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Organic Chemistry

AN INTERMEDIATE TEXT

SECOND EDITION

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Robert V. Hoffman

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ORGANIC CHEMISTRY

SECOND EDITION

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SECOND EDITION

Robert V. Hoffman

New Mexico State University



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To Rose

CONTENTS

Preface	xiii
Preface to the First Edition	xv
1 Functional Groups and Chemical Bonding	1
Functional Groups / 1 Orbitals / 5 Bonding Schemes / 7 Antibonding Orbitals / 13 Resonance / 18 Conjugated π Systems / 21 Aromaticity / 23 Bibliography / 26 Problems / 27	
2 Oxidation States of Organic Compounds	32
Oxidation Levels / 32 Oxidation States in Alkanes / 34 Oxidation States in Alkenes / 34 Oxidation States in Common Functional Groups / 35 Oxidation Level Changes During Reactions / 35 Bibliography / 41 Problems / 41	
3 Acidity and Basicity	47
Bronsted and Lewis Acids and Bases / 47 Acid Strength / 49 Acid–Base Equilibria / 53	

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VIII CONTENTS

Amphoteric Compounds / 56 Structural Effects on Acidity / 56 Electronegativity / 58 Inductive Effects / 59 Resonance Effects / 61 Bibliography / 63 Problems / 63

4 Curved-Arrow Notation

Electron Movement / 69 Heterolytic Bond Cleavages / 70 Heterolytic Bond Formation / 71 Homolytic Bond Making and Bond Breaking / 73 Resonance Structures / 75 Depiction of Mechanism / 76 Bibliography / 82 Problems / 82

5 Mechanisms of Organic Reactions

Activation Energy / 87 Activated Complex / 88 Reaction Energetics / 89 Structure of the Activated Complex / 91 Hammond Postulate / 96 Reaction Kinetics / 99 Determining Activation Energies / 104 Isotope Effects / 105 Electronic Effects / 110 Hammett Equation / 111 Bibliography / 118 Problems / 118

6 Stereochemical and Conformational Isomerism

Stereochemical Structures / 125 Chirality / 128 Configuration of Chiral Centers / 129 Multiple Stereocenters / 132 Optical Activity / 137 Absolute Configuration / 138 Physical Properties of Enantiomers / 139 Resolution of Enantiomers / 140 Stereoselective Reactions / 144 69

86

CONTENTS ix

Formation of Enantiomers / 144 Formation of Diastereomers / 146 Stereochemistry to Deduce Mechanism / 152 Conformational Analysis / 157 Conformational Energies / 164 A Values / 166 Strain in Ring Systems / 167 Stereoelectronic Effects / 172 Bibliography / 176 Problems / 176

7 Functional Group Synthesis

Functional Group Manipulation / 183 Carboxylic Acids / 185 Esters / 188 Amides / 190 Acid Chlorides / 191 Aldehydes / 192 Ketones / 194 Imines and Imine Derivatives / 197 Alcohols / 198 Amines / 201 Alkenes / 203 Alkanes / 207 Bibliography / 208 Problems / 209

8 Carbon–Carbon Bond Formation between Carbon Nucleophiles and Carbon Electrophiles

Synthetic Strategy / 217 Nucleophilic Carbon / 218 Electrophilic Carbon / 220 Reactivity Matching / 223 Generation of Nucleophilic Carbon Reagents / 224 Generation of Electrophilic Carbon Reagents / 227 Matching Nucleophiles with Electrophiles / 227 Enolates / 228 Enolate Regioisomers / 234 Diastereoselection in Aldol Reactions / 236 Organometallic Compounds / 239 Neutral Carbon Nucleophiles / 239 C=C Formation / 242 Cyclopropanation Reactions / 244 183

X CONTENTS

Metal-Catalyzed Carbon–Carbon Bond Formation / 246 Pd(0)-Catalyzed Carbon–Carbon Bond Formation / 247 Heck Reaction / 251 Suzuki Coupling / 253 Stille Coupling / 254 Olefin Metathesis / 256 Bibliography / 261 Problems / 262

9 Carbon–Carbon Bond Formation by Free-Radical Reactions

Free-Radical Reactions / 272 Free-Radical Polymerization / 277 Nonpolymerization Reactions / 278 Free-Radical Initiation / 280 Free-Radical Cyclization / 283 Bibliography / 288 Problems / 288

10 Planning Organic Syntheses

Retrosynthetic Analysis / 292 Carbon Skeleton Synthesis / 296 Umpolung Synthons / 302 Acetylide Nucleophiles / 305 Ring Construction / 306 Robinson Annulation / 310 Diels–Alder Reaction / 312 HOMO–LUMO Interactions / 313 Stereoelectronic Factors / 316 1,3-Dipolar Cycloadditions / 319 Bibliography / 323 Problems / 324

11 Structure Determination of Organic Compounds

Structure Determination / 332 Chromatographic Purification / 333 Instrumental Methods / 335 Nuclear Magnetic Resonance / 336 Chemical Shift / 338 Spin–Spin Coupling / 344 Descriptions of Spin Systems / 350 272

292

CONTENTS **Xİ**

Second-Order Splitting / 354 Structure Identification by ¹H NMR / 355 Carbon-13 NMR / 360 Infrared Spectroscopy / 366 IR Stretching Frequencies / 367 Use of IR Spectroscopy for Structure Determination / 371 Mass Spectrometry / 377 Fragmentation Processes / 384 Bibliography / 388 Problems / 388

395

Index

PREFACE

In keeping with a mechanistic emphasis, the book was reorganized. The chapter on mechanism is now Chapter 5 instead of Chapter 10. Thus the first six chapters focus on the mechanistic and structural underpinnings of organic chemistry. Synthetic aspects of organic chemistry are then discussed from a mechanistic and structural point of view. Several new sections have been added and others expanded. An expanded discussion of resonance and aromaticity is found in Chapter 1. A section on organopalladium chemistry and olefin metathesis has been added to Chapter 8 as they relate to current methods of carbon–carbon bond formation. Chapter 9 on free-radical reactions for carbon–carbon bond formation has been revised. The discussion of Diels–Alder chemistry has been moved to Chapter 10 and expanded. A number of new problems have been added which serve to further illustrate the principles developed in each chapter. Finally, thanks to input from many people who have read this text and taught from it, the discussion has been further honed and errors corrected.

What has evolved is a greater initial emphasis of the mechanistic and structural approach to organic chemistry. The application of these principles in a discussion of modern synthetic methodology (functional group manipulation, carbon–carbon bond formation, retrosynthetic analysis) provides a new organizational framework for understanding many of the most common and most important synthetic reactions.

What has not changed is the premise that this text is meant to provide the tools students need to master the material in advanced courses or compete successfully in the workplace.

ROBERT V. HOFFMAN

PREFACE TO THE FIRST EDITION

This text was inspired by two observations. The first is that many entering graduate students took organic chemistry as sophomores but since that time have had little exposure to organic chemistry in a formal sense. Because of this time lapse in their organic preparation, they often have difficulty performing well when placed directly into mainstream graduate level organic courses. What is much more effective is to first place them in a course which will bring them back up to speed in basic organic chemistry and at the same time introduce many of the advanced topics which are crucial to understanding current advances in the field. A course well suited for this purpose is a one-semester, advanced organic course at the senior undergraduate/beginning graduate level. Most departments, including ours, have such a course in place. Textbook selection for this course is problematic, however. If one of the standard advanced texts is used, only a small part is actually covered and students are not prepared to master the complexities, whereas if an undergraduate text is used, it often fails to push the students to the next level. Consequently, there is a real need for a one-semester text which gives a review of basic principles in addition to an exposure to the ideas which are currently of great importance in organic chemistry. This text was written to fill this need.

A second observation instrumental in shaping the approach of this text was made during group discussions of the organic faculty and students. One common exercise is to present practice cumulative exam problems to the group and discuss ways in which they might be solved. It is very common for the students to analyze the question in terms of reactions and transformations and try to arrive at a solution based on the question as written. On the other hand, it is very common for the faculty to ask very simple questions first—for example, "What is the oxidation change?" "What is the pK_a of the acid and what is the base?" and "What stereochemical changes occur?" It is clear that more experienced organic chemists begin from a very basic point of view and progress to a more complex solution, whereas novice organic chemists tend to jump in at a much more difficult level. It thus appears very important to initially emphasize the basic principles on which organic chemistry depends and then progress to more specialized topics, all the while emphasizing their relationship to the basic principles. This text utilizes this organizational approach.

XVI PREFACE TO THE FIRST EDITION

The result is a textbook designed for a one-semester advanced organic chemistry course. First and foremost it is a textbook and not a reference text. There is plenty of material to fill a semester, but it is not comprehensive in its coverage. Topics were chosen to provide a basic and well-rounded discussion of ideas important in modern organic chemistry and to provide students with the necessary tools to succeed in more specialized advanced courses. It is a book to be taught from; thus instructors should take the opportunity to include special or favorite topics at appropriate points. References to alternative textbook and literature reviews of the subjects are included so that students can go to the library and get a different explanation. This is important for encouraging students to do library work as a means to independently gain insight and understanding. Finally, there are abundant problems included at the end of each chapter so that students can practice applying what they are learning. Working problems is the single most effective way to learn and organize the large amount of information that is encountered in organic chemistry, so there are a large number of practice problems available at all levels of difficulty.

The goal of this text is to provide senior undergraduate students the organic background required to move on successfully in their careers. For beginning graduate students lacking this background, it provides a succinct yet rigorous preparation for advanced organic courses.

R.V.H.

1

FUNCTIONAL GROUPS AND CHEMICAL BONDING

Functional Groups	1
Orbitals	5
Bonding Schemes	7
Antibonding Orbitals	13
Resonance	18
Conjugated π Systems	21
Aromaticity	23
Bibliography	26
Problems	27

FUNCTIONAL GROUPS

There are over 12 million known compounds of which more than 80% are organic compounds. To make sense out of the nearly 9 million organic compounds and be able to manipulate them and make new compounds, there must be some system of organization whereby organic compounds can be categorized by a particular property or group of properties. A natural method utilized by early practitioners was to group organic compounds by the reactions that they underwent. Thus there developed a whole variety of qualitative tests called classification tests which

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could be used to systematically categorize the reactivity of a compound and thus allow it to be grouped with others of similar chemical reactivity. These tests are still very useful to practicing organic chemists and collectively are known as organic qualitative analysis.

Classification tests are used to distinguish organic compounds and segregate them into different functional classes based on their chemical properties. Originally a group of compounds that showed similar chemical behavior based on the classification tests were named for a property or behavior (e.g., acids from acer meaning "sour," aromatic compounds from their odors). With the evolution of the science of chemistry and the development of more modern views of atoms and molecules, a different definition of functional classes is possible. The behavior of organic compounds is now organized into patterns that are based on recurrent groups of atoms-functional groups. The sites in molecules at which chemical reactions occur are localized at the functional groups in the molecule; the rest of the molecule is the same after the reaction as before. Thus, instead of thinking of the whole molecule in terms of its chemical reactivity, it is only necessary to recognize what functional group or groups are present in the molecule. It is then possible to predict the chemical behavior of the molecule based on the known chemistry of the functional groups that it contains.

This turns out to be a huge simplification. Since the numbers of functional groups are relatively small, it is possible to classify a very large number of individual compounds by a relatively small number of functional groups. So the first step to enlightenment in organic chemistry is to realize the key role that functional groups play in simplifying the subject, and the second step is to learn the functional groups by name, structure, and formula. While a great number of them may have already been encountered in the introductory organic course, it is helpful to review them. Table 1.1 is a list of the most common functional groups. While there are quite a few other functional groups that are not shown, those found in Table 1.1 are the most common and are present in the vast majority of organic compounds. Notice that not all functional groups contain only carbon atoms (e.g., the nitro group and the carbodiimide groups), and some functional groups differ at atoms other than carbon (compare the nitro and nitroso groups and the sulfoxide and sulfone groups). Since functional groups are reference points for predicting and understanding the reactions of individual organic molecules, it is very important to be able to recognize these functional groups (and others that might be encountered in the future). It is also useful to learn normal structural abbreviations that are used to indicate functional groups that are present in chemical structures. The abbreviations in Table 1.2 correspond to the groups that are shown in Table 1.1.

A major reason that the behavior of organic compounds can be generalized in terms of the functional groups they contain is because the bonds holding a given functional group together are the same regardless of the compound which contains that functional group. The four compounds shown below all contain the carboxylic acid functional group, which is highlighted within the boxes. Thus all

