



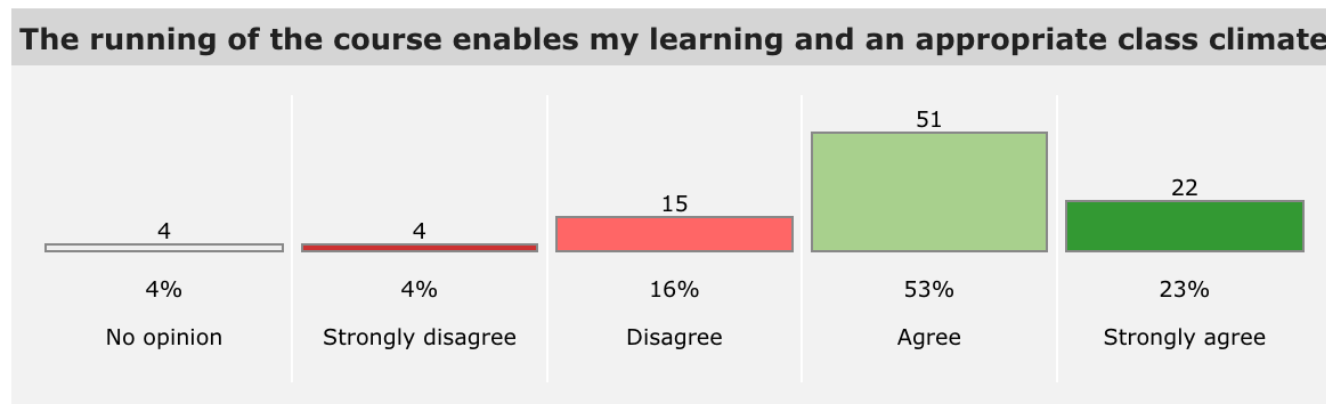
# CH-110 Advanced General Chemistry I

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# Housekeeping notes

Indicative feedback:

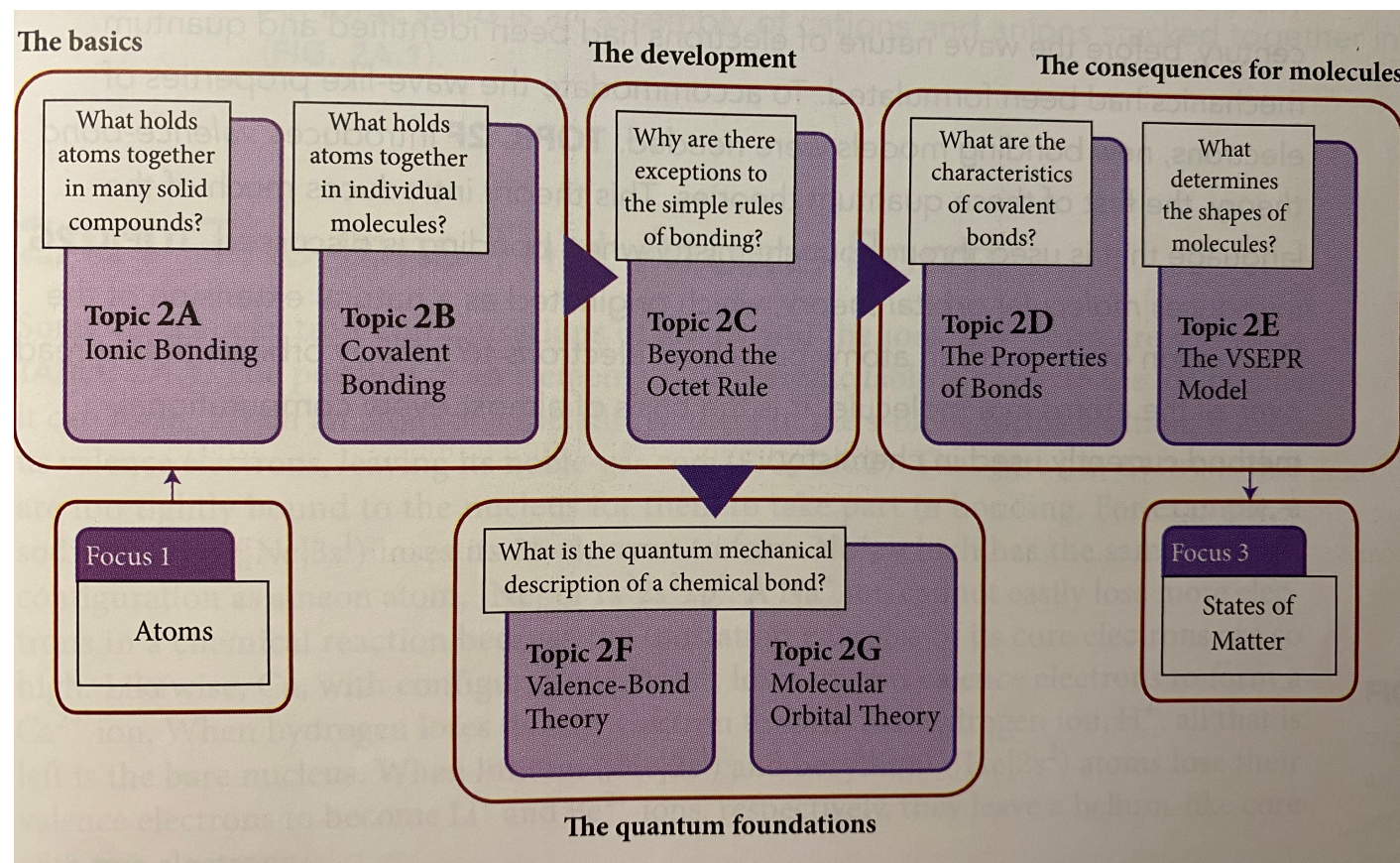
- Too many slides, not clear what is important.
- No French translation for week 3 (now available).
- Too much noise in the classroom.



