

review of physiological chemistry

16th
EDITION

h.a. harper
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physiological chemistry

Preface

This *Review*, which first appeared in 1939, has continued to be prepared through the years with the intention of supplying a reasonably concise presentation of those aspects of chemistry that are most relevant to the study of biology and medicine. Through the past 15 editions, as in this one, a “whole organ” or systemic concept of biochemical phenomena has been favored, while still giving due regard to the burgeoning information on the subcellular and molecular aspects of biologic material. It is hoped that such an approach will continue to maintain the book as a direct service to students and practitioners of the health sciences related to medicine without neglecting fundamental advances in modern molecular chemistry and biology.

In the 16th edition, it was decided to reorganize the textual material completely and to recognize as co-authors the 2 colleagues who have for many years served as major contributors to the work, Professors Victor Rodwell and Peter Mayes. In addition, we have enlisted the services of several contributors who have written chapters in their specialized areas as identified in the table of contents. Laurel V. Schaubert has continued to exert her considerable artistic talents in the preparation of illustrations, structural formulas, and metabolic schemes.

It cannot be a surprise to those who have used this book over the years that it now contains substantially more pages than we started with. It can only be hoped that we have reached a satisfactory compromise between an adequate presentation of an ever-growing body of knowledge and our desire to maintain a concise presentation.

Harold A. Harper
Victor Rodwell
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San Francisco
June, 1977

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1...

Introduction

The purpose of this chapter is (1) to review certain aspects of organic chemistry relevant to the understanding of physiologic chemistry and (2) to provide certain guidelines designed to assist the learning and integration of the information presented in this book.

The early chapters of this book deal with the structures and properties of chemical compounds important in physiologic chemistry. Some of these structures will be familiar from the study of organic chemistry, but many are highly complex structures (eg, heterocyclic structures*) perhaps not previously encountered. The chemistry and the physiologic chemistry of unfamiliar molecules are largely predictable from those of structurally similar molecules as well as from the structure of molecules that possess identical functional groups.† In general, each functional group in a molecule will behave in a predictable way with respect to the reactions it will undergo. This will be a valuable guide also to the kinds of enzyme-catalyzed transformations that the group undergoes in living cells. The chemical elements which comprise functional groups will first be considered.

THE ELEMENTS OF THE SECOND & THIRD PERIODS OF THE PERIODIC TABLE

With the exception of certain metal ions, physiologic chemistry is, for the most part, related to the chemistry of the elements of the second and third periods of the periodic table.

In 1976, instruments designed to detect either new or the known forms of life were landed on the planet Mars. The experiments that were conducted assumed the existence of certain probable similarities

***Hetero atoms** (Greek *heteros* = "other") such as O, N, and S also form covalent bonds with carbon, eg, in ethylamine, $C_2H_5NH_2$, ethyl alcohol, C_2H_5OH , and ethyl mercaptan, C_2H_5SH . Hetero atoms have one or more pairs of electrons not involved in covalent bonding. Since these unshared electrons have a negative field, compounds with hetero atoms attract protons, ie, they act as **bases** (see Chapter 2). Heterocyclic structures are cyclic structures that contain hetero atoms.

†A **functional group** (eg, $-NH_2$, $-COOH$, $-OH$) is a specific arrangement of linked chemical elements that has well-defined chemical and physical properties.

Table 1-1. The elemental composition of living cells.

Element	Composition by Weight (%)	Element	Composition by Weight (%)
O	65	Cu, Zn	0.70
C	18	Se, Mo	
H	10	F, Cl, I	
N	3	Mn, Co, Fe	
Ca	1.5	Li, Sr Al, Si, Pb V, As Br	Tracest
P	1.0		
K	0.35		
S	0.25		
Na	0.15		
Mg	0.05		
Total	99.30		

†Variable occurrence in cells. No known function in most cases.

between terrestrial life and hypothetical life elsewhere in the universe. One central assumption was that extra-terrestrial life would use some or all of the same elements used by terrestrial life.

On earth, all cells, regardless of their origin (animal, plant, or microbial), contain the same elements in approximately the same proportions (Table 1-1). Thus, of the more than 100 known elements, only 19 are essential for terrestrial life. Perhaps there is some logical chemical explanation for their selection.