Table 1.1 Common Functional Groups		
~~~~	C=C	—C≡C—
alkane	alkene	alkyne
<u> </u>	)C=O	O ∥ ∕C∼OH
alcohol (1°, 2°, 3°)	aldehyde	carboxylic acid
C-0-C	C=0	O C OR
ether	ketone	ester
C−N	C=N H	O C_N
amine (1°, 2°, 3°)	aldimine	amide (1°, 2°, 3°)
- C-X	C=N	—C≡N
alkyl halide (1°, 2°, 3°)	ketimine	nitrile
R∽ ^{N=O}	R−N_0-	O C C Cl
nitroso	nitro compound	acid chloride
→C-SH	CS<	$-\frac{\mathbf{C}}{\mathbf{C}} -\mathbf{O} - \mathbf{S} - \mathbf{R}$
mercaptan, thiol	sulfide	sulfonate ester
disuinde	suitoxide	suirone
		RO C=C
acetal	aromatic	enol ether
	C=C=C	C=C=N
epoxide	anene	Ketenennine

carbodiimide

Table 1.1 Common Functional Groups

C=C=O N=C=N

ketene

∕N=C=O

isocyanate

Alkane	R
Alkene	$R_2C = CR_2$
Alkyne	RC≡CR
Alcohol	ROH
Aldehyde	RCHO
Carboxylic acid	RCO ₂ H
Ether	ROR
Ketone	RC(O)R
Ester	RCO ₂ R
Amine	$RNH_2$ , $R_2NH$ , $R_3N$
Aldimine	RHC=NR
Amide	RC(O)NH ₂ , RC(O)NHR, RC(O)NR ₂
Alkyl halide	RX
Ketimine	$R_2C=NR$
Nitrile	RCN
Nitroso	RNO
Nitro	RNO ₂
Acid chloride	RC(O)Cl
Mercaptan, thiol	RSH
Sulfide	RSR
Sulfonate ester	RSO ₃ R′
Disulfide	RSSR
Sulfoxide	RS(O)R
Sulfone	RSO ₂ R
Acetal	$(RO)_2CR_2$
Aromatic	Ar–X
Enol ether	$ROCH=CR_2$
Allene	$R_2C = C = CR_2$
Ketene	$R_2C=C=O$
Keteneimine	$R_2C = C = NR$
Carbodiimide	RN=C=NR
Isocyanate	RNCO

 Table 1.2
 Common Functional Group Abbreviations

four contain the bonding pattern characteristic of the –COOH functional group which is *independent of the bonds found in rest of the molecule!* 



Since most organic reactions involve the conversion of one functional group to another, it follows that most organic reactions quite simply involve bond changes involving functional groups. If one knows the bonds found in the reactant functional group and the bonds found in the product functional group, then one automatically knows what bonding changes are required to effect the desired chemical change. Thus, in addition to being able to recognize functional groups, it is also important to be able to describe the numbers and types of bonds found in functional groups.

Bonds in functional groups can first be described by Lewis structures, which are merely formalisms for denoting numbers of shared and unshared electron pairs, formal charges, and types of bonds (numbers of shared pairs, single, double, and triple). Chemistry students learn to write Lewis structures in virtually all of their early chemistry courses. How to write Lewis structures will not be reviewed here, but knowing the correct Lewis structures for molecules and functional groups in molecules is an indispensable first step in being able to describe the structure and bonding of functional groups.

The next level of insight into functional groups comes from the translation of Lewis structures into more accurate bonding descriptions based on modern bonding theories. Structural details including geometries also result from the proper description of the bonding in the functional group. The ideas of structure and bonding currently in use had their origins in the late 1920s. It is again beyond the scope of this book to trace the developments which were seminal in the development of current theories; however, early studies were all rooted in the quest to understand and be able to describe the behavior of electrons in atoms. The development of quantum mechanics and the particle–wave duality of the electron and the uncertainty principle led to mathematical descriptions of the behavior of electrons in the electric field of the nucleus. The solution of those equations resulted in a new conceptual framework for understanding chemical bonding.

#### ORBITALS

The theory suggests that the behavior of each electron in an atom can be described by a wave function  $(\psi)$ , which is a function of the space coordinates of the electron and thus has spatial characteristics. These one electron wave functions are called atomic orbitals (AOs). Atomic orbitals describe electron densities in the atom at various distances and directions from the nucleus. By choosing a low constant absolute value for the wave function, a contour surface can be constructed. The probability of finding an electron  $(\psi^2)$  is highest inside the contour surface.

Thus instead of thinking of where an electron *is*, it is more correct to think about where the electron is *likely to be*. Orbitals are thus regions of space where an electron is more likely to be found. These regions of space, which have a significant electron population (orbitals), have shape, size (distance from the nucleus), and energy. Familiar examples of s, p, and d AOs are shown in Figure 1.1. The most common elements present in organic compounds are first-row elements (C, H, N, O); therefore 1s, 2s, and 2p AOs are most commonly encountered. The concept of AOs was a breakthrough in understanding the properties of atoms.

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#### 6 FUNCTIONAL GROUPS AND CHEMICAL BONDING



Figure 1.1 Depiction of s, p, and d atomic orbitals.

In molecules, the problem of understanding the interactions of electrons with the nuclei is more complicated because there are more nuclei and more electrons that interact. Imagine, however, the situation that occurs when two nuclei approach one another. If the two atoms come close enough together, an AO of one atom which contains a single electron will occupy to some extent the same region of space as an AO of the second atom which contains a single electron. When those AOs overlap, an electron from one atom shares a region of space with the electron from the other atom. When such an event occurs, each electron is no longer influenced by just one nucleus but by two. This requires a new mathematical description of the behavior of electrons influenced by two nuclei. Again the solution to those equations defines a new region of space where there is a high probability of finding both electrons. Furthermore, only two electrons can occupy any particular region of space. This new region of space is called a molecular orbital (MO), the electrons in the MO are of lower energy than when they were in their separate AOs, and the lowered energy gives rise to a chemical bond between the atoms. This process is shown in Figure 1.2.

In other words, chemical bonds result from the overlap of singly occupied AOs to give a doubly occupied MO (called a bonding MO) in which each electron of the pair interacts with both nuclei. Because each of the electrons interacts



Figure 1.2 Cartoon version of the overlap of 1s atomic orbitals to give a new bonding molecular orbital.

with two nuclei, they are more tightly bound (i.e., they are of lower energy) than they were in the separated atoms and are more likely to be found between the two nuclei.

Because the total number of interacting orbitals is conserved, the interaction of two AOs gives rise not only to the bonding MO of lower energy but also to an MO of higher energy called an antibonding MO. This orbital is normally unfilled by electrons; however, it can play a role in chemical reactions. For now we will concentrate on bonding MOs formed by the overlap of atomic orbitals.

#### **BONDING SCHEMES**

Bond formation between atoms occurs primarily to enable each atom to achieve an inert gas electron configuration in the valence level (a valence octet for all elements except hydrogen which requires only two electrons to achieve the electronic configuration of helium). An atom can achieve an inert-gas electronic configuration by giving up electrons, accepting electrons, or sharing electrons with another atom. An ionic bond is formed when one atom gives up one or more electrons to reach an octet electronic configuration (as a positively charged ion) and a second atom accepts one or more electrons to reach an octet electronic configuration (as a negatively charged ion). For example, the reaction of a cesium atom with a fluorine atom occurs by the transfer of an electron from the cesium atom to the chlorine atom. By doing so, both cesium and chlorine have reached a valence octet electron configuration. The cesium atom has been converted to a positively charged cesium ion with the octet electronic configuration of xenon, and the chlorine has been converted to a negatively charged chloride ion with the octet electronic configuration of argon. The "bond" between cesium and chlorine is due to the electrostatic attraction of the cesium and chloride ions.

The reaction of potassium metal with *tert*-butanol gives an ionic bond between the *tert*-butoxy anion and a potassium cation by transfer of electrons from potassium to the hydroxyl functional group. Hydrogen is evolved as a by-product. By losing an electron, potassium gains the octet electronic configuration of argon, oxygen has an octet structure (three lone pairs and one pair of shared electrons), and hydrogen has the electronic configuration of helium. (Based on functional group behavior, any other alcohol is predicted to react with potassium in the same way—and they do!)

$$---- \overset{\cdots}{\underset{\cdots}{\text{oH}}} + K \cdot \longrightarrow --- \overset{\cdots}{\underset{\cdots}{\text{o:}}} \overset{\cdots}{\underset{-}{\text{c}}} K^{+} + \frac{1}{2} H - H$$

Most bonds in organic molecules, however, are covalent bonds in which electrons are shared between two atoms. Sharing electrons is a way to enable each

atom of the bonded pair to reach an octet electronic configuration without having to give up or gain an electron. Covalent bonds are formed by the overlap of singly occupied AOs to form new MOs that contain a pair of electrons. Each atom in essence gains an electron by sharing. The reaction of a chlorine atom with a fluorine atom occurs by the overlap of a singly occupied 3p orbital of chlorine with a singly occupied 2p orbital of fluorine to give a bond between the two atoms that contains two electrons. This is shown both by using Lewis structures and by using orbital pictures. The type of bond formed is called a  $\sigma$ bond because the region of greatest electron density falls on the internuclear axis.



This simple picture is adequate for many diatomic molecules with univalent atoms, but it is not sufficient to describe the bonding in most polyatomic molecules. In addition to electron sharing to reach octet electronic configurations, other considerations such as the number of bonds to an atom, the number of electron pairs that are shared between two bonded atoms, and repulsion energies that are present between electron pairs require some modification of the picture. These factors can be rationalized by the idea that valence shell atomic orbitals (2s and 2p's) can combine to form hybrid AOs. These hybrid AOs overlap with AOs of other atoms in the usual fashion to form covalent bonds. Hybrid AOs have energies, shapes, and geometries which are intermediate between the atomic orbitals from which they are formed. Hybridization of AOs is an outgrowth of bond formation that enables atoms to derive the greatest amount of bond energy from electron sharing and to allow bonded atoms to achieve octet electronic configurations.

If four single bonds and/or electron pairs originate from a single atom, then the s orbital and the three p orbitals of the valence shell combine to form four equivalent sp³ hybrid orbitals that are then used in bond formation to other atoms. Depending on the number of electrons in the valence shell of the atom, these sp³ hybrid orbitals can contain either a single unpaired electron which can be shared with another atom by overlap and bond formation or an unshared pair of electrons which is normally not involved in bond formation. Thus alkanes, which have all single bonds, have carbon atoms which are sp³ hybridized. For example, methane has four single C–H bonds originating at carbon, and these bonds are  $\sigma$  bonds produced by the overlap of four sp³ hybrid orbitals of carbon with four 1s AOs of four hydrogens to give four sp³-1s  $\sigma$  bonds from carbon to hydrogen. The geometry of the four equivalent sp³ hybrid orbitals (and hence the compound produced by overlap with these orbitals) is tetrahedral. Thus methane has four equivalent C–H  $\sigma$  bonds which point toward the corners of a regular tetrahedron and have H–C–H bond angles of 109.5°:

#### BONDING SCHEMES 9



In a similar fashion each carbon of propane is sp³ hybridized and tetrahedral since each carbon has four single bonds to other atoms originating from it. For example, the central carbon of propane has two equivalent sp³-1s C-H  $\sigma$  bonds and two equivalent sp³-sp³ C-C  $\sigma$  bonds. (Note that sp³ orbitals from one carbon can overlap with sp³ orbitals from another carbon to produce carbon-carbon bonds.) The geometry is very close to tetrahedral, but the C-C-C bond angle is slightly larger (111°) to accommodate the bigger CH₃ groups.



Other first-row elements can also be sp³ hybridized. The only requirement is that they have a combination of four single bonds and/or electron pairs originating from a single element. Ammonia, which has three N–H bonds and a lone pair on nitrogen, is thus sp³ hybridized and has three equivalent sp³–1s N–H  $\sigma$  bonds and a lone pair which occupies an sp³ hybrid orbital. The geometry is close to tetrahedral with an H–N–H bond angle of 107°. Other amines also have sp³-hybridized nitrogen and are close to a tetrahedral geometry around the nitrogen atom.

The oxygen atom in the water molecule has two bonds and two lone pairs so it too is sp³ hybridized. There are two equivalent sp³-1s O-H  $\sigma$  bonds and two lone pairs occupying sp³-hybridized orbitals. Electron–electron repulsions of the lone pairs cause greater distortions from a true tetrahedral geometry so that the H–O–H bond angle is 105°. Other singly bonded oxygen functional groups such as alcohols, ethers, and acetals have sp³-hybridized oxygens and nearly tetrahedral geometries.

Second-row elements such as silicon, phosphorus, and sulfur can also have sp³ hybridization of the valence shell orbitals, although hybridization is not necessarily required for second-row elements. When second-row elements do hybridize,

however, the 3s and 3p AOs combine to form the sp³ hybrid orbitals. Tetramethylsilane, the standard reference for nuclear magnetic resonance (NMR) spectra, has tetrahedral geometry and thus sp³ hybridization of the 3s and 3p valence shell orbitals of silicon. Dimethyl sulfone has nearly tetrahedral bond angles, indicating that the sulfur is sp³ hybridized. Although formal charges are present, the two bonds to oxygen can be thought to arise by the overlap of a filled sp³ orbital on sulfur with an unfilled sp³ orbital on oxygen. The resulting  $\sigma$  bond is called a coordinate covalent, or dative, bond because both of the shared electrons in the bond come from only one of the bonded elements. Hydrogen sulfide has an H–S–H bond angle of 92°, which indicates that sulfur is not hybridized in this compound.



When two pairs of electrons are shared between two elements, a different bonding arrangement is required to enable the atoms to reach valence octet electron configurations. Because of the Pauli exclusion principle, only one sigma bond is possible between any two atoms because only one pair of electrons can occupy the space along the internuclear axis. The second pair of electrons that is shared by the two atoms must therefore be located in space someplace other than along the internuclear axis. The second pair of shared electrons is located in a different type of covalent bond, a  $\pi$  bond, which has electron density found on either side of the internuclear axis. The  $\pi$  bonding results from the parallel overlap (or sideways overlap) of atomic p orbitals. To accommodate the need for a singly occupied atomic p orbital available for the formation of a  $\pi$  bond, hybridization of the valence AOs takes place between the s orbital and two of the three p atomic orbitals. Hybridization of one s and two p AOs produces



three equivalent sp² hybrid AOs and a p orbital remains unhybridized in order to produce a  $\pi$  bond.

This bonding scheme permits two pairs of electrons to be shared between two atoms so that each pair occupies a different region of space and does not violate the Pauli exclusion principle. Since only two p orbitals are used in the hybridization and they are orthogonal and define a plane, the sp²-hybridized carbon is planar with bond angles of 120°. The remaining p orbital, which is left unhybridized to form the  $\pi$  bond, is perpendicular to the molecular plane. Once formed, the  $\pi$  bond keeps the entire system rigid and planar, because rotation of one end of the  $\pi$ -bonded system relative to the other end requires that the  $\pi$ bond be broken.

Elements other than carbon are also  $sp^2$  hybridized if they share two electron pairs with another atom. Thus imines have  $sp^2$ -hybridized nitrogen (and carbon) to account for formation of the C–N double bond. The lone pair on nitrogen occupies an  $sp^2$  hybrid orbital. The bond angles are all 120° around both carbon and nitrogen since both are  $sp^2$  hybridized. Similar considerations hold for the oxygen atom of carbonyl groups of all kinds. The two unshared pairs of electrons on oxygen both occupy  $sp^2$  orbitals. The interorbital angle is 120°, as expected for trigonal hybridization.



The sharing of three pairs of electrons between two atoms can be accomplished by extrapolation of the above considerations. That is, since there can only be one  $\sigma$  bond connecting the atoms, then the other two pairs of shared electrons must be in two different  $\pi$  bonds, each of which is formed by the parallel overlap of a p orbital. Furthermore the  $\pi$  bonds must be mutually orthogonal so as not to violate the Pauli exclusion principle. Hybridization of one s orbital and one p orbital gives two equivalent sp hybrid AOs which are linearly opposite to one another.



The two remaining p atomic orbitals, which are mutually orthogonal, are used to produce two orthogonal  $\pi$  bonds. The geometry of triply bonded systems is thus linear about the triple bond.

Similar considerations apply to the triply bonded nitrogen found in nitriles. The sp-hybridized carbon and nitrogen atoms form an sp-sp  $\sigma$  bond and two 2p-2p  $\pi$  bonds between carbon and nitrogen. The unshared pair on nitrogen occupies an sp hybrid orbital.



Another instance where sp hybridization is required occurs in molecules with cumulated double bonds such as allenes, ketenes, and carbodiimides. The end atoms of the cumulated units are sp² hybridized because each shares two electron pairs with another element (the central carbon) and there is a  $\sigma$  and a  $\pi$  bond. The structure, however, requires that two  $\pi$  bonds originate from the central carbon—one  $\pi$  bond going toward one end of the cumulated system, the other  $\pi$  bond going toward the other end. Thus two 2p AOs are required for  $\pi$  bonding from the central carbon and sp hybridization is appropriate. Consequently the geometry is linear at the middle atom and trigonal at the end atoms. A further consequence of the orthogonal  $\pi$  bonds is that planar bonds originating at the end carbons lie in two orthogonal planes with a dihedral angle of 90°. (A dihedral angle is the angle made by two intersecting planes.)



Besides providing a theoretical framework by which the structure, geometry, and octet structure of bonded elements can be explained and understood, the concept of hybridization also predicts the ordering of stabilities and energies of bonds and the energy of lone pairs of electrons in hybrid orbitals. Because s AOs are of lower energy than p AOs, hybrid orbitals with a greater proportion of s character should be more stable and thus form stronger bonds. Unshared pairs of electrons in hybrid orbitals with greater s character should also be of lower energy (more stable). As the percentage of s character of hybrid orbitals increases in the order  $sp^3 - 25\%$  s character  $< sp^2 - 33\%$  s character < sp - 50%s character, it is found that the strength of bonds formed by overlap with those orbitals increases in a parallel fashion. For example, the bond dissociation energies of primary C-H bonds have been measured and fall in the order that is predicted by the percentage of s character of the hybrid orbitals on carbon: sp³ C-H, 105 kcal/mol; sp² C-H, 111 kcal/mol; and sp C-H, 133 kcal/mol. Electron pairs are more stable in orbitals with more s character; thus the acidities of primary C-H bonds are found to be sp³ C-H,  $pK_a = 50$ ; sp² C-H,  $pK_a = 44$ ; and sp C-H,  $pK_a = 25$ . This is due to the fact that the anions formed by proton removal give carbanions that have the negative charge in  $sp^3$ ,  $sp^2$ , and sporbitals, respectively. Because the lone pair is more stable in an orbital of greater s character, the anion formed by removal of an sp C-H proton is more stable (and hence the proton is more easily removed) than the anion formed by removal of an sp² C-H proton, which in turn is more stable (and hence the proton is more easily removed) than the anion formed by removal of an sp³ C-H proton. Other examples of the effects of greater s character in orbitals are encountered routinely.

The concept of hybridization of AOs to give new hybrid AOs involved in the bonding patterns of atoms is a useful and practical way to describe the way in which functional groups are constructed. It provides a rationale for the structure as well as the geometry and electron distribution in functional groups and molecules in which they are found. It can also be used to predict reactivity patterns of functional groups based on these considerations.

#### ANTIBONDING ORBITALS

The overlap of AOs to give a new MO in which an electron pair is shared by the interacting atoms was illustrated in Figure 1.2. The new MO, which contains the shared electron pair, is of lower energy than the AOs from which it was produced by overlap. This energy change ( $\Delta E$ ) is illustrated in Figure 1.3 (N represents the *nucleus* of some element in the bond formation process). The  $\Delta E$ is related closely to the bond energy of the bond produced. The same model holds irrespective of the type of AOs which overlap (simple AOs or hybrid AOs) or the type of bond formed ( $\sigma$  or  $\pi$ ).

While this model is easy to visualize and understand, it is actually only half of the story. When AOs interact, the number of new MOs which are produced



Figure 1.3 Energy changes that occur during the overlap of AOs to form covalent bonds.



Figure 1.4 Formation of bonding and antibonding MOs from the overlap of AOs.

from that interaction must equal the number of AOs which initially interact. Furthermore, for each MO produced which is lower in energy than the energy of the interacting AOs, there will be one produced which will be *higher* in energy by the same amount (Figure 1.4). So when two half-filled AOs interact, there will be two MOs produced, one of lower energy which will contain the electron pair and is termed the *bonding* MO. The second molecular orbital is of higher energy, is unfilled, and is termed the *antibonding* MO.

For each bond in a molecule which is described by the overlap of AOs, there will be a bonding MO which is of lower energy and when filled with an electron pair gives rise to a stable bond between elements. There will also be an antibonding MO which is of higher energy and thus unfilled. Antibonding orbitals correspond to the situation where nuclei are moved to within the bonding distance of one another but there is *no* electron sharing; in fact the electrons and nuclei actually repel one another. This electronic and nuclear repulsion is what increases the energy of the antibonding level. Because the bonding MO is filled and the antibonding MO is unfilled, the system is at a lower net energy than the individual AOs and bond formation takes place. This occurs for both  $\sigma$  and  $\pi$  bonds as shown in Figure 1.5 (the antibonding orbitals are indicated by the asterisk). Overlap of an  $sp^3$  AO on a carbon with a 1s AO on a hydrogen gives a  $\sigma$ -bonding MO that is filled with two electrons and an unfilled, higher energy, antibonding MO termed a  $\sigma^*$  MO. Likewise, overlap of two 2p AOs on carbon gives a  $\pi$  MO which contains a shared pair of e⁻ and a  $\pi^*$  MO which is of higher energy and is unfilled.

#### ANTIBONDING ORBITALS 15



**Figure 1.5** Formation of  $\sigma$  and  $\pi$  bonding orbitals and  $\sigma^*$  and  $\pi^*$  antibonding orbitals in a double bond between two atoms (N).



Thus far it would appear that antibonding orbitals are real orbitals, but they seem to be merely mathematical artifacts since they are unfilled and thus do not enter into bonding or energy considerations. For ground-state molecules this is actually true—all of the electrons are found in bonding orbitals. Why, then, should we even concern ourselves with their existence?

The answer lies in the realization that antibonding orbitals are still, in fact, orbitals. They are regions of space where one could have electrons. In ground-state molecules, electrons fill the lower energy bonding orbitals. Suppose, how-ever, you wished to take an electron out of a bonding orbital and move it to a higher level. Where would it go? Or suppose you wished to add electrons to a molecule which already had its bonding orbitals filled. Where would the

electrons go? Suppose an electron-rich reagent were to donate electrons to a molecule. Where would the electrons go?

In these examples the electrons could go into a higher energy, unfilled MO which could be a nonbonding orbital (when one is present) or an antibonding orbital (which is always present). Thus it is most common to have the electrons go into an antibonding MO. Although they are of high energy, antibonding orbitals are usually unfilled and can accept electrons from several sources if sufficient energy is available to promote electrons into the antibonding energy level. Absorption of light energy can cause an electron to be promoted from the highest occupied molecular orbital (HOMO), which is usually a bonding MO, to the lowest unoccupied molecular orbital (LUMO), which is most often an antibonding MO. For example, if an olefin which contains a carbon–carbon  $\pi$ bond is exposed to ultraviolet light of the correct frequency (and hence energy), the molecule can absorb the energy of the light by promoting a  $\pi$  electron from the bonding MO into the antibonding MO. This new electronic state is termed an excited state and is higher in energy than the initial electron-paired state called the ground state. (The electron spins can be paired in the singlet excited state or unpaired in the triplet excited state.) Excited states of molecules are high-energy states which are much more reactive than ground states and can be described in terms of the population of antibonding orbitals. Consequently, almost all photochemical reactions which occur by the reactions of excited-state species are intimately dependent on the existence of and population of antibonding orbitals.



The reduction of organic molecules by the addition of electrons can take place by chemical reagents or at the surface of electrodes. In either case electrons are added to the organic compound, thus reducing it. Now electrons cannot just go anywhere; they must go into an unfilled orbital. Thus, during a reduction, electrons are injected into the LUMO of the molecule, which is often an antibonding orbital. Population of the antibonding orbital raises the total energy of the molecule and subsequent reactions follow. The electrochemical reduction of alkyl bromides illustrates the process well. An electron is added into the  $\sigma^*$ orbital of the carbon-bromine bond, which is the LUMO of a saturated alkyl bromide. Population of the antibonding orbital raises the energy of the molecule and weakens the carbon-bromine bond, which then dissociates to give bromide ion and a carbon-centered free radical which has an unpaired electron in a hybrid AO (nonbonded energy level).

#### ANTIBONDING ORBITALS 17



Almost all dissolving metal and electrochemical reductions follow this same general sequence. An electron is donated into an unfilled orbital which is usually an antibonding MO, the energy of the molecule is raised, and chemical change ensues.

When a nucleophile attacks an electrophile, it donates a pair of electrons to the electrophile. Electron donation must take place by an overlap interaction between a filled orbital on the nucleophile which contains the electron pair to be donated and an unfilled orbital (LUMO) on the electrophile, which is usually an antibonding orbital. Population of the LUMO by electron donation raises the energy of the system leading to bonding change and new bond formation. Addition of an alkoxide to a ketone is a typical example of the process. The electron pair to be donated is in a hybrid AO and therefore is at a nonbonding energy level (*n*). Overlap with the  $\pi^*$  orbital of the carbonyl group starts to populate the  $\pi^*$  orbital. This weakens the  $\pi$  bond, and the carbon-oxygen  $\pi$ bond of the carbonyl group is broken and a new lower energy  $\sigma$  bond is formed between the oxygen of the alkoxide and the carbonyl carbon. The electrons of the  $\pi$  bond end up in a nonbonding AO on oxygen in the product. This process is shown schematically.



Nucleophilic additions and substitutions are the most widespread of all organic reactions, and all have the same general orbital requirements. An orbital containing an electron pair of the nucleophile overlaps with an antibonding orbital of the electrophile, which leads to population of the antibonding level (in most cases). This raises the energy of the system and bond and electron reorganization follows to give products. The electron pair must be able to be donated (i.e., not tightly bound or of higher energy) and the antibonding orbital be of sufficiently low energy to ensure effective overlap.

Thus it is seen that, although antibonding orbitals are not a major factor in describing the bonding of ground-state molecules, they can play a pivotal role in the reactions of molecules. Therefore it is important to keep in mind the existence of antibonding orbitals and their ability to accept electrons and control the reactivity of molecules.

#### RESONANCE

Valence shell electrons of the atoms in a molecule are either shared or unshared. The shared electrons are found in either  $\sigma$  or  $\pi$  bonds. Unshared electrons are found in AOs (usually hybrid AOs for first-row elements). Lewis structures provide a way to indicate the shared and unshared pairs of electrons in molecules. Sometimes, however, it is possible to indicate the electron distribution in molecules by more than one Lewis structure. For example, a carboxylate anion can be represented by two equivalent but different Lewis structures.



These structures are equivalent because they have the same numbers of bonds, unshared pairs of electrons, and the same charge. They are different because the negative charge is located on different oxygen atoms. Moreover the bonds from carbon to a particular oxygen are double in one structure and single in the other. When more than one correct Lewis structure can be written for a molecule, each structure is a resonance form of the molecule. The actual molecule is a resonance hybrid of the contributing resonance forms, and its properties result from a combination of the properties of the contributing resonance forms. Thus each oxygen atom carries a  $-\frac{1}{2}$  charge, and the bonds between carbon and each oxygen atom have a bond order of 1.5 and are of the same length.

A very good analogy is a mule. A mule is a hybrid of a horse and a donkey. A mule is neither a horse nor a donkey but it has properties of each. The resonance hybrid of the carboxylate anion is a resonance hybrid of the contributing resonance forms and has properties of each.

Another classic example of resonance is the benzene molecule. The localized resonance forms are termed Kekulé forms (after Friedrich August Kekulé, who first deduced the structure of benzene) and have alternating single and double bonds between carbon atoms. The actual benzene molecule is a resonance hybrid of the contributing resonance forms as the bond lengths are equal (single and double bonds have different lengths).



The bond order is between one (single) and two (double). The resonance hybrid is often pictured with a circle in the ring to indicate the delocalized electron distribution in the molecule.

Double-headed arrows are used to indicate resonance forms. It is important to note that resonance forms are not in equilibrium, just as a mule is not a horse part of the time and a donkey the rest of the time.

The presence of resonance forms means that the electrons are not localized between two nuclei but are delocalized over more than two nuclei. The result of electron delocalization is that electrons are attracted by a greater number of nuclei, which leads to a lower energy for the molecule and hence greater stability. Simply put, resonance delocalization is a stabilizing feature of molecules.

A molecule for which resonance forms can be written is more stable than any of the contributing resonance forms. Thus the carboxylate ion (a resonance hybrid) is more stable than either of the contributing resonance forms. The difference in energy between the energy of the molecule and the energy of the most stable resonance form is the resonance energy (RE) of the molecule. The resonance energy represents the stabilization of the molecule due to the delocalization of electrons.



The amount of resonance energy is related to the relative energies of the contributing resonance forms. The greatest resonance stabilization is found when the contributing resonance forms are degenerate (equal) in energy. Thus molecules such as the carboxylate ion, benzene, the allyl anion, and the allyl cation all have significant resonance stabilization because the main resonance contributors are of the same energy.





In contrast, resonance stabilization is less in an amide because the resonance forms  $A_1$  and  $A_2$  given below are very different in energy. Nevertheless, because an amide is a resonance hybrid of  $A_1$  and  $A_2$ , it is predicted that there should be some double-bond character in the bond between carbon and nitrogen. This is in fact the case since many amides show restricted rotation around the C–N bond (typical of a  $\pi$  bond). Moreover, the nitrogen atom in amides is nearly planar and not very basic, also indicating that the lone pair is delocalized.



It is also generally true that the greater the number of contributing resonance forms, the greater will be the resonance stabilization. For this reason the enolate of a  $\beta$ -diketone has much more resonance stabilization than the enolate of a simple ketone (three resonance forms versus two). The electrons are delocalized over five atoms in the former versus three atoms in the latter. In addition, the electron density on the carbon atom is less in the diketone enolate than in a simple methyl ketone enolate.



Resonance has a significant influence on the electron distributions and energies of molecules. The delocalization of electrons is described by the contributions of resonance forms, which are themselves localized structures with discrete bonds. Such structures are known as valence bond (VB) structures, and this approach to the description of bonding in molecules is called the valence bond approach. As long as one keeps in mind that resonance forms are limiting VB structures and that the actual molecule is a resonance hybrid of these VB structures, a great deal of insight into the structure and properties of molecules can be gained.

#### CONJUGATED $\pi$ SYSTEMS

Another way to describe delocalized bonding uses the MO approach. The same principles of overlap of AOs can be applied to systems where more than two p AOs overlap to form  $\pi$  systems. First, the number of MOs produced by the overlap will be the same as the number of atomic p orbitals which interact. Thus for the allyl system where three contiguous p orbitals interact, there will be three MOs produced from the interaction of three 2p AOs. For the butadienyl system where there are four contiguous p orbitals interacting, four MOs will result, and so on.



Second, the energy distribution of the MOs will be disposed symmetrically about the energy of the AOs before they interact (nonbonded energy level). This means that the total energy of the bonding MOs is offset by the total energy of the antibonding MOs. For example, if one MO is of lower energy by  $-\Delta E$  due to overlap, then there must be an antibonding MO raised to higher energy ( $+\Delta E$ ). Molecular orbitals which are lower in energy than the nonbonding energy are bonding MOs, ( $-\Delta E$ ), those which are higher in energy than the nonbonding energy are antibonding MOs ( $+\Delta E$ ), and those at the same energy as the nonbonding energy are nonbonding MOs ( $\Delta E = 0$ ).

For the allyl system which has 3 MOs from the overlap of three 2p AOs, one MO will be lowered in energy  $(-\Delta E)$  and so one MO will be raised by the same amount. The remaining MO must stay at the nonbonding level  $(\Delta E = 0)$  to maintain energy symmetry around the nonbonding level.

$$2p - - - - - - - - \Delta E \begin{cases} - - \pi_3 \text{ (antibonding MO)} \\ - \Delta E \begin{cases} - - \pi_2 \text{ (nonbonding MO)} \\ - - \Delta E \end{cases}$$

What is interesting is that this overlap model allows the orbital diagram to be constructed without concerning itself with electrons. The MOs produced by the

interaction of AOs can each hold two paired electrons, and these can be filled in depending on the number of electrons present in the  $\pi$  system. Thus the bonding diagrams for the allyl cation, allyl radical, and allyl anion can be constructed by merely filling the orbitals with the number of  $\pi$  electrons present in these species (two, three, and four  $\pi$  electrons, respectively). This orbital picture also demonstrates that all three intermediates in the allyl system are stabilized because each contains two electrons in the  $\pi_1$ -bonding MO and any remaining electrons are in the nonbonding orbital.



Two of the four MOs of the butadienyl system are at lower energy than the nonbonded energy level  $(-\Delta E_1, -\Delta E_2)$ , and two are at higher energy than the nonbonded energy level  $(+\Delta E_1, +\Delta E_2)$ . The four  $\pi$  electrons of butadiene fill the two bonding MOs and give a stable molecule. It should also be obvious that butadienyl species with less than or more than four  $\pi$  electrons should be significantly less stable than butadiene itself. Removal of an electron requires energy because the electron would have to come from a relatively stable bonding MO. Addition of an electron to the butadienyl  $\pi$  system requires that it be put into an antibonding MO which is also energetically unfavorable.



A great many  $\pi$  systems have been examined by this approach and the orbital diagrams understood. As seen before, the antibonding orbitals are often unfilled in the ground state but play an important part in the excited states and reactions of these compounds.

#### AROMATICITY 23

#### AROMATICITY

A special type of orbital interaction occurs when a conjugated  $\pi$  system is in a ring. The  $\pi$  system of benzene is a classic example of this behavior. In benzene, the carbons of the six-membered ring are sp² hybridized, and so each has a singly filled 2p orbital to interact with the others of the conjugated system. The six 2p orbitals interact, giving rise to six new MOs. Three are bonding MOs and three are antibonding MOs. Because of the symmetry properties of the six-membered ring, the six MOs are distributed energetically, as shown below. The six available  $\pi$  electrons completely fill the bonding levels, leading to an enhanced stability of the  $\pi$  system, which is termed aromatic stabilization or aromaticity.



This "extra" stability of benzene and other aromatic compounds was a wellknown phenomenon. In fact, aromaticity was first described as chemical stability (unreactivity) toward reagents that normally attack double bonds and  $\pi$  systems. Moreover, reagents which did attack the aromatic ring gave substitution products in which the aromatic ring was retained; the same reagents usually give addition products with typical double bonds and conjugated  $\pi$  systems.

Since stability refers to energy level, aromaticity was later defined as the energy difference between an aromatic  $\pi$  system and a model  $\pi$  system in which there is no aromatic stabilization. The aromatic stabilization of benzene was taken as the difference between the heat of hydrogenation of benzene ( $\Delta H_{hyd} = -49.8 \text{ kcal/mol}$ ) and the heat of hydrogenation of the hypothetical molecule cyclohexatriene ( $\Delta H_{hyd} = -85.8 \text{ kcal/mol}$ ), which has three noninteracting double bonds in a six-membered ring. The heat of hydrogenation of cyclohexatriene was estimated as being three times the heat of hydrogenation of cyclohexene. Since both give cyclohexane upon hydrogenation, a difference in the heats of hydrogenation must be due to a difference in the energies of the starting materials. This difference amounts to 36 kcal/mol (it is termed the RE of benzene), and it corresponds to the extra stability of benzene due to aromatic stabilization. The same approach can be used to estimate the resonance energy of other aromatic molecules.

A physical distinction between benzene and the hypothetical model compound is that benzene has equal bond lengths and bond angles and is planar, whereas the hypothetical model would have localized bonds and unequal bond lengths (double
## 24 FUNCTIONAL GROUPS AND CHEMICAL BONDING



bonds are shorter than single bonds). Thus the resonance energy determination is only as good as the model system that is used.

Aromaticity was found to be a general property of many (but not all) cyclic, conjugated  $\pi$  systems. Moreover, it was found that aromaticity in molecules can be predicted by Huckel's rule. The structural requirements implicit in Huckel's rule are that there be 4n + 2 (*n* is an integer)  $\pi$  electrons in a cyclic, conjugated  $\pi$  system. Obviously benzene, which has six  $\pi$  electrons (4n + 2, n = 1) in a conjugated  $\pi$  system, is aromatic. However, Huckel's rule predicts that molecules such as cyclodecapentaene 4n + 2 = 10 (n = 2) and [18]-annulene 4n + 2 = 18 (n = 4) should be aromatic, have equal bond lengths, and be planar—and they are.

A further manifestation of aromaticity is the presence of ring current in aromatic molecules. When aromatic compounds are placed in the magnetic field of an NMR instrument, a ring current is induced in the  $\pi$  system. The ring current results in an induced magnetic field which causes the protons attached to the aromatic ring to absorb nearly 2 ppm downfield from simple olefinic protons. Aromatic character can thus be detected by a downfield shift of protons attached to the aromatic ring.

In contrast to aromatic molecules which have  $4n + 2\pi$  electrons, cyclobutadiene and cyclooctatetraene do not have  $4n + 2\pi$  electrons and are not aromatic. In fact, these molecules, which contain  $4n\pi$  electrons (*n* is an integer), are *less* stable than the planar model compounds and are termed antiaromatic. Both of these molecules adopt shapes that *minimize* interactions of the  $\pi$  orbitals.



Cyclobutadiene is an antiaromatic 4n = 4 (n = 1) system, and it is quite unstable and can only be observed at very low temperatures. Although it must be planar

(accounting for its instability), it distorts to a rectangular geometry with unequal bond lengths to minimize  $\pi$ -bond interactions. Planar cyclooctatetraene would be an antiaromatic 4n = 8 (n = 2) system, and thus it adopts a boat shape so that the  $\pi$  bonds are orthogonal and cannot interact!

Huckel's rule is more than an operational way to identify aromatic molecules. Its origins are in MO theory and its applicability is general, regardless of ring size or charge. In terms of Huckel's rule, the requirement for aromatic stabilization is that there is a cyclic system with all atoms having a p orbital available for interaction. The array of MOs produced from this interaction is populated by the total number of electrons that are present in the interacting p orbitals. If that number of electrons is 4n + 2, then the molecule will have aromatic stabilization. It turns out that the above requirements lead to a situation where the bonding MOs are completely filled, the nonbonding orbitals are either completely filled or completely empty, and antibonding levels are unfilled.

As seen above in the MO description of benzene, there are three bonding MOs that are filled by the six electrons of the  $\pi$  system. In another example, the tropylium ion is known to be aromatic. The interaction of seven 2p orbitals leads to an MO array with three bonding MOs and four antibonding MOs. The six electrons fill the bonding MOs and give an aromatic system, irrespective of the fact that, to do so, one of the seven interacting p orbitals must be unfilled, leading to a net positive charge on the delocalized aromatic ion.



It is also clear why cyclooctatetraene is not aromatic. The interaction of eight contiguous 2p AOs in a planar ring gives rise to an MO array which has three occupied bonding MOs and two nonbonding MOs which are degenerate and thus singly occupied. Since this is an unstable bonding situation, the molecule distorts to the shape of a boat so that interactions are avoided and four isolated  $\pi$  bonds can form. It is clear that either by removing two electrons ( $8 \rightarrow 6 \pi$  electrons) or by adding two electrons ( $8 \rightarrow 10 \pi$  electrons), one could reach an aromatic system. It turns out that cyclooctatetraene is easily reduced by the addition of two electrons which fill the nonbonding MOs and give a planar, aromatic dianion.



#### 26 FUNCTIONAL GROUPS AND CHEMICAL BONDING

Examples of simple aromatic molecules and ions which have been studied are shown below.



Other elements can also participate in the formation of aromatic species. Furan, pyrrole, and thiophene are all aromatic molecules. This is due to the fact that if the heteroatom is sp² hybridized, then a doubly occupied p orbital interacts with the carbon 2p orbitals to give an MO array which contains six  $\pi$  electrons and is aromatic. Note that in the development of the MO diagram for these systems the identity of the heteroatom is not important. It is only important in determining the magnitude of the aromatic stabilization.



The added stability of an aromatic system is a significant energetic feature of molecules. Reactions which occur with the formation of an aromatic system are generally facile, while reactions in which an aromatic system is disrupted are generally very difficult. Thus aromaticity can dramatically influence the reactivity of compounds and should be kept in mind.

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# PROBLEMS

**1.1.** Excluding alkyl groups, name and point out the functional groups in the following molecules:



# 28 FUNCTIONAL GROUPS AND CHEMICAL BONDING



- **1.2.** Give the bonding scheme (orbitals, etc.) and geometry for the following functional groups:
  - (a) alkyl nitrile (use R for alkyl group)
  - (b) alkyl azide (use R for alkyl group)
  - (c) nitro alkane (use R for alkyl group)
  - (d) *N*-methyl pyrrole (it is aromatic)
- **1.3.** For the following compounds, give the approximate bond angles around the atoms indicated by an arrow:



**1.4.** For the following compounds add all lone pairs of electrons to the structures and then specify the type of orbital in which they are located:

CH ₃ OCH ₃	CH ₃ CHO	$CH_3O^-$	$CH_3C \equiv O^+$
CH ₃ CH=NCH ₃	CH ₃ CH ₂ NH ₂	$CH_2 = N^-$	CH ₃ CN
CH ₃ F	CH ₃ Cl	CH ₃ Br	
CH ₃ CNO	CH ₃ NC	CH ₃ SCN	CH ₃ NCS

- **1.5.** On the basis of electronic structure and orbital energies, supply predictions for the following and explain your answer:
  - (a) Which will be more nucleophilic towards methyl iodide?



(b) Which will be more basic?

(c) Which anion will be more stable?

$$CH_3CH = CH \stackrel{\bigcirc}{\longrightarrow} or CH_3C \equiv C \stackrel{\bigcirc}{\longrightarrow}$$

**1.6.** Which of the following compounds or ions are aromatic? Draw orbital diagrams to demonstrate why.



**1.7.** Consider the tropanyl anion **T** and the cyclopentadienyl anion **C**. Which one is more stable and why? Predict the structure of each based on your analysis.

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**30** FUNCTIONAL GROUPS AND CHEMICAL BONDING



**1.8.** Compound **K** is found almost entirely in its enol form **E**. Why?



**1.9.** Explain why compound **B** can be considered a doubly aromatic molecule.



**1.10.** It is found that bromodiazirine **A** undergoes loss of bromide to produce a cation much more easily than bromocyclopropane **C**. Can you think of a reason why?



**1.11.** Compounds  $L_1$  and  $L_2$  are both lactones. Can you think of a reason why it is much more difficult to remove the  $\alpha$  proton of  $L_1$  than to remove the  $\alpha$  proton of  $L_2$ ?



**1.12.** Explain why squaric acid ionizes completely (dissociates two protons) in water and is nearly as strong as sulfuric acid.



**1.13.** Explain why guanidine is one of the strongest noncharged organic bases known.



**1.14.** Using resonance arguments, explain why cyclopentadiene is more acidic than indene.



**1.15.** It has been found by NMR measurements that the  $\alpha$ -methylene groups (*) of *N*-acetylpyrrolidine **1** are not equivalent whereas the  $\alpha$ -methylene groups in *N*-(2-propenyl)-pyrrolidine **2** are equivalent. Provide an explanation based on resonance.



# 2

# OXIDATION STATES OF ORGANIC COMPOUNDS

Oxidation Levels	32
Oxidation States in Alkanes	34
Oxidation States in Alkenes	34
Oxidation States in Common Functional Groups	35
Oxidation Level Changes During Reactions	35
Bibliography	41
Problems	41

# **OXIDATION LEVELS**

Besides bonding patterns, functional groups also vary with respect to the oxidation states of carbon in those functional groups. Thus another way to classify functional groups is by the carbon oxidation level. Correspondingly, organic reactions can be categorized as to whether an oxidation, a reduction, or no change in oxidation level has occurred in the organic reactants in going from reactants to products. This is a very useful distinction since the reagents used in a given transformation must be compatible with the oxidation change that occurs in the reaction. It is important to remember this fundamental truth—that no oxidation can occur without a corresponding reduction and no reduction can occur without a corresponding oxidation. As a consequence, if a transformation of an organic compound involves a change in its oxidation level, then the reagents necessary to cause that change must be able to undergo the complementary change in

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oxidation level. To have an oxidation, an oxidizing agent is required which itself gets reduced in the process. Similarly, to carry out the reduction of an organic compound, a reducing agent is needed which itself gets oxidized. Reagents can thus be categorized on the basis of their oxidizing or reducing properties. If no change in oxidation state occurs during a chemical reaction, then reagents used to effect the transformation should themselves undergo no oxidation or reduction. Moreover, if a reagent is not normally an oxidizing agent, then it is not easily reduced and cannot be used to oxidize something else. Conversely, if a reagent is not normally a reducing agent, then it is not easily oxidized and cannot be used to reduce something else.

Oxidation is defined as the loss of electrons. This concept is very straightforward when dealing with metal ions. Thus the change  $Mg^0 \rightarrow Mg^{2+}$  is an oxidation because magnesium has lost two electrons in going from the element to the positive ion. Similarly, oxidation of  $Cu^{1+} \rightarrow Cu^{2+}$  involves loss of an electron from  $Cu^{1+}$  to give the  $Cu^{2+}$  species.

Reduction is defined as the gain of electrons. The conversion of  $Ag^{1+} \rightarrow Ag^{0}$  involves a gain of an electron by the silver ion and thus silver is reduced. Likewise, the permanganate ion  $MnO_4^-$  has Mn[VII] but  $MnO_2$  manganese dioxide has Mn[IV]. Thus the gain of three electrons by manganese causes a reduction in oxidation level of from +7 to +4.

Because organic compounds have an overwhelming preponderance of covalent bonds, changes in oxidation state of carbon are not as easily determined by inspection as they are for metal ions. While the definitions of oxidation and reduction for organic compounds are the same as for metal ions (i.e., gain or loss of electrons), the oxidation state of a carbon atom is determined by the types of covalent bonds originating from it. A set of rules has been developed to assign numerical values for the contributions of atoms covalently bonded to carbon to the oxidation state of that carbon. Summation of the contributions of its covalently bonded substituents gives the oxidation state of a particular carbon in a molecule. Furthermore the oxidation levels of various carbons can be compared just as +2 or +3 oxidation states in metal ions can be compared.

These rules are simple and are summarized as follows:

- 1. Bonds to hydrogen or other elements more electropositive than carbon contribute -1 to the oxidation level.
- 2. Bonds to other carbon atoms contribute 0 to the oxidation level.
- 3. Bonds to oxygen or other elements more electronegative than carbon contribute +1 to the oxidation level.
- 4. Multiple bonds to an element count as multiple single bonds to that element. That is, the carbon-oxygen double bond of carbonyl group (C=O) is oxidatively equivalent to a carbon atom with two single bonds to oxygen originating from it (-O-C-O-).
- 5. A pair of electrons on carbon contribute -1 to the oxidation level.
- 6. A positive charge on carbon contributes +1 to the oxidation level.

#### 34 OXIDATION STATES OF ORGANIC COMPOUNDS

Given these contributions, the oxidation level of a given carbon can be determined by adding together the contributions of the four attached bonds.

### **OXIDATION STATES IN ALKANES**

Considering the various alkanes shown below, one sees that alkanes can have several different oxidation levels for carbon. Oxidation levels can range from -4 for methane and -3 for the carbon atom of methyl groups all the way to 0 for the quaternary carbon of neopentane. In spite of the several oxidation levels possible in alkanes, the functional group approach tells us that all are saturated alkanes and thus have the same functional equivalency and similar reactivity patterns.



This conclusion is made from the fact that all the carbons in alkanes have four single bonds originating from them and those  $\sigma$  bonds go to either carbon or hydrogen. Thus one cannot automatically assign a molecule to a functional class based solely on a certain oxidation level of the carbons that it contains.

# **OXIDATION STATES IN ALKENES**

For alkenes, several carbon oxidation levels are again possible. Furthermore, *both* carbon atoms must be considered as part of the same alkene functional group. While the total oxidation level can go from -4 for ethylene (as the sum of the oxidation level of both carbon atoms in the functional group) to 0 for a tetrasubstituted alkene, we again recognize that all are of the same functional class.



Furthermore it is evident that because the lowest possible oxidation level of a single carbon atom in an alkene is -2 while the lowest possible oxidation level of a carbon atom in an alkane is -4, alkenes are thus oxidized relative to alkanes.

Alkynes	Alcohols	Aldehydes and Ketones	Acids and Derivatives
H−C≡C−H -1 -1	CH ₃ CH ₂ - OH -1	O H C H	$\begin{array}{c} O \\ \parallel \\ C \\ +2 \end{array} OH (OR, NH_2, etc.) \end{array}$
$\begin{array}{c} R-C \equiv C-H \\ 0 & -1 \end{array}$	$(CH_3)_2CH - OH_0$	$\mathbf{R} \overset{\mathbf{O}}{\overset{\mathbf{I}}{\overset{\mathbf{C}}{\overset{\mathbf{C}}{\overset{\mathbf{C}}{\overset{\mathbf{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\overset{\mathcal{H}}{\ove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II_
$\begin{array}{c} R-C \equiv C-R \\ 0 & 0 \end{array}$	(CH ₃ ) ₃ C – OH +1	$\mathbf{R} \stackrel{\mathbf{O}}{\overset{\mathbf{I}}{\underset{+2}{}}} \mathbf{R}$	$R_{+3}^{C}$ OH (OR, NH ₂ , etc.)

# OXIDATION STATES IN COMMON FUNCTIONAL GROUPS

The same process can be carried out to determine the oxidation levels of carbon atoms in several common functional types. It is clear that by using these procedures we can assign oxidation levels to carbon atoms in a wide variety of compounds. It is also clear that knowing the oxidation level is insufficient to assign the functional group present. For example, the alkane neopentane, the alkene isobutylene, the alkyne propyne, the alcohol isopropanol, and formaldehyde all have a carbon with an oxidation level of 0 yet all belong to completely different functional classes and have different physical and chemical characteristics.

Thus the oxidation level of a given carbon is dependent only on the groups which are attached to it, not on the functional group to which it belongs.

# **OXIDATION LEVEL CHANGES DURING REACTIONS**

Comparing the oxidation levels of various carbon atoms is excellent for illustrating what oxidation state change must occur at a particular carbon in a given reaction of that compound. For example,



#### 36 OXIDATION STATES OF ORGANIC COMPOUNDS

The oxidation level of the primary alcohol (-1) is less than the aldehyde product (+1); thus this conversion requires an oxidation of the alcohol function to the aldehyde. Any reagent capable of effecting this change must necessarily be an oxidizing agent which is itself reduced. In the preceding reaction the need for an oxidant is noted above the arrow by the [O]. The reverse process, conversion of an aldehyde to a 1° alcohol, is a reduction. Any reagent capable of effecting this change must be a reducing agent and is itself oxidized. A reduction is commonly indicated by a bracketed [H] above the arrow.

By the same analysis conversion of an aldehyde to an acetal involves neither oxidation nor reduction. As a consequence no oxidant or reductant is necessary to carry out this reaction.



Similarly, for the process shown below, the oxidation state of C-1 remains at -1 throughout the sequence; thus the overall sequence involves no change in oxidation level at C-1, nor does either step. Modifications of substituents and substitution of one electronegative group for another are generally not redox processes.

$$CH_3CH_2OH \xrightarrow{TsCl} CH_3CH_2OTs \xrightarrow{NaI} CH_3CH_2I$$

One often must consider the balanced reaction in order to be certain of any net changes in oxidation state, and similar procedures for determining the oxidation level can be followed for other covalently bound elements. For example, the conversion of methane into ethane is an oxidation of the carbon atoms since the carbons in methane are at the -4 level while in ethane they are at the -3 oxidation state.

$$2 \text{ CH}_4 \longrightarrow \text{ CH}_3\text{CH}_3 + \text{H}-\text{H}$$
  
 $2 \times -4 = -8 \qquad -3 -3 + -1 -1 = -8$ 

However, the oxidation level of hydrogen changes from being bound to carbon (0) in the reactant to being bound to another hydrogen (-1) in the product. Thus hydrogen is formally reduced. The sum of oxidation levels in the reactants (-8) is the same as that in the products (-8), and the overall process is neither an oxidation nor reduction. This transformation can be thought of as an internal redox process since part of the reactant (carbon) is oxidized and part (hydrogen) is reduced. Generally, such internal redox processes require only a catalyst, not an oxidant or reductant.

On the other hand, if the by-product of the conversion of methane to ethane is  $H^+$ , then the balanced reaction is written as shown below and a net oxidation is required. An oxidizing agent is thus needed to effect this process. Again the recognition that the organic reactant (methane) and product (ethane) are both alkanes is not sufficient to determine that an oxidant is necessary.

 $2 \text{ CH}_4 \longrightarrow \text{ CH}_3 \text{ CH}_3 + 2 \text{ H}^+$  $2 \times -4 = -8 \qquad -3 -3 + (2 \times +1) = -4$ 

The Grignard reaction is often one of the first reactions encountered for the preparation of organometallic compounds. As such it provides a method for the conversion of an alkyl bromide to an alkane. From the example shown below it is seen that the overall oxidation level change from the organic reactants to the products is from 0 to -2, so a reduction has occurred. Magnesium is the reductant and is itself oxidized from 0 to +2 oxidation state. The actual reduction takes place in the first step of the process in which the C–Br bond is converted to a C–Mg–Br bond. The reaction with water is merely a hydrolysis that does not change the oxidation state of carbon.

$$(CH_3)_2CH - Br \xrightarrow[ether]{Mg} (CH_3)_2CH - Mg - Br \xrightarrow[H_2O]{H_2O} (CH_3)_2CH_2 + HOMgBr -2 -2$$

Reactions of olefins and acetylenes illustrate that the overall change in oxidation level of an organic functional group must be considered when deciding if a net oxidation level change has occurred in a chemical reaction. For example, addition of hydrogen across an acetylene gives a net reduction of each carbon and thus is a reductive process with respect to the alkyne. The same is true for the hydrogenation of an alkene. From the point of view of the alkyne and the alkene, the hydrogen can be considered a reducing agent since it undergoes oxidation during the process.

$$R-C \equiv C-R \xrightarrow{H_{2}} R \xrightarrow{R} C = C \xrightarrow{R} \xrightarrow{H_{2}} R \xrightarrow{H_{2}} R \xrightarrow{H_{2}} R \xrightarrow{H_{2}} R \xrightarrow{H_{2}} R \xrightarrow{H_{1}} R \xrightarrow{H_{$$

The conversion of an alkyne to a *trans*-alkene can be accomplished by heating with lithium aluminum hydride (LAH), by reaction with lithium in liquid ammonia (Li, NH₃). Thus all of these reagents (H₂/P-2 Ni, LAH, and Li, NH₃) are reducing agents for alkynes and give alkenes as the reduced products.

In general, any reaction which results in the addition of two hydrogen atoms across a  $\pi$  bond of any type are reductions. Conversions of aldehydes and ketones

## 38 OXIDATION STATES OF ORGANIC COMPOUNDS

to alcohols are reductions; thus any reagents which are capable of effecting that conversion must function as reducing agents. Thus NaBH₄, LAH, and a large variety of other reagents *reduce* aldehydes and ketones to alcohols by the net addition of hydrogen across the C–O  $\pi$  bond. By the same logic, conversion of a primary alcohol to an aldehyde (the reverse process) must be an oxidation, and reagents which are capable of effecting this conversion, such as dimethyl sulfoxide (DMSO) and acetic anhydride (Swern oxidation) or pyridinium chlorochromate (PCC), are oxidants. Similar considerations hold for other  $\pi$ -bonded functional groups, including acid derivatives and nitriles.

Alkenes also undergo a variety of other addition reactions in which a reagent is added across the double bond. Hydration and hydrohalogenation are classic examples.

$$\begin{array}{c} H \\ R \\ R \\ -1 \\ -1 \end{array} \stackrel{H^+, H_2O}{\xrightarrow{(\text{or } HX)}} R \xrightarrow{\text{HO}(X) H} \\ R \\ 0 \\ H \\ H \\ -2 \end{array}$$

Consideration of the oxidation level reveals that while one carbon is reduced (the one to which hydrogen adds), the other is oxidized (the one to which the oxygen adds). There is no net change in oxidation level of the alkene functional group. Likewise the reverse processes of these addition reactions, namely, elimination of HX from alkyl halides and dehydration of alcohols to give alkenes, are not redox processes. Additions of water to alkynes is analogous. In this case, however, the product is a ketone, the oxidation level of the ketone is seen to be the same as the alkyne, and so no net change in oxidation level has occurred.

$$\begin{array}{c} R - C \equiv C - R \xrightarrow{H^+, H_2O} & H & O \\ I & I & I \\ 0 & 0 & H_{gSO_4} \end{array} \rightarrow \begin{array}{c} R - C - C - R \\ I & I \\ H \\ -2 & +2 \end{array}$$

The conversion of alkenes to 1,2-diols by osmium tetroxide is also an olefin addition reaction. In this case a hydroxy group is added to each carbon of the olefin group, and the addition is termed an oxidative addition since the diol product is at a higher oxidation level than the alkene reactant. Oxidation of the carbon atoms of the alkene takes place in the first step, which is the reaction with  $OsO_4$  to produce the intermediate osmate ester.



Zinc serves to further reduce osmium and free the diol product. Similar oxidative additions to alkenes occur with bromine, chlorine, IN₃, peracids, and many other electrophiles.



Peracids such as *m*-chloroperbenzoic acid (MCPBA) clearly illustrate the redox nature of oxidative addition. In this reaction the olefin is oxidized and the MCPBA is reduced to *meta*-chlorobenzoic acid, which precipitates slowly from solution.



Another common reaction process is one in which one atom or group replaces another atom or group. These are known as substitution reactions. When one electronegative group is substituted for another, no change in oxidation level occurs; thus the reagents which carry out such substitutions are neither oxidants nor reductants.

$$\xrightarrow{-1}_{OH} \xrightarrow{HBr} \xrightarrow{-1}_{Br} \xrightarrow{CH_3O^-} \xrightarrow{-1}_{OCH_3O^+}$$

Such substitutions in saturated compounds can be carried out by a variety of strategies involving different nucleophiles and leaving groups, but the oxidation states remain the same. Acyl substitutions are analogous. For this reason carboxylic acid derivatives are treated as a common family of compounds. All have the same oxidation level and all can be converted from one to another by substitution reactions not requiring oxidation or reduction.



## 40 OXIDATION STATES OF ORGANIC COMPOUNDS

Many useful functional group transformations occur in more than one step, and it is not uncommon to find that different redox processes can be found in different steps of the process. However, from the methods of determining oxidation states it is clear that substitution of an electronegative group by a carbon group or a hydrogen atom is a reduction and requires a reducing agent. For example, conversion of an acid chloride to a ketone by a lithium organocuprate reagent involves a reduction of the acid chloride to the ketone oxidation level.

$$R \xrightarrow{O}_{+3} Cl + LiCuR_{2}' \longrightarrow R \xrightarrow{O}_{+2} R' + LiCl + CuR'$$

Consequently the copper is oxidized from a cuprate species to an organocopper. By classifying organocuprates as reducing agents toward acid chlorides, we should expect that they could act as reducing agents toward other functional groups. It is not surprising therefore that as Michael addition reagents they can be used to give net reduction of an  $\alpha$ ,  $\beta$ -unsaturated ketone.



Reaction of the organocuprate intermediate with water gives the fully reduced product. If the organocuprate intermediate is reacted with bromine, the  $\alpha$ -brominated product is formed. This product has the equivalent oxidation level as the starting enone but differs in that an additional carbon substituent is present. Functionally this is equivalent to the addition of HBr to an enone. Thus functionally no net redox has taken place. If individual steps are considered, it is clear that the first step (addition of the organocuprate to the enone) is a reduction and the second step (reaction of the cuprate with bromine) is an oxidation.



No net change in the oxidation level has occurred for the overall process; however, each step in the sequence can involve an oxidation or reduction. This is

an important idea to keep in mind—even though no net change in the oxidation level occurs, individual steps in the sequence may have an oxidation or reduction and thus would require oxidants or reductants consistent with the individual step being undertaken.

The realization that many reactions or steps in reactions involve an oxidation or reduction is an important consideration when these reactions are being studied and learned. The change in oxidation level produced is indicative of the transformation and provides an additional organizational category by which reactions can be classified. Reagents can also be classified by their ability to cause oxidation or reduction. For example, from its addition reactions with alkenes and alkynes, bromine can be considered an oxidizing reagent for organic molecules. It is not surprising, therefore, to find that bromine also serves as an oxidant toward other functional groups such as enols, hydrocarbons, aldehydes, and organometallic compounds. Lithium aluminum hydride is well known as a reductant; thus, if it reacts with an organic compound, it is a good bet that some functional group is being reduced by the addition of a hydride. By applying the concepts of oxidation and reduction, a new view of organic reactions is possible which is often neglected but extremely important.

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#### PROBLEMS

**2.1.** Give the oxidation level of the indicated carbons in the following compounds:



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42 OXIDATION STATES OF ORGANIC COMPOUNDS



**2.2.** For the following reactions (1) write a balanced equation; (2) determine if an oxidation, reduction, or no change has occurred for the organic substrate(s); (3) if a redox process has occurred, indicate the oxidized and reduced products; (4) indicate the reagent responsible for the change in oxidation level of the organic component; and (5) name the functional group in the reactant and indicate what functional group it has been converted to in the product (if possible).





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44 OXIDATION STATES OF ORGANIC COMPOUNDS



**2.3.** The following conversions require the use of some reagent to effect the chemical change. Indicate whether an oxidizing agent, a reducing agent, or simply a catalyst is needed.





**2.4.** Based on the following reactions, classify the reagent over the arrow as an oxidant, a reductant, or neither.



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46 OXIDATION STATES OF ORGANIC COMPOUNDS







# 3

# ACIDITY AND BASICITY

Bronsted and Lewis Acids and Bases	47
Acid Strength	49
Acid–Base Equilibria	53
Amphoteric Compounds	56
Structural Effects on Acidity	56
Electronegativity	58
Inductive Effects	59
Resonance Effects	61
Bibliography	63
Problems	63

# **BRONSTED AND LEWIS ACIDS AND BASES**

The notion of acids and bases is one of the first and most important ideas we are exposed to in chemistry, but it is one of the things that is often poorly understood. The concept is actually very simple if a few basic ideas are *always* kept in mind. The first things that must be remembered are the definitions of acids and bases and what general structural features make compounds react as acids or bases.

Bronsted acids are defined as proton donors. The dissociation of a Bronsted acid yields a proton *and* the conjugate base (an anion, if the acid is a neutral compound) of the acid.

$$HA \longrightarrow A^- + H^+$$

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# 48 ACIDITY AND BASICITY

Bronsted bases are proton acceptors. The only way that a Bronsted base can "accept" a proton is to supply an electron pair and form a bond to the proton. Thus Bronsted bases often are compounds with unshared pairs of electrons that can be donated to a proton to form a bond. Upon reaction with a proton, a base is converted to its conjugate acid.

# $B: +H^+ \longrightarrow B-H^+$

So that charge is conserved, anionic bases upon reaction with a proton give neutral conjugate acids; neutral bases upon reaction with a proton give positively charged conjugate acids. Occasionally shared pairs of electrons can be given up to a proton such as when olefins react with acids. In such cases  $\pi$  electrons are the electron pair which forms a bond to the proton.

The above two equations describing the behavior of Bronsted acids and bases are not strictly correct because a Bronsted acid does not just dissociate, it donates a proton *to* something which accepts a proton. The proton does not just dissociate and float around in solution but is always attached to something. Furthermore a Bronsted base does not just find a proton to accept; it accepts a proton from a Bronsted acid. Thus acidity and basicity are paired behavior—"you can't have one without the other." This is the most common misconception about acids and bases and leads to the greatest amount of difficulties when trying to apply the principles of acidity and basicity to real reactions.

A second description of acidity and basicity is the Lewis definition. A Lewis acid is an electron acceptor and a Lewis base is an electron donor. The definition of Lewis acids includes the proton since the proton can accept electron pairs. Thus Bronsted acids are a subset of Lewis acids since all Bronsted acids yield a proton. However, other common Lewis acids (BF₃, TiCl₄, SnCl₄, AlCl₃) are routinely used as organic reagents and all function by accepting unshared or shared electron pairs. In general the electron pairs that are accepted are readily available; that is, they are not tightly bound, so they are usually lone pairs or  $\pi$  electrons.

Lewis bases are electron pair donors and the electrons are given up to Lewis acids (electron acceptors). Unshared electron pairs are the more common type of electron pairs to be donated to Lewis acids, although on occasion shared pairs can be donated to the Lewis acid. In the case of Lewis acid–base reactions, the product is termed a *Lewis acid–base complex*.

Given these definitions, it is crucial to emphasize again that no compound is inherently an acid or a base. A compound only functions as an acid (in the Bronsted sense) if it donates a proton *to* something; that is, there must be a base present (a proton acceptor) to have an acid–base reaction. Thus a compound acts as an acid only in the presence of a base, and a substance can only act as a base in the presence of an acid. A good example of this concept is the fact that HCl in the vapor phase is an undissociated, covalent molecule. This is because there are no molecules present capable of accepting a proton from HCl. As a result, HCl does not function as an acid in the gas phase. If  $H_2O$  is added, however, an immediate acid-base reaction takes place in which HCl donates a proton to water, which is capable of accepting a proton from HCl. In this reaction an unshared pair of electrons on the oxygen atom of water is donated to the proton and an O-H bond is formed. In its reaction with water, HCl can function as an acid only because water can function as a base to accept the proton.

HCl 
$$\implies$$
 H⁺ + Cl⁻ vapor phase — no reaction!

but

$$HCl + H_2O \longrightarrow H_3O^+ + Cl^-$$

While the necessity of having *both* and acid and a base present in order to have an acid-base reaction is axiomatic, it is surprising how often this concept is neglected. However, if these conditions are met, then a wide variety of organic compounds can donate protons to appropriate bases (they are deprotonated) and a wide variety of compounds can accept protons from appropriate acids (they are protonated).

The same considerations are true for Lewis acids and bases. Pure boron trifluoride does not act as a Lewis acid because there is nothing present capable of donating electrons to it. If diethyl ether is added, boron trifluoride etherate, a stable Lewis acid-base complex, is produced. Obviously the electron pairs on the oxygen atom of diethyl ether can be donated to boron trifluoride. When a Lewis base is present, then boron trifluoride functions very effectively as a Lewis acid.

$$BF_3 + O(Et)_2 \longrightarrow F_3B - O(Et)_2$$

Most organic compounds do not act as Lewis acids because they are generally closed-shell molecules with filled valence levels and no unfilled orbitals capable of accepting electrons. By virtue of the unshared pairs of electrons on oxygen and nitrogen atoms, organic compounds which contain these elements can often function as Lewis bases toward many Lewis acids. Lewis acids such as BF₃, TiCl₄, or SnCl₄ are commonly used to react with oxygen-containing Lewis bases such as carbonyl compounds, alcohols, or ethers.

# ACID STRENGTH

Long before we ever took a chemistry course we all had some knowledge of the relative strengths of acids and bases. If you asked any of your friends if they

## 50 ACIDITY AND BASICITY

would rather spill battery acid or vinegar (acetic acid) on their skin, all would choose vinegar. (Even those who have never studied chemistry would make the same choice.) And if you asked them why, they would tell you that battery acid is "stronger" than vinegar. The same people would also prefer to ingest a solution of baking soda (NaHCO₃) rather than lye (KOH). Again they know that lye is stronger than soda (although they may not know both are bases). Thus the concept of strong or weak acids and bases is known to more people than you might suspect.

Chemists, however, would rather be a little more specific than describing acids and bases as either strong or weak. They need to be able to put acid or base strength on a quantitative basis. One way to measure the strength of an acid would be to place your index finger into the acid. The ability of an acid to transfer a proton to your index finger, which would act as a base and accept a proton, would result in an audible response—the pain index. From our previous discussion, if battery acid and vinegar were compared by placing the index finger in each, the pain index for battery acid would be much greater than for vinegar because battery acid is "stronger" and can thus transfer a proton to your finger more effectively than vinegar. Other acids could be tested in a similar fashion and their acid strength could be ranked by the audible signal. Since the same base, an index finger, is used to test every acid, the pain index is a direct and quantitative measure of the strength of a given acid compared to the other acids that have been tested. Unfortunately the method is not very reproducible and the sensor can only be used for a limited number of measurements without degradation.



But the idea is clear! The strength of an acid can be quantitated by knowing how well it transfers a proton to some standard base. Moreover the extent to which it transfers a proton to the standard base could be compared with the extent to which other acids transfer a proton to the same standard base. Thus, by using a single base, we could not only measure the strength of an acid but also compare acids quantitatively.

To make it simple, let us choose a molecule as the standard base. The acid strength will be a measure of how well the acid transfers a proton to that molecule acting as a base, and we can quantitate the acid strength by measuring the amount of the acid that is ionized in the presence of that base. Thus the ratio of the amount of acid which has transferred a proton to the base compared to the amount of acid which has not transferred a proton to the base is a direct measure of the strength of that acid. Water has been chosen as the standard base molecule (in place of a finger) and the *ratio* of ionized acid to undissociated acid is called  $K_{eq}$  (the equilibrium constant).

HA + H₂O 
$$\xrightarrow{K_{eq}}$$
 H₃O⁺ + A⁻

The equilibrium constant  $K_{eq}$  is expressed as the amount of products divided by the amount of reactants present at equilibrium; or

$$K_{\rm eq} = \frac{[{\rm H}_3{\rm O}^+][{\rm A}^-]}{[{\rm H}{\rm A}][{\rm H}_2{\rm O}]}$$

(Since two product molecules are produced from two reactant molecules, the *ionization ratio* of the acid is expressed as the product of the product concentrations divided by the product of the reactant concentrations). This numerical ratio can be used to compare the strengths of acids. It is very important to realize just how simple this notion is. The equilibrium constant (in this case the ionization constant of an acid) is merely a ratio of the molecules which have donated a proton to water to those which have not.

To further reduce the complexity, since the concentration of water,  $[H_2O]$ , is a constant in dilute solutions where this treatment is valid, this fraction can be simplified to

$$K_{\rm eq}[{\rm H}_2{\rm O}] = K_{\rm a} = \frac{[{\rm H}_3{\rm O}^+][{\rm A}^-]}{[{\rm H}{\rm A}]}$$

which is again essentially a ratio relating the ionized acid molecules to the unionized acid molecules. For example, if  $K_a$  is  $1/10^5$ , or  $10^{-5}$ , then one of every  $10^{2.5}$ acid molecules is ionized. On the other hand, if  $K_a$  is  $10^6/1$ , then for every  $10^3$ molecules ionized, only one is not ionized. The latter acid is a much stronger acid than the former because a much higher fraction is ionized and donates a proton to water.

We further simplify the comparison by defining  $pK_a = -\log K_a$  because this gives us small numbers to deal with rather than exponentials. It must be noted, however, that  $pK_a$  units are logarithmic units so one  $pK_a$  unit represents a change of 1 in 10 in the ratio of acid molecules which are ionized to those which are not. By the use of  $pK_a$ 's, the larger the number, the weaker the acid. For example, if one compares acetic acid  $pK_a = 4.75 = -\log K_a (K_a = 1.8 \times 10^{-5})$  with hydrofluoric acid  $pK_a = 3.2 = -\log K_a (K_a = 6.3 \times 10^{-4})$ , it is seen that HF is the stronger acid by about 1.5 orders of magnitude. Likewise, HCl, which has  $pK_a = -6.6 (K_a = 3.98 \times 10^{-5})$ , by more than 10 orders of magnitude ( $10^{10}$  stronger). By using a standard, constant molecule as a base, the measure of the fraction of proton transfer to that base becomes a meaningful and quantitative

# 52 ACIDITY AND BASICITY

Acid	Conjugate Base	pK _a
$H_2SO_4$	$\mathrm{HSO}_4^-$	-9
HCl	Cl-	-6
ArSO ₃ H	$ArSO_3^-$	-6 to $-7$
OH+ 	o ↓	-6 to $-7$
RY	RY	
Y = H, R, OR		
$H_3O^+$	$H_2O$	-1.74
HF	$F^-$	3.22
ArNH ₃ ⁺	$ArNH_2$	5
pyridine·H ⁺	pyridine	5.2
RCO ₂ H	$RCO_2^-$	4-6
$H_2CO_3$	HCO ₃ ⁻	6.4
RC(O)CH ₂ C(O)R	$RC(O) \overset{\leftrightarrow}{C}HC(O)R$	9
$HCO_3^-$	$CO_3^-$	10.33
ArOH	ArO ⁻	10
RNH ₃ ⁺	$RNH_2$	10
$R_2 N H_2^+$	$R_2NH$	11
$R_3NH^+$	R ₃ N	12
$RC(O)CH_2CO_2R$	$RC(O) \stackrel{\leftrightarrow}{C} HCO_2 R$	11
RO ₂ CCH ₂ CO ₂ R	$RO_2 CCHCO_2 R$	13
H ₂ O	OH-	15.74
RCH ₂ OH	$RCH_2O^-$	16
R ₂ CHOH	$R_2 CHO^-$	17
R ₃ COH	$R_3CO^-$	18
RC(O)NH ₂	RC(O)NH ⁻	18-19
RC(O)CH ₂ R	RC(O) [⊖] CHR	19-20
RO ₂ CCH ₂ R	RO₂CCHR	24
RC≡CH	RC≡C⊖	25
NH ₃	$\rm NH_2^-$	33
R ₂ NH	$R_2N^-$	35
$R_2C = CHCH_3$	$R_2C = CHCH_2^-$	43
$R_2C=CH_2$	$R_2C = CH^-$	44
RH	$R^{-}$	50

 Table 3.1
 pK_a Values for Common Organic Acids

way to compare acid strengths. Table 3.1 is a compilation of  $pK_a$ 's of various acids. It is only a representative group of  $pK_a$  values; literally thousands have been precisely measured. It is evident that, although the acid strengths of organic compounds can vary over 50 orders of magnitude, quantitative comparisons can be made between various acids with good accuracy.

# ACID-BASE EQUILIBRIA

Since  $pK_a$  values always refer to the ionization of an acid in water under standard conditions (dilute aqueous solution at 25°C), they can be used to predict the positions of acid–base equilibria. The position of the following equilibrium ( $pK_{eq}$ ) can be calculated by noting that in going from left to right isopropanol is acting as an acid.

$$\rightarrow$$
 OH + CH₃CH₂NH₂  $\rightarrow$  O⁻ + CH₃CH₂NH₃⁺

In going in the reverse direction, from right to left, the ethyl ammonium ion is acting as an acid. Having identified the compounds acting as acids on either side of the equilibrium, the  $pK_a$  values for those acids are found and added to the equation.

$$\begin{array}{c} \searrow \text{OH} + \text{CH}_3\text{CH}_2\text{NH}_2 \xrightarrow{pK_{eq}} \\ & \searrow \text{O}^- + \text{CH}_3\text{CH}_2\text{NH}_3^+ \\ pK_a = 17 \\ & pK_a = 10 \end{array}$$

To evaluate the position of the equilibrium,  $pK_{eq}$  is determined by the equation  $pK_{eq} = pK_a$  (reactant acid)  $-pK_a$  (product acid). For the above example  $pK_{eq} = 17 - 10 = 7$ , and based on the definition of pK,  $K_{eq} = 10^{-7}$  for this equilibrium. Thus the equilibrium lies far to the reactant side; that is, very little isopropoxide ion or ethyl ammonium ion is present at equilibrium. This technique is applicable for virtually any acid-base equilibrium. The three required steps are to (a) write a balanced equation that describes the equilibrium to be analyzed, (b) identify the species which is acting as an acid on each side of the equilibrium and write down its  $pK_a$ , and (c) subtract the  $pK_a$  of the product acid from the  $pK_a$  of the reactant acid to give the  $pK_{eq}$  for the equilibrium in question. It is a requirement that the  $pK_a$ 's of the acids on each side of the equilibrium are known or can be estimated reasonably well. Furthermore, the  $pK_{eq}$  that is determined refers to the equilibrium *in the direction it is written*. It is therefore important to write the chemical equilibrium as you wish to analyze it.

The reaction of methyl lithium with water is an acid-base reaction. Going from left to right, water donates a proton to CH₃Li so it functions as the acid on the left. Going from right to left, methane donates a proton to LiOH so it functions as the acid on the right. The  $pK_a$ 's of the acids on either side of the equilibrium reaction are subtracted and  $pK_{eq} = -34.3$ . Thus the equilibrium constant is  $K_{eq} = 10^{34.3}$ , which shows that the equilibrium lies very far to the right—so far to the right that for all practical purposes the conversion is quantitative.

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54 ACIDITY AND BASICITY

CH₃Li + H₂O 
$$\xrightarrow{K_{eq}}$$
 CH₄ + LiOH  $K_{eq} = 10^{34.3}$   
p $K_a = 15.7$  p $K_a = 50$ 

It is now common experimental practice to react ketones with lithium diisopropyl amide (LDA) in order to generate the enolate of the ketone. This methodology has largely replaced the older approach to enolates, which employed alkoxide bases to remove a proton alpha to the carbonyl group. Comparison of the equilibrium constants for these two acid-base reactions reveals why the LDA method is preferable. The use of the amide base leads to essentially complete conversion of the ketone to its enolate ( $K_{eq} \approx 10^{16}$ ). At equilibrium, there is virtually no unreacted ketone present in solution, only diisopropyl amine and the enolate, which can subsequently react cleanly and controllably with electrophiles which are added to the reaction mixture. In contrast, the use of an alkoxide base results in only partial conversion of the ketone to its enolate  $(K_{eq} \approx 10^{-2})$  so that at equilibrium the enolate, even greater amounts of the unreacted ketone, and some unreacted alkoxide are present in the reaction mixture. It is therefore difficult to react the enolate with an added electrophile while avoiding competing reactions of the enolate with the unreacted ketone and the alkoxide with the added electrophile. Thus control of the reaction is difficult and consequently yields are decreased and the product mixture is more complex.

$$(i-Pr)_{2}^{\Theta} \overset{\bigoplus}{\text{Li}} + \text{RC}-\text{CH}_{2}\text{R'} \overset{\bigoplus}{\text{(iPr)}_{2}\text{NH}} + \text{RC}=\text{CHR'} \quad K_{eq} = 10^{16}$$

$$\downarrow \\ O \\ DLi \\ OLi \\$$

The ability to make good estimates of acid–base equilibrium constants is an invaluable aid in thinking about organic reactions and processes. Moreover, experimental workup procedures often require pH control that can be easily understood on the basis of  $pK_a$  considerations. Thus the concept of acid strength is exceedingly important and should be mastered.

A similar development can be used for the quantitation and comparison of the base strengths of organic bases (the ability to accept protons from acids). To do this,  $K_b$  is defined as

$$K_{\rm b} = \frac{[\rm OH^-][\rm BH^+]}{[\rm B]}$$

the fraction of organic base which removes a proton from the standard acid—water. A scale of  $K_b$ 's was developed with which to compare base strengths quantitatively.

The situation can become quite confusing since two sets of constants are defined,  $K_a$  and  $K_b$ . It turns out that acidities and basicities are inversely related by the ionization constant of water. That is, the stronger an acid is in donating a proton to water, the weaker its conjugate base is in removing a proton from water. This seems eminently reasonable since if something gives up a proton easily, then it should not take protons back easily. Put in more chemical terms, strong acids have weak conjugate bases, and weak acids have strong conjugate bases. It has become the convention to list only  $pK_a$ 's as a measure of *both* acidity and basicity. The only thing to remember is to assign the appropriate  $K_a$  to the reaction in the "acidic" direction.

For example, the base strengths of NH₃ and CH₃O⁻ can be compared in two different ways. First, the reactions of these two bases with water can be written as follows, and by the preceding analysis the two equilibrium constants can be estimated by identifying the acids on either sides of the two equilibria and subtracting their  $pK_a$ 's.

$$CH_{3}O^{-} + H_{2}O \xrightarrow{K_{eq}} CH_{3}OH + OH^{-} pK_{eq} = -1, K_{eq} = 10$$

$$pK_{a} = 15 \qquad pK_{a} = 16$$

$$NH_{3} + H_{2}O \xrightarrow{K_{eq}} NH_{4}^{+} + OH^{-} pK_{eq} = 5.8, K_{eq} \approx 10^{-6}$$

$$pK_{a} = 15 \qquad pK_{a} = 9.2$$

It is seen that the equilibrium for the reaction of methoxide with water lies much farther to the right ( $K_{eq} = 10$ ) than the reaction of ammonia with water ( $K_{eq} = 10^{-6}$ ). Clearly methoxide is much better at removing a proton from water than is ammonia by about  $10^7$ . Therefore methoxide is a stronger base by about  $10^7$  than ammonia.

Alternatively it is noted that the conjugate acids of ammonia and methoxide are the ammonium ion and methanol, respectively, and the equations for their ionization in water are

$$\begin{array}{rcl} \mathrm{NH}_{4}^{+} &+ &\mathrm{H}_{2}\mathrm{O} & & & & \mathrm{NH}_{3} &+ &\mathrm{H}_{3}\mathrm{O}^{+} & & pK_{\mathrm{eq}} = 11, \ K_{\mathrm{eq}} = 10^{-11} \\ pK_{\mathrm{a}} = 9.2 & & & pK_{\mathrm{a}} = -2 \end{array}$$

$$\begin{array}{rcl} \mathrm{CH}_{3}\mathrm{OH} &+ &\mathrm{H}_{2}\mathrm{O} & & & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ &\mathrm{H}_{2}\mathrm{O} & & & & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ &\mathrm{H}_{2}\mathrm{O} & & & & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ & \mathrm{H}_{2}\mathrm{O} & & & & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ & \mathrm{H}_{2}\mathrm{O} & & & & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ & \mathrm{H}_{2}\mathrm{O} & & & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ & \mathrm{H}_{2}\mathrm{O} & & & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ & \mathrm{H}_{2}\mathrm{OH} &+ & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ & \mathrm{H}_{2}\mathrm{OH} & & & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ & \mathrm{H}_{2}\mathrm{OH} & & & & \\ \mathrm{CH}_{3}\mathrm{OH} &+ & \mathrm{H}_{2}\mathrm{OH} &+ & \\ \mathrm{CH}_{3}\mathrm{OH} &+ & \mathrm{H}_{3}\mathrm{OH} &+ & \\ \mathrm{CH}_{3}\mathrm{OH} &+$$

Comparing these two equilibria, it is seen that methanol is a much weaker acid than ammonium ion by about  $10^7$  (i.e., the  $pK_a$  of methanol is larger by about seven logarithm units than the  $pK_a$  of ammonium); thus its conjugate base methoxide should be a much stronger base than ammonia by  $10^7$ . Either method of analyzing the situation is acceptable, and each gives the same relative basicities without the need for two sets of ionization constants,  $K_a$  and  $K_b$ .

# 56 ACIDITY AND BASICITY

# AMPHOTERIC COMPOUNDS

A glance at the  $pK_a$  values in Table 3.1 reveals that many classes of compounds can act either as acids or as bases, depending on the reaction environment. Such materials are termed *amphoteric*. They must have an acidic proton (i.e., a proton attached to an electronegative element or group) and unshared pairs of electrons that can be donated to a proton. For example, water, alcohols, and other hydroxylic compounds as well as amines and amides are all amphoteric materials. Comparing the  $pK_a$ 's of these materials permits an assessment of the predominant behavior in a given environment. For example, if an amine is dissolved in water, it could function as an acid or a base. To determine which behavior will predominate, the position of the equilibrium can be determined for each process. Comparison of these values will indicate which will be the principal behavior. Thus, as an acid, the amine would donate a proton to water to give an amide anion and the hydronium ion.

RNH₂ + H₂O 
$$\xrightarrow{K_{eq}}$$
 RNH⁻ + H₃O⁺  $K_{eq} = 10^{-35}$   
p $K_a = 33$   $pK_a = -2$ 

As a base the amine would accept a proton from water to give an ammonium ion and hydroxide.

RNH₂ + H₂O   

$$pK_a = 15$$
 RNH₃⁺ + OH⁻  $K_{eq} = 10^{-6}$ 

Since the equilibrium constant for the amine acting as an acid is  $10^{-35}$  and that for the amine acting as a base is much larger at  $10^{-6}$ , reaction as a base will be the main behavior in aqueous solution. The magnitude of the equilibrium constant ( $10^{-6}$ ) indicates that it is only a weak base.

On the other hand, if you treat an amine such as diisopropyl amine with *n*-butyl lithium in tetrahydrofuran (THF), then a different behavior is indicated. In this case the equilibrium lies far to the right so as to be virtually irreversible. The amine therefore functions as an acid and donates a proton to the much stronger base butyl lithium. This is the standard method for the preparation of the versatile base LDA.

$$(i-\Pr)_2 NH + n-BuLi$$
  $(i-\Pr)_2 N^- LI^+ + n-BuH$   $K_{eq} = 10^{+13}$   
 $pK_a = 35$   $pK_a = 48$ 

# STRUCTURAL EFFECTS ON ACIDITY

Now that a method is in hand to compare acid strengths quantitatively and predict the position of acid-base equilibria, a look at Table 3.1 reveals that organic compounds have an enormous range of acidities from very strong acids such as arenesulfonic acids ( $pK_a = -6.5$ ) and protonated carbonyl compounds ( $pK_a = -7$  to  $pK_a = -10$ ) to very weak acids such as alkanes ( $pK_a = 50$ ) and alkenes ( $pK_a = 45$ ). This huge range of acidity of  $\sim 10^{60}$  is reflective of the huge diversity of structural elements present in organic compounds.

How the structure of an acid influences its  $pK_a$  provides a quantitative way to compare the structure of a compound with its reactivity (in this case acidity). Such structure–reactivity correlations are crucial for our understanding of how reactions take place and for being able to predict how a structural change will affect the outcome of a reaction. The ability to predict how a reaction will respond to changes in structure (or other variables) takes us out of the realm of trial and error and into the realm of rational approaches to chemical transformations. Let us examine briefly some of the structural features which are major influences on acidity.

Returning to the dissociation of an acid in water, it is seen that the process has some energy costs and some energy gains. It is this energy balance  $(\Delta G)$  which determines the equilibrium constant and hence the strength of an acid. The dissociation of an acid in water has an energy cost from the breaking of a bond to hydrogen and the separation of charges produced by the ionization.

$$HA + H_2O \implies A^- + H_3O^+$$

However, there is an energy gain from the formation of the OH bond of the hydronium ion and the solvation of the anion by hydrogen bonding to the solvent and solvation of the hydronium by its hydrogen bonding to the solvent. If a series of different acids is now compared, it becomes clear that a major energetic difference in the dissociation of various acids in water is the *stability of the conjugate base* and its interaction with the aqueous solvent.

This is because the other energetic factors which influence the equilibria are similar for different acids. Bonds from hydrogen to the first-row elements have similar bond strengths ( $\pm$ 5 kcal/mol) so the energy cost of breaking the bond to hydrogen is relatively constant for most acids of first-row elements. This analysis is especially true for acids with the proton bonded to the same element. Moreover, since the solvent is always water, the energy required to separate charges is about the same. Finally, the H–O bond of the hydronium ion is the same, regardless

# 58 ACIDITY AND BASICITY

of which acid supplies the proton. Consequently, the principal differences in the  $\Delta G$ 's of ionization for various acids are due to the differences in stability of their conjugate bases in the reaction mixture.

This analysis suggests that structural features which stabilize the conjugate base (often an anion) will therefore increase the acidity of an acid. While there are exceptions to this general approach (e.g., comparison of the acidities of acids in the second and third rows of the periodic table), it provides a sound basis for predicting what structural factors can increase or decrease the acidity of organic acids.

There are three principal factors that lead to increased stability of anions: (a) the electronegativity of the atom carrying the negative charge, (b) inductive effects which can stabilize negative charge, and (c) resonance effects which delocalize the negative charge over several atoms and hence stabilize the anion.

# Electronegativity

Increased electronegativity of an atom allows it to carry negative charge more readily, and the stability of the anion is increased. It is for this reason that the order of acidity of first-row hydrides is C-H < NH < -OH < FH. Transfer of a proton from these substances to water yields a series of anions whose stabilities are ordered according to the electronegativity of the negatively charged atom.



Thus

-C-H N-H  $-\ddot{O}-H$  F-Hincreasing acidity  $-\dot{O}$ 

Such ordering is valid only for elements in the same row in the periodic table. Comparisons between acids in which the proton is lost from elements from different rows is not valid because the bond strength to the acidic hydrogen changes greatly from row to row. In the above analysis bond strength is assumed to be relatively constant; thus it cannot be neglected when significant bond strength differences between acids are present.

The effective electronegativity of the atom carrying the charge is also dependent on the hybridization of that atom. As the s character of an orbital increases, electrons in that orbital are more stable due to greater attraction to the nucleus. Thus the effective electronegativity of the atom increases. This effect is clearly seen in the relative acidities of hydrocarbons. Removal of a proton from alkanes, alkenes, and alkynes produce conjugate bases with electron pairs in sp³, sp², and sp orbitals, respectively. As the amount of s character increases from 25 to 33 to 50% in this series, the stability of the conjugate base increases and accounts for the marked increase in acidity in the series. Based on these data, it is expected that cyclopropane, which because of ring strain has the hydrogens bonded to carbons which are hybridized at about an sp^{2.5} level (29% s character), should have a p $K_a$  between that of an alkane and an alkene. In fact, the p $K_a$  of cyclopropane is 46 as predicted.

Acid	Conjugate Base	Percents Character	pK _a
R–CH ₃	$R-CH_2^-$	25	$\approx 50$
$RCH=CH_2$	RCH=CH ⁻	33	44
R–C≡CH	$R-C\equiv C^{-}$	50	25

The increase in acidity by 25 orders of magnitude between  $sp^3$ - and sphybridized carbon acids is similar to that found for the difference in acidity between an ammonium ion ( $sp^3$  hybridization) and a protonated nitrile (sp hybridization). It is clear that the hybridization of the orbital they occupy can play a major role in stabilizing electron pairs and thus influencing the effective electronegativity of an atom.

$$R-C \equiv \stackrel{+}{N}-H + H_2O \implies R-C \equiv N: + H_3O^+ \quad pK_a = -10$$
$$R-NH_3^+ + H_2O \implies R-\dot{N}H_2 + H_3O^+ \quad pK_a = 10$$

## **Inductive Effects**

The inductive effect is the ability of a substituent or group near the acidic proton to alter the electron distribution at the reaction center by through-bond displacement of electrons. The result is that substituents which withdraw electrons from the reaction center by the inductive effect stabilize anions and thus increase the acidity of the conjugate acids of those anions. Conversely, groups which donate electrons make the reaction center more electron rich and thus make the formation of the anion at that center more difficult. The conjugate acid is thus a weaker acid.

This is easily demonstrated by considering a group of substituted acetic acids (Table 3.2). Compared to acetic acid (X = H), replacement of a hydrogen by more electron-withdrawing groups [Cl, F,  $(CH_3)_3N^+$ ] leads to an increase in the acidity. Replacement of hydrogen with an electron-donating *t*-butyl group decreases the acidity. We can understand these changes in terms of the inductive effect. If we compare the conjugate bases of acetic acid and chloroacetic acids, it is seen that the carbon–chlorine bond has a dipole moment associated with it.
#### 60 ACIDITY AND BASICITY

x	pK _a
t-Bu	5.05
Н	4.75
Cl	2.86
F	2.66
$(CH_3)_3N^+$	1.83

Table 3.2 Acidities of Substituted Acetic Acids X–CH₂CO₂H

This bond dipole induces smaller dipole moments in adjacent bonds, which in turn induces ever smaller dipole moments in adjacent bonds.

$$\begin{array}{ccc} & \longleftarrow & O \\ C_1 - CH_2 - C - O^- \end{array} \qquad \begin{array}{ccc} & \longleftarrow & \rightarrow & O \\ (CH_3)_3 C - CH_2 - C - O^- \end{array}$$

The result of this inductive effect is that the electron density on the carboxylate anion is reduced, the negative charge is distributed over more atoms, and the chloroacetate anion is stabilized relative to acetate. Because the chloroacetate anion is more stable than the acetate ion, its conjugate acid, chloroacetic acid, is a stronger acid than the conjugate acid of the acetate ion, acetic acid (Table 3.1).

As is expected, groups with higher electronegativity (X = F) or electron deficiency result in greater inductive electron withdrawal, the anion is more stable, and the acidity is increased. Conversely, a group such as *t*-butyl is electron donating relative to hydrogen. Its inductive effect serves to increase the electron density on the carboxylate group, destabilize the anion, and thus decrease the acidity of its conjugate acid.

As mentioned, inductive effects operate through bonds by successive bond polarizations. As such, they diminish rapidly with distance so that very little effect results if an inductive effect must be transferred through more than four bonds. As seen in Table 3.3, placement of a chlorine substituent next to the carboxyl group causes a hundredfold increase in acidity. Moving it to the  $\beta$  position reduces the effect significantly, while a  $\gamma$ -chloro substituent causes almost no acidity increase.

Table 3.3Acidities of ChlorobutanoicAcids as Position of Chlorine AttachmentIs Varied		
Acid	pK _a	
CH ₃ CH ₂ CH ₂ CO ₂ H CH ₃ CH ₂ CHClCO ₂ H CH ₃ CHClCH ₂ CO ₂ H ClCH ₂ CH ₂ CH ₂ CO ₂ H	4.88 2.80 4.06 4.52	

Inductive effects serve to alter the electron distributions in molecules, and consequently they are very important influences on many types of reactions—not just acidity and basicity. To the extent that electronic changes occur during the conversion of reactants to products, inductive effects can facilitate or impede those electronic changes and thus change the rates of conversion. It is important then to keep them in mind when other examples of reactivity changes are discussed.

#### **Resonance Effects**

A final structural effect which influences acidity is the delocalization of electrons via resonance. In terms of acid-base behavior, resonance delocalization can stabilize the conjugate base of an acid, thus making the acid a stronger acid. For example, alcohols have  $pK_a$ 's of ~16 whereas carboxylic acids have  $pK_a$ 's of ~5. In each case the acidic proton is lost from oxygen. The bond strength to the proton and the electronegativity of the atom carrying the charge (oxygen) are identical; thus these factors cannot account for the large difference in acidity. On the other hand, the alkoxide ion is a localized anion with the oxygen atom carrying a full negative charge while the carboxylate ion is resonance delocalized. In the carboxylate ion the electron pair and negative charge are distributed between both oxygens so that each oxygen carries only a partial negative charge (actually about  $-\frac{1}{2}$ ) and the anion is greatly stabilized. Thus carboxylic acids are more acidic than alcohols by  $\approx 10^{11}$  or so.

$$ROH + H_2O \xrightarrow{pK_a=16} RO^- + H_3O^+$$

$$R \xrightarrow{O}_{OH} + H_2O \xrightarrow{pK_a=5} H_3O^+ + R \xrightarrow{O}_{O^-} \xrightarrow{R} R \xrightarrow{O^-}_{O}$$

The following groups of compounds illustrate the profound effect that resonance delocalization has on the stability of anions and hence the acidity of the conjugate acids. To compare the acidities of these acids, the conjugate bases can be ranked according to their resonance stabilization and that ranking of anion stabilization is predictive of the acidity orders.

R-OH
 
$$\bigcirc$$
 OH
 R-NH2

  $pK_a = 16$ 
 $pK_a = 10$ 
 $pK_a = 5$ 
 $pK_a = 33$ 

 1
 2
 3
 4

 O
 R-OH
  $\bigcirc$ 
 CH2
 OH
 R-NH2

  $R$ 
 $\square$ 
 NH2
 R-OH
  $\bigcirc$ 
 $>$ 
 $\bigcirc$ 
 $>$ 

#### 62 ACIDITY AND BASICITY

While resonance stabilization is greatest for those compounds which have more electron density distributed to more electronegative elements (compare 1, 2, and 3), delocalization of charge over any elements results in significant anion stabilization and a corresponding increase in acidity of the conjugate acid of that anion (e.g., 6, 7, 8). Moreover electronegativity effects can be considered in addition to resonance effects where applicable. Both amides 5 and methyl ketones 8 have resonance stabilization, but in amide anions the negative charge is shared between nitrogen and oxygen, while in ketone enolates the negative charge is shared between carbon and oxygen. Due to the greater electronegativity of nitrogen over carbon, the amide anion is more stable and hence amides ( $pK_a \approx$ 18–19) are somewhat more acidic than ketones ( $pK_a \approx 20-21$ )



A particularly strong type of resonance stabilization is found for those compounds which form an aromatic ring upon removal of a proton. The enhanced aromatic stability of the conjugate base translates into a large increase in acidity of the acid. Whereas the doubly allylic proton of 1,4-pentadiene is predicted to have a  $pK_a \approx 40$  due to resonance stabilization of the anion, the doubly allylic proton in cyclopentadiene has a  $pK_a = 16$  because the resulting anion produces an aromatic  $\pi$  system.