# Bonds Between Atoms

Focus 2

# Overview Chapter 2 (Focus 2: Bonds Between Atoms)



# Covalent Bonding

Topic 2B

**Last week:** Topic 2B.1 Lewis Structures

Topic 2B.2 Resonance

Topic 2B.3 Formal Charge

WHY DO YOU NEED TO KNOW THIS MATERIAL?

- Understanding the nature of covalent bonding is essential for interpreting structures, properties and reactions of molecules.

WHAT DO YOU NEED TO KNOW ALREADY?

- Electron configurations of many-electron atoms (Topic 1E)
- Lewis symbols (Topic 2A)
- Nomenclature for different types of compounds (Fundamentals D and C)
- Concept of oxidation number (Fundamentals K)
- Concept of electronegativity (Topic 1F)

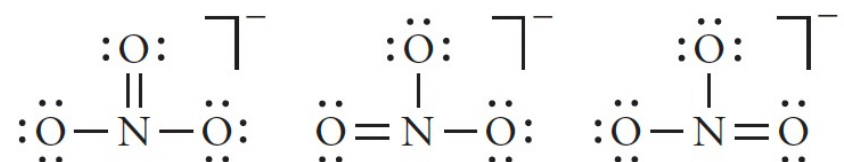
# Resonance

Topic 2B.2

## 2B.2: Resonance

### Resonance and bond order

- Some molecules are not represented adequately by a single Lewis structure.
- Nitrate,  $\text{NO}_3^-$ , has three valid Lewis structures of equal energy:

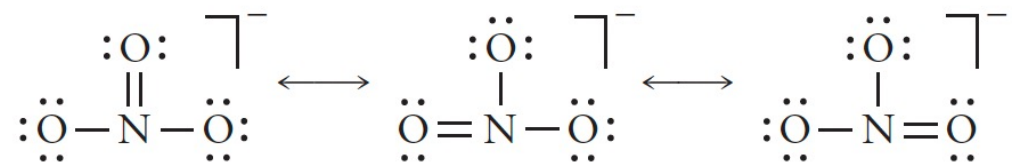


- Double bonds are shorter than single bonds, they hold atoms together more tightly.
- Observed bond length is the same for all three bonds: 124 pm
- Typical N-O single bond: 140 pm, typical N=O double bond: 120 pm
- **Bond order** is between 1 and 2 for nitrate NO bond

## 2B.2: Resonance

### Resonance structures

- All three bonds are identical in  $\text{NO}_3^-$
- Better model is to **blend all three Lewis structures**, each bond has intermediate properties between a single and a double bond