Six nonmetals (O, C, H, N, P, and S), which contribute almost 98% of the total mass of cells, provide the structural elements of protoplasm. From them the functional components of cells (walls, membranes, genes, enzymes, etc) are formed. These 6 elements all occur in the first 3 periods of the periodic table (Table 1-2).

The relative abundance of these 6 elements in the seas, crust, and atmosphere of earth does not by itself explain their utilization for life. Aluminum is more abundant than carbon but performs no known func-

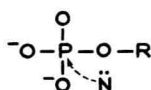
Table 1-2. The structural elements of protoplasm.

Period	Group							
	I	II	III	IV	V	VI	VII	VIII
1	H							He
2	Li	Be	B	C	N	O	F	Ne
3	Na	Mg	Al	Si	P	S	Cl	Ar

tion essential to life. By contrast, the intrinsic chemical properties of these 6 elements suggest their unique suitability as building blocks for life. Desirable features for structural elements apparently are as follows: (1) Small atomic radius. (2) The versatility conferred by the ability to form 1-, 2-, 3-, and 4-electron bonds. (3) The ability to form multiple bonds.

Small atoms form the tightest, most stable bonds—a distinct advantage for structural elements. H, O, N, and C are the **smallest atoms capable of forming 1-, 2-, 3-, and 4-electron bonds**, respectively. Utilization of all possible types of electron bonds permits maximum versatility in molecular design. So also does the ability to form multiple bonds, a property confined almost entirely to P, S, and the elements of period 2. Advantages of C- versus Si-based life include: (1) Greater chemical stability of C—C versus Si—Si bonds. (2) The ability of C, but not of Si, to form multiple bonds (eg, the oxides of C are diffusible, monatomic gases, whereas the oxide of Si is a viscous polymer). (3) The stability of C—C bonds, but not of Si—Si bonds, to rupture by nucleophilic reagents* such as O_2 , H_2O , or NH_3 .

Similar factors uniquely qualify P and S for utilization in energy transfer reactions. Energy transfer is facilitated by bonds susceptible to nucleophilic attack† (eg, nucleophilic attack of the 6-OH of glucose on the terminal P—P bond of ATP, forming ADP plus glucose-6-phosphate). P and S resemble Si in that P—P or S—S bonds, like Si—Si bonds, are susceptible to nucleophilic rupture by virtue of their unoccupied third orbitals. However, unlike Si, P and S form multiple bonds (more versatile), a consequence of their smaller atomic diameters. Most energy transfer reactions in biochemistry may be visualized as resulting from attack of a nucleophile (N) on the unoccupied third orbital of a phosphorus atom:



The characteristic chemical and physical properties of the chemical elements of life are the same throughout the known universe. It thus seems probable that if life exists elsewhere, the same elements are employed for the same or similar reasons. Taking this one step further, it seems likely that the kinds of biologic molecules formed from these elements and the kinds of reactions they might undergo would bear strong similarities to those on earth. For this reason, a biochemist is probably the scientist most likely to recognize and understand extraterrestrial life in whatever size or physical shape it might occur.

*Electron-rich elements or compounds.

†Attack of an electron-rich center upon an electron-deficient center.

REVIEW OF ORGANIC CHEMISTRY

It is believed that a sound understanding of organic chemistry is an essential prerequisite to the study of physiologic chemistry. Satisfactory knowledge of organic chemistry will enhance an understanding of the reactions of chemical compounds that are catalyzed in cells by the class of proteins known as enzymes.

This section is not intended as a complete review of organic chemistry but rather as a summary of the main points. The material should be quite familiar to those who have only recently completed the study of this branch of chemistry.

The Covalent Bond

The region in space where an electron is most likely to be found is termed an **orbital**. The sizes and shapes of different orbitals may be thought of as determining the spatial arrangements of atoms in molecules. The most fundamental of the “rules” that describe the electronic configurations of atoms is the **Pauli exclusion principle: only 2 electrons can occupy any given orbital, and these must have opposite spins**. Electrons of like spin tend to get as far away from each other as possible. Electrons in molecules occupy orbitals in accordance with similar rules.

