Chapter 1

Introduction

The study of organic chemistry focuses on the chemistry of elements and materials essential for the existence of life. In addition to carbon, the most common elements present in organic molecules are hydrogen, oxygen, nitrogen, sulfur, and various halogens. Through the study of organic chemistry, our understanding of the forces binding these elements to one another and how these bonds can be manipulated are explored. In general, our ability to manipulate organic molecules is influenced by several factors that include the nature of functional groups near sites of reaction, the nature of reagents utilized in reactions, and the nature of potential leaving groups. In addition, these three factors impart further variables that influence the course of organic reactions. For example, the nature of the reagents used in given reactions can influence the reaction mechanisms and ultimately the reaction products. By recognizing the interplay between these factors and by applying principles of arrow-pushing, which in reality represents bookkeeping of electrons, reasonable predictions of organic mechanisms and products can be realized without the burden of committing to memory the wealth of organic reactions studied in introductory courses. In this chapter, the concept of arrow-pushing is defined in context with various reaction types, functional groups, mechanism types, reagents/nucleophiles, and leaving groups.

1.1 DEFINITION OF ARROW-PUSHING

Organic chemistry is generally presented through a treatment of how organic chemicals are converted from starting materials to products. For example, the Wittig reaction (Scheme 1.1) is used for the conversion of aldehydes and ketones to olefins, and the Diels–Alder reaction (Scheme 1.2) is used for the formation of six-membered ring systems and treatment of alkyl halides with reagents such as tributyltin hydride (Scheme 1.3),
resulting in removal of the associated halides. However, by presenting these reactions as illustrated in Schemes 1.1, 1.2, and 1.3, no explanation is provided as to how the starting materials end up as their respective products.

By definition, the outcome of any chemical reaction is the result of a process resulting in the breaking and formation of chemical bonds. Referring to material covered in most general chemistry courses, bonds between atoms are defined by sets of two electrons. Specifically, a single bond between two atoms is made of two electrons, a double bond between atoms is made of two sets of two electrons, and a triple bond between atoms is made of three sets of two electrons. These types of bonds can generally be represented by Lewis structures using pairs of dots to illustrate the presence of an electron pair. In organic chemistry, these dots are most commonly replaced with lines. Figure 1.1 illustrates several types of chemical bonds using both electron dot notation and line notation. The list of bond types shown in Figure 1.1 is not intended to be exhaustive with respect to functional groups or potential combinations of atoms.

While chemical bonds are represented by lines connecting atoms, electron dot notation is commonly used to represent lone pairs (nonbonding pairs) of electrons. Lone pairs are found on heteroatoms (atoms other than carbon or hydrogen) that do not require bonds with additional atoms to fill their valence shell of eight electrons. For example, atomic carbon possesses four valence electrons. In order for carbon to achieve a full complement of eight valence electrons in its outer shell, it must form four chemical bonds, leaving no electrons as lone pairs. Atomic nitrogen, on the other hand, possesses five valence electrons. In order for nitrogen to achieve a full complement of eight valence electrons, it must form three chemical bonds, leaving two electrons as a lone pair. Similarly, atomic oxygen
possesses six valence electrons. In order for oxygen to achieve a full complement of eight valence electrons, it must form two chemical bonds, leaving four electrons as two sets of lone pairs. In the examples of chemical bonds shown in Figure 1.1, lone pairs are not represented in order to focus on the bonds themselves. In Figure 1.2 the missing lone pairs are added where appropriate. Lone pairs are extremely important in understanding organic mechanisms because they frequently provide the sources of electron density necessary to drive reactions as will be discussed throughout this book.

As organic reactions proceed through the breaking and subsequent formation of chemical bonds, it is now important to understand the various ways in which atomic bonds can be broken. In general, there are three ways in which this process can be initiated. As shown in Scheme 1.4, the first is simple separation of a single bond where one electron from the bond resides on one atom and the other electron resides on the other atom. This type of bond cleavage is known as homolytic cleavage because the electron density is equally shared between the separate fragments and no charged species are generated. It is this process that leads to free radical reactions.

Unlike homolytic cleavage, heterolytic cleavage (Scheme 1.5) of a chemical bond results in one species retaining both electrons from the bond and one species retaining no electrons from the bond. In general, this also results in the formation of ionic species where
the fragment retaining the electrons from the bond becomes negatively charged while the other fragment becomes positively charged. These charged species then become available to participate in ion-based transformations governed by the electronic nature of reactants or adjacent functional groups.

