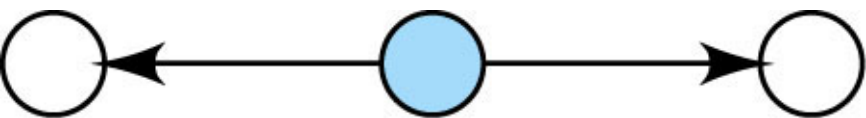
ال CO2 حكيما انو بس حركة بال stretch وحكيما ال bending بظهر
كمان فيه مع انو لو اجيت حسبت على القانون اللي حكيماه قبل شوي
لازم يظهر 4 حركات لكن بالواقع بس حركتين وحدة wave number
إلها 2330 والثانية 667 وهون بنعرف انو الأعلى طاقة هي
asymmetric وال bending الأقل... في ال H2O على القانون لازم
يظهر 3 حركات وفعليا هداول بظهروا بالواقع ال stretch ال wave
number الهم 3650 والثانية 3760 وال bending 1595 وبال
stretch بنعرف انو الأعلى طاقة هو ال asymmetric والاقل بشوي
symmetric

CO₂ Molecule

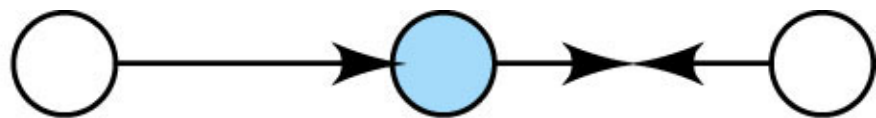
Let us consider the infrared spectrum of carbon dioxide. If no coupling occurred between the two C=O bonds, an absorption peak would be expected at the same wavenumber as the peak for the C=O stretching vibration in an aliphatic ketone (about 1700 cm⁻¹). Experimentally, carbon dioxide exhibits two absorption peaks, the one at 2330 cm⁻¹ and the other at 667 cm⁻¹. Carbon dioxide is a linear molecule and thus has $3 \times 3 - 5 = 4$ normal modes. Two stretching vibrations are possible. The symmetric vibration causes no change in dipole. Thus, the symmetric vibration is infrared inactive.

The asymmetric vibration produce a change in dipole moments, so absorption at **2330** cm^{-1} results.

The remaining two vibrational modes of carbon dioxide involve scissoring. The two bending vibrations are the resolved components at 90 deg to one another of the bending motion in all possible planes around the bond axis. The two vibrations are identical in energy and thus produce a single peak at **667** cm^{-1} .



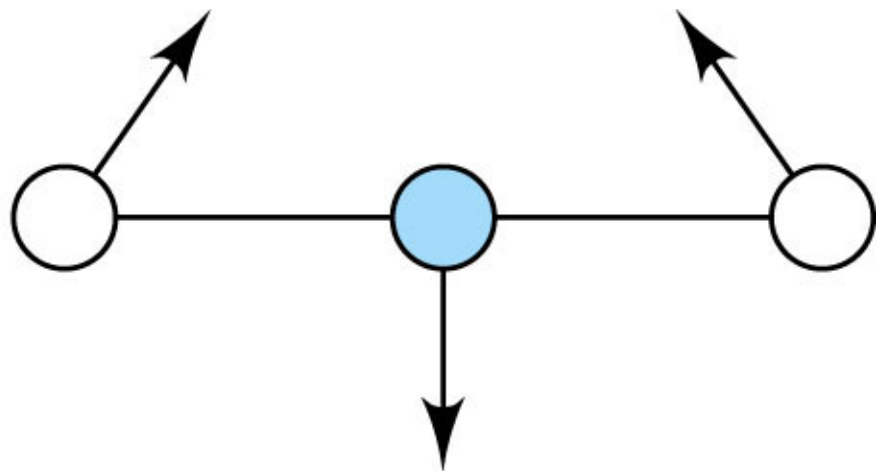
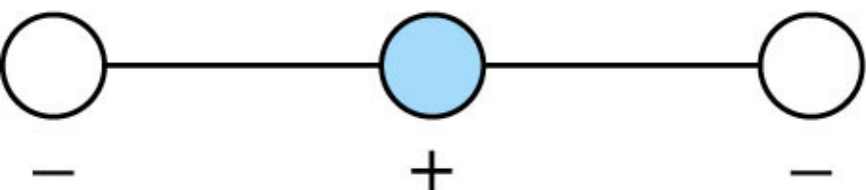
Symmetric



