Applications of Group Theory in Chemistry

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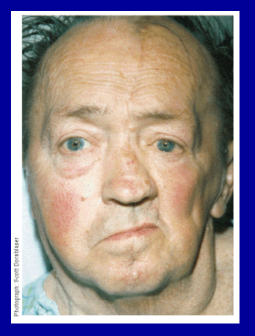
He hath made every thing beautiful in his time; also He hath set the world in their heart, so that no man can find out the worl that God maketh from the beginning to the end

Ecclesiastes 3:11

Symmetry and Periodicity – the two great powers of nature

Appreciating symmetry in a scientific way

Facial symmetry





Invariance to transformation as an indicator of facial symmetry





Mirror image

Symmetry - beautiful

Flower is beautiful

Leaves are beautiful

All beauty is due to symmetry

Even molecules are beautiful

Symmetry Elements

Symmetry elements are mirror planes, axis of rotation, centers of inversion, etc.

A molecule has a given symmetry element if the operation leaves the molecule <u>appearing</u> as if nothing has changed (even though atoms and bonds may have been moved.)

Symmetry Elements

Element

n-fold axis
Mirror plane
Center of inversion *n*-fold axis of
improper

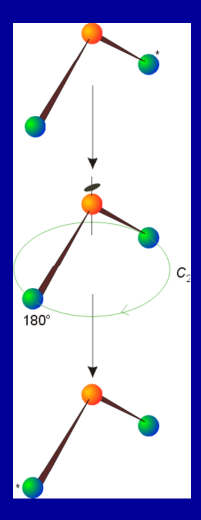
Symmetry OperationSymbolIdentityERotation by $2\pi/n$ C_n Reflection σ Inversioni

Rotation by $2\pi/n$ S_n rotation followed by reflectionperpendicular to the axis of rotation

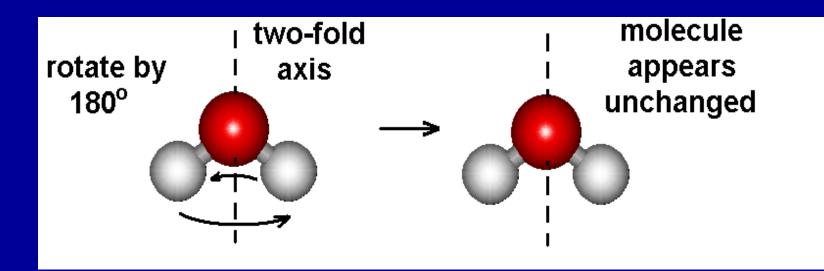
Identity, E

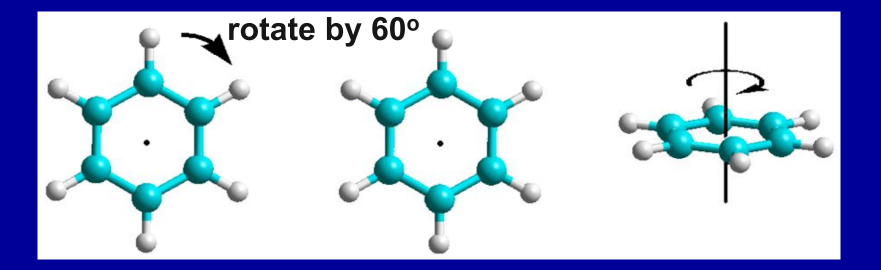
All molecules have Identity. This operation leaves the entire molecule unchanged. A highly asymmetric molecule such as a tetrahedral carbon with 4 different groups attached has only identity, and no other symmetry elements.

n-fold Rotation



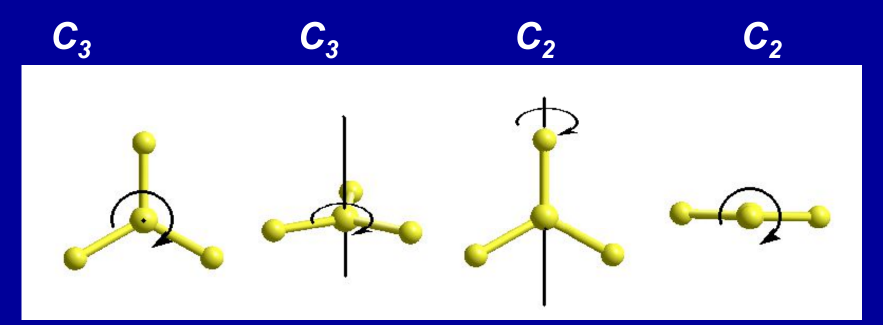
Water has a 2-fold axis of rotation. When rotated by 180°, the hydrogen atoms trade places, but the molecule will look exactly the same.





Rotational axes of BF₃

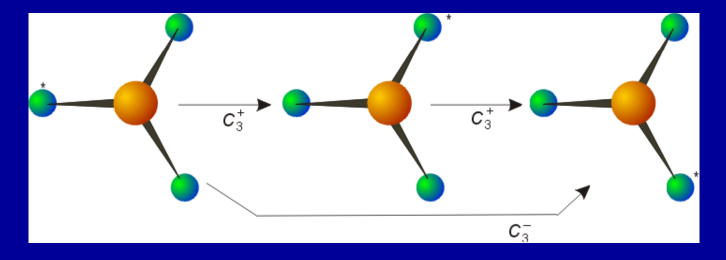
principal axis (highest value of *C_n*)



three-fold axis viewed from above

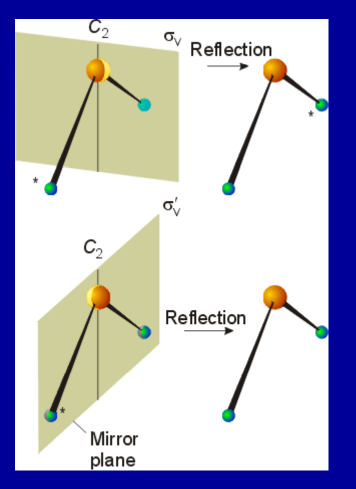
three-fold axis viewed from the side two-fold axis viewed from the side two-fold axis viewed from above

n-fold Axis of Rotation



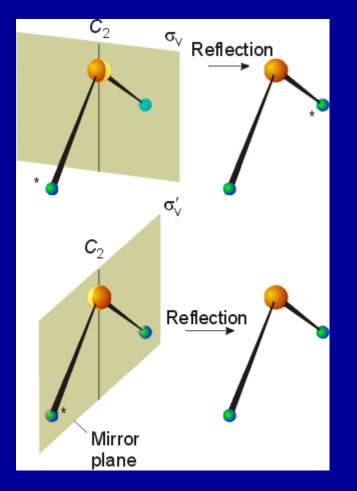
Ammonia has a C_3 axis. Note that there are <u>two</u> operations associated with the C_3 axis. Rotation by 120° in a clockwise or a counterclockwise direction provide two different orientations of the molecule.

Mirror Planes



The reflection of the water molecule in either of its two mirror planes results in a molecule that looks unchanged.

Mirror Planes

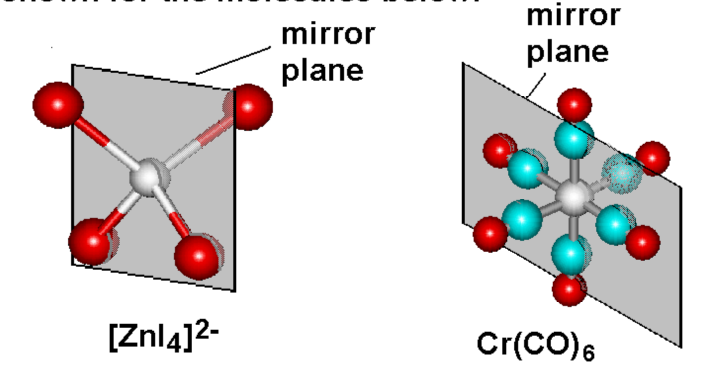


The subscript "v" in σ_v , indicates a vertical plane of symmetry. This indicates that the mirror plane includes the principal axis of rotation (C_2) .

1) Mirror Planes.

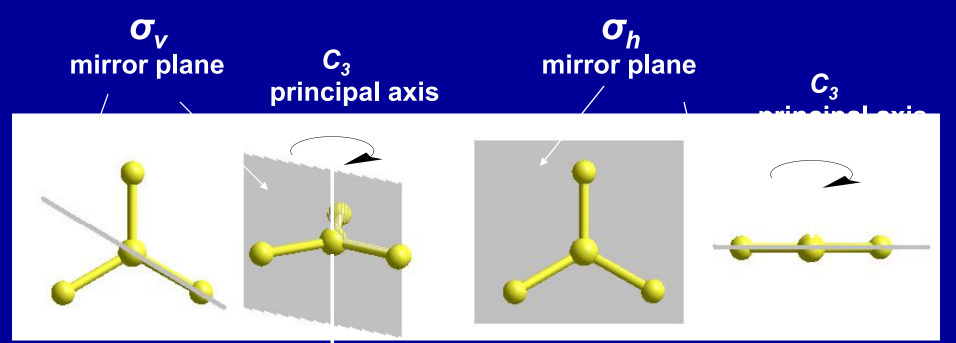


Mirror symmetry means invariance to reflection, as shown for the molecules below:



Mirror planes (σ) of BF₃:

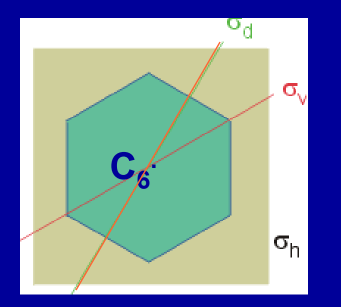
Mirror planes can contain the principal axis (σ_v) or be at right angles to it (σ_h) . BF₃ has one σ_h and three σ_v planes: (*v* = vertical, *h* = horizontal)



 σ_v mirror plane contains the C_3 axis

 σ_h mirror plane is at right angles to the C_3 axis

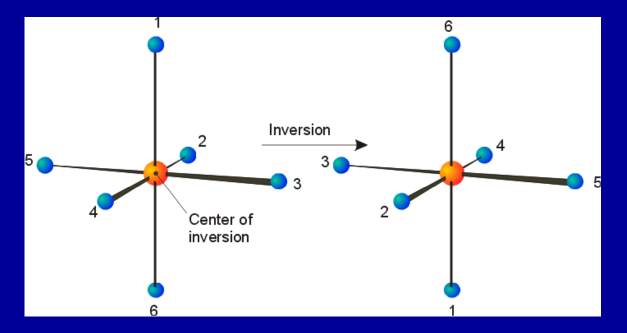
Mirror Planes



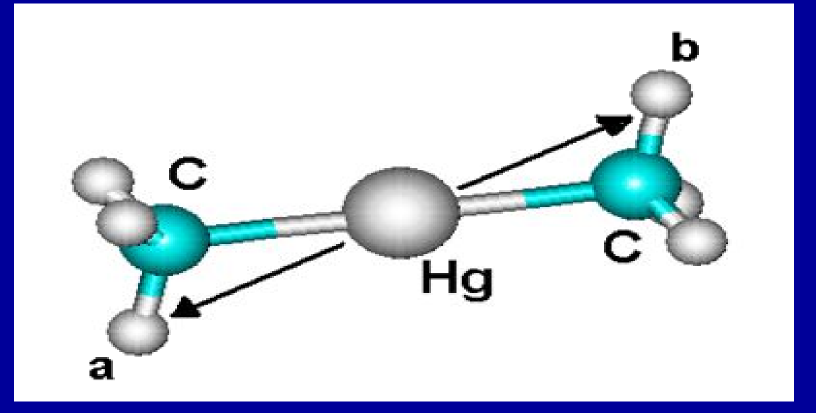
The vertical planes, σ_v , go through the carbon atoms, and include the C_6 axis. The planes that bisect the bonds are called *dihedral* planes, σ_{d}

Inversion

The inversion operation projects each atom through the center of inversion, and across to the other side of the molecule.

