Chapter 10

Chemical Bonding I: Basic Concepts

What is a "Chemical Bond"?

A "strong" attractive force between atoms

How strong is "strong"?

- Strong enough to stop the "bonded" atoms from moving away completely when agitated by thermal motion
- Strong enough to compete with attractions to <u>other</u> atoms

2

An attraction is not much of a bond if it cannot compete with other forces, regardless of how strong it is.

To be a "bond", it needs to be the dominant force between the atoms.

3

1

An atom can be bonded to more than one atom at the same time, but each bond <u>remains intact</u> over time. Na⁺ is bonded to 6 Cl⁻ ions (and each Cl⁻ to 6 Na⁺) in a NaCl crystal.





In solution, the force between Na⁺ and Cl⁻ is overwhelmed by the forces between the ions and the water molecules, and there is no "bond" between Na⁺ and Cl⁻. They diffuse away from each other.

4

Any attractive force represents a lowered energy (more stability), whether it is a bond or not.

So, it's definitely true that:

Atoms in a bond have lower energy compared with the same atoms separated from each other.

It may not be the strongest, lowest-energy bond an atom could make, but each atom has lower energy than when it is alone.



A useful way to classify chemical bonds

Ionic

- Due to the electrostatic attraction between a cation
 (+) and an anion (-).
- No sharing of electrons

Covalent

• Due to the attraction of the nuclei of two atoms to a higher density of electrons between them (therefore "shared" by both atoms.)

7



8

What about intermediate cases?

- Most bonds actually fall somewhere in between
- Still , classifying bonds as "ionic" or "covalent" is very useful

We will not consider other kinds of bonding that cannot be easily understood in terms of these two modes of bonding.

9



10

lonic bonds do not lead to molecules.

They lead to "ionic compounds".

In solid form, ionic compounds form regular crystals where the (+) and (-) charges stack in a way that maximizes the force of attraction (and minimizes the potential energy).



But:

The ionic bond doesn't need to be formed as a result of a direct transfer of an electron between two atoms.

All we need are oppositely charged ions. They may be brought together long after they were formed quite separately from one another.



When we melt an ionic substance, individual ions start floating away separately. They **<u>don't</u>** melt as neutral units where the cation and the anion are stuck together.



13



15

Polar covalent bonds and Electronegativity

We learned earlier that some elements "like" electrons more than others, as indicated by their larger ionization energy or electron affinity.

- > It costs more to remove an electron from them
- More energy is released when an electron is added to them

When an element is making a covalent bond, its relative attraction to electrons doesn't go away.

If it attracted electrons strongly alone, it will still do that when bonded.

Covalent bonding

Another way two atoms can be more stable together than apart is by offering the valence electrons the chance to spend more time between the nuclei.

The (–) electron would be near not one but <u>two</u> (+) <u>nuclei</u>, making the system more stable (lower in potential energy)

When this opportunity arises, there is no reason for only one atom's electron to do it. <u>An electron from both atoms</u> will want to concentrate in the space between them.

14



If the atoms are farther than the "bond length", potential energy of the electrons are higher.

The energy it takes to pull the atoms from "bond length" to completely apart is called "bond dissociation energy".



One intuitive way to define electronegativity for an element is to average the <u>ionization energy</u> and <u>electron</u> <u>affinity</u> (which measure how much it "likes" electrons)

> and then scale the set of numbers obtained for all elements appropriately.

That is different from how Linus Pauling* defined it, but that's ok; there are many ways to define the idea of electronegativity mathematically.

*Linus Pauling was a famous Nobel laurate chemist

19

The "shared" bonding electrons spend more time near the more electronegative atom.

Increased electron density around the more electronegative atom gives it an extra "partial" negative charge (shown as δ -).

Conversely, the less electronegative atom develops a partial positive charge (shown as δ +).



21



F, O, N, and Cl have the highest electronegativity values.