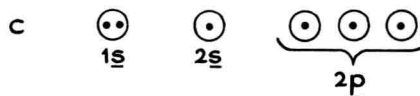
To form a covalent bond, 2 atoms must be positioned so that an orbital of one overlaps an orbital of the other. Each orbital must contain a single electron, and these must have opposite spins. The 2 atomic orbitals merge, forming a single **bond orbital** containing both electrons. Since this new arrangement contains less energy (ie, is more stable) than that of the isolated atoms, **energy is evolved when bonds are formed**. The amount of energy (per mol) given off when a bond is formed is called the **bond dissociation energy**. For a given pair of atoms, the greater the overlapping of atomic orbitals, the stronger the bond.

The carbon atom (atomic number = nuclear charge = 6) has 6 electrons, 2 of which are unpaired and occupy separate $2p$ orbitals:

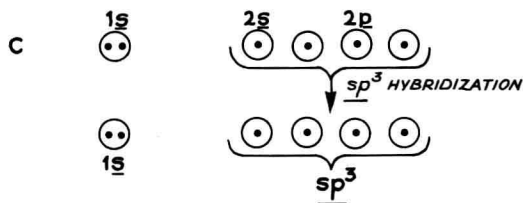


Although this suggests that C should form 2 bond orbitals with H, 4 bonds are formed, giving CH_4 . Since bond formation is an exergonic (stabilizing) process, as many bonds as possible tend to be formed. This occurs even if the resulting bond orbitals bear little resemblance to the original atomic orbitals.

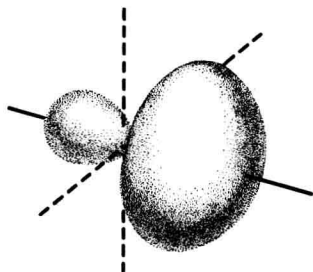
To produce a tetravalent C atom, mentally “promote” one of the $2s$ electrons to the empty p orbital:



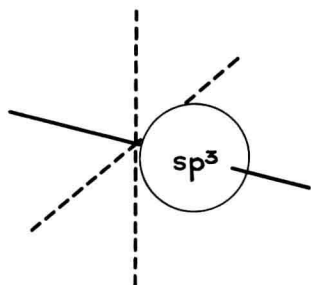
While this representation suggests C should form 3 bonds of one type (using the p orbitals) and a fourth of another type (using the s orbital), the 4 bonds of methane are known to be equivalent. The **molecular orbitals** have a mixed or hybridized character and are termed sp^3 orbitals since they are considered to arise from mixing of one s and 3 p orbitals:



sp^3 Orbitals have the following shape:



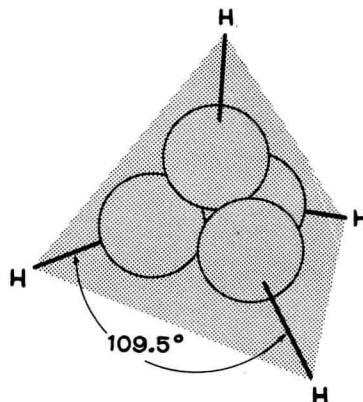
We shall neglect the back lobe and represent the front lobe as a sphere:



Concentrating atomic orbitals in the direction of a bond permits greater overlapping and strengthens the bond. The most favored hybrid orbital is therefore much more strongly directed than either s or p orbitals, and the 4 orbitals are exactly equivalent. Most important, these hybrid orbitals are directed toward the corners of a regular **tetrahedron**. This permits them to be as far away from each other as possible (recall Pauli exclusion principle).

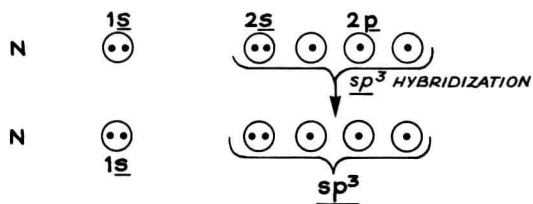
Bond Angle

For maximum overlapping of the sp^3 orbitals of C with the s orbitals of hydrogen, the 4 H nuclei must be along the axes of the sp^3 orbitals and at the corners of a tetrahedron. The angle between any 2 C–H bonds must therefore be the **tetrahedral angle** 109.5° :