- **Blending of structures is called resonance depicted by double-headed arrows**
- The real structure is a **resonance hybrid**, the molecule does not flicker between structures, but is in a constant in-between state (analogy: mule, not horse, not donkey)

## 2B.2: Resonance

### Delocalization

- Electrons that are shown in different positions in a set of resonance structures are said to be **delocalized**.
- Delocalization means that a **shared electron pair is distributed over several pairs of atoms** and cannot be identified with just one pair of atoms.
- The three resonance structures of nitrate do not exist as actual molecules, they are simply a way of showing that the **electrons are spread across the molecule**.
- Delocalization **lowers the energy** of a molecule.

## 2B.2: Resonance

### Rules to write appropriate resonance structures

- In each contributing structure, **the nuclei of the atoms are in the same positions**; only the locations of lone pairs and bonding pairs are different.
- Equivalent structures (structures that differ only in the positions of multiple bonds and lone pairs) **contribute equally to the resonance**.
- **Low-energy structures contribute more** to the resonance mixture than high-energy structures. (You will see how to judge relative energies in the next section.)

Note: resonance does NOT occur between structures with atoms in different arrangements. For example, no resonance occurs between hypothetical structures NNO and NON.

## 2B.2: Resonance

### **Example 2B.3: Writing a resonance structure (blackboard)**

Suggest two Lewis structures that contribute equally to the resonance structure for the  $\text{O}_3$  molecule. Experimental data show that the two bond lengths are the same.

## 2B.2: Resonance

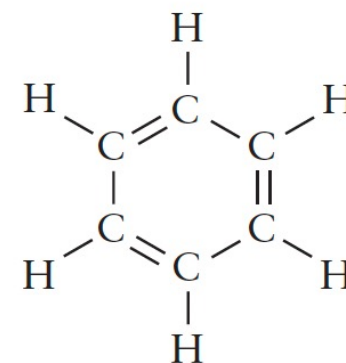
### **Example 2B.3: Writing a resonance structure (blackboard)**

Write the Lewis structures contributing to the resonance hybrid for the (a) acetate ion,  $\text{CH}_3\text{CO}_2^-$  and (b) nitrite ion,  $\text{NO}_2^-$ .

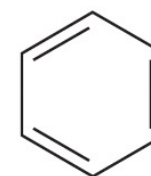
## 2B.2: Resonance

### Benzene, C<sub>6</sub>H<sub>6</sub>

- Benzene is also a resonance hybrid
- Planar hexagonal ring of six C with each an H attached
- One Lewis structure shown (**11**): **Kekulé structure**
- Equivalent line or stick form shown in (**12**)



11 Kekulé structure



12 Kekulé structure, stick form

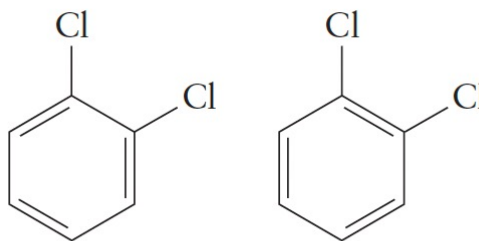
## 2B.2: Resonance

### Benzene's properties do not align with Kekulé structure

One Kekulé structure does not fit all the experimental evidence:

- 1. Reactivity:** benzene does not undergo reactions typical of compounds with double bonds
- 2. Bond lengths:** All carbon-carbon bonds in benzene are the same length
- 3. Structural evidence:** only one version of 1,2-dichlorobenzene (in which the chlorine atoms are attached to two adjacent carbon atoms) exists.

If the Kekulé structure were correct, there would be two distinct versions of 1,2-dichlorobenzene:

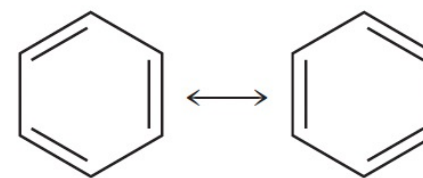


However, only one known to exist!

