Chapter 22 Peptides, Proteins, and α -Amino Acids

from Organic Chemistry

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Chapter Outline of the Book

- 1. Organic Molecules and Chemical Bonding
- 2. Alkanes and Cycloalkanes
- 3. Haloalkanes, Alcohols, Ethers, and Amines
- 4. Stereochemistry
- 5. Organic Spectrometry

II. Reactions, Mechanisms, Multiple Bonds

- 6. Organic Reactions *(*Not yet Posted*)
- 7. Reactions of Haloalkanes, Alcohols, and Amines. Nucleophilic Substitution
- 8. Alkenes and Alkynes
- 9. Formation of Alkenes and Alkynes. Elimination Reactions
- 10. Alkenes and Alkynes. Addition Reactions
- 11. Free Radical Addition and Substitution Reactions

III. Conjugation, Electronic Effects, Carbonyl Groups

- 12. Conjugated and Aromatic Molecules
- 13. Carbonyl Compounds. Ketones, Aldehydes, and Carboxylic Acids
- 14. Substituent Effects
- 15. Carbonyl Compounds. Esters, Amides, and Related Molecules

IV. Carbonyl and Pericyclic Reactions and Mechanisms

- 16. Carbonyl Compounds. Addition and Substitution Reactions
- 17. Oxidation and Reduction Reactions
- 18. Reactions of Enolate Ions and Enols
- 19. Cyclization and Pericyclic Reactions *(*Not yet Posted*)

V. Bioorganic Compounds

- 20. Carbohydrates
- 21. Lipids
- 22. Peptides, Proteins, and α -Amino Acids
- 23. Nucleic Acids

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22: Peptides, Proteins, and α-Amino Acids

•Peptides

Protein Structure and Organization
Properties of α-Amino Acids
Enzymes and Enzyme Catalysis

Preview

Proteins are a major class of bioorganic molecules present in all organisms. They contain one or more **polypeptide** chains with the repeating general structure -(NH-CHR-C(=O))-. These repeating units come from 20 different chiral α -amino acids with the general structure H₂N-CHR-CO₂H. The R groups play a major role in determining conformations of the peptide chains and the shapes of proteins. Free α -Amino acids are polyprotic acids because they have at least two functional groups (CO₂H and NH₂) with acid and conjugate base forms. Some of these 20 α -amino acids also have an acid-base functional group in R. *Enzymes* are proteins that catalyze biochemical reactions. These enzyme-catalyzed reactions take place in specific regions of the enzymes called **active sites.**

22.1 Peptides

Peptides are polymers of α -amino acids joined by *amide* or *peptide bonds*.

Figure 22.1



Peptide Structure (22.1A)

Peptides with two α -amino acid components are **dipeptides**, those with three are **tripeptides**, and so on. Peptides with 3-10 α -amino acid components are called **oligopeptides**, while those with many α -amino acid components are **polypeptides**. These terms are similar to the terms *disaccharide*, *oligosaccharide*, and *polysaccharide* that we learned when we studied carbohydrates (Chapter 20).

 α -Amino Acids in Peptides. Hydrolysis of peptides cleaves amide bonds (Chapters 15 and 16) releasing the individual α -amino acids.

Figure 22.2



 α -Amino acids in naturally occurring peptides generally have one R group and one H on C $_{\alpha}$ (H₂N-C $_{\alpha}$ HR-CO₂H). We will see later in this chapter that the NH₂ and CO₂H groups of free α -amino acids exist as NH₃⁺ and CO₂⁻ so the general structure of these amino acids is actually ⁺H₃N-CHR-CO₂⁻.

Figure 22.3



 α -Amino Acids Can be D or L. When R is a group other than H, these α -amino acids are chiral compounds with two enantiomeric forms because C_{α} is chiral (Chapter 4). We can identify the configuration of C_{α} as R or S if we know the structure of the C_{α}-R group, but we usually describe the two enantiomers as D or L as we show using Fischer projections in Figure 22.4. Most α -Amino acids in naturally occurring peptides have L-configurations.



The Definition of D and L for α -Amino Acids. Definitions of D and L for enantiomers of α -amino acids are based on those of D and L-glyceraldehyde (Chapter 20) (see Figure 22.5)[next page]. We define C(=O)O⁻ of the amino acid as equivalent to C(=O)H of glyceraldehyde (both have C=O) and place it at the top of the α -amino acid Fischer projection. We define R of the amino acid as equivalent



to CH₂OH of glyceraldehyde and place it at the bottom of the Fischer projection. The result is a pair of enantiomers with NH₃⁺ to the right (defined as D) or to the left (defined as L) on C_{α}. Although we will see below in Figure 22.6 that one α -amino acid (*proline*) has an additional bond between the aminium group and R (⁺H₂N-R), its D and L assignments similarly depend on the "right" or "left" orientation of its C_{α}-N bond.

The R Groups. While α-amino acids have many different R groups, we focus here on the 20 R groups of the "**standard**" α-amino acids (Figure 22.6) in naturally occurring peptides. Figure 22.6 (continued next page)

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Figure 22.6 (continued)
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Unchanged Pola R Grou	T-(6)	
HO-CH2-C2H-C02 NH3+	serine Ser	5
HO-CT(-0H)H-CL	threevine Th	n T
40-0-04-Cx	typosine Ti	n y
HS-CH2-CJ	Cepteine Cy	• C
H_N(O=)C - CH-C~	appropriagine As	n N
H2N(0=)C-CH2CH1-C2	glutamine Gl	n Q
charged Polo R Group	5(5)	
(Do (0=) C - CH2-C+	aspertracid A	P D
OO(O=) C CH CH C	glutanic acid G	lu E
$+H_3N-(OH_3)-C_4$	lepine l	up K
HN(H, N=)C -NH-(CH)- Ca	Orginino F	hg R
HT CH - Ca	histidens t	Lis H

You can see that these "standard" R groups are diverse. They include alkyl and aryl groups, heterocyclic rings, sulfur containing groups, alcohols, aminium ions, carboxylates, and amides. The R group of *proline* differs from the others because it chemically binds to its α -amino group. Also note in Figure 22.6 that both *isoleucine* and *threonine* each have an additional chiral C* (C3) in their R groups. The naturally occurring stereoisomer of *L-isoleucine* is *S* at C3, while C3 of *L-threonine* is *R* (Figure 22.7)

Figure 22.7 [previously Fig. 22.9]



Nonstandard R Groups. There are a few R groups in naturally occurring peptides beyond those shown in Figure 22.6. These "**nonstandard**" groups (Figure 22.9) arise from biosynthetic modification of a *"standard"* R group (shown in Figure 22.9) that is already present in a peptide. [There is no Figure 22.8]

Figure 22.9 [previously Fig. 22.7]

Categories of "Standard" Amino Acids. In order to learn their names and R group structures, it is helpful to group the "standard" amino acids in the three categories shown in Figure (graphic 22.6). R groups of the 5 "**charged polar**" amino acids are electrically charged (- or +) at physiological pH values (about pH 7), while those of the remaining 15 amino acids are electrically neutral under the same conditions. R groups of the 5 "**uncharged polar**" amino acids form hydrogen bonds to each other and to water, while R groups of the "**nonpolar**" amino acids do not form hydrogen bonds.