20



22



To compare the polarities of bonds

- We don't always need the exact electronegativity values
- We don't need the exact values of electronegativity differences
- We can use the relative locations in the periodic table
- The farther the two elements are from each other in the direction from the top-right corner (high electronegativity) of the periodic table to the bottom left (low electronegatiovity), the more polar the bond will be

He

Ne

Ar

Kr

CI

Br

Ĩ. Xe

Å ċ Ń ò Ē

AI Si

Ga

11/22 1/

Zn

Cd Hg

32 Ge

Pb Bi

P S

As

Sb Sb

Te

Po At





In cases when it's too difficult to use the periodic table to compare electronegativities, we can use a table of electronegativities.

- But be aware that there is no precise relationship between electronegativity difference and bond polarity
- Even equal electronegativities can go with a nontrivial bond polarity
- Purely quantum mechanical factors can be as important as electronegativity difference

27



- Larger the separated charges, larger the dipole moment
- Farther the separation, larger the dipole moment

We use an arrow to represent a dipole moment.

- Tail of the arrow at the positive center of charge.
- Point to the negative charge center

"Chemist's dipole"

Unfortunately the opposite of the convention used in Physics, or by whomever does any actual calculations with dipoles 🙃



When we are using just the periodic table, we will avoid cases

N being higher than P in the periodic table makes its EN go up, and

But O is so much higher (both in position and electronegativity)

than Te that EN difference between O and I is greater than

between Te and I (O is a lot more electronegative than I)

Mo Tc Ru Rh

when the relative electronegativities are ambiguous.

be closer to that of Cl.

Sc Martine Y

Lu

67-78 ¥

Ťi v CI

Žr

Hf

de

Ta

w Re 76 OS n Ir

Ĥ

Li Be

Na Mg

Ca

Sr

ĸ

Rb

Cs Ba

26

Bond polarity and molecular polarity

- A molecule cannot be polar without polar bonds
- But the presence of polar bonds doesn't guarantee that the molecule is polar because bond dipoles can cancel out
- If the bond dipoles of the polar bonds cancel then the molecule is non-polar. Otherwise it's polar.

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Polar Molecules, Non-polar Molecules

• <u>In practice</u>, if a molecule's dipole moment is "small enough" it would be called "nonpolar"

<u>But</u>

- For our purposes, we will call a molecule "nonpolar" only if its dipole moment is <u>exactly</u> zero
 - It's impractical to calculate the molecular dipole moment at this stage
 - \geq But we can determine if the dipole moment is zero or not
- And we will call a molecule "polar" if its dipole moment is nonzero (no matter how small)

33

Molecular polarity and symmetry A simple way to judge polarity: Symmetric I on the symmetric (lopsided) Polar

We learned that larger electronegativity difference between bonded atoms leads to more polarity in the bond

- If bond polarity is high enough, then the bond is "ionic"
- If not declared "ionic", we will call the bond "polar covalent" as long as there is an electronegativity difference between the bonded atoms



- 100% covalent bond is possible In <u>any diatomic molecule</u>, atoms share the bond electrons equally
- Even the largest electronegativity difference <u>cannot</u> give a 100% ionic bond
 - CsF is less than 75% ionic

Practice:

Arrange the following bonds from most to least polar:

a) N–F	O–F	C–F
b) C—F	N–O	Si–F
c) Cl–Cl	B–Cl	S–Cl

39

Practice:

Which of the following bonds would be the most polar without being considered ionic?

Mg-0 C-0 0-0 Si-0 N-0

To not be considered ionic, the bond needs to be between two nonmetals.

The most polar bond between the nonmetals occurs with the bond that has the greatest difference in electronegativity.



Simple general rule for binary compounds:

nonmetal-nonmetal bond: Covalent Never ionic metal-nonmetal bond*: Ionic * In some cases it may be polar <u>covalent</u> But then we will be told that we have a <u>molecule</u>

38

Practice:

Which of the following bonds in a binary compound would be the least polar yet still be considered polar covalent?