Having introduced homolytic cleavage and heterolytic cleavage as the first two ways in which bonds are broken at the initiation of organic reactions, attention must be drawn to the possibility that bonds can rearrange into lower energy configurations through concerted mechanisms where bonds are simultaneously broken and formed. This third process, associated with pericyclic reactions, is illustrated in Scheme 1.6 using the Cope rearrangement and does not involve free radicals or ions as intermediates. Instead, it relies on the overlap of atomic orbitals, thus allowing the transfer of electron density that drives the conversion from starting material to product. Regardless, whether reactions rely on free radicals, ions, or concerted mechanisms, all can be explained and/or predicted using the principles of arrow-pushing.

Arrow-pushing is a term used to define the process of using arrows to conceptually move electrons in order to describe the mechanistic steps involved in the transition of starting materials to products. An example of arrow-pushing is illustrated in Scheme 1.7 as applied to the Cope rearrangement introduced in Scheme 1.6. As the Cope rearrangement proceeds through a concerted mechanism, the movement of electrons is shown in a
single step. As will become apparent, **arrow‐pushing** is broadly useful to explain even very complex and multistep mechanisms. However, while **arrow‐pushing** is useful to explain and describe diverse mechanistic types, it is important to note that different types of arrows are used depending on the type of bond cleavage involved in a given reaction. Specifically, when **homolytic cleavage** is involved in the reaction mechanism, **single‐barbed arrows** are used to signify the movement of **single electrons**. Alternatively, when **heterolytic cleavage** or **concerted** steps are involved in the reaction mechanism, **double‐barbed arrows** are used to signify the movement of **electron pairs**. Schemes 1.8 and 1.9 illustrate the use of appropriate arrows applied to **homolytic cleavage** and **heterolytic cleavage**, respectively.

### 1.2 FUNCTIONAL GROUPS

Having presented the concept of **arrow‐pushing** in context of the steps that initiate chemical reactions, some factors impacting the flow of electrons leading from starting materials to products can now be explored.

As a general rule, **electrons** will flow from atomic centers **high in electron density** to atomic centers **low in electron density**. This dependence on **polarity** is similar to the way that electricity flows in an electrical circuit. If there is no difference in **electrical potential** between the ends of a wire, electricity will not flow. However, if a **charge** is applied to one end of the wire then the wire becomes **polarized** and electricity flows. If we imagine a simple **hydrocarbon** such as ethane, we can analogously relate this system to a **non-polarized wire**. Both carbon atoms possess the same density of electrons and thus ethane has no polarity. However, if functionality is added to ethane through the introduction of groups bearing **heteroatoms**, the **polarity** changes and electron flow can be used to induce chemical reactions. These heteroatom‐bearing groups are known as **functional groups** and serve to donate or withdraw electron density.

While **functional groups** can be either **electron donating** or **electron withdrawing**, these properties rely upon the specific heteroatoms the functional group is composed of and the configuration of these heteroatoms relative to one another. With respect to the specific heteroatoms, **electronegativity** of the heteroatoms is the driving force influencing **polarity**. Thus, the more **electronegative** the atom, the greater the affinity the atom has for electrons. As a calibration for **electronegativity**, the **Periodic Table of the Elements** serves as an excellent resource. Specifically, moving from left to right and from bottom to
top, electronegativity increases. For example, nitrogen is more electronegative than carbon, and oxygen is more electronegative than nitrogen. Likewise, fluorine is more electronegative than chlorine, and chlorine is more electronegative than bromine. It is important to note that the influence of electronegativity on polarity is so strong that simply replacing a carbon atom with a heteroatom is enough to impart strong changes in polarity compared to the parent structure. Figure 1.3 illustrates common organic functional groups as components of common organic molecules.

Polarity in organic molecules is generally represented as partial positive (δ+) charges and partial negative (δ−) charges. These partial charges are induced based on the presence of heteroatoms either by themselves or in groups. These heteroatoms, as described in the previous paragraph and in Figure 1.3, define the various functional groups. Returning to the example of ethane as a nonpolar parent, Figure 1.4 illustrates how polarity changes as influenced by the introduction of heteroatoms and functional groups. As shown, heteroatoms such as nitrogen, oxygen, and halogens, due to their increased electronegativities compared to carbon, adopt partial negative charges. This causes adjacent carbon atoms to take on partial positive characteristics. As illustrated in Figure 1.4, charges on carbon atoms are not limited to positive. In fact, when a carbon atom is adjacent to a positive or partial positive center, it can adopt partial negative characteristics. This ability to control the charge characteristics of carbon atoms leads to the ability to create reactive centers with a
diverse array of properties. By taking advantage of this phenomenon of *induced polarity*, we are able to employ a multitude of chemical transformations, allowing for the creation of exotic and useful substances relevant to fields ranging from material science to food science to agriculture to pharmaceuticals.

