## **Structure Determines Properties**

## SUMMARY

This chapter sets the stage for all of the others by reminding us that the relationship between structure and properties is what chemistry is all about. It begins with a review of Lewis structures, moves to a discussion of the Arrhenius, Brønsted–Lowry, and Lewis pictures of acids and bases, and concludes with the effects of structure on acidity and basicity.

Section 1.1 A review of some fundamental knowledge about atoms and electrons leads to a discussion of **wave functions**, **orbitals**, and the **electron configurations** of atoms. Neutral atoms have as many electrons as the number of protons in the nucleus. These electrons occupy orbitals in order of increasing energy, with no more than two electrons in any one orbital. The most frequently encountered atomic orbitals in this text are *s* orbitals (spherically symmetrical) and *p* orbitals ("dumbbell"-shaped).



- Section 1.2 An **ionic bond** is the force of electrostatic attraction between two oppositely charged ions. Atoms at the upper right of the periodic table, especially fluorine and oxygen, tend to gain electrons to form anions. Elements toward the left of the periodic table, especially metals such as sodium, tend to lose electrons to form cations. Ionic bonds in which carbon is the cation or anion are rare.
- Section 1.3 The most common kind of bonding involving carbon is **covalent bonding.** A covalent bond is the sharing of a pair of electrons between two atoms. Lewis structures are written on the basis of the octet rule, which limits second-row elements to no more than eight electrons in their valence shells. In most of its compounds, carbon has four bonds.



Each carbon has four bonds in ethyl alcohol; oxygen and each carbon are surrounded by eight electrons.

Section 1.4 Many organic compounds have **double** or **triple bonds** to carbon. Four electrons are involved in a double bond; six in a triple bond.



Ethylene has a carbon-carbon double bond; acetylene has a carbon-carbon triple bond.

Section 1.5 When two atoms that differ in **electronegativity** are covalently bonded, the electrons in the bond are drawn toward the more electronegative element.



Section 1.6 Table 1.5 in this section sets forth the procedure to be followed in writing Lewis structures for organic molecules. It begins with experimentally determined information: the **molecular formula** and the **connectivity** (order in which the atoms are connected).

H :O: H C C C Ö H H H H The Lewis structure of acetic acid

Different compounds that have the same molecular formula are called **isomers**. If they are different because their atoms are connected in a different order, they are called **constitutional isomers**.



Formamide (*left*) and formaldoxime (*right*) are constitutional isomers; both have the same molecular formula (CH<sub>3</sub>NO), but the atoms are connected in a different order.

Section 1.7 Counting electrons and assessing charge distribution in molecules is essential to understanding how structure affects properties. A particular atom in a Lewis structure may be neutral, positively charged, or negatively charged. The **formal charge** of an atom in the Lewis structure of a molecule can be calculated by comparing its electron count with that of the neutral atom itself.

Formal charge 5 Group number in periodic table 2 Electron count

where

Electron count 5 Number of unshared electrons 1 (Number of shared electrons)

Section 1.8 Many molecules can be represented by two or more Lewis structures that differ only in the placement of electrons. In such cases the electrons are delocalized, and the real electron distribution is a hybrid of the contributing Lewis structures, each of which is called a **resonance** form. The rules for resonance are summarized in Table 1.6.



Two Lewis structures (resonance forms) of formamide; the atoms are connected in the same order, but the arrangment of the electrons is different.

Section 1.9 The shapes of molecules can often be predicted on the basis of **valence shell electron-pair repulsions.** A tetrahedral arrangement gives the maximum separation of four electron pairs (*left*); a trigonal planar arrangement is best for three electron pairs (*center*), and a linear arrangement for two electron pairs (*right*).

Section 1.10 Knowing the shape of a molecule and the polarity of its various bonds allows the presence or absence of a **molecular dipole moment** and its direction to be predicted.



Section 1.11 Curved arrows increase the amount of information provided by a chemical equation by showing the flow of electrons associated with bond making and bond breaking. In the process:

Both water and carbon dioxide have polar bonds, but water is a polar molecule and carbon dioxide is not.

an electron pair of nitrogen becomes the pair of electrons in a CON bond. The COBr bond breaks, with the pair of electrons in that bond becoming an unshared pair of bromide ion.

Section 1.12 According to the **Arrhenius** definitions, an acid ionizes in water to produce protons (H<sup>1</sup>) and a base produces hydroxide ions (HO<sup>2</sup>). The strength of an acid is given by its equilibrium constant  $K_a$  for ionization in aqueous solution:

$$K_{a} = \frac{[\mathrm{H}^{+}][:\mathrm{A}^{-}]}{[\mathrm{HA}]}$$

or more conveniently by its  $pK_a$ :

$$pK_a 5 2\log_{10}K_a$$

Section 1.13 According to the **Brønsted–Lowry** definitions, an acid is a proton donor and a base is a proton acceptor.

 $\mathbf{B} \not + \mathbf{H} \not - \mathbf{A} \iff \mathbf{B} - \mathbf{H} + \mathbf{A}^{-}$ Base Acid Conjugate Conjugate acid base

The Brønsted-Lowry approach to acids and bases is more generally useful than the Arrhenius approach.

- Section 1.14 **Basicity constants** are not necessary in the Brønsted–Lowry approach. Basicity is measured according to the  $pK_a$  of the conjugate acid. The weaker the conjugate acid, the stronger the base.
- Section 1.15 The strength of an acid depends on the atom to which the proton is bonded. The two main factors are the strength of the HOX bond and the electronegativity of X. Bond strength is more important for atoms in the same group of the periodic table, electronegativity is more important for atoms in the same row. Electronegative atoms elsewhere in the molecule can increase the acidity by **inductive effects**.

**Electron delocalization** in the conjugate base, usually expressed via resonance between Lewis structures, increases acidity.

Section 1.16 The position of equilibrium in an acid–base reaction lies to the side of the weaker acid.

This is a very useful relationship. You should practice writing equations according to the Brønsted–Lowry definitions of acids and bases and familiarize yourself with Table 1.8 which gives the  $pK_a$ 's of various Brønsted acids.

Section 1.17 The Lewis definitions of acids and bases provide for a more general view of acid-base reactions than either the Arrhenius or Brønsted-Lowry picture. A **Lewis acid** is an electron-pair acceptor. A **Lewis base** is an electron-pair donor. The Lewis approach incorporates the Brønsted-Lowry approach as a subcategory in which the atom that accepts the electron pair in the Lewis acid is a proton.