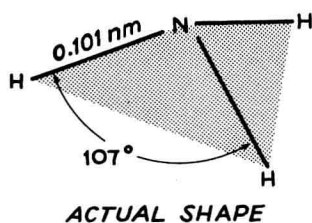
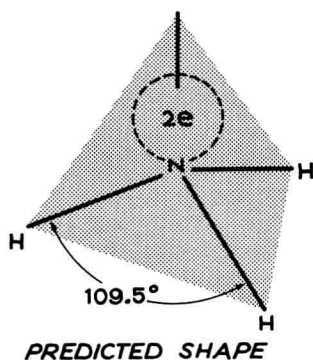


Methane has been shown experimentally to conform to this model. Each C–H bond has exactly the same length (0.109 nm) and dissociation energy (102 kcal/mol), and the angle between any pair of bonds is 109.5° . **Characteristic bond lengths, bond energies, and bond angles thus are associated with covalent bonds.** Unlike the ionic bond, which is equally strong in all directions, **the covalent bond has directional character.** Thus, the chemistry of the covalent bond is much concerned with molecular size and shape. Three kinds of C atom are encountered: **tetrahedral** (sp^3 hybridized), **trigonal** (sp^2 hybridized), and **digonal** (sp hybridized).

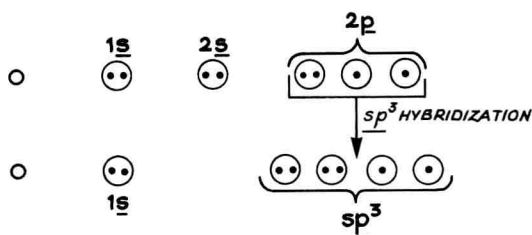
In ammonia (NH_3), nitrogen (atomic number = 7) has a valence state similar to that described for carbon: 4 sp^3 orbitals directed to the corners of a tetrahedron.



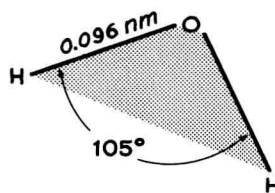
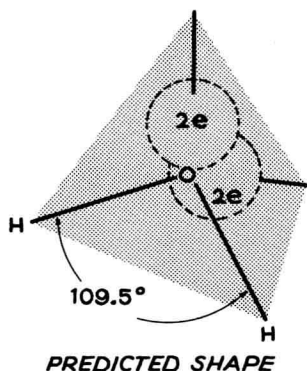
Each of the unpaired electrons of N occupying one of the sp^3 orbitals can pair with that of a H atom, giving NH_3 . The fourth sp^3 orbital contains an unshared electron pair. The unshared electron pair appears to occupy more space and to compress the bond angles slightly to 107° . It is a region of high electron density and confers on NH_3 its basic properties (attracts protons).



In H_2O , the O (atomic number = 8) has only 2 unpaired electrons and hence bonds to only 2 hydrogens.



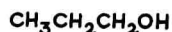
Water also is tetrahedral. The 2 hydrogens occupy 2 corners of the tetrahedron and the 2 unshared electron pairs the remaining corners. The bond angle (105°) is even smaller than that in NH_3 .



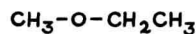
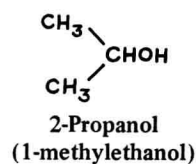
ACTUAL SHAPE

Isomers

Isomers (Greek *isos* = same; *meros* = part) are chemical compounds that have identical elemental compositions. For example, for the empirical formula $\text{C}_3\text{H}_6\text{O}$, three isomers are possible.



1-Propanol

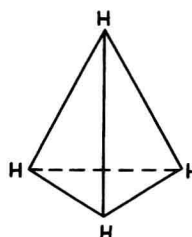


Methylethyl ether

The chemical properties of compounds having the same empirical formula are frequently quite different (eg, 1-propanol and methylethyl ether). Occasionally, they are quite similar (eg, 1-propanol and 2-propanol), and in certain special cases discussed below they are identical.

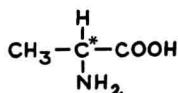
Stereoisomers

Stereoisomers differ only in the way in which the constituent atoms are oriented in space; they are like one another with respect to which atoms are attached to which other atoms. In methane, CH_4 , the 4 hydrogen atoms are at the vertices of an imaginary equilateral tetrahedron (4-sided pyramid) with the carbon atom at the center.