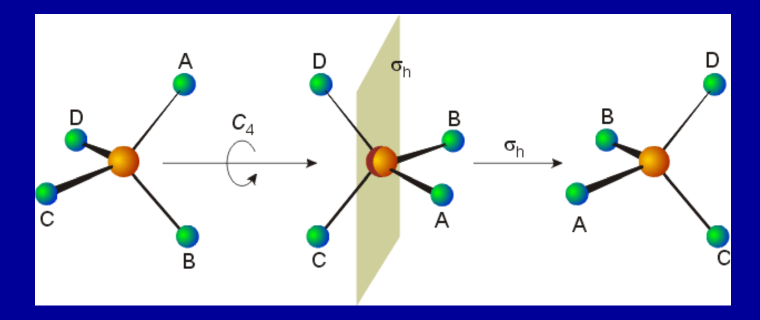


Inversion or center of symmetry



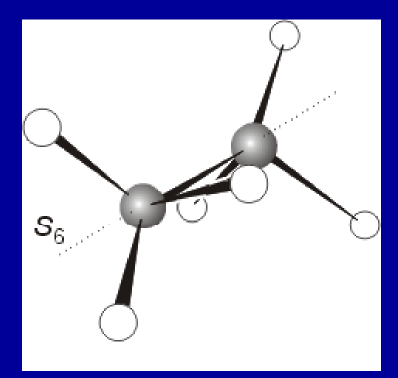
Improper Rotation

An improper rotation is rotation, followed by reflection in the plane perpendicular to the axis of rotation.



Improper Rotation

The staggered conformation of ethane has an S_6 axis that goes through both carbon atoms.



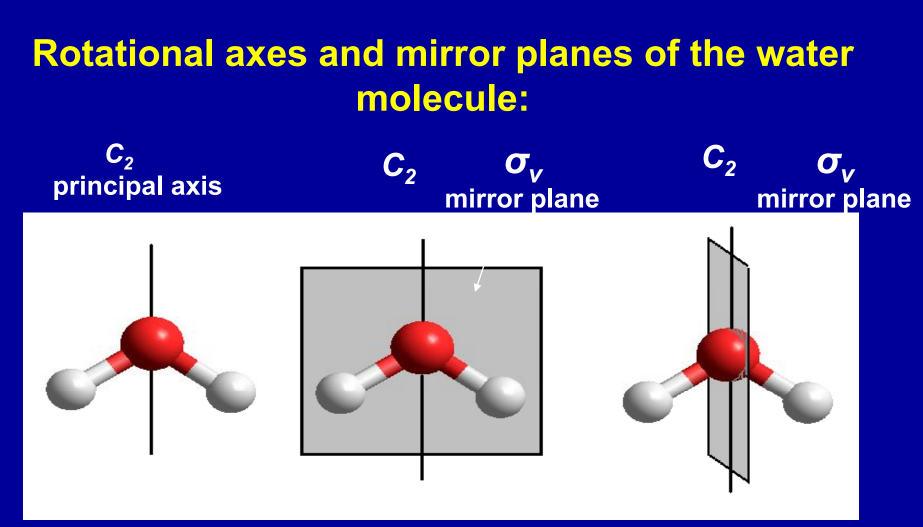
Symmetry Elements

Element

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Mirror plane
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Symmetry OperationSymbolIdentityERotation by $2\pi/n$ C_n Reflection σ Inversioni

Rotation by $2\pi/n$ S_n rotation followed by reflectionperpendicular to the axis of rotation



The water molecule has only one rotational axis, its C_2 axis which is also its principal axis. It has two mirror planes that contain the principal axis, which are therefore σ_v planes. It has no σ_h mirror plane, and no center of symmetry.

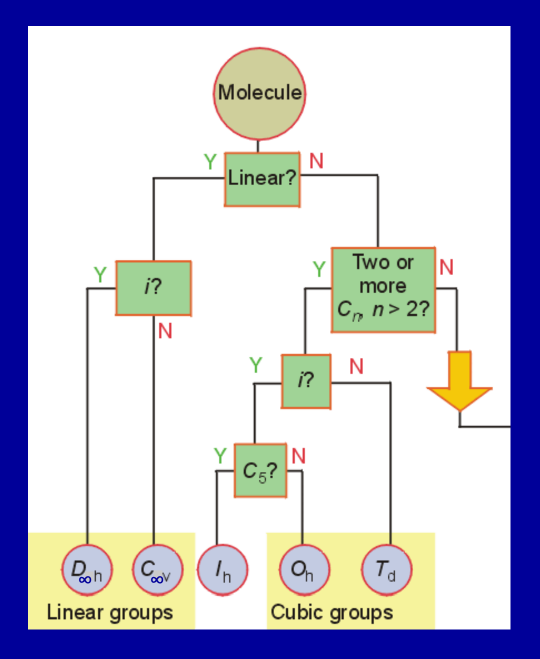
Group Theory

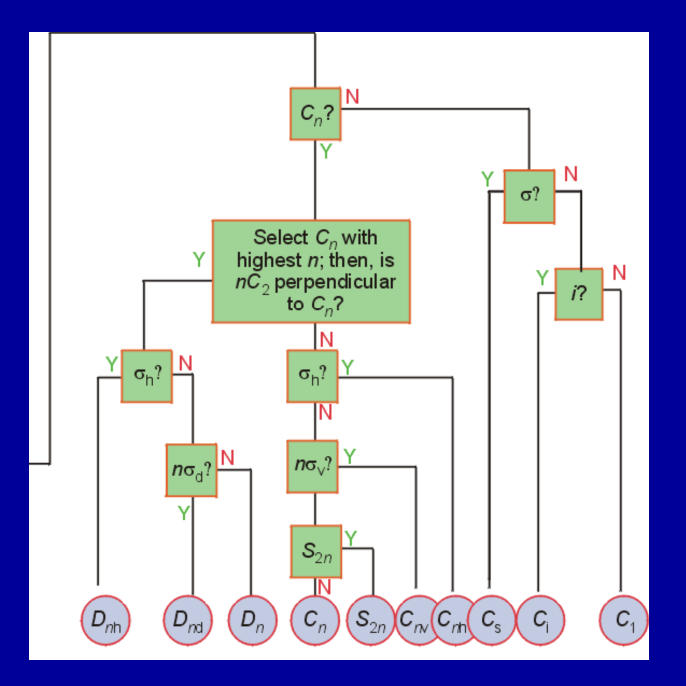
- A group is a mathematically defined collection of (symmetry) operations that have a specific set of mathematical properties.
 - Identity element
 - Inverse operation for each member of the group
 - Closure
 - Associative

To find out whether the symmetry elements form a group multiplication table is constructed.

Point Groups

Molecules with the same symmetry elements are placed into *point groups*.





Chemical Applications of Group Theory

Chirality

- Dipolemoment
- To predict hybridisation scheme
- vibrational spectroscopy
 - ir and Raman
 - # allowed stretching & bending frequencies

Chirality – optical activity

Allenes – disymmetry Lactic acid – Assymmetry

Chiral molecules lack an improper axis of rotation (S_n), a center of symmetry (*i*) or a mirror plane (σ).

Dipolement

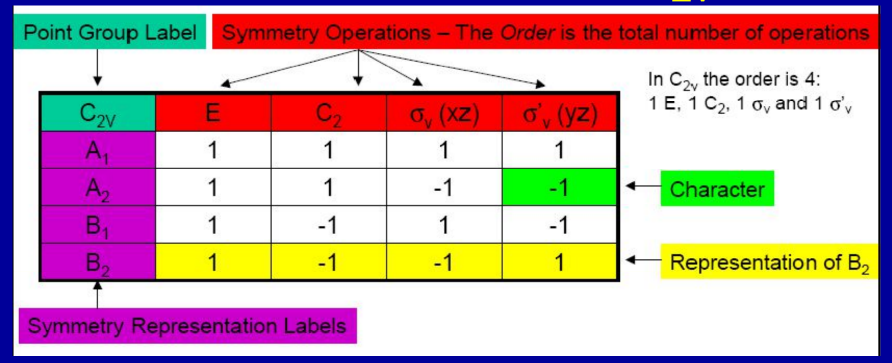
- Predicting polarity of molecules. A molecule cannot have a permanent dipole moment if it
- a) has a center of inversion
- b) belongs to any of the D point groups
- c) belongs to the cubic groups Tor O
 - Compounds having many C₂ axes
 - Compounds having inversion centre
 - Dipolemoment is along C_n axis and in the reflection plane.

Symmetry operations – matrices (Representation) – reducible and irreducible Matrices have character – character table Characted Table – **Hybridisation Scheme** Vibrational modes

Character Tables

The symmetry properties of each point group are summarized on a *character table*. The character table lists all of the symmetry elements of the group, along with a complete set of *irreducible representations*.

Character Table (C_{2v})



"A" means symmetric with regard to rotation about the principle axis. "B" means anti-symmetric with regard to rotation about the principle axis. Subscript numbers are used to differentiate symmetry labels, if necessary. "1" indicates that the operation leaves the function unchanged: it is called "symmetric". "-1" indicates that the operation reverses the function: it is called "anti-symmetric".

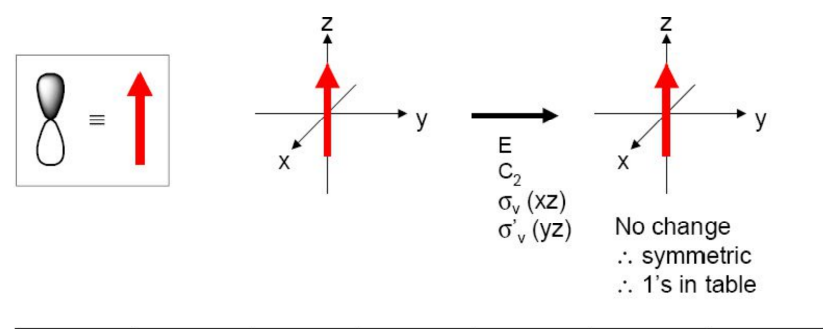
Character Table (C_{2v})

| | | | | | Symmetry of Functions | | |
|-----------------|---|----------------|------------------|----------------------|-----------------------|--|--|
| | | | | | | | |
| C _{2V} | E | C ₂ | $\sigma_{v}(xz)$ | σ' _v (yz) | | | |
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² | |
| A ₂ | 1 | 1 | -1 | -1 | R _z | ху | |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | XZ | |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz | |

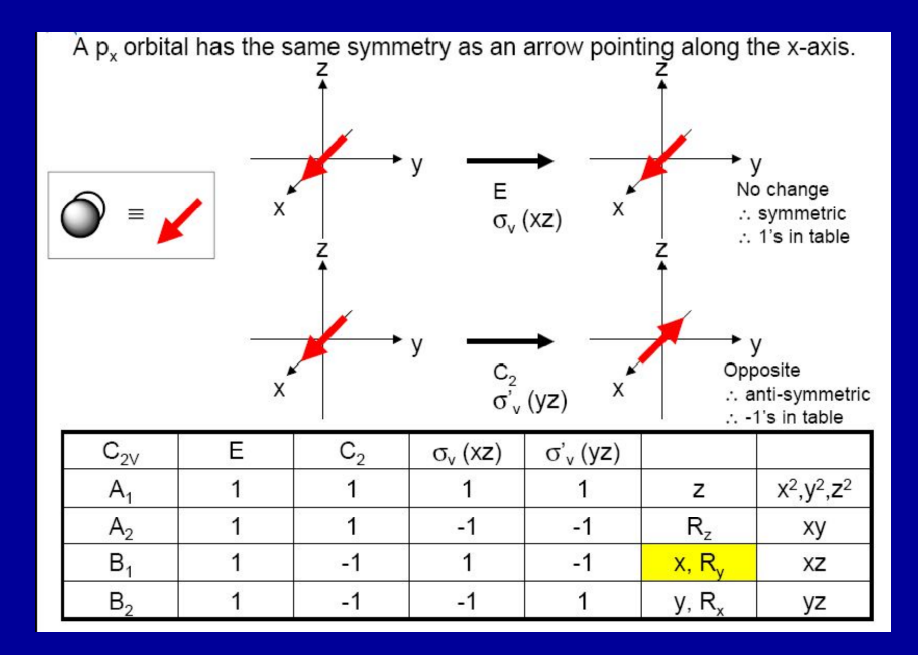
The functions to the right are called *basis functions*. They represent mathematical functions such as orbitals, rotations, etc.

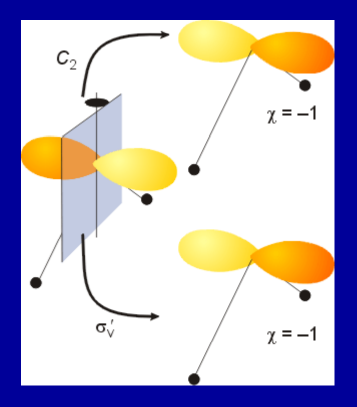


A p_z orbital has the same symmetry as an arrow pointing along the z-axis.



| C _{2V} | E | C ₂ | σ _v (xz) | σ' _v (yz) | | |
|-----------------|---|----------------|---------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

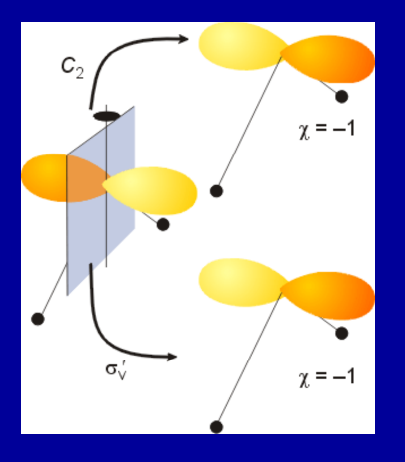




If a p_x orbital on the central atom of a molecule with C_{2v} symmetry is rotated about the C_2 axis, the orbital is reversed, so the character will be -1.

If a p_x orbital on the central atom of a molecule with C_{2v} symmetry is rotated about the C_2 axis, the orbital is reversed, so the character will be -1.

| C _{2V} | E | C ₂ | $\sigma_{v}(XZ)$ | σ' _v (yz) | | |
|-----------------|---|----------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² |
| A ₂ | 1 | 1 | -1 | -1 | R _z | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | XZ |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | уz |



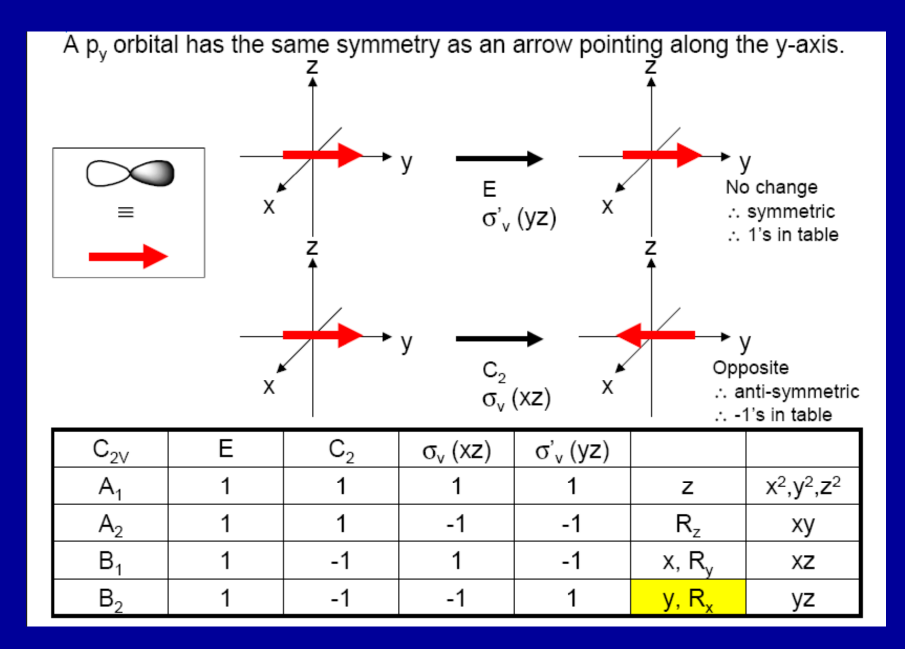
If a p_x orbital on the central atom of a molecule with C_{2v} symmetry is reflected in the yz plane, the orbital is also reversed, and the character will be -1.

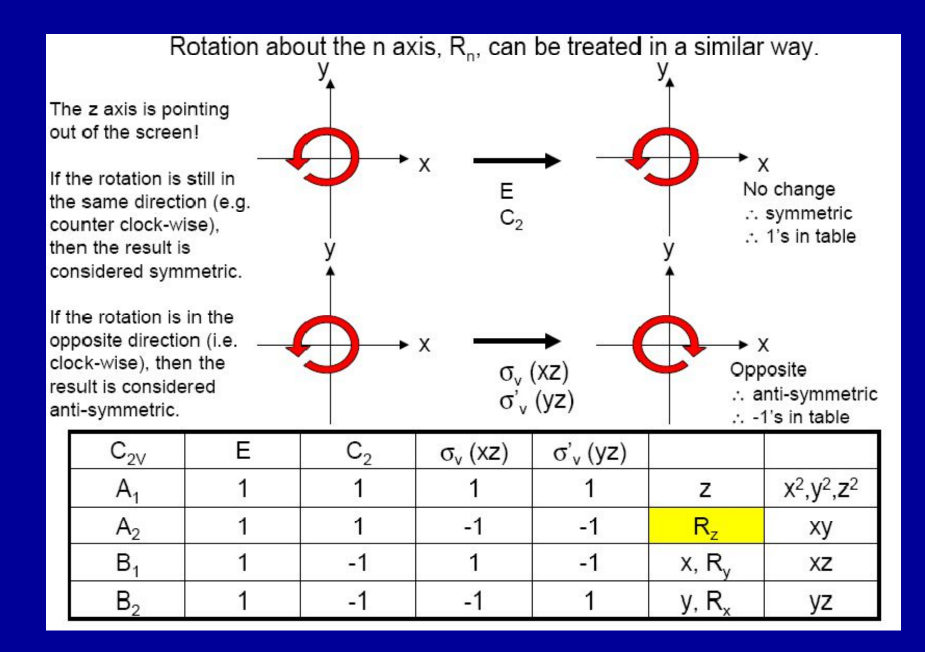
If a p_x orbital on the central atom of a molecule with C_{2v} symmetry is reflected in the yz plane, the orbital is also reversed, and the character will be -1.

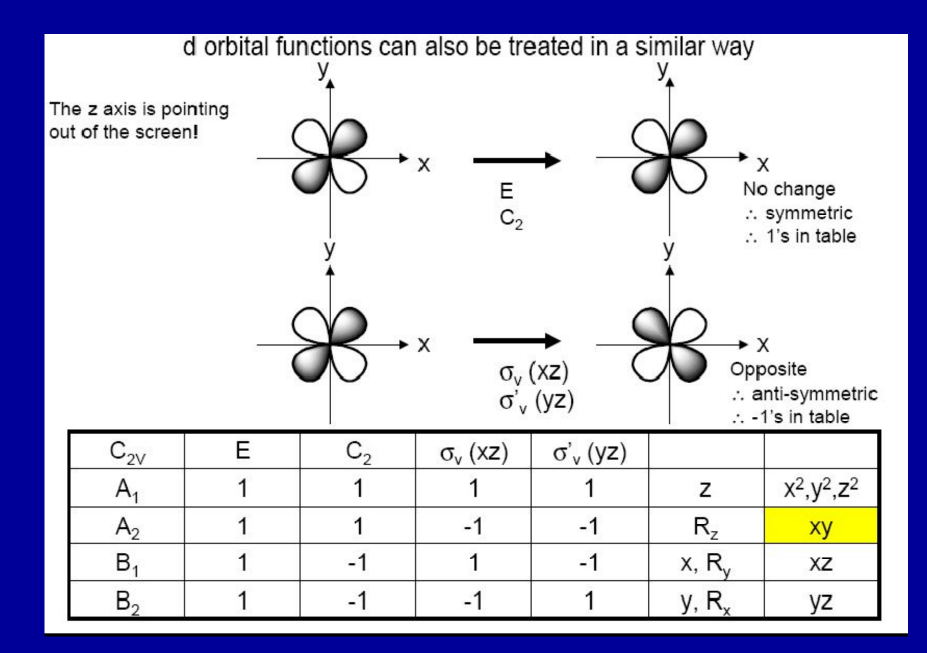
| C _{2V} | E | C ₂ | $\sigma_{v}(XZ)$ | σ' _v (yz) | | |
|-----------------|---|----------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² |
| A ₂ | 1 | 1 | -1 | -1 | R _z | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | ХZ |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | уz |

If a p_x orbital on the central atom of a molecule with C_{2v} symmetry is reflected in the xz plane, the orbital is unchanged, so the character is +1.

| C _{2V} | E | C ₂ | $\sigma_{v}(XZ)$ | σ' _v (yz) | | |
|-----------------|---|----------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² |
| A ₂ | 1 | 1 | -1 | -1 | R _z | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | ХZ |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | уz |







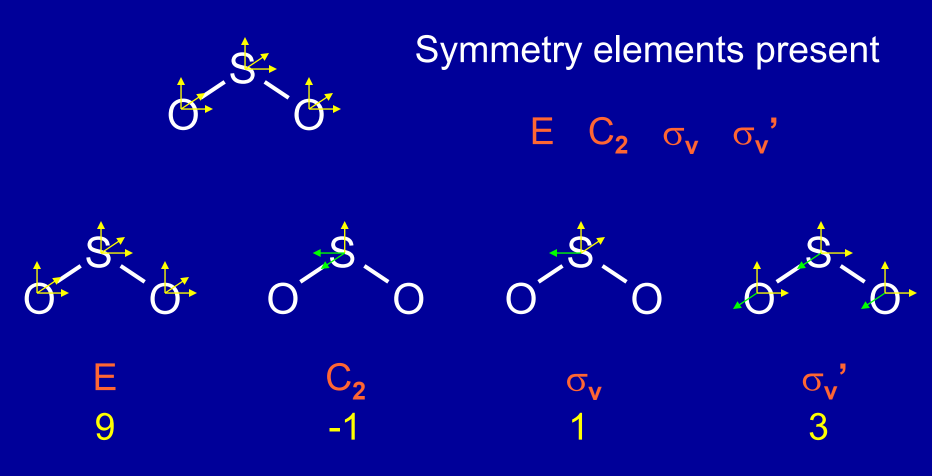
Character Table Representations

- 1. Characters of +1 indicate that the basis function is unchanged by the symmetry operation.
- 2. Characters of -1 indicate that the basis function is reversed by the symmetry operation.
- 3. Characters of 0 indicate that the basis function undergoes a more complicated change.

Character Table Representations

- 1. An *A* representation indicates that the functions are symmetric with respect to rotation about the principal axis of rotation.
- 2. *B* representations are asymmetric with respect to rotation about the principal axis.
- 3. *E* representations are doubly degenerate.
- 4. T representations are triply degenerate.
- 5. Subscrips *u* and *g* indicate asymmetric (*ungerade*) or symmetric (*gerade*) with respect to a center of inversion.

Determination of the Reducible Representation of a Molecule



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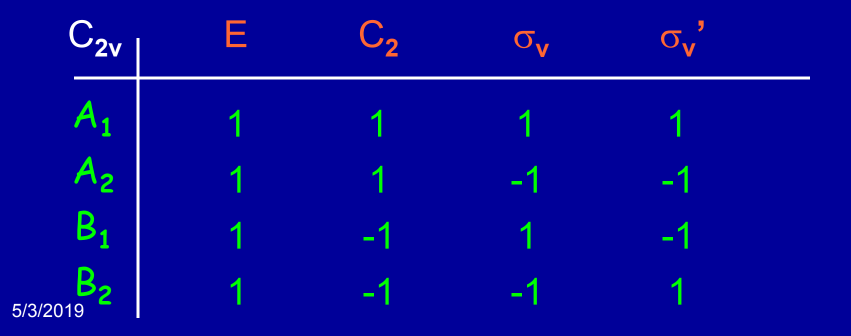
Reducible Representation

Symmetry operations that are applied to the each atom of the molecule that does not move produce the reducible representation



Character Table

A reducible representation can be decomposed into a set of irreducible representations using a character table



Decomposing a Reducible Representation to Irreducible Representations

 $a_i = (1/h) \sum (X^R \cdot X_i^R \cdot C^R)$

 $aA_1 = (1/4) [(9 \cdot 1 \cdot 1) + (-1 \cdot 1 \cdot 1) + (1 \cdot 1 \cdot 1) + (3 \cdot 1 \cdot 1)]$ = 3

 $aA_{2} = (1/4) [(9 \cdot 1 \cdot 1) + (-1 \cdot 1 \cdot 1) + (1 \cdot -1 \cdot 1) + (3 \cdot -1 \cdot 1)]$

 $aB_1 = (1/4) [(9 \cdot 1 \cdot 1) + (-1 \cdot -1 \cdot 1) + (1 \cdot 1 \cdot 1) + (3 \cdot -1 \cdot 1)]$

 $aB_2 = (1/4) [(9 \cdot 1 \cdot 1) + (-1 \cdot -1 \cdot 1) + (1 \cdot -1 \cdot 1) + (3 \cdot 1 \cdot 1)]$

Decomposing a Reducible Representation to Irreducible Representations

 $a_i = (1/h) \sum (X^R \cdot X_i^R \cdot C^R)$

 $aA_1 = (1/4) [(9 \cdot 1 \cdot 1) + (-1 \cdot 1 \cdot 1) + (1 \cdot 1 \cdot 1) + (3 \cdot 1 \cdot 1)]$ = 3

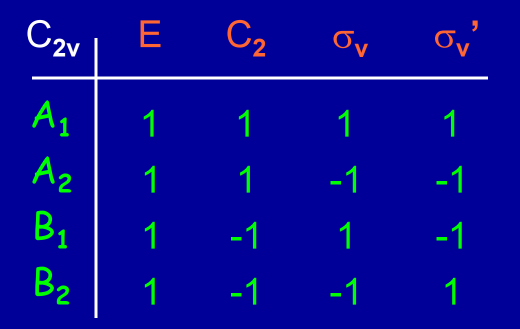
 $aA_{2} = (1/4) [(9 \cdot 1 \cdot 1) + (-1 \cdot 1 \cdot 1) + (1 \cdot -1 \cdot 1) + (3 \cdot -1 \cdot 1)]$

 $aB_1 = (1/4) [(9 \cdot 1 \cdot 1) + (-1 \cdot -1 \cdot 1) + (1 \cdot 1 \cdot 1) + (3 \cdot -1 \cdot 1)]$

 $aB_2 = (1/4) [(9 \cdot 1 \cdot 1) + (-1 \cdot -1 \cdot 1) + (1 \cdot -1 \cdot 1) + (3 \cdot 1 \cdot 1)]$

$$\begin{array}{c|c} C_{2v} & E & C_2 & \sigma_v & \sigma_v' \\ \hline \Gamma_{red} & 9 & -1 & 1 & 3 \end{array}$$

 $\Gamma_{\text{irred}} = 3 A_1 + A_2 + 2 B_1 + 3 B_2$



Degrees of Freedom

Every molecule has 3n energy modes

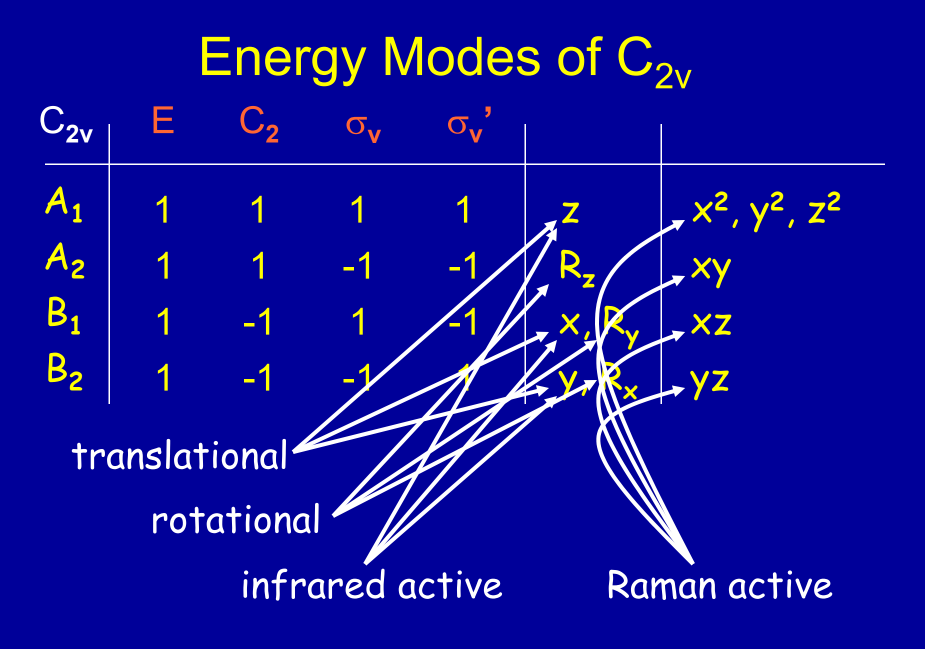
| atoms | degrees of freedom | translation | rotation | vibration |
|--------------|-----------------------|-------------|----------|-----------|
| 1 | 3 | 3 | 0 | 0 |
| 2 | 6 | 3 | 2 | 1 |
| 3 (linear) | 9 | 3 | 2 | 4 |
| 3 (nonlinear |) 9 | 3 | 3 | 3 |

Irreducible Representations $3 A_1 + A_2 + 2 B_1 + 3 B_2$

Represents the energy modes of the molecule



How many vibrational modes does SO_2 have?

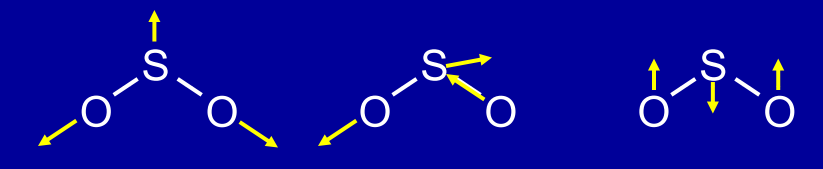


Energy Modes of SO₂

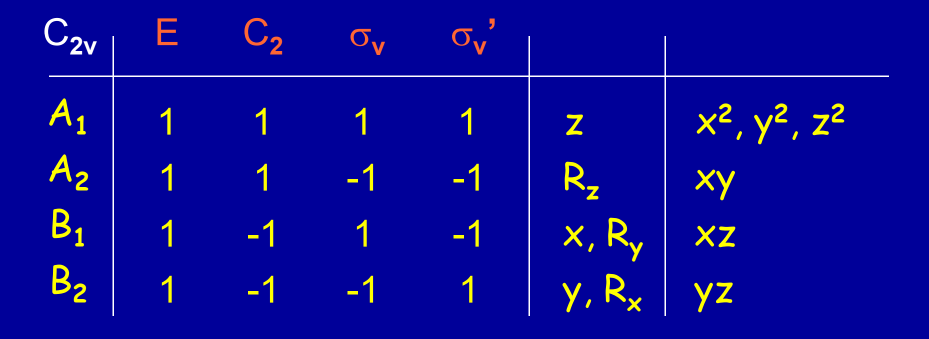
 $\Gamma_{\rm irred} = 3 A_1 + A_2 + 2 B_1 + 3 B_2$

Assign the 9 energy modes of SO₂ Are all the vibrational modes infrared active? Are all the vibrational modes Raman active?

Vibrational Modes of SO₂



symmetric stretch A₁ asymmetric stretch B₂ symmetric bend A₁



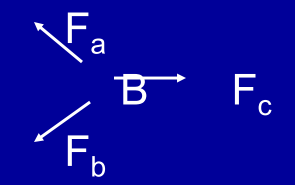
Determining Hybridization

- 1. Determine the point group of the molecule.
- 2. Consider the σ bonds as vectors, and determine how they are transformed by the symmetry operations of the group.
- Obtain the characters for the bonds. For each symmetry operation, a bond which remains in place contributes a value of +1. If the bond is moved to another position, it contributes a value of 0.
- 4. *Reduce* the set of characters to a linear combination of the character sets of the point group.

Hybridization

 Determine the hybridization of boron in BF₃. The molecule is trigonal planar, and belongs to point group D_{3h}.

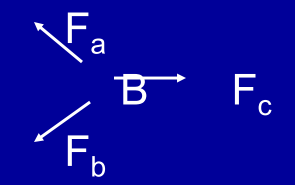
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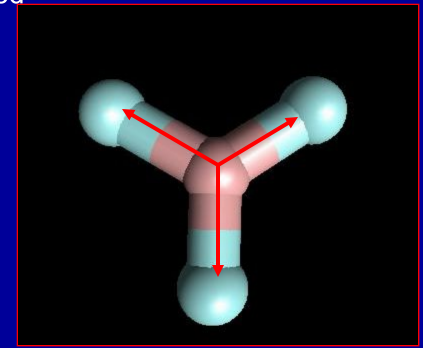
Hybridization

Determine how each vector (σ bond) is transformed by the symmetry operations of the aroup.

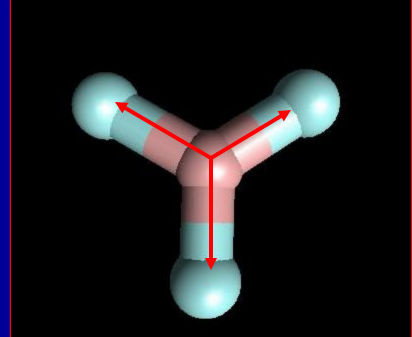
| D_{3h} | E | 2C3 | 3C2 | $\sigma_{\rm h}$ | 2S3 | $3\sigma_v$ | | |
|------------------|---|-----|-----|------------------|-----|-------------|-------------|---|
| A _l ' | 1 | 1 | 1 | 1 | 1 | 1 | | x ² +y ² , z ² |
| A2' | 1 | 1 | -1 | 1 | 1 | -1 | R., | |
| E' | 2 | -1 | 0 | 2 | -1 | 0 | (x,y) | (x^2-y^2, xy) |
| A1" | 1 | 1 | 1 | -1 | -1 | -1 | | |
| A1" A2" | 1 | 1 | -1 | -1 | -1 | 1 | z | |
| E'' | 2 | -1 | 0 | -2 | 1 | 0 | (R_v,R_v) | (xz,yz) |

Determining Hybridization $2C_3$ $3C_2$ σ_h $2S_3$ $3\sigma_v$ E **Fred**

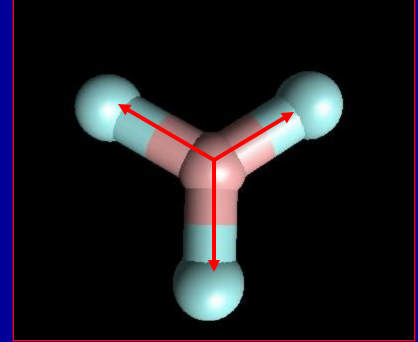
Determining Hybridization $E \quad 2C_3 \quad 3C_2 \quad \sigma_h \quad 2S_3 \quad 3\sigma_v$ $\Gamma_{red} \quad 3$



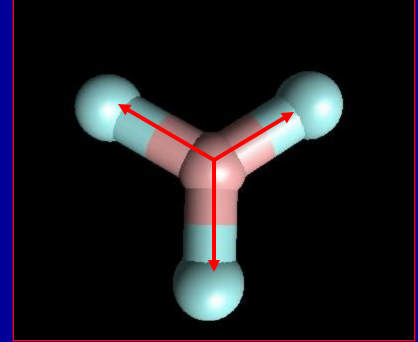
Determining Hybridization $E \quad 2C_3 \quad 3C_2 \quad \sigma_h \quad 2S_3 \quad 3\sigma_v$ $\Gamma_{red} \quad 3 \quad 0$



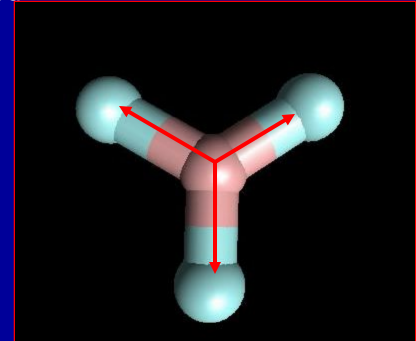
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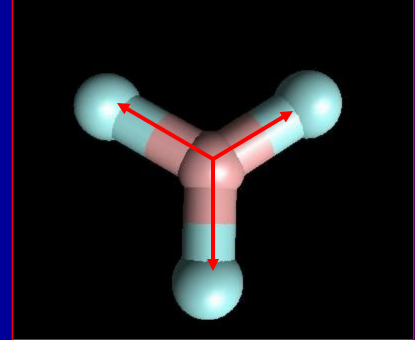
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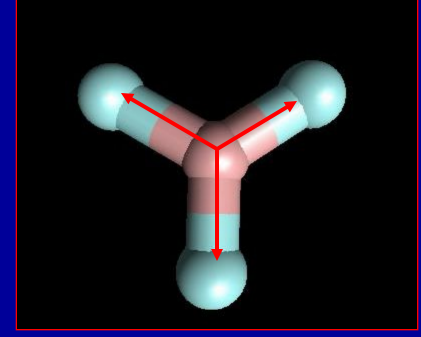


Determining Hybridization $E \quad 2C_3 \quad 3C_2 \quad \sigma_h \quad 2S_3 \quad 3\sigma_v$ $\Gamma_{red} \quad 3 \quad 0 \quad 1 \quad 3$



Determining Hybridization $E \quad 2C_3 \quad 3C_2 \quad \sigma_h \quad 2S_3 \quad 3\sigma_v$ $\Gamma_{red} \quad 3 \quad 0 \quad 1 \quad 3 \quad 0$





Reducing a Representation

$$n_i = \frac{1}{h} \sum_c g_c \chi_i \chi_r$$

n_i = the number of times an irreducible representation *i* occurs in the reducible representation

- h = the order of the group (the total number of operations in the point group)
- c = the class (type) of operation
- g_c = the number of operations in the class
- χ_i = the character of the irreducible representation

 χ_r = the character of the reducible representation

Reducing a Representation

$$n_i = \frac{1}{h} \sum_{c} g_{c} \chi_i \chi_r$$

| D _{3h} | E | 2C3 | 3C2 | $\sigma_{\rm h}$ | 2S3 | $3\sigma_{\rm V}$ | | |
|-----------------|---|-----|-----|------------------|-----|-------------------|--------------|---|
| A1' | 1 | 1 | 1 | 1 | 1 | 1 | | x ² +y ² , z ² |
| A2' | 1 | 1 | -1 | 1 | 1 | -1 | R, | |
| E' | 2 | -1 | 0 | 2 | -1 | 0 | (x,y) | (x ² -y ² , xy) |
| A1" | 1 | 1 | 1 | -1 | -1 | -1 | | |
| A1" A2" | 1 | 1 | -1 | -1 | -1 | 1 | z | |
| E'' | 2 | -1 | 0 | -2 | 1 | 0 | (R_v, R_v) | (xz,yz) |

The order of the group, h, is the total number of operations.

h = 1+2+3+1+2+3=12

Hybridization of BF₃

 Γ_{red} reduces to $A_1' + E'$. The orbitals used in hybridization must have this

| D_{3h} | Е | 2C3 | 3C2 | $\sigma_{\rm h}$ | 2S3 | $3\sigma_v$ | | |
|------------------|---|-----|-----|------------------|-----|-------------|--------------|---|
| A ₁ ' | 1 | 1 | 1 | 1 | 1 | 1 | | x ² +y ² , z ² |
| A2' | 1 | 1 | -1 | 1 | 1 | -1 | R., | |
| E' | 2 | -1 | 0 | 2 | -1 | 0 | (x,y) | (x ² -y ² , xy) |
| A ₁ " | 1 | 1 | 1 | -1 | -1 | -1 | | |
| A1" A2" | 1 | 1 | -1 | -1 | -1 | 1 | z | |
| E'' | 2 | -1 | 0 | -2 | 1 | 0 | (R_v, R_v) | (xz,yz) |

Hybridization of BF_3 Γ_{red} reduces to $A_1' + E'$. The orbitals used in hybridization must have this symmetry.

| D _{3h} | E | 2C3 | 3C2 | $\sigma_{\rm h}$ | 2S3 | $3\sigma_v$ | | |
|-----------------|---|-----|-----|------------------|-----|-------------|----------------|---|
| Al, | 1 | 1 | 1 | 1 | 1 | 1 | | x ² +y ² , z ² |
| A2' | 1 | 1 | -1 | 1 | 1 | -1 | R ₇ | |
| E' | 2 | -1 | 0 | 2 | -1 | 0 | (x,y) | (x ² -y ² , xy) |
| A1" | 1 | 1 | 1 | -1 | -1 | -1 | | |
| A2" | 1 | 1 | -1 | -1 | -1 | 1 | z | |
| E" | 2 | -1 | 0 | -2 | 1 | 0 | (R_v, R_v) | (xz,yz) |

The *s* orbital and the d_{z^2} orbitals on boron have A_1 ' symmetry. The 3 d_z^2 orbital is too high in energy to hybridize.

Hybridization of BF_3 Γ_{red} reduces to $A_1' + E'$. The orbitals used in hybridization must have this symmetry.

| D _{3h} | E | 2C3 | 3C2 | $\sigma_{\rm h}$ | 2S3 | $3\sigma_v$ | | |
|-----------------|---|-----|-----|------------------|-----|-------------|----------------|---|
| Al, | 1 | 1 | 1 | 1 | 1 | 1 | | x ² +y ² , z ² |
| A2' | 1 | 1 | -1 | 1 | 1 | -1 | R ₇ | |
| E' | 2 | -1 | 0 | 2 | -1 | 0 | (x,y) | (x ² -y ² , xy) |
| A1" | 1 | 1 | 1 | -1 | -1 | -1 | | |
| A2" | 1 | 1 | -1 | -1 | -1 | 1 | z | |
| E" | 2 | -1 | 0 | -2 | 1 | 0 | (R_v, R_v) | (xz,yz) |

The p_x and p_y orbitals and the $d_x^2_{-y}^2$ and d_{xy} orbitals have E ' symmetry. Since the *d* orbitals on boron are too high in energy, they will not be used.

Hybridization of BF_3 Γ_{red} reduces to $A_1' + E'$. The orbitals used in hybridization must have this symmetry.

| D_{3h} | Е | 2C3 | 3C2 | $\sigma_{\rm h}$ | 2S3 | $3\sigma_v$ | | |
|------------|---|-----|-----|------------------|-----|-------------|----------------|---|
| Al' | 1 | 1 | 1 | 1 | 1 | 1 | | x ² +y ² , z ² |
| A2' | 1 | 1 | -1 | 1 | 1 | -1 | R ₇ | |
| E' | 2 | -1 | 0 | 2 | -1 | 0 | (x,y) | (x ² -y ² , xy) |
| A1" | 1 | 1 | 1 | -1 | -1 | -1 | | |
| A1" A2" | 1 | 1 | -1 | -1 | -1 | 1 | z | |
| E" | 2 | -1 | 0 | -2 | 1 | 0 | (R_v, R_v) | (xz,yz) |

The hybridization of boron will sp^2 or, more specifically, sp_xp_{y} .

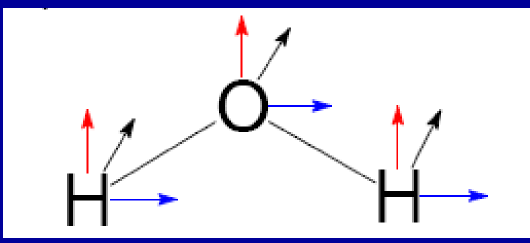
Molecular motion includes translations, rotations and vibrations. The total number of degrees of freedom (types of molecular motion) is equal to 3N, where N is the number of atoms in the molecule.

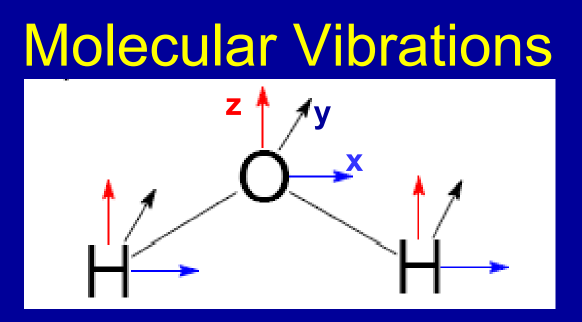
Of the 3N types of motion, three represent molecular translations in the x, y or z directions. Linear molecules have two rotational degrees of freedom, and non-linear molecules have three rotational degrees of freedom.

For linear molecules, the number of molecular vibrations = 3N-3-2 = 3N-5.

For non-linear molecules, the number of molecular vibrations = 3N-3-3=3N-6.

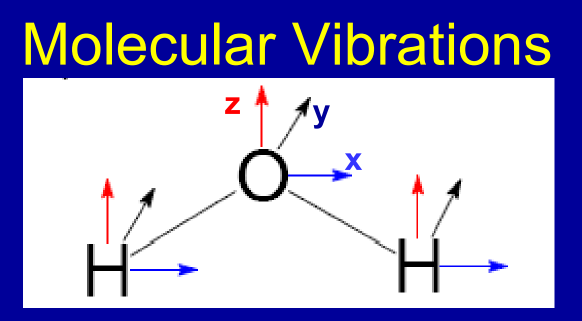
To obtain Γ_{red} for <u>all</u> molecular motion, we must consider the symmetry properties of the three cartesian coordinates on <u>all</u> atoms of the molecule.



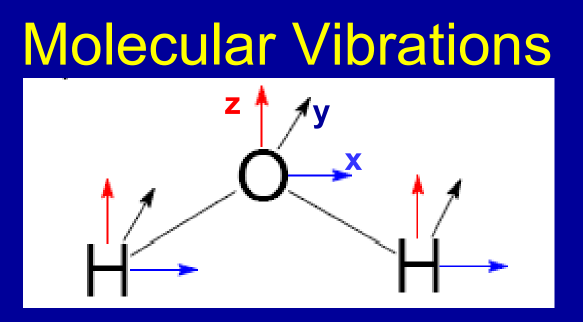


The molecule lies in the xz plane. The x axis is drawn in blue, and the y axis is drawn in black. The red arrows indicate the z axis.

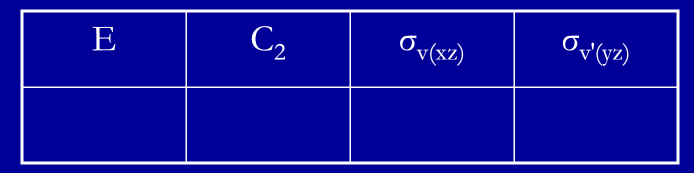
The molecule lies in the xz plane. The x axis is drawn in blue, and the y axis is drawn in black. The red arrows indicate the z axis.

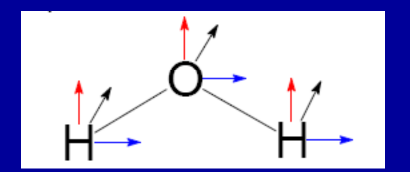


If a symmetry operation changes the position of an atom, all three cartesian coordinates contribute a value of 0.

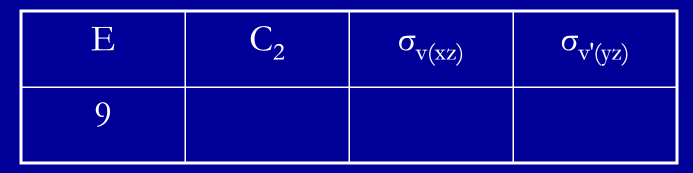


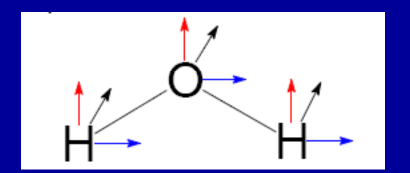
For operations that leave an atom in place, the character is +1 for an axis that remains in position, -1 for an axis that is reversed, and 0 for an axis that has been moved.



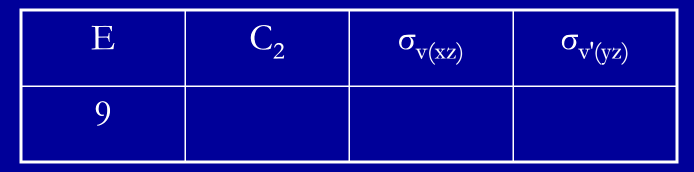


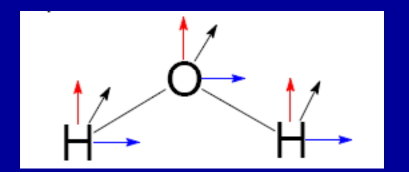
Identity leaves all 3 atoms in position, so the character will be 9.



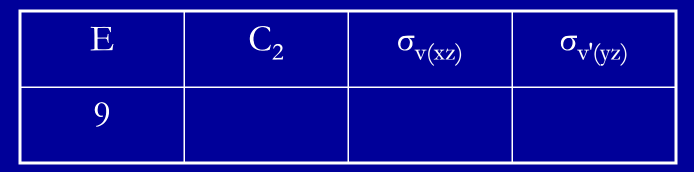


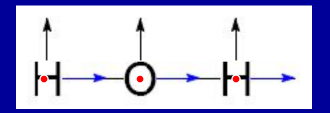
Identity leaves all 3 atoms in position, so the character will be 9.



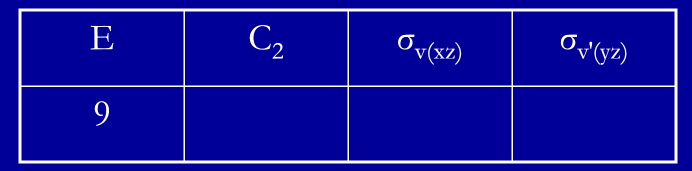


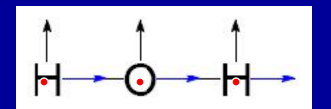
The C_2 axis goes through the oxygen atom, and exchanges the hydrogen atoms.

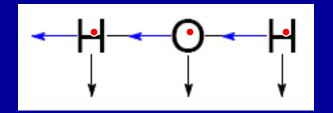




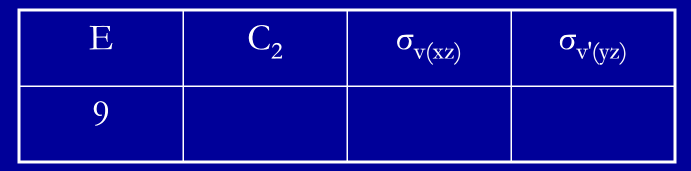
The z axis (red) on oxygen stays in position. This axis contributes +1 towards the character for C_2 .

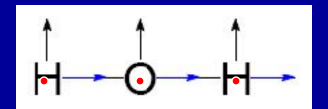


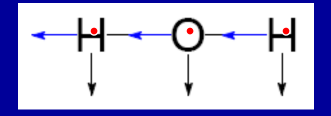




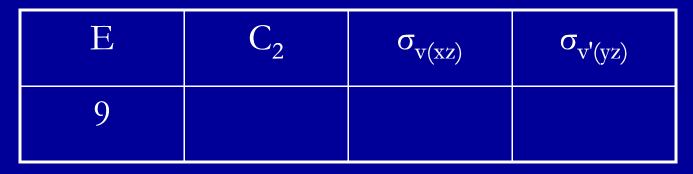
The y axis (black) on oxygen is rotated by 180°. This reverses the axis, and contributes -1 to the character for C_2 .



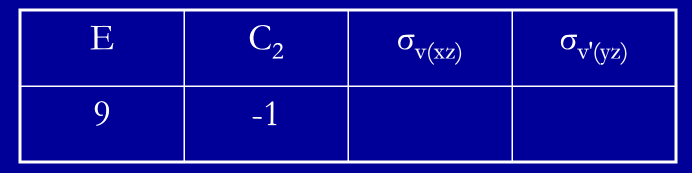




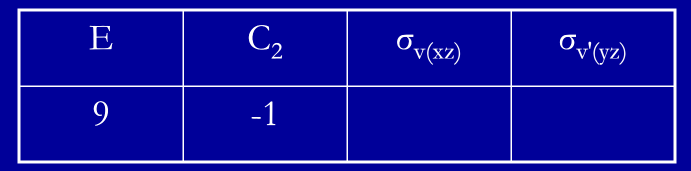
The x axis (blue) on oxygen is also rotated by 180° . This reverses the axis, and contributes -1 to the character for C₂.

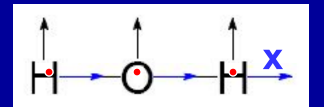


The character for the C_2 operation will be +1 (z axis on oxygen) -1 (y axis on oxygen) -1 (x axis on oxygen) = -1

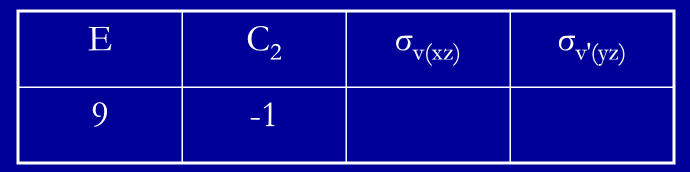


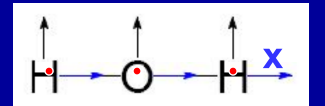
The character for the C_2 operation will be +1 (z axis on oxygen) -1 (y axis on oxygen) -1 (x axis on oxygen) = -1



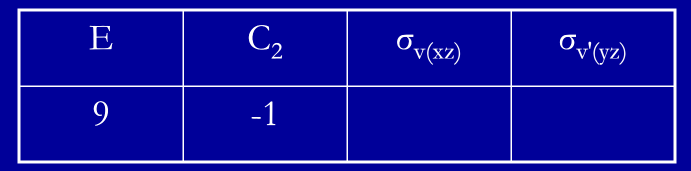


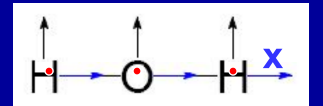
The xz mirror plane is the molecular plane, and all three atoms remain in position.



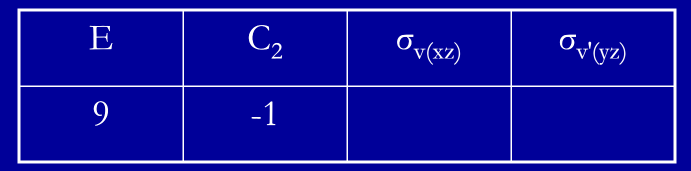


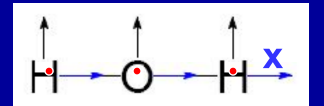
The z axis and the x axis both lie within the xz plane, and remain unchanged.



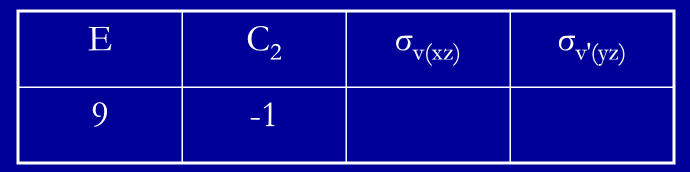


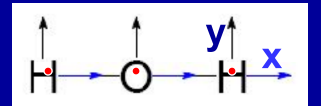
Each unchanged axis contributes +1 to the character for the symmetry operation.





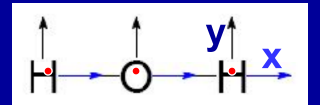
For 3 atoms, the contribution to the character will be: 3(1+1) = 6





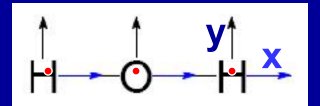
The y axis will be reversed by the mirror plane, contributing a value of -1 for each of the three atoms on the plane.

| E | C ₂ | $\sigma_{ m v(xz)}$ | $\sigma_{ m v'(yz)}$ |
|---|----------------|---------------------|----------------------|
| 9 | -1 | 3 | |



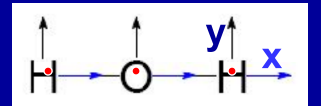
The character for the xz mirror plane will be: 6-3 = 3

| E | C ₂ | $\sigma_{ m v(xz)}$ | $\sigma_{ m v'(yz)}$ |
|---|----------------|---------------------|----------------------|
| 9 | -1 | 3 | |



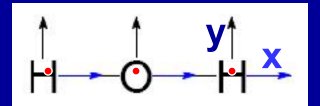
The yz mirror plane bisects the molecule. Only the oxygen atom lies in the plane.

| E | C ₂ | $\sigma_{ m v(xz)}$ | $\sigma_{ m v'(yz)}$ |
|---|----------------|---------------------|----------------------|
| 9 | -1 | 3 | |



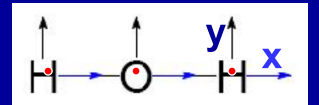
The y and z axis lie within the yz plane, and each contributes +1 to the character.

| E | C ₂ | $\sigma_{ m v(xz)}$ | $\sigma_{ m v'(yz)}$ |
|---|----------------|---------------------|----------------------|
| 9 | -1 | 3 | |



The x axis on oxygen is reversed by the reflection, and contributes a -1 towards the character.

| Е | C ₂ | $\sigma_{ m v(xz)}$ | $\sigma_{ m v'(yz)}$ |
|---|----------------|---------------------|----------------------|
| 9 | -1 | 3 | 1 |



The character for reflection in the yz plane is: 1+1-1=1

| Е | C ₂ | $\sigma_{ m v(xz)}$ | σ _{v'(yz)} |
|---|----------------|---------------------|---------------------|
| 9 | -1 | 3 | 1 |

The above reducible representation is sometimes called Γ_{3N} , because it reduces to all (3N) modes of molecular motion. Γ_{3N} for water reduces to: $3A_1 + A_2 + 3B_1 + 2B_2$

 Γ_{3N} for water = $3A_1 + A_2 + 3B_1 + 2B_2$

Note that there are 9 modes of motion. These include vibrations, rotations and translations.

Molecular Vibrations Γ_{3N} for water = $3A_1 + A_2 + 3B_1 + 2B_2$

Translations have the same symmetry properties as x, y and z.

| C _{2V} | E | C ₂ | σ _v (x z) | σ' _v (yz) | | |
|-----------------|---|-----------------------|------------------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations Γ_{3N} for water = $3A_1 + A_2 + 3B_1 + 2B_2$

| C _{2V} | E | C ₂ | $\sigma_{v}(xz)$ | σ' _v (yz) | | |
|-----------------|---|-----------------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations 2 Γ_{3N} for water =/3A₁ + A₂ + 3B₁ + 2B₂

| C _{2V} | E | C ₂ | σ _v (x z) | σ' _v (yz) | | |
|-----------------|---|-----------------------|------------------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations 2 2 Γ_{3N} for water = $3A_1 + A_2 + 3B_1 + 2B_2$

| C _{2V} | E | C ₂ | σ _v (x z) | σ' _v (yz) | | |
|-----------------|---|-----------------------|------------------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations 2 2 Γ_{3N} for water =/3A₁ + A₂ + 3B₁ + 2B₂

| C _{2V} | E | C ₂ | σ _v (x z) | $\sigma'_{v}(yz)$ | | |
|-----------------|---|-----------------------|------------------------------|-------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations 2 2 1 Γ_{3N} for water = $3A_1 + A_2 + 3B_1 + 2B_2$

| C _{2V} | E | C ₂ | σ _v (x z) | $\sigma'_{v}(yz)$ | | |
|-----------------|---|-----------------------|------------------------------|-------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations $\Gamma_{rot \& vib} = 2A_1 + A_2 + 2B_1 + 1B_2$

| C _{2V} | Е | C ₂ | $\sigma_{v}(xz)$ | σ' _v (yz) | | |
|-----------------|---|-----------------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations $\Gamma_{rot \& vib} = 2A_1 + A_2 + 2B_1 + 1B_2$

| C _{2V} | E | C ₂ | $\sigma_{v}(xz)$ | σ' _v (yz) | | |
|-----------------|---|----------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations $\Gamma_{rot \& vib} = 2A_1 + A_2 + 2B_1 + 1B_2$

| C _{2V} | E | C ₂ | $\sigma_{v}(xz)$ | σ' _v (yz) | | |
|-----------------|---|----------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | xy |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations $\Gamma_{rot \& vib} = 2A_1 + 2B_1 + 1B_2$

| C _{2V} | E | C ₂ | $\sigma_v (xz)$ | σ' _v (yz) | | |
|-----------------|---|-----------------------|-----------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | R _z | xy |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations $\Gamma_{rot \& vib} = 2A_1 + 1B_1 + 1B_2$

| C _{2V} | E | C ₂ | σ _v (x z) | $\sigma'_{v}(yz)$ | | |
|-----------------|---|-----------------------|------------------------------|-------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Rotations and Translations

Trans R_z Ζ Trans У Trans R_y X

Molecular Vibrations $\Gamma_{vib} = 2A_1 + B_1$ The three vibrational modes remain. Two have A_1 symmetry, and one has B_1 symmetry.

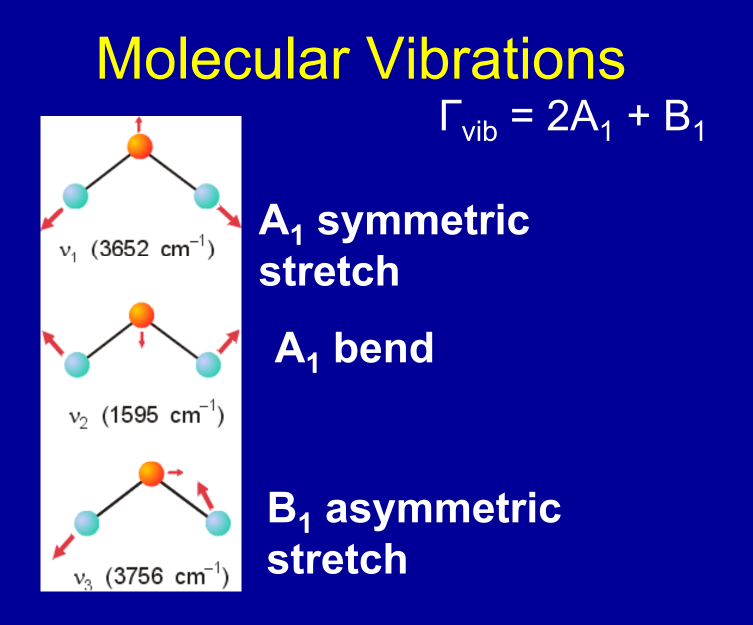
| C _{2V} | E | C ₂ | $\sigma_{v}(xz)$ | $\sigma'_{v}(yz)$ | | |
|-----------------|---|-----------------------|------------------|-------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations $\Gamma_{vib} = 2A_1 + B_1$ Two vibrations are symmetric with respect to all symmetry operations of the group.

| C _{2V} | E | C ₂ | σ _v (x z) | σ' _v (yz) | | |
|-----------------|---|-----------------------|------------------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Molecular Vibrations $\Gamma_{vib} = 2A_1 + B_1$ One vibration is asymmetric with respect to rotation and reflection perpendicular to the molecular plane.

| C _{2V} | Е | c ₂ | σ _v (x z) | σ' _v (yz) | | |
|-----------------|---|-----------------------|------------------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

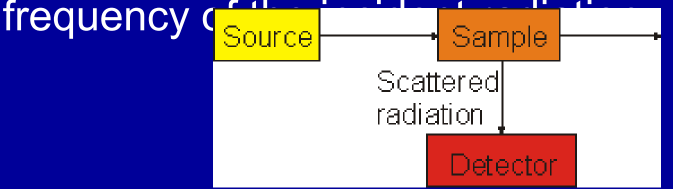


Molecular Vibrations

For a molecular vibration to be seen in the infrared spectrum (IR active), it must change the dipole moment of the molecule. The dipole moment vectors have the same symmetry properties as the cartesian coordinates x, y and z.

