THE ELECTRONIC THEORY OF ACIDS AND BASES
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BY W. F. LUDER
Associate Professor of Chemistry
Northeastern University

AND SAVERIO ZUFFANTI
Associate Professor of Chemistry
Northeastern University

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To
GILBERT N. LEWIS
The theory of relativity in physics has led to an increased amount of correlation in physical phenomena and to a keener insight into the fundamental problems of physics. More recently, another kind of relativity theory has made its appearance in chemistry. We owe this theory to the brilliant intuition of G. N. Lewis, by whom it was first proposed in 1923. Unfortunately, it was largely overlooked until about 1940. Since then, numerous papers have been published supporting and developing the theory, which is becoming known as the electronic theory of acids and bases.

According to the Lewis theory, acidity and basicity have nothing to do with the presence of any one element or specific group of elements. Nor is any particular substance to be considered an acid or a base in an absolute sense. Acidity and basicity are relative terms indicating certain types of experimental behavior which may be exhibited by the vast majority of chemical compounds. Most of them may behave either as acids or as bases depending upon conditions.

This theory makes possible a great increase in the amount of correlation possible in all phases of chemistry. It casts a flood of illumination over a wide area of apparently unrelated phenomena, from the corrosion of metals to catalysis in organic chemistry. In making such a statement, no claim of finality for the electronic theory of acids and bases is intended. No doubt the theory will be modified and improved. Nevertheless it seems evident that it can perform a useful function in chemistry. What this function is we hope to set forth more definitely in our conclusion (Chapter 13).

This book attempts a systematic presentation of the relationship of the electronic theory of acids and bases to chemistry as a whole. Much of the material has been published by us previously, in a series of ten papers in the Journal of the American Chemical Society, the Journal of Chemical Education, and Chemical Reviews. These papers have aroused so much interest that we have been encouraged to reorganize and combine them with some new material to make this book.
We have tried to present the electronic theory of acids and bases and its ramifications in an elementary manner. The theory is of interest to all types of chemists, and the book has been written with that in mind. Three chapters (10, 11, and 12) contain details on catalysis which are of most interest to organic chemists, but if necessary they can be skipped without losing continuity; Chapter 9 gives the essence of their contents.

One test of any new theory is: Does it offer a usable explanation which succeeds in correlating a larger number of experimental facts? We believe that according to this test the Lewis theory of acids and bases is successful. It does correlate a much larger body of experimental facts. On the other hand, a new theory in order to be convincing must not depart too far from previously accepted ideas. The wide interest in the Lewis theory indicates that it does not.

At the present time, then, the electronic theory of acids and bases seems to be the most satisfactory available. This is not to say that it is a final explanation. Like any other scientific theory, it is not to be believed in, but merely to be used until a more inclusive theory appears.

In conclusion, we express our gratitude to Professor G. Albert Hill for reading our manuscript and for his advice and encouragement, to Miss Amy F. Meserve for her assistance in the preparation of the manuscript, to Dean William C. White for his interest, and to the editors of the *Journal of Chemical Education*, the *Journal of the American Chemical Society*, and *Chemical Reviews* for their permission to use material first published in their journals.

We wish especially to acknowledge our great debt to Gilbert Newton Lewis, who shortly before his death read our manuscript and not only gave us the benefit of his suggestions and criticisms but granted us the privilege of dedicating our book to him.

W. F. Luder
Saverio Zuffanti

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CONTENTS

CHAPTER  PAGE
1. HISTORICAL BACKGROUND            1
2. ATOMIC ORBITALS AND VALENCE      18
3. THE ELECTRONIC THEORY OF ACIDS AND BASES  43
4. ELECTROPHILIC AND ELECTRODOTIC REAGENTS  69
5. ACIDIC AND BASIC RADICALS        80
6. NEUTRALIZATION                   87
7. TITRATIONS WITH INDICATORS      91
8. DISPLACEMENT                     98
9. CATALYSIS                        107
10. ACID CATALYSIS                   114
11. BASE CATALYSIS                   137
12. ALKOXIDES AS CATALYSTS         150
13. CONCLUSION                      157

INDEX                                   159
Chapter 1

HISTORICAL BACKGROUND

1. Introduction

The fundamental nature of acids and bases has been an intriguing problem for chemists since the time of Robert Boyle. The answer to the problem has been revised many times and is still the subject of considerable controversy. At present, there are no less than five competing theories of acids and bases! They may be listed as follows:

1. The water theory (sometimes called the Arrhenius theory).
2. The theory of solvent systems (Franklin-Germann).
3. The proton theory (Brønsted-Lowry).
4. The positive-negative theory (Usanovich).
5. The electronic theory (G. N. Lewis).

The most familiar of the five probably are the first three. The water theory of acids and bases was very widely accepted early in the century, though perhaps not so generally as we are inclined to think. For example, some organic chemists recognized substances like pyridine to be bases regardless of solvent, and the metallurgists spoke of acidic and basic linings for refining processes. More recently the proton theory of acids and bases has been gaining adherents rapidly, while a few chemists have preferred the theory of solvent systems.

But the fact is that none of these three theories satisfactorily explains more than a limited portion of the experimental behavior of acids and bases. For a long time an increasing amount of data has compelled realization of the fact that acid-base phenomena are far more widespread than is generally acknowledged. Those who have grasped this fact have already abandoned the water
theory. Yet neither the theory of solvent systems nor the proton theory is inclusive enough to cover all the data.

The theory of solvent systems conforms to the experimental fact that there are many other substances besides those containing hydrogen which exhibit typical acid properties. But it makes the definitions of acid and base as rigidly dependent upon the solvent as does the water theory.

The proton theory emphasizes the important fact that acid-base phenomena can be observed in any solvent or even in the absence of a solvent. It also takes into account the experimental fact that there are many other substances besides the hydroxyl ion which exhibit typical basic properties. Yet it does not recognize the complementary data with regard to acids. The followers of Brønsted have maintained that only substances capable of giving up protons can be called acids.

We have, then, two independent and, in important features, contradictory theories of acids and bases. This situation seems to be due to the neglect of two factors: first, a portion of the experimental data, and second, the electronic theory of the covalent bond. Probably such neglect was natural and even necessary in the early stages of the development of each theory. Possibly neither would have accomplished as much as it has without some such limitation. However, as a result of this neglect, neither theory gives an insight into the fundamental nature of acids and bases.

One of the most powerful theoretical tools available to the chemist is the electronic theory of the covalent bond, which we owe primarily to G. N. Lewis. Lewis was also the first to suggest an electronic explanation of acid-base phenomena.\(^1\) The theory of the covalent bond is now almost universally accepted among chemists, but the electronic theory of acids and bases until quite recently has been ignored where it has not been actively opposed.\(^2\)

Lewis's definitions of acids and bases do not depend upon the presence of any particular element. As he himself points out: "To restrict the group of acids to those substances which contain hydrogen interferes as seriously with the systematic understanding of chemistry as would the restriction of the term oxidizing

---


agent to substances containing oxygen." Another theory which starts from the same point is that of Usanovich, which will be considered below.

From an experimental standpoint, a substance that exhibits the properties of an acid should be called an acid, regardless of preconceived notions about the dependence of acidic properties on a particular element. Those properties have been agreed upon from the beginning. Yet many substances possessing them are not yet generally recognized as acids. This situation persists in spite of the fact that some of these substances were once called acids, when the principal criterion of an acid was its experimental behavior.

Some of the properties by which acids were recognized when the term first came into use were listed by Boyle as follows: they dissolve many substances; they precipitate sulfur from its solution in alkalis; they change blue plant dyes to red; they lose all these properties on contact with alkalis. These were recognized as the properties of aqueous solutions of acids. If the solution of a substance in water had these and other typical acid properties, the substance itself was known as an acid. Thus carbon dioxide and sulfur trioxide were called acids because their solutions exhibited the properties common to all aqueous solutions of acids.

This strictly experimental approach was largely abandoned during the series of controversies which began with Lavoisier's attempt to make oxygen the necessary constituent of all acids. After Davy had shown that some acids do not contain oxygen, hydrogen became the "acidifying principle." Davy himself wrote in 1814 that "Acidity does not depend upon any particular elementary substance, but upon peculiar arrangement of various substances." We shall see how nearly correct Davy was. But Liebig successfully maintained the hydrogen theory against Berzelius by defining an acid as any substance that contains easily replaceable hydrogen atoms.

With the advent of the Arrhenius theory of ionization, an acid was defined as a hydrogen compound ionizing in water solution to give hydrogen ions. A base was a hydroxyl compound which would give hydroxyl ions in water solution. The reaction between an acid and a base—i.e., neutralization—would produce a salt and water:

\[
\text{HA} + \text{BOH} \rightarrow \text{BA} + \text{H}_2\text{O}
\]

\[\text{acid} \quad \text{base} \quad \text{salt} \quad \text{water}\]

3 Lewis, J. Franklin Inst., 228, 293 (1938).
These definitions were quite generally adopted in spite of the efforts of several investigators to show how inadequate they were. The emphasis upon ions in chemical reactions was at its height. A physical chemistry textbook of the period, as quoted by Folin and Flanders, contained the statement, "We have already reached a point where we can say that nearly all, if not all, chemical reactions are due to ions, molecules as such not entering into chemical action." In this atmosphere the work of Collie and Tickle, Hantzsch, Folin and Flanders, and Lapworth was largely overlooked.

Collie and Tickle, in their paper published in 1899, suggested that oxonium salts are similar to ammonium salts. They referred to "such bases as those of the pyridine series" and even spoke of the "hypothetical base oxonium hydroxide, OH$_3$OH." Hantzsch noted the basic action of water, methyl alcohol, and dimethyl ether in anhydrous sulfuric acid. Folin and Flanders, in a paper published in 1912, reported the titration of a large number of acids in such solvents as benzene, toluene, chloroform, and carbon tetrachloride. They used sodium ethoxide and sodium amoxide as bases and phenolphthalein as an indicator. They noted that weak acids which cannot be titrated in water give excellent results in organic solvents. Even hydrogen sulfide was titrated. They found their solutions of acids practically non-conducting and concluded that very few ions were present. (This conclusion has been supported by the work of Fuoss and Kraus.) Carbon dioxide could not be titrated in either chloroform or benzene, a result which seems to agree with Lewis's statement that carbon dioxide is a "secondary" acid (Chapter 3). The most striking thing about their paper appears in a footnote in which the authors mention that mercuric chloride can be titrated with phenolphthalein and sodium ethoxide in the same manner as any other acid, but they failed to draw the logical conclusion that mercuric chloride might be an acid. Lapworth was one of the first to attack the Arrhenius-

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$^5$ Folin and Flanders, J. Am. Chem. Soc., 34, 774 (1912).
$^7$ Hantzsch, Z. physik. Chem., 61, 257 (1908); Hantzsch, ibid., 65, 41 (1909); Hantzsch, Z. Electrochem., 24, 201 (1918); Hantzsch, ibid., 29, 221 (1923).
$^9$ Fuoss and Kraus, J. Am. Chem. Soc., 55, 2387 (1933); Fuoss and Kraus, ibid., 65, 3614 (1933); Luder, P. B. Kraus, C. A. Kraus, and Fuoss, ibid., 58, 255 (1936).
Ostwald theory of the catalytic activity of acids. However, these and similar investigations failed to make much impression until the concept of the covalent bond began to reduce the overemphasis upon ionic reactions.

With the growing clarity of the electronic theory of valence, the inadequacy of the water theory of acids and bases was increasingly recognized. Even more important was the large amount of work undertaken with solvents other than water. It soon became evident that acid-base phenomena could also be observed in non-aqueous solvents. Many non-aqueous solvents are incapable of ionizing as water does, and many of the so-called organic bases are also non-ionic, e.g., pyridine. Even when ions are produced in non-aqueous solvents by the reaction of compounds that had gradually come to be recognized as bases, the anion, of course, is not the hydroxyl ion. Two alternatives seemed available: (1) to make the definitions independent of solvent, but to retain hydrogen as the one essential element; or (2) to make the definitions independent of any particular element, but to tie them very closely to the solvent in which the phenomena were observed. We shall consider the proton theory first.

2. One-Element Theories of Acids and Bases.

Before discussion of the Brønsted-Lowry theory a few words about an earlier one-element theory may not be amiss. The first one-element theory was apparently due to Lavoisier. Oxygen was supposed to be necessary for the manifestation of acidic properties. Later it became evident that some substances that do not contain oxygen are nevertheless acids. Instead of concluding with Davy that no particular element is responsible for acidity, most chemists of the time, in spite of the opposition of Berzelius, gradually swung over to the view that hydrogen was essential to an acid. Now, one hundred years later, we find ourselves in the same position of having to give up an element, except that it is hydrogen we must give up instead of oxygen.

In spite of the overwhelming trend toward hydrogen as the "acidifying principle," some remnants of the theory of Berzelius persisted—for example, the use by metallurgists of the term "acid" to denote oxides such as silicon dioxide and the term "base" for oxides like calcium oxide. Berzelius, as a consequence of his theory of valence, classified oxides as acids or bases. Furthermore, the theory provided a method of predicting the relative
strengths of acids or bases.\textsuperscript{10} Bases like sodium oxide were supposed to be strong because of the high positive charge on the metal of the combination. The negative charge in oxides was supposed to be due to oxygen, and the acidity of acids was due to the high negative charge of the oxygen compared to the weaker positive charge of the non-metallic constituent, such as sulfur or carbon. Therefore, acids with a high proportion of oxygen should be strong. For example, sulfur trioxide should be a stronger acid than sulfur dioxide. This prediction is in accord with the experimental facts, as we shall see later, but development along this line was halted because of the inadequacy of the theory of valence upon which it depended.

When Berzelius' theory of valence was discarded, primarily as the result of Faraday's discovery of the laws of electrolysis, the theory of acids and bases founded upon it also had to be abandoned. Doubt had already been cast on it by Davy's demonstration that hydrochloric acid did not contain oxygen. There was no doubt of the acidity of hydrochloric acid; so, still thinking in terms of an elementary acidifying principle, many chemists came to regard hydrogen as the one element necessary for acid properties. Ideas regarding bases did not become so definite until the advent of the proton-donor theory.

The modern one-element theory of acids and bases is usually credited to Brønsted and Lowry. They proposed the proton theory independently in 1923. But G. N. Lewis, who set forth his electronic definitions of acids and bases in the same year, also explained the proton-donor concept as a special case of his broader theory.\textsuperscript{1} According to the proton theory, an acid donates a proton to a base, and a base accepts a proton from an acid. The acid and base may be either compounds or ions, as shown in the following examples:

\[
\begin{align*}
\text{HCl} + \text{NH}_3 & \rightarrow \text{NH}_4\text{Cl} \\
\text{NH}_4^+ + \text{OH}^- & \rightarrow \text{NH}_3 + \text{H}_2\text{O}
\end{align*}
\]

This picture of acid-base reactions makes them independent of solvent, as is shown in the first equation, and eliminates the arti-

ficial connection between acids, bases, and salts, as is shown in the second equation.

Since any group of atoms that gives up a proton is called an acid, acids may be cations, anions, or neutral molecules:

\[
\begin{align*}
\text{acid} & : \quad \text{NH}_4^+ + \text{C}_2\text{H}_3\text{O}_2^- \rightarrow \text{HC}_2\text{H}_3\text{O}_2 + \text{NH}_3 \\
\text{acid} & : \quad \text{HSO}_4^- + \text{OH}^- \rightarrow \text{HOH} + \text{SO}_4^{2-} \\
\text{base} & : \quad \text{HCl} + \text{C}_5\text{H}_5\text{N} \rightarrow \text{C}_5\text{H}_5\text{NHCl}
\end{align*}
\]

Bases also may be cations, anions, or molecules:

\[
\begin{align*}
\text{acid} & : \quad \text{Co(OH)(NH}_3)_5^{+2} + \text{H}_3\text{O}^+ \rightarrow \text{Co(H}_2\text{O)(NH}_3)_5^{+3} + \text{H}_2\text{O} \\
\text{base} & : \quad \text{C}_2\text{H}_3\text{O}_2^- + \text{H}_2\text{O} \rightarrow \text{HC}_2\text{H}_3\text{O}_2 + \text{OH}^- \\
\text{acid} & : \quad \text{C}_2\text{H}_5\text{OH} + \text{HCl} \rightarrow \text{C}_2\text{H}_5\text{OH}_2^{+1} + \text{Cl}^{-1}
\end{align*}
\]

When an acid gives up its proton, the residue must be a base, since it can retrieve a proton. The acid and base thus related are called a conjugate system:

\[
\begin{align*}
\text{acid} & : \quad \text{NH}_4^+ & \rightarrow & \text{NH}_3 + \text{H}^+ \\
\text{base} & : \quad A_1 & \rightarrow & B_1
\end{align*}
\]

This equation does not represent an actual reaction, since none takes place unless a base is present to accept the proton, e.g.:

\[
\text{base} \quad \text{B}_2 \quad \text{acid} \quad \text{A}_2
\]

Addition of these two equations gives

\[
\begin{align*}
\text{acid} & : \quad \text{NH}_4^+ + \text{OH}^- \rightarrow \text{HOH} + \text{NH}_3 \\
\text{base} & : \quad A_1 & \rightarrow & B_1
\end{align*}
\]

The reaction involves two conjugate pairs. Probably the most direct way of describing what happens in such a reaction is to say that the two bases (\(\text{NH}_3\) and \(\text{OH}^-\)) are competing for the proton.
HISTORICAL BACKGROUND

When we consider the familiar types of acid-base phenomena, such as ionization and hydrolysis, we find them all described by the type equation

\[ A_1 + B_2 \rightleftharpoons A_2 + B_1 \]

as shown in the following equations:

**Ionization:**

- Strong acid in water:
  \[ \text{HCl} + \text{H}_2\text{O} \rightarrow \text{H}_3\text{O}^{\text{+}} + \text{Cl}^{-} \]
- Weak acid in water:
  \[ \text{HC}_2\text{H}_3\text{O}_2 + \text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^{\text{+}} + \text{C}_2\text{H}_5\text{O}_2^{-} \]
- Weak base in water:
  \[ \text{HOH} + \text{NH}_3 \rightleftharpoons \text{NH}_4^{\text{+}} + \text{OH}^{-} \]

**Hydrolysis:**

- Hydrolysis of sodium acetate:
  \[ \text{H}_2\text{O} + \text{C}_2\text{H}_3\text{O}_2^{-} \rightleftharpoons \text{HC}_2\text{H}_3\text{O}_2 + \text{OH}^{-} \]
- Hydrolysis of ammonium chloride:
  \[ \text{NH}_4^{\text{+}} + \text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^{\text{+}} + \text{NH}_3 \]

One method of comparing the strengths of acids and bases is by means of the equilibrium constant:

\[ K' = \frac{[A_2] \times [B_1]}{[A_1] \times [B_2]} \]

For example, for molecular acids in water the equation reduces to an expression equivalent to that of the water theory:

\[ A + \text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^{\text{+}} + B \]

\[ K = K' \times [\text{H}_2\text{O}] = \frac{[\text{H}_3\text{O}^{\text{+}}] \times [B]}{[A]} \]

The equation gives the strength of A relative to \( \text{H}_3\text{O}^{\text{+}} \). For a cation acid such as the ammonium ion, the equilibrium constant is seen to be identical with what was previously called the hydrolysis constant:

\[ \text{NH}_4^{\text{+}} + \text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^{\text{+}} + \text{NH}_3 \]

\[ K = \frac{[\text{H}_3\text{O}^{\text{+}}] \times [\text{NH}_3]}{[\text{NH}_4^{\text{+}}]} \]
A strong acid (HCl) must have a weak conjugate base (Cl⁻); conversely, a weak acid (HC₂H₃O₂) must have a strong conjugate base (C₂H₅O₂⁻). The relationship between an acid and its conjugate base makes it unnecessary to list basic dissociation constants. For example, for ammonia

\[
K_b = \frac{[\text{NH}_4^+\text{I}] \times [\text{OH}^-\text{I}]}{[\text{NH}_3]}
\]

but

\[
K_b = \frac{[\text{NH}_4^+\text{I}] \times [\text{OH}^-\text{I}]}{[\text{NH}_3]} \times \frac{K_w}{[\text{H}_3\text{O}^+\text{I}] \times [\text{OH}^-\text{I}]} = \frac{K_w[\text{NH}_4^+\text{I}]}{[\text{H}_3\text{O}^+\text{I}] \times [\text{NH}_3]} = \frac{K_w}{K_a}
\]

where \(K_a\) refers to the ammonium ion.

This method of comparing the strengths of acids according to the extent of reaction with the solvent cannot be used for all solvents. If the solvent is inert, e.g., a hydrocarbon, other methods must be adopted. Some of them will be discussed later.

The proton theory of acids and bases recognizes the existence of a large number and variety of bases, both molecular and ionic: hydroxyl ion, amide ion, ethoxide ion, piperidine and other amines, alcohols, ethers, acetate ion, hydrosulfide ion, cyanide ion, bisulfate ion, ketones, and many others. It also recognizes that acid-base phenomena do not depend upon the solvent. But its great weakness is that it ignores a large body of experimental data by restricting the use of the word *acid* to proton donors. For a time the only alternative to the proton theory seemed to be the theory of solvent systems.


The development of the theory of solvent systems was begun by Franklin in 1905.¹¹ Reasoning from formal analogy to the hydrogen ion-hydroxyl ion theory he defined acids and bases in liquid ammonia. According to his theory, if water ionizes into hydronium and hydroxyl ions, liquid ammonia must ionize into ammonium and amide ions:

\[
2\text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^+ + \text{OH}^-\text{I}
\]

\[
2\text{NH}_3 \rightleftharpoons \text{NH}_4^+ + \text{NH}_2^-\text{I}
\]

Substances like ammonium chloride are acids, and substances like sodium amide are bases, in liquid ammonia. Ammonia solutions of acids and bases neutralize each other just as aqueous solutions do. For example:

\[
\begin{align*}
\text{acid} & \quad \text{base} & \quad \text{salt} & \quad \text{solvent} \\
\text{OH}_3\text{Cl} + \text{NaOH} & \rightarrow \text{NaCl} + 2\text{H}_2\text{O} \\
\text{NH}_4\text{Cl} + \text{NaNH}_2 & \rightarrow \text{NaCl} + 2\text{NH}_3
\end{align*}
\]

Other characteristic properties of acids and bases, such as the reaction of acids with metals and of bases with non-metals, were observed. The similarity between ammonia and water solutions was demonstrated very widely.\(^\text{12}\) Acids like \(\text{B(OH)}_3\) and \(\text{B(NH}_2\text{)}_3\) were compared, and the latter was called an ammono acid. \(\text{BiN}\) reacts with \(\text{NH}_4\text{I}\) in ammonia to form \(\text{BiI}_3\) and \(\text{NH}_3\), just as in water \(\text{Bi}_2\text{O}_3\) reacts with \(\text{HCl}\) to give \(\text{BiCl}_3\) and \(\text{H}_2\text{O}\). The behavior of \(\text{Zn(NH}_2\text{)}_2\) in ammonia is analogous to that of \(\text{Zn(OH)}_2\) in water. Both are insoluble, but the addition of \(\text{KNH}_2\) and \(\text{KOH}\), respectively, brings them into solution. \(\text{Zn(NH}_2\text{)}_2\) is amphoteric in the same way that \(\text{Zn(OH)}_2\) is:

\[
\begin{align*}
\text{Zn(OH)}_2 + 2\text{KOH} & \rightarrow \text{K}_2\text{Zn(OH)}_4 \\
\text{Zn(NH}_2\text{)}_2 + 2\text{KNH}_2 & \rightarrow \text{K}_2\text{Zn(NH}_2\text{)}_4
\end{align*}
\]

Since metals evolve hydrogen when reacting with either hydronium ions in water or ammonium ions in ammonia,

\[
\begin{align*}
2\text{H}_3\text{O}^{+1} + \text{Mg} & \rightarrow \text{Mg}^{+2} + 2\text{H}_2\text{O} + \text{H}_2 \\
2\text{NH}_4^{+1} + \text{Mg} & \rightarrow \text{Mg}^{+2} + 2\text{NH}_3 + \text{H}_2
\end{align*}
\]

it appeared that the acid properties of both the hydronium and the ammonium ions must be due to the proton. The question then arose whether the idea of the solvent system could be applied to systems in which no protons were present.

Germann, Cady and Elsey, Jander, Wickert, and Smith have extended the solvent system theory of acids and bases to include non-protonic systems. Germann\(^\text{13}\) showed that aluminum chloride in phosgene has typical acid properties. The solution dissolves metals with evolution of carbon monoxide gas and is neutralized


\(^{13}\) Germann and Timpany, J. Am. Chem. Soc., 47, 2275 (1925); Germann, ibid., 47, 2461 (1925); Science, 61, 70 (1925).
by metallic chlorides such as calcium chloride. Germann assumed that the aluminum chloride forms with the solvent a complex which he called a solvo acid:

\[
\text{COCl}_2 \rightleftharpoons \text{COCl}^+ + \text{Cl}^-
\]

ionization of the solvent

\[
\text{AlCl}_3 + \text{COCl}_2 \rightleftharpoons \text{COCl}^+ + \text{AlCl}_4^-
\]

The concentration of the solvent cations has been increased by the addition of the acid (aluminum chloride); hence the reaction of the solution with a metal is more rapid than that of the solvent alone:

\[
\text{Ca}^+ + \text{COCl}^+ \rightarrow \text{Ca}^{2+} + \text{CO} + \text{Cl}^-
\]

This is analogous to the reaction in water: 14

\[
\text{Ca} + 2\text{H}^+ \rightarrow \text{Ca}^{2+} + \text{H}_2
\]

Germann's definitions of acids and bases were simplified by Cady and Elsey, 15 who defined an acid as a solute that gives rise to a cation characteristic of the solvent, and a base as a solute that gives rise to an anion characteristic of the solvent.

Jander 16 and coworkers used these last definitions to interpret the results of their work with liquid sulfur dioxide as solvent. Thionyl chloride is acidic, tetramethylammonium sulfite is basic, and aluminum sulfite is amphoteric in liquid sulfur dioxide. Ionization of this solvent gives thionyl and sulfite ions:

\[
2\text{SO}_2 \rightleftharpoons \text{SO}^{2+} + \text{SO}_3^{2-}
\]

The neutralization of thionyl chloride by tetramethylammonium sulfite in this solvent is represented as follows (assuming that neutralization is the formation of salt and solvent):

\[
\text{SOCl}_2 + [\text{(CH}_3)_4\text{N}]_2\text{SO}_3 \rightarrow 2\text{(CH}_3)_4\text{NCl} + 2\text{SO}_2
\]

The amphoteric behavior of Al\(_2\)(SO\(_3\))\(_3\) in sulfur dioxide is analogous to that of Al(OH)\(_3\) in water. Each contains the negative ion of the respective solvent. Aluminum sulfite, which is insoluble in sulfur dioxide, dissolves on addition of either thionyl chloride

14 No doubt both positive ions (H\(^{+1}\) and COCl\(^{+1}\)) are solvated, but we assume this to be understood. Therefore we shall indicate this solvation only when showing it is necessary for clarity in our discussion.


16 Jander and Mesech, Z. physik. Chem., A188, 255 (1939); also see preceding papers in this series.
or tetramethylammonium sulfite, just as aluminum hydroxide, which is insoluble in water, dissolves on addition of either nitric acid or potassium hydroxide.

Smith 17 changed the definitions somewhat in reviewing his work with selenium oxychloride as a solvent. He found that stannic, ferric, silicon, arsenious, and titanium chlorides and sulfur trioxide displayed typical acid properties in selenium oxychloride. He defined an acid as an electron-pair acceptor toward the solvent, and a base as an electron-pair donor toward the solvent. These definitions, suggested in 1938, were the first to show the influence of the proposals made by Lewis in 1923.

The height of formalism was reached by Wickert 18 in his definitions of acids and bases in terms of the solvent system. He overlooks such experimental behavior as amphoteric in order to state his definitions wholly in terms of ions. Shatenstein 19 also has pointed out one of the several inconsistencies in Wickert's presentation. Although Wickert defines an acid as an ionic compound the cation of which has an incomplete electronic configuration, he recognizes that ammonium salts are acids in ammonia. Another contradiction of the experimental facts is his listing of antimony trichloride but not aluminum chloride as an acid.

The essential ideas of the theory of solvent systems are summarized in Table 1. In the first three examples, it is obvious enough that the acid that has reacted with each solvent is hydrogen bromide. But Smith 17 seems to be the only adherent of the theory of solvent systems who recognized that in examples 5 and 6 aluminum chloride and stannic chloride are true acids.

### Table 1

Neutralization Reactions According to the Theory of Solvent Systems

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<thead>
<tr>
<th>Solvent</th>
<th>Acid</th>
<th>+ Base</th>
<th>→ Salt</th>
<th>+ Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>H₂O</td>
<td>H₃O⁺, Br⁻¹</td>
<td>K⁺¹⁺, OH⁻¹</td>
<td>K⁺¹⁺, Br⁻¹</td>
<td>2H₂O</td>
</tr>
<tr>
<td>NH₃</td>
<td>NH₄⁺, Br⁻¹</td>
<td>K⁺¹⁺, NH₂⁻¹</td>
<td>K⁺¹⁺, Br⁻¹</td>
<td>2NH₃</td>
</tr>
<tr>
<td>C₂H₅OH</td>
<td>C₂H₅OH₂⁺, Br⁻¹</td>
<td>K⁺¹⁺, OC₂H₅⁻¹</td>
<td>K⁺¹⁺, Br⁻¹</td>
<td>2C₂H₅OH</td>
</tr>
<tr>
<td>SO₂</td>
<td>SO⁺², Br₂⁻¹</td>
<td>K₂⁺², SO₄⁻²</td>
<td>2K⁺²⁺, Br⁻¹</td>
<td>2SO₂</td>
</tr>
<tr>
<td>COCl₂</td>
<td>COCl⁺¹, AlCl₄⁻¹</td>
<td>K⁺¹⁺, Cl⁻¹</td>
<td>K⁺¹⁺, AlCl₄⁻¹</td>
<td>COCl₂</td>
</tr>
<tr>
<td>SeOCl₂</td>
<td>(SeOCl)₂⁺¹⁺, SnCl₆⁻²</td>
<td>2K⁺²⁺, Cl⁻¹</td>
<td>K₂⁺²⁺, SnCl₆⁻²</td>
<td>2SeOCl₂</td>
</tr>
<tr>
<td>SbCl₅</td>
<td>Sb⁺³⁺, Br₈⁻¹</td>
<td>3K⁺³⁺, Cl⁻¹</td>
<td>3K⁺³⁺, Br⁻¹</td>
<td>SbCl₅</td>
</tr>
</tbody>
</table>

The strength of the solvent-systems theory lies in its emphasis upon the fact that acid behavior is not confined to solutions containing proton donors. The advocates of the theory have demonstrated that, although their acid solutions do not contain protons, they show all the typical experimental behavior of aqueous solutions of hydrogen acids. The weaknesses of this theory are two: first, the limitation of acid-base phenomena to solvent systems; and second, the emphasis upon ionization as the most important factor in acid-base properties. Probably the first followed from the second; at any rate, many investigators have shown that ionization plays a far less important role than is indicated by the solvent-systems theory. It would appear that this theory describes only one aspect of the nature of acids and bases: namely, their reactions with amphoteric solvents and the properties of the resulting solutions. We are most familiar with these properties since they are most easily observed. A reluctance to go beyond them is readily understood, but for many chemists the Brønsted theory has overcome this reluctance, at least with respect to bases. But the Brønsted theory admits of no acids other than proton donors. As the proponents of the solvent-systems theory have shown, this limitation does not correspond to the experimental facts. If the experimental approach is to prevail, it is improper to go on saying—as Meerwein, Shatenstein, and others do—that certain substances are “acid-analogous” because of their properties but are not acids because they do not contain hydrogen. Brønsted is undoubtedly correct in attributing acid-base properties to the molecules themselves rather than to their solutions. In this respect the Brønsted theory, as far as it goes, is closer to the experimental facts than the theory of solvent systems. Just as important is Brønsted’s view that acids and bases are not necessarily ionic.

4. The Positive-Negative Theory.

Any attempt to reconcile these two contradictory theories of acids and bases must involve a deeper insight into their fundamental nature. One such attempt has been made by Usanovich.


Usanovich has defined an acid as any substance capable of giving up cations or of combining with anions, and a base as any substance capable of giving up anions or of combining with cations. He also suggests that oxidation-reduction reactions are a special case of acid-base phenomena. Acids combine with electrons as well as with anions, and bases give up electrons to acids. Oxidizing power is a limited phase of acidity, and both are due to attraction for negative particles. Some examples of neutralization according to Usanovich are given in Table 2.

**TABLE 2**

**Neutralization Reactions according to the Theory of Usanovich**

<table>
<thead>
<tr>
<th>Acid</th>
<th>Base</th>
<th>Salt</th>
</tr>
</thead>
<tbody>
<tr>
<td>SO₃</td>
<td>Na₂O</td>
<td>Na₂SO₄</td>
</tr>
<tr>
<td>Sb₂S₅</td>
<td>3(NH₄)₂S</td>
<td>2(NH₄)₂SbS₄</td>
</tr>
<tr>
<td>Fe(CN)₅</td>
<td>3KCN</td>
<td>K₃Fe(CN)₆</td>
</tr>
<tr>
<td>CH₃I</td>
<td>(CH₃)₄N</td>
<td>(CH₃)₄NI</td>
</tr>
<tr>
<td>Cl₂</td>
<td>2Na</td>
<td>2NaCl</td>
</tr>
</tbody>
</table>

Sulfur trioxide is an acid because it combines with the anion O⁻²:

\[
SO₃ + O^{-2} \rightarrow SO_4^{-2}
\]

Antimony pentasulfide is an acid because it combines with the sulfide ion:

\[
Sb₂S₅ + 3S^{-2} \rightarrow 2SbS₄^{-3}
\]

Ferric cyanide combines with the cyanide ion. Methyl iodide gives up the cation CH₃⁺¹. Chlorine combines with two electrons from two sodium atoms.

This theory is general and covers more of the experimental behavior, but objections may be raised to it. Certain inconsistencies have been pointed out by Shatenstein, among them the emphasis upon salt formation and the formal reasoning involved in making ions so important in the scheme. In addition, one might mention the lack of correlation between the definitions and the degree of "coordination-unsaturation" which Usanovich recognizes is of great importance in determining acidity and basicity. In one sense the positive-negative theory is a retrogression to the early ideas of Berzelius on valence. Furthermore, the inclusion of oxidation-reduction as a special case of acid-base phenomena does not seem to be justified. The relationship is close but, as we shall see, not quite as Usanovich presents it.
5. The Electronic Theory of Acids and Bases.

An earlier attempt to reconcile the proton and the solvent-systems theories was made by G. N. Lewis in 1923. Strictly speaking, it was not an attempt at reconciliation, since both theories were proposed independently by Lewis as special cases of his more general and more fundamental theory. Brønsted and Lowry presented their ideas in the same year, but the general form of the solvent-systems theory came several years later. The conflict between the two theories has gone on, although the solution to the problem has been at hand since 1923. Each theory was consistent as far as it went, but each was merely part of the whole picture. The inclusion of both into one clear, simple, and fundamental theory was accomplished by Lewis largely because he founded his theory firmly upon experimental facts, allowing no preconceived notion of the necessity of a particular element or the desirability of emphasizing ions or the solvent to interfere.

Lewis chose four familiar experimental criteria as the basis of his definitions of acids and bases. The four "phenomenological criteria" are:

I. Neutralization. Acids and bases may combine more or less rapidly with each other.

II. Titration with Indicators. Acids and bases may be titrated against each other by the use of substances, usually colored, known as indicators.

III. Displacement. An acid or base will in general replace a weaker acid or base from its compounds.

IV. Catalysis. Acids and bases frequently act as catalysts.

All four of these characteristics must be displayed by a particular substance for classification as an acid or a base. However, in order to make the definitions as concise as possible, the first and second properties, neutralization and titration with indicators, are chosen for specific mention. Acids are substances which, like hydrogen chloride, neutralize sodium hydroxide or any other base. Bases are substances which, like sodium hydroxide, neutralize hydrogen chloride or any other acid. Many substances are capable of acting in either way and are called amphoteric.

These definitions include both the Brønsted and the solvent-systems theories as well as the water theory. The list of bases according to the Lewis theory is identical with that according to
the Brønsted theory, and all the substances recognized as acids by the solvent-systems theory are included. A simple experiment, first suggested by Lewis,\(^3\) strikingly illustrates the application of these definitions. It is only one of many that can easily be performed (see Chapter 7). Crystal violet is an indicator which gives the same color change in different solvents. When sodium hydroxide is titrated against hydrochloric acid in water, with crystal violet as the indicator, the solution is yellow when strongly acidic and violet when basic. Pyridine and triethylamine can be titrated in a similar manner against hydrochloric acid and are therefore bases. If pyridine is dissolved in some comparatively inert solvent, such as chlorobenzene, the same violet color is observed when crystal violet is added. Now, if boron trichloride or stannic chloride is added to the basic pyridine solution, the color changes to yellow. Thus, boron trichloride and stannic chloride are acids. If triethylamine, acetone, or any other fairly strong base is added, the color changes back to violet. Similar titrations can be performed in other solvents with other indicators and with many other acids and bases none of which contains hydrogen or hydroxyl ions. With the proper choice of solvent and indicator, all the substances that are acids according to the solvent-systems theory can be shown to be acids on this experimental basis (neutralization and titration). Displacement and catalysis reactions support this conclusion. Especially striking is the similarity in the action of acid catalysts such as aluminum and boron trichlorides, sulfur trioxide, hydrogen fluoride, and sulfuric acid. (Both displacement and catalysis will be considered in some detail in subsequent chapters.) All four of these experimental criteria provide ample evidence that there is an inherent difference between acids and bases. The difference is not dependent upon the solvent or upon the presence of a particular element. Therefore it would seem to involve a contrast in electronic configuration.

Searching for the property common to all acids, or that common to all bases, Lewis concluded that acids and bases correspond respectively to what Sidgwick later called acceptor and donor molecules.\(^2\) Neutralization is the formation of the coordinate covalent bond between the acid and the base. Such a definition


\(^3\) The coordinate covalent bond will be discussed in the next chapter.
holds when the acid and base are either hydrogen ion \(^2\) and hydroxyl ion

\[
\text{acid} \quad \text{base} \\
H^{+1} + \ddot{O}H^{-1} \rightarrow \dddot{O}H:O \dddot{H}
\]

or boron trichloride and ammonia

\[
\text{acid} \quad \text{base} \\
\text{Cl} \quad 
\downarrow \quad \downarrow \\
\text{Cl} \quad \text{B} + \arctan \quad :\text{N}H \rightarrow \arctan \quad \dddot{\text{B}:\text{N} \rightarrow} \arctan
\]

The base donates a share in a lone pair of electrons to the acid to form the coordinate covalent bond between the two. Formation of the coordinate bond is always to be considered the first step, even though ionization may subsequently take place. Examples of ionization will be considered in later chapters.

Before discussing the consequences of these ideas further, let us now turn to a consideration of some of the more recent developments in the theory of valence inasmuch as the "rule of eight" in valence theory still has not been discarded by many chemists. For example, in order to understand why the covalent compound stannic chloride, a liquid much like carbon tetrachloride in physical properties, is such a strong acid, we must become familiar with the limitations of the octet theory. This can be accomplished without reference to any theory of acids and bases. The following chapter deals with the rule of two which replaces the rule of eight. It also contains suggestions for the modification in writing electronic formulas which must be made in adopting the newer ideas.

\(^2\) In general we shall refer to the "hydronium ion" as the hydrogen ion and write its formula as \(\text{H}^{+1}\) wherever use of the formula \(\text{H}_2\text{O}^{+1}\) does not add to the discussion.
Chapter 2

ATOMIC ORBITALS AND VALENCE

1. Introduction ............................................. 18
2. The Rule of Eight ......................................... 19
3. Atomic Orbitals ........................................... 21
4. The Origin of Valence ..................................... 27
5. Electrovalence ............................................. 28
6. Covalence ................................................... 29
7. Electron Affinity .......................................... 36
8. The Coordinate Covalent Bond ............................ 38

1. Introduction.

One of the most powerful tools available to the chemist is the electronic theory of valence. Still it is not yet being used generally to full advantage. The two principal factors responsible for the delay seem to be reluctance to abandon the rule of eight and failure to make use of the concept of atomic orbitals.

Ever since the proposal of the theory of the covalent bond by G. N. Lewis in 1916, it has been obvious that there are numerous compounds which do not "obey" the rule of eight, which was strongly emphasized by Kossel and Langmuir. However, elementary textbooks have stressed it so thoroughly that the rule of eight is still generally accepted by those unfamiliar with the recent literature in the field.

The idea of atomic orbitals from elementary quantum theory is now often presented as an aid to the understanding of atomic structure but is still rarely applied to the problem of valence. It is the purpose of this chapter to show in an elementary way how abandoning the rule of eight and applying the theory of atomic orbitals can lead to fuller understanding of valence. In order to do this, some of the details, such as are involved in hybridization of orbits, for example, will be either omitted or modified for the sake of simplicity. Some of the modifications explicitly proposed or implicit in the discussion may give rise to objection on the ground that they do not give the complete picture. But, after all, we do not yet understand all the phenomena of valence by any means.
2. The Rule of Eight.

The familiar dictum that in forming chemical bonds atoms gain, lose, or share electrons to make the outermost shell of each atom contain eight electrons probably gained such general acceptance because the majority of known chemical compounds is made up primarily of elements in the first two rows of the periodic table. With but few exceptions, these elements do obey the rule of eight. The presence of carbon with its thousands of compounds seems to lend added support to the rule. Actually, of course, the sheer number of compounds of any one element has no bearing whatever on the rule of eight. The carbon atom counts merely as one atom in the ten. But, even in these ten, two of the eight elements which do not already have complete valence shells refuse to obey the rule of eight, namely, beryllium and boron.

There is little doubt that boron trichloride is a covalent compound and, therefore, that the boron atom in the compound has only six electrons in its valence shell (crosses represent original boron electrons):

\[
\text{Cl} : \quad \text{B} : \text{Cl} : \quad \text{Cl} : \quad \text{Cl} : \\
\quad \text{Cl} : \quad \text{B} : \text{Cl} : \quad \text{Cl} : \\
\quad \text{Cl} : \\
\]

Beryllium chloride also seems to be covalent, as shown by its high solubility in benzene. If the compound is covalent the beryllium atom has four outer electrons in beryllium chloride (the two crosses represent the two electrons from the beryllium atom):

\[
\text{Cl} : \quad \text{Be} : \text{Cl} : \quad \text{Cl} : \\
\quad \text{Cl} : \quad \text{Be} : \text{Cl} : \quad \text{Cl} : \\
\quad \text{Cl} : \\
\]

When the third period is considered, three out of seven atoms show "exceptions" to the rule of eight: aluminum, phosphorus, and sulfur. Aluminum chloride, phosphorus pentfluoride, and sulfur hexafluoride are typical covalent compounds. They must, therefore, contain six, ten, and twelve bonding electrons, respectively.
To be sure, AlCl₃ is associated to Al₂Cl₅ at low temperatures, but we shall adopt the custom of not showing association where it does not add to the clarity of the presentation. There is no more reason for habitually indicating the association of AlCl₃ than for continually writing dimeric formulas for those carboxylic acids that associate to double molecules. We should not permit irrelevant details to confuse the chemical significance of an electronic formula. The formula should correspond as closely as possible with chemical behavior. The dimeric formula for AlCl₃ obscures the principal feature of its chemistry. Confirmation of this statement will be evident in subsequent chapters.

Upon proceeding further in the periodic table, many other examples of covalent compounds containing six, ten, and twelve electrons in the valence shell may be found. Most of these are in groups III, V, and VI of the representative elements (Fig. 2). Several occur in group VII but will be discussed later. The existence of iodine heptafluoride shows that fourteen electrons are possible in the valence shells of some atoms.

Other violations occur also in the "related metals" (Fig. 2). Tungsten, in addition to the di-, tri-, and tetrachlorides, forms covalent penta- and hexachlorides, WCl₅ and WCl₆. WBr₅ and WBr₆ are also known. The low boiling point (19.5°C) of WF₆ indicates that this compound is hexacovalent. In addition to the di-, tri-, and tetrachlorides, molybdenum forms covalent compounds having the formulas MoCl₅ and MoF₆. Osmium forms a covalent hexafluoride (b.p. 205°C) and also an octafluoride, OsF₈, a stable typically covalent compound having a boiling point of 47.3°C. Here is one compound at least having twice the number of bonding electrons permitted by the rule of eight.

The examples given above have included only compounds containing simple covalent bonds. Coordination compounds and metal carbonyls offer many more "violations" of the rule of eight. Other examples in which some electrons are not used as bonding electrons (analogous to the lone pair in ammonia) will be discussed subsequently. Turning to a consideration of the valences

¹In view of the rarity of pentacovalency, doubt may arise concerning its existence in WCl₅ and WBr₅. However, the low boiling points of these compounds, together with the fact that the boiling point of the bromide is the higher of the two, may be taken as sufficient evidence for covalency until crystal-structure measurements are made. If the compounds were ionic the reverse order of boiling points would be expected.
of the “related metals” (Fig. 2) when acting as positive ions, we find that not one of the many positive ions formed conforms to the rule of eight. In none of them does the loss of electrons leave a completed shell of eight electrons. This fact is largely obscured in the older type of eight-column periodic chart but is very obvious in Fig. 2.

Most of the evidence illustrated by the compounds just discussed has been largely neglected in most textbooks. A few such compounds have been considered by them, e.g., phosphorus pentachloride and sulfur hexafluoride, and various devices have been adopted in the attempt to force compliance with the rule of eight. One was the hypothesis of single-electron bonds in such compounds as phosphorus pentachloride and sulfur hexafluoride. It is no doubt true that one-electron bonds do exist in a very few compounds. The hydrogen-molecule ion is evidently held together by one electron, but the bond is much weaker than the electron pair even in this, the most favorable case. If one-electron bonds exist in the boron hydrides, they are still weaker, as shown by the instability of their compounds. To maintain eight valence electrons for the sulfur atom in the hexafluoride would require four one-electron bonds which (because of the large size of the sulfur atom compared to hydrogen or boron) would be far more unstable than those in boron. Yet sulfur hexafluoride is one of the most stable compounds known.\(^3\) In spite of this conclusive evidence, it has been claimed\(^3\) that parachor measurements indicate the presence of one-electron bonds in such compounds. But, as pointed out by R. Samuel, the parachor may lend itself equally well to different schemes of valence, depending upon the original assumptions.\(^4\) Additional exceptions to the rule of eight will be considered in more detail below.

Abandoning the rule of eight would seem to be the first step in acquiring the fuller understanding of valence made possible by the progress in our knowledge of atomic structure.

### 3. Atomic Orbitals.

According to the quantum theory, the periodic system can be regarded as being built up from hydrogen by adding one proton


at a time (plus the required number of neutrons) to the nucleus and one electron at a time to the atom outside the nucleus. From the periodic table and spectroscopic data it is deduced that the electrons arrange themselves in main levels and sublevels in a regular way, as shown in Fig. 1. Figure 1 is a plot of approximate relative energy levels. Each circle represents one atomic orbital. On paper each circle may be regarded as a box that can be occupied

either by one electron or by two electrons spinning in opposite directions.

The numbers in Fig. 1 are different values of the principal quantum number \( n \). Each main level is designated by one value of the principal quantum number. The letters \( s, p, d, f \) correspond to the values 0, 1, 2, 3 for the orbital quantum number \( l \), which determines the "eccentricity" of the orbital. The difference in eccentricity leads to sublevels in the main levels. Both types of numbers must be used in the equations for the energy of the orbitals. The orbital quantum number has a series of values for each value of \( n \):

\[
l = 0, 1, 2, 3 \ldots n - 1
\]

For example, in the third main level \((n = 3)\) there are three sublevels \((l = 0, 1, 2)\) designated \(3s, 3p, 3d\). From spectroscopic data obtained when the atoms are placed in a magnetic field it is found that \( s \) sublevels always contain one orbital, \( p \) sublevels always contain three orbitals, \( d \) sublevels contain five orbitals, and \( f \) sublevels contain seven orbitals. It is obvious from Fig. 1 that in the upper energy levels there is considerable overlapping of main levels. Furthermore, some of the closer levels apparently interchange occasionally. The interchange is indicated by vertical lines joining the sublevels involved.

In general, the electrons occupy the lowest energy levels first. One electron goes into each orbital in a given sublevel (one line in Fig. 1) until each of the orbitals in that sublevel is occupied by one electron. When every orbital in a given sublevel contains one electron, then, and only then, they begin filling up with two electrons in each orbital. Two electrons in an orbital are said to be coupled or paired. Supposedly, the opposite spins of coupled electrons result in electromagnetic attraction more than sufficient to overcome the repulsion of like charges.

The way in which electrons are added to the orbitals can be made clearer by considering some examples from Fig. 2. From the chart the electron configuration of aluminum is seen to be

\[
^{13}\text{Al} 1s^2 2s^2 2p^6 3s^2 3p
\]

where the superscripts represent the number of electrons in each kind of orbital. If we wish to use the superscripts to represent the number of electrons in each orbital, the three \( 2p \) orbitals must be separated as follows:

\[
^{13}\text{Al} 1s^2 2s^2 2p^2 2p^2 2p^2 3s^2 3p
\]
## THE REPRESENTATIVE ELEMENTS
(Differentiating Electron in Outermost Shell)

<table>
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<tr>
<th>n</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
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<tr>
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<td>2</td>
<td>3</td>
<td>4</td>
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</tr>
<tr>
<td>P</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
<td>6</td>
</tr>
</tbody>
</table>

## THE RELATED METALS
(Differentiating Electron in Second from Outermost Shell)

<table>
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<th>5</th>
<th>6</th>
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<tr>
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<td>2</td>
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<td>10</td>
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<td>12</td>
<td>13</td>
<td>14</td>
<td>15</td>
<td>16</td>
</tr>
</tbody>
</table>

## THE RARE EARTHS
(Differentiating Electron in Third from Outermost Shell)

<table>
<thead>
<tr>
<th>n</th>
<th>19</th>
<th>20</th>
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</thead>
<tbody>
<tr>
<td>f</td>
<td>17</td>
<td>18</td>
<td>19</td>
<td>20</td>
<td>21</td>
<td>22</td>
<td>23</td>
<td>24</td>
</tr>
</tbody>
</table>

---

**Fig. 2.**

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Remembering that the orbitals take electrons one at a time until each orbital in a given sublevel has one electron in it before coupling begins, we take the electron configurations from the chart and write them as follows (writing the $2p$ orbitals together since there is no change in them):

Silicon

\[ ^{14}\text{Si} \quad 1s^2 \quad 2s^2 \quad 2p^6 \quad 3s^2 \quad 3p \]

Phosphorus

\[ ^{15}\text{P} \quad 1s^2 \quad 2s^2 \quad 2p^6 \quad 3s^2 \quad 3p^3 \]

Sulfur

\[ ^{16}\text{S} \quad 1s^2 \quad 2s^2 \quad 2p^6 \quad 3s^2 \quad 3p^2 \]

Chlorine

\[ ^{17}\text{Cl} \quad 1s^2 \quad 2s^2 \quad 2p^6 \quad 3s^2 \quad 3p^3 \quad 3p \]

Note that phosphorus has three unpaired $p$ electrons, but that, since there can be only three $p$ orbitals, sulfur has only two unpaired electrons and chlorine only one.

We shall now make a slight modification in the usual way of writing electronic symbols. The five atoms just discussed may be written as follows:

:Al· :Si· :P· :S· :Cl·

Written in this way, the symbols indicate the paired and unpaired electrons of the valence shell. For example, the symbol for phosphorus now shows the two paired $3s$ electrons and the three unpaired $3p$ electrons.

For those elements which have two incomplete shells (main levels), i.e., most of the "related metals" (Fig. 2), valence bonds to other atoms often involve both shells. The original Lewis conventions provided no way to represent such atoms, but if the rule of eight is abandoned and the convention just proposed is adopted, the method is obvious. Dots representing the electrons in both shells should be used, regardless of how many may be required. No distinction between the two energy levels need be made since they are so close together, even closer than indicated in Fig. 1. The vertical lines on the right of Fig. 1 joining the $4s$ with the $3d$, the $5s$ with the $4d$, and the $6s$ and $4f$ with the $5d$ sublevels are used to indicate that there is practically no difference. Actually, these levels may interchange in different atoms.

From Fig. 2, we see that manganese has the electron configuration:

\[ ^{25}\text{Mn} 1s^2 2s^2 2p^6 3s^2 3p^6 3d^3 3d^3 3d^4 4s^2 \]

The valence electrons are obviously the two paired 4s electrons and the five single or unpaired electrons, with practically no difference in energy among the seven. The electronic symbol would be

\[ \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \]

with two dots close together for the paired 4s electrons and the other five far enough apart so as to represent five unpaired electrons. For iron

\[ ^{26}\text{Fe} 1s^2 2s^2 2p^6 3s^2 3p^6 3d^2 3d^2 3d^2 3d^2 4s^2 \]

the electronic symbol would be

\[ \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \]

showing eight electrons, four paired and four single. For cobalt

\[ ^{27}\text{Co} 1s^2 2s^2 2p^6 3s^2 3p^6 3d^2 3d^2 3d^2 3d^2 3d^2 4s^2 \]

the electronic symbol would be

\[ \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \]

showing nine electrons, six paired and three single. For nickel

\[ ^{28}\text{Ni} 1s^2 2s^2 2p^6 3s^2 3p^6 3d^2 3d^2 3d^2 3d^2 3d^2 4s^2 \]

the electronic symbol would be

\[ \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \cdot \]

showing ten electrons, eight paired and two single. Copper will be taken up later, since its consideration involves a discussion of its valence. With zinc there is again only one level in the valence shell, consisting of the two 4s electrons. The 3d orbitals are now fully occupied.

The advantages of writing electronic symbols in the manner suggested will become more fully apparent with the discussion of valence.

To summarize: (1) The order in which atoms are built up by the addition of electrons to the corresponding nucleus is: 1s, 2s, 2p, 3s, 3p, (4s, 3d), 4p, (5s, 4d), 5p, (6s, 5d, 4f), 6p, 7s, 6d, accord-
ing to Fig. 1. (The parentheses correspond to the short vertical lines joining the same levels on the right side of the figure. Both devices emphasize that these levels have about the same energy and may even change their order in different elements.) (2) All the orbitals in a given sublevel must contain one electron apiece before pairing of spins begins to complete the orbitals with two electrons in each. (3) Electronic symbols should be written showing all paired and unpaired electrons in the valence shell. Since in many atoms the valence shell includes electrons from two energy levels, as many as twelve dots may often be required. Occasionally more will be needed.

4. The Origin of Valence.

When we turn to a consideration of chemical compounds, a natural assumption to make as a starting point is: Uncompleted orbitals in one atom may be filled by electrons from another atom. This filling of orbitals may be done by loose electrons (resulting in electrovalence) or by electrons that remain in the other atoms (resulting in covalence).

The rule of eight is replaced by the rule of two. Both electrovalence and covalence are due to pairing of electrons to fill incomplete orbitals. Since it is only for \( n = 2 \) that the maximum number of electrons is eight, the two rules are equivalent only for the second period. For the third period, \( n = 3 \) giving nine orbitals, holding a maximum of eighteen electrons. Considerations of charge and space, in addition to an insufficient number of electrons, show that this number is not approached in the third period except by compounds such as phosphorus pentafluoride and sulfur hexafluoride and by ions like \( \text{AlF}_6^{-3} \) and \( \text{SiF}_6^{-2} \). In this period and the later ones, the number of electrons in the valence shells of the atoms in molecules may seem at first glance to be quite unpredictable, but the rule of two in conjunction with a table of electron configurations often will solve the problem. Several examples will now be considered under the separate headings of electrovalence and covalence.

The only rule to be followed is that uncompleted orbitals in one atom may be filled by electrons from another atom. It is called the rule of two because a completed orbital contains two electrons.

Many readers will immediately recognize that the discussion owes its origin primarily to London (Z. Physik, 46, 455 [1928]). However, they will also note certain modifications as the argument proceeds.
5. Electrovalence.

Since substitution of the rule of two for the rule of eight makes no difference as far as the electrovalences of the representative elements (Fig. 2) are concerned, they will not be discussed here. Neither will the rare earths be considered. Readers interested in the rare earths may consult the Journal of Chemical Education for an article by Pearce and Selwood.\(^8\)

One fact should be mentioned before we proceed with a brief discussion of the related metals (Fig. 2). Practically no monatomic negative ions have a charge greater than \(-2\). Very few monatomic positive ions have a charge greater than \(+3\). For example, we are accustomed to think of "stannic ion" as having a valence of \(+4\), but actually, there is no such thing as a stannic ion in pure stannic chloride, a typical covalent compound. Even in water, the stannic ion has no existence as a monatomic ion with four positive charges on it. Apparently, insufficient energy is involved in ordinary chemical reactions to build up charges greater than \(-2\) or \(+3\). According to Coulomb's law, the size of the atoms involved must also have an important bearing upon the formation of ions, so that beryllium, a small atom, does not show much tendency to form an ion.

The rule of two centers attention on those atoms in a molecule that are completing their orbitals in the combination, and so it does not add much to a discussion of the related metals as positive ions. However, it might be well to point out that the related metals do not violate the rule of two as they do the rule of eight. As mentioned previously, none of the many positive ions of these metals conforms to the rule of eight, but the rule of two does not specify the nature of the positive ion. The number of electrons removable to complete the orbitals of some other atom is determined by size and charge as well as by the numbers of unpaired electrons.

For example, the electronic configuration of iron is

\[
26\text{Fe } 1s^2 2s^2 2p^6 3s^2 3p^6 3d^6 3d 3d 3d 3d 4s^2
\]

and the electronic symbol showing electrons from both incomplete shells as valence electrons is

\[
\text{Fe}^2+
\]

It is easy to remove two electrons (suppose we say two of the unpaired $3d$ electrons) to give the ferrous ion, a little more difficult to remove a third one, and impossible because of the large electrical field to remove the fourth electron by ordinary chemical means. Most of the ions of these metals are left with unpaired electrons, as is shown by their color and paramagnetism.

6. Covalence.

When atoms complete one or more orbitals to form covalent bonds, the number of pairs shared between them depends upon various factors, including: (1) the number of unpaired electrons in each atom, (2) spatial considerations. Bearing the second factor in mind, we shall emphasize the first. We shall do so by first applying the rule of two to each compound chosen as an example as if it were formed from neutral atoms whose electron configurations are given in Fig. 2.

Considering the representative elements first, we find that the maximum number of valence electrons for elements in the second period according to Fig. 1 is eight. Beryllium and boron chlorides have already been discussed sufficiently to illustrate compounds in which the maximum of eight is not reached. The properties of boron trichloride as a strong acid make it apparent that the one empty orbital of the boron atom can still be filled by two paired electrons from an atom in a basic molecule. The remaining four active elements in the second period require no further discussion, since for them the rule of two becomes equivalent to the rule of eight.

In the third period (only the representative elements in that period being considered for the time being) the number of orbitals is nine. The tendency of magnesium to fill up its orbitals seems to be the reason for the activity of Grignard reagents. The tendency of aluminum chloride to fill up an empty orbital in the aluminum atom results in its familiar activity as an acid catalyst. The formation of $\text{SiF}_6^{2-}$ ion apparently involves the filling of two orbitals of silicon in silicon tetrafluoride by paired electrons.

---

9 R. Samuel (see note 4) mentions the possible existence of nitrogen pentfluoride. Such a compound if it does exist would involve promotion from $n = 2$ to $n = 3$.

10 Chapter 3.

11 Chapters 9 and 10.
from two fluoride ions to give a total of twelve electrons. Carbon in the preceding period cannot do this because in its compounds all its orbitals are filled by eight electrons.

The phosphorus atom has two paired and three single electrons in its valence shell, and so phosphorus trifluoride is written (using crosses for phosphorus electrons):

\[
\begin{align*}
\text{:F:} \\
\text{\textcross{X}} \text{\textcross{X}} \text{\textcross{X}} \text{\textcross{X}} \text{\textcross{X}} \text{\textcross{X}} \text{\textcross{X}} \text{\textcross{X}} \\
\text{\textcross{P} \text{x} \text{F:}} \\
\text{:F:} \\
\end{align*}
\]

Phosphorus pentafluoride must be formed by uncoupling the lone pair of 3s electrons. This reaction may be formally represented as follows:

\[
\begin{align*}
\text{:F:} \text{\textcross{X}} \text{\textcross{X}} \text{\textcross{X}} \\
\text{\textcross{P} \text{x} \text{F:}} \\
\text{:F:} \\
\rightarrow 2 \text{\textcross{X}} \text{\textcross{X}} \text{\textcross{X}} \\
\text{\textcross{P} \text{x} \text{F:}} \\
\end{align*}
\]

where the small arrow indicates that the two paired electrons are uncoupled during the reaction. Actually, the two fluorine atoms are combined in a molecule and the paired electrons between them must also be uncoupled, but they are written separately for simplicity. This uncoupling of paired electrons requires energy, but not more than the reaction can amply supply, since the compound is formed and is fairly stable. The extreme instability of nitrogen pentafluoride, if it exists at all, must be due to the fact that the nitrogen atom has no more orbitals available for which \( n = 2 \), and promotion of electrons from \( n = 2 \) to \( n = 3 \) would require more energy than its synthesis can furnish.

The question which arises at this point, namely, what determines in detail the number of atoms grouped about the central atom in such cases, is beyond the scope of the discussion in this chapter. Suffice it to say here that completion of orbitals may be opposed by the repulsion of built-up charges or hindered by lack of space for enough atoms to gather around the central atom.

The pentachloride (Powell, Clark, and Wells, J. Chem. Soc., 1942, 642) and pentabromide (Powell and Clark, Nature, 145, 971 [1940]) of phosphorus have been the subject of crystal structure investigations which have shown them to be ionic in the solid state. The ions are \( \text{PCl}_4^{+1} \) and \( \text{PCl}_6^{-1} \) for the chloride and \( \text{PBr}_4^{+1} \) and \( \text{Br}^{-1} \) for the bromide. This fact does not mean that they are not covalent in the gaseous state, nor does it imply that \( \text{PF}_5 \) is not pentacovalent in all three states. The boiling point for the fluoride is far below the boiling points of the chloride and bromide. If the fluoride were ionic, its boiling point would be higher.
The series of sulfur compounds $S_2F_2$, $SF_2$, $SF_4$, $S_2F_{10}$, $SF_6$ offers an interesting application of the rule of two. The sulfur atom has two electron pairs and two single electrons:

$$\text{SF}_2\text{ must be formed by direct pairing of the two unpaired electrons in the sulfur atom:}$$

$$\text{SF}_2$$

The formation of $SF_4$ involves the uncoupling of one pair, leading to ten electrons in the sulfur valence shell. Both pairs are uncoupled in $SF_6$, giving a total of twelve:

$$\text{SF}_6$$

Two different electronic formulas have been suggested for $S_2F_2$:  

$$\text{S}_2\text{F}_2$$

The chemical evidence for a choice between the two seems inconclusive. In the second formula, two of the paired electrons on the upper sulfur atom have been uncoupled to pair off with the two unpaired electrons of the lower sulfur atom. Thus the upper sulfur atom has ten electrons in its valence shell.

$S_2F_{10}$ resembles $SF_6$ to some extent in its chemical properties but is more reactive. The electronic formula is similar in that in both molecules the sulfur atoms have twelve valence electrons:

$$\text{S}_2\text{F}_{10}$$

The interhalogen compounds of the types $AB_3$, $AB_5$, and $AB_7$ are also interesting examples of the rule of two. $ICl_3$ must involve

the uncoupling of two of the paired electrons in the iodine atom of iodine monochloride:

\[ \text{I}_2 \times \text{Cl}_2 \xrightarrow{\text{uncoupling}} \text{I}_2^+ \text{Cl}_2^- + 2 \text{Cl}^- \xrightarrow{\text{coupling}} \text{Cl}_2^+ \text{I}_2^- \text{Cl}_2^- \]

In this formula there are ten electrons in the valence shell of iodine. In iodine pentachloride there must be twelve, one more pair being uncoupled:

Iodine heptafluoride should be written with fourteen valence electrons, all seven of the original iodine electrons being shared with the seven fluorine atoms:

From these examples, chosen from the representative elements, it is apparent that the rule of two (more often than the rule of eight) automatically gives the correct result when molecules are considered as if formed in the same way as the periodic system is built up. A few more examples taken from the related metals will be considered next.

The older eight-column periodic charts obscured the similarities observed in the chemical properties of the related metals. The type of chart illustrated by Fig. 2 clears up this source of confusion. All these atoms have either one or two electrons in the outermost shell and therefore behave primarily as metals. Most of them have two electrons in the last shell, and from the chemist's point of view most of the remainder could be considered as having two also. Their similarities when acting as cations have already been considered.

In spite of their primary characteristics as metals, some of them in one respect do behave as non-metals. Chromium and manganese, for example, form with oxygen negative ions that correspond to sulfate and perchlorate ions. Yet neither chromium nor manganese form negative “chromide” or “manganide” ions, com-
parable to sulfide and chloride ions. The answer to this apparent contradiction lies in the fact that most of the related metals have two incomplete shells of practically the same energy. Therefore, all the electrons in both shells should be considered valence electrons in writing electronic formulas.

As an example, let us compare the behavior of sulfur and chromium. Sulfur and oxygen atoms both have six valence electrons, two electron pairs and two single electrons in each:

\[
\begin{array}{ccc}
\text{S} & \text{O} \\
\text{H} & \text{H} \\
\end{array}
\]

Applying the rule of two, we may imagine sulfuric acid as being formed in the following way: First, the two incomplete orbitals in each oxygen can be filled with two electrons from one sulfur atom and two electrons from two hydrogen atoms:

\[
\begin{array}{ccc}
\text{H} & \text{O} & \text{S} & \text{O} & \text{H} \\
\end{array}
\]

Two more oxygen atoms, each with two unpaired electrons, can then complete their orbitals by uncoupling the two electron pairs on the sulfur atom and forming double bonds:

\[
\begin{array}{ccc}
\text{H} & \text{O} & \text{S} & \text{O} & \text{H} \\
\end{array}
\]

Note that there are twelve electrons in the valence shell of the sulfur atom according to the rule of two. The customary way of writing the formula with only eight electrons was adopted primarily to force compliance with the rule of eight.\(^{15}\)

The chromate is formed in a similar way. Representing the five unpaired 3d electrons and the one 4s electron as being in the same valence shell, we find that chromium has the same number of valence electrons as sulfur:

\[
\begin{array}{ccc}
\text{Cr} & \text{S} \\
\end{array}
\]

The one electron on the left may be considered the 4s electron, but this distinction is not important, since all are practically equal.

\(^{15}\) This return to "classical" formulas will be considered further in section 7.
equivalent (Fig. 1). The first step in the formation of chromic acid can be formulated in exactly the same way as before:

\[
\text{H:O:Cr:O:H (hypothetical)}
\]

The next step is different only in that the four remaining electrons are already single; hence no uncoupling by the oxygen atoms is necessary:

\[
\begin{array}{c}
\text{H:O:Cr:O:H} \\
\end{array}
\]

The reason that the \( \text{Cr}^{-2} \) ion does not exist is that such an ion would still contain four unpaired electrons and would be too unstable. To fill all six incomplete orbitals to form a monatomic ion would give it a charge of \(-6\), which is much too great a charge to build up against the Coulomb repulsion.

Similar analysis shows that fourteen electrons are shared in perchloric and permanganic acids:

\[
\begin{array}{c}
\text{H:O:Cl:O:} \\
\text{H:O:Mn:O:}
\end{array}
\]

All four of these formulas are equivalent to the old formulas used before electrons were discovered, e.g.,

\[
\begin{array}{c}
\text{H--O--S--O--H} \\
\text{H--O--Cl--O}
\end{array}
\]

showing a valence of two for oxygen, six for sulfur, and seven for chlorine. Anticipating objections to the idea of fourteen electrons for chlorine, we may recall the interhalogen compounds \( \text{AB}_5 \) and \( \text{AB}_7 \). In these compounds, the relative sizes of the two atoms seem to determine whether the compounds will be formed. So far, there is no known chlorine pentafluoride, but \( \text{BrF}_5 \) and \( \text{IF}_7 \) do exist. Apparently there is not room enough for five fluorine atoms around one chlorine atom. But there are only four oxygen
atoms involved in perchloric acid. If fourteen electrons are holding seven other atoms to iodine in the heptafluoride, it does not seem unreasonable to find that fourteen electrons are holding four oxygen atoms to chlorine.

The series of osmium halides OsCl$_3$, OsCl$_4$, OsF$_5$, OsF$_6$ is interesting. The electronic symbol for the osmium atom is similar to that of iron, with two 6s electrons and two 5d electrons paired, leaving four 5d electrons unpaired:

$$\overset{\text{x}}{\text{Os}}\overset{\text{x}}{\text{Cl}}$$

OsCl$_3$ is ionic, affording one example of the relatively small number of trivalent ions. Three of the four electrons can be removed, but not a fourth, since OsCl$_4$ is not ionic but covalent

$$\begin{array}{c}
    \text{Cl} \\
    \text{O} \\
    \text{Cl}
\end{array}$$

with a total of twelve electrons for the osmium valence shell. The osmium atom in OsF$_5$ must have fourteen electrons

$$\begin{array}{c}
    \text{F} \\
    \text{O} \\
    \text{F}
\end{array}$$

and the osmium atom in OsF$_6$ must have sixteen electrons:

$$\begin{array}{c}
    \text{F} \\
    \text{O} \\
    \text{F}
\end{array}$$

The failure of the related metal carbonyls and nitrosyls to obey the rule of eight was recognized very early. So it is not surprising that our present concepts of their valence relationships already are so close to the ideas presented here that no further discussion of them is necessary.

Discussion of other covalent compounds of the related metals would be interesting, but in regard to most of them it would be merely speculation, since insufficient experimental data are available. However, the above examples of covalent compounds,
together with those considered in the previous section, should be sufficient to illustrate the application of the rule of two to the problem of valence. It is evident that proper use of the rule of two demands both application of our knowledge of atomic orbitals and modification of our present method of writing electronic symbols so as to represent two partly filled energy levels of practically the same energy as one valence shell.


Though substituting the rule of two for the rule of eight effects some improvement in the theory of valence, it is still far from being a completely satisfactory theory. So far in our discussion reference to any tendency opposing the filling of atomic orbitals has been omitted deliberately. That there is such a tendency becomes obvious upon considering the following example. The formation of sulfur monoxide might be represented as due to the sharing of the two unpaired electrons in each atom:

\[
\text{\textit{S}} + \text{\textit{O}} \rightarrow \text{\textit{S}}:\text{\textit{O}}
\]

But the attraction of the oxygen atom for electrons is greater than that of the sulfur atom. We are all familiar with the qualitative manifestation of this fact in the representative elements (Fig. 2). A more exact comparison is given in Table 1. (The larger numbers represent greater attraction for electrons.)

**TABLE 1**

**First Ionization Potentials (in Volts)**

(Data from Sisler and Vanderwerf, *J. Chem. Education*, 22, 390 [1945].)

<p>| | | | | | | |</p>
<table>
<thead>
<tr>
<th></th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>H</td>
<td>He</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>13.53</td>
<td>24.48</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Li</td>
<td>Be</td>
<td>B</td>
<td>C</td>
<td>N</td>
<td>O</td>
<td>F</td>
</tr>
<tr>
<td>5.37</td>
<td>9.28</td>
<td>8.33</td>
<td>11.22</td>
<td>14.48</td>
<td>13.55</td>
<td>18.6</td>
</tr>
<tr>
<td>Na</td>
<td>Mg</td>
<td>Al</td>
<td>Si</td>
<td>P</td>
<td>S</td>
<td>Cl</td>
</tr>
<tr>
<td>5.12</td>
<td>7.61</td>
<td>5.96</td>
<td>8.12</td>
<td>11.1</td>
<td>10.31</td>
<td>12.96</td>
</tr>
</tbody>
</table>

How to take account of this difference in attraction for electrons is a difficult and still unsettled problem. For a single bond, the difference might be represented by writing the electrons closer to the more electrophilic (electron-attracting) atom. But for a
double bond, such as we have tentatively written for sulfur monoxide, it might better be pictured as follows:

\[ \ddots \vdots \ddots \leftrightarrow \ddots \vdots \ddots \]

This formula corresponds, perhaps, a little better to the instability of sulfur monoxide. But probably neither the double-bond (as first written) nor the single-bond formula represents the actual molecule. This state of affairs apparently holds for many other molecules. It has led to the theories of resonance and of molecular orbitals. For reasons discussed below, neither of these theories will be extensively used in this book. At any rate, the condition of "strain" in molecules like sulfur monoxide, as the formula is written on the basis of our knowledge of atomic orbitals, cannot be ignored. A redistribution of electrons among all orbitals to give the most stable arrangement in view of the two opposing tendencies must be contemplated. A possible way out of the dilemma is to write formulas generally according to the rule of two but modified where necessary to show as far as possible the potentialities of the compounds for chemical reaction. (This would mean usually ignoring association between identical molecules.)

To illustrate this principle the following formulas may be considered:

\[ \ddots \vdots \ddots \quad \ddots \vdots \ddots \quad \ddots \vdots \ddots \quad \ddots \vdots \ddots \quad \ddots \vdots \ddots \]

Note that the first three formulas show six electrons, the fourth eight, and the fifth twelve for the sulfur atom. The number increases from eight to twelve (from formulas IV to V) because of the electron affinity of the protons. Consider that the two protons added to a sulfate ion cause the redistribution of electrons. We imagine the positive protons shifting all electrons toward themselves. This makes the sulfur atom more positive, as shown below:
Thus the number of electrons on the sulfur atom is increased to twelve. This return to the "classical" structures for such oxy-acids as

\[
\begin{align*}
\text{H}_2\text{O} & \quad \text{O} \\
\text{H} & \quad \text{O} \\
\text{S} & \quad \text{O} \\
\text{H} & \quad \text{O} \\
\end{align*}
\]

with twelve and fourteen electrons, is supported by Phillips, Hunter, and Sutton.\(^{16}\) They also suggest that the stability of such acids is actually due to the formation of the double bonds. This accounts for the fact that fluorine does not form such acids.\(^{16}\) Since fluorine has only four orbitals, no double bonds can be formed.

If we regard each electronic formula as dynamic rather than static, there is little to be gained by emphasizing the concept of resonance hybrids. Furthermore, the detailed theory of resonance has come under heavy fire.\(^{17}\) It is claimed by A. Burawoy\(^{17}\) that "... the interpretation of polyatomic structures does not require the hypothesis of resonance among several idealized valence-bond structures, a speculative application of the quantum-mechanical conception of resonance, which is in disagreement with numerous facts." In view of this controversy, we shall avoid detailed application of the theory of resonance. The theory of molecular orbitals, an alternative treatment, is too complex for the present discussion.

It is important to bear in mind the dynamic nature of electronic formulas, which arises from the two opposing tendencies which must be balanced in the electron distribution of many molecules. We shall encounter more examples in later chapters, but we mention only two additional examples here, carbon dioxide and aldehydes:

\[
\begin{align*}
\text{H}_2\text{O} & \quad \text{O} \\
\text{H} & \quad \text{C} \\
\text{O} & \quad \\
\end{align*}
\]

8. The Coordinate Covalent Bond.

There is another way in which orbitals in one atom may be filled by electrons from another atom. Consider the molecule of

\(^{17}\) Burawoy, Nature, 165, 328 (1945); Burawoy, Chemistry & Industry, 51, 434 (1944).
the covalent compound boron trichloride. The maximum number of orbitals available for compound formation in atoms of the second period is four, one 2s and three 2p orbitals. In boron trichloride, only three are filled:

\[
\begin{align*}
\cdot\text{Cl} & \\
\cdot\text{Cl} & \text{B} \\
\cdot\text{Cl} & \\
\end{align*}
\]

The fourth may be filled by a lone pair of electrons from another molecule, e.g., ammonia, to form an addition compound:

\[
\text{Cl} - \text{B} + \text{N} - \text{H} \rightarrow \text{Cl} - \text{B} : \text{N} - \text{H}
\]

In the past, this reaction has been regarded as another confirmation of the rule of eight. Actually the limitation of eight electrons exists only for the second period, as we have already pointed out. In the third period, five more orbitals are available, more in the later periods. In a large number of compounds, some of the additional orbitals are used for coordinate covalent bond formation.

The electrical "strain" involved in the formation of a coordinate covalent bond may result in subsequent ionization to form such ions as

\[
\text{AlF}_6^{\text{3-}}, \text{SnCl}_6^{\text{2-}}, \text{ZrF}_7^{\text{3-}}, \text{Mo(CN)}_8^{\text{4-}}
\]

all of which fail to obey the rule of eight. The reaction between stannic chloride and hydrogen chloride may be represented formally as follows:

\[
\begin{align*}
\text{Cl} & \quad \text{Cl} & \quad \text{Cl} & \quad \text{Cl} & \quad \text{Cl} & \quad \text{H} \\
\text{Sn} + 2\text{H} - \text{Cl} & \rightarrow & \rightarrow & \rightarrow & \rightarrow & \rightarrow
\end{align*}
\]

Subsequent ionization may depend upon the environment; nevertheless, the tin atom is using six of its orbitals (containing twelve electrons in all) for bonding purposes.
At one time, the hydrogen bridge was regarded by some chemists as an example of coordination. Chelation and association, such as often occur between carboxyl groups, were thought to indicate the existence of 2-covalent hydrogen. The formulas were written to show the hydrogen bonded by two electron pairs, e.g., in the following formula to show the dimeric association of acetic acid:

```
O-H-O
```

CH₃-C
```
O-H-O
```

Two-covalent hydrogen is now regarded as unlikely, in view of the large energy difference between 1s and 2s orbitals (as shown in Fig. 1). Two electrons are the maximum for the hydrogen atom. The forces holding the hydrogen atoms to the carbonyl oxygens are probably electrostatic and should be represented by dotted lines:

```
O···H-O
```

CH₃-C
```
O···H-O
```

The original definition ¹⁸ of the coordinate covalent bond apparently was not intended to imply any fundamental difference between it and any other covalent bond. The word "coordinate" was used merely to indicate an imaginary difference in the way the covalent bond is formed.

In view of the controversies evident ¹⁹ in the literature, it seems best to retain the original definition. A coordinate covalent bond is merely any covalent bond both electrons of which were originally supplied by one of the two atoms involved. Once such a bond is formed, there may be no way of distinguishing it from any other covalent bond. (One advantage of this definition is that it might render meaningless such controversies as the one between Samuel and Wheland. ¹⁰)


Three examples should be sufficient to illustrate this inability to tell the difference between a covalent bond and a coordinate covalent bond after they have been formed, as we imagine their formation. When we write the (imaginary) equation

\[
\begin{array}{c}
\text{H} \\
\text{H:} \overset{\bullet}{\text{N}}: + \text{H}^+ \rightarrow \left[ \begin{array}{c}
\text{H} \\
\text{H:} \overset{\bullet}{\text{N}}: \text{H} \\
\text{H}
\end{array} \right]^+ \\
\end{array}
\]

we see in the ammonia molecule a potential coordinate bond. The nitrogen atom is going to supply both electrons to form a covalent bond. For our convenience, we call this bond a coordinate covalent bond. But upon considering the ammonium ion, we find that we are unable to distinguish this bond from the other bonds, which previously have been called simple covalent bonds (because each atom supplied one electron to the bond). This example is a familiar one, but, in view of the tendency of some authors to write coordinate covalent bonds with charges on each atom, it deserves stronger emphasis.

The second example is more striking. When we write the formula for water as

\[
\text{H:O:H}
\]

it rarely occurs to anyone to consider that the two H–O bonds are anything but covalent. Yet, if we write the abbreviated equation for neutralization in water,

\[
\text{H}^+ + \overset{\bullet}{\text{O}}: \text{H} \rightarrow \text{H:O:H}
\]

we see that one of the bonds is a coordinate covalent bond. Actually, either bond or both could be regarded as either covalent or coordinate. Since our choice depends only upon the immediately previous history of the molecule, it is obvious that there can be no real distinction between the two types of bonds. This is very well expressed in Gilman by the following graphic illustration:

\[
\overset{\text{X}}{\text{H}} \overset{\text{X}}{\text{Cl}} \quad \text{electron pairing} \quad \overset{\bullet}{\text{H}} \overset{\bullet}{\text{Cl}} \quad \text{coordination} \quad \text{H}^+ \quad \overset{\text{X}}{\text{Cl}}^{-1}
\]

Of course many compounds do exist in which the location of a coordinate covalent bond can be established. But in these also

---

we find that there is no inherent difference in the two types of bonds except in the way they are formed. In the compound

\[
\begin{array}{c}
\text{Cl} \\
\text{B-N-H} \\
\text{Cl}
\end{array}
\]

the coordinate covalent bond is located between the boron and nitrogen. But it is no different from the other covalent bonds. A large amount of experimental evidence \(^{16,19}\) supports this view. In summarizing some of it, obtained from dipole-moment measurements, Phillips, Hunter, and Sutton \(^{16}\) conclude, “True coordinate links appear, therefore, to be nearly of the same length to be expected for a normal, single covalency between the same two elements.” Furthermore, it is well known that, while coordinate covalent bonds are often highly polar, many ordinary covalent bonds are also highly polar.\(^{19}\)

In view of this brief summary of the present situation, there seems to be no justification for the designation of coordinate covalent bonds by means of plus and minus signs. There is no difference warranting such a distinction between “normal” and “coordinate” covalent bonds. The only difference is as stated in the original definition. A coordinate covalent bond is merely a covalent bond in which both electrons are thought of as being supplied by the same atom.

The chemistry of the coordinate covalent bond is largely the chemistry of acids and bases, the subject of this book.
Chapter 3

THE ELECTRONIC THEORY OF ACIDS AND BASES

1. The Theoretical Interpretation.

The experimental criteria of acid-base phenomena as listed in Chapter 1 are: (1) neutralization, (2) titration with indicators, (3) displacement, and (4) catalysis. When the chemical reactions between substances thus classified as acids or bases are examined in detail, the theoretical explanation of their fundamental nature becomes apparent. An acid is capable of accepting a share in a lone electron pair from a base to form a coordinate covalent bond. A base donates a share in a lone electron pair to the acid. The formation of the coordinate bond is the first step in neutralization reactions:

\[ \text{H}^+ + \text{O}^2- + \text{H}_{2}\text{O} \rightarrow \text{H}_2\text{O} + \text{O}^2- \]

Sometimes the product is a covalent compound. At other times, formation of the coordinate bond may be followed—or accompanied—by ionization, so that the product is a salt. For example, the compound \( \text{C}_5\text{H}_5\text{N}:\text{AlBr}_3 \) seems to be a typical salt. Both aluminum bromide and pyridine are covalent compounds. When
they are mixed, the white precipitate appears to have the characteristics (e.g., relatively high melting point) of a salt. The electrical "strain" produced by the formation of the coordinate bond may result in the ionization of one of the bromine atoms:

\[ \text{Br}^- + \text{Br}^- \rightarrow \text{Br}^- + \text{Br}^- + \text{Cl}^- \]

Sometimes formation of the coordinate bond and ionization may be regarded as simultaneous. When the hydrogen bridge was regarded as involving 2-covalent hydrogen, the reaction between hydrogen chloride and water might have been written in similar fashion:

\[ \text{H}_2\text{O}^- + \text{H}_2\text{O}^- \rightarrow \text{H}_2\text{O}^- + \text{H}_2\text{O}^- + \text{Cl}^- \]

The existence of the hypothetical intermediate addition compound in which the hydrogen bridge between the hydrogen chloride and water molecules involves 2-covalent hydrogen is now regarded as unlikely. Probably it would be better to represent the formation of the coordinate bond as taking place simultaneously with ionization, as follows:

\[ \text{H}_2\text{O}^- + \text{H}_2\text{O}^- \rightarrow \text{H}_2\text{O}^- + \text{H}_2\text{O}^- + \text{Cl}^- \]

Thus the electronic theory pictures the reaction in exactly the same manner as the proton-donor theory.

This example is only one of many which could be given to show that the electronic theory of acids and bases includes the proton theory as a special case. The simultaneous coordination and ionization pictured by the electronic theory is equivalent to the proton-transfer mechanism of the Brønsted theory. The bases of the
Brønsted theory accept protons because they have lone electron pairs that can be used in coordinate-bond formation. All these bases can, therefore, combine with other acceptor molecules in the same way as they do with the proton:

\[
\text{from a proton donor: } \quad H^+ + :N \rightarrow [H:N]^{+1}
\]

When we find that aluminum chloride and the proton behave in the same manner in titrations, displacement, and catalysis, as well as in these examples of neutralization, it is clear that we can no longer restrict the name acid to proton donors. To designate acids which owe their acidity to the proton, Lewis suggested the term hydrogen acid (or H-acid).

The electronic theory of acids also includes the theory of solvent systems as a special case. The following reactions are only two of many which could be listed to illustrate this statement.

1. Aluminum chloride and phosgene:

\[
\begin{align*}
:Cl: & + :Cl:Al + :Cl:C:Cl: \rightarrow :Cl:Al:Cl:C:Cl: \\
:Cl: & + :Cl:C:Cl: + :Cl: \rightarrow :Cl:Al:Cl:C:Cl: + :Cl:Al:Cl:C:Cl: \rightarrow [\ldots]^{-1} + :Cl:Al:Cl:C:Cl:^{+1}
\end{align*}
\]

2. Pyridine and water:

\[
\begin{align*}
\text{base: } & + \text{ acid: } \rightarrow \text{ solvent cation: } \\
N \rightarrow H:O:H & \rightarrow N:H^{+1} + O:H^{-1}
\end{align*}
\]

The addition of the acid in reaction 1 increases the concentration of solvent cations. The addition of the base in reaction 2 increases

\[1\text{ Lewis, } J. \text{ Franklin Inst., } 226, 293 (1938).\]
the concentration of solvent anions. The electronic theory gives a fundamental reason for this increased concentration of the ions of the solvent. Acids coordinate with the negative ion of the solvent because the negative ion has lone electron pairs to offer the acid. Bases coordinate with the positive ion of the solvent because the positive ion has a vacant orbital which can accept a lone electron pair from the base.

2. Reactions of Acids and Bases with the Solvent.

The properties of acids and bases with which we are most familiar from the study of water solutions depend to a great extent upon the presence of the solvent. For example, magnesium reacts slowly with hot water, liberating hydrogen. The reaction is much more rapid in acid solution. The difference must be due to the increased concentration of the solvent cation, the hydrogen ion. At first glance, Lewis’s theory seems to have little relation to this large body of experimental behavior with which we are so familiar.

For example, Walden has presented the following objections based upon this apparent lack of relationship: (1) the significance of dissociation constants and conductivity measurements would be destroyed; (2) the part played by the solvent would be eliminated; (3) the oppositeness of acids and bases toward indicators would appear to be “purely incidental observations.” We have already seen that the oppositeness of acids and bases toward indicators is by no means a purely incidental observation, and we proceed to show that the other two objections are as groundless.

Water may be regarded as the product of the neutralization of hydrogen ion by hydroxyl ion. The proton is an acid because it tends to accept an electron pair from a base to fill its orbital. The hydroxyl ion is a base because the oxygen atom can donate an electron pair to an acid. The formation of the coordinate bond between the proton and the hydroxyl ion is neutralization. The question whether the product is actually neutral, in the sense that the donor and acceptor properties of the oxygen and hydrogen atoms are balanced, is probably not of great importance. (Sidgwick believes that the oxygen is “more powerful” as a donor than the hydrogen is as an acceptor.) What is more important is that

\[ \text{References:} \]
\[ ^2 \text{Walden, } \textit{Salts, Acids, and Bases}, \text{ McGraw-Hill Book Company, New York, 1929.} \]
\[ ^3 \text{Sidgwick, } \textit{The Electronic Theory of Valency}, \text{ Oxford University Press, New York, 1927.} \]
the relative acidity of water can be compared with that of other solvents. For example, glacial acetic acid is more acidic and liquid ammonia more basic than water. In terms of the electronic theory, this means that acetic acid has a greater tendency to accept an electron pair than water has and that ammonia has a greater tendency to donate an electron pair.

When an acid is dissolved in a solvent, the initial reaction between the acid and the solvent depends primarily upon two factors: the strength of the acid (its tendency to accept an electron pair), and the basic strength of the solvent (its tendency to donate an electron pair). In a given solvent, the strength of the acid can be measured, within the limits of the "leveling effect" of Hantzsch (to be discussed later), by means of the equilibrium constant of the reaction with the solvent. For example, if glacial acetic acid, a typical covalent liquid which conducts an electric current poorly, reacts with water according to the equation

\[ \text{HC}_2\text{H}_5\text{O}_2 + \text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^{+1} + \text{C}_2\text{H}_3\text{O}_2^{-1} \]

the equilibrium constant,

\[ K = \frac{[\text{H}_3\text{O}^{+1}] \times [\text{C}_2\text{H}_3\text{O}_2^{-1}]}{[\text{HC}_2\text{H}_5\text{O}_2]} \]

serves as a measure of acid strength when compared with similar constants for other acids. This is true only if the acid is not too strong. For strong acids like hydrogen chloride, also a typical covalent compound, the reaction proceeds completely to the right in a solvent as basic as water.

Similar conclusions apply to acids that do not contain protons. If the reaction occurring when carbon dioxide, a weak acid, is dissolved in water is represented by the equation (the mechanism will be discussed below)

\[ \text{CO}_2 + 2\text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^{+1} + \text{HCO}_3^{-1} \]

the equilibrium constant,

\[ K = \frac{[\text{H}_3\text{O}^{+1}] \times [\text{HCO}_3^{-1}]}{[\text{CO}_2]} \]

may serve as a measure of the acid strength of the carbon dioxide. Strong acids like sulfur trioxide act in the same manner as hydrogen chloride. Sulfur trioxide accepts an electron pair from the
THE ELECTRONIC THEORY OF ACIDS AND BASES

Oxygen atom in the water molecule just as does the hydrogen in hydrogen chloride. The reaction

\[
\text{SO}_3 + 2\text{H}_2\text{O} \rightarrow \text{H}_3\text{O}^{+1} + \text{HSO}_4^{-1}
\]

proceeds extensively toward the right. The same considerations hold for such acids as boron chloride, aluminum chloride, or stannic chloride. The boron and aluminum atoms tend to accept an electron pair to complete their s and p orbitals. The tin atom tends to gain two electron pairs as in \(\text{H}_2\text{SnCl}_6\) to complete two of its d orbitals. There is no valid reason for calling the same type of reaction by two different terms: namely, ionization in the reactions of hydrochloric acid and acetic acid, and hydrolysis in the reactions of sulfur trioxide, carbon dioxide, and stannic chloride. The net result in both types of reaction is an increase in the concentration of the solvent cations. We shall see that this increased concentration of the solvent cations is responsible for most of the familiar properties of acids and bases in water and similar solvents. It is due to the tendency of an acid to accept an electron pair from a base in order to complete the characteristic stable electron configuration of the acid.

The actual mechanism may be regarded in either of two ways, as represented by simplified equations for the reaction between sulfur trioxide and water

\[
\begin{align*}
\text{O} &\quad :\text{O}\ldots\text{S} + :\text{O}\ldots\text{H}\rightarrow[ :\text{O}\ldots\text{S}:\text{O}\ldots\text{H}]^{-1} + \text{H}^{+1} \\
\text{O} &\quad :\text{O}\ldots\text{H}\rightarrow :\text{O}\ldots\text{S}:\text{O}\ldots\text{H} \rightarrow[ :\text{O}\ldots\text{S}:\text{O}\ldots\text{H}]^{-1} + \text{H}^{+1}
\end{align*}
\]

a direct reaction between the sulfur trioxide molecule and the hydroxyl ion; or

\[
\begin{align*}
\text{O} &\quad :\text{O}\ldots\text{H}\rightarrow :\text{O}\ldots\text{S}:\text{O}\ldots\text{H} \rightarrow[ :\text{O}\ldots\text{S}:\text{O}\ldots\text{H}]^{-1} + \text{H}^{+1}
\end{align*}
\]

a direct reaction between the sulfur trioxide molecule and the water molecule, followed by ionization. Either way the result is the same. If the acid is strong enough and if the solvent is basic enough, the concentration of the cation characteristic of the solvent is increased.
In order to show the reaction between carbon dioxide and water, it is necessary to write the formula of the carbon dioxide molecule in its "activated" form (discussed later in connection with secondary acids):

\[
\begin{align*}
\ce{::O:} & \quad \ce{::O:} \\
\ce{C} + \ce{::O:H} & \rightarrow \ce{C::O:H} \\
\ce{::O:} & \quad \ce{::O:}
\end{align*}
\]

\[[-1]
\ce{\cdot H} \quad \ce{\cdot H}
\]

\[\text{acid} \quad \text{base}
\]

Apparently the small number of ions formed is due to the small proportion of activated carbon dioxide molecules. Nevertheless, the concentration of solvent cations is increased in this case also.

The corresponding conclusion holds for bases dissolved in ionizable solvents. The solution contains a greater concentration of anions than the pure solvent. The strength of the base in a given solvent can be estimated from the equilibrium constant. For example, when pyridine is dissolved in water, the pyridine molecule acts as a base in donating an electron pair to the water molecule:

\[
\ce{C_6H_5N: + HOH \rightleftharpoons C_6H_5N:H^+ + OH^-}
\]

The equilibrium constant,

\[
K = \frac{[\ce{C_6H_5N:H^+}] \times [\ce{OH^-}]}{[\ce{C_6H_5N}]}.
\]

serves to measure the basic strength of the pyridine.

These examples, purposely chosen with water as the solvent, are enough to show that the part played by the solvent is not "deliberately eliminated." Dissociation constants and conductivity measurements still have as much significance as ever. Walden's objections obviously do not apply. When a sufficiently strong acid reacts with water, the concentration of the hydrogen ion is increased; when a sufficiently strong base reacts with water, the concentration of the hydroxyl ion is increased. In a given solvent the strength of an acid or a base, within limits, can be measured by its dissociation constant. There are also many examples in the literature to support this conclusion for solvents other than water.

The typically acid properties of aluminum chloride in phosgene are due to this increased concentration of solvent cations. Ger-
mann found that the conductivity of the aluminum chloride solution was less than that of the solution of Ca(AlCl₄)₂, and concluded that the acid is rather weak in phosgene. Aluminum chloride is an acid because it accepts an electron pair to fill the vacant p orbital of the aluminum atom. Phosgene is amphoteric and in this reaction is a base. No doubt the resulting cation will be solvated because of the strong tendency of the carbon atom to fill its valence shell. But showing this solvation adds nothing to the discussion, so we may write the equation as

\[
\text{AlCl}_3 + \text{COCl}_2 \rightarrow \text{AlCl}_4^{-1} + \text{COCl}^{+1}
\]

The equilibrium constant,

\[
K = \frac{[\text{COCl}^{+1}] \times [\text{AlCl}_4^{-1}]}{[\text{AlCl}_3]}
\]

will serve as a measure of acid strength when compared with the dissociation constants of other acids in phosgene. According to Germann's conductivity measurements \( K \) is small, so aluminum chloride is a fairly weak acid in phosgene as a solvent. Similar treatment can be given the results of other investigators.

Meerwein has shown that aluminum alkoxides, when dissolved in alcohols, increase the concentration of the solvent cation in the same manner as aluminum chloride does in phosgene:

\[
\begin{align*}
\text{R} & \quad \text{R} \\
\text{O} & \quad \text{O} \\
\text{R-O-Al} + & \quad \text{O-R} \\
& \quad \text{R-O-Al-O-R} \\
\end{align*}
\]

\[
\begin{align*}
\text{R} & \quad \text{R} \\
\text{O} & \quad \text{O} \\
\text{R-O-Al-O-R} & \quad + \text{H}^{+1}
\end{align*}
\]

\[\text{Germann, Science, 61, 70 (1925).}\]

\[\text{Meerwein, Ann. 455, 227 (1927).}\]
The concentration of the hydrogen ion in the solvent is increased. Other acids, such as boron trifluoride, also increase the hydrogen-ion concentration in organic acids. The work of Jander with sulfur dioxide, that of Smith and others with selenium oxychloride, and some of the work in liquid ammonia can be interpreted in a similar way, not only for acids but for bases as well.

The reason for the solvent-system definitions of Cady and Elsey (Chapter 1) is clear. Acids often do increase the concentration of solvent cations; bases often increase the concentration of solvent anions. However, this does not always happen. When acids react with solvents like ether and pyridine, ionization to give a cation characteristic of the solvent is unlikely. Usanovich has shown the similarity in electrical conductivity of solutions of such acids as the arsenic and antimony trichlorides in ether to a solution of sulfuric acid in ether. The conductance of the solutions is greater if the ether is replaced by a stronger base like pyridine. Arsenic trichloride reacts with pyridine with liberation of a large amount of heat, forming after evaporation of the excess pyridine a crystalline compound C$_5$H$_5$N$:$AsCl$_3$. It is a familiar fact that pyridine forms crystalline compounds with those salts which, according to Lewis's theory, are fairly strong acids, e.g., zinc chloride. In these examples ionization of the solvent is impossible. The reaction may be represented as follows:

\[ \text{C}_5\text{H}_5\text{N}$:$ + \text{AsCl}_3 \leftrightarrow \text{C}_5\text{H}_5\text{N}$:$\text{AsCl}_3 \leftrightarrow \text{C}_5\text{H}_5\text{N}$:$\text{AsCl}_2^{+1} + \text{Cl}^{-1} \]

The arsenic atom becomes more negative by gaining a share in the lone electron pair of the nitrogen atom. The resulting electrical strain can be relieved by the ionization of a chlorine atom. A similar situation holds for ether. An oxonium salt is formed in solution. If we regard the ether and pyridine as solvents, we see that no cation characteristic of the solvent is formed.

We might have drawn the same conclusion from the previously considered reaction between pyridine and water:

\[ \text{C}_5\text{H}_5\text{N}$:$ + \text{HOH} \leftrightarrow \text{C}_5\text{H}_5\text{N}$:$\text{H}^{+1} + \text{OH}^{-1} \]

If pyridine is considered the basic solvent and water the acid dissolved in it, there is again no splitting of the solvent to give a

---

6 Jander and Mesech, *Z. physik. Chem.*, A183, 255, 277 (1939); also preceding papers in this series.
cation characteristic of the solvent. The above examples illustrate the inadequacy of the idea that acids and bases can be defined in terms of ions. In certain solvents, acids increase the concentration of solvent cations and bases increase the concentration of solvent anions, but in other solvents they do not. These experimental facts do not affect the electronic theory of acids and bases because it is not stated in terms of ions.

3. Typical Reactions of Acids and Bases.

Typical properties of acids and bases are usually taken to be those that are observed in water solutions of acids and bases. Most of them are due to the increased concentration of solvent cation or solvent anion caused by the presence of the acid or base. The most familiar reactions dependent upon this effect probably are the reactions between the free elements and solutions of acids and bases; electrolysis; and the reactions of amphoteric substances. The first is the only one that requires further discussion before similar reactions in other solvents are considered.

Active metals like sodium and calcium react with pure water. Active non-metals like chlorine and sulfur also react with water, but the reactions are more complex. Chlorine reacts to give hydrochloric and hypochlorous acids. Sulfur reacts slowly, when heated with water, to give several products. These reactions may be considered due to the presence of hydrogen ions and hydroxyl ions. The hydrogen ions oxidize active metals and become free hydrogen. The reaction of hydroxyl ions with active non-metals is not so simple. This is probably due to the fact that the oxygen atom has a greater attraction for electrons than any other atom except fluorine. Many metals will reduce hydrogen ion, but only one non-metal, fluorine, will oxidize hydroxyl ion under ordinary circumstances. Chlorine and sulfur are unable to remove electrons completely from the hydroxyl ion.

These reactions proceed much more rapidly when the hydrogen-ion or hydroxyl-ion concentration is increased by the addition of an acid or base. The increase in rate seems to be a mass action effect; e.g., in the reaction

\[ \text{Mg} + 2\text{H}^+ \rightarrow \text{Mg}^{+2} + \text{H}_2 \]

increasing the concentration of the hydrogen ion will have the same effect whether it is done "directly" by adding hydrogen chloride or "indirectly" by adding sulfur trioxide to the water.
In like manner, it makes no difference how the hydroxyl-ion concentration is increased. It may be brought about by adding the ions directly through the addition of sodium hydroxide or it may be done indirectly by adding triethylamine to the water:

\[(C_2H_5)_3N + HOH \rightleftharpoons (C_2H_5)_3N:H^+ + OH^-\]

It is worth noting at this point that, in these typical reactions with metals (discussed further in Chapter 4), hydrogen ion and hydroxyl ion are not acting strictly as acid and base, respectively. The hydrogen ion acts as an oxidizing agent, removing electrons completely from the metals which react with it. The hydroxyl ion acts as a reducing agent toward the only element capable of removing electrons from it:

\[2F_2 + 4OH^- \rightarrow 4F^- + O_2 + 2H_2O\]

Such reactions, as well as those of electrolysis and of amphoteric behavior, have been observed in other solvents. Reactions that occur in ammonia, sulfur dioxide, acetic acid, hydrogen sulfide, hydrogen fluoride, phosgene, selenium oxychloride, alcohols, and sulfuric acid are analogous to those that take place in water. Some of them have been interpreted according to the solvent-systems theory; others, according to the proton theory. All of them may be understood more clearly on the basis of the electronic theory of acids and bases. Only a few examples will be discussed here.

A solution of aluminum chloride in phosgene dissolves metals with the liberation of carbon monoxide. According to Lewis's theory, the aluminum chloride is an acid and accepts an electron pair from the solvent. The resulting electrical "strain" favors the ionization which increases the concentration of solvent cation. The solvent cation oxidizes the metal, and carbon monoxide is produced

\[COCl^{+1} + Ca \rightarrow CO + Cl^{-1} + Ca^{+2}\]

or

\[CO(AlCl_4)_2 + Ca \rightarrow CO + Ca(AlCl_4)_2\]

This behavior is analogous to that of sulfur trioxide when dissolved in water. The sulfur trioxide accepts an electron pair from the solvent, and the concentration of the solvent cation is greatly increased. The solvent cation oxidizes the metal, and hydrogen is produced.

Similar reactions are observed in the other solvents listed above. Those in selenium oxychloride are particularly interesting since,
as with phosgene, no protons are involved. Although the reported value of its solvent conductance\(^9\) seems to be too high, the ion concentrations in pure selenium oxychloride are likely to be relatively great. When an acid like stannic chloride is dissolved in selenium oxychloride, the result is as expected. The solution reacts more vigorously with metals than the pure solvent does because of the increased concentration of solvent cation:

$$\text{SnCl}_4 + 2\text{SeOCl}_2 \rightarrow (\text{SeOCl}^{+1})_2, \text{SnCl}_6^{-2}$$

A similar effect has not yet been investigated in sulfur dioxide. One would expect that the same behavior would be observed. Sulfur monoxide should be produced when metals react with acid solutions of sulfur dioxide. One method of preparing sulfur monoxide is by the action of thionyl chloride on metals.\(^{10}\) The reactions occur at high temperatures (100° C. or above) and may be compared with the displacement of hydrogen from water by hot iron. At lower temperatures it is necessary to add acids (e.g., either sulfur trioxide or sulfuric acid) to the water before the iron will displace hydrogen from it. In the same way, we may venture to predict that sulfur monoxide will be displaced by metals from either thionyl chloride or sulfur dioxide at much lower temperatures if strong acids such as sulfur trioxide and boron trifluoride are added first.

The simplified notation (ignoring solvation and stages of ionization) for the dissociation of such solvents emphasizes the similarity in their reactions with metals:

\[
\begin{align*}
\text{HOH} & \rightleftharpoons \text{H}^{+1} + \text{OH}^{-1} \quad (\text{H}_2 \text{evolved}) \\
\text{COCl}_2 & \rightleftharpoons \text{CO}^{+2} + 2\text{Cl}^{-1} \quad (\text{CO} \text{evolved}) \\
\text{SOCl}_2 & \rightleftharpoons \text{SO}^{+2} + 2\text{Cl}^{-1} \quad (\text{evolution of SO predicted}) \\
2\text{SO}_2 & \rightleftharpoons \text{SO}^{+2} + \text{SO}_3^{-2} \quad (\text{evolution of SO predicted})
\end{align*}
\]

Corresponding reactions for bases, i.e., the action of the solvent anion as a reducing agent, are observed in liquid ammonia. Whereas, in water, fluorine is the only active non-metal which can oxidize hydroxyl ion, in ammonia the other halogens can oxidize the amide

\(^{11}\) Work to test these predictions will be undertaken soon.
ion. Iodine reacts with amide ion, just as fluorine reacts with hydroxyl ion:  

$$3\text{I}_2 + 6\text{NH}_2^- \rightarrow 6\text{I}^- + \text{N}_2 + 4\text{NH}_3$$

The reaction is more vigorous in a solution of potassium amide in ammonia than in ammonia alone. Other reactions with non-metals like sulfur, involving an increased concentration of solvent anion, are more complex. They are similar to those in water, since, like the hydroxyl ion, the amide ion does not readily lose electrons completely.

When electrolysis reactions in various solvents are considered, the conditions that determine which ion is to be discharged at either electrode are such that no definite conclusions can be drawn with regard to acid-base phenomena. For example, hydrogen is discharged at the cathode when an aqueous solution of an acid is electrolyzed, but hydrogen is also produced when an aqueous solution of sodium sulfate or of sodium hydroxide is electrolyzed. The most that can be said for the results of electrolysis is that they are consistent with what has just been said concerning cations and anions characteristic of the solvent. Carbon monoxide is discharged at the cathode when a solution of aluminum chloride in phosgene is electrolyzed. Selenium dioxide and selenium monochloride are produced at the cathode upon electrolysis of a solution of stannic chloride in selenium oxychloride.

Interpretation of the work of Bagster and Cooling according to the electronic theory of acids and bases would indicate the possibility that they unknowingly produced sulfur monoxide by electrolysis. Their interest in demonstrating the existence of the hydronium ion apparently caused them to overlook indications which might have led to the discovery of sulfur monoxide. When water was added to liquid sulfur dioxide and gaseous hydrogen bromide was passed in, two liquid layers were formed. Electrolysis of the sulfur dioxide layer yielded hydrogen at the cathode and bromine at the anode. Water collected at the cathode in proportion to the amount of silver deposited in a coulometer, but the amount of hydrogen discharged was less than expected, if the only ion being discharged was the $\text{H}_3\text{O}^{+1}$ ion. These facts indicate that hydrated $\text{SO}^{+2}$ ions were being discharged as well as hydronium ions. Bagster and Cooling were not able to account for the

smaller amount of hydrogen, but they did siphon off the sulfur dioxide layer from the water layer and try electrolysis of the sulfur dioxide alone. The conductance fell rapidly, and sulfur was deposited at the cathode, but no water. This behavior seems to indicate that the (solvated) $\text{SO}^{+2}$ ion was being discharged to form sulfur monoxide. Sulfur monoxide decomposes readily to form sulfur and sulfur dioxide.

Amphoteric reactions such as those of the hydroxides of aluminum and zinc also occur in other solvents. When potassium hydroxide is added to insoluble aluminum hydroxide in water, the following reaction takes place:

\[
\text{HO-Al} + \text{:O-H}^{-1} \rightarrow \text{HO-Al-OH}^{-1}
\]

The $\text{Al(OH)}_4^{-1}$ ion is soluble. It is formed because the aluminum hydroxide is acidic, in that the aluminum atom accepts an electron pair to fill its empty $p$ orbital. An analogous reaction takes place in liquid ammonia, when insoluble aluminum amide is dissolved by potassium amide:

\[
\text{Al(NH}_2)_3 + \text{NH}_2^{-1} \rightarrow \text{Al(NH}_2)_4^{-1}
\]

The zinc ion is also a fairly strong acid, and its insoluble compounds with solvent anions are often amphoteric. For example, in water, liquid ammonia, and glacial acetic acid the following reactions occur:

\[
\text{Zn(OH)}_2 + 2\text{OH}^{-1} \rightarrow \text{Zn(OH)}_4^{-2} \quad \text{(in water)}
\]

\[
\text{Zn(NH}_2)_2 + 2\text{NH}_2^{-1} \rightarrow \text{Zn(NH}_2)_4^{-2} \quad \text{(in ammonia)}
\]

\[
\text{Zn(C}_2\text{H}_3\text{O}_2)_2 + 2\text{C}_2\text{H}_3\text{O}_2^{-1} \rightarrow \text{Zn(C}_2\text{H}_3\text{O}_2)_4^{-2} \quad \text{(in acetic acid)}
\]

All the complex anions formed are soluble. Such reactions occur because zinc and aluminum ions are fairly strong acids, having considerable tendency to fill their empty orbitals.


15 Davidson, *Chem. Revs.*, 8, 175 (1931).
All the reactions so far discussed in this section have involved amphoteric solvents which, it is assumed, may ionize. Most of the reactions involve the same effect of acids and bases on the concentration of solvent cations or solvent anions that is observed in water. They are "typical" reactions only because they resemble those in water, with which we are more familiar. We would expect, therefore, that acid-base properties in inert solvents such as benzene, or in non-amphoteric solvents like pyridine, would not be "typical." In these solvents, there is no possibility of a "typical" increase in solvent cations or anions upon the addition of acids or bases. Ions are produced when acids and bases react with non-amphoteric solvents like pyridine and ether, but the ions formed are not "characteristic" of the solvent.

As a rule, no ions are produced when acids and bases are dissolved in relatively inert solvents such as benzene. The assumption of "protective coatings" to explain the absence of a reaction between metals and a solution of hydrogen chloride in benzene is unnecessary. In water, metals react more rapidly when an acid is present because of the increased concentration of solvent cations. The metal reacts slowly with water, even in the absence of the acid, but no such reaction takes place in benzene because there are no solvent cations with which the metal can react. The statement is sometimes made that acids dissolved in inert solvents do not react with carbonates. But the reaction with carbonates is one that does not necessarily depend on solvent cations. It depends on the strength of the acid required to displace the weaker acid, carbon dioxide, from its compound. Lewis has shown that a strong acid like boron trichloride will displace carbon dioxide from sodium carbonate in a mixture of carbon tetrachloride and acetone.

The fact that acids are not usually ionized in inert solvents permits a better determination of their relative strengths than is possible in water. In water, perchloric, hydriodic, hydrobromic, hydrochloric, and nitric acids are all practically 100 per cent ionized, thus appearing to be of equal strength. They all have such a strong tendency to accept an electron pair that the reaction with water goes to completion. Because of this "leveling effect," as Hantzsch called it, there can be no stronger acid than hydrogen ion in water. Any acid much stronger than hydrogen ion will displace it completely. One way in which the strengths of such acids can be compared is by measuring them in some inert solvent.
This has been done by Hantzsch, who found a great difference in the strengths of several strong hydrogen acids. Perchloric acid is the strongest, followed in order by hydriodic, hydrobromic, hydrochloric, and nitric acids.

We have seen that Walden's fears that Lewis would deliberately eliminate the important part played by the solvent in acid-base properties were groundless. Lewis's acids and bases, dissolved in suitable amphoteric solvents, have the "typical" properties of acids and bases. These "typical" properties are the properties with which we are familiar from our study of water chemistry. Now that we are beginning to branch out into other fields, we may expect to find increasingly that the electronic theory of acids and bases is the only one so far proposed that is at all adequate.

4. The Extent of Acid-Base Phenomena.

The measure of correlation to be effected by the electronic theory of acids and bases can be surmised by recalling that all the substances that Sidgwick called electron-pair acceptors and donors are really acids and bases. There is no need for any other name for them. The list of bases compiled by Brønsted and his followers is identical as far as it goes with that of Lewis, but their list of acids is limited because of their insistence upon the criterion of proton exchange. "To restrict the group of acids to those substances which contain hydrogen interferes as seriously with the systematic understanding of chemistry as would the restriction of the term oxidizing agent to substances containing oxygen." There is little doubt that the recognition of acids as electron-pair acceptors and bases as electron-pair donors will lead to as great a degree of systematization as did the recognition of oxidizing agents as electron acceptors and reducing agents as electron donors. Furthermore, the possibility of correlating the two types of phenomena now appears for the first time. This correlation will be attempted in the next chapter.

The rapid growth of the list of basic substances evident since the advent of the Brønsted theory has already made acceptable to most chemists the idea that almost any atom or group of atoms may act as a base if a strong enough acid is used. The electronic theory supplies the explanation. Most atomic groupings are surrounded by some lone pairs of electrons which can be utilized by

---

a sufficiently strong acid to form a coordinate bond. Even the "inert" gases can act as bases by donating one or more pairs of their outer octets to strong acids. Booth and Wilson,\(^1\) who demonstrated the existence of the boron trifluoride compounds with argon, suggested that more stable compounds would be obtained with krypton and xenon. Lewis\(^1\) also suggests that, since sulfur trioxide is in general a stronger acid than boron fluoride, even more stable addition compounds will be formed between sulfur trioxide and the "inert" gases.

The list of substances which can, under proper circumstances, behave as acids is perhaps not quite so extensive, but for practical purposes can be regarded so. Now that the octet theory has been abandoned, we are not so reluctant to admit that hundreds of compounds may act as acids by accepting shares in electron pairs even though they already have eight bonding electrons. Two examples of such very reactive compounds (strong acids) are stannic chloride and silicon tetrafluoride:

\[
\begin{align*}
\text{SnCl}_4 & \quad \text{SiF}_4 \\
\begin{array}{c}
\text{Cl} \\
\text{Cl} \\
\text{Si} \\
\text{F}
\end{array} & \quad \begin{array}{c}
\text{Cl} \\
\text{Cl} \\
\text{Si} \\
\text{F}
\end{array}
\end{align*}
\]

which form ions:

\[
\begin{align*}
\left[ \begin{array}{c} 
\text{Cl} \\
\text{Cl} \\
\text{Cl} \\
\text{Cl} \\
\text{Sn}
\end{array} \right]^{-2} & \quad \left[ \begin{array}{c} 
\text{F} \\
\text{F} \\
\text{F} \\
\text{F} \\
\text{Si}
\end{array} \right]^{-2}
\end{align*}
\]

In some ways we might think of acidic or basic tendencies as being due to the properties of atoms themselves, but this idea may be misleading, as the following example shows. Sulfur trioxide is a strong acid because in it the sulfur atom has a great tendency to accept an electron pair, but the sulfide ion is a fairly strong base. Davy was not so far from the truth when he said that acidity does not depend upon a particular element, but upon the arrangement of atoms.

\(^{17}\) Booth and Wilson, \textit{J. Am. Chem. Soc.}, 57, 2273 (1935).
The two preceding paragraphs have implied that many substances may act either as acids or as bases, depending upon the particular reaction under consideration. Amphoteric behavior seems to be much more widespread than was previously supposed.

Aluminum hydroxide

\[
\begin{array}{c}
\text{H} \\
\vdots \text{O} \\
\text{H} : \text{O} : \text{Al} \\
\vdots \vdots \text{O} \\
\vdots \text{H}
\end{array}
\]

is obviously amphoteric because the aluminum atom can accept a share in an electron pair from a base, or the hydroxy radical can donate a share in one of their electron pairs to an acid. But even strong acids like hydrogen chloride must also be considered amphoteric from the electronic viewpoint. The electronic formula

\[
\begin{array}{c}
\text{H} \\
\vdots \text{Cl}
\end{array}
\]

shows the possibility of basic behavior because of the three lone electron pairs on the chlorine atom. This prediction is borne out experimentally by the reactions of hydrogen chloride with the strong acids sulfur trioxide and stannic chloride to form addition compounds.\(^{13}\)

Frequently, consideration of the electronic structure will reveal whether a molecule is primarily acidic or basic and sometimes will give an idea as to its strength as an acid or base. It is found experimentally that these acids and bases, given sufficient difference in strength, combine "without impediment." Lewis calls such acids and bases primary.\(^1\) No heat of activation is required for the neutralization of a primary acid by a primary base. On the other hand, certain substances that experimentally behave like acids, e.g., carbon dioxide and organic acid halides, have electronic formulas which, as usually written, do not show the possibility of their acting as electron-pair acceptors. Such acids and bases are called secondary by Lewis. Neutralization of some of these substances is measurably slow. Their neutralization apparently requires heat of activation. Lewis suggests that such second-

ary acids and bases are not acids or bases in their normal states of lowest energy, as represented by the electronic formulas usually written, but may become acids and bases through excitation. These substances act like acids and bases, except that their neutralization is slow, but their electronic formulas do not indicate such behavior. Perhaps some of the difficulty lies in the formulas.

It has been known for some time that the properties of the carbon-oxygen double bond are not well represented by the usual formula. Lowry suggested in 1923 that, instead of representing the carbonyl group as

\[ R_2C::O: \]

the formula

\[ R_2C:O: \]^{(+) \cdot (-)} \]

should be used. (The plus and minus signs in parentheses indicate electrical dissymmetry; they are not intended to represent unit charge or to indicate valence.) Since the carbon atom in the second formula has only six electrons the possibility of acidic behavior is indicated. Basic behavior should be evident also through the electron pairs on the oxygen. Compounds containing such groups should be amphoteric. The familiar addition reactions of aldehydes confirm this conclusion. The first step in such reactions can be represented by the following equations:

\[
\begin{align*}
H & \quad :\text{Cl}: & H & \quad :\text{Cl}: \\
\text{R}:\text{C}:O: + \text{Al}:\text{Cl}: & \rightarrow \text{R}:\text{C}:O:\text{Al}:\text{Cl}: \\
& \quad :\text{Cl}: & \quad :\text{Cl}: \\
\text{base} & \quad \text{acid} & \\
\text{H} & \quad \text{H} & \quad \text{H} & \quad \text{H} \\
\text{R}:\text{C} + :\text{N}:\text{H} & \rightarrow \text{R}:\text{C} : \text{N}:\text{H} \\
& \quad :\text{O}: & \quad :\text{O}: & \quad \text{H} \\
\text{acid} & \quad \text{base} & 
\end{align*}
\]

The acid halides exhibit more definite "typical" acidic behavior in one respect, because they can react with water to increase the concentration of hydrogen ions:
Carbon dioxide is another example of a "secondary" acid. Assuming that the activated molecule can be represented by a formula similar to that just used for the carbonyl group, the reaction between carbon dioxide and water could be written:

\[
\begin{align*}
\text{R}-\overset{\cdot}{C} + \overset{\cdot}{:O}:H & \rightarrow \text{R}-\overset{\cdot}{C}:\overset{\cdot}{:O}:H + H^+ + \overset{\cdot}{:Cl}^- \\
\overset{\cdot}{:O} & \quad \overset{\cdot}{:O} & \quad \overset{\cdot}{:O} \\
\text{acid} & \quad \text{base} & \\
\downarrow & \\
\text{R}:C:\overset{\cdot}{:O}:H & \\
\overset{\cdot}{:O} & \\
\end{align*}
\]

Another type of "secondary" behavior is evident when amphoteric molecules partially or wholly neutralize each other, or when molecules are neutralized by the solvent in which they are placed. For example, the acid A may be dissolved in the weak base B and be neutralized, in that the stable electron configuration of the acidic atom has been attained by acceptance of electron pairs from the base. Yet, if a stronger base B\(^1\) is added, the stronger base will replace the weaker one in combination with the acid:

\[
B^1 + AB \rightarrow AB^1 + B
\]

Such reactions will be discussed in greater detail in Chapter 8. They usually require activation, but in many reactions the activation energy is small enough to be ignored. This seems to be true of the reactions of most hydrogen acids.

Another extension of the idea of acids and bases involves acidic and basic radicals in organic compounds. For example, the familiar ortho-para-directing groups for substitutions in the benzene ring are basic. They have electron pairs that they can share with a neighboring atom. The meta-directing groups are acidic. They can share an electron pair possessed by a neighboring atom.
The action of the acid and basic radicals in directing substituents will be considered in Chapter 5.

Many compounds which at first glance seem to offer no possibility of behaving as acids or bases may be forced to do so by reaction with strong bases or acids. For example, in the sulfonation of benzene by sulfur trioxide, the simplest explanation would seem to be that the $\text{SO}_3$ molecule coordinates with two of the electrons from the benzene molecule as follows:

Thus a very strong acid can force even benzene to behave as a base. Similar behavior induced by other strong acids like aluminum chloride and boron trifluoride can be postulated to clear up a great deal of the mystery which until quite recently surrounded catalysis of the type observed in Friedel-Crafts reactions. This kind of catalysis will be dealt with in the concluding chapters.

One striking illustration of the simplification of chemistry made possible by the Lewis theory occurs in the hydrolysis of salts. According to the electronic theory, any sufficiently strong acid or base will increase the concentration of cation or anion in an ionizing solvent. When a test with litmus shows a salt solution to be acidic or basic, a single simple equation is often all that is necessary by way of explanation. For example, the hydrolysis of zinc chloride may be represented by the equation

$$\text{Zn}^{+2} + \text{HOH} \rightarrow \text{ZnOH}^{+1} + \text{H}^{+1}$$

The explanation is simply that the zinc ion in zinc chloride is acidic.

The solubility of some salts in pyridine and other amines of low dielectric constant is more readily understood in terms of acid-base behavior. The metallic ion, being acidic, coordinates molecules of solvent about it. Silver salts in pyridine, ethylenediamine,\(^\text{19}\) and other amines\(^\text{20}\) show "abnormal" conductance curves.


A possible explanation lies in the fact that the silver ion is a fairly strong acid. For example, Bromley and Luder \(^{10}\) measured the conductance of potassium iodide, silver nitrate, and silver iodide in ethylenediamine. It was hoped that potassium nitrate could also be measured to permit comparison of \(\Lambda_0\) values, but it proved too insoluble. Figure 3 and Table 1 summarize the data (\(a\) is the "ion size").

**TABLE 1**

<table>
<thead>
<tr>
<th>SALT</th>
<th>(\Lambda_0)</th>
<th>(K \times 10^4)</th>
<th>(a \times 10^8)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KI</td>
<td>69.2</td>
<td>6.02</td>
<td>5.34</td>
</tr>
<tr>
<td>AgNO(_3)</td>
<td>61.4</td>
<td>8.07</td>
<td>5.98</td>
</tr>
<tr>
<td>AgI</td>
<td>48.8</td>
<td>0.431</td>
<td>3.84 *</td>
</tr>
</tbody>
</table>

* This value has little meaning, since other than coulomb forces are involved.

It is not surprising that the solubility of potassium nitrate is too low for conductance measurements. The potassium ion has very little tendency to coordinate with bases. Silver nitrate (be-
cause of the acidic silver ion) and potassium iodide (because of the amphoteric nature of the iodide ion\textsuperscript{21}) behave about as expected. Both the silver and the iodide ions are probably solvated by the highly basic solvent. "Normal" conductance behavior is observed in that greater ion size as indicated by lower $\Lambda_0$ is associated with a larger value of $K$ (Table 1).

At first glance, the behavior of silver iodide seems rather surprising. The low $\Lambda_0$ value indicates larger ions, as would be expected, since both ions should be solvated rather than only one, as in potassium iodide and silver nitrate. But $K$, instead of being larger, is actually only about one-fifteenth the values for the other two salts. The explanation may lie in the fact that the model, upon which the expectation of a high $K$ value accompanying low mobility is based, assumes only the presence of coulomb forces between the ions.\textsuperscript{22} In silver iodide additional forces are present which make $K$ less than predicted when coulomb forces alone are involved. These forces are of the acid-base type. The amphoteric nature of the iodide ion is beginning to be appreciated and is here substantiated. The results for silver nitrate and potassium iodide indicate that both silver and iodide ions are acting as acids in their association with the highly basic solvent. But toward the acidic silver ion the iodide ion acts as a base, tending to donate a pair of electrons to give covalent character to the bond between the two ions. The presence of these additional forces decreases $K$ to a value far below that expected from the low value of $\Lambda_0$. This conclusion is supported by the fact that, while AgCl and AgBr crystallize in ionic lattices, crystalline AgI has the zinc blende structure, in which the atoms are held together by covalent bonds.\textsuperscript{23}

Tables 2 and 3 summarize some of the simpler fundamentals of the electronic theory of acids and bases.

\textsuperscript{21} The basic nature of the iodide ion has long been accepted by followers of Brønsted. The acidic nature of the iodide ion is due to its ability to take on additional electron pairs. The ion itself contains only eight electrons, but the iodine atom is capable of holding more valence electrons, as shown by the formulas of the covalent compounds $\text{ICl}_5$ and $\text{IF}_7$ discussed in Chapter 2. Additional information on this point may be obtained from the review by R. B. Sandin (Chem. Revs., 32, 249 [1943]).

\textsuperscript{22} Fuoss and Kraus, J. Am. Chem. Soc., 55, 1019 (1933).

### TABLE 2

**ACID-BASE REACTIONS IN GENERAL**

(H⁺ used for simplicity)

<table>
<thead>
<tr>
<th>Description</th>
<th>Acid</th>
<th>Base</th>
<th>Product</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Neutralization</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>The water definition is a special case.</td>
<td>H⁺⁺¹</td>
<td>OH⁻⁻¹</td>
<td>HOH</td>
</tr>
<tr>
<td>The solvent-systems definition is a special case.</td>
<td>(COCl)⁺⁺¹</td>
<td>Cl⁻⁻¹</td>
<td>COCl⁻⁻¹</td>
</tr>
<tr>
<td>The Brønsted definition is a special case.</td>
<td>HCl</td>
<td>NH₃</td>
<td>NH₄Cl</td>
</tr>
<tr>
<td>The electronic theory includes all three.</td>
<td>Electrón-pair acceptor</td>
<td>Electron-pair donor</td>
<td>Neutralization—formation of a coordinate bond</td>
</tr>
<tr>
<td>And many others, e.g.</td>
<td>BCl₃</td>
<td>(C₂H₅)₃N</td>
<td>(C₂H₅)₃N → BCl₃</td>
</tr>
<tr>
<td></td>
<td>AlCl₃</td>
<td>(C₂H₅)₂O</td>
<td>(C₂H₅)₂O → AlCl₃</td>
</tr>
<tr>
<td></td>
<td>SO₂</td>
<td>C₂H₅N</td>
<td>C₂H₅N → SO₃</td>
</tr>
<tr>
<td></td>
<td>CO₂</td>
<td>CaO</td>
<td>CaCO₃</td>
</tr>
<tr>
<td></td>
<td>Ca⁺⁺²</td>
<td>OH⁻⁻¹</td>
<td>Ca(OH)₂</td>
</tr>
<tr>
<td></td>
<td>Ag⁺⁺¹</td>
<td>NH₃</td>
<td>Ag(NH₃)₂⁺⁺¹</td>
</tr>
<tr>
<td><strong>Displacement</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A strong acid may displace a weaker one from combination with a base.</td>
<td>BCl₃</td>
<td>Na₃CO₃</td>
<td>CO₃ + [Na₃O → BCl₃]</td>
</tr>
<tr>
<td></td>
<td>H⁺⁺¹</td>
<td>Al(OH)₃</td>
<td>Al⁺⁺³ + HOH</td>
</tr>
<tr>
<td></td>
<td>H⁺⁺¹</td>
<td>Cu(NH₃)₄⁺⁻²</td>
<td>Cu⁺⁺² + NH₃⁺⁻¹</td>
</tr>
<tr>
<td></td>
<td>C₂H₅N → SO₃</td>
<td>NH₃</td>
<td>C₂H₅N + [H₂N → SO₃]</td>
</tr>
<tr>
<td></td>
<td>H₂C₂H₅O₂</td>
<td>OH⁻⁻¹</td>
<td>C₂H₅O₂⁻⁻¹ + HOH</td>
</tr>
<tr>
<td></td>
<td>NH₄⁺⁺¹</td>
<td>OH⁻⁻¹</td>
<td>NH₃ + HOH</td>
</tr>
<tr>
<td><strong>Catalysis</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Many acids catalyze organic reactions.</td>
<td>AlCl₃, BF₃, HF, SO₃</td>
<td>E.g., reaction between alcohols and benzoyl chloride more rapid in pyridine</td>
<td></td>
</tr>
<tr>
<td>Bases also catalyze organic reactions.</td>
<td></td>
<td>Acid increases concentration of solvent cations</td>
<td></td>
</tr>
<tr>
<td>The presence of acids increases speed of reaction of ionizing solvents with metals.</td>
<td></td>
<td>SO₃ + HOH → H⁺⁺¹ + HSO₄⁻⁻¹; AlCl₃ + COCl₂ → COCl⁺⁺¹ + AlCl⁻⁻¹</td>
<td></td>
</tr>
<tr>
<td>Bases also may speed solvent reactions similarly.</td>
<td></td>
<td>Base increases concentration of solvent anions</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>NH₃ + HOH → OH⁻⁻¹ + NH₄⁺⁺¹; (C₂H₅)₃N + COCl₂ → (C₂H₅)₃NCOCl⁺⁺¹ + Cl⁻⁻¹</td>
</tr>
</tbody>
</table>
### TABLE 3

**Acid-Base Reactions in Ionizing Solvents**

*(Hydration of the solvent cation usually not shown)*

<table>
<thead>
<tr>
<th>Description</th>
<th>Acid</th>
<th>Base</th>
<th>Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ionization</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>According to the theory of solvent systems</td>
<td>AlCl₃</td>
<td>COCl₂</td>
<td>COCl⁺⁺ + AlCl₄⁻⁻</td>
</tr>
<tr>
<td>According to the Brønsted theory</td>
<td>HCl</td>
<td>HOH</td>
<td>H₃O⁺⁺ + Cl⁻⁻</td>
</tr>
<tr>
<td>Both are special cases of the electronic theory</td>
<td>Electron-pair acceptor</td>
<td>Electron-pair donor</td>
<td>Coordinate bond formation involving ionization</td>
</tr>
<tr>
<td>Other examples:</td>
<td>CO₂</td>
<td>HOH (s)</td>
<td>H⁺⁺ + HCO₃⁻⁻</td>
</tr>
<tr>
<td>(s denotes solvent.) Note that an acid reacting with an ionizing solvent increases the concentration of solvent cations; a base increases the anion concentration.</td>
<td>HOH (s)</td>
<td>NH₃</td>
<td>OH⁻⁻ + NH₄⁺⁺</td>
</tr>
<tr>
<td>Hydrolysis of Salts</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reaction of cation acid with solvent (when anion is weak base, e.g., Cl⁻⁻)</td>
<td>NH₄⁺⁺</td>
<td>HOH</td>
<td>H₃O⁺⁺ + NH₃</td>
</tr>
<tr>
<td>Reaction of anion base with solvent (when cation is weak acid, e.g., Na⁺⁺)</td>
<td>Zn⁺⁺⁺</td>
<td>HOH</td>
<td>H⁺⁺ + ZnOH⁻⁻</td>
</tr>
<tr>
<td>Amphoteric Behavior</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water is amphoteric (it can donate or accept an electron pair).</td>
<td>HOH</td>
<td>NH₃</td>
<td>NH₄⁺⁺ + OH⁻⁻</td>
</tr>
<tr>
<td>Aluminum hydroxide is amphoteric (donates or accepts an electron pair).</td>
<td>HCl</td>
<td>HOH</td>
<td>H₅O⁺⁺ + Cl⁻⁻</td>
</tr>
<tr>
<td>Complex Ions</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Al(OH)₃⁺⁺</td>
<td>OH⁻⁻</td>
<td>Al(OH)₃⁻⁻</td>
</tr>
<tr>
<td></td>
<td>H⁺⁺⁺</td>
<td>Al(OH)₃⁺⁺</td>
<td>Al⁺⁺⁺ + HOH</td>
</tr>
<tr>
<td></td>
<td>Ag⁺⁺⁺</td>
<td>CN⁻⁻</td>
<td>Ag(CN)₃⁻⁻</td>
</tr>
<tr>
<td></td>
<td>Cu⁺⁺⁺</td>
<td>NH₃</td>
<td>Cu(NH₃)₄⁺⁺</td>
</tr>
<tr>
<td></td>
<td>SnCl₄⁺⁺</td>
<td>Cl⁻⁻</td>
<td>SnCl₄⁻⁻²</td>
</tr>
<tr>
<td></td>
<td>As₂S₅⁺⁺</td>
<td>S⁻⁻</td>
<td>As₂S₅⁻⁻³</td>
</tr>
</tbody>
</table>
Although consistent application of the electronic theory of acids and bases simplifies and systematizes a great deal of chemistry, the amount of correlation can be increased still further by developing the relationship between acids and bases on the one hand and oxidizing and reducing agents on the other. This will be done in the following chapter.
Chapter 4

ELECTROPHILIC AND ELECTRODOTIC REAGENTS

1. The Resemblance of Oxidation-Reduction to Acid-Base Reactions

The experimental relationship between acids and oxidizing agents is a close one. In fact, as has been pointed out in Chapter 3, the “typical” effect of an acid solution on metals is due to the oxidizing action of the solvent cation; for example:

\[ \text{Mg} + 2\text{H}^+ \rightarrow \text{Mg}^{2+} + \text{H}_2 \] (in water)

and

\[ \text{Ca} + \text{CO}^+ \rightarrow \text{Ca}^{2+} + \text{CO} \] (in phosgene)

The acidic cation of the solvent in this particular type of reaction is behaving not as an acid but as an oxidizing agent. The action of basic anions upon non-metals (already referred to) is comparable. Hydroxyl ions in water, and amide ions in liquid ammonia, reduce active non-metals, for example:

\[ 2\text{F}_2 + 4\text{OH}^{-1} \rightarrow 4\text{F}^{-1} + \text{O}_2 + 2\text{H}_2\text{O} \] (in water)

and

\[ 3\text{I}_2 + 6\text{NH}_2^{-1} \rightarrow 6\text{I}^{-1} + \text{N}_2 + 4\text{NH}_3 \] (in ammonia)

These experimental data show the close relationship between acids and oxidizing agents on the one hand and bases and reducing agents on the other, but occasionally a comparison has been attempted which, at first glance, seems to lead to an opposite conclusion.

A formal analogy is worked out as follows: A reducing agent is an electron donor. An acid is a proton donor. An acid, therefore,
resembles a reductant in that both are donors. An acid cannot
donate protons unless a base is present, nor can a reducing agent
donate electrons unless an oxidizing agent is present to accept
them. Both types of reactions are mutual affairs between the
acid and the base, or between the oxidant and the reductant. For
example, the equation
\[
\begin{align*}
\text{reductant}_1 + \text{oxidant}_2 & \rightleftharpoons \text{oxidant}_1 + \text{reductant}_2 \\
\text{Zn}^0 + \text{Cu}^{+2} & \rightleftharpoons \text{Zn}^{+2} + \text{Cu}^0
\end{align*}
\]
is similar to the type equation of the Brønsted theory:
\[
\begin{align*}
\text{acid}_1 + \text{base}_2 & \rightleftharpoons \text{base}_1 + \text{acid}_2 \\
\text{HCl} + \text{H}_2\text{O} & \rightleftharpoons \text{H}_3\text{O}^{+1} + \text{Cl}^{-1}
\end{align*}
\]
This mutuality is regarded as a confirmation of the analogy.

Actually the analogy between the two equations depends upon
the fact that the typical acid-base reaction of the Brønsted theory
represents only one of the four experimental criteria of acid-base
phenomena—displacement. (Displacement will be considered in
detail in Chapter 8.) The whole analogy is only a formal com­
parison, and, of course, its advocates do not claim that it shows
any definite relationship between acid-base and oxidation-reduc­
tion reactions.

Usanovich attempted to explain the relationship by including
oxidation as a special case of acidic behavior. Reduction was
considered a special case of basic behavior. Chlorine was listed as
an acid and sodium as a base. This classification has more experi­
mental justification than would appear at first glance. Sodium,
when reacting with water, increases the concentration of solvent
anions as do bases when dissolved in many amphoteric solvents:
\[
2\text{Na} + 2\text{H}_2\text{O} \rightarrow 2\text{OH}^{-1} + 2\text{Na}^{+1} + \text{H}_2
\]
The corresponding property of acids in amphoteric solvents,
namely, the increase in solvent cation concentration, is exhibited
by chlorine:
\[
\text{Cl}_2 + \text{H}_2\text{O} \rightarrow 2\text{H}^{+1} + \text{Cl}^{-1} + \text{OCl}^{-1}
\]
On the other hand, some oxidizing and reducing agents do not
have such properties of acids and bases. For example, the per­
manganate ion is a strong oxidizing agent, but it is not an acid.

It seems, therefore, that a more general classification is necessary, one that will include both types of behavior.

2. Electronic Reactions.

An inclusive classification can be made by dividing the reagents that take part in acid-base and oxidation-reduction reactions into two types. Electrophilic reagents are those that gain electrons in the reaction under consideration. Electrodotic reagents are those that lose electrons in the reaction under consideration. The term *electrophilic* has been in use for some time to refer to reagents such as chlorine molecules and hydrogen ions which have an attraction for electrons. Those reagents which, like sodium, give up electrons readily have been called *nucleophilic*. *Electrophilic* seems to be a good word to retain, since it graphically describes the theoretical action of the reagents to which it applies. But *nucleophilic* is an unfortunate choice. It is difficult to picture sodium or other strong reducing agents as actually nucleophilic. A term that indicates ability to give up electrons readily is more appropriate. *Electrodotic* (Gr. διδομεν, to give) was the term originally suggested, but, as pointed out by Professor N. F. Hall, *electrodotic* seems better etymologically. Some reagents may be electrophilic under one set of conditions but electrodotic under another set. This double behavior is already familiar as far as acid-base phenomena are concerned; the word used to describe it is “amphoteric.” For lack of a better one, we might as well use the same word to apply to oxidizing and reducing agents. When a reagent is acting as an acid or an oxidizing agent, it is electrophilic. When it is acting as a base or a reducing agent, it is electrodotic.

Both acids and oxidizing agents are electron acceptors. An acid accepts a share in an electron pair held by a base; an oxidizing agent takes over completely the electrons donated by a reducing agent. Both are electrophilic and in many cases may act either as acids or as oxidizing agents:

\[
\begin{align*}
\text{acid} & : & H^+1 + :O:H^{-1} \rightarrow H:O:H \\
\text{oxidant} & : & H^+1 + \text{Na} \rightarrow \text{Na}^+1 + \frac{1}{2}\text{H}_2
\end{align*}
\]


Both bases and reducing agents are electron donors. A base donates a share in an electron pair to an acid; a reducing agent loses electrons completely to an oxidizing agent. Both are electrodotic and may act either as bases or as reducing agents:

\[ \text{H}_2\text{O}^+: -1 + \text{H}^+ \rightarrow \text{H}_2\text{O}:H \]

\[ 2\text{H}_2\text{O}^+: -1 + \text{I}_2 \rightarrow 2\text{I}^+-1 + \frac{1}{2}\text{O}_2 + \text{H}_2\text{O} \]

A few reagents are listed in Table 1 according to some manifestations of their electrophilic and electrodotic tendencies. The most active oxidizing agent, fluorine, is placed at the top of the list of electrophilic reagents and one of the strongest acids, sulfur trioxide, is at the bottom. The order is roughly that of increasing acid strength downward. The list of electrodotic reagents is arranged correspondingly, with the strongest reducing agent, cesium, at the top and the strongest bases toward the bottom. Apparently some of the strongest oxidizing agents do not act as
acids, nor do some of the strongest acids act as oxidizing agents. A similar relationship appears to hold for electrodotic reagents.

Discussion of a few examples from Table 1 should help to clarify its significance. As shown above, hydrogen ion acts as an acid in accepting a share in an electron pair from the hydroxyl ion, but toward an active reducing agent such as sodium it behaves not as an acid but as an oxidant. Silver ion acts as an acid in accepting a pair of electrons from hydroxyl ion to form silver hydroxide

$$\text{Ag}^{+1} + \text{H}^{-1} \rightarrow \text{AgO} : \text{H}$$

which decomposes to silver oxide. It also acts as an acid in reacting with ammonia or cyanide ion to form complex ions:

$$\text{Ag}^{+1} + 2\text{NH}_3 \rightarrow \text{Ag(NH}_3)_2^{+1}$$
$$\text{Ag}^{+1} + 2\text{CN}^{-1} \rightarrow \text{Ag(CN)}_2^{-1}$$

Toward reducing agents the silver ion acts not as an acid but as an oxidizing agent:

$$2\text{Ag}^{+1} + \text{Cu}^{0} \rightarrow 2\text{Ag}^{0} + \text{Cu}^{+2}$$

Hydroxyl ion acts as a base in sharing a pair of electrons with hydrogen ion to form water, but it acts as a reducing agent in giving up electrons to the oxidant fluorine. Amide ion behaves as a base toward ammonium ion

$$\text{NH}_2^{-1} + \text{NH}_4^{+1} \rightarrow 2\text{NH}_3$$

but as a reductant toward iodine:

$$6\text{NH}_2^{-1} + 3\text{I}_2 \rightarrow 6\text{I}^{-1} + 4\text{NH}_3 + \text{N}_2$$

Cyanide and sulfide ions react as bases toward water

$$\text{CN}^{-1} + \text{H}_2\text{O} \rightarrow \text{HCN} + \text{OH}^{-1}$$
$$\text{S}^{-2} + 2\text{H}_2\text{O} \rightarrow \text{H}_2\text{S} + 2\text{OH}^{-1}$$

but as reductants toward oxidizing agents. For example, the ease with which sulfide ion is oxidized by ferric ion is well known:

$$\text{S}^{-2} + 2\text{Fe}^{+3} \rightarrow \text{S}^{0} + 2\text{Fe}^{+2}$$
Iodide ion is a very weak base but does react with some acids to give stable complex ions, e.g.,

$$4I^- + Hg^{+2} \rightarrow HgI_4^{-2}$$

On the other hand, it is a good reducing agent, being easily oxidized to free iodine:

$$2I^- + Cl_2 \rightarrow 2Cl^- + I_2$$

3. Amphoteric Behavior.

Four of the substances included in Table 1—water, hydrogen chloride, hydroquinone, and stannous ion—appear in both columns. This is entirely in accord with the familiar behavior of acids and bases described by the term "amphoteric." We propose to use the same term to apply to the corresponding behavior when it is encountered in oxidation-reduction reactions.

It is an experimental fact that the same substance may under properly chosen conditions act as an acid, a base, an oxidizing agent, or a reducing agent. For example, water acts as an acid toward ammonia, as a base toward sulfur dioxide, as an oxidizing agent toward active metals, and as a reducing agent toward fluorine (Table 2). Water is not only amphoteric in the usual sense but also is both an oxidizing agent and a reducing agent. The same is true of many other substances besides water. Most reagents cannot be arbitrarily classified as acid or base, or as oxidizing or reducing agent. Their behavior depends upon the specific circumstances of a particular reaction. Therefore, the words electrophilic and electrodotic are relative terms, depending upon conditions. They refer to the behavior of a substance as it acts in the particular reaction under consideration.

Continuing with the other substances listed on both sides of Table 1, hydrogen chloride is ordinarily thought of as a strong acid, but it behaves as a base toward sulfur trioxide or stannic chloride (Chapter 3). On the other hand, the hydrogen ion acts

Note that, as shown in Table 2, a reducing agent as well as a base may increase the concentration of solvent anions. Also an oxidizing agent as well as an acid may increase the concentration of solvent cations. Thus, according to the theory of solvent systems, sodium would be a base and fluorine an acid. This conclusion is one more illustration of the inadequacy of the idea that acids and bases can be defined in terms of ions.
Odd-electron molecules (often called free radicals) can behave either as electrophiles or electrodotes or even as both simultaneously. Probably the outstanding examples are the triaryl methyl
compounds. For example, hexaphenylethane is partially dissociated in benzene solution to give triphenylmethyl:

\[
\begin{array}{c}
\text{C} \\
\text{phenyl} \\
\end{array}
\]

in which the central carbon atom has seven electrons. Nevertheless, it is an electrically neutral molecule comparable to another odd-electron molecule, namely, nitrogen dioxide. Probably triphenylmethyl should no more be called a "free radical" than nitrogen dioxide. They are both similar in associating into double molecules and for the same reason. Each molecule has one unshared electron.

When two triarylmethyl molecules associate, each one is acting simultaneously as an electrophilic and as an electrodotic reagent to form a covalent bond:

\[
\begin{array}{c}
\text{C} \\
\text{phenyl} \\
\end{array} + \begin{array}{c}
\text{C} \\
\text{phenyl} \\
\end{array} \rightarrow \begin{array}{c}
\text{C} \\
\text{phenyl} \\
\end{array} \begin{array}{c}
\text{C} \\
\text{phenyl} \\
\end{array}
\]

On the other hand, the triarylmethyl molecule may act as a reducing agent (an electrodot) to form a carbonium ion:

\[
\begin{array}{c}
\text{C} \\
\text{phenyl} \\
\end{array} \rightarrow \begin{array}{c}
\text{C} \\
\text{phenyl} \\
\end{array} + 1e
\]

or it may act as an oxidizing agent (an electrophile) to form a carbanion:

\[
\begin{array}{c}
\text{C} \\
\text{H} \\
\text{H} \\
\end{array} + 1e \rightarrow 
\begin{array}{c}
\text{C} \\
\text{H} \\
\text{H} \\
\end{array}^{-1}
\]

Obviously odd-electron molecules must also be classified as amphoteric.

5. Classification of Chemical Reactions.

In view of the amount of systematization and simplification in chemistry made possible through the classification of substances as electrophilic or electrodotic according to the way they behave in specific reactions, a natural question to bring up is: How does this new classification of reactions affect the older one? In the past, chemical reactions have been listed as (1) combination, (2) decomposition, (3) displacement, (4) metathesis, (5) neutralization, and (6) oxidation-reduction. This scheme has already been criticized by Hazlehurst. With the Brønsted theory in mind, Hazlehurst suggests using only three types in place of the familiar six: acid-base, oxidation-reduction, combination-decomposition. His third classification seemed necessary because of the limitation by the Brønsted theory of acids and bases to proton donors and proton acceptors. For example, since the reaction

\[\text{SO}_3 + \text{H}_2\text{O} \rightarrow \text{H}_2\text{SO}_4\]

is considered neither an acid-base nor an oxidation-reduction reaction, the third classification is required. According to the Lewis theory this is an acid-base reaction. From the broader viewpoint it is apparent that most of the other reactions included in the combination-decomposition type by Hazlehurst can be assigned to either the first or second. However, there is one class of reactions which cannot be reassigned in this way, namely, reactions involving combination of odd-electron molecules or the decomposition of the product of such combination.

\[\text{Hazlehurst, J. Chem. Education}, 17, 466 (1940).\]
We suggest the following modifications in Hazlehurst's classification of chemical reactions: (1) broadening the first class to include all acids and bases as well as proton donors and proton acceptors; (2) limiting the third class to odd-molecule reactions. The three classes then become:

I. Acid-base reactions.
II. Oxidation-reduction reactions.
III. Odd-molecule reactions.

More than one class may be involved in a given reaction.

One advantage of such a classification is that, while it is based upon experimental data, it also has a simple theoretical basis. Acid-base reactions involve coordinate covalent bonds; oxidation-reduction reactions involve changes in valence; odd-molecule reactions involve simple covalent bonds. The following comparison illustrates the differences between the suggested classification and the older one:

1. *Combination* may involve neutralization, oxidation, or combination of odd-electron molecules, e.g.,

   \[ \text{BCl}_3 + \text{NH}_3 \rightarrow [\text{H}_3\text{N}\rightarrow\text{BCl}_3] \]  (acid-base)

   \[ 2\text{Na} + \text{Cl}_2 \rightarrow 2\text{NaCl} \]  (oxidation-reduction)

   \[ \text{NO}_2 + \text{NO}_2 \rightarrow \text{N}_2\text{O}_4 \]  (odd-molecule combination)

2. *Decomposition* may be merely the reverse of the above reactions.

3. *Displacement* is ordinarily associated with either acid-base or oxidation-reduction phenomena. For example:

   \[ \text{Zn} + \text{CuSO}_4 \rightarrow \text{ZnSO}_4 + \text{Cu} \]  (oxidation-reduction)

   \[ \text{HCl} + \text{H}_2\text{O} \rightarrow \text{H}_3\text{O}^{+1} + \text{Cl}^{-1} \]  (acid-base)

   (Displacement reactions involving acids and bases will be considered in Chapter 8.)

4. *Metathesis* in the usual sense, according to our present understanding of electrolytic reactions, does not occur at all. When the term was first proposed for such reactions, their true nature was not understood. The reaction which then was written

   \[ \text{NaCl} + \text{AgNO}_3 \rightarrow \text{NaNO}_3 + \text{AgCl} \]

   is now written

   \[ \text{Ag}^{+1} + \text{Cl}^{-1} \rightarrow \text{AgCl} \]
and is usually regarded as combination. But even this classification is unnecessary, since the formation of a precipitate depends upon solubility. Even the "reaction"

$$\text{Na}^{+1} + \text{Cl}^{-1} \rightarrow \text{NaCl}$$

will take place if the concentrations of the two ions in solution are high enough, for example, upon evaporation of the solvent. Evidently "metathesis" is merely a matter of solubility, not a class of chemical reactions.
1. Introduction.

The classification of chemical substances as electrophilic and electrodotic according to their behavior in their reactions with other substances can be extended to radicals in a molecule. When this is done, a flood of light is thrown upon the nature of acid-base catalysis. Ingold, Robinson, and others have already done much to clarify many organic reaction mechanisms, but the electronic theory of acids and bases provides a measure of correlation and insight which so far is unobtainable by any other method. Generalized acid-base catalysis will be considered later. In this chapter we shall deal primarily with the effect of acidic and basic radicals in the benzene ring.

Various terms have been proposed to describe what we here call acidic and basic radicals. The principal ones are (1) electrophilic and cationoid for acid radicals and (2) nucleophilic and anionoid for basic radicals. The reasons for preferring electrophilic and electrodotic to describe general behavior have been given in the preceding chapter. In this chapter we are concerned with the specific acidic and basic nature of the radicals, so the terms acidic and basic will be used.

The electronic theory of acids and bases is very helpful in explaining the orienting effects produced by many functional groups attached to benzenoid compounds. The acidic radicals (electro-
philic or electron-withdrawal groups) are those that produce mainly a meta-directing influence on subsequent ring substitutions.

\[
\text{O} \quad \hat{\longrightarrow} \quad \text{O}
\]

Such groups as carbonyl (\(-\text{C} \equiv \)), sulfo (\(-\text{SO}_2\text{OH}\)), cyano (\(-\text{CN}\)), and nitro (\(-\text{NO}_2\)) fall in this category. The basic radicals (electrotopic or electron-release groups) are those that produce mainly an ortho- and para-directing influence. Some of these groups are the amino (\(-\text{NH}_2\)), substituted amino (\(-\text{NHR}\) or \(-\text{NR}_2\)), hydroxyl (\(-\text{OH}\)), and alkoxyl (\(-\text{OR}\)) groups.


The acidic characteristics of these radicals are often ascribed to a displacement of an electron pair (\(\hat{\longrightarrow}\)) which results in an electron deficit on one atom (Chapter 2). This atom thus becomes acidic (electrophilic).

\[
\begin{align*}
\text{O} & \quad \hat{\longrightarrow} \quad \text{O} \\
\text{O} & \quad \hat{\longrightarrow} \quad \text{O} \\
\text{O} & \quad \hat{\longrightarrow} \quad \text{O} \\
\end{align*}
\]

The sulfur atom in the sulfo group is also made electrophilic (to realize its covalence maximum, as discussed in Chapter 2), by the electron displacements (\(\hat{\longrightarrow}\)) shown.

The presence of these acidic radicals in a benzenoid system results in a neutralization within the molecule. For example, the electrophilic nitrogen of the nitro group behaves as an acid and accepts a share in an electron pair belonging to the phenyl group. The phenyl group thus behaves as a base by furnishing an electron pair to the nitrogen atom.

\(^4\) Relative to hydrogen.

In the following formulas the unshared pairs of electrons are represented by writing a dash (for each electron pair) parallel, rather than perpendicular, to the symbol of the atom.

Whether or not a double bond is actually formed by this neutralization is unimportant. The acidic nitro group would, nevertheless, cause the ortho carbon to become electrophilic because of the resulting shift of an electron pair toward the acidic nitrogen. These electronic shifts, and those shown below, would produce low electron densities at the ortho and para carbons and relatively high electron densities at the meta carbons.

The acidic functional groups produce two main effects on the phenyl groups: (1) relatively increased electron densities at the meta positions, and (2) reduced overall electron density of the ring due to electron withdrawal from the ring.

3. Basic Radicals.

The basic radicals behave as bases because they contain unshared pairs of electrons which they can share with any electrophilic group. The phenyl group may behave as an "electron sink"; it can accept a share in a pair of electrons supplied by basic (electrodotic) groups.
The basic groups are thus neutralized by the acidic phenyl group. The result of these electron shifts is an increase in electron density on the ortho carbon atom. These changes, in turn, increase the electron density on the other ortho and para carbon atoms.
The basic functional groups produce two main effects on the phenyl group: (1) increased electron densities at the ortho and para positions, and (2) increased electron density of the ring due to electron release toward the ring.

4. Mechanism of Ring Substitution.

A discussion of the effect of acidic and basic functional groups on the benzenoid ring would not be complete without a brief mention of their relationship to the mechanism of substitution on the ring.

Ingold, Robinson, and Price have laid the foundation of, and also elaborated on, an electronic interpretation of ring substitutions. The orientating influence of acidic and basic radicals can be adequately explained if it is assumed that the substituent is an acid radical.

The experimental evidence indicates that the substitution is probably produced by the action of "positive fragments" on the benzenoid system. These electrophilic (acidic) fragments are either produced by catalysts or if already present their concentration will be increased by the presence of a catalyst (Table 1).

<table>
<thead>
<tr>
<th>REAGENT</th>
<th>CATALYST</th>
<th>POSITIVE FRAGMENT (STRONG ACID)</th>
<th>NEGATIVE FRAGMENT (WEAK BASE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH₃Br</td>
<td>FeBr₃</td>
<td>CH₃⁺⁺</td>
<td>FeBr₄⁻⁻</td>
</tr>
<tr>
<td>:O</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CH₃—C—Cl</td>
<td>AlCl₃</td>
<td>[ :O⁻⁻⁺⁺]</td>
<td>AlCl₄⁻⁻</td>
</tr>
<tr>
<td>:Cl:Cl:</td>
<td>BCl₃</td>
<td>[ :Cl⁻⁻⁺⁺]</td>
<td>BCl₄⁻⁻</td>
</tr>
</tbody>
</table>

These strongly acidic "positive fragments" will attack the benzenoid rings at the points of high electron density. In the previous section we have shown that acidic radicals will increase the electron density at the meta positions, whereas the basic functional

---

10 Bodendorf and Bohme, Ann., 516, 1 (1935).
11 Wertyporosh and Firna, Ann., 500, 287 (1933).
12 Ramser and Wiberg, Ber., 63B, 1136 (1930).
groups will increase the electron density at the ortho and para positions.

The resulting intermediate products of these reactions are thermodynamically unstable compounds that become stabilized by proton expulsion to return to a resonating benzenoid structure:

It is to be kept in mind that the benzenoid formula shown for the final compound represents only one of many possible resonance structures. The formula is not presented as being the one and only correct electronic configuration. To represent the mechanism of a further substitution we would have to show the electronic effect produced on the benzenoid ring by the two groups

\[ \text{Pfeiffer and Wizinger, } \textit{Ann.}, 461, 132 (1928). \]
already present. In this manner we would know the points of high electron density and thus be able to predict the point of attack in the ring.

5. Ease of Ring Substitution.

Holleman $^{12}$ has made a careful study of the role played by a large number of acidic and basic radicals in the nitration of benzenoid compounds.

The presence of an acidic radical, on a benzenoid ring, causes an overall decrease in electron density. The result is a decrease in activity in substitution reactions (since the entering group is electrophilic). Organic chemists have long been familiar with the difficulty of nitrating a compound such as nitrobenzene or benzenesulfonic acid. Concentrated nitric acid is required for the nitra­tions.

Basic radicals, on the other hand, cause an overall increase in electron density on the benzenoid ring. The increase in the number of electrons in the ring would tend to produce increased activity. Thus a substance such as phenol can be nitrated very easily; dilute nitric acid will produce a mixture of ortho and para nitrophenols. Also toluene can be nitrated fourteen times as fast as benzene.$^{13}$

$^{12}$ Holleman, Chem. Revs., 1, 187 (1925).
$^{13}$ Wibaut, Rec. trav. chim., 34, 241 (1915).
Chapter 6

NEUTRALIZATION

This chapter and those following it are concerned with the four experimental criteria of acids and bases adopted by Lewis: neutralization, titration with indicators, displacement, and catalysis.

Neutralization is the formation of a coordinate covalent bond between a primary acid (see Chapter 3) and a primary base, e.g.,

\[ \text{H}^+ + \text{O} : \text{H} \rightarrow \text{H} : \text{O} : \text{H} \quad (1) \]

\[ \text{Ag}^{+1} + 2 \text{NH}_3 \rightarrow \text{Ag} : (\text{NH}_3)^{+1}_{\text{base}} \quad (2) \]

It is customary to consider that the "hydrogen ion" occurs as a solvated proton, for example, the hydronium ion in water or the ammonium ion in liquid ammonia:

\[ \begin{array}{c}
\text{H} \\
\text{H} : \text{O} : \text{H}^{+1} \\
\ddash \text{H} \\
\text{H} : \text{N} : \text{H}^{+1} \\
\text{H} \\
\end{array} \]

These cations are both secondary acids, and their reactions would involve displacement, not neutralization (Chapter 8). It is customary to disregard solvation of metallic ions such as the acidic silver ion in equation 2. Yet there is no doubt that the silver ion and other acidic ions in water solution are solvated as is the proton. Apparently we may speak either of neutralization or of displacement, depending upon our point of view. This question will be discussed further in Chapter 8.

However, when there seems to be no doubt that a primary acid (e.g., boron trichloride in the absence of a solvent) is involved, there is no difficulty in using the term neutralization. At any rate, the electronic theory of acids and bases gives definite meaning to the word. The type equation of the Brønsted theory

\[ \text{acid}_1 + \text{base}_2 \rightarrow \text{acid}_2 + \text{base}_1 \quad (3) \]

is often assumed to dispose of the concept of neutralization. We shall see in Chapter 8 that this is not the case. The equation may
neutralization

represent the fact that acids or bases will replace weaker acids or bases from their compounds. It does not abolish neutralization.

The acid boron trichloride is neutralized by the base triethylamine when both substances are in the pure liquid or gaseous state:

\[
\text{Cl}: \text{Et} + \text{Cl}: \text{Et} \rightarrow \text{Cl}: \text{B} + \text{N}: \text{Et} 
\]

\[
\begin{align*}
\text{Cl}: \text{Et} + \text{Cl}: \text{Et} & \rightarrow \text{Cl}: \text{B} + \text{N}: \text{Et} \\
\text{Cl}: \text{Et} + \text{Cl}: \text{Et} & \rightarrow \text{Cl}: \text{B} + \text{N}: \text{Et}
\end{align*}
\]

The product is usually called a molecular or addition compound. In some cases it may be a salt. In this particular example the possibility of ionization of one of the chlorine atoms ought to be considered. When the boron chloride gains a share in the lone pair of electrons on the nitrogen atom, the boron atom is made more negative, thus repelling the chlorine electrons. It may not occur to a great extent in this particular compound, but where sufficient electrical "strain" is set up, upon the acceptance of a share in another electron pair by the acid, one would expect ionization to be favored. Such ionization in solution would, of course, be greatly affected by the dielectric constant of the solvent. Three examples of ionization, of the many that could be given, are the following:

\[
\begin{align*}
\text{C}_6\text{H}_5\text{N} + \text{HCl} & \rightarrow \text{C}_6\text{H}_5\text{NH}^+ + \text{Cl}^- \\
\text{AlCl}_3 + \text{COCl}_2 & \rightarrow \text{COCl}^+ + \text{AlCl}_4^- \\
\text{SnCl}_4 + 2\text{SeOCl}_2 & \rightarrow 2\text{SeOCl}^+ + \text{SnCl}_6^{2-}
\end{align*}
\]

In these and similar examples the product is usually considered a salt. There seems to be no need for the name "pseudo salt." It would appear that when such compounds are crystallized from the solvent the ions may persist in the solid state. (We use the word salt to mean an ionic compound.) Hence, we would call the compounds salts.

However, the formation of a salt is not always the result. According to the water theory of acids and bases, a salt invariably resulted from neutralization; but salt formation is obviously unnecessary, as the following examples show:

\[
\begin{align*}
\text{SO}_3 + \text{HOH} & \rightarrow \text{H}_2\text{SO}_4 \\
\text{BCl}_3 + (\text{CH}_3)_2\text{O} & \rightarrow [(\text{CH}_3)_2\text{O} \rightarrow \text{BCl}_3]
\end{align*}
\]
The essential thing in neutralization is the formation of the coordinate covalent bond.

The neutralization product is not necessarily neutral in the sense that it is no longer either acidic or basic. Unfortunate as this may seem, it is a familiar fact. For example, in equation 8 the neutralization product, sulfuric acid, is still strongly acidic. In equation 9 the product is definitely amphoteric in subsequent displacement reactions with either acids or bases.

Neutralization does not necessarily involve the solvent. Some discussion of the role of the solvent will be given in Chapter 8. Here we shall consider a few reactions which do not involve the solvent. Some have already been mentioned, e.g., the gaseous phase reaction between boron fluoride and amines or ammonia. Others occurring at high temperatures have been listed by Audrieth and Moeller.¹ The formation of certain electrolytic melts, borax and metaphosphate bead tests, the manufacture of glass and cement, and the formation of slag in the blast furnace may all be classified as neutralization reactions.²

Some of them might also be classified as acid-base displacement, depending upon how the choice of base is made, as the following examples indicate:

\[
\begin{align*}
\text{FeS} + \text{S}^2^- & \rightarrow \text{FeS}_2^{2-} & (10) \\
\text{AlF}_3 + 3\text{F}^- & \rightarrow \text{AlF}_6^{3-} & (11) \\
\text{SiO}_2 + \text{O}^{2-} & \rightarrow \text{SiO}_3^{2-} & \text{acid} & \text{base} & (12)
\end{align*}
\]

Equation 12 might represent displacement rather than neutralization if the oxide ion comes from an oxide which is not ionic. (Note that in equations 11 and 12 the central acidic atom of the product in each reaction has twelve valence electrons—two more violations of the old "rule of eight" discussed in Chapter 2.)

The effect of the solvent upon neutralization will be treated in Chapter 8 since a previous discussion of displacement is necessary. The type equation of the Brønsted theory eliminates neutralization from all acid-base reactions covered by the theory. This is because the equation represents displacement reactions of secondary acids and bases. Since the Lewis theory includes the Brønsted

² An interesting application to glass manufacture is given in a paper by Sun and Silverman, J. Am. Ceramic Soc., 28, 8 (1945).
theory we reach the same conclusion as far as hydrogen acids are concerned. However, the Lewis theory does give meaning to the term neutralization as applied to reactions between primary acids and bases. Neutralization is the combination of a primary acid with a primary base to form a coordinate covalent bond between the two.
Chapter 7

TITRATIONS WITH INDICATORS

1. Introduction

The second of the experimental criteria of acids and bases is their behavior in titrations against each other with the aid of indicators. Such titrations may be performed in a great variety of solvents.

Lewis\(^1\) titrated bases like pyridine and triethylamine with acids such as solutions of BCl\(_3\) and SnCl\(_4\) in carbon tetrachloride, and AgClO\(_4\) dissolved in benzene. These solutions were titrated back and forth with the use of indicators such as thymol blue, butter yellow, and crystal violet. Crystal violet is an especially convenient indicator because of its solubility in a variety of solvents and because it usually gives the same color change in different solvents.

Comparison by means of a series of titrations is one method of establishing whether certain substances are acids or bases. One example of the titration procedures is given in the next section. Titrations of the above solutions make it obvious that there are many acids other than H-acids and many bases other than HO-bases. This fact is not surprising when the fundamental electrophilic and electrodotic characteristics of such substances are considered.

\(^1\) Lewis, J. Franklin Inst., 226, 293 (1938).
2. Experimental Titrations.

The following experiments demonstrate only a few of the many titrations that can be carried out to show the acidic and basic characteristics of compounds.

**WATER USED AS SOLVENT.** The indicator used in this series of experiments is crystal violet dissolved in water.

*Acids* (HCl, H$_2$SO$_4$, HC$_2$H$_3$O$_2$, HNO$_3$). Ten milliliters of water is poured into each of four 6-inch test tubes, and 3 or 4 drops of the indicator is added. The violet color that results can be changed to yellow by the addition of acid. When dilute acid is added slowly, the intermediate green color may easily be observed. The HCl, H$_2$SO$_4$, HC$_2$H$_3$O$_2$, and HNO$_3$ are used, one in each tube, to show this color change.

The reaction is explained by the fact that these acid solutions contain the hydronium ion, H$_3$O$^{+}$1, which is electrophilic.

*Bases* (NH$_3$, C$_5$H$_5$N, C$_2$H$_5$OH, 1,4-dioxane). The basic characteristics of these compounds can be demonstrated by the addition of a few drops of each to the four yellow solutions obtained in the above experiment. In each solution the color is changed back from yellow to violet on the addition of the base. These compounds are bases because they are electron-pair donors and can form coordinate covalent bonds with the protons of the above-mentioned acids.

Example:

\[
\begin{align*}
\text{dioxane} & \\
\begin{array}{c}
\text{CH}_2-\text{CH}_2 \\
\text{O:} \\
\text{CH}_2-\text{CH}_2 \\
\text{base}
\end{array} & \\
\text{O:} + \text{H}_3\text{O}^{+1} \rightarrow \begin{array}{c}
\text{CH}_2-\text{CH}_2 \\
\text{O:} \\
\text{CH}_2-\text{CH}_2 \\
\text{acid}
\end{array}^{+1} \\
& + \text{H}_2\text{O}
\end{align*}
\]

**CHLOROBENZENE USED AS SOLVENT.** The indicator used in these experiments is crystal violet dissolved in chlorobenzene.

*Acids* (HCl (gas) dissolved in C$_6$H$_5$Cl, BCl$_3$ dissolved in C$_6$H$_5$Cl, and fuming SnCl$_4$). Ten milliliters of chlorobenzene is poured into each of three 6-inch test tubes, and 3 or 4 drops of the indicator is added to each tube. The violet color that results can be changed to yellow by the addition of a few drops of an acid. The above acids are used to show this color change, one in each test.

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Experiment 93

Here again we have substances that are acidic because they are electron-pair acceptors.

*Bases* (ethers, carboxylic acid anhydrides, alcohols).

![Chemical structures]

All these substances can be demonstrated to be bases by allowing them to react with the acids mentioned above. If these bases are added to the yellow solutions formed in the first part of this experiment, the violet color of the indicator quickly reappears.

Example:

**Phthalic acid anhydride**

\[
\begin{align*}
\text{base} & : \text{Cl} : \\
\text{O} & : \text{Cl} : \\
\text{C} & : \text{Cl} : \\
\text{C} & : \text{Cl} : \\
\text{O} & : \text{Cl} : \\
\text{C} & : \text{Cl} : \\
\end{align*}
\]

**Carbon Tetrachloride Used as Solvent.** The indicator used is crystal violet dissolved in chlorobenzene rather than in carbon tetrachloride. (The indicator is quite insoluble in carbon tetrachloride, but when it is added in chlorobenzene solution no precipitation takes place; a clear solution results.)

**Acids** (BCl₃, HCl (gas) dissolved in CCl₄, and AlCl₃ in CCl₄). Ten milliliters of anhydrous carbon tetrachloride is poured into each of three 6-inch test tubes and 3 or 4 drops of the indicator is added to each tube. The violet color that results can be changed to yellow by the addition of some of these acids. The AlCl₃ is not very soluble in CCl₄. Shaking the mixture for a considerable length of time and then filtering gives a clear filtrate which does not behave as an acid. If any hydrolysis should take place, due to the presence of traces of moisture, the resulting HCl gas would dissolve in the solvent and an acid reaction would be observed.
If some of the original mixture of AlCl₃ suspended in CCl₄ is used, however, the violet color of the indicator disappears and the resulting yellow color shows that the solid AlCl₃ is behaving as an acid. This experiment helps to prove that the AlCl₃ itself is acting as an acid, not any HCl resulting from hydrolysis.

Enough AlCl₃ can be dissolved in the CCl₄ by warming so that a clear solution of it may be used as an acid solution.

As the BCl₃ and the HCl gas are readily soluble in the solvent, no trouble is encountered in the preparation and use of these acidic solutions.

Bases (pyridine, amines, esters).

\[
\begin{align*}
\text{pyridine} & : \text{CH} = \text{CH} & \text{tripropylamine} & : \text{CH}_3\text{CH}_2-\text{N} : \\
\text{base} & : \text{CH} = \text{CH} & \text{acid} & : \text{CH} = \text{CH}
\end{align*}
\]

These substances can be demonstrated to be bases by the fact that when they are added to the yellow solutions resulting in the first part, the violet color of the indicator reappears.

Example:

\[
\begin{align*}
\text{pyridine} & : \text{CH} = \text{CH} & \text{AlCl} : \text{Cl} : & \rightarrow & \text{CH} = \text{CH} & \text{AlCl} : \text{Cl} : \\
\text{base} & : \text{CH} = \text{CH} & \text{acid} & : \text{CH} = \text{CH}
\end{align*}
\]

Discussion. In all these experiments the acidic substances are those that contain atoms capable of accepting a share in an electron pair and the bases are those substances containing atoms capable of donating a share in an electron pair.

It is obvious that hydrogen chloride behaves as an acid regardless of whether it is dissolved in water, chlorobenzene, carbon tetrachloride, or any other solvent. (In water solutions of the common acids it is the H⁺ or H₃O⁺ that behaves as an acid, whereas in the organic solvents it is the molecules of the compounds that behave as acids.) The same may be said for all other compounds that are capable of accepting electron pairs.
It is apparent that compounds like pyridine, alcohols, ethers, and many other substances act as bases whether dissolved in water, chloroform, carbon tetrachloride, chlorobenzene, or any other solvent that will dissolve them, because fundamentally these bases are compounds that are capable of donating an electron pair.

3. Indicator Color Changes.

The observation that different acids cause the same color change is readily explained as follows.

The addition of increasing amounts of an acid to a solution of crystal violet will cause the violet color to change to green and then to yellow. Rosenstein and Adams explained these changes in water solution as due to the successive addition of $H^+$ to the dimethylamino groups of the indicator.

The above formulas represent two of the several possible resonance structures that can be written to show the violet form of the indicator. It is obvious that the unshared electron pair on the nitrogen atom of each $N(CH_3)_2$ group can form a quinoid structure as shown in ring c of the left-hand formula. Other resonance structures would have ring a or b as the quinoid ring. If $H^+$ or

---

BF$_3$ or any other electrophilic substance is added to the indicator solution, coordination with one of the unshared pairs of electrons will result.

![Chemical structure](image)

This addition removes one of the N(CH$_3$)$_2$ groups (ring c) from the resonating system and produces a green-colored substance similar to malachite green.

![Chemical structure](image)

Further color change to yellow is produced by the addition of a second H$^+$ to another N(CH$_3$)$_2$ group, leaving only one N(CH$_3$)$_2$ group in the resonating system.$^5$

It is obvious that the presence of any strongly electrophilic substance such as BCl₃, AlCl₃, AgClO₄, SO₃, and SnCl₄ could result in a coordination in a similar manner to the behavior of the H-acids. The color changes are the same regardless of the acid or solvent used. Lewis and Bigeleisen⁶ give a quantitative demonstration that H⁺⁺ or SnCl₄ added to one —NR₂ of methylene blue results in almost identical spectra.

The addition of basic (electrodotic) substances would, of course, neutralize any of the above acids in the solution and also remove from the dye molecule the coordinated electrophiles. This reaction would again return the N(CH₃)₂ groups to the resonating system of the dye molecule, and the green and then the violet colors would again reappear.

Thus the electronic theory provides a simple and consistent explanation of the experimental behavior of a great variety of acids and bases toward indicators.

Chapter 8

DISPLACEMENT

1. Introduction.

The third experimental criterion of acids and bases is displacement. There are two types of acid-base displacement: (1) a strong acid in general will displace a weaker one from combination with a base; (2) a strong base will displace a weaker one from combination with an acid. A weaker coordinate bond is broken to form a stronger one.

A familiar example of the first type of displacement (of a weaker acid by a stronger one) is the liberation of carbon dioxide from sodium carbonate solution upon the addition of hydrogen chloride:

$$2H^+ + CO_3^{2-} \rightarrow H_2O + CO_2$$

The hydrogen ions displace the weak acid, carbon dioxide, from combination with the oxide ion. Another example described by G. N. Lewis is the displacement of carbon dioxide when finely divided sodium carbonate is warmed with boron trichloride or stannic chloride in a mixture of carbon tetrachloride and acetone. The other product was not analyzed, but the reaction might be written tentatively as:

$$BCl_3 + Na_2CO_3 \rightarrow [Na_2O→BCl_3] + CO_2$$

Another example of the displacement of one acid by another is the high-temperature reaction (favored by the volatility of the CO_2):

$$SiO_2 + Na_2CO_3 \rightarrow Na_2SiO_3 + CO_2$$

Examples of the second kind of displacement (of a weak base by a stronger one) are more familiar, since the type equation of the Brønsted theory represents the displacement of one base by a stronger base in competition for the proton, for example:

$$\text{H}_2\text{O} + \text{HCl} \rightarrow \text{H}_3\text{O}^{+} + \text{Cl}^{-}$$

Water acting as a base displaces the weaker base, chloride ion, from combination with the proton. Another example that does not involve the proton is the displacement of ammonia from combination with trimethylboron by trimethylamine:

$$\text{(CH}_3\text{)}_3\text{N}^- + \text{H}_3\text{N}:\text{B(CH}_3\text{)}_2 \rightarrow \text{(CH}_3\text{)}_3\text{N}:\text{B(CH}_3\text{)}_2 + :\text{NH}_3^-$$


Some writers have indicated that hydrogen acids require special consideration in the Lewis terminology. No doubt such a misunderstanding arises from the emphasis of the Brønsted theory upon displacement of one base by another as the only criterion of acid-base phenomena. Actually displacement is only one of the four “phenomenological criteria” of acids and bases, and the Brønsted type equation is only one of the two types of acid-base displacement. The Brønsted theory is thus included in the Lewis theory and requires no special consideration.

In order to show that special treatment is unnecessary for hydrogen acids, we need only demonstrate that the displacement reaction represented by the typical Brønsted equation is not unique. The same kind of reaction may take place between many other acids and bases. A detailed consideration of one example should be sufficient illustration. When the addition compound formed by the neutralization of boron trichloride by acetone is added to pyridine, displacement of the weaker base (acetone) by the stronger base (pyridine) takes place. This reaction is exactly analogous to the displacement of the weakly basic chloride ion from combination with the proton when hydrogen chloride is added to water:

Both boron trichloride and the proton are primary acids which, to form the acid1 compounds shown above, have been neutralized by the weak bases acetone and chloride ion, respectively. However, in both the resulting compounds, stronger bases are able to displace the weaker ones from combination with the two acids. All four acids (in both equations) are secondary acids.

Many more examples of displacement reactions of exactly the same type could be given. Some of them are responsible for catalysis and will be considered in the following chapters. One familiar example is interesting since it has been the subject of some misunderstanding:

\[
\text{Ag}^{+} + \text{NH}_3 \rightleftharpoons \text{Ag}
\]

The silver ion is a fairly strong acid which will combine with many bases, including ammonia and hydroxyl ion. The reaction represented by the above equation is analogous to any Brønsted type equation, for example:

\[
\text{NH}_4^{+} + \text{OH}^{-} \rightleftharpoons \text{H}_2\text{O} + \text{NH}_3
\]
Silver ions and ammonia molecules combine until equilibrium is reached when their concentrations are low. Silver ions and hydroxyl ions also combine until equilibrium is reached when the concentrations of both ions are low. If the two ions have been added in equal amounts, equilibrium is attained when each has a concentration equal to $1.4 \times 10^{-4} M$. Now if in the other solution the concentrations of ammonia and silver ion in equilibrium with the complex ion are adjusted to have comparable molarities, that is, $1.4 \times 10^{-4} M$, the concentration of the complex is $0.47 \times 10^{-4} M$. In such a solution only about one-fourth of the silver ions have combined to form the complex. Thus under comparable conditions a large proportion of the silver ions do not combine with ammonia molecules.

Perhaps the cause of the misunderstanding lies in the method by which we are accustomed to observe the preparation of the silver-ammonia complex. When we add excess ammonia solution to silver nitrate solution the initial precipitate dissolves, forming the complex ion. This apparently means that ammonia is a much stronger base toward silver ion than hydroxyl ion is. But we must remember that the silver hydroxide precipitate is formed first. When the first few drops of ammonia solution are added the ammonia is so dilute that it is largely ionized to form hydroxyl ions which coordinate with the silver ions to form the hydroxide. But, by the time the precipitate is completely dissolved, the amount of ammonia added is enough so that if the silver ion were not present the concentration of $\text{NH}_3$ would be roughly 100 times the concentration of $\text{OH}^{-1}$ ion. Actually, if concentrated sodium hydroxide solution is now added to the solution, the ammonia is displaced and the precipitate returns. This is what one would expect when mass-action effects are taken into account. The effects observed are similar whether the acid is the proton or the silver ion. The displacement of one base by another depends not only upon their relative strengths, but upon concentration factors also.

Since silver hydroxide actually appears as silver oxide when attempts are made to isolate it, we cannot be positive that a coordinate bond is formed between silver and hydroxyl ions. However, there are two reasons for believing that one is formed: (1) other hydroxides of similar nature (small positive ion) are often not ionic even in the solid state; (2) silver iodide does not crystallize into an ionic lattice although the chloride and bromide do.
An idea of the relative basic strengths of ammonia and hydroxyl ions toward silver ion can be obtained from equation 7. The equilibrium constant for equation 7 is

$$K = \frac{[NH_3]^2}{[Ag(NH_3)_{2}^{+}] \times [OH^{-}]}$$  \hspace{1cm} (9)

which may be evaluated by combining

$$\frac{[Ag^{+}] \times [NH_3]^2}{[Ag(NH_3)_{2}^{+}]} = 6 \times 10^{-8}$$  \hspace{1cm} (10)

with

$$[Ag^{+}] \times [OH^{-}] = 2 \times 10^{-8}$$  \hspace{1cm} (11)

We see that the constant has a value of 3, indicating that ammonia is about as strong a base as hydroxyl ion toward silver ion.

It is worth while noting at this point that in equations 5, 7, and 8 the acid$_1$ and acid$_2$ compounds are amphoteric in exactly the same sense. The addition compounds of boron trichloride with acetone and with pyridine are amphoteric because they contain both acidic and basic constituents. They can react both ways because stronger bases will displace the weaker bases and stronger acids will displace the boron trichloride. In the same way silver hydroxide in equation 7 and water in equation 8 are also amphoteric. They appear as acids in these equations, but both may act as bases because of the presence of the hydroxyl group.


When the solvent takes part in acid-base phenomena, the reactions are usually displacement reactions. Considering only acid-base reactions, solvents may be divided into three classes: (1) those that are ordinarily inert toward acids and bases, e.g., benzene, carbon tetrachloride, and chlorobenzene; (2) those that are ionizable, e.g., water, ammonia, sulfur dioxide, phosgene, and selenium oxychloride; (3) those that do not ionize but do react with acids and bases, e.g., ether and pyridine. If we consider the neutralization of boron trichloride by triethylamine in the three types of solvents, we find that the net result may be the same as when the neutralization occurs in the absence of a solvent:

$$(C_2H_5)_3N: + BCl_3 \rightarrow (C_2H_5)_3N:BCl_3 \text{ (neutralization)}$$  \hspace{1cm} (12)

When the solvent is inert it is merely a diluent and the neutralization product is obtained directly. When the solvent is ionizable,
intermediate displacement reactions with the solvent may be ob­
served. If either the acid or the base, or both, are strong enough,
they will at least partially displace cations or anions of the solvent;
e.g., if selenium oxychloride is chosen as the solvent:

\[
\begin{align*}
\text{BCl}_3 + \text{SeOCl}_2 & \rightarrow \text{SeOCl}^{+1} + \text{BCl}_4^{-1} \quad \text{(displacement)} \quad (13) \\
(\text{C}_2\text{H}_5)_3\text{N}: + \text{SeOCl}_2 & \rightarrow \quad (\text{C}_2\text{H}_5)_3\text{N}:\text{SeOCl}^{+1} + \text{Cl}^{-1} \quad \text{(displacement)} \quad (14)
\end{align*}
\]

When the two solutions are mixed, the following reaction takes
place:

\[
\begin{align*}
\text{SeOCl}^{+1}, \text{BCl}_4^{-1} + (\text{C}_2\text{H}_5)_3\text{N}:\text{SeOCl}^{+1}, \text{Cl}^{-1} & \rightarrow \\
(\text{C}_2\text{H}_5)_3\text{N}:\text{BCl}_3 + 2\text{SeOCl}_2 
\end{align*}
\]

The same neutralization product is obtained as when the reaction
is carried out directly or in an inert solvent.

When the solvent reacts without ionizing, it reacts with either
the acid or the base, but not ordinarily with both. Such solvents
are usually not amphoteric. For example, if boron trichloride
reacts with triethylamine in ether, the boron trichloride would
react with the ether, but the triethylamine would not. The oxy­
gen atom in the ether can donate an electron pair to form a coordi­
nate bond, but the hydrogen atoms in ether have little tendency
to form hydrogen bridges:

\[
(\text{C}_2\text{H}_5)_2\text{O} + \text{BCl}_3 \rightarrow [(\text{C}_2\text{H}_5)_2\text{O} \rightarrow \text{BCl}_3] \quad (16)
\]

When triethylamine is added, it merely displaces the weaker base
and the resulting product is the same as before:

\[
[(\text{C}_2\text{H}_5)_2\text{O} \rightarrow \text{BCl}_3] + (\text{C}_2\text{H}_5)_3\text{N} \rightarrow \\
[(\text{C}_2\text{H}_5)_3\text{N} \rightarrow \text{BCl}_3] + (\text{C}_2\text{H}_5)_2\text{O} \quad (17)
\]

In all four cases the final neutralization product is the same.

These examples help to clarify the relationship of neutralization
to displacement mentioned at the beginning of Chapter 6. Neu­
tralization is the formation of the coordinate covalent bond be­
tween a primary acid and a primary base. The reaction repre­
sented by equation 12 is neutralization, but we note that when a
solvent is involved the same result may be obtained through a
series of displacements. Equations 13 and 14 are both displace­
ment reactions. Equation 15 represents two simultaneous dis­
placements. At the same time equation 15 also represents neutralization as far as the original primary acid and primary base are concerned. Whether or not a solvent is used as a medium for this reaction, the final product is the result of neutralization—the formation of the coordinate covalent bond between the boron chloride and triethylamine.

4. The Leveling Effect.

In comparing the relative strengths of acids and bases in a particular solvent such as water, liquid ammonia, or glacial acetic acid, the range of comparison is limited by the strength of the solvent as an acid or base.

For example, the difference between perchloric, hydrobromic, sulfuric, hydrochloric, and nitric acids is obscured in water because they all react completely with the solvent in dilute solution. The reaction

$$\text{HCl} + \text{H}_2\text{O} \rightarrow \text{H}_3\text{O}^{+1} + \text{Cl}^{-1}$$

proceeds completely to the right because the water molecule is so much stronger as a base than chloride ion that it displaces it completely from combination with the proton. The perchlorate, bromide, bisulfate, and nitrate ions are all similar to the chloride ion in acting as extremely weak bases.

Since such acids react completely with water, only one acid, the hydronium ion, is left in solution. Consequently the acids all appear equally strong. This is known as the leveling effect.

A solvent which is more basic than water, e.g., liquid ammonia, will level the strength of a larger group of acids. For example, in water, acetic acid is much less ionized than hydrochloric acid because the acetate ion formed by the reaction

$$\text{HC}_2\text{H}_3\text{O}_2 + \text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^{+1} + \text{C}_2\text{H}_3\text{O}_2^{-1}$$

is a much stronger base than the chloride ion. Water is not able to displace it to a very great extent from its combination with the proton. But when acetic acid is added to liquid ammonia as a solvent the reaction

$$\text{HC}_2\text{H}_3\text{O}_2 + \text{NH}_3 \rightarrow \text{NH}_4^{+1} + \text{C}_2\text{H}_3\text{O}_2^{-1}$$

goes all the way to completion.\(^5\) Ammonia is a stronger base than water and is so much stronger than the acetate ion that it

is able to displace it completely. Thus in liquid ammonia all acids stronger than acetic acid will appear to have the same strength, that of the ammonium ion. Since the ammonium ion is weaker than the hydronium ion, we say that the strength of acids is leveled down farther in ammonia than in water.

Obviously a solvent more acidic than water would have a similar leveling effect on bases such as ammonia and the amines.

The leveling effect is one phenomenon that must be taken into account when comparing relative acid or base strengths. Since water levels downward the strengths of the strong acids like hydrogen chloride, it is necessary to use a less basic solvent such as acetic acid or even an “inactive” one like benzene. In acetic acid the reaction

\[ \text{HCl} + \text{HC}_2\text{H}_3\text{O}_2 \rightleftharpoons \text{H}_2\text{C}_2\text{H}_3\text{O}_2^{+1} + \text{Cl}^{-1} \]

has little tendency to proceed to the right because acetic acid is so weakly basic. Hence a comparison with other acids is possible. It has been found that \( \text{HClO}_4 > \text{HBr} > \text{H}_2\text{SO}_4 > \text{HCl} > \text{HNO}_3 \) gives the relative strengths of these strong acids in acetic acid as a solvent.⁶

5. The Strengths of Acids and Bases.

In the preceding section we have tacitly assumed that the familiar method of comparing the strengths of acids and bases by displacement reactions yields values which are independent of the reference base or acid. This is not the case, as several investigators including Lewis have shown. However, the exceptions are usually obvious and as yet probably do not warrant discarding the idea of monotonic series for the confusion that would thereupon result. To quote Lewis: “The fact that we can thus get a satisfactory acidity function over the whole range from pure water to pure sulfur trioxide shows the desirability of attempting to construct a monotonic series of acids even though we have pointed out that it can never be exact and that there may be some very large specific variations.”⁷

One of the exceptions referred to by Lewis occurs in the reversal of the relative strengths of triethylamine and ammonia. Ammonia is the weaker base toward the proton, but, when the reference acid is \( m\)-dinitrobenzene, ammonia appears to be much stronger than

---

triethylamine. Double chelation is the explanation advanced by Lewis and Seaborg.\(^8\)

```
O
O
O
N
O

\[ \text{Formula I represents one of the resonating structures of } m \text{-dinitrobenzene. Formula II represents the compound formed upon the addition of ammonia. Obviously such double chelation is not to be expected from triethylamine.} \]

Another interesting example of reversal from the expected order of basic strength is given by H. C. Brown.\(^2\) Using displacement reactions of the type

\[
(CH_3)_3N + H_3N:B(CH_3)_3 \rightleftharpoons (CH_3)_3N:B(CH_3)_3 + :NH_3
\]

he found, for example, that when trimethylboron is used as the reference acid, trimethylamine is a stronger base than ammonia, as it is toward the proton. (Note that the above reaction is a displacement reaction entirely analogous to the type equation of the Brønsted theory.) But when the reference acid is changed to tri-\(t\)-butylboron, the order is reversed. The explanation advanced is given in terms of "\(F\)-strain." "\(F\)-strain is that strain which is caused by steric interference of the atoms or groups which are attached to different atoms resulting in a force which tends to separate the two atoms."\(^2\)

Chapter 9

CATALYSIS

1. Nature of Catalysis .......................................................... 107
2. Acid Catalysis ................................................................. 108
3. Base Catalysis ............................................................... 111

1. Nature of Catalysis.

The fourth criterion of acid and base behavior suggested by G. N. Lewis is catalysis. The following chapters represent an attempt to discuss this criterion systematically. Familiar examples of acid-base catalysis are considered from the Lewis viewpoint in order to show that there is no essential difference in catalytic behavior whether protons are involved or not.

"Catalysis" as usually defined has implied that the catalyst is recovered unchanged. There seems to be a tendency to broaden the definition. For example, Hammett ¹ points out that there is no important distinction between catalysis in the older sense and the kind of acceleration produced by a base which appears as its "conjugate acid" at the end of the reaction. A few of the examples presented here are of the conjugate type, but most of them conform to the older definition.

Since it is already generally recognized that basic behavior is displayed by a very large number of substances,²⁻⁸ only enough examples of base-catalyzed reactions are included here to show that when a base acts as a catalyst the substance attacked by the base is behaving as an acid.

³ Brønsted, Rec. trav. chim., 42, 718 (1923).
⁴ Lowry, Chemistry & Industry, 42, 43 (1923).
However, in spite of the fact that numerous examples of typically acidic properties displayed by substances which do not contain protons have already been listed by many writers,\textsuperscript{9, 10} the idea that acidic behavior does not depend upon protons and, in fact, is as widespread as basic behavior is not yet widely accepted. So in our discussion of catalysis, the principal emphasis is upon systematic application of the new theory to acidic catalysis rather than to basic catalysis.

2. Acid Catalysis.

Evidence has been cited previously to show that the common Friedel-Crafts type catalysts such as aluminum chloride, ferric chloride, stannic chloride, and zinc chloride are acids. Their catalytic activity is well known in connection with organic reactions, but it is not usually recognized that they also catalyze such common reactions as those of metals with water. The reaction of iron with pure water to give hydrogen is a very slow one because the concentration of hydrogen ions is so low. Any acid sufficiently strong to increase the hydrogen-ion concentration substantially will accelerate the reaction. It is immaterial whether the acid is \textit{molecular} hydrogen chloride, acetic acid, sulfur trioxide, carbon dioxide, stannic chloride, or \textit{ionic} zinc chloride or cupric sulfate. The rate of reaction depends upon increasing the hydrogen-ion concentration, which in turn depends upon the concentration and strength of the acid which is displacing protons from the water. We shall see that this \textit{increase in positive-ion concentration is typical of acid catalysis}. Obviously these ideas may have a far-reaching effect on the study of corrosion. Corrosion of metals is a complex subject, but it would seem that application of the electronic theory of acids and bases might be helpful in its investigation.

Since the Friedel-Crafts catalysts are usually thought of as halides of a few metals, a natural question to raise might be: If these halides are really acids differing in no fundamental manner from H-acids in their behavior, why do not H-acids catalyze reactions of the Friedel-Crafts type also? The answer is that they do. Hydrogen fluoride,\textsuperscript{11} phosphoric acid, and sulfuric acid\textsuperscript{12, 13} have

\textsuperscript{9} Lewis, \textit{J. Franklin Inst.}, 226, 293 (1938).
\textsuperscript{10} Luder, \textit{Chem. Revs.} 27, 547 (1940). See Chapter 10 for additional references.
\textsuperscript{11} Simons and Archer, \textit{J. Am. Chem. Soc.}, 60, 2953 (1938).
\textsuperscript{12} Burwell and Archer, \textit{J. Am. Chem. Soc.}, 64, 1032 (1942).
been established as catalysts for Friedel-Crafts and similar reactions.

The catalytic activity of sulfuric acid seems especially interesting. Depending on the concentration of water or of sulfur trioxide, either sulfur trioxide or the H-acid or both may be responsible for the catalysis. Lewis and Bigeleisen 14 have shown 15 that the very rapid increase in the acidity function of Hammett and Deyrup as 100 per cent sulfuric acid is approached is probably due to sulfur trioxide. This might indicate that in the sulfonation of benzene the sulfur trioxide is the principal agent in a direct electrophilic attack.

![Chemical diagram](image)

Bordwell, Suter, and Webber 16 have found this concept 17 useful in explaining the mechanism for the reaction of dioxane sulfotrioxide with olefins.

The Friedel-Crafts reactions are acid-catalyzed and can be taken as typical examples of this type of catalysis. For example, the alkylation of benzene by an alcohol may be represented as follows:

```
H   H
R-C:O:H + B:F ⇌ R-C:O:B:F
base    acid
```

```
H   H
R-C:O:B:F ⇌ [H] + [H:F]
acid    base
```

The similarity of the absorption curves for methylene blue in stannic chloride and in sulfuric acid, given by the same authors, is a striking demonstration of the essential likeness in the behavior of these two acids toward indicators.

The first step involves the typically acidic behavior of boron trifluoride in accepting a share in a pair of electrons from the alcohol-oxygen $^{18,19}$ to form a coordinate bond. The alcohol is then behaving as a base just as it does according to the Brønsted theory when hydrogen chloride is added to it. The ionization that occurs produces a "positive fragment" $^{20}$ which is itself acidic, since it has a great tendency to accept a share in an electron pair. The formation of this acid is exactly analogous to any acid-base displacement in which one acid displaces another from combination with a base. The catalytic effect of boron trifluoride in producing this positive fragment is also analogous to the catalytic effect of similar acids in increasing the "positive fragment" concentration in water, i.e., the hydrogen-ion concentration. As pointed out above, the acceleration of the reaction between iron and water by hydrogen chloride, sulfur trioxide, aluminum chloride, zinc chloride, etc., depends upon the increase in the concentration of hydrogen ions or "positive fragments" in the water. This formation of "positive fragments" seems to be typical of acid catalysis. The acid catalyst increases the speed of the reaction by increasing, by displacement, the concentration of the acid group involved in the reaction.

The use of hydrogen fluoride as an acid catalyst in the alkylation

of benzene and phenol with alcohols has also been reported.21, 22, 23 This reaction again demonstrates the similarity between boron trifluoride and H-acids. It is interesting to note that hydrogen fluoride does not effect methylation, although aluminum chloride does.23 This reaction seems to indicate that aluminum chloride is a stronger acid than hydrogen fluoride.

The use of an acid catalyst does not always result in a displacement reaction and the formation of a positive fragment. For example, olefins will react with an acid catalyst 24, 26 in such a manner that a "positive center" is created as shown in the following equation:

\[
\text{R-} \text{C} \text{CH}_2 + \text{BF}_3 \rightleftharpoons \text{R-} \text{C} \text{CH}_2
\]

The addition compound formed can thus behave in a manner similar to that of the "positive fragments." Obviously, both the carbonium ion (equation 3) and the molecule of the addition compound (equation 7) are electrophilic (acidic) and can force benzene to behave as an electrodote (base).

3. Base Catalysis.

In all the examples of basic catalysis considered in these chapters, the characteristic effect of the basic catalyst is the increase in concentration of the basic group involved in the reaction. In most reactions this is accomplished by the familiar displacement of one base by another, for example, the displacement of the ester anion 7, 26, 27 from an ester:

\[
\text{CH}_3\text{C} \text{O} \text{O-R} + [\text{O} : \text{R}]^{-1} \rightleftharpoons \text{CH}_3\text{C} \text{O} \text{O-R} + \text{O} : \text{R}^{-1}
\]

27 Arndt and Eistert, Ber., 69, 2384 (1936).
The increase in the concentration of the ester anion accounts for the increase in reaction rate of the base-catalyzed condensation of the ester,

\[
\text{CH}_3\text{CH}_2\text{C}^{(+)} + \left[ \text{CH}_3\text{C}^\text{O}^- \right]^{-1} \rightarrow \text{CH}_3\text{CH}_2\text{C}^\text{O}^- + \text{CH}_3\text{C}^\text{O}^- \text{R}
\]

to form the \( \beta \)-keto ester.

In some reactions (equation 11) displacement of a base apparently does not result, but the combination of the basic catalyst with the reacting substance merely localizes the electron excess on one atom, which can then act as a base. For example:

\[
\text{CH}_3\text{CH}_2\text{C}^{(+)} + \left[ \text{CH}_3\text{C}^\text{O}^\text{H}^- \right]^{-1} \rightarrow \text{CH}_3\text{CH}_2\text{C}^\text{O}^- + \text{CH}_3\text{C}^\text{O}^- \text{R}
\]

In the above reaction, the basic catalyst (OH\(^{-1}\)) coordinates with the carbonyl carbon and causes a shift of the C—H electron


\(^{29}\) Muller, Gawlick, and Kreutzmann, *Ann.*, 515, 97 (1934).
pair,\(^{30}\) producing C—H. The product of this reaction can behave as a base because it can furnish a hydride ion \(^{31}\) to an acid:

\[
\begin{array}{c}
\text{base} \\
\begin{array}{c}
\text{(a)} \\
\text{H}
\end{array}
\end{array} + \begin{array}{c}
\text{acid} \\
\begin{array}{c}
\text{(b)} \\
\text{H}
\end{array}
\end{array} \rightarrow \begin{array}{c}
\text{product}
\end{array}
\]

\[(12)\]

The examples of basic substances used as catalysts in these and the following illustrations do not differ fundamentally from those considered in illustrations of the Brønsted concept of a base. Admittedly, these substances are proton acceptors, but they can and do react with acid substances other than protons. The proton is only one of many acid groups or compounds having similar electrophilic characteristics. Brønsted's bases are proton acceptors because they can furnish lone pairs of electrons to form a coordinate bond with an acid whether the acid is a proton, a hydronium ion, a neutral H-acid, or any other electron-pair acceptor.


In this chapter it will be demonstrated that acid catalysis is due to the increased concentration of the acidic group involved in the reaction, caused by the addition of the acid catalyst.

In the preceding chapters it has been shown that the behavior of substances like sulfur trioxide, boron trifluoride, aluminum chloride, stannic chloride, and silver perchlorate is analogous to the behavior of H-acids in neutralization, in displacement, and in titrations with indicators. Many acid-catalyzed reactions will be discussed here in order to emphasize the fact that acidity depends on the electrophilic nature of the reagent and not on the presence of any particular element.
The mechanisms of Friedel-Crafts reactions have been studied extensively. The alkylations involve the use of olefins, alkyl halides, alcohols, ethers, and esters. The acylations make use of acids, esters, acid halides, and acid anhydrides. This type of reaction is acid-catalyzed, using such compounds as boron, aluminum, iron, tin, and other metallic halides, as well as sulfuric acid, phosphorus pentoxide, orthophosphoric acid, and hydrogen fluoride. These act as acids in the catalytic activity described because they all have a strong tendency to accept a share in an electron pair as the first step in the reaction.

2. Alkylations.

Alkyl Halides. Conductance and dielectric-constant measurements indicate the formation of ionic complexes between the catalyst and the alkyl halide:

$$
\text{R:}^+ \text{X:} + \text{Al:Cl:} \rightleftharpoons \text{R:}^+ \text{X:Al:Cl:} \rightleftharpoons [\text{R}]^{+1} + \text{Cl:} \text{Cl:} \rightleftharpoons \text{(1)}
$$

According to the Lewis concept of acids and bases the catalyst in this reaction behaves as an acid and the halide reacts as a base. Many other metallic halides catalyze this reaction in a similar manner. All of them can be shown to be acids by titrating with indicators in the proper solvent.

Simons and Archer report the use of hydrogen fluoride as the catalyzing agent. They found that the tertiary halides react readily at 0°C, the secondary halides require a temperature of 25°C, but the primary halides react only at higher temperatures (80°C), thereby affording an approximation of the relative acid characteristics of the alkyl groups. The reactivity of the above halides is in agreement with other data on the electrophilic properties of these alkyl groups.

3 Wertyporoch and Firla, Ann., 600, 287 (1933).
7 Fittig, Ann., 283, 85 (1894).
The “positive fragments” formed as a result of these acid-base reactions alkylate aromatic compounds through an electrophilic mechanism involving the electron-deficient carbonium ion (R\(^+\)).

\[
\begin{align*}
\text{acid} & & \text{base} \\
[R]+H^+ & \iff & +1 \\
\begin{array}{c}
\text{H} \\
\text{R}
\end{array} & \iff & \begin{array}{c}
\text{H} \\
\text{R}
\end{array}
\]
\end{align*}
\] (2)

Wohl and Wertyporoch have reported that boron trifluoride will not catalyze the alkylation of benzene by alkyl chlorides. Hennion and Kurtz have found that the presence of water or alcohol helps the reaction as follows:

\[
\begin{align*}
\text{base} & & \text{acid} \\
\text{H}_2\text{O} & + & \text{BF}_3 \iff \text{H}_2\text{O} + \text{BF}_3 \\
\text{acid} & & \text{base} \\
\text{H}^+ & + & \text{RCl} \iff \text{RCl} + \text{H}^+
\end{align*}
\] (3)

Burwell and Archer report that, although the alkyl chlorides and bromides are not catalyzed by boron trifluoride, the alkyl fluorides will react vigorously. They attribute this reactivity to the greater stability of the BF\(_4^-\) ion as contrasted with the BF\(_3\)Br\(^-\) ion or the BF\(_3\)Cl\(^-\) ion.

**OLEFINS.** Olefins can be employed for the alkylation of aromatic compounds by using acid catalysts such as hydrogen fluoride, sulfuric acid, phosphoric acid, phosphorus pentoxide, boron trifluoride, and aluminum chloride.

---

10 Wohl and Wertyporoch, *Ber.*, 64, 1360 (1931).
The condensation of olefins with aromatic compounds can be explained through an electrophilic mechanism involving the acid-base concept. The electrophilic (acidic) catalyst reacts with the electrodotic (basic) olefin to form an acidic intermediate which can react with the aromatic compound which is electrodotic. The reactions seem to be:

\[
\begin{align*}
\text{H} & \quad \text{Cl} \\
\text{C} & \quad \text{Al} \cdot \text{Cl} \\
\text{H} & \quad \text{Cl} \\
\text{base} & \quad \text{acid} \\
\end{align*}
\]

\[
\begin{align*}
\text{H} & \quad \text{H} \\
\text{C} & \quad \text{C} \\
\text{base} & \quad \text{acid} \\
\end{align*}
\]

Alcohols. Toussaint and Hennion alkylated benzene with alcohols, using boron trifluoride as the acid catalyst and phosphorus pentoxide, sulfuric acid, and benzenesulfonic acid as assistants. They explained the mechanism on the basis of a dehydration of the alcohol to an olefin:

\[
\begin{align*}
\text{CH}_3 & \quad \text{CH}_3 \\
\text{CH}_2 & \quad \text{C} \\
\text{H} & \quad \text{H} \\
\end{align*}
\]

The electrophilic catalyst \( \text{BF}_3 \) coordinates with the electrophilic olefin:

\[
\begin{align*}
\text{CH}_3 & \quad \text{F} \\
\text{C} & \quad \text{H} + \text{BF}_3 \quad \rightleftharpoons \quad \text{CH}_3 \\
\text{H} & \quad \text{C} \\
\end{align*}
\]

(9)

The addition compound that results is an acid because of the low electron density on the center carbon of the olefin. This strongly acidic coordination compound can make a substance like benzene behave as a base. The alkylation would next involve a mechanism \(^{23}\) such as that shown in equations 6 and 7. The fact that benzyl alcohol can be used to give diphenylmethane,\(^{24}\) however, seems to indicate that the mechanism involving olefin formation as a preliminary step is not likely.

A preferable mechanism \(^{25},^{26},^{27}\) would seem to be as follows:

\[
\begin{align*}
\text{CH}_3 & \quad \text{F} \\
\text{CH}_2 & \quad \text{F} \\
\text{CH}_2 & \quad \text{O} : \text{H} \\
\end{align*}
\]

\[
\begin{align*}
\text{CH}_3 & \quad \text{F} \\
\text{CH}_2 & \quad \text{F} \\
\text{CH}_2 & \quad \text{O} : \text{H} \\
\end{align*}
\]

\[
\begin{align*}
\text{base} & \quad \text{acid} \\
\text{base} & \quad \text{acid} \\
\text{base} & \quad \text{acid} \\
\end{align*}
\]

(10)

(11)

\(^{21}\) Pfeiffer and Wizinger, \textit{Ann.}, 461, 132 (1928).
\(^{27}\) Welch and Hennion, \textit{J. Am. Chem. Soc.}, 63, 2063 (1941).
The electrodotic (basic) alcohol reacts with the electrophilic (acidic) catalyst to form an addition compound that is capable of ionizing. The \( \text{n-propyl carbonium ion} \) (positive fragment) isomerizes to form the isopropyl ion. The function of the catalyst is to produce a relatively high concentration of these positive fragments which can react with a benzenoid compound to produce an alkylated product as follows:

\[
\begin{align*}
\text{CH}_3 & \quad \text{CH}_3 \\
\text{acid} & \quad \text{base}
\end{align*}
\]

The catalyst is regenerated by the following reaction:

\[
\begin{align*}
\left[ \begin{array}{c}
\text{F} \\
\text{H:O:B:F}
\end{array} \right]^{-1} + H^{+1} & \rightleftharpoons H:O:B:F \\
\text{acid} & \quad \text{H:F}
\end{align*}
\]

ETHERS. The catalytic effect of boron trifluoride on alkylations with ethers involves an acid-base reaction between boron trifluoride and ether. That this is a true acid-base reaction can be readily demonstrated by titrations in different solvents and with various indicators. The positive fragment that results from the ionization then reacts with the benzene, as previously described.

\[
\begin{align*}
\text{R:O:F} & \quad \text{F} \\
\text{R}_1 & \quad \text{R}_1:O:B:F
\end{align*}
\]


Ramser and Wiberg, Ber., 63B, 1136 (1930).


Hydrogen fluoride \(^{24}\) has also been used to catalyze these reactions. Whether or not the acid contains hydrogen, the behavior is essentially the same. The acid accepts a share in a pair of electrons from the base to form a coordinate bond.Ionization into single ions may then occur if the dielectric constant is high enough and the ions can be solvated. In media of very low dielectric constant considerable association into higher ionic multiples is to be expected, \(^{32-36}\) but this usually does not have to be indicated in the equations.

**Esters.** The behavior of esters toward acids may result in an alkylation or an acylation, depending on the relative tendencies of acyl, alkyl, and aryl groups to become acidic ions in the presence of aluminum chloride, boron trifluoride, metallic halides, or \(\text{H-acids}.\) \(^{36-38}\) The alkylation mechanism can be represented as:

\[
\begin{align*}
\text{O} & \quad \text{F} \\
\text{R} - \text{C} : \text{O} : + \text{B} : \text{F} & \Rightarrow \text{R} - \text{C} : \text{O} : \text{B} : \text{F} \\
\text{R}' & \quad \text{F} \\
\text{base} & \quad \text{acid}
\end{align*}
\]

The acyl group, being more electrophilic than the alkyl \((R')\) group, will not be released, \(^{36-38}\) and, therefore, in the presence of an aromatic compound an alkylation will result.

However, if \(R'\) is an aryl group which is more strongly electrophilic, the acyl group will be released as a carbonium ion \(^{39,40}\) and acylation will result. This reaction is discussed further under the section on acylation.

The following mechanism has been suggested for the hydro­fluoric acid-catalyzed \(^{41}\) alkylation:

\[
\text{O} \quad \overset{\text{base}}{\begin{array}{c}
\text{R-C:O:} \\
\text{R'}
\end{array}} + \overset{\text{acid}}{\begin{array}{c}
\text{H:}\overset{\text{.:}}{}: \\
\text{R'}
\end{array}} \Rightarrow \begin{array}{c}
\left[ \overset{\text{base}}{\text{O}} \quad \overset{\text{acid}}{\text{R-C:O:H}} \right]^{+1} \\
\left[ \text{R'} \right]^{-1}
\end{array}
\]  

(17)

Again the action of the catalyst is essentially the same whether hydrogen is present or not. The same positive fragments are produced by the catalyst.

**ALLYLIC GROUPS.** Simons and Archer \(^{42}\) investigated the alky­lations of benzene using benzyl chloride, cinnamic acid, and allyl alcohol as the alkylating agents and hydrogen fluoride as the acid catalyst. The products they obtained are in agreement with those that might be predicted on the basis of the polarization or polariz­ability of the alkylating agents. Boron trifluoride, stannic chlo­ride, and other acid catalysts also will catalyze these alky­lations.

Possible mechanisms might be as follows:

1. **Benzyl chloride.**

\[
\text{base} \quad \begin{array}{c}
\text{O} \\
\text{CH}_{3}
\end{array} \quad \overset{\text{base}}{\begin{array}{c}
\text{Cl:} \\
\text{H:}\overset{\text{.:}}{}
\end{array}} + \overset{\text{acid}}{\text{H:}\overset{\text{.:}}{}} \Leftrightarrow \begin{array}{c}
\left[ \overset{\text{base}}{\text{O}} \quad \overset{\text{acid}}{\text{CH}_{3}} \right]^{+1} \\
\text{Cl: H:}\overset{\text{.:}}{}
\end{array}
\]  

(19)

\[
\left[ \overset{\text{acid}}{\text{CH}_{3}} \right]^{+1} \quad \overset{\text{base}}{\text{C}} \quad \Leftrightarrow \overset{\text{base}}{\text{O}} \quad \overset{\text{acid}}{\text{CH}_{3}} \quad + \overset{\text{acid}}{\text{H}^{+2}}
\]

(20)


2. Cinnamic acid.

\[
\text{base} \quad \text{acid} \quad \text{base} \quad \text{acid}
\]

\[
\text{\(\text{+H}^+\)} \quad \text{\(\text{+H}^+\)} \quad \text{\(\text{+H}^+\)} \quad \text{\(\text{+H}^+\)}
\]

3. Allyl alcohol.

\[
\text{base} \quad \text{acid} \quad \text{base} \quad \text{acid}
\]

\[
\text{\(\text{+H}^+\)} \quad \text{\(\text{+H}^+\)} \quad \text{\(\text{+H}^+\)} \quad \text{\(\text{+H}^+\)}
\]

With allyl alcohol the first point of electrophilic attack is at the carbon-oxygen bond, thus producing an allyl carbonium ion which will attack a benzene molecule to produce the 3-phenyl-1-propene. This product is very susceptible to electrophilic attack by the catalyst because of the polarization of the molecule. Thus a car-
bonium ion is produced which can react with benzene to form 1,2-diphenylpropane. The polarization indicated is supported by the fact that practically no 1,3-diphenylpropane is formed in the reaction.

The use of sulfuric acid as an electrophilic catalyst produces much larger amounts of the 1,2-diphenylpropane and less of the 3-phenyl-1-propene, probably because $\text{H}_2\text{SO}_4$ is a stronger acid.

3. Acylations.

Esters. Acylation by aromatic esters,30, 40, 43, 44 mentioned in a preceding section, can be catalyzed by aluminum chloride, boron trifluoride, metallic halides, and $\text{H}$-acids as catalysts. The possible mechanism is represented by the following equations:

$$
\begin{align*}
\text{base} & \quad \text{acid} \\
\text{acid} & \quad \text{base}
\end{align*}
$$

The positive fragment released from the ester, by reaction with the acid catalyst, is the $\left(\text{R}-\text{C}^{+}\right)$ group.39, 49 The formation of the fragment represents a displacement of one acid by another. The $(\text{C}_6\text{H}_5)^{+1}$ group is not split off because it is a stronger acid than the acyl carbonium ion. Equation 29 shows the usual electrophilic attack that results in substitution in the benzenoid compound.

Aromatic esters will undergo a rearrangement (Fries reaction) to produce ortho- and para-hydroxyketones45 when treated with

acidic catalysts such as AlCl₃ or ZnCl₂. In view of our preceding discussion, this reaction is not surprising and can be explained readily as an acid-base reaction:

\[
\text{R-C:O} + \text{Al:Cl} \rightleftharpoons \text{R-C:O:Al:Cl} \rightleftharpoons \left[ \text{R-C}^{+} \right] + \left[ \text{O:Al:Cl}^{-} \right]
\]

In this special example of an acylation there is an intramolecular reaction between the acidic acyl group and the electrodotic aromatic nucleus. The fact that ortho and para derivatives are obtained can readily be explained on the basis of increased electron density at these positions, an increase that facilitates the electrophilic attack of the carbonium ion.

**Carboxy Compounds.** Carboxy compounds have been employed for acylations using hydrogen fluoride as a catalyst:

\[
\text{R-C-O-H} \xrightarrow{\text{HF}} \text{R-C-CH₃} + \text{H₂O + HF}
\]

Simons reports that carboxylic acids dissolved in liquid hydrogen fluoride produce conducting solutions:

\[
\text{R-C:O:H} + \text{HF} \rightleftharpoons \left[ \begin{array}{c} \text{O} \\ \text{R-C:O:H} \end{array} \right]^{+1} + \text{F}^- \quad (33)
\]

The resulting cation will then dissociate to form water and the acyl carbonium ion, which will acylate an aromatic compound through an electrophilic attack:

\[
\left[ \begin{array}{c} \text{O} \\ \text{R-C:O:H} \end{array} \right]^{+1} \rightleftharpoons \left[ \begin{array}{c} \text{O} \\ \text{R-C}^{(+)} \end{array} \right]^{+1} + \text{O:H} \quad (34)
\]

It will be noted that these weak carboxylic acids are being forced to act as bases by the much stronger acids used as catalysts.

**ACYL HALIDES.** The mechanism of the acylation reaction of the acid halides can be explained on the basis of the following equation:

\[
\text{R-C:Cl} + \text{H}^{+1} \rightleftharpoons \left[ \begin{array}{c} \text{O} \\ \text{R-C}^{(+)} \end{array} \right]^{+1} + \text{H:Cl} \quad (35)
\]

**CARBOXY ANHYDRIDES.** The mechanism of the acid-catalyzed acylations using a carboxy anhydride as the acylating agent could be written as follows:

\[
\begin{array}{c}
\text{R-C}^{(+)} + \text{Al:Cl} \rightleftharpoons \text{O:Al:Cl} \\
\text{R-C}^{(+)} + \text{Cl} \rightleftharpoons \text{R-C}^{(+)} \end{array}
\]

\[
\left[ \begin{array}{c} \text{O} \\ \text{R-C}^{(+)} \end{array} \right]^{+1} + \left[ \begin{array}{c} \text{O} \\ \text{R-C:O:AlCl}_3 \end{array} \right]^{-1} \quad (36)
\]

The positive fragments formed in these reactions will acylate benzenoid systems through a reaction mechanism such as that shown in equation 29.

4. Esterification of Aldehydes (Cannizzaro Reaction).

A special example of the Cannizzaro reaction results in the direct formation of an ester from two molecules of an aldehyde.\textsuperscript{48, 49} The catalytic agent used here is an acid (aluminum alkoxide) according to the Lewis theory. This acid coordinates with a molecule of the aldehyde, acting as a base, and produces an acidic intermediate, as follows:

\[
\begin{align*}
\text{R} - \text{C}:=\cdot: + \text{Al} - \text{O} &= \text{R} - \text{C}:=\cdot: \text{Al} - \text{O} - \text{R} \\
\text{base} &\quad \text{acid} \\
\text{acid} &\quad \text{base}
\end{align*}
\]  

A small amount of the aldehyde is probably already present in its reactive form:

\[
\begin{align*}
\text{R} - \text{C}:=\cdot: &\quad (+) \\
\text{H} &\quad (-)
\end{align*}
\]

The function of the catalyst seems to be to increase the concentration of the reactive form that has an electrophilic carbonyl carbon as shown in equation 37.

The next step has been considered the removal of a hydride ion from a second molecule of the aldehyde as follows:

\[
\begin{align*}
\text{R} - \text{C} - \text{O} &\rightarrow \text{Al(OR)}_3 + \text{R'} - \text{C} \\
\text{acid} &\quad \text{base}
\end{align*}
\]

\[
\begin{align*}
\left[ \begin{array}{c}
\text{R} - \text{C} - \text{O} - \text{Al(OR)}_3 \\
\text{base}
\end{array} \right]^{-1} + \left[ \begin{array}{c}
\text{R'} - \text{C} \\
\text{acid}
\end{array} \right]^{+1}
\end{align*}
\]


\textsuperscript{49} Tischtschenko, \textit{Chem. Zentr.}, 77, 1309, 1552 (1906).
The final step involves the displacement of one acid by another.

\[
[R-CH_2-O-Al(OR)_3]^{-1} + \left[ \begin{array}{c} \text{O} \\ \text{base} \end{array} \right]^{+1} \rightleftharpoons \left[ \begin{array}{c} \text{R'-C} \\ \text{acid} \end{array} \right] 
\]

\[\text{R}-\text{CH}_2-\text{O}-\text{C}-\text{R'} + \text{Al}(OR)_3 \] (39)

5. Condensation Reactions of Carbonyl Compounds.

The reactions of aldehydes with hydroxylamine, hydrazines, and semicarbazides are catalyzed by acids. These reaction mechanisms can be explained in terms of the new terminology.

The acid catalyst coordinates with the carbonyl oxygen, which is electrodotic:

\[
\text{R} \cdot \text{C} + \text{HA} \rightleftharpoons \left[ \begin{array}{c} \text{O} \\ \text{R} \end{array} \right]^{+1} \] (40)

The resulting carbonium ion behaves as an acid and thus coordinates with the electrodotic nitrogen of the above-mentioned compounds. For example, with the semicarbazide the following reactions will take place:

\[
\left[ \begin{array}{c} \text{R} \cdot \text{OH} \\ \text{C}(+) \end{array} \right]^{+1} + \text{H}_2\text{NNHCNH}_2 \rightleftharpoons \left[ \begin{array}{c} \text{R} \cdot \text{OH} \\ \text{C:N-NH-C-} \end{array} \right]^{+1} \] (41)

\(^{50}\) Conant and Bartlett, J. Am. Chem. Soc., 54, 2881 (1932).

It OR O 0 0 C-N-NH-C-NH2 + A−1 ⇄

\[
\begin{align*}
&\text{R OH} & \text{O} \\
&\text{C-N-NH-C-NH2} & + \text{HA} \\
&\text{R} & \text{H}
\end{align*}
\]

\[
\begin{align*}
&\text{R OH} & \text{O} \\
&\text{C-N-NH-C-NH2} & \rightarrow \\
&\text{R} & \text{H}
\end{align*}
\]

\[
\begin{align*}
&\text{R} & \text{O} \\
&\text{C=N-NH-C-NH2} & + \text{H2O} \\
&\text{R}
\end{align*}
\]

6. Halogenation.

**Aromatic Compounds.** The halogenation\(^{52-58}\) of aromatic compounds probably has a mechanism similar to that of the Friedel-Crafts reactions:

\[
\begin{align*}
&\text{Br} \\
&\text{Br:Br:} + \text{Fe:Br:} \rightleftharpoons \left[ \text{Br:} \right]^{+1} + \left[ \text{Br:Fe:Br:} \right]^{-1} \\
&\text{base} & \text{acid}
\end{align*}
\]

The positive fragment formed in the reaction will attack the aromatic compound and cause substitution\(^9\) to take place in the manner previously described:


Other acidic compounds like antimony pentachloride, stannic chloride, and iodine can catalyze the reaction through the heterofission of the bromine molecule and the formation of the polar complex.

Benzene can be iodinated under anhydrous conditions with silver perchlorate as a catalyst. One suggested mechanism is:

$$\text{Br}_3 + \text{AgClO}_4 \rightarrow \text{AgBr} + \text{ClO}_4^-$$

Such a simple ionic equation is probably inadequate because of the high ionic association observed in benzene. However, there is no doubt that silver ion is acidic, as shown by its behavior in forming complexes in water and by its effect on indicators in other solvents.

**Alcohols.** The catalytic action of zinc chloride on the conversion of alcohols to halides is well known. The catalyst used is an acid which coordinates with the basic alcohol. An $\alpha, \gamma$-shift of the chlorine atom accounts for the formation of the organic halide.

$$\text{R:O:} + \text{H:Cl:} \rightleftharpoons \left[ \text{R:O:H} \right]^{+1} + \left[ \text{Cl:} \right]^{-1}$$

---

60 Lewis, *J. Franklin Inst.*, 226, 293 (1938).
7. Hydrolysis.

The hydrolysis of benzyl chloride is catalyzed by the presence of Hg\(^{+2}\) salts.\(^{62}\) The electrophilic Hg\(^{+2}\) salts are here behaving as acids and react with the organic halide to form a carbonium ion:

\[
\begin{align*}
\text{CH}_2\text{Cl} & \quad + \quad \text{Hg}^{+2} \\
\begin{array}{c}
\text{base} \\
\text{acid}
\end{array}
\end{align*}
\]

\[
\begin{align*}
\text{CH}_2\text{Cl} & \quad + \quad \text{Hg}^{+2} \\
\begin{array}{c}
\text{acid} \\
\text{acid}
\end{array}
\end{align*}
\]  

\[
\begin{align*}
\text{CH}_2\text{Cl} & \quad + \quad \text{H}^{+1} \\
\begin{array}{c}
\text{base} \\
\text{base}
\end{array}
\end{align*}
\]

We see here the familiar reaction of the acid catalyst, i.e., the displacing of one acid by another. The reaction rate is increased by the catalyst because the Hg\(^{+2}\) ions increase the concentration of benzyl carbonium ions. The result is a speeding up of the reaction shown in equation 53.

8. Sulfonation and Nitration.

Boron trifluoride has been reported as an acid catalyst in sulfonation and nitration reactions.\(^{63}\) The amounts of catalyst required

---


indicated that the reaction proceeds as follows:

\[ \text{RH} + \text{HOSO}_2\text{OH} + \text{BF}_3 \rightarrow \text{R-SO}_2\text{OH} + \text{H}_2\text{O}:\text{BF}_3 \] (54)

\[ \text{RH} + \text{HONO}_2 + \text{BF}_3 \rightarrow \text{R-NO}_2 + \text{H}_2\text{O}:\text{BF}_3 \] (55)

Since the rate of sulfonation increases with increasing acid concentration in sulfuric acid of less than 100 per cent concentration and with the concentration of sulfur trioxide in fuming acid,\textsuperscript{64-66} the sulfonation probably takes place largely through the typically acidic displacement of a proton by the acid, \( \text{SO}_3 \). (See page 63.)

The catalytic function of the boron trifluoride may be to increase the concentration of the sulfur trioxide by combining with the water to form \( \text{H}_2\text{O}:\text{BF}_3 \).

\[ \text{H}_2\text{SO}_4 + \text{BF}_3 \rightarrow \text{SO}_3 + \text{H}_2\text{O}:\text{BF}_3 \] (57)

The actual mechanism of nitration is not known, but it may possibly involve the \( \text{NO}_2^+ \) ion\textsuperscript{67} as follows:

\[ \text{H} : \text{O} : \text{N} : \text{O}^- + \text{F} : \text{B} : \text{F} \rightleftharpoons \text{H} : \text{O} : \text{N} : \text{O}^- + \text{F} \text{B}:\text{F} \rightarrow \left[ \text{H}:\text{O} : \text{F} : \text{B} : \text{F} \right]^{-1} + \left[ \text{N} : \text{O} : \text{O} \right]^{-1} \] (58)

\[ \text{base} \quad \text{acid} \]

The catalytic function of the boron trifluoride may be to produce a higher concentration of the \( \text{NO}_2^+ \) ions according to equation 58.

\textsuperscript{64} Martens, Z. physik. Chem., 62, 713 (1908).

\textsuperscript{65} Pinnow, Z. Elektrochem., 21, 380 (1915).

\textsuperscript{66} Pinnow, Z. Elektrochem., 23, 243 (1917).

\textsuperscript{67} Price, Chem. Revs., 29, 51 (1941).
Hydrogen fluoride has been reported as having a strong catalytic effect on sulfonation and nitration reactions.\(^\text{68}\)

\[
\begin{align*}
&\text{\includegraphics[width=1cm]{reaction1.png}} + \text{H}_2\text{SO}_4 \xrightarrow{\text{HF}} \text{85-95°C} \\
&\text{\includegraphics[width=1cm]{reaction2.png}} \overset{\text{HF}}{\xrightarrow{0^\circ\text{C.}}}
\end{align*}
\]

\(\text{(75 per cent yield)} \quad (60)\)

\[
\begin{align*}
&\text{\includegraphics[width=1cm]{reaction3.png}} + \text{HNO}_3 \xrightarrow{0^\circ\text{C.}} \\
&\text{\includegraphics[width=1cm]{reaction4.png}} \overset{\text{HF}}{\xrightarrow{0^\circ\text{C.}}}
\end{align*}
\]

\(\text{(83 per cent yield)} \quad (61)\)

9. **Cyclic Ketone Formation.**

Aromatic carboxylic compounds can be readily cyclized to form cyclic ketones. Many acid catalysts such as AlCl\(_3\),\(^\text{69-72}\) SnCl\(_4\),\(^\text{73-76}\) HF,\(^\text{77}\) H\(_2\)SO\(_4\),\(^\text{78, 79}\) and P\(_2\)O\(_5\)\(^\text{80}\) can be used.

The catalytic effects observed in these reactions can be explained readily as instances of simple acid-base catalysis if one adopts the conception of acids and bases proposed by G. N. Lewis. One specific example is the condensation of \(\alpha\)-benzoylbenzoic acid to anthraquinone, which has been studied by several authors without arriving at a completely satisfactory explanation of the effects observed.

\[
\begin{align*}
\text{\includegraphics[width=3cm]{anthraquinone.png}} & \rightarrow \text{\includegraphics[width=3cm]{anthraquinone.png}} + \text{H}_2\text{O} \\
& (62)
\end{align*}
\]


\(^{69}\) Mayer and Stamm, *Ber.*, 56, 1424 (1923).


Deane and Huffman \textsuperscript{81} have extended rate measurements of the \textit{o}-benzoylbenzoic acid condensation into oleum of concentrations up to 29 per cent $\text{SO}_3$ at 75 and 85°. They found that the velocity constant increased steadily up to their maximum sulfur trioxide percentage.

This increase in reaction rate with increased concentration of sulfur trioxide in sulfuric acid is readily understood when considered from the Lewis viewpoint. It is merely due to increased overall acidity. Sulfur trioxide is a stronger acid than sulfuric acid. Lewis and Bigeleisen \textsuperscript{82} have shown that the acidity of sulfuric acid solutions increases with concentration of sulfur trioxide. Close correlation is evident in comparing the Lewis and Bigeleisen plot of Hammett's acidity function with the plot of variation in reaction rate with concentration of sulfur trioxide given by Deane and Huffman.\textsuperscript{81} This close correlation indicates that the condensation of \textit{o}-benzoylbenzoic acid is an acid-catalyzed reaction, and that the reaction rate does not depend upon a particular species but rather upon the acid strength. Probably $\text{SO}_3$, $\text{H}_2\text{SO}_4$, and $\text{H}_3\text{O}^{+1}$ are the three principal species involved, but the question of the exact proportion is unimportant, since the overall acidity can be measured experimentally. Of course it is obvious that in 65 per cent oleum the ratio of $\text{SO}_3$ to $\text{H}_3\text{O}^{+1}$ is very great.

According to this viewpoint, the inhibiting effect of the product anthraquinone and of other substances can readily be explained. Anthraquinone is weakly basic. Each oxygen atom can donate a share in a pair of electrons to a sufficiently strong acid.\textsuperscript{83} The presence of the anthraquinone in such a highly acidic medium simply decreases its acidity and, therefore, its effectiveness as a catalyst. A similar explanation applies to the other inhibitors tested by Deane and Huffman. Several of these substances such as $\text{AlCl}_3$, $\text{HCl}$, and $\text{CuCl}_2$ are typical generalized acid catalysts. Even these acids, strong enough to catalyze Friedel-Crafts and similar reactions, are forced to behave as weak bases by the stronger mixture of sulfur trioxide and sulfuric acid.

In view of this very great acidity of fuming sulfuric acid, the statement of Deane and Huffman that "The effect of definite added

\textsuperscript{81} Deane and Huffman, \textit{Ind. Eng. Chem.}, \textbf{35}, 684 (1943).
\textsuperscript{82} Lewis and Bigeleisen, \textit{J. Am. Chem. Soc.}, \textbf{65}, 1144 (1943).
\textsuperscript{83} Ebert, \textit{Z. Elektrochem.}, \textbf{31}, 113 (1925).
amounts of concentrated hydrofluoric acid will more fully indicate
the status of this reaction as a case of acid-base catalysis" is not
likely to be upheld. They themselves have already tried a
stronger acid than HF, namely, AlCl₃, and found that it did
not increase the reaction rate in fuming sulfuric acid but actually
decreased it.

Newman has postulated the formation of the cyclic carbonium
ion

\[
\begin{align*}
\text{O} & \quad \text{C} = \text{O}^- + 1 \\
\end{align*}
\]

as the first step in the condensation. This suggestion is based on
the formation of a quantity of pseudo ester when methyl alcohol
was present. The cyclic ion is considered to be formed by the
reaction of sulfuric acid on the o-benzoylebenzoic acid with H₃O⁺ and
2HSO₄⁻ as the other products. This explanation conforms
to the Brønsted theory of acids and bases, but obviously the
Brønsted theory cannot explain the large increase in velocity con­
stant as the amount of sulfur trioxide is increased. On the other
hand, the observed behavior is exactly what is expected on the
basis of the electronic theory of acids and bases. Sulfur trioxide
is a much stronger acid (i.e., has a greater tendency to appropriate
lone electron pairs) than H₂SO₄ or H₃O⁺.

However, it is not assumed that the hypothesis of a cyclic ion
is incorrect. The cyclic ion could be formed just as well by sulfur
trioxide as by sulfuric acid

\[
o\text{-benzoylbenzoic acid} + \text{SO}_3 \rightleftharpoons [\text{cyclic ion}]^+ + \text{HSO}_4^- \]

And so, whether the cyclic ion or some other explanation is adopted,
the ability of the Lewis theory to explain the observed catalytic
effects is not affected.

An alternative mechanism, which we prefer for reasons dis­
cussed subsequently, is as follows:

The use of \( \text{H}_2\text{SO}_4 \), \( \text{H}_3\text{O}^{+} \), or any other acid species present, in place of the sulfur trioxide, in the above equations would serve equally well since according to the Lewis theory acids coordinate with lone electron pairs. In equation 63 the sulfur trioxide behaves as a typical acid by coordinating with the oxygen electron pair to split off the \( \text{OH}^{-1} \) group, thus leaving the carbon atom electron-deficient. The coordination makes the carbon atom acidic, since it now has a great tendency to appropriate a lone electron pair to complete its octet again. The two possibilities are: the formation of the cyclic ion, and the formation of anthraquinone according to equation 64. Both reactions undoubtedly take place, since Newman seems to have demonstrated conclusively the existence of a large proportion of the cyclic ion at low temperatures. But, since the reaction under discussion is the formation of anthraquinone, it seems unnecessary to involve the cyclic ion. We prefer to regard its formation as a side reaction to, rather than the first step in, the production of anthraquinone.
The reasons for this preference are: (1) it fits the experimental facts equally as well as Newman's explanation; (2) it is simpler and more direct; (3) it is in accord with the newer explanation of the mechanism of acylations; (4) when the structural formulas of all three species are considered there seems to be no way of getting a cyclic ion from o-benzoylbenzoic acid except through the open-chain ion as an intermediate step.

Gleason and Dougherty showed that the condensation is actually first order and not merely an apparent first-order reaction because of an excess of sulfuric acid. No explanation of this order of reaction has been offered. There is a rapid evolution of large quantities of heat when oleum is added to the o-benzoylbenzoic acid. We have observed that this rapid evolution of heat on mixing is many times greater at high sulfur trioxide concentrations, corresponding roughly to the much greater velocity constant. A logical explanation of these two facts in keeping with the other aspects of the Lewis interpretation is that the carbonium ion is formed rapidly according to equation 63 upon mixing. The rate-determining step would then be equation 64, which is obviously first order. The difference in rate with increasing amounts of SO₃ is due to the shifting of the equilibrium in equation 63, so that the initial concentration of carbonium ion is greater as the acidity of the solution is increased.

10. Summary.

In all the examples of acid catalysis considered in this chapter the characteristic effect of the acid catalyst is the increase in concentration of the acid group involved in the reaction. This is usually accomplished by the familiar displacement of one acid by another. In a few reactions displacement does not result, but the combination of the acid with the reacting substance merely localizes the electron deficiency on one atom, which can then behave as an acid. One example is the alkylation by olefins.

Either way, the net result is the characteristic increase in the concentration of the acid group. The effect is similar whether the catalyst is an H-acid or any other acid.

# Chapter 11

**BASE CATALYSIS**

1. Introduction. .............................................. 137

2. Reactions of Carbonyl Compounds Containing α-H Atoms 138
   - Self-Condensation ........................................ 138
   - Aldehydes .................................................. 138
   - Ketones ..................................................... 139
   - Esters ...................................................... 140
   - Condensation with Other Carbonyl Compounds .......... 141
   - Carboxy Anhydrides ..................................... 141
   - Malonic Ester Derivatives .............................. 143

3. Reactions of Carbonyl Compounds Containing No α-H Atoms 145
   - Self-Condensation ........................................ 145
   - Use of Electron-Release Catalyst ..................... 145
   - Use of Electron-Withdrawal Catalyst ................. 147
   - Condensation with Other Carbonyl Compounds ........ 149

4. Summary ..................................................... 149

## 1. Introduction.

In this chapter, it will be shown that the characteristic effect of the basic catalyst is the increase in concentration of the basic group involved in the reaction. The mechanisms that bring about these changes will be discussed by means of well-known reactions.

The bases used as catalysts are electrodotic reagents that may be ions or molecules, e.g.,

![Chemical Structures]

and :C≡N:. These reagents are also recognized as bases, by the Brønsted theory, because they are proton acceptors. In this chapter, however, it will be shown that these bases can and do combine with many electrophilic (acidic) substances other than the proton. It will be evident that the proton is only one of many different electrophilic substances that exhibit acidic behavior.
2. Reactions of Carbonyl Compounds Containing α-H Atoms.

\[
\begin{array}{c}
\text{SELF-CONDENSATION. Aldehydes (aldol condensations). The} \\
\text{synthetic importance of these condensations is known to all organic} \\
\text{chemists. A wide variety of compounds can be prepared through} \\
\text{the condensation of two carbonyl compounds. The catalysts used} \\
\text{are bases such as acetates, carbonates, pyridine, and amines.}^1 \\
\text{These catalytic agents are bases according to both the Lewis and} \\
\text{the Brønsted theories. The mechanism}^2 \text{ of the reaction may be} \\
\text{represented as follows:}
\end{array}
\]

\[
\begin{array}{c}
\text{One base (pyridine) tends to displace the other from combination} \\
\text{with the proton. This base (the carbanion) can then react with} \\
\text{another molecule of the aldehyde which is thus behaving as an} \\
\text{acid:}
\end{array}
\]

\[
\begin{array}{c}
\text{Bell, J. Chem. Soc., 1937, 1637.} \\
\text{York, 1940, p. 343.}
\end{array}
\]
When the reaction is carried out in deuterium water no deuterium hydrogen becomes attached to the carbon, indicating that water takes no direct part in the mechanism of the reaction.

Ketones (aldolization). The effect of basic catalysts on the active $\alpha$-H atoms of ketones is the same as that produced in aldehydes. The result is the displacement of one base by another base. The preparation of diacetone alcohol from acetone using Ba(OH)$_2$ as a catalyst, is familiar to organic chemists:

$$\text{CH}_3\text{-C-CH}_3 + \left[ :\dd{O}:H \right]^{-1} \rightleftharpoons \left[ :\dd{C}:-\text{CH}_3 \right]^{-1} + \text{H}:\dd{O}:\text{H}$$

(4)

$$\text{CH}_3 :\dd{O}:(-) + \left[ :\dd{C}:-\text{CH}_3 \right]^{-1} \rightleftharpoons \left[ \text{CH}_3 :\dd{C}:(-) \:\dd{O}:-\text{CH}_3 \right]^{-1} + \text{H}:\dd{O}:\text{H}$$

(5)

$$\left[ \text{CH}_3 :\dd{O}:(-) \:\dd{C}:-\text{CH}_2:-\text{C}-\text{CH}_3 \right]^{-1} + \text{H}:\dd{O}:\text{H} \rightleftharpoons \left[ \text{CH}_3 :\dd{O}:\:\dd{C}:-\text{CH}_2:-\text{C}-\text{CH}_3 \right]$$

(6)


The basic catalysts used in this type of reaction are OH⁻, ammonia, primary amines, and secondary amines. The basic catalyst displaces a basic anion from the ketone as shown in equation 4. The result is an increase in the reaction rate for the reaction shown by equation 5, i.e., the rate-determining step in this mechanism. Equation 6 shows the regeneration of the catalyst.

Esters (Claisen reaction). The Claisen reaction probably involves an ionic mechanism similar to that shown for the aldehydes and ketones. The first step is the displacement of the ester anion by the basic alkoxide ion:

\[
\text{CH}_3\text{C}^\text{OC}_2\text{H}_5 + \text{[O:C}_2\text{H}_5\text{]}^{-1} \rightleftharpoons \text{[CH}_2\text{C}^\text{OC}_2\text{H}_5\text{]}^{-1} + \text{H:C}_2\text{H}_5
\]  

Evidence of ester anion formation according to equation 7 has been reported in the literature. Other strongly basic catalysts such as the triphenylmethyl carbanion can be used.

The rate-determining step in this condensation is probably the following:

CARBONYL COMPOUNDS CONTAINING $\alpha$-H ATOMS 141

\[
\begin{align*}
\text{CH}_3&\text{C} + \left[ \begin{array}{c} O \\
\text{acid} \\
\text{base}
\end{array} \right]^{-1} \\
\text{CH}_3&\text{C}:\text{CH}_2\text{C}-\text{OC}_2\text{H}_5 \\
\text{base}
\end{align*}
\]

\[
\begin{align*}
\text{CH}_3&\text{C} + \left[ \begin{array}{c} O \\
\text{base}
\end{array} \right]^{-1} \\
\text{CH}_3&\text{C}:\text{CH}_2\text{C}-\text{OC}_2\text{H}_5 \\
\text{base}
\end{align*}
\]

Condensation with Other Carbonyl Compounds. Carboxy Anhydrides (Perkin reaction). Carboxy anhydrides, such as acetic acid anhydride, are usually condensed with aldehydes which do not contain active $\alpha$-H atoms. Although other aldehydes could be used, in this type of reaction, the formation of undesirable by-products restricts the choice of aldehydes. With an aldehyde containing no $\alpha$-H atoms, the basic catalyst will react with the $\alpha$-H atoms of the carboxy anhydride and produce a basic anion, as shown in equation 11.

The Perkin reaction, therefore, is fundamentally the reaction of an aldehyde (with no $\alpha$-H atoms), an acid anhydride, and the sodium salt of its acid. After much controversy in regard to the function of the anhydride and its salt, it is now generally accepted that the anion of the salt serves the function of a basic catalyst. Many other bases such as alkali metal acetates, carbonates, sul-

fates, phosphates, pyridine, quinoline, and triethylamine have been used for the condensation.\textsuperscript{17-19}

Although the overall reaction is represented as

\[
\text{CHO} \xrightarrow{(\text{CH}_2\text{CO})_2\text{O}, \text{CH}_3\text{COONa}} \text{CH=CH-COOH} \quad (10)
\]

the mechanism\textsuperscript{2} of the reaction can be shown as follows:

\[
\begin{align*}
\text{CH}_3\text{C}-\text{O}-\text{C}-\text{CH}_3 + \left[ \begin{array}{c}
:O\text{O} \\
\text{base}
\end{array} \right]^{-1} & \rightleftharpoons \\
\left[ \begin{array}{c}
: \text{CH}_2\text{C}-\text{O}-\text{C}-\text{CH}_3 \\
\text{base}
\end{array} \right]^{-1} + H:O\text{C}-\text{CH}_3 & (11)
\end{align*}
\]

\[
\begin{align*}
\text{acid} & \quad \text{base} \\
\text{acid} & \quad \text{base}
\end{align*}
\]

\[
\begin{align*}
\text{H} \quad \text{H} & \quad \text{H} \\
\text{acid} & \quad \text{base}
\end{align*}
\]

\[
\begin{align*}
\text{acid} & \quad \text{base}
\end{align*}
\]

\[
\begin{align*}
\text{base} & \quad \text{base} \\
\text{base} & \quad \text{base}
\end{align*}
\]

\[
\begin{align*}
\text{H} \quad \text{H} & \quad \text{H} \\
\text{acid} & \quad \text{base}
\end{align*}
\]

\[
\begin{align*}
\text{base} & \quad \text{base}
\end{align*}
\]

\textsuperscript{17} Bakunin and Peccerillo, \textit{Gazz. chim. ital.}, \textit{65}, 1145 (1935).
\textsuperscript{18} Kuhn and Ishikawa, \textit{Ber.}, \textit{64}, 2347 (1931).
\textsuperscript{19} Muller, \textit{Ann.}, \textit{491}, 251 (1931).
CARBONYL COMPOUNDS CONTAINING $\alpha$-H ATOMS

\[
\begin{align*}
\text{H} & \quad :O: \\
\text{C} & \quad - \quad \text{CH}_2 - \quad \text{C} - \quad \text{O} - \quad \text{C} - \quad \text{CH}_3 \quad \rightleftharpoons \\
\text{H} & \quad :O: \\
\end{align*}
\]

\[
\begin{align*}
\text{H} & \quad :0: \\
\text{C} & \quad - \quad \text{CH}_2 - \quad \text{C} - \quad \text{O} - \quad \text{C} - \quad \text{CH}_3 \quad + \quad \text{H}_2\text{O} \quad (14)
\end{align*}
\]

\[
\begin{align*}
\text{H} & \quad :0: \\
\text{C} & \quad - \quad \text{CH}_2 - \quad \text{C} - \quad \text{O} - \quad \text{C} - \quad \text{CH}_3 \quad + \quad \text{H}_2\text{O} \quad \rightarrow \\
\text{H} & \quad :0: \\
\text{C} & \quad - \quad \text{CH}_2 - \quad \text{C} - \quad \text{O} - \quad \text{C} - \quad \text{CH}_3 \quad + \quad \text{H}_2\text{O} \quad \rightarrow \\
\end{align*}
\]

\[
\begin{align*}
\text{H} & \quad :0: \\
\text{C} & \quad - \quad \text{CH} = \text{CH} - \quad \text{C} - \quad \text{O} - \quad \text{C} - \quad \text{CH}_3 \quad + \quad \text{H}_2\text{O} \quad \rightarrow \\
\text{H} & \quad :0: \\
\text{C} & \quad - \quad \text{CH} = \text{CH} - \quad \text{C} - \quad \text{OH} \quad + \quad \text{CH}_3\text{COOH} \quad (15)
\end{align*}
\]

Malonic Ester Derivatives (Knoevenagel reaction). Malonic acid esters can be condensed with aldehydes through the use of a basic catalyst. The very labile $\alpha$-H atom of the malonic ester reacts with the catalyst as follows:

\[
\begin{align*}
\text{COOC}_2\text{H}_5 & \quad + \quad \text{H} : \quad \text{C} - \quad \text{H} \quad \rightleftharpoons \quad \left[ \text{COOC}_2\text{H}_5 \right]^{-1} \quad + \quad \left[ \text{H} : \quad \text{N} - \quad \text{H} \right]^+ \quad (16)
\end{align*}
\]

The anion, thus formed, will undergo an acid-base reaction with the carbonyl compound:

\[
\begin{align*}
\text{COOC}_2\text{H}_5 & \quad + \quad \text{H} : \quad : \quad \text{C} - \quad \text{H} \quad \rightleftharpoons \quad \left[ \text{COOC}_2\text{H}_5 \right]^{-1} \quad + \quad \left[ \text{COOC}_2\text{H}_5 \right]^{-1} \quad (17)
\end{align*}
\]

\[\text{Knoevenagel, } \text{Bcr.}, \text{31, 2598 (1898)}.\]
The catalyst is regenerated in the following reaction:

\[
\begin{align*}
\text{base} & \quad \text{COOC}_2\text{H}_5 \quad \text{H} \\
\text{acid} & \quad \text{H} : \text{N} - \text{H} \\
\end{align*}
\]

\[\Rightarrow\]

The $\alpha,\beta$-unsaturated acid is produced by loss of water, hydrolysis of the ester groups, and decarboxylation:

\[
\begin{align*}
\text{OH} \quad \text{COOC}_2\text{H}_5 & \quad \Rightarrow & \quad \text{COOC}_2\text{H}_5 \\
\text{H} \quad \text{COOC}_2\text{H}_5 & \quad \text{H} \quad \text{COOC}_2\text{H}_5 \quad + \text{H}_2\text{O} \\
\text{CH} = \text{CH} \quad \text{COOC}_2\text{H}_5 & \quad + 2\text{H}_2\text{O} \quad \Rightarrow \\
\text{COOH} & \quad \text{COOH} \\
\text{CH} = \text{CH} \quad \text{COOH} & \quad + 2\text{C}_2\text{H}_5\text{OH} \quad (20) \\
\text{CH} = \text{CH} \quad \text{COOH} & \quad \rightarrow \quad \text{CH} = \text{CH} \quad \text{COOH} + \text{CO}_2 \quad (21)
\end{align*}
\]

Equation 16 shows the basic catalyst removing an active $\alpha$-H atom from the malonic ester. It is apparent that, if any other carbonyl compound with active $\alpha$-H atoms is present, a similar type of reaction would take place with the catalyst.\textsuperscript{21} The best

results (fewer by-products) are obtained, therefore, when the malonic acid ester is condensed with carbonyl compounds containing no active $\alpha$-H atoms.

The presence of two carbonyl groups in malonic ester makes the $\alpha$-H atoms so reactive that it is possible to condense it with aldehydes, ketones, and esters containing only one carbonyl group, without getting too high a percentage of by-products.\textsuperscript{22} This reaction is feasible because in a competitive reaction of the carbonyl compounds containing $\alpha$-H atom with the basic catalyst, the malonic ester having the most labile hydrogens will win out.

In the previously discussed Perkin reaction, limiting the choice of aldehyde to one containing no $\alpha$-H atoms, i.e., benzaldehyde, is more important because of the less-labile H-atoms present in the aliphatic anhydrides.

3. Reactions of Carbonyl Compounds Containing No $\alpha$-H Atoms.

\[
\begin{align*}
\text{SELF-CONDENSATION. Use of Electron-Release Catalyst (Cannizzaro reaction). Fundamentally, the base-catalyzed Cannizzaro reaction involves the reaction between two molecules of an aldehyde containing no $\alpha$-hydrogen atoms.} & \\
\text{The overall result is the transfer of one aldehyde hydrogen to another aldehyde group, thus producing two radicals that could combine to form an ester.} & \\
\text{The presence of the basic catalyst prevents the ester formation, and an alcohol and a metal carboxylate result.} & \text{24, 25 The mechanism can be shown as follows:}
\end{align*}
\]

\begin{align*}
\text{acid} & \quad \text{base} \\
\begin{bmatrix}
\text{CHO} \\
\text{H}
\end{bmatrix} & + \begin{bmatrix}
\text{H} \\
\text{O}
\end{bmatrix}^{-1} \leftrightarrow \begin{bmatrix}
\text{C} \\
\text{O}
\end{bmatrix}^{-1}
\end{align*}

\text{von Auwers et al., Ann., 432, 46 (1923).}
\text{Pomeranz, Monatsh., 21, 389 (1900).}
\text{Euler and Lovgren, Z. anorg. allgem. Chem., 147, 123 (1925).}
\text{Geib, Z. physik. Chem., A169, 41 (1934).}
The amphoteric nature of benzaldehyde is shown by this base-catalyzed reaction and by the previously discussed (page 126) acid-catalyzed reaction to form an ester. In one reaction the carbonyl oxygen offers a share in an electron pair to the acid catalyst, and in the other reaction the carbonyl carbon accepts a share in an electron pair from the basic catalyst. In the base-catalyzed reactions of aldehydes containing no α-hydrogen the effect of the entering basic catalyst is to shift the carbon-hydrogen electron pair toward the hydrogen, thus favoring C—H rather than C—H (equation 22). An electron shift in this direction is caused only with electron-release groups such as OH⁻ and O₂C₂H₅⁻ ions. This change enables a hydride ion to be split off more easily than it would be if the basic catalyst were an electron-withdrawal group, such as the cyanide ion. The latter group would cause the carbon-hydrogen electron pair to shift C—H rather than C—H and thus favor the removal of a proton rather than a hydride ion. Thus we can explain the specificity of the cyanide ion as a basic catalyst in the benzoin condensation, which will be discussed later.

The reaction in deuterium water results in an alcohol containing no deuterium hydrogen attached to the carbon. Thus the transfer of a hydrogen from one aldehyde molecule to another does not involve water in the transfer mechanism.

Another example of the acidic behavior of the amphoteric aldehyde and the basic action of a catalyst in this type of reaction is the formation of benzyl benzoate under anhydrous conditions:

A second molecule of aldehyde reacts as an acid with this intermediate and removes a hydride ion, thus forming the neutral ester molecule and at the same time regenerating the basic catalyst, i.e., the benzylate ion:

\[
\begin{align*}
&\text{acid} \\
&\text{base}
\end{align*}
\]

Use of the Electron-Withdrawal Catalyst (benzoin reaction). The specificity of the basic cyanide ion used as a catalyst in the formation of benzoin has been considered puzzling, and apparently no adequate explanation has yet been suggested. Hydrogen cyanide, mercuric cyanide, and sodium hydroxide have no effect on the rate of benzoin formation, whereas sodium cyanide, potassium cyanide, and barium cyanide have a powerful catalytic effect.

The kinetics of the reaction, however, leaves little doubt that the following mechanism exists:

\[
\begin{align*}
&\text{acid} \\
&\text{base}
\end{align*}
\]

In the product formed, the cyanide group, which is an electron-withdrawal group, greatly activates the \(\alpha\)-hydrogen by causing


an electron displacement in the C—H bond which results in $\ce{C-H}$ rather than $\ce{C-H}$. Therefore, instead of a hydride ion, a proton is split off by the oxygen of a similar molecule. The net result can be considered a shift of a proton from the carbon to the oxygen of the carbonyl group. It seems reasonable to suspect that the specificity of the cyanide ion as a catalyst for this reaction is due to the fact that it has electron-withdrawal properties. The other common basic catalysts, such as $\ce{OH^-}$, $\ce{OC2H5^-}$, and the amines, all have electron-release properties, which would cause the electron shift to $\ce{C-H}$ and thus produce a hydride ion rather than a proton:

$$\begin{bmatrix}
\text{base} \\
\ce{H} \\
\text{O} \\
\text{C:C:N} \\
\text{H}
\end{bmatrix}^{-1} \Rightarrow \begin{bmatrix}
\text{base} \\
\text{H} \\
\text{O} \\
\text{C:C:N} \\
\text{H}
\end{bmatrix}^{-1}$$

(28)

A second molecule of the benzaldehyde reacts as an acid with this basic complex as follows:

$$\begin{bmatrix}
\text{base} \\
\text{H} \\
\text{O} \\
\text{C:C:N} \\
\text{H}
\end{bmatrix}^{-1} + \begin{bmatrix}
\text{acid} \\
\text{H} \\
\text{O} \\
\text{C:C:N} \\
\text{H}
\end{bmatrix} \rightarrow \begin{bmatrix}
\text{base} \\
\text{H} \\
\text{O} \\
\text{C:C:N} \\
\text{H}
\end{bmatrix}^{-1}$$

(29)

$$\begin{bmatrix}
\text{base} \\
\text{H} \\
\text{O} \\
\text{C:C:N} \\
\text{H}
\end{bmatrix}^{-1} \rightarrow \begin{bmatrix}
\text{acid} \\
\text{H} \\
\text{O} \\
\text{C:C:N} \\
\text{H}
\end{bmatrix}^{-1} + [\text{C:C:N}]^{-1}$$

(30)
**SUMMARY**

**Condensation with Other Carbonyl Compounds.** The condensation of aldehydes having no α-H atoms with other carbonyl compounds has already been discussed. The Perkin and Knoevenagel reactions fall in this category.

In discussing the above reactions, we emphasized the effect of the basic catalyst on the α-H atoms of the carbonyl compounds (i.e., carboxy anhydrides and malonic esters). The resulting anions accelerated the formation of the end products of these reactions. The anions were shown to react with an aldehyde containing no α-H atoms (equations 12 and 17).

It is obvious, from the reactions already discussed, that the basic catalyst will have practically no effect on carbonyl compounds having no α-H atoms, if other carbonyl compounds containing α-H atoms are present.

4. Summary.

In all the examples of basic catalysis considered above, the characteristic effect of the basic catalyst is the increase in concentration of the basic group involved in the reaction. This is accomplished by the familiar displacement of one base by another: for example, the displacement of the ester anion from ethyl acetate by the ethoxide ion catalyst in the Claisen reaction. In some reactions, however, displacement apparently does not result, but the combination of the basic catalyst with the reacting substance merely localizes the electron excess on one atom, which can then act as a base. An example is the shifting of the electron pair of the C—H linkage toward the hydrogen in the base-catalyzed Cannizzaro reaction, thus enabling a second molecule of aldehyde to remove a hydride ion.
Chapter 12

ALKOXIDES AS CATALYSTS

1. Introduction.

The importance and widespread use of many alkoxides as catalysts warrants a brief discussion of the role they play in reaction mechanisms. These catalysts may often lead to "surprising" results if the acidic, basic, or amphoteric nature of the alkoxide is not considered.

The alkoxides may be amphoteric, highly acidic, or highly basic. Alkali metal alkoxides such as sodium ethoxide are basic in reactions because of the ethoxide ion \( \text{OC}_2\text{H}_5^- \). Aluminum alkoxide, \( \text{Al(OR)}_3 \), would be primarily acidic in behavior because of the strong tendency of the electrophilic (i.e., acidic) aluminum atom to accept a share in a pair of electrons from an electrophilic compound (i.e., a base). Toward another, stronger acid, however, aluminum alkoxide would react as a base as does aluminum hydroxide. A survey of the literature shows several very interesting investigations in the field of catalytic condensations of aldehydes in which acidic, basic, and amphoteric catalysts are used. Kulpinski and Nord have reported the use of an alkoxide ("complex"), \( \text{Mg[Al(OR)}_4]_2 \), which, according to their experimental results, seems to exhibit amphoteric behavior.

With these facts clarified by the application of the electronic theory of acids and bases, a much better understanding of aldehyde reactions seems possible.

4 Tischtschenko, Chem. Zentr., 77, 1309, 1552 (1906).
2. Basic Alkoxide Catalysts.

Aldehydes having a $\text{--CH}_2\text{--}$ group in the alpha position (such as propanal) will behave as acids in the presence of a strongly basic catalyst like $\text{OH}^-\text{1}$ or $\text{OC}_2\text{H}_5^-\text{1}$ ions. Thus, $\text{NaOC}_2\text{H}_5$ would react with an aldehyde as follows:

$$
\text{R--C--C} + [\text{O:C}_2\text{H}_5^-\text{1}] \Rightarrow [\text{R--C--C}]^{-1} + \text{H:O:C}_2\text{H}_5\text{(1)}
$$

The actual mechanism that results in this proton removal by the basic catalyst is not particularly important. It may be a coordination of the base with the proton accompanied by an ionization, or a coordination of the basic catalyst with the carbonyl carbon followed by an $\alpha,\gamma$-shift. In either mechanism the catalytic effect is due to the increase in concentration of the carbanion that results from this first step.

This type of reaction leads to an aldolization (formation of a compound similar to aldol)

$$
\text{R'--CH}_2\text{--C} + [\text{O:C}_2\text{H}_5^-\text{1}] \Rightarrow [\text{R'--C--C}]^{-1} + \text{H:O:C}_2\text{H}_5\text{(2)}
$$

or, if a molecule of water is subsequently eliminated on acidification, crotonization (formation of a compound similar to crotonaldehyde) will result:

$$
\begin{align*}
\text{R'--CH}_2\text{--C} & \xleftarrow{\text{H}^+\text{3}} \text{R'--CH}_2\text{--C} \\
\text{H & H & H} & \text{H & H & H}
\end{align*}
$$

Aldehydes that have available only one hydrogen in the alpha position, such as the α-alkyl aldehydes (—CHR—CHO), are less acidic \(^7\) than those containing a —CH\(_2—\) group in the alpha position. This is due to the electron-release effect of the alkyl group which displaces the C—H electron pair nearer to the proton and makes the proton removal by the basic catalyst more difficult. Aldolization of such aldehydes would, therefore, require strongly basic catalysts.

It is apparent from this discussion that alkali metal alkoxides, such as NaOC\(_2\)H\(_5\), which furnish a high concentration of OC\(_2\)H\(_5—1\) ions, will behave as strongly basic (electrodotic) catalysts. Their use with aldehydes would produce an aldolization or crotonization if active α-hydrogens are available. Therefore, the experimental observation \(^5\) that alpha —CH\(_2—\) and —CHR— aldehydes behave similarly toward alkali metal alkoxides is to be expected, since these strongly basic catalysts will cause both types of aldehydes to behave as acids to produce similar types of compounds.

When no α-hydrogen is available in the aldehyde (such as benzaldehyde), a stronger base, like OH\(^−1\), is needed to catalyze the reaction by coordinating with the carbonyl carbon:

\[
\begin{align*}
\text{acid} & \quad \begin{array}{c}
\text{O} \\
\text{C} \\
\text{H}
\end{array} + \begin{array}{c}
\text{O} \\
\text{H}
\end{array}^{-1} \quad \Leftrightarrow \quad \begin{array}{c}
\text{a} \\
\text{C} \\
\text{H}
\end{array} & \quad \text{base}
\end{align*}
\]

This coordination leads to a Cannizzaro type of reaction (see Chapter 11). The effect of the coordinated OH\(^−1\) group is to displace the C—H electron pair nearer to the hydrogen C—H so that a hydride ion may be split off by another aldehyde group acting as an acid:

\[
\begin{align*}
\text{acid} & \quad \begin{array}{c}
\text{O} \\
\text{C} \\
\text{H}
\end{array} + \begin{array}{c}
\text{a} \\
\text{C} \\
\text{H}
\end{array} : \text{O} : \text{H}^{-1} \quad \Leftrightarrow \quad \begin{array}{c}
\text{b} \\
\text{C} \\
\text{H}
\end{array} & \quad \text{base}
\end{align*}
\]

Benzyl alcohol and sodium benzoate are formed in the presence of sodium hydroxide. This change has often been referred to as an oxidation-reduction reaction, but experimental evidence has shown that oxidizing agents have no catalytic effect. This result would support the conclusion that the reaction is of the generalized acid-base type.

3. Acidic Alkoxide Catalysts.

The amphoteric nature of carbonyl compounds is shown by the fact that their reactions can be catalyzed by acids as well as bases. In the preceding section it has been shown that aldehydes behave as acids in the presence of alkali metal alkoxides.

The basic behavior of carbonyl compounds such as aldehydes is illustrated by their condensation reactions in the presence of aluminum alkoxides (acids).

Alkoxides like Al(OR)$_3$ will behave as fairly strong acid catalysts, I-3 owing to the tendency of the aluminum atom in these compounds to accept a share in a pair of electrons. In view of this fact it is to be expected that aluminum alkoxides will cause the amphoteric aldehydes to behave as bases, so that a simple ester results. As this behavior involves only the basic characteristics of the carbonyl group itself it makes little difference whether we have one, two, or no $\alpha$-hydrogen atoms in the aldehyde; an ester is the result. The first step is a typical acid-base neutralization reaction with the formation of a coordinate covalent bond:

\[
\begin{align*}
R & \quad R \\
\cdot & \quad \cdot \\
:O: & \quad :O: \\
R-C:O: + Al:O:R & \rightleftharpoons R-C:O:Al:O:R \\
\downarrow & \quad \downarrow \\
H & \quad H \\
\cdot & \quad \cdot \\
:O: & \quad :O: \\
R & \quad R \\
base & \quad acid & \quad acid
\end{align*}
\]

The acidic (electrophilic) aluminum atom makes the carbonyl carbon so positive that it can remove a hydride ion from another aldehyde molecule:

---

154 ALKOXIDES AS CATALYSTS

Thus a simple ester is formed:

\[
\begin{align*}
\text{acid} & \quad \left[ R' - O - Al(OR)_3 \right]^{-1} \\
\text{base} & \quad \left[ R' - C - O - Al(OR)_3 \right]^{-1}
\end{align*}
\]

(7)


Kulpinski and Nord\textsuperscript{6} describe the use of "complex" alkoxide catalysts for the condensation of aldehydes to glycol esters. These catalysts are of a type represented by Mg[Al(OR)\textsubscript{4}]\textsubscript{2}, a substance that evidently consists of Mg\textsuperscript{2+} ions and Al(OR)\textsubscript{4}\textsuperscript{-1} ions. With aldehydes having —CH\textsubscript{2}— in the alpha position, this catalyst produces mainly glycol esters. However, with aldehydes having —CHR— in the alpha position, mainly simple esters are formed. Although these reactions may seem puzzling at first, they are readily understood when the amphoteric nature of the aldehydes and the catalysts is taken into consideration, from the viewpoint of the electronic theory of acids and bases.

These "complex" catalysts can react as weak acids (because of Mg\textsuperscript{2+}) or as weak bases (because of Al(OR)\textsubscript{4}\textsuperscript{-1}). The manner in which the catalyst reacts depends on the relative acidity or basicity of the aldehyde involved.

The aldehydes having —CH\textsubscript{2}— in the alpha position are more acidic than the aldehydes having —CHR— in the alpha position.\textsuperscript{7}
Apparently the former behave as acids toward the Al(OR)₄⁻¹ part of the catalyst, because aldolization is the result:

\[
R'-CH_2-C + [\text{Al(OR)}_4]^{-1} \rightleftharpoons R'-CH-C + H\text{Al(OR)}_4 (9)
\]

The aldol-type compounds formed in such reactions are really aldehydes having \(-\text{CHR-}\) groups in the alpha position (except acetaldehyde). These aldehydes are more basic than the aldehydes containing a \(-\text{CH}_2-\) group in the alpha position, and therefore the acidic Mg²⁺ ion rather than the basic Al(OR)₄⁻¹ might be expected to acid-catalyze the reaction in the typical way to produce an ester (glycol ester):

\[
R'-CH_2-C-CH-C + A \rightleftharpoons R'-CH_2-C-CH-C(O-) + A (11)
\]

9 Kulpinski and Nord (see note 5) report a very low yield of glycol ester with acetaldehyde. This is probably due to the fact that no alpha-alkyl aldehyde is produced here, and, therefore, practically no acidic catalysis follows the aldolization to produce an ester.
Whether the catalyst is Mg\(^{+2}\) is questionable in view of the presence of HA1(OR)\(_4\), another and probably stronger acid\(^{10}\) formed during the first step. The production of the ester, however, is obviously the result of acid catalysis. The removal of hydride ion in the second step is probably facilitated by the basic catalytic effect that the “complex” catalyst has on this \(\alpha-\text{CH}_2\) aldehyde (see equations 4 and 5).

The esters thus produced are glycol esters and involve a reaction of the aldehyde in two stages, the first of which is base-catalyzed (aldolization) and the second is acid catalyzed (ester formation). These glycol esters will be formed\(^8\) with an amphoteric catalyst or a complex catalyst such as the “complex” alkoxides.

5. Summary.

An understanding of the electronic theory of acids and bases is of great assistance in the interpretation of catalytic condensations of aldehydes. The reactions are easily explained when the acid-base characteristics of both the aldehydes and the catalysts are understood.

Strongly basic alkoxide catalysts result in aldolization (and crotonization) only; acidic alkoxides produce simple esters only. Amphoteric or “complex” alkoxides, however, may combine both reactions and produce a glycol ester.

\(^{10}\) Meerwein, *Ann.*, 455, 227 (1927).
Chapter 13

CONCLUSION

In the Preface we postponed a more detailed discussion of the real function of the electronic theory of acids and bases in chemistry until now. More discussion of the theory itself was necessary before its significance could be presented.

As first proposed by Lewis, the electronic theory of acids and bases was not directly related to the phenomena of oxidation and reduction. But we have tried to bring out the relationship between these two aspects of chemical behavior and to place them both in a larger setting. Both are manifestations of the relative attraction of two chemical substances for electrons.

The electrophilic nature of a chemical substance is not an absolute property like its density. Its action as an electrophile is relative to any number of other substances. If it is reacting with a base it gains only a share in a pair of electrons; if it is reacting with a reducing agent it gains electrons outright. Furthermore, the same substance may also behave as an electrodote toward another set of substances, i.e., as a base toward stronger acids and as a reducing agent toward oxidizing agents.

At first this relativity may seem a bit disconcerting. But it is a part of the experimental facts, however we interpret them. We have only to remember water as an example (as discussed in Chapter 4) to see how true this is. The water molecule by itself is neither electrophile (acid or oxidant) nor electrodote (base or reductant). But it behaves as an acid toward ammonia, as an oxidant toward sodium, as a base toward sulfur dioxide, as a reductant toward fluorine. All these words—acid, oxidant, base, reductant—describe the behavior of water toward other substances. This behavior does not depend upon any particular element, but upon the relative attraction of the different pairs of substances for electrons. The result is most simply expressed as the achievement of a more stable arrangement of electrons.

This more stable overall arrangement of electrons can be achieved in three ways: (1) outright transfer of electrons (oxida-
tion-reduction); (2) formation of new coordinate covalent bonds (acid-base phenomena); (3) formation of covalent bonds (odd-molecule reactions). The second of the three is the only one needing any further discussion here. New coordinate covalent bonds may be formed in two ways: (a) neutralization as discussed in Chapters 3 and 6, and (b) displacement as presented in Chapter 8 and in the chapters on catalysis. A more stable overall arrangement of electrons in the molecules concerned is brought about by either reaction.

The one unifying principle underlying all chemistry is simply this: those reactions occur which lead to the formation of molecules having the most stable electron configuration.

We now see more clearly the function of the Lewis theory of acids and bases. The elimination of the idea that acidity depends upon the presence of a particular element permits a much greater systematization of chemical reactions than has been possible in the past.

Much remains to be done in the way of quantitative measurement as suggested by this new relativity in chemistry. No doubt, as more and more experimental results accumulate, the theory will undergo modification and refinement. Eventually it may give way entirely to another, but the least that can be said for it now is that it can rid chemistry forever of one-element theories of the behavior of matter. This in itself is an accomplishment of considerable magnitude. It opens up our minds as widely as any previous idea in chemistry has ever done.
INDEX

Acceptor molecules, 16
Acetate ion, 7, 8
Acetates, basic behavior of, 138
Acetic acid, 8, 9, 47, 56
  association of, 40
  titration of, 92
Acetone, as a base, 16
  displaced by pyridine, 99
Acid, 43
Acid-analogous, 13
Acidic radicals, 62, 63
Activation, 60, 62
Acylation, mechanism of, 123
  with carboxy anhydrides, 125
  with carboxy compounds, 124
  with esters, 123
Addition compounds, 88
Alcohols, basic behavior of, 109, 117
  halogenation of, 129
  titration of, 92, 93
Aldehydes, acidic behavior of, 112
  amphoteric behavior of, 61
  basic behavior of, 126, 127
  reaction with semicarbazide, 127
Aldol condensations, 138, 151
Alkali metal acetates, catalytic action of, 141
Alkoxides, alkali metal, 151
  aluminum, 50
  basic behavior of, 111
  catalytic effect of, 111, 126, 140
Alkyl halides, alkylation with, 115
  basic behavior of, 115
Alkylation, mechanism of, 110, 116, 119
  with alcohols, 109, 117
  with alkyl halides, 115
  with allyl alcohol, 122
  with benzyl chloride, 121
  with cinnamic acid, 122
  with esters, 120
Alkylation, with ethers, 119
  with olefins, 111, 116
  with 3-phenyl-1-propene, 122
Allyl alcohol, alkylation with, 122
Aluminum alkoxides, acidic behavior of, 50, 126, 153
  catalytic effect of, 126
Aluminum amide, 56
Aluminum bromide, 43, 44
Aluminum chloride, 10, 11, 12, 19, 20, 45
  catalytic effect of, 110, 115, 117, 124
  titration of, 93
Aluminum hydroxide, 11, 56, 60
Aluminum sulfite, 11
Amide ion, 9, 55
Amino radicals, 81, 83
Ammonia, as a base, 6, 7, 9, 17, 39, 41, 42, 43
  displaced by triethylamine, 99
  solvent, 10, 56
  titration of, 92
Ammonium chloride, 60
Ammonium iodide, 10
Ammonium ion, 6, 7, 8, 9, 41
  as an oxidizing agent, 10
Ammonium sulfide, 14
Ammonium, 10
Amphoteric behavior of, aluminum amide, 56
  aluminum hydroxide, 56
  benzaldehyde, 146
  hydrogen chloride, 74
  iodide ion, 65
  quinhydrone, 75
  quinone, 75
  water, 74, 75
  zinc acetate, 56
  zinc amide, 10, 56
  zinc hydroxide, 10, 56
Anhydrides, titration of, 93
Aniline, substitution in, 83
Anion, increased concentration of, by
bases, 11, 49–52
reducing action of, 53, 54, 55, 69
Anionoid radicals, 80
Antimony pentasulfide, 14
Antimony trichloride, 12
conductance in ether, 51
Argon, boron trifluoride compound of,
59
Arrhenius, 1, 3
Arsenic trichloride, conductance of in
ether, 51
Arsenious chloride, 12
Atomic orbitals, 18, 21, 22, 23, 25
Atomic structure chart, 24
Barium hydroxide, basic behavior of,
139
Base, 43
Basic radicals, 62, 63
Benzenaldehyde, acidic behavior of, 112
Benzene, as a base, 63
solvent, 4, 57
Benzoin reaction, 147
o-Benzoylbenzoic acid, in preparation
of anthraquinone, 132
Benzyl alcohol, preparation of, 130
Benzyl chloride, alkylation with, 121
hydrolysis of, 130
Benzylate ion, catalytic effect of,
147
Beryllium chloride, 19
Berzelius, theory of acids and bases,
5, 6
Bismuth chloride, 10
Bismuth iodide, 10
Bismuth nitride, 10
Bismuth oxide, 10
Bisulfate ion, 7
Boric acid, 10
Boron amide, 10
Boron hydrides, 21
Boron trichloride, 16, 17, 19, 29, 39,
42, 43, 47
titrations of, 91, 92, 93
Boron trifluoride, 51
argon compound of, 59
Boron trifluoride, catalyst in alkyla-
tions, 109, 111, 118
catalyst in sulfonation and nitra-
tion, 130
Boyle, properties of acids, 3
Bromination of phenanthrene, 129
Bromine, basic behavior of, 128
Bromine pentafluoride, 34
Brønsted-Lowry theory, 5, 6, 7, 8, 9
Cady and Elsey, 11
Calcium chloride, 11
Calcium oxide, 5
Cannizzaro reaction, acid catalyzed,
126
base catalyzed, 112, 145, 152
Carbonanions, 77, 111, 112
Carbon dioxide, 3, 4, 6, 47, 49, 62
displaced by boron trichloride, 98
displaced by silicon dioxide, 98
displaced by stannic chloride, 98
electronic formula of, 38
secondary acid, 62
Carbon monoxide, 10
Carbon tetrachloride, as a solvent, 4
titrations in, 93
Carbonates, basic behavior of, 138,
141
Carbonium ions, 76, 84, 109, 110, 115,
118, 120, 121, 122, 123, 125, 130
Carbonyl compounds, reaction with
hydrazine, 127
reaction with hydroxylamine, 127
reaction with semicarbazide, 127
Carbonyl group, amphoteric behavior
of, 61
Carboxyls, 20
Carboxy anhydrides, acylation with,
125
Carboxy compounds, acylation with,
124
Catalysis, 15
definition, 107
Catalysts, acidic, 108
basic, 111
electron-release, 145, 152
electron-withdrawal, 145, 152
Catalytic action, of acetates, 138
of alkali metal acetates, 141
Catalytic action, of alkali metal alkoxides, 151
of alkoxide, 111
of aluminum alkoxides, 126, 153
of aluminum chloride, 110, 115, 117, 124
of barium hydroxide, 139
of benzylate ions, 147
of boron trifluoride, 109, 111, 118
of carbonates, 138, 141
of “complex” alkoxides, 154
of cyanides, 147
of hydrogen chloride, 110
of hydrogen fluoride, 108, 110, 115, 121, 122, 125
of hydroxyl ions, 112
of magnesium aluminum alkoxides, 150
of mercuric ions, 130
of phosphates, 142
of pyridine, 138, 142
of quinoline, 142
of sulfates, 142
of sulfur trioxide, 110
of triphenylmethyl carboxonium, 140
of zinc chloride, 110
Cation, increased concentration of,
by acids, 11, 47–52
oxidizing action of, 10, 11, 53, 54, 69
Chelation, 105
Chloride ion displaced by water, 99
Chlorine, 14
Chlorobenzene solutions, titrations in, 92
Chloroform as a solvent, 4
Chromic acid, 34
Cinnamic acid, alkylation with, 122
preparation of, 142
Claisen condensation of esters, 111
Claisen reaction, 140
Classification of chemical reactions,
77, 78, 79
Combination, 77, 78
Complex ions, of aluminum, 27, 39
of molybdenum, 39
of silicon, 27, 29, 39, 59
of silver, 100
of tin, 59
Complex ions, of zirconium, 39
Concentration effects, anion, 52, 70
cation, 52, 70
Condensation, of aldehydes, 138
of carboxy anhydrides, 141
of esters, 140
of ketones, 139
Conductance, antimony trichloride in ether, 51
arsenic trichloride in ether, 51
in ethylenediamine, 64
potassium iodide in ethylenediamine, 64
silver salts in ethylenediamine, 64
sulfuric acid in ether, 51
Conjugate acid, 9
Conjugate base, 9
Conjugate pairs, 7
Conjugate system, 7
Coordinate covalent bond, see Valence
Coordination compounds, 20
Covalent bond, see Valence
Crotonization, 151
Crystal violet, indicator in titrations,
16, 91, 92, 93
quinoid structure, 95
Cyanide ion, 9
Cyanides, catalytic action of, 147
Cyanate radicals, 81
Cyclic ion, in anthraquinone formation, 134
Davy, 3, 6
Decomposition, 77, 78
Deuterium water in Cannizzaro reaction, 146
Diaacetone alcohol, formation of, 139
Dimethyl ether as a base, 4
1,4-Dioxane, basic behavior of, 92
Dioxane sulfotrioxide, reaction with olefins, 109
Diphenylmethane, preparation of, 121
1,2-Diphenylpropane, preparation of, 122
3,3-Diphenylpropanoic acid, preparation of, 122
Displacement, 77, 87, 98, 99, 100
Dissociation constants, 46, 47, 48, 49,
50, 64
Donor molecules, 16
Electrostatic reagents, 71, 72
Electrolysis, 55
Electron affinity, 36, 37, 38
Electron configuration, 23, 24, 25,
26
Electronic symbols, 25, 26, 27
Electron-release catalysts, 145, 152
Electron-release effect of hydroxyl
ions, 146
Electron-withdrawal catalysts, 146
Electron-withdrawal effect of cyanide
ions, 146
Electrophilic reagents, 71, 72
Electrovalence, see Valence
Energy levels, 22, 23
Equilibrium constant, 8
Ester anion formation, 140
Esters, acidic behavior of, 111
acetylation with, 123
alkylation with, 120
basic behavior of, 120, 123
preparation from aldehydes, 126
titration of, 94
Ethers, alkylation with, 119
basic behavior of, 119
titration of, 93
Ethoxide ion, 9
Ethyl alcohol, 7, 12
Ethylenediamine, solvent, 63, 64, 65
Excitation of secondary acids and
bases, 61
Experimental criteria of acid-base
properties, Boyle, 3
Lewis, 15, 43
F-strain, 106
Ferric bromide, acidic behavior of,
128
Ferric chloride, 11
Ferric cyanide, 14
Fluorine, 30
Franklin, 9
Free radicals, 75, 76, 77
Friedel-Crafts catalysts, boron tri-
fluoride, 109, 111
Friedel-Crafts catalysts, hydrogen
fluoride, 108, 110, 115, 121, 122,
125
phosphoric acid, 108
sulfuric acid, 108
Friedel-Crafts reactions, 63
Fries reaction, 124
Germann, 10, 11
Glycol ester formation, 155
H-acids, see Hydrogen acid
Halogenation, see Hydrogen acid
Halogenation, mechanism of, 128
of alcohols, 129
of aromatic compounds, 128
of phenanthrene, 129
Heat of activation in neutralization,
60
Hexaphenylethane, 76
Hydrazines, preparation of, 127
Hydride ion removal, in carbonyl re-
actions, 153
Hydrochloric acid, 6, 7, 15
titration of, 92
Hydrogen acid, 45, 99
titration of, 92
Hydrogen as a constituent of acids,
5, 6
Hydrogen bridge, 40, 44
Hydrogen chloride, as a base, 60
catalytic activity of, 110
Hydrogen fluoride, 16
catalyst in sulfonations and nitra-
tions, 132
catalytic effect of, 108, 110, 115,
121, 122, 125
Hydrogen ion, 3
as an oxidizing agent, 10, 11
Hydrogen sulfide, as an acid, 4
Hydrogen-molecule ion, 21
Hydrolysis, 8, 63
of benzyl chloride, 130
Hydronium ion, 7, 8, 9, 10
Hydroquinone, amphoteric behavior
of, 75
Hydrosulfide ion, 9
Hydroxy ketones, preparation of, 124
Hydroxyl ion, 3, 7, 9
catalytic effect of, 112
INDEX

Hydroxylion, reducing action, 53
Hydroxylamine, reaction with carbonyl compounds, 127

Indicators, 15, 16, 95, 96
Inert solvents, 4, 57
Iodination with silver perchlorate catalyst, 129
Iodine, 55
   basic behavior of, 129
Iodine heptafluoride, 20, 32, 34
Iodine monochloride, 32
Iodine pentachloride, 32
Iodine triehloride, 32
Ionic complexes, in Friedel-Crafts reactions, 115
   in halogenations, 128
Ionization potentials, 96

Jander, 11

β-Keto esters, formation of, 112
Knoevenagel reaction, 143

Lavoisier, 3
Leveling effect, 57, 104, 105
Liebig, 3

Magnesium, reaction with water, 10, 29, 46
Magnesium aluminum alkoxides, catalytic action of, 150
Malachite green, 96
Mechanism, of acylations, 123
   of aldolization, 138, 151
   of alkylation, 110, 116, 119
   of base-catalyzed Cannizzaro reaction, 146
   of benzoin reaction, 147
   of carbonyl condensations, 138, 139, 140, 141, 143
   of Claisen reaction, 140
   of crotonization, 151
   of cyclic ketone formation, 132
   of glycol ester formation, 155
   of halogenation, 129, 130
   of Knoevenagel reaction, 143
   of nitration, 131
   of Perkin reaction, 141
   of sulfonation, 131

Mercuric chloride, as an acid, 4
Mercuric salts, acidic behavior of, 130
   catalytic effect of, 130
Meta-directing groups, 62, 81
Metals, reactions of acids with, 10, 11, 46, 53, 69
Metathesis, 77, 78
Methyl alcohol, 4
Methyl iodide, 14
Methylene blue, absorption curve, 109
Molecular compounds, 88
Molybdenum chlorides, 20

Negative fragments, 84
Neutralization, and Brønsted theory, 87, 99
   and displacement, 87, 103
   electronic theory, 15, 16, 17, 43
   Lewis theory, 15, 16, 17, 43
   positive-negative theory, 14
   primary acids and bases, 15, 16, 17, 43, 87
   secondary acids and bases, 60, 87
   solvent effect in, 102, 103
   solvent-systems theory, 10, 11, 12
   water theory, 3
Nitration mechanism, 131
Nitric acid, titration of, 92
Nitro radicals, 81, 82, 85
Nitrobenzene, substitution in, 82, 85
Nitrogen dioxide, 76
Non-metals, reactions of bases with, 53, 69
Nucleophilic radicals, 80

Octet theory, 19, 20
Odd-electron molecules, 75, 76, 77
Olefins, alkylation with, 111
Oleum, catalyst in anthraquinone formation, 133
One-electron bonds, 21
One-element theories, 5
Orbitals, atomic, 13, 21, 22, 23, 25
Ortho- and para-directing groups, 62, 81
Osmium chlorides, 35
Osmium fluorides, 20, 35
Oxidation-reduction, 69, 70, 77, 78
in positive-negative theory, 14
Oxidizing agents, ammonium ion, 10
cation of solvent, 10, 11, 53, 54, 69
hydrogen ion, 10, 11, 69
silver ion, 73
Oximes, formation of, 127
Oxonium hydroxide, 4
Oxonium salts, 3
Oxyacids, 38
Oxygen as a constituent of acids, 5
Parachor, 21
Perchloric acid, 34, 38
Periodic chart, 24
Perkin reaction, 141
Permanganic acid, 34
Phenanthrene, halogenation of, 129
Phenol, substitution in, 83, 85
Phenolphthalein, 4
3-Phenyl-1-propene, alkylation with, 122
3-Phenyl-1-propene, preparation of, 122
Phosgene, 10, 11, 12, 45, 49, 50
Phosphates, catalytic action of, 142
Phosphorus pentabromide, 30
Phosphorus pentachloride, 30
Phosphorus pentfluoride, 19, 27, 29, 30
Phosphorus trifluoride, 30
Piperidine, 9
Positive fragments, 84, 108, 109, 110, 111, 128
Positive-negative theory, 13, 14
Potassium amide, 10, 12
Potassium chloride, 12
Potassium cyanide, 14
Potassium ethoxide, 12
Potassium sulfate as a base, 12
Primary acids and bases, 60
Protective coatings, 57
Proton-donor theory, 6; see also
Brønsted-Lowry theory
Pyridine, 16, 43, 44, 45, 49, 51, 63
basic behavior of, 138
catalytic action of, 142
titration of, 91, 92, 94
Quantum numbers, 23
Quinoline, catalytic action of, 142
Radicals, acidic, 81, 82
basic, 81, 83
electron-release characteristics, 81, 83, 84
electron-withdrawal characteristics, 81, 82
meta-directing, 62, 81
nucleophilic, 80
ortho- and para-directing, 62, 81
Rare earths, 24, 28
Reducing agents, amide ion, 73
anion of solvent, 53, 54, 55, 69
hydroxyl ion, 53, 69
iodide ion, 74
sulfide ion, 73
Related metals, 21, 24, 25, 26
Representative elements, 20, 24
Resonance, 37, 38
Rule of eight, 19, 20
Rule of two, 27, 28, 32
Salts, 88
Secondary acids and bases, 60
Selenium oxychloride, 12, 51, 53, 54
Semicarbazones, formation of, 127
Sharing of electrons, 23
Sidgwick, 16
Silicon dioxide, 5
Silicon tetrachloride, 12
Silicon tetrafluoride, 29, 59
Silver hydroxide, 101
Silver perchlorate, acidic behavior of, 129
titration of, 91
Smith, 12
Sodium, 14
Sodium amide, 10
Sodium amoxide, 4
Sodium ethoxide, 4
Sodium hydroxide, 15, 16
Sodium oxide, 6, 14
Solvent, inert, 4, 57, 102
ionizable, 102
reaction of acids and bases with, 48–58
INDEX

Solvent-systems theory, 9, 10, 11, 12, 13, 45
Solvo acid, 11
Stannic chloride, 12, 16, 17, 23, 39, 54, 59
absorption curve, 109
titration of, 91, 92
Strength of acids and bases, 47, 49, 57, 58, 105, 106
Substitution mechanism, acidic radicals in, 84, 85
basic radicals in, 84, 85
Sulfates, catalytic action of, 142
Sulfide ion, 9
Sulfo radicals, 81
Sulfonation, mechanism of, 131
of benzene, 63, 109
Sulfur dioxide, electronic formula, 37
solvent, 54
Sulfur fluorides, 19, 20, 27, 31
Sulfur hexafluoride, 19, 20, 27
Sulfur monoxide, 36, 37, 54, 55
Sulfur trioxide, as an acid, 12, 14, 16, 43, 46, 47, 63, 109
catalytic activity of, 110
electronic formula of, 37
Sulfuric acid, conductance in ether, 51
electronic formula of, 33, 37, 38
solvent, 4
titration of, 92
Tetramethylammonium sulfite, 11
Thionyl chloride, 11, 54
Titanium chloride, 12
Titration, 15, 16, 92
of acetic acid, 92
of alcohols, 92, 93
of aluminum chloride, 93
of ammonia, 92
of anhydrides, 93
of boron trifluoride, 91, 92, 93
of 1,4-dioxane, 92
of esters, 94
of ethers, 96
of H-acids, 92
Titration, of hydrochloric acid, 92
of nitric acid, 92
of pyridine, 91, 92, 94
of silver perchlorate, 91
of stannic chloride, 91, 92
of sulfuric acid, 92
of triethylamine, 91, 94
Titrations, in carbon tetrachloride solutions, 93
in chlorobenzene solutions, 92
in water solutions, 92
Toluene as a solvent, 4
Tri-t-butylboron, 106
Triethylamine, 16, 53
titration of, 91, 94
Trimethylamine, 14, 16, 106
Trimethylboron, 106
Triphenylmethyl carbanion, 140
Tungsten chlorides, 20
Usanovich, 13, 14
Valence, coordinate covalence, 38, 39, 40, 41, 42
covalence, 29, 30, 31, 32, 33, 34, 35, 36
electrovalence, 28
origin of, 27
Walden, 46
Water, as an acid, 7, 45, 72, 75
as a base, 4, 8, 43, 44, 70, 72, 75
as an oxidizing agent, 75
as a reducing agent, 75
as a solvent, 12
Water solutions, titrations in, 92, 93
Water theory, inadequacy of, 3, 5
Wickert, 12
Zinc acetate, 56
Zinc amide, 10, 56
Zinc chloride, catalytic effect of, 110, 130
Zinc hydroxide, 10, 56
Zinc ion, hydrolysis, 63