Asymmetric

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Stretching

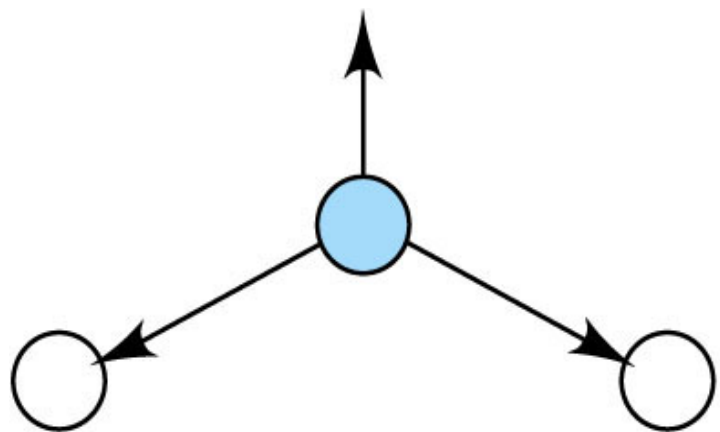


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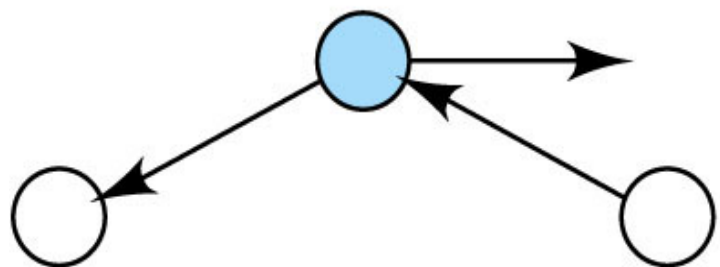
Bending

H₂O molecule

Triatomic molecule such as water, sulfur dioxide, or nitrogen dioxide have $3 \times 3 - 6 = 3$ vibrational modes. The central atom is not in line with the other two, a symmetric stretching vibration will produce a change in dipole and will thus be responsible for infrared absorption. Stretching peaks at **3650** and **3760** cm⁻¹ appear in the infrared spectrum for the symmetric and asymmetric vibrations of the water molecule. There is only one component to the scissoring vibration for this nonlinear molecule. For water, the bending vibration cause absorption at **1595** cm⁻¹.

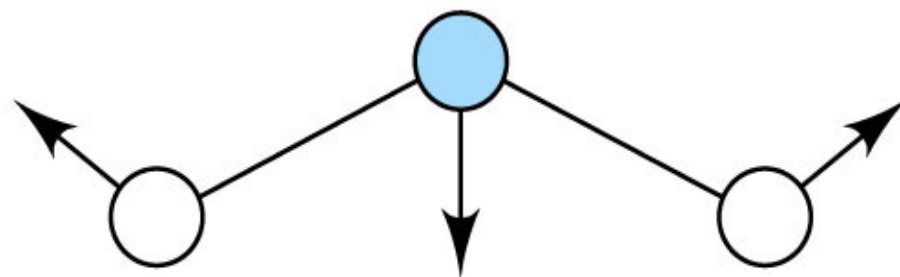


Symmetric stretching



Asymmetric stretching

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Scissoring

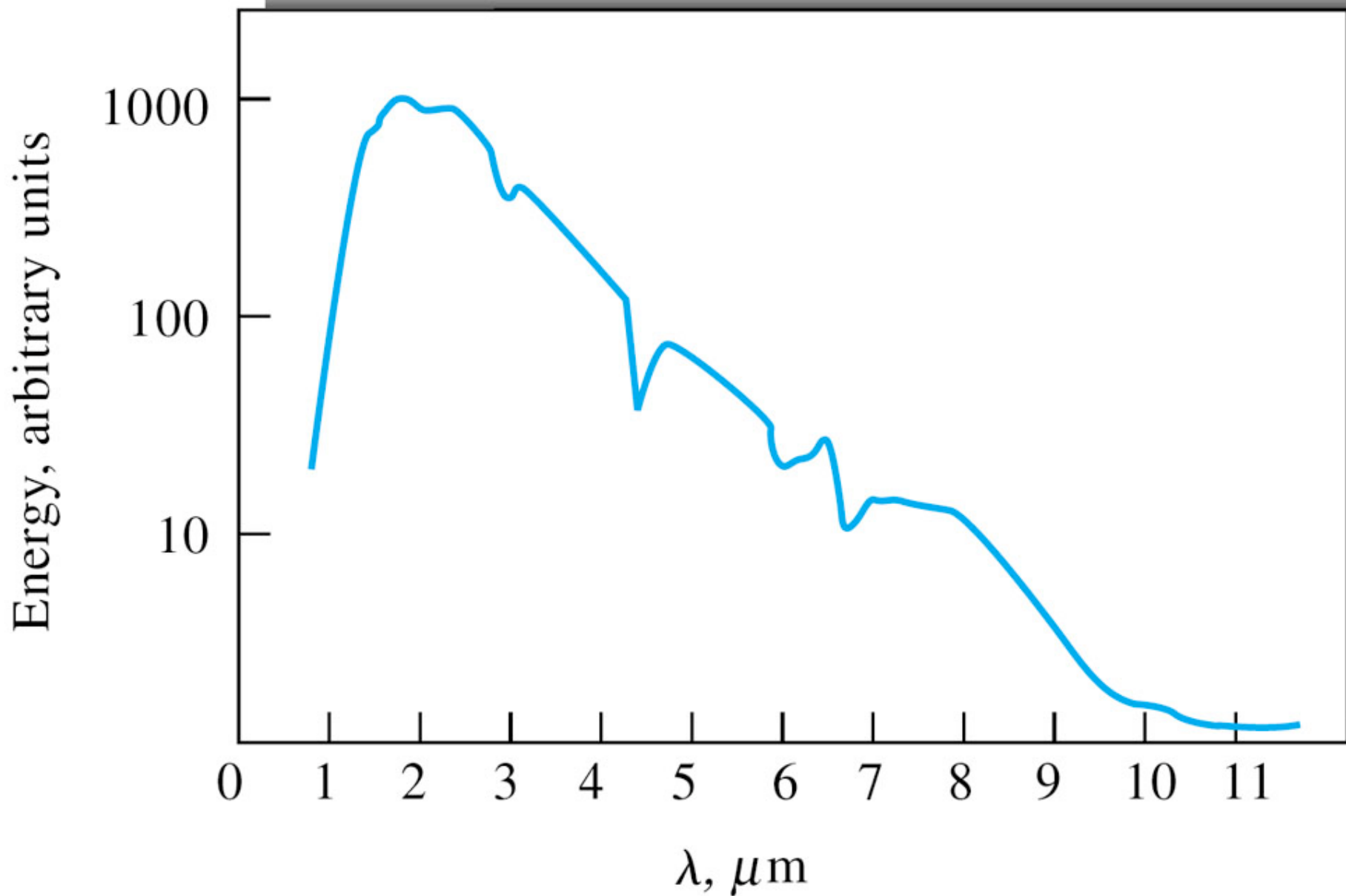
INFRARED SOURCES

- Sources معنا source مركبات اذا سخناها بعطوني طاقة ب IR range

Infrared sources consist of an **inert solid** that is heated electrically to a temperature between 1500 and 2200 K. Continuum radiation approximating that of a blackbody results. The maximum radiant intensity at these temperatures occurs between **5000 to 5900 cm⁻¹**.

- The Nernst Glower: The Nernst glower is composed of **rare earth oxides** formed into a cylinder having a diameter of 1 to 2 mm and a length of perhaps 20 mm. Platinum leads are sealed to the ends of the cylinder to permit electrical connection to what amounts to a resistive heating element. As current is passed through the device, temperature between 1200 K and 2200 K result.

سبيكة من الرصاص والبلاتينيوم اذا سخناها بتعطي طاقة ب range ال 1200k ال 2200



ال global source مركب بتكون من سيلاكون وكربون وشكلو
tetrahedral لما تتسخن بتعطي طاقة بحدود 1300-1500K ..ال
incandescent بتتكون من نيكل وكروم ال mercury عبارة عن غاز
مضغوط ... ال tungsten زي اللي بكون بالمصابيح بعطي اللون الاحمر
او الضوء اللي مائل على احمر لما أعطيه الكهرباء وال carbon
dioxide عبارة عن laser

- The Global Source: A Global is a silicon carbide rod, usually about 50 mm in length and 5 mm in diameter. It also is electrically heated (1300 to 1500 K). Spectral energies of the Global and the Nernst glower are comparable except in the region below 5 m, where the Global provides a significantly greater output.
- Incandescent Wire Source: A source of somewhat lower intensity but longer life than the Global or Nernst glower is a tightly wound spiral of nichrome wire heated to about 1100 K by an electrical current.

- The Mercury Arc: For the far-infrared region of the spectrum ($\lambda > 50 \text{ } \mu\text{m}$), none of the thermal sources just described provides sufficient radiant power for convenient detection. Here, a high-pressure mercury arc is used. This device consists of a quartz-jacketed tube containing mercury vapor at a pressure greater than one atmosphere. Passage of electricity through the vapor forms an internal plasma source that provides continuum radiation in the far-infrared region.

- The Tungsten Filament Lamp: An ordinary tungsten filament lamp is a convenient source for the near-infrared region of 4000 to 12,800 cm^{-1} .
- The Carbon Dioxide Laser Source: A tunable carbon dioxide laser is used as an infrared source for monitoring the concentrations of certain atmospheric pollutants and for determining absorbing species in aqueous solutions. A carbon dioxide laser produces a band of radiation in the 900 to 1100 cm^{-1} range.

Sources

Nernst Glower	heated rare earth oxide rod (~1500 K)	1-10 μm
Globar	heated SiC rod (~1500 K)	1-10 μm
W filament lamp	1100 K	0.78-2.5 μm
Hg arc lamp	plasma	>50 μm
CO ₂ laser	stimulated emission lines	9-11 μm

حان ليس

الافندسان

والا

منا مكي