

**Course Title:**

**Chemistry of Amino Acids ,**  
**Protein and Lipids**

الكلية : العلوم

الشعبة : Biochemisty

الفرقة : الثانية

اعداد:

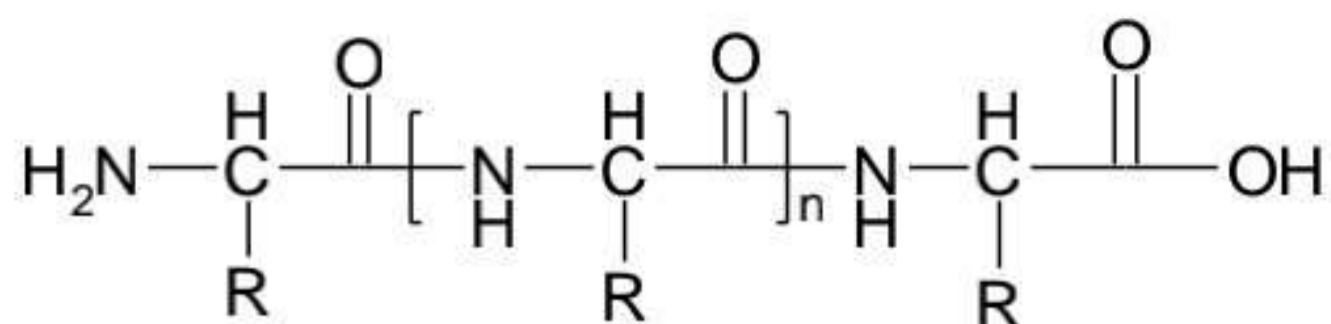
**Ass. Prof. Dr.Hussien Temerik**

## Amino Acids, Peptides, Proteins, and Nucleic Acids

### 5.1 INTRODUCTION:

Three types of organic polymers are essential to the life processes of every living cell. One of these, the polysaccharides, was discussed in Chapter 3. In this chapter we shall discuss the remaining two types. These are the nucleic acids and the proteins. The nucleic acids may be simply viewed as templates, from which the proteins are constructed. We shall return to them in Section 5.10. The proteins are polymers of amino acids. These ubiquitous macromolecules constitute nearly three-fourths of the dry weight of most animal tissues. and, indeed, they are involved in the structure and function of every living organism. Some proteins have a purely structural function (skin, hair, and muscle fiber, for example). Many others have a catalytic function (enzymes), which permits reactions to take place in living systems that would proceed so slowly in the absence of enzymes that the life could not be maintained. Other proteins have a regulatory function (hormones), and still others participate in the immunological defense mechanism of the organism (antibodies). In a human there are estimated to be about 5 million different proteins present, each of which is performing a function necessary for the well-being of the human. Other species of higher animals have similar numbers of proteins, and most of these proteins differ from one species to another. Some of them even differ from one individual to another.

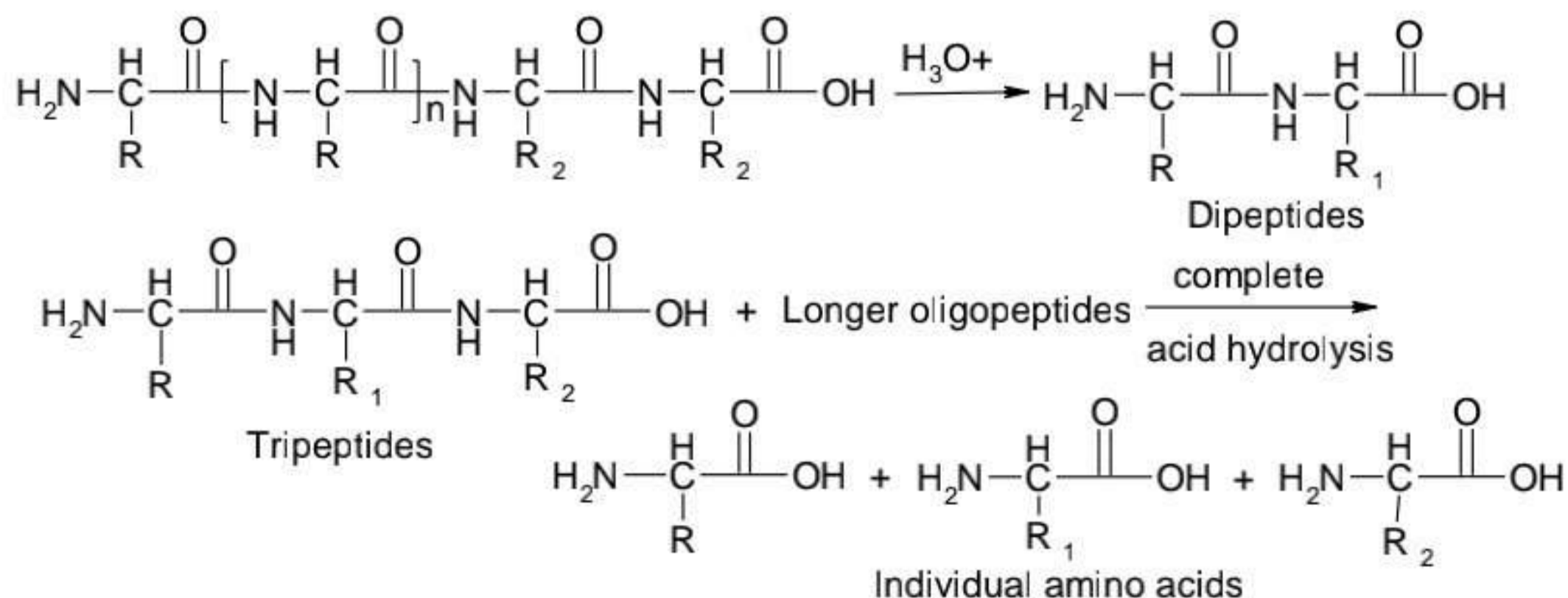
Proteins are composed of  $\alpha$ -amino acids joined together through amide linkages called peptide bonds:



Partial hydrolysis of proteins by acids, bases, or enzymes yields smaller polyamides. Complete hydrolysis can be accomplished to give the individual amino acid components.

The molecular weights of proteins range from 6000 for insulin to 41,000,000 for the protein portion of tobacco mosaic virus. Polyamides of molecular weight less than 5000 are usually termed polypeptides. The large proteins are highly organized complexes of smaller subunits. In the case of tobacco mosaic virus protein, many identical subunits, each with a molecular weight of 17,500, are associated by noncovalent interactions. It seems likely that very few proteins of molecular weight greater than 100,000 will be found to consist of only one continuous polypeptide backbone.

Natural polypeptides achieve their ability to carry out biological functions by virtue of their specific sequences of amino acids and their exact three-dimensional arrangement of these amino acids. The first step in the study of a protein is determination



of the amine acid science, called the primary structure. With the advent of more sophisticated techniques increasingly detailed aspects of protein structure are being investigated. These include the nature of the spatial relationship of near neighbors, sometimes called the secondary nurture; the gross folding of one chain, tertiary structure and the spinal relationship of one polypeptide chain to another quaternary structure.

## **5.2 NATURALLY OCCURRING AMINO ACIDS:**

From all natural sources over 100 amino acids have been isolated and identified to date. The great majority of amino acids have the amino group attached to the carbon  $\alpha$  to the carboxylic acid. With very few exceptions, the  $\alpha$  carbon also bears a hydrogen atom. The fourth bond of the  $\alpha$  carbon is joined to a group which has over 100 variations. Thus, most of the naturally occurring amino acids differ only in the structure of the organic residue attached to the  $\alpha$ -carbon. An interesting and important fact is that almost all amino acids isolated from proteins have the L-configuration at the  $\alpha$ -carbon, although some amino acids isolated from microbiological sources are the mirror image isomers i.e., in the D-configuration (see p.5).



L-Configuration

Of the amino acids isolated from living material, only about 20 are naturally occurring components of proteins. The remainder are found as intermediates or end products of metabolism. All living species are able

to synthesize amino acids. Many species, however, are deficient in their ability to synthesize within their own metabolic system all the amino acids necessary for the life of their species. The eight amino acids with this special significance for the human species are called essential amino acids (Table 5.1). They are essential not because they are the only amino acids required for human functioning but because they are essential in the diet of the human species

Table 5.1: Essential Amino Acids:


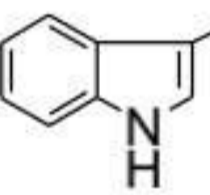
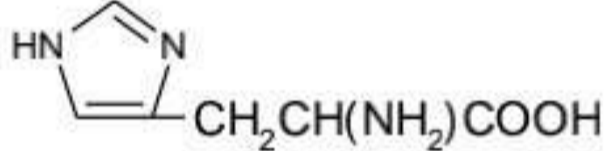
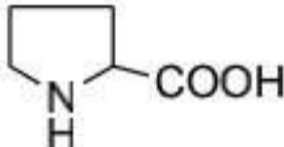
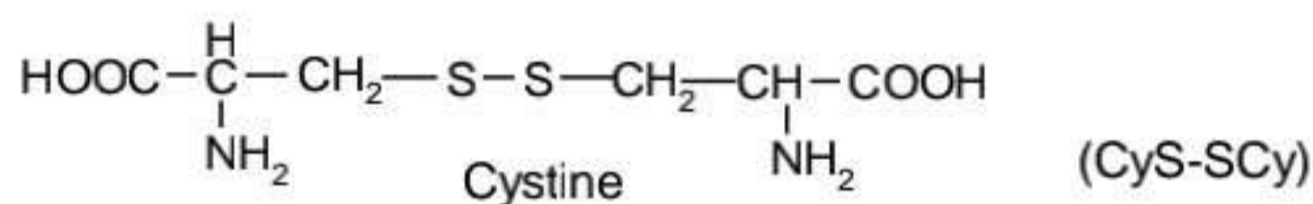
Structure	Name	Abbreviation
$\begin{array}{c} \text{CH}_3\text{CHCH}(\text{NH}_2)\text{COOH} \\   \\ \text{CH}_3 \end{array}$	L-(+)- Valine	Val
$(\text{CH}_3)_2\text{CHCH}_2\text{CH}(\text{NH}_2)\text{COOH}$	L-(-)- Leucine	Leu
$\begin{array}{c} \text{CH}_3\text{CH}_2\text{CHCH}(\text{NH}_2)\text{COOH} \\   \\ \text{CH}_3 \end{array}$	L-(+)- Isoleucine	Ile
$\begin{array}{c} \text{CH}_3\text{CHCH}(\text{NH}_2)\text{COOH} \\   \\ \text{OH} \end{array}$	L-(-)- Threonine	Thr
$\text{CH}_3\text{S}(\text{CH}_2)_2\text{CH}(\text{NH}_2)\text{COOH}$	L-(-)- Methionine	Met
 $\text{C}_6\text{H}_5\text{CH}_2\text{CH}(\text{NH}_2)\text{COOH}$	L-(-)- Phenylalanine	Phe
 $\text{Indole-3-CH}_2\text{CH}(\text{NH}_2)\text{COOH}$	L-(-)- Tryptophan	Trp
$\text{NH}_2(\text{CH}_2)_4\text{CH}(\text{NH}_2)\text{COOH}$	L-(+)- Lysine	Lys

Table 5.2: Other Common Amino Acids<sup>a</sup>:

Structure	Name	Abbreviation
$\text{CH}_3\text{CH}(\text{NH}_2)\text{COOH}$	alanine	Ala
$\text{HN}=\underset{\text{NH}_2}{\text{C}}\text{NH}(\text{CH}_2)_3\text{CH}(\text{NH}_2)\text{COOH}$	Arginine	Arg
$\text{HOOCCH}_2\text{CH}(\text{NH}_2)\text{COOH}$	Aspartic acid	Asp
$\text{HSCH}_2\text{CH}(\text{NH}_2)\text{COOH}$	Cysteine	CySH
$\text{HOOC}(\text{CH}_2)_2\text{CH}(\text{NH}_2)\text{COOH}$	Glutamic acid	Glu
$\text{H}_2\text{NCH}_2\text{COOH}$	Glycine	Gly
	Histidine	His
	Proline	Pro
$\text{HOCH}_2\text{CH}(\text{NH}_2)\text{COOH}$	Serine	Ser
$\text{HO}-\text{C}_6\text{H}_4-\text{CH}_2\text{CH}(\text{NH}_2)\text{COOH}$	Tyrosine	Tyr
$\text{NH}_2\text{COCH}_2\text{CH}(\text{NH}_2)\text{COOH}$	Asparagine	Asn
$\text{NH}_2\text{CO}(\text{CH}_2)_2\text{CH}(\text{NH}_2)\text{COOH}$	Glutamine	Gln

<sup>a</sup>One of the amino acids commonly found in protein hydrolysates has the name cystine, and has the following structure:



It is clearly a dimer of cysteine. Where the thiol groups of the latter have been oxidized to form a disulfide linkage. The dimer actually results because of two monomers at widely spaced intervals in the polypeptide are joined together by a disulfide bridge. Thus the basic amino acid is cysteine: consequently, the dimer is not included here.

since our cells cannot synthesize them. The other 12 amino acids (Table 5.2) found in the biochemicals derived from human beings can be synthesized in individual cells from simpler starting materials that contain

carbon, hydrogen, oxygen, and nitrogen.

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### EXERCISE 5.1

Glycylglycine is a dipeptide composed of two molecules of glycine. Write its structure.

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The various coexisting species have different sets of amino acids which they require-but are unable to synthesize. However, all creatures contain within themselves all amino acids, so that any animal may normally acquire needed amino acids from others. Man may acquire his daily minimum of essential amino acids by eating such things as filet mignon from the cow, Dover sole from the fish, and eggs Benedict from the chicken. Vegetarians can survive because there are sources of plant protein that contain all the essential amino acids. Rice has a high protein content, as do legumes (peas, beans), Corn, wheat, and rye are other grains that have a significant quantity of plant protein which includes the essential amino acids. Proteins eaten by humans (and other animals) are completely hydrolyzed to amino acids, and these are then used as building blocks to construct the proteins of the individual.

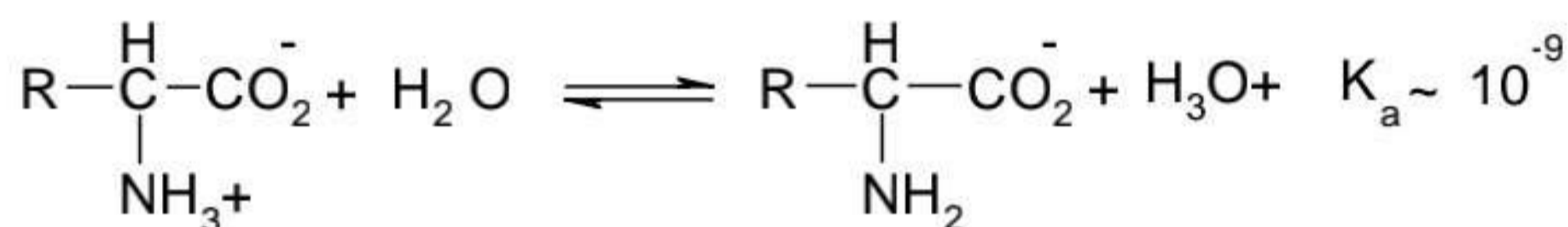
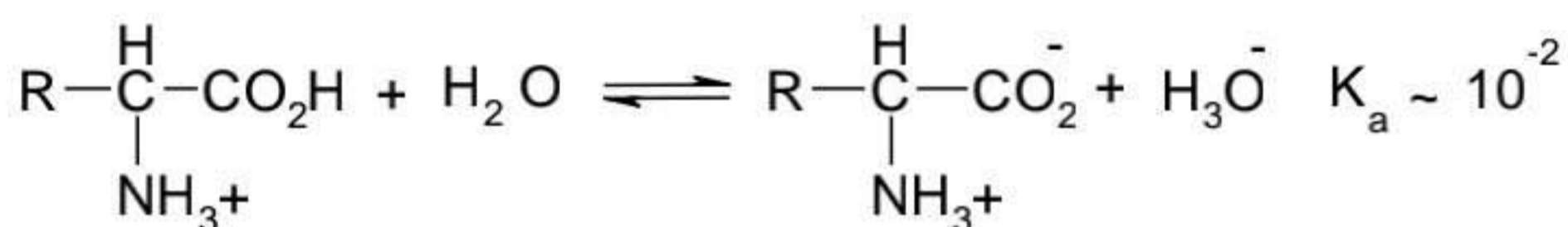
### CHEMICAL AND PHYSICAL PROPERTIES OF AMINO ACIDS:

Amino acids are high-melting solids which, because of their two polar groups, would be expected to be insoluble in organic solvents but soluble in water. Since the carboxylic acid function is acidic and the amino group basic, the amino acids actually exist as dipolar ions

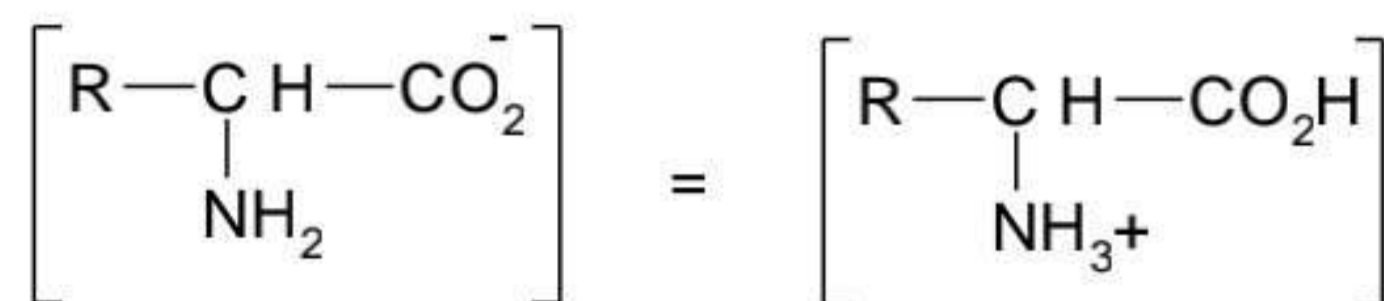
(zwitterions), rather than in the un-ionized forms shown in the previous section.



Amino acids with no ionizable side chains have two ionization constants with  $\text{pK}_a$ 's of about 2 and 9.



if an electrical potential is placed across two electrodes in a solution of an amino acid, the amino acid will migrate to the anode or the cathode, depending upon the pH. At one pH, called the isoelectric point, there is no *net* migration of the amino acid because the concentration of anion is the same as the concentration of the cation:

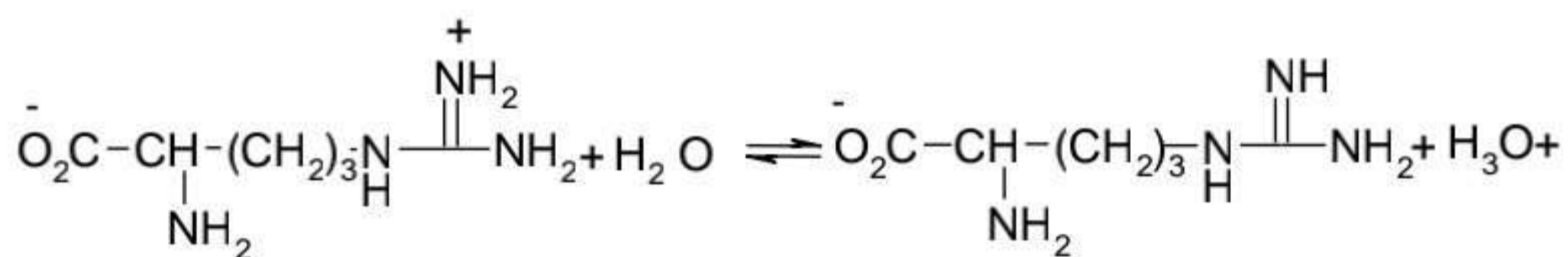
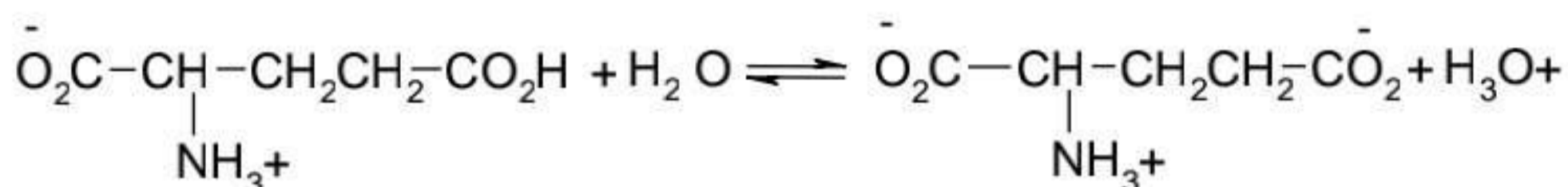


The isoelectric point is an individual characteristic of an amino acid; for example, it is pH 6.0 for glycine, pH 5.5 for phenylalanine, pH



11.2 for arginine, and pH 3.2 for

The amino acids with functional groups that are ionizable have ionization constants characteristic of those functional groups. For example, the side chain of glutamic acid has a  $pK_a$  of 4.3 and that of arginine has a  $pK_a$  of 13.2.



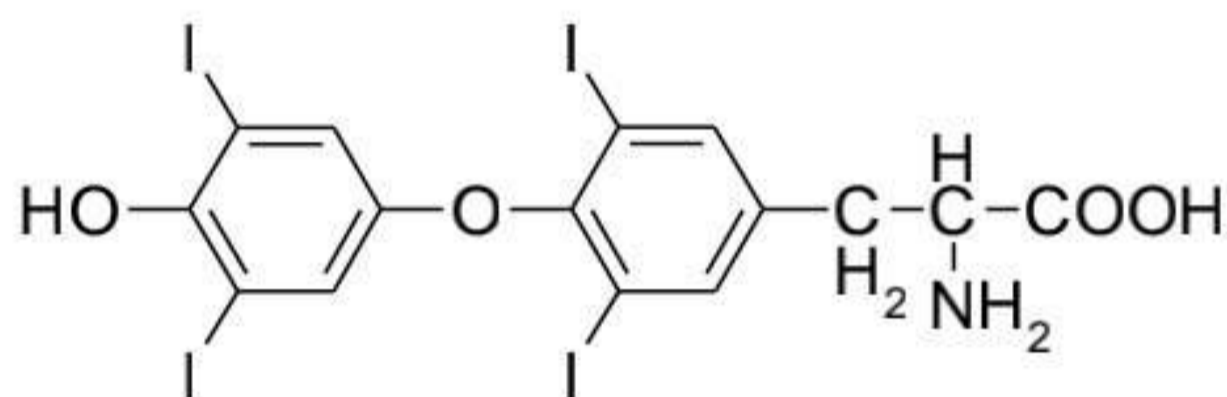
Saturated carboxylic acids absorb in the infrared at  $1725-1700\text{ cm}^{-1}$ . Amino acids, however, absorb at  $1400$  and  $1600\text{ cm}^{-1}$ , absorption frequencies characteristic of the carboxylate ion. When a neutral amino acid solution is made acidic, the  $1720\text{ cm}^{-1}$  carbonyl stretching frequency of the carboxylic acid appears. This is consistent with the proposed dipolar structures.

Amino acids undergo most of the reactions characteristic of carboxylic acids and aliphatic amines. Amino acid esters are relatively unstable, and they are usually obtained as hydrochloride salts. The amino group reacts with nitrous acid, as do other aliphatic amino groups. The accompanying evolution of nitrogen is, in fact, often used to analyze for free amino groups in amino acids and their derivatives.

### **5.4 Primary structure and biological activity of polyamides**

The number of possible random combinations of the 20 or so amino acids found in hormones, enzymes, and all other proteins is almost infinite. However, biological activity is not achieved by randomness but by a very precise ordering of the combined amino acids. Many scientists are now studying the primary structure-the amino acid sequence-of the polypeptides of biological importance for human beings. In this section we shall give some examples of the relationship between sequence and activity for polypeptides of varying chain length.

Even a single amino acid may exhibit potent biological activity: for example, thyroxine is a hormone that is an active principle for those animals which have a thyroid gland.



Thyroxine

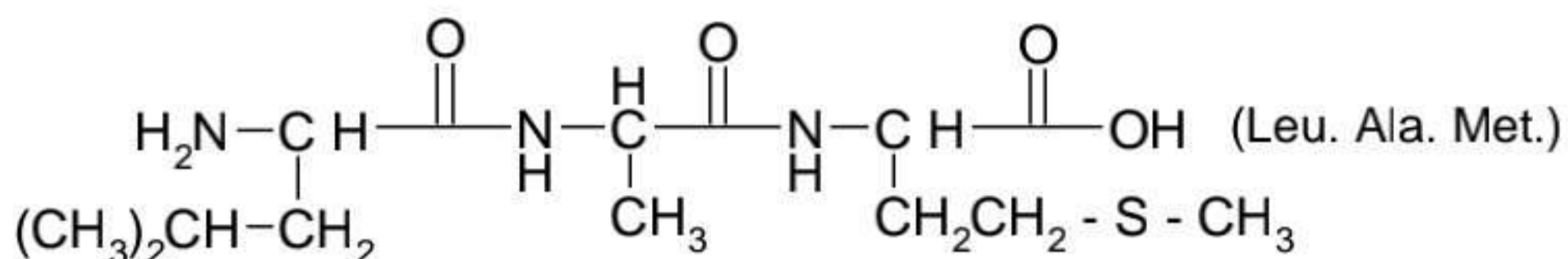
This relatively simple molecule exerts a profound effect upon the metabolism of almost every cell in the body.

### **Nomenclature:**

In naming peptides, amino or N-terminal end refers to the end with the free amino group and carboxy or C-terminal end refers to the end with the free carboxyl group.

By convention, the N-terminal end is written to the left and the C-terminal end to the right. The amino acids are then named left to right, replacing

ine with *yl*, except for the C-terminal amino acid, as in the example shown.



Leucylalanylmethionine

Sometimes a polypeptide will be found as a simple derivative, and this can also be indicated in the structure. For example, the formula Leu. Ala. Met(NH<sub>2</sub>) would represent the above polypeptide, in which the free carboxyl group had been converted into an amide. Similarly, the formula (Ac) Leu. Ala. Met would indicate the acetyl derivative of the amino end of the molecule.

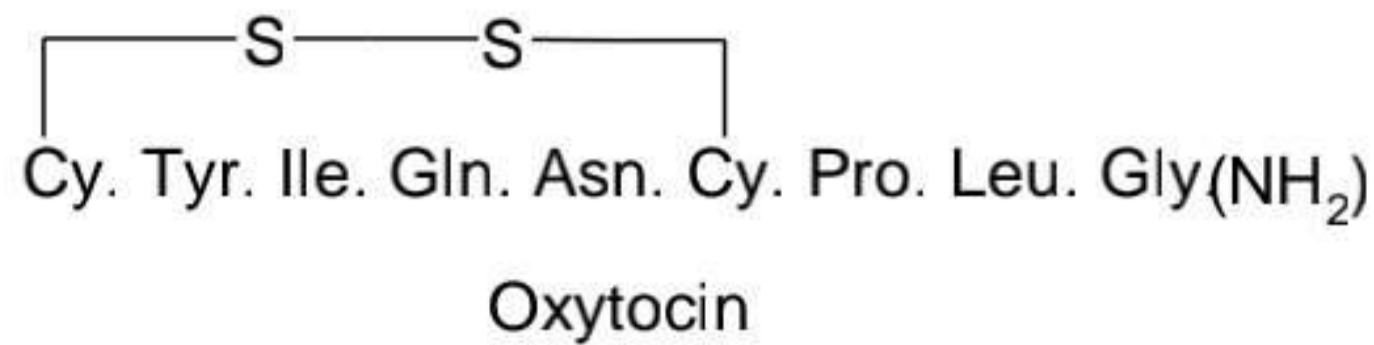
### EXERCISE 5.2:

Write out the full structures that correspond to the following formulas:

- Pro.Val. Glu(NH<sub>2</sub>)
- Arg. Gly. Phe. Ser
- Ser. Tyr. Arg. Asp

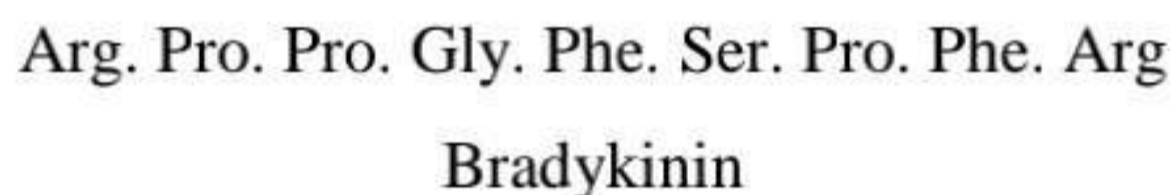
Oxytocin and bradykinin are both polypeptides composed of nine amino acids, but different acids and in different sequences. Their resulting biological functions are strikingly different.

*Oxytocin* is one of the most physiologically active compounds known.



Oxytocin is responsible for uterine contractions during childbirth and acts upon lactating mammary glands to stimulate the ejection of milk. It is interesting that only the female of the species produces this relatively simple polypeptide. It is even more interesting that this specific chemical is equally effective in causing a chicken to lay an egg or a cow to give down her milk to a farmer on a cold morning, or in causing a pregnant female human to give birth to a child. Chemically the oxytocin from chickens, cows, and hogs is identical. Oxytocin obtained from chickens is used clinically for the induction of labor.

*Bradykinin* is also a very active substance. It is released by blood plasma globulins in response to a wasp sting, and is a potent pain-causing agent.



Obviously any change in the amino acid sequence of bradykinin or oxytocin, whether it was substitution, deletion, or addition, would result in a profound modification of the biological activity.

Larger natural polypeptides may perform the same function in different species without being identical in primary structure. For example, insulin, a hormone that controls carbohydrate metabolism, differs in the arrangement of 4 of its 51 amino acids -in each of many different species. However, bovine insulin may be used to compensate for

the insulin deficiency of human beings suffering from diabetes.

Proteins of more than 100 amino acids are transferred between species with difficulty because of immunological problems (discussed in Section 5.5). For example, enzymes that perform identical functions in different species cannot simply be transferred between these species. There are differences in the amino acid sequences which result in a recognizably changed three-dimensional structure (Section 5.5). Very large proteins such as those in hair, muscle, and skin are present in such a complex arrangement that, except for identical twins, no individual of a species will accept and use the hair, muscle, or skin of another.

A more dramatic example of the importance of the amino acid sequence is provided by the polypeptide globin, the protein moiety of hemoglobin (see elsewhere for a discussion of the structure and function of hemoglobin). Globin has 146 amino acid residues in a very specific order, and for the human being, a substitution, a deletion, or an addition of even one amino acid to the number or the sequence may result in serious disease or possibly death. The disease called sickle-cell anemia is a molecular disease suffered by people whose globin differs from normal only in that the sixth amino acid in the series of 146 is valine rather than glutamic acid. Victims of this disease, which is hereditary, are unable to utilize oxygen at the normal rate and therefore must avoid high, oxygen-rare altitudes and any exercise that is physically taxing on their red blood cells. The formula below shows the substitution that distinguishes the globin of a normal human from one who has sickle-cell anemia.

**Normal Globin:**

Val. His. Leu. Thr. Pro. Glu. Glu. Lys.

**Sickle-Cell Globin:**

Val. His. Leu. Thr. Pro. Val. Glu. Lys

Under conditions of oxygen deficiency or abnormal physical activity requiring rapid oxygen metabolism, the red blood cells of people with this disease take the shape of a "sickle" or quarter-moon, and they completely cease to function unless oxygen is administered effectively and immediately.

Enzymes are even more complex polypeptide material. Every living cell contains thousands of enzymes, each of which is responsible for catalyzing a single specific chemical reaction. The complete chemical structure has been elucidated for several enzymes. Recently, one enzyme, ribonuclease, which contains 124 amino acids in a specific sequence, has been synthesized (Section 5.9). Chymouypsin has been purified to the point where it has crystallized and the precise number of amino acid residues (246) and their precise sequence have been determined. X-ray structure determination has established its three-dimensional structure at a resolution of better than 3 Å. At this resolution individual atoms cannot be discerned, but the overall shape, and the twists and turns of various segments, can be seen (Section 5.7).

**5.5 IMMUNOCHEMISTRY:**

The human body reacts immediately whenever it is subjected to the introduction of any foreign substance, including larger polypeptides. It

examines alien material very carefully for unfamiliar chemical structural characteristics and, should it recognize any, it causes the production of the specific polypeptide which is able to specifically bind to the foreign matter, so precipitating it from the surrounding medium.

In immunological terms, any such foreign substance is an antigen. The main encasing protein is gamma globulin of the blood of the host and is called an antibody. By a mechanism not completely understood, this initial antigen-antibody reaction elicits the production of greater amounts of the specific antibody required. Excess antibodies remain in the bloodstream, where they afford the body a specific type of immunity for as long as they remain in excess. Should the offending antigen return while they are present, it will immediately be precipitated and the body will suffer no harmful effects. Unfortunately, immunity is not necessarily a permanent condition; its duration may range from several hours to a lifetime, depending on the nature of the antigen. Thus immunity from smallpox is normally long term, whereas that from the common cold lasts only a matter of days or weeks.

It is important to note that immunity is highly specific for a given antigen. Each new foreign cell that invades the body elicits a new supply of specific antibodies, which are stored in the blood as gamma globulins. Quite obviously, the gamma globulin blood fractions of individuals will vary to the extent to which they have been exposed to different foreign cells.

In this antigen-antibody phenomenon, we have a more precise explanation for the failure of skin and kidney and heart transplants among individual members of the same species. The body of the receptor looks

upon the skin or kidney cells of the donor as foreign matter and immediately sets up an antibody-type rejection mechanism. Only between identical siblings, whose bodies are made up of protein materials that have identical structures and are therefore not "foreign," can such transplants be successful. One approach used today is to attempt to suppress production of gamma globulins. Drugs are known that accomplish this purpose. The patient is then able to receive a foreign substance, for example a heart, but has lost immunity and is therefore susceptible to many diseases from which he had previously been cured. Further, globulin synthesized in the body after the heart transplant still does not accept the heart as nonforeign material and finally rejects it by the antibody-antigen reaction.

## **5.6 DETERMINATION OF THE STRUCTURES OF PEPTIDES:**

The first step in determining the primary structure of a peptide is to hydrolyze it to its individual amino acids and to assess which ones are present and how many of each. Although these analyses are somewhat complicated, the details for such procedures have been very thoroughly worked out, and the analyses themselves are now highly automated.

### **OPTIONAL MATERIAL:**

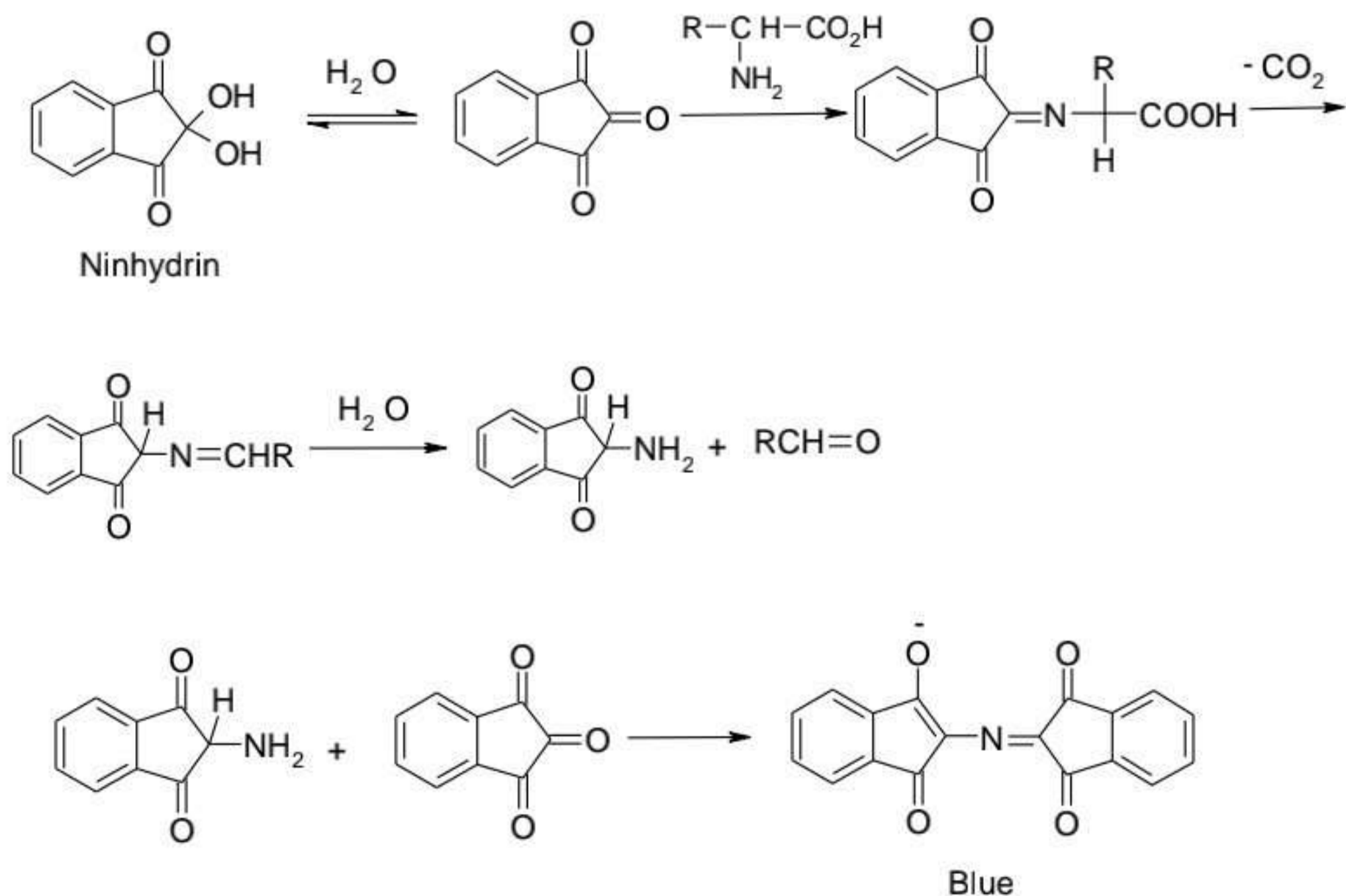
*Analysis of Amino Acids.* Peptides are hydrolyzed in 6 N HCl at 105 °C; base cannot be used because it racemizes the  $\alpha$  carbons. Tryptophan, is sensitive to acid and is partially destroyed in the hydrolysis, which can be corrected for in quantitative studies. Instead of



glutamine and asparagine, the corresponding acids and ammonia are isolated.

The mixture of amino acids obtained upon hydrolysis can be separated and analyzed by use of an "amino acid analyzer." In this automatic equipment, aliquots of the mixture of amino acids are placed on separate columns of a sulfonic acid ion-exchange resin. One column is held at pH 5.3 and used for basic amino acids, ammonia, and tryptophan. A second is held at pH 3.25 for the other amino acids while 0 to 250 ml of elutant pass through the column: the pH is then increased to 4.25 while 250-500 ml of elutant passes through. Sodium citrate buffer solutions elute the acids, and the eluted solutions are mixed with ninhydrin and heated.

The reagent ninhydrin produces a blue color with primary  $\alpha$ -amino acids by the following series of reactions:

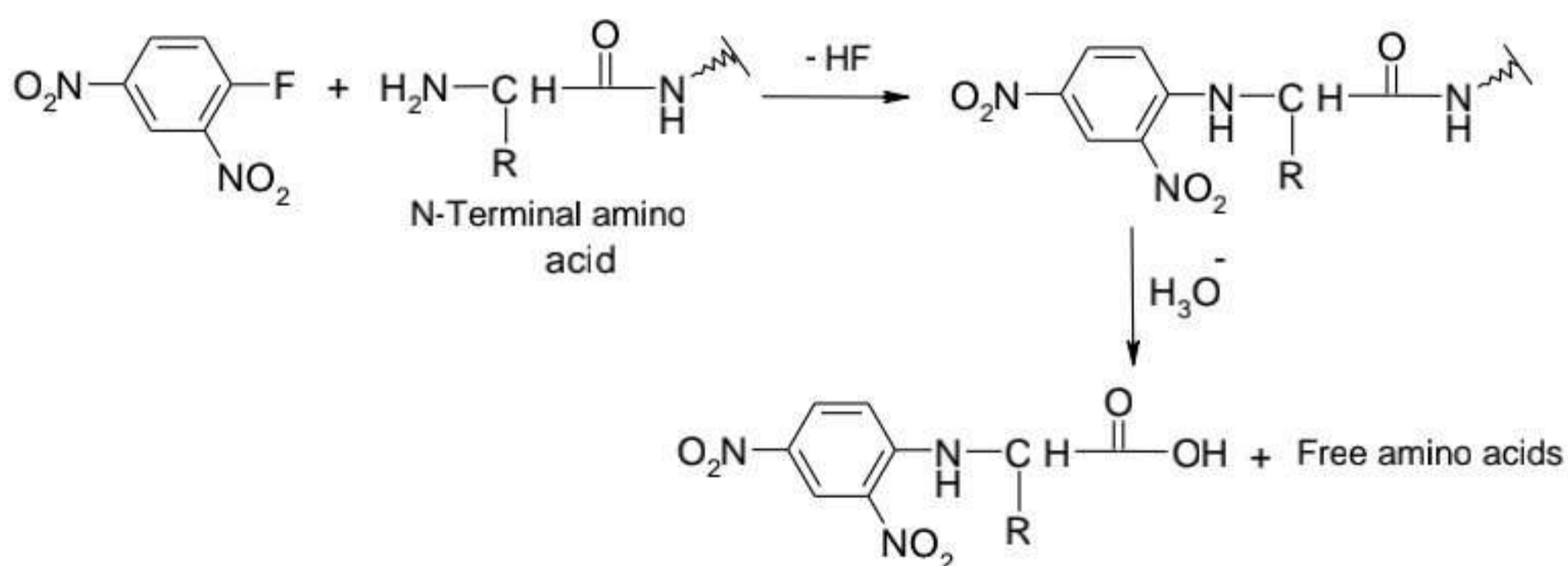


A spectrophotometer measures the optical absorption of the products of the ninhydrin reaction, and the recorder continuously plots the milliliters of eluate versus intensity of the ninhydrin color. The position of the absorption peak (in ml), which depends on the volume of buffer needed to elute a particular amino acid, is characteristic for each amino acid; the quantity of the acids is obtained from the areas under their peaks.

Gas chromatography is also useful for analyzing mixtures of small amounts of amino acids. Amino acids are too nonvolatile to be studied directly; their esters, for example trimethylsilyl esters, have sufficient volatility, however.

After determining the identities of the amino acids present, and their ratios, the next big problem is to determine the sequence of the amino acids in a peptide.

One common technique used to determine the N-terminal amino acid is to allow the amino group to react with 1-fluoro-2,4-dinitrobenzene (nucleophilic aromatic substitution) and then to hydrolyze the peptide:

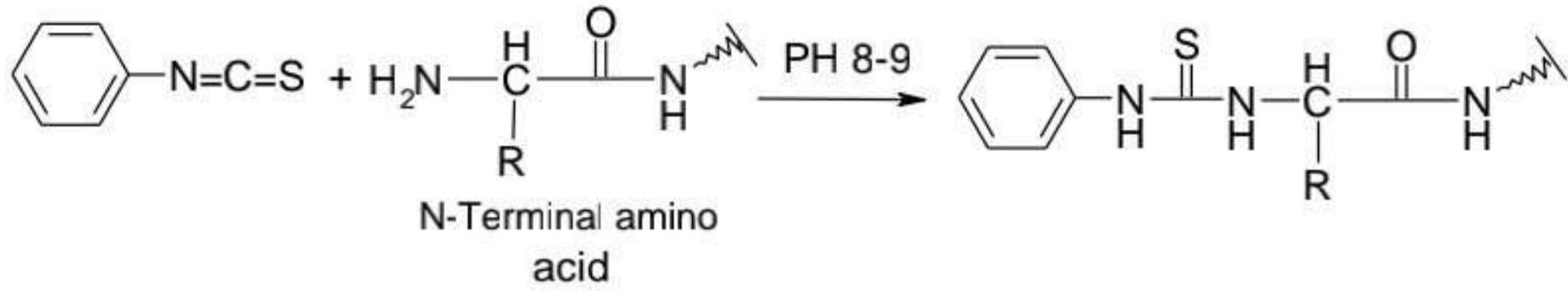


The N-(2,4-dinitrophenyl) derivative is isolated and identified.

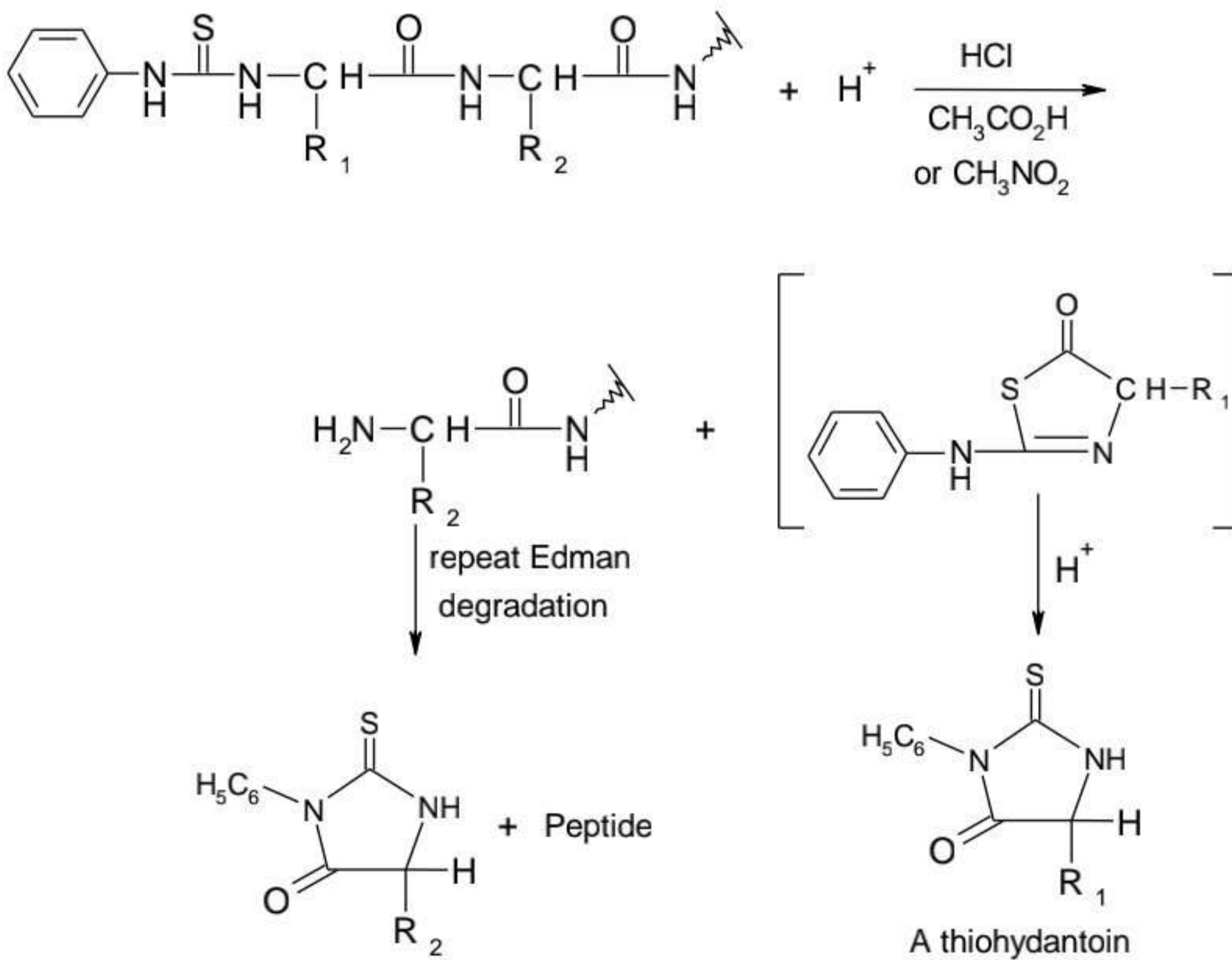
An alternative procedure for determination of the N-terminal amino acid, which does not hydrolyze the peptide, is called Edman degradation.

Chapter 5 Amino Acids, Peptides, Proteins, and Nucleic Acids

An N-phenylthiocarbamyl derivative is prepared with phenyl isothiocyanate.

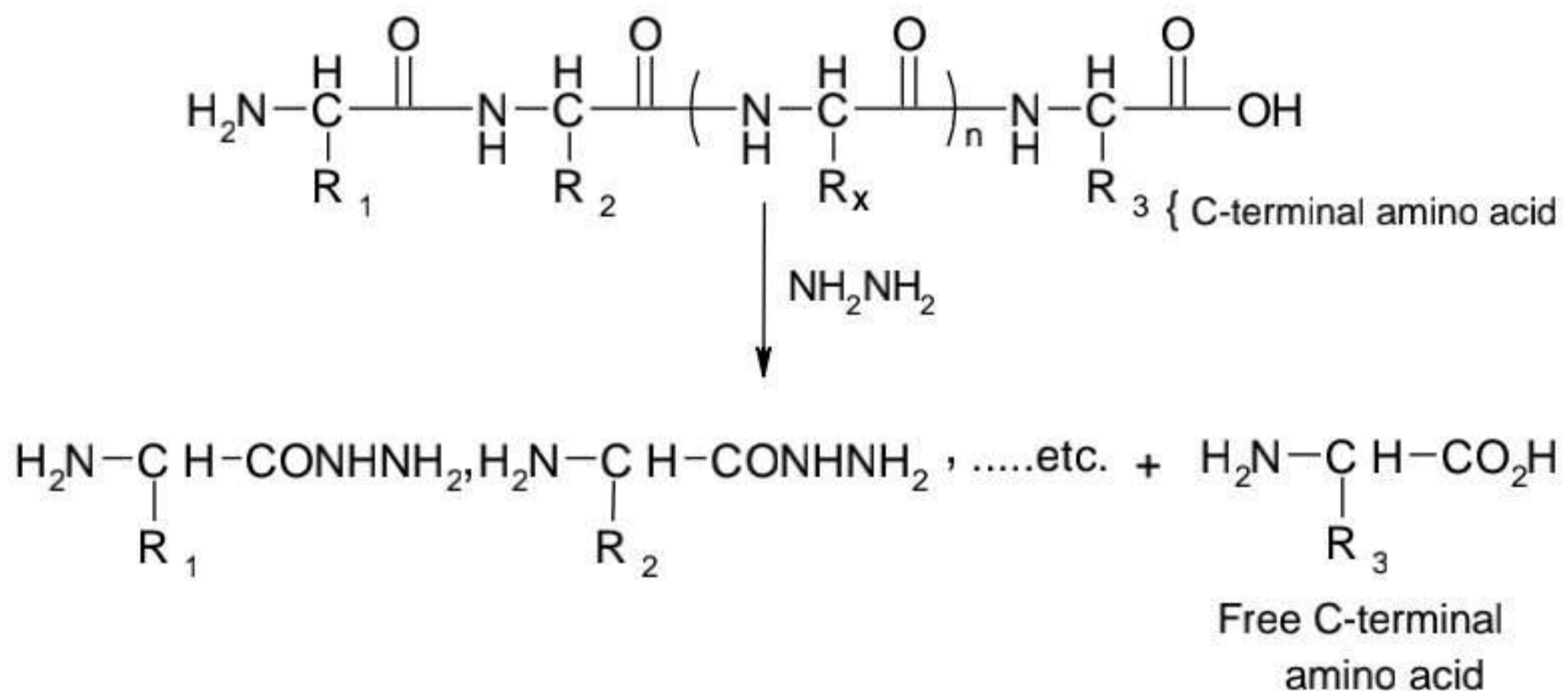


When this derivative is treated with hydrogen chloride in nitromethane or acetic acid, a thiohydantoin forms without destroying the remaining linkages.



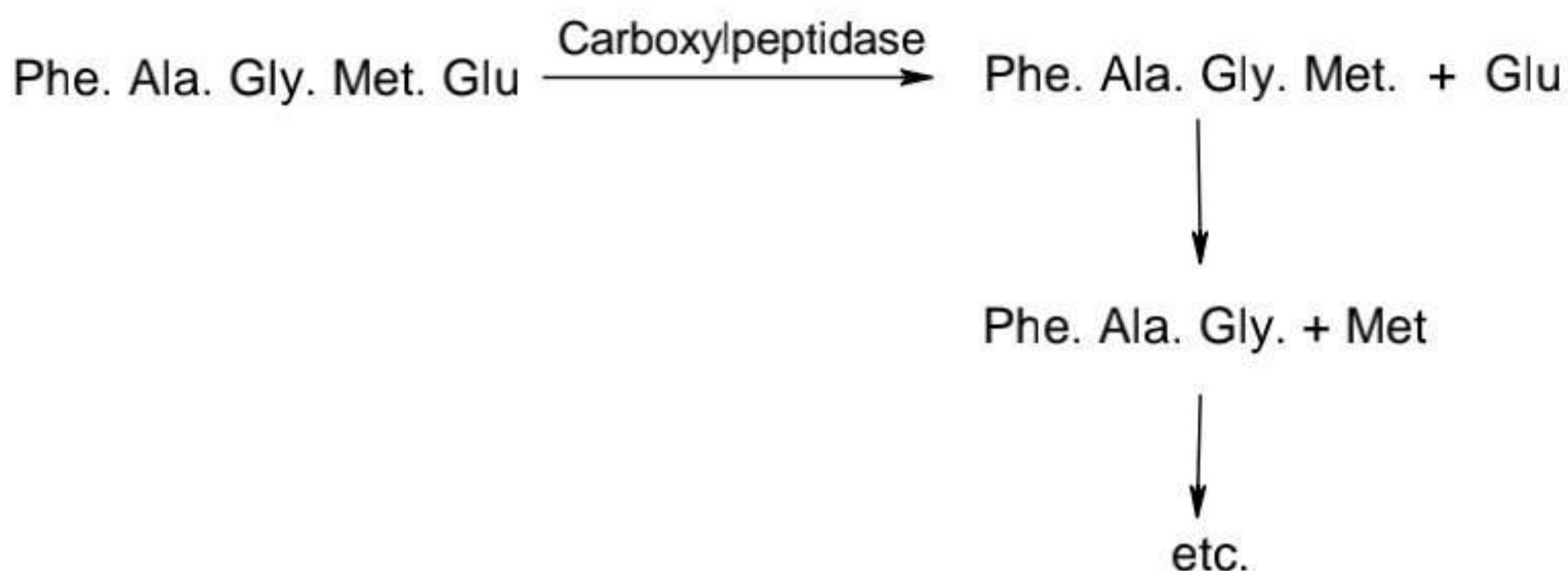
The thiohydantoin is identified and the nature of its R group characterizes the N-terminal amino acid. This procedure can be continued on the remaining chain; there is now available commercially an automated procedure which the manufacturer claims will perform 30 successive Edman degradations on a purified protein.

To determine the C-terminal end, the peptide can be heated with anhydrous hydrazine to convert the amide linkages in the chain into hydrazides:



The C-terminal amino acid is identified as the free acid, whereas the others in the chain are obtained as hydrazides.

Certain carboxypeptidases, which are enzymes obtained from the pancreas, attack C-terminal peptide bonds and free the C-terminal amino acid. However, they continue acting on the remaining peptide and systematically liberate the newly formed C-terminal acids; for example,



The action of the enzyme cannot be stopped after it has released Gln. The enzyme proceeds to attack the tetrapeptide and release Met, and then to attack the tripeptide. and so on. Thus, the sequence of only a limited number of units can be obtained before the situation gets too mixed up to sort out. The identities of the amino acids that are freed are determined as a function of incubation time, and as much sequencing is done as is possible from the data. If the C-terminal end is in the form of the amide, carboxy-peptidase does not act to free it.

To illustrate some of the techniques employed in amino acid sequence determination, we will follow an example of such a determination. Using an amino acid analyzer and a crude molecular-weight determination,  $\alpha$ -MSH, a melanocyte-stimulating hormone from pituitary glands, was found to have the molecular formula: (Arg, Glu, Gly, His, Lys, Met, Phe, Pro, Ser<sub>2</sub>, Trp, Tyr, Val, NH<sub>3</sub>) The commas between the abbreviations indicate that the sequence is unknown or unspecified. The acids were present in equimolar quantities, except for serine.

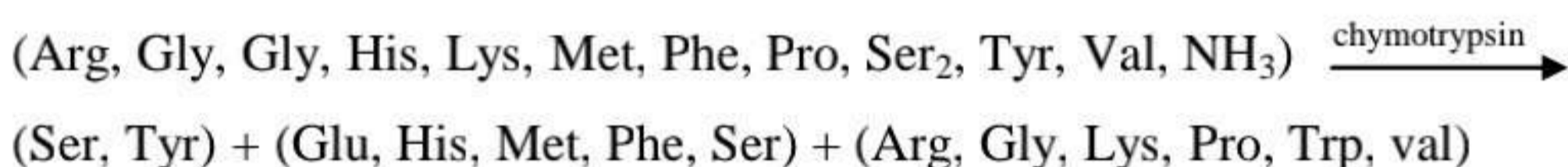
No N-terminal amino acid was found in the Edman degradation or dinitrophenylation reaction and no C-terminal amino acid was liberated

by carboxypeptidase. As we will see later, this is due to tie-up of the N-terminal end as the N-acetyl derivative and the C-terminal as the amide:

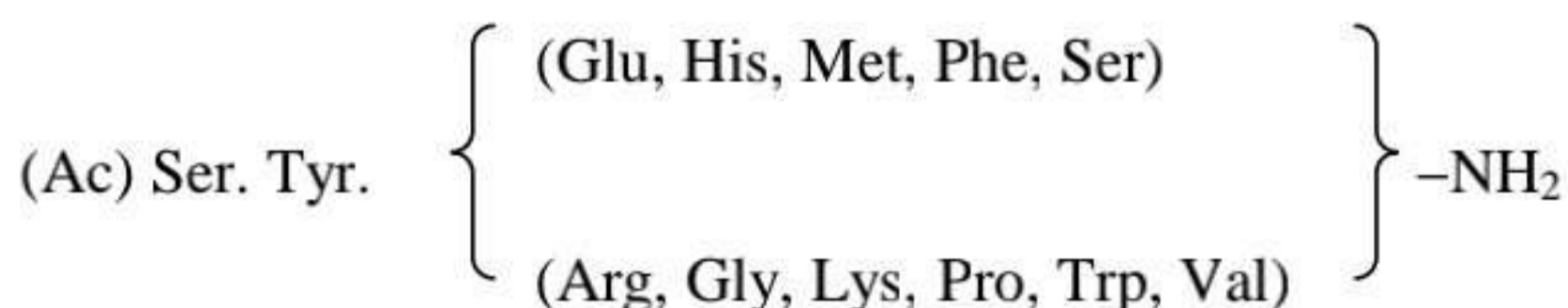


It could also have meant that the peptide chain was cyclic. Thus the amino group and the carboxyl group must be free if these procedures are to work.

Chymotrypsin, another pancreatic enzyme, preferentially attacks peptide bonds whose carbonyl function is furnished by one of the aromatic amino acids tyrosine, tryptophan, and phenylalanine. although it will also catalyze the hydrolysis of bonds with leucine, methionine, asparagine, and glutamine. After chymotryptic hydrolysis of  $\alpha$ -MSH, three peptide fragments were isolated:

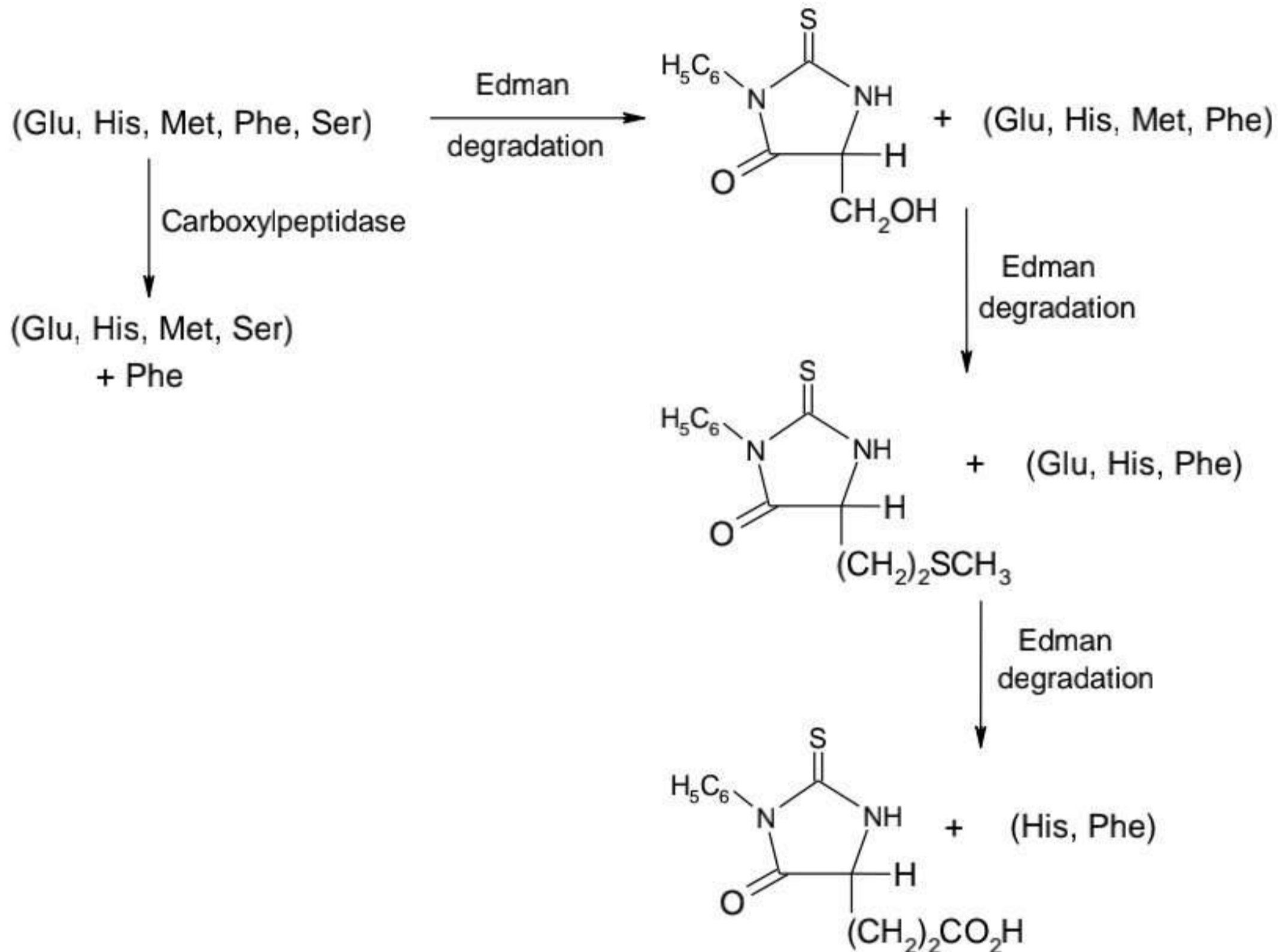


Analysis of the serine in the dipeptide (Ser.Tyr) showed it to be in the form of the N-acetyl derivative. Thus Ser. Tyr is the N-terminal end of a  $\alpha$ -MSH, and a partial structure of  $\alpha$ -MSH is



The pentapeptide fragment gave phenylalanine as the C-terminal

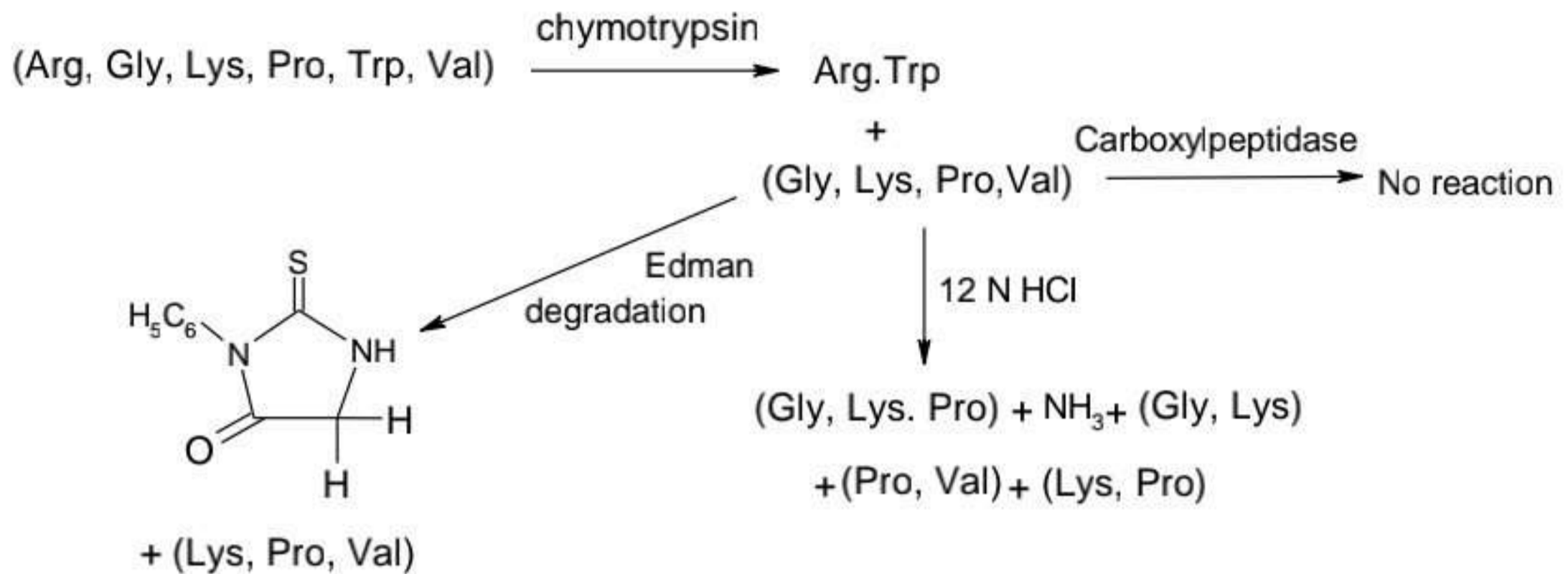
amino acid upon carboxypeptidase action, leading to the partial structure (Glu, His, Met, Ser). Phe. From an Edman degradation, it was found that serine is N-terminal: Ser. (Glu, His, Met). Phe. Two more Edman degradations gave, first, methionine, and second, glutamic acid:



Thus the pentapeptide is Ser. Met. Glu. His. Phe. Histidine is placed fourth in line because that is the only slot left.

The hexapeptide (Arg, Gly, Lys, Pro, Trp, Val) was inert to carboxypeptidase, and, therefore, represents the C-terminal end of  $\alpha$ -MSH. Further chymotryptic hydrolysis of this hexapeptide gave (Arg, Trp) and (Gly, Lys, Pro, Val). The dipeptide must be Arg. Trp because

chymotrypsin attacks the carbonyl function of tryptophan and not of arginine:



The tetrapeptide (Gly, Lys, Pro, Val) was subjected to the Edman degradation, and glycine was found to be the N-terminal amino acid. It was partially hydrolyzed in 12 N HCl at 37 °C for 120 hours. The following peptides were obtained, along with ammonia: (Gly, Lys, Pro) + (Gly, Lys) + (Pro, Val) + (Lys, Pro) + NH<sub>3</sub>. The ammonia is formed because the C-terminal end is in the amide form. Since the Edman reaction established Gly as the amino terminal residue of this peptide, the isolation of the dipeptides (Gly, Lys), (Lys, Pro), and (Pro, Val) establishes the obligatory sequence Gly Lys. Pro' Val(NH<sub>2</sub>). The presence of tripeptide containing Gly, Lys, and Pro adds strength to the sequence assignment. The complete hexapeptide must, therefore, be Arg. Trp. Gly. Lys. Pro. Val(NH<sub>2</sub>).

At this point, the information known about  $\alpha$ -MSH is

(Ac)Ser. Tyr + Ser. Met. Glu. His. Phe + Arg. Trp. Gly. Lys. Pro. Val (NH<sub>2</sub>)



If Tyr and Ser, and Phe and Arg, which were cleaved by chymotrypsin, are joined. there is only one way to put the sequence together:

This sequence is consistent with all data concerning the primary structure of  $\alpha$ -MSH. A method for determination of the primary structures of proteins which is currently under investigation involves the use of a mass spectrometer to fragment the molecule and a computer to sort out and interpret the results.

---

### EXERCISE 5.3

A hexapeptide gave upon hydrolysis the following amino acids: Glu, Gly, Glv, Lvs, Ser, Tvr. When a partial hydrolysis was carried out, there were isolated two dipeptides and a tripeptide that had the following structures: Gly. Ser, Tyr. Lys and Gly. Glu. Gly. It was found that tyrosine occupied the N-terminal position. by an appropriate test. What is the structure of the hexapeptide?

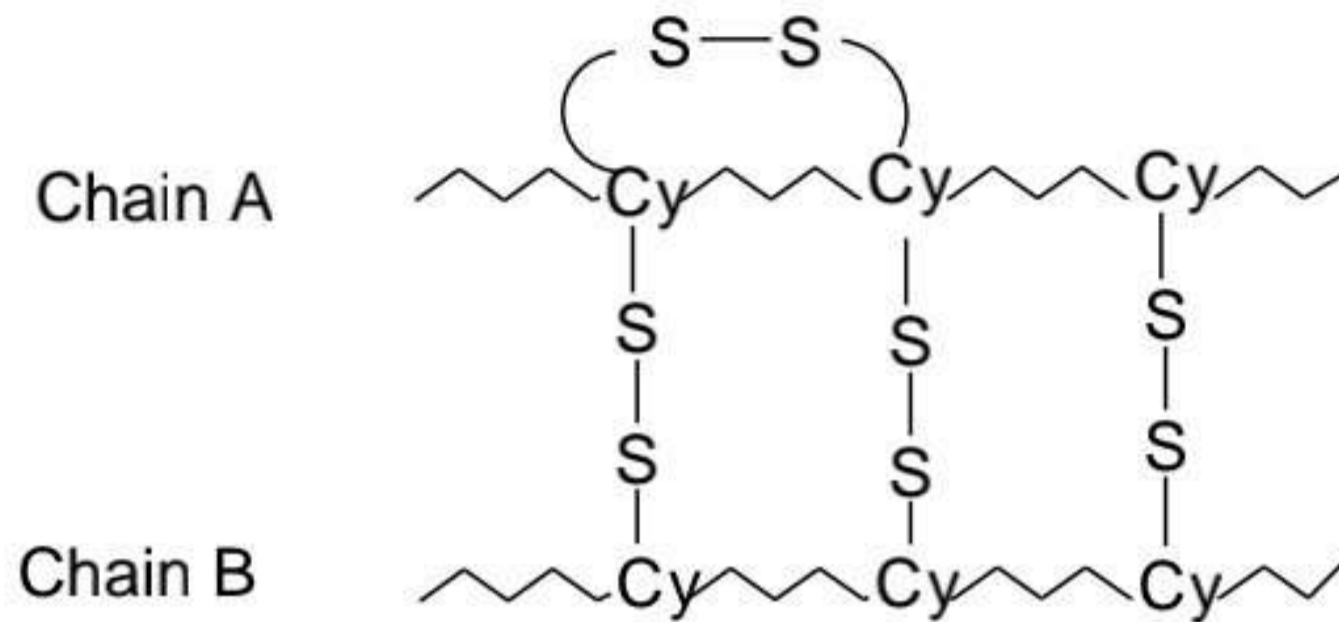
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### **5.7 PROTEINS:**

In spite of the fact that their molecular weights range up to millions, many proteins have been crystallized or at least purified until they behave as homogeneous substances. Care must be taken in the investigation of proteins, because they can be altered quite easily by changes in pH, by uv radiation, by heat, and by organic solvents. Such alteration is generally referred to as denaturation. A denatured protein, while very similar in its gross chemical structure to a native protein, no longer performs its important biological function. Simple proteins, for example the enzyme lysozyme, are hydrolyzed only to amino acids. Others contain non-amino acid portions, called prosthetic groups, and were originally referred to as conjugated proteins. In nucleoproteins (from cell nuclei), the prosthetic groups are nucleic acids: mucoproteins contain complex polysaccharides. Some prosthetic groups are much simpler, as exemplified by the oxidation-reduction enzymes known as flavoproteins, which contain bound derivatives of the vitamin riboflavin.

Proteins are amphoteric dipolar ions which migrate in an electric field and have characteristic isoelectric points. Even though the chain composing the backbone of the protein is comprised of relatively stable amide linkages, proteins are reactive and exhibit highly specific behavior. This reactivity is associated with the free active groups on the side chains, for example amino groups from lysine, guanido groups from arginine, or sulfhydryl groups from cysteine. Many proteins contain several peptide chains held together by cross linkages. Disulfide bonds between cysteines can link two chains, or even remote parts of the same chain: for example, beef insulin contains an A chain of 21 amino acids connected via two

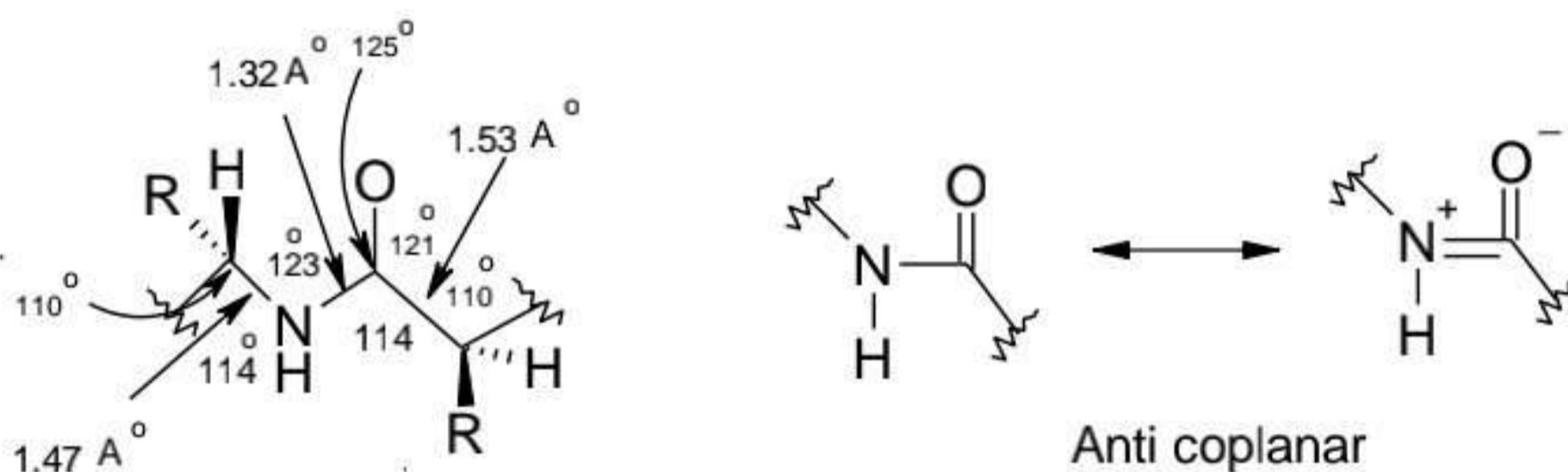
disulfide linkages to a B chain of 30 amino acids, forming a cyclic protein:



The discussion thus far has been involved with the characterization of polypeptides as a linear array of amino acids, that is, primary structure. One must not neglect, however, the manner in which these chains are arranged three-dimensionally. For instance, the finding that a particular amino acid side chain of an enzyme is involved in the catalysis of some reaction tells nothing about the details of its involvement; that is, we do not know whether it actually participates in the reaction, or is involved in the binding of the reactant (substrate) to the enzyme, or is merely necessary for maintaining the overall three-dimensional structural integrity of the enzyme molecule by interactions among the side chains of the constituent amino acids. All these roles, however, stipulate that the amino acid must be located very exactly. This spatial organization of proteins, as mentioned in the introduction to this chapter, is currently a topic of intense investigation in many laboratories.

An invaluable technique for studying three-dimensional protein structure is X-ray crystallography. An X-ray diffraction pattern is obtained from a crystal, and a structure is proposed, if possible, which

would be expected to give such a pattern. From X-ray determinations of amino acid and peptide structures, the amide portion of the chain has been found to be planar and, anti. The following representation shows bond lengths and angles of a unit in the peptide chain:

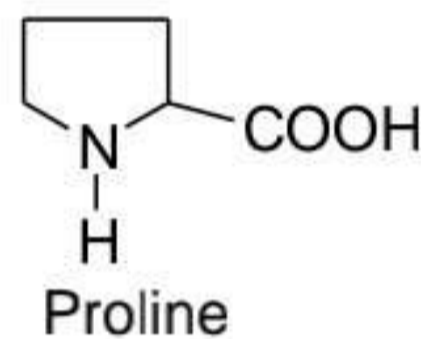


The carbon-nitrogen bond of the amide linkage has approximately 40 percent double-bond character, as a result of resonance. This resonance interaction strongly hinders rotation about that bond. Very importantly, however, rotations are free for bonds between the amide groups and the  $\alpha$  carbons as well as for the  $\alpha$  and the carbonyl carbons, thus permitting many conformations for the protein.

X-ray techniques were also instrumental in elucidating the two major ways in which the peptide backbone can interact with itself. This level of organization is referred to as secondary structure. The first of these two types of interaction is shown in Figure 5.1 and is known as the  $\alpha$  helix. Note that each amide group is hydrogen-bonded to the amide group, which is the third one from it in either direction along the chain. There are 3.6 amino acid units per turn of the helix. The side chains extend away from the axis of the helix. All natural amino acids are of the L-configuration and to date, all protein helices have been found to be right-handed. This is a very common structural component of proteins; an

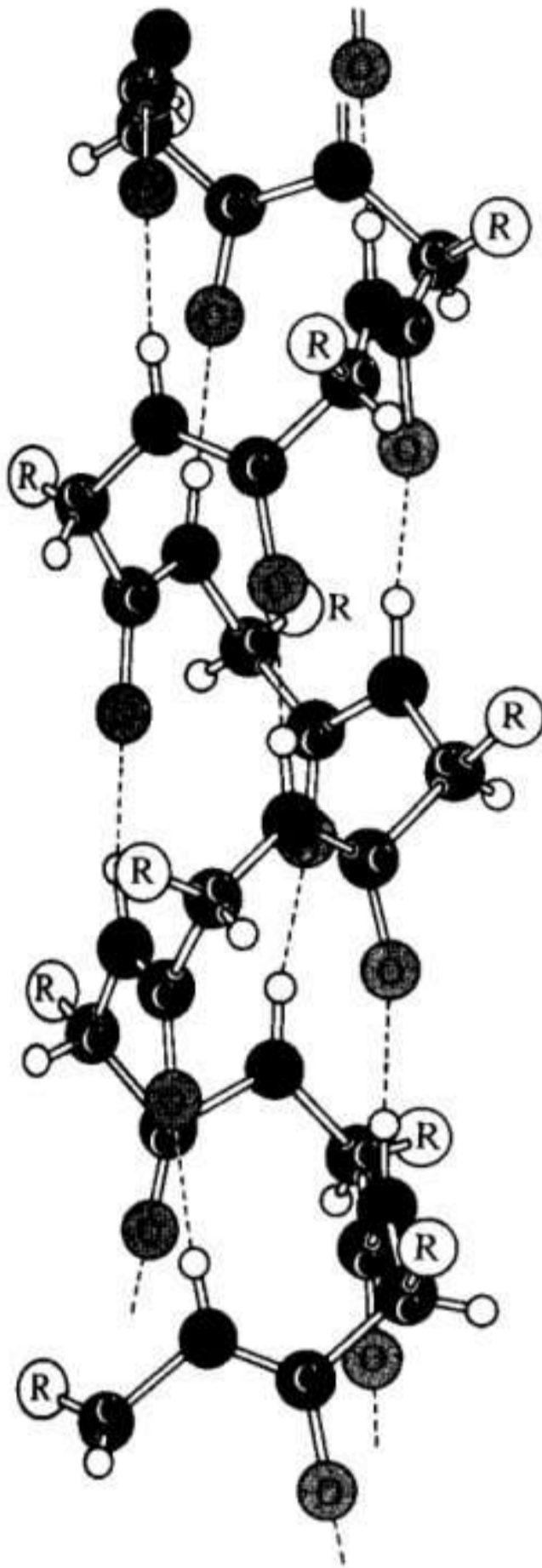
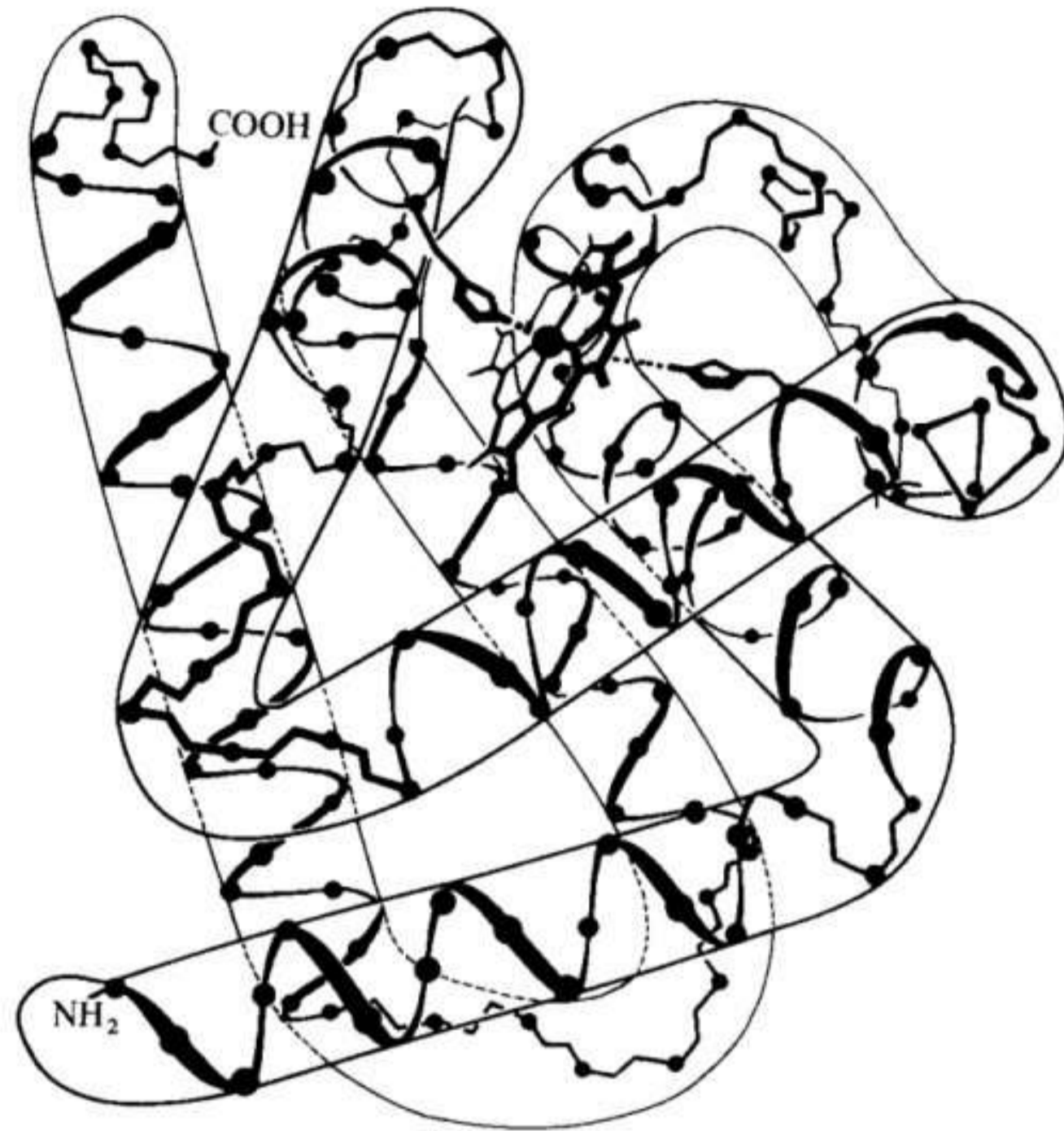
extreme example is the oxygen-carrying protein hemoglobin, which is about 75 percent a helix.

The cyclic nature of the amino acid proline forces the carbonyl group, which is attached to the proline nitrogen, to assume a conformational arrangement different from the one required for helix formation (which all the other amino acids can assume). Thus many proteins are found to consist of helical segments of different lengths, interrupted from time to time where the helix “goes a round corners.” The latter often occurs at the point where a proline residue is found in the amino acid sequence. In most proteins there are also regions where the amino acids are not ordered in any way that is simple to describe.



The interaction of various parts of the protein with each other via the amino acid side chains determines the tertiary structure of the protein. The bonds involved might be salt linkages, such as between an  $-NH_2$  of Lys and a carboxyl of Asp; hydrogen bonds, such as between Ser and His; or van der Waals forces, such as between Tyr

Figure 5.1: Alpha helix.

Figure 5.2: X-ray structure of myoglobin. (from R.E. Dickerson in H. Neurath<sup>(ed)</sup> The proteins, Vol. 2, Academic Press, New York, 2<sup>nd</sup> ed., 1964, P 634).

and Phe. Disulfide bonds between cysteine residues on adjacent chains often help to stabilize the tertiary structure. X-ray crystallography has been used to determine the actual three-dimensional configurations of a number of proteins. Myoglobin, a protein similar to hemoglobin, has the shape shown in Figure 5.2. as determined by Perutz and Kendrew. The resolution of this X-ray study (and several similar studies) is not good enough to be able to see individual atoms: only gross shapes are discernible. The orientation of the peptide chain within this gross

structure can be deduced since the primary structure of the protein is completely known. One of the consistent features of all protein structures studied so far is the presence of large numbers of polar amino acid residues on the surface of the molecule, with large clusters of nonpolar residues in the interior in contact with each other. Note the presence of several helical segments.

Quantitatively of less importance is the  $\beta$ , or pleated-sheet, structure, commonly seen in the fibrous proteins such as those found in silk, hair, and feathers. This is depicted in Figure 5.3. Notice that the chains are antiparallel. You can easily appreciate that steric crowding between R groups would make the straight-chain representation unfavorable. Thus, while silk, with a high percentage of Gly and Ala, could assume this configuration, bulky side chains would prohibit its formation. It is., of course, not obligatory that all parts of a protein molecule have either of these configurations.

There is one further level of organization in proteins, the quaternary structure, which describes the way multiple subunits (not always identical) can aggregate to form large complexes. As mentioned before, tobacco mosaic virus protein is actually a multiple of small subunits. As is usually the case with viruses, the protein complex forms a

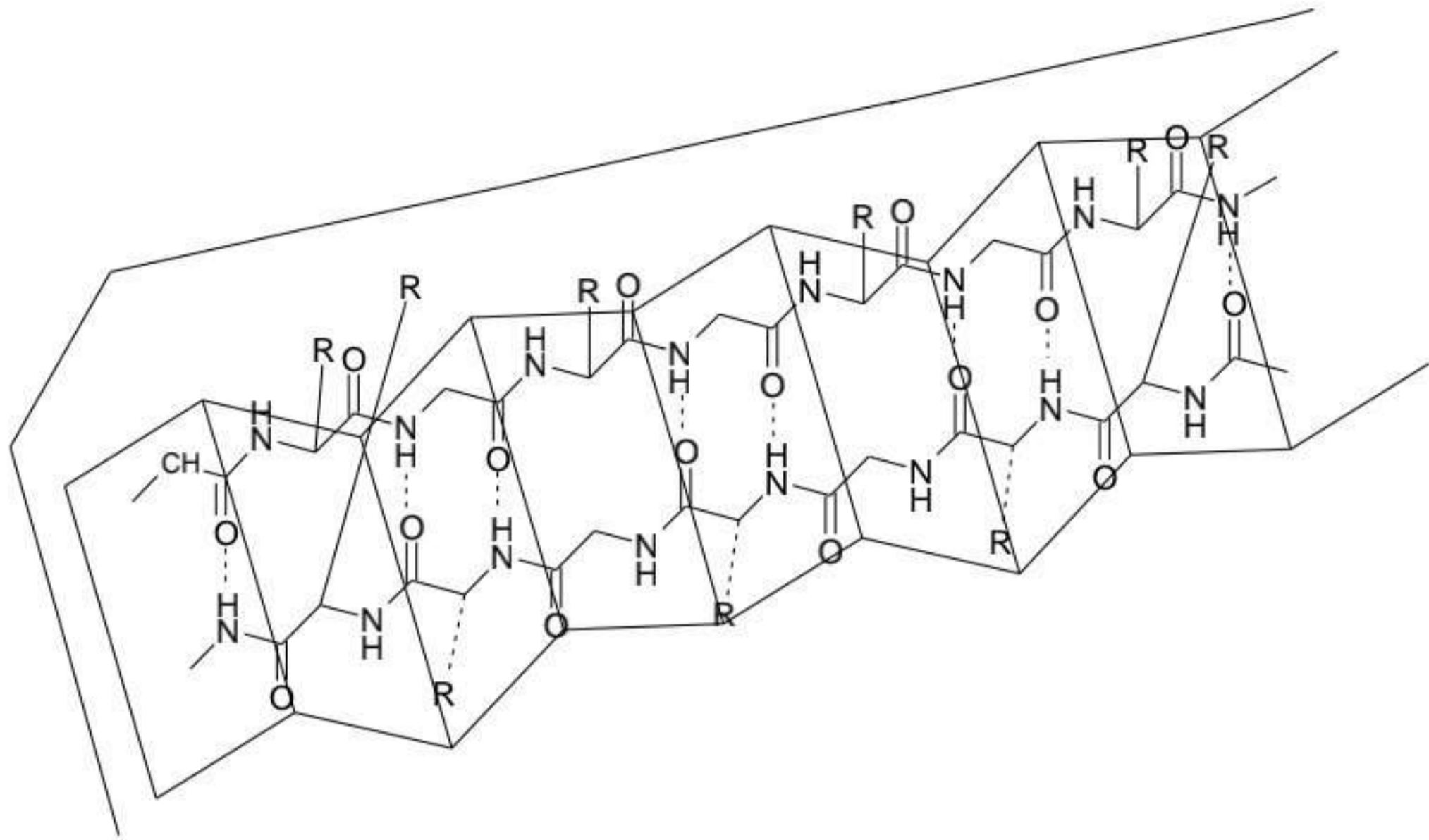


Figure 5.3: Beta, or Pleated-sheet, structure of proteins.

protective sheath around the nucleic acid core of the virus, which, of course, contains the genetic information required for the production of more virus particles.

Many sequential reactions in metabolism are efficiently catalyzed by well-organized complexes of enzymes which obviate the necessity of having the product of one enzymatic reaction float free in the cell waiting until it randomly collides with the enzyme required for its next transformation. Several of these complexes have been broken down into their individual enzyme components. The separated components can subsequently be reassociated in Vitro, and they will exhibit the original overall metabolic transformations. This experimental result shows that after synthesis of the individual enzymes, spontaneous assembly can produce the efficient complex observed in the cell.



**5.8 STRUCTURAL BASIS OF ENZYME CATALYSIS:**

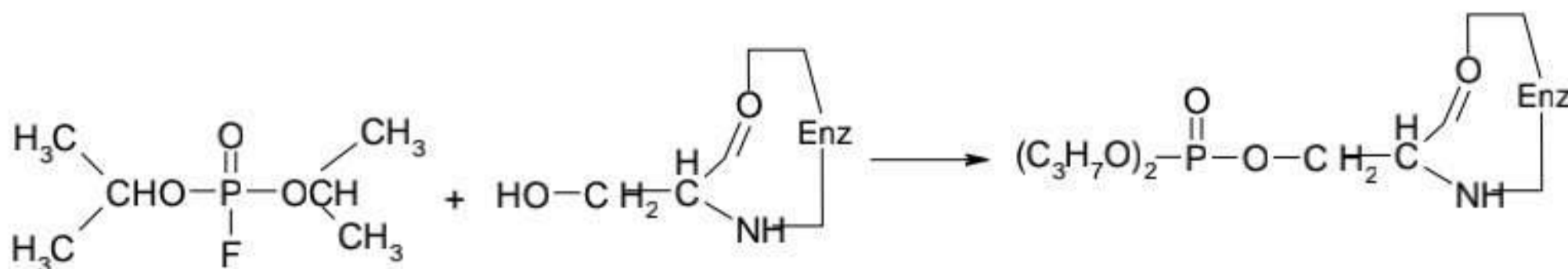
Having now looked at various factors that influence the overall structures of proteins, we are in a position to establish in a more meaningful way the correlation of structure with function which imparts to enzymes their extraordinary specificity and catalytic power. These characteristics (at least in enzymes that require no prosthetic groups for activity) must be determined solely by the specific spatial relationships among individual amino acid side chains of the polypeptide. No types of enzyme catalysis have yet been found that are mechanistically different from reactions carried out in test tubes, and similarly, no enzyme-catalyzed reactions have ever been documented that would not occur (eventually) without catalysis. It is noteworthy, however, that some enzymes have the capacity to speed up reactions by a factor of 10<sup>10</sup> beyond their rates without catalysis.

As an example to illustrate a structure-function relationship, we will use the enzyme chymotrypsin, which was mentioned in Section 5.6. As was described there, this proteolytic (peptide-hydrolyzing) enzyme preferentially attacks peptide bonds whose carbonyl function is furnished by an aromatic amino acid.

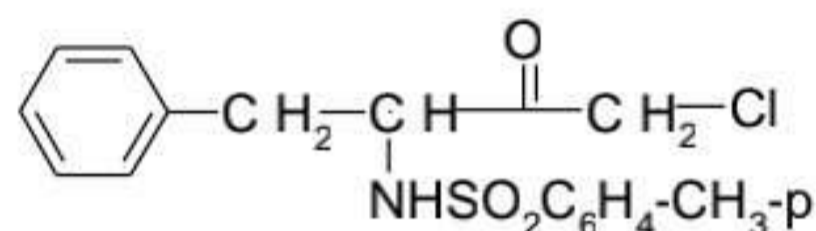
Some time before the three-dimensional structure of chymotrypsin had been established, a number of amino acids had been suspected by being components of the “active site” of the enzyme. A short description of several experiments that led to these suspicions will illustrate the type of approach used to investigate the mechanism of action of an enzyme.

- 1- Reaction of the enzyme with low concentrations of diisopropyl fluorophosphate led to rapid inactivation of the enzyme. Upon

hydrolysis, the diisopropyl phosphate group was found to be covalently linked to a serine residue.



- 2- The pseudo-substrate 1-chloro-4-phenyl-3-(*p*-toluenesulfonamido)-2-butanone,



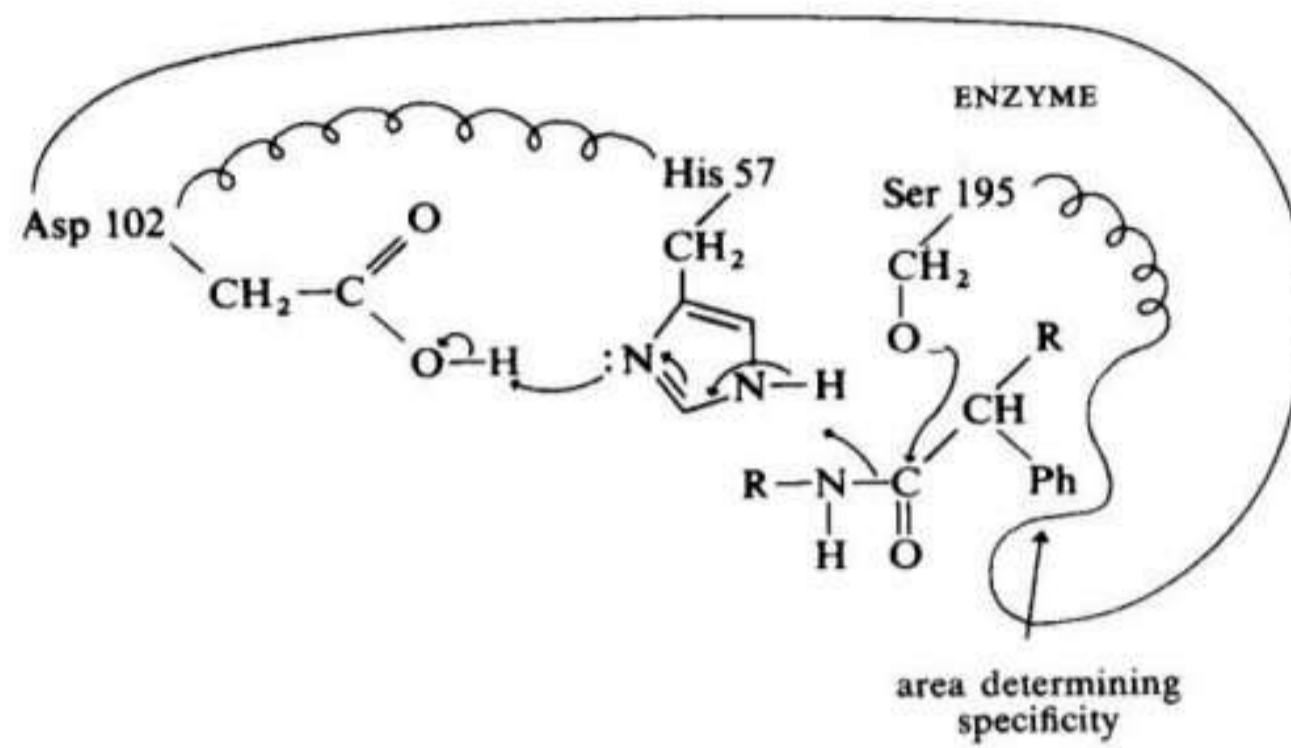
was found to react covalently with the enzyme in 1:1 stoichiometry, leading to complete loss of activity. Notice the designed similarity of the compound to a phenylalanine--containing peptide. suggestive that this compound is brought specifically into the region of the active site by the specificity-determining portions of the enzyme. Upon hydrolysis of the enzyme, the reagent was found to have reacted with a histidine residue.

- 3- Further evidence implicating histidine as a component of the active site involved the drastic change in the enzyme's activity as the pH of the reaction was varied near the pK of an imidazole nitrogen.
- 4- When the enzyme reaction was carried out in D<sub>2</sub>O rather than in water, the rate of hydrolysis decreased by over half, thus implicating a proton transfer (general acid-base catalysis) in at least the rate-determining step.

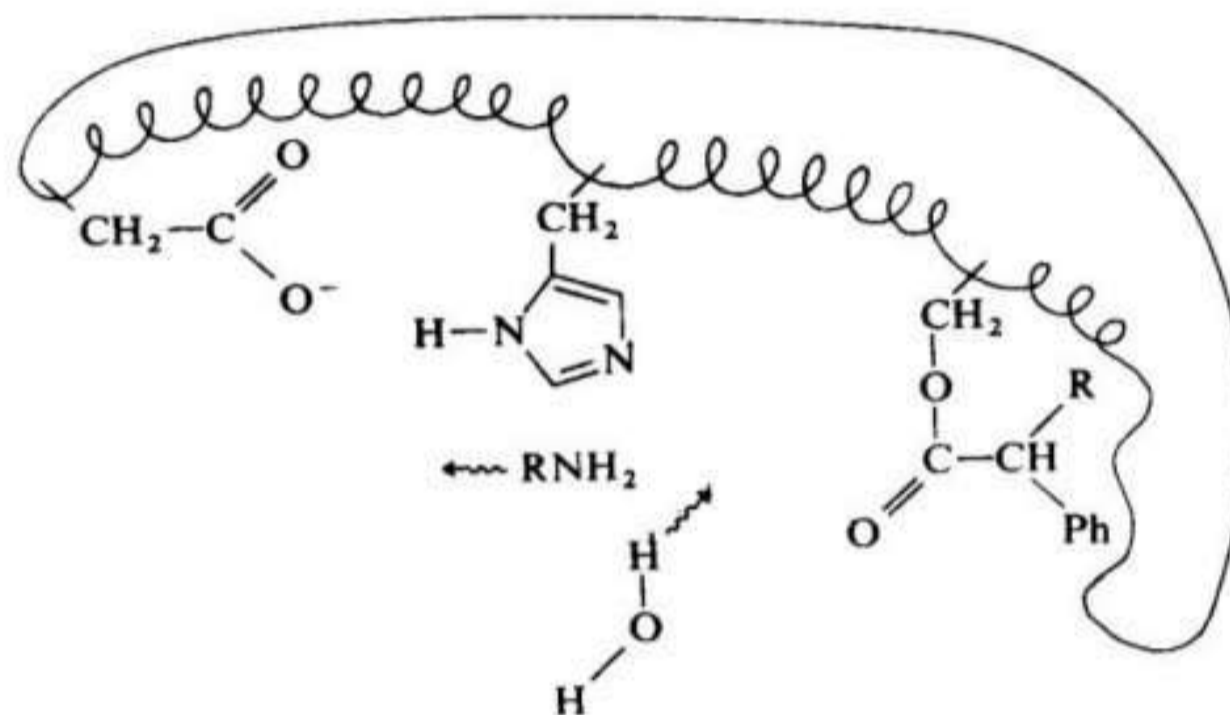
Subsequent X-ray analysis of the crystalline enzyme revealed that what would appear to be the active site is rather close to the surface of the molecule, as if designed to approach a rather large substrate. The high specificity of the enzyme would seem to be determined by two relatively short peptides within the enzyme which consist entirely of small nonpolar amino acids forming a kind of pocket into which the aromatic substrate would be held by van der Waals forces. Nearby were histidine and serine residues, and also in the same area was found an aspartic acid that could facilitate protonation of the histidine ring. Very importantly, these amino acids are not consecutive as one might at first think, but are actually separated from each other linearly by many other amino acids. Histidine is number 57, aspartic acid 102, and serine 195. The remainder of the protein is presumably involved in holding these catalytic and specific segments in the proper neighboring relationship.

The overall mechanism of action of chymotrypsin, shown in the following figures, was postulated before X-ray data were available (with the exception of the initial proton donation by aspartic acid). The confirmation by X-ray analysis of the feasibility of the proposed chemical mechanism is a striking example of the mutual benefit that can be derived from different but complementary lines of investigation of a single problem. The proposed mechanism of action of chymotrypsin is illustrated as follows:

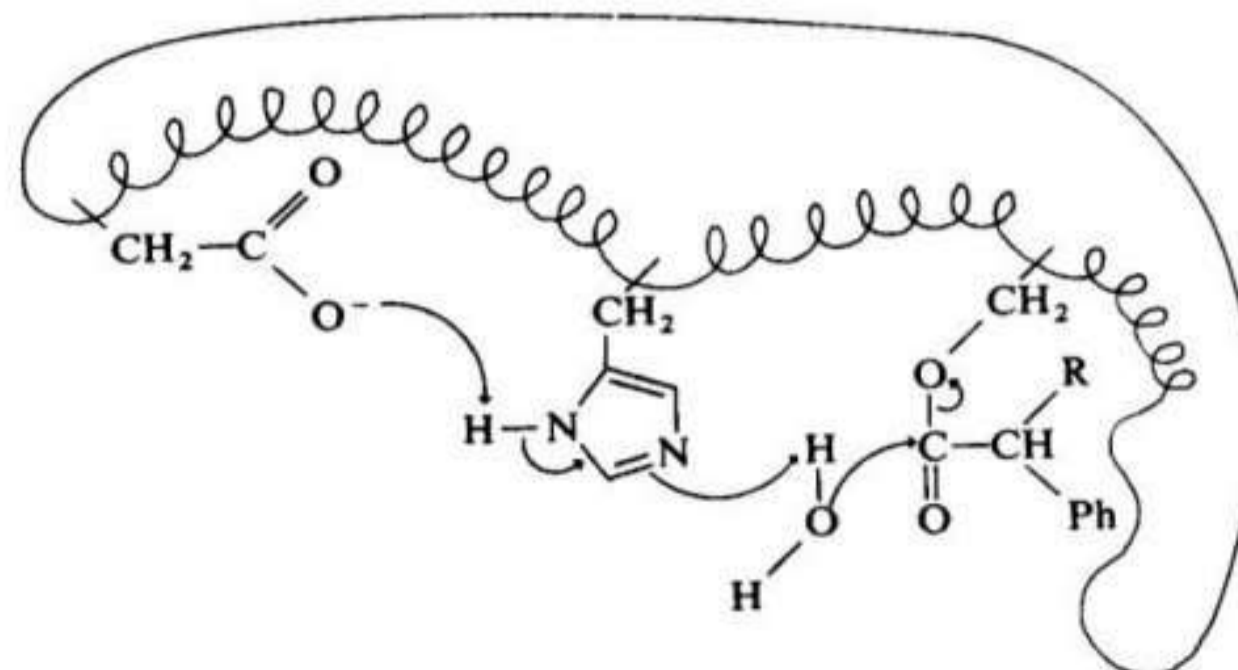
1. *Initial Binding of the Substrate* (after proton transfer from Ser to Asp via His)



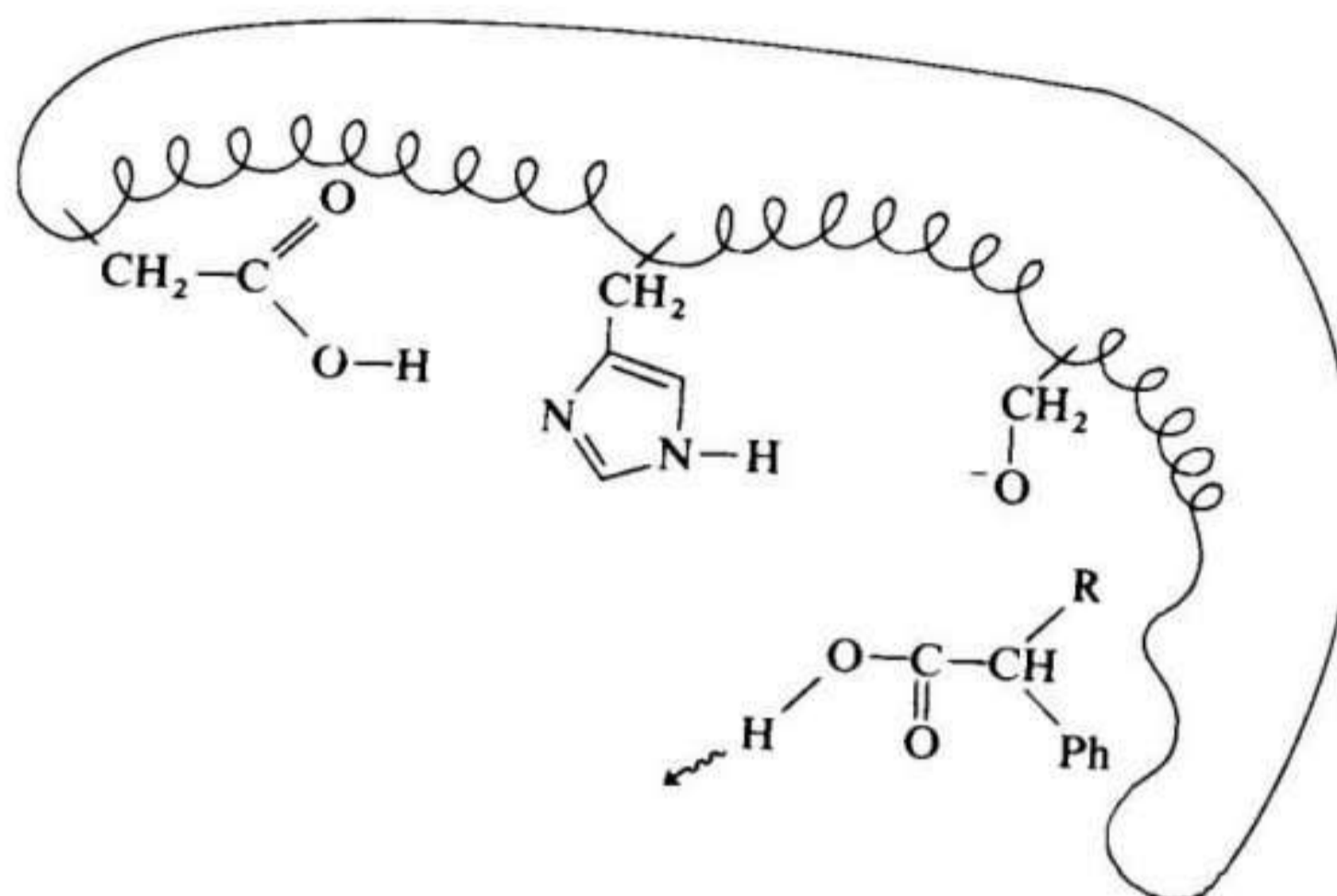
2. *Formation of an Acyl Enzyme Intermediate and Release of Amine*



3. *Deacylation*

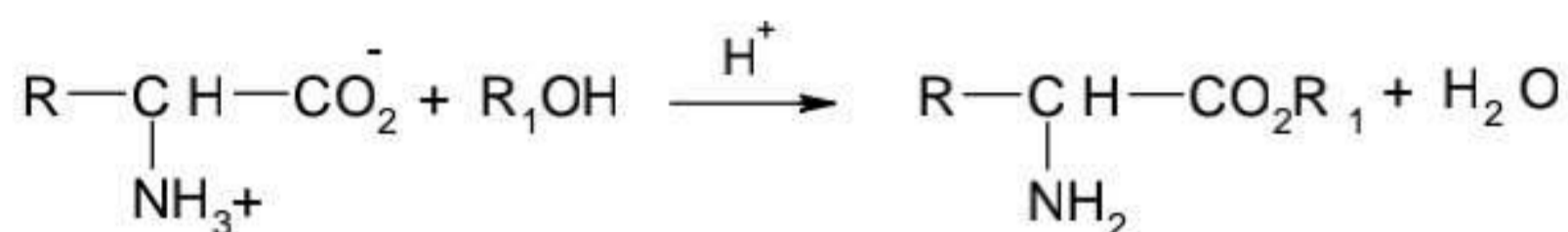


## 4. Release of Carboxylic Acid

**5.9 PEPTIDE SYNTHESIS:**

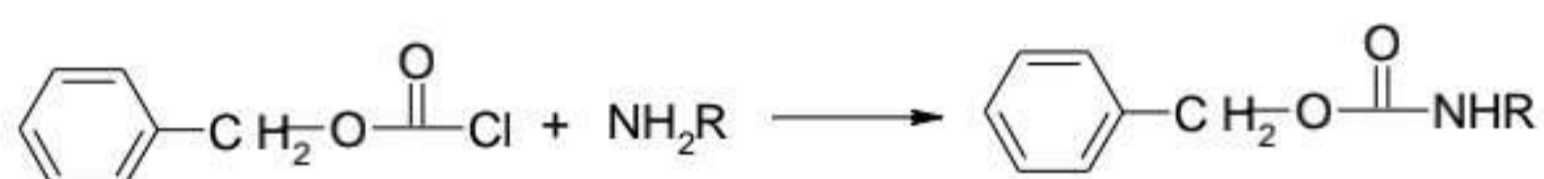
Peptide synthesis has long been an intriguing area of organic chemistry. The ability to duplicate in the laboratory one of the most complex processes of nature holds a fascination for the organic chemist. In any synthetic sequence directed toward the preparation of a particular polypeptide, there are difficulties to be encountered characteristic of the side-chain functional groups of the individual amino acids. These problems are too numerous and varied to go into here. However, some problems are common to all peptide syntheses. The two main difficulties seem to be (1) blocking the amino group of an amino acid while activating the acid group of the same molecule; and (2) blocking the acid group of an amino acid while leaving the primary amino group free to react.

The second of the two problems is taken care of quite effectively by treating the amino acid with an alcohol to give an ester.

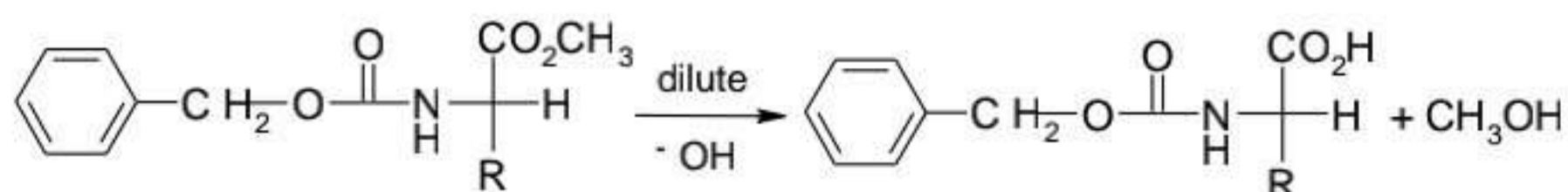
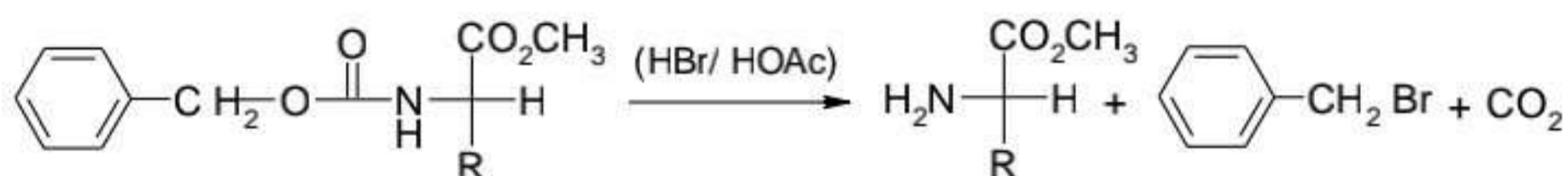
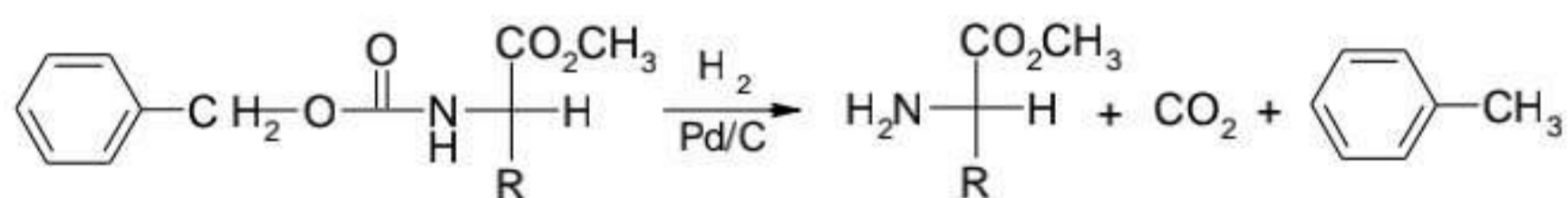


The first problem, that of blocking the amino group while activating the carboxyl group, is somewhat more involved and has been dealt with in many ways. The carbobenzoxy and t-butoxycarbonyl groups are widely used as amino blocking groups.

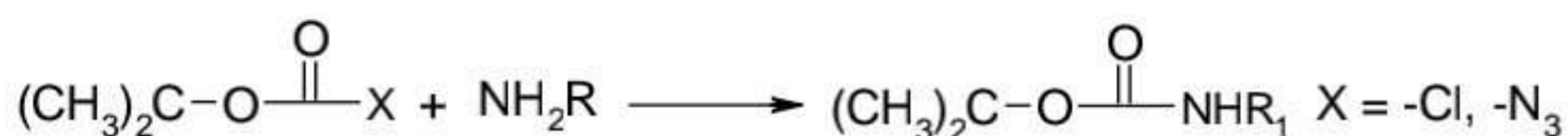
1- Carbobenzoxy. The N-carbobenzoxy (N-benzyloxycarbonyl) group is formed by treating an amino acid with benzyl chloroformate.



This protecting group has the advantage that it can be removed by hydrogenolysis or by acid hydrolysis, thereby generating the free amine, but it is relatively stable to dilute alkali.

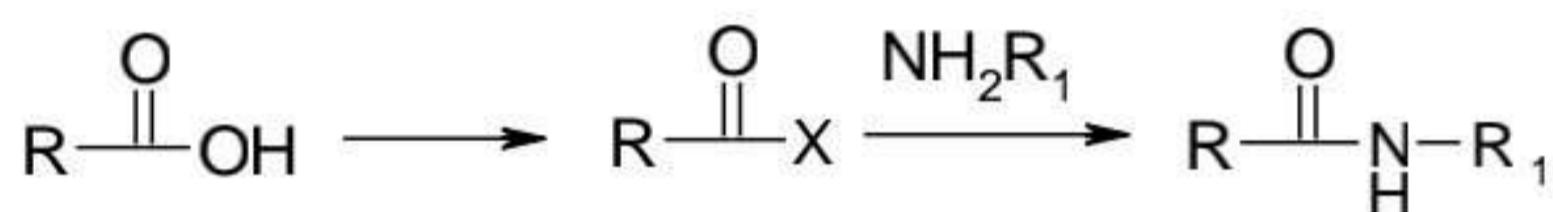


2- Butoxycarbonyl The N-t-butoxycarbonyl group is formed by treatment of the amino acid with t-butoxycarbonyl chloride (t-butyl chloroformate at 0 °C or by treatment of the amino acid with the more stable t-butoxycarbonyl azide at slightly higher temperatures.

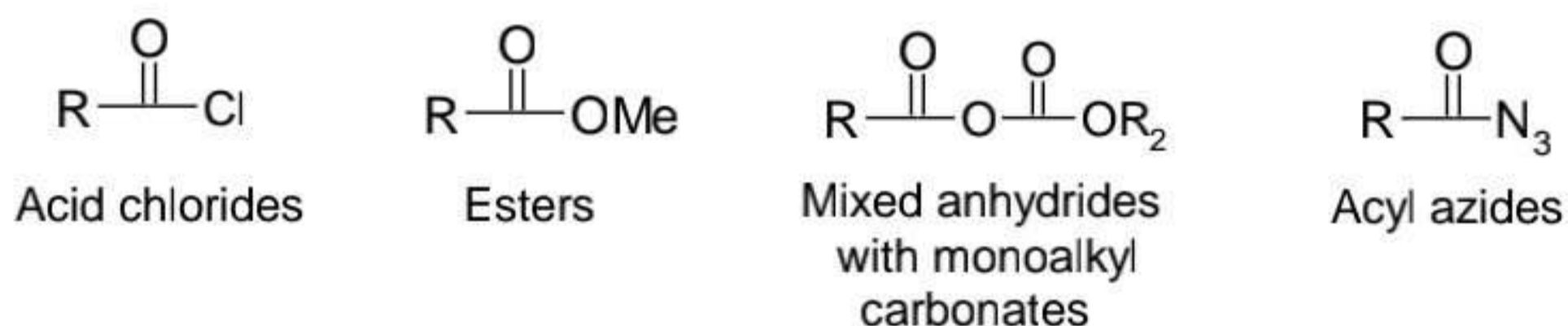


The N-t-butoxycarbonyl group can be removed by dilute acid, leaving the free amine. but it is unaffected by hydrogenolysis or dilute base.

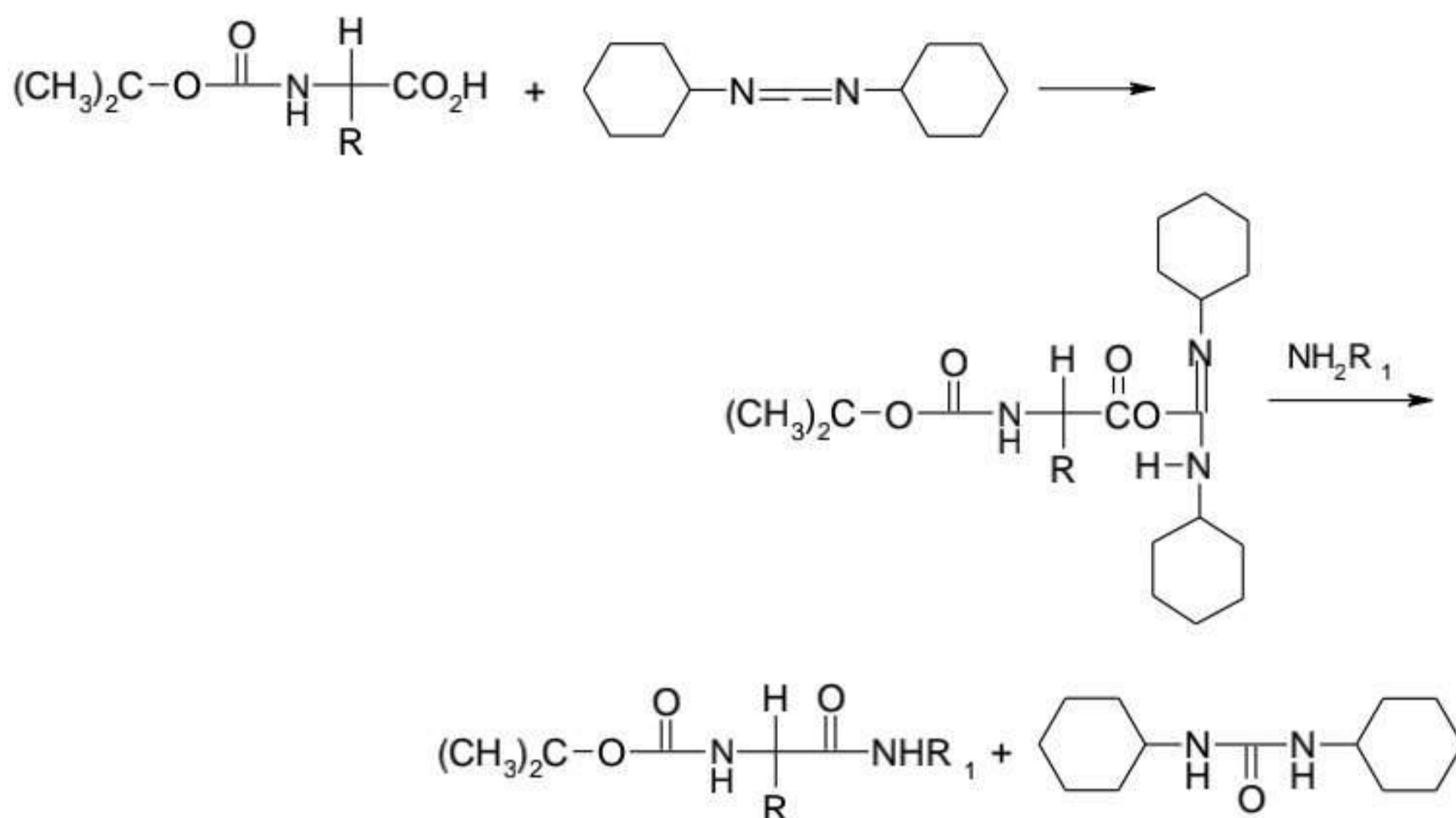
As far as activating the acid function is concerned, it is necessary to convert the -OH of the acid to a better leaving group.



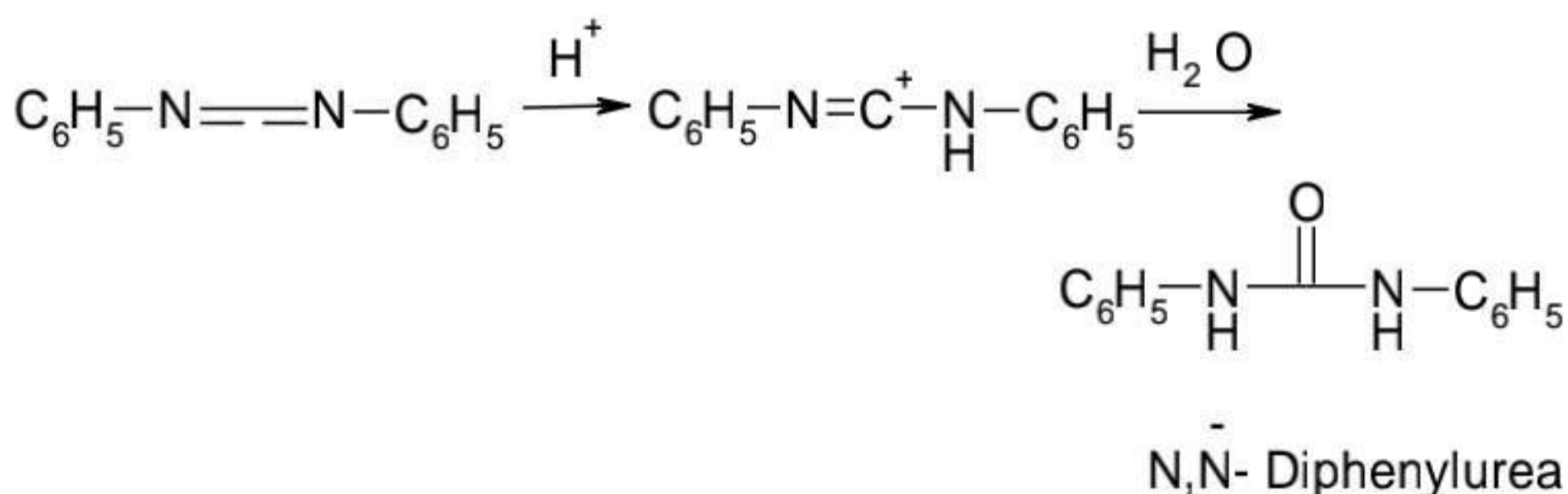
Many such groups have been used and include the following:



The use of these groups has been largely supplanted by that of dicyclohexylcarbodiimide-a reagent that in one step activates the carbonyl group and effects the coupling between an amino group and an acid with the removal of water.



Carbodiimides. These are a special class of imines of the general structure  $\text{R}-\text{N}=\text{C}=\text{N}-\text{R}$ : they are formally diimides of carbon dioxide. They add nucleophiles readily at the central carbon atom. For example, water adds to give substituted ureas; the reaction is acid-catalyzed.



Carbodiimides can be prepared by dehydration of ureas, the reverse of the preceding reaction.

One can well imagine how it would be possible to build a chain of any length with amino acids in any desired sequence by adding on one at a time. In the laboratory, however, this is not the usual method. Normally small (two-, three-, or four-unit) polypeptides are constructed and then



these units are coupled as if they were themselves amino acids. Du Vigneaud,' in fact, employed this method in his landmark synthesis of oxytocin in 1954. The main disadvantage of this synthetic approach is the racemization, which is nearly always a problem in each step; the purification of intermediates of two, three, or four polypeptide units to optical purity is consequently very difficult.

A recent and rather novel approach to polypeptide synthesis is that of Merrifield. The method, essentially, is to bind an amino acid through the carboxyl group to a highly porous polymeric resin. The t-butoxycarbonyl protection can then be removed from the amino group by merely washing the resin with acid. By then washing the resin with an activated acyl compound, the polypeptide chain is built up. In this method, the purification of smaller intermediates is eliminated, but, depending upon the accumulated amount of racemization, the problem of purification of the final polypeptide product still remains. By this method biologically active ribonuclease was synthesized by assembling the 124 amino acids using 369 chemical reactions and 11,931 steps of the "peptide synthesis machine," which carries out the operations with a minimum of human intervention.

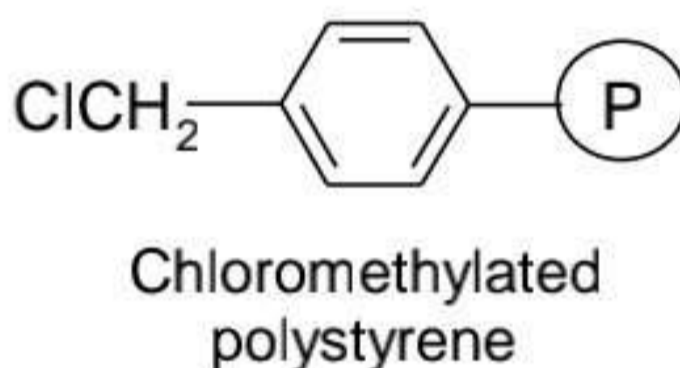
The purity of the final protein presents a difficult synthetic problem, which can be understood using the ribonuclease synthesis as an example. Suppose that a 90 percent yield of pure product is obtained when each amino acid is added, the other 10 percent of the product being racemic or other impurity. As most of us know from our laboratory work, a 90 percent yield is usually pretty good. After 124 steps, each with a 90 percent yield, the overall yield is  $(90 \text{ percent})^{124} = 0.0002 \text{ percent}$ . A 95

percent yield on each step will give an overall yield of 0.2 percent, while if the yield can be raised to 99 percent on each step, the overall yield can be raised to 30 percent. Clearly, yield and purity are extremely critical problems in protein synthesis.

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EXERCISE 5.4:

The polymeric resin used in the Merrifield “solid-phase” synthesis can be designated as:



In the first step of a solid-phase synthesis, a N-t-butoxycarbonyl-protected amino acid (*t*-BOC-amino acid), as its sodium salt, is allowed to react with the polymer. The product thus obtained is then treated with dilute acid. Draw a diagram that would be representative of the solid phase at this stage, if phenylalanine were used as the amino acid.

---

EXERCISE 5.5:

Draw the structure of the material which would be obtained by allowing the product produced in Exercise 5.4 to react with *t*-BOC-glycine in the presence of dicyclohexyl-carbodiimide.

EXERCISE 5.6:

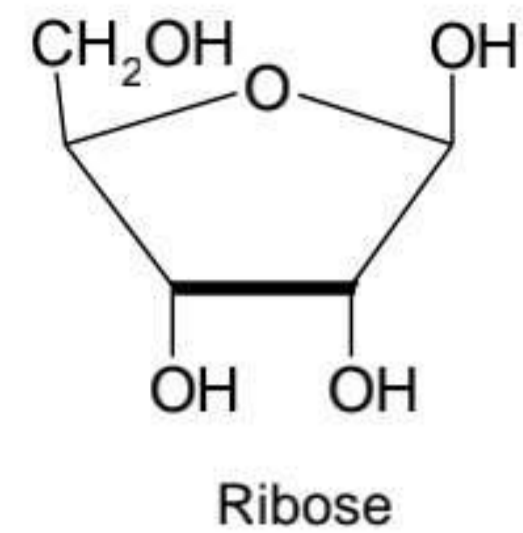
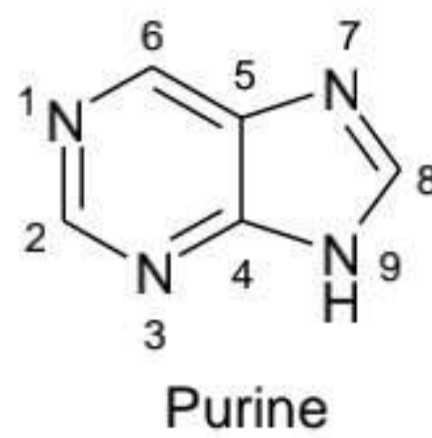
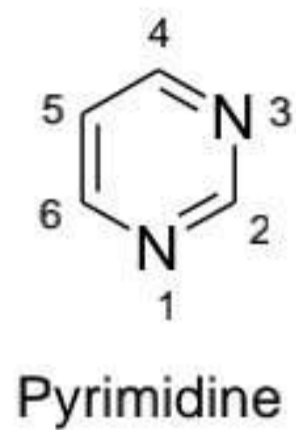
Beginning with the material obtained in Exercise 5.5, outline the steps necessary for the synthesis of the tetrapeptide Pro. Ala. Gly. Phe. Note that the completed peptide can be released from the polymer support by treatment with hydrogen bromide in trifluoroacetic acid.

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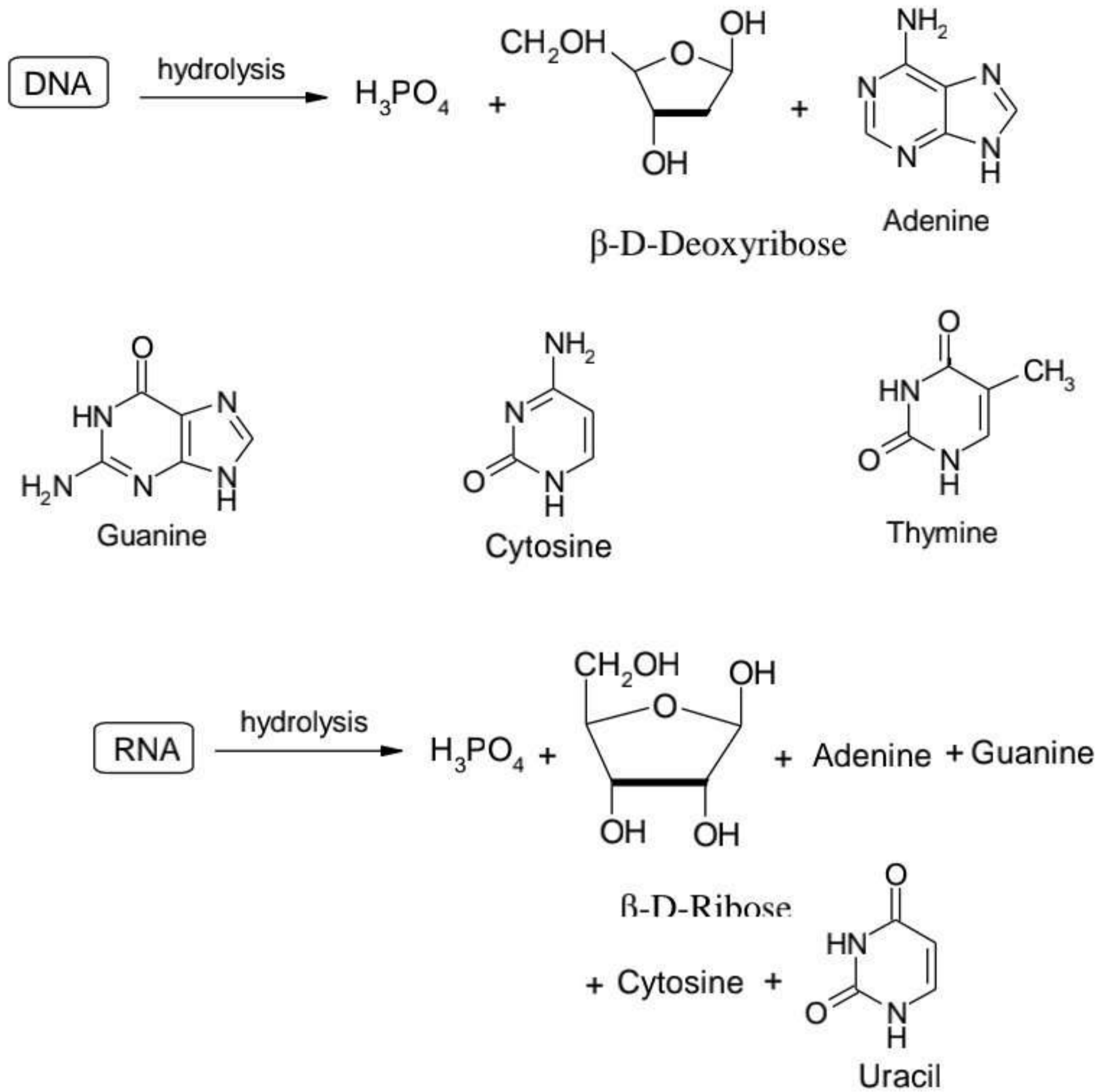
**5.10 THE NUCLEIC ACIDS:**

Of the three types of vitally important biopolymers, polysaccharides (Chapter 4) and proteins (Section 5.7) have already been discussed. The nucleic acids constitute the remaining type. Two varieties of nucleic acids are found in cells, ribonucleic acids (RNA) and deoxyribonucleic acids (DNA). As discussed below, DNA constitutes the genetic material of cells. Both RNA and DNA are essential for the biosynthesis of proteins. Like the proteins themselves, RNA and DNA are very large molecules: molecular weights of up to 2,800,000.000 have been estimated for some DNAs.

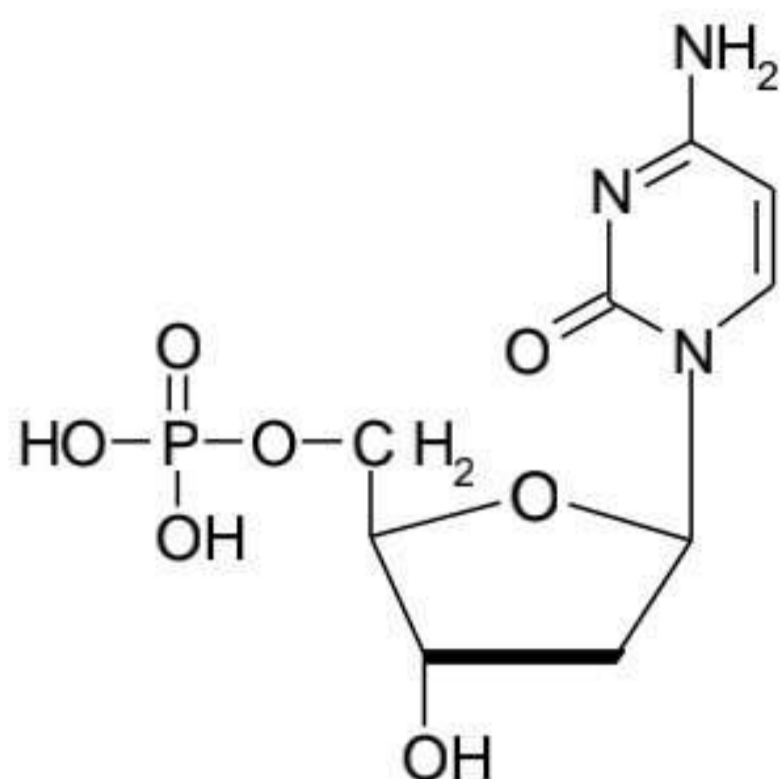
Following are shown the structures of some compounds we shall need to have in mind before going further with the discussion of nucleic acids. These compounds have all been mentioned in earlier chapters, but their structures will be repeated here for convenience. They are pyrimidine and purine and ribose.



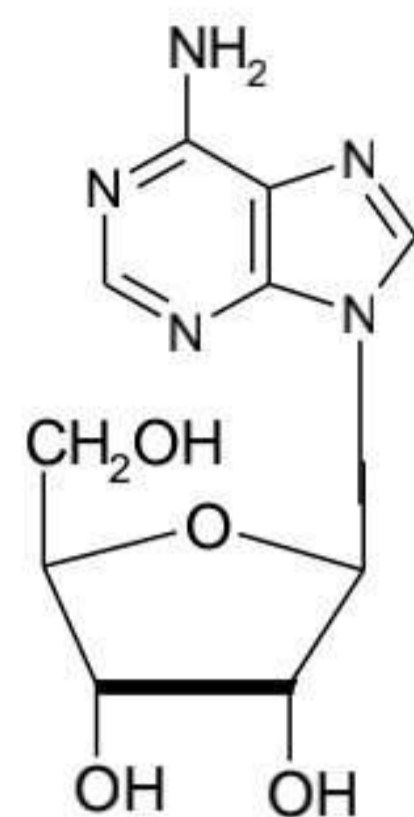
On hydrolysis, both types of nucleic acids yield phosphoric acid, a sugar, and a mixture of purine and pyrimidine bases. The sugar from RNA is ribose, that from DNA is deoxyribose. The major bases from DNA are the purines adenine and guanine and the pyrimidines cytosine and thymine. RNA yields mainly adenine, guanine, cytosine, and another pyrimidine base, uracil.



Mild degradation of a nucleic acid yields a mixture of acids known as nucleotides. Each nucleotide contains the elements of one purine or pyrimidine base, one phosphate unit, and one pentose unit. The phosphate unit may be selectively removed by further careful hydrolysis to convert a nucleotide into a nucleoside, a molecule built up of a pentose joined to a purine or pyrimidine base. In a nucleotide, C-1 of the sugar is joined to N-1 of a pyrimidine or N-9 of a purine; the phosphoric acid unit is present as an ester at C-5 of the sugar.

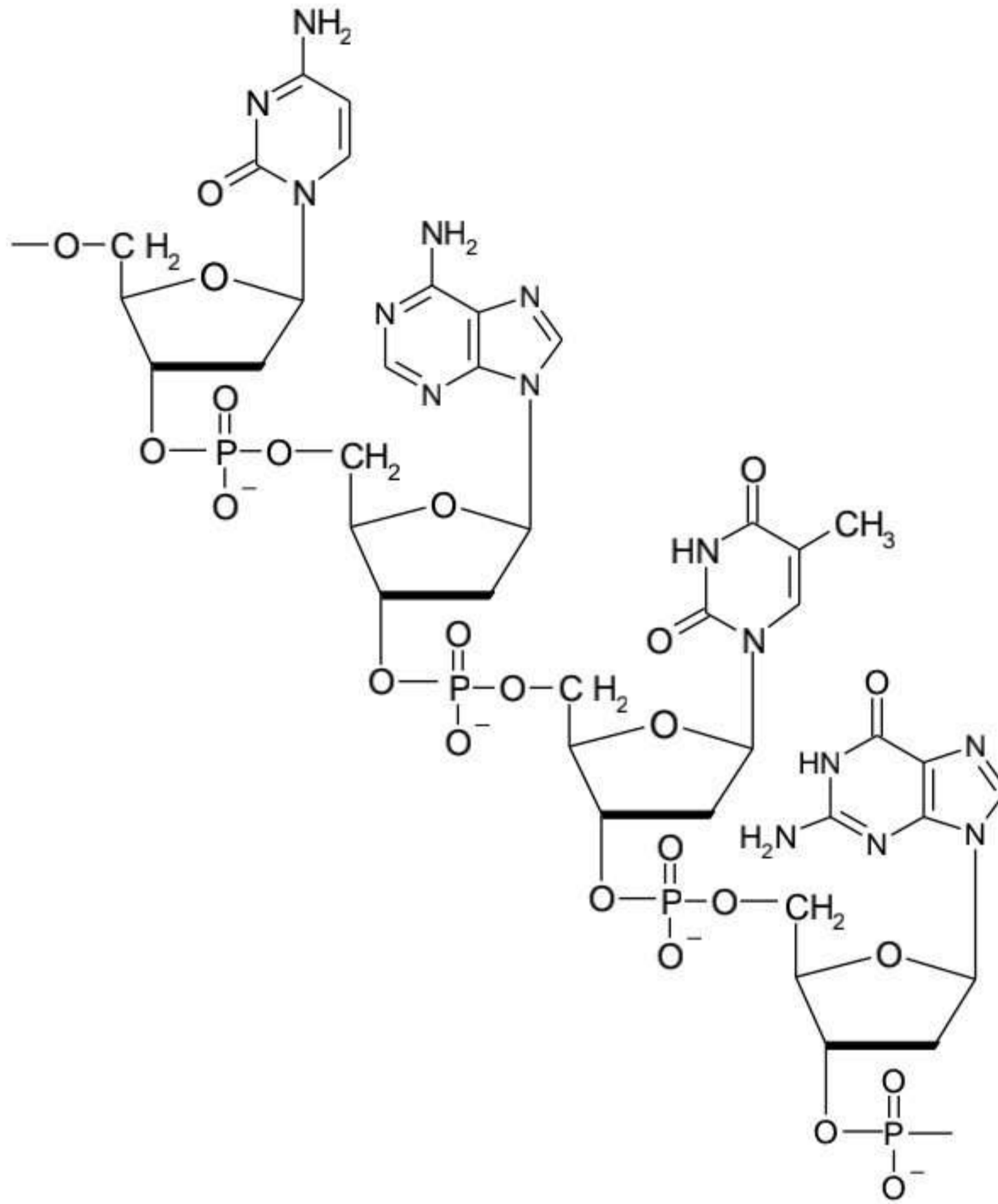


Deoxycytidylic acid  
(a nucleotide)

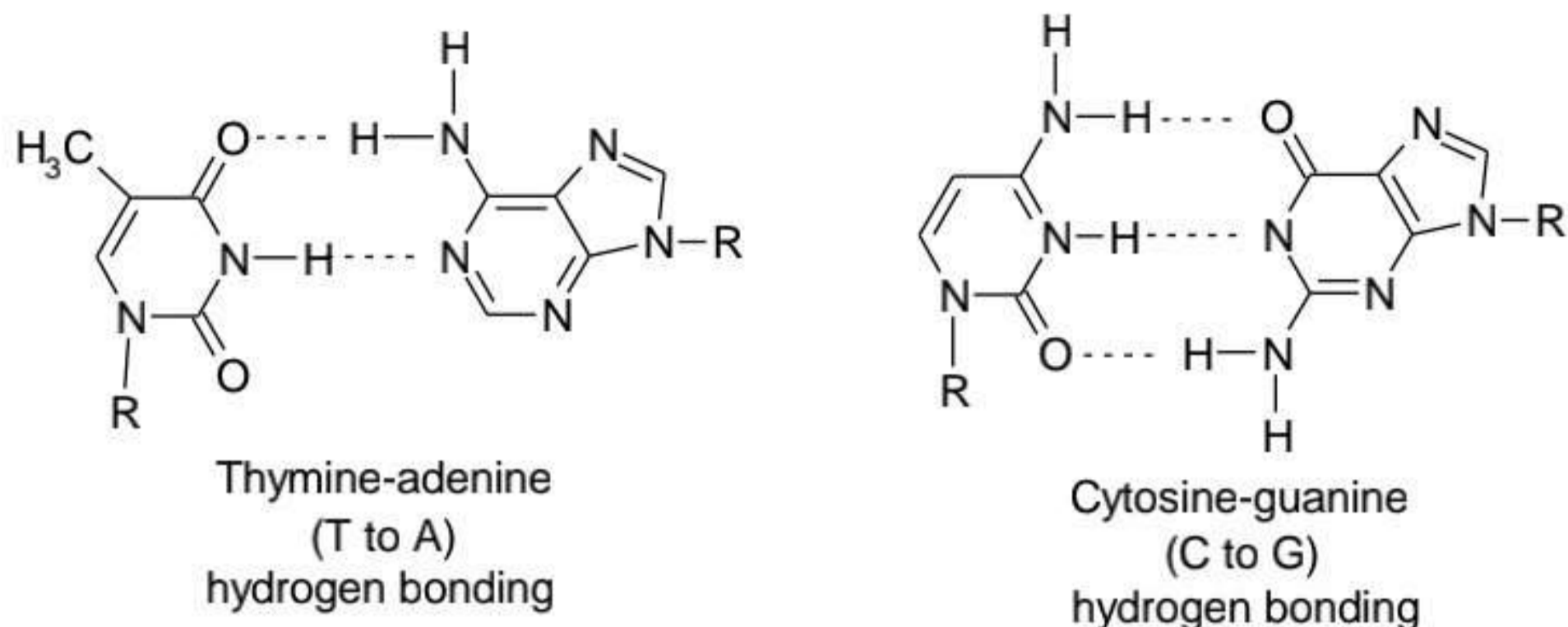


Adenosine  
(a nucleoside)  
(A)

In a nucleic acid chain the phosphoric acid is esterified to form a bridge between C-5 of the sugar of one nucleoside and C-3 of the sugar of another nucleoside. In this way, the sugar-phosphate units can form a long backbone or framework, which bears purine and pyrimidine base substituents at regular intervals. A typical segment of a DNA chain is shown.



Watson and Crick in 1953 proposed the now-accepted double-helical structure of DNA. According to their analysis, the DNA molecule actually consists of two complementary strands that are twisted about a common axis as helices having the same chirality (handedness). Each adenine unit of one chain is specifically hydrogen-bonded to a thymine of the opposite chain, and each guanine of one chain is similarly bonded to a complementary cytosine unit.

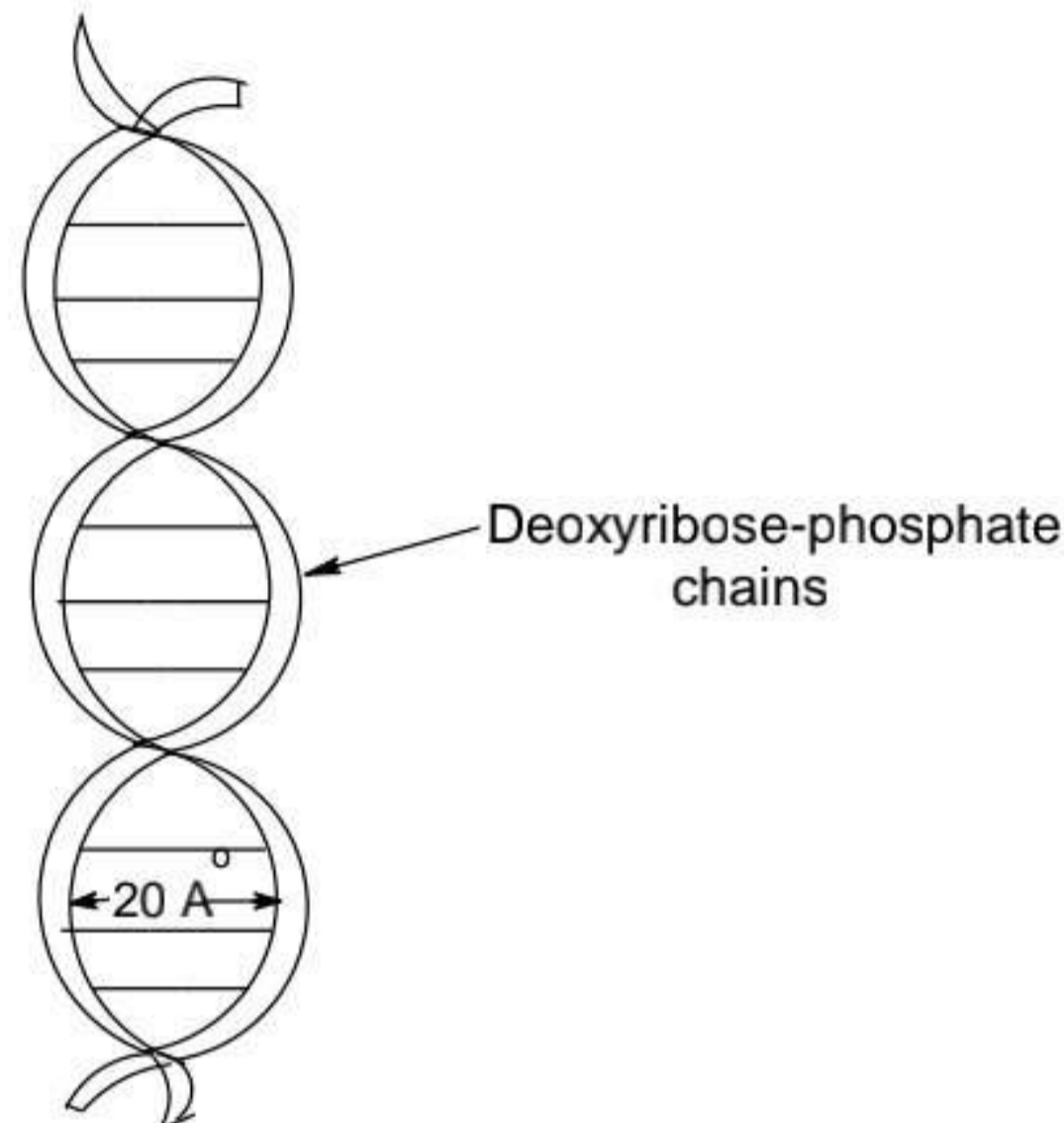


The double-helical structure of DNA is shown schematically in Figure 5.4. The helical strands represent the sugar-phosphate backbones, which are held nicely in place by hydrogen bonding between the complementary base units. The order of the bases on the chain of the DNA molecule is extremely significant biologically; it is the fundamental source of the hereditary information of the genes.

A molecule of DNA reproduces itself by a remarkably simple mechanism. The two strands of the DNA molecule dissociate, and then free nucleotides hydrogen-bond with the nucleotides of the dissociated strands. An enzyme catalyzes the polymerization of these free nucleotides in an order complementary to that of the original strands, producing two new double-stranded DNA molecules identical to the original one. It



Figure 5.4: Double-helix structure of DNA.



has been said that this simple process is the “secret of life.” If not the secret of life, it is at least the secret of why children look much more like their relatives than like elephants or oak trees.

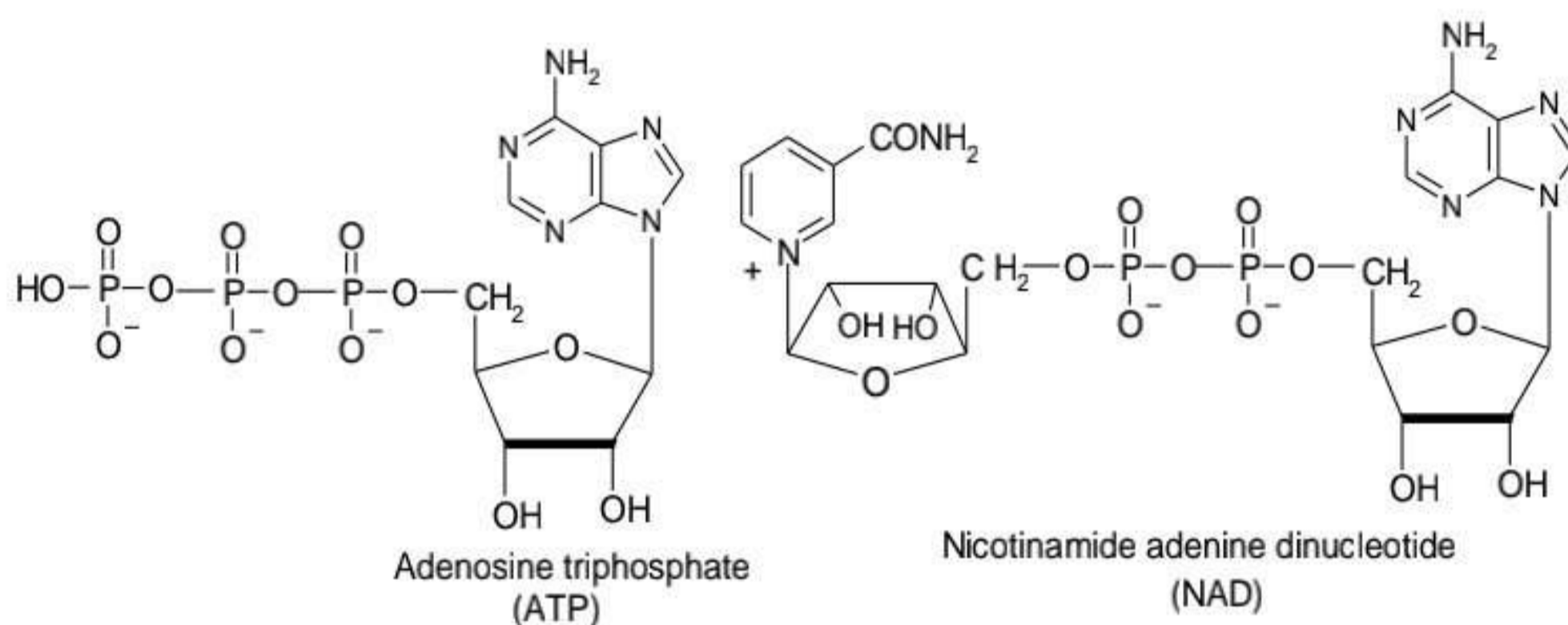
The DNA has two important functions. First it is the reference material or "book of instructions" as to how a given plant or animal is to be constructed, and it reproduces itself as explained above. Its second function is to act as a template in producing RNA. RNA is the material that actually carries out the synthesis of proteins.

Nucleosides and nucleotides serve other very important biochemical roles as portions of essential biological catalysts (coenzymes). Adenine units are most frequently encountered in these compounds.

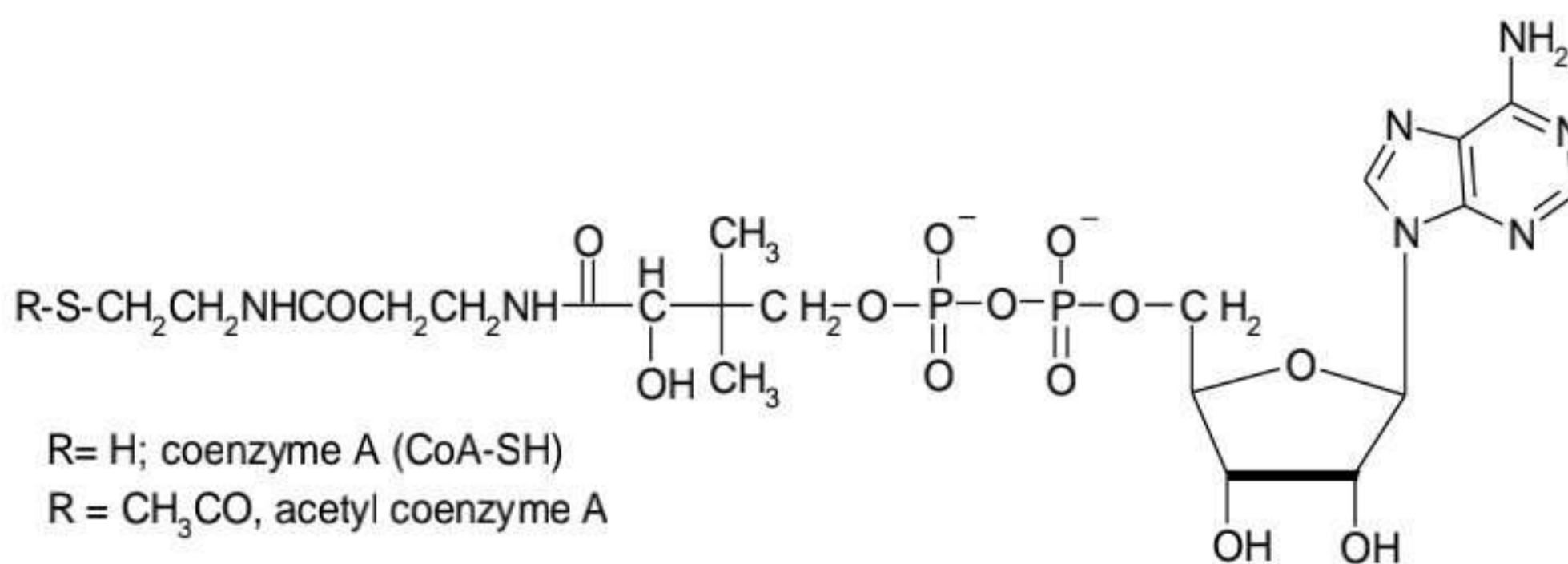
Numerous enzymes require the presence of a small nonprotein moiety more or less tightly bound to the protein for efficient performance of catalytic function. Since their nonprotein moieties are intimately

involved with the overall reaction, they are termed coenzymes.

Examples of such nucleotides are nicotinamide adenine dinucleotide (NAD), and adenosine triphosphate (ATP), which functions as a pool for chemical energy in cells because of its energy-rich triphosphate unit.

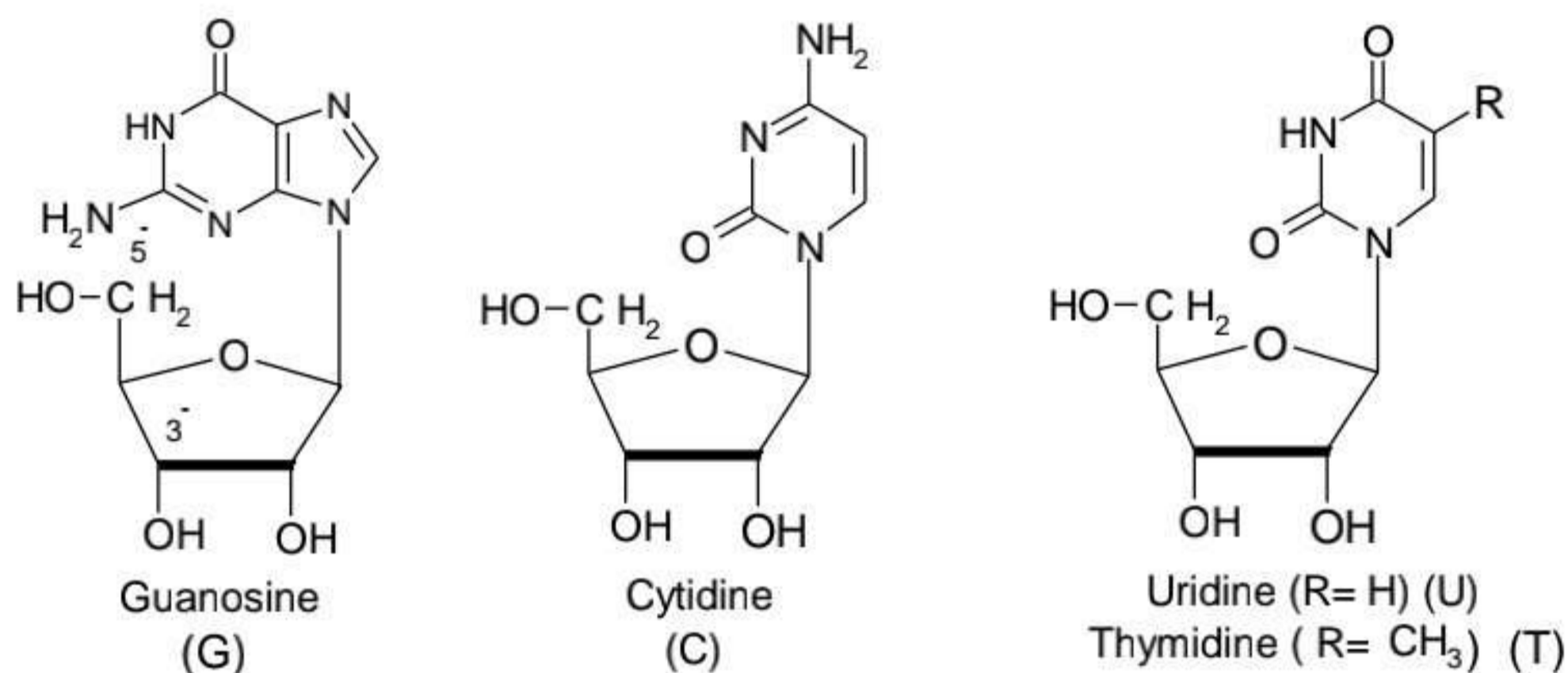


A third very important adenine nucleotide derivative is coenzyme A (CoA-SH), which plays a very significant role in biosynthesis, especially in acyl group-transfer reactions. Acyl derivatives of CoA-SH are effective acylating agents because of diminished resonance interaction between the sulfur atom and the carbonyl group, as compared with oxygen-carbonyl delocalization in ordinary esters.



**5.11 PROTEIN BIOSYNTHESIS:**

Ribonucleic acids have the general formula shown in the previous section. Note that the nitrogen bases cytosine, uracil, guanine, and adenine occur in a precise and characteristic sequence in any given ribonucleic acid chain. A more wieldy general formula results from the following abbreviations. The letters U, G, A, C, and T represent the nucleosides (ribose-nitrogen base conjugates) uridine, guanosine, adenosine (Section 5.10), cytidine, and thymidine, respectively.



The letter  $p$  represents the phosphate unit. Notice (Section 5.10) that the phosphate esters link at 5', 3', which gives a directionality to the polymer. Thus, by convention the 5' end of the chain is written to the left in our abbreviation and the 3' end to the right. So the segment of DNA chain shown on page 712 can be abbreviated:

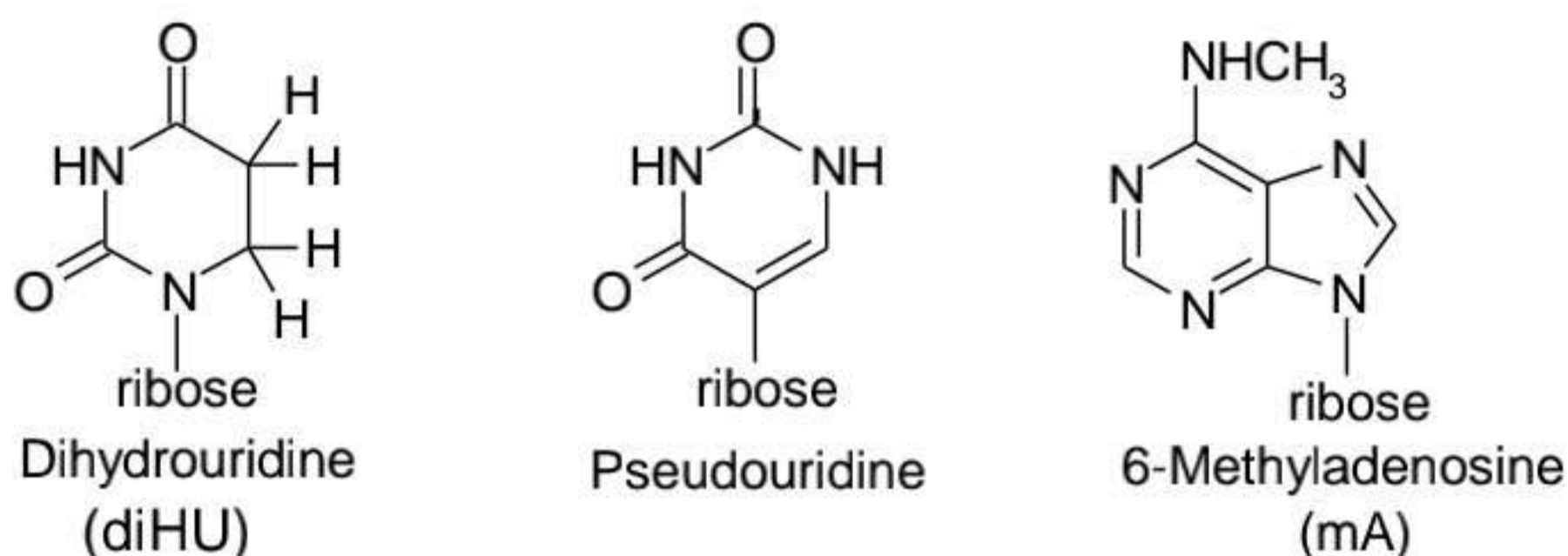


There are three major classifications of RNA; they are designated according to their functional properties. Messenger RNAs (mRNA) carry the genetic information from DNA for subsequent translation into specific protein sequences. Transfer RNAs (tRNA), previously called soluble RNAs, have molecular weights of 25,000-30,000, corresponding to 75-90 nucleotides. Ribosomal RNAs (rRNA) are much larger, having z molecular weights of 0.5-1 million. The two latter types of RNA will be discussed below.

The mechanisms by which the ribonucleic acids perform their varied and complicated functions are subject to intensive study among biochemists today. One key theory has emerged (mentioned in Section 5.10), which has defied attempts at disproof. It is the very specific hydrogen bonding that exists between AU base pairs and between GC base pairs, which is responsible for determining the eventual amino acid sequence in proteins. The diagram presented on page 12 actually depicts an AT pair, but inspection of the structures of T and U will show that both have the same hydrogen-bonding properties. No such analogous complementary hydrogen bonding can exist between AC, AG, UG, or LC pairs. Thus, a trinucleotide such as CpUpC will be strongly attracted to its complementary trinucleotide GpApG, less strongly to one such as CpApC, and virtually not at all to one such as ApCpU. This binding specificity controls the process of protein synthesis in the living cell.

How does this synthesis take place? First, the free amino acids which have been synthesized by the body or derived from ingested nutrients become esterified to the 3'-OH end of a specific tRNA. This molecule is currently the subject of much attention and a short look at its

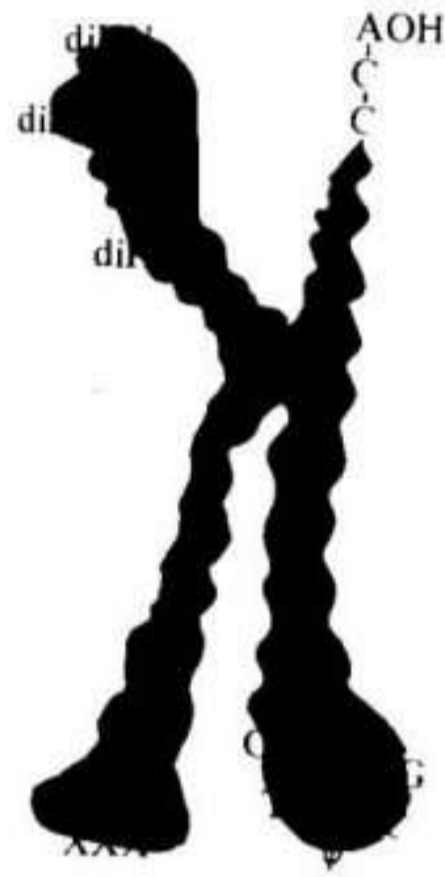
structure as it is understood today will be profitable. The exact nucleotide sequence of a number of tRNAs has now been established, and preliminary structural characteristics have been described on the basis of X-ray data. As stated above, all tRNA molecules are approximately the same size and share other common features. Invariably, they terminate on the 3' end with the sequence -pCpCpA-OH. Also, they all undergo a certain amount of chemical modification after their initial synthesis, leading to a number of modified bases. such as



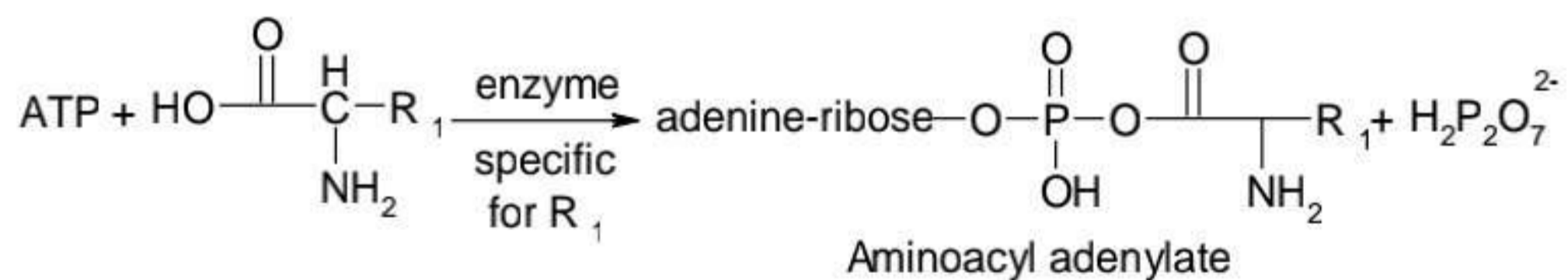
These unusual bases are present in every tRNA in the form of (1) a looped-out region with a sequence locally rich in diHU, and (2) another loop sequence, which is invariably —pCipTpΨpCpGp—. With all these similarities, where, then, are tRNAs different, and how does each amino acid invariably become attached to only one species of tRNA? The one answer with which all can agree involves a region very near the center of the chain and consists of a triplet of nucleotides whose sequence is different for each of the amino acid-specific tRNAs studied thus far. This trinucleotide sequence is called the anticodon region, for reasons soon to become apparent. There are also localized heterogeneities in the diHU loop from one species of tRNA to another.

A number of three-dimensional models of tRNAs have now been proposed. A tRNA is schematically represented in Figure 5.5, where XXX represents the anticodon, hydrogen bonding is indicated by dotted lines, and p's have been omitted for the sake of brevity.

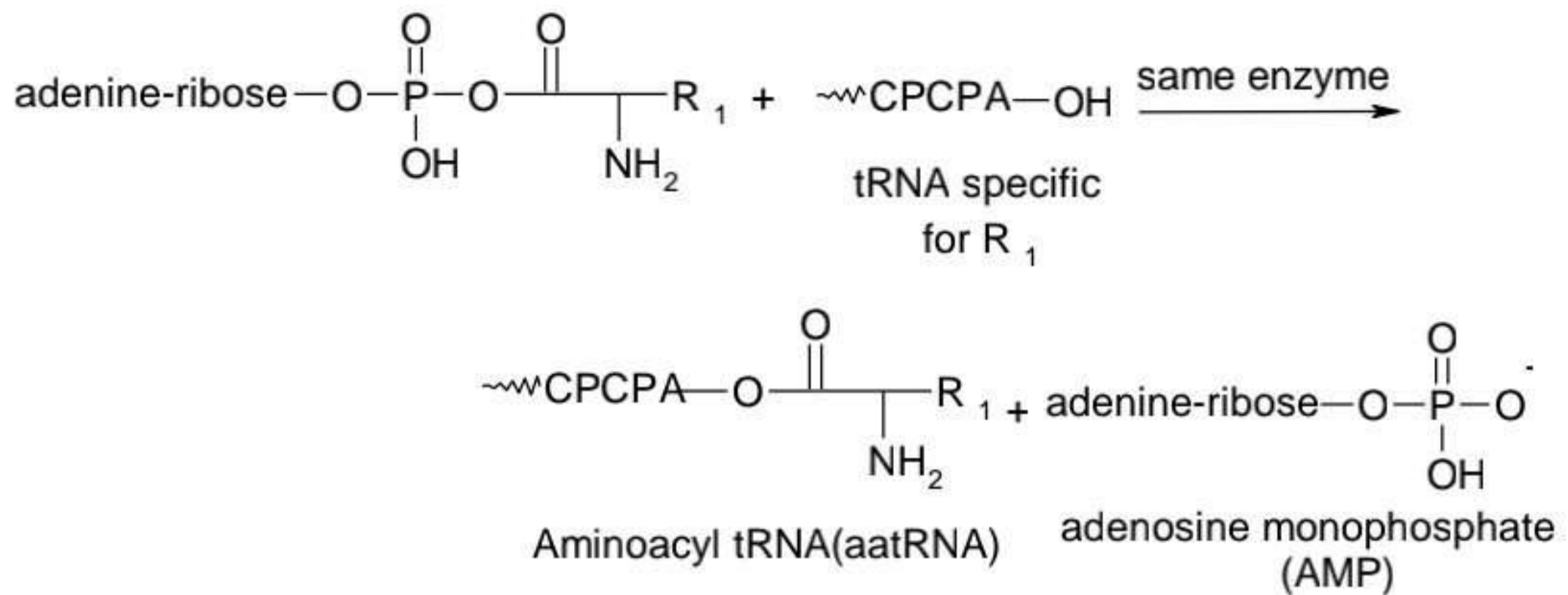
Figure 5.5 Schematic diagram of transfer ribonucleic acid.



The enzyme that joins the amino acid to the tRNA catalyzes first the "activation" of the amino acid with the high-energy biochemical adenosine triphosphate (ATP) (Section 5.10).



The second step involves the actual esterification:



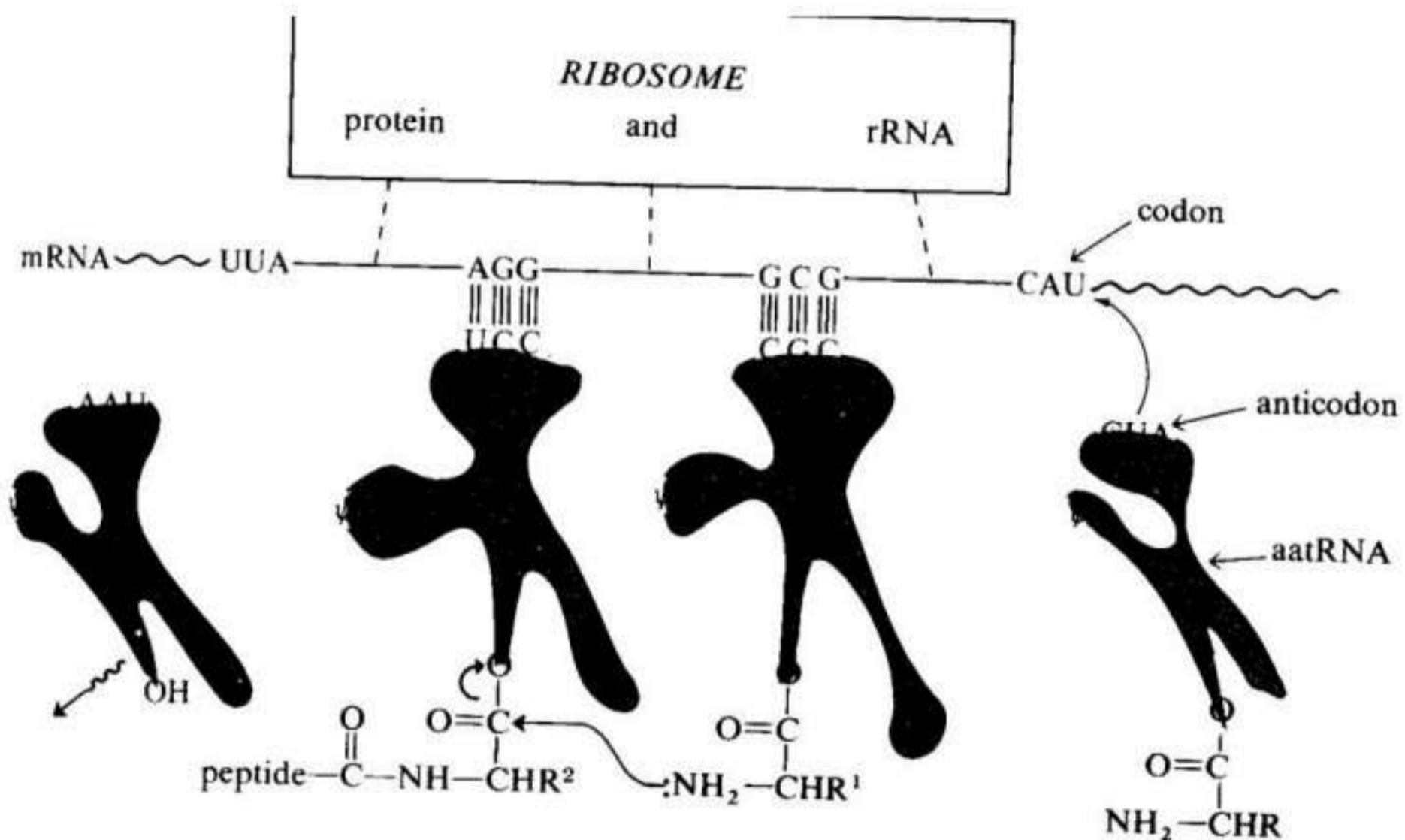
This reaction is carried out with each amino acid, each time utilizing an amino acid specific enzyme and tRNA, until all amino acyl tRNA esters have been produced, each with a specific anticodon. It is in this form that the amino acids are ready to be polymerized into polypeptide linkages.

The actual formation of the peptide bond occurs on a nucleoprotein particle called the ribosome. Every living cell capable of synthesizing protein contains many ribosomes, and even though there are minor differences depending on the source, they are always composed of one small and two large strands of RNA (rRNA), with which are associated some 60 or 80 smaller proteins. The functional nucleoprotein particle is roughly pear-shaped with a diameter of about 200 Å. It is with the ribosome that the mRNA interacts prior to protein synthesis. After formation of the mRNA-ribosome complex, different amino acyl tRNA (aatRNA) molecules in the surrounding medium come into contact with it. If the exposed anticodon loop is exactly complementary to the specific triplet (the codon) on the ribosome-stabilized mRNA, it will form a

hydrogen-bonded triplet of base pairs. Although the exact sequence of events that follows is not known with certainty, strong evidence indicates that the  $\Psi$ -containing loop of the tRNA becomes attached to the ribosome and further stabilizes the ternary aminoacyl-tRNA-mRNA-ribosome complex. If only two or three nucleotides are capable of pairing, the association will not be strong enough for the 41 loop to "lock" the complex, and the "wrong" aminoacyl-tRNA will diffuse away and be replaced by another until a correct match is made. After this happens, the following codon on the mRNA (the next three nucleotides) is in a position to react with its complementary anticodon on another tRNA. When two such alignments have been made, the carboxyl group of the first amino acid participates in amide bond formation with the free amino group of the recently incoming aminoacyl tRNA. This enzyme-catalyzed reaction, of course, frees the first amino acid from its tRNA, which by unknown means senses this change and diffuses away from the complex, leaving behind a dipeptidyl-tRNA-mRNA-ribosome complex. The ribosome then moves "down" the mRNA just enough to bring the next (third) codon into a position in which it can react with its own anticodon on yet another aminoacyl tRNA. This sequence of reactions is repeated until some chain-termination signal on the mRNA causes synthesis to stop and the finished protein to be released for use as a hormone, enzyme, or structural protein. An intermediate stage in the process is depicted schematically in Figure 5.6.



Figure 5.6 Schematic diagram of polypeptide biosynthesis. The mRNA has become associated with the large ribosome. The codons of the mRNA are shown interacting with the complementary anticodons of the aatRNAs.



Using known values of hydrogen-bond energies, theorists have calculated that if the genetic code were a doublet or quadruplet one, life as we know it would not be possible. The attractive force between two hydrogen-bonded base pairs is inadequate to keep the large tRNA molecule in place on the ribosome sufficiently long for peptide bond formation to occur. Conversely, four base pairs would be so strongly bonded that proteins might require months or more to be synthesized. The actual measured rate of protein synthesis in the living cell is about two amino acids per second.

The next several years should see a tremendous increase in the knowledