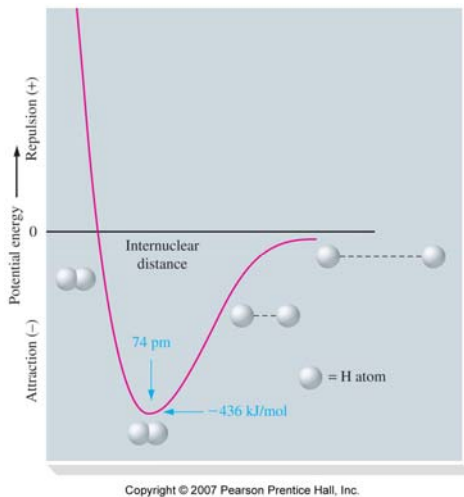


Chemical Bonding and Molecular Structure

What are bonds? Why do bonds form?

Valence Electrons:

- Number and distribution of valence e⁻ determines reactivity
- Main Group Elements (s and p block):
- Transition Metals (d block):
- EXAMPLES: Bromine and Iron



Bonding: Ionic vs Covalent Two Extremes

Ionic Bonding:

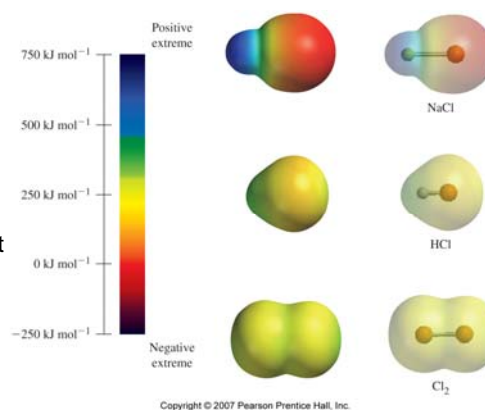
- Electrostatic attraction
 - Discrete charges... “transferred” electrons
- Modeled using Coulomb’s Law

$$F = \frac{Q_1 Q_2}{\epsilon r^2}$$

Q = charge
 r = separation
 ϵ = dielectric constant

Covalent Bonding:

- Also an electrostatic attraction
 - No charge separation... “shared” electrons

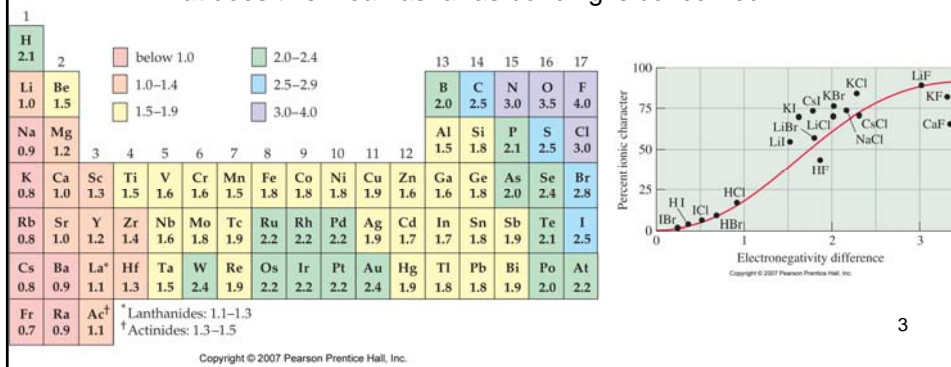


Electronegativity: Essential in understanding bonding and predicting structure.

Electronegativity: ability of an atom in a bond (molecule) to attract electrons to itself

- Trends in electronegativity
- Related to Ionization Energy and Electron Affinity

$$EN \propto (IE - EA)$$
- What does this mean as far as bonding is concerned?



3

Bonding Models

- Attempts to explain real life (shape, reactivity, properties). Sometimes they work better than others. REAL LIFE WINS!

Using Lewis Dot Structures to Model Molecules

- Localized electron bonding model
 - Valence electron bookkeeping
- Primary goal in bond formation:
- Representing bonding using Lewis structures has two key features:
 - 1.
 - 2.
- Best suited for s and p block compounds
 - Particularly up to Ne

4

Guidelines for Drawing Lewis Structures

Underlying criteria: "Octet Rule": Kind of like the Pirate Code

General Scheme: (CH₂Cl₂ and CO₂ as examples)

1. Determine arrangement of atoms (skeletal structure)...HOW?
 - Central vs peripheral (terminal) atoms
2. Determine total # of valence e⁻...HOW?
3. Draw single bonds between central atom and each peripheral atom
4. Distribute remaining e⁻ as lone pairs around peripheral atoms until all have an octet
5. Add multiple bonds to central atom if necessary until all atoms have filled octets.
6. Double-check that all e⁻ have been used and that all atoms have filled octets!