### 1.3 NUCLEOPHILES AND LEAVING GROUPS

As discussed in Section 1.2, *polarity* is the key to the ability to initiate most chemical reactions. However, this is not the only factor influencing the ability to initiate reactions. In fact, the type of reaction on a given molecule is often dependent on the nature of the *solvent* and the *reagents* used. For example, solvent *polarity* can influence the *reaction rate* and the *reaction mechanism*. Furthermore, the nature of the chemical *reagents* used can affect the *reaction mechanism* and the identity of the final product. The following definitions will be key to understanding the terminology used in the following chapters.

**Nucleophiles** are reagents that have an affinity for positively charged species or electrophiles. In organic reactions, *nucleophiles* form chemical bonds at sites of partial positive charge through donation of their electrons. This generally results in the need for the starting compound to release a *leaving group*. An example of a *nucleophilic reaction* is shown in Scheme 1.10 where Nu: represents the nucleophile and L: represents the leaving group. *Arrow-pushing* is used to illustrate the movement of the electron pairs.

**Leaving groups** are the components of chemical reactions which detach from the starting material. Referring to Scheme 1.10, the *leaving group*, L:, ends up separate from the product while the *nucleophile*, Nu:, becomes incorporated into the product. Furthermore, while an initial evaluation of the material covered in an introductory organic chemistry course may seem overwhelming, the majority of the material covered can be reduced to the principles illustrated in the single reaction shown in Scheme 1.10.
In this chapter, the basic principle of arrow‐pushing was introduced in the context of organic reactions driven by homolytic cleavage, heterolytic cleavage, or concerted mechanisms. Furthermore, the concept of polarity was introduced using heteroatoms and common organic functional groups. This discussion led to the definitions of nucleophiles and leaving groups in the context of simple nucleophilic reactions. Finally, by pulling these ideas together, the concept of approaching the study of mechanistic organic chemistry from a simplified perspective of understanding the principles of arrow‐pushing was introduced.

While characteristics such as homolytic cleavage, heterolytic cleavage, and concerted mechanisms were discussed, the principles of arrow‐pushing apply equally to all. However, with respect to heterolytic cleavage, an understanding of the properties of organic acids and bases is essential in order to understand underlying organic mechanisms. These concepts are introduced in Chapters 3 and 4.
PROBLEMS

1. Add arrow-pushing to explain the following reactions:

a. \[ \text{N}=\text{C}^\ominus + \text{H}_2\text{C}^-\text{I} \rightarrow \text{N}^\ominus \text{C}^-\text{H}_3 + \text{I}^\ominus \]

b. \[ \text{H}_3\text{C}^-\text{NH} + \text{H}_3\text{C}^-\text{OCH}_3 \rightarrow \text{H}_3\text{C}^-\text{NH}\text{OCH}_3 \]

c. \[ \text{H}_3\text{C}^-\text{NH} \text{OCH}_3 \rightarrow \text{H}_3\text{C}^-\text{NHCH}_3 + \text{O}^-\text{OCH}_3 \]
d. \[ \text{H}_3\text{C}-\text{NH}_2 + \text{H}_2\text{C}-\text{Cl} \rightarrow \text{H}_3\text{C}-\text{N}_2\text{CH}_3 + \text{Cl}^- \]

e. \[ \text{H}_3\text{C}-\text{CH}_2 + \text{H}_2\text{CH}_3 \rightarrow \text{H}_3\text{C}-\text{CH}_2\text{H}_3 \]

f. \[ \text{H}_3\text{C}-\text{CH}_3 + \text{H}^\oplus \rightarrow \text{H}_3\text{C}-\text{CH}_3 \]
g. \[ \text{H}_3\text{C}^- \cdot \text{CH}_3 \text{O}^- \cdot \text{H} \rightarrow \text{H}_3\text{C}^- \cdot \text{CH}_3 + \text{H}_3\text{C}^- \cdot \text{CH}_3^- \cdot \text{H} \]