Abbreviated Names. Each "standard" amino acid name has a *three-letter abbreviation* as well as a *one-letter designation* (Figure 22.6). The three-letter abbreviations are usually the first three letters of the full name, while one-letter designations correspond to the first letter of the name where possible. You will often see three-letter abbreviations used interchangeably with full names.

Glx and Asx. The abbreviation *Glx* refers to both *Glu* and *Gln*, and *Asx* refers to both *Asp* and *Asn*. Amide groups of *Gln* and *Asn* sometimes hydrolyze to carboxylate groups of *Glu* and *Asp* during amino acid analysis (described later in the chapter).

Figure 22.10

As a result, the relative amounts of carboxylate and amide side chains in a naturally occurring peptide may be uncertain so they are grouped together as *Glx* or *Asx*.

Peptide Synthesis (22.1B)

Laboratory syntheses of peptides make use of familiar reactions illustrated below. *Biosynthesis* of peptides involves *nucleic acids* so we defer this topic until Chapter 23.

General Considerations. If we wish to make a dipeptide from two amino acids $(AA_x \text{ and } AA_y)$, we must recognize that it can have two possible structures $(AA_x-AA_y \text{ and } AA_y-AA_x)$ (Figure 22.11)[next page]. Chemists write peptide formulas so that the first amino acid in the sequence is **N-terminal** (it has the unreacted NH₂ group), while the last one is **C-terminal** (it has the terminal CO₂H group). As a result, AA_x is the *N-terminal* amino acid in AA_x -AA_y while AA_y is the *N-terminal* amino acid in AA_x -

Each amino acid has an amino group and a carboxylic acid group, so we might expect that direct reactions of AA_x and AA_y (or of appropriate derivatives) may give not just AA_x - AA_y



and AA_y - AA_x , but AA_x - AA_x and AA_y - AA_y as well (Figure 22.12). Since the number of possible combinations of amino acids increases rapidly as the size of the desired peptide increases, chemists do not use direct reactions of amino acids to make peptides.

Figure 22.12

Automated Peptide Synthesis. We can avoid the problem of multiple products by using **automated peptide synthesis**. We illustrate its general features here for the synthesis of the tripeptide AA_z - AA_y - AA_x :

Figure 22.13

AAX + Nesin
$$\longrightarrow$$
 AAX-Resin \xrightarrow{AAy} AAy-AAX-Resin
Step(1) Step(1) Step(3) \downarrow AAZ
AAZ-AAy-AAX + Resin $\xrightarrow{Step(4)}$ AAZ-AAY-AAX-Resin

(1) Chemically bind AA_x to a solid insoluble resin.

(2) Couple AA_v to AA_x -Resin.

(3) Couple AA_z to AA_y - AA_x -Resin.

(4) Remove $AA_z - AA_y - AA_x$ from the resin.

To accomplish these general steps we **protect** and **deprotect** NH₂ groups and **activate** CO₂H groups. One way chemists *protect* NH₂ groups is with *tert*-butyloxycarbonyl (*t*-Boc) groups from *tert*-butyloxycarbonyl chloride (t-BocCl).

$$(CH_{3}C-O-E-CH+H_{1}N-CH-CEO)OH \longrightarrow (CH_{3}C-O-E-NH-CH-C(=0)OH$$

$$\xrightarrow{-HCP} Deprotection \int CF_{3}CO_{1}H \\ CH_{2}C=CH_{2} + CO_{2} + H_{1}N-CH-C(=0)OH$$

Trifluoroacetic acid in methylene chloride *deprotects* NH_2 groups by removing the *t*-Boc groups. We *activate* CO_2H groups on N-protected amino acids (*t*-Boc-AA) by treating them with **dicyclohexylcarbodiimide** (**DCCD**). The resulting carboxylic acid derivatives (C(=O)-O-DCCD) react readily with NH_2 groups to form amide bonds (C(=O)-NH).

Figure 22.15



The detailed steps of automated peptide synthesis are:

(1) Attach AA_x to the Solid Support

Couple CO₂H groups of N-protected AA_x (*t*-Boc-AA_x) with benzyl chloride groups (Ar-CH₂-Cl) on the *resin solid support* to form esters. Deprotect *t*-Boc-AA_x-Resin to give reactive NH₂ group on each AA_x.

Figure 22.16



(2) Couple AA_y to AA_x

Activate N-protected AA_y (*t*-Boc-AA_y) using DCCD and couple it with AA_x -Resin. (Figure 22.17)[next page] An amide bond forms between AA_y and AA_x leading to *t*-Boc-AA_y-AA_x-Resin. Subsequent deprotection gives AA_y -AA_x-Resin with a reactive NH₂ group on each AA_y.



(3) Couple AA_z to AA_y

Activate N-protected AA_z (*t*-Boc-AA_z) using DCCD and couple it with AA_y -AA_x-Resin (Figure 22.18). An amide bond forms between AA_z and AA_y leading to *t*-Boc-AA_z-AA_y-AA_x-Resin.

Figure 22.18

(4) Isolate AA_z - AA_v - AA_x

Treat *t*-Boc-AA_z-AA_y-AA_x-Resin with liquid HF cleaving *t*-Boc from AA_z, and the ester bond between AA_x and the resin, giving free AA_z-AA_y-AA_x (Figure 22.19)[next page].

Protection of Functional Groups on R. We must also protect functional groups such as NH₂, OH, SH, and CO₂H on amino acid side chains (R) during peptide synthesis by converting them into derivatives such as benzyl groups (Figure 22.20)[next page]. These benzyl groups cleave along with the *t*-Boc group and the ester linkage to the resin during treatment with liquid HF in Step 4.

Figure 22.19

$$\frac{1}{4} - Boc - AA_2 - AA_y - AA_x - Resin \\
HF \downarrow$$

$$\begin{pmatrix}
(H_3)_2 C = CH_1 + Co_1 + (AA_2 - AA_y - AA_x) + Resin \\
HN - CH - C(=0) - NH - CH_2(=0) - NH - CH_2($$

22.2 Protein Structure and Organization

Proteins are biological molecules made up of one or more polypeptides. The amino acids of the polypeptides, and their relative order in the polypeptide, are the protein's **primary** (1°) structure. The 3-dimensional structure of localized segments of the protein's polypeptides is **secondary** (2°) structure, while the overall shape of a protein is its **tertiary** (3°) structure. **Quaternary** (4°) structure describes the way that the individual polypeptides interact with each other in a protein with multiple polypeptides.