A carbon atom to which 4 different atoms or groups of atoms are attached is known as an asymmetric carbon

atom. For example, in the formula for alanine, the asymmetric (alpha) carbon atom is starred (*).



Alanine

Many carbohydrates, peptides, steroids, nucleic acids, etc contain 2 or more asymmetric C atoms. A thorough understanding of the stereochemistry of systems with more than one asymmetric center is therefore essential.

Representations of Spatial Relationships Between Atoms

Certain spatial relationships are readily visualized using ball-and-stick atomic models. A compound having asymmetric carbon atoms exhibits **optical isomerism**. Thus, lactic acid has 2 nonequivalent optical isomers, one being the mirror image or **enantiomer** of the other (Fig 1-1).

The reader may convince himself that these structures are indeed different by changing the positions of either enantiomer by rotation about any axis and attempting to superimpose one structure on the other.

Although enantiomers of a given compound have the same chemical properties, certain of their physical and essentially all of their physiologic properties are different. Enantiomers rotate plane-polarized light to an equal extent but in opposite directions. Since enzymes act on only one of a pair of enantiomers, only half of a **racemic mixture** (a mixture of equal quan-

tities of both enantiomers) generally is physiologically active.

The number of possible different isomers is 2^n , where n = the number of different asymmetric carbon atoms. An aldotetrose, for example, contains 2 asymmetric carbon atoms; hence, there are $2^2 = 4$ optical isomers (Fig 1-2).

To represent 3-dimensional molecules in 2 dimensions, **projection formulas**, introduced by Emil Fischer, are used. The molecule is placed with the asymmetric carbon in the plane of the projection. The groups at the top and bottom project **behind** the plane of projection. Those to the right and left project **equally above** the plane of projection. The molecule is then projected in the form of a cross (Fig 1-3).

Unfortunately, the orientation of the tetrahedron differs from that of Fig 1-1. **Fischer projection formulas may never be lifted from the plane of the paper and turned over.** Since the vertical bonds are really **below** the projection plane while the horizontal bonds are **above** it, it also is not permissible to rotate the Fischer projection formula **within the plane of the paper** by either a 90-degree or a 270-degree angle, although it is permissible to rotate it 180 degrees.

A special representation and nomenclature for molecules with 2 asymmetric carbon atoms derives from the names of the 4-carbon sugars erythrose and threose. If 2 like groups (eg, 2 -OH groups) are on the same side, the isomer is called the "**erythro**" form; if on the opposite side, the "**threo**" isomer. Fischer projection formulas inadequately represent one feature of these molecules. Look at the models from which these formulas are derived. The upper part of Fig 1-2 represents molecules in the "**eclipsed**" form in which the groups attached to C_2 and C_3 approach each other as

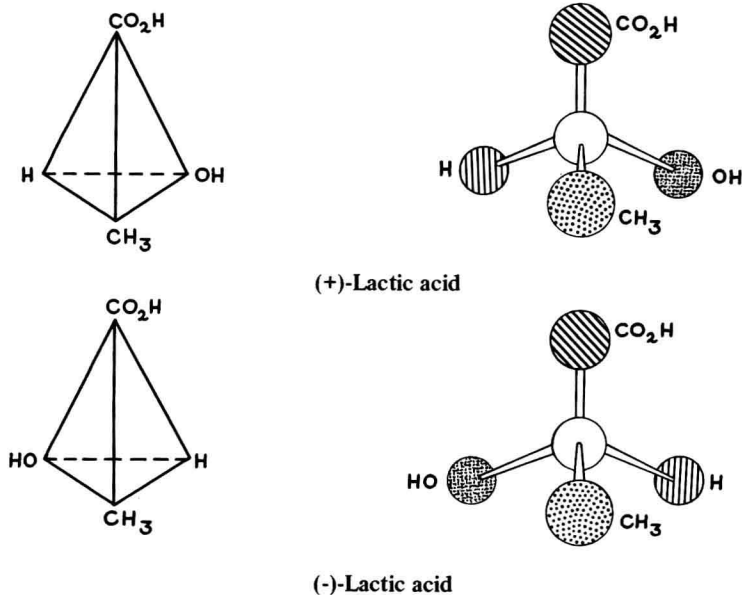


Figure 1-1. Tetrahedral and ball-and-stick model representation of lactic acid enantiomers.