Aromaticity also explains why tropolone  $(pK_a \approx -5)$  is slightly more basic than a normal ketone  $(pK_a \approx -7)$ . The conjugate acid is stabilized upon protonation by the formation of an aromatic tropylium ion.



Inductive and resonance effects described above can significantly alter the electron distributions in molecules and can influence not only acidity but many

other reactions as well. A general understanding of these effects will be important in many different transformations we will encounter.

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### PROBLEMS

**3.1.** Using the  $pK_a$  data from Table 3.1, predict the position of the following equilibria and give an estimate of the equilibrium constant:

(a) 
$$NH_3 + OH^- \implies NH_2^- + H_2O$$

(b)  $CH_3CO_2H + CO_3^{-2} \longrightarrow CH_3CO_2^{-} + HCO_3^{-}$ 

(c) 
$$\operatorname{RCO}_2H + \operatorname{R}_2NH \Longrightarrow \operatorname{RCO}_2^- + \operatorname{R}_2NH_2^+$$

- (d)  $R_3COH + RC(O)CHR' \implies R_3CO^- + RC(O)CH_2R'$
- (e) NaF + HCl  $\implies$  NaCl + HF
- (f)  $ArOH + RNH_2 \implies ArO^- + RNH_3^+$
- (g)  $RC(O)OR' + H_2SO_4 \implies RC(OH+)OR' + HSO_4^-$
- (h)  $RC(O)CH_2C(O)R' + R_3N \longrightarrow RC(O)CHC(O)R' + R_3NH^+$
- (i)  $RC \equiv C^- + RCH_2OH \implies RC \equiv CH + RCH_2O^-$
- (j)  $CH_3Li + (CH_3)_2C=CH_2 \iff CH_4 + (CH_3)_2C=CHLi$
- **3.2.** Predict the positions of the following equilibria (left, right, or center) and give an approximate pK value for the equilibrium. You may need to look

#### 64 ACIDITY AND BASICITY

up or estimate  $pK_a$  values using structural effects on the  $pK_a$  values of known compounds. If you do estimate the  $pK_a$ 's, rationalize your method of estimation.



3.3. List (by number) in order of increasing acidity of the underlined protons.

(a) 1.  $CH_{3}CH_{2}-\underline{H}$  2.  $CH_{3}O-\underline{H}$ 3.  $\underline{H}-F$  4.  $CH_{3}CH_{2}NH-\underline{H}$ (b) 1.  $\underline{H}-Cl$  2.  $\underline{H}-F$ 3.  $\underline{H}-Br$  4.  $\underline{H}-I$ (c) 1.  $CH_{3}CHCH_{3}$  2.  $Cl-O-\underline{H}$ 

3. 
$$H_3C - C - O\underline{H}$$
  
 $\parallel$   
 $O$   
 $H_3C - C - \underline{H}$   
 $\parallel$   
 $O$   
 $O$ 

( <b>d</b> )	1. О    СН ₃ СН ₂ С—О <u>Н</u>	2. O $H_3CH_2C-CH_2\underline{H}$
	3. O $H_3CH_2C - NH H$	4. CH ₃ CH ₂ OCH ₂ — <u>Н</u>
(e)	1. СH ₃ CH ₂ C−О <u>Н</u> ∥ О	2. F $H_3$ CHC $-O\underline{H}$ H O
	3. $Cl$ $CH_3CHC - OH$ O	4. Cl $\stackrel{ }{\underset{CH_2CH_2C}{\overset{ }{\underset{O}{\underset{O}{}}}}}$
( <b>f</b> )	1. CH ₃ O— <u>Н</u>	2. (CH ₃ ) ₂ CHO— <u>Н</u>
	3. CH ₃ CH ₂ O <u>H</u>	4. (CH ₃ ) ₃ CO− <u>H</u>
( <b>g</b> )	10 _{`<u>Н</u>}	$2. \underbrace{\qquad }_{F} O_{\underline{H}}$
	3. F O <u>H</u>	$\overset{4.}{\underset{F}{\bigvee}}^{0}\underline{H}$
( <b>h</b> )	$1. \underbrace{O O O}_{H_3C CH_2 O\underline{H}}$	2. 0
	3. O $H_{3}CCH_{2}CH_{2}-\underline{H}$	4. O Cl ₃ C O <u>H</u>
(i)	<u>1.</u> CH ₃ CH ₂ CH ₂ — <u>Н</u>	2. O →−CH ₂ <u>H</u>
	3. ^H ₂ C=CH-CH ₂ - <u>H</u>	4. о н о <u>н</u>

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66 ACIDITY AND BASICITY



- **3.4.** Give explanations for the following observations:
  - (a) Amides are protonated on oxygen rather than on nitrogen.
  - (b) Ethers are better Lewis bases than ketones.
  - (c) Tetramethyl guanidine (p $K_a \approx 12$ ) is a much stronger base than N,N-dimethylacetamide (p $K_a \approx -0.5$ ).



- (d) Boron trifluoride  $(BF_3)$  is a stronger Lewis acid than trimethyl borate  $[(CH_3O)_3B]$ .
- (e) Piperidine is a much stronger Lewis base than pyridine.





pyridine

(f) *p*-Nitrophenol has  $pK_a \approx 8$  whereas phenol has  $pK_a \approx 10$ .



(g) *o*-Chloroaniline is a weaker base than *p*-chloroaniline.



(h) Sodium borohydride in alcohol does not reduce imines effectively. If BF₃ is added to the mixture, however, the reduction proceeds much more rapidly and efficiently.



- **3.5.** Provide a rationale for the fact that hydrogen sulfide  $(H_2S)$  is more acidic than water  $(H_2O)$ .
- **3.6.** It has been found that benzenesulfonic acid  $(pK_a \approx -6)$  is a much stronger acid than benzoic acid  $(pK_a \approx 5)$ . Discuss the factors that can account for this.



**3.7.** The reaction of pyrrole with a strong acid leads to protonation on C-2, not on nitrogen. Explain why.

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68 ACIDITY AND BASICITY



**3.8.** By  $pK_a$  measurements it can be shown that 1,1,1-bicyclopentanecarboxylic acid is 10 times more acidic than 2,2-dimethylpropanoic acid, even though both have a quaternary carbon attached to the carboxylic acid group. Account for this difference in  $pK_a$ .



**3.9.** It is observed that the rate of ionization of allylic benzoates is, in most cases, inversely proportional to the  $pK_a$ 's of the corresponding benzoic acids. (i.e., the lower the  $pK_a$ , the faster the rate of ionization.) Account for this correlation.



Z = various substituents

# 4

# **CURVED-ARROW NOTATION**

Electron Movement	69
Heterolytic Bond Cleavages	70
Heterolytic Bond Formation	71
Homolytic Bond Making and Bond Breaking	73
Resonance Structures	75
Depiction of Mechanism	76
Bibliography	82
Problems	82

# **ELECTRON MOVEMENT**

In a very simple sense, most organic reactions are accomplished merely by the movements of electrons. Since molecules are composed of atoms held together by bonds and covalent bonds are merely shared pairs of electrons, the conversion of one molecule to another by changes in chemical bonds between reactants and products can be described simply as changes in electron pairs that are shared between the various nuclei in the reactants. While this is a gross oversimplification, it nevertheless provides us with a very important tool with which to keep track of bonding changes that occur during the transformation of one molecule into another.

We keep track of electron pairs by noting changes in their location in molecules by means of curved arrows. The curved arrow depicts movement of an electron

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### 70 CURVED-ARROW NOTATION

pair *from* the tail of the arrow *to* the head of the arrow. For example, in the reaction of a proton with water to produce the hydronium ion, a new bond is formed from oxygen to hydrogen. The electrons in that bond start out as a lone pair on oxygen and are donated to and shared with the proton. We can easily show this electron movement with a curved arrow from the lone pair on oxygen to the proton.



In a similar sense the Lewis acid-base reaction between ammonia and boron trifluoride can be depicted by a curved arrow from the lone pair of electrons on nitrogen to the boron atom. This electron movement creates a new bond between nitrogen and boron, but the curved-arrow notation clearly illustrates that the electron pair of that bond is supplied by the nitrogen. An additional consequence is that the nitrogen atom gains a formal positive charge and the boron atom gains a formal negative charge.



Before further applications of electron movement using curved-arrow notation are presented, it is important to recognize just why it is such a useful tool for describing reactions. First it permits us to keep track of valence shell electrons during a chemical reaction and thus serves as a method for electronic bookkeeping. Second it shows how *changes* in bonding result from changes in electron distribution. Third it can show likely mechanisms for chemical reactions in terms of the breaking and making of chemical bonds.

For curved-arrow notation to be used correctly, however, the structural and bonding principles which we have already learned must be adhered to. Thus donor-acceptor properties, oxidation states, hybridization and the octet structure of atoms, normal valences, formal charges, reactive intermediates, and so on, must all be taken into account as curved-arrow notation is used to track changes in electron distribution that occur during chemical reactions. Within the framework of these principles, however, curved-arrow notation can be a powerful and effective tool for depicting bonding changes during chemical reactions. To do this, it is necessary to first understand the general processes of bond formation and bond cleavage that are commonly encountered.

# HETEROLYTIC BOND CLEAVAGES

Simple bond cleavages can proceed with the shared pair ultimately residing on either of the previously joined elements. In either case one atom has a sextet electronic structure and a positive charge. The other has a valence octet with at least one lone pair (the pair that was formerly shared) and a negative charge. Such bond cleavage, in which the previously shared pair goes with one of the bonded atoms, is termed *heterolytic cleavage* and necessarily results in the formation of charged species.



In general, the movement of electrons during heterolytic cleavage follows the direction of bond polarity. In a polar covalent bond the shared pair is displaced toward the more electronegative element. Upon cleavage the pair of bonded electrons are transferred completely to the more electronegative element, which becomes negatively charged, and the more electropositive element loses the bonded electron pair and becomes positively charged.



If one of the bonded elements is positively charged to begin with, it can gain the bonded pair upon bond cleavage and become neutral. Note, however, that net charge is always conserved in any reaction. Moreover, bond cleavages are depicted the same whether they involve  $\sigma$  or  $\pi$  bonds.



# HETEROLYTIC BOND FORMATION

The formation of a bond between two atoms can proceed by one of the atoms donating an electron pair and the other atom accepting the electron pair. As before, charge must be conserved and the loss and gain in electrons by the donor

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#### 72 CURVED-ARROW NOTATION

and acceptor, respectively, must be accompanied by a corresponding change in formal charge.



The atom donating the electron pair must obviously have an electron pair that is not tightly bound and thus is available for donation. Commonly, lone pairs and  $\pi$ -bonded electron pairs can be most easily donated, but occasionally electron pairs in  $\sigma$  bonds can be donated if the  $\sigma$  bond is weak or electron rich.

The atom accepting the electron pair must have an unfilled orbital available which the donated electron pair can populate. This can be an unfilled valence shell orbital, as is the case if the acceptor atom has a valence sextet, or it can be an accessible antibonding orbital, either  $\sigma^*$  or  $\pi^*$ .

Thus the reaction of acetone with  $BF_3$  is a Lewis acid–base reaction in which a lone pair of the ketone oxygen atom is donated to an unfilled valence orbital of  $BF_3$ . Bond formation is accompanied by the development of formal charges on both oxygen and boron.



The capture of carbocations by alcohols involves a similar donation of a lone pair of electrons on oxygen to the vacant 2p atomic orbital of the sp²-hybridized, sextet carbocation. Note that charge must be conserved so the first formed product is a positively charged oxonium ion.



Reaction of an alkene with a nitronium ion involves donation of the  $\pi$ -electron pair of the alkene into a  $\pi^*$  orbital of the nitronium ion. Donation of a bonded electron pair necessarily means that the bond from which it comes is broken. Likewise population of an antibonding level by electron donation generally results in breaking of the bond to which the antibonding orbital corresponds. In this case electron donation of the olefinic  $\pi$ -electron pair results in the rupture of the olefinic  $\pi$  bond and acceptance into the N–O  $\pi^*$  orbital results in breakage of the N–O  $\pi$  bond as well. Note that a new bond is formed and net charge is conserved.



Addition of an alkyl lithium reagent, which has a very electron rich  $\sigma$  bond between carbon and lithium, with a ketone involves electron donation of the  $\sigma$ electrons to the  $\pi^*$  orbital of the ketone to give a new carbon-carbon bond. The lithium counterion also plays a role in the addition by complexing with the lone pairs of the carbonyl oxygen, thus making the carbonyl group more electron deficient. Effectively this lowers the energy of the  $\pi^*$  orbital so its energy better matches the energy of the electron donor, and donation of electrons into that orbital is facilitated.



It is important to remember the requirements for any two-electron bondmaking process: first, there must be an available pair of electrons to be donated and, second, there must be an unoccupied orbital of suitable energy available into which the electrons can be donated.

#### HOMOLYTIC BOND MAKING AND BOND BREAKING

Heterolytic processes make up a large proportion of organic transformations because most bonds are somewhat polarized. Heterolytic cleavage is merely an increase of this polarity to the limit at which there is no bond remaining; that is, electron movement follows in the direction established by the bond polarity to give a cation–anion pair.

If a bond is particularly weak and/or nonpolar, bond cleavage can occur by a nonpolar or homolytic process. One electron of the shared pair goes with each of the two bonded atoms. Bond breaking then is the movement of single electrons rather than electron pairs and is indicated in curved-arrow notation as "halfheaded" arrows. Homolytic cleavage of a bond does not result in the formation of charge but does result in the formation of unpaired electron intermediates called free radicals. Free radicals normally have seven electrons in the valence

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#### 74 CURVED-ARROW NOTATION

shell and as a consequence are very reactive intermediates. Common examples of compounds which undergo homolytic bond cleavages include halogens ( $Br_2$ ,  $Cl_2$ ,  $F_2$ ), peroxides (R–O–O–R), and azocompounds (R–N=N–R).



All of these free-radical precursors are characterized by relatively weak, nonpolar bonds which, upon heating, break to give free-radical intermediates. Free radicals are very reactive and proceed to products by a variety of oneelectron, or homolytic, reactions.

Homolytic bond formation can occur when two free-radical species contact each other. Each has an available unpaired electron, and if these two electrons are shared, a new bond will result.

$$H_3C \cdot \cap \cap Cl \longrightarrow H_3C - Cl$$

This is simply the reverse of the homolytic cleavage. It is a very exothermic process (by the amount equal to the energy of the bond being formed), and it occurs at a *very* fast rate.

Homolytic bond formation can also occur by the reaction of a free radical with a bonded pair of electrons. Two common examples of this behavior are hydrogen (or other atom) abstraction reactions and free-radical addition to double bonds. Atom abstraction reactions take place by the interaction of a free radical with a  $\sigma$ -bonded atom. One electron of the  $\sigma$  bond pairs with the unpaired electron of the free radical to produce a new bond. The remaining electron of the  $\sigma$  bond remains on the fragment from which the atom has been abstracted and produces a new free-radical species. This process is energetically driven by bond strengths; that is, atom abstraction only occurs if the bond that is formed is stronger than the one that is broken. In the example of hydrogen abstraction shown below, a phenyl radical readily abstracts a benzylic hydrogen from toluene to give benzene plus the benzyl free radical because the aromatic C–H bond (103 kcal/mol) that is formed is appreciably stronger than the benzylic C–H bond (85 kcal/mol) that is broken.

Hydrogen Abstraction



Addition to  $\pi$  bonds is a second very common reaction of free radicals. Interaction of the free radical with the  $\pi$ -electron pair causes one of the  $\pi$  electrons to pair up with the unpaired electron of the free radical to produce a new bond to one of the  $\pi$ -bonded atoms. The remaining  $\pi$  electron is now unpaired and thus forms a new free-radical species. The process is often very favorable since the new  $\sigma$  bond (70–90 kcal/mol) formed in the addition process is normally much stronger than the  $\pi$  bond (60 kcal/mol) which is broken in the reaction. In the above example a new carbon–carbon  $\sigma$  bond is formed by free-radical addition to produce a new carbon–centered free radical; however, a wide variety of other free-radical species add readily to olefins.

# **RESONANCE STRUCTURES**

Curved-arrow notation is also a very useful device with which to generate resonance structures. In this application it is truly a bookkeeping system. Since individual canonical forms do not exist but are only thought of as resonance contributors to the description of a real molecule, the use of curved-arrow notation to convert one canonical form to another is without physical significance. Nevertheless it provides a useful tool to keep track of electrons and bonds in canonical structures. For example, the structures of carboxylate resonance contributors can be interconverted as follows:



#### 76 CURVED-ARROW NOTATION

Likewise the Kekulé forms of benzene can be shown:



Allyl cations, for example, can be shown nicely while keeping track of charges, electrons, and bonds:



Similar considerations can be used for a variety of intermediates and structures. In using curved-arrow notation to generate contributing resonance structures, the same rules of valence, charge, bonding, and so on, must be applied. Given these criteria, however, it is a straightforward exercise to generate complete sets of resonance structures which can then be evaluated.

# **DEPICTION OF MECHANISM**

Use of curved-arrow notation to depict the mechanisms of organic reactions requires that appropriate mechanistic principles be superimposed on the correct use of curved arrows to denote movement of electrons. The mechanism of a reaction is the stepwise process by which reactants are converted to products, and generally each step involves bond making and/or bond breaking that can readily be depicted by curved-arrow notation.

Simple substitution reactions are shown in curved-arrow notation as



In this example a nucleophile donates electrons to the electrophile, in this case a carbon with a leaving group attached, to produce a new  $\sigma$  bond. As the new  $\sigma$  bond is formed, the bond to the leaving group breaks and the substitution of one group for another is completed. The iodide nucleophile has unshared pairs of electrons which are donated. The  $\sigma^*$  orbital of the C–Br bond is the acceptor orbital. In any donor–acceptor interaction which leads to a chemical change, identification of both the donor and acceptor is very crucial in predicting what change will occur and determining how the change might occur. Addition of Grignard reagents to carbonyl groups involves donation of the electrons of an electron-rich carbon-metal  $\sigma$  bond to the  $\pi^*$  orbital of the carbonyl group. As shown below, carbon-carbon bond formation is accompanied by carbon-oxygen  $\pi$  bond cleavage, oxygen-metal bond formation, and a corresponding change in geometry from trigonal to tetrahedral.



Ring opening of epoxides by alkoxides is used to emphasize that charge must be conserved during each mechanistic step. Because the reactants as written (neglecting spectator ions) have a net negative charge, the products *must* have a net negative charge.



The conservation of charge is a fundamental law for all processes, such as the addition of nucleophiles to  $\pi$  systems or acid-base reactions. The first step of the basic hydrolysis of nitriles has the hydroxide ion adding to the  $\pi$  bond of the nitrile. For the purposes of mechanistic discussion, the hydroxide is shown without its counterion and the net charge on the reactant side of the equation is -1. Consequently, the product of this first step (and each subsequent step) must also have a net negative charge.



In aqueous solution, proton transfer to the first formed intermediate is *very* rapid. However, again for illustrating the stepwise changes that must occur on the way from reactants to products using curved-arrow notation, these steps are shown independently.

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#### 78 CURVED-ARROW NOTATION

Similarly, the production of enolates from carbonyl compounds involves base removal of a proton from the  $\alpha$  position. The enolate is negatively charged and has delocalized electrons.



Although this process can be written to give a single canonical form  $\mathbf{A}$ , it must be realized that the enolate is a delocalized species and resonance forms  $\mathbf{A}$  and  $\mathbf{B}$  can be generated as discussed previously using curved-arrow notation. This is *not* a mechanistic step since the delocalized product is a resonance hybrid of  $\mathbf{A}$  and  $\mathbf{B}$ —that is  $\mathbf{A}$  is not converted to  $\mathbf{B}$ , but rather the curved arrows merely indicate the changes in electron distribution that must be used to describe the canonical form  $\mathbf{B}$ .

These previous examples are reactions where the electron donor (nucleophile) supplies the "electronic push" to accomplish bond breaking. Many nucleophiles, either neutral or anionic, have lone pairs of electrons that are easily donated. They can be donated to even weak electron acceptors (electrophiles).

There are other reactions, also easily describable by electron movement (curved arrows), in which  $\pi$  electrons are donated. In such reactions the  $\pi$  electrons are bonded electrons and hence the  $\pi$ -donor nucleophile is a weak electron donor. Consequently, a much stronger electron acceptor (stronger electrophile) is required for the "electronic pull" for electron donation to occur successfully. However, such descriptions are simply a matter of semantics because curved-arrow notation only shows changes in electrons, it does not indicate driving force. For a donor–acceptor interaction to occur productively, there has to be an energetic driving force for the process, and the energy levels of the donor and acceptor must be matched so that electron movement from the donor to the acceptor can occur.

For example, the protonation of a double bond has a proton as the electrophile and the  $\pi$  bond as the electron donor.



When bonded electrons are donated, it is important to remember that one of the bonded atoms which shared that pair is now left without a valence octet. That is, removal of a bonded pair must result in a sextet atom. Such is the case after protonation of a double bond as seen above. Because of the instability of a sextet electronic configuration, several strategies are available to stabilize it.

Neighboring atoms having unshared pairs of electrons can undergo bridging interaction with the cationic center to give structures in which all atoms have valence octets. An archetypical example is the bromination of olefins. Electrophilic addition of bromine to the double bond is predicted to give an  $\alpha$ -bromocarbocation. However, formation of the bridged bromonium ion avoids a sextet configuration of carbon and thus is formed preferentially. Bridging interactions occur when a carbocation is generated vicinal to substituents such as -OR, -Cl, -F, -SR,  $-NR_2$ , and so on, all of which have lone pairs capable of bridging interactions.



Resonance stabilization can also make  $\pi$ -electron donation much more effective by avoiding the formation of a sextet carbocation. Lone-pair donation from the oxygen of enol derivatives is very important to the good donor ability of these compounds. The resulting oxonium ion has all valence octets (although positively charged) and is thus stabilized over sextet canonical forms.



Resonance stabilization is important in electrophilic aromatic substitution as well. While each of the canonical forms of the Wheland intermediate has a sextet carbon atom, the charge is distributed over the remaining five atoms of the ring by resonance and is thus greatly stabilized.



The reactions of nucleophiles with electrophiles also relates to the overall oxidative change of a reaction. As is expected, nucleophilic atoms which are

#### 80 CURVED-ARROW NOTATION

more electronegative than carbon are not reductants and usually give no change in oxidation state, for example,



Conversely nucleophiles which are carbanion or hydride equivalents are reductants,



Carbocation or proton electrophiles give no change in oxidation level whereas electrophiles which are electronegative elements ( $Br_2$ ,  $Cl_2$ , NBS, peracids, etc.) are oxidants,



Besides intermolecular reactions, curved-arrow notation is also useful in indicating bonding changes in intramolecular reactions and rearrangement. For example, Cope-type rearrangements are seen to involve changes in three pairs of bonded electrons.



The arrows can be written in either directional sense since these reactions are concerted rearrangements with all bond making-bond breaking taking place at the same time. This example emphasizes the fact that curved-arrow notation is merely an electron bookkeeping method.

Cationic rearrangements are also handled easily by keeping track of where electron pairs come from and where they go. For example, the neophyl-type rearrangement below leads to skeletally rearranged products.



The curved-arrow notation clearly shows the electron flow needed to effect the rearrangement. What curved-arrow notation does not show is the timing of these events—that is, whether loss of a leaving group precedes or is concerted with 1,2-phenyl migration or if a bridged ion is an intermediate. Such considerations, if known, can be included in more detailed mechanistic sequences.



With these considerations, then, the steps one goes through to use electron movement to generate a possible reaction mechanism are as follows:

1. Write a balanced equation for the reaction. While spectator ions may be neglected, it is imperative to write correct Lewis structures for reactants and products. This step is very important but often neglected.

# 82 CURVED-ARROW NOTATION

- 2. Note the connectivity changes that occur, changes in oxidation level that occur, and the reagents or reactant types necessary for the conversion.
- 3. Write a stepwise process for the reaction using curved arrows to account for bonding changes. The use of curved arrows for electron movement should be guided by bond polarities, donor-acceptor properties, electronegativities, and structural factors and should result in a reasonable series of bonding changes from reactant to product.
- 4. Evaluate intermediates for stability and valence. If they fit normal chemical expectations, then the mechanism is potentially correct. There may be other mechanisms operating or the timing of individual steps (synchronous, concerted, etc.) may be different, but the above process can be used to generate them as well.

Thus we see that, used properly, curved-arrow notation for electron movement is indispensable to the organic chemist as a way to depict chemical change in complex molecules. Furthermore, it can be extended to include a method for showing the mechanism if the ground rules are understood and followed carefully.

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- An excellent discussion with many examples is found in F. M. Menger and L. Mandell, *Electronic Interpretation of Organic Chemistry*, Plenum, New York, 1980, pp. 47–116.

### PROBLEMS

**4.1.** For the following reactions, show the complete structures of the reactants and products of the step shown, point out the bonds which have been made and/or broken, identify the electron donors and acceptors, and use curved-arrow notation to indicate electron flow.

(a) 
$$CH_3 - S^{\ominus} + CH_2CH_3 \longrightarrow OTs$$



**4.2.** For the following transformations, give an acceptable mechanism as indicated by electron movement:



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84 CURVED-ARROW NOTATION



**4.3.** Using curved-arrow notation show how to derive three principal resonance structures for the following:



PROBLEMS 85



# 5

# MECHANISMS OF ORGANIC REACTIONS

Activation Energy	87
Activated Complex	88
Reaction Energetics	89
Structure of the Activated Complex	91
Hammond Postulate	96
Reaction Kinetics	99
Determining Activation Energies	104
Isotope Effects	105
Electronic Effects	110
Hammett Equation	111
Bibliography	118
Problems	118

A very common way for students to try to learn organic chemistry is to commit a vast amount of material to memory and then try to retrieve the correct information when it is needed. The organization of organic transformations around the chemistry of functional groups is an important tool in this learning process. Thus selected functional group transformations provide the initial body of knowledge that must be mastered. This is why organic chemistry students typically carry around piles of study cards, each with a reaction written on it that is to be committed to memory. This time-honored exercise allows students to learn *what* happens during a particular functional group reaction. Even with the

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simplifications of the functional group approach, there are still large numbers of reactions to deal with. The task is to learn them and, at the same time, organize them into coherent patterns that allow them to be retrieved when needed.

As a consequence beginning students typically concentrate on the beginning and end of a chemical reaction, that is, what reactants are converted to what products. There is also the need to learn what reagents cause these conversions. Such knowledge is an indispensable first step in the study of organic chemistry. In addition to learning *what* happens, however, it is also very interesting and important to understand *how* and *why* these transformations proceed.

The objective of this enlightenment is to expand our knowledge base so that it is possible to make reasonable predictions about what might happen in a reaction for which there is little precedence. Furthermore, it allows us to make educated guesses as to how a reaction might respond to changes in the structure of the starting material, or reagents, or reaction conditions. This jump in logic, from learning what happens during a reaction to predicting what will happen in a new reaction, requires a very clear understanding of mechanistic principles and the means by which they can be determined.

As we have seen, the mechanism of a reaction is the stepwise process by which reactants are converted to products. Moreover most steps in a reaction mechanism involve the movement and redistribution of electrons in the reactants or intermediates until the electronic configuration of the product is obtained. The electronic changes which are often depicted by curved-arrow notation result in bond making and/or bond breaking needed to get from the reactant to the product.

# **ACTIVATION ENERGY**

The first and most important fact about any chemical reaction or any elementary step in a chemical process is that there is an energy barrier separating the products from the reactants (Figure 5.1). For the reactants to be converted to products, the



Reaction coordinate (r)

Figure 5.1 Energy changes during a chemical reaction.

#### 88 MECHANISMS OF ORGANIC REACTIONS

energy of the system must be raised until the energy of the reactant(s) is equal to the height of energy barrier. Then and only then can the reactants change into products by passing over the barrier and relaxing into the product state. The energy barrier is of prime importance because (a) the height of the barrier determines how fast reactants can be converted to products at a given temperature and (b) at the maximum height of the energy barrier, which is called the transition state, there is a collection of atoms, called the activated complex, that is the *structural bridge* between reactants and products.

### ACTIVATED COMPLEX

Both of the considerations given above are related because the activated complex is an unstable and high-energy collection of atoms which has bonds and electron distributions distorted relative to those in both the reactants and products. The structure of the activated complex is determined by the bonding and structural changes that must occur in order for the reactants to be converted to products. Moreover the more the structure of the activated complex is distorted from normal geometries and bond distances, the higher is its energy. Consequently more energy must be added to the system to distort the reactants to the structure of the activated complex.

As a structural bridge between reactants and products, the activated complex has structural features of both. For example, consider a very simple reaction, the ionization of a tertiary bromide to a carbocation and a bromide ion in methanol. The reactant is a fully covalent molecule with a complete, yet polarized, bond between the tertiary carbon and the bromine atom.



The products contain a trivalent, positively charged carbon atom surrounded by a solvent shell and a bromide ion which carries a negative charge and is surrounded and hydrogen bonded to a shell of solvent molecules. The distance between the carbon and the bromine atom in the products is much greater than in the reactants. The bond between carbon and bromine is broken as the bromide moves away from the carbon and the pair of bonding electrons ends up in the valence shell of the bromide ion. We can depict this reaction using curved-arrow notation which tracks electron movement.



However, between reactants and products is an activated complex which must have structural features common to both. Qualitatively the activated complex  $(\ddagger)$  can be pictured as a structure in which the bromine has started to move away from carbon, lengthening the C–Br bond.



It has begun to develop a partial negative charge since the bonded pair of electrons is even more displaced onto the bromine. For the same reason, the tertiary carbon has begun to develop partial positive charge. The bond connecting carbon and bromine is longer and weaker than in the reactant, but there is still some bonding energy between carbon and bromine that is completely absent in the products. In addition, due to the greater amount of negative charge on bromine in the activated complex than in the reactant, the solvent starts to organize around the bromine, but the interaction is not nearly so strong as in the bromide ion product. The solvation of the carbocation product is much weaker than the solvation of the bromide in a hydroxylic solvent and is often disregarded; however, to the extent that the carbocation product is solvated, the solvent will begin to solvate the partially charged carbon in the activated complex.

The foregoing has been a rather microscopic description of the activated complex (perhaps too detailed), but the idea is quite clear. At whatever level the activated complex is described, it must contain structural features of both the reactants and the products it connects; that is, it can only be described in terms of the stable structures of reactants and products.

This then is the dilemma. The activated complex occurs at the transition state and has a vanishingly short lifetime ( $\sim 1$  bond vibration or  $\sim 10^{-13}$  sec) yet its structure and energy must be described and changes in its structure and energy must be evaluated if we are to compare different reactants in a predictive way.

### **REACTION ENERGETICS**

To attack this problem, we have to begin with those species whose structures we know—the reactants and products (Figure 5.2). Each is a stable collection of atoms characterized by a free energy. The difference in free energy between the reactants and products is  $\Delta G$ —the free energy of reaction which indicates whether the reaction takes place with evolution of energy (exergic) or with the uptake of energy (endergic). Connecting reactants and products is an energy profile which contains the activation barrier  $\Delta G^{\ddagger}$  which must be surmounted if the reaction is to proceed.

#### 90 MECHANISMS OF ORGANIC REACTIONS



Figure 5.2 Free energy changes going from reactants to products.



Figure 5.3 Free energy changes during a multistep reaction.

Now, for a given chemical transformation, there may be more than one step. In such cases each step of the transformation can be thought of as a discrete reaction and will have reactants and products and an activation barrier. Furthermore, each minimum on the energy curve between reactants and products is a more or less stable collection of atoms with a finite lifetime. These species are usually higher in energy than either the reactants or products and are called reaction intermediates. We have encountered many common reaction intermediates in reactions we have studied. These include carbocations, carbanions, enolates, free radicals, carbenes, and so on.

A multistep process involving intermediates still has an overall free energy of reaction ( $\Delta G$ ) (reactants vs. products), but each elementary step is a separate

#### STRUCTURE OF THE ACTIVATED COMPLEX 91



**Figure 5.4** Energy profile of a multistep transformation showing the rate determining step. Intermediate  $I_1$  is the reactant in the rate determining step and intermediate  $I_2$  is the product.

chemical reaction and therefore each step has a free energy, an activation barrier  $(\Delta G^{\pm})$ , a transition state, and an activated complex corresponding to the chemical changes occurring in that step of the process (Figure 5.3). The step with the transition state of highest free energy is called the rate-determining step because only by passing over the highest barrier can reactants proceed to products.

The rate-determining step thus defines how fast reactants can be converted to products (Figure 5.4). Steps in the process before the rate-determining step contribute to the rate law for the reaction by contributing to the overall barrier height. Steps after the rate-determining step have no bearing on the rate of reaction.

# STRUCTURE OF THE ACTIVATED COMPLEX

Even multistep processes can be simplified to a consideration of the energy and structure of the activated complex of the rate-determining step. One very common and successful approach for describing the structure of the activated complex is based on the notion that it must have structural features of both the reactant and product. Moreover it is generally assumed that the energy curve in the vicinity of the transition state is reasonably symmetrical; thus the relative energies of the reactants and products give some indication of the structure of the activated complex. This is a very important concept since it relates energy terms to structural characteristics. Free-energy changes associated with a chemical reaction can be measured fairly easily, whereas structural information about the activated complex cannot be directly measured at all. Thus the connection between energy and structure is unique in providing insight into the structure of the activated complex.

In a typical reaction coordinate diagram which describes energy changes as the reactants progress to products, the ordinate is calibrated in energy units and the abscissa (reaction coordinate) describes structural changes that occur on going

# 92 MECHANISMS OF ORGANIC REACTIONS



Figure 5.5 Reaction coordinate diagram for the ionization of a tertiary bromide.

from reactant to product. The reaction coordinate (r) is an arbitrary axis which cannot easily be defined in simple units but corresponds to structural changes in many dimensions. These structural changes must, however, all be related to the *differences* in structure between reactants and products. Recalling the earlier example of the ionization of a tertiary bromide to a carbocation, it is seen that the reaction coordinate corresponds to several distinct types of changes—breaking of the C–Br bond, flattening of the carbocationic carbon, charge development on carbon and bromine, and change in the solvent shell around the bromide (Figure 5.5). Thus the reaction coordinate cannot represent a single type of change, except by extreme oversimplification.

The connection between the energy and structure of the activated complex can be illustrated in the following way. If the energy barrier connecting reactants and products is symmetric and the reactants and products are of equal energy, then it is easy to see that the transition state will lie halfway along the reaction coordinate, and thus the structure of the activated complex is "halfway" between reactants and products. To illustrate this situation, consider the substitution of one iodide for another in the reaction of methyl iodide with iodide ion. (We can be sure this reaction occurs by using radioactive iodide.)

$$^{*}I^{-} + CH_{3} - I \longrightarrow ^{*}I - CH_{3} + I^{-}$$

The products are identical to the reactants and thus have the same energies (Figure 5.6). Consequently this substitution reaction has a transition state which falls midway between the reactants and products along the reaction coordinate. Thus the activated complex has a structure midway between that of reactants and products (Figure 5.7). This corresponds to a collection of atoms having a pentavalent carbon with trigonal bipyramidal geometry (the carbon is half inverted in geometry) with a half bond between each iodine and carbon and half of a full

#### STRUCTURE OF THE ACTIVATED COMPLEX 93



Figure 5.6 Reaction coordinate diagram for a reaction in which the reactants and products are of equal energy.

negative charge on each iodine. As shown below, this activated complex is a simplified but reasonable depiction of the structure of the activated complex.



If the products are lower in energy than the reactants in a reaction step, the symmetry of the activation barrier causes the transition to lie less than halfway along the reaction coordinate (Figure 5.8). The structure of the activated complex is more closely related to the structure of the reactants than to the structure of the products.



Figure 5.7 Reaction coordinate diagram for the reaction of methyl iodide with iodide ion.

#### 94 MECHANISMS OF ORGANIC REACTIONS



Figure 5.8 Reaction coordinate diagram for an exergic reaction in which the products are lower in energy than the reactants.

For example, the reaction of methyl triflate with cyanide ion is also a bimolecular substitution reaction, and it is a very exothermic process. Thus the transition state lies more toward the reactants (Figure 5.9), and the structure of the activated complex will be more like the structure of the reactant than the structure of the products. Thus in the activated complex the cyanide–methyl bond will be little formed ( $<\frac{1}{2}$ ) and the carbon–trifloxy bond will be largely intact ( $>\frac{1}{2}$ ). Most of the negative charge will remain on the cyanide nucleophile and little will have developed on the triflate leaving group. The geometry at the carbon will still be tetrahedral-like, although some flattening will have occurred.



For a reaction step in which the products are less stable than the reactants (endergic), the transition state will lie farther along the reaction coordinate toward the products and the activated complex will have a structure more similar to the products than the reactants (Figure 5.10).

If we consider reaction of mesylate ion with methyl bromide, we find that this is an endergic reaction; thus the transition state lies along the reaction coordinate farther toward the products than the reactants (Figure 5.11). The activated complex will therefore have a structure more resembling the products. There will be significant carbon–oxygen bond formation between the mesylate group and carbon and only a weak residual bond between carbon and bromine. The bromine

#### STRUCTURE OF THE ACTIVATED COMPLEX 95



Figure 5.9 Reaction coordinate diagram for the reaction of methyl triflate with cyanide ion.



**Figure 5.10** Reaction coordinate diagram for an endergic reaction in which the products are of higher energy than the reactants.

will have acquired significant negative charge and the carbon will be partially inverted in geometry.



It is possible, therefore, to gain significant insight into the structural characteristics of the activated complex from the structures of the reactants and products


Figure 5.11 Reaction coordinate diagram for the reaction of methyl bromide with mesylate ion.

and their relative energies. For an exothermic reaction step, the activated complex more resembles the reactants and is described as early. For an endothermic reaction step the activated complex more resembles the products and is described as late. The more exothermic is a process, the earlier is the transition state, while the more endothermic is a process, the later is the transition state.

These connections between energy and structure provide a powerful method for characterizing the activated complex. Furthermore changes made in reactants and products produce energy changes which can be translated into changes in structure of the activated complex; hence it is possible to predict how changes in structure will influence changes in energy and rates of reaction. These considerations are central to understanding the mechanisms of chemical reactions, and they permit us to make the best structural and reagent choices for a particular conversion.

#### HAMMOND POSTULATE

The relationship between energy of the transition state and the structure of the activated complex is summarized by the Hammond postulate, which states that the structure of the activated complex for any reaction step is most similar to the species (reactant or product) to which it is most similar in energy. As seen previously in Figure 5.8, exergic reactions (where the reactants are higher in energy than the products) have early transition states and activated complexes that resemble the reactants. Moreover, endergic reactions (in which the products are higher in energy than the reactants) have late transition states and activated complexes that resemble the products, as in Figure 5.10.

A most useful application of the Hammond postulate involves reactions which proceed by the formation of unstable intermediates, such as the carbocations,

#### HAMMOND POSTULATE 97



**Figure 5.12** Factors which lower the energy of a product lower the energy of the transition state leading to that product.

carbanions, carbenes, free radicals, and so on. The rate-determining step of such reactions is necessarily endothermic, and the Hammond postulate serves as a useful tool for identifying structural characteristics of the activated complex leading to that intermediate. The logical next step is to ask how structural features in reactants change the structure and thus energy of the activated complex.

The Hammond postulate states that in endergic reactions, features which stabilize and thus lower the energy of a product lower the energy of the transition state leading to that product. This is shown in Figure 5.12. If product 2 (P₂) is lower in energy than product 1 (P₁), then transition state 2 ( $\pm_2$ ) will be lower than transition state 1 ( $\pm_1$ ). It will also be earlier. As a consequence, P₂ will have a lower activation barrier and be formed faster than P₁. A simplified restatement of the Hammond postulate is that more stable products are formed faster. It must be remembered that this analysis is for endothermic reactions and assumes that the reactants have the same or similar energies.

The ionization of alkyl tosylates to give carbocations is an endothermic reaction. Knowing that  $3^{\circ}$  carbocations are more stable than  $2^{\circ}$  carbocations, we would conclude that the activation barrier for ionization of the  $3^{\circ}$  tosylate is lower than that of the  $2^{\circ}$  tosylate, and thus the  $3^{\circ}$  tosylate should ionize faster (Figure 5.13).

We would also predict that the transition state for ionization of the  $3^{\circ}$  tosylate would be earlier, so there should be less C–O bond breaking and less charge development than in the activated complex for ionization of the  $2^{\circ}$  tosylate.

The Hammond postulate provides a key relationship between the rate of reaction and the activated complex of that reaction (Figure 5.14). In practice, structural changes are made in the reactant(s) and the influence of those changes on the rate of reaction is measured. If the reaction is faster, then the change in the



Figure 5.13 Transition state energy differences for the ionization of 2° and 3° tosylates.



**Figure 5.14** Relationship between product energy changes and transition states for endergic reactions.

reactant has led to a lower product energy and hence a lower activation energy and an earlier transition state (one which has more reactant character). If the reaction is slower, then the change in the reactant has led to a higher product energy and hence a higher activation energy and a later transition state (one with more product character). The results of rate studies can thus be translated into structural changes (bonding, charge distribution, geometry) in the activated complex, which further translates into the mechanistic information about the reaction.

The Hammond postulate is best applied to reactions with unstable intermediates, such as the carbocations in the above example. In such cases the transition state is late and the activated complex more resembles the intermediate. Thus changes in the energy of the intermediate have the greatest effect on the energy of the transition state and thus the rate of the reaction.

However, the Hammond postulate also holds for exergic reactions where the transition state is early and the activated complex more resembles the reactant. For such reactions changes in the energy of the reactants have the greatest effect on the energy of the transition state and thus the rate of the reaction.

#### **REACTION KINETICS**

The first step in delineating the mechanism of a reaction is to determine what reactant species must come together to produce the activated complex of the rate-determining step. This can be done by determining the order of the reaction with respect to each of the reactants in the process. If the rate of a reaction is dependent on the concentration of a particular reactant, then that reactant is involved in the transition state of the rate-determining step of that reaction. This provides important structural information about the activated complex since it reveals which chemical species are present in the activated complex.

The order of a reaction is found by determining the relationship between rate and concentration for each reactant. Thus for the elementary process  $A \rightarrow B$ , the rate of reaction  $\nu$  can be expressed as the decrease in the concentration of reactant A with time or the increase in the concentration of product B with time:

$$v = \frac{-d[\mathbf{A}]}{dt} = +\frac{d[\mathbf{B}]}{dt}$$

Further, the rate of the reaction can be expressed as a function of the concentration of A, where k is the rate constant for the process:

$$\frac{-d\mathbf{A}}{dt} = +\frac{d[\mathbf{B}]}{dt} = k[\mathbf{A}]$$

This differential rate expression shows that the reaction rate is directly dependent on the concentration of A—the greater is [A], the faster is A converted to B. The reaction is said to be first order with respect to A because the exponent of [A] is 1.

The above expression is the first-order differential rate law for the conversion of A to B. The change in concentration of A over the complete course of the reaction is given by the integrated rate law, which is found by solving the differential rate law:

$$-\frac{dA}{dt} = k[A]$$
$$\frac{d[A]}{[A]} = k \ dt$$
$$-\ln\frac{[A_0]}{[A]_t} = kt \quad \Rightarrow \ln[A]_t = -kt + \ln A_0$$

The integrated rate law shows that the natural logarithm of the concentration of the starting material A decreases linearly with time. By determining the concentration of A at various times  $[A_t]$  and plotting  $\ln[A]_t$  versus *t*, a straight line with slope -k will be obtained *if the reaction is first order in A*. (Since the concentration of product at any time  $[B_t] = [A_0] - [A_t]$ , a plot of the increase in the concentration of B with time in the form of  $\ln([A_0] - [B_t])$  versus time would also give a straight line whose slope is +k.) If such linear dependence is observed, then the reaction is first order with respect to A and the rate constant for the reaction can be determined. If the rate plot is not linear, then the reaction is not first order with respect to A; that is, a first-order rate law does not correctly describe the kinetic behavior.

Second-order reactions occur when two reagents must collide in solution to produce the activated complex. Thus for the reactions

$$2A \longrightarrow B \qquad A + B \longrightarrow C$$

each reaction is second order because the sum of the exponents of species in the rate law is 2:

$$-\frac{dA}{dt} = k[A]^2 \qquad \frac{dC}{dt} = k[A][B]$$

This means that in the first instance two molecules of A must collide to produce the activated complex. In the second case a molecule of A and B collide to produce the activated complex. In each case the second-order dependence requires that *both* of the colliding molecules are a part of the activated complex of the rate-determining step.

Integration of these differential rate laws gives

$$\frac{1}{A_t} - \frac{1}{A_0} = kt \qquad \text{and} \qquad \frac{1}{A_0 - B_0} \ln \left[ \frac{B_0 A_t}{A_0 B_t} \right] = kt$$

Again plotting concentration versus time using these integrated second-order rate laws gives linear plots *only if the reaction is a second-order process*. The rate constants can be determined from the slopes. If the concentration-time plots are not linear, then the second-order rate equations do not correctly describe the kinetic behavior. There are integrated rate laws for many different reaction orders.

It is often possible to simplify the rate law of a second-order process by employing pseudo-first-order conditions. For a second-order reaction where

$$A + B \longrightarrow P$$

and

$$\frac{-dA}{dt} = \frac{dP}{dt} = k[A][B]$$

the integrated rate law for a second-order reaction can be used to plot the kinetic data. Alternatively, if the concentration of one of the reactants, for example A, is much larger than the other (10-fold excess minimum, 20-fold better), then its concentration does not change significantly over the course of the reaction; thus  $[A]_t \approx [A]_0$ . The differential rate law can be approximated as

$$\frac{d\mathbf{P}}{dt} = k[\mathbf{A}][\mathbf{B}] = k[\mathbf{A}_0][\mathbf{B}] = k'[\mathbf{B}]$$

where  $k' = k[A]_0$ . This new rate law is a first-order equation which is much easier to deal with and plot. A plot of ln[B] versus *t* will give a straight line of slope *k'*. From *k'* and [A]₀ (which is known), the rate constant can easily be determined. This is a much simpler method for determining the rate constant than the normal second-order treatment. The order of the reaction with respect to A can be checked by using several different initial concentrations of [A]₀ and the relationship

$$k' = k[\mathbf{A}]_0$$

Therefore

$$\log k' = \log[A]_0 + \log k$$

Plots of log k' versus log[A]₀ will have unit slope if the reaction is first order in A. Alternatively, doubling [A]₀ should double the rate if the reaction is first order in A.

In a multistep process involving a reactive intermediate, the rate law for the overall reaction cannot be written down a priori because the step in which the reactants disappear is different than the step in which the products are formed (Figure 5.15). In a large number of cases, the intermediate is of high energy and reacts very rapidly—either returning to reactant or going on to product.

In such cases the steady-state approximation can be used to derive a rate expression that can be tested. Thus for a reaction process involving an intermediate [I]

$$A \stackrel{k_1}{\underset{k_{-1}}{\longrightarrow}} I \stackrel{k_2}{\longrightarrow} P$$

and

$$v = \frac{d\mathbf{P}}{dt} = k_2[\mathbf{I}]$$

The concentration of the intermediate [I] which gives product is given by the difference between the rate of its formation from A and the rate of its conversion back to reactant A or forward to product P

$$\frac{d\mathbf{I}}{dt} = k_1[\mathbf{A}] - k_{-1}[\mathbf{I}] - k_2[\mathbf{I}] = 0$$



Figure 5.15 Reaction coordinate diagram for a multistep process involving a reactive intermediate.

The steady-state approximation assumes that since I is very reactive, its concentration will be very low at any time during the reaction and it will not change appreciably. Therefore dI/dt = 0. Solving the above expression for the concentration of I and substitution into the rate law for the formation product gives

$$[I] = \frac{k_1[A]}{k_{-1} + k_2}$$

and

$$\frac{d\mathbf{P}}{dt} = \frac{k_2 k_1}{k_{-1} + k_2} [\mathbf{A}] = k_{\text{obs}} [\mathbf{A}]$$

where the observed rate constant  $k_{obs} = k_2 k_1 / k_{-1} + k_2$ . This is a first-order rate expression in reactant A and can be integrated and plotted normally.

Several limiting cases can be envisioned for such a multistep process. If  $k_2 \gg k_{-1}$  (the intermediate goes on to product more rapidly than it returns to reactant),

$$\frac{d\mathbf{P}}{dt} = \frac{k_2 k_1}{k_2 + k_{-1}} [\mathbf{A}]$$

simplifies to

$$\frac{d\mathbf{P}}{dt} = k_1[\mathbf{A}] \quad \text{for } k_2 \gg k_{-1}$$

This represents the case where the first step is rate determining. (If this situation is known beforehand, one need not work through the steady-state approximation but merely write down  $dP/dt = k_1A$  since the first step is rate limiting and irreversible.)

REACTION KINETICS 103

If  $k_{-1} \gg k_2$ , then

$$\frac{d\mathbf{P}}{dt} = \frac{k_1}{k_{-1}}k_2[\mathbf{A}] = K_{\rm eq}k_2[\mathbf{A}]$$

This represents the case where there is a fast preequilibrium preceding the rate-determining step. It is again not necessary to work through the steady-state approximation if this situation is known to exist. If the intermediate I is in equilibrium with the reactant A, then

$$K_{\text{eq}} = \frac{[\mathbf{I}]}{[\mathbf{A}]}$$
 and  $[\mathbf{I}] = K_{\text{eq}}[\mathbf{A}]$ 

substitution for [I] in the rate expression gives

$$\frac{d\mathbf{P}}{dt} = k_2[\mathbf{I}] = K_{\rm eq}k_2[\mathbf{A}]$$

This is actually the correct way to think about this case since the steady-state approximation requires that the concentration of I is low, which it may not be if there is a fast preequilibrium.

If  $k_2 \approx k_{-1}$ , then the full steady-state rate expression is needed to describe the rate of reaction:

$$\frac{d\mathbf{P}}{dt} = \frac{k_2 k_1}{k_2 + k_{-1}} [\mathbf{A}]$$

Consider once again the solvolysis of a tertiary bromide in methanol.



This reaction can be reduced to the kinetic scheme.

$$A \underset{k_{-1}}{\overset{k_1}{\longleftarrow}} I + Br \overset{\bigoplus}{\longrightarrow} P$$

The rate of product formation is given by the pseudo first-order expression in which  $k_2 = k$ [CH₃OH] (normally  $k_2$  is taken as the rate constant because the concentration of methanol is constant),

$$\frac{d\mathbf{P}}{dt} = k_2[\mathbf{I}]$$

and the steady-state approximation is written as

$$\frac{dP}{dt} = O = k_1[A] - k_{-1}[I][Br^-] - k_2[I]$$

Since the return of intermediate I to reactants is a second-order reaction, solving for [I] and substitution into the rate law give

$$\frac{d\mathbf{P}}{dt} = \frac{k_2 k_1 [\mathbf{A}]}{k_{-1} [\mathbf{B}\mathbf{r}^-] + k_2}$$

If  $k_2 \gg k_{-1}[Br^-]$ , then  $\nu = k_1[A]$  and simple first-order behavior is found. If  $k_{-1}[Br] \gg k_2$ , then there is a rapid ionization preequilibrium and

$$K_{\text{eq}} = \frac{[\mathrm{I}][\mathrm{Br}^-]}{[\mathrm{A}]}$$
 and  $[\mathrm{I}] = \frac{K_{\text{eq}}[\mathrm{A}]}{[\mathrm{Br}^-]}$ 

Then

$$\frac{d\mathbf{P}}{dt} = k_2[\mathbf{I}] = \frac{k_2 K_{\rm eq}[\mathbf{A}]}{[\mathbf{Br}^-]}$$

The rate of product formation will be directly proportional to [A] but inversely proportional to [Br⁻]. By using an excess of Br⁻ so that its initial concentration [Br⁻]₀ does not change appreciably over the course of the reaction, pseudo-first-order behavior can be achieved with  $k_{obs} = k_2 K_{ea}/[Br^-]_0$ .

If  $k_{-1}[Br^-] \approx k_2$ , then the full rate expression will be needed to describe the kinetic behavior. The rate will be first order in [A] and the rate will slow down in the presence of added bromide but not in a simple inverse relationship.

#### DETERMINING ACTIVATION ENERGIES

By using the above methods, the rate constants for most organic reactions can be obtained. Rate constants, by virtue of the fact that they relate directly to the passage of reactants over the barrier of the rate-determining step, can be used to probe the energy and structure of the activated complex. The energy of the activated complex corresponds to the height of the activation barrier for the ratedetermining step. The barrier height can be calculated by the Arrhenius equation

$$\ln k_{\text{rate}} = -\frac{E_{\text{a}}}{RT}$$

where  $k_{\text{rate}}$  is the rate constant of a reaction, R is the ideal gas constant, and T is the absolute temperature. The Arrhenius activation energy  $E_a$  is determined from plots of ln k versus 1/T at various temperatures and largely corresponds to the enthalpy of activation since  $E_a = \Delta H^{\pm} + RT$ . The enthalpy and entropy

of activation and hence the free energy of activation are determined by the Eyring equation

$$\ln \frac{k_{\text{rate}}}{T} = -\frac{\Delta H^{\pm}}{RT} + \frac{\Delta S^{\pm}}{R} - \ln \frac{k}{h}$$

where k is Boltzmann's constant and h Planck's constant. Plots of  $\ln(k_{\text{rate}}/T)$  versus 1/T give straight lines with slope  $-\Delta H^{\pm}/R$  and intercept  $\Delta S^{\pm}/R - \ln(k/h)$ . Each can be numerically evaluated to give  $\Delta H^{\pm}$ ,  $\Delta S^{\pm}$ , and finally  $\Delta G^{\pm}$  by  $\Delta G^{\pm} = \Delta H^{\pm} - T \Delta S^{\pm}$ .

#### **ISOTOPE EFFECTS**

Besides the energy of the activated complex, structure and bonding in the activated complex can be probed in several other ways using rate constant data. One very powerful way to investigate bonding in the activated complex is to use kinetic isotope effects. Isotope effects derive from the fact that a heavier isotope of an element has a lower zero-point energy and hence more energy is required to break a bond to a heavier isotope than a bond to a lighter isotope (i.e., the activation energy is greater). At a given temperature this means that the rate of reaction for a compound containing a heavy isotope is slower than the rate of reaction for the compound with a lighter isotope. This is *only* true if breaking of that bond is involved at the transition state of the rate-determining step. If breaking of this bond occurs prior to or after the rate-determining step, isotopic substitution does not give a large change in the rate. This effect is most pronounced for hydrogen/deuterium, which has the largest mass difference of any isotopic pair and thus the largest difference in zero-point energies. If a bond to hydrogen (or deuterium) is being broken in the rate-determining step, then  $k_{\rm H}/k_{\rm D}$  values of 2–8 are typical. These are termed *primary* kinetic deuterium isotope effects.

If the C–H(D) bond is not being broken in the rate-determining step, there are sometimes smaller effects on the rate resulting from isotopic substitution that are termed *secondary* kinetic deuterium isotope effects. They result from zeropoint energy differences in deformation modes but they are small and typically  $k_{\rm H}/k_{\rm D}$  values are 1–1.3 for these effects. If a kinetic deuterium isotope effect is found to be greater than about 1.5, it is a primary kinetic deuterium isotope effect and C–H(D) bond breaking is occurring in the rate-determining step. If a kinetic deuterium isotope effect and C–H(D) bond breaking is not occurring in the rate-determining step.

The largest values of primary kinetic deuterium isotope effects are found for reactions where the bond to hydrogen is about one-half broken  $(k_{\rm H}/k_{\rm D})$  values are 6–8). Smaller values are found in reactions in which the bond to hydrogen is less than or more than one-half broken. Normally,  $k_{\rm H}/k_{\rm D}$  values less than maximum correspond to bond cleavage of  $<\frac{1}{2}$ . Primary kinetic deuterium



Figure 5.16 Transition states for the bromination of toluene and isopropylbenzene by NBS.

isotope effects thus provide insight into the extent of C-H bond cleavage in the activated complex.

For example, the free-radical bromination of toluene by *N*-bromosuccinimide (NBS) proceeds with  $k_{\rm H}/k_{\rm D} = 4.9$ , while for the same bromination of isopropyl benzene,  $k_{\rm H}/k_{\rm D} = 1.8$  (Figure 5.16). Both are primary kinetic deuterium isotope effects, indicating that hydrogen abstraction by a bromine atom is the rate-determining step. The much lower value of the isotope effect for isopropyl-benzene suggests that the transition state is much earlier than for toluene. The lesser extent of hydrogen transfer in isopropylbenzene is due to the more stable radical being produced, resulting in an earlier transition state.

Therefore



later transition state

earlier transition state

The electrophilic nitration of benzene using acetyl nitrate involves the replacement of a hydrogen on the benzene ring by a nitro group. The reaction is second order overall, first order in benzene, and first order in the nitrating agent— $v = k[C_6H_6]$ [acetyl nitrate].



Use of fully deuterated benzene gave  $k_{\rm H}/k_{\rm D} = 1$ . These data suggest that the nitrating agent attacks the benzene ring in the rate-determining step, but C–H

bond breaking is not involved in the rate-determining step. These observations are consistent with an electrophilic attack of the nitrating agent of the  $\pi$  system. The proton is lost in a subsequent fast step, after the rate-determining step.



Thus, even though loss of hydrogen is required for the product to be formed, its removal is not taking place in the rate-determining step of the reaction—it must take place after the rate-determining step.

The base-promoted bromination of ketones is a second-order process, first order in ketone and first order in base; thus v = k[ketone][base]. The bromine concentration does not appear in the rate law; that is, the reaction is zero order in [Br₂].

$$R_{CH_{2}} \stackrel{O}{\underset{CH_{3}}{\overset{[OH^{-}]}{\longrightarrow}}} R_{CH_{2}} \stackrel{O}{\underset{CH_{2}}{\overset{Br}{\longrightarrow}}} R_{H_{2}O + Br^{-}}$$

Use of deuterated substrates gives  $k_{\rm H}/k_{\rm D} = 6.5$ . This is a primary kinetic deuterium isotope effect, indicating that proton removal is an essential component of the rate-determining step. The lack of rate dependence on bromine requires that bromine is added to the molecule after the rate-determining step. A mechanism consistent with these facts has proton removal and enolate formation rate determining.

$$R \xrightarrow{O}_{CH_{2}} CH_{3} + OH^{-} \xrightarrow{k_{slow}} R \xrightarrow{O}_{CH_{2}} CH_{2} + H_{2}O$$

$$R \xrightarrow{O}_{CH_{2}} CH_{2} + Br_{2} \xrightarrow{k_{fast}} R \xrightarrow{O}_{CH_{2}} CH_{2} + Br^{-}$$

If we now take this basic scenario and add our notions of electron movement to the picture, we can construct a detailed picture of electronic change that is consistent with the observed facts. 108

MECHANISMS OF ORGANIC REACTIONS



The activated complex for proton removal, the rate-determining step, can be envisioned as having a partial charge from proton removal delocalized into the carbonyl group (as it is in the product enolate). This also requires that the proton being removed has a dihedral angle of 90° with the plane of the carbonyl group so that the developing charge can overlap with the carbonyl  $\pi$  bond.



Base-promoted elimination in the two  $\beta$ -phenethyltrimethylammonium derivatives shown below is found to be second order overall, first order in substrate, and first order in base; that is,  $\nu = k[C_6H_5CH_2CH_2N^+(CH_3)_3][CH_3CH_2O^-].$ 

This means that both the substrate and the ethoxide base are present in the transition state of the rate-determining step. The rate constants for the deuterated and protio substrates were measured. The magnitudes  $(k_{\rm H}/k_{\rm D} = 3-4)$  of the kinetic deuterium isotope effects for both substrates are typical primary kinetic deuterium isotope effects, which means that C–H bond breaking is involved at the transition state of the rate-determining step. This suggests that proton removal

by a base in the activated complex is an essential element of the rate-determining step and is a key feature in the mechanism of the elimination reaction.

The difference between the  $k_{\rm H}/k_{\rm D}$  values, however, means that the extent of C–H bond breaking at the transition state in the second substrate is different from the first. (The transition state of the second substrate is actually earlier in terms of proton removal by the base because in both cases proton transfer is greater than half completed.) Thus a change in structure of the substrate leads to a distinct change in the structure of the activated complex which can be detected and described by kinetic isotope effects.

From the above examples it is clear that kinetic deuterium isotope effects are a powerful way to probe bonding changes in the activated complex. The magnitude of the isotope effect indicates whether bonds to hydrogen are being made or broken in the rate-determining step. Differences in kinetic isotope effects in closely related precursors can also be used to pinpoint whether one transition state is earlier than another—a direct measure of the effect of the substrate structure on the structure of the transition state.

Other elements can be used to measure isotope effects; however, the magnitudes of these isotope effects are much smaller than primary kinetic deuterium isotope effects. Substitution of  ${}^{13}C$  for  ${}^{12}C$  in a reaction could lead to a *maximum* kinetic isotope effect of  $k{}^{12}C/k{}^{13}C = 1.05$  for a reaction in which a bond to carbon is broken in the rate-determining step. (Recall that maximum  $k_H/k_D$ 's are 8–10.) Most standard kinetic methods are not capable of distinguishing such small rate differences reproducibly, and so kinetic isotope effects for elements other than hydrogen (deuterium) are not very abundant in the literature. In some instances, isotopic abundances determined by mass spectrometry can be used to measure such differences accurately and isotope effects can be informative. The decarboxylation of malonic acid proceeds with  $k{}^{12}C/k{}^{13}C = 1.045$ . This large primary-isotope effect (for carbon) indicates that C–C bond breaking is well developed in the transition state. This detailed information about the structure of the activated complex permits a shift in focus from a curved-arrow type of mechanism to a real structure of the activated complex.



#### **ELECTRONIC EFFECTS**

Besides bond breaking, another common feature of many reactions is the formation of charged species as intermediates. Carbocations, carbanions, oxonium ions, and so on, are all commonly encountered intermediates formed in the ratedetermining step of multistep reactions. As a consequence, charge development in the activated complex is expected. In terms of the reaction mechanism, it is very important to know the charge type (positive, negative, or none) and the extent of charge development in the activated complex.

The use of rate constants can provide a clue to charge development as well. Changes in the rate constant of a reaction due to changes in structure can be indicative of the charge distributions present in the activated complex. For example, rate constants are much larger for the base-promoted deuterium exchange of phenylacetone than for acetone itself because the phenyl group stabilizes the negative charge on the enolate ion (and the transition state leading to it). Hence the  $\alpha$  proton is removed more rapidly and deuterium exchange is speeded up correspondingly. This behavior is entirely consistent with an increase of electron density on the  $\alpha$  carbon during the rate-determining step.



The hydration of styrene,  $\alpha$ -methylstyrene, and  $\alpha$ -trifluoromethylstyrene gives a benzylic alcohol product; however,  $\alpha$ -methylstyrene reacts  $10^5$  more rapidly than styrene itself, while  $\alpha$ -trifluoromethylstyrene reacts  $10^7$  less rapidly than styrene. This behavior is consistent with the rate-determining step being protonation of the double bond to give a carbocation. The developing positive charge in the transition state is stabilized by the inductive effect of the methyl group in  $\alpha$ -methylstyrene, the transition state is of lower energy, and it thus reacts faster. The developing positive charge in the transition state is destabilized by the electron-withdrawing inductive effect of the trifluoromethyl group in  $\alpha$ -trifluoromethylstyrene, the transition state is of higher energy, and thus it reacts more slowly.

While changes in rate constants in response to changes in structure are extremely valuable for indicating the type of charge development occurring in the activated complex, the actual extent of charge development in the activated complex is an additional structural descriptor that would be very useful.

#### HAMMETT EQUATION 111



#### HAMMETT EQUATION

The Hammett equation is a very useful tool for monitoring the extent of charge development in the activated complex. If a substituent is attached to the meta or para position of a benzene ring, it will change the ability of the aromatic ring to donate or withdraw electrons relative to benzene itself. Thus a p-methylphenyl group should be electron donating relative to phenyl while a p-bromophenyl group should be electron withdrawing relative to phenyl. Furthermore, placement of the bromo group in the meta position places it closer to the point of attachment of the phenyl ring so the electron-withdrawing ability of the m-bromophenyl group is greater than that of the p-bromophenyl group. Substituents are normally attached only to the meta or para positions of the phenyl so that they do not sterically interfere with the site of attachment of the phenyl group.



By the use of a model reaction (ionization of benzoic acids), the ability of a substituent to modify the electron-donating or electron-withdrawing ability of the phenyl group and thus influence that reaction can be defined quantitatively by the Hammett equation,

$$\log \frac{K_{\rm Z}}{K_{\rm H}} \equiv \sigma$$

where  $K_Z$  is the acidity constant for a benzoic acid with some substituent-Z attached and  $K_H$  is the acidity constant for benzoic acid itself.

The result is a substituent constant ( $\sigma$ ) which is a numerical description of the electronic effect of a substituent relative to a hydrogen atom on the model reaction. Stated a different way, a substituent constant  $\sigma$  is a quantitative way to

Substituent	$\sigma_m$	$\sigma_p$	$\sigma_m^+$	$\sigma_p^+$	$\sigma_p^-$
NH ₂	-0.09	-0.57	-0.16	-1.3	
OCH ₃	0.10	-0.28	0.05	-0.78	
CH ₃	-0.06	-0.14	-0.10	-0.31	
H (reference substituent)	0	0	0	0	0
Cl	0.37	0.24	0.40	0.11	
Br	0.37	0.26	0.41	0.15	
$CO_2R$	0.35	0.44	0.37	0.48	0.68
CF ₃	0.46	0.53	0.57		_
CN	0.62	0.70	0.56	0.66	1.00
NO ₂	0.71	0.81	0.73	0.79	1.27

Table 5.1  $\sigma$  Values for Common Aromatic Substituents

describe the electron-donating or electron-withdrawing properties of a substituent when it is attached to a benzene ring. Different  $\sigma$  values are obtained for a given substituent if it is in the meta or the para position so  $\sigma$  values are positionally dependent. Table 5.1 lists some common  $\sigma$  values.

The model reaction used to evaluate  $\sigma$  constants is an ionization equilibrium in which a negative charge is produced upon going from reactants to products. Electron-withdrawing groups have positive  $\sigma$  values because they increase the ionization relative to hydrogen,  $K_Z/K_H > 1$ , and  $\log(K_Z/K_H) > 0$ . Electrondonating substituents have negative  $\sigma$  values because they decrease ionization relative to hydrogen,  $K_Z/K_H < 1$ , and  $\log(K_Z/K_H) < 0$ . Hydrogen itself is treated as a substituent with  $\sigma = 0$  because  $K_Z = K_H$  and thus  $K_Z/K_H = 1$  and  $\log(K_Z/K_H) = 0$ .

Furthermore the absolute magnitude of the  $\sigma$  value provides a quantitative measure of the relative electron-donating or electron-withdrawing ability. Thus a *m*-CF₃ group ( $\sigma = 0.46$ ) is a stronger electron-withdrawing group than *m*-Cl ( $\sigma = 0.37$ ) but is a less strong electron-withdrawing group than *m*-CN ( $\sigma = 0.62$ ). On the other hand, a *p*-methyl group ( $\sigma = -0.14$ ) is a weaker electron donor than *p*-OCH₃ ( $\sigma = -0.28$ ) but a better electron donor than *p*-Si(CH₃)₃ ( $\sigma = -0.07$ )

These substituent constants can be used with rate data to evaluate the type and extent of charge development in the activated complex of the rate-determining step for a wide variety of chemical reactions. The rates of reaction for a particular transformation are measured using a series of compounds which differ only by the phenyl substituents present; for example,



#### HAMMETT EQUATION 113

Then the differences in rate caused by the electronic effect of the substituent are correlated by the Hammett equation  $\log(k_z/k_H) = \rho \sigma_z$ , where  $k_z$  is the rate constant obtained for a compound with a particular meta or para substituent,  $k_H$  is the rate constant for the unsubstituted phenyl group, and  $\sigma_z$  is the substituent constant for each substituent used. The proportionality constant  $\rho$  relates the substituent constant (electron donating or withdrawing) and the substituent's effect on rate. It gives information about the type and extent of charge development in the activated complex. It is determined by plotting  $\log(k_z/k_0)$  versus  $\sigma_z$  for a series of substituents. The slope of the linear plot is  $\rho$  and is termed the reaction constant. For example, the reaction shown above is an elimination reaction in which a proton and the nosylate group are eliminated and a C–N  $\pi$ bond is formed in their place. The reaction is second order overall, first order in substrate, and first order in base. The rate constants were measured for several substituted compounds:

Substituent	$k_{ m Z}$	$\sigma_{\rm Z}$
Н	$8.69 \times 10^{-3}$	0
p-CH ₃	$7.39 \times 10^{-3}$	-0.14
m-CH ₃	$7.94 \times 10^{-3}$	-0.06
p-Cl	$1.17 \times 10^{-2}$	0.24
m-Cl	$1.50 \times 10^{-2}$	0.37
m-CF ₃	$1.54 \times 10^{-2}$	0.46

These data were plotted according to the Hammett equation to give the plot in Figure 5.17. The first thing to note is that the Hammett plot is linear. The linearity of the plot implies that the substituent constants determined for the ionization of



Figure 5.17 Hammett plot of rates versus substituent constants for an imine-forming elimination.

benzoic acids ( $\sigma$ ) are correctly modeling electronic changes taking place in the reaction under consideration; that is, the influence of a substituent on the model reaction is of the same type as for the reaction under investigation.

Next the sign and absolute magnitude of the  $\rho$  value determined from a Hammett plot give information about charge development at the transition state. The sign of  $\rho$  tells whether a positive or negative charge is being developed in the activated complex relative to the reactants. A positive  $\rho$  value means that electron density is increased (negative charge is being produced) in the activated complex. A negative  $\rho$  value means that electron deficiency is being produced (often a positive charge) in the activated complex. Generally  $\rho$  values have absolute magnitudes between 0 and 3, but values as high as 10 or 12 are known. A value of  $\rho = 0$  means that substituents have no electronic effect on the reaction rate, and thus no charge is being developed at the transition state. Large absolute values of  $\rho$  mean that substituents influence the rate greatly, and thus the amount of charge developed in the activated complex is large and influenced significantly by the electronic properties of the substituents. For example,

$$C_{6}H_{5} \underbrace{\bigcup_{CH_{2}}^{O}}_{CH_{2}} \underbrace{\bigcup_{i}^{N}}_{CH_{3}} O \underbrace{\bigcup_{Ei_{3}N}^{Z}}_{CH_{3}} O \underbrace{\bigcup_{Ei_{3}N}^{O}}_{N \underbrace{CH_{3}}} O \underbrace{\bigcup_{i}^{O}}_{N \underbrace{CH_{3}}} O \underbrace{\bigcup_{i}^{O}}_{N \underbrace{CH_{3}}} O = +1.4$$

This  $\rho = 1.4$  means that negative charge is being developed on the arenesulfonate group at the transition state, consistent with this group departing as an anionic leaving group as part of the rate-determining step. The magnitude of 1.4 means that a significant amount of charge is developed on the leaving group. This interpretation is possible only by comparing  $\rho = +1.4$  with  $\rho$  values for sulfonate leaving groups in other reactions in which a range of normal values is known. The normal range is about +0.8 to +1.65; therefore the value of 1.4 indicates significant charge development at the transition state. In addition to the positive  $\rho$  value, it is also known that the reaction is second order overall, first order in substrate, and first order in triethylamine. Moreover, there is a primary kinetic deuterium isotope effect for the benzylic position. These observations allow a detailed structure to be drawn of the activated complex in which concerted 1,3 elimination results in formation of the three-membered ring product.



The reaction of N,N-dimethylbenzylamines with methyl iodide is found to have  $\rho = -1.0$ . The negative sign indicates that partial positive charge is being

produced in the transition state. The modest size of  $\rho$  is consistent with the charge being developed on the nitrogen, which is insulated from the aromatic ring by the saturated CH₂ group. The ability of substituents to influence rate is reduced by the insulating methylene group, and therefore  $\rho$  is smaller. In comparison the methylation of *N*,*N*-dimethylaniline has  $\rho = -3.3$  because the nitrogen atom, on which positive charge is developed in the transition state, is directly attached to the phenyl ring and the substituents have a greater influence on the stability of the charge being developed. Hence the magnitude of  $\rho$  is much larger.

$$Z \xrightarrow{CH_2 - N} CH_3 + CH_3 I \longrightarrow Z \xrightarrow{H_2 - N(CH_3)_3} \rho = -1.0$$

$$Z \xrightarrow{CH_3} + CH_3 I \longrightarrow Z \xrightarrow{H_2 - N(CH_3)_3} \rho = -3.3$$

For similar reactions, comparison of the  $\rho$  values can be used to determine which reaction has a greater charge development. Comparison of the olefinforming eliminations below reveals which reaction has greater charge development at the benzylic position and thus which has a greater degree of proton removal in the activated complex.

$$Z \longrightarrow CH_2 - CH_2 \xrightarrow{I} Eto^- Z \longrightarrow CH = CH_2 + EtOH + I^- \rho = +2.07$$

$$Z \longrightarrow CH_2 - CH_2 \xrightarrow{CI} Eto^- Z \longrightarrow CH = CH_2 + EtOH + CI^- \rho = +2.58$$

The larger value of  $\rho = 2.58$  for the chloride leaving group than for the iodide leaving group ( $\rho = 2.07$ ) suggests that greater negative charge is developed on the benzylic position when chloride is the leaving group than when iodide is the leaving group. Base removes the proton to a greater extent for the poorer chloride leaving group than for the better iodo leaving group. Greater electron density on the benzylic position is required to expel the chloride leaving group than the iodide leaving group (i.e., chloride needs a greater "push" than iodide). The transition state for proton removal is later with the chloride leaving group and earlier with the iodide leaving group. These differences in  $\rho$  values are thus very useful for discussing changes in the structures of activated complexes and reaction mechanisms caused by structural changes in the reactants.

Sometimes Hammett plots of rates versus  $\sigma_z$  values are nonlinear. When this occurs, it usually indicates that the model reaction from which  $\sigma_z$  values were determined (the ionization of benzoic acids) does not accurately model the electronic changes occurring in the reaction being studied. In most cases this happens when a positive or negative charge is being developed at a position where direct resonance interactions with the substituent magnify the electronic effect of substituents. Usually this happens when charge is developed at a benzylic position or directly on the aromatic ring.



For these cases, new model reactions were developed and the electronic effects of substituents were obtained as  $\sigma^+$  and  $\sigma^-$  substituent constants. The use of  $\sigma^+$  constants is applicable for reactions in which a positive charge is generated on or in direct conjugation with the aromatic ring, while  $\sigma^-$  constants are used for reactions in which a negative charge is generated on or in direct conjugation with the some starts is found in Table 5.1. These constants are used in the same way as  $\sigma_z$  constants. Rates are measured and correlated with the appropriate set of  $\sigma$  values ( $\sigma$ ,  $\sigma^+$ , or  $\sigma^-$ ) in typical Hammett fashion:

$$\log \frac{k_{\rm Z}}{k_{\rm H}} = \rho^+ \sigma_{\rm Z}^+$$
 and  $\log \frac{k_{\rm Z}}{k_{\rm H}} = \rho^- \sigma_{\rm Z}^-$ 

As before, the  $\sigma$  constants are positionally dependent, that is,  $\sigma^+$  meta is different than  $\sigma^+$  para.

A better linear correlation of the rates with a particular set of  $\sigma$  values means that the set of  $\sigma$  values used is a better model for the electronic character of the

reaction being studied. Knowing that a better correlation is found for  $\sigma^+$  than  $\sigma$  means that the reaction probably involves positive charge development which can be delocalized directly onto the aromatic ring. For example, the hydration of substituted styrenes gives a much better linear correlation of  $\log(k_Z/k_H)$  versus  $\sigma^+$  than  $\sigma$ . This is consistent with the rate-determining step being protonation of the double bond, giving a benzylic cation. At the transition state, resonance delocalization of the developing positive charge into the ring accounts for the better correlation with  $\sigma_Z^+$ . The value  $\rho^+ = -3.57$  means that a significant amount of positive charge is developed at the transition state.



The rates of rearrangement of aromatic amine oxides is found to be correlated much better by  $\sigma^-$  than with  $\sigma$  and has  $\rho^- = 3.6$ . A mechanism consistent with this finding has the oxygen attacking the substituted aromatic ring, thus increasing the electron density on the ring. This charge is delocalized over the aromatic ring by resonance and thus is in direct resonance interaction with substituents in the para position. The process is much better modeled by  $\sigma^-$  than by  $\sigma$  constants.



Because of the way in which they were developed, only  $\sigma_p^-$  substituent constants are available. If a substituent is meta, then normal  $\sigma_m$  values are used because there is no direct resonance interaction with a substituent in the meta position.

By a variety of techniques, it is possible to gain detailed insight into the structure and properties of the activated complex. This is a remarkable ability considering that the lifetime of the activated complex is on the order of a bond vibration and thus cannot be observed directly. This ability to determine the energy of the transition state and structurally characterize the activated complex is a cornerstone of organic chemistry. It allows us to determine mechanisms of reactions, provides a way to evaluate and predict the effects of structural changes on the reactivity of molecules, and enables us to devise new reactions and processes based on new mechanistic principles that are discovered. The result is an ever-increasing arsenal of reactions than can be used most effectively for carrying out needed organic transformations.

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#### PROBLEMS

**5.1.** The base-promoted elimination of quaternary ammonium ions (Hofmann elimination) has been proposed to proceed by an E2-like mechanism. Tell how each of the following observations supports this mechanistic classification. Be specific about exactly what each piece of information reveals.

(a)

$$Z \xrightarrow{\bigoplus} -CH_2CH_2 \xrightarrow{\oplus} N(CH_3)_3 \xrightarrow{EtO^-} Z \xrightarrow{\bigoplus} -CH = CH_2 + EtOH + N(CH_3)_3$$

 $\rho = +3.58$  for a series of Z substituents when plotted against  $\sigma_Z$ 

(b) 
$$\bigoplus$$
 CH₂CH₂-N(CH₃)₃ vs.  $\bigoplus$  CD₂CH₂-N(CH₃)₃  
 $\frac{k_{\rm H}}{k_{\rm D}} = +3.23$ 

(c)  

$$12 \oplus L^{14} \oplus L^{14} \oplus L^{12} \oplus$$

(**d**)



5.2. Explain the mechanistic significance of the  $\rho$  values for the elimination of 1 under the two different sets of conditions.



**5.3.** Propose a mechanism for the following reaction given that  $k_{\rm H}/k_{\rm D} = 6.1$  and  $\rho^- = 2.02$  (there is a better correlation with  $\sigma^-$  substituent constants than with  $\sigma$  values).



**5.4.** Tell how you could use isotopes to distinguish between the following two mechanisms:



**5.5.** The following transformations occur by different mechanisms. Show them and predict what a Hammett study would show for each of them.



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- 120 MECHANISMS OF ORGANIC REACTIONS
- **5.6.** The base hydrolysis of a series of substituted phenoxy esters gave a much better correlation when the rate constants were plotted against  $\sigma_Z^-$  than against  $\sigma_Z$ . Give the mechanistic significance of this behavior.



5.7. Based on the better correlation of rates versus  $\sigma_Z^+$  and  $\rho^+ = -1.03$ , give a likely mechanism for the following substitution:



**5.8.** When  $17\alpha$ -methyl- $5\alpha$ -androstan- $3\beta$ ,  $17\beta$ -diol and its deuterated analog were tested, it was found that the protio compound is three times more active than the deutero analog. It is also known that the diol itself is not biologically active. What is the likely metabolically active material? Explain.



**5.9.** The oxidation of substituted benzaldehydes to substituted benzoic acids by pyridinium fluorochromate (PFC) has been studied, and it has been found that the reaction is first order with respect to pyridinium fluorochromate but is of complicated order with respect to the aldehyde. The following scenario was proposed to account for this behavior:

Using the steady-state approximation, derive the rate law for this mechanistic scenario.

It was further found using  $\alpha$ -deutero benzaldehyde that  $k_{\rm H}/k_{\rm D} = 5.33$  and using substituted benzaldehydes that  $\rho^+ = -2.2$ :



Use these data to delineate the nature of the rate-determining step and propose a plausible mechanism for the reaction. Explain how you used the data to arrive at the mechanism.

**5.10.** The rate of reaction of substituted aromatic chlorides with methoxide to produce substituted anisoles is found to give a linear correlation with  $\sigma^-$  but not with  $\sigma_Z$ . Explain in terms of a reaction mechanism.

$$Z \longrightarrow Cl + CH_{3}O^{\ominus} \longrightarrow Z \longrightarrow OCH_{3} + Cl^{\ominus} \rho^{-} = +8.47$$

**5.11.** Derive the rate law that would describe the rate of product formation for the following reaction assuming that the cationic intermediate is a steady-state intermediate:



**5.12.** Derive the rate law that would describe the rate of product formation for the following reaction where no assumptions are made as to the relative magnitudes of  $k_1$ ,  $k_{-1}$ , or  $k_2$ :

$$\begin{array}{c} O \\ \parallel \\ PhCCH(CH_3)_2 \end{array} \xrightarrow{k_1} PhC = C(CH_3)_2 \\ OH \\ \parallel \\ PhC = C(CH_3)_2 + Cr(VI) \xrightarrow{k_2} products \end{array}$$

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- 122 MECHANISMS OF ORGANIC REACTIONS
- **5.13.** Benzaldehyde cyanohydrin formation shown below may involve ratedetermining attack by either H⁺ or CN⁻. A  $\rho$  value of +2.3 was found for the rate of formation of cyanohydrins from a series of substituted benzaldehydes. Which step is rate determining? Based on the fact that cyanohydrin formation does not occur in basic solution, write a complete mechanism for the process.



**5.14.** For the following reactions tell which should have  $k_{\rm H}/k_{\rm D}$  values greater than 1.5 for isotopic substitution at the starred hydrogens:

(a) 
$$CH_2^* \subset CH_2^* \subset CH_2^* $

(c) 
$$(CH_3)_2CH - CH - CH_2CH_3 \xrightarrow{EtOH} (CH_3)_2C = CH - CH_2CH_3 + ArSO_3H \\ OSO_2Ar$$

- **5.15.** For the solvolyses of **A** and **B**, rates of ionization were found to correlate best with  $\rho^+$  and gave the  $\rho^+$  values shown.
  - (a) How do the transition states for the solvolyses of A and B vary?
  - (b) How do you know this and why?



**5.16.** Draw mechanisms for the following transformations using curved-arrow notation based on the data you are given. There may be more than one step.



 $\rho^+ = -4.2$  (poor correlation vs.  $\sigma$ )

# 6

# STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

Stereochemical Structures	125
Chinality	123
Chirality	128
Configuration of Chiral Centers	129
Multiple Stereocenters	132
Optical Activity	137
Absolute Configuration	138
Physical Properties of Enantiomers	139
Resolution of Enantiomers	140
Stereoselective Reactions	144
Formation of Enantiomers	144
Formation of Diastereomers	146
Stereochemistry to Deduce Mechanism	152
Conformational Analysis	157
Conformational Energies	164
A Values	166
Strain in Ring Systems	167
Stereoelectronic Effects	172
Bibliography	176
Problems	176

The functional groups present in a molecule provide a focal point for chemical transformations of the molecule and define the types of reactions that will normally

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take place. The functional groups also affect the types and strengths of intermolecular interactions in which a molecule engages. Thus functional groups are important determinants of both the chemical and physical properties of molecules. In addition to the functional groups present in a molecule, the structure of a molecule can also influence its chemical and physical properties significantly.

The term *stereochemistry* refers to the spatial arrangement of atoms and groups in a molecule. Organic molecules are, for the most part, three-dimensional objects. The shapes of molecules are the result of the hybridization and bonding geometries of the atomic constituents of the molecule. Since carbon atoms normally define the backbone of the molecule, they play a major role in defining the ultimate shape the molecule takes. Thus familiarity with the principles of hybridization and bonding is indispensable in developing an understanding of the stereochemistry of organic molecules.

Because the shape of a molecule can be crucial to its function and properties, the stereochemical outcome of a chemical reaction may be just as important as the chemical outcome. Therefore we must also learn how to deal with stereochemical issues in molecules and how to understand the stereochemistry of chemical processes.

#### STEREOCHEMICAL STRUCTURES

To discuss the spatial relationships of groups in molecules, we first have to be able to draw structures in such a way that the stereochemical features will be represented unambiguously. Thus a system is needed to depict in two dimensions the spatial relationships between groups in molecules that occur in three dimensions.

An early method to picture the three-dimensional properties of molecules was the use of Fischer projections. In Fisher projections, bonds are drawn either vertically or horizontally. Bonds which are vertical project into the space behind the plane of the paper (blackboard, computer screen). Bonds which are horizontal project into the space in front of the plane of the paper (blackboard, computer screen).



This provides a perfectly good way to denote the stereochemical structure of a molecule if a few rules are followed.

While it is a planar figure, a Fischer projection can only be rotated  $180^{\circ}$  in the plane of the paper, and it may not be taken out of the plane of the paper

#### 126 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

and flipped over. As shown below, rotating the molecule  $180^{\circ}$  gives the identical molecule whereas rotating the molecule  $90^{\circ}$  gives a nonsuperimposable isomer (the enantiomer). Within these constraints, however, Fischer projections are quite valid for showing the stereochemistry of a molecule.



Furthermore,



A more recent approach to the three-dimensional depiction of molecules is the use of wedged and dashed lines. Simple lines are used to denote bonds in the plane of the paper (blackboard, computer screen); wedged lines denote bonds projecting into the space in front of the plane; dashed lines denote bonds projecting into the space behind the plane. These bonds are not restricted to any particular orientation but are general. Bearing in mind the tetrahedral geometry of saturated carbon atoms, these figures can be turned and rotated at will as long as you are able to keep track of what happens to each of the four valences as the molecule is tumbled.

STEREOCHEMICAL STRUCTURES 127



It may help to make models and practice manipulating these structures. An added benefit of depicting molecules using wedged and dashed figures is that other types of molecules can be depicted using this convention. For example, the planar geometry of olefins and the twisted geometry of allenes is readily pictured using wedged and dashed bonds.

A further simplification of stereochemical notation for saturated carbon centers is to stretch out the carbon skeleton in the plane of the paper (blackboard, computer screen). Valences of atoms or groups other than hydrogen are indicated by a bold line if they project into the space in front of the plane and with a dashed line if they project into the space behind the paper (blackboard, computer screen).

$$\bigcup_{i=1}^{OH} = \bigcup_{i=1}^{OH} \underbrace{\underset{i=1}{H}}_{ClCH_2CH_2CH_2CH_2} \underbrace{\underset{i=1}{CH_2CH_2CH_2}}_{CH_3CH_2} \underbrace{\underset{i=1}{Cl}}_{Cl} = \bigcup_{i=1}^{Cl} \underbrace{\underset{i=1}{Cl}}_{Cl}$$

Other common methods for representing the three-dimensional structures of molecules include Newman projections for showing conformational relationships and sawhorse figures. Newman projections look down a carbon–carbon bond so that the front carbon, designated by a circle, obscures the carbon directly behind it. Valences (bonds) to the front carbon extend to the center of the circle, while bonds to the rear carbon stop at the circle. Sawhorse projections have the carbon–carbon bond at oblique angles, which attempts to represent a perspective drawing of the molecule. Thus for 2-chloro butane, if one chooses to examine the 2,3 bond, then the sawhorse and Newman projections would be



Keeping in mind the three-dimensional properties of molecules, Newman projections can be converted to wedged-dashed structures or Fischer projections as desired. It is important to develop facility for manipulating structures and

#### 128 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

visualizing the three-dimensional properties of molecules from various stereostructures. The use of molecular models in conjunction with two-dimensional structures is often helpful in making the visual connections.

#### CHIRALITY

Because of the tetrahedral geometry of saturated carbon and the associated three-dimensional properties, molecules can have chirality as one stereochemical feature. Any object is chiral if it is different (nonsuperimposable) than its mirror image. Likewise a molecule is chiral if it is nonsuperimposable on its mirror image. This requirement does not consider conformational changes (rotations about single bonds) as valid conditions for nonsuperimposability. Thus, for the molecules below, the first is achiral (not chiral) because it is superimposable on its mirror image and the second is chiral because it is not superimposable on its mirror image.



When a molecule is chiral, then it will have two isomeric forms called enantiomers, each of which is the nonsuperimposable mirror image of the other. Enantiomers are distinct stereoisomers because they are compounds that have the same molecular formula and sequence of bonded elements but which differ in the spatial arrangement of groups in the molecule. If a molecule is chiral, and thus has two enantiomers, it usually (but not always) contains at least one chiral center. In organic compounds a chiral center usually corresponds to an asymmetric tetrahedral carbon atom.



The most common bonding motif which results in an chiral center is a tetrahedral carbon atom which is bonded to four different groups. A tetrahedral carbon with four different groups attached is described variously as a chiral center, a chiral carbon, or an asymmetric center because that carbon lacks symmetry elements. On the other hand, a tetrahedral carbon with two or more of the same groups attached automatically has a plane or axis of symmetry associated with it and so it is achiral (not chiral). When a molecule contains a chiral center, that molecule lacks symmetry elements and thus is chiral, and it can have two enantiomers which are nonsuperimposable mirror images of each other.

A tetrahedral carbon with four different groups attached is also one type of stereogenic center. A stereogenic center is an atom at which the interchange of two ligands results in a stereoisomer. It turns out that the interchange of two groups bonded to a chiral carbon results in a stereoisomer of that molecule. If there is only a single chiral center in the molecule, then the interchange of two groups leads to the enantiomer of that molecule.



The four different groups attached to a chiral carbon can be different elements, isotopes, or functional groups, and chiral centers can be present in both open-chain molecules or cyclic compounds. The recognition of chirality and chiral centers in molecules is an important step in determining the numbers of stereoisomers that are possible for a given compound.

#### **CONFIGURATION OF CHIRAL CENTERS**

As noted above, when a single chiral center is identified in a molecule, then there will be two stereoisomers (enantiomers) of that molecule. Enantiomers differ only by the spatial arrangements of groups around the chiral center; however, they are distinct isomers. To discuss enantiomers, it is necessary to have some way to name or denote which one is which; that is, we have to convert the configuration (spatial arrangement) of groups around the chiral center into a name or designation. An early method was to designate enantiomers as either D or L based on the relationship of their chiral center to the chiral center in D- or L-glyceraldehyde. This system of nomenclature was difficult to apply to larger molecules and particularly ones with more than one chiral center.

#### **130** STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

In response to this nomenclature dilemma, the Cahn–Ingold–Prelog (IUPAC, International Union of Pure and Applied Chemistry) system of nomenclature was developed and is now the standard method to specify the relative configuration of chiral centers in molecules. Each chiral center will have two possible mirrorimage configurations, which are designated as either R or S.

The strategy for determining whether the chiral center has an R or S configuration is based on the symmetry properties of a tetrahedral carbon. First one assigns priorities to the four groups attached to the chiral center. Next one orients the molecule so as to sight directly down the bond from the chiral carbon to the group of lowest priority. The remaining three bonds will form a trigonal array (as in Newman projections).



clockwise : R configuration

(If it is not apparent from stereostructures, build a model and satisfy yourself that this is so.)

If you then follow these three groups around a circle from the group of highest priority to the group of lowest priority, you must proceed either clockwise (right), which is designated as the R configuration, or counterclockwise (left), which is designated as the S configuration. This provides a general method for assigning the configuration to any chiral center.

In summary:

- 1. Assign priorities to the groups attached to the chiral center.
- 2. Orient the molecule so the group of lowest priority points directly away from your eye.
- 3. Follow the direction of the remaining groups from the highest to lowest priority. If the procession is clockwise, the configuration is designated R; if the procession is counterclockwise, the configuration is designated S.



highest to lowest priority is counterclockwise  $\therefore$  S

The other enantiomer will obviously have the R configuration.



The only remaining task is to assign priorities to the groups attached to the chiral center. This is done by the following rules:

1. Priority is first assigned on the basis of the atomic number of the atoms attached directly to the chiral center. Atoms of higher atomic number are given higher priorities. Thus for 1-bromo-1-fluoroethane, the ordering of priorities is Br > F > C > H on the basis of their respective atomic numbers of 35 > 9 > 6 > 1.



2. When assignment of priority cannot be made on the basis of atoms attached directly to the chiral center, proceed away from the chiral center and examine the next sets of atoms for differences in atomic numbers of attached atoms. Thus for 2-chlorobutane, two carbon atoms are attached to the chiral center. To establish which carbon group takes priority, note that the next atoms are H,H,H for the methyl group and H,H,C for the ethyl group. Thus the latter has higher priority because of the greater atomic number of carbon.



3. Groups containing multiple bonds are assigned priority as if both atoms were doubled or tripled. Thus a vinyl group is equivalent to a 2-butyl group by so-called phantom atoms. The phantom atoms do not include the requisite number of hydrogen atoms to complete the valences.
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#### **132** STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM



These rules and applications are summarized completely in most introductory organic texts. It is important to be able to assign R,S configurations to stereogenic centers in molecules and to construct chiral molecules given the R or S configuration. In this way it is very easy to determine the stereochemical relationships between stereoisomers.

# MULTIPLE STEREOCENTERS

When there is more than one chiral center in a molecule, the number of possible stereoisomers increases. Since each chiral center can have either the R or S configuration, for a molecule of n chiral centers, there will be  $2^n$  possible stereoisomers. Thus 3-phenyl-2-butanol has two stereogenic centers and four possible stereoisomers. These are shown below with the configuration of each chiral center designated.