Primary (1°) Structure (22.2A)

Amino acid content and amino acid sequence are our first concern in elucidation of the structure of a protein.

Content. We identify and quantify the amino acids in a polypeptide using an **automated amino acid analyzer** that performs the following steps illustrated in Figure 22.21 [next page].

(1) Hydrolysis of the polypeptide to its component amino acids by heating in strong acid (6 M HCl) or strong base (2 to 4 M NaOH).

(2) Separation of the resulting mixture of amino acids into its individual amino acids by column chromatography. [continued on page 14]



(3) Derivatization of the amino acids so they are detectable by spectroscopic methods such as the examples we show in Figure 22.22 (Sometimes derivatization is done before chromatographic separation (step 2)).

Figure 22.22



(4) Identification of each spectral signal in the spectrum as that of a specific amino acid and determination of the amount of amino acid present from the signal intensity.

Sequence. We determine the *sequence* of amino acids in a polypeptide by identifying its component amino acids one at a time starting at its N-terminus or C-terminus. An important **end-group analysis** procedure for N-terminal amino acids is **Edman Degradation** that first derivatizes and then cleaves the N-terminal amino acid of a polypeptide leaving the rest of the chain intact (Figure 22.23).



Automated sequenators carry out N-terminal amino acid derivatization, cleavage, and analysis in a repetitive stepwise fashion allowing accurate determination of as many as 100 amino acid units in a peptide chain.

We can analyze C-terminal amino acids using **carboxypeptidase enzymes**. Such enzymes specifically hydrolyze (cleave) the peptide bond between the C-terminal amino acid and the rest of the chain.

Figure 22.24

 $\frac{-1}{2} = \frac{1}{2} = \frac{1$

They are useful for analyzing only the first few amino acids at the C-terminal end of a peptide because they cleave different C-terminal amino acids at different rates. As a result, C-terminal amino acids of shortened chains formed during the analysis procedure may cleave more rapidly than those of the original polypeptide. This can quickly complicate the reaction mixture with a variety of individual amino acids and shortened peptide chains. **Aminopeptidase enzymes** similarly operate on the N-terminal end of a peptide and have the

same limitations as *carboxypeptidases*.

When peptide chains are too long for accurate determination of their full sequence by a method such as Edman degradation, we can cleave them into smaller peptides using **specific peptide cleavage reactions**. We determine the sequences of these smaller peptides and use them to reconstruct the original sequence of the complete peptide. One of the most specific peptide cleavage reactions uses the enzyme **trypsin** to cleave the peptide bond in $CH(R_x)C(=O)$ —NHCH(R_y) where R_x is from *Lys* or *Arg*, while R_y is from any amino acid except *Pro*. As a result, the C-terminus of each cleaved segment is *Lys* or *Arg*, with the exception of the segment with the original C-terminus (Figure 22.25) [next page].

More about Enzymatic Peptide Cleavage. The *carboxypeptidases* and *aminopeptidases* that we have just described are **exopeptidases** because they cleave the *terminal* amide bonds connecting N-terminal or C-terminal amino acids to the rest of the peptide chain. In contrast, *trypsin* is one of several enzymes that cleave amide bonds "inside" the peptide chain (**endopeptidases**). Others are α -

chymotrypsin, elastase, and pepsin. *Enzymes* are *proteins* that catalyze specific types of reactions.

We will discuss enzyme catalysis in the last section of this chapter.

Figure 22.25



Separation of Individual Peptide Chains. Polypeptide chains of proteins with two or more peptide chains often covalently bind to each other with disulfide bonds (S-S) formed from neighboring SH groups of the amino acid *Cys*.

Figure 22.26



In order to determine content and sequence of individual polypeptide chains of such proteins, we must break these disulfide bonds by *reduction* or *oxidation*. We then separate the chains using **denaturing agents** and isolate them using chromatography. We describe *denaturing agents* later in this chapter.

Secondary (2°) Structure (22.2B)

Secondary (2°) protein structure includes such localized structural features of peptide chains as the α -helix, the β -pleated sheet, and the *planarity* of amide groups.

Planarity of Amide Groups. We learned in Chapter 15 that amide groups have significant electron delocalization (Figure 22.27)[next page]. This causes them to be planar since rotational barriers about their C-N bonds are 60 to 75 kJ/mol (rotational barriers about C-N single bonds are 12 to 20 kJ/mol).





While substituents on amide bonds are *cis* or *trans*, the most favorable conformation in peptides is that where C_{α} 's of the chain are *trans*. As a result, we can represent peptide chains as a series of "planes", containing the amide groups, that connect at the C_{α} 's (Figure 22.28). Since these planes have a restricted range of relative orientations, amide groups impart significant rigidity to a polypeptide.

Figure 22.28



Helical Structures. The α -*helix* structure that Pauling and Corey proposed in 1951 is a result of repeating planar *trans* amide groups in peptides. The C=O of each amide group in the α -helical region hydrogen bonds to an N-H of another amide group. Two intervening amide groups separate the hydrogen bonded amide groups leading to a favorable distance of 2.8 Å between O and N in the resulting C=O·····H-N hydrogen bonds (Figure 22.29)[next page]. Since C $_{\alpha}$'s have L-configurations, the α -helix adopts a **right-handed** twist so the C $_{\alpha}$ -R groups point away from the chain. The α -Helical regions contain ~12 amino acid residues.

β-Pleated Sheets. Segments of peptide chains separated by many intervening amino acids also hydrogen bond to each other to give *β-pleated sheets* (Figure 22.30)[next page]. These *β-pleated sheets* proposed by Pauling and Corey in 1951, are **parallel** or **antiparallel** and the hydrogen bonded segments can be from the same or different peptide chains. A "sheet" includes 2 to 15 peptide chain segments that average about 6 amino acid residues in length. You can see from the figure that C=O····H-N bonds in *parallel* β-pleated sheets are distorted and they are less stable than those in *antiparallel* sheets where there is no distortion.







Other Structures. α -Helical regions and β -pleated sheets connect using other peptide structures called **coils** or **loop conformations**. α -Helices and β -pleated sheets make up about 50% of peptide chains.

Tertiary (3°) Structure (22.2C)

We generally describe proteins at the 3° structural level as **globular** or **fibrous**. Proteins of both types contain all of the 2° structural elements we have just described.

Fibrous Proteins. The *fibrous* protein of skin, bone, and muscle has an elongated "fiber-like" shape. An example is α -keratin of hair, horns, and nails.



Two α -helical polypeptide chains wind together and group with other polypeptide dimers to form **protofilaments**. Two **protofilaments** comprise a **protofibril** and four **protofibrils** wind together in a **microfibril**. Microfibrils cluster to form **macrofibrils** which come together in groups to form **cells**. A strand of hair includes many cells grouped in a sheath.