Mg-0 C-0 0-0 Si-0 N-0

To be considered polar covalent, unequal sharing of electrons must still occur. We choose the bond with the least difference in electronegativity, but still with some unequal sharing of electrons.

40

Electron Configurations in Stable Compounds

Considering only compounds of <u>main-group elements</u>:

Covalent bond:

- Shared electrons fill up the valence shells of <u>both</u> atoms.
- Both atoms achieve the electron configuration of the <u>next</u> noble gas.

Ionic bond:

- The valence shell of the **nonmetal** achieves the electron configuration of the <u>next</u> *noble gas*.
- The valence shell of the metal is emptied, and it achieves the electron configuration of the previous noble gas.

In other words:

For <u>main group elements</u> (i.e. elements in the leftmost 2 and rightmost 6 columns of the periodic table; i.e. outside of transition block), <u>bond formation</u> almost always does this:

Attain the electron configuration of the closest noble gas.

This is true both for ionic and covalent bonds.

43

Localized Electron Model

In this model:

- A molecule is composed of atoms that are bound together by sharing pairs of electrons using the atomic orbitals of the bound atoms.
- But we still consider the electrons as contributing to the valence shell of individual atoms, rather than to the entire molecule (or the polyatomic ion)

45

Modeling Bonding

- The reality of what keeps atoms in compounds together is complex.
- But it would be wasteful, impractical, and to solve the quantum mechanical equations every time we ask a question about molecules.
- The complexity can be reduced by using <u>models</u> that are good enough to be useful and simple enough to think about compounds and molecules in an intuitive, tangible way.

44

Localized Electron Model does three things:

- Description of valence electron arrangement (Lewis structure). Only 2-dimensional description here.
- 2. Description of atomic orbital types used by atoms to share electrons or hold lone pairs. What happens to atomic orbitals when they form bonds
- 3. Prediction of geometry (via the VSEPR model)
 3-Dimensional description of molecular structure
 > We will see it at the end of this chapter

46

Lewis Structures

A simple and surprisingly powerful way to do "quantum chemistry" for molecules and polyatomic ions

Uses the "Octet Rule":

Each atom in a covalently bonded entity "wants":

8 electrons (an "octet") in its valence shell (except for H, which always gets 2 electrons) Octet rule is applied through electron pairs

$\label{eq:valence} Valence \ electron \ \underline{pairs} \ making \ octets \ everywhere$

- Each atom has 4 pairs of electrons, making an "octet"
 Hydrogen attains only one pair (a "duet")
- Each bond is a pair of electrons ("bonding pair")
- Non-bonding electrons also form pairs ("lone pair")
- Exceptions to the "octet rule" exist, and can be handled

General steps for writing Lewis structures:

- 1. Calculate the number of valence electrons.
- 2. Draw the "skeletal bonds" that show which atom is connected to which. Each bond is a pair of electrons.
- 3. Complete the octets of atoms (except H) by creating lone pairs, starting with the peripheral atoms.
- If the central atom doesn't have an octet when you run out of your valence electron budget, turn a peripheral atom's lone pair into an extra bond with the central atom.
- 5. Repeat Step 4 if needed, using a lone pair of another (rather than the same one as before) peripheral atom whenever possible.
- Admire your work. You just did "quantum chemistry" in a very quick way that is unfairly powerful!

49

Step 1: Calculate the number of valence electrons Sum up the valence electrons brought by each atom • Subtract the charge (so, a positive charge reduces the number of electrons, and a negative charge increases it) • This is our "budget" of electrons Example: H₂O 1 valence electron from each H (2)(1) + (1)(6) = 8 val. e⁻ 6 valence electrons from O Example: H₂O⁺ Each H: 1 val. e⁻; each O: 6 val. e⁻ (3)(1) + (1)(6) - 1 = 8 val. e⁻ 1 (+) charge to subtract Example: OH⁻ 1 val. e⁻ from H, 6 val. e⁻ from O (1)(1) + (1)(6) - (-1) = 8 val. e 1 (–) charge to subtract