## 2B.2: Resonance

### Concept of resonance resolves contradictions

- **Two Kekulé structures with exactly the same energy:** they only differ in position of double bonds (**14**)
- Electrons shared in C=C double bonds are **delocalized over the entire molecule**
- Bond lengths intermediate between single and double bond and identical (circle in **15**)
- Resonance stabilizes a molecule → benzene less reactive than expected



14 Benzene resonance structure



15 Benzene, C<sub>6</sub>H<sub>6</sub>

## 2B.2: Resonance

### Summary

Resonance is a blending of structures with the same arrangement of atoms but different arrangements of electrons. Resonance results in electron delocalization; it spreads multiple-bond character over a molecule and results in a lower energy.

# Formal Charge

Topic 2B.3

## 2B.3: Formal charge

### Nonequivalent Lewis structures

**Nonequivalent** Lewis structures are Lewis structures that **do not correspond to the same energy**.

- They do NOT in general make the same contribution to a resonance structure.

How do you decide which resonance structures has the lowest energy?

- Compare the **number of valence electrons distributed around each atom** in a structure with the number of **valence electrons on the free atom**. The smaller the differences are, the lower is its energy and the greater is its contribution to a resonance hybrid.

## 2B.3: Formal charge

### Formal charge as a measure of electron redistribution

- **Formal charge:** the charge an atom would have if the bonding were perfectly covalent (each atom had exactly a half-share in each pair of bonding electrons).

#### How many electrons does an atom "own" in a molecule?

- It "owns" all its lone pairs and half of each shared bonding pair.
- The formal charge is calculated as the difference between this number and the number of valence electrons in the free atom:

$$\text{Formal charge} = V - \left( L + \frac{1}{2} B \right)$$

$V$  is the number of valence electrons,  $L$  is the number of electrons present on the bonded atom as lone pairs, and  $B$  is the number of bonding electrons on the atom.

## 2B.3: Formal charge

### TOOLBOX 2.2

### HOW TO USE FORMAL CHARGE TO DETERMINE THE MOST LIKELY LEWIS STRUCTURE

#### CONCEPTUAL BASIS

To assign a formal charge, we establish the “ownership” of the valence electrons of an atom in a molecule and compare that ownership with the free atom. An atom owns one electron of each bonding pair attached to it and owns its lone pairs completely. The most plausible Lewis structure will be the one in which the formal charges of the atoms are lowest.

#### PROCEDURE

*Step 1* Decide on the number of valence electrons ( $V$ ) possessed by each free atom by noting the number of its group in the periodic table.

*Step 2* Draw the Lewis structures.

*Step 3* For each bonded atom, count each electron in its lone pairs ( $L$ ), plus one electron from each of its bonding pairs ( $B$ ).

*Step 4* For each bonded atom, subtract the total number of electrons it “owns” from  $V$ , as in Eq. 4.

Each equivalent atom (the same element, the same number of bonds and lone pairs) has the same formal charge. A check on the calculated formal charges is that their sum is equal to the overall charge of the molecule or ion. For an electrically neutral molecule, the sum of the formal charges is zero. Compare the formal charges of each possible structure. The structure with the lowest formal charges represents the least disturbance of the electronic structures of the atoms and is the most plausible (lowest energy) structure.

*This procedure is illustrated in Example 2.6.*

## 2B.3: Formal charge

### **Example 2B.4: Selecting the most likely arrangement of atoms (blackboard)**

Write three Lewis structures with different atomic arrangements for the thiocyanate ion ( $\text{SCN}^-$ ) and select the most likely structure by identifying the structure with formal charges closest to zero. For simplicity, consider only structures with double bonds between neighbouring atoms.

Reminder: Electronegativity TREND:  $\text{F} > \text{O} > \text{Cl} > \text{N} > \text{Br} > \text{I} > \text{S} > \text{C/Se} > \text{H} > \dots$

## 2B.1: Lewis structures

### Student question (blackboard)

The Lewis structure of chlorite ion could also have a double bond, why is the structure shown in class the preferred structure?

## 2B.3: Formal charge

### Formal charge to predict atom arrangement in molecules

- If the atom has **more electrons in the molecule** than when it is a free, neutral atom, then the atom has a **negative formal charge**, like a monatomic anion.
- Of the assignment of electrons **leaves the atom with fewer electrons** than when it is free, then the atom has a **positive formal charge**, as if it were a monatomic cation.

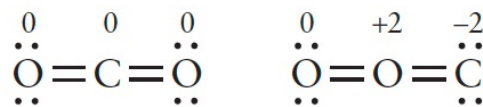
Formal charge can be used to predict the most favorable arrangement of atoms in a molecule and most likely Lewis structure for that arrangement:

- A Lewis structure in which the **formal charge of the individual atoms are closest to zero** typically represents the lowest-energy arrangement of the atoms and electrons.
- The structure of lowest energy is typically the one with a **positive formal charge on the least-electronegative atoms**.

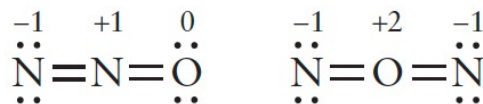
## 2B.3: Formal charge

### Formal charge to predict atom arrangement in molecules

- A **low formal charge** (as close to zero as possible) indicates that an atom has undergone only a small redistribution of electrons relative to the free atom.
- For carbon dioxide, the formal charge predicts that the structure OCO is more likely than COO.
- Similarly, it predicts that NNO is more likely than NON for N<sub>2</sub>O.



17



18

## 2B.3: Formal charge

### Formal charge vs. oxidation number

Formal charge and oxidation number both provide information about the number of electrons around an atom in a compound, but are distinct in origin and significance:

- Formal charge **exaggerates the covalent character** of bonds by assuming that the electrons are shared equally.
- Oxidation number **exaggerates the ionic character of bonds**. It represents the atoms as ions, and **all the electrons in a bond are assigned to the more electronegative atom** (the atom with the greater attraction for electrons).
- For CO<sub>2</sub>: formal charge is zero. Oxidation number of carbon is +IV. 
$$\overset{0}{\underset{\cdot\cdot}{\text{O}}}=\overset{0}{\text{C}}=\overset{0}{\underset{\cdot\cdot}{\text{O}}}$$
- Formal charges **depend on the particular Lewis structure** drawn, oxidation numbers **do not**.
- Formal charges are used to assess **relative energies of Lewis structures**, oxidation numbers are used when assessing the **oxidizing and reducing properties** of molecules.

## 2B.3: Formal charge

### Reminder: How to assign oxidation numbers

#### TOOLBOX K.1 HOW TO ASSIGN OXIDATION NUMBERS

##### CONCEPTUAL BASIS

To assign an oxidation number, we imagine that each atom in a molecule, formula unit, or polyatomic ion is present in ionic form (which it might not be). The oxidation number is then taken to be the charge on each “ion.” The “anion” is usually oxygen as  $\text{O}^{2-}$  or the element farthest to the right in the periodic table (actually, the most *electronegative* element; see Section 2.12). We then assign to the other atoms charges that balance the charge on the “anions.”

##### PROCEDURE

To assign an oxidation number  $N_{\text{ox}}(\text{E})$  to an element E, we start with two simple rules:

- 1 The oxidation number of an element uncombined with other elements is 0.
- 2 The sum of the oxidation numbers of all the atoms in a species is equal to its total charge.

The oxidation numbers of elements in most of the compounds in this text are assigned by using these two rules along with the following specific values:

- The oxidation number of hydrogen is +1 in combination with nonmetals and  $-1$  in combination with metals.
- The oxidation number of elements in Groups 1 and 2 is equal to their group number.
- The oxidation number of all the halogens is  $-1$  unless the halogen is in combination with oxygen or another halogen higher in the group. The oxidation number of fluorine is  $-1$  in all its compounds.
- The oxidation number of oxygen is  $-2$  in most of its compounds. Exceptions are its compounds with fluorine (in which case, the previous statement takes precedence) and its occurrence as peroxides ( $\text{O}_2^{2-}$ ), superoxides ( $\text{O}_2^-$ ), and ozonides ( $\text{O}_3^-$ ).

This procedure is illustrated in Example K.1.

## 2B.3: Formal charge

### Summary

The formal charge gives an indication of the extent to which atoms have gained or lost electrons in the process of covalent bond formation; atom arrangements and Lewis structures with the lowest formal charges are likely to have the lowest energy.

## The skills you have mastered are the ability to

- Draw the Lewis structures of molecules.
- Write the resonance structures for a molecule.
- Use formal charge calculations to select the most likely atom arrangement.

**Summary: You have learned that a covalent bond consists of a shared electron pair and that atoms typically acquire a complete octet (or duplet for hydrogen) of electrons. The patterns of electron-pair sharing in covalent compounds are represented by drawing Lewis structures. You have seen that in some cases, it is necessary to represent a molecule as a resonance hybrid that spreads multiple-bond character over the entire molecule. You have seen that the lowest-energy Lewis structure can often be identified by calculating the formal charges of the atoms.**

# Beyond the Octet Rule

Topic 2C

Topic 2C.1 Radicals and biradicals

Topic 2C.2 Expanded valence shells

Topic 2C.3 Incomplete octets

WHY DO YOU NEED TO KNOW THIS MATERIAL?

- Octet rule is a good starting point, there are exceptions that go beyond the octet.

WHAT DO YOU NEED TO KNOW ALREADY?

- Lewis structures
- Concept of resonance
- Assign formal charges in Lewis structure

## 2C Beyond the octet rule

### Three types of exceptions to the octet rule

1. Molecules with an odd number of electrons
2. Certain elements are able to accommodate more than eight electrons in valence shell
3. Atoms in compounds may have incomplete octets

# Radicals and Biradicals

Topic 2C.1

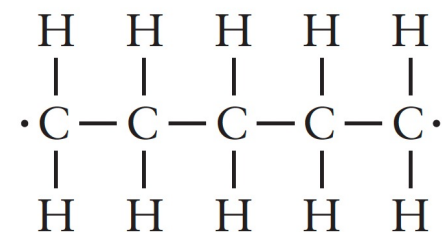
## 2C.1 Radicals and biradicals

### 1. Molecules with an odd number of electrons

- Species with at least one unpaired electron: **radicals**

E.g. methyl radical,  $\cdot\text{CH}_3$  or nitric oxide, NO (draw on **blackboard**)

- Radicals are highly reactive, cannot be stored, fleeting in nature
- Involved in ozone formation and decomposition
- Antioxidants prevent damage by radicals
- Biradical**: molecule with at least two unpaired electrons, usually on different atoms.
- Oxygen is a **biradical**:  $[\text{He}]2s^22p_x^22p_y^12p_z^1$



## 2C.1 Radicals and biradicals

### Self-test 2C.1B (blackboard)

Write the Lewis structure of nitrogen dioxide,  $\text{NO}_2$ .

## 2C.1: Radicals and biradicals

### Summary

A radical is a species with an unpaired electron; a biradical has two unpaired electrons on either the same or different atoms.

# Expanded Valence Shells

Topic 2C.2

## 2C.2: Expanded valence shells

### Hypervalent compounds

A compound consisting of molecules that contain an atom with more than eight electrons is called **hypervalent**.

**Variable covalence:** when an element is able to form compounds with different numbers of attached atoms, e.g.  $\text{PCl}_3$  vs.  $\text{PCl}_5$ .

What determines hypervalency?

- **Size of atom:** P is large enough to fit as many as six Cl atoms around it ( $\text{PCl}_5$  known), N is too small ( $\text{NCl}_5$  unknown)

## 2C.2: Expanded valence shells

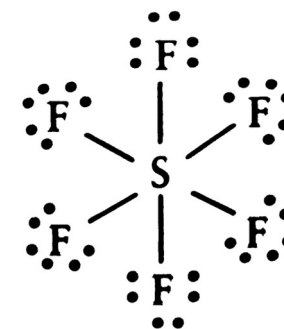
### Where are the extra electrons found?

#### Two explanations:

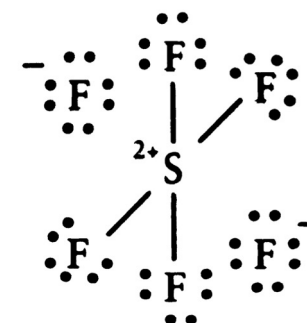
**1. Occupation of d-orbitals:** Valence shell expands by using d-orbitals as well as the s- and p-orbitals. Hypervalence is characteristic of Period 3 and beyond.

E.g. sulfur with  $[\text{Ne}]3s^23p^4$  configuration ( $\text{SF}_6$ ) has empty 3d-orbitals.

**2. Ionic-covalent resonance:** This explanation preserves the octet rule. It treats structures as **resonance hybrid of ionic structures**, e.g.  $\text{SF}_6$  is composed of  $(\text{SF}_4)^{2+}$  and  $(\text{F}^-)_2$ . The cation has an octet, as do all the fluorine atoms and fluoride ions.



Sulfur hexafluoride,  $\text{SF}_6$



Sulfur hexafluoride,  $\text{SF}_6$

## 2C.2: Expanded valence shells

**Where are the extra electrons found?**

**Which explanation is better?**

This depends on the molecule. To decide, one has to run detailed calculations as will be described in Topics 2F and 2G.

Important distinction: The first explanation makes use of d-orbitals, the second does not.

In this topic, the choice of model is **unresolved**, but we can still draw Lewis structures.

## 2C.2: Expanded valence shells

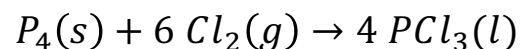
### **Example 2C.1: Writing a Lewis structure with an expanded valence shell (blackboard)**

- (a) Write the Lewis structure of sulfur tetrafluoride on the basis that the sulfur atom can expand its valence shell and give the number of electrons in that shell.
- (b) Propose an ionic-covalent resonance structure for  $\text{SF}_4$  in which the octet rule is obeyed for all atoms.

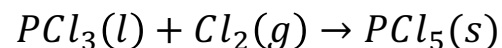
## 2C.2: Expanded valence shells

### Variable valence of phosphorous

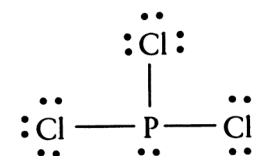
- Phosphorous has variable valence, e.g. forms toxic, colorless liquid phosphorous trichloride with a limited supply of chlorine:



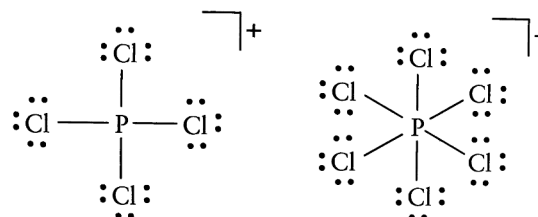
- $PCl_3$  obeys the octet rule
- When  $PCl_3$  reacts with additional chlorine:



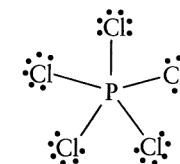
- At RT,  $PCl_5$  is an ionic solid made of  $PCl_4^+$  and  $PCl_6^-$
- At 160 °C: a gas composed of  $PCl_5$



Phosphorus trichloride,  $PCl_3$



9 Phosphorus pentachloride,  $PCl_5(s)$

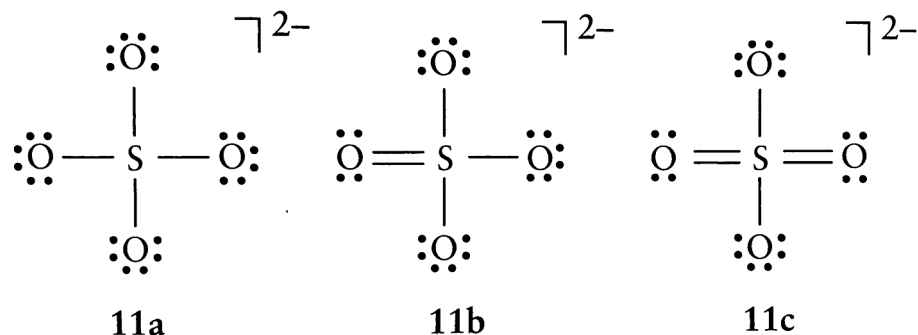


10 Phosphorus pentachloride,  $PCl_5$

## 2C.2: Expanded valence shells

### Example 2C.2: Selecting the dominant resonance structure for a molecule (blackboard)

Identify the dominant resonance structure of the sulfate ion ( $\text{SO}_4^{2-}$ ) from the three structures shown (**11a-11c**) by calculating the formal charges on the atoms in each structure.



## 2C.2: Expanded valence shells

### Summary

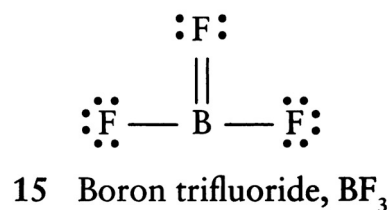
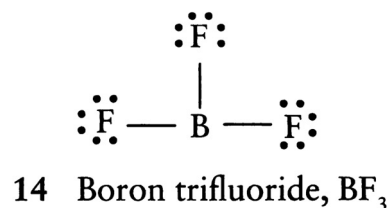
Apparent expansion of the valence shell can take place in elements of Period 3 and later. These elements can exhibit variable covalence and be hypervalent. Formal charge helps to identify the dominant resonance structure.

# Incomplete Octets

Topic 2C.3

## 2C.3: Incomplete octets

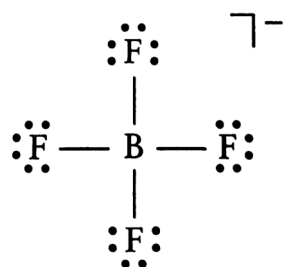
### Boron may have incomplete octets



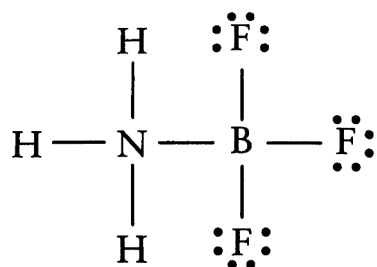
- Boron trifluoride only has six valence electrons at the boron atom (structure **14**)
- Structure **15** does not exist due to F's high ionization energy: It does not like to share its electrons.
- Experimental evidence: B-F bond length is a resonance hybrid between **14** and **15**, with single bonds making the major contribution

## 2C.3: Incomplete octets

### Coordinate covalent bonds



16 Tetrafluoroborate,  $\text{BF}_4^-$



17

- A special way for boron (and similar atoms) to complete their octets: an additional atom or ion with a lone pair of electrons might form a bond providing *both* electrons.
- A bond in which both electrons come from one of the atoms is called a **coordinate covalent bond**.

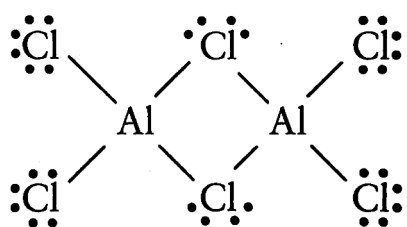
Examples:

$\text{BF}_4^-$  forms when  $\text{BF}_3$  is passed over metal fluoride (**16**)

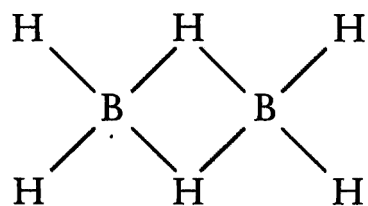
Or:  $\text{BF}_3(g) + \text{NH}_3(g) \rightarrow \text{NH}_3\text{BF}_3(s)$  (**17**)

## 2C.3: Incomplete octets

### Aluminum can also have unusual Lewis structures



18 Aluminum chloride,  $\text{Al}_2\text{Cl}_6$



19 Diborane,  $\text{B}_2\text{H}_6$

- **Formation of dimers** is another way to complete octets using coordinate covalent bonds.
- Linked **pairs of molecules**
- Examples: aluminum chloride (**18**)
- Dimer does not exist for aluminum trichloride but not boron trichloride (Al atom bigger)
- Also exists for diborane ( $\text{B}_2\text{H}_6$ ) (**19**)

## 2C.3: Incomplete octets

### Summary

Compounds of boron and aluminum may have unusual Lewis structures in which boron and aluminum have incomplete octets or atoms that act as bridges.

The skills you have mastered are the ability to

- Draw the Lewis structures of molecules and ions with either expanded or incomplete valence shells.
- Use formal charge calculations to evaluate alternative Lewis structures.

**Summary: You have learned that molecules with an unpaired electron and therefore having incomplete octets are called radicals. These compounds are normally very reactive. When drawing Lewis structures for molecules that contain Period 3 and later elements, you have seen that there are two possible explanations, one of which is to allow for the expansion of the valence shell by using d-orbitals and the other in which octets are preserved and there is ionic-covalent resonance.**