Molecular Vibrations

Raman spectroscopy measures the wavelengths of light (in the IR range) scatted by a molecule. Certain molecular vibrations will cause the frequency of the scattered radiation to be less than the





Molecular Vibrations

For a molecular vibration to be seen in the Raman spectrum (Raman active), it must change the polarizability of the molecule. The polarizability has the same symmetry properties as the quadratic functions:

xy, yz, xz, x^2 , y^2 and z^2

| C _{2V} | Е | C ₂ | σ _v (xz) | σ' _v (yz) | | |
|-----------------|---|----------------|---------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² |
| A ₂ | 1 | 1 | -1 | -1 | R, | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

The two vibrations with A_1 symmetry have z as a basis function, so they will be seen in the infrared spectrum of water. This will result in two peaks (at different frequencies) in the IR spectrum of

| C _{2V} | Е | C ₂ | $\sigma_{v}(xz)$ | σ' _v (yz) | | |
|-----------------|---|----------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² |
| A ₂ | 1 | 1 | -1 | -1 | R, | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

The two vibrations with A₁ symmetry also have quadratic basis functions, so they will be seen in the Raman spectrum of water as well.

| C _{2V} | Е | C ₂ | $\sigma_{v}(xz)$ | σ' _v (yz) | | |
|-----------------|---|----------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² |
| A ₂ | 1 | 1 | -1 | -1 | R, | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

The two vibrations with A_1 symmetry will appear as two peaks in both the IR and Raman spectra. The two frequencies observed in the IR and Raman for these vibrations will be the same in both

| C _{2V} | Е | C ₂ | $\sigma_{v}(xz)$ | σ' _v (yz) | | |
|-----------------|---|----------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² |
| A ₂ | 1 | 1 | -1 | -1 | R, | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

The vibration with B_1 symmetry has x and xz as basis functions. This vibration will be both IR active and Raman active. This vibration will appear as a peak (at the same frequency) in both spectra.

| C _{2V} | Е | C ₂ | $\sigma_{v}(xz)$ | σ' _v (yz) | | |
|-----------------|---|----------------|------------------|----------------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² ,z ² |
| A ₂ | 1 | 1 | -1 | -1 | R, | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | xz |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

Both the IR and Raman spectra should show three different peaks.

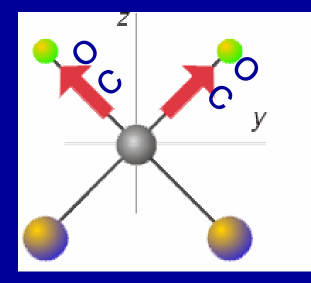
Summary

- 1. Obtain the point group of the molecule.
- 2. Obtain Γ_{3N} by considering the three cartesian coordinates on all atoms that aren't moved by the symmetry operation.
- 3. Reduce Γ_{3N} .
- 4. Eliminate translations and rotations.
- 5. Determine if remaining vibrations are IR and/or Raman active.

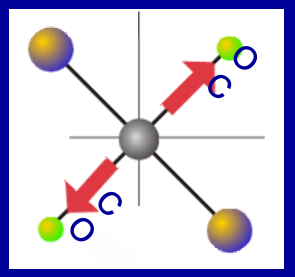
Application: Carbonyl Stretches

 Can IR and Raman spectroscopy determine the difference between two square planar complexes: *cis*-ML₂(CO)₂ and *trans*-ML₂(CO)₂?

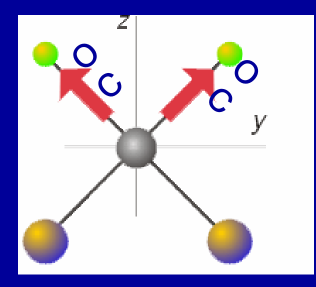
cis and trans $ML_2(CO)_2$



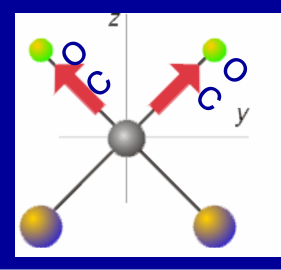
cis isomer – C_{2v}



trans isomer – D_{2h}

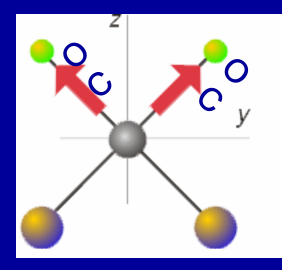


C_{2v} :E C_2 σ_{xz} σ_{yz} Γ_{CO} :2020



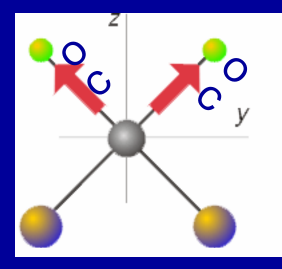
 Γ_{CO} reduces to A_1 + B_1 . A_1 is a symmetric stretch, and B_1 is an asymmetric

| C _{2V} | Е | C ₂ | $\sigma_v(xz)$ | $\sigma'_v(yz)$ | | |
|-----------------|---|----------------|----------------|-----------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | XZ |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |



 Γ_{CO} reduces to A_1 + B_1 . The symmetric stretch (A_1) is IR and Raman active.

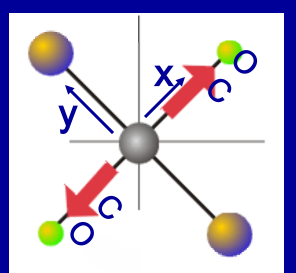
| C _{2V} | E | C ₂ | $\sigma_v(xz)$ | $\sigma'_v(yz)$ | | |
|-----------------|---|----------------|----------------|-----------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | ХZ |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |



Γ_{CO} reduces to A₁ +
 B₁.
 The asymmetric stretch (B₁) is both
 IR and Raman

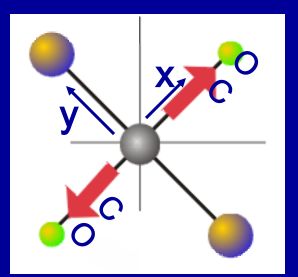
| C _{2V} | E | C ₂ | $\sigma_v(xz)$ | $\sigma'_v(yz)$ | | |
|-----------------|---|----------------|----------------|-----------------|-------------------|--|
| A ₁ | 1 | 1 | 1 | 1 | z | x ² ,y ² , z ² |
| A ₂ | 1 | 1 | -1 | -1 | Rz | ху |
| B ₁ | 1 | -1 | 1 | -1 | x, R _y | XZ |
| B ₂ | 1 | -1 | -1 | 1 | y, R _x | yz |

The *trans* isomer lies in the xy plane. The point group D_{2h} has the following symmetry elements:



trans isomer –

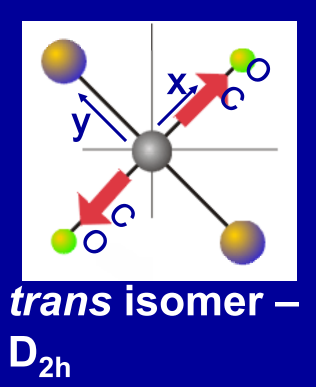
The *trans* isomer lies in the xy plane. Γ_{CO} is obtained by looking only at the two C-O bonds.



trans isomer –

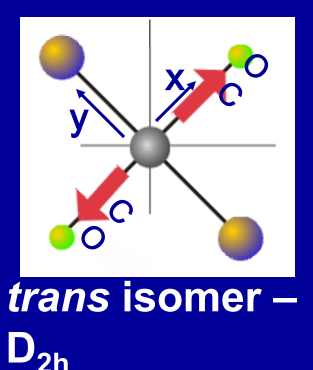
| D _{2h} | Е | $C_2(z)$ | $C_2(y)$ | $C_2(x)$ | i | σ_{xy} | $\sigma_{_{ m XZ}}$ | $\sigma_{ m yz}$ |
|-------------------|---|----------|----------|----------|---|---------------|---------------------|------------------|
| $\Gamma_{\rm CO}$ | 2 | 0 | 0 | 2 | 0 | 2 | 2 | 0 |

 Γ_{co} reduces to A_g (a symmetric stretch) and B_{3u} (an asymmetric stretch).



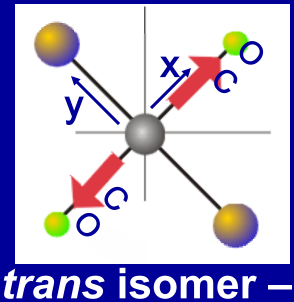
 Γ_{co} reduces to A_g (a symmetric stretch) and B_{3u} (an asymmetric stretch).

 A_g has x^2 , y^2 and z^2 as basis functions, so this vibration is Raman active.



 A_g has x^2 , y^2 and z^2 as basis functions, so this vibration is Raman active.

 B_{3u} has x as a basis function, so this vibration is IR active.



D_{2h}

 A_g has x², y² and z² as basis functions, so this vibration is Raman active.

 B_{3u} has x as a basis function, so this vibration is IR active.

The IR and Raman spectra will each show one absorption at different frequencies.

Exclusion Rule

If a molecule has a center of symmetry, <u>none</u> of its modes of vibration can be both infrared and Raman active.

Exclusion Rule

If a molecule has a center of symmetry, none of its modes of vibration can be both infrared and Raman active The cis and trans isomers of square planar ML₂(CO)₂, can be easily distinguished using spectroscopy. The *cis* isomer has absorptions that are seen in both the IR and Raman spectra, whereas the *trans* isomer