51



A couple of terms to clarify

"Central atom"

- The atom at the center of the molecule or ion
- In some cases there may be more than one "central atom", and we treat each one the same way
- They may violate the octet rule

"Peripheral atoms"

- Atoms bonded to the central atom
- They don't violate octet rule

50





"Rules of thumb" for Lewis structures

If the molecular formula contains a single atom of an element (and there is only one such element in the formula) it is safe to designate it as "central atom"

e.g.
$$NH_3$$
, H_2O , CO_2
central central central

55



56

"Rules of thumb" for Lewis structures

If the chemical formula is written symmetrically, the skeletal structure is likely symmetric. The atoms in the middle are "central" (more than one "central" atom is possible) and will be treated as such separately.

Even if the formula isn't written symmetrically, symmetric structures are often favored in nature.

First see if you can draw a symmetric skeletal structure before attempting asymmetric ones.

57





- Hydrogen is never the central atom
- Hydrogen only makes single bonds
- Hydrogen never has a lone pair
- Halogens only make single bonds as peripheral atoms
- Carbon almost never has lone pairs (except in CO and CN⁻)
- Carbon never violates the octet rule
 - It may have an incomplete octet in a reactive cation or a radical, but you should give it an incomplete octet only if absolutely must









Each structure has 2 N-O single bonds and 1 N=O double bond But the double bond is with a different O in each.



62





When the resonance structures are equivalent, they become important. They imply that:

- The electrons of the extra bond(s) cannot "decide" which bond distribution is best
- The electrons in the extra bonds(s) get distributed, i.e. "delocalized"
- Those electrons add to the bonding of more than one skeletal bond.
- The extra bonding they bring to each skeletal connection is only partial

Actual structure is an <u>average</u> of the resonance structures.

The electrons in the extra bond are **<u>delocalized</u>**.



The "moving" bond is taking the place of a lone pair, which must also delocalize. But it's harder to show. So we don't show lone pairs when we show delocalized bonds

- · One extra bond is spread over 3 skeletal bonds
- It adds 1/3 bond to each bond
- Each N—O bond is 1+1/3 = 1.33 bond
- Slightly stronger and shorter than a single bond
- But not quite a double bond

67

Even though formal charges aren't completely "real", they do

have enough reality to give good guidance on the local charges of a covalent entity (molecule or a polyatomic ion).

They can explain the polarity of a bond you would expect to be nonpolar, or the non-polarity of a bond you would expect to be polar.

They are useful in developing intuition on the reaction mechanisms.

For example, if an atom has a positive formal charge, it may be "vulnerable to an attack" by an electron-rich reactant.

69



Formal Charge

The charge an atom **would** have **if** electrons in bonds were shared **equally**.

Electronegativities are completely ignored.

(Exact opposite of the pretense in assigning oxidation numbers where each atom's electronegativity was exaggerated as if each atom in a compound was an ion)

Formal charge is due to the **electron surplus or deficit** compared with the number of valence electrons in the non-bonded atom.

68

In calculating formal charges, we assume:

- Both electrons of a lone pair of an atom belong to that atom.
- Only one electron of a bond belongs to the atom.

Formal charges must add up to the net charge of the species.





In the preferred resonance structure:

- Formal charges should be as close to zero as possible.
- Formal charges match the atom's electronegativity
 (-) formal charge on the more electronegative atom
 (+) formal charge on the less electronegative atom

















If you find it difficult to understand or remember the "usual number of bonds" for a given element, the shortcut is probably not useful for you, and you can ignore it.

79

Consider the Lewis structure for POCl₃. Assign the formal charge for each atom in the molecule. P: 5 - 4 - 0 = +1O: 6 - 6 - 1 = -1Consider the Lewis structure for POCl₃. Assign the molecule. Consider the Lewis structure for POCl₃. Assign the molecule.

Seeing that the octet rule is not violated (we only need to check the central atom), we could also use the shortcut for formal charges.

Cl: 7 - 6 - 1 = 0

P: 4-3 = +1 O: 1-2 = -1 Cl: 1-1 = 0 If you visually get used to the usual number of bonds an element makes, you can basically "see" the formal charge by the number of bonds.

81



- Boron (B) and Aluminum (AI) often have only 6 electrons in their covalent compounds. Can have more if both of a bonding pair is brought by another atom.
- Peripheral atoms never violate the octet rule



shortcut Oxygen (Group 6A) normally makes 2 bonds Sulfur (Group 6A) normally makes 2 bonds



80









87



Even though it's pretty standard practice to reduce formal charges in oxyanions by making double bonds to the central atom, we won't do that.

Do not violate the octet rule just to reduce formal charges.

It's extra work to do something that has little evidence from nature, and predicts nothing new about the ion.

If an oxyanion has 4 oxygens, giving each one a single bond and a -1 formal charge will give you all the predictions you need!

86





of Lone

Pairs

Structure

How a Lewis Structure leads to Molecular Geometry: VSEPR Model

"Valence Shell Electron-Pair Repulsion"

- Predicts the 3-dimensional shape of the molecule
- The structure around a given atom is determined by minimizing electron pair repulsions.
- Electron groups locate and orient themselves as far apart from one another as possible

91

Steps to Apply the VSEPR Model

- 1. Draw the Lewis structure for the molecule.
- 2. Count the electron groups and arrange them in a way that minimizes repulsion (put the groups as far apart as possible.) You get the geometry of the electron groups.
- 3. Put the peripheral atoms at the end of bonding groups Choose the location of the bonds and lone pairs if necessary.
- Determine the name of the <u>molecular structure</u> from positions of the <u>atoms</u>.

If you find it hard to think in 3D, you can memorize the table(s) at: https://www.papazyan.org/MolecularGeometry,Polarity,Hybridization.pdf

93



VSEPR Model

Actually it would be more accurate to call it VSEGR

Valence Shell Electron-Group Repulsion?

It's the electron group that counts.

A double-bond (2 pairs) or a triple bond (3 pairs) is 1 group.

A lone pair is also one group

92







Ideal Bond Angles & Actual Bond Angles

- Lone pairs are "fatter" and closer to the central atom than bonding pairs
- They push the bonds towards one another
- When the push from lone pairs do not cancel one another:
 >Bond angles are smaller than ideal
- If there is only one lone pair, no cancellation possible
 Bond angles are smaller than ideal
- When lone pairs are symmetrically placed and cancel one another's effects:
 - Bond angles are ideal

99



- In XeF₂ the lone pairs press on the Xe-F bonds <u>symmetrically</u> (no net push from any one side)
- F-Xe-F bond angle is equal to the ideal linear angle of 180°



98



100

Practice

Determine the shape for each of the following molecules, and include bond angles:

HCN PH₃

SF₄

Practice

Determine the shape for each of the following molecules, and include bond angles:



103



105

Guessing symmetry from bond-angle distortion

- Imagine the lone pairs as huge, and pushing on the bonds
- Are the lone pairs completely cancelling each other's push?
- If yes, the bond angles remain ideal, and also the molecule is symmetric
- If no, the bond angles get "squished" to be smaller than ideal, and also the molecule is asymmetric



Bond angle = 180° (ideal)

Nonpolar molecule





Bond angle < 109.5° Polar molecule

Polar Molecules, Non-polar Molecules

Remember:

• In practice, if a molecule's dipole moment is "small enough" it would be called "nonpolar"

But

- For our purposes, we will call a molecule "nonpolar" only if its dipole moment is exactly zero
- And we will call a molecule "polar" if its dipole moment is nonzero (no matter how small)

104

How "symmetric" does a molecule need to be in order to be "nonpolar" (have a zero dipole moment)? If we see a T-shaped or a V-shaped (bent) molecule, we

might perceive it as "pretty symmetric"



But they are not symmetric enough to be non-polar There is a trick for figuring out symmetry ...

106

If all the peripheral atoms are the same, the "bond-angle intuition" works simply:

- ➢ If bond angles are ideal → zero dipole
- ➢ If bond angles are not ideal → nonzero dipole

But please remember:

Polarity is not caused by the non-ideal angles Both polarity and non-ideal bond angles are caused by asymmetry

What if there are no lone pairs, and there are more than one kind of peripheral atoms?

If one or more of the peripheral atoms are different from the rest, <u>treat them as if they are lone pairs</u> and see if they would distort the bond angles.

As with real lone pairs:

- If they would distort the angles, the molecule is asymmetric, therefore polar
- If they would cancel each other's push and the angles would remain ideal, then the molecule is symmetric, therefore nonpolar.

109

Additional online resources on molecular geometry <u>They are linked under Ch. 8 resources</u> **Two very cool and useful tools:** 3D VSEPR simulation phet.colorado.edu/sims/html/molecule-shapes/latest/molecule-shapes_en.html

Molecular Polarity simulation phet.colorado.edu/sims/html/molecule-polarity/latest/molecule-polarity_en.html

A table summarizing VSEPR (plus hybridizations covered in Ch. 9): www.papazyan.org/MolecularGeometry,Polarity,Hybridization.pdf

111



Imagine the tiny hydrogen in CHCl₃ as a huge lone pair. Would it push on the other bonds and reduce the bond angles?

Yes. It would.

Therefore CHCl₃ is asymmetric enough to be polar.

110

Bond lengths and bond strengths

Many factors contribute to bond strength, or whether a bond forms at all, but:

- <u>All else being equal</u>, <u>shorter</u> bond length goes with <u>stronger</u> bond
- Both for covalent and ionic bonds, the sizes of the atoms (or ions) making the bond limit how short the bond can be.
 - $\ensuremath{\mathsf{Therefore}}\xspace$ how strong the bond can be
- When the bond is made by large atoms (or ions):
 Shared electrons can't be very close to the nuclei
 >lons can't get close to each other

112

Bond Enthalpies (Bond Energies)

"Bond Energy" and "Bond ethalpy" often used interchangeably It's the energy needed to break a bond

Also known as "bond dissociation energy"

To break bonds:

Energy must be *added* to the system Endothermic

To <u>form</u> bonds: energy is *released* Exothermic

A small bond enthalpy (BE) means:

- The bond is relatively weak.
- The bond released little energy when formed.
- The atoms forming the bond did not reduce their energy much.
- The atoms are relatively high in energy.

A large bond enthalpy (BE) means:

- The bond is relatively strong.
- The bond released a lot of energy when formed.
- The atoms forming the bond reduced their energy by a lot.
- The atoms are relatively low in energy.

115





If a reaction breaks weak bonds (small BE) and forms strong bonds (large BE):

- The energy released by forming strong bonds is larger than the energy needed to break the weak bonds.
- There will be a net energy release as a result.
- Exothermic reaction.

If a reaction breaks strong bonds (large BE) and forms weak bonds (small BE):

- The energy released by forming weak bonds is less than the energy needed to break the strong bonds.
- There will be a net energy absorption as a result.
- Endothermic reaction.

116



118

- Figuring out the bonds broken and the bonds formed can be an error-prone process
- Instead, we can simply sum over <u>all</u> the bonds in the reactants and do the same for products. Bonds that remain intact during the reaction will appear in both sums and will cancel out
- More numbers to add, but less attention is required, less error-prone

$$n = \begin{pmatrix} Sum \text{ of } \\ bond \text{ BEs in} \\ reactants \end{pmatrix} - \begin{pmatrix} Sum \text{ of } \\ bond \text{ BEs in} \\ products \end{pmatrix}$$

Caution: This subtraction is in the opposite order compared with the formula for obtaining ΔH_{rxn} from ΔH_f values.

 ΔH_{rx}