The configuration of each chiral center can be determined in the usual way, but there is a much faster way to draw the four stereoisomers. First draw a stereostructure of the molecule such as **A** and then go through the process of determining the R or S configuration. Each chiral center is a stereogenic center in that the interchange of two ligands results in a stereoisomer. Since each chiral center can be changed merely by exchanging any two valences, the remaining stereoisomers of **A** can now be generated easily by switching valences of one stereocenter (**B**), then the other (**C**), then both (**D**). Furthermore, if the relative configurations (R,S) are

known for **A**, then the configurations of the other stereoisomers are immediately known since exchanging any two valences inverts the configuration (R goes to S). For 3-phenyl-2-butanol, structure **A** has the configuration (2R,3S)-3-phenyl-2butanol. (The stereochemical information is denoted by the number of the carbon and its configuration enclosed in parentheses before the name of the compound.) By exchanging the hydrogen and methyl groups C-2 of **A**, the configuration of C-2 is inverted and isomer **B** is produced, the 2S,3S isomer. Exchange of the hydrogen and phenyl groups at C-3 of **A** inverts the configuration of C-3 and isomer **C** is produced, the 2R,3R isomer. Exchanging the hydrogen and methyl group at C-2 and the hydrogen and phenyl group at C-3 of **A** gives isomer **D**, the 2S,3R isomer.

While the exchange of the methyl and hydrogen groups at C-2 of A was used to invert the configuration, the same configurational change at C-2 of A could be accomplished by exchanging any two valences, such as hydrogen and hydroxyl or hydroxyl and methyl. The same is true for configurational changes at C-3. Exchange of any two ligands will invert the configuration of a tetrahedral stereogenic center.

Thus, given the configurations for the three chiral centers of an aldopentose such as D, one can rapidly write down the configurations of the chiral centers of E, F, or G,

which are stereoisomers of **D**, without having to apply the Cahn–Ingold–Prelog rules for each isomer. This is done by simply noting which substituents are switched relative to those in **D**. If the configuration is switched from that in **D**, then the designation (R,S) will also be switched from that in **D**. Isomers **D**–**G** are only four stereoisomers out of eight possible stereoisomers for an aldopentose which has three stereocenters  $(2^3 = 8)$ .

Now that we can generate the stereoisomers of compounds with more than one chiral center, it is appropriate to ask what are the relationships between these isomers. Thus 2-bromo-3-acetoxy butane has two chiral centers and four stereoisomers, shown below.

Since each carbon of **1** is the mirror-image configuration of the carbons in **4** (i.e., C-2 of **1** is R and C-2 of **4** is S; C-3 of **1** is S and C-3 of **4** is R), then the molecules themselves are mirror images, but they are nonsuperimposable. They are thus enantiomers. This relationship can also be shown by reorienting the molecules to see that they are mirror images,



but nonsuperimposable and therefore are enantiomers.



A similar analysis reveals that 2 and 3 are also enantiomers. Comparison of any other pairs of stereoisomers, 1 and 2, for example, shows that they are not mirror images: The C-2 of 1 is R and C-2 of 2 is S *but* C-3 of both 1 and 2 is S. Isomers 1 and 2 are also not superimposable. So 1 and 2 are a second type of stereoisomer and are nonsuperimposable, non-mirror images called diastereomers. Diastereomers have the same molecular formula and sequence of bonded elements but different spatial arrangements and are nonsuperimposable, non-mirror images.

A third type of stereoisomer occurs when a molecule with several stereogenic centers contains an internal plane of symmetry. This usually happens when two of the stereogenic centers are attached to the same four different valences. For example, 2,4-dibromopentane has two stereogenic centers and thus four stereoisomers, 5-8. It is easily seen that 6 and 7 are enantiomers, 5 and 6are diastereomers, and so on.



However, **5** and **8** are identical. Although there are two chiral centers in **5** (and **8**), the molecule itself is achiral because it contains an internal mirror plane. Thus it has a plane of symmetry. Structure **8** is superimposable on **5** by a  $180^{\circ}$ 

rotation and thus is the same compound. This molecule is called a meso isomer, a compound which contains chiral centers but itself has a plane of symmetry. Even though 2,4-dibromopentane has two stereogenic centers, there are really only three stereoisomers, a pair of enantiomers and a meso compound which is diastereomeric with the enantiomeric pair.

It is clear from the above examples that the presence of chiral centers in molecules leads to stereoisomers. There is another type of molecule which itself is chiral but has no chiral center. The molecular chirality arises from the presence of a screw axis in the molecule. Allenes and biphenyls are common examples of such compounds, and because they are chiral, they exist as enantiomers.



We have seen that when more than one stereocenter is present in a molecule, both enantiomers and diastereomers are possible. Distinguishing between enantiomers requires the relative configurations of each stereogenic center to be specified. However, to distinguish diastereomers, only the relative spatial orientation of groups needs to be specified. For example, aldotetroses have two stereocenters and the four stereoisomers are shown below:



The enantiomeric relationship between D-threose and L-threose is specified by the 2S,3R and 2R,3S configurations (each stereocenter is the mirror image of the other). Moreover the enantiomeric relationship between D-erythrose and L-erythrose is clear from the 2R,3R and 2S,3S configurations. However, threose and erythrose are diastereomers. The different spatial orientation of the –OH groups extending from the chain in the Fisher projections makes the diastereomeric relationship obvious without the need for specifying the configuration; that is, they are clearly nonsuperimposable and non-mirror images.

By extension, other diastereomeric pairs of molecules which contain two adjacent stereogenic centers can be designated as threo or erythro depending on whether substituents extend to opposite (threo) or the same (erythro) sides of the Fisher projection of the molecule. For example,



The three and erythre designation denotes a diastereomeric relationship of the isomers. Each three and erythre isomer will also have enantiomers which will also have a three-erythre diastereomeric relationship to each other.

More recently a new method for designating the stereochemical relationship of diastereomers has been developed. In this method the carbon backbone is extended in the plane of the paper, blackboard, or computer screen in the horizontal direction. Groups will extend from this backbone either in front of the plane or behind it and are designated by bold or dashed bonds, respectively. If two substituents extend in the same direction, their spatial relationship is designated syn; if they extend in opposite directions, their spatial relationship is designated anti.



Molecules which are syn-anti isomers of each other are diastereomers, and there will be two syn enantiomers and two anti enantiomers. The syn-anti designation is not restricted to substituents on vicinal carbon atoms as is the threo-erythro designation and is thus more versatile.

# **OPTICAL ACTIVITY**

We have seen how stereochemical relationships can be designated and distinguished. Now let us see how the stereochemistry influences the chemical and/or physical properties of molecules.

Individual pure enantiomers are identical to each other in most respects in that they have the same physical properties, melting point (m.p.), boiling point (b.p.), refractive index, polarity, and solubility. The only difference between individual enantiomers is that they behave differently in chiral environments. For example, each enantiomer of an enantiomeric pair produces a rotation of the plane of plane-polarized light to an equal but opposite extent. This is because plane-polarized light is itself chiral and each enantiomer interacts differently with the light (Figure 6.1).

If a compound rotates plane-polarized light, it is termed *optically active*. To be optically active, a compound must be chiral and one enantiomer of the compound must be present in excess over its mirror image. The enantiomer which produces clockwise rotation of plane-polarized light is designated the positive enantiomer, and the enantiomer which produces counterclockwise rotation of plane-polarized light is designated the negative enantiomer. At a given wavelength under standard conditions of concentration (1 g/mL) and pathlength (10 cm), a pure enantiomer will give the maximum rotation in degrees. Its pure mirror image will give an equal but opposite rotation.

The rotation of plane-polarized light by a pure enantiomer is an inherent property of that enantiomer. However, the amount of rotation actually measured is dependent on the concentration of molecules in the light beam, the pathlength, and the wavelength of the light used. To account for these variables, the observed rotation is converted to the specific rotation [ $\alpha$ ], which is defined as the rotation observed for a solution of 1 g/mL concentration in a 10-cm-pathlength cell. Furthermore a wavelength of 589 nm, the D line of sodium, is normally the standard wavelength for measuring the specific rotation and 25°C is the standard temperature of the measurement. The specific rotation [ $\alpha$ ]_D²⁵, in which the superscript indicates the temperature of the measurement and the subscript D (or number) indicates the wavelength of light used to measure the optical rotation, is calculated from



Figure 6.1 Rotation of plane polarized light by enantiomers.

where  $\alpha_{obs}$  is the observed rotation, *l* is the pathlength of the cell used for the measurement, and *C* is the concentration of the sample in grams per milliliter.

When both enantiomers are present in solution, the observed rotation will reflect the enantiomeric composition of the mixture. If equal amounts of enantiomers are present, the solution will not exhibit optical activity, because for each molecule that rotates light in one direction there will be another molecule that rotates light in the opposite direction and the net rotation is zero. Such a mixture is called a racemic mixture and is indicated by  $(\pm)$ . Thus  $(\pm)$ -2-butanol is an equal mixture of the R and S enantiomers of 2-butanol. In the liquid state, racemic mixtures have the same physical properties as the individual enantiomers.

If one enantiomer is present in excess over the other, then the solution will have a net rotation corresponding in sign (+ or -) to that of the more abundant enantiomer. The composition of the mixture is denoted by the optical purity or the percent enantiomeric excess (ee%). The enantiomeric excess is defined as ee = % major enantiomer - % minor enantiomer and is a measure of the optical purity of the sample. Values range from 100% (pure enantiomer, ee = 100% - 0%) to 0% (racemic mixture or ee = 50% - 50%). A sample which has an optical purity of 92% is thus a mixture of 96% of one enantiomer and 4% of the other enantiomer.

# ABSOLUTE CONFIGURATION

The observation of optical activity and the measurement of optical rotation distinguish one enantiomer (+) from the other (-). The sign of rotation (+ or -)is thus an experimental way to differentiate enantiomers. A second way to designate these stereoisomers is to assign the configurations as R or S based on stereostructures (for enantiomers which contain a single chiral center). The R and S configurations are relative configurations based on three-dimensional structures which are drawn on paper. However, it is not possible to predict a priori whether the R enantiomer (for example) will be dextrorotatory (+) or levorotatory (-).

The absolute configuration of an enantiomer is determined only when the optical rotation of an enantiomer (+ or -) can be matched with its configuration (R or S). For example, the absolute configuration of lactic acid has been found to be R-(-) in that the R enantiomer is levorotatory.



Conversion to the methyl ester does not change the configuration of the stereocenter, which remains R. However, the rotation is found to be positive so the absolute configuration is R-(+). This illustrates that while the relative configuration (R,S) can be used to show the structures of stereocenters, the absolute configuration must be known to show the changes in configuration that occur during a chemical sequence. That is, just knowing that a chemical reaction changes the optical rotation in one direction or another is not sufficient to indicate whether a change in configuration at a chiral center has occurred.

# PHYSICAL PROPERTIES OF ENANTIOMERS

If both enantiomers are present in a solid sample, the melting point and the solubility of the solid mixture are often found to be different than those of the pure enantiomers. This is due to the fact that the solid-state interaction of two R enantiomers or two S enantiomers is often different than the solid-state interaction of an R and an S enantiomer. (In fact, these interactions are diastereomeric.) The result is that three different scenarios are possible when a racemic mixture is crystallized from solution:

- 1. If an enantiomer has a greater affinity for molecules of like configuration, then two sets of crystals will be formed, one set composed only of the (+) form and the other composed only of the (-) form. This racemic mixture is called a conglomerate because it is a mixture of two different types of crystals. Moreover it behaves as a typical mixture—the melting point is lower than the pure enantiomeric components and the solubility is higher (Figure 6.2). This is a relatively rare situation.
- 2. If an enantiomer has a greater affinity for molecules of opposite configuration, then crystals are produced which contain equal numbers of the (+)



Figure 6.2 Enantiomers which crystallize as a conglomerate.



Figure 6.3 Enantiomers which crystallize as a racemate.



Figure 6.4 Enantiomers which crystallize as s solid solution.

and (-) forms. The solid compound which has properties different than either pure enantiomer and exists only in the solid state is called a racemate or a racemic compound or a racemic mixture. A racemate is often higher melting and less soluble than a pure enantiomer and behaves as a mixture in the presence of either pure enantiomer (Figure 6.3). This is the most common situation and allows an unequal mixture of enantiomers to be purified. Upon crystallization, the racemate will precipitate first, leaving behind the enantiomer in excess.

3. If one enantiomer has similar affinity for molecules of either configuration, then the enantiomers are randomly distributed in the crystal and the solid is a "racemic solid solution" or mixed crystal. Such solids are identical with either enantiomer (Figure 6.4).

# **RESOLUTION OF ENANTIOMERS**

We have seen that individual enantiomers have identical physical properties and only can be distinguished in a chiral environment. Plane-polarized light is such a chiral environment, and one enantiomer is dextrorotatory and one is levorotatory. Another way to distinguish enantiomers is to allow them to react (or interact) with other chiral molecules. The interaction of a mixture of enantiomers with a single enantiomer of a chiral molecule produces a mixture of diastereomers as illustrated.



Since diastereomers have different physical properties, they can be separated on the basis of those physical properties. After separation of the diastereomers, the individual enantiomers are reclaimed, and in this way the two enantiomers will have been separated. Such interactions form the basis for all separations of racemic mixtures into pure enantiomers, which is termed "resolution" of enantiomers. There are several experimental techniques used to resolve enantiomers, but all utilize a chiral reagent of some type to furnish the chiral environment needed to distinguish the enantiomers.

Crystallization has been a traditional method for separating the diastereomers produced from a racemic mixture and a chiral resolving agent. For example, racemic carboxylic acids can be treated with an optically active alkaloid (which is basic) and the resulting diastereomeric salts are separated by crystallization. The individual enantiomeric acids are then regenerated from the salts. A variety of alkaloids have been used as resolving agents for racemic acids. They include brucine strychnine, ephedrine, quinine, morphine, and  $\alpha$ -phenyl ethylamine, among others.



Racemic bases can be resolved by treating them with an optically active acid and separating the resulting diastereomeric salts by fractional crystallization. The individual enantiomeric bases are regenerated from the salts. Common acid-resolving agents include camphorsulfonic acid and derivatives of it, tartaric acid, malic acid, and pyroglutamic acid, among others.



Alcohols are often resolved by conversion to half esters of phthalic acid or succinic acid, which are then resolved as typical acids. The alcohol is then regenerated from the resolved half ester by hydrolysis or reductive cleavage with LAH. A second method for resolving alcohols is to convert them to esters of optically active acids. This gives a mixture of diastereomeric esters which are separated by fractional crystallization and the alcohol is recovered by hydrolysis or reductive cleavage.



In recent times chromatography has become a major technique for separations and has increasingly supplanted fractional crystallization as a way to separate diastereomeric compounds. Not only do diastereomers have different solubilities, they also interact with surfaces such as silica gel or alumina differently. The mixture of diastereomeric esters obtained by coupling a racemic alcohol to optically active acid can often be separated by high-performance liquid chromatography (HPLC), radial chromatography, or flash chromatography. Chromatography is often much faster and more efficient than crystallization. The individual alcohols can be regenerated in the usual fashion.

The preceding methods for the resolution of enantiomers rely on the formation of strongly bound diastereomers (ionic or covalent) which are then separated. It has become more and more common to use weak interactions as a means of resolving enantiomers. Chiral chromatography columns are useful for the separation of a variety of compounds, including amino acids. A chiral substance is permanently attached to the column surface. If a mixture of enantiomers is passed over the surface, the individual enantiomers will interact with the chiral surface differently and thus will elute along the column at different rates. (The enantiomer which interacts with the surface more strongly will elute more slowly). They can thus be collected individually. A variety of chiral stationary phases are available to separate an ever increasing number of examples.



The use of enzymes to resolve enantiomers has become an extremely popular method only recently. Enzymes are chiral catalysts which often exhibit very high selectivity for one enantiomer of a racemic mixture. Since enzymes are soluble in aqueous solution, it was often impossible to get sufficiently high concentrations of organic substrates in the aqueous medium to achieve conversion at any reasonable rate. The finding that a variety of esterases (lipases) can function very well in organic solvents has removed this major stumbling block to the practical utilization of biochemical transformations for resolution. In addition, a variety of enzymes are available commercially. Moreover a variety of other enzymes are now available and can be used to resolve enantiomers effectively. For example, some amidases and peptidases (amide bond hydrolysis) can be applied to the resolution of enantiomers.

Thus it is very easy to acetylate a racemic alcohol and treat the racemic mixture of acetates with a lipase. One enantiomer is hydrolyzed to the alcohol and the other remains as the ester. These are separated chromatographically and each component is obtained with high optical purity. This technique is becoming more important and could be the most general technique for resolution in the future.

ROH	$\xrightarrow{Ac_2O}$	ROAc	lipase	R*OH	+	R*OAc
R, S mixture		R, S mixture		resolved alcohol R or S		resolved ester S or R

The use of kinetic resolution to obtain a single enantiomer from a mixture of enantiomers is often useful for particular functional groups. Since individual enantiomers react at different rates with chiral reagents, treatment of a racemic mixture with a limited amount (0.5 equiv.) of a chiral reagent will convert one of the enantiomers to product in preference to the other. After workup one enantiomer will be recovered unchanged while the other will have been converted to a new product. The efficiency of the kinetic resolution will depend on the relative rates of reaction of the two enantiomers. If rates of reaction (selectivity factor) vary by >100, then the recovered enantiomer will be >99% optically pure. Lower selectivity factors will lead to less pure enantiomers.

Optically active diisopinocamphenylborane can be used to resolve racemic olefins. The reagent adds to one enantiomer, and the other is unchanged. Optical purities on the order of  $\sim$ 37–65% are possible. Chiral allylic alcohols can be resolved with chiral epoxidizing agents derived from tartrate complexes of titanium. One enantiomer is epoxidized and the other is not. Thus, the two alcohol enantiomers can be separated, one as the unsaturated alcohol and one as the epoxy alcohol. Use of the other tartrate isomer reverses the stereoselectivity. Selectivities on the order of >100 are possible with this method. As in any kinetic resolution, however, only one enantiomer can be recovered. The other is converted to a different chiral product.

Even if the separation of enantiomers by any of the above methods is not completely successful, it is often possible to further raise the enantiomeric excess by crystallization or chromatography. In this way many pure enantiomers are now available.

# STEREOSELECTIVE REACTIONS

Chiral compounds are very important substances. Many natural products, medicinal compounds, and biomolecules exist as a single, optically active stereoisomer. Furthermore the opposite enantiomer or diastereomer may not have any physiological activity or may, in fact, have a detrimental physiological effect. There is therefore great interest in reactions in which only one stereoisomeric form of a compound is produced by a particular synthetic sequence.

The simplest approach is to use a starting material of known absolute configuration and manipulate it to the final product using reactions whose stereochemical outcome at the chiral center is known and/or predictable. For example, the synthesis of unusual amino acids as single enantiomers can begin with a "normal" amino acid from the chiral pool. (The "chiral pool" is a large group of molecules which are readily available and whose absolute configurations are known with certainty. Often these molecules are or are derived from natural products which can be isolated from natural sources in enantiomerically pure form.)



Since the stereochemical changes in each reaction of the sequence are known, a particular amino acid starting material (R or S) will give a particular configuration in the product. In this strategy of asymmetric synthesis, all or part of the final molecular skeleton is derived from the chiral precursor. While simple, this strategy is limited by the size of the chiral pool and by the types of reactions which occur stereospecifically at tetrahedral centers.

It is much more common for reactions to produce new chiral centers from achiral starting materials. Consequently, if we are to use the whole arsenal of synthetic methods available to us and at the same time produce single stereoisomers, then we must be able to control (or at least understand) the stereochemistry of reactions occurring at achiral centers.

# FORMATION OF ENANTIOMERS

Since chiral centers are most commonly tetrahedral, the conversion of trigonal centers to tetrahedral centers by some type of addition process is the most common way in which new chiral centers are created. The reaction of carbonyl groups with nucleophiles is a classic example. If substituents on the carbonyl group and the nucleophile are all different, then a new chiral center is produced, as in the reaction of acetophenone with sodium borohydride to produce 1-phenylethanol.



The carbonyl group is trigonal and planar and can be thought of as having two faces. Addition of hydride to one face gives one enantiomer while addition to the opposite face gives the opposite enantiomer. As rewritten below, attack from above gives the R enantiomer while attack from below gives the S enantiomer. The faces are stereochemically nonequivalent since different stereoisomers are produced.



To differentiate the faces of a carbonyl group, the Re–Si nomenclature has been developed. The groups around the carbonyl carbon are given priorities by the same rules used in the Cahn–Ingold–Prelog system for R,S nomenclature. Then going from the group of highest priority to the group of lowest priority around the face of a carbonyl group, proceeding in the clockwise direction defines the Re face and proceeding in the counterclockwise direction defines the Si face.



The Re–Si nomenclature enables the faces of a carbonyl group to be differentiated stereochemically; however, the carbonyl group itself is achiral. Moreover, the Re–Si designation is not indicative of the stereochemistry of the chiral center produced by addition. In the above example hydride addition to the Si face gives

the R enantiomer while hydride addition to the Re face gives the S enantiomer. If ethyl lithium were added, the stereochemistry would be reversed, that is,  $Si \rightarrow S$  and  $Re \rightarrow R$ .

New chiral centers are produced by addition reactions to other trigonal centers as well. Hydrogenation of 3-methyl-3-hexene gives 3-methylhexane. Clearly the addition of hydrogen to one face of the planar olefinic system gives one enantiomer and addition to the opposite face gives the opposite enantiomer. Likewise reaction of styrene with chlorine or bromine ( $X_2$ ) or potassium permanganate produces products with a new chiral center. Formation of the two possible enantiomers results from addition to either face of the olefin.



Reactive intermediates which are planar can also produce enantiomers. The acid-catalyzed addition of water to 1-pentene proceeds via a secondary carbocation. Because the carbocation is a trigonal, planar intermediate, water can add to either face to give the R or S enantiomers.



In reactions in which neither the reactants (C=O, C=C, C⁺) nor the reagents  $(BH_4^-, EtMgBr, Br_2, H_2O, etc.)$  are chiral, there is no possibility for controlling which face undergoes addition (in fact, addition to either face is equivalent); thus a racemic mixture will be produced. Such processes are described as having no enantioselectivity.

# FORMATION OF DIASTEREOMERS

Diastereomers are defined as compounds which have the same molecular formula and sequence of bonded elements but which are nonsuperimposable, non-mirror

images. While Z,E isomers are one subclass of diastereomers which are achiral, the majority of diastereomeric compounds are chiral compounds which have more than one chiral center. Furthermore it is important to recall that for a compound with *n* chiral centers there will be  $2^n$  stereoisomers. These will be divided into  $2^n/2$  pairs of enantiomers, and each pair of enantiomers will be diastereomeric with the other pairs of enantiomers. This was reviewed earlier in this chapter.

One of the most direct ways to produce diastereomers is by addition reactions across carbon–carbon double bonds. If the structure of the olefin substrate is such that two new chiral centers are produced by the addition of a particular reagent across the double bond, then diastereomers will result. For example, the addition of HBr to Z-3-chloro-2-phenyl-2-pentene produces 2-bromo-3-chloro-2-phenylpentane as a mixture of four diastereomers. Assuming only Markovnikov addition, the diastereomers are produced by the addition of a proton to C-3 followed by addition of bromide to the carbocation intermediate at C-2. Since the olefin precursor is planar, the proton can add from either face, and since the carbocation intermediate is also planar and freely rotating, the bromide can add to either face to give diastereomeric products. The possibilities are delineated schematically (but not mechanistically) below.



Now even though there are four possible stereoisomers that can be produced, they will not necessarily be formed in equal amounts. Diastereomers are not equal in terms of their energies; consequently, reactions which produce diastereomers reflect these energy differences in the various transition states and therefore proceed at different rates. Thus diastereomers are normally formed in unequal amounts. This is a very important concept since it provides the kinetic basis for the stereoselectivity found in many different organic reactions. Restating this idea, if reactions produce diastereomers and thus proceed via diastereomeric transition states, then the energy barriers for the formation of individual diastereomers will be different, the rates of formation of individual diastereomers will be different, and they will be formed in unequal amounts. The greater are the differences in the energy barriers, the greater will be the differences in rates and the more stereoselective will be the reaction. In contrast to HBr addition, which gives a mixture of diastereomers, there are a variety of other olefin addition reactions which yield a single diastereomer from a starting olefin of defined stereochemistry. Furthermore a starting olefin with a different stereochemistry will give a

different single diastereomer. Such reactions are described as being stereospecific or highly stereoselective.

The diastereoselectivity for any process is often reported as a diastereomeric excess (de%), which is analogous to the optical purity reported for mixtures of enantiomers. The de% is given by de% = % major diastereomer – % minor diastereomer. For diastereospecific reactions in which a single diastereomer is produced, de = 100%, while for reactions in which there is no selectivity and diastereomers are produced in equal amounts, de = 0%.

A typical example of a stereospecific olefin addition reaction is the addition of bromine to olefins. If *cis*-2-pentene is used as the substrate, only the 2R,3R and 2S,3S pair will be produced (they are enantiomers).

$$\underset{H_{3}C}{\overset{H}{\longrightarrow}} \underbrace{\overset{H}{\longrightarrow}}_{CH_{2}CH_{3}} \underbrace{\overset{Br_{2}}{\xrightarrow{CCl_{4}}}}_{H_{3}C} \underbrace{\overset{Br}{\xrightarrow{R}}}_{H_{3}C} \underbrace{\overset{H}{\xrightarrow{R}}}_{Br} + \underbrace{\overset{H}{\xrightarrow{H_{3}C}}}_{Br} \underbrace{\overset{H}{\xrightarrow{S}}}_{S} \underbrace{\overset{H}{\xrightarrow{S}}}_{Et} (6.2)$$

Because the addition of bromine is stereospecifically trans or anti, one bromine atom adds to each face of the olefin and can go to either carbon. If *trans*-2-pentene is used as the substrate, then only the 2R,3S and 2S,3R pair is produced (they are also enantiomers.). However, the pair from *cis*-2-pentene is diastereomeric with the pair from *trans*-2-pentene.

The stereospecificity observed in olefin bromination is only possible if the inherent facial relationship of the olefinic bond is maintained throughout the addition process *and* only one bromine atom adds to each face. In bromination, the electrophilic addition leads to a bridged bromonium ion which not only maintains the initial olefin geometry but also forces the second bromine to add from the opposite direction (anti).



(Contrast this to the addition of HCl or water to a double bond where the intermediate is free to rotate so that the olefin geometry is lost and both the proton and the nucleophile can add to either face).

Other olefin additions which proceed via bridged intermediates should show similar stereospecificity and addition should occur anti. Chlorination of olefins is an obvious analogy to bromination, but the addition of sulfenyl chlorides, oxymercuration, and expoxidation/hydrolysis all give stereospecific anti addition across the double bond because bridged intermediates are involved.



There are other stereospecific olefin addition processes which occur with cis or syn stereochemistry. Common examples include catalytic hydrogenation, hydroboration/oxidation, and dihydroxylation using osmium tetroxide. The stere-ospecificity of these syn additions requires that the facial properties of the olefinic bond be maintained throughout the addition process *and* that both new bonds are formed to the same face of the olefin. This is normally accomplished by a concerted syn addition to the  $\pi$  system.



Stereospecificity in hydrogenation is gained by a surface-mediated delivery of the hydrogen atoms to one face of the olefin. Stereospecificity in both hydroboration/oxidation and osmium tetroxide/reduction results from a concerted addition to one face of the  $\pi$  system. This mode of addition guarantees that both new bonds are formed on the same face of the olefin. Although the reagents can add to either face of the olefin, this leads only to enantiomers of a single diastereomer. The concerted addition is the key feature which assures syn selectivity.

Another type of stereoselectivity is possible when a new chiral center is produced in a molecule which already contains one or more chiral centers. A typical example of such a process would be addition to an aldehyde or ketone which already contains a chiral center.



To understand the stereoselectivity that might be observed, it is first necessary to delineate the stereochemical possibilities. The existing chiral center can be either R or S or both. Addition to the carbonyl group can potentially occur from either face since it is planar. Thus, if the existing chiral center is of the R configuration, the products of addition will have either the R,R or R,S configurations and are diastereomers. If the starting material has only the R configuration, each diastereomer is optically active because only one enantiomer is produced. If the existing chiral center is of the S configuration, the products of addition can be either S,S or S,R diastereomers. Each diastereomer will be optically active, and they are enantiomers of the diastereomeric pair formed from the R configuration of the precursor. If the starting material is a racemic mixture, then all four stereoisomers will be produced—two sets of enantiomeric diastereomers. (That is, the R will give R,R and R,S and the S will give S,S and S,R.) The product mixture will be optically inactive.

A given chirality of the starting material gives two diastereomers, and it is normal to find that these two diastereomers are not produced in equal amounts. Because the two diastereomeric products are of different energies, the diastereomeric transition states leading to them will be of different energies, the rates of their formation will be different, and they will be produced in unequal amounts. If one diastereomer is produced in excess of the other, the reaction is diastereoselective. If only one diastereomer is produced, the reaction is diastereospecific or highly diastereoselective. The same analysis would apply if the starting material is racemic. The reaction would still produce two diastereomers and each would be formed as a pair of enantiomers, and the same diastereoselectivity would be observed. As stated previously, the addition of nucleophiles to chiral carbonyl compounds is a very common type of reaction which produces diastereomeric mixtures. The diastereoselectivity varies with the reagents and conditions. Some examples are



By analogy, the formation of diastereomers is observed for additions to other trigonal systems, such as olefins, which have a chiral center elsewhere in the molecule. In these cases, if optically active starting materials are used, then the diastereomers will be optically active. If racemic starting materials are employed, the diastereomeric mixture will be optically inactive. In either case it is common to find different amounts of the two diastereomers.



The formation of diastereomers is also possible when two new chiral centers are produced from achiral starting materials. A pertinent example is found in aldol-type reactions between enolates and carbonyl compounds. The achiral enolate and the achiral aldehyde or ketone gives a product with two new

chiral centers. Thus there can be two diastereomers produced, syn and anti, and because there is no initial chirality, each diastereomer will be produced as a racemic mixture of enantiomers. The syn and anti diastereomers will usually not be produced in equal amounts.



Factors which influence the stereoselectivity of organic reactions have been under intense investigation recently because of the increasing requirement and profitability of producing stereoisomerically pure compounds. A great deal of progress has been made, but even more remains to be accomplished. The specific contributors to stereoselectivity in individual reactions will be discussed as they are encountered. At this point it is important to be aware of the stereochemical variations that are possible.

# STEREOCHEMISTRY TO DEDUCE MECHANISM

In the above discussion of stereoselectivity the mechanisms of various reactions have been used to rationalize why some are stereoselective and some are not. Thus the bromination of olefins proceeds via a bridged bromonium ion intermediate and gives only trans addition across the double bond [reactions (6.2) and (6.3)]. In contrast, the addition of HBr across a double bond gives a carbocation intermediate that does not maintain the facial integrity of the olefin and is thus much less stereoselective [reaction (6.1)]. In these examples the mechanism of the reaction is used to explain and understand the diastereoselectivity that is observed. There are many other examples (usually in textbooks) where the mechanism of a reaction is used to rationalize the stereoselectivity of the process. To do this requires that the mechanism be known with certainty.

In most cases in the real world of chemistry both currently and historically, the reverse order is followed; that is, the mechanism is deduced with certainty only after the stereoselectivity has been determined. Because the stereochemical outcome of a chemical reaction is experimentally determined, it provides a powerful tool for examining the intimate spatial details of transition states and for determining how a reaction takes place at the molecular level. As such stereochemical studies have had a huge impact on the elucidation of reaction mechanisms.

In addition to bond breaking and charge buildup in the transition state, stereochemical changes during a reaction provide insight into the structural requirements of the activated complex of the rate-determining step. If a reaction is stereospecific, that is, if only one stereoisomer is formed in a reaction, then there is likely to be a particular spatial relationship between groups that is required for efficient product formation. (The key here is the term "efficient" because reactions can sometimes proceed if the correct spatial relationship is not obtainable, but they will go much more slowly.) If a reaction is stereoselective, that is, if one stereoisomer is the major but not exclusive product, then one particular spatial relationship is favored over another in the product-forming step. If the stereoselectivity of a reaction can be understood, then key structural elements in the activated complex can often be identified.



Several common examples show the power of this reasoning. The reaction of osmium tetroxide with olefins followed by reduction gives diols resulting exclusively from syn addition to the double bond. The reaction is hence stereospecific The addition stereochemistry is clearly seen in cyclic olefins, but it is also seen in acyclic olefins where single diastereomers are produced [reaction (6.4)].

The results from the cyclic series show that both oxygens come from the same side of the double bond, probably from a single  $OsO_4$  molecule. The results in the acyclic series demonstrate that both oxygens add to the ends of the double bond at the same time. If one oxygen added first, an intermediate with a single carbon–carbon bond would be formed which could isomerize by rotation around that bond. The observation of complete diastereoselectivity requires that both C–O bonds be formed simultaneously. Thus a concerted addition across the double bond is the most reasonable pathway consistent with these results. The stereochemical analysis is shown below for the cis starting material.



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#### 154 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

but



An analogous analysis for the trans olefin would predict only the d,l diastereomer for concerted addition of  $OsO_4$  to the  $\pi$  bond, but the same meso-d,l mixture would be obtained if the addition were stepwise.

Contrast the above syn addition of osmium tetroxide with the well-known anti stereochemistry found in the addition of bromine to alkenes. Cyclic systems give only trans addition in most cases, and acyclic olefins give single diastereomers that depend on the geometry of the starting olefin. These results are consistent with one bromine adding to one face of the olefin to give a bridged ion which maintains the stereochemistry of the original olefin. Bromide ion adds from the opposite face to give a single diastereomeric dibromide product.



*Walden inversion* was the term given to the change in stereochemistry observed in bimolecular nucleophilic substitutions. For example, reaction of (2S)-2-triflyloxyesters with sodium azide gives (2R)-2-azidoesters.



Inversion of configuration requires that the nucleophile adds electrons to the  $\sigma^*$  orbital of the carbon–triflate bond from the side opposite that bond. As required by the stereochemistry, formation of the bond from azide to carbon is concurrent with cleavage of the carbon leaving group bond.



If racemization were observed, it could only be due to a cleavage of one of the bonds to the chiral center prior to carbon-nitrogen bond formation or subsequent to it. This could occur by (a) enolization of the starting triflate, (b) an ionization of triflate to a carbocation and then nucleophilic attack by the azide, or (c) enolization in the azido product. The fact that clean inversion occurs means not only that the substitution by azide occurs with inversion but also that none of these other processes is significant under the reaction conditions since they would lead to racemized product.

(a)



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156 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM



The stereoelectronic requirements of groups undergoing base-promoted elimination is also easily seen by stereochemical studies. Treatment of *trans*-2-methylcyclohexyl tosylate gives 3-methylcyclohexene as the major product while treatment of *cis*-2-methylcyclohexyl tosylate gives the more stable 1-methylcyclohexene as the only product.



These data are consistent with the favored transition state having an antiperiplanar relationship between the proton being removed and the leaving group. In a sixmembered ring, this can only occur when they both are diaxial. In the trans isomer, the conformation in which the tosylate is axial only has a proton at C-6 axial and antiperiplanar. Thus elimination occurs across C-6 and C-1 to give only 3-methyl cyclohexene. In the cis isomer, the conformation in which the tosylate group is axial has antiperiplanar hydrogens at *both* C-2 and C-6. Elimination could proceed in either direction; however, removal of the proton at C-2 is favored because the more stable olefin product is produced.

These examples show the power of stereochemical information in pinpointing structural elements of activated complexes. Combined with other types of mechanistic information, even the most intimate mechanistic details can be clarified in many cases. For example, consider the solvolysis in ethanol of 3-phenyl-2-tosyloxy butane in which the replacement of the tosylate group by a solvent nucleophile is noted.

While this appears to be a simple substitution reaction, the details can be further explored. It was found that this reaction proceeded by a first-order rate law, which suggests an ionization pathway (Sn1) for the substitution. However, when a group of substituted aromatic compounds were investigated, plots of the rate constants (log  $k_Z/k_H$ ) gave a much better correlation with  $\sigma_Z^+$  than with  $\sigma_z$  and  $\rho^+ = -1.3$ . This Hammett study reveals that a positive charge is developed on the aromatic ring in the transition state of the rate-determining step. About the only way for this to happen is for the phenyl ring to interact with the positive charge produced by the ionization of the leaving group.

The use of the 2R,3R isomer led to formation of only 2R,3R-2-ethoxy-3phenylbutane. Thus the configuration at each chiral center was retained in the product. These stereochemical data rule out simple ionization and solvent capture as a reaction mechanism since this would lead to a mixture of 2R and 2S configurations. From these observations it has been postulated that the phenyl group assists ionization of the leaving group by electron donation to produce a bridged ion.



The bridged ion has a positive charge delocalized over the aromatic ring as required by the  $\sigma_Z^+$  correlation. Furthermore the solvent nucleophile can only add from the side opposite the bridging phenyl group, leading to retention of configuration as the stereochemical results demand.

Stereochemical studies can be an indispensable adjunct to other types of mechanistic investigations for unraveling the details of reaction processes. They allow the positions of atoms or groups in a molecule to be tracked through a reaction, thereby revealing the spatial requirements of the reaction.

# CONFORMATIONAL ANALYSIS

In the most basic sense chemical reactions are really only changes in the distribution of electrons. Such changes result in the breaking and making of chemical bonds and cause reactants to be converted to products. However, before such

electronic changes can take place, molecules taking part in the reaction must approach each other within bonding distance or they must undergo a change in geometry which permits overlap between the necessary orbitals to take place which results in electron redistribution. Restated another way, for chemistry to occur, molecules must first interact in a spatial sense. Consequently the shapes of molecules and the surface features they display are an important influence on their interactions with other molecules.

The large number of  $\sigma$  bonds present in organic molecules has a direct bearing on their shapes. Since a  $\sigma$  bond is axially symmetric along the bond, rotations of the groups connected by a  $\sigma$  bond do not cause it to break. (Such cannot be said of  $\pi$  bonds!) Thus molecules with many  $\sigma$  bonds are capable of large numbers of internal rotational motions which largely determine the shape, size, and surface characteristics of the molecule. Conformational analysis is the study of rotational motions in molecules and how they affect molecular properties.

The simplest hydrocarbon capable of internal rotational motion is ethane. Ethane has two tetrahedral methyl groups connected by a carbon–carbon  $\sigma$  bond. As such the methyl groups are free to rotate one relative to the other. However, it is found that the various rotational positions are not equivalent spatially or energetically. In fact, there are two limiting rotational positions for ethane. The lowest energy conformation (synonymous with a conformational isomer) is the one in which the C–H bonds of each methyl group are staggered between the C–H bonds of the other methyl group across the  $\sigma$  bond. This is the lowest energy conformation because the electron clouds of the bonds are the farthest distance apart, and their repulsions are minimized.



The highest energy conformation of ethane is the one in which the C–H bonds of each methyl groups are eclipsed with the C–H bonds of the other methyl group across the  $\sigma$  bond. This is the highest energy because the electron clouds of the C–H bonds are as close as they can be, and their repulsions raise the energy of the molecule.

The staggered and eclipsed forms of ethane are conformational stereoisomers (conformational isomers, conformers) because they have the same molecular formulas and sequences of bonded elements but different spatial arrangements due to rotations around single bonds. (Actually there are an infinite number of conformational isomers (also called conformations) because there are an infinite number of degrees of rotation around the bond, but normally one only needs to be concerned with energy minima and maxima.)

The difference in energy between the higher and lower energy forms of ethane is only 2.9 kcal/mol (12 kJ/mol); thus rotations around the bond are very rapid at

#### CONFORMATIONAL ANALYSIS 159



Figure 6.5 Energy changes during the rotation of ethane.



Figure 6.6 Energy changes during the rotation of propane.

room temperature (about  $10^{11} \text{ sec}^{-1}$ ). However, if one plots the change in energy as ethane rotates between the staggered and eclipsed forms, a periodic behavior is seen (Figure 6.5). Moreover, if a large number of snapshots of ethane were taken, they would show that most of the time ethane is found in the staggered conformation. The equilibrium between the staggered and eclipsed conformations favors the staggered by 99.2% to 0.8%.

A similar analysis of propane reveals analogous behavior with two major conformations—staggered and eclipsed—and periodic energy changes as rotation about a sigma bond occurs (Figure 6.6). There is a difference from ethane, however, in that the energy difference between the staggered and eclipsed conformations is now 3.3 kcal/mol (14 kJ/mol). This increase means that a hydrogen and methyl group eclipsed across a carbon–carbon bond repel each other more than two hydrogen atoms. This suggests that the electron cloud of the methyl group comes closer to the electron cloud of the C–H bond so the repulsion is greater. Since the electron clouds associated with the methyl group occupies more space than a hydrogen, that is, it is "larger." Thus, it follows that groups in molecules have definite sizes, and the size of these groups is one factor which contributes to the overall shape of the molecule because of its influence on the preferred conformation of the molecule.

Conformational isomerism around the central bond in butane is more complex because the various staggered and eclipsed conformations are not equivalent as they are in ethane and propane (Figure 6.7). Starting with the eclipsed

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## 160 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM



Figure 6.7 Conformational energy changes during rotation around the 2,3 bond of butane.

conformation with the dihedral angle between the two methyl groups at  $0^{\circ}$ , rotation around the central bond leads to two different staggered conformations and one additional eclipsed conformation.

The most stable staggered conformation is that in which the methyl groups are antiperiplanar (dihedral angle of  $180^{\circ}$ )—called the anti conformation. The other staggered conformation is that in which the dihedral angle between the methyl groups is  $60^{\circ}$ —called the gauche conformation (there are two of them for rotations of  $60^{\circ}$  or  $300^{\circ}$ , respectively). The other eclipsed conformation is that in which the two methyl groups each eclipse a hydrogen (there are two of them for rotations of  $+120^{\circ}$  and  $240^{\circ}$ , respectively).

From the energy diagram it is seen that the gauche conformation is 0.9 kcal/mol (3.7 kJ/mol) higher than the anticonformation. This must be due to some residual repulsion between the methyl groups when the dihedral angle is only 60° between them. Also the energy difference between the anti conformer and the highest energy eclipsed conformer is 4.5 kcal/mol (18.8 kJ/mol). Thus the greater effective size of the methyl groups results in increased repulsion when they are eclipsed.



In addition to conformational isomerism about the 2,3 bond in butane, rotations about the 1,2 bond and the 3,4 bond are possible. The energy changes here are much smaller and are comparable to those found in propane.

The importance of conformational isomerism lies in the fact that the predominant shape that molecules adopt is dependent on the energies of the various staggered and eclipsed conformations. In combination they can be used to predict the probable shapes the molecule normally assumes, and these shapes are those which are presented to reagents in solution.

In contrast to open-chain systems in which groups can rotate through  $360^{\circ}$  around  $\sigma$  bonds, cyclic systems can undergo conformational change through only limited ranges. Like open-chain systems, however, conformational changes in rings minimize eclipsing interactions across  $\sigma$  bonds. Cyclopropane is a flat ring without conformational motion. Cyclobutane is not planar because, if it were, all the C–H bonds around the ring would be eclipsed. The molecule undergoes a conformational change that bends the molecule out of planarity by about  $35^{\circ}$ . This reduces eclipsing and leads to a lower overall energy. A similar situation is found in cyclopentane, which adopts an envelope conformation (one ring apex out of plane) which is in equilibrium with four other envelope conformations (each apex up) to avoid the 10 C–H eclipsing interactions that would be present if the molecule were planar.



Saturated six-membered rings are the most common ring systems in nature because they present an optimal conformational situation. As seen in cyclohexane, the molecule adopts a puckered shape called a *chair conformation* in order to avoid angle strain in the ring bonds. In the chair form, all bond angles are  $109^{\circ}$  and all the bonds are staggered.

This results in a molecule whose energy is comparable to a completely staggered, open-chain alkane. This is easy to see by viewing the molecule in a Newman projection. Viewing the molecule through the ring so that the two side carbon–carbon bonds are seen head on, as in a Newman projection, generates the view in (6.5).

The chair conformer can undergo conformational isomerism to a second chair conformer which is degenerate in energy with the first. Cyclohexane is thus a dynamic molecule which exists largely in one of two chair isomers. These are the lowest energy conformations. Other higher energy conformations of cyclohexane include the boat form, which is 10.1 kcal/mol (42.3 kJ/mol) above the chair form, and the twist boat form, which lies 3.8 kcal/mol (15.9 kJ/mol) above the chair form.



Although these are well-defined conformational isomers, their energies are such that they are virtually unpopulated at room temperature. (The twist boat is an intermediate in the conversion of one chair form to the other.) At the same time the conversion of one chair form to the other occurs rapidly at room temperature, and both chair forms are in rapid equilibrium.

Because cyclohexane exists in the chair form, the C–H bonds of the methylene groups are nonequivalent. There are two types of valences on each  $CH_2$  group. One type is perpendicular to a plane loosely defined by the ring carbons and is called axial. The second type falls generally in the plane loosely defined by the ring carbons and is termed equatorial. These are shown both in combination and individually.



Three axial valences on alternate (1,3) carbons point to one side of the ring (up) and the other three axial valences on alternate (1,3) carbons point to the other side of the ring (down). The same is true for equatorial valences; while the directionality is not so obvious for equatorial valences; they still point toward one side of the ring (up) or the other (down).

There are two chair forms and two types of valences (axial and equatorial). The conversion of one chair form to the other interconverts the axial and equatorial valences (i.e., a valence which is axial in one chair form is equatorial in the other chair form and vice versa). In the structures below one of the carbons is indexed with a star ( $\star$ ) to help keep track of it.

#### CONFORMATIONAL ANALYSIS 163



In cyclohexane the chair forms have equal energy, but if groups other than hydrogen are attached to the cyclohexane ring, the two chair forms are no longer equivalent. In one chair isomer the group is equatorial and in the other chair isomer it must be axial. This is shown for methylcyclohexane.



Both conformations have the methyl group staggered between the vicinal protons. When the methyl group is axial, it is sufficiently close to the syn-axial protons to undergo 1,3 diaxial interactions and be repelled by them. This raises the energy of the axial conformer relative to the equatorial conformer. For a methyl group, the energy difference is about 1.8 kcal/mol. (Actually, the relationship of an axial methyl group to the ring bonds is a gauche conformational relationship. Thus the value of 1.8 kcal/mol for an axial methyl group is the value of two gauche butane interactions with the ring bonds!)



Other groups would behave similarly, with the axial isomer being higher in energy (less stable) than the equatorial isomer because of 1,3 diaxial interactions. These two isomers are conformational isomers because they are interconvertible by rotations about C–C single bonds, but they are also called conformational diastereomers since they have different physical properties and are nonsuperimposable, non-mirror images.

When more than one group is attached to cyclohexane, the stereoisomeric possibilities increase. First, structural isomers of the 1,2, 1,3, or 1,4 type are possible.

Next relative configurations (R,S) are possible for 1,2- or 1,3-disubstituted isomers. (The 1,4 isomer has a plane of symmetry.) The relative stereochemistry can be denoted as cis or trans, depending on whether the substituents point toward the same side or opposite sides of the ring. Finally, the cyclohexane ring can undergo chair–chair interconversion leading to different conformational isomers. These possibilities are shown for methylcyclohexanol in (6.6).

The first three types of isomerism are familiar and have been discussed previously. The conformational isomerism is very understandable if it is remembered that axial and equatorial valences exchange upon chair-chair interconversion. For example, to draw the trans isomer of 3-methylcyclohexanol, one of the groups must be equatorial and the other axial. The other chair form *must* have the groups in opposite valences. Similarly *trans*-2-methyl-cyclohexanol has both groups equatorial in one chair form. The other chair form must therefore have both groups axial.

# **CONFORMATIONAL ENERGIES**

The energetic consequences of chair-chair interconversions in substituted cyclohexanes are related to the interconversion of axial and equatorial valences. Because axial groups undergo 1,3 diaxial interactions which increase the energy of the molecule, the obvious energy preference is for groups larger than hydrogen to occupy equatorial positions. The relative energies of various chair conformers of multisubstituted cyclohexanes can thus be evaluated by noting the numbers of 1,3 diaxial interactions in each of the conformers.

Structural isomers:



#### 165 CONFORMATIONAL ENERGIES

Cis-trans isomers:



cis

cis





Conformational isomers:



For example, trans-1,2-dichlorocyclohexane has diaxial and diequatorial chair forms. The diaxial conformer should be less stable because it has two sets of 1.3 diaxial interactions between the chlorines and the axial protons. Knowing the equatorial-axial preference for a single chlorine substituent on a cyclohexane is 0.52 kcal/mol (2.2 kJ/mol), one can predict that the diequatorial isomer is favored by 1.04 kcal/mol (4.4 kJ/mol). cis-1,3-Dichlorocyclohexane has two chair forms with two equatorial chlorines and two axial chlorines, respectively. The diaxial isomer should be more than 1 kcal/mol higher in energy than the diequatorial isomer because the 1,3 diaxial interactions between two axial chlorines should be more severe than the 1,3 diaxial interactions between an axial chlorine and axial hydrogens. In contrast, trans-1,3-dichlorocyclohexane has a single axial chlorine in either chair conformer; thus the two chair forms are of the same energy.



Similarly trans-4-methylcyclohexanol has diequatorial and diaxial substituted chair forms. It is thus predicted that the former should be about 2.7 kcal/mol (11 kJ/mol) more stable than the latter because an axial methyl is less stable by

1.8 kcal/mol (7.3 kJ/mol) and an axial OH group is less stable by 0.9 kcal/mol (3.8 kJ/mol) than their equatorial counterparts.



Similar qualitative assessments can be made for more highly substituted cyclohexanes. It is found that *cis,cis*-1,3,5-trimethyl cyclohexane exists in only one chair form. This must be the triequatorial isomer because the other chair form has three axial methyl groups interacting on the same side of the ring. This should cause severe steric interactions and be much less stable than the all-equatorial isomer. In fact, the energy difference between the all-equatorial and the all-axial isomer should be greater than 5.4 kcal/mol ( $3 \times 1.8$  kcal/mol). This can be estimated by noting that an axial methyl group is less stable by 1.8 kcal/mol when the other axial valences with which it interacts are protons. The 1,3 diaxial interactions should be even greater if the other axial valences hold groups larger than protons; thus the energy difference should be greater than  $3 \times 1.8 = 5.4$  kcal/mol.

Since 1,3 diaxial interactions are the major factor which increases the energy of conformations with axial substituents, it is reasonable to expect that larger groups would have more severe steric interactions, causing the energy difference between the axial and equatorial position to increase. That is, the larger the group, the larger is the 1,3 diaxial interactions with axial protons and the greater is the energy difference between the axial and equatorial and equatorial forms.

This effect is clearly seen by comparing methylcyclohexane and t-butylcyclohexane. The axial-equatorial energy difference is 1.8 kcal/mol for the methyl group while it is 4.9 kcal/mol for the t-butyl group. This is because the t-butyl group is much larger than a methyl group, and 1,3 diaxial interactions are much stronger. In fact, these interactions are so large that the t-butyl group has been employed to anchor the particular chair conformation that has the t-butyl group and virtually unpopulated; thus the chemistry that is observed arises from reactions of a single conformation of the t-butyl-substituted ring.

# A VALUES

Viewed another way, if the axial-equatorial energy difference is mainly a function of steric bulk, then it might be used to assess the relative size of various groups. That is, if the energy difference between the two chair conformational isomers of a monosubstituted cyclohexane were measured, it might serve as a

Group	Α	Group	Α	Group	Α
HgCl	-0.25	Cl	0.52	$CH = CH_2$	1.7
HgBr	0	OAc	0.71	CH ₃	1.8
D	0.008	OMe	0.75	$C_2H_5$	1.8
CN	0.2	OH	0.94	<i>i</i> -Pr	2.1
F	0.25	$NO_2$	1.1	$C_{6}H_{11}$	2.1
$C \equiv CH$	0.41	COOEt	1.15	SiMe ₃	2.5
I	0.46	COOMe	1.3	$C_6H_5$	2.7
Br	0.55	COOH	1.4	t-Bu	4.9
OTs	0.52	NH ₂	1.4		

 Table 6.1
 A Values: Free-Energy Differences between Axial and Equatorial Conformations of Monosubstituted Cyclohexanes (kcal/mol)

quantitative measure of the effective steric bulk of a particular group. Table 6.1 is a collection of such data. The free-energy differences between equatorial and axial substituents on a cyclohexane ring are called A values and are a quantitative measure in kilocalories per mole of the effective steric bulk of a substituent. An important point is that these A values are not a measure of the physical size of a group, but rather are a measure of its steric interactions. Thus the *t*-butyl group (A = 4.9) is seen to be significantly more bulky than the trimethylsilyl group (A = 2.5), yet physically the trimethylsilyl group occupies more volume. The difference is that the carbon-silicon bond is longer than the carbon-carbon bond so that the trimethylsilyl group is farther away from the ring. Thus its effective bulk, which is the strength of 1,3 diaxial interactions, is actually less than the *t*-butyl group. The same trend is seen in the halides, where the A values for chloride (A = 0.52), bromide (A = 0.55), and iodide (A = 0.46) decrease even though the size of these atoms increase: Cl < Br < I. Thus A values are related to the effective steric bulk and not the actual physical size of substituents. In this respect they are very useful since it is clearly the *effective* size of substituents which gives rise to steric effects in chemical reactions. Thus A values can be used to predict steric changes resulting from the introduction of a group into a molecule. While the effective steric bulk of a group in a different molecule may not be quite the same as when it is attached to the cyclohexane ring, the trends should be parallel. In this way A values provide a useful way to evaluate steric effects semiguantitatively.

# **STRAIN IN RING SYSTEMS**

Molecules which contain rings comprise an important subset of organic compounds. In many respects their behavior and properties are identical to those of functionally analogous open-chain systems. In other respects, however, the greater structural order imposed by a ring causes the properties of cyclic compounds to be quite different than analogous open-chain systems. These differences
#### 168 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

arise from the need to accommodate preferred bond angles and conformations in structures which restrict the adoption of optimal geometries.

As we have seen previously, molecules with saturated carbon atoms have preferred bond angles of about  $109^{\circ}$  around the carbons and they are found primarily in staggered, anti conformations. This geometry allows the optimal overlap of all the interacting orbitals and yields the greatest amount of bond energy to be gained. Consequently the molecule has a lower energy than any other geometry and is thus the most stable.

It is possible to measure the heats of combustion for a series of saturated hydrocarbons and thereby determine how much energy is released  $(-\Delta H_{comb})$  when a -CH₂-group in a saturated hydrocarbon reacts with oxygen. Thus the heat of combustion of an open-chain methylene group is -157.4 kcal/mol. (The heat of combustion is negative because heat is evolved.)

$$-CH_2 - + \frac{3}{2}O_2 \longrightarrow CO_2 + H_2O \quad \Delta H_{comb} = -157.4 \text{ kcal/mol}$$

By a similar method, the heats of combustion of methylene groups in various ring compounds can be determined and the results are presented in Table 6.2. These data show that only cyclohexane and rings larger than  $C_{13}$  have methylene groups of the same energy as methylene groups in open-chain systems. Since in every case the products of combustion (1 CO₂ and 1 H₂O) are the same, any differences in the heats of combustion must reside in the methylene groups themselves. The value of 157.4 kcal/mol represents the least amount of energy that can be released by combustion since an open-chain methylene group is at the lowest energy possible for that unit. All other values are equal to or higher than 157.4 kcal/mol.

A value of  $-\Delta H_{\text{comb}}/-\text{CH}_2$ -unit = 157.4 kcal/mol (as in cyclohexane) indicates that a methylene group in that compound is as stable as one in an open-chain system and thus it is at the lowest possible energy. A value of  $\Delta H_{\text{comb}}/-\text{CH}_2$ -unit >157.4 mkcal/mol means that a methylene group in that ring system is *less stable* and therefore *of higher energy* than a methylene group in an open-chain compound by an amount equal to the difference between its  $\Delta H_{\text{comb}}$  and 157.4 [ $\Delta(\Delta H_{\text{comb}})$ ]. These differences are shown in the third column of Table 6.2. Finally, since each ring system contains a different number of methylene groups, the total increase in energy of the ring system due to all the methylene groups is found by multiplying the  $\Delta(\Delta H_{\text{comb}})$  by the number of methylene groups in the molecule. This is shown in the last column as total strain.

Strain can be defined as an increase in energy of a molecule (making it *less* stable) which results from any structural feature which causes nonoptimal overlap of atomic orbitals. Imperfect overlap leads to weaker bonds so less energy is derived from bond formation. Strain energy is the increase in energy due to imperfect overlap compared to the energy of an analogous system which has optimal overlap and the most bonding energy possible. This is shown for small rings in Figure 6.8.

Hydrocarbon	$-\Delta H_{\rm comb}$ /-CH ₂ -unit (kcal/mol)	$\Delta(\Delta H_{\rm comb})$ vs. Open Chain (kcal/mol)	Total Strain (kcal/mol)
Open chain	157.4	0	0
Cyclopropane	166.5	9.1	27.3
Cyclobutane	163.8	6.4	25.6
Cyclopentane	158.7	1.3	6.5
Cyclohexane	157.4	0	0
Cycloheptane	158.3	0.9	6.3
Cyclooctane	158.6	1.2	9.6
Cyclononane	158.8	1.4	12.6
Cyclodecane	158.6	1.2	12.0
Cycloundecane	158.4	1.0	11.0
Cyclododecane	157.8	0.4	4.8
Cyclotridecane	157.7	0.3	3.9
Cyclotetradecane	157.4	0	0

Table 6.2 Heats of Combustion of Methylene Groups in Cyclic Hydrocarbons

The methylene groups of cyclopropane are less stable by 9.1 kcal/mol than the methylene groups in open-chain hydrocarbons, and cyclopropane itself contains 27.3 kcal/mol of strain energy because there are three such methylene groups. Cyclopropane is a strained molecule because the three-membered ring requires that the bond angles at carbon be  $60^{\circ}$  when saturated carbon normally has bond angles of  $109^{\circ}$ . The distortion of bond angles from the normal tetrahedral geometry leads to increased energy or strain.

The smaller bond angles prevent normal overlap along the internuclear axis (dotted line). Overlap occurs outside the internuclear axis and a "bent" bond is formed (also referred to as a  $\tau$  bond). The  $\tau$  bond is weaker than a  $\sigma$  bond due to nonoptimal overlap and thus is of higher energy. This type of strain is termed *angle strain* (sometimes called *Baeyer strain*). It comes about because normal bond angles are not possible and thus nonoptimal overlap results. Angle strain



Figure 6.8 Strain in the methylene groups of small ring compounds.

#### 170 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

in cyclopropane is manifested chemically by greater chemical reactivity of the  $\tau$  bonds. Thus cyclopropane reacts with bromine, acids, and can be hydrogenated in stark contrast to an unstrained alkane, which is inert to these reagents.



Cyclobutane also exhibits angle strain since 90° bond angles are required by the four-membered ring instead of the preferred tetrahedral geometry. As expected, however, the amount of angle strain is less than in cyclopropane. The ring bonds of cyclobutane are still  $\tau$  bonds, but they are closer to the internuclear axis and the overlap is better. Consequently they are more stable (stronger) than the  $\tau$  bonds in cyclopropane. The greater bond strength is manifested by a lower chemical reactivity. Cyclobutane is more like an unstrained alkane and does not react with bromine or acids, although it can be hydrogenated under high pressure.



To achieve a bond angle of  $90^{\circ}$ , cyclobutane must be planar. This would force the adjacent C–H bonds to be eclipsed and would also raise the energy of the system. As a compromise, cyclobutane folds diagonally by  $35^{\circ}$ . While this raises the angle strain somewhat, it decreases the eclipsing interactions so that the lowest possible energy is attained. While each methylene group is less strained than one in cyclopropane, the total molecular strain is similar to that of cyclopropane.

If planar, cyclopentane should have minimal strain energy since the internal angles of a regular pentagon  $(108^{\circ})$  are very close to the normal tetrahedral angle. In fact, cyclopentane is a strained by 6.5 kcal/mol. If it were planar, all the hydrogen atoms around the cyclopentane ring would be eclipsed. This results in a second type of strain called *torsional strain*. Torsional strain is an increase in energy due to enforced eclipsing interactions. To reduce the torsional strain, cyclopentane adopts an "envelope" conformation in which one apex is bent out of the molecular plane by some 20°. This introduces some angle strain but

reduces the torsional strain and is the lowest energy structure that the molecule can achieve.



Any one of the five methylene groups of cyclopentane can be bent out of plane; thus cyclopentane is a dynamic system which undergoes a series of conformational changes between different envelope conformations.

The methylene groups of cyclohexane have the same  $\Delta H_{\rm comb}$  as methylene groups in open-chain systems and thus cyclohexane is unstrained. It was originally predicted by Baeyer that cyclohexane should have angle strain due to required bond angles of 120° for a planar structure. In fact, cyclohexane adopts a puckered conformation in which all the bond angles are 109° and all the hydrogens are perfectly staggered. The chair conformation of cyclohexane and its conformational ramifications have been discussed earlier.

It is also expected that rings larger than  $C_6$  could adopt puckered structures and thus maintain normal bond angles and staggered groups. As such they should be unstrained. In fact, it is seen in Table 6.2 that ring systems between 7 and 13 members are all strained to some degree. This strain results from the fact that puckering causes some hydrogens across the ring to approach each other closer than the sum of their van der Waals radii. These transannular interactions cause an adjustment in the conformational structure and raises the energy of the ring system. This increase in energy seen in medium-sized rings ( $C_7-C_{13}$ ) is called transannular strain. After the ring size has reached  $C_{14}$ , the interior of the ring is large enough to minimize transannular strain. Consequently the  $\Delta H_{comb}$  returns to the normal value of 157.4 kcal/mol found in unstrained systems.



It must be emphasized that *angle strain, torsional strain*, and *transannular strain* are simply terms invented by chemists to categorize structural features in rings which lead to increased energy (strain). The molecules themselves adopt structures which are the most stable and have the greatest amount of bonding energy. Nevertheless, strain is an important feature in molecules to keep in mind since it is manifested by altered chemical reactivity.

#### 172 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

For example, molecules which have significant angle strain such as cyclopropanes, epoxides, and aziridines undergo ring cleavage reaction very readily. Such reactions are very exergic because the strain energy is released upon ring opening. The exergic nature of these ring-opening reactions means that they have early transition states and generally have low activation energies. Conversely the formation of small rings requires that the strain energy must be added to the products in order for them to form.

Medium rings are difficult to form by intramolecular ring-closing reactions because the transannular strain in the products increases the energy of the transition state leading to those products. Thus the energy barrier to ring closure is raised and the intramolecular ring closure slows down significantly. Consequently intermolecular reactions (and not intramolecular ring-closing reactions) predominate. Transannular strain is not the only factor influencing these processes, but it is a major factor.

Saturated six-membered rings are easy to produce because they are unstrained. They are also resistant to ring-opening reactions for the same reason. Consequently saturated six-membered rings are the most common ring systems in the natural world.

## STEREOELECTRONIC EFFECTS

In addition to determining its shape and surface characteristics, the conformational preferences of a molecule can also contribute to its chemical reactivity. Many reactions require that reacting groups achieve a particular spatial relationship so that overlap of appropriate orbitals leading to the needed electron redistribution can take place. These geometry-dependent orbital interactions which influence chemical reactivity are described generally as stereoelectronic factors. A very well known example of a reaction with distinct stereoelectronic requirements is the Sn2 reaction. Here the incoming nucleophile must approach from the side opposite the leaving group. This permits electron donation into the  $\sigma^*$ antibonding orbital and results in the inversion stereochemistry found for these processes. Structural features which prevent the stereoelectronic requirements of the reaction from being met, such as the cage structure of the norbornyl skeleton which prevents the incoming nucleophile from approaching from in back of the leaving group, will slow or prevent the reaction entirely.

$$Nu: \longrightarrow_{U'} C - X \rightarrow \left[ Nu - C - X \right]^{\pm} \longrightarrow Nu - C'_{U_1} + X \quad but \qquad X \\ unreactive$$

In other reactions a particular disposition of groups in a molecule is required for the reaction to proceed efficiently. Moreover it is often found that the proper

#### STEREOELECTRONIC EFFECTS 173

stereoelectronic requirements of that reaction can be met only if a particular conformation of the molecule is populated. If the needed conformation is energetically accessible and thus populated, the reaction can proceed normally; if not, it is very slow and alternate processes might intervene. Classic examples are E2 reactions which require an anti-periplanar relationship between the proton being removed and the leaving group which departs. In open-chain systems this rarely presents a problem since the barriers to rotations about single bonds are low and reactive conformers are easily populated. Nevertheless this stereoelectronic requirement can have stereochemical consequences. d.l-Stilbene dibromide undergoes dehydrohalogenation in hot pyridine whereas the meso isomer reacts much more slowly. As seen below, the reactive conformation of the d.l isomer has the bulky phenyl groups anti to each other and is energetically favored, whereas the reactive conformation of the meso isomer has the phenyl groups gauche to each other and is energetically unfavorable. Thus the reactive conformer of the d,l compound is populated and reacts effectively; the reactive conformer of the meso compound is not populated significantly and the rate of elimination is much slower. (Eclipsing of the phenyl groups in the transition state also slows the reaction of the meso isomer.)



In cyclic systems where conformational motions are more restricted, stereoelectronic effects can play a much larger role in determining the outcomes of reactions. For example, base-promoted elimination in *cis*-4-(*t*-butyl)-cyclohexyl tosylate occurs 70 times faster than in the trans isomer. The reason is that the *t*-butyl group controls the conformation of the cyclohexane ring (it occupies an equatorial valence nearly exclusively); consequently, the cis isomer has an axial tosylate group which is antiperiplanar to axial hydrogens in the  $\beta$  positions. The trans isomer has an equatorial tosylate group which has no antiperiplanar hydrogens. The cis isomer meets the stereoelectronic requirements for basepromoted elimination and thus reacts significantly faster than the trans isomer, which does not.



The pyrolysis of acetate esters yields olefins by a concerted syn elimination of acetic acid. In open-chain systems where rotations are facile, it is possible for the

#### 174 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

acetate group to achieve a syn relationship with any of the vicinal protons, and the major product primarily reflects the energies of the products (trans is favored).



In contrast, it is found that *trans*-1-acetoxy-2-phenylcyclohexane gives 1-phenylcyclohexene as the major product (86.5%) upon heating, whereas *cis*-1-acetoxy-2-phenylcyclohexane gives 3-phenylcyclohexene as the major product (93%). Comparing these two isomers, it is calculated from *A* values that the trans diequatorial isomer is favored by 3.4 kcal/mol over the trans diaxial isomer. The diequatorial conformer has a proton at C-2 syn to the acetate group in an axial–equatorial disposition which undergoes elimination to produce the more stable conjugated product. (In this case, even the less stable diaxial conformer has a syn proton available in an axial–equatorial relationship with the acetate group and gives the same product.) In contrast, the cis isomer has only one conformation with a syn proton in an axial–equatorial relationship with the acetate group, and that elimination gives the less stable, nonconjugated 3-phenyl cyclohexene as the major product.



but



A more striking example of the influence of conformation on the reaction outcome is seen in the nitrous acid deamination of 2-aminocyclohexanols which takes place by rearrangement of a group on the carbinol carbon that is anti to the developing carbocation. The deamination reaction is very fast and the products reflect the population of the chair conformers. The trans isomer exists mainly in the diequatorial conformer; thus the only group anti to the amino group is a ring bond. Indeed ring contraction is the only process observed. In the cis isomer, however, both chair forms are populated (the A values of OH and NH₂ are similar), so products of *both* ring contraction and hydride migration are obtained.



but



A particularly revealing example is seen in the reaction of 2-bromo-4phenylcyclohexanols with Ag[I]. In the secondary carbinol, reaction takes place from the higher energy conformer because, even though its population is low, hydride migration by the hydrogen anti to the bromide assists the loss of axial bromide so it is the fastest reaction. Placement of a methyl group at the carbinol carbon raises the energy of the axial bromide conformation even higher, and its population is further reduced. Now because of its greater relative concentration, the conformer with bromide in the equatorial position reacts faster and leads to ring contraction.



The examples presented above point out some important features about organic reactions. First, many have distinct stereoelectronic requirements that must be met

176 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM

if the reaction is to proceed efficiently. Second, the correct stereoelectronic relationships are primarily dependent on the conformations of the substrate. Finally, the populations of various conformers determine if stereoelectronic requirements can be satisfied and thus play a significant role in product partitioning.

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#### PROBLEMS

**6.1.** Indicate the chiral centers in the following molecules and give the relative configuration (R,S) of each:





**6.2.** The following structures are representations of 3-fluoro-2-phenyl-2-pentanol. Give the stereochemical relationship of each structure to (2R,3R)-3-fluoro-2-phenyl-2-pentanol:



- **6.3.** (a) Draw the four stereoisomers of 4-methyl-2-hexanol and give the relationship of each to the others.
  - (b) Draw all the stereoisomers of 3-bromo-4-methylhexane, give the R,S designation of each chiral center, and give the relationship of each to the others.
  - (c) Using Fischer projections, draw all of the stereoisomers of 2-fluoro-3-methyl-1,4-pentanediol and give the relationship of each to the others.
  - (d) Draw all of the stereoisomers of 1,4-diphenyl-1,4-dibromobutane, give the R,S designation of each chiral center, and give the relationship of each to the others.
- **6.4.** Give the stereochemical relationship between the following pairs of compounds:

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#### 178 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM



- **6.5.** For the following compounds, show two chair conformations, indicate which is more stable, and give an estimate of the energy difference between the two:
  - (a) *trans*-1-ethyl-3-phenylcyclohexane
  - (b) *cis*-1-(*tert*-butyl)-4-isopropylcyclohexane
  - (c) *trans*-2-amino-1-cyanocyclohexane
  - (d) (2R,6S)-1-bromo-2,6-dimethylcyclohexane
  - (e) *cis*-4-(*tert*-butyl)-2-methylcyclohexanone
  - (f) *cis*-1,1,3,4-tetramethylcyclohexane
- **6.6.** Estimate the difference in energy between the chair conformations of *trans*-2-methoxycyclohexanol. The actual value is about 3.1 kcal/mol. Can you explain this?
- **6.7.** Show all of the staggered conformers of 2,3-dimethylbutane and estimate the energy differences between them.
- **6.8.** The Beckman rearrangement could occur by either a stepwise or a concerted mechanism.

$$R \xrightarrow{N^{\circ}OH}_{R'} \xrightarrow{I. TsCl, py, } R \xrightarrow{O}_{H} R' + R' \xrightarrow{H} R \xrightarrow{O}_{H} R \xrightarrow{CH_3CH_2}_{H^{\circ}CH_3} \xrightarrow{H}_{CH_3CH_3}^{N^{\circ}OH}_{CH_3}$$

- (a) Show both mechanisms using curved-arrow notation.
- (b) Suppose you had made oxime 1.
  - 1. Would it rotate plane-polarized light?
  - 2. Label the configurations of the chiral centers in 1.
  - **3.** Show how **1** could be used to help distinguish the mechanisms you have given.
- **6.9.** Explain why (1S,3R)-3-*tert*-butylcyclohexyl tosylate undergoes E2 elimination with potassium *tert*-butoxide very slowly while the (1R,3R) reacts much more rapidly.
- **6.10.** The reaction of *cis*-2-pentene with iodine azide (IN₃) in dichloromethane gives (2S,3S)-3-azido-2-iodopentane and (2R,3R)-3-azido-2-iodopentane but not any other diastereomers. What is the stereochemistry of the addition and give a curved-arrow mechanism to account for it.

- 180 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM
- **6.11.** The reaction of *trans*-2-hexene with aqueous peracetic acid gives (2S,3R)-2,3-hexane diol and (2R,3S)-2,3-hexanediol but not any other diastereomers. What is the stereochemistry of the addition?
- **6.12.** Heating (2S)-3-methyl-3-phenyl-2-butyl tosylate in ethanol leads to skeletal rearrangement and the formation of (3S)-2-ethoxy-2-methyl-3-phenylbutane. What does this information tell you about the stereoelectronic course of the skeletal rearrangement?
- **6.13.** Treatment of *trans*-2-phenylcyclohexyl tosylate with potassium *tert*-butoxide gives mainly 3-phenylcyclohexene in a fairly slow process, whereas under the same conditions *cis*-2-phenylcyclohexyl tosylate gives 1-phenylcyclohexene in a much shorter reaction time. Explain this difference.
- **6.14.** What do the following strain energies suggest about the origin of strain in three-membered rings?



**6.15.** Although cyclobutane is a puckered molecule (by about 25°), its oxygen analog oxetane is virtually flat. Give a rationale for this behavior.



**6.16.** The strain energy of spiropentane (62.5 kcal/mol) is more than twice that of cyclopropane (27.3 kcal/mol). Suggest an explanation.



**6.17.** Based on the properties of the cyclohexane ring, which of these isomers is predicted to have a larger dipole moment? Explain your choice.



- **6.18.** Draw the conformational isomers of *cis*-1,2-dimethylcyclohexane and *cis*-3,4-dimethylcyclohexanone. While the cyclohexane conformers are of equal energy, the cyclohexanone conformers are not. Indicate which conformer is favored and explain why.
- 6.19. Addition of osmium tetroxide to norbornene 2 followed by reductive cleavage with sodium sulfite gives the exo,exo diol 3. The same reaction sequence carried out on 7,7-dimethylnorbornene 4 gives endo,endo diol 5. From these results deduce the mechanism of the addition and facial selectivity for these two substrates.



**6.20.** Treatment of *E*-1-phenyl-2-butene **5** with  $I_2$  and silver benzoate (1:2) followed by saponification gives an equal mixture of [2S,3R]-1-phenyl-2,3-butanediol and [2R,3S]-1-phenyl-2,3-butanediol. Treatment of **5** with  $I_2$  and silver acetate in the presence of water followed by saponification gives an equal mixture of [2R,3R]-1-phenyl-2,3-butanediol and [2S,3S]-1-phenyl-2,3-butanediol. Determine the stereochemistry for these two processes. Can you account for the difference mechanistically?



**6.21.** Treatment of [4S]-4-*t*-butyl-1-methylcyclohexene with borane–THF followed by oxidation with  $H_2O_2$ –NaOH gives a mixture of [1S, 2S, 5S]-5-*t*-butyl-2-methylcyclohexan-1-ol and [1R, 2R, 5S]-5-*t*-butyl-2-methylcyclohexan-1-ol and no other diastereomers. There are two steps in this process—addition to give an organoborane and oxidation which cleaves the carbon–boron bond to an alcohol.

182 STEREOCHEMICAL AND CONFORMATIONAL ISOMERISM



[4S]-4-t-butyl-1-methylcyclohexen

- (a) Is the overall process stereospecific?
- (b) What are the stereoselectivities of each step (mode of addition and stereochemistry of cleavage) which are consistent with the observed products?
- (c) It is known that the oxidation reaction occurs with retention of configuration. What then must be the stereochemistry of the hydroboration step?
- **6.22.** Treatment of [2R,3R]-2,3-dibromo-3-methylpentane with Zn gives (Z)-3-methyl-2-pentene as the only product. What is the stereochemistry of the reduction? Based on this result could the reaction of a trans olefin with Br₂ and then Zn be used as a way to convert trans olefins to cis olefins?

# 7

# FUNCTIONAL GROUP SYNTHESIS

Functional Group Manipulation	183
Carboxylic Acids	185
Esters	188
Amides	190
Acid Chlorides	191
Aldehydes	192
Ketones	194
Imines and Imine Derivatives	197
Alcohols	198
Amines	201
Alkenes	203
Alkanes	207
Bibliography	208
Problems	209

# FUNCTIONAL GROUP MANIPULATION

As we have seen in earlier undergraduate organic coursework, organic chemistry is often organized around the chemistry of functional groups. Functional groups are recurring sequences of bonded elements which give typical and characteristic chemical reactions. Recall that there are characteristic reactions of ketones, esters,

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#### 184 FUNCTIONAL GROUP SYNTHESIS

secondary alcohols, and so on. Consequently, it is very important to be able to introduce a particular functional group into a molecule or to interconvert functional groups in a molecule in order to utilize particular reactions available to them.

To manipulate functional groups, it is first necessary to be able to identify them, understand their bonding and oxidation levels, and recognize the bonding changes that are needed to convert one functional group into another. A variety of functional group reactions are available, and much of the introductory organic chemistry course is devoted to learning reagents and reactions for carrying out functional group interconversions.

Very often these reactions are traditional and illustrative, but they are not necessarily the best way to manipulate a particular functional group. Many traditional methods have been replaced, in practice, by newer reactions or reagents which offer certain advantages over older methods. In general, these advantages have to do with mild conditions, selectivity, generality, and/or experimental simplicity. Nevertheless all types of functional group interconversions, new or old, are still based fundamentally on the ideas that have been developed earlier in this book.

This chapter is not meant to serve as a surrogate for an undergraduate text as it does not cover all the functional groups covered in the introductory course. Instead it concentrates on the most common ones. It is assumed that the functional group transformations taught at the undergraduate level have already been mastered. If this material has not been learned, it should be reviewed using a standard undergraduate text.

The discussion which follows is organized in the following fashion. First a common undergraduate text was used to provide a list of "standard" or "traditional" preparations for the major functional groups dealt with herein. Most of these reactions should be familiar because they are the ones learned (or not learned) in the undergraduate course in organic chemistry. Although these methods will be listed and perhaps discussed briefly, the discussion in no way serves as a review of these methods.

The main focus of this chapter will be to introduce the most widely used and practical ways (or "real" ways) to introduce the major functional groups. These latter methods have practical synthetic value and are usually the first choices in real laboratory situations, but often they differ from the standard list of preparations. What is important is that these first-choice methods must be integrated into the methods previously encountered so that a wider view of how to manipulate functional groups is achieved.

The functional group order goes from highest oxidation level (carboxylic acid) to lowest oxidation level (alkanes). As might be anticipated, carboxylic acids are most often prepared oxidatively and alkanes are most often prepared reductively. Functional groups of intermediate oxidation level can be accessed either reductively or oxidatively. Furthermore the methods discussed for each functional group are organized by the oxidation level change that is used, that is, oxidative methods followed by methods requiring no change in oxidation level followed by reductive methods.

#### CARBOXYLIC ACIDS

Carboxylic acids have a relatively high oxidation level (+3), and thus a majority of synthetic methods to access carboxylic acids are oxidative in nature. Traditional preparations include the following:

Oxidation of olefins

$$R-CH=CH-R' \xrightarrow[b]{a. KMnO_4, heat} RCO_2H + R'CO_2H$$
(7.1)

Oxidation of primary alcohols

$$R-CH_2OH \xrightarrow[b. Na_2Cr_2O_7, H_2SO_4]{a. KMnO_4, OH} RCO_2H$$
(7.2)

Oxidation of alkylbenzenes

$$R \xrightarrow{KMnO_4, OH^-} CO_2^{\Theta} \xrightarrow{H^+} CO_2H (7.3)$$

Hydrolysis of nitriles

$$R-C \equiv N \xrightarrow[b. OH^-, H_2O, heat]{a. conc. HCl, heat} RCO_2H + NH_3$$
(7.4)

While the above reactions will provide carboxylic acid products, each has problems associated with it. The cleavage of olefins to carboxylic acids [reaction (7.1)] can be carried out using potassium permanganate or by ozonolysis at low temperature followed by oxidative workup with hydrogen peroxide. Neither of these methods is very useful since only symmetric olefins provide a single carboxylic acid product. Unsymmetrical olefins give a mixture of two acids which must be separated. Furthermore the most useful synthetic processes are those which build up structures, whereas these reactions are degradative in nature.

Primary alcohols can be oxidized to carboxylic acids by a variety of reagents [reaction (7.2)]. Often potassium permanganate or sodium dichromate were given as reagents to use in this transformation. These are powerful oxidants, and many other functional groups that might be present cannot survive the reaction conditions. Milder oxidants are preferred and the best of these is chromic acid in acetone (Jones reagent). Jones reagent is a mixture of chromic acid and a stoichiometric amount of sulfuric acid which is needed in the redox process to keep the solution at near a pH of 7. This technique is fast, easy, and efficient and the

#### **186** FUNCTIONAL GROUP SYNTHESIS

reagent solution is easily prepared from chromium trioxide and sulfuric acid in acetone. The oxidation can be carried out by adding the Jones reagent by burette to the alcohol. Oxidation is instantaneous and the addition can be stopped precisely when all the alcohol has been consumed. Using a stoichiometric amount of chromic acid usually leaves other functional groups untouched. This is the method of choice for the synthesis of carboxylic acids from primary alcohols.

The oxidation of alkyl benzenes to benzoic acids [reaction (7.3)] is still carried out occasionally, and this oxidation is most likely the only one where potassium permanganate is the reagent of choice. Any carbon group attached to the aromatic ring is degraded to the carboxylic acid group under the very vigorous conditions of this oxidation.

An interesting twist on the oxidation of aromatic compounds forms the basis of a new and very useful synthesis of carboxylic acids. Normally the aromatic ring is resistant to oxidation and the side chains are oxidatively degraded to carboxylic acids, as in reaction (7.3). It has been found that ruthenium tetroxide is a mild and selective oxidant of aromatic rings and completely degrades the ring to the carboxylic acid but leaves aliphatic groups unoxidized. This is essentially the reverse of the chemoselectivity seen in potassium permanganate oxidations of arenes, where the side chains are oxidized but the aromatic ring is left intact. The selectivity and mildness is seen in the following example in which no amide or silyl ether (OTMS) oxidation was observed and there was no epimerization of either chiral center:



Another common way to install a carboxylic acid group is to hydrolyze a carboxylic acid derivative. Such hydrolyses do not require a change in oxidation level (carboxylic acid derivatives are at the oxidation level of the acid itself), but they do normally require acid or base catalysis. Nitriles [reaction (7.4)] often require vigorous conditions for hydrolysis mainly because they are only weakly electrophilic. Either concentrated hydrochloric acid or sodium hydroxide can be used to hydrolyze nitriles. The first stage of the hydrolysis produces an amide and the amide is subsequently hydrolyzed to the acid. Each step of the hydrolysis requires strenuous conditions and is useful mainly for nitriles that lack other functional groups that would be destroyed by the stringent conditions. It has been found that a mixture of sodium hydroxide and hydrogen peroxide can be used to hydrolyze nitriles more efficiently than sodium hydroxide alone and is the reagent of choice, although many other reagent combinations have been reported.

Amides are stabilized by resonance and are thus difficult to hydrolyze. Like nitriles they can be hydrolyzed by concentrated hydrochloric acid or concentrated sodium hydroxide. These are powerful reagents that tend to react with many other functional groups as well. Thus hydrolysis is not a satisfactory method for amides with many other functional groups present. As with nitriles, a mixture of sodium hydroxide and hydrogen peroxide is one of the more effective reagents for hydrolysis.

Esters, on the other hand, are very common hydrolytic precursors to carboxylic acids. The traditional reaction for the hydrolysis of esters is basic saponification using sodium hydroxide or potassium hydroxide. While acid catalysis can also be employed, preparative methods usually use base catalysis because formation of the carboxylate salt drives the reaction to the right and gives high yields of products.

$$R \xrightarrow[]{O} OR' \xrightarrow[]{a. NaOH, H_2O} R-CO_2 \stackrel{\bigoplus}{M} + R'OH$$

While esters are much more easily hydrolyzed than amides, traditional saponification suffers from the fact that most esters are not soluble in aqueous base and so the rate of the hydrolysis is limited by the solubility, not by the reactivity. This limitation is overcome by the use of lithium hydroxide in aqueous THF, the reagent of choice for basic hydrolysis of esters. Methyl and ethyl esters are cleaved readily by this combination as most esters are soluble in this solvent mixture.

R-CO₂R' 
$$\xrightarrow{\text{LiOH}}$$
 R-CO₂ Li + R'OH  
(R' = Me, Et)  $\xrightarrow{1-2 \text{ hr}, 25 \,^{\circ}\text{C}}$  90–100%

If structural constraints prevent the use of basic hydrolysis of the ester group, acid hydrolysis must be used. Nowadays it is much more common, in such instances, to use *tert*-butyl esters because they are cleaved rapidly and efficiently by trifluoroacetic acid (TFA) to the carboxylic acid and isobutylene. This cleavage is different from the normal acid-catalyzed hydrolysis of esters in that the alkyl–oxygen bond is broken rather than the acyl–oxygen bond. This change in mechanism is brought about by the stability of the *tert*-butyl cation which is produced upon alkyl–oxygen cleavage. As an added benefit, the isobutylene by-product is a gas which escapes from the reaction mixture.



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#### **188** FUNCTIONAL GROUP SYNTHESIS

If neither acidic nor basic conditions are compatible with other groups present in the ester to be hydrolyzed, then  $\beta$ -trimethylsilylethyl esters are often prepared. Trimethylsilylethyl esters are cleaved easily by fluoride under mild, neutral conditions. Typical sources of fluoride are cesium fluoride (CsF) or the more soluble tetrabutylammonium fluoride (TBAF).

$$R \xrightarrow{O} O CH_2CH_2 \xrightarrow{Me} Me + F^- \rightarrow R \xrightarrow{O} O + CH_2 = CH_2 + F \xrightarrow{K} O + Me$$

Current methods for the hydrolysis of esters are fast, efficient, and sufficiently mild that they are compatible with the presence of a variety of other functional groups and/or stereocenters in the molecule. For example, protected amino acid esters are hydrolyzed quantitatively without racemization or deprotection by LiOH in aqueous THF.



Carboxylic acid groups can also be installed in molecules using the reaction of an organometallic compound with carbon dioxide. This is a reductive method since the carbon dioxide is reduced to a carboxylic acid by formation of a new carbon–carbon bond. Both Grignard reagents and organolithium compounds work well in this reaction.

#### ESTERS

Derivatives of carboxylic acids are generally made from the carboxylic acid. The traditional routes to esters are as follows:

1. Acid-catalyzed reaction between an acid and an alcohol (Fischer esterification):

$$R \stackrel{O}{\longrightarrow} OH + R'OH \stackrel{H^+}{\longleftarrow} R \stackrel{O}{\longrightarrow} OR' + H_2O$$

2. Reaction of an acid chloride (or acid anhydride) with an alcohol:

$$R - CO_2H \longrightarrow RCCI \xrightarrow{R'OH} R - C - OR'$$

While still useful for large-scale esterification of fairly robust carboxylic acids, Fischer esterification is generally not useful in small-scale reactions because the esterification depends on an acid-catalyzed equilibrium to produce the ester. The equilibrium is usually shifted to the side of the products by adding an excess of one of the reactants—usually the alcohol—and refluxing until equilibrium is established, typically several hours. The reaction is then quenched with base to freeze the equilibrium and the ester product is separated from the excess alcohol and any unreacted acid. This separation is easily accomplished on a large scale where distillation is often used to separate the product from the by-products. For small-scale reactions where distillation is not a viable option, the separation is often difficult or tedious. Consequently Fischer esterification is not widely used for ester formation in small-scale laboratory situations. In contrast, intramolecular Fischer esterification is very effective on a small scale for the closure of hydroxy acids to lactones. Here the equilibrium is driven by the removal of water and no other reagents are needed. Moreover the closure is favored entropically and proceeds easily.



A second very common way to convert carboxylic acids to esters is by the reaction of the corresponding acid chloride with an alcohol. A tertiary amine such as pyridine or triethylamine is used to scavenge the HCl by-product. It has also been found effective to add small amounts of N,N-dimethyl-4-aminopyridine (DMAP) to the reaction mixture in order to promote efficient product formation. If the acid chloride is readily available, this is a very satisfactory preparation. If the acid chloride is not available, a disadvantage to this method is that a carboxylic acid must first be converted to the acid chloride, which must be isolated and purified prior to the formation of the ester. If the chlorinating agent is not separated from the acid chloride, the alcohol will also react with the chlorinating agent leading to a mixture of products that may be difficult to separate.

$$R - CO_{2}H \longrightarrow \bigcap_{\substack{||\\ RCCl}}^{O} \xrightarrow{R'OH}_{\substack{R'OH\\ DMAP(caL)\\ CH_{2}Cl_{2}}} O \\ R - C - OR' + R_{3}NH^{\oplus}Cl^{\ominus}$$

#### 190 FUNCTIONAL GROUP SYNTHESIS

For small-scale esterification reactions (<500 mg), the best methods should occur rapidly under mild conditions and only produce by-products which are easily separated from the reaction products. Under these criteria an extremely efficient way to convert acids to methyl esters is to titrate the carboxylic acid with an ethereal solution of diazomethane. The methyl ester is produced rapidly and quantitatively, and the by-product of the esterification is nitrogen. Although diazomethane is a reactive and explosive compound, solutions of diazomethane can be prepared from readily available reagents and used safely in the laboratory.

$$R \xrightarrow{O} OH \xrightarrow{CH_2N_2} R \xrightarrow{O} OCH_3 + N_2$$
  
guantitative

A second method to efficiently produce methyl esters of carboxylic acids is to treat the acid with potassium carbonate and methyl iodide. The methyl ester is produced under mild conditions and is easily separated from the reaction byproducts. This method is somewhat different in that the ester is formed by a nucleophilic displacement of iodide by the carboxylate ion. Normally carboxylates are not thought of as good nucleophiles—and they are not—but methyl iodide is a quite reactive electrophile which matches the poor nucleophilicity of the carboxylate satisfactorily.

$$R \xrightarrow{O} OH \xrightarrow{K_2CO_3} R \xrightarrow{O} OCH_3 + KI$$

Besides the above methods, many other satisfactory ways to convert acids to esters are commonly encountered.

#### AMIDES

Amides are usually obtained from carboxylic acids or their derivatives. The traditional method of preparation of amides is to react the corresponding acid chloride with an amine.

$$R - CO_2 H \longrightarrow \begin{array}{c} O \\ \parallel \\ RCCl \end{array} \xrightarrow{R'NH_2} \begin{array}{c} O \\ R - C \\ - CH_2 Cl_2 \end{array} \xrightarrow{R'NH_2} \begin{array}{c} O \\ R - C \\ - NHR' \end{array} + R'NH_3^{\oplus} Cl^{\ominus}$$

This substitution process replaces the chloride with an amine without a change in the oxidation level. This remains an excellent and efficient method. However, excess amine or another base is required to neutralize the equivalent of HCl produced by the substitution. In this approach, formation of an acid chloride is required to activate the carbonyl group and make it more electrophilic. This activation is required in most transformations of carboxylic acids because carboxylic acids themselves are not sufficiently electrophilic to react with most nucleophiles. Furthermore, since many nucleophiles are also basic, they can react with carboxylic acids to give the carboxylate ion, which is an even poorer electrophile than the carboxylic acid itself.

Carbodiimides are increasingly being used to promote the conversion of carboxylic acids to amides. This method was originally developed for creating the amide bonds in peptides. In this method the carboxylic acid is treated with a carbodiimide [a very common one is dicyclohexylcarbodiimide (DCC), although many others have been developed]. An activated acylating agent is produced which reacts with the amine present in the mixture to produce an amide. The advantage of this approach is that the acid is activated to a reactive electrophile in situ so the activated species need not be isolated. Yields are normally high and the urea by-product from the carbodiimide can be separated and removed from the amide by one of several methods.

$$\begin{array}{c} O \\ H \\ R - C - OH \end{array} \xrightarrow{ \begin{array}{c} C_6H_{11}N = C = NC_6H_{11} \\ (DCC) \end{array}} \begin{array}{c} O \\ H \\ R'NH_2 \end{array} \xrightarrow{ \begin{array}{c} O \\ H \\ R'NH_2 \end{array}} \begin{array}{c} O \\ R - C - NHR' + C_6H_{11}NHCNHC_6C_{11} \\ urea \end{array}$$

Amides are also available from nitriles, which have the same oxidation level. Direct acid or base hydrolysis of a nitrile usually requires fairly severe conditions and often does not stop at the amide stage but goes on the carboxylic acid. Treatment of nitriles with a solution of HCl in ethanol furnishes an imidate ester which is hydrolyzed in aqueous acid to the amide. Because a nitrile is the starting material, only primary amides can be produced by this process.

$$R - C \equiv N \xrightarrow{HCI}_{EtOH} \begin{array}{c} OEt \\ R - C \equiv NH \end{array} \xrightarrow{H_{3}O} \begin{array}{c} O \\ H_{3}O \\ R - C = NH \end{array} \xrightarrow{H_{3}O} \begin{array}{c} R \\ R - C - NH_{2} \end{array}$$
imidate ester

#### ACID CHLORIDES

Traditional textbook preparations of acid chlorides from carboxylic acids include

$$\begin{array}{ccccccc} O & O & O \\ R - C - OH & \xrightarrow{SOCl_2} & R - C - CI + HCI + SO_2 \\ \end{array}$$

$$\begin{array}{cccccccc} O & O \\ 3 & R - C - OH & \xrightarrow{PCl_3} & 3 & R - C - CI + H_3PO_3 \\ \end{array}$$

$$\begin{array}{ccccccccc} O & O \\ R - C - OH & \xrightarrow{PCl_5} & R - C - CI + POCI_3 + HCI \end{array}$$

#### 192 FUNCTIONAL GROUP SYNTHESIS

The traditional methods utilize sulfur or phosphorous halides to convert the acid to the acid chloride. Of these methods, thionyl chloride [often with a catalytic amount of dimethyl formamide (DMF)] is the most useful since the by-products of the reaction are gases (SO₂, HCl) which can be easily purged from the reaction mixture with a stream of nitrogen. The acid chloride product can then be purified on a small scale by bulb-to-bulb distillation or crystallization. Because an excess of thionyl chloride is usually used, there must be a purification step to remove the excess reagent.

Another superior reagent for the preparation of acid chlorides is oxallyl chloride in methylene chloride. Addition of a carboxylic acid leads to the smooth evolution of gas (CO₂, CO, HCl) which can be used as a crude monitor of the reaction progress. The acid chloride is very easily purified since oxallyl chloride boils at  $62^{\circ}$ C and is easily evaporated from the product. In many instances, the crude product is sufficiently pure to be used directly.

$$R \xrightarrow{O}_{Cl} + \underbrace{O}_{Cl} \xrightarrow{O}_{Cl} + \underbrace{O}_{Cl} \xrightarrow{CH_2Cl_2}_{4-5 \text{ hr}} R \xrightarrow{O}_{C-Cl} + CO_2 + CO + HCI$$

#### ALDEHYDES

The aldehyde functional group is a very reactive functional group; thus methods to prepare it must be mild and allow the aldehyde group to survive the reaction conditions. Traditional methods for introduction of the aldehyde functional group include



Aldehydes are intermediate in oxidation level, and thus the aldehyde functional group can be installed by either reduction of carboxylic acid derivatives or oxidation of alcohols. Aldehydes are rarely installed without a change of oxidation level. One difficulty is that they undergo both oxidation *and* reduction readily. Special methods are required to stop at the aldehyde stage rather than proceeding by further reduction or oxidation.

Reductive methods utilize carboxylic acid derivatives as starting materials, and the trick is to stop the reduction at the aldehyde stage, which is normally more easily reduced than the starting material. While there are a variety of reducing systems known and many employ acid chlorides as precursors, the most effective reduction method for the preparation of aldehydes is the diisobutylaluminum hydride (DIBAH) reduction of either esters or nitriles using a single equivalent of the reducing agent. By using low temperatures, the intermediate anions produced by hydride addition are at the aldehyde oxidation level, but they are resistant to further reduction. Hydrolysis delivers the aldehyde. Care must be taken to maintain low temperature during both the reaction and the hydrolysis.



The oxidation of primary alcohols to aldehydes also suffers from the problem of overoxidation of the aldehyde to a carboxylic acid. Mild methods capable of stopping the oxidation at the aldehyde oxidation level are required if aldehydes are to be obtained. The most common and effective reagent for this purpose is pyridinium chlorochromate (PCC), produced by the reaction of pyridinium hydrochloride with chromium trioxide. This reagent is soluble in dichloromethane and smoothly oxidizes primary alcohols to aldehydes in high yields. Because of the mild, neutral reaction conditions and the use of stoichiometric amounts of oxidant, the aldehyde product is not oxidized further.

The activation of DMSO by electrophilic reagents such as oxallyl chloride or trifluoroacetic anhydride (TFAA) (among many others) produces an oxidant capable of oxidizing primary alcohols to aldehydes in high yields. This oxidation is called the Swern oxidation and yields the aldehyde (oxidized product) by reductive elimination of dimethylsulfide (reduced product) and proceeds under mild, slightly basic conditions. It is a second widely used and effective oxidative method for the production of aldehydes from primary alcohols.

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**194** FUNCTIONAL GROUP SYNTHESIS

$$R-CH_{2}OH + \bigcup_{H_{3}C} S \bigoplus_{CH_{3}} \underbrace{\xrightarrow{TFAA}}_{Et_{3}N} \xrightarrow{R} R \xrightarrow{CH_{3}} O - S \bigoplus_{CH_{3}} \underbrace{\xrightarrow{Et_{3}N}}_{H} \xrightarrow{R} C = O + :S : CH_{3}$$

A different oxidative approach toward the preparation of aldehydes uses the ozonolysis of vinyl groups. If a vinyl group is present in a molecule, it can be oxidatively cleaved to an aldehyde by ozonolysis. This process cleaves the carbon–carbon double bond, but it is mild and very successful in many cases.



The formation of aldehydes without a change in oxidation level is not a common synthetic approach because most compounds that can be hydrolyzed to aldehydes without change in the oxidation level are formed from aldehydes in the first place. Thus acetals can be hydrolyzed rapidly to aldehydes by acidic water, but they are normally prepared from aldehydes. As such this is a very common protection strategy for aldehydes wherein they are first converted to an acetal and later hydrolyzed back to the aldehyde when the time is right.



#### KETONES

Ketones have the same oxidation level as aldehydes, but their preparation poses far fewer problems. Most importantly they are very resistant to oxidation so they can be prepared by any number of oxidative routes without difficulty. Textbook preparations of ketones are listed below and many of these traditional methods remain the methods of choice for the preparation of ketones:

Ozonolysis



Friedel–Crafts acylation



Oxidation

$$\begin{array}{c} OH \\ \downarrow \\ CH \\ R_1 \\ R_2 \end{array} \xrightarrow{H_2 CrO_4} \begin{array}{c} O \\ II \\ \sigma r [0] \\ R_1 \\ R_2 \end{array} \xrightarrow{R_1} \begin{array}{c} R_2 \\ R_2 \\ R_1 \\ R_2 \end{array}$$

Alkyne hydration

$$R - C \equiv CH \qquad \xrightarrow{H_3 O^{\oplus}} \qquad \begin{array}{c} O \\ Hg^{+2} \end{array} \qquad \begin{array}{c} O \\ Hg^{+2} \end{array} \qquad \begin{array}{c} O \\ C \\ C \\ CH_3 \end{array}$$

Addition to acid chlorides

$$\mathbf{R} \xrightarrow{\mathbf{O}} \mathbf{Cl} \xrightarrow{(\mathbf{R}')_2 \mathrm{CuLi}} \mathbf{R} \xrightarrow{\mathbf{O}} \mathbf{R} \xrightarrow{\mathbf{O}} \mathbf{R}' + \mathbf{R}' \mathrm{Cu}$$

Addition to acids

$$R \xrightarrow{O} OH \qquad \frac{1.2 \text{ R}'\text{Li}}{2.\text{ H}_3 O^+} \xrightarrow{O} OH \qquad R \xrightarrow{C} R'$$

One of the most common methods for the preparation of ketones is by the oxidation of secondary alcohols. The use of chromic acid (Jones reagent) is easy, safe, and effective for the oxidation of secondary alcohols to ketones. Furthermore Jones reagent gives a nearly neutral solution and thus can be used with a variety of acid-sensitive functional groups.



Sodium hypochlorite (household bleach) and acetic acid offers a very cheap and effective alternative to Jones reagent for the oxidation of secondary alcohols to ketones and has been widely used for the synthesis of ketones.

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196 FUNCTIONAL GROUP SYNTHESIS



If very mild and/or basic conditions are required, PCC is the reagent of choice and works very well.



There are *many* other reported methods for the oxidation of secondary alcohols to ketones; in fact, over 140 different methods are listed in *Comprehensive Organic Transformations* by Larock. However, few are as versatile and useful as those listed above.

Conversion of carboxylic acid derivatives to ketones requires a net reduction of oxidation level. Furthermore, since the two groups attached to the carbonyl group are carbon-containing groups, it follows that a carbon nucleophile must be the reductant, usually an organometallic reagent. Carboxylic acid derivatives such as esters, acid chlorides, and acid anhydrides do not stop at the ketone oxidation level upon reaction with most organometallic reagents but are further reduced to tertiary alcohols. (This same problem of overreduction was seen for aldehyde preparation.) However, carboxylic acids themselves react smoothly with organolithium reagents to furnish ketones upon hydrolytic workup. This method is an effective way to produce ketones. A key to the success of this reaction is the fact that the tetrahedral intermediate is a dianion which is stable to further addition. Only organolithium reagents are useful in this process for only they are powerful enough nucleophiles to add to the very weakly electrophilic carbonyl group of the carboxylate anion.



As with aldehydes, production of ketones by nonredox processes is not a common synthetic approach. Ketone derivatives having the same oxidation level are usually produced from ketones themselves. Several examples of enol and acetal ketone derivatives are shown below. All are prepared from ketones, all can be readily hydrolyzed back to the ketone in the presence of acidic water, and, with the exception of vinyl acetates, all are very stable to strong bases and nucleophiles. Acetals are often used as ketone (and aldehyde) protecting groups while enol derivatives are versatile synthetic intermediates.

#### IMINES AND IMINE DERIVATIVES 197



One hydrolytic method that is useful for the preparation of ketones is the hydrolysis of dithianes. 1.3-Dithiane can be alkylated by treatment with butyl lithium followed by an alkylating agent. The two sulfurs flanking the acetal carbon acidify the protons on that carbon such that butyl lithium can remove one giving a sulfurstabilized anion. This anion reacts with alkyl halides or sulfonates to give alkylated products. This sequence can be repeated to give a bis-alkylated product. Hydrolysis then yields a ketone. Dithioacetals are much more resistant to hydrolysis than acetals and thus  $Hg^{2+}$  is often used to promote efficient hydrolysis.

$$S \xrightarrow{1. \text{ BuLi}}_{2. \text{ R'-X}} \xrightarrow{S}_{R} \xrightarrow{1. \text{ BuLi}}_{2. \text{ R'-X}} \xrightarrow{S}_{R} \xrightarrow{H_20}_{Hg^{2+}} \xrightarrow{H_20}_{Hg^{2+}} \xrightarrow{R}_{R} \xrightarrow{R}_{R}$$

The acidifying effect of the sulfur atoms is an interesting phenomenon. It is not due to electronegativity since the corresponding 1,3-dioxane, which has even more electronegative oxygen atoms flanking the acetal carbon, cannot be converted to an anion by butyl lithium. The sulfur atoms have unfilled valence level d orbitals available that can accept electron density and thus stabilize an adjacent anion. One can consider the anion as being resonance stabilized with the negative charge being delocalized into the flanking sulfur atoms.



## **IMINES AND IMINE DERIVATIVES**

Nitrogen analogs of both aldehydes and ketones which have the same oxidation level are imines and imine derivatives. In almost every instance these compounds

#### **198** FUNCTIONAL GROUP SYNTHESIS

are prepared by an exchange reaction between an amine derivative and a carbonyl compound. For simple imines (R = alkyl, aryl) water removal by either a dehydrating agent (KOH or molecular sieves) or azeotropic distillation is often employed to drive the reaction to completion.

$$\mathbf{R} - \mathbf{\ddot{N}} \mathbf{H}_{2} + \mathbf{\ddot{N}}_{2} \mathbf{\ddot{C}} \mathbf{R}_{1} \longrightarrow \mathbf{R} - \mathbf{\ddot{N}} = \mathbf{\dot{C}}_{\mathbf{R}_{2}}^{'} \mathbf{R}_{1} + \mathbf{H}_{2}\mathbf{O}$$

A wide variety of other R groups attached to nitrogen can be used and all give the C=N bond formation with loss of water. However, most of these substitutions on the nitrogen atom reduce the nucleophilicity of the amino group so acid catalysis is used to facilitate the reaction. Many of these nitrogen derivatives of aldehydes and ketones have very interesting chemistry in their own right. These derivatives also serve as a classic way to identify aldehydes and ketones. Many of these derivatives tend to be solids with characteristic melting points. Thus conversion of an aldehyde or ketone to a derivative, such as oxime or a 2,4dinitrophenylhydrazone (2,4-DNPH) derivative, produces a solid with a distinct melting point. Comparison of the melting point with the known value can be used to confirm the structure of the aldehyde or ketone.



#### ALCOHOLS

The alcohol functional group is a very important functional group in organic chemistry. Not only do many important compounds and pharmaceuticals contain the alcohol group, but the alcohol group can be prepared *from* many other

groups and converted *to* many functional groups. Thus alcohols occupy a central position in functional group manipulations. In addition, most of the traditional methods for the preparation of alcohols are still among the most useful methods.

$$R_{1} \xrightarrow{O} OR_{2} \xrightarrow{1. \text{ LAH}} R_{1}CH_{2}OH + R_{2}OH$$

$$R_{2} = H, \text{ alkyl, aryl}$$

$$R_{1} \xrightarrow{O} OR_{2} \xrightarrow{1. \text{ BH}_{3}, \text{ THF}} R_{1}CH_{2}OH$$

$$R_{2} = H, \text{ but not alkyl, aryl}$$

Alcohols are at a fairly low oxidation level compared to other oxygencontaining functional groups and consequently are readily prepared by reduction. Large numbers of reductive methods have been reported for the preparation of alcohols. Carboxylic acids and esters react vigorously with lithium aluminum hydride (LAH) to produce primary alcohols. Carboxylic acids, but not esters, are also reduced easily by borane, which is the only reducing agent that reacts faster with carboxylic acids than with esters or other acid derivatives.



Its unique reactivity comes from the fact that borane first forms a Lewis acid-base complex with the acid and then a boron-carboxylate intermediate which increases the reactivity of the boron hydride and delivers the hydride by an intramolecular reaction. As such it provides a selective way to reduce acids and produce alcohols in the presence of most other functional groups.

Aldehydes and ketones are conveniently reduced by sodium borohydride, which is much milder than LAH and does not require aprotic conditions (an alcohol is often the preferred reaction solvent). Aldehydes give primary alcohols while ketones give secondary alcohols. 200 FUNCTIONAL GROUP SYNTHESIS



Alcohols and olefins are at the same oxidation level and are interconvertible without a change in oxidation level. Addition of water across an olefinic double bond is thus a common method for the preparation of alcohols. However, simple acid-catalyzed addition of water is often the least desirable alternative. Instead methods are used which permit much greater regiochemical control and are milder. For example, the preferred way to produce the Markovnikov alcohol from an olefin is to use hydroxymercuration for the addition step followed by reductive removal of mercury with NaBH₄. In this process mercury serves as a surrogate for the proton during the addition step and is replaced by hydrogen in the reduction. The advantage is that the mercuric electrophile forms a bridged or partially bridged intermediate that is stabilized and hence gives higher yields and cleaner product mixtures because rearrangements are suppressed. Other nucleophiles such as alcohols, acids, and hydrogen peroxide can also be employed as the cation trap.



Hydroboration is widely employed to obtain an anti-Markovnikov alcohol from an olefin. Addition of diborane to the double bond produces an organoborane intermediate. Three equivalents of the olefin are needed to consume the  $BH_3$  and a trialkylborane is produced. Reaction with basic  $H_2O_2$  converts the carbon-boron bond to a carbon oxygen bond. This process is effective and widely used.

3 
$$R_1 - CH = CH_2 \xrightarrow{BH_3 \cdot THF} (R_1 - CH - CH_2 - )_3 B \xrightarrow{NaOH} R_1 CH_2 CH_2 OH$$
  
via  $R_1 - CH \xrightarrow{CH_2} BH_2 \longrightarrow R_1 - CH_2 \xrightarrow{CH_2} BH_2$ 

The initial addition step occurs in a concerted fashion so that the hydrogen and boron are added to the same side of the planar double bond. Furthermore,

#### AMINES 201

the cleavage of the carbon-boron bond occurs with retention of configuration. These features can be used advantageously to prepare stereochemically pure alcohols.



# AMINES

Amines are saturated, nitrogen-containing functional groups that are widely encountered. Because the nitrogen atom of amines is basic, nucleophilic, and oxidizable, some constraints on the preparation of amines result. A collection of textbook amine preparations includes the following:

Amonolysis

$$NH_3 + R-X \longrightarrow R-NH_2 + polyalkylation$$

Azide substitution

$$N_3^{\ominus}$$
 + R-X  $\longrightarrow$  R-N₃  $\xrightarrow{Na/EtOH}$  R-NH₂ R-NH₂

Gabriel synthesis



Nitro group reduction

Ar-NO₂ 
$$\xrightarrow{H_2/Pd}$$
 Ar-NH₂

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#### 202 FUNCTIONAL GROUP SYNTHESIS

Reductive amination



Reduction

$$R \xrightarrow{O} R \xrightarrow{NR_2'} \text{ or } R - C \equiv N \xrightarrow{LAH} R - CH_2 - NR_2'$$
  
R' = H, alkyl, aryl

Amines are at the same low oxidation level as alcohols and consequently are easily prepared by reduction. Amides and nitriles are reduced efficiently by LAH to amines. Nitriles give only primary amines while amides give  $1^{\circ}$ ,  $2^{\circ}$ , or  $3^{\circ}$  amines depending on the number of carbon substituents on the amide nitrogen. The advantage of this method is that amides are easy to prepare from acid chlorides and amines while nitriles are available by displacement reactions.



One problem with this method is that the workup must be done carefully as the amine products tend to complex tenaciously with the aluminum salts formed from the LAH upon workup and thus are not recovered easily. There are standard workups which avoid these issues, but these should be followed carefully. Reduction of azides by catalytic reduction, phosphine or phosphite reagents, or Sn(II) chloride are all effective methods. The azides are also available from displacement reactions and give primary amines upon reduction.



Aromatic amines are readily prepared by the reduction of aromatic nitro groups by Fe/HCl or Sn/HCl or by catalytic hydrogenation.



Displacement reactions are rarely used for the preparation of amines as polyalkylation reduces yields and makes product mixtures more complex. However, reaction of primary amines with primary and secondary sulfonates can provide good yields of monoalkylated product if care is taken to control the conditions and mode of addition. Benzylamine is particularly common as a primary amine nucleophile since the benzyl group can be removed by hydrogenolysis to give a primary amine.

$$\begin{array}{c} R & \bigcirc \\ CH & OCH_2 & \xrightarrow{2 C_6 H_5 CH_2 NH_2} \\ OSO_2 C_6 H_4 NO_2 & \xrightarrow{CH_2 Cl_2} \end{array} \begin{array}{c} R & \bigcirc \\ CH & OCH_3 \\ NHCH_2 C_6 H_5 \end{array} \xrightarrow{H_2/Pd} \\ R & \bigcirc \\ NHCH_2 C_6 H_5 \end{array}$$

#### ALKENES

Traditional preparations of alkenes include the following:

Elimination



Dehydration



Dibromide reduction


204 FUNCTIONAL GROUP SYNTHESIS

Alkyne reduction



Alkenes are relatively low oxidation level hydrocarbons. The most common way to prepare alkenes is to carry out the elimination of a small molecule from between vicinal carbon atoms. However, this is only a viable strategy if the regiochemistry of elimination can be controlled. That is, traditional dehydrohalogenations or dehydrations often are regioselective but not regiospecific, so that mixtures of structurally isomeric olefins are formed. For example,



The formation of regioisomers is due to the presence of several sets of nonequivalent vicinal hydrogens of similar but not identical reactivity. The resulting mixture of similar products must be separated if only one of the regioisomers is desired. Since the alkene isomers are very similar in physical properties, such separations can be very difficult and certainly are not practical.

Several strategies to control the elimination regiochemistry have been developed. These include placement of the leaving group, steric bulk of the base, and/or establishment of thermodynamic control. By placing the leaving group at the end of a chain, only terminal olefins can be produced by elimination because there is only one set of vicinal hydrogens that can be removed by the base. Diazabicycloundecane (DBU) is the base used in the example below. It is very useful for promoting olefin-forming eliminations since it is a strong nitrogen base which is also relatively nonnucleophilic.



but

By using very bulky alkoxide bases (*t*-butoxide or amyloxide), attack of the base occurs at the least hindered position—usually at the end of chains if possible. In this way the regioselectivity of elimination is controlled by steric factors so that one isomer is produced nearly exclusively.



Finally, when eliminations which give conjugated systems are possible, they are favored significantly by the greater stability of the conjugated  $\pi$  system.



Dehydrations produce olefins from alcohols by the acid-catalyzed elimination of a water molecule from between two carbons. Acid-catalyzed dehydrations often give mixtures of products because the intermediate carbocation is prone to cationic rearrangements to more stable carbocations prior to formation of the olefin product. Moreover, even when the intermediate carbocation is not subject to skeletal rearrangement, as in the case of tertiary alcohols, mixtures of regioisomers are often produced during the loss of a proton from the carbocation. As a consequence, the acid-catalyzed dehydration of alcohols is generally not a viable synthetic method.



There are many other methods for carrying out 1,2 eliminations to give olefins. Several are particularly useful and widely used. Selenoxide eliminations are frequently used to install the double bond of  $\alpha$ ,  $\beta$ -unsaturated carbonyl compounds. They occur by concerted, cyclic, syn processes



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#### 206 FUNCTIONAL GROUP SYNTHESIS

Silyloxide eliminations (Petersen olefination) also proceed readily and regiospecifically to give olefins. When base is used to produce the oxyanion, the elimination occurs with syn stereochemistry. If an acid is used to promote the elimination, it occurs in an anti fashion, leading to the opposite olefin stereochemistry. This is a very useful way to generate either a Z or E olefin from the same starting material.



Similarly phosphine oxide eliminations (Wittig reaction) occur very readily to give olefins.

Both of the latter two methods of elimination are part of a longer sequence of reactions that produce olefins. Initial formation of a single bond to a carbonyl carbon is followed by elimination to an alkene. Thus the alkene is a condensation product of two smaller units. Schematically,

 $\begin{array}{c} O \\ R_1 \\ \hline \\ R_2 \end{array}^{} + \begin{array}{c} O \\ CH \\ R_3 \end{array} \\ \hline \\ R_3 \end{array} \\ \hline \\ R_1 \end{array} \\ \begin{array}{c} O \\ R_2 \end{array} \\ \hline \\ R_1 \end{array} \\ \begin{array}{c} O \\ R_1 \end{array} \\ \hline \\ \\ R_3 \end{array} \\ \begin{array}{c} Syn \\ R_3 \end{array} \\ \begin{array}{c} Syn \\ elimination \end{array} \\ \begin{array}{c} R_2 \\ R_1 \end{array} \\ \begin{array}{c} R_2 \\ R_1 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \end{array} \\ \begin{array}{c} R_2 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \\ R_3 \end{array} \\ \begin{array}{c} R_1 \\ R_3 \\ R_3 \\ \\ \end{array} \\ \begin{array}{c} R_1 \\ R_3 \\ R_3 \\ R_3 \\ R_3 \\ \\ \end{array}$ 

where X is an element (Si or P) which can remove oxygen to the alkene. It should also be noted that both anionic versions of these eliminations proceed with syn stereochemistry between the oxyanion and the heteroatom; thus the stereochemistry of the intermediate dictates the geometry of the olefin product.

Alkenes can also be produced effectively by the reduction of alkynes. The reduction can be carried out stereospecifically to give either cis or trans olefins as desired. This is a very useful method because of the stereocontrol. The P-2 nickel catalyst for the cis hydrogenation is produced in situ by the reduction

ALKANES 207

of Ni[II] acetate with sodium borohydride and the reaction is carried out at atmospheric pressure making this a very simple method for the preparation of cis olefins. The lithium in liquid ammonia reduction of alkynes to the trans olefin is also very straightforward experimentally.



#### ALKANES

Alkanes are the most highly reduced of all organic compounds. As a consequence, virtually all preparations of alkanes are reductive. Alkenes and alkynes can both be reduced to alkanes by catalytic hydrogenation. While many catalysts can be employed, palladium on carbon is by far the most common.

$$\begin{array}{c} R_1 & - \overline{\phantom{aaaa}} \\ \text{or} & R_2 \\ R_1 & - \overline{\phantom{aaaaaaaaaa}} \\ R_2 & R_1 - CH_2CH_2 - R_2 \end{array}$$

Primary and secondary alcohols can be converted to alkanes by conversion to tosylates followed by reduction with LAH. This reduction is valuable because deuterium can be easily introduced into the alkane by the use of lithium aluminum deuteride (LAD) instead of LAH.

$$R_1$$
-CH₂-OH  $\longrightarrow$   $R_1$ -CH₂-OTs  $\xrightarrow{LAH}$   $R_1$ -CH₃

Ketones can be reduced directly to alkanes by the Wolff-Kishner reduction. In this reduction, the ketone is converted to the hydrazone, which is treated in situ with sodium hydroxide. An internal redox reaction occurs in which the carbon is reduced and the hydrazine is oxidized to nitrogen. The best experimental conditions include the use of NaOH and ethylene glycol as solvent to carry out the reduction.

$$\begin{array}{c} R_1 \\ R_2 \end{array} = 0 \xrightarrow{H_2 N N H_2} \begin{array}{c} R_1 \\ R_2 \end{array} = N N H_2 \xrightarrow{NaOH} \begin{array}{c} R_1 \\ HO \end{array} OH, \Delta \end{array} \xrightarrow{R_1} C H_2 + N_2$$

#### 208 FUNCTIONAL GROUP SYNTHESIS

The reduction of ketones to alkanes can also be done by the Clemmensen reduction using zinc and HCl. This reaction is specific for aromatic ketones, however.

$$Ar \xrightarrow{O}_{HC1} \xrightarrow{Z_n} Ar-CH_2-R_1$$

Alkyl halides (Cl, Br, I) can be converted to alkanes by two types of reactions. The halogen can be reduced off most effectively using lithium or zinc metal. This procedure works best with bromides and iodides.

$$RX \xrightarrow{Zn/HOAc} RH$$

Alternatively alkyl halides undergo coupling reactions with lithium organocuprates (which are prepared from alkyl halides) to give alkanes by carbon–carbon bond formation. Other metals can be used to promote the same kind of coupling, but the use of cuprates is the most efficient and general.

$$R_1X + (R_2)_2CuLi \longrightarrow R_1-R_2 + R_2-Cu$$

It is clear that there are many different ways to carry out the installation of a particular functional group in a molecule. The ones discussed here are often the most general and practical, and they are often the first ones tried in the laboratory. However, it is also common that a particular substrate will not give good results with any of the common reagents. For this reason new methods of functional group manipulation are constantly being sought that are even more general, milder, more selective, cheaper, easier, and use more readily available starting materials than other methods.

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- M. B. Smith and J. A. March, Advanced Organic Chemistry, Reactions, Mechanism, and Structure, 5th ed., Wiley, New York, 2001, can also be used in the same manner.
- The best comprehensive listing of functional group preparations is in R. C. Larock, *Comprehensive Organic Transformations*, 2nd ed., Wiley VCH Publishers, 1999.

#### PROBLEMS

**7.1.** Show how you would prepare each of the following products from the given starting material. For each *overall* transformation indicate (1) the starting and ending functional group and (2) what change in oxidation level (if any) must be accomplished. Where more than one step is required, show each step distinctly, including reagents and conditions needed to effect the conversion



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210

FUNCTIONAL GROUP SYNTHESIS





**7.2.** Show *two* different starting materials from which you could obtain each of the following compounds. Show the reagents and conditions necessary to convert each of the starting materials into the desired product.

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FUNCTIONAL GROUP SYNTHESIS



**7.3.** Give the final products of the following reaction sequences and show the intermediate products as well. If mixtures are expected, indicate which will be the major product.



²¹² 

#### PROBLEMS 213



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#### PROBLEMS 215



# 8

# CARBON-CARBON BOND FORMATION BETWEEN CARBON NUCLEOPHILES AND CARBON ELECTROPHILES

Synthetic Strategy	217
Nucleophilic Carbon	218
Electrophilic Carbon	220
Reactivity Matching	223
Generation of Nucleophilic Carbon Reagents	224
Generation of Electrophilic Carbon Reagents	227
Matching Nucleophiles with Electrophiles	227
Enolates	228
Enolate Regioisomers	234
Diastereoselection in Aldol Reactions	236
Organometallic Compounds	239
Neutral Carbon Nucleophiles	239
C=C Formation	242
Cyclopropanation Reactions	244
Metal-Catalyzed Carbon–Carbon Bond Formation	246
Pd(0)-Catalyzed Carbon-Carbon Bond Formation	247
Heck Reaction	251
Suzuki Coupling	253
Stille Coupling	254

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	SYNTHETIC STRATEGY	217
Olefin Metathesis		256
Bibliography		261
Problems		262

#### SYNTHETIC STRATEGY

The object of the game in organic synthesis is to assemble a particular molecule, the target, which has the correct carbon skeleton and the proper array of functional groups distributed on that skeleton. There are two general strategies for accomplishing this synthetic challenge. The first is to start with a molecule having the desired carbon skeleton and then manipulate the functionality on that skeleton to that of the desired compound. The previous chapter as well as many texts deals with functional group preparations and thus provides methods for the installation and/or manipulation of functional groups on the skeleton.

The second general synthetic strategy is to assemble the proper carbon skeleton and then adjust the functionality in the resulting molecule to that of the target. Obviously assembling the carbon skeleton from smaller building blocks is a more versatile and convergent approach because the carbon skeleton of the target may not be available (or at least not readily available). Moreover, once the synthetic sequence needed to assemble the carbon skeleton has been developed, it can potentially be adapted to produce a series of structurally related molecules. This is particularly useful when such a series of compounds is needed to test for particular physical, chemical, or biological properties.

To assemble needed carbon skeletons from smaller units, it is absolutely crucial to be able to form carbon–carbon bonds between them. Consequently carbon–carbon bond-forming reactions are among the most important organic reactions. Since any particular carbon–carbon bond is merely a covalent link between two skeletal fragments, synthesis of a target such as  $R_3C-CR'_3$  can be accomplished by forming a carbon–carbon bond between the two skeletal fragments  $R_3C$  and  $CR'_3$ . Since the bond to be made contains two electrons that are shared, there are three modes by which these electrons can be distributed between the two fragments to be joined.

$$\left.\begin{array}{c} R_{3}C \oplus \text{ and } \oplus CR'_{3} \\ R_{3}C \oplus \text{ and } \oplus CR'_{3} \\ R_{3}C \cdot \text{ and } \cdot CR'_{3} \end{array}\right\} \longrightarrow R_{3}C - CR'_{3}$$

There are two ionic modes of bond formation where one carbon fragment is nucleophilic ( $e^{-}$  rich) and the other is electrophilic ( $e^{-}$  poor). There is one free-radical mode in which each fragment has a single unpaired electron which becomes

shared upon bond formation. To understand carbon–carbon bond-forming reactions that take place by ionic or two-electron processes, we need to know what species or compounds can serve as carbon-centered nucleophiles and what species or compounds can serve as carbon-centered electrophiles. (Free-radical reactions will be considered later.)

#### NUCLEOPHILIC CARBON

Carbon-centered nucleophiles are those compounds or intermediates which contain an electron-rich carbon atom and thus are capable of donating an electron pair from that carbon atom to an electrophile. The electron pair that is donated is found in a filled orbital in the nucleophilic carbon and the electrons are not tightly bound. Donation to the electrophilic carbon occurs by overlap of the filled orbital of the donor with an unfilled orbital of the acceptor. The most common carbon nucleophiles fall into three main classes:

(a) Organometallic molecules contain a carbon-metal bond which is polarized toward carbon. As a result the carbon is very electron rich and reacts vigorously with electrophiles.

$$\begin{array}{c} | \checkmark \\ \hline C & M \\ \delta - | & \delta + \end{array}$$
 M = MgX, Li, Cu, Zn, etc.

Examples of organometallic compounds most commonly used as carbon nucleophiles include Grignard reagents (RMgX), organolithiums (RLi), organocuprates (R₂CuLi), and occasionally organocadmiums (R₂Cd) and organozincs (RZnBr).

(b) Enolates are anionic derivatives of carbonyl compounds formed by proton removal from a position adjacent to the carbonyl group. Resonance delocalization of the negative charge with the carbonyl group stabilizes enolate anions and makes them somewhat less reactive than organometallic compounds.



They are, however, reactive carbon nucleophiles. Examples of enolates include anionic derivatives of aldehydes, ketones, acid derivatives, and dicarbonyl compounds.

#### NUCLEOPHILIC CARBON 219



Structurally related to enolates are anionic derivatives of imines and nitro compounds. The former are less stable (more reactive) than enolates because nitrogen cannot support a negative charge as well as oxygen and thus resonance stabilization is diminished compared to enolate anions. Nitronate anions are much more stable (less reactive) than enolates because resonance with the nitro group transfers the negative charge to oxygen where it is stabilized by the formal positive charge on nitrogen.



(c) A third type of molecule capable of functioning as a nucleophilic carbon equivalent is one that contains an electron-rich  $\pi$  bond. Such species are usually uncharged and function as carbon nucleophiles because the  $\pi$  electrons are less tightly bound and can consequently be donated to good electron acceptors. Enol derivatives are neutral derivatives of carbonyl compounds which have an oxygen or nitrogen substituent attached to a carbon–carbon double bond. They are covalent analogs of enolate ions, and because they neutral, they are much less reactive electron donors than enolate anions. In fact, most enol derivatives are stable compounds which can be isolated. In spite of being neutral, however, resonance interaction of the lone pairs of the heteroatom with the  $\pi$  system increases the electron density at the  $\beta$  position of the double bond and the double bond is electron rich. Besides the parent enol, examples of enol derivatives of carbonyl compounds include those shown below.



Besides a heteroatom substituent, which renders a double bond electron rich by resonance interaction of the lone pairs with the  $\pi$  system, other substituents can

also result in  $\pi$  bonds being electron rich and thus reacting as electron donors. Attachment of substituents less electronegative than carbon to the double bond increases its  $\pi$ -electron density significantly by an inductive effect. Vinylsilanes and vinyl stannanes can both be considered to have electron-rich  $\pi$  bonds and have been used as  $\pi$ -electron donors. Allylsilanes, by virtue of hyperconjugation between the allylic carbon–silicon bond and the  $\pi$  system, also function as good  $\pi$ -electron donors in many reactions.



#### **ELECTROPHILIC CARBON**

Carbon-centered electrophiles are compounds or intermediates which are electron poor and thus capable of accepting electrons from electron donors. To be an electron acceptor, an electrophile must have an unfilled orbital on carbon available for overlap with a filled orbital of the donor. Unfilled atomic p orbitals or antibonding orbitals (both  $\sigma^*$  and  $\pi^*$ ) are the most common types of acceptor orbitals. The most common carbon electrophiles fall into four major categories:

(a) Cationic carbon electrophiles are the most reactive because of the positive charge they carry. They can, however, have a variety of structures depending on the hybridization of the carbon acceptor. Trialkyloxonium tetrafluoroborates (Meerwein salts)  $R_3O^+$  BF₄⁻, for example, are sp³-hybridized carbon electrophiles and are extremely reactive toward nucleophiles. The acceptor orbital is an antibonding C–O  $\sigma^*$  orbital which is low in energy because of the positive charge on oxygen.

Triphenylmethyl (trityl) tetrafluoroborate, on the other hand, is  $sp^2$  hybridized but it is also extremely reactive toward electron donors. The acceptor orbital of the trityl cation is an unfilled 2p atomic orbital on the charged carbon. These carbon electrophiles are isolable compounds, but they are extremely reactive with any sort of electron donor (H₂O vapor is a common culprit).



Many other cationic carbon electrophiles cannot be isolated but can be generated insitu in a reaction mixture by Bronsted or Lewis acid-base reactions. In the presence of Bronsted acids, carbonyl compounds are protonated and produce positively charged oxonium ions. Compared to the carbonyl compound itself, the  $\pi^*$  orbitals of oxonium ions are much stronger electron acceptors.



Protonation of the carbonyl group is an equilibrium process, and the extent of protonation (the position of the protonation equilibrium) is dictated by the  $pK_a$ 's of the Bronsted acid and the oxonium ion. While this equilibrium usually lies far to the left, the reactivity of the oxonium ion is often sufficiently great that only small amounts of the oxonium ion are needed to react effectively with the electron donor.

Addition of a strong Lewis acid such as TiCl₄, BF₃, or SnCl₄ to a carbonyl compound is another common method to produce a very powerful cationic electrophile in solution. Complexation between the Lewis acid and the lone pairs of electrons on the carbonyl oxygen give a species which, although formally neutral, behaves as a cationic carbon electrophile in the same fashion as a protonated carbonyl group. These are strong electrophiles that react with many types of nucleophiles. While aldehydes and ketones are common carbonyl components which are activated with Lewis acids, esters and amides also yield strongly electrophilic species with Lewis acids.

(b) Aliphatic compounds with good leaving groups attached to primary or secondary carbon atoms are very commonly used as carbon electrophiles. The leaving group is an electronegative group attached by a polarized  $\sigma$  bond.



The bond polarity makes the carbon atom electron deficient and capable of accepting electrons from carbon electron donors (carbon-centered nucleophiles) into the  $\sigma^*$  antibonding orbital. Population of the  $\sigma^*$  orbital by electron donation weakens the bond to the leaving group. Ultimately the leaving group is cleaved from the

molecule and retains the pair of electrons from the connecting bond. Examples of such compounds include alkyl halides, alkyl sulfonates, and alkyl sulfates.

The electrophilicity of such compounds is largely related to the leaving ability of the leaving group. The leaving ability of a group is in turn related to (1) its bond strength to carbon and (2) its ability to accept the bonded pair of electrons and become electron rich (most often negatively charged). Leaving abilities range from excellent (triflate) to moderate (chloride). The electrophilicity of the acceptor molecule thus can be adjusted by changing the leaving group.



(c) Carbonyl compounds are very common carbon electrophiles by virtue of the polarized carbon–oxygen  $\pi$  bond. Electron donation into the  $\pi^*$  orbital of the carbonyl carbon breaks the C–O  $\pi$  bond and produces a tetrahedral adduct which can then proceed to products.



This is a very general process for carbonyl compounds; however, the electrophilic reactivity of the carbonyl group is very dependent on the groups attached to it. The reactivity is ranked in the following order:



Electron-withdrawing groups (Cl,  $RCO_2$ ) increase the electrophilicity while resonance-donating groups -OR,  $-NR_2$  decrease the reactivity toward electron donors. Steric effects are also a significant influence on carbonyl reactivity. The trigonal carbonyl reactant goes to a more crowded tetrahedral intermediate upon addition of the nucleophile; thus bulky groups attached to the carbonyl carbon lead to more crowded transition states and result in much slower addition reactions. This steric rationale is one explanation for the greater reactivity of aldehydes over ketones. Extremely sterically hindered ketones such as di-*tert*butyl ketone undergo carbonyl addition by nucleophiles at negligible rates for most nucleophiles.

(d)  $\alpha,\beta$ -Unsaturated carbonyl compounds can act as electrophiles under certain conditions and are bidentate in that both the carbonyl carbon and the  $\beta$ carbon are electron deficient. Thus nucleophiles can attack at either position.



The regioselectivity of nucleophilic addition is a function of the type of nucleophile employed and in many instances can be controlled to give Michael addition to the  $\beta$  carbon.



#### **REACTIVITY MATCHING**

Having defined the types of commonly used carbon nucleophiles and carbon electrophiles, it would seem that if you react any of the carbon nucleophiles (electron donors) with any of the carbon electrophiles (electron acceptors), then a carbon–carbon bond should be formed. While this is theoretically true, it is unworkable from a practical point of view. If, for example, a carbanion nucleophile was reacted with a cationic electrophile, it is unlikely that the desired carbon–carbon bond formation would be detected, even after the smoke cleared. Or if a silyl enol ether nucleophile was reacted with an  $\alpha$ ,  $\beta$ -unsaturated ester, no reaction could be observed to take place in any reasonable time frame.



For carbon-carbon formation to be successful, the reactivities of the nucleophile and electrophile must be matched so that reaction occurs at a reasonable

and controllable rate. Thus we must be able to easily generate both carbon nucleophiles and carbon electrophiles to be able to choose the appropriate partners for successful C–C bond formation.

#### **GENERATION OF NUCLEOPHILIC CARBON REAGENTS**

The three major classes of nucleophilic carbon species are organometallic compounds, enolate derivatives and related carbanionic compounds, and neutral enol derivatives:

1. Organometallic compounds which contain a carbon-metal bond are the most reactive carbon nucleophiles. In most cases they are also powerful bases and must be prepared and used under strictly anhydrous and aprotic conditions. A very common way to produce organometallic compounds is to reduce alkyl halides with active metals. Grignard reagents and organolithium compounds are routinely produced in this manner. The transformation is a two-electron reduction of the alkyl halide to a carbanion equivalent; the metal is oxidized.



This procedure works well for alkyl, vinyl, and aryl halides and provides a convenient source of organomagnesium halides and organolithium compounds. In addition, a variety of other metals can be exchanged for lithium in organolithium compounds to give different organometallic compounds of modified reactivity. Reaction of two equivalents of an organolithium compound with a cuprous halide gives a lithium organocuprate in which the carbon–lithium bonds of the organolithium reactant are converted to carbon–copper bonds in the anionic organocuprate. Lithium merely serves to balance the charge of the organocuprate. By a similar exchange, dialkylmercury compounds can be prepared from organolithiums and Hg[II] halides.

> $2 \text{ RLi} + \text{CuX} \longrightarrow \text{R}_2\text{CuLi} + \text{LiX}$ lithium organocuprate  $2 \text{ RLi} + \text{HgX}_2 \longrightarrow \text{R}_2\text{Hg} + 2\text{LiX}$ organomercury compound

A second way to make organometallic compounds for use as carbanion nucleophiles is to use halogen-metal exchange. In this process an alkyl halide and an organometallic compound undergo a metathesis reaction to give a new organometallic compound and a new alkyl halide. This process is thought to take place by nucleophilic attack on the halogen atom by the organometallic reagent.

$$R-Li + R'-Br \longrightarrow R'-Li + R-Br$$

One requirement is that the  $pK_a$  of the new organometallic compound is lower than the  $pK_a$  of the starting organometallic. This in essence means that the equilibrium is driven to products by the formation of a more stable anion. This method is commonly used to make vinyl lithiums from vinyl halides and alkyl lithiums and aryl lithiums from aryl halides and alkyl lithiums because the electron pair in an sp² orbital of a vinyl or aryl lithium compound is more stable than the electron pair in an sp³ orbital of an alkyl lithium.

 $= \underbrace{}_{Br} + BuLi \longrightarrow = \underbrace{}_{Li} + BuBr$ 

A method often employed to drive the halogen-metal exchange equilibrium to completion is to employ *tert*-butyl lithium as the organolithium component. In addition to being the most basic organolithium compound because of the tertiary substitution, conversion of the *tert*-butyl bromide by-product to isobutylene also occurs under the reaction conditions and drives the exchange equilibrium to completion. Note that two equivalents of *tert*-butyl lithium are required as one equivalent is used in the halogen-metal exchange and one equivalent is consumed in converting *tert*-butyl bromide to isobutylene.

$$Br + 2 - Li \longrightarrow Li + - LiBr$$

2. Enolates and related carbanionic nucleophiles are routinely generated by removal of an acidic proton in a molecule with a base. Carbonyl groups acidify their  $\alpha$  protons somewhat and make their removal by a base a common process. However, structural features other than carbonyl groups can also acidify protons bound to carbon and thus facilitate their removal by bases. For example, p $K_a$  values for structurally acidified C–H protons include the ones given below.



The  $pK_a$ 's of commonly used bases are as follows:

$$CO_3^{\ominus}$$
  $RO^{\ominus}$   $N \ominus$  NaH, KH, R-Li  
 $pK_a = 12$   $pK_a = 15-19$   $pK_a = 35$   $pK_a > 35$ 

By knowing (or estimating) the  $pK_a$  of a proton to be removed, it is possible to choose a base with a higher  $pK_a$  in order to have essentially complete conversion to the anionic carbon nucleophile. When these conditions are met, proton exchange occurs readily and a carbon nucleophile is produced. It must be remembered, however, that many bases can serve as nucleophiles. If the structural feature which acidified the C–H proton is an electrophile, then a nucleophilic base cannot be used. For example, butyl lithium ( $pK_a > 45$ ) converts phenylacetylene ( $pK_a \sim 25$ ) smoothly to its conjugate base by proton removal, whereas it reacts as a nucleophile with the carbonyl group of acetophenone in spite of the fact that the  $\alpha$  protons of acetophenone have  $pK_a = 21$  and are thus more acidic than the terminal proton in phenylacetylene.



To circumvent problems of nucleophilicity, lithium diisopropylamide (LDA), potassium hexamethyldisilylamide (KHMDS), and KH are often employed for proton removal since they are very strong bases ( $pK_a > 35$ ) but relatively poor nucleophiles. Hence they remove protons from acidic C–H bonds but normally do not attack carbonyl groups or other electrophilic centers.



If the C–H proton is highly acidified as in a  $\beta$ -dicarbonyl compound (p $K_a \sim 10-14$ ) or nitro compounds (p $K_a = 9-12$ ), weaker bases such as alkoxides

 $(pK_a \sim 17)$  can be used to convert the material completely to its conjugate base, and thus aprotic conditions are no longer required. However, a common protocol to convert dicarbonyl compounds to their enolates in a clean, controllable manner is to use sodium hydride in dry THF.



3. A third major class of carbon nucleophiles is enol derivatives. In general, these are stable compounds that are prepared by one of the functional group transformations outlined in the previous chapter.

#### **GENERATION OF ELECTROPHILIC CARBON REAGENTS**

Electrophilic carbon species are most often stable compounds with an electrophilic functional group present. Since they are stable molecules, they need not be generated as transients in the reaction mixture. The functional types which are good electrophiles have been defined earlier in this chapter and, the preparations of these functional groups were outlined in the previous chapter.

#### MATCHING NUCLEOPHILES WITH ELECTROPHILES

Having defined nucleophilic and electrophilic carbon species and having learned to produce a variety of them, the next step is to match the reactivities of the nucleophiles and electrophiles so that carbon–carbon bond formation can occur in a controllable and selective fashion. Qualitatively the order of reactivities for nucleophiles and electrophiles used in carbon–carbon bond-forming reactions are shown below. In general, many of the most useful carbon–carbon bond-forming reactions take place with nucleophiles and electrophiles in the middle ranges of reactivity. Highly reactive electrophiles and nucleophiles are often difficult to control while nucleophiles and electrophiles of low reactivity often fail to react effectively. Nevertheless it is reactivity matching that is most important in producing useful reactions.

When stabilized (and consequently less reactive) anions are employed as the nucleophile, more reactive electrophiles are needed for successful carbon–carbon bond formation. Nitronate anions, which are highly resonance stabilized, fail to react with simple alkyl halide electrophiles. On the other hand,  $\beta$ -dicarbonyl compounds react effectively with primary and some secondary alkyl bromides and iodides to give monoalkylated products.



Under the same conditions simple enolates react vigorously with alkyl halides (which must be primary) to give mono- and polyalkylated products. The reactivity of the simple enolate is greater and cannot be controlled at room temperature. However, if the alkylation is carried out at low temperature, the reaction can be controlled and smooth monoalkylation of simple enolates can be achieved. The same is true for the alkylation of acetylide anions, which must be carried out at low temperature for successful alkylation.

$$R-C \equiv C \ominus + CH_3CH_2-I \xrightarrow{-78^\circ C} R-C \equiv C-CH_2CH_3$$

#### ENOLATES

Enolates are important nucleophiles which react nicely with a variety of carbonyl compounds. In this case, the nucleophilic reactivity of the enolate and the electrophilic reactivity of the carbonyl group are well matched and a wide variety of products can be made. The type of enolate (ketone, ester, etc.) and the type of carbonyl electrophile (aldehyde, ketone, ester, etc.) determine the structure of the final product. Furthermore these reactions are often named according to the two partners that are reacted and the type of product produced from them.

The aldol condensation is the reaction of an aldehyde or ketone enolate with an aldehyde or ketone to give a  $\beta$ -hydroxy aldehyde or ketone. A simple aldol reaction is one in which the enolate nucleophile is derived from the carbonyl electrophile. Very often the  $\beta$ -hydroxy carbonyl product dehydrates to give an

ENOLATES 229

 $\alpha$ , $\beta$ -unsaturated carbonyl compound; however, the aldol nature of the dehydration product can be discerned by disconnection of the double bond of the unsaturated product.



If the enolate nucleophile is derived from an aldehyde or ketone different than the carbonyl electrophile, a crossed-aldol condensation results. Normally best success is achieved if the carbonyl electrophile employed for the crossedaldol condensation is more reactive than the carbonyl electrophile from which the enolate is derived. For example, ketone enolates react with aldehydes effectively, but aldehyde enolates do not give the crossed aldol with most ketones but selfcondense instead.



The Claisen condensation is the reaction of the enolate of an ester with an ester electrophile. The product is a  $\beta$ -keto ester since the tetrahedral intermediate collapses by expulsion of an alkoxide.



A crossed Claisen is the reaction of an ester enolate with an aldehyde or ketone to produce a  $\beta$ -hydroxy ester. This works well because aldehydes and ketones are more reactive electrophiles than esters; thus the ester enolate reacts faster with the aldehyde or ketone than it condenses with itself, avoiding product mixtures. Moreover, the aldehyde or ketone should not have  $\alpha$  hydrogens so that proton transfer to the more basic ester enolate is avoided. This would lead to the formation of an aldehyde or ketone enolate in the mixture, and an aldol reaction would be a major competing reaction.



For the same reason it is generally not feasible to carry out a crossed-Claisen reaction between the enolate of one ester and a second ester which has  $\alpha$  protons. This is due to the fact that if nucleophilic addition to the carbonyl group is not fast, proton exchange can occur, giving a mixture of enolates and thus a mixture of products.



There are many other named reactions that follow the same general features but differ as to the type of enolate or the carbon electrophile. These include the Reformatski reaction, the Darzens reaction, and the Dieckmann ring closure. They were in widespread use for many years and were named as a convenient way to characterize the reactants employed and type of product which results. The reason that there are so many variations on the same theme is that control of the reaction products depends on the ability to generate a particular enolate nucleophile and react it with a particular carbonyl electrophile. In earlier times alkoxide bases were the strongest bases routinely available to synthetic chemists. Since alkoxides have  $pK_a = 15-19$  while protons  $\alpha$  to carbonyl groups have  $pK_a = 20-25$ , the reaction of an alkoxide base with a carbonyl compound produces only a small amount of the enolate at equilibrium, and it is produced in the presence of the unreacted carbonyl compound, which is an electrophile. For simple aldol and Claisen reactions, this is the ideal situation for self-condensation.



If, however, it is necessary to generate a crossed product by the reaction of an enolate derived from one carbonyl compound with a second carbonyl compound as the electrophile, things can go bad rapidly. Because both carbonyl groups must be present in solution at the same time and each can form enolates to some extent, there can be four possible products from the various combinations of enolates and carbonyl compounds. This problem was illustrated for the crossed-Claisen condensation above. The number of products can be minimized if one carbonyl component lacks  $\alpha$  protons and cannot form an enolate and is also a more reactive electrophile than the second carbonyl component. If these conditions are met, then crossed condensations can be carried out successfully using alkoxide bases. Many of the named reactions were developed so that product mixtures could be avoided.

Today reactions of enolates are usually carried out much differently by utilizing very strong, nonnucleophilic bases for generating the enolate nucleophile. Instead of having only small equilibrium concentrations of an enolate produced in solution, the use of strong, nonnucleophilic bases like LDA, KHMDS, and KH that have  $pK_a$ 's >35 permits carbonyl compounds, whose  $\alpha$  protons have  $pK_a$ 's of 20–25, to be converted completely to enolate anions. Doing so completely converts the carbonyl compound into a nucleophile which cannot condense with itself and is stable in solution. This enolate can then be reacted with a second carbonyl compound in a subsequent step to give product:

Step 1



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232 CARBON-CARBON BOND FORMATION



As long as nucleophilic addition of the preformed enolate to the second carbonyl component is rapid and the carbonyl electrophile is added *after* the enolate is formed, the product is predictable and is not a mixture. The rule of thumb to ensure success is that the carbonyl electrophile should be more reactive than the carbonyl compound from which the enolate is derived. If this condition is met, the carbonyl electrophile can have  $\alpha$  protons and the structural possibilities are increased tremendously. Typical enolate–carbonyl pairs that have been condensed by this methodology include the following:

Enolate	Carbonyl Compound	Product Type
Ester enolate	Aldehydes, ketones	$\beta$ -Hydroxy ester
Ester enolate	Acid chlorides or carbonylimidazole	$\beta$ -Ketoester
Ketone enolate	Aldehyde	$\beta$ -Hydroxyketone
Ketone enolate	Acid chloride	$\beta$ -Diketone

Acetylides can also react as nucleophiles toward aldehydes and ketones to give propargylic alcohols, which provides a simple way to install the triple bond in molecules.

 $R-C \equiv C \ominus + \underset{R_1}{\overset{O}{\amalg}} \underset{R_2}{\overset{O}{\longrightarrow}} R-C \equiv C \overset{OH}{\underset{R_2}{\longleftarrow}} R_1$ 

Esters, amides, and nitriles are relatively weak electrophiles. They react sluggishly or not at all with enolates. Esters are more electrophilic than amides and nitriles and react readily with carbanionic-type reagents such as organolithiums or Grignard reagents. As seen previously, two equivalents of the organometallics are added and tertiary alcohols are produced. Tertiary amides and nitriles react with organolithiums (but not Grignard reagents) to give ketones after hydrolytic workup. A single nucleophilic addition occurs to give an anionic intermediate which is stable to further nucleophilic addition. The oxidation level is that of a ketone which is unmasked upon hydrolysis.

#### ENOLATES 233



Carbonyl electrophiles are obviously a very important group of electrophiles that react successfully with a spectrum of carbon nucleophiles. Among carbonyl electrophiles, however, large differences in reactivity are observed. Acid chlorides are very reactive electrophiles whereas esters and amides are much weaker and fail to react with several classes of carbon nucleophiles. Aldehydes and ketones are probably the most widely utilized groups of carbonyl electrophiles and exhibit moderate electrophilic reactivity. No matter what carbonyl electrophile is used, however, it reacts by nucleophilic addition to the carbonyl carbon to produce a tetrahedral intermediate. The ultimate reaction product reflects subsequent chemistry of the tetrahedral intermediate.

While the use of strong bases has changed the way in which many condensation reactions are carried out, it is important to remember the types of products that are produced from them. Recall that the aldol condensation yields  $\beta$ -hydroxy aldehydes or ketones which are easily dehydrated to  $\alpha$ , $\beta$ -unsaturated aldehydes or ketones.

$$\begin{array}{c} O \\ R \xrightarrow{} CH_2 R' \xrightarrow{aldol} R \xrightarrow{} R \xrightarrow{} R' \xrightarrow$$

It is thus possible to look at a molecule such as **A** below and recognize that it is a  $\beta$ -hydroxy ketone and thus could be formed in a crossed-aldol reaction between enolate **B** and aldehyde **C**. Likewise **D** could potentially be produced by dehydration of the aldol product of cyclohexanone

In addition to these intermolecular processes, intramolecular versions of the Claisen (Dieckmann) reaction and the mixed Claisen and the aldol reaction (Robinson annulation) are also well known. In all cases the same structural classes of products are formed.



#### **ENOLATE REGIOISOMERS**

Enolates are commonly used as the nucleophilic component in carbon–carbon bond-forming reactions. By using strong, nonnucleophilic bases, both esters and ketones are easily converted to their enolates. Ketones, however, are problematic with regard to the regioselectivity of enolate formation if they are unsymmetric. As seen in the following example, two regioisomeric enolates can be produced by the removal of the nonequivalent  $\alpha$  protons of 2-pentanone by base, and thus two regioisomeric aldol products are possible.



Regiochemical control of enolate formation is thus an important consideration when planning ways to construct a carbon–carbon bond using a ketone enolate. There are several strategies for controlling the regiochemistry of proton removal

The first is to take advantage of the fact that the less-substituted  $\alpha$  position has slightly more acidic protons. If a ketone is added *slowly* to a cold solution of LDA, the more acidic proton will be removed preferentially. The resulting enolate is termed the *kinetic enolate* because the more acidic proton is removed *faster* than the less acidic proton. Both steric and electronic factors contribute to the more rapid removal of protons from the less highly substituted  $\alpha$  carbon.



The enolate that is the most stable usually has the most highly substituted double bond and is called the thermodynamic enolate. If a slight excess of the ketone is used or a trace of protic impurities is present, equilibrium between the enolates is established and isomerization to the more highly substituted enolate occurs.



The thermodynamic enolate is lower in energy so it is the one favored if equilibrium is achieved. For this reason, great care must be taken in the preparation and reaction of the kinetic enolate so that equilibration does not occur. On the other hand, preparation and reaction of the thermodynamic enolate is much easier and demands less rigorous reaction conditions.

Besides the direct formation of kinetic or thermodynamic enolates of ketones, other strategies can be employed to produce regiospecific products. An older and extremely valuable strategy for making the synthetic equivalent of a particular regiospecific enolate utilizes some group (G) to acidify a proton  $\alpha$  to a ketone so that it is removed preferentially by base. The resulting enolate is used as a carbon nucleophile and then the group (G) is removed. In this way it appears that one  $\alpha$  position of a ketone has been regioselectively transformed when, in fact, the group G has guided the chemistry in the reactant but is not present in the product.

$$\begin{array}{c} \begin{array}{c} O \\ R_1 \\ CH_2 \\ H \\ H \\ H \\ H \end{array} \xrightarrow{G} \begin{array}{c} :_B \\ CH_2 \\ H \\ H \end{array} \xrightarrow{C} CH_2 \\ H \\ H \\ CH_2 \\ H \\ H \\ CH_2 \\ H \\ H \\ CH_2 \\$$

The most common group G is an ester function (although many other groups have been employed as well). The starting  $\beta$ -ketoester, which can be prepared easily by a Claisen-type reaction of an ester enolate and an acid chloride, has a very acidic  $\alpha$  proton (p $K_a \sim 9-10$ ) which is easily removed (i.e., 1 equiv. NaH or 1 equiv. EtO⁻).



The resulting enolate is used as a nucleophile to form a new carbon-carbon bond. The ester is then hydrolyzed and  $CO_2$  is thermally ejected to provide an  $\alpha$ -substituted ketone. This strategy is simple, efficient, and convenient and is widely used. This synthesis is commonly referred to as the acetoacetic ester synthesis since the most simple starting material is an ester of acetoacetic acid if  $R_1 = H$ .

A third strategy for controlling enolate formation is to convert the carbonyl group to a N,N-dimethylhydrazone. The hydrazone is less reactive than the carbonyl group, and removal of an  $\alpha$  proton by a strong base takes place at the least hindered  $\alpha$  position. Alkylation followed by hydrolysis gives back carbonyl product that is the same as the result of kinetic control of enolate

formation. However, this method does not have the problems of equilibration as found for simple enolate formation. The regioselectivity of proton removal from the hydrazone is probably related to the geometry of the hydrazone. The dimethylamino group is pointed toward the least hindered  $\alpha$  position for steric reasons and directs the base to that position by coordination with the lone pairs on nitrogen.



The use of hydrazones is particularly important to form the enolate equivalents of aldehydes. Aldehydes are quite reactive as electrophiles, so as soon as some enolate has been formed, it reacts with the unreacted aldehyde present in solution. Conversion of the aldehyde to its N,N-dimethylhydrazone (=NNMe₂) lowers the electrophilicity so that  $\alpha$ -proton removal can take place and then the electrophile of choice can be added. Hydrolysis gives back the aldehyde. In this case the geometry of the hydrazone is unimportant since aldehydes have only one  $\alpha$  position from which protons can be removed by base.



#### DIASTEREOSELECTION IN ALDOL REACTIONS

The reaction of enolates with aldehydes or ketones to produce  $\beta$ -hydroxy carbonyl derivatives is a very common and a very useful way to make carbon–carbon bonds. A fundamental stereochemical feature of the reaction is that two new chiral centers are produced from achiral starting materials. Hence syn and anti diastereomers will be produced, each as a pair of enantiomers. This is shown schematically for the reaction of a propionate enolate with isobutyraldehyde. Because they have different energies, the syn and anti diastereomers will be

produced in unequal amounts, but each will be produced in racemic form because both starting materials are achiral.



The diastereoselectivity of the reaction results from a combination of three factors. First the carbonyl electrophile can undergo addition on either its Re or Si face. Second, the enolate nucleophile is planar and can attack the carbonyl group from either of its faces. Third, the enolate geometry can be either Z or E. To control the diastereoselectivity, it is first necessary to use a single isomer of the enolate. In general, the E enolate is the kinetic enolate and the Z enolate is thermodynamically favored. Methods are available to produce either as the major isomer by  $\alpha$ -proton removal from carbonyl compounds with strong bases. This is particularly true of esters and amides. Pure Z and E enolates can also be prepared by first converting the carbonyl compound to a Z and E mixture of silyl enol ethers, separating these isomers, and regenerating the Z and E enolates with methyl lithium. Suffice it to say that there are known ways to produce either Z or E enolates in pure form.



The stereoelectronic requirements for carbonyl addition are that electron donation occurs by interaction of the donor with the  $\pi^*$  orbital of the carbonyl group. To meet the stereoelectronic requirements and explain the diastereoselectivity, the Zimmerman–Traxler model is used. Interaction of the lithium cation with the oxygen of the enolate and of the carbonyl electrophile leads to a six-membered

chairlike transition state. If the geometry of the enolate is fixed, the only variable is the orientation of the electrophile. The preferred orientation has the larger substituent in a pseudoequatorial position. This preferred orientation produces the major diastereomer. An example is shown for the Z enolate of ethyl propionate reacting with isobutyraldehyde, which predicts that the anti diastereomer should be favored (and it is!). A similar analysis predicts that the E enolate should give the syn diastereomer as the major product (and it does!).



This model is extremely useful in understanding the stereochemical outcomes of aldol processes. It also provides a framework for influencing the diastereoselectivity in a rational way. For instance, if the ethoxy group in the above example is changed to a much bulkier group, increased transannular interactions in the pseudoaxial transition state would make it even higher in energy and result in increased selectivity for the anti isomer (and it does!)

Even greater diastereoselectivity in the aldol reaction can be achieved using boron enolates as the carbon nucleophile. Boron enolates are easily prepared from aldehydes and ketones, and the syn and the anti isomers can be separated as pure compounds. They react with aldehydes and ketones to give aldol products by a similar transition state. The difference is that boron oxygen bonds are shorter than lithium oxygen bonds, and thus steric interactions in the transition state are magnified and result in greater diastereoselectivity.



#### ORGANOMETALLIC COMPOUNDS

Organometallic compounds such as Grignard reagents and organolithium reagents are very powerful nucleophiles which react with a wide variety of carbonyl compounds. In general, organolithium reagents are more reactive than Grignard reagents. Both types of reagents react rapidly with aldehydes and ketones and yield secondary and tertiary alcohols after aqueous workup. The new carbon–carbon bond joins the organometallic fragment with the carbonyl carbon.

$$\begin{array}{ccc} R-M & + & \bigcap_{M=MgX, Li}^{O} & & R \xrightarrow{OH} \\ R & & R_1 & R_2 \end{array} \xrightarrow{R_2} & R \xrightarrow{Q} \\ \end{array}$$

The tetrahedral intermediate produced by addition of organolithiums or Grignard reagents to esters collapses to a ketone which is more reactive than the original ester electrophile. A second equivalent adds to give a tertiary alcohol as the product. This process cannot be controlled by temperature or by mode of addition since the intermediate product is more reactive than the starting material.

Collapse of the tetrahedral intermediate can be prevented, however, by reaction of a carboxylic acid with two equivalents of an organolithium (or a carboxylate salt with one equivalent of the organolithium). Addition to the carboxylate gives a dianionic tetrahedral intermediate which has the ketone oxidation state but is stable to further addition under the reaction conditions. The ketone is revealed only upon hydrolysis. This is a widely used method for the preparation of ketones by the formation of a carbon–carbon bond, but it is restricted to organolithium reagents as the carbon nucleophile. Grignard reagents do not react with carboxylates, illustrating they have a slightly reduced nucleophilicity relative to organolithiums.

$$R_1CO_2H \xrightarrow{R_2-Li} R_1CO_2^{\ominus} Li^{\oplus} \xrightarrow{R_2-Li} R_1 \xrightarrow{OLi} R_2 \xrightarrow{H_2O} R_1 \xrightarrow{OLi} R_2$$

#### **NEUTRAL CARBON NUCLEOPHILES**

Other types of carbon nucleophiles such as uncharged enol derivatives and other  $\pi$ -bonded systems are only weakly nucleophilic and consequently unreactive
toward normal carbon electrophiles like carbonyl compounds or alkyl halides or sulfonates. For them to be used effectively as nucleophiles, strong electrophiles must be used to match this reduced nucleophilicity. This can be accomplished by increasing the reactivity of a normal electrophile. Typically this is done by treating an electrophile with a Lewis acid. Coordination of lone pairs of electrons with the Lewis acid increases the electrophilicity markedly.

Examples are



A variety of Lewis acids can be used. Among the more commonly used ones are AlCl₃, TiCl₄, SnCl₄, ZnCl₂, BF₃, and TMSOTf (trimethylsilyl triflate). The choice of Lewis acid is often critical to the success of the reaction and is usually made by referring to similar transformations that have been successfully reported in the literature. Often it is not possible to rationalize why one Lewis acid works and another one does not, so the initial choice of catalyst is normally made by literature precedent. With the arsenal of Lewis acids available a suitable catalyst can usually be found.

By this strategy, reactive carbon electrophiles can be generated for successful reaction with a variety of weak carbon nucleophiles. More important examples include the Friedel–Crafts reaction in which aromatic compounds (nucleophiles) react with alkyl and acyl halides (electrophiles in the presence of Lewis acids).



#### NEUTRAL CARBON NUCLEOPHILES 241

The Mukaiyama reaction is a versatile crossed-aldol reaction that uses a silyl enol ether of an aldehyde, ketone, or ester as the carbon nucleophile and an aldehyde or ketone activated by a Lewis acid as the carbon electrophile. The product is a  $\beta$ -hydroxy carbonyl compound typical of an aldol condensation. The advantages to this approach are that it is carried out under acidic conditions and elimination does not usually occur.



The transition state is thought to be an open structure. Assuming that a particular silyl enol ether geometry is used, the substituents will tend to occupy opposite faces of the transition state and thus give a particular diastereomer (syn-anti) preferentially. Because of the open transition state geometry, the diastereoselectivity is not high.

The reaction of allyl silanes with aldehydes and ketones activated as electrophiles by Lewis acids is a very useful method for preparing homoallylic alcohols. Since allyl silanes are only modestly nucleophilic, strong electrophiles are needed to ensure a good reactivity match.



The same considerations apply to intramolecular versions. For example, although epoxide **E** is a stable compound, treatment with a Lewis acid activates the epoxide as an electrophile and cyclization with the olefinic  $\pi$  system occurs to give the steroid ring system. Note that the silyl enol ether group of **E** functions as the terminating group. This polyene cyclization is similar to the way in which steroids are biosynthesized. The cyclization is triggered by the generation of an electrophile sufficiently strong to react intramolecularly with the weakly nucleophilic  $\pi$  bond.

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#### **C=C FORMATION**

In addition to connecting skeletal fragments by formation of carbon–carbon single bonds, it is also possible to utilize reactions which give carbon–carbon double bonds to assemble carbon skeletons. It should be recognized that while the final products of such reactions contain a carbon–carbon double bond, they are generally sequential processes in which a single carbon–carbon bond is formed first and the  $\pi$  bond is formed in a subsequent elimination step.

An elementary example of this process is the reaction of an organometallic reactant with a ketone (or aldehyde) followed by dehydration of the resulting alcohol to the olefin. This is truly a sequential process in that the product alcohol is dehydrated in a second, independent reaction step. It suffers as a useful synthetic method because regioisomers are often formed in the elimination step.

$$\overset{O}{\longleftarrow} + \text{R-CH}_2-\text{M} \longrightarrow \overset{OH}{\swarrow}_{CH_2}^{R} \overset{-\text{H}_2O}{\longrightarrow} \overset{R}{\longrightarrow}^{R}$$

Alternatively it is possible to have both steps, addition and elimination, occur spontaneously if appropriate reagents are employed. There are two common strategies in use: the Wittig reaction and the Wittig–Horner reaction. The Wittig olefination uses a phosphorus-stabilized carbanion (ylid) as a nucleophile and a carbonyl compound as an electrophile. Typically the ylid is generated in situ from a triphenylphosphonium salt and a strong base such as LDA or an alkyl lithium.

$$\begin{array}{ccc} \text{R-CH}_2\text{-Br} &+ & : P(C_6H_5)_3 \longrightarrow & \text{RCH}_2 \longrightarrow & P(C_6H_5)_3 \xrightarrow{\text{LDA}} & \bigoplus & \bigoplus & \bigoplus & \bigoplus & \text{RCH} \longrightarrow $

The ylid is a neutral compound which is resonance stabilized by phosphorus. The phosphorus atom, being a second-row element, has unfilled d orbitals in the valence shell that can accept electrons from carbon. Consequently a major resonance contributor is a structure without formal charges which has a carbon-phosphorus double bond. Nevertheless in the resonance hybrid the carbon atom next to the phosphorus is electron rich and is a good carbon nucleophile which can add to carbonyl groups to form new carbon-carbon bonds. The cyclic intermediate (oxaphosphetane) spontaneously loses triphenylphosphine oxide at room temperature to give an olefin.



In sum, a new olefinic link is produced, but by an addition–elimination sequence. In this reaction a stronger C–O double bond in the starting material is replaced by a weaker C–C double bond in the product. The thermodynamic driving force for the reaction is the formation of the P–O bond, which is very strong.

The Wittig reaction is a very important method for olefin formation. The stereochemistry about the new carbon–carbon double bond is the Z (or less stable) isomer. This unusual stereoselectivity indicates that product formation is dominated by kinetic control during formation of the oxaphosphetane.

By adding a strong base to the cold solution of the oxaphosphetane before it eliminates, the oxaphosphetane equilibrates to the more stable anti isomer and the E olefin is produced upon elimination. This so-called Schlosser modification in conjunction with the normal Wittig reaction enables either the Z or E isomer of the olefin to be prepared selectively.

The Wittig-Horner reaction is the Wittig process applied to carbonyl-activated ylids and uses trimethylphosphite as the phosphorous reagent. Reaction with a bromoester gives a phosphate intermediate. Deprotonation with a base such as sodium hydride and addition of an aldehyde or ketone gives, after elimination of a phosphonate, an  $\alpha$ , $\beta$ -unsaturated ester. In this case the intermediate betaine is acidic and undergoes equilibration prior to elimination so that only the more stable E regioisomer is produced.



A recent alternative to the Wittig reaction uses silicon as the atom which promotes oxygen loss. This reaction, called Peterson olefination, uses an  $\alpha$ -silyl anion as the carbon nucleophile and a carbonyl compound (aldehyde or ketone) is the electrophile. Thus ethyl  $\alpha$ -trimethylsilylacetate can be converted to an enolate and reacts with an aldehyde to give an  $\alpha,\beta$ -unsaturated ester. The driving force for elimination is the formation of an extremely strong silicon–oxygen bond, which converts the oxygen atom into a much better silyloxy leaving group. Only the more stable olefin isomer is produced since equilibration occurs in the enolate intermediate.



Another common  $\alpha$ -silyl anion is produced by the halogen exchange from a methyl (but not other group) attached to silicon. Other  $\alpha$ -silyl carbanions can be generated by other processes. Such anions lack the resonance stabilization of an ester group seen in the previous example. They are consequently less stable and must be generated under carefully controlled conditions. They are good nucleophiles and add effectively to aldehydes and ketones.

$$(C_{6}H_{5})_{3}Si - CH_{2}Br \xrightarrow{n-BuLi} (C_{6}H_{5})_{3}Si - CH_{2}Li + \bigwedge_{R_{1}}^{O} \xrightarrow{O} H_{2}C \xrightarrow{R_{1}} (C_{6}H_{5})_{3}Si - OLi$$

$$\equiv Si \longrightarrow + R-Li \longrightarrow \equiv Si \longrightarrow Li + \bigwedge_{R_{1}}^{Li} R_{2} \xrightarrow{O} R \xrightarrow{H} \xrightarrow{R_{1}} R_{2}$$

#### CYCLOPROPANATION REACTIONS

Formation of two single bonds from a carbon atom is also well known for building up carbon skeletons. In this process, a three-membered ring is formed by reaction of a difunctional carbon atom with an olefin. Because of the strain of threemembered rings, their synthesis is not trivial and a small number of reactions which effectively append three-membered rings to molecules are important and widely used.



The Simmons–Smith cyclopropanation utilizes methylene diiodide and a zinc–copper couple to produce a carbenoid intermediate. This intermediate reacts with olefins to give cyclopropanes. The geometry of the double bond is preserved in the cyclopropane.



This addition is sensitive to steric biases in the olefin, and the methylene group will enter from the least hindered side of the molecule. Alcohol substituents in the olefin will facilitate the reaction and guide the methylene group syn to the alcohol.



The base-promoted  $\alpha$  eliminations of chloroform or bromoform provide a simple method for the production of dihalocarbenes. These add readily to olefinic double bonds to give 1,1-dihalocyclopropanes. The halogens can be removed (one or both) by reduction; the most common method is to use tri-*n*-butyltin hydride.

Another common method for forming cyclopropanes is to react  $\alpha$ -diazoketones or esters with olefins under the influence of copper or, better yet, rhodium or ruthenium catalysis. Again a metal carbenoid intermediate is produced which reacts with the olefin.



The importance of this strategy is that functionalized cyclopropanes are produced which can be further manipulated. The process can also be carried out intramolecularly with high efficiency.



#### METAL-CATALYZED CARBON-CARBON BOND FORMATION

Most of the examples of carbon-carbon bond-forming reactions discussed earlier utilize stoichiometric quantities of the carbon nucleophile and the carbon electrophile. Most commonly the nucleophilic species is formed in solution (organometallic, enolate, phosphorane, etc.) and the electrophile is added. Thus stoichiometric quantities of reagents such as bases or metals are used to form the reactive nucleophile. These are, for the most part, classic processes that work well and give predictable results. Many have been used quite successfully for well over one hundred years for assembling organic structures.

In the last several decades there has been ever-increasing interest in discovering new methods for the formation of carbon–carbon bonds utilizing transition metal complexes as catalysts. The reasons for doing this are many:

- 1. Many new reactions are possible which might overcome the functional group requirements and/or limitations known for many of the older methods. The metal complex provides a structurally defined catalytic center at which reagents are brought together and reacted. This contrasts with the random collisions characteristic of many traditional uncatalyzed processes. Moreover the multiple oxidation states available to transition metals provide new mechanisms for the activation of reagents involved in the bond-forming process.
- Different ligands complexed to the metal might be used to modulate the reactivity of the catalyst and thus achieve increased selectivity. This would increase the yields and efficiencies of the reaction and lead to single products rather than mixtures.
- 3. Chiral ligands would offer the opportunity to introduce chirality into the transition state between two achiral reaction partners. In this way stereos-election could be tuned via the choice of catalyst and ligands.
- 4. High turnover numbers would allow very small amounts of catalyst to be used, thus simplifying the workup and purification of reaction products.
- 5. Finally, such processes would have high atom economy. This means that most of the atoms put into the reaction mixture end up in the products and not the by-products.

As one might expect, the fairly large number of transition metals has resulted in a very large number of organic transformations. Virtually all of the transition metals have been used to create new carbon-carbon bonds in various reactions. However, two metals in particular have had the greatest impact on preparative organic chemistry. The use of palladium and ruthenium complexes for the formation of carbon–carbon (and other) bonds has led to profound changes in synthetic planning. The reactivity patterns of complexes of these metals are quite general and quite distinct from the reactivity patterns of traditional carbon–carbon bond-forming reactions. Moreover they are extraordinarily tolerant of diverse functional groups. As a result completely new strategies for the construction of carbon skeletons have emerged which complement older methods. What results is a much richer synthetic toolkit for building molecules.

The following discussion will focus on the most common and general reactions these two metals catalyze. Given the tremendous amount of information that has been gathered, it comes as no surprise that it is impossible to cover this chemistry fully in this text. Instead the general principles will be developed and some common applications will be illustrated. The idea is to provide a foundation for understanding the reactivity patterns of these complexes. Further insight can be gained from a number of specialized monographs and review articles.

#### Pd(0)-CATALYZED CARBON-CARBON BOND FORMATION

The most important catalytic reactions for the formation of carbon–carbon bonds involve the chemistry of Pd(0). Complexes of zero-valent palladium such as Pd(PPh₃)₄ are available commercially or can be prepared in situ by the reaction of Pd(II) salts [e.g., Pd(OAc)₂, PdCl₂, etc.] with phosphines or other reductants. The most stable complexes are those in which the sum of d electrons from the metal and electrons donated by ligands totals 18. Palladium(0) has a d¹⁰ electron configuration and thus normally coordinates with four ligands which each donate a pair of electrons to the metal. Such complexes are said to be coordinatively saturated and tend to be stable and relatively unreactive. However, dissociation of one or two ligands in solution produces a 16- or 14-electron complex which is reactive and seeks to regain the 18-electron configuration.

Palladium(0)-catalyzed transformations generally involve three steps: oxidative addition, insertion or transmetallation (really a special type of insertion), and reductive elimination. Together they comprise a pathway for the formation of new carbon–carbon bonds. Oxidative addition takes place when a coordinatively unsaturated Pd(0) species cleaves a covalent bond to give a new complex in which the palladium is oxidized to Pd(II). Typically dissociation of two phosphine ligands to a 14-electron complex is the first step followed by oxidative addition to give a 16-electron Pd(II) complex.

Oxidative addition

 $Pd(PPh_{3})_{4} \longrightarrow Pd(PPh_{3})_{2} + 2 PPh_{3}$  14 electron, Pd(0)  $Pd(PPh_{3})_{2} + R-X \longrightarrow R-Pd-X(PPh_{3})_{2}$  16 electron, Pd(II)

This is quite analogous to the formation of a Grignard reagent by the oxidative addition of Mg(0) to an alkyl halide. What is remarkable is the generality and functional tolerance of the palladium process. A variety of bonds undergo oxidative addition with Pd(0). Bonds from carbon to halogen and other good leaving groups such as sulfonates, esters, and phosphonates are used most often (and often referred to as C–X or R–X bonds), but many other bond types are known to react. Even though the product of oxidative insertion has a carbon–palladium bond, this bond is unaffected by most functional groups. Thus alcohols, amines, amides, esters, ketones, aldehydes, and even carboxylic acids can be present in the substrate without interfering with the addition reaction or subsequent reactions. This is a truly phenomenal tolerance for functionality!

Another interesting facet of Pd(0) oxidative insertion is the chemoselectivity of the process. The most reactive bonds are vinyl and aryl C–X bonds, whereas with most other metals these are the least reactive bond types. Palladium(0) also inserts into allylic halides and esters, acid halides, and several other bonds but reacts only sluggishly with C–X bonds to saturated carbon. Taken together these characteristics make Pd(0) chemistry nearly unique.

The second step is insertion or transmetallation. An insertion reaction occurs when the palladium-carbon bond adds across a  $\pi$  bond to give a new organopalladium species. The types of  $\pi$  bonds normally reactive include alkenes, dienes, alkynes, carbon monoxide, and sometimes carbonyl  $\pi$  bonds. By far the most common reactions use alkenes and alkynes for the insertion reaction. This step results in a new carbon-carbon bond.

Insertion

 $R-Pd-X(PPh_3)_2 + A=B \longrightarrow R-A-B-Pd-X(PPh_3)_2$ 

for example,

 $R-Pd-X(PPh_3)_2 + CH_2 = CH_2 \longrightarrow R-CH_2-CH_2-Pd-X(PPH_3)_2$ 

The regiochemistry of the insertion results from a combination of factors which are still being sorted out. It is possible to think of the carbon attached to palladium as electron rich, and it tends to attack the  $\pi$  system at the least electron rich position. Thus alkenes with electron-withdrawing groups react faster than alkenes with electron-donating groups. It is quite paradoxical, however, that alkenes, dienes, and alkynes react much more readily than carbonyl compounds, even though the latter are much more electron deficient.

Moreover there appears to be a steric bias which causes the R group to attack the least hindered end of the  $\pi$  system. In cases where the two ends of the  $\pi$  bond are similar or where electron-donating groups are attached, the regiochemistry can be very sensitive to the reaction conditions and the ligands that coordinate to palladium. At this point controlling the regiochemistry in such systems is more art than science! Nevertheless, in most cases it is possible to predict the regiochemistry with good success. Finally the stereochemistry of the insertion is syn; thus the insertion appears to be a concerted 1,2 addition across one face of the  $\pi$  system.



Transmetallation occurs when compounds with bonds from carbon to several main-group elements (e.g., B, Al, Sn, Si, Hg) are present in the reaction mixture. The palladium intermediate from oxidative addition can undergo exchange of palladium and the main-group element. This essentially yields a second carbon ligand bonded to palladium. The most common compounds used for transmet-allation are tributyl tin compounds (R–SnBu₃) and boronic acids [R–B(OH)₂]. Again the most common and successful examples have the main-group element bonded to an aromatic ring or an alkene.

Transmetallation

 $R-Pd-X(PPh_3)_2 + R'-M \longrightarrow R-Pd-R'(PPh_3)_2 + MX$ M = main-group element

for example,

$$\begin{array}{c|c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & & \\ & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & &$$

The last step is reductive elimination in which the organic product is liberated and Pd(0) is regenerated to begin the catalytic cycle again. When there are two carbon ligands attached to palladium, as is the case when a transmetallation has occurred, the two carbon fragments couple with the expulsion of Pd(0). This occurs rapidly after a transmetallation and in these instances is the step in which carbon–carbon bond formation occurs.

Reductive elimination

 $R-Pd-R'(PPh_3)_2 \longrightarrow R-R' + Pd(PPh_3)_2$ 

for example,

A second common reductive elimination process termed  $\beta$ -hydride elimination occurs when there is a hydrogen atom  $\beta$  to the carbon–palladium bond, as in the case where an insertion reaction has taken place. The palladium atom inserts into the  $\beta$  carbon–hydrogen bond to give a palladium hydride species coordinated to the alkene. This is a reversible reaction and is akin to the process of alkene hydrogenation catalyzed by palladium. Dissociation of the alkene and elimination of HX gives back the Pd(0) catalyst. Since a strong acid is liberated in the  $\beta$ elimination, a base such as triethylamine is usually added to the reaction mixture to scavenge this acid. Although the formation of alkenes by  $\beta$ -hydride elimination is a facile process, it is not possible to form an alkyne or allene by  $\beta$ -hydride elimination from a vinyl palladium species.

 $\beta$ -Hydride elimination

$$\begin{array}{c} \text{R-CH}_2\text{-}\text{CH}_2\text{-}\text{Pd-X}(\text{PPH}_3)_2 \longrightarrow \text{H-Pd-X}(\text{PPH}_3)_2 \longrightarrow \text{R-CH} = \text{CH}_2 + \text{Pd}(0) + \text{HX} \\ | \\ \text{R-CH} = \text{CH}_2 \end{array}$$

for example,

$$R \xrightarrow{Pd-X(PPh_3)_2} R \xrightarrow{Pd-X(PD$$

While the above reactions represent only a small fraction of the reactions known for palladium, they form the basis of a powerful methodology for building carbon structures. Several variations have been developed which utilize certain types of reactants and give particular types of products. All these variations, however, contain a common theme. In each case an electron-deficient reagent (e.g., a vinyl halide or aromatic triflate) reacts with an electron-rich reagent (e.g., an alkene, an organoborane, or an organotin) with the formation of a new carbon–carbon bond. In that sense these reactions are related to the reactions between carbon nucleophiles and carbon electrophiles discussed previously in this chapter. They are quite different, however, because they proceed *only* in the presence of Pd(0). In fact they proceed *only* in the coordination sphere of Pd(0). The ability of Pd(0) to catalyze these reactions is nearly unique! We will now examine some of the more common processes.

#### **Heck Reaction**

The Heck reaction involves the coupling of an organopalladium species formed by oxidative addition to an alkene followed by  $\beta$ -hydride elimination. The product is an alkene in which a vinyl hydrogen on the original alkene is replaced by the organic group on palladium. Thus aryl and alkenyl halides can be coupled to alkenes.

$$\begin{array}{ccc} \text{R-X} + \text{Pd}(0) & \xrightarrow{\text{oxidative}} & \text{R-Pd-X} + & & & & & \\ \hline \text{R} = \text{aryl, alkenyl} & & & & & & \\ \end{array} \xrightarrow{\text{R-Pd-X}} & & & & & & & & \\ \hline \text{R}' & & & & & & & \\ \hline \text{R}' & & & & & & & \\ \hline \text{R}' & & & & & & & \\ \hline \text{R}' & & & & & & \\ \hline \text{R}' & & & & & & \\ \hline \text{R}' & & & & & & \\ \hline \text{R}' & & & & & & \\ \hline \text{R}' & & & & & & \\ \hline \text{R}' & & \\ \hline \text{R}$$

Because the by-product of the coupling is a strong acid, bases are usually added to the reaction mixture to scavenge it. For example, 4-iodobromobenzene can be coupled with methyl acrylate to give the 4-bromocinnamate ester in >68% yield. This reaction takes advantage of the faster oxidative addition to the carbon–iodine bond to give a single product.



The Heck reaction was discovered in the early 1970s and is extremely useful for rapidly assembling carbon skeletons. This reaction is unique to palladium! A great deal of information is known about the reaction. For example, the success of the reaction depends on each of the three steps involved. Electron-donating groups decrease the reactivity of alkenyl halides and triflates toward Pd(0), whereas electron-withdrawing group increase the rate of oxidative addition. In cases where Pd(II) salts are used, it is assumed that they are converted to Pd(0) by some redox process.

The insertion reaction is stereospecific and syn. Moreover the  $\beta$ -hydride elimination is also syn. For acyclic alkenes there is free rotation in the organopalladium intermediate so that the more stable *trans*-alkene is formed. Electron-withdrawing groups in the alkene also increase the rate of the insertion reaction and give higher yields generally, but the reaction is limited to relatively sterically unhindered alkenes. In general, polar solvents such as DMF or acetonitrile are most commonly used. There are several common additives which aid in the reaction. These include lithium or tetraalkylammonium chlorides and bromide, silver salts, or cuprous iodide, but exactly how they function is unknown at present.

The conversion of carbonyl compounds to their enol triflates provides a very simple way to couple the carbonyl carbon to an alkene. In general, however, aryl

and vinyl iodides are the preferred substrates because of their ease of oxidative addition. Terminal alkynes are also good coupling partners.



Intramolecular versions of the Heck reaction are very useful for the construction of ring systems. The entropic advantage of having both coupling partners present in the same molecule increases the efficiency of the insertion reaction and leads to efficient reactions. Moreover the intramolecular version can be carried out on hindered substituted alkenes, whereas the intermolecular Heck reaction is largely restricted to monosubstituted alkenes. These reactions illustrate the syn stereochemistry of both the insertion reaction and the elimination. A number of multicyclic natural products have been synthesized using intramolecular Heck reactions to assemble the skeletons, and this has become a powerful synthetic tool for such compounds.



#### Suzuki Coupling

The coupling of organoboron compounds with aryl or alkenyl halides is called the Suzuki reaction and was discovered in the early 1980s. This is a tremendously versatile method for joining two carbon fragments and is widely used in the commercial manufacture of pharmaceuticals, in the synthesis of compound libraries, and in drug discovery. After oxidative addition to the halide, the organopalladium intermediate undergoes transmetallation with the boronic acid or ester. The new carbon–carbon bond is formed in the reductive elimination which produces the product and regenerates the Pd(0) catalyst. A base must be present for the transmetallation to proceed, and oxybases such as alkoxides, carbonates, or hydroxide are most commonly employed. The reaction is highly tolerant of a wide variety of functional groups and thus extremely versatile.

$$R-X + Pd(0) \xrightarrow{\text{oxidative} \\ \text{addition}} R-Pd-X + R'-B(OH)_2 \xrightarrow{\text{transmetallation}} R-Pd-R' \xrightarrow{\text{reductive} \\ \text{elimination}} R-R' + Pd(0)$$

As noted for the Heck reaction, aryl, alkenyl, and alkynyl bromides, iodides, and triflates are best for the oxidative addition. However, aromatic, heteroaromatic, alkenyl, and even alkyl boronic acids and esters can be coupled effectively. The reaction appears almost oblivious to other functional groups present!



Since the oxidative addition occurs with retention of configuration and the transmetallation is also stereospecific with retention, the method is extremely valuable for the stereoselective synthesis of conjugated dienes. The

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#### 254 CARBON-CARBON BOND FORMATION

stereochemistry of the products is determined by the stereochemistry of the coupling precursors.



The required vinyl boranes and vinyl iodides can both be easily made by the hydroboration of alkynes with disiamyl borane (Sia). Thus the Suzuki reaction is an important methodology for the synthesis of conjugated polyene natural products.



#### Stille Coupling

Stille coupling was also developed in the early 1980s and is similar to Suzuki coupling in its sequence. It is used to couple aryl or vinyl halides or triflates with organotin compounds via oxidative addition, transmetallation, and reductive elimination. The oxidative addition reaction has the same requirements and preferences as discussed earlier for the Heck and Suzuki reactions. The reductive elimination results in formation of the new carbon–carbon bond. The main difference is that the transmetallation reaction uses an organotin compound and occurs readily without the need for an oxygen base. Aryl, alkenyl, and alkyl stannanes are readily available. Usually only one of the groups on tin enters into

the coupling reaction, and different groups transfer to palladium with different selectivities. Since simple alkyl groups have the lowest transfer rate, the most common tin reagents have three simple alkyl groups (usually methyl or *n*-butyl). The fourth group which is transferred is alkynyl, aryl, alkenyl, benzyl, or allyl.

$$R-X + Pd(0) \xrightarrow{\text{oxidative}} R-Pd-X + R'-SnR''_{3} \xrightarrow{\text{transmetallation}} R-Pd-R' \xrightarrow{\text{reductive}} R-R' + Pd(0)$$

Coupling of an aryl triflate with an arylstannane is a good method for the preparation of biaryls and other bis-aromatic species of all types. Coupling of vinyl groups takes place with retention of stereochemistry. Furthermore transfer of the allyl group occurs smoothly.



This is very robust chemistry that works very well with enol triflates. Intramolecular reactions have been used to close rings of many sizes, including large rings.



The use of  $(Me_3Sn)_2$  provides a unique way to convert vinyl and aryl halides into the very tin reagents needed for subsequent Stille couplings!



#### **OLEFIN METATHESIS**

In the last 10 years or so an exciting new strategy has emerged for the formation of carbon–carbon double bonds, namely olefin metathesis. This work grew out of the development of Ziegler–Natta catalysts for the polymerizarion of cyclic olefins. It was found that when 2-pentene was treated with a catalyst prepared from tungsten hexachloride and ethylaluminum dichloride, a mixture of 2-pentene, 2-butene, and 3-hexene was produced in minutes at room temperature (rt)!



It was shown that the mixture was an equilibrium mixture. Thus it appears that the alkenes are being broken apart at the double bonds and the pieces reassembled randomly. This process was termed olefin metathesis because the ends of the carbon–carbon double bonds are being interchanged.

It was subsequently shown that the active catalytic species is a metallocarbene complex which contains a carbon-metal double bond. This catalyst undergoes a 2+2 addition with an alkene to give two metallocyclobutanes 1 and 2 (below) which depend on the regiochemistry of addition. Metallocyclobutane 1 is a non-productive intermediate since cleavage across the ring by A or B gives back the reactants. Metallocyclobutane 2 can cleave across the ring A' to give the original alkene and metallocarbene or it can cleave in the opposite direction B' to give a new alkene and a new metallocarbene. The new metallocarbene can add the starting material to give two metallocyclobutanes 3 and 4. It is seen that 3 is a nonproductive intermediate whereas 4 can give another new alkene by cleavage in the B' direction. Since each step is reversible, it can be seen that ultimately the ends of the double bond will be exchanged to give a mixture of olefins.



Olefin metathesis is a unique reaction and is only possible by transition metal catalysis. In fact only complexes of Mo, W, Re, and Ru are known to catalyze olefin metathesis. Once it was known that metallocarbenes were the actual catalytic species, a variety of metal carbene complexes were prepared and evaluated as catalysts. Two types of catalysts have emerged as the most useful overall. The molybdenum-based catalysts developed by Schrock and ruthenium-based catalysts developed by Grubbs.



Both are stable metallocarbene complexes, but they have very different reactivity profiles. The molybdenum catalyst is highly reactive and is effective with sterically demanding olefins. Its drawbacks are that it is not highly tolerant of diverse functional groups and has high sensitivity to air, moisture, and solvent impurities. The ruthenium system, on the other hand, is catalytically active in the presence of water or air, and it exhibits a remarkable functional group tolerance. It is not a reactive as the molybdenum catalyst, particularly toward sterically bulky substrates. However, it is readily available and is the reagent of choice for all but the most difficult substrates.

At present the most synthetically important metathesis process is ring-closing metathesis (RCM) as it can be used for the preparation of a wide variety of medium to large ring compounds from acyclic diene precursors. For such dienes there are two competing processes. The first is RCM, in which the two olefinic units undergo intramolecular reaction to give a cyclic product. The second is intermolecular metathesis, which yields open-chain dimers or oligomers.



The rate of formation of the RCM product is given by a first-order rate law since the metathesis is intramolecular,  $rate_{RCM} = k_1$  [catalyst][diene]. The rate of dimer formation is a second-order rate law because the reaction is intermolecular,  $rate_{dimer} = k_2$  [catalyst][diene]². Although these two rate laws are not strictly comparable because of their different orders, at normal concentrations (0.25–1 M) the difference in rates is mostly due to the difference in the rate constants  $k_1$  and  $k_2$ . Given that the double-bond energies in five- and six-membered rings are very similar to the bond energy of an open-chain double bond, the  $\Delta H^{\pm}$  for the two processes are similar. Thus  $\Delta H^{\pm}$  cannot be responsible for a significant difference in the rates of the two reactions.

The rate constant  $k_1$  for the RCM process is greater than the dimerization rate constant  $k_2$  because of a distinct entropic advantage. In RCM the two reacting bonds are present in the same molecule and two molecules (the product and ethylene) are formed from one; thus RCM proceeds with a gain in entropy. The dimerization process causes loss of translational degrees of freedom because one molecule is formed from two and thus occurs with a loss of entropy. As a result the formation of five- and six-membered rings have larger rate constants and thus proceed at faster rates than dimerization under normal conditions.

The use of the Grubbs catalyst is largely restricted to terminal dienes. In these cases the by-product is ethylene. Since this is an equilibrium process, the ethylene diffuses out of solution, which helps drive the reaction to completion. Nevertheless the catalyst is highly tolerant of functional groups and generally gives good yields.



The formation of medium rings presents challenges for this and most ringforming reactions. There are two major reasons for this. The first is that the molecule must be able to adopt a reactive conformation in which the two double bonds are in close proximity for reaction to occur. For larger ring sizes where the distance between double bonds increases, the population of reactive conformations decreases so the rate slows as well. The second factor is the increased ring strain in medium rings which develops at the transition state and thus raises  $\Delta H^{\pm}$ and slows the rate of the intramolecular process. The rate of the intermolecular dimerization remains roughly the same. Thus the rate of the intramolecular process for medium rings slows dramatically while the dimerization rate remains the same (for a given concentration).

This behavior is easily demonstrated by the following examples. Formation of a five-membered ring takes place easily under mild conditions. Formation of a similar eight-membered ring from an open-chain precursor gave no RCM product but only dimerization. In contrast bis-allylcatechol gives the eight-membered ring RCM product efficiently. The phenyl ring provides a conformational constraint which keeps the olefins in reasonably close proximity and the lack of C–H bonds in the catechol fragment reduces transannular strain in the eight-membered ring. These factors allow the RCM to be favored over dimerization.



A second way to increase the proportion of RCM is to keep the concentration of the starting diene as low as is feasible. Because dimerization is a secondorder reaction, decreasing the concentration of the diene decreases the rate of the second-order reaction much faster than the rate of the first-order RCM reaction decreases. Thus carrying out the reaction at very low concentrations can actually result in the RCM becoming significantly faster than dimerization. There are several ways to do this experimentally. The first is to simply dilute the reaction mixture to the desired concentration and then run the reaction normally. Although simple, this approach requires a lot of solvent which must be paid for and then removed. A simpler process is to use very slow addition of a solution of the diene to a solution of the catalyst. If the diene reacts as it is added, then the concentration of the diene remains very low over the course of the addition without having to use large volumes of solvent to achieve it.

For large rings where ring strain is again minimal, only the effective concentration of reactive conformers limits the RCM process. In these cases, running the reaction at high dilution slows the dimerization sufficiently to allow successful formation of large rings. This is truly a remarkable ability since large rings are difficult to access by other reactions.



The effect of conformation on the efficiency of RCM is extremely important. The RCM of the polyether diene shown below gives only a 39% yield under normal conditions. Addition of lithium ions to the reaction mixture causes the ether oxygens to wrap around the lithium cation. This forces the olefinic bonds into close proximity and the RCM product is now formed in >95% yield. Other structural features such as hydrogen bonding, restricted rotation (as in amides), or steric effects which cause particular conformations to be favored are very important in the formation of large ring systems. Such effects can affect RCM either positively or negatively depending on whether the olefinic bonds are held proximal or distal to each other.

#### BIBLIOGRAPHY 261



The development of RCM has dramatically changed synthetic planning for ring systems, particularly medium and large ring systems. It represents a new paradigm for assembling carbon skeletons of ring compounds. Further advances will only increase its importance as a method for carbon–carbon bond formation.

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#### PROBLEMS

**8.1.** Give the products of the following reactions. Indicate the new carbon–carbon bond that has been formed and identify the carbon nucleophile and electrophile.



PROBLEMS 263



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**8.2.** Give reactions which would produce the indicated bond in the following compounds. Give the reacting partners and tell which is the electrophile and nucleophile. Also tell how you would generate any reactants which are not stable compounds.



264 CARBON-CARBON BOND FORMATION

PROBLEMS 265



**8.3.** Show how you would prepare each of the following molecules from the indicated starting materials. Where more than one step is required, show each step clearly.



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**8.4.** Give the products of the following reactions. [The reagent Pd(0) is a generic term referring any one of a number of possible zerovalent palladium reagents. The reagent referred to a Grubbs catalyst is the ruthenium carbenoid species referred to in the chapter.]





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#### CARBON-CARBON BOND FORMATION



PROBLEMS 269



# iranchembook.ir/edu

**270** CA

#### CARBON-CARBON BOND FORMATION



**8.5.** Show a method for constructing the following compounds from the indicated starting materials.



**8.6.** It was observed that when urea **6A** was treated with Grubbs catalyst under high dilution, only dimers and oligomers were produced. However, when



urea 6B was reacted under the same conditions, cyclic urea 6C was produced in 61% yield. Provide a rationalization for these results.

# 9

# CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS

Free-Radical Reactions	272
Free-Radical Polymerization	277
Nonpolymerization Reactions	278
Free-Radical Initiation	280
Free-Radical Cyclization	283
Bibliography	288
Problems	288

#### FREE-RADICAL REACTIONS

In the previous chapter the formation of carbon–carbon bonds was discussed in terms of polar or two-electron processes. In such reactions one carbon serves as an electron donor (nucleophile) and a second carbon serves as an electron pair acceptor (electrophile). The result of the donor–acceptor interaction of these two species is a new carbon–carbon bond in which the electron pair is shared by the donor and acceptor.

 $\begin{array}{cccc} R_3 C \oplus & and & \ominus C R'_3 \\ & & or & & \longrightarrow & R_3 C - C R'_3 \\ R_3 C \ominus & and & \oplus C R'_3 \end{array}$ 

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Another way to form a bond between two carbons is for each carbon atom to supply one electron. In this case interaction between two carbons which each has a single, unshared electron would result in formation of a carbon–carbon bond. Species with unshared electrons are called free radicals, and thus formation of carbon–carbon bonds by this strategy requires carbon-centered free radicals as reactants.

 $R_3C \cdot \text{and} \cdot CR'_3 \longrightarrow R_3C - CR'_3$ 

Carbon-centered free radicals are carbon atoms which have three bonds and seven valence shell electrons, with the unshared electron occupying a valence orbital. They are generally thought to be planar (sp² hybridized) and the unshared electron is in a 2p atomic orbital. They are not rigidly planar but are easily deformed to a pyramidal geometry. Because they have only seven valence level electrons, free radicals are very reactive intermediates and they rapidly undergo a variety of reactions. Because of their high reactivity, the formation of bonds by the combination of two free radicals is actually rare because the free-radical species must survive long enough to encounter another free radical with which to react. Normally free radicals undergo other reaction processes before they encounter a second free radical with which they can combine.



The reactivity of carbon-centered free radicals results from their drive to achieve an octet electronic configuration, which they do by two principal reaction processes. The first is atom transfer. This process is one in which an atom with one electron is transferred from a closed-shell molecule (fully paired, valence octets) to the free radical.



#### 274 CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS

Due to the conservation of spin, a new radical species is formed. If the atom that is transferred is a hydrogen, then the process is called hydrogen abstraction and is the most common atom transfer reaction; however, other atoms can be transferred to free radicals as well. The driving force for atom transfer (abstraction) reactions is usually the formation of a stronger bond and/or a more stable free radical.

A second very common free-radical reaction is addition to  $\pi$  systems to give a new bond and a new free radical. In this process the  $\pi$  bond is broken.



This process is quite common for carbon-centered free radicals because the carbon–carbon  $\sigma$  bond which is formed is stronger by about 30 kcal than the  $\pi$  bond which is broken. Other radical species, however, are well known to undergo olefin additions as well. The addition of bromine to olefins is the key step in the anti-Markovnikov addition of HBr to olefins.

Another common feature of free-radical reactions is that they tend to be chain processes. Since any chemical reaction must exhibit conservation of spin, the reaction of a free radical with a closed-shell (fully electron paired) molecule must result in the production of a new free-radical species which can participate in subsequent free-radical reactions. The series of free-radical reactions leading to product is often a cyclic process in which the initial free radical is produced once again in the last step of the cycle so that the reaction sequence starts over again. The process is termed a *chain reaction* because each step of the process is linked directly to the preceding step.

Free-radical chain reactions can generally be divided into three phases:

- 1. *Initiation* is the phase of the process in which free radicals are produced that can start the chain reaction.
- 2. *Propagation* is the phase of the process in which free radicals undergo reactions which form products and produce new free radicals which can continue the chain.
- 3. *Termination* is the phase of the process in which free radicals are removed from the system by recombination or other reactions, thus interrupting the chain reaction.

A classic example is the free-radical addition of chloroform to olefins initiated by benzoyl peroxide.



Initiation normally requires molecules with weak bonds to undergo homolytic cleavage to produce free radicals. Since bond homolysis even of weak bonds is endothermic, energy in the form of heat ( $\Delta$ ) or light ( $h\nu$ ) is usually required in the initiation phase. However, some type of initiation is *required* to get any free-radical reaction to proceed. That is, you must first produce free radicals from closed-shell molecules in order to get free-radical reactions to occur. Benzoyl peroxide contains a weak O–O bond that undergoes thermal cleavage and decarboxylation (probably a concerted process) to produce phenyl radicals which can initiate free-radical chain reactions.



Azobisisobutyronitrile (AIBN) is perhaps the most widely used initiator. It undergoes either thermolytic ( $\Delta$ ) or photolytic cleavage ( $h\nu$ ) to give isobutyronitrile radicals which can initiate free-radical reactions.



Hexa-*n*-butylditin can be photolyzed to two tri-*n*-butyltin radicals which are initiators for tin-based free-radical reactions.
276 CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS

$$(n-\mathrm{Bu})_3\mathrm{Sn}-\mathrm{Sn}(n_\mathrm{Bu})_3 \xrightarrow{hv} 2 (n-\mathrm{Bu})_3\mathrm{Sn}$$

Many other free-radical initiators are available as well, and the choice of initiator is normally based on literature precedent and ease of use.

The propagation phase of a free-radical chain reaction is usually a cyclic sequence in which a molecule of product is produced and the propagating radical is regenerated by the sequence. In the above example, the trichloromethyl radical adds to the double bond to give a new carbon-centered radical which abstracts a hydrogen from chloroform to produce a molecule of product and another trichloromethyl radical that continues the chain. Notice that this stage is cyclic and infinite. If a single •CCl₃ were generated, it would continue to form one molecule of product and another •CCl₃ until the olefin was converted completely to product—one molecule at a time. Clearly this would be very slow, but because a single initiation event can lead to many molecules of product, one needs very little initiation. If, for example, the "chain length" of the propagation cycle is 200-300 (a common value), then one would need only a 1/200-1/300 ratio of initiator to olefin to convert the olefin completely to product. Thus initiation at 0.5-0.3% would suffice. If chain lengths are longer, then less initiation is required; if they are shorter, then more initiation would be required.

Termination reactions, while rare, do occur and serve to interrupt propagation cycles by removing propagating radicals from the system. Often these reactions are radical recombinations, but termination reactions also include the reaction of propagating free radicals with other species in solution (called scavengers) to give radicals incapable of participating in the propagation cycle. In the above example, a scavenger could react with either the trichloromethyl radical or the trichloromethyl addition product to give an unreactive free radical and thus interrupt the chain process.



The more effective a termination step is, the shorter will be the propagation cycle and the less product will be produced per initiation event. In the limiting case, if each initiation event was terminated, then *no* product would be produced. This is the role of "antioxidants" added to many products and most processed food. These additives scavenge free radicals produced by the reaction of oxygen with C–H bonds and prevent them from participating in oxidation propagation cycles—thus oxidative degradation is stopped or slowed markedly.

It is the reactivity of free radicals which has made them difficult to understand and control. Because of their great reactivity, they are quite unselective and tend to react with anything in solution, and hence multiple pathways and many products are often the rule. Moreover many initiation methods fail to produce single free-radical species in a controlled and efficient fashion. As a result of these factors, the use of free radicals in preparative organic chemistry has seen two distinct phases: first free-radical polymerization and then nonpolymerization reactions.

#### FREE-RADICAL POLYMERIZATION

The first major use of free radicals was in olefin polymerization reactions. Polymerization reactions are amenable to free-radical initiation for several reasons. First the olefin is the only reagent present so as to minimize competing reactions. Second the initiator radical is produced by heat or light or catalysis in the presence of a huge excess of the olefin. Under these conditions free-radical addition to the double bond is virtually the only process that occurs. Moreover, the new radical species resulting from olefin addition is also produced in the presence of a huge excess of olefin so that it adds to another olefin molecule to give a larger free radical. The process continues. By controlling the purity of the starting olefin and the reaction conditions so that terminations are rare, chain lengths in the tens of thousands can be achieved. This leads to the formation of thousands of carbon–carbon bonds per polymer molecule and extremely long polymer chains.



Termination

As a consequence of the fact that free-radical reactions are chain processes, they are very well suited for the preparation of polymers rather than single products. That is, products are obtained whose size is determined by the number of propagation cycles that occur before a termination event stops the growing chain.

#### 278 CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS

If the number of propagation cycles is between 200 and 300, then the product mixture will contain molecules which contain between 200 and 300 monomers. It is more reasonable to describe the product mixture in terms of the "average molecular weight" rather than a single product with a discrete molecular weight. The physical properties reported for a polymer are those of a mixture of polymeric molecules rather than of a single polymeric compound.

Free-radical polymerization was a mainstay of the plastics industry for many years. While new and better methods have been developed for the polymerization of many substrates, free-radical polymerization is still used for the preparation of many plastics and composites. The success of these methods is based on an understanding of the process. Huge amounts of effort have been expended in finding initiation reactions that produce free radicals controllably and reproducibly. The reaction environment has been studied intensively so that propagation reactions are maximized and termination events minimized. Finally the rational control of termination reactions, which are necessary to control the chain length and thus the average size of the polymer molecules produced, has been successfully developed. It is important to emphasize that the properties of the product mixture were the gauge by which the understanding and control were measured.

Thus it is not necessary to produce one product molecule with a defined molecular weight. It is only necessary to obtain a product mixture whose average molecular weight and physical properties fall within a defined range. Now it is true that obtaining a more narrow range of molecular weights in a polymer leads to much more consistent physical properties. This explains why so much effort was made to control free-radical polymerization. Those efforts played a large role in the creation of the "plastic society" in which we live.

#### NONPOLYMERIZATION REACTIONS

The use of free-radical reactions for the preparation of single molecules as products requires greater control of the various steps in the process. Traditional free-radical addition reactions carried out in solution (to minimize polymerization) often gave low yields and mixtures of products and thus were not of real synthetic value. In the ideal case one would want to be able to generate a specific free-radical species which would undergo a single reaction process and then be terminated. The termination step should give a single product plus the reactive radical to continue the chain. Thus very selective methods of initiation are needed, and a clear understanding of propagation and termination steps is required in order to control product formation more closely.

Initiator 
$$\xrightarrow{\Delta}$$
 2 In •  
In • + M  $\longrightarrow$  M •  
M •  $\longrightarrow$  P + In •

Based on these requirements for the controlled production of single molecule products from free-radical reactions, the second phase of free-radical chemistry began about 20–30 years ago when fast kinetic methods were developed to measure the rate constants for known free-radical processes in solution. Thus it became possible to measure the comparative rates of hydrogen abstraction versus olefin addition and the relative rates of intermolecular versus intramolecular reactions. The results of these investigations led to the realization that intramolecular olefin additions which produce rings are often much faster than other reaction pathways, especially in dilute solutions. Moreover, the formation of five-membered rings by intramolecular olefin addition is much faster than the formation of rings of other sizes. It is this kinetic selectivity which can be used as the basis for efficient carbon–carbon bond-forming processes.



The identification of this single reaction process which occurs much faster than others can be used as a focal point for selective carbon–carbon bond construction. That is, if a radical can be produced on a carbon which is five carbons away from a double bond, then the fastest reaction which occurs is cyclization to a five-membered ring. Because free radicals are uncharged, nonpolar entities, such cyclizations are also found to be largely unaffected by inductive effects or solvents or substitution patterns. As a consequence protecting groups are normally not needed for free-radical cyclizations and a wide range of reaction conditions are compatible with an efficient reaction.



Moreover, if this cyclization can be incorporated effectively in a chain process, the cyclized radical would be trapped and yield a single product.

#### 280 CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS

#### FREE-RADICAL INITIATION

Besides new insight into the reactivity of free radicals, methods for the production of carbon-centered free radicals have also seen major improvements in the last several years. One very common new method is to use tin-based reagents as radical chain carriers. Trialkyltin radicals readily abstract bromine or iodine from carbon to produce a carbon-centered free radical. Placement of a bromide or iodide substituent on a substrate thus permits formation of a carbon-centered free radical at that position using tin-based methodology. This process was initially developed for the reduction of alkyl halides, and it remains an excellent synthetic method for that purpose. The complete chain mechanism for the reduction is shown.

One equivalent of tributyltin hydride is required to reduce one equivalent of alkyl halide. Each step in the sequence is energetically favored so that side reactions are minimized. That is, the tributyltin radical does not abstract hydrogen from the alkyl; it only attacks the halide (Br or I).



Likewise the carbon radical abstracts the hydrogen from tin much more readily than other C–H hydrogens. Thus the process is clean from a mechanistic point of view and consequently leads to clean reduction products.

If this method of free-radical generation is applied to an unsaturated alkyl halide, the free radical has two pathways available. It can abstract a hydrogen from tributyltin hydride and give a reduced product as in the reduction process described above. Alternatively, it can cyclize to give a new cyclic radical which can abstract hydrogen from tributyltin hydride to give a cyclized product. These competing propagation steps are shown.



Now the ratio of products obtained depends on the competing rates of the two processes. The rate of reduction of the radical is given by  $k_a[\mathbf{R} \cdot]$  [Bu₃SnH] while the rate of cyclization is given by  $k_c[\mathbf{R} \cdot]$ . As was mentioned previously, it has been found that  $k_c$  is sufficiently large that cyclization is often the major process. However, reduction is always possible. One way to control reduction is to note that it is a second-order process which depends on both the concentration of the radical R• and tributyltin hydride. Thus the rate of reduction can be lowered simply by running the reaction under dilute conditions. As the concentration goes down, the rate of reduction is favored.

After cyclization has taken place, the only reaction available to the cyclopentyl methyl radical is hydrogen abstraction. Thus it remains in solution until it reacts with tributyltin hydride to produce methylcyclopentane and a tributyltin radical which continues the chain.

A second common way to produce free radicals for use in carbon-carbon bond-making reactions is to use esters of N-hydroxypyridine-2-thione. This method is also tin based and relies on the propensity for tin radicals to add to carbon-sulfur double bonds. Subsequent fragmentation reactions lead to free radicals that are ultimately used in the propagation steps.



The free radicals  $R_{\bullet}$  participate in a product producing a chain process that regenerates  $Bu_3Sn_{\bullet}$  and thus continues the chain. While there are many steps in this overall sequence, it is a chain process. If each step is exothermic and kinetically favored, then each step will proceed selectively and the product will be formed efficiently and reproducibly, irrespective of the number of the steps

#### 282 CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS

in the chain. This turns out to be the case, and the above method is an effective and reliable sequence for initiating free-radical reactions.

It is also known that once the tributyltin radical adds to the sulfur of *N*-hydroxypyridine-2-thione, breakage of the N–O bond and decarboxylation are very fast and probably concerted. They are driven energetically by formation of the very stable carbon dioxide molecule. It comes as no surprise that if atoms other than carbon are attached to the carboxyl group, then they would also end up as free radicals after decarboxylation. This is shown for a urethane analog as a source of nitrogen-centered free radicals. A wide variety of other free-radical species can be produced by this strategy, and it is thus quite useful.



A third tin-based method of free-radical production also utilizes tin radical addition to a carbon–sulfur double bond as a key reaction. In this case a thio-noester (usually a thiono carbonate) is the reactant. As in the previous method, addition of tin to the sulfur atom is followed by fragmentation to a carbon-centered radical.



The driving force for the fragmentation is formation of the C=O double bond. If R• reacts with  $Bu_3SnH$ , a tributyltin radical is produced which continues the chain. Carried out in this manner this reaction is called the Barton deoxygenation of alcohols, since alcohols are precursors for the thiono esters.

#### FREE-RADICAL CYCLIZATION 283



If, on the other hand,  $R_{\bullet}$  is unsaturated and can undergo cyclization rapidly, it will do so. This competition between reduction of the first formed radical  $R_{\bullet}$ and its cyclization to a new cyclic radical  $R'_{\bullet}$  is the same as discussed for the formation of free radicals from alkyl halides and tributyltin radicals. The only difference is in the way in which the carbon-centered radical is produced.

A number of other strategies have been developed for the production of free-radical intermediates for carbon–carbon bond construction; however, the tin-based methods described above are by far the most common. Thus alkyl halides, carboxylic acids, and alcohols are all excellent precursors for free radicals by these methods. The choice of method can be made on the basis of which substrate fits into the synthetic scheme most efficiently or which precursor is most readily available.



#### FREE-RADICAL CYCLIZATION

With good methods available for producing carbon-centered free radicals, the cyclization process can be examined in greater detail. Cyclization involves the intramolecular addition of a free-radical to a double bond. Of course, this requires that the two reacting parts of the molecule, the free-radical center and the  $\pi$  bond, come within bonding distance of one another.



#### 284 CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS

It is quite easy for open-chain systems to undergo intramolecular cyclization because of their many rotational degrees of freedom. More rigid systems undergo efficient cyclization only if the free-radical center and the  $\pi$  system are held in close proximity, as in the first example below. Where the molecular geometry is fixed in such a way as to prevent effective interaction between the free-radical center and the  $\pi$  system, cyclization is inefficient and reduction predominates. Cyclization in the second example is an obvious impossibility!



Other cases are not always so obvious, yet any structural or steric feature which influences the close approach of the  $\pi$  bond and the free-radical center will influence the rate of cyclization and hence the yield of cyclized product. For example, trans-fused cyclopentyl systems are much higher in energy than cis-fused ones; thus the trans-fused cyclopentyl compound does not cyclize effectively and gives only reduction,



whereas the cis-fused cyclizes efficiently with little reduction,



The cyclization itself can produce two different ring sizes depending on which carbon of the double bond is attacked. Of the two possibilities, it is seen that one mode of cyclization gives a secondary radical while the other mode produces a primary free radical.

$$(9.1)$$

$$(9.2)$$

Since the order of free-radical stabilities falls in the order  $3^{\circ} > 2^{\circ} > 1^{\circ}$ , product stability would dictate that cyclization should preferentially occur to give the more stable secondary radical—a six-membered ring in reaction (9.1) (path *a*) and a seven-membered ring in reaction (9.2)(path *a*).

In contrast, is known that the rates of ring-forming free-radical cyclizations are 5 > 6 > 7. Experimentally it was found that reaction (9.1) gives the fivemembered ring product (path *b*) exclusively, and reaction (9.2) gives the sixmembered ring product (path *b*). Thus the regioselectivity of ring formation is controlled not by thermodynamic considerations but by kinetic control of the cyclization. It turns out that bond formation between a radical and a  $\pi$  system stereoelectronically requires an approach angle of about 110° between the freeradical center and the olefinic plane. (This is due to the fact that free-radical addition results from donation of the unpaired electron on the radical into the  $\pi^*$ antibonding orbital of the olefin, which coincidentally makes an angle of about 110° with the olefinic plane.)

In an intramolecular cyclization, attack on the end of the double bond closest to the radical center (an exocyclic cyclization) achieves the proper approach angle. Attack on the other olefinic carbon requires that the radical reach across the double bond to achieve the proper approach angle. This is a higher energy path and is kinetically disfavored. The same arguments hold for cyclizations which can produce six- or seven-membered rings.



#### 286 CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS

A final feature of radical cyclizations is that they are mainly influenced by steric factors and are practically insensitive to inductive effects. Since free radicals are charge neutral, their reactivity is not greatly influenced by either electrondonating or electron-withdrawing groups. For instance, the following cyclizations occur with similar efficiencies even though the electronic character of the cyclizing radicals are vastly different:



It has also been shown that the electronic character of the olefin to which the radical adds has little influence on the efficiency of the intramolecular cyclization. Intramolecular competition between addition to an electron-rich enol ether or a simple double bond gives a 1 : 1 ratio of products, demonstrating that free-radical cyclizations have a remarkable insensitivity to inductive effects.



Resonance effects, on the other hand, can significantly affect the regiochemistry of the cyclization. Resonance delocalization of the unpaired electron of a free radical stabilizes that radical. This is why the allyl radical is much more stable than the *n*-propyl radical. Thus, if a double bond is substituted with a group capable of providing resonance stabilization to a free radical, it undergoes free-radical addition much more readily than a double bond which cannot provide such resonance stabilization.

#### FREE-RADICAL CYCLIZATION 287



Steric effects can also influence the cyclization process markedly. Bulky substituents which hinder the approach of the free radical to the  $\pi$  system can prevent cyclization altogether and give only reduced product.



Below are shown a few examples of the types of complex structures that can be assembled by intramolecular free-radical cyclization. Note the presence of a great many polar functional groups present in the cyclization substrates which are compatible with the process. While the examples shown do not need protecting groups, a great number of other free-radical cyclizations are known which have unprotected alcohols, carbonyl groups, and carboxylic acids in the cyclization precursor.



#### 288 CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS

Free-radical cyclization reactions nicely complement the Pd(0)-catalyzed intramolecular Heck reaction, which also provides cyclic products from unsaturated halides. Free radicals can be generated easily at saturated carbons from saturated alkyl bromides, and the products are reduced relative to the reactants. In contrast, intramolecular Heck reactions work best for vinyl and aryl bromides (in fact they do not work for alkyl halides), and the products are at the same oxidation level as the reactants. Moreover, free radicals attack the double bond at the internal position, whereas palladium insertion causes cyclization to occur at the external carbon.

The advances made in using free radicals as synthetic intermediates in the last 10-20 years have been extraordinary due to new methods to effectively generate free radicals and new insights into their reactivity patterns which allow them to be controlled. As a consequence, the construction of ring systems has been tremendously facilitated.

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#### PROBLEMS

**9.1.** Give balanced chemical equations for each mechanistic step for the following transformations:



#### PROBLEMS 289



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**9.2.** For each of the following reactions, the structures of possible isomeric products are shown. Predict which product will be favored and give the reasons for your prediction.



#### CARBON-CARBON BOND FORMATION BY FREE-RADICAL REACTIONS





**9.3.** Give the products of the following reactions. Where more than one product is likely to be formed in significant yield, indicate which will be the major product.



# 10

# PLANNING ORGANIC SYNTHESES

Retrosynthetic Analysis	292
Carbon Skeleton Synthesis	296
Umpolung Synthons	302
Acetylide Nucleophiles	305
Ring Construction	306
Robinson Annulation	310
Diels-Alder Reaction	312
HOMO-LUMO Interactions	313
Stereoelectronic Factors	316
1,3-Dipolar Cycloadditions	319
Bibliography	323
Problems	324

# **RETROSYNTHETIC ANALYSIS**

The last several chapters have been concerned with learning how to manipulate functional groups, how to make carbon–carbon bonds, and ways to put pieces of molecules together. Next all of these ideas must be integrated into the general idea of synthetic planning. If we are given a particular molecule to synthesize (target), we must be able to plan the actual chemical route to be used in the preparation of the target. The task is to devise a strategy whereby a particular starting material is converted by a series of steps (reactions) to the desired target.

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One fact must be recognized at the outset—that the target is the compound which must be produced from the starting material. This rather obvious statement of the problem is often overlooked by students, but it lies at the heart of synthetic planning. Targets are chosen in order to achieve some purpose. A particular target might be chosen as a drug candidate, or as a potential insecticide, or as a motor oil additive. Whatever its purpose, a target will have particular structural features that must be produced by the synthetic sequence. Getting close won't do!

Starting materials can be chosen by a variety of criteria. For example, a particular starting material might have the same carbon skeleton as the target, it might be a by-product of a chemical plant and therefore be readily available, it might be very cheap and easily obtained from a specialty chemical company, or it might contain a chiral center necessary in the target. Whatever the reason, a particular starting material imposes certain requirements on the synthetic route that must be taken to produce the target.

To transform the starting material into the target, reactions must be chosen which will accomplish the conversion efficiently and with the correct selectivity. Reactions which do not or cannot deliver the target are without merit for the synthesis of that particular target. This relatively simple idea is also often forgotten, and it is common to force a particular reaction to give a product even though the oxidation level, reactivity, regioselectivity, and/or stereochemistry of the reaction are wrong for the target being considered.

Herein lies one of the difficulties of synthetic planning. We tend to learn organic reactions in the forward direction—that is, reactants A and B give product C. This type of information handling is a convergent process in that a set of conditions is imposed which leads to a limited number of potential outcomes—in most cases a single product!

#### $A + B \rightarrow C$

Yet synthetic planning asks the opposite question—what reactants are necessary to give product C? This requires that we think backward from products to reactants. This type of problem solving is a divergent process in that a great many potential reactants are possible and many possibilities must be explored before any one set of reactants is chosen as the solution.

Let's look at a very simple one-step synthesis to illustrate this difference. Given the following set of reactants, methyl cyclohexanol and chromic acid in acetone, it is very easy to write the product as 4-methylcyclohexanone:



This is because we have learned that secondary alcohols are oxidized efficiently to ketones by chromic acid and recognize this as a Jones oxidation. If

#### 294 PLANNING ORGANIC SYNTHESES

4-methylcyclohexanol is recognized as a secondary alcohol, then it must give a ketone with Jones reagent.

In contrast, if we are asked to synthesize the target bicyclic ketone **K** shown below, we must work backward. If we had an alcohol **A** of the same carbon skeleton, then we could oxidize it to the ketone by a Jones oxidation. This step is indicated by a double arrow  $(\Rightarrow)$  and is called a retrosynthetic step. It shows how we work backward ("retro") from the target **K** to a given starting material **A**. (Of course, the actual synthesis would be carried out in the opposite direction—from the starting material **A** to the product **K**.) Alternatively oxidative cleavage (O₃) of olefin **O** would furnish the target, as would the periodate cleavage of exocyclic diol *x*-**D**.



Pinacol rearrangement of endocyclic diol n-D and hydrolysis of dibromide B would also furnish the target K. Thus each retrosynthetic step in our backward analysis corresponds to a synthetic step to the target in the forward direction.



Although it is possible to generate a variety of potential solutions to the synthetic task at hand, the problem is not yet solved. The validity of each step must be checked. For each retrosynthetic step to be valid, there must be a reaction or reagent capable of effecting the transformation with needed chemospecificity, regiospecificity, and stereospecificity to give the target compound as the major product. In the example written above, oxidation of an alcohol (**A**) to a ketone (**K**) is a very facile step, and there are many ways to do this conversion so it is a valid retrosynthetic step. Likewise the olefin cleavage (**O**) or the diol cleavage (x-**D**) are also well known to be clean and would yield the target. Conversely the pinacol rearrangement of n-**D** would probably not be regiospecific, and the hydrolysis of dibromide **B** is a very messy reaction which often gives complex

mixtures rather than single products. These latter two retrosynthetic solutions are therefore invalid solutions to the synthetic problem.

Moreover the following retrosynthetic analysis for the preparation of  $\mathbf{K}$  is not valid either because there is no good reaction to directly convert an olefin to a ketone:



The olefin could be converted to an alcohol and then the alcohol could be oxidized to the target ketone, but this is actually two sequential reactions, not one. They must be written as



Here we see that the first retrosynthetic step, the preparation of a ketone from an alcohol, is valid as discussed above; however, the second retrosynthetic step is not. Although the addition of water across a double bond is a straightforward, common reaction, its use in the present example requires that hydration take place regiospecifically even though there is no significant control element present to ensure the needed regioselectivity. Thus a mixture of alcohols is expected from the hydration reaction rather than the single-alcohol regioisomer needed for oxidation to the target ketone.



no regiochemical control

For each retrosynthetic step to be a valid solution, the reactants must give the appropriate product with needed structural features and control in the forward direction, that is, the direction taken in the actual synthesis. Thus each retrosynthetic step must be checked both forward and backward to effectively plan workable synthetic routes to new molecules.

To carry out an effective retrosynthetic analysis, one must keep in mind certain basic features which must be dealt with during the synthesis. These

#### 296 PLANNING ORGANIC SYNTHESES

include functional groups, oxidation levels, and stereochemistry. In addition, it is often necessary to construct the carbon skeleton so carbon–carbon bond-forming processes must be integrated into the sequence at the best juncture all the while keeping in mind the above considerations.

#### CARBON SKELETON SYNTHESIS

In general, it is most efficient to construct the carbon skeleton first and then adjust the functionality to give the target. Thus in retrosynthetic analysis we most often move backward from the target to compounds which contain functional groups important in carbon–carbon bond-making reactions. As a consequence carbonyl groups play a very important role in retrosynthetic analysis. They are very useful sources of both electrophilic and nucleophilic carbon that can be used in making carbon–carbon bonds. Based on our earlier discussions, a carbonyl group can be seen to influence the polarity of nearby carbons as shown.



By bond polarity and resonance, the carbonyl carbon and a carbon  $\beta$  to the carbonyl carbon can be utilized as electrophilic centers—the carbonyl group by direct nucleophilic addition and the  $\beta$  carbon by Michael addition to an  $\alpha,\beta$ -unsaturated ketone. By resonance interaction, the  $\alpha$  position in carbonyl compounds and  $\gamma$  positions in  $\alpha,\beta$ -unsaturated carbonyl compounds can be converted to nucleophilic centers by proton removal. These "normal" polarities are used frequently in retrosynthetic planning as points of disconnection to establish potential bond-forming steps using carbonyl groups.

As an initial exercise consider the synthesis of C from cyclohexyl bromide. First one has to note relevant facts about the target, such as (a) it contains an ester and (b) it has two more carbons than the starting material so a two-carbon fragment will have to be attached by a new carbon–carbon bond.

#### CARBON SKELETON SYNTHESIS 297



Looking at the starting material, it is noted that it contains an electrophilic carbon; thus the needed carbon–carbon bond could be formed by reaction of a two-carbon nucleophile with the electrophilic center of bromocyclohexane. Noting that the ester group needed in the product acidifies the  $\alpha$  position, it could be used to make the nucleophilic carbon required for carbon–carbon bond formation. Retrosynthetically this can be written as



So the synthesis could be done in one step by making the anion of methyl acetate and reacting it with bromocyclohexane. The polarities of the reaction partners match nicely, but the problem is that alkylations of secondary bromides with enolates often give poor yields. The enolate is a strong base, which promotes elimination in the secondary bromide rather than giving the substitution product needed in the synthesis. Thus elimination from cyclohexyl bromide to cyclohexene would be a major process if the reaction were attempted. While the retrosynthetic step seems reasonable, the synthetic step has known difficulties. It is important to work backward in the retrosynthetic analysis and then check each forward step for validity.

What is needed in the synthesis of C is a two-carbon nucleophile (or its equivalent) which is less basic than an enolate so elimination is not competitive. If product C is recognized as an acetic acid derivative, then the following analysis can be made. A malonate ion used as the carbon nucleophile is much less basic than a simple ester enolate and hence undergoes substitution readily but does not promote elimination effectively, particularly in secondary systems.



Now clearly there will have to be some functional group adjustment in the synthesis because a hydrolysis of the alkylated malonate must be carried out in order to give decarboxylation to the acetic ester derivative. Either an unsymmetric

#### 298 PLANNING ORGANIC SYNTHESES

malonate must be used that can be differentially hydrolyzed or both ester functions of the malonate could be hydrolyzed and after decarboxylation the acid could be reesterified. The actual synthesis could be planned as shown in (10.1) or (10.2):



Synthesis (10.2) contains an extra step but uses very cheap and available starting materials. Synthesis (10.1) is shorter and goes in high yield but requires anhydrous conditions for the alkylation and a more expensive malonate starting material.

The use of Pd(0) to form carbon–carbon bonds was shown to be very effective in many cases. Could one of the coupling methods catalyzed by Pd(0) be used to attach the two-carbon fragment needed to construct the skeleton of **C**? Since we are restricted to cyclohexyl bromide as the starting material, direct reaction with Pd(0) is not feasible because Pd(0) does not give oxidative addition with saturated bromides. Moreover saturated bromides do not undergo transmetallation with Pd(0), so it could not serve as the second component in a Pd(0)-catalyzed coupling. Thus the reactivity requirements of Pd(0)-catalyzed coupling reactions are incompatible with the starting material and thus are not usable for the present construction.

Next consider the synthesis of  $\mathbf{M}$  from "readily available" starting materials. The relevant facts about  $\mathbf{M}$  are that (a) it contains an aromatic ring, an acetate ester, and a vinyl group; (b) it has a straight chain attached to the ring; and (c) all the functional groups are isolated.



To begin the retrosynthetic analysis, note that the acetate ester is easily produced from the corresponding alcohol **A**. Therefore conversion of **A** to **M** using acetic anhydride/pyridine could be used in the synthetic step. (Remember: For each retrosynthetic step, a reaction must be available to accomplish the synthetic step.)

Now the alcohol functional group in A is a natural point for bond disconnection to take place since alcohols are the products of carbon nucleophiles and

carbonyl groups. If we consider bond  $\mathbf{a}$  in our retrosynthetic analysis, then the next retrosynthetic step would be



Since the anion N is a nonstabilized carbanion, an organometallic nucleophile such as an organolithium or a Grignard reagent could be prepared from the corresponding bromide.



The bromide could be prepared from 3-phenyl-1-propanol (\$59/kg). The unsaturated aldehyde **O** can be made by oxidation (CrO₃.py) of 4-penten-1-ol (\$41.80/10 g). A cheaper way is to make ethyl 4-pentenoate from ethyl acetate (\$15/ga) and allyl bromide (\$19/100 g) and reduce it to the aldehyde **O** with DIBAH (\$19/ 0.1 mo).



Thus a synthesis of  $\mathbf{M}$  based on this retrosynthetic analysis would start with ethyl acetate, allyl bromide, and 3-phenyl-1-propanol.



#### 300 PLANNING ORGANIC SYNTHESES

Now we go back to A and consider disconnection at a different bond. Suppose we recognize that alcohol A could easily come from reduction of ketone K.



Now considering the polarities possible, a great number of disconnections can be envisioned. Choosing bond **b** means that polarity (with respect to the carbonyl group) would be



and thus an enolate reacting with a carbon electrophile would be appropriate. A valid retrosynthetic step would be



Because the tosylate is primary, substitution should be the major pathway (although in this case elimination could be problematic because of conjugation with the phenyl ring). We note, however, that the enolate needed is the kinetic enolate of 5-hexen-2-one. This poses a regiochemical control problem which can be solved by making the N,N-dimethylhydrazone of the ketone. The ketone 5-hexen-2-one is available (\$46.20/25 g) or can be made by allylation of acetone.



Thus a synthesis based on this retrosynthetic analysis starts with  $\beta$ -phenylethanol (\$35.60/kg), acetone, and allyl bromide. This route is comparable to the first in both number of steps and cost. It differs in that regiochemical control of enolate formation is a crucial feature. Several other syntheses of **K** can be devised by other disconnections suggested by the natural polarities engendered by the ketone group.

#### CARBON SKELETON SYNTHESIS 301



Next consider compound  $\mathbf{R}$ . When the relevant facts are considered, we see that  $\mathbf{R}$  is merely an olefin with a saturated ring present. Because of the fivemembered ring and because the target has 12 carbon atoms, it is unlikely that compounds with the carbon skeleton of  $\mathbf{R}$  will be available commercially; hence carbon–carbon bond-forming reactions will be needed to assemble the carbon skeleton.



Moreover the carbon–carbon double bond is a natural starting point for bond disconnection. A logical retrosynthetic step would be disconnection to a ketone because a Wittig reaction could be used to convert the ketone to the ethylidene product. (Note that dehydration of an alcohol to the olefin is not a viable synthetic step because dehydration would lead to a mixture of trisubstituted olefins.)



Once the ketone is recognized as a useful intermediate, normal polarities can be used to disconnect it retrosynthetically. For example, a good disconnection could be as shown, where Michael addition to ethyl vinyl ketone by a cyclopentyl anion would give the needed ketone.



#### 302 PLANNING ORGANIC SYNTHESES

The cyclopentyl nucleophile, which should be an organocuprate to ensure Michael addition, could be produced from cyclopentyl bromide. The synthetic sequence consistent with the retrosynthetic analysis turns out to be a rather simple synthesis of what at first sight is a more difficult molecule.



Now there are a variety of other ways to disconnect  $\mathbf{R}$  in the retrosynthetic analysis. As long as each synthetic step is valid and the target can be produced by the proposed synthetic route, then it is a correct solution. There can be many correct synthetic solutions for a given target and the "best" one may depend on factors other than those related strictly to the synthetic viability. Availability of starting materials, disposal of reaction by-products, number of steps, reagent sensitivity, expected yields, number of purifications, and the stereochemistry (among others) all contribute to the evaluation of a synthetic route.

#### UMPOLUNG SYNTHONS

Because of the polarities associated with carbonyl groups, some difunctional compounds are much easier to produce than others. For example, 1,3-dicarbonyl compounds and 1,5-dicarbonyl compounds are easy to produce using standard retrosynthetic steps with normal polarities induced by the carbonyl group.



In contrast, 1,2-dicarbonyl compounds or 1,4-dicarbonyl compounds are more difficult to disconnect by valid retrosynthetic steps. Consider a 1,2-diketone. Disconnection of the bond between the carbonyl groups requires that one of the carbonyl groups has the normal electrophilic character, but the other carbonyl carbon must have nucleophilic character (an acyl anion or its equivalent), which is not the normal polarity of a carbonyl group.

#### UMPOLUNG SYNTHONS **303**



In the same way, disconnection of a 1,4-diketone requires either an acyl anion equivalent reacting with a normal  $\beta$ -carbonyl electrophile or a normal  $\alpha$ -carbonyl nucleophile reacting with an abnormal  $\alpha$ -carbonyl electrophile. These abnormal or reversed-polarity reagents are said to have umpolung reactivity.



There is consequently a need for synthetic equivalents (synthons) of these reversed-polarity (umpolung) reagents. The development of reagents with umpolung reactivity has been an important addition to modern synthetic methodology. Acyl anion equivalents, among the most common umpolung synthons, can be produced by many strategies. For instance, nitroalkanes can be used as nucleophiles and the nitro function can be cleaved to the carbonyl group. Thus nitronates can be thought of as acyl anion equivalents.



Likewise 1,3-dithianes can be deprotonated by alkyl lithium bases and the resulting anions are strong nucleophiles. The dithiane group can be hydrolyzed back to the carbonyl group. Thus the dithiane serves as a synthon for the acyl anion.



#### 304 PLANNING ORGANIC SYNTHESES

Cyanohydrin derivatives have also been widely used as acyl anion synthons. They are prepared from carbonyl compounds by addition of hydrogen cyanide. A very useful variant is to use trimethylsilyl cyanide with an aldehyde to produce a trimethylsilyloxy cyanide. The cyano group acidifies the  $\alpha$  position (p $K_a \approx 25$ ) and the  $\alpha$  proton can be removed by a strong base. Alkylation of the anion and unmasking of the hydroxy group cause elimination of cyanide and re-formation of the carbonyl group.



These are only three of many ways that have been reported for the formation of acyl anion equivalents, which are among the most common umpolung synthons to be found in the literature. All are prepared by a similar strategy in that they contain functional groups which can sustain a negative charge on an adjacent carbon *and* can be converted back to a carbonyl group.

Another common umpolung synthon is a homoenolate. Normally the  $\beta$  position of a carbonyl compound is an electrophilic center (by Michael addition to an  $\alpha,\beta$ -unsaturated carbonyl derivative). To make it a nucleophilic center, an organometallic is needed since it is unactivated and nonconjugated. A common way to do this is to use a  $\beta$ -bromo acetal.



The bromine substituent can be metallated to give a carbanion equivalent  $\beta$  to the acetal group. Now since the acetal is easily hydrolyzed to the ketone, it is

a synthon for a  $\beta$ -carbonyl anion—an umpolung reagent. So it is important to recognize normal and reversed polarities when doing retrosynthetic analysis of a target in order to use umpolung synthons when they are needed.

For the lactone target **L** shown below, cleavage of the lactone ring gives a  $\gamma$ -hydroxy acid. This can be disconnected at any one of the three intervening bonds between the hydroxyl group and the carbonyl group (**a**, **b**, **c**).



If each of these is considered independently, it is easily seen that none of the disconnections has normal carbonyl polarities. (The same conclusion could be reached by merely noting that the hydroxy group and the carboxyl group have a 1,4 relationship. It was seen above that normal carbonyl-based polarities are not suited to the formation of 1,4 difunctional systems.) Thus one either has to use an umpolung synthon or go to functional groups other than carbonyl groups to guide the reactivity. For example, one could use an epoxide electrophile rather than a carbonyl electrophile for the bond-forming reaction.



#### ACETYLIDE NUCLEOPHILES

While the carbonyl group is a very common starting point for bond disconnections in retrosynthetic analysis, an olefinic or acetylenic unit is also a useful reference point in many instances. This is because a terminal acetylene can be used as an

#### **306** PLANNING ORGANIC SYNTHESES

effective nucleophile to install the triple bond into molecules and it can be reduced stereospecifically to either the cis or trans olefin. Thus for the cis-olefin target  $\mathbf{T}$ , the following retrosynthetic analysis leads to an efficient synthetic pathway which uses a nucleophilic displacement by an acetylide anion as a key carbon–carbon bond-forming step.



The synthetic sequence would be



#### RING CONSTRUCTION

The use of carbonyl groups to set the polarity of bond disconnections in retrosynthetic analysis is useful for the construction of rings as well. If a carbon electrophile and a carbon nucleophile are connected by a carbon chain, they can react with each other to form a carbon–carbon bond. This is an absolutely normal type of carbon–carbon bond-forming process, but the fact that the carbon nucleophile and carbon electrophile are connected by a chain means that the new carbon–carbon bond closes up the ends of the chain, forming a ring.

For example, the Claisen reaction is a reaction of an ester enolate with an ester to produce a  $\beta$  ketoester. We learned this reaction earlier.



If both ester groups are in the same molecule and are connected by a chain, then a Claisen-type reaction between the  $\alpha$  position of one ester and the carbonyl

group of the other gives a new carbon–carbon bond and closes up the ring. (This reaction is actually called the Dieckmann condensation, but it is nothing more than an intramolecular Claisen reaction.)



Ring-forming reactions are very important in retrosynthetic analysis because many interesting targets are cyclic compounds and often rings must be installed rather than being present in the starting materials. From a retrosynthetic point of view, there is really no difference between ring-forming reactions and other carbon–carbon bond-forming reactions. One looks for the same polarities and functional group features as in acyclic systems.

The only thing that is different is that some rings are more easily formed than others. The rule of thumb is that rings of three, five, and six members are routinely formed, while rings of four or more than six members are formed with greater difficulty. For example, reaction of diethyl malonate with 1,2-dibromoethane and two equivalents of base gives diethyl cyclopropane-1,1-dicarboxylate in high yield. Ring formation occurs by a double-displacement sequence.



Likewise reactions with 1,3-dibromopropane, 1,4-dibromobutane, or 1,5dibromopentane give the corresponding cyclobutyl-, cyclopentyl-, and cyclohexyl-1,1-dicarboxylates.



The yields of these reactions are not the same, however, and reactions which produce three-, five-, and six-membered rings are generally more effective. Use of 1,6-dibromohexane fails to give the cycloheptyl product.

Ring closure requires that a reactive center at one end of the chain encounters a reactive center at the other end *of the same chain* in the bond-forming process. The alternative is for the reactive center on one chain to react with a

#### 308 PLANNING ORGANIC SYNTHESES

reactive center of a different chain. The first case produces a ring, the second case a polymer. An analogy is a chain with complementary hooks at each end representing electrophilic and nucleophilic carbons at the ends of the chain. If two hooks on the same chain link up, a ring is formed, whereas if hooks from different chains link up, a larger molecule is formed with hooks remaining on each end. These can link up further to form progressively larger molecules.



For short chains which would give three- to six-membered rings upon ring closure, there is a higher probability that one end of the chain will encounter the other end of the same chain and react *intra*molecularly before it will encounter the end of another chain and react *inter*molecularly. Thus ring closure is normally favored over oligomerization for smaller rings of three to six members. On the other hand, as the chains become longer, it becomes less likely that the end of a chain will encounter the other end of the same chain before it encounters and reacts with the end of another chain. The break point is between six-membered rings, which are formed readily, and seven-membered rings, which are not easily formed. While this reasoning is a great simplification, it suffices to provide a good working model to predict the success for ring-forming reactions.

We can use this model in retrosynthetic analysis quite successfully. Suppose one were asked to produce cyclohexanone C from acyclic starting materials.



In this monofunctional compound, the ketone could serve as an electrophilic center in a cyclization step. Disconnection at the indicated bond leads to the polarity shown; however, it is immediately obvious that the carbon nucleophile occurs at an unactivated position, and there is no good way to produce it there without a control element at that position.



However, use of an ester group could activate this position toward anion formation and thus we could write instead



Now all is well in terms of polarity and we recognize this as a Dieckmann reaction followed by hydrolysis and decarboxylation of the  $\beta$ -ketoester product. Proceeding backward we write



Alkylation of diethyl suberate with benzyl iodide would produce the  $\alpha$ -benzylated product which would cyclize in the presence of base. This particular target does not present a regiochemical difficulty. Base could pull off either  $\alpha$  proton and two different enolates would be produced; however, both enolates cyclize to give the same product after decarboxylation.



If molecular symmetry were included in the cyclization precursor, then we would not have to worry about regiochemistry in the ring closure. Noting that the product of ring formation is a  $\beta$  ketoester, which itself is a good carbon nucleophile, an alternate retrosynthesis (which actually is much better) is the following in which the benzyl group is added after the ring is formed (in the forward synthesis):





#### **ROBINSON ANNULATION**

Another example of a very common ring-forming sequence is the Robinson annulation. This sequence allows a six-membered ring to be appended to an existing carbonyl group.



The strategy of the sequence is a Michael addition to an  $\alpha,\beta$ -unsaturated ketone followed by an intramolecular aldol reaction. Treatment of a ketone enolate with a Michael acceptor gives a diketone intermediate which is poised to produce a six-membered ring if an enolate is produced and it intramolecularly adds to the carbonyl group.



This process nicely accounts for formation of product, but if we consider intermediate **I**, we see that there are several different  $\alpha$  protons that could be removed by base, H_a, H_b, and H_c. Furthermore the acidities of these various  $\alpha$ protons should be comparable so all should be removed to similar extents under the reaction conditions. If we sequentially remove each proton and write the product from an intramolecular carbonyl addition, the following products could be produced. The fact is that only **P** is produced to any extent. This is due to the preference of six-membered ring formation over the formation of the more strained four-membered ring product or the more strained bridged product. Thus the enolate formed by the removal of  $H_a$  closes faster than the enolates formed by removal of  $H_b$  or  $H_c$ .



Furthermore, since the aldol reaction is reversible, if any of these higher energy products were formed, they could open back up under the reaction conditions. The exclusive formation of  $\mathbf{P}$  is an example of kinetic as well as thermodynamic control as the more stable product is formed fastest.

For the purposes of retrosynthetic analysis, a six-membered ring in a target can be related to a Robinson annulation of an existing ketone with an  $\alpha,\beta$ -unsaturated ketone. Normally  $\alpha,\beta$ -unsaturated methyl ketones are used to facilitate the ring closure, but this is not an absolute requirement. Thus the target steroid **S** could potentially be constructed by a series of Robinson annulations as shown. The last retrosynthetic step (the first synthetic step) could be problematic as a mixture of regioisomers would be formed.



Furthermore the bicyclic starting material could also be constructed by a Robinson annulation on a cyclopentanedione. In this case the final functionality must be achieved by selective reductions of the olefin and ketone functions at appropriate stages in the synthesis.
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A great many complex multicyclic targets have been synthesized by the Robinson annulation attesting to its generality and versatility.

# DIELS-ALDER REACTION

Another approach for the construction of rings is to use reactions which start with two acyclic compounds and produce cyclic products. There are many of these processes, but the most used and most useful is the Diels–Alder reaction. This is a reaction between a diene and an olefin to give a new six-membered ring. It is also termed a 4 + 2 cycloaddition because one partner (the diene) containing four  $\pi$  electrons adds to a two-electron fragment (the olefin) containing two  $\pi$  electrons to yield a ring.



The Diels–Alder reaction is one type of a much larger class of ring-forming reactions termed *cycloaddition reactions*. In most instances, new carbon–carbon bonds are formed during cycloaddition processes. Such reactions are neither polar reactions (although the reactants might be polar molecules) nor free-radical reactions. Rather cycloadditions are usually concerted reactions in which all bond making and bond breaking are taking place at the same time. Such reactions have been described as having a "no-mechanism mechanism." Curved arrows can be used for keeping track of electrons, but they are by no means of mechanistic significance. Alternatively dotted lines are sometimes used to describe the transition state, implying that all bonds are being made and broken simultaneously. No matter which mechanistic formalism is used to describe the process, the product contains a six-membered ring which has a double bond.



Concerted cycloaddition reactions result from the interaction between  $\pi$  systems in two different molecules. The  $\pi$  system in one molecule reacts with the  $\pi$  system in a second molecule to produce new bonds. Now since both the diene

and the olefin are closed-shell, ground-state molecules, the first question that comes to mind is "How do such systems react?"

The formation of new bonds in the Diels–Alder reaction requires that the  $\pi$  electrons in the individual diene and olefin  $\pi$  systems become reorganized and shared in the new bonding pattern of the cyclic product. It follows that for this bonding change to occur, the two  $\pi$  systems must overlap so that electrons can move into new orbitals. The most straightforward way the needed orbital overlap can occur is for one  $\pi$  system to function as an electron donor and the other  $\pi$  system to function as an electron acceptor. Therefore the bonding changes in the Diels–Alder reaction result from a donor–acceptor interaction between the diene and olefin  $\pi$  systems.

It is well known that  $\pi$  electrons are less tightly bound than  $\sigma$  electrons, and consequently they can readily be donated to a variety of electrophiles (e.g., bromine, protons, mercuric ion), so it is easy to imagine a  $\pi$  system acting as an electron donor. In contrast,  $\pi$  systems, being electron rich, are not often thought of as electron acceptors (which would make them even more electron rich).

For a  $\pi$  system to function as an electron acceptor, it must have unfilled orbitals available to accept electrons. In the case of olefins or dienes those are  $\pi^*$  antibonding molecular orbitals. Thus interaction of the HOMO of one  $\pi$  system with the LUMO of a second  $\pi$  system produces a donor-acceptor pair (HOMO donating to LUMO) enabling electrons to be transferred from one  $\pi$  system to another with resulting bond formation.

In picture form this can be drawn as



# **HOMO-LUMO INTERACTIONS**

One requirement for a successful HOMO–LUMO interaction is that the symmetry of the HOMO must match the symmetry of the LUMO (either both symmetric or both antisymmetric). If so, then the interaction is symmetry "allowed" and will lead to productive cycloaddition. If the symmetries do not match, then the HOMO–LUMO overlap is symmetry "forbidden" and cycloaddition will not proceed. Molecular orbitals can be classified by their phase symmetry with respect to a plane normal to the  $\pi$  system. The symmetry is related to the number of nodal planes which occur in each individual molecular orbital. For the olefin component, the  $\pi$  orbital (HOMO) is symmetric with respect to this plane and the  $\pi^*$  orbital

# 314 PLANNING ORGANIC SYNTHESES

(LUMO) is antisymmetric with respect to this plane. For the diene component, the HOMO is antisymmetric and the LUMO is symmetric. Based on these symmetries, it is seen that the HOMO–LUMO interaction between butadiene and ethylene is symmetry allowed and thus can proceed productively to product.



It turns out that the orbitals for any diene and any olefin have the same symmetry properties so that all Diels-Alder reactions are symmetry allowed.

While symmetry requirements dictate whether a cycloaddition can occur, they do not determine the strength of the HOMO–LUMO interaction. The strength of the donor–acceptor interaction and the rate of cycloaddition is inversely related to the difference in energy between the HOMO and LUMO which are interacting. If the HOMO–LUMO energy gap is small, the interaction is strong and the reaction is rapid, whereas if the HOMO–LUMO energy gap is large, the interaction is weak and the reaction is slow. The Diels–Alder reaction between butadiene and ethylene is very slow, meaning that the donor–acceptor interaction is very weak because the HOMO–LUMO energy gap is large.



It is also pertinent that there are two HOMO-LUMO interactions possible between butadiene and ethylene, one in which the HOMO is that of the diene, which acts as the electron donor, and one in which the HOMO is that of the olefin, which would be the electron donor. A "normal" Diels-Alder reaction is one in which the diene is electron rich and acts as the electron donor and the olefin (dienophile) is electron poor and acts as the electron acceptor. In such a case the diene HOMO and the dienophile LUMO are closer in energy, the donor acceptor interaction between them is strong, and the reaction takes place at a more rapid rate.



Examples of such partners are dienes substituted with alkyl groups or other electron-donating groups and dienophiles having carbonyl groups attached, which makes them electron deficient. The dienophile need only be a two-electron  $\pi$  system so olefin, acetylene, and azo  $\pi$  systems can all serve effectively as dienophiles so long as they have electron-deficient  $\pi$  bonds. Furthermore the dienophile may be symmetrically or unsymmetrically substituted with electron-withdrawing groups.



# 316 PLANNING ORGANIC SYNTHESES

# STEREOELECTRONIC FACTORS

The interaction between the diene HOMO and the dienophile LUMO takes place when the ends of the two  $\pi$  systems overlap to permit the transfer of electrons from the HOMO into the LUMO. This requirement of overlap imposes stereoelectronic constraints on the two reaction partners. First, the diene must be able to adopt an s-cis conformation so the ends of the diene can contact and overlap with the ends of the dienophile  $\pi$  system. For acyclic dienes, even though the s-trans conformer is favored, rotation about the central carbon–carbon bond is rapid and there will be a steady-state population of the required s-cis form present so that the cycloaddition can occur.



However, when the diene system is constrained to the s-cis conformation by a cyclic framework, the effective concentration of the s-cis diene is much higher than for acyclic dienes, which have the s-cis conformer as a minor component of the rotomeric equilibrium. Such conformationally constrained dienes react much more easily and are excellent Diels–Alder dienes. Examples are cyclopentadienes, 1,3-cyclohexadienes, and furans.



s-cis Diels-Alder dienes

Second, substituents on the dienophile (olefinic or azo) can adopt a position in the transition state either exo or endo to the diene system. It has been found that the endo transition state is favored significantly over the exo transition state. This preference has been attributed to secondary orbital interactions (attraction) between the diene and polar substituents on the dienophile.



This distinction is important because exo and endo transition states lead to different diastereomers. Control of diastereoselection is extremely important to the utility of the Diels-Alder reaction since mixtures of diastereomers are avoided and control of multiple stereogenic centers is achieved.



For example, reaction of (E,E)-2,4-hexadiene with methyl crotonate gives a single product in which the relative stereochemistry of four contiguous stereogenic centers is explicitly defined by the geometry of the starting materials and the endo transition.



A third consideration of the Diels-Alder reaction is the regiochemistry of the products. If either the diene or the dienophile is symmetric, then only a single regioisomer is possible. If both the diene and the dienophile are unsymmetric, however, two regioisomers are possible depending on the relative orientation of the substituents at the transition state. Usually one of these regioisomers is favored over the other.



318 PLANNING ORGANIC SYNTHESES



The control of regiochemistry has been rationalized on the basis of the orbital coefficients of the HOMOs and LUMOs, but in fact, it is not well understood. In most cases such cycloadditions are not regiospecific and isomeric mixtures are formed, although one regioisomer usually predominates. Qualitative estimation of the electron distributions in the diene and dienophile can often be used to predict the major product. For example, C-1 of siloxy diene (**A**) should be much more electron rich than C-4. In addition C-3 of acrylate (**B**) should be more electron deficient than C-2.



Thus the most favorable donor-acceptor interaction should occur between C-1 of the diene and C-3 of the dienophile. This interaction would favor 1,4 orientation of the substituents in the Diels-Alder product, as is observed.

In spite of the fact that the major product is often predictable, such systems are rarely regiospecific. Because mixtures of regioisomers which must be separated are the rule, either a symmetric diene or a symmetric dienophile is usually employed to avoid such regiochemical issues.

The Diels-Alder reaction can be used to create rings in many situations that would be difficult to accomplish by ring closing approaches. Consider product **P**. This product can be made by a Diels-Alder reaction between diene **D** and acrylate **A**.



Because the diene is acyclic, it can achieve the required s-cis conformation by rotation. The polarity is correct because the diene is electron rich and the

dienophile is electron poor. The stereochemistry is fine because the endo transition state gives the correct stereochemical relationship of the groups around the cyclohexyl ring. The stereochemistry can be seen more clearly by a drawing of the transition state.



# **1,3-DIPOLAR CYCLOADDITIONS**

The essential features of the Diels–Alder reaction are a four-electron  $\pi$  system and a two-electron  $\pi$  system which interact by a HOMO–LUMO interaction. The Diels–Alder reaction uses a conjugated diene as the four-electron  $\pi$  system and a  $\pi$  bond between two elements as the two-electron component. However, other four-electron  $\pi$  systems could potentially interact with olefins in a similar fashion to give cycloaddition products. For example, an allyl anion is a fourelectron  $\pi$  system whose orbital diagram is shown below. The symmetry of the allyl anion nonbonding HOMO matches that of the olefin LUMO (as does the olefin HOMO and the allyl anion LUMO); thus effective overlap is possible and cycloaddition is allowed. The HOMO–LUMO energy gap determines the rate of reaction, which happens to be relatively slow in this case.



Molecules isoelectronic with the allyl anion but which are neutral and have at least one resonance form with formal positive and negative changes in a 1,3 relationship are called 1,3 dipoles.

Azides 
$$R-N=N=N \ominus \longrightarrow R-N=N \ominus$$

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320 PLANNING ORGANIC SYNTHESES



All have an orbital diagram analogous to the allyl anion in which three interacting p orbitals give rise to three molecular orbitals containing a total of four  $\pi$ electrons. For example, a nitrone is seen to have a C–N  $\pi$  bond interacting with a filled orbital on the oxygen atom to define a new  $\pi$  system containing four  $\pi$ electrons.



Interaction of the HOMO of the 1,3 dipole with the LUMO of a simple  $\pi$  bond (called a dipolarophile in this process) leads to bond formation between the ends of the 1,3 dipole and the olefin, producing a new five-membered ring.



The process is a concerted  $4_{\pi} + 2_{\pi}$  cycloaddition and is related electronically to the Diels–Alder reaction. The formal charges are destroyed during the cyclization and a wide variety of heteroatom components are possible in the 1,3 dipole. Moreover other  $\pi$  bonds besides alkenes and alkynes can be used as dipolarophiles. As a result, 1,3 dipolar cycloadditions have been used to make a large number of heterocyclic compounds.

Since the 1,3 dipolar cycloaddition is concerted, the reaction is stereospecific and the geometry of the olefin is maintained in the cyclic product.



If a symmetric dipolarophile is used, then no regioisomers are possible. If, however, the dipolarophile is unsymmetric, then regioisomers are possible.



As in the case of the Diels–Alder reaction, the regioselectivity can be understood in terms of the electron distribution in the 1,3 dipole and the dipolarophile. For example, a nitrile oxide should have a relatively electron deficient carbon and a relatively electron rich oxygen. Reaction with propene, which has greatest electron density at C-2 because of the inductive effect of the methyl group, gives the regioisomer C. Matching the polarity of the dipole and the dipolarophile predicts this product. Conversely reaction with methyl acrylate, which because of conjugation has electron deficiency at C-3, gives regioisomer D as the major product.



# 322 PLANNING ORGANIC SYNTHESES

Polarity matching to predict the major product of 1,3 dipolar cycloadditions is qualitative only and frequently fails to predict the major product correctly. This is because each 1,3 dipole tends to exhibit a characteristic regioselectivity toward particular dipolarophiles that may be modified by steric and/or strain effects. In fact there is still some uncertainty as to just what factors do influence the regioselectivity in these systems.

Nevertheless 1,3 dipolar cycloadditions are an important method for the synthesis of a wide variety of heterocyclic compounds. Furthermore they illustrate the generality of 4 + 2 cycloaddition reactions as a means to prepare cyclic products efficiently from acyclic precursors.

To use 1,3 dipolar cycloadditions in a retrosynthetic sense, it is necessary to know what 1,3 dipoles are available. The list on pages 319–320 is representative of the more common and useful examples, although many others have been reported. Azides, diazo compounds, and nitrones are normally isolable compounds which can be added to a solution of an olefin. Other 1,3 dipolar species such as nitrile oxides and azomethine ylides are not stable molecules; they must be generated in the reaction mixture in the presence of the olefin. As might be expected, many different ways to generate 1,3 dipoles have been developed.



Nitrile oxides are often generated by the dehydration of nitro compounds by reagents such as phenyl isocyanate. Azomethine ylides can be generated by the pyrolysis of aziridines or by the prototopic isomerization of imines upon heating.

The next step is to identify the five-membered ring which could be assembled by a 1,3 dipolar cycloaddition and then identify the  $\pi$  system and 1,3 dipole needed to give the proper array of heteroatoms. Thus if the pyrrolidine **H** is needed, it is clear the alcohol could be made by reducing the ester function of **E**. Also important is the issue of the all-cis stereochemistry. One way to ensure the all-cis stereochemistry is to do a catalytic hydrogenation of the olefin **O**. The delivery of hydrogen would come from the less hindered face of O and would give the all-cis product. The needed olefin O could be made by a 1,3 dipolar addition between an azomethine ylide and diphenyl acetylene.



The only concern is the cis stereochemistry of the cycloadduct **O**. If the planar azomethine ylide adopts the least sterically hindered "W" geometry, then the cis isomer will be produced as a pair of enantiomers. The use of *cis*-stilbene as the dipolarophile to obtain the all-cis geometry in one step would require that only the endo transition state produces product. Although endo transitions are favored in 1,3 dipolar cycloadditions, mixtures of diastereomers from the exo and endo transition states are usually formed. Catalytic hydrogenation has a higher facial selectivity and is much more likely to give a single diastereomer.



These are but a few examples of how retrosynthetic analysis can be used to develop one or more synthetic routes to a target. Developing synthetic strategies is one of the most creative activities that organic chemists perform. It requires that many different inputs and conditions be cohesively merged into a single thematic development that contains elements of texture and beauty, proportion and balance, and risk and reward. The process is every bit as creative as painting, sculpting, or writing the great American novel!

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# PROBLEMS

**10.1.** The following group of syntheses are fundamentally functional group manipulations. Give a retrosynthetic pathway from the target on the left to the starting material on the right. Then provide a synthetic pathway with proper reagents and conditions for each step.



PROBLEMS 325



**10.2.** Show a retrosynthetic pathway to the following targets which involves the formation of the bond or bonds indicated by an arrow:



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OCH₃

³²⁶ 

PROBLEMS 327



**10.3.** Provide retrosynthetic pathways for the following targets from "simple, readily available" starting materials:



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PLANNING ORGANIC SYNTHESES



³²⁸ 



- **10.4.** Using curved-arrow notation, show how the following 1,3 dipolar compounds are formed from the indicated starting materials:
  - (a)  $\underset{l_1}{\overset{0}{\text{Ph}-\text{CH}_2-N}} \stackrel{0}{\overset{1}{\text{Ph}}} + 2 \text{Ph} \text{N}=\text{C}=\text{O} \longrightarrow \text{Ph}-\text{C}\equiv N-\text{O}$



**10.5.** Give the products of the following reactions. Where possible indicate the regioselectivity and stereoselectivity of the reaction.



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PLANNING ORGANIC SYNTHESES



**10.6.** Provide a synthesis of the following compounds from the indicated starting materials. Give reagents and conditions needed to carry out each step.



PROBLEMS 331



# 11

# STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS

Structure Determination	332
Chromatographic Purification	333
Instrumental Methods	335
Nuclear Magnetic Resonance	336
Chemical Shift	338
Spin-Spin Coupling	344
Descriptions of Spin Systems	350
Second-Order Splitting	354
Structure Identification by ¹ H NMR	355
Carbon-13 NMR	360
Infrared Spectroscopy	366
IR Stretching Frequencies	367
Use of IR Spectroscopy for Structure Determination	371
Mass Spectrometry	377
Fragmentation Processes	384
Bibliography	388
Problems	388

# STRUCTURE DETERMINATION

Because of the wide variety of reactions available to synthetic chemists, it is possible to devise synthetic strategies for just about any target that we wish.

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Nevertheless the planning and execution of any synthesis must be verified by showing that the product of each step is in fact the predicted compound and that the target compound was actually obtained. Thus a critical part of any synthesis involves determining and proving the structures of synthetic intermediates and final products. While the majority of the time careful planning will result in the formation of the expected product, there are always enough exceptions to make structure proof an imperative step.

In earlier times, proof of structure was based largely on wet methods. The first step was to rigorously purify the compound by crystallization, distillation, sublimation, and so on. The functional groups present in the material were established by classification tests. The elemental analysis gave the molecular formula, and from a knowledge of the starting materials, a tentative structure could be written. Confirmation of the structure was obtained by either degradation to known compounds or an alternative synthesis of the compound from known starting materials. Thus, while the presence of functional groups could be determined rather straightforwardly, the connectivity of atoms and groups in a molecule were much more difficult to establish.

Today structure proof involves the same components—purification, functional group identification, and establishment of atom and group connectivity; however, the ways in which these are accomplished are more efficient, sensitive, and reliable. They are also much faster. The ability to run reactions, purify products, and determine structures on milligram scales, often in a matter of hours, has caused a huge increase in the rate at which structural information can be obtained. This has resulted in an exponential growth of chemical knowledge and is directly responsible for the explosion of information being continually published in the chemical literature.

# CHROMATOGRAPHIC PURIFICATION

The first step in the identification of any compound is to obtain that material in pure form. The most common way to achieve this goal today is to use chromatography. While a discussion of the many separation and purification techniques which utilize some form of chromatography are outside the focus of this book, all rely in one way or another on the interaction of molecules with a surface. Such interactions depend much more on the chemical properties of a molecule (functional groups, polarity, unsaturation, etc.) than on physical properties of the bulk substance (boiling point, vapor pressure, etc.). Furthermore the interactions of a compound with a surface allow it to be resolved (separated) from other molecules by placing it in a flowing system (mobile phase). When the molecule is not adsorbed to the surface, it moves over the surface at the same velocity as the mobile phase (Figure 11.1).

When it is adsorbed to the surface, it does not advance with the mobile phase. Since adsorption is an equilibrium process, those compounds which are only weakly adsorbed (M in Figure 11.1) to the surface spend a greater portion of

#### 334 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS



Figure 11.1 Interaction of molecules with a surface and a flowing mobile phase.

time in the mobile phase and thus move over the surface faster than compounds which are more strongly adsorbed ( $M^*$  in the figure) and thus spend more time immobilized on the surface. Because different compounds are adsorbed differently on the surface, each can travel at a different rate over the surface. By collecting the effluent from the surface as a series of fractions, individual compounds can be separated cleanly from other components in the original mixture because each component exits the surface at a different time.

Many different mobile phases have been utilized to provide the forward velocity for nonadsorbed molecules. If the mobile phase is a gas, then the technique used is gas chromatography (GC). In GC, the surface to which the molecules adsorb can be a wide variety of materials which are often prepared by coating an inert surface with a polymer whose properties are related to its structure. In this way the surface properties and hence adsorption of the solid surface can be varied to give the best chromatographic resolution.

If a liquid is used as the mobile phase, the technique used is liquid chromatography (LC). The solid adsorbent is constrained in a tube or column through which the liquid mobile phase flows. Any number of solvents, buffer solutions, or supercritical fluids can be used as liquid mobile phases. High-pressure liquid chromatography (HPLC) is used if pressure is needed to force the liquid phase through the tube. If the liquid phase moves over a thin adsorbent surface propelled by capillary action, the technique used is thin-layer chromatography (TLC). In general, two types of surfaces are used as the solid phase.

In "normal"-phase LC systems, the solid phase is a polar solid such as silica gel (most common) or alumina and the liquid is generally an organic solvent of low polarity. In such a case, polar compounds bind more strongly to the polar silica gel surface and thus travel more slowly along the surface, whereas

nonpolar components have a lower affinity for the polar surface and a greater affinity for the nonpolar eluting solvent. They consequently elute from the column more rapidly. In *reversed-phase* systems, the surface of silica gel is modified to produce a nonpolar hydrocarbon-derivatized surface, and the mobile phase often is a polar, aqueous solvent mixture. In this case polar compounds have a low affinity for the nonpolar surface; they remain dissolved in the polar mobile phase and elute more rapidly. Nonpolar components have a higher affinity for the nonpolar surface than the polar mobile phase and elute more slowly. Using various chromatographic techniques, it is possible to separate most mixtures into the individual components efficiently and very rapidly.

In addition to chromatographic techniques, traditional purification methods such as recrystallization, distillation, or sublimation are also employed. Such methods often require much more material than chromatographic techniques.

# **INSTRUMENTAL METHODS**

When the reaction product(s) is obtained in pure form, modern instrumental methods of structure determination, rather than traditional wet methods, provide the fastest way to determine the functionality and connectivity present. Today's chemist has a large number of tools available with which to probe the structure of molecules, but for determining the structures of organic molecules the "big three" are nuclear magnetic resonance (NMR) spectroscopy, infrared (IR) spectroscopy, and mass spectrometry (MS). The frequency of use of these techniques generally falls in the same order (NMR > IR > MS).

The first two of these methods (NMR and IR) are spectroscopic techniques in which the molecule is interrogated with electromagnetic radiation in a particular range of frequencies (NMR uses radio frequencies and IR spectroscopy uses infrared radiation). Only certain frequencies will be absorbed by a particular compound, and those frequencies that are absorbed can be used to infer structural details about the compound.

Mass spectrometry is not a spectroscopic technique because absorption of electromagnetic energy is not involved in any way. In MS the molecule is fragmented into ions and these charged pieces are separated on the basis of their mass-to-charge ratio. Since the usual charge is +1, the masses of the pieces are determined. Knowledge of the masses of the pieces allows the structure of the compound to be reconstructed.

Another distinction between these techniques is the structural information they are capable of revealing. Both NMR spectroscopy and MS establish connectivity between atoms and groups in a molecule (albeit slightly differently). In addition the functional groups present are suggested. Infrared spectroscopy does not generally establish connectivity but is unmatched for identifying functional groups present in the molecule. By combining these three major methods, the functional groups, the molecular weight, and the connectivity of the atoms and groups can be established rapidly and efficiently. It is normally a trivial step then to write the structure of the molecule.

# 336 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS

# NUCLEAR MAGNETIC RESONANCE

The speed with which NMR spectroscopy has been incorporated into scientific inquiry is truly amazing. The first commercial spectrometers became available in the 1950s. By the middle 1980s whole bodies could be placed in the probes of NMR spectrometers (magnetic resonance imaging) and the structures of body parts could be determined in exquisite detail. Today structures of proteins and other macromolecules in solution or in the solid state are determined routinely. What was unthinkable in the 1960s is routinely practiced today even by undergraduates! The power of the method and the structural detail it provides have no doubt fueled its rapid development.

Nuclear magnetic resonance spectroscopy is possible due to the absorption of energy at particular frequencies by atomic nuclei when they are placed in a magnetic field. Most atomic nuclei are characterized by a property termed *spin*, and this gives rise to a magnetic moment associated with that nucleus. The magnitude of the magnetic moment of the nucleus, which is also quantized by the spin quantum number, is characteristic of that nucleus. Nuclei such as ¹²C, ¹⁶O, and ³²S have nuclear magnetic moments of zero. Other nuclei such as ¹H, ¹¹B, ¹³C, ¹⁵N, ¹⁷O, ¹⁷O, ¹⁹F, and ³¹P have finite magnetic moments and spin quantum numbers of  $I = \frac{1}{2}$  and are most useful in NMR measurements. Still other nuclei such as ²D and ¹⁴N have finite magnetic moments but spin numbers  $I > \frac{1}{2}$  and are much more difficult to deal with, although today's NMR instruments handle these elements routinely as well.

Fortunately for organic chemists, hydrogen and carbon are the most common nuclei found in organic compounds, and the ability to probe these nuclei by NMR is invaluable for organic structure determination. Since proton magnetic resonance (PMR) is the most common type, the behavior of ¹H nuclei in magnetic fields will serve as a model for other nuclei which have spin quantum numbers  $I = \frac{1}{2}$  and thus behave similarly (¹³C, ¹⁹F, etc.).

The proton has a nuclear magnetic moment (denoted as a vector quantity) which under normal circumstances can adopt any spatial orientation. Since this magnetic moment is a nuclear property, each hydrogen in a molecule has an identical nuclear magnetic moment. When placed in a strong magnetic field, the magnetic moment of the nucleus interacts with the magnetic field. The strength of the interaction depends on the strength of the applied field ( $H_0$ ) and the nuclear magnetic moment characterized by the magnetogyric ratio  $\gamma$  (the same for all hydrogens but different for other nuclei).

In a strong magnetic field the nuclear magnetic moment is no longer free to adopt just any orientation. Instead the spin quantum number of  $I = \frac{1}{2}$  for the hydrogen nucleus results in only two allowed orientations (2I + 1) of the nuclear moment relative to the direction of the applied field—either aligned with  $H_0$ (lower energy) or opposed to it (higher energy.) The difference in energy ( $\Delta E$ ) between the two states is given by  $\Delta E = \gamma h H_0/2\pi$  and is dependent on the cross product of the strength of the applied field  $H_0$  and the magnetic moment of the hydrogen ( $\gamma$ ). Since  $\gamma$  is the same for all hydrogen nuclei, the energy difference between the two allowed orientations is proportional only to the strength of the applied field (Figure 11.2).

If the magnetic field  $H_0$  is fixed and held very constant, then the energy gap between the two spin states of the hydrogen nuclei will remain constant. Irradiation of the system with radiation of the appropriate frequency ( $\Delta E = h\nu$ ) will cause the energy to be adsorbed and the spin of the nucleus will flip from the low-energy state (aligned) to the higher energy state (opposed). It is this absorption of energy which is used to probe the structural features of the molecule (Figure 11.3).

Now since the magnetic moment of a nucleus ( $\gamma$ ) is an atomic property, for a given magnetic field  $H_0$ , all hydrogens should absorb energy at the same frequency. However, examination of a molecule such as 1,2,2-trichloropropane (see Fig. 11.4) reveals that the two different types of hydrogens (H₁ and H₃) absorb at two different frequencies ( $\nu_1$  and  $\nu_3$ ) (Figure 11.4).



Figure 11.2 Energy of the spin states of a hydrogen atom in a magnetic field.



Figure 11.3 Energy absorption by a hydrogen atom in a magnetic field.



Figure 11.4 Different absorption frequencies for the protons of 1,2,2-trichloropropane.

# 338 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS

Since the applied field  $H_0$  is constant and all hydrogen nuclei have the same magnetic moment  $\gamma$ , the fact that H₁ and H₃ absorb at two different frequencies requires that the magnetic field that is actually experienced by each set of nuclei  $(H_{\text{eff}})$  is different. Stated differently, even though a constant magnetic field  $H_0$ is applied to the sample, each type of hydrogen H₁ and H₃ experiences a unique magnetic field  $H_{\text{eff}}$  (where  $H_{\text{eff}} \neq H_0$ ) and consequently absorbs energy at a unique frequency  $\nu_1$  and  $\nu_3$ . Thus the different types of protons are distinguished by different frequencies at which they absorb energy. Furthermore, integration of the absorption intensities of the two signals gives a 3 : 2 ratio which corresponds to the smallest whole-number ratio of each type of proton present. (The integrated area of the peak is given by the step height on the integration curve.) Thus ¹H NMR is able to distinguish different types of protons in a molecule and tell how many there are (at least by ratio).

# CHEMICAL SHIFT

The range of frequencies over which protons absorb in most organic molecules depends on the applied field. For example, for an applied field of 14,000 G, most protons will absorb over a range of 600 Hz beginning at the value of  $60 \times 10^6$  Hz (60 MHz), or from 59,999,400 to 60,000,000 Hz. At 23,486 G this range is 1000 Hz near the value of 100 MHz or from 99,999,000 Hz to 100,000,000 Hz. Thus the actual range of frequency of absorption depends on the magnetic field of the instrument. (This is exactly as expected since the energy gap between the spin states and hence the frequency of absorption are dependent on the applied field.) To compare absorption values from different instruments, a dimensionless scale must be devised that is independent of the magnetic field of the instrument. This is accomplished by using the absorption of a given set of protons is measured relative to the frequency of absorption of TMS. This absorption frequency difference  $\Delta \nu$  in hertz (cps) is expressed as  $\delta$ , the chemical shift of the protons in ppm, where

$$\delta = \frac{\Delta \nu \text{ (Hz)}}{\text{operating frequency of spectrometer (Hz)}} \times 10^6 \text{ ppm}$$

The chemical shift  $\delta$  is dimensionless and independent of the spectrometer. Since normal absorption ranges  $\Delta \nu$  are about 0–600 Hz for an operating frequency of  $60 \times 10^{-6}$  Hz, or 0–1000 Hz at  $100 \times 10^{6}$  Hz, and so on, chemical shifts range from 0 to 10 ppm for most protons.

In practice a small amount of TMS (<1%) is added to the NMR sample, the TMS signal is set at 0 ppm, and the protons of the sample are then measured in parts per million relative to TMS. The choice of TMS as a standard is useful because nearly all other protons absorb at frequencies lower than TMS. It is routine to present NMR spectra with low frequency on the left and high frequency on the right (Figure 11.5). Thus the TMS signal defines  $\delta = 0$  ppm on the right side



Figure 11.5 Typical NMR spectrum from 0 to 10 ppm.



Figure 11.6 Shielding of the nucleus by an electron cloud.

of the spectrum and other proton signals are found to the left or downfield from TMS from 0 to about 10 ppm. It is also normal to describe signals having larger chemical shifts as being downfield from protons with smaller chemical shifts. The left side of the spectrum is termed *low field* and the right side *high field*.

With a method available to measure differences in chemical shifts between protons, it is appropriate to ask why different protons experience different  $H_{\text{eff}}$ 's even though a single  $H_0$  is applied to the sample. The explanation lies in the fact that nuclei are surrounded by electron clouds (Figure 11.6). In the applied field  $H_0$ , electron pairs in bonds surrounding the hydrogens act to counter the applied field by induced fields ( $H_{\text{ind}}$ ). The result is that the nucleus is shielded from the applied field by its electron cloud. (Nuclei which are more shielded come at higher fields and have lower chemical shifts.)

Thus it is the electron density around the nucleus which shields the nucleus from the applied field. It follows that the greater the electron density around a proton, the larger will be the induced field  $H_{ind}$  and that proton will be more shielded. It will appear more upfield and will have a smaller chemical shift ( $\delta$  value). Conversely the lower the electron density around a proton, the less shielded it will be, the more downfield it will be, and it will have a larger  $\delta$  value (Figure 11.7).

Structural features which withdraw electrons from protons cause downfield shifts and larger  $\delta$  values, while structural features which increase electron density around protons cause upfield shifts and lower  $\delta$  values. For example, chemical

#### 340 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS



Figure 11.7 Changes in chemical shift due to electron density around the proton.

shifts for methyl chloride, dichloromethane, and chloroform are  $\delta = 3.0$ ,  $\delta = 5.5$ , and  $\delta = 7.1$ , respectively. The inductive effects of increasing numbers of chlorine atoms decrease the electron density about the hydrogens and result in increasing chemical shifts.

 $Cl \xrightarrow{H}_{H} \xrightarrow{Cl}_{\delta = 3.0} \xrightarrow{Cl}_{C} \xrightarrow{H}_{\delta = 5.5} \xrightarrow{Cl}_{\delta = 7.1}$ 

Likewise 1,2,2-trichloropropane discussed previously has the two-proton signal downfield from the three-proton signal. This is because the methylene protons are influenced by the inductive effects of three chlorine atoms, two vicinal and one geminal, while the methyl group is influenced by only two vicinal chlorine atoms. The electron density is higher at the methyl hydrogens, which are more shielded and occur at higher fields than the two protons of the methylene group.



Consideration of a series of compounds containing methyl groups illustrates clearly the influence of the electron density on chemical shift. As the electron-withdrawing ability of groups attached to the methyl group increase, progressive downfield shifts are evident and  $\delta$  values increase. Conversely TMS comes very far upfield because

silicon-carbon bonds are polarized toward carbon and result in very high electron density about the methyl hydrogens of TMS.



Although the influence of electron density on chemical shift is clear, it is not the only factor which determines the chemical shift, as seen from the following series of compounds:



Comparing the methyl groups, we find that typical saturated aliphatic methyl groups come at 0.9-1.1 ppm. However, attaching a methyl group to a double bond gives a change to  $1.8\delta$ . Attaching the methyl group to an aromatic ring moves it further downfield to  $2.4\delta$ . Attachment to a triple bond moves it back upfield to  $1.3\delta$ . Analogous but even larger changes in chemical shift are seen for protons directly attached to double bonds, aromatic rings, and triple bonds. Simple electron density shielding arguments cannot satisfactorily account for these large changes in chemical shift.

For example, the greater s character of  $sp^2$  orbitals and hence greater effective electronegativity of  $sp^2$ -hybridized carbon might account for the downfield shift of the protons of a methyl group when it is attached to an olefinic carbon rather than a saturated  $sp^3$  carbon; however, the  $sp^2$  carbons of aromatic rings should induce the same downfield shift. In fact, aromatic methyl groups are shifted significantly further downfield. By the same argument, attachment of a methyl group to the sp-hybridized carbon of an acetylene, which has even greater s character, should cause the chemical shift to move even further downfield. In fact, propargylic methyl groups are found at higher field than allylic methyl groups.

# 342 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS

It is clear that there are other factors at work which influence the chemical shifts of different types of protons.

Simple shielding of the hydrogen nucleus by its surrounding cloud of electrons is *isotropic* in that the induced magnetic field is the same for any orientation of the hydrogen relative to the magnetic field. This is due to the fact that the electron cloud around the hydrogen nucleus behaves as though it is spherical (or nearly so). Other types of electron clouds (double bonds, aromatic clouds, triple bonds) are not spherically symmetric. As a consequence, the induced fields for these types of bonds are not the same at different orientations of the functional group in the magnetic field. This *anisotropic shielding*, or *anisotropy*, leads to regions of shielding and deshielding around the functional group that are averages of the orientations possible.

Aromatic rings have among the strongest anisotropy of any group. Above and below the ring there is a strong shielding region ( $H_{ind}$  is in opposition to the applied field) while in the plane of the ring there is a strong deshielding region ( $H_{ind}$  is in the same direction as the applied field). This phenomenon is termed *ring current* and has been used as a criterion to establish whether a compound is aromatic (Figure 11.8). Consequently protons and groups attached to the ring are in the plane of the ring and thus are strongly deshielded and come at low fields relative to a comparable proton in a nonaromatic compound. Aromatic protons normally come at  $\delta > 7$  ppm and benzylic methyl groups come at  $\delta \approx 2.4$ , which are both significantly shifted downfield due to the anisotropy of the aromatic ring. (The shift of benzylic protons is less than the shift of aromatic protons because they are further from the aromatic ring than the protons directly attached to the ring.)



If protons could be positioned in the center of or above the aromatic ring, they would fall in the shielding region and should come at high field. For example, 18-annulene is an aromatic compound (4n + 2, n = 4). The protons on the outside of the ring lie in the deshielding region and have  $\delta = 9.3$  ppm while those on the inside of the ring fall in the shielding region and have  $\delta = -3.0$ . They come at higher field than TMS due the anisotropic shielding from the ring current. For the same reason, the central protons in *p*-cyclophanes come at higher fields because they are placed over the aromatic ring in the shielding region.

Double bonds contain one  $\sigma$  bond and one  $\pi$  bond, which results in anisotropic shielding, as shown in Figure 11.9. There is a conical shielding region normal to the molecular plane and a deshielding region in the molecular plane. This is







Figure 11.9 Anisotropic shielding by a double bond.



Figure 11.10 Anisotropic shielding by a triple bond.

true for all double-bonded functional groups such as olefins, carbonyl groups, and imines, and it explains why olefinic protons ( $\delta \approx 5$ ) and aldehyde protons ( $\delta = 9-10$ ) absorb at such low fields.

Acetylene (and nitriles), because of their cylindrical symmetry, have shielding regions along the triple-bond axis (Figure 11.10). Thus groups attached to the triple bond are constrained to the shielding region and are shifted upfield relative to similar vinyl protons. Thus acetylenic protons come at  $\delta = 2-3$  and propargylic methyl groups are upfield from allylic methyl groups.

The chemical shift of a given proton is thus determined by a combination of isotropic shielding by the electron cloud surrounding the proton and by

## 344 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS

anisotropic shielding due to the presence of nearby functional groups which are strongly anisotropic. These factors are usually sufficient to give unique chemical shifts for most protons in a molecule, and they can normally be distinguished using modern high-field NMR spectrometers (200–300 MHz). Furthermore the integration of these signals gives the numbers of the different types of protons.

# SPIN-SPIN COUPLING

The determination of the numbers of different kinds protons in a molecule is a very important use of NMR spectroscopy, but it does not establish the connectivity of the carbons bearing those protons and the connectivity is crucial in correct structure determination. However, NMR spectroscopy can also give insight into the connections between functional groups by the presence of spin-spin coupling in the NMR spectrum.

Consider the NMR spectrum of 1,1-dichloro-2,2-dibromoethane (Figure 11.11). Based on the different electronegativities of chlorine and bromine, the two protons in the molecule are nonequivalent and should thus give signals at different chemical shifts with the same integrated areas. The dichloromethyl proton should appear downfield relative to the dibromomethyl proton. The actual NMR spectrum indeed shows two different signals, one for  $H_a$  and one for  $H_b$ , but each absorption consists of two lines and is termed a doublet. The signal for each proton is thus "split" into two resonances. This splitting is due to the fact that each proton can sense the spin state of the neighboring proton and is called spin–spin splitting.

In the magnetic field of the NMR spectrometer, both  $H_a$  and  $H_b$  have distinct absorption frequencies based on the  $H_{eff}$  each proton experiences. This gives rise to individual signals for  $H_a$  and  $H_b$ . Focus now on one hydrogen,  $H_a$ . The hydrogen which is next to  $H_a$  (namely  $H_b$ ) has two spin states (aligned or opposed to  $H_o$ ) that are nearly equally populated (actually there are slightly more in the lower energy spin state than the upper, but the difference is very small). Thus the magnetic moment of the neighboring proton  $H_b$  ( $\mu_{H_b}$ ) either adds or subtracts an incremental amount ( $\mu_{H_b}$ ) to  $H_{eff}$ —the field experienced by  $H_a$ . As a consequence  $H_a$  will experience two distinct magnetic fields  $H_{eff} - \mu_{H_b}$  and  $H_{eff} + \mu_{H_b}$ . Consequently  $H_a$  will absorb energy at two distinct frequencies, and



Figure 11.11 NMR spectrum of 1,1-dichloro-2,2-dibromoethane.

#### SPIN-SPIN COUPLING 345



Figure 11.12 Splitting diagram for 1,1-dichloro-2,2-dibromoethane.

the signal for H_a will be split into two lines—a doublet—due to the presence of the adjacent proton H_b. Focusing now on H_b, the same analysis leads to the prediction that H_b will also experience two distinct magnetic fields,  $H_{\text{eff}} - \mu_{\text{H}_a}$ and  $H_{\text{eff}} + \mu_{\text{H}_a}$ ; absorb energy at two different frequencies; and thus will be split into a doublet by the presence of H_a (Figure 11.12).

The middle of the doublet corresponds to the actual chemical shift of the proton due to  $H_{\text{eff}}$ , the total integrated area under both lines of the doublet corresponds to the signal intensity of one proton, and the width between the two lines in hertz (cps) is called J, the coupling constant. The coupling constant is a measure of the strength of the interaction between the coupled nuclei that leads to spin-spin splitting. The J values for proton-proton coupling can range from 0 to 20 Hz, but most commonly coupling constants fall in the range of 0–10 Hz.

A value of J = 0 means that there is no significant interaction with neighboring protons, and thus the absorption is not affected by the spin states of neighboring protons. This normally occurs when there are *more than three bonds* separating different types of protons.



Geminal coupling (two bonds,  $J_{gem}$ ) and vicinal coupling (three bonds,  $J_{vic}$ ) are the types of spin-spin splitting normally encountered. In addition, the interaction between protons is reciprocal—if two protons are coupled, they are coupled equally and  $J_{1,2} = J_{2,1}$ . That is, if  $H_a$  is split by  $H_b$  by some amount, say J =6 Hz, then  $H_b$  must be split by  $H_a$  by J = 6 Hz. Finally equivalent protons do not split each other; thus the *t*-butyl hydrogens of *t*-butanol are a singlet (Figure 11.13).

#### 346 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS



Figure 11.13 NMR signal for the methyl protons of *t*-butanol.



**Figure 11.14** Proton coupling in 1,1,2-trichloroethane.



If there are more than one neighboring hydrogen atom, then different splitting patterns are observed. For example, 1,1,2-trichloroethane has two signals, a two-proton doublet (J = 6 Hz) upfield (H_a) and a one-proton triplet (J = 6 Hz) downfield (H_b) (Figure 11.14). The equivalent protons H_a of the CH₂–Cl group give the upfield absorption, which is split into a doublet by the single vicinal proton H_b. The single methine proton H_b gives the downfield absorption, which is split into a triplet by the two adjacent, equivalent methylene protons H_a. The triplet splitting is due to the three spin distributions possible for the two equivalent CH₂ protons, both aligned, one aligned (two possibilities), both opposed. The H_b triplet has three lines in a 1 : 2 : 1 ratio, which reflects the numbers of spin states of the neighboring CH₂ group. Because the two sets of protons are coupled, the spacing between each line of the triplet (J = 6 Hz) must be the same as the doublet splitting (J = 6 Hz). The middle line of the triplet corresponds to the chemical shift of the  $CH_2$  group, whereas the middle of the doublet corresponds to the chemical shift of the methine proton  $H_b$  (Figure 11.15).

Diethyl ether (Figure 11.16) has a three-proton triplet at  $1.2\delta$  (J = 7 Hz) for the methyl protons, which are split by the two protons of the CH₂ group. The methylene protons absorb at 3.3 $\delta$  and are split into four lines (quartet) in a 1:3:3:1 ratio (Figure 11.17). This splitting occurs because the three equivalent protons of the methyl group can have four possible spin distributions which are nearly equally populated. They are three aligned; two aligned, one opposed (three possibilities); one aligned, two opposed (three possibilities); and all opposed. The center of the quartet is the chemical shift of the CH₂ group and the coupling constant of the quartet (J = 7 Hz) must be the same as the coupling constant of the methyl triplet (J = 7 Hz) since the two sets of protons are coupled. (When protons are coupled, the signal for each set is split by the same coupling constant.)

If these considerations are generalized, it is seen that the signals for protons coupled equally to n equivalent vicinal protons will be split in to multiplets



Figure 11.16 NMR spectrum of diethyl ether.



Figure 11.17 Splitting diagram for diethyl ether.
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#### 348 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS

having n + 1 lines. The intensities of the individual lines in the multiplets follow Pascal's triangle:

Nearest Neighbors	Lines/Intensity	Splitting
0	1	Singlet
1	11	Doublet
2	1 2 1	Triplet
3	1331	Quartet
4	1 4 6 4 1	Pentet
5	1 5 10 10 5 1	Sextet
6	1 6 15 20 15 6 1	Septet
	Pascal's triangle	

The middle of the multiplet is the chemical shift of the protons responsible for that absorption, and the total integrated area under the multiplet corresponds to the total number of protons of the signal; however, the integrated area of the individual lines of the multiplet are in the ratio of Pascal's triangle. Several examples of simple splitting patterns are shown:



The n + 1 rule for predicting the multiplicity of a given proton signal holds when the coupling constants with all of the nearest neighbors are the same. For example, the multiplicity of the central methylene group of 1-bromo-3chloropropane (Figure 11.18) is a pentet which requires that  $J_{12} = J_{23}$ . That is, the central methylene group has the same coupling constant to the protons of the bromomethyl group ( $J_{12}$ ) as to the protons of the chloromethyl group ( $J_{23}$ ). Those groups are not equivalent and have different chemical shifts, but each signal is split into a triplet by the C-2 methylene group by the same J value.

If a proton or set of protons is not coupled equally to neighboring protons, then the n + 1 rule is not adequate to describe the multiplicity of the absorption. Instead one observes multiplets of multiplets as the splitting pattern (e.g., doublet of doublets or triplet of doublets). The multiplicity can be understood by carrying out sequential splitting diagrams. For example, consider a proton H_b split by two neighboring vicinal protons H_a and H_c by  $J_{ab} = 2$  Hz and  $J_{bc} = 7$  Hz. This is

#### SPIN-SPIN COUPLING 349



**Figure 11.19** Splitting pattern when  $J_{ab} \neq J_{bc}$ .

shown schematically in Figure 11.19 where the H_b signal is split into a doublet by H_c ( $J_{bc} = 7$  Hz) and each line of that doublet is split into a doublet by H_a ( $J_{ab} = 2$  Hz). The result is a doublet of doublets. The spacing between the small doublet splitting is J = 2 Hz and the splitting between the centers of the two doublets is J = 7 Hz. The same diagram is produced by first splitting the H_b signal by  $J_{ab} = 2$  Hz and then splitting each line into a doublet by  $J_{bc} = 7$  Hz.

Because of the requirement that  $J_{ab} = J_{ba}$ ,  $H_a$  will be split into a doublet (J = 2 Hz) by  $H_b$  and  $H_c$  will also be split into a doublet (J = 7 Hz) by  $H_b$ . Taking into account these different splitting patterns, the connectivity relationships between  $H_a$ ,  $H_b$ , and  $H_c$  are clear. Because  $H_a$  and  $H_c$  are both doublets but they are split by different coupling constants, they cannot be coupled to each other. The signal for  $H_b$ , however, is seen to be a doublet of doublets with J = 2 Hz and J = 7 Hz. Since these values are the same as the couplings of  $H_a$  and  $H_c$ ,  $H_b$  is coupled to both  $H_a$  and  $H_c$  and the connectivity is thus between  $H_a$ ,  $H_b$ , and  $H_c$ . Splitting patterns are thus powerful ways to establish connectivity in molecules. The patterns seen in Figures 11.20 and 11.21 are typical of the types of connections encountered in various organic compounds.

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## 350 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS



## **DESCRIPTIONS OF SPIN SYSTEMS**

It is often helpful to categorize spin systems in terms of the chemical and magnetic equivalence of coupled protons. Protons are chemically equivalent if they have the same chemical environment and thus the same chemical shift. Chemical equivalence can result from either identical environments or rapid rotations which yield an "average" environment for a group of protons. Considering toluene, it

is seen that the two meta protons are found in the plane of the ring between the ortho and para protons.



They have the same chemical environment and thus absorb at the same frequency. The methyl group is a singlet indicating that the three methyl protons absorb at the same frequency and thus are chemically equivalent, yet in the conformation shown it is clear that the environment of each proton is not the same. One is found in the plane of the ring while a second is above and the third is below the plane of the ring. However, due to rapid rotation of the methyl group, these hydrogens rapidly exchange positions and thus all absorb at an "average" frequency and all are chemically equivalent by rotation.

Protons are magnetically equivalent if they have the same chemical shift and are coupled equally to other equivalent nuclei in the molecule. This is similar to chemical equivalence but is a more rigorous definition of equivalence. For example, the methyl protons of isobutane are chemically and magnetically equivalent since they absorb at the same frequency and are all coupled equally to the methine proton (which should be split into a 10-line multiplet!). Likewise the two methyl groups of *p*-xylene are chemically and magnetically equivalent because they are coupled equally (J = 0) to the aromatic protons both ortho and meta to them.



On the other hand, the ortho protons of *o*-dibromobenzene are chemically equivalent because they have the same environment, but they are magnetically nonequivalent since a given ortho proton is not coupled equally to the two meta protons (there is a 1,2 interaction with one and a 1,3 interaction with the other and  $J_{1,2} \neq J_{1,3}$ ). By the same arguments the methylene groups of 1,4-dibromo-*cis*-2-butane are chemically equivalent but magnetically nonequivalent because each methylene group is not coupled equally to the chemically equivalent vinyl hydrogens (again  $J_{1,2} \neq J_{1,3}$ ). Of course this also means that the vinyl hydrogens are

magnetically nonequivalent since a given vinyl hydrogen is not equally coupled to the two methylene groups.



We would expect that the spectrum of the latter compound would consist of two signals: a two-proton triplet in the vinyl region and a four-proton doublet in the allylic region. This is because the coupling constant  $J_{1,3}$  is zero. It if were not zero, then a more complicated spectrum would result. Thus magnetic nonequivalence can lead to much more complicated spectra.

Using chemical and magnetic equivalence, it is possible to designate the number and type of different protons in a spin system. This is done by choosing letters of the alphabet to indicate protons of similar chemical shift: ABC or MNO or XYZ. For only two types of protons, letters from the first part and last past of the alphabet are chosen (A, X). If three types are present, then letters from the middle group are also used (e.g., A, M, X). A subscript is used to indicate how many of each type of protons is present in the spin system. Several spin systems are shown below with their designations. All are examples of groups of chemically and magnetically equivalent protons. A molecule can contain more than one spin system if they are isolated from each other.



Protons that are chemically equivalent but magnetically nonequivalent are indicated by, for example, AA'. The examples of such systems given below illustrate the method. This system for designating spin systems is merely a labeling device. The appearance of actual spectra will depend on the magnitude of the various J values. Nevertheless this is a convenient and common way of categorizing coupled proton systems.



Another structural factor which can lead to nonequivalence of aliphatic protons is the symmetry properties of protons:

1. Aliphatic protons which are interconvertible by a rotational axis are termed homotopic and are chemically and magnetically equivalent. For example, the methylene protons of diphenylmethane are homotopic, as are the methylene protons and the methyl protons of propane.



2. Methylene protons which are not interconvertible by rotation but are interconvertible by reflection through a plane of symmetry are enantiotopic and are chemically and magnetically equivalent in an achiral environment. Alternatively protons are enantiotopic if sequential replacement by a different group gives a pair of enantiomers. The methylene protons of methyl propionate are enantiotopic because they are interchangeable by reflection but not rotation. (The protons of both methyl groups are interchangeable by rotation and are thus homotopic.) Replacement of  $H_a$  and  $H_b$  by another group such as hydroxy gives R- and S-methyl lactate, respectively. The benzylic protons of benzyl alcohol are enantiotopic by the same criteria.



3. Methylene protons which are not interconvertible by either reflection or rotation are diastereotopic and are chemically and magnetically



Figure 11.22 Diastereotopic protons H_a and H_b of ethyl 4-oxo-2-azidopentanoate.

nonequivalent. The presence of one or more chiral centers in a molecule leads to diastereotopic methylene groups since the replacement of each proton by another group gives a pair of diastereomers. Since diastereotopic protons are not related by symmetry, they have unique environments and thus unique chemical shifts and coupling constants.

The C-2 methylene protons  $H_a$  and  $H_b$  in ethyl-3-azido-4-oxopentanoate are diastereotopic because of the chiral center at C-3 (Figure 11.22). The protons  $H_a$ and  $H_b$  have slightly different chemical shifts and split each other, and they are not coupled equally to the C-3 methine proton  $H_c$ . Thus  $H_a$  and  $H_b$  split each other into an AB quartet, which is further split into doublets by  $H_c$ . Note that the splitting for each proton of the AB quartet has a different coupling constant with  $H_c$ . Although  $H_c$  is slightly obscured by the CH₂ protons of the ethyl group, it can be seen that the signal for this proton is not a triplet but rather looks like a doublet of doublets, as expected from the fact that  $J_{ac} \neq J_{bc}$ .

### SECOND-ORDER SPLITTING

Our discussions of spin-spin splitting and multiplicity have been based on *first-order or weakly coupled spectra* which are spin systems where  $\Delta \nu/J \ge 10$ . The difference in chemical shift in hertz of coupled protons divided by the coupling constant is 10 or more. In such a case clean 1 : 1 doublets, 1 : 2 : 1 triplets, and so on, are observed and coupling constants and chemical shifts can be read directly from line positions in the spectrum.

As  $\Delta v/J$  decreases, the simple multiplets observed in weakly coupled spectra become increasingly distorted; new lines can appear and others merge or disappear. Such spectra are termed *second-order or strongly coupled spectra*. In these cases the chemical shift does not lie in the center of the multiplet and coupling constants are not always obvious. A simple example of such a change is seen



Figure 11.23 The change from a weakly coupled AX system to a strongly coupled AB system.

when the chemical shifts of a first-order AX system become much closer and the spectrum becomes a second-order AB system (Figure 11.23). This is not a 1:3:3:1 quartet but an AB quartet in which the intensities of the inner and outer lines depend on the difference in chemical shifts.

The treatment of such systems is outside the scope of this book, but it is possible to calculate the chemical shifts and coupling constants from line positions and intensities. There are also experimental methods by which chemical shifts and coupling constants can be determined in complex spectra. These include isotope exchange, decoupling techniques, lanthanide shift reagents, and the use of higher field NMR spectrometers. Since  $\Delta v$  increases with the strength of the magnetic field while J values do not change with magnetic field strength, the ratio  $\Delta v/J$  increases as the field strength increases. Thus the higher the field strength, the larger is the ratio  $\Delta v/J$  and the greater is the chance to observe first-order coupling. In recent times spectrometers of 300–500 MHz are routinely accessible so that the problems of second-order spectra are becoming much less common. With the advent of even higher field instruments, first-order spectra will be available for most compounds. (The first commercial 750-MHz spectrometer was delivered in 1994.)

# STRUCTURE IDENTIFICATION BY ¹H NMR

Generally the NMR spectrum of a compound is used in conjunction with other available information for identification purposes. The reactants and the reagents and reaction conditions can serve as a guide to the types of products that might be expected. Structure identification often merely confirms the structures of products that were predicted from the chemistry employed in the synthesis. In other cases products are obtained whose spectra do not match the predicted products. In such cases more information is usually required to solve the structure. Thus while NMR is an extraordinarily powerful tool, it is not sufficient to solve all structural problems. This latter fact must be kept in mind.

The reaction between 1-phenyl-1-propanol and chromic acid gives a liquid product  $\mathbf{P}_1$  with the ¹H NMR spectrum shown in Figure 11.24. The spectrum of the reactant 1-phenyl-1-propanol contains a five-proton broad singlet for the

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356 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS



Figure 11.24 NMR spectrum of 1-phenyl-1-propanol.

aromatic protons. The methine proton  $H_1$  is split into a triplet by  $H_{2,2'}$  and is further split into doublets by the OH proton, which must be exchanging slowly in this sample to give splitting. The OH proton is split by  $H_1$  into the doublet at 2.1 $\delta$ . Protons  $H_2$  and  $H_{2'}$  are a multiplet rather than a pentet as expected by the n + 1 rule. This is due to the fact that they are diastereotopic. Thus they have different chemical shifts, split each other, and each are split into pentets by the n + 1 rule. The multiplet at 1.75 $\delta$  is the result.

The NMR spectrum of the product  $P_1$  (Figure 11.25) shows the five-proton aromatic signal at 7.3 $\delta$  in the reactant has been converted to a three-proton multiplet over the range of 7.35–7.6 $\delta$  and two-proton doublet at 7.94 $\delta$  (5H total). The product also has an  $A_2X_3$  system (an ethyl group). The chemical shift of the methylene group at 2.99 $\delta$  in the product is reasonable for a methylene group next to an aromatic carbonyl group. Furthermore the one-proton multiplet at 4.55 $\delta$  for the methine proton and the one-proton doublet for the OH proton of the starting material are not present in the product. The NMR is consistent with an oxidation of the alcohol to propiophenone as predicted by the chemistry.

When 1-methylcyclohexanol is heated with anhydrous copper sulfate, two products  $P_2$  and  $P_3$  are isolated in a 85:15 ratio. The NMR spectra of these products are shown in Figures 11.26 and 11.27. The appearance of signals in the olefinic region of both products indicates that both are elimination products. Furthermore the major product has an intact methyl group (s, 1.63 $\delta$ , 3 H) whose chemical shift indicates it is likely allylic. The vinyl signal integrates for a single



Figure 11.27 Methylenecyclohexane.

proton. The remaining protons are multiplets that have an integrated area corresponding to 8 H. From these data it is clear that the major product  $\mathbf{P}_1$  is 1-methyl cyclohexene.



The minor product  $\mathbf{P}_2$  does not have a methyl signal, and the vinyl signal is integrated for 2H. The remaining protons have a total integrated area of 10H and there is a 4H multiplet downfield from the remaining 6H multiplet. These data are consistent with *exo*-methylenecyclohexane as the minor product. The 4H signal is due to the allylic protons of the ring

Treatment of 3-pentanone with isopropenyl acetate is reported to give 3acetoxy-2-pentene. The product isolated from the reaction has the ¹H NMR spectrum shown in Figure 11.28.



Preliminary examination shows the isolated product to be a mixture; however, the products appear to be similar. The spectrum includes two singlet methyl groups (3H) and two  $A_2X_3$  shown by overlapping triplets at 1.05 $\delta$  (Can you pick them out?) and two  $AX_3$  groups shown by the allylic methyl doublets at 1.5 $\delta$  and 1.65 $\delta$ . Particularly revealing is the vinyl signal at 5.1 $\delta$ . Its relatively high field results from the fact that enol derivatives are electron rich by resonance interaction of the oxygen lone pairs with the olefinic  $\pi$  system, which causes the vinyl proton  $\beta$  to the oxygen group to be shielded. Furthermore it is not a simple



Figure 11.28 Vinyl acetate.

quartet but is actually overlapping quartets due to splitting by a methyl group. This is indicative of a partial structure.



This partial structure along with the ethyl groups and acetate methyl singlets confirms the structure assignment. The allylic methylene group is at 2.1-2.3 and is overlapped by the two acetate singlets. The product is a mixture of the Z and E isomers in the ratio of 2:5 as determined by the integrated areas of the methyl doublets. It is not possible to unambiguously assign the isomers from the NMR, but it is likely that the minor isomer is the Z isomer since the allylic methyl group would be sterically deshielded by the acetate group and absorb downfield from the E isomer. The smaller allylic methyl doublet is found downfield from the major isomer

The carbodiimide coupling of *N*-methylphenylglycine with benzylamine gives a product whose ¹H NMR is shown in Figure 11.29. The expected product is the amino amide. The NMR spectrum shows first that both reactants are incorporated in the product.



The methyl singlet is indicative of the  $-NHCH_3$  group, and the aromatic signal has increased to 10H, indicating that two phenyl rings are present in the product.





The signal at  $4.4\delta$  is proper for the benzyl group, but the splitting pattern is problematic until it is recognized that because there is a chiral center at C-2, the benzyl protons are diastereotopic and thus nonequivalent. They are part of an ABX spin system and thus give the complex splitting pattern seen—actually a two-proton multiplet that looks like a doublet or a very close AB quartet.

The two N–H protons in this compound illustrate different exchange behavior. The N–H proton at C-2 comes upfield at  $1.74\delta$  as a broadened singlet due to fairly rapid exchange and does not to split either the C-2 proton at  $4.07\delta$  or the *N*-methyl group. Conversely the amide N–H proton is a much broader singlet at 7.55 $\delta$  and splits the benzylic protons by a small amount because the exchange is slower. It turns out that when protons exchange rapidly, as they do on the NH of the amino group, the spin state of the proton is blurred and coupling information is lost. The neighboring proton cannot actually feel one spin state or the other because the protons with different spin states are exchanging rapidly.

When the proton does not exchange rapidly as on the N–H of the amide group, normal coupling is observed. Since the rates of proton exchange are often critically dependent on the solution conditions, coupling to acidic protons is variable and thus may or may not be observed.

The above examples illustrate how NMR spectra are routinely used to answer questions about reactions and products. Spectra are usually examined in conjunction with other information that permits a broad-based structure identification to be carried out. Outside of structure questions in texts and on exams, one is almost never handed an NMR spectrum and asked to identify the compound in the absence of other supporting information.

### CARBON-13 NMR

While proton magnetic resonance (PMR) is the most common type of NMR, it is also possible to observe other nuclei which have spin quantum numbers not equal to zero. Of greatest interest to organic chemists is ¹³C NMR spectroscopy. Carbon-13 has a spin quantum number  $I = \frac{1}{2}$ , the same as a proton, so that when placed in a magnetic field, two possible orientations with respect to the field are possible—one of lower energy and one of higher energy. Transitions between these two spin states occur at discrete frequencies in the radio frequency region. Absorption of energy at the resonance frequency causes nuclei in the lower energy level (aligned) to undergo a transition to the higher energy level (opposed). This process is the same as discussed previously for protons, and the equations which govern the absorption are the same and will not be repeated.

There are significant differences between a  13 C nucleus and a proton which must be dealt with:

- 1. Low (~1%) natural abundance of  13 C.
- 2. Lower magnetogyric ratio of ¹³C, making the signal for ¹³C much lower than that of a proton.

3. Strong coupling to protons, although first order, gives complex multiplets which often overlap, making peak assignments difficult.

These limitations made the development of  13 C NMR spectroscopy lag significantly behind the development of  1 H NMR. In the earliest work the relatively weak sensitivity of  13 C was a major stumbling block and compounds specifically labeled with  13 C had to be prepared in order to obtain usable spectra. Today it is possible to obtain excellent  13 C spectra on natural abundance samples of <25 mg in less than 30 min. The hardware and software advances which have enabled such progress to be made lie in three areas:

- 1. Improved signal detection
- 2. Fourier transform techniques
- 3. Digital signal averaging

Suffice it to say that modern NMR spectrometers are capable of obtaining ¹³C spectra quickly and easily so that ¹³C NMR is now a routine tool for structure identification.

These instrumental improvements do not solve the problems of coupling between protons and carbon that complicate ¹³C spectra, but other techniques have. The proton-coupled ¹³C spectrum of 3-octanone demonstrates that proton-carbon coupling significantly complicates the spectrum due to the large number of lines produced. From this spectrum, it is impossible to tell how many carbons are present or what are their chemical shifts because of overlapping multiplets in the spectrum. To solve this problem, broad-band proton decoupling is used to remove all proton-carbon couplings, and one is left with proton decoupled or fully decoupled spectra which have only singlet absorptions for each carbon present. For example, 3-octanone has eight lines in the fully decoupled ¹³C spectrum, as predicted by the fact that each of the carbons is in a unique chemical environment and thus has a unique chemical shift. A distinct advantage of ¹³C NMR is that ¹³C absorbs over a range of  $\sim$ 250 ppm (compared to 10 ppm) for ¹H). This means that each carbon can be distinguished by a unique chemical shift. Thus it is possible to tell exactly how many nonequivalent carbons are present in a molecule merely by counting the lines in the fully decoupled spectrum (Figure 11.30). (The small three-line signal at 77.3 $\delta$  is from the solvent CDCl₃. It appears in all ¹³C spectra run in CDCl₃ and is normally ignored.)

In addition to the number of nonequivalent carbons present, the chemical shifts of the carbons can reveal a great deal about the types of bonding patterns and substituents which are present. Because of the great range of chemical shifts observed for ¹³C ( $\approx$ 250 ppm), even small changes in the environment around carbon can result in a significant change in chemical shift. Figure 11.31 is a brief compilation of ¹³C chemical shifts for representative classes of organic compounds. By assigning the chemical shifts in many series of compounds, it has been possible to develop correlation equations for calculating ¹³C chemical shifts based on structural features present in the molecule. These correlation



Figure 11.31 Representative ¹³C chemical shifts for various classes of organic compounds.

equations generally provide excellent agreement between calculated and observed ¹³C chemical shift values. Thus it is now routine to test possible structures by calculating ¹³C chemical shifts and comparing them with the observed spectra. For example, the calculated and observed ¹³C chemical shift values for cocaine are seen to be in remarkable agreement for most of the carbons in this reasonably complex compound.



The value of fully decoupled ¹³C NMR spectra is primarily tied to determining how many nonequivalent carbons are present and their chemical shifts. Unfortunately, the integrated areas of ¹³C signals are not directly proportional to the numbers of carbons responsible for those signals under most circumstances. Thus both *n*-heptane and 4-(1-propyl) heptane have four signals in their ¹³C NMR spectra, but it is not possible to determine if the ratio of different carbon types is 1:2:2:2 as expected for *n*-heptane or the 1:3:3:3 expected for the branched compound.



It *is* possible to distinguish them based on the chemical shift of C-4, which is calculated to be 36.5 ppm in *n*-heptane but 53.1 ppm in the branched compound.

A second difficulty of fully decoupled ¹³C NMR spectra is that the connectivity in the molecule is difficult to establish (except by chemical shift correlation) because coupling patterns are absent. This dilemma is partially resolved by the use of a technique called off-resonance decoupling. In off-resonance decoupled ¹³C spectra, the carbons are coupled only to those protons directly attached to them and the coupling is first order. Thus quaternary carbons are singlets, methine carbons are doublets, methylene carbons are triplets, and methyl carbons are quartets. It is possible to use this information to establish proton–carbon connectivity,

which can be used to add protons to partial structures determined by ¹³C chemical shift data.

The carbons of 1,2-epoxy-5-hexene can be assigned from the off-resonance decoupled spectrum (Figure 11.32). In the fully decoupled spectrum it is clear that the olefinic carbons ( $\approx$ 115 and 138 $\delta$ ) are distinct from the epoxide carbons ( $\approx$ 47 and 52 $\delta$ ) and from the methylene carbons ( $\approx$ 30 and 32 $\delta$ ), but it is not possible to assign which is which. In the off-resonance decoupled spectrum, both the olefinic and epoxide carbons are distinguished by their splitting patterns from the numbers of directly attached protons. The methylene carbons, however, are both triplets and cannot be distinguished.

A final feature of importance in ¹³C NMR spectra is the notion of equivalency. Because some type of decoupling is normally done, either broad band or off resonance, magnetic equivalency is not an issue in ¹³C NMR, but chemical equivalence remains an issue. If two carbon atoms share the same chemical environment, then of course they will have the same chemical shift. Thus it is important to recognize local or molecular symmetry elements. In a previous



Figure 11.32 Fully decoupled and off-resonance decoupled spectra of 1,2-epoxy-5-hexene.

example *n*-heptane is seen to have four signals. The internal plane of symmetry results in three equivalent pairs of carbons in addition to the unique central carbon. Toluene (or any monosubstituted benzene) has four signals for the aromatic protons in addition to the methyl carbon signal. The xylenes offer another example of equivalency.



*o*-Xylene has four signals, *m*-xylene has five signals, and *p*-xylene has only 3. In general, the more symmetric is a molecule, the fewer ¹³C signals it will have. For example adamantane has only 2 absorptions and buckminsterfulluene ( $C_{60}$ ) has only a single line in its ¹³C spectrum.



Thus when the number of ¹³C signals is less than the number of carbon atoms present in the molecule, there must be symmetry elements present that make some carbon atoms equivalent. The pyrolysis of 2-acetoxy-2.3-dimethylbutane in a hot tube at 200°C gives two products which are both found to have the formula  $C_6H_{12}$ . The major product has only two ¹³C absorptions while the minor product has five ¹³C signals (Figure 11.33). Thus the major product is likely to



Figure 11.33 Major and minor butene products.

be the symmetric olefin while the minor product is the less symmetric olefin. Note that even the minor product has a pair of equivalent carbons giving rise to five rather than six lines, but the symmetry is still significantly less than that of the major olefin.



The foregoing has been a brief introductory discussion of NMR which has concentrated on some basic principles that are very useful in understanding the technique. The actual practice of NMR today is much more advanced. The incorporation of Fourier transform techniques has revolutionized NMR spectroscopy. All types of pulse sequences and two-dimensional (2D) techniques have been developed to provide even greater structural detail than has been discussed above. A discussion of such techniques belongs in a more specialized text, but it must be remembered that while these techniques are faster, more sensitive, and much more sophisticated, they are still largely based on the principles presented here, as is the interpretation of the results.

## INFRARED SPECTROSCOPY

Infrared spectroscopy is a very useful spectroscopic tool for determining the presence of functional groups and bonding sequences in a compound by the absorption of light in the IR region of the electromagnetic spectrum. The IR region comprises light with wavelengths from about  $1 \times 10^{-4}$  to  $8 \times 10^{-7}$  m (100–0.8 µm) and lies between the microwave region and the visible region of the spectrum. The wavelengths of greatest interest to organic chemists range from 2.5 to 15 µm, the so-called mid-IR region, because the greatest amount of structural information can be obtained by spectroscopy in this spectral region.

Infrared radiation, like any electromagnetic radiation, is characterized by properties of frequency ( $\nu$ ) and wavelength ( $\lambda$ ) that are related by the speed of light *c*:

$$c = \lambda v$$
 or  $v = \frac{c}{\lambda}$ 

To scale frequency to a more convenient range, IR spectroscopists have defined a frequency unit called wave number  $\overline{\nu}$  given by  $\overline{\nu} = 1/\lambda$ , where  $\lambda$  is the wavelength in centimeters. The units of  $\overline{\nu}$  are reciprocal centimeters (cm⁻¹). The wave number is the number of vibrations which occur over a 1-cm distance. Thus the higher the wave number, the more vibrations occur in a 1-cm distance and thus the higher the frequency. Normally IR spectra are recorded between 4000 and 650 cm⁻¹ (2.5 and 15  $\mu$ m). The energy of IR radiation is given by E = hv and thus

$$E = h/c\overline{\nu}$$

The energy of the IR light absorbed by molecules during IR spectroscopy is typically in the range of 9-2.5 kcal/mol (for 4000-650-cm⁻¹ light). This amount of energy is not enough to break bonds in molecules, but it is enough to cause transitions in vibrational modes in the molecule. Thus IR spectroscopy is best described as nondestructive, vibrational spectroscopy.

As was discussed in the chapter on chemical bonding, a molecule can simplistically be thought of as a collection of bonds which hold the nuclei together in certain spatial relationships so that the lowest possible energy for the molecule is achieved. Deformations from these optimal angles and distances correspond to bond stretchings and bendings and these are examples of vibrational motion in the molecule. Since vibrational motions, which include both bond-stretching and bond-bending modes of vibration, are quantized, each vibrational mode in the molecule absorbs energy at a particular frequency (which happens to fall in the IR region). This provides the basis for IR spectroscopy. By determining which frequencies of IR radiation are absorbed by a molecule, it is possible to conclude what types of vibrational modes are absorbing energy in the molecule and consequently what atoms and bonds (functional groups) are present in the molecule which give rise to these vibrational modes.

A molecule which contains *n* atoms will have 3n - 6 fundamental modes of molecular vibration. These 3n - 6 fundamental vibrational motions can be divided into two types—stretching modes, of which there are n - 1, and bending modes, of which there are 2n - 5. Stretching vibrations are those in which the internuclear distances between bonded elements change. Bending vibrations are those in which bond angles change. In general, it takes more energy to stretch a bond than to deform bond angles. Therefore absorption frequencies which correspond to bond-stretching modes are often higher (higher energy) than absorption frequencies which correspond to bending modes (lower energy). Stretching frequencies are normally found in the higher frequency portion of the spectrum (4000–1200 cm⁻¹) while bending frequencies are found in the lower frequency region (~1200–600 cm⁻¹). Furthermore, it is the stretching frequencies which give the most clearcut structural information about a compound.

# **IR STRETCHING FREQUENCIES**

As mentioned above, stretching modes are those in which internuclear distances change. For such a distance change to occur, the bond between the nuclei must be stretched or compressed. One way to think about this is to consider a chemical bond as a spring connecting two masses (Figure 11.34). Each spring has a particular force constant which corresponds to the force required to



Figure 11.34 Analogy between a spring connecting two masses and a chemical bond.

compress or stretch that spring. Furthermore the frequency of vibration is dependent on the masses that the spring connects. For a given spring, heavy masses lead to lower frequency vibrations while light masses lead to higher frequency vibration.

This analogy is actually very close to correct in describing stretching vibrations of bonds because the potential curves for springs and bonds have the same general shape and behavior at lower energies. The major difference is that the vibrational energies of a bond are quantized while those of a spring are not. As a result there are discrete vibrational energy levels that are allowed for a bond in a molecule. At room temperature, the vast majority of molecules are in the lowest vibrational energy levels.

To go from the lowest vibrational level  $v_1$  to the next higher vibrational level  $v_2$  (which is the only type of vibrational transition that is allowed), energy must be absorbed by the molecule which corresponds exactly to  $\Delta E$ , the energy gap between the two vibrational levels (Figure 11.35). It follows that if the energy required to cause a vibrational transition is  $\Delta E$ , then only light of the particular frequency and hence energy which corresponds to the energy of the vibrational transition will be absorbed. Moreover the frequency of light which causes the transition is given by



Figure 11.35 Potential curves for springs (left) and bonds (right).

Functional Group	Frequency (cm ⁻¹ )	
Alcohol O-H (free)	3640-3610	
Alcohol O-H (H bonded)	3500-3200 (variable)	
Amine N–H	3500-3300	
	$(1^{\circ} \text{ doublet}, 2^{\circ} \text{ singlet})$	
Terminal alkyne C-H	3315-3270	
Olefinic and aromatic C-H	3080-3020	
Aliphatic C–H	2990-2850	
Aldehyde C–H	2900-2700	
Nitrile −C≡N	2300-2200	
Terminal –C≡C	2260-2210	
Internal –C≡C	2140-2100 (weak)	
Ester C=O	1750-1740	
Aldehyde C=O	1740-1720	
Ketone C=O	1700-1720	
Amide C=O	1715-1650	
Unsaturated ketone C=O	1680-1660	
Alkene C=C	1675-1640	
Aliphatic C–O	1280-1000 (strong)	

 Table 11.1
 Functional Group Stretching Frequencies

For most bonds, these frequencies occur in the IR region. Each stretching mode in a molecule has its own potential curve and associated energy levels. Thus each stretching mode in a molecule will absorb IR energy at the particular frequency required to cause the transition from the lowest energy level to the next higher energy level. Since each of these energy levels is dependent upon the force constants of the bonds and the masses of the atoms they connect, each type of bond has a characteristic IR absorption corresponding to the stretching frequency of that bond.

Because functional groups are recurring groups of atoms connected by similar bonding patterns, a given functional group tends to give characteristic IR absorptions due to the vibrational frequencies of bonds present in that functional group. Infrared spectroscopy thus provides a fast and effective way to identify functional groups present in a molecule by noting the presence of absorptions corresponding to the bond types present in those functional groups. Table 11.1 is a compilation of IR absorptions for commonly encountered bonds and functional groups. Normal ranges are given since the absorption frequency of a given bond type can vary somewhat depending on the structure. The frequencies given are all stretching frequencies; bending frequencies are much more numerous and usually harder to interpret and are not included in this work.

It is also possible to identify structural features in molecules which strengthen or weaken bonds and thus lead to shifts in IR frequencies. Several general effects on IR frequencies are summarized as follows:

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- 370 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS
  - 1. Multiple bonds are stronger than single bonds and thus have larger force constants and absorb at higher frequencies than single bonds. For example,



2. More polar bonds are generally stronger than less polar bonds and consequently absorb at higher frequencies.

С-Н	N-H	О-Н	F - H
2862	3300	3650	$4138 \text{ cm}^{-1}$
-C-H $sp^3$	=	н Н	<del>≡</del> −H
2890	3040	0	$3300 \text{ cm}^{-1}$

3. Conjugation lowers the absorption frequency of each conjugated group due to a lowering of the bond order due to the contributions of resonance forms with lower bond orders.



4. Hydrogen bonding causes the absorption frequency of acidic protons to vary widely depending on the solution environment. In general, the greater is the H bonding, the lower is the absorption frequency. For example, normal alcohol OH groups in dilute, nonbasic solvents come at 3610–3650 cm⁻¹ (termed the free OH stretch). As the concentration is increased and H bonding increases, the OH absorption becomes broad and moves to lower frequencies.

#### USE OF IR SPECTROSCOPY FOR STRUCTURE DETERMINATION 371



The very strongly H-bound carboxylic acid dimer has a very broad absorption at  $\sim 2500 \text{ cm}^{-1}$  for the OH bond. Because H bonding is dependent on concentration and on the polarity and H-bonding properties of the solvent, frequency shifts due to H bonding are quite variable.

While other structural effects on absorption frequencies are known, the above factors are the ones most commonly encountered in routine organic structure determination.

# USE OF IR SPECTROSCOPY FOR STRUCTURE DETERMINATION

As seen above, IR spectroscopy is most commonly used to identify functional groups and bonding patterns in molecules from the higher energy portion of the spectrum  $(1200-4000 \text{ cm}^{-1})$  where absorptions are primarily due to bond-stretching vibrations. Some information on atom connectivity in the molecule can also be deduced from the frequency shifts caused by structural factors. In general, however, it is not possible to completely deduce the structure of a molecule by examination of its IR spectrum. However, IR spectroscopy is a powerful complement to NMR spectroscopy for structure determination.

For example, the reaction of 3-chlorocyclohexanone with DBU in toluene gives a product which is seen to have two vinyl protons by NMR and thus is an elimination product, probably either A or B. Now while one might rationalize by both chemical intuition and by the splitting pattern that conjugated isomer A is the product, examination of the IR spectrum shows a carbonyl group (at 1680 cm⁻¹) and an olefin band (at 1630 cm⁻¹). A typical cyclohexanone comes at 1710 cm⁻¹ and cyclohexene comes at 1643 cm⁻¹.



Clearly the observed frequencies of the product are at lower frequencies than a simple ketone or olefin and are indicative of a conjugative interaction between these two functions. Thus A and not B is the product.

Treatment of propiophenone with *m*-CPBA (*m*-chloroperbenzoic acid) in dichloromethane gives a single product. The carbonyl absorption of propiophenone is at 1695 cm⁻¹ whereas the product has a carbonyl absorption at 1745 cm⁻¹. This information reveals that the carbonyl group is intact but is no longer a ketone. The shift to higher frequency is consistent with the conversion to an ester, and thus the product could be either **C** or **D**.



Since ethyl benzoate **C** has a carbonyl stretch at 1725 cm⁻¹, the likely product is **D**, phenyl propionate. This is confirmed by the NMR spectrum, which has the methylene group as a quartet at 2.42 $\delta$ . This chemical shift is typical for a methylene group next to an ester carbonyl but is much too high field for the –CH₂–group in ethoxy ester **C**, which comes at about 3.6 $\delta$ .

Reaction of cinnamic acid, which has the IR and ¹H NMR spectra shown in Figures 11.36–11.38, with BH₃·THF gives rapid consumption of the starting material. A single product  $P_4$  is formed that has the spectra shown. Comparison of the IR spectra shows that the double bond in the reactant (1630 cm⁻¹) is intact in the product (1654 cm⁻¹) but moved to higher frequency. The product  $P_4$  also has both vinylic (3026 cm⁻¹) and saturated (2861 cm⁻¹) C–H bonds. Both reactant and product have an O–H adsorption, but the strong H bonding in



Figure 11.36 IR spectrum of cinnamic acid.



Figure 11.38 NMR spectra of cinnamic acid (upper) and product P₄ (lower).

the acid (broad absorption at 3200–2600 cm⁻¹) is replaced by a shift to higher frequency in the product (3349 cm⁻¹) indicative of weaker H bonding found in an alcohol. Furthermore the carbonyl group in the reactant acid (1681 cm⁻¹) is missing in the product. The IR data suggest that the carboxylic acid has been reduced by  $BH_3$ -THF in preference to hydroboration of the double bond.

The ¹H NMR corroborates this conclusion since two vinyl protons are observed both in the reactant and product; however, a new two-proton doublet appears at 4.15 $\delta$  for the newly produced allylic methylene group. The acid O–H proton is moved far upfield as well. The coupling constants of the vinyl protons (J = 16 Hz) show the starting compound to be trans, and the large splitting for the downfield vinyl doublet of the product (J = 16 Hz) shows the trans stereochemistry to be maintained in the unsaturated alcohol product. Moreover the splitting between the methylene group and the upfield vinyl proton clearly supports its allylic position.

The ¹³C spectrum (Figure 11.39) is consistent with these structural assignments as the carbonyl carbon in the reactant (172.5 $\delta$ ) is gone and the product contains a new signal at 63.3 $\delta$ , typical for a change to sp³ hybridization and an allylic group.



In the preceding example several types of spectroscopy are brought to bear. While the product structure could probably be deduced from IR spectroscopy or NMR (either ¹H or ¹³C), the use of all three methods confirms the assignment. It is often prudent to use more than a single technique for structure determination so that the results reinforce each other. If a structure assignment is not consistent with *all* the data, the structure is probably incorrect.

This lesson is brought home in the following example taken from a recent experiment. Ample chemical precedent suggested that the treatment of E with methyl amine should give F:



The spectra of the product are shown in Figures 11.40–11.42. As is clear, all of the appropriate resonances are present. In the ¹H NMR (Fig. 11.40) the amide *N*-methyl group is a doublet (J = 4 Hz) at 2.76 $\delta$  due to weak splitting by the amide NH proton and the amino N–CH₃ group is the sharp singlet at 2.89 $\delta$  which is not split by the amine N–H proton due to rapid exchange. The C–H methine proton is a singlet at 4.03 $\delta$  and the ethoxy group is evident by the quartet–triplet A₂X₃ pattern.

However, several pieces of data just did not seem to fit. First the chemical shift of the amino N-methyl group is at 2.90 $\delta$  whereas several known compounds



Figure 11.40 Product F¹H NMR spectrum.

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#### 376 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS



of similar structure had the amino *N*-methyl group at 2.48 $\delta$ . The 0.5-ppm shift might be due to the electron-withdrawing properties of the carbethoxy group, but that shift appears to be too large. In fact the chemical shift of 2.9 $\delta$  is exactly that expected for an amide *N*-methyl, but in this case it is not split and there is already an amide *N*-methyl signal at 2.76 $\delta$ , where it should be. Next, the integrated area of the N–H peak at 4.6 $\delta$  is only 1H whereas it should account for two N–H's which exchange. Moreover, the C–H signal at 4.03 $\delta$  integrates for two protons rather than one. While one might discount the discrepancies in the integrated areas as being due to the fact that the sample was not analytically pure and while one might force fit the change in chemical shift of the amine N–CH₃ group, the compound did not dissolve in acid, as was expected for the amine. Adding to the difficulty was the IR spectrum of **F** (Fig. 11.41). In addition to the normal ester C=O stretch at 1745 cm⁻¹, the IR had a C=O stretch at 1636 cm⁻¹, which is at a much lower frequency than a normal secondary amide ( $\approx$ 1685 cm⁻¹).

Based on these discrepancies, the assigned structure had to be discarded. Starting once again from the beginning, the carbethoxy group (**a**) is set by the  $A_2X_3$ spin system and the ester C=O stretch. The 2H singlet at 4.03 $\delta$  is assigned as a two-proton CH₂ group next to the ester group (**b**) since it has the right chemical shift and integrates for two protons. Another fragment which is indicated is the C(O)NHCH₃ group by the chemical shift of the methyl group, its small splitting by the N–H amide proton, and the rather low C=O stretching frequency which suggests an amide group (**c**). What remains is an N–CH₃ fragment, which can only be placed between fragments **b** and **c**. That gives the unsymmetric



Figure 11.42 Product F ¹³C NMR off-resonance.

urea as the assigned structure of **F**. This structure fits the data in every way. The methylene group (2H) and the single N–H proton fit the integration. Both N–CH₃ groups are amide types rather than one amino and one amide type, and both should appear at  $\sim$ 2.8–3.0. The urea group has greater resonance stabilization, and thus the carbonyl group has more single-bond character and comes at a lower IR frequency than an amide. Finally **F** is *not* an amine and should not behave chemically like one, that is, it should not dissolve in acid.

$$CH_{3}CH_{2}O \xrightarrow{O} CH_{3}CH_{2}O \xrightarrow{O} CH_{2} + \underbrace{O}_{CH_{2}}CH_{3} \Rightarrow \underbrace{O}_{EtO} \underbrace{CH_{2}}_{O} \xrightarrow{V}_{O} \xrightarrow{V}_{CH_{3}}H_{V}$$

$$a \qquad b \qquad c \qquad F$$

The structure is confirmed by the off-resonance decoupled ¹³C spectrum (Figure 11.42), which shows the CH₂ group as a triplet  $61.7\delta$  in addition to the other expected carbon resonances.

It is clear that all the data must fit the structure and vice versa. If not, the assigned structure is probably not correct. It is important to let the data indicate the structure and not make the data fit a preconceived structure. Often the chemistry will suggest a structure and the data will support that structure. However, that need not always be so, and it is necessary to always check that the data fit the structure. New chemistry is often discovered precisely because expected products are not supported by structural data. Consequently new structures resulting from new chemistry are revealed.

## MASS SPECTROMETRY

Mass spectrometry is not a true spectroscopic technique that involves absorption of energy at particular frequencies. Rather one excites the molecule as a whole

and then observes its subsequent reactions. The use of MS for structure determination might be described as chemical archeology. A real archaeologist picks up fragments of a pot or other vessel, identifies them, fits them back together, and can tell what the original object was in great detail. In MS a molecule is purposely broken into pieces, the pieces are identified by mass, and the original structure is then inferred from the pieces.

To measure the mass spectrum, a molecule is bombarded with a stream of energetic electrons (70 eV) and one of the electrons of the molecule is ejected from one of the orbitals after a collision.

$$\mathbf{M} \stackrel{e^-}{\longrightarrow} \left[ \mathbf{M}^+ \right] + 2e^-$$

This produces a charged species with an unpaired electron called a radical cation. The first formed radical cation is called the molecular ion because it contains all of the atoms present in the starting molecule. The molecular ion contains a large amount of excess energy deposited by the collision which dislodged the electron. Considering that electrons of 70 eV energy contain 1613.5 kcal/mol energy, the molecular ion usually contains energies far in excess of bond dissociation energies (80–100 kcal/mol) so it dissociates into fragments.

Fragmentation processes must conserve both charge and spin; thus a radical cation can undergo the following types of cleavages. The fragment products also contain excess energy since fragmentations are adiabatic, and they can themselves undergo further fragmentations to smaller pieces.

$$\begin{bmatrix} M^{+} \end{bmatrix} \longrightarrow F^{+} + \text{ neutral molecule}$$

$$F_{1}^{+} + F_{2}^{+}$$

$$F_{1}^{+} + F_{2}^{+}$$

Generally fragmentations occur very rapidly in the region where the initial ionization takes place called the source. The packet of ions, which includes the molecular ion and the fragment ions, is then accelerated by an electric field, focused through slits, and the beam of ions travels at high speed down a curved tube toward the detector. At this point one can think of the ion beam as a stream of charged projectiles having different masses. Now moving charges are subject to the influences of electric and/or magnetic fields so if a magnetic field (or electric field) is applied to the ion beam, the path of the particles will curve. When the trajectory of a particle matches the curve of the tube, the particle will reach the detector. If not, it will hit the wall and be annihilated (Figure 11.43).

The path of the moving ion will curve according to its speed, mass-to-charge ratio (m/e), and the strength of the electric or magnetic field through which it passes. Fragments of low m/e will curve more than fragments of higher m/e. Thus if the ion packet is accelerated uniformly, the magnetic or electric field can



Figure 11.43 Path of ions in mass spectrometer.



Figure 11.44 Molecular ion and fragment ions.

be varied so that fragments of different m/e values can be curved to strike the detector and counted in turn. What results are a series of signals corresponding to different m/e values for the various charged fragments produced from the molecular ion (Figure 11.44).

The vast majority of fragments will have only a single positive charge (i.e., e = 1); thus the m/e of a given ion corresponds to the mass of the ion in atomic mass units. It is very important to remember that only ions (cations or radical cations) are detected—neutral species (closed-shell molecules or radicals) are not detected because they are not accelerated and they are not influenced by the applied field. Thus MS yields information about the mass of the molecular ion and the masses of fragment ions produced from the molecular ion. This so-called cracking pattern provides information about connectivity in the molecule that can be used to reconstruct the intact precursor molecule.

The molecular ion is one of the most important ions in the mass spectrum of a compound for the following reasons:

1. The *m/e* value of the molecular ion is equal to the molecular weight (MW) of the compound. This gives a rough estimate of the number of carbon atoms. Furthermore a knowledge of the history of the sample and the reagents used often permits the molecular formula to be deduced.

For example, treatment of *p*-toluic acid with ethyl iodide and potassium carbonate gave an oil whose molecular ion is at m/e = 164. This molecular ion corresponds to the addition of 28 mass units to the starting material, consistent with the formation of an ethyl ester by displacement of iodide.

$$H_{3}C - \underbrace{\bigcirc}_{MW=136} CO_{2}H \xrightarrow[K_{2}CO_{3}]{CH_{3}CH_{2}L} (M^{+} m/e = 164) \implies H_{3}C - \underbrace{\bigcirc}_{O} CH_{2}CH_{3}$$

2. High-resolution mass spectrometers which can measure *m/e* values to four decimal places are capable of confirming the molecular formula of the molecular ions. These so-called exact mass measurements can be used because the atomic weights of the elements are not *exactly* whole numbers (except for ¹²C, which is the standard at 12.0000 amu). The exact masses of some elements and their most abundant isotopes are given in Table 11.2. To find the exact mass of a molecule, the atomic mass of the most abundant isotope for each element is used to calculate the exact mass of the compound. This is compared to the exact mass measured on a high-resolution

Element	Isotope	Natural Abundance (%)	Exact Mass
Hydrogen	$^{1}\mathrm{H}$	100	1.00783
	² H (deuterium)	0.016	2.01410
Carbon	$^{12}C$	100	12.0000 (standard)
	¹³ C	1.08	13.0034
Nitrogen	$^{14}N$	100	14.0031
C	¹⁵ N	0.38	15.0001
Oxygen	¹⁶ O	100	15.9949
	¹⁷ O	0.04	16.9991
	¹⁸ O	0.20	17.9992
Fluorine	¹⁹ F	100	18.9984
Silicon	²⁸ Si	100	27.9769
	²⁹ Si	5.10	28.9765
	³⁰ Si	3.35	29.9738
Phosphorus	³¹ P	100	30.9738
Sulfur	$^{32}S$	100	31.9721
	³³ S	0.78	32.9715
	³⁴ S	4.40	33.9679
Chlorine	³⁵ Cl	100	34.9689
	³⁷ Cl	32.5	36.9659
Bromine	⁷⁹ Br	100	78.9813
	⁸¹ Br	98	80.9163
Iodine	$^{127}I$	100	126.9045

Table 11.2 Exact Masses of Elements and Their Common Isotopes

mass spectrometer. If the two values agree to the third decimal point, then it is certain that the molecular formula used to calculate the exact mass is correct.

Consider the three compounds  $C_8H_{16}N_2$ ,  $C_9H_{18}N$ ,  $C_9H_{16}O$ . All would give a molecular ion of m/e = 140 in the low-resolution mass spectrum. Using the elemental exact masses in Table 11.2, the molecular exact masses are calculated:

 $\begin{array}{ll} C_8 H_{16} N_2 \Rightarrow (8 \times 12.000) + (16 \times 1.00783) + (2 \times 14.0031) = 140.13148 \\ C_9 H_{18} N \Rightarrow (9 \times 12.000) + (18 \times 1.00783) + (14.0031) = 140.14404 \\ C_9 H_{16} O = (9 \times 12.000) + (16 \times 1.00783) + 15.9949 = 140.12018 \end{array}$ 

It is clear that if the mass of the molecular ion can be determined to 0.001 amu, these three compounds can be distinguished clearly. Instead of having to calculate exact masses, there are many published tables of exact masses for any elemental composition and there are many computer programs that calculate the exact mass after input of the molecule formula.

3. The analysis of isotopic clusters of the molecular ion can be used to infer the presence of elements based on their isotopes. The molecular ion corresponds to the m/e for the ion corresponding to some molecular formula. Examination of the molecular ion (Figure 11.45) reveals that in addition to the expected molecular ion, there is normally a smaller peak at M + 1 and an even smaller one at M + 2. These are due to the fact that there are naturally occurring isotopes of higher mass that, if present in a given molecule, cause its mass to be higher than for the lighter isotopes.

The most obvious example of such behavior is for molecules which contain chlorine or bromine. The two isotopes of chlorine occur naturally in the ratio of  ${}^{35}\text{Cl} : {}^{37}\text{Cl} = 100 : 32.7$  (3.058 : 1). A molecule such as chlorobenzene would







Figure 11.46 Mass spectrum of chlorobenzene.



Figure 11.47 Mass spectrum of ethyl bromoacetate.

exhibit two distinct molecular ions—one at m/e = 112 for those molecules which have the ³⁵Cl isotope and one at m/e = 114 for those molecules which contain the ³⁷Cl isotope. The intensities of these peaks should be 3.058:1, reflecting the probability that a molecule has one or the other of the isotopes (Figure 11.46).

A similar situation is seen for molecules containing bromine. The isotopes ⁷⁹Br and ⁸¹Br occur in a ratio ⁷⁹Br/⁸¹Br = 100:97.5 (1.026:1). Thus a molecule such as ethyl bromoacetate will exhibit molecular ions at m/e = 166 and m/e = 168 for molecules which contain ⁷⁹Br and ⁸¹Br, respectively. The ratio of peak intensities will be 1.026:1 because this is the relative abundance of the two bromine isotopes present in the molecule (Figure 11.47).

#### MASS SPECTROMETRY 383



Figure 11.48 Molecular ion of o-dichlorobenzene.

Now if a molecule contains more than one chlorine atom, the appearance of isotope clusters can be calculated by the probabilities of isotope distributions and the natural abundances of the isotopes. For example, if a molecule contains two chlorine atoms such as *o*-dichlorobenzene, then there will be peaks at M, M + 2, and M + 4 for molecules which have two ³⁵Cl, one ³⁵Cl and one ³⁷Cl, and two ³⁷Cl (Figure 11.48).

The relative intensities of these peaks can be calculated by taking into account the number of combinations that can give the required isotopic substitution and the probability of an isotope being present. An M + 2 peak in the above example will result if either of the chlorine atoms is ³⁷Cl; thus the intensity of an M + 2 peak will be  $2 \times (1/3.058)$ . An M + 4 peak will occur only if both chlorine atoms are ³⁷Cl; thus the intensity of the M + 4 peak will be  $2 \times (1/3.058)^2$ . The squared term follows from the necessity that both chlorine atoms must be ³⁷Cl. Thus the intensities of the peaks in the isotopic cluster of the molecular ion are (approximately) given as

$$M: M + 2: M + 4 = 1: (2 \times \frac{1}{3}): 1 \times (\frac{1}{3})^2 = 1: 0.66: 0.111$$

The presence of isotopic clusters is particularly clear for molecules containing chlorine or bromine because of the abundance of two isotopes. The same considerations are applicable, however, for other elements that have smaller abundances of higher isotopes. These natural isotopic abundances are given in Table 11.2. As can be seen, ¹³C is present to the extent of 1.08% of ¹²C while ²H is present only to the extent of .016% of ¹H. Now if a molecule such as benzene is examined,
#### 384 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS

the molecular ion is found at m/e = 78 but there are an M + 1 peak and an M + 2 peak whose intensities are 6.58% and 0.22%. The M + 1 peak is due to the probability that one of the six carbons will be ¹³C or one of the six hydrogens will be deuterium. The M + 2 peak is due the probability that two of the carbons in the same molecule will be ¹³C (the probability of two deuterium atoms in the same molecule is exceedingly small) or that one ¹³C and one deuterium are present in the same molecule. The intensities of M + 1 and M + 2 peaks can be calculated for various molecular formulas based on these probabilities and they have been tabulated in several texts.

This information can be used to deduce the elemental composition of a compound. For example, the oxidation of 1,2-diazacyclohexane was carried out in cyclohexane. A product was isolated and was found to have a molecular ion of m/e = 84. At this point the experimenter realized that both the expected product and the reaction solvent have a molecular weight of 84.



Measurement of the isotopic cluster of the molecular ion showed an M + 1 peak of 5.30% and an M + 2 peak of 0.15% of the molecular ion. From tables of isotopic abundance ratios it was found that the expected product  $C_4H_8N_2$  should give M + 1 and M + 2 peaks of 5.21 and 0.11%, respectively, while cyclohexane  $C_6H_{12}$  should give M + 1 and M + 2 peaks of 6.68 and 0.19%, respectively. It is clear that the isolated product is most likely the expected cyclic azo compound and not cyclohexane.

Of course nowadays exact mass measurement could also distinguish these two molecules, as could a variety of other instrumental techniques. The analysis of isotopic clusters is most useful for detecting the presence of halogens, sulfur, and silicon, all of which have abundant isotopes of two atomic weight units higher, thus leading to relatively large M + 2 peaks.

### FRAGMENTATION PROCESSES

Besides the molecular ion, fragmentation processes can be used to infer groups present in the molecule and the connectivity of those groups. The requirement of spin and charge conservation in any fragmentation means that both cations and radical cations can be produced as ions by fragmentation. Because of the great amount of energy deposited in the molecular ion, there is sufficient energy to break any of the bonds in the molecule. It has been found, however, that fragmentations tend not to be random but occur in such a way that the most stable ions are produced. Normally the most stable ion is the most abundant ion in the mass spectrum. The most abundant ion is called the *base peak* of the spectrum and is arbitrarily scaled at 100%, and the abundances of other ions are given as percentage relative to the base peak. Several examples of very stable ions are as follows:



Fragmentations often occur from the molecular ion by loss of neutrals or radicals to give more stable ions or radical ions. The differences in mass correspond to the mass of the uncharged fragment that has been expelled. The mass spectrum of ethane has a molecular ion at m/e = 30 and a major peak at m/e = 15. This corresponds to the loss of a fragment of 15 amu from the molecular ion. Thus the ethane molecular ion undergoes fragmentation of the C–C bond to give a methyl cation which is detected at m/e = 15 and a methyl radical which is not detected as it is uncharged. This very simple example is indicative of the process.

$$H_{3}C - CH_{3} \xrightarrow{-e^{-}} [H_{3}C - CH_{3}]^{+} \longrightarrow CH_{3}^{\oplus} + CH_{3}^{\bullet}$$
$$m/e = 30 \qquad m/e = 15$$

Ethyl benzoate (Figure 11.49) has a molecular ion at m/e = 150 and a base peak at m/e = 105 (M - 45) and a smaller peak at m/e = 77. The base peak at m/e = 105 corresponds to loss of the ethoxy radical from the molecular ion



Figure 11.49 Mass spectrum of ethyl benzoate.

#### 386 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS

to give the very stable phenylacylium ion. Loss of CO from the phenylacylium ion gives the phenyl cation, but due to the instability of the phenyl cation, this pathway is minor. Also observed is a peak at m/e = 122 due to the benzoic acid radical cation resulting from loss of the neutral ethylene molecule from the molecular ion by a different fragmentation process.



1,3-Diphenylpropanone has a molecular ion at m/e = 210 and significant fragment ions of m/e = 119 (M - 15) and m/e = 65. The base peak is m/e = 91. In this example, loss of a benzyl radical from the molecular ion produces an acylium ion (m/e = 119) which rapidly loses CO because the resulting benzyl cation is extremely stable—one of the most stable ions normally encountered. Examination of the mass spectrum (Figure 11.50) shows that there are many additional small peaks present other than those just discussed. Their presence is indicative of the high energy deposited in the molecular ion upon ionization which permits a large number of fragmentations to occur. Nevertheless the fragmentations which occur most often and lead to the most intense peaks are those



Figure 11.50 Mass spectrum of 1,3-diphenylpropanone.

that follow common ideas about reactivity and ion stability.



Both ethers and alcohols readily undergo loss of groups next to the oxygen so as to produce an oxonium ion. Thus *tert*-butyl ethyl ether m/e = 102 has a very large M - 15 peak due to loss of a methyl radical.



The methyl group could be lost from either the *t*-butyl group (path a) or the ethyl group (path b) to give two different oxonium ions with the same m/e value. The base peak at m/e = 57 is the *t*-butyl cation and indicates that at least part of the time the methyl group is lost from the ethyl group (path b) because subsequent loss of formaldehyde from the oxonium ion gives the *t*-butyl cation. The *t*-butyl cation can also be produced by a single fragmentation of the molecular ion by loss of the ethoxy radical. The stability of the *t*-butyl cation makes it the base peak and ensures its production by a variety of routes. This is not to say that all of the M - 15 peak comes from path b, and most likely there is some contribution to the m/e = 87 peak from path a; however, the oxonium ion thus produced is unlikely to fragment into the very unstable ethyl cation.



#### 388 STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS

By working with mass spectral fragmentation patterns, it is possible to develop very keen insight into the ways that molecules disintegrate under high-energy conditions. This permits identification of the structure from the pieces and insight into how they were produced. In conjunction with other structural tools, MS provides invaluable insight into molecular formula and connectivity issues in a molecule and is thus an important tool in structure elucidation.

The foregoing discussion has been a very elementary introduction into MS as a tool for structure identification. Advances in sample introduction, methods of ionization, and ion collection and detection have been remarkable, and today the mass spectra of peptides, nucleic acids, proteins, and other biopolymers are routinely obtained. Using known cracking patterns, MS is the method of choice for identifying drugs and drug testing since it requires only minute quantities (micrograms). It has been sent on the Mars probe to look for amino acids as an indication of life forms on Mars. One goal of current research efforts is to use MS as a method for sequencing peptides and oligonucleotides by their fragmentation patterns. Mass spectrometry is thus an important analytical and structural tool whose evolution continues at a rapid pace. It remains an important component of structural investigation.

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### PROBLEMS

**11.1.** For the following compounds, label the spin systems present, label the symmetry properties of protons where possible (homotopic, enantiotopic,

diastereotopic), and predict the splitting pattern for the proton(s) indicated by an arrow.



**11.2.** Tell how you could use ¹H NMR to distinguish the following pairs of compounds. Be specific as to what data you would look for and how you would interpret it. There might be more than one feature in the ¹H NMR that could be used so give a complete answer.



#### **390** STRUCTURE DETERMINATION OF ORGANIC COMPOUNDS



**11.3.** Tell how you could use ¹³C NMR to distinguish the following pairs of compounds. Be specific as to what data you would look for and how you would interpret it. There might be more than one feature of ¹³C NMR that could be used so give a complete answer.





**11.4.** Tell how you could use IR spectroscopy to distinguish the following pairs of compounds. Be specific as to what data you would look for and how you would interpret it. There might be more than one way to distinguish them by IR so give a complete answer.



392



**11.5.** Tell how you could use mass spectrometry (MS) to distinguish the following pairs of compounds. Be specific as to what data you would look for and how you would interpret it. There might be more than one way to distinguish them by MS so give a complete answer.



**11.6.** Rationalize the major fragmentation pathways observed for ethyl phenylacetate **A** and diethyl phenylmalonate **B**.







**11.7.** Give two instrumental methods that would permit you to distinguish the following. Explain what data you would use and how it would allow you to make the distinction.



# SOLUTIONS TO CHAPTER PROBLEMS

# **CHAPTER 1**



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#### **396** SOLUTIONS TO CHAPTER PROBLEMS



internal bond angles =  $108^{\circ}$ 



### 398 SOLUTIONS TO CHAPTER PROBLEMS

Because the electrons are in an  $sp^2$  orbital which has less s character than the  $e^-$  in the sp orbital in the other compound.

(b)  $H_3CCH_2$ - $\ddot{O}H$ 

Because of being in an orbital of 25% s character compared to 33% s character in the ketone, the e⁻ are less tightly bound and more easily donated to a proton.

(c)  $H_3CC \equiv C^{\Theta}$ 

Because the  $e^-$  are in an sp orbital with greater s character than the lone pair in the sp² orbital of the other compound.



**1.8.** Tautomer **K** is a resonance-stabilized structure as shown; however, tautomer **E** has a principal resonance contributor which is an aromatic species. The need to separate charges decreases the stability somewhat, but the aromatic system is a large stabilizing feature. Thus **E** is of much lower energy than **K** and the equilibrium between **K** and **E** is shifted to **E**.





By using the lone pair on nitrogen, each ring can have six  $\pi$  electrons and thus satisfy Huckel's rule. Thus each ring is an aromatic ring and the molecule can be considered doubly aromatic. The need to separate charges to accomplish this decreases the aromatic stabilization somewhat. Nevertheless the aromatic stabilization makes **B'** a significant resonance contributor. As a consequence **B** has a great deal of charge separation and thus a large molecular dipole moment of 9.6D.



Both bromides give benzylic cations which are resonance stabilized by the benzene rings. The nitrogens of **A** are more electronegative than the carbons of **C** which should destabilize the ion somewhat. The greatest stabilization comes from the fact that the three-membered cationic ring of **A** is a  $2\pi$  aromatic system. This aromatic stabilization of the cation makes its formation much more rapid than the cation from **C**.

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400 SOLUTIONS TO CHAPTER PROBLEMS



Removal of a proton from  $L_1$  gives an anion whose lone pair is orthogonal to the  $\pi$  bond of the carbonyl group by virtue of the rigid geometry of the bicyclic system. Consequently the lone pair cannot overlap with the carbonyl  $\pi$  bond and delocalization via resonance is not possible—it is effectively a localized anion. Removal of a proton from  $L_2$  gives rise to a lone pair in a p orbital which can overlap with the carbonyl  $\pi$  bond and thus resonance delocalization is possible. Thus the anion from  $L_2$  is resonance stabilized and is thus formed more easily.



The bis anion of squaric acid is highly resonance stabilized by four equivalent resonance contributors as well as an aromatic resonance contributor. This results in both protons being easily removed.

0e



Protonation of guanidine gives an ion that is stabilized by four resonance forms, three of which are equivalent. This gives a large amount of resonance stabilization to the protonated form, making guanidine easily protonated and thus a good base.







The cyclopentadiene anion is stabilized by five equivalent resonance structures. The anion is an aromatic anion by virtue of it being a six- $\pi$ -electron system. The indenyl anion is stabilized by a total of seven resonance contributors. However, they are nonequivalent and all but one require that the aromatic cloud of the benzene ring is disrupted. Thus, while the negative charge is well delocalized, the resonance stabilization is less than that of the cyclopentadiene system. Thus the proton is not as easily removed, making indene a weaker acid.

**1.15.** If the CH₂ groups of **1** are not equivalent, then the rate of rotation of the acetyl group is slow so that one methylene group is in the vicinity of the C=O group and the other is in the vicinity of the CH₃ group. Such is not the case for **2**. The rotation of the isopropenyl group is rapid and the methylene groups experience an averaged environment. Because of the electronegativity of oxygen, contributions of **1a** to **1** are more important than the contributions of **2a** to **2**. Thus **1** has greater C-N double-bond character and it is difficult to rotate about that bond. The C-N double-bond character is small in **2**, so rotation is facile, leading to an averaged environment.



### CHAPTER 2

2.1.



402

#### SOLUTIONS TO CHAPTER PROBLEMS







### CHAPTER 2 **405**



















### CHAPTER 3 407



# **CHAPTER 3**

3.1.		
(a)	$NH_3 + OH^- \longrightarrow NH_2^- + H_2O$ 33 $K_{eq} = 10^{-17.3} \qquad 15.7$	Lies to the left
(b)	$CH_{3}CO_{2}H + CO_{3}^{-} \longrightarrow CH_{3}CO_{2}^{-} + HCO_{3}^{-}$ 5 $K_{eq} = 10^{+5.3} \qquad 10.3$	Lies to the right
( <b>c</b> )	$\begin{array}{cccc} \text{RCO}_2\text{H} + \text{R}_2\text{NH} & & & \\ 5 & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & & \\ & & & & \\ & & &$	Lies to the right
(d)	$R_{3}COH + RC(O)CHR' \xrightarrow{\Theta} R_{3}CO^{-} + RC(O)CH_{2}R'$ $18 \xrightarrow{K_{eq} = 10^{+2}} 20$	Lies to the right
(e)	NaF + HCl NaCl + HF -6 $K_{eq} = 10^{+2.8}$ -3.2	Lies to the right
( <b>f</b> )	$\begin{array}{rrr} \text{ArOH} + \text{RNH}_2 & \xrightarrow{K_{eq}=1} & \text{ArO}^- + \text{RNH}_3^+ \\ 10 & 10 \end{array}$	Lies in middle
(g)	$RC(O)OR' + H_2SO_4 \longrightarrow RC(OH^+)OR' + HSO_4^-$ -9 -6.5	Lies to the right
( <b>h</b> )	$\begin{array}{c} \text{RC}(\text{O})\text{CH}_2\text{C}(\text{O})\text{R}' + \text{R}_3\text{N} & & \\ 9 & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ $	Lies to the right
(i)	$RC \equiv C^{-} + RCH_{2}OH  RC \equiv CH + RCH_{2}O^{-}$ $16 \xrightarrow{K_{eq} \equiv 10^{+9}} 25$	Lies to the right

408 SOLUTIONS TO CHAPTER PROBLEMS

(j) 
$$CH_3Li + (CH_3)_2C=CH_2 \longrightarrow CH_4 + (CH_3)_2C=CHLi$$
  
44 50 Lies to the right

3.2. (a)

**(b)** 

$$\begin{array}{c} O \\ CH_3 \\ H_2 \\ H_3 \\ H_4 \\ H_2 \\ H_4 \\ H_4 \\ H_5 \\$$



$$(CH_3)_2CH_2NH_3^+ + H_2O \xrightarrow{K_{eq} = 10^{-11.7}} (CH_3)_2CH_2NH_2 + H_3O^+$$
Left  
10 -1.7

(c) 
$$O^- + CH_3 \longrightarrow O^+ \xrightarrow{K_{eq} = 10^{-6}} OH + CH_2 \longrightarrow OH^+ CH_2 \longrightarrow$$

(d)  

$$CH_{3}NO_{2} + -CH_{2} \stackrel{O}{\coprod}_{H} \stackrel{K_{eq} = 10^{+9}}{\longleftarrow} -CH_{2}NO_{2} + O_{CH_{3}} \stackrel{O}{\coprod}_{H}$$
Right

(e)  
F-CH₂CO₂⁻ + Cl-CH₂CO₂H 
$$\xrightarrow{K_{eq} = 10^{-0.2}}$$
  
2.86 F-CH₂CO₂H + Cl-CH₂CO₂⁻  
2.66 Left-very slig

Left—very slightly Fluorine has a larger inductive effect than chlorine.

(f)  

$$\begin{array}{c}
H \\
NH_{3}^{+} + HCO_{3}^{-} \xrightarrow{K_{eq} = 10^{-3.6}} \\
10 \\
\end{array} \begin{array}{c}
H \\
NH_{2} + H_{2}CO_{3} \\
6.4 \\
\end{array}$$
Left

(g) 
$$\begin{array}{c} + & O \\ & \downarrow \\ C_{2}H_{5} \\ \hline -6 \\ \end{array} \\ \begin{array}{c} H \\ -6 \\ \end{array} \\ \end{array} \\ \begin{array}{c} H \\ + \\ C_{15} \\ -6 \\ \end{array} \\ \begin{array}{c} H \\ + \\ C_{10}H_{10} \\ \hline \\ C_{2}H_{5} \\ \end{array} \\ \begin{array}{c} O \\ C_{2}H_{5} \\ \hline \\ C_{2}H_{5} \\ \end{array} \\ \begin{array}{c} O \\ C_{15} $

(h) 
$$O$$
  
 $+ (C_6H_5)_3C^ \xrightarrow{K_{eq}=10^{+23}}$   $O^-$  Left  
17 (conjugation in enolate)  $40$ 

CHAPTER 3 409

- **3.3.** (a) 1 < 4 < 2 < 3
  - **(b)** 2 < 1 < 3 < 4
    - (c) 4 < 1 < 2 < 3
    - (d) 4 < 2 < 3 < 1
    - (e) 1 < 4 < 3 < 2
    - (f) 4 < 2 < 3 < 1
    - (g) 1 < 4 < 3 < 2
    - (h) 3 < 2 < 1 < 4(i) 1 < 3 < 2 < 4
    - (I) 1 < 3 < 2 < 2
    - (j) 4 < 1 < 2 < 3</li>
      (k) 4 < 2 < 3 < 1</li>
- **3.4.** (a) Amides are protonated on oxygen because a resonance-stabilized cation is produced. Protonation on nitrogen gives a localized cation.



- (b) The lone pairs on the ether oxygen are in sp³ while those of the ketone are in sp² orbitals. Thus the e⁻ pairs of the ketone are more tightly bound and less able to be donated, making the ketone the weaker base.
- (c) This is because the conjugate acid of tetramethylguanidine has much greater resonance stabilization than the N,N-dimethyl amide (more forms and equivalent forms.)



(d) The amount of e⁻ donation to boron by resonance is much greater for oxygen than it is for fluorine because of fluorine's greater electronegativity. Therefore the boron of trimethyl borate is more e⁻ rich and it is a poorer Lewis acid



#### 410 SOLUTIONS TO CHAPTER PROBLEMS

- (e) The lone pair of piperidine is in an sp³ orbital while the lone pair of pyridine is in an sp² orbital. The greater s character causes the e⁻ to be more tightly bound and less able to be donated to H⁺.
- (f) The conjugate base of *p*-nitrophenol is resonance delocalized into the nitro group. The negative charge is thus more stable and the conjugate acid is more acidic.



- (g) The inductive effect of the chlorine in the ortho position is greater than that of the para chloro group because it is closer to the amino group. The greater inductive effect removes more e⁻ density from nitrogen and makes the ortho isomer a weaker base.
- (h) BF₃ complexes with the lone pair of the imine and makes it much more electron deficient and thus electrophilic. In the complexed form it reacts with NaBH₄ because it is a much better electrophile

**3.5.** 
$$H_2S \xrightarrow{H_2O}_{pK_a=7.0} HS^- + H_3O^+ H_2O \xrightarrow{H_2O}_{pK_a=15.74} HO^- + H_3O^+$$

Oxygen is more electronegative than sulfur so the hydroxide ion should be more stable than the hydrosulfide anion. This should result in water being more acidic. However, the hydrogen-sulfur bond is nearly 40 kcal/mol weaker than the hydrogen-oxygen bond. Thus the energy cost of bond breaking for hydrogen sulfide is much less than for water. This overshadows the effect of electronegativity and makes hydrogen sulfide more acidic.

Compare the conjugate bases. Since sulfur is slightly more electronegative than carbon and is bonded to two other oxygens rather than one, the inductive effect predicts benzenesulfonic acid should be stronger. But the huge difference in acidity is much more than can be explained by an inductive effect argument. The sulfonate ion has three equivalent resonance forms while the carboxylate group has only two. This could explain the difference. However, the sulfur in such systems is actually tetrahedral so any resonance argument must involve overlap with the d orbitals on sulfur. Another bonding picture is two coordinate covalent bonds from sulfur to oxygen. This leaves sulfur with a formal +2 charge right next to the anionic oxygen. Since these are merely a different electron, this could simply be considered an additional resonance form. Thus the greater resonance stabilization of the sulfonate anion makes the conjugate acid very acidic.



Protonation at C-2 gives a resonance-stabilized (although nonaromatic) cation. Protonation at nitrogen destroys the aromaticity of the pyrrole ring by taking the lone pair out of the aromatic system. At the same time the cation produced is not resonance stabilized. Thus protonation on C-2 is favored.

3.8.



3.9. Compare with



In each case a benzoate ion is produced. Substituents which stabilize the benzoate ion would make the benzoic acid more acidic and the  $pK_a$  decreases. Substituents which stabilize the benzoate ion should also cause it to ionize from the allyl group more easily (faster), and thus the rate of ionization would go up. Thus substituents which make the benzoate ion more stable cause the  $pK_a$  to go down and the rate of ionization to go up. The reverse is also true—substituents which make the benzoate ion less stable would raise the  $pK_a$  but lower the rate of ionization. The two are thus inversely proportional.

412 SOLUTIONS TO CHAPTER PROBLEMS

### **CHAPTER 4**









### CHAPTER 5

- 5.1. (a) The positive sign of  $\rho$  shows that an increase in electron density occurs during the transition state of the rate-determining step. The magnitude shows that there is a considerable electron density increase. Since the reagent responsible is the base ethoxide, it is pretty clear that significant benzylic proton removal by the base is occurring in the activated complex of the rate-determining step. Moreover proton removal is well advanced.
  - (b) This is a primary kinetic deuterium isotope effect, consistent with the notion that removal of the benzylic proton by base is taking place in the rate-determining step.
  - (c) This is also a primary kinetic isotope effect; only this is a carbon isotope effect. This means that cleavage of the carbon-nitrogen bond is occurring at the transition state of the rate-determining step. Since both proton removal and loss of the leaving group are occurring at the transition state, this then seems to be a concerted E2-type elimination.
  - (d) These results show that there is a distinct stereoelectronic requirement for this process—that the proton and the leaving group are antiperiplanar. This is typical of an E2-type base-promoted elimination. The major product is the result of the deuterium isotope effect, which results in faster removal of the proton than the deuterium.
- **5.2.** The positive value of  $\rho$  is indicative of increasing electron density on the benzylic carbon at the transition state of the rate-determining step. This is consistent with a mechanism in which the methoxide is acting as a base to remove the benzylic proton in the rate-determining step. The small value suggests that proton removal is not far advanced.

#### 416 SOLUTIONS TO CHAPTER PROBLEMS

The second case with the negative value of  $\rho$  indicates that in this system a decrease in electron density occurs at the transition state. Ionization of the chloride leaving group to give a carbocation is consistent with this  $\rho$  value. The modest magnitude is consistent with the fact that charge formation occurs at the  $\beta$  position relative to the ring so the effect of substituents on charge formation is small. Also the correlation with  $\sigma$  is consistent since there is no direct resonance interaction of the developing charge with the aromatic ring due to the insulating methylene group.



5.3.



The positive sign, magnitude, and correlation with  $\sigma^-$  indicate that there is a significant increase in electron density on an atom in conjugation with the aromatic ring. That would be the sulfur atom attached to the ring that is substituted. The primary kinetic deuterium isotope effect shows that cleavage of the C–H bond occurs in the rate-determining step. The best mechanism is a concerted, base-promoted elimination





Since the lower pathway has proton removal as an integral part of the mechanism, you could use isotopic substitution in two ways. First make the deuterated compound **D**. The kinetic deuterium isotope effect should be primary for the lower path but very near 1 for the upper path. Moreover, if you looked at the product, there should be no deuterium left in the product if the lower path is followed, whereas deuterium should be retained in the product if the addition elimination path is followed.

#### 5.5.

Addition-elimination



Ionization-addition



In the first case the  $\rho$  value will be positive because of the increase in electron density on the nitrogen next to the aromatic ring and may correlate better with  $\sigma^-$  than with  $\sigma$  because of conjugation with the ring. In the second case, ionization would lead to a decrease in electron density on the nitrogen attached to the ring and would thus give a negative r value.

The better correlation with  $\sigma^-$  indicates that increasing electron density in conjugation with the aromatic ring is involved in the rate-determining step. For the two mechanistic steps of the hydrolysis, both should have positive  $\rho$  values but only the second has charge development in conjugation with the ring. Thus for this hydrolysis breakdown of the tetrahedral intermediate is rate determining.

5.7.

5.6.

$$Z \xrightarrow{N} N \xrightarrow{N} Z \xrightarrow{N} N \xrightarrow{NaN_3} Z \xrightarrow{N} N \xrightarrow{NaN_3} Z \xrightarrow{N} N \rho^+ = -1.03$$

The correlation with  $\sigma^+$  and the negative slope indicate charge deficiency formed in conjugation with the ring. This in turn suggests an ionization mechanism in which ionization of bromide gives a cation. This cation might be particularly favored because it is an aromatic cation (Huckel) and would also explain the relatively low  $\rho^+$  value. The  $\rho^+$  is low because the product cation is stabilized; thus the transition state is early with little charge development.

418 SOLUTIONS TO CHAPTER PROBLEMS





The difference in activity for the protio and deutero compounds is likely a kinetic isotope effect of the transformation into the active compound. Since that involves removal of the methine proton, a reasonable deduction is that the ketone is the active species.

**5.9.** Steady state assumption:

$$\frac{dP}{dt} = k_2 \text{ [complex]}$$

$$\frac{d[\text{complex}]}{dt} = 0 = k_1 [\text{PhCHO}][\text{PFC}] - k_1 [\text{complex}] - k_2 [\text{complex}]$$

$$[\text{complex}] = \frac{k_1 [\text{PhCHO}][\text{PFC}]}{k_1 + k_2}$$
This should be first order in aldehyde also.

The kinetic deuterium isotope effect indicates that C–H bond cleavage is part of the rate-determining step and the negative  $\rho$  value shows that there is electron deficiency being produced on the benzylic carbon in the activated complex. Therefore



The correlation with  $\sigma^-$ , the positive slope, and the relatively large magnitude of  $\rho^-$  means that a large charge density is built up on the ring in the activated complex of the rate-determining step. Addition of methoxide to give the Meisenheimer complex could account for this.

5.12.

5.11. Using the symbols A, I, and P,

$$\frac{d\mathbf{P}}{dt} = k_2[\mathbf{I}]$$

$$\frac{d\mathbf{I}}{dt} = k_1[\mathbf{A}][\mathbf{B}\mathbf{r}_2] - k_{-1}[\mathbf{I}][\mathbf{B}\mathbf{r}^{-1}] - k_2[\mathbf{I}] = 0 \quad (\text{steady-state approximation})$$

$$[\mathbf{I}] = \frac{k_1[\mathbf{A}][\mathbf{B}\mathbf{r}_2]}{k_{-1}[\mathbf{B}\mathbf{r}^{-1}] + k_2}$$

$$\frac{d\mathbf{P}}{dt} = \frac{k_2k_1[\mathbf{A}][\mathbf{B}\mathbf{r}_2]}{k_{-1}[\mathbf{B}\mathbf{r}^{-1}] + k_2}$$

$$\begin{array}{c} OH\\ PhCCH(CH_3)_2 & \xrightarrow{k_1} & OH\\ PhC = C(CH_3)_2 & K_{eq} = \frac{k_1}{k_{-1}} \end{array}$$

$$\begin{array}{c} \mathbf{C} = \mathbf{O} & \mathbf{E}\\ OH\\ PhC = C(CH_3)_2 & + Cr(VI) & \xrightarrow{k_2} \end{array} \text{ products}$$

E

Using the above abbreviations

$$\frac{d\mathbf{P}}{dt} = k_2[\mathbf{E}][\mathrm{Cr}(\mathrm{VI})] \qquad \frac{d\mathbf{E}}{dt} = 0 = k_1[\mathbf{C}=\mathbf{O}] - k_{-1}[\mathbf{E}] - k_2[\mathbf{E}][\mathrm{Cr}(\mathrm{VI})]$$
$$[\mathbf{E}] = \frac{k_1[\mathbf{C}=\mathbf{O}]}{k_2[\mathrm{Cr}(\mathrm{VI})] + k_{-1}}$$
$$\frac{d\mathbf{P}}{dt} = \frac{k_2k_1[\mathbf{C}=\mathbf{O}][\mathrm{Cr}(\mathrm{VI})]}{k_2[\mathrm{Cr}(\mathrm{VI})] + k_{-1}}$$
$$k_{-1} >>> k_2, \text{ preequilibrium: } \frac{d\mathbf{P}}{dt} = k_2K_{\mathrm{eq}}[\mathbf{C}=\mathbf{O}][\mathrm{Cr}(\mathrm{VI})]$$

Р

If  $k_{-1} >>> k_2$ , preequilibrium:  $\frac{d\mathbf{I}}{d\mathbf{t}} = k_2 K_{eq}[\mathbf{C=O}][Cr(VI)]$ If  $k_2 >>> k_{-1}$ , first step rate determining:  $\frac{d\mathbf{P}}{dt} = k_1[\mathbf{C=O}]$ If  $k_1 \approx k_{-1} \approx k_2$ , then the overall expression describes the rate.

5.13. Mechanism could be



If the protonation were rate determining, then a decrease in electron density at the carbonyl carbon occurs. The  $\rho$  value should be negative. If the second step is rate determining, then nucleophilic addition should result in an increase in electron density at the carbonyl group. the  $\rho$  value should be positive. The data show that the second step is probably rate determining. Moreover the lack of reaction under basic conditions means that the carbonyl group must be activated by protonation for the cyanide to add. Thus the protonation step is probably an equilibrium step.
#### 420 SOLUTIONS TO CHAPTER PROBLEMS

- **5.14.** (a) Since a base is used in the reaction and the base apparently removes the  $\alpha$  proton, there should be a primary kinetic deuterium isotope effect of >1.5. For this to be true, proton removal must occur in the rate-determining step.
  - (b) In this case the base removes a proton from the molecule but the proton that is removed is not the one for which isotopic substitution has been done. Thus there should be only secondary isotope effects for this reaction of <1.5 (actually about 1.1).
  - (c) In this reaction a proton is lost from the molecule in order to form the product and that proton is one for which isotopic substitution is indicated. However, since there is no good base present to cause its removal, it is likely that the rate-determining step is ionization of the leaving group to give a carbocation. The proton is lost from this cation in a subsequent step. Since proton loss occurs after the rate-determining step, no primary kinetic deuterium isotope effect will be seen. Thus the isotope effect will be <1.5. Actually there is a secondary kinetic deuterium isotope effect of about 1.08.
- **5.15.** (a) Since both correlate with  $\sigma^+$ , there is a development of positive charge on the benzylic carbon which can be delocalized into the aromatic ring. The lower  $\rho$  value for **B** means that less electron deficiency is produced on the benzylic carbon of **B** than **A**.
  - (b) Since both of these tertiary, benzylic substrates solvolyze by ionization as the rate-determining step, the difference in the transition state parameters suggests that the transition state for the *tert*-butyl case **B** is much earlier. One reason could be that the product ion from **B** is more stable than the product ion from **A**, which would result in an earlier transition state by the Hammond postulate. This is not a likely explanation since both are tertiary benzylic carbocations as mentioned earlier. The difference in  $\rho$  values is probably related to steric factors. Since the bulky *t*-butyl groups tend to flatten out the normal tetrahedral geometry, **B** is strained relative to **A** and thus of higher energy. (This effect is actually termed B strain.) Thus the products are of similar energy but the reactants are not. The Hammond postulate predicts that the reactant of higher energy, **B**, will have an earlier transition state, as is seen.

Small composite  $\rho$  value of some (–) and some (+) steps.



Large positive  $\rho$  value means that enolate formation is rate-detemining step.



Positive  $\rho$  value consistent with nucleophilic addition to epoxide which increases electron density on oxygen. Small magnitude consistent with negative charge being insulated from the ring.



Large negative  $\rho^+$  value consistent with formation of positive charge on the benzylic carbon. Thus epoxide ring opening (of protonated epoxide) is the rate-determining step.





#### 422 SOLUTIONS TO CHAPTER PROBLEMS



6.4. (a) Both 1S, 2R, conformational isomers(c) Enantiomers

- (e) Identical
- (g) Conformational diastereomers
- (i) Enantiomers

- (b) Syn-anti, diastereomers
- (d) Enantiomers
- (f) Diastereomers
- (h) Enantiomers
- (j) Conformational isomers





 $\Delta G = -0.7$  kcal/mol

CHAPTER 6 **423** 



Because of H bonding in the diequatorial form,  $\Delta G$  includes the energy required to break the H bond. Thus the actual  $\Delta G$  is greater than that calculated.



424

SOLUTIONS TO CHAPTER PROBLEMS



If it is stepwise, then the amide product would be racemic; if concerted, then optically active.

6.9.









only anti elimination possible — much faster

6.10.





For anti elimination, the trans compound only has a single trans diaxial conformer and that is very little populated. Elimination is slow and gives nonconjugated product



The cis compound has a much more populated trans diaxial conformer and the elimination can give the more stable conjugated product. Elimination is much faster



#### 426 SOLUTIONS TO CHAPTER PROBLEMS

- **6.14.** The origins of strain in three-membered rings could be both angle strain and torsional strain. Since lone pairs are effectively smaller than bonds to hydrogen, replacing the C–H bonds in cyclopropane by one lone pair in aziridine or by two lone pairs in oxirane should reduce torsional strain. Since these changes do not change the strain, it is clear that strain in three-membered rings is due entirely to angle strain.
- **6.15.** Puckering in four-membered rings is due to the molecule relieving torsional strain. In doing so, some additional angle strain is introduced but the net result is the most stable structure. Replacing the C–H bonds by lone pairs which are smaller than C–H bonds lessens the torsional strain so the molecule flattens to reduce angle strain.
- **6.16.** Normally the exocyclic bonds (i.e., the C–H bonds) of cyclopropane are greater than  $109^{\circ}$  and in fact approach  $120^{\circ}$ . This is because the ring bonds have greater p character to accommodate the smaller angle and the external bonds have greater s character. The greater s character causes the angles to be greater than  $109^{\circ}$ . Adding a spiro ring now forces the spiro carbon to have exocyclic bond angles of less than  $109^{\circ}$ . This adds increased strain to the system such that the strain is greater than to single cyclopropyl rings (2 × 27.3 kcal/mol = 54.6 kcal/mol).

6.17.



Inspection of the two sets of chair structures reveals that in one compound the allequatorial conformer is overwhelmingly favored. In the other compound both chair structures have comparable energies so both will be populated significantly. In the all-equatorial isomer, the carbon-chlorine bond dipole moments reinforce one another leading to a large molecular moment. In the other compound the chlorines are both equatorial part of the time but part of the time they are trans diaxial where the carbon-chlorine bond dipole moments tend to cancel one another. Thus the average dipole moment of these two conformations will be less than the first compound, which exists virtually completely in the all-equatorial conformer.

6.18.



In the cyclohexane there is one axial methyl in either conformation; thus the two conformations are of equal energy and will be equally populated. Conformer **1** has a 1,3 diaxial methyl-proton interaction and a 1,3 interaction between the methyl group and the carbonyl group. Conformer **2**, on the other hand, has two 1,3 diaxial methyl-proton interactions. Since the carbonyl group is somewhat smaller than a

 $CH_2$  group, conformer 1 has slightly smaller 1,3 diaxial interactions and therefore will be favored slightly.

**6.19.** The fact that only cis product is produced from either 2 or 4 means that both oxygens in the product come from the osmium reagent. This could result from either a concerted, 3 + 2 cycloaddition of  $OsO_4$  to the double bond or a sequential addition of one then another oxygen to the same side of the double bond.



Comparison of 2 and 4 shows that the facial preference is for exo attack. Only when the exo face is blocked by a methyl group does the reagent attack the endo face.

**6.20.** Knowing that ultimately two hydroxyl groups are added to each end of the double bond, one can construct the possible isomers easily.





By inspection one can see that the the 2R,3S and 2S,3R isomers are enantiomers and the 2R,3R and 2S,3S pair are enantiomers. As they are drawn, one can see the original E stereochemistry of the double bond. Thus the first pair comes from a stereospecific trans addition across the double bond while the second pair comes from a stereospecific cis addition. Clearly the stereochemistry of the double bond is maintained throughout the addition. Since iodine is the electrophile, a bridged iodonium ion is likely responsible. The initial addition gives only a single pair of enantiomers since the only nucleophile is benzoate. Also noted is that iodine is not in the product so something must replace the iodine. The silver ion would help to remove the iodine. Since iodine would cause a trans addition and the product has oxygens added trans, there must be a replacement of iodine by an

### 428 SOLUTIONS TO CHAPTER PROBLEMS

oxygen ligand with retention of configuration at the iodine center. Neighboring group participation is responsible.



The other set of enantiomers must result from inversion of configuration. The presence of water must prevent the benzoyl group from acting as a neighboring group.



- (a) From the products it is clear that a hydrogen and a hydroxyl have added in a syn fashion to either face of the olefin.
- (b) Since a hydrogen and a boron add in the first step and then the boron ligand is converted to an oxygen ligand in the second step, the possibilities are (a) mode of addition *syn-anti* and (b) stereochemistry of cleavage *retention-inversion*. Picking only the lower face for the initial addition step, the four stereochemical possibilities are



(c) Since the actual product has the hydroxyl group and methyl groups cis to one another, the syn retention and anti inversion sequences are the only possible ones. If it is known that cleavage occurs with retention, then the hydroboration reaction must occur by syn addition across the double bond.



From the stereochemistry shown, the zinc reduction is a trans elimination of two bromines. Since the addition of bromine to an olefin is also trans, you could not use this sequence to isomerize an olefin.

### **CHAPTER 7**

- **7.1.** (a)  $1^{\circ}$  Alcohol  $\rightarrow$  aldehyde, [O] = +2, PCC or Swern oxidation
  - (b) Ester  $\rightarrow$  ether, [H] = -4 i. LAH ii. NaH, CH₃1
  - (c) Acid → amide, no change
    i. SOCl₂, DMF
    ii. CH₃NHCH₂CH₃

```
430
          SOLUTIONS TO CHAPTER PROBLEMS
          or
          i. DCC, Et<sub>3</sub>N, CH<sub>3</sub>NHCH<sub>2</sub>CH<sub>3</sub>
    (d) Ketone \rightarrow methylene compound, +2 \rightarrow 0 [H]
          i. Ph<sub>3</sub>P=CH<sub>2</sub> (Wittig) or Me<sub>3</sub>SiCH<sub>2</sub>-Li (Peterson)
    (e) 1^{\circ} Alcohol \rightarrow aldehyde, -1 \rightarrow +1, PCC or Swern oxidation
     (f) Lactone \rightarrow lactol (hemiacetal); +3 \rightarrow +1; DIBAH, toluene, -78^{\circ}
    (g) 2^{\circ} Alcohol \rightarrow 2^{\circ} amine, no change
          i. H<sub>2</sub>CrO<sub>4</sub>, acetone
          ii. CH<sub>3</sub>NH<sub>3</sub>, NaBH<sub>3</sub>CN, H<sup>+</sup>
    (h) Acid \rightarrow ester, +3 \rightarrow +3
          i. DCC, Et<sub>3</sub>N, EtOH
          or
          i. oxallyl chloride
          ii. EtOH, py
     (i) Alkene \rightarrow alcohol; 0, -2 \rightarrow +1, -3 no change
          i. HgOAc<sub>2</sub>, H<sub>2</sub>O
          ii. NaBH<sub>4</sub>
     (i) Alkene \rightarrow alcohol; -1, -2 \rightarrow -2, -1 no change
          i. BH3-THF
          ii. NaOH, H<sub>2</sub>O<sub>2</sub>
    (k) Ketone \rightarrow alcohol + cyclization, +2 \rightarrow 0 [O]
          i. NaBH<sub>4</sub>
          ii. TsOH, benzene, \Delta
     (I) Ketone \rightarrow acetal; no change; HOCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>OH, TsOH, toluene, \Delta
   (m) Acid \rightarrow alcohol \rightarrow acetylation, +3 \rightarrow -1 [H]
          i. BH<sub>3</sub>-THF
          ii. Ac<sub>2</sub>O, py, DMAP
    (n) Alkene \rightarrow alcohol, no change
          i. HgOAc2, H2O
          ii. NaBH4, MeOH
    (o) Ketone \rightarrow \alpha, \beta-unsaturated ketone; -2, -3, \rightarrow -1, -2, [O]
          i. LDA
          ii. PhSeSePh
          iii. H<sub>2</sub>O<sub>2</sub>
          or
          i. py^-H^+ Br_3^-
          ii. K<sub>2</sub>CO<sub>3</sub>
    (p) 2^{\circ} Alcohol \rightarrow acetal, 0 \rightarrow +2, [O]
          i. H<sub>2</sub>CrO<sub>4</sub>
          ii. CH<sub>3</sub>OH, H<sup>+</sup>, \Delta
    (q) trans-Alkene \rightarrow alkyne; -1, -1 \rightarrow 0, 0 [O]
          i. Br<sub>2</sub>, CCl<sub>4</sub>
          ii. NaNH<sub>2</sub>
```









7.3.







### 436 SOLUTIONS TO CHAPTER PROBLEMS



## **CHAPTER 8**

8.1.





#### 438











**8.2.** For each part the first structure is nucleophile and the second is electrophile.



CHAPTER 8 441







**(p)** E





8.3.



**(b)** 

CH₃O



CH₃O













SOLUTIONS TO CHAPTER PROBLEMS



**8.6.** The formation of medium rings requires that the two reacting double bonds be in proximity to one another and that dimerization is controlled by high dilution. Since **6A** does not cyclize under high dilution, the rate of cyclization is slow compared to dimerization. Placement of a much larger cyclohexyl group on nitrogen causes cyclization to proceed much faster, assuming that the dimerization rate is similar for both, which is a fair assumption. The cyclohexyl group must cause a conformational change about the urea linkage that causes the two double bonds to be in close proximity, thus increasing the cyclization rate greatly.

### **CHAPTER 9**

9.1.



CHAPTER 9 447





SOLUTIONS TO CHAPTER PROBLEMS



#### 450 SOLUTIONS TO CHAPTER PROBLEMS

- 9.2. (a) exo-5-membered cyclization favored over endo-6.
  - (b) Tough call, but hindered nature of the internal olefin position where cyclization must occur should retard cyclization. Reduction prevails.
  - (c) Approach to molecule by the tin radical and hydrogen donor takes place from the least hindered side away from the methyl groups.
  - (d) Cyclization occurs preferentially to the unsaturated ester because the product radical is resonance stabilized.
  - (e) Exo-5 cyclization is favored because it is faster than exo-6 cyclization because the olefin is closer to the radical center.
  - (f) Five-membered ring formation is faster than six and six-membered ring formation will also be more sterically hindered due to the carboethoxy group. The product radical is resonance delocalized so a mixture is probably formed. However, the conjugated product would probably be favored slightly.
  - (g) This transition state for ring closure minimizes steric interactions and is favored. It leads to the trans, trans stereochemistry in the product radical. The trans stereochemistry between the radical and the double bond prevents further cyclization and reduction takes place to give the monocyclic product.



### **CHAPTER 10**

(There are many correct alternate answers)

10.1.





(c)







(**d**)















SOLUTIONS TO CHAPTER PROBLEMS



⁴⁵⁴ 

0

Η





 $\Rightarrow$ 

CH₃C





(**t**)







10.3.



Epoxide opening gives the needed trans isomer. Reduction of an alkylated cyclohexanone gives mostly the cis.
#### 456

SOLUTIONS TO CHAPTER PROBLEMS



(c)









EtO



(i)

















- -

460

SOLUTIONS TO CHAPTER PROBLEMS















CHAPTER 11 463



464

#### SOLUTIONS TO CHAPTER PROBLEMS



The benzylic protons of the ABX signal should be downfield at  $\approx 2.5$ . The methine signal should be further split by the amide NH.

The methylene protons of the ABX signal should be <2.4 and the methine signal should not be split by the amino NH.

465 CHAPTER 11



There should be a 4H signal for the  $\alpha$  protons at  $\approx 2.2$  ppm.



This has the A2X3 ethyl group prominent and the methine proton will be a pentet by equivalent splitting by the equivalent neighboring protons.



for the axial  $\alpha$  protons at  $\approx 2.4$  ppm and a 2H multiplet further upfield.

and

This has no ethyl signals and the methine signals will be a multiplet due to nonidentical splitting by the ring H's.



**(b)** 

(c)



 $CH_3$ 

Off-resonance line methine signal is a doublet.

H₃C_{CH2}O

be more upfield.



The methylene group (determined by off resonance) will be further upfield and the methyl group will be more downfield.



The methylene group (determined

downfield and the methyl group will

by off resonance) will be further







Acetal carbon should be further downfield and give singlet in offresonance decoupled expt.



Ether methine carbons should be further upfield and give doublets in off resonance.

#### 466

#### SOLUTIONS TO CHAPTER PROBLEMS



Unsaturated ketone 1690 cm⁻¹



468

#### SOLUTIONS TO CHAPTER PROBLEMS



(**d**)





Would have a big M - 44 peak

due to O-O cleavage and loss

Would have big M – 31 peak. Base peak is 105

and



Would have big M-15 peak

due to 3• cation

and very large peak at m/e = 98



Would have very small M - 15 peak, large M - 45 peak, and the *m/e* peak at 98 would be very small.



C1



No  $Cl_2$  isotope pattern in parent ion, M – 28 should be

large, and a M-15 should be

CH₃

Very large parent ion and M - 35would be about it. Also the typical pattern for two-chlorine isotope would be very apparet in the parent ion.





and

Base peak at m/e = 91.

Smaller m/e = 91, large M – 1 peak.

apparent.





A



and



¹H NMR: no methyl triplet upfield.

(b) H O H₃C H and

MS: large M-1 peak but little M-29 peak.

IR: conjugated so aldehyde and olefin are at lower frequency.

¹H NMR: 2H vinyl signals with J > 12 due to trans.

(c)

and

¹³C NMR: 4 lines.

¹H nmr: normal methylene resonances.

IR has higher C=O frequency due to e⁻ withdrawing group.

MS: much more M-29 due to formation of allylic cation.

IR: isolated so normal aldehyde and olefin stretches.

¹H NMR: 3H vinyl signal which is ABX.

¹³C NMR: 6 lines.

¹H NMR: this has a 2H methylene signal downfield due to steric deshielding.

(e)

#### 470 SOLUTIONS TO CHAPTER PROBLEMS



 $\bigcirc O_{\mathbb{N}} \longrightarrow O$  CH₂CH₃

IR: carbamate C=O 1680 cm⁻¹.

¹H NMR: methoxy singlet at 3.5 ppm.

¹³C NMR: C=O carbon 160 ppm.



IR: no terminal CH at ≈3300 cm⁻¹ conjugated ester and acetylene.

¹H NMR: isopropyl septet at 3.6 ppm. MS: large M-59, M-39.



¹³C NMR: no C=O signal.



IR: terminal CH at  $\approx 3300 \text{ cm}^{-1}$ , normal ester at 1735 cm⁻¹.

¹H NMR: isopropyl septet at 2.3 ppm. MS: large M-43, M-55.

# INDEX

AB quartet, 355 Acetoacetic ester synthesis, 235 Acetylide nucleophiles, 232, 305 Achiral, 128 Acid-base equilibrium constants, 51 Acid chlorides, preparation from acids, 191 Acidity electronegativity effect, 58 inductive effect, 59 resonance effect, 61 structure reactivity effects, 57 Acid strength, 50 Activated complex, 88 charge development, 89, 110, 112 energy, 97 lifetime, 89 structure, 91 Activation energy, 87 Activation enthalpy,  $\Delta H^{\ddagger}$ , 105 Activation entropy,  $\Delta S^{\ddagger}$ , 105 Acyl anion equivalent, 302, 303 Acyl anion synthon 1,3-dithiane anions, 303 cyanohydrins, 304 nitronate anion, 304 Acyl substitution, 39 Addition reaction, 37 AIBN, see Azobisisobutyronitrile Alcohols by hydroboration, 200 by hydroxymercuration, 200 from aldehydes and ketones, 199

from carboxylic acids, 199 preparation, 198 Aldehyde derivatives, 198 Aldehydes from Swern oxidation, 193 preparation, 192 Aldol condensation, 228 diastereoselection, 236 Alkanes by Clemmensen reduction, 208 by Wolff-Kishner reduction, 207 from lithium organocuprate coupling, 208 oxidation level, 34 preparation, 207 Alkenes by 1,2-eliminations, 204 oxidation level, 34 preparation, 203 regioisomers, 204  $\alpha,\beta$ -unsaturated carbonyl electrophiles, 223  $\alpha$ -diazoketones for cyclopropanation, 245 Amides, preparation, 190 Amines, preparation, 201 Amphoteric compounds, 56 Angle strain, 169 Anisotropy, 342 anti addition, 148 anti conformation, 160 Antiaromatic compounds, 24 Antibonding molecular orbitals, 14

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#### 472 INDEX

Antibonding orbitals, 14 Aromatic stabilization, 23 Aromaticity, 23 Arrhenius activation energy,  $E_a$ , 104 Arrhenius equation, 104 Asymmetric center, 129 Atom transfer to free radicals, 74, 273 Atomic mass units, 379 Atomic orbitals (AO), 5 overlap, 6 Atomic spin. 336 A-values, 166 table, 167 Axial position, 162 Azobisisobutyronitrile (AIBN), 275, 280 Baeyer strain, 169 Barton deoxygenation, 282 Boat conformation, cyclohexane, 162 Bond breaking, homolytic, 275 Bond disconnection in retrosynthetic analysis, 296 Bond force constant, 368 Bond making, homolytic, 273 Bond order, 19 Bond polarity, 71, 370 Bond strength, 71, 370 Bonding molecular orbital, 6 Boron enolate, 238 Boronate esters, Suzuki coupling, 253 Bridged intermediates, 81 Bridging interaction, 79 Broad-band proton decoupling, 361 Bronsted acid, 47 Bronsted base, 48 Cahn-Ingold-Prelog nomenclature, 130 Carbon-centered electrophiles, 220 Carbon-centered free radicals, 273, 274 properties, 273 Carbon-centered nucleophiles, 218 Grignard reagents, 218 organolithiums, 218 ¹³C chemical equivalency, 364 ¹³C chemical shifts, 361 table, 362 Carbon electrophiles, by Lewis acid complexation, 221 ¹³C NMR, 360

Carbon oxidation level, 33 Carbonyl electrophiles, 222 Carboxylic acids, preparation, 185 Cationic carbon, as electrophile, 220 Cationic rearrangements, 81 Chair conformation, cyclohexane, 161 Chemical equivalence, 350 Chemical shift.  $\delta$ , 338 Chemospecificity, 294 Chiral center, 128, 129 Chiral chromatography, 142 Chiral object, 128 Chiral pool, 144 Claisen condensation, 229 Classification tests, 1 Concerted reactions, 312 Configuration absolute, 138 relative (R,S), 130 Configuration of enantiomers, 129 Conformational analysis, 157 Conformational diastereomers, 163 Conformational energies, 158, 160, 163 Conformational isomer, 158 Conformer, 158 Conjugated  $\pi$  systems, 21, 370 Conservation of charge, 77 Cope rearrangement, 81 Coupling constant, J, 345 Covalent bond, 7 Crossed-aldol condensation, 229 Crossed Claisen condensation, 230 Curved arrow notation, 69 Cycloaddition reactions, 312 4+2 Cycloaddition, 312  $4_{\pi} + 2_{\pi}$  Cycloaddition reactions, 312 Cyclopropane ring, formation, 308

Diastereomeric excess, 148 Diastereomeric transition states, 147, 150 Diastereomers, 134 formation of, 146 Diastereoselectivity, aldol reaction, 236 Diastereotopic protons, 353 1,3-Diaxial interactions, 163 Diazabicycloundecane (DBU), 204 Dieckmann condensation, 307 Diels–Alder cycloaddition, 312 Diels–Alder dienophiles, 315

Diels-Alder reaction diastereoselectivity, 316 endo transition state, 316 regioselectivity, 317 s-cis conformation, 316 stereoelectronic factors, 316 Dienophile, 315 Differential rate expression, 99 Dihedral angle in allenes, 12 1,3-Dipolar cycloaddition, 319 regioselectivity, 321 1.3-Dipoles, 319 preparation, 322 1,3-Dithianes, 197 Donor-acceptor interaction, 76, 313 E2 reactions, 173 Early-late transition states, 96 Eclipsed conformation, 158 Effective size, 167 Electron density and chemical shift, 340 Electron-electron repulsion, 9 Electron movement, 70 Electron sharing, 8 Electrophile, 17 Electrophilic aromatic substitution, 79 Enantiomeric excess, 137 Enantiomers, 128, 129 formation of, 144 Enantiotopic protons, 353 Endergic reaction, 94 Energy barrier, 87 Enolate formation, 225 preparation, 231 Enolate regioisomers, 234 Enolates as nucleophiles, 218, 228 Enol triflates Heck reaction. 251 Stille reaction, 255 Envelope conformation, cyclopentane, 161.171 Enzymes as resolving agents, 142 Equatorial position, 162 Ester preparation, 188 saponification, 187 Exact mass measurement, 380

Excited state, 16

Exergic reaction, 94 Eyring equation, 105 First order rate law, 99 First order spectra, 354 Fischer esterification, 189 Fischer projections, 125 Free energy of activation,  $\Delta G^{\ddagger}$ , 89 Free energy of reaction,  $\Delta G$ , 89 Free radical, 73 initiation, 274, 280 initiators, 275, 280  $\pi$ -bond addition, 274 propagation, 276 termination, 276 Free radical addition, 75 Free radical chain reactions, 274 Free radical cyclization, 283 approach angle, 285 kinetic control, 285 resonance effects, 286 steric control. 287 Free radical polymerization, 277 Free radical scavengers, 276 Functional groups, 2, 86, 183, 369 table, 3 Gauche conformation, 160 Geminal coupling, 345 Grignard reaction, 37, 77 Grignard reagents, preparation, 224 Heff, 338, 339, 344 Halogen-metal exchange, 224 Hammett equation, 111 Hammett plot, nonlinear, 116 Hammond postulate, 96 H-bonding, 370 Heat of combustion, 168 Heck reaction, 251 intramolecular, 252 Heterolytic bond cleavage, 71 Heterolytic bond formation, 71 HOMO, 16 Homoenolate synthon,  $\beta$ -bromo acetals, 304HOMO-LUMO energy gap, 314 HOMO-LUMO interaction, 313 strength, 314 symmetry, 314

474 INDEX

Homolytic cleavage, 73 Homotopic protons, 353 Huckel's rule, 24 Hybrid atomic orbitals, 8 Hybridization, 8 Hydrazones as enolate equivalents, 236 Hydrogen abstraction, 74

Imines, preparation, 197 Inert gas configuration, 7 Integrated rate law, 99 Intramolecular radical addition, 279 Inversion of configuration, 172 Ionic bond, 7 Ionization ratio, 51 IR bending frequencies, 367 IR spectroscopy, 366 IR stretching frequencies, 367 table, 369 IR in structure determination, 371 Isotope effects, 105 Isotopic clusters, 381

Jones reagent, 185, 195

 $K_{a}$ , acidity constant, 51  $K_{b}$ , base constant, 54 Kekulé forms, 18 Ketone derivatives, 198 Ketone enolate, formation, 234 Ketones, preparation, 194 Kinetic deuterium isotope effects primary, 105 secondary, 105 Kinetic enolate, formation, 234 Kinetic resolution, 143

Lewis acid, 48 Lewis base, 48 Lewis structures, 5 Linear geometry, 12 Lithium diisopropyl amide (LDA), 54, 226 Low field, high field, 339 LUMO, 16

Magnetic equivalence, 351 Magnetogyric ratio, 336 Markovnikov addition, 200 Mass spectrometry, 377 fragmentation, 384 Mass to charge ratio, *m/e*, 378 *m*-Chloroperbenzoic acid (MCPBA), 39 meso isomer, 135 Metallocarbene intermediate, 256 Metallocyclobutane, 256 Michael addition, 223 Mobile phase, 333 Molecular ion, 378, 379 Molecular orbital (MO), 6 Mukaiyama reaction, 241 Multiplicity, 348

Neutral carbon nucleophiles, 239 Newman projections, 127 *N*-Hydroxypyridine-2-thione, 281 Nickel P-2 catalyst, 206 NMR, 336 Normal phase chromatography, 334 Nuclear magnetic moment, 336 Nucleophile, 17

Off-resonance decoupling, 363 Olefin metathesis, 256 medium ring formation, 259 molybdenum catalysis, 257 ruthenium catalysis, 257 Optical activity, 137 Orbital diagrams, 22 Orbitals, 5 Order of reaction, 99 Organolithiums, preparation, 224 Organometallic nucleophile, 218, 239 Organotin compounds, Stille coupling, 254 Osmium tetroxide, 38, 153 Oxaphosphetane intermediate, 243 Oxidation. 33 Oxidizing agent, 33 Palladium(0), see Pd(0) Pascal's triangle, 348 Pauli exclusion principle, 11

Pd(0)  $\beta$ -hydride elimination, 250 catalysts, 247 insertion, 248 oxidative addition, 247

INDEX 475

oxidative addition chemoselectivity, 248reductive elimination, 249 transmetallation, 249 Petersen olefination, 206, 244  $\pi$ -bond, 10  $\pi$ -bonds as nucleophiles, 78, 219  $\pi^*$  orbitals, 14 Phosphorous ylid, 242  $pK_a, 51$  $pK_a$  values, table, 52  $pK_{eq}$ , 53 Plane polarized light, 137 Polarized  $\sigma$ -bond as electrophiles, 221 Priority rules, 131 Proton exchange in NMR, 360 Pseudo-first-order rate law, 100 Purification, chromatographic, 333 Pyridinium chlorochromate (PCC), 193 R,S-configuration, 130 Racemate, 140 Racemic mixture, 138 Racemization, 155 Radical cation, 378 Rate-determining step, 91 Rate expression, 99 RCM, see Ring closing metathesis Reaction constant o. 113 sign of  $\rho$ , 114 Reaction coordinate, 91 Reaction intermediate, 90 Reaction mechanism, 76 Reaction rate, electronic effects, 110 Reaction stereoselectivity, 144 Reactive intermediates, 96, 110 Reactivity matching, 223 Rearrangement, cationic, 81 Redox, internal, 36 Reducing agent, 33 Reduction, 33 Reductive amination, 202 Regiospecificity, 294 Re-Si nomenclature, 145 Resolution of enantiomers, 140 Resolving agents, 141 Resonance, 18 Resonance delocalization, 20

Resonance energy, 19, 23 Resonance form, 18 Resonance hybrid, 18 Resonance stabilization, 20 Resonance structures by curved arrow notation, 75 Retrosynthetic analysis, 292 Retrosynthetic step, 294 Reversed phase chromatography, 335 Ring closing metathesis (RCM), 258 conformational effects, 259 Ring closure vs. polymerization. 306 - 308Ring current, 24, 342 Ring formation, relative rates, 307 Robinson annulation, 310 Ruthenium tetroxide, 186 Sawhorse projections, 127 s-character, 13 Second order rate law, 100 Second order spectra, 354 Shielding, 339  $\sigma$ -bond, 8  $\sigma$ -constants, 111  $\sigma^+$ -constants, 116  $\sigma^{-}$ -constants, 116  $\sigma^*$  orbitals, 14 Simmons-Smith cyclopropanation, 245 sp hybridization, 11  $sp^2$  hybrid orbitals, 11 sp³ hybrid orbitals, 8 Specific rotation, 137 Spin quantum number, 336 Spin-spin splitting, 344 Spin system designation, 352 Spin systems, 350 Staggered conformation, 158 Steady-state approximation, 101 Stereoelectronic effect, 172 Stereogenic center, 129 Stereoisomers, 129 Stereospecific, 148 Stereospecificity, 294 Stille coupling, 254 intramolecular, 255 Strain, 168 Strain energy cycloalkanes, table, 169

#### 476 INDEX

Strain energy (continued) cyclobutane, 170 cyclopropane, 169 Strongly coupled spectra, 354 Structure determination, 332 Structure reactivity effects, 110 Substituent constant,  $\sigma$ , 111 table, 112 Substitution reaction, 172 Suzuki coupling, 253 Swern oxidation, 193 Symmetry allowed cycloaddition, 313 syn addition, 149 syn-anti diastereomers, 136 Synthetic planning, 293 Synthon, 303

τ-bond, 169
t-Butyl ester, trifluoroacetic acid (TFA) cleavage, 187
Tetrahedral geometry, 8
Thermodynamic enolate, 234
Thiono ester, 282
Threo-erythro diastereomers, 136
Torsional strain, 170
Transannular interactions, 171 Transannular strain, 171 Transition state, 88 Trigonal geometry, 11 Twist boat conformation, cyclohexane, 162

Umpolung reactivity, 303 Umpolung reagents, 303

Valence bond structures, 20 Valence shell electrons, 7 Vibrational spectroscopy, 367 Vicinal coupling, 345

Walden inversion, 154 Wave number, 366 Weakly coupled spectra, 354 Wedge-dash structures, 126 Wheland intermediate, 79 Wittig-Horner reaction, 243 Wittig reaction, 206

Zero-point energy, 105 Ziegler–Natta catalysts, 256 Zimmerman–Traxler model, 237