Globular Proteins. The shapes of *globular* proteins also reflect their name. Most *enzymes* are *globular* proteins, and there are many others such as the familiar oxygen transport proteins **myoglobin** and **hemoglobin**.

Figure 22.32 (Schematic Representations)



Myoglobin has one peptide chain with 153 amino acid residues surrounding a heme group containing an Fe(II) atom. In muscle tissue, it accepts O_2 from oxygenated *hemoglobin* in blood. *Hemoglobin* has four polypeptides with heme groups that have globular tertiary structures like *myoglobin*. *Hemoglobin* binds oxygen in the lungs, transports it to myoglobin, and returns to the lungs with CO₂ formed as a metabolic product in muscle tissue.

Factors that Determine Protein Shape (22.2D)

The 3° structure of a protein depends on the interactions of the amino acid side chains (the R groups) with each other and with the medium surrounding the protein. These interactions involve *electrostatic forces, hydrogen bonding*, **hydrophobic bonding**, and *disulfide bonds*.



We will see below that the same forces also bind together the multiple peptide chains of a protein in its *quaternary* (4°) structure.

Hydrophobic Bonding. A major influence on the shape and stability of a protein is the desire of *nonpolar* amino acid side chains to minimize their exposure to H_2O . This leads them to the interior of a protein where they interact with each other by *hydrophobic bonding*. For example, nonpolar alkyl groups induce complementary dipoles in each other. While they are individually weak, these attractive induced dipolar interactions taken together have a major effect on the shape and stability of a protein (Figure 22.33). Side chains with permanent dipolar groups also interact attractively with each other and induce dipoles in nonpolar groups as well.

Thermodynamics of Hydrophobic Bonding. You may be surprised to learn that the enthalpy (H) of a hydrocarbon in water is more favorable (lower) than the enthalpy of the same hydrocarbon in a hydrocarbon medium. In spite of this, hydrocarbons prefer to dissolve in hydrocarbons and not water because the entropy (S) of a hydrocarbon in water is less favorable (much lower) than it is in a hydrocarbon medium. Water molecules form highly structured cages around hydrocarbon molecules leading to a decrease in volume of the aqueous solution and a decrease in the entropy of the system. The net result of these effects on H and S is that the free energy (G) for the hydrocarbon in water is greater (less favorable) than in a hydrocarbon medium. The unfavorable $T\Delta S_{(water \rightarrow hydrocarbon)}$ term overwhelms the favorable $\Delta H_{(water \rightarrow hydrocarbon)}$ term in the equation $\Delta G = \Delta H - T\Delta S$ so that $\Delta G_{(water \rightarrow hydrocarbon)}$ is unfavorable.

Electrostatic Interactions and Hydrogen Bonding. In contrast to individually weak *hydrophobic* interactions, *ionic* interactions between *charged polar* side chains within a protein are individually strong (Figure (graphic 22.33)). However, since charged polar groups readily hydrogen bond to H₂O on the exterior of a protein, these groups predominate on the exterior surface of a protein. Uncharged polar groups such as OH and C(=O)NH₂ form hydrogen bonds with each other in the interior of a protein, but they also hydrogen bond to H₂O molecules on the surface of proteins and are present in both locations.

Disulfide Bonds. The SH groups of *Cys* stabilize and determine the shape of proteins by forming disulfide bonds (S-S). Their effect on the shapes of proteins is the basis of the "**permanent wave**" process that straightens or curls hair (Figure 22.34)[next page]. Application of a reducing agent to the hair breaks S-S bonds to give individual SH side chains. With disulfide bonds broken, curlers or straighteners mechanically modify the shape of the hair. Treatment with an oxidizing agent reforms S-S bonds that hold the hair in a new shape.

Disulfide bonds periodically cleave and reform over time allowing the strands of hair to slowly resume their natural configuration.



Quaternary (4°) Structure. Most proteins have more than one polypeptide chain and the ways that these multiple chains interact with each other determine the quaternary (4°) structure of the protein. The forces that hold these individual polypeptide chains together are the same as those that determine protein 3° structure. Hemoglobin (Figure 22.32) has four peptide subunits that come together to form its 4° structure.

Denaturation. We often refer to a protein with the same shape and activity that it has in an organism as the **native state** or **native form** of the protein. **Denaturation** of a protein involves a change in the 4° , 3° , and/or 2° structure of the *native state*.

Figure 22.35



While some proteins denature and then return to their native state (**reversible denaturation**), 2°, 3°, and/or 4° structures of many proteins are sufficiently fragile that denaturation is irreversible. Biochemists studying protein structure and function try to avoid protein denaturation, except to separate individual peptide chains of a protein for content and sequence determination.

Native proteins are often just slightly more thermodynamically stable than *denatured* proteins, so denaturation frequently occurs with relative ease. Denaturating agents, or conditions, include (a) temperature increases, (b) pH changes, (c) detergents, (d) water soluble

organic compounds such as alcohols and urea, and (e) ions such as ClO₄⁻, SCN⁻, Ca⁺², Ba⁺², and guanidinium $(C(NH_2)_3^+)$.

A temperature increase provides energy to overcome attractive forces between R groups, while a change in pH alters the charge on acidic or basic side chains (next section) and the attractive interactions between R groups. The chemical agents provide *intermolecular* competition with the *intramolecular* attractive forces between R groups in the native form.

22.3 Properties of *α*-Amino Acids

We have learned the structures of the 20 "standard" amino acids that organisms use to make peptides and proteins, and have just seen how these R groups influence the structures of peptides and proteins. In this section we examine the properties of α -amino acids as acids and bases as well as their synthetic and biosynthetic origins.

α-Amino Acids Are Polyprotic Acids (22.3A)

The "standard" α -amino acids have fully protonated forms that are diprotic ("H₂A") or triprotic acids ("H₃A").

The Standard Diprotic α -Amino Acids. Most "standard" α -amino acids when fully protonated are diprotic acids that we can represent by the general formula (" H_2A "). Their "H₂A" forms are positively charged, and have the specific formula H_2A^{+1} .

$$\begin{array}{cccc} -\mathrm{H}^{+} & -\mathrm{H}^{+} \\ \mathrm{H}_{2}\mathrm{A}^{+1} \xrightarrow{\rightarrow} & \mathrm{H}\mathrm{A}^{0} \xrightarrow{\rightarrow} & \mathrm{A}^{-1} \\ \leftarrow & \leftarrow \\ +\mathrm{H}^{+} & +\mathrm{H}^{+} \end{array}$$

 H_2A^{+1} is in equilibrium with the uncharged monoprotonated form HA^0 and it is in equilibrium with the unprotonated form A^{-1} . An example is *glycine* (Figure 22.37) where H₂A⁺¹ is ⁺H₃N-CH₂-CO₂H, and A⁻¹ is H₂N-CH₂-CO₂⁻ (NH₃⁺ and CO₂H have each lost a proton).

Figure 22.37 [there is no Figure 22.36]



While these are the only possible structures for H_2A^{+1} and A^{-1} , HA^0 could conceivably be either the *zwitterion* form ⁺H₃N-CH₂-CO₂⁻, or the *uncharged* form H₂N-CH₂-CO₂H.

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As we stated at the beginning of the chapter, it turns out that the structure of HA^0 is the dipolar *zwitterion* form. This seems reasonable since we would expect the *basic* NH₂ group and *acidic* CO₂H group in the *uncharged* structure to react with each other to form the NH₃⁺ and CO₂⁻ groups of the *zwitterion* form.

Figure 22.38

Uncharged HN-CH-Cat -> HAT-CH- core Ewitherion

But we can also support this with the more detailed analysis that follows.

Justification of the Zwitterion Structure of HA^0 . Quantitative support for the zwitterion structure of HA^0 comes from the respective pK_a values for the equilibria between H_2A^{+1} and HA^0 , and between HA^0 and A^{-1} (Figure 22.39).

Figure 22.39



If HA⁰ is the *zwitterion*, $pK_{a1} = 2.3$ is that for ionization of CO₂H in H₂A⁺¹. However, if HA⁰ is the *uncharged* form, $pK_{a1} = 2.3$ is that for ionization of the NH₃⁺ group in H₂A⁺¹.

Simple carboxylic acids (RCH₂CO₂H) have pK_a 's ≈ 5 , while simple aminium ions (RCH₂NH₃⁺) have pK_a 's between 10 and 11. As a result, we can see that $pK_{a1} = 2.3$ is much closer to that for CO₂H ionization ($pK_a \approx 5$) than that for NH₃⁺ ionization ($pK_a \approx 10$ to 11). This also means that $pK_{a2} = 9.6$ corresponds to ionization of NH₃⁺ in the zwitterion form of HA⁰ consistent with the $pK_a \approx 10$ to 11 for ionization of NH₃⁺ in a simple aminium ion (RCH₂NH₃⁺).

The lower pK_a for ionization of CO₂H in ⁺H₃N-CH₂-CO₂H, compared to ionization of CO₂H in RCH₂CO₂H, results from the *electron withdrawing inductive effect* of the positively charged NH₃⁺ group (Chapter 14) that stabilizes ⁺H₃N-CH₂-CO₂⁻.

Figure 22.40a

e wethrawly HAV-CH2-CO2H = R-CH2CO2 plass undertwee ______HAV-CH2-CO2H = R-CH2-CO2 plass undertwee ______ undertwee ______ effective CRO

This effect causes the energy difference between $^{+}H_3N-CH_2-CO_2H$ and $^{+}H_3N-CH_2-CO_2^{-}$ to be less than that between RCH₂CO₂H and RCH₂CO₂⁻ resulting in a lower pK_a for the amino acid.

Figure 22.40b



Besides *Gly*, there are 12 other diprotic "standard" amino acids (Table 22.1)[next page]. Their values of pK_{a1} (C_{α} -CO₂H) and of pK_{a2} (C_{α} -NH₃⁺) are remarkably similar to those of *Gly* (an exception is pK_{a2} for *Pro* that is structurally different from the others). The average value of pK_{a1} is about 2.2 while that of pK_{a2} (excluding *proline*) is about 9.3. We discuss the pI values later in this section.

Diprotic Amino Acid Forms at Different pH Values. The pK_a values for the H₂A⁺¹,

HA⁰, and A⁻¹ forms of the 13 diprotic amino acids allow us to predict the pH dependence of their concentrations.

Figure 22.41



(1) H_2A^{+1} (⁺ H_3N -CHR-CO₂H) is the major form in solution at pH values smaller than pK_{a1} (below approximately 2.2).

(2) The concentrations of H_2A^{+1} [+H₃N-CHR-CO₂H] and HA^0 [+H₃N-CHR-CO₂-] are equal when the solution pH is equal to pK_{a1} (at approximately 2.2).

(3) HA^0 (+H₃N-CHR-CO₂-) is the major form when the solution pH is between pK_{a1} and pK_{a2} (between approximately 2.2 and 9.3)

(4) The concentrations of HA⁰ [⁺H₃N-CHR-CO₂⁻] and A⁻¹ [H₂N-CHR-CO₂⁻] are equal when the solution pH is equal to pK_{a2} (approximately 9.3).

(5) A^{-1} (H₂N-CHR-CO₂⁻) is the major form when the solution pH is greater than pK_{a2} (above approximately 9.3).

Name	pK_{a1} (C _{α} -CO ₂ H)	pK_{a2} (C _{α} -NH ₃ ⁺)	pI
Nonpolar R			
alanine	2.3	9.7	6.0
glycine	2.3	9.6	6.0
isoleucine	2.4	9.6	6.0
leucine	2.4	9.6	6.0
methionine	2.3	9.2	5.7
phenylalanine	1.8	9.1	5.5
proline	2.0	10.6	6.3
tryptophan	2.8	9.4	5.9(?)
valine	2.3	9.6	6.0
Uncharged Polar R			
asparagine	2.0	8.8	5.4
glutamine	2.2	9.1	5.7
serine	2.2	9.2	5.7
threonine	2.1	9.1	5.6
Average (*w/o proline)	(2.2)	(9.3)*	(5.8)*

Table 22.1. Acid Dissociation Constants for Diprotic Amino Acids

The Standard Triprotic α -*Amino Acids.* The remaining 7 "standard" amino acids (Table 22.2) are *triprotic* acids ("H₃A") because their R's have acidic or basic functional groups in addition to the C_{α}-CO₂H and C_{α}-NH₃⁺ groups. [Table 22.2] The charges on these forms depend on the specific R group.

"H3A"	-H ⁺ →	"H2A"	-H ⁺ →	"HA"	-H ⁺ →	"A"
	←		←		←	
	$+H^+$		$+H^+$		$+H^+$	

Table 22.2. Acid Dissociation Constants for Triprotic Amino Acids					
Name	pK_{a1} (C _{α} -CO ₂ H)	$pK_{a2} (C_{\alpha}-NH3^+)$	рК _R	pI	
Charged Polar R					
aspartic acid	1.9	9.6	3.7	2.8	
glutamic acid	2.2	9.7	4.3	3.2	
arginine	2.2	9.0	12.5	10.8	
histidine	1.8	9.2	6.0	7.6	
lysine	2.2	9.0	10.5	9.7	
Uncharged Polar R					
cysteine	2.0	10.3	8.2	5.1	
tyrosine	2.2	9.1	10.1	5.7	
Överall Average	(2.1)	(9.4)			

We continue to use pK_{a1} and pK_{a2} for ionization of C_{α} -CO₂H and C_{α} -NH₃⁺, and introduce **pK**_{aR} as the acid dissociation constant of the acid-base group in R. In spite of major differences in R, the average values of pK_{a1} (2.1) and pK_{a2} (9.4) for these 7 triprotic amino acids are very similar to those of the diprotic amino acids (Tables 22.1 and 22.2). The pK_{a1} (C_{α}-CO₂H) is always the lowest acid dissociation constant for all the standard amino acids

(Tables 22.1 and 22.2), but pK_{aR} (C_{α} -R) is smaller or larger than pK_{a2} (C_{α} -NH₃⁺) (Table 22.2) depending on the structure of the acidic R group. We will discuss pI values later.

Once we specify R for a triprotic amino acid, we can write unique structures and electrical charges for "H₃A" and "A". However we also need to know relative values of pK_{aR} and pK_{a2} in order to write structures and electrical charges for "H₂A" and "HA". Because their structures and charges depend on R, we will examine the triprotic amino acids in three separate categories based on their R groups.

Aspartic and Glutamic Acid. Glu and Asp each have an acidic CO₂H group in R. Since they differ only in the number of CH₂ groups connecting that CO₂H to C_{α}, their acid dissociation equilibria and associated pK_a values are very similar. We show here all of the different protonated forms for aspartic acid, but just the triprotonated form of glutamic acid. It's remaining forms are analogous to those shown for aspartic acid.

Figure 22.42



The pK_{aR} values (≈ 4) of these two amino acids are closer to pK_a 's of simple carboxylic acids (≈ 5) than pK_{a1} values (≈ 2) because the one or two CH₂ groups that separate the CO₂H groups in R from NH₃⁺ decrease the inductive influence of the NH₃⁺ described earlier.

The relative concentrations of the four protonated forms of *Glu* or *Asp* depend on pH as we show in Figure 22.43, and their specific electrical charges are H_3A^{+1} , H_2A^0 , HA^{-1} , and A^{-2} .



At physiological pH (approximately pH 7), their major forms are HA⁻¹ where the R groups $(CH_2CO_2^- \text{ or } CH_2CO_2^-)$ have an electrical charge of -1.

Lysine, Arginine, and Histidine. In contrast to *Asp* and *Glu*, the three amino acids *Lys*, *Arg*, and *His* all have *basic* functional groups in R. That group in *Lys* is NH₂, *Arg* has a **guanidino** group (NHC(=NH)NH₂), while *His* has an **imidazole** ring.

Figure 22.44



These groups are all positively charged when they are protonated so the four forms of these amino acids are H_3A^{+2} , H_2A^{+1} , HA^0 , and A^{-1} .

Although each R is basic, the pK_{aR} values are significantly different (Table 22.2). The pK_a order for *Lys* and *Arg* is $pK_{a1} < pK_{a2} < pK_{aR}$, while that for *His* is $pK_{a1} < pK_{aR} < pK_{a2}$. We illustrate the relative concentrations of their four forms as a function of pH in Figure 22.45 [next page]. At approximate physiological pH 7, the electrical charge on the R groups of *Arg* and *Lys* is +1. In contrast, the R of *His* is predominantly neutral at pH 7, but a significant amount of the protonated +1 form is also present. (Remember that R in *Glu* and *Asp* is -1 at pH 7.)

Cysteine and Tyrosine. The remaining triprotic amino acids are *Cys* and *Tyr*. While their R groups are acidic, they are much weaker acids than the CO₂H groups of *Glu* and *Asp*.

Figure 22.46

$$\begin{array}{c} Cyp & H_{N} - CH_{2} - SH - \left[pkan = 8:2 \right] \\ CorH \\ Typ & H_{N} - CH - CH_{2} - OH - \left[pkan = 10.1 \right] \\ CaH \end{array}$$

The SH group of *Cys* has an approximate pK_{aR} of 8, while the phenol group of *Tyr* has an approximate pK_{aR} of 10. As a result, the pK_a order for *Cys* is $pK_{a1} < pK_{aR} < pK_{a2}$ while that for *Tyr* is $pK_{a1} < pK_{a2} < pK_{aR}$. In spite of the differences in pK_a order for these two



amino acids, the net electrical charges on the four forms of each of them are H_3A^{+1} , H_2A^0 , HA^{-1} , and A^{-2} . We show their relative concentrations as a function of pH in Figure 22.47. You can see that at pH 7 their R groups predominantly exist in their uncharged protonated forms (CH₂-SH or CH₂-Ph-OH).



Isoelectric Points (22.3B)

The **isoelectric point** (**pI**) for any amino acid is the pH value where the electrically neutral form has its highest concentration. pI values allow us to predict the behavior of amino acids in electrical fields. When we place a solution of amino acids between positive and negative electrodes, amino acids with a net (+) *charge* migrate toward the negative electrode, those with a net (-) *charge* migrate toward the positive electrode, while those that are *uncharged* do not migrate. We can adjust the direction of migration of amino acids in an electric field by raising or lowering pH with respect to their pI values in order to facilitate their separation in solution.

pI Values of Diprotic Amino Acids. We have seen that the electrically neutral form of all diprotic amino acids is HA⁰. It is the major form at pH values between pK_{a1} and pK_{a2} (see Figure 22.41) and its maximum concentration is at a pH (its pI value) midway between these two pK_a values. For example, the pI for *glycine* (R = H) is 6.0 (pI_{Gly} = (pK_{a1} + pK_{a2})_{Gly}/2 = (2.3 + 9.6)/2 = 6.0). pI's of all the other diprotic amino acids are also about 6 since their pK_{a1} and their pK_{a2} values are very similar (see Table 22.1).

pI Values of Triprotic Amino Acids. In contrast to diprotic amino acids, the electrically neutral forms and pI values of *triprotic* amino acids depend on the structure of R (Table 22.2). To a good approximation, the highest concentration of the electrically neutral form (H_nA^0) of any triprotic acid occurs at a pH midway between the two pK_a's associated with that neutral form $(pK_{a(n+1)} \text{ and } pK_{a(n)})$.

Figure 22.48

As a result, *Asp* and *Glu* have relatively low pI values of about 3 (see Figure 22.43), *Arg*, *Lys*, and *His* have relatively high pI values ranging from 8 to 11 (see Figure 22.45), and *Cys* and *Tyr* have pI values between 5 and 6 (see Figure 22.47).

pI Values of Proteins. A protein also has a pI value equal to the pH where the number of its positively charged (+) R groups (from *Arg*, *Lys*, and *His*) is exactly equal to the number of its negatively charged (-) side chains (from *Asp*, *Glu*, *Cys* and *Tyr*). In contrast to amino acids, protein pI values cover the broad range from <1 to >12 because pKa's of their R groups depend on whether R is folded inside or outside the protein and how it interacts with other R's. Proteins often have their lowest solubility at pH = pI and biochemists use this behavior to facilitate their isolation and purification.

Laboratory Synthesis of Amino Acids (22.3C)

Chemists use the following reactions to synthesize α -amino acids in the laboratory. They give racemic mixtures of the D and L enantiomers, so resolution is required to obtain individual enantiomers.

Amination of α -Bromo acids (Chapter xx). Br₂ NH₃ RCH2CO2H \rightarrow $R\underline{C}H(Br)CO_2H \rightarrow$ RCH(NH2)CO2H PBr₃ Strecker Synthesis (Chapter xx). $H_{3}O^{+}$ KC≡N $R\underline{C}H(=O)$ **→** $R\underline{C}H(NH_2)C=N \rightarrow$ RCH(NH2)CO2H NH_4^+ **Reductive Amination** (Chapter xx). NH3 R<u>C</u>(=O)CO₂H RCH(NH2)CO2H NaBH₄ Diethylacetamidomalonate Synthesis. (1) EtO⁻ CH₃C(=O)-HN-<u>C</u>H(CO₂Et)₂ CH₃C(=O)-HN-<u>C</u>(R)(CO₂Et)₂ **→** (2) R-Br $H_{3}O^{+}$ $CH_3C(=O)OH + {}^+H_3N-\underline{C}(R)(CO_2H)_2$ $CH_3C(=O)-HN-C(R)(CO_2Et)_2$ \rightarrow $^{+}\text{H}_3\underline{N}$ -<u>C</u>H(R)CO₂H + CO₂ $^{+}\text{H}_{3}\underline{\text{N}}-\underline{C}(R)(CO_{2}\text{H})_{2}$ \rightarrow

Biosynthesis of α -Amino Acids (22.3D)

Humans and other mammals biosynthesize just 10 of the "standard" amino acids in amounts necessary for biosynthesis of proteins. They are called "**non-essential**" amino acids to contrast with the 10 "**essential**" amino acids that we cannot biosynthesize and must obtain directly or indirectly from plants or other sources (Table 22.3) [next page].

Essential and	1 Iton-Essential	A MILLIO A	icius
-Essential		Essentia	ıl
Gln		Arg*	Met
Gly		His	Phe
Pro		Ile	Thr
Ser		Leu	Trp
Tyr**		Lys	Val
	-Essential Gln Gly Pro Ser Tyr**	-Essential Gln Gly Pro Ser Tyr**	-Essential Essential Essential Gln Arg* Gly His Pro Ile Ser Leu Tyr** Lys

Table 22.x. Essential and Non-Essential Amino Acids

* We biosynthesize Arg in small quantities. ** Tyr requires Phe for biosynthesis.

Non-Essential Amino Acids. Biosynthesis of *Ala*, *Asp*, *Glu*, and *Ser* occurs by NH₂ transfer to α -ketocarboxylates (RC(=O)CO₂⁻) from α -amino acids (R'CH(NH₃⁺)CO₂⁻)

R- <u>C(</u> =O)CO ₂ -	+	R'-CH(NH3 ⁺)CO2 ⁻	\rightarrow	$R-\underline{C}H(NH3^+)CO_2^-$	+	$R'-C(=O)CO_2^-$
existing		existing		new		new
ketocarboxylate		amino acid		amino acid		ketocarboxylate

already present in an organism.

Figure 22.49



These "amino transfers" are redox reactions that reduce the <u>C</u> of the <u>C</u>=O group to <u>C</u> of the new <u>C</u>H(NH₃⁺) group. *Asp*, *Glu*, and *Ser* serve as biosynthetic precursors to the other non-essential amino acids except *Tyr* which forms from *Phe* (Figure 22.50)[next page].

Essential Amino Acids. The flow charts in Figures 22.51-22.54 [next page] outline the biosynthetic origins of the essential amino acids in plants and microorganisms. The atoms in the starting materials and products have marks so that you can trace their participation in the



overall transformations. Each overall transformation we show in Figures 22.51-22.54 includes many intermediate steps that biochemistry texts describe in detail.











22.4 Enzymes and Enzyme Catalysis

Enzymes are proteins that catalyze biochemical reactions in organisms. We will first examine general aspects of enzymes and enzyme catalysis and then the specific mechanistic details of catalysis by α -chymotrypsin.

General Features (22.4A)

All enzymes and enzyme catalyzed reactions share a number of general features.

Active Sites. In enzyme catalyzed reactions, the reactant biomolecule (**substrate**) binds to a region of the enzyme called its **active site**. The *active site* is an indentation or cleft in the enzyme where R groups on amino acid residues interact with the *substrate* by noncovalent attractive forces.

Figure 22.55



These attractive forces include *hydrophobic bonding*, *hydrogen bonding*, and *electrostatic interactions*. They are the same as those we described earlier for interactions between R groups of polypeptides.

Enzyme Catalysis Mechanism. Once in the *active site*, a series of reactions transforms the *substrate* into product. These reactions may use amino acid R groups in the active site as reagents, as well as other reactants that diffuse into the active site. The general scheme involves reversible formation of an enzyme-substrate complex (ES) from enzyme (E) and substrate (S), followed by its conversion into product (P) and regeneration of the enzyme.

$$E + S \xleftarrow{k_1 k_2} E + S \xleftarrow{k_2 K_2} E + P \xrightarrow{k_1} k_2$$

The k_2 step is generally not a single reaction, but includes a number of sequential molecular transformations. We examine such a mechanism for α -chymotrypsin catalyzed hydrolysis of peptides at the end of this section.

Substrate Specificity. Enzyme catalyzed reactions have stereochemical and geometric specificity. Enzyme active sites have specific stereochemical configurations because their peptide chains contain only L-amino acids. As a result, active sites only interact with specific stereoisomers of chiral substrates, or they only catalyze stereospecific reactions on achiral substrates. As an example, the enzyme **yeast alcohol dehydrogenase** exclusively removes H_a from the CH₂ group of ethanol giving acetaldehyde containing only H_b (Figure 22.56) [next page]. Besides their *stereospecificity*, enzymes often catalyze reactions on one or

Figure 22.56



only a few specific members of a general class of compounds. This *geometric specificity* varies from enzyme to enzyme. Although *yeast alcohol dehydrogenase* slowly dehydrogenates (oxidizes) a number of simple primary alcohols to aldehydes, it overwhelmingly favors *ethanol* as its substrate. In contrast, α -chymotrypsin effectively hydrolyzes amide bonds of peptides, amide bonds of simple amides, and ester bonds.

Types of Enzymes. Enzymes often have common names with the ending *ase* added to the name of a substrate, or the name of the reaction, that they catalyze. In addition, systematic names classify them by the general type of process they catalyze. **Oxidoreductases** oxidize or reduce substrates, **transferases** catalyze functional group transfers, **hydrolases** hydrolyze functional groups, **lyases** form double bonds, **isomerases** cause isomerization reactions, and **ligases** make chemical bonds.

α-Chymotrypsin (22.4B)

 α -*Chymotrypsin* is one of several *hydrolase* enzymes (commonly called **proteases**) that catalyze hydrolysis of amide bonds of peptides to give smaller peptide fragments. It is a globular enzyme composed of 241 amino acid residues.

Figure 22.57



 α -Chymotrypsin Active Site. The active site of α -chymotrypsin contains the R groups of its amino acids His 57, Asp 102, and Ser 195. Amino acid residues in polypeptides have sequential numbers, so many other amino acids separate His 57, Asp 102, and Ser 195. In spite of this, their R groups are close neighbors in the active site. They form hydrogen bonds with each other because of the folded 3° structure of the protein (Figure 22.58) [next page].

General Hydrolysis Mechanism. The OH group of *Ser 195* adds to the C=O group of a peptide bond initiating the series of reactions that leads to hydrolysis of the peptide (amide)





bond (Figure 22.59). We represent the enzyme schematically as "E-OH" where OH is that of Ser 195 and R-C(=O)-NHR' represents the peptide. Hydrolysis cleaves the peptide into a new N-terminal peptide fragment H_2NR' and new C-terminal peptide fragment R-CO₂H.



Detailed Hydrolysis Mechanism. The detailed mechanism in Figure 22.60 [next page] shows how the other two R groups in the active site participate.

(1) The peptide (S) forms a complex (ES) with α -chymotrypsin (E) in which an amide bond is close to the OH of Ser 195.

(2) The OH of *Ser 195* attacks C=O of the amide bond giving a tetrahedral intermediate.

(3) The C-N bond of the tetrahedral intermediate breaks giving the N-terminal peptide fragment (R'NH₂) and an acylated enzyme. *His 57* activated by *Asp 102* provides acid catalysis.

(4) $R'NH_2$ diffuses from the active site and is replaced by H_2O .

(5) H₂O activated by hydrogen bonding to His 57, nucleophilically attacks C=O of the acyl-enzyme intermediate giving a new tetrahedral intermediate.

(6) The C-O-E bond of the tetrahedral intermediate breaks giving the C-terminal peptide fragment RCO_2H that diffuses out of the active site of the enzyme.



Chapter Review

Peptides

(1) Peptides contain α -amino acids (⁺H₃N-C_{α}HR-C(=O)O⁻) joined by amide bonds. (2) When R \neq H, C_{α}'s are chiral and have L-configurations. (3) There are 20 "standard" α -amino acids biosynthetically incorporated into naturally occurring polypeptides. (4) We can classify R groups of the 20 "standard" α -amino acids as "nonpolar", "uncharged polar", and "charged polar". (5) Names of amino acids have three-letter abbreviations

and one-letter designations. (6) Peptides are synthesized in the laboratory by automated peptide synthesis that starts with the *C-terminal* amino acid bound to a solid resin support and sequentially adds amino acids, using N-protection and carbonyl activation.

Protein Structure and Organization

(1) *Primary (1°)* protein structure includes amino acid *content* and *sequence*. (2) Content is determined using automated amino acid analyzers that hydrolyze proteins, chromatographically separate the individual amino acids, and provide a spectral display of their derivatives. (3) Sequence is determined using Edman degradation (N-terminal), and carboxypeptidases (C-terminal), in conjunction with peptide cleavage reactions catalyzed by endopeptidases. (4) Before determining content or sequence, disulfide bonds are cleaved by reduction or oxidation. (5) *Secondary (2°)* protein structure includes planar electron delocalized amide groups, as well as α -helices and β -pleated sheets resulting from hydrogen bonding between separated amide groups. (6) The *fibrous* and *globular tertiary (3°)* structures of proteins result from interactions of the amino acid R groups with each other and with water. (7) R group interactions include hydrophobic bonding, electrostatic interactions, hydrogen bonding, and disulfide bond formation. (8) Protein quaternary (4°) structure is the result of interactions between individual polypeptides in a protein with two or more peptide chains. (9) Native states of proteins are denatured by heat, pH changes, certain organic compounds, and ions, because they disrupt favorable interactions between R groups.

Properties of α -Amino Acids

(1) Depending on their R group, α -amino acids are diprotic or triprotic acids. (2) For the 13 *diprotic* α -amino acids, pK_{a1} for C α -NH₃⁺ \approx 2.2, while pK_{a2} for C α -CO₂H \approx 9.3. (3) For diprotic acids, H₂A⁺ predominates below pH \approx 2.2, HA predominates between pH \approx 2.2 to 9.3, while A⁻ predominates above pH \approx 9.3. (4) Values of pK_{a1} (C α -NH₃⁺) and pK_{a2} (C α -CO₂H) for triprotic amino acids are about the same as those of diprotic acids, but pK_{aR} values depend on R. (5) Fully protonated forms of the triprotic acids *Asp* and *Glu* have the formula H₃A⁺ and their R groups are (-) at physiological pH, those of *Lys*, *Arg*, and *His* are H₃A⁺² and their R groups are (+) at physiological pH, while those of *Cys* and *Tyr* are H₃A⁺ and their R groups are uncharged at physiological pH. (6) pI values for diprotic amino acids are approximately 6, those of the triprotic acids, *Asp* and *Glu* are about 3, those for *Lys*, *Arg*, and *His* are between 8 and 11, while those of *Cys* and *Tyr* are between 5 and 6. (7) Racemic mixtures of α -amino acids arise from (a) amination of α -bromocarboxylic acids, (b) the Strecker synthesis, (c) reductive amination of α -ketocarboxylic acids, and (d) the diethylacetamidomalonate synthesis. (8) Humans and other animals biosynthesize "non-essential" "standard" α -amino acids *Ala*, *Asp*, *Glu*, and *Ser*, by amino transfer reactions to α -ketocarboxylates, and these in turn serve as biosynthetic precursors for *Asp*, *Glu*, and *Ser*. (9) Plants and microorganisms, but not humans or other animals, biosynthesize "essential" "standard" α -amino acids.

Enzymes and Enzyme Catalysis

(1) Enzymes are proteins that catalyze biochemical reactions. (2) Substrates bind to the active sites of enzymes and the resulting enzyme-substrate complexes undergo reactions leading to the final product and regenerate active enzyme. (3) Enzyme-catalyzed reactions are stereochemically and geometrically specific. (4) Enzymes are classified as *oxidoreductases, transferases, hydrolases, lyases, isomerases,* and *ligases.* (5) α -chymotrypsin is a hydrolase for peptide and amide bonds. (6) In the active site of α -chymotrypsin, *Ser 195* nucleophilically attacks the C=O of an amide bond leading to formation of a tetrahedral intermediate, C-N cleavage is catalyzed by *His 57* giving an N-terminal peptide fragment, H₂O attacks the acyl-enzyme, and the resulting tetrahedral intermediate decomposes to give a C-terminal peptide fragment and active enzyme.