5

Guidelines for Drawing Lewis Structures

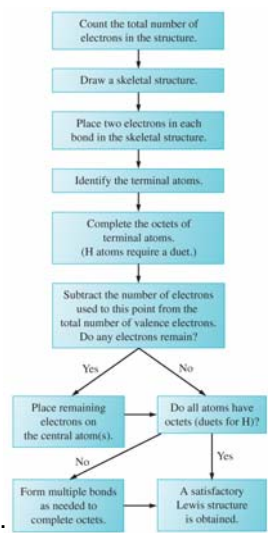
Formal Charge (FC)

- Calculated for a **specific** atom in a molecule
$$FC = \text{group \#} - [\text{\# of e}^- \text{ in lone pairs} + \frac{1}{2}(\text{\# of bonding e}^-)]$$
- **IMPORTANT!** The sum of all the FC for a species or ion **MUST** equal the net charge on the species!

Example: Calculate the FC on each atom in CN⁻:

FC can help when drawing Lewis Structures

1. FC on each atom should be as small as possible
2. (-) FC should appear on the most electronegative atoms, (+) FC on least electronegative.
3. FC of the same sign on adjacent atoms is unlikely.



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Things to Keep in Mind

Molecules are THREE DIMENSIONAL and real life wins!

- Not everything satisfies the octet rule!
 - Hydrogen is different! (so is boron)
 - Some atoms can “expand their octets”
 - Odd electron species exist! (NO, for example)
 - Transition metals and the octet rule.
- Carbon forms four bonds...usually.
- Isoelectronic Species (i.e. NO^+ , N_2 , CO , CN^-)
- Resonance (Section 10.5)

7

Bonding, Lewis Structures, and Molecular Shape

Valence Shell Electron Pair Repulsion (VSEPR) model:

- Bonding and lone pairs of electrons repel each other and try to get as far apart as possible

What shapes would we predict? AX_n or AX_nE_m models. **(It all starts with Table 10.1)**

- X = terminal atoms, E = unshared electron pairs on central atom.
- Suggestion: think of “things” around the central atom.
- Those “things” want to get as far apart in space as possible.
- Unshared pairs take up more room than bonding pairs.
- Electron-pair geometry may be different than molecular geometry

8

Bonding, Lewis Structures, and Molecular Shape

Case 1: No lone pairs around central atoms

- (don't worry about lone pairs on peripheral atoms)
- Electron-pair and molecular geometry are the same.

Case 2: Lone pairs on central atom:

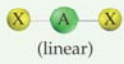
- Two things to remember
 1. Lone pairs occupy space (*more space than bonding pairs*)
 2. Lone pairs repel electrons
- BUT: You predict shape by predicting electron-pair geometry and converting to molecular geometry.
 - Electron-pair and molecular geometry may (and typically are) different.

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Bonding, Lewis Structures, and Molecular Shape

- Families evolve based on total number of "Things" around central atom ($n + m$)
 - *A multiple bond counts as one "Thing"*
- # Things = 2

TABLE 10.1 Molecular Geometry as a Function of Electron-Group Geometry

| Number of Electron Groups | Electron-Group Geometry | Number of Lone Pairs | VSEPR Notation | Molecular Geometry | Ideal Bond Angles | Example |
|---------------------------|-------------------------|----------------------|----------------|---|-------------------|----------|
| 2 | linear | 0 | AX_2 |  (linear) | 180° | $BeCl_2$ |

 VSEPR Basic Molecular Configurations models



^aFor a discussion of the structure of SO_2 , see page 402.

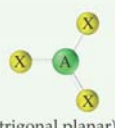
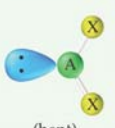
^bFor a discussion of the placement of the lone-pair electrons in this structure, see page 401.

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Bonding, Lewis Structures, and Molecular Shape

- # Things = 3

TABLE 10.1 (Continued)

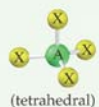
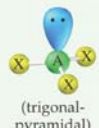
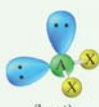
| Number of Electron Groups | Electron-Group Geometry | Number of Lone Pairs | VSEPR Notation | Molecular Geometry | Ideal Bond Angles | Example |
|---------------------------|-------------------------|----------------------|-------------------|--|-------------------|------------------------------|
| 3 | trigonal-planar | 0 | AX ₃ |  (trigonal planar) | 120° | BF ₃ |
| | trigonal-planar | 1 | AX ₂ E |  (bent) | 120° | SO ₂ ^a |

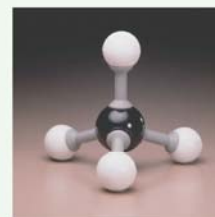


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Bonding, Lewis Structures, and Molecular Shape

- # Things = 4






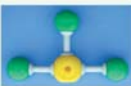

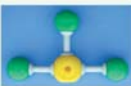
| Number of Electron Groups | Electron-Group Geometry | Number of Lone Pairs | VSEPR Notation | Molecular Geometry | Ideal Bond Angles | Example |
|---------------------------|-------------------------|----------------------|--------------------------------|---|-------------------|-----------------|
| 4 | tetrahedral | 0 | AX ₄ |  (tetrahedral) | 109.5° | CH ₄ |
| | tetrahedral | 1 | AX ₃ E |  (trigonal-pyramidal) | 109.5° | NH ₃ |
| | tetrahedral | 2 | AX ₂ E ₂ |  (bent) | 109.5° | OH ₂ |



Bonding, Lewis Structures, and Molecular Shape

- # Things = 5

TABLE 10.1 (Continued)




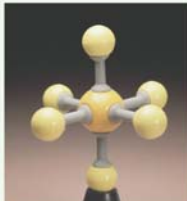

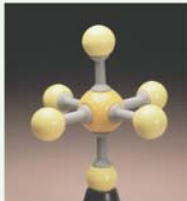
| Number of Electron Groups | Electron-Group Geometry | Number of Lone Pairs | VSEPR Notation | Molecular Geometry | Ideal Bond Angles | Example |
|---------------------------|-------------------------|----------------------|----------------|---|-----------------------|--|
| 5 | trigonal-bipyramidal | 0 | AX_5 |  (trigonal-bipyramidal) | $90^\circ, 120^\circ$ | PCl_5  |
| | trigonal-bipyramidal | 1 | AX_4E |  (seesaw) | $90^\circ, 120^\circ$ | SF_4  |
| | trigonal-bipyramidal | 2 | AX_3E_2 |  (T-shaped) | 90° | ClF_3  |
| | trigonal-bipyramidal | 3 | AX_2E_3 |  (linear) | 180° | XeF_2  |

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Bonding, Lewis Structures, and Molecular Shape

- # Things = 6

TABLE 10.1 (Continued)

| Number of Electron Groups | Electron-Group Geometry | Number of Lone Pairs | VSEPR Notation | Molecular Geometry | Ideal Bond Angles | Example |
|---------------------------|-------------------------|----------------------|----------------|---|-------------------|---|
| 6 | octahedral | 0 | AX_6 |  (octahedral) | 90° | SF_6  |
| | octahedral | 1 | AX_5E |  (square-pyramidal) | 90° | BrF_5  |
| | octahedral | 2 | AX_4E_2 |  (square-planar) | 90° | XeF_4  |

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Bond and Molecular Polarity

Polar bonds result from unequal sharing of electrons

- Due to electronegativity differences
- Produced bond dipoles.

Just as bonds can be polar, molecules can also be polar

- Molecular polarity plays major role in:
 - solubility (like dissolves like),
 - reactivity (biochemical interactions)

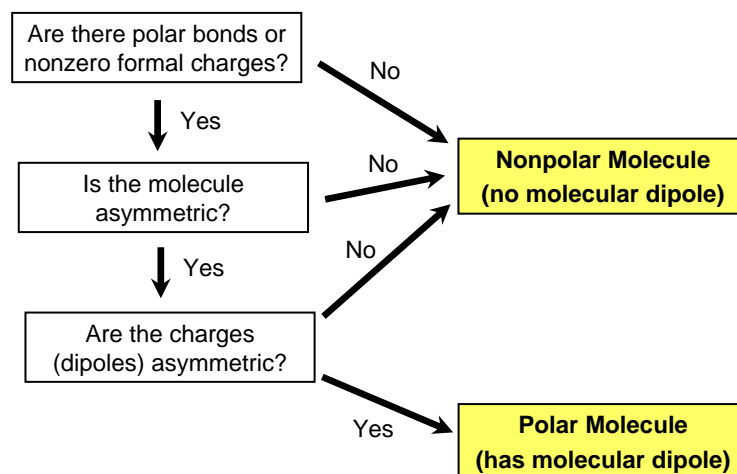
Molecular polarity is determined by polarity and *orientation* of bonds in the molecule

- Need structure first!

Examples: H₂O, CO₂, CCl₄

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Predicting Molecular Polarity



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Bond Characteristics

Bond Order:

Bond Length (Table 10.2):

TABLE 10.2 Some Average Bond Lengths^a

| Bond | Bond Length, pm | Bond | Bond Length, pm | Bond | Bond Length, pm |
|------|-----------------|------|-----------------|-------|-----------------|
| H—H | 74.14 | C—C | 154 | N—N | 145 |
| H—C | 110 | C=C | 134 | N=N | 123 |
| H—N | 100 | C≡C | 120 | N≡N | 109.8 |
| H—O | 97 | C—N | 147 | N—O | 136 |
| H—S | 132 | C=N | 128 | N=O | 120 |
| H—F | 91.7 | C≡N | 116 | O—O | 145 |
| H—Cl | 127.4 | C=O | 143 | O=O | 121 |
| H—Br | 141.4 | C=O | 120 | F—F | 143 |
| H—I | 160.9 | C—Cl | 178 | Cl—Cl | 199 |
| | | | | Br—Br | 228 |
| | | | | I—I | 266 |

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Bond Characteristics

Bond Strength (Bond Dissociation Energy, Table 10.3): Useful in predicting thermodynamics

- Bond breaking: _____ energy
- Bond formation: _____ energy

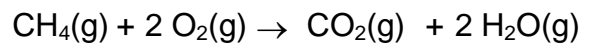
TABLE 10.3 Some Average Bond Energies^a

| Bond | Bond Energy, kJ/mol | Bond | Bond Energy, kJ/mol | Bond | Bond Energy, kJ/mol |
|------|---------------------|------|---------------------|-------|---------------------|
| H—H | 436 | C—C | 347 | N—N | 163 |
| H—C | 414 | C=C | 611 | N=N | 418 |
| H—N | 389 | C≡C | 837 | N≡N | 946 |
| H—O | 464 | C—N | 305 | N—O | 222 |
| H—S | 368 | C=N | 615 | N=O | 590 |
| H—F | 565 | C≡N | 891 | O—O | 142 |
| H—Cl | 431 | C=O | 360 | O=O | 498 |
| H—Br | 364 | C=O | 736 ^b | F—F | 159 |
| H—I | 297 | C—Cl | 339 | Cl—Cl | 243 |
| | | | | Br—Br | 193 |
| | | | | I—I | 151 |

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Calculating $\Delta H^\circ_{\text{rxn}}$ for gas phase reactions:

- Why Gas Phase?



Lewis
Structure

Bonds
Broken

Bonds
Formed

Energies
(kJ)

$$\Delta H_{\text{rxn}} = \Delta H_{\text{bonds broken}} - \Delta H_{\text{bonds formed}}$$

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Isomers

Isomers: different compounds with same molecular formula

Stereoisomers: isomers where connections are same, but arrangement in space different

Constitutional Isomers: isomers where atoms are connected differently

Enantiomers: stereoisomers that are mirror images

Diastereomers: stereoisomers that are not mirror images

EXAMPLE: $\text{C}_2\text{H}_2\text{Cl}_2$ has 3 possible structures, only 2 are diastereomers

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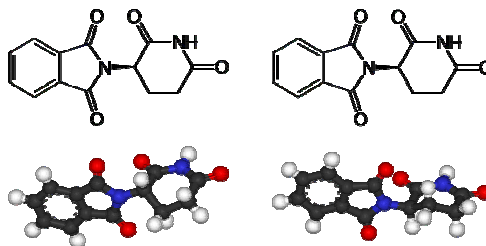
Enantiomers

Molecules that have Enantiomers are Chiral

- Amino Acids are one Enantiomer
- Some bacteria use the amino acids of the other chirality to trick their hosts

Enantiomers have similar physical properties (nearly identical)

- Interact differently with polarized light
- May have dramatically different reactivity
 - Thalidomide



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