h. \[
\begin{align*}
\text{H}_3\text{CO}^- & \cdot \text{OCH}_3 + \text{Cl}^- \\
\text{H}_3\text{C}^- \cdot \text{O}^- & \cdot \text{CH}_3 \\
\text{H}_3\text{C}^- \cdot \text{O}^- & \cdot \text{CH}_3
\end{align*}
\rightarrow
\begin{align*}
\text{H}_3\text{CO}^- & \cdot \text{OCH}_3 + \text{Cl}^- \\
\text{H}_3\text{C}^- \cdot \text{O}^- & \cdot \text{H} \\
\text{H}_3\text{C}^- \cdot \text{O}^- & \cdot \text{CH}_3
\end{align*}
\]

i. \[
\text{H}_3\text{CO}^- \cdot \text{OCH}_3 + \text{Cl}^- \rightarrow \text{H}_3\text{CO}^- \cdot \text{OCH}_3 + \text{Cl}^-
\]
j. \[ \text{Br}-\text{Br} \rightarrow \text{Br} \cdot + \text{Br} \cdot \]

k. \[ \text{Br} \cdot + \text{H}_2\text{C}=\text{CH}_2 \rightarrow \text{Br}-\text{H} \cdot \text{CH} \cdot \text{CH}_3 \]

l. \[ \begin{array}{c}
\text{Br}-\text{H} \cdot \\
\text{H} \cdot \text{H} \\
\text{H} \cdot \text{H}
\end{array}
+ \begin{array}{c}
\text{H} \cdot \text{H} \\
\text{H} \cdot \text{H} \\
\text{H} \cdot \text{H}
\end{array}
\rightarrow \begin{array}{c}
\text{Br} \cdot \\
\text{H} \cdot \\
\text{H} \cdot \text{H} \\
\text{H} \cdot \text{H} \\
\text{H} \cdot \text{H}
\end{array} \]
2. Place the partial charges on the following molecules:

a. \[ \begin{array}{c}
\text{H} & \text{H} \\
\text{H} & \text{H}
\end{array} \]

b. \[ \begin{array}{c}
\text{H}_3\text{C} & \text{H} \\
\text{H} & \text{H}
\end{array} \]
INTRODUCTION

c. 
\[
\begin{array}{c}
\text{O} \\
\text{H}_3\text{C}^-\text{CH}_3
\end{array}
\]

d. 
\[
\begin{array}{c}
\text{O} \\
\text{H}_3\text{C}^-\text{O}^-\text{CH}_3
\end{array}
\]

e. 
\[
\begin{array}{c}
\text{N}^-\text{C}^-\text{O}^-\text{CH}_3 \\
\text{H}_2
\end{array}
\]
g. \[ \text{苯} - \text{CH}_3 \]

h. \[ \text{苯} - \text{O} - \text{CH}_3 \]

i. \[ \text{苯} - \text{Cl} \]

j. \[ \text{苯} - \text{NO}_2 \]
k. \[
\begin{array}{c}
\text{O} \\
\text{\text{Ph}} \\
\text{\text{H}} \\
\text{\text{H}} \\
\text{\text{OH}}
\end{array}
\]

l. \[
\begin{array}{c}
\text{O} \\
\text{\text{Ph}} \\
\text{\text{CH}_3} \\
\text{\text{OH}}
\end{array}
\]

m. \[
\begin{array}{c}
\text{O} \\
\text{\text{Ph}} \\
\text{\text{CH}_3} \\
\text{\text{O}} \\
\text{\text{OH}}
\end{array}
\]

n. \[
\begin{array}{c}
\text{Cl} \\
\text{\text{Ph}} \\
\text{\text{OH}}
\end{array}
\]
o. \[
\begin{align*}
\text{CH}_3 & \quad \text{O} \\
\text{N} & \quad \text{O} \\
\text{C} & \quad \text{O} \\
\text{H}_2 & \quad \text{C} \\
\end{align*}
\]

p. \[
\text{CH}_3 \quad \text{O}
\]

q. \[
\text{CH}_3 \quad \text{O} \\
\text{C} & \quad \text{O} \\
\text{H}_2 & \quad \text{C} \\
\text{O} & \quad \text{O} \\
\]

r. \[
\text{Cl} & \quad \text{O} \\
\text{C} & \quad \text{O} \\
\text{H}_2 & \quad \text{C} \\
\text{O} & \quad \text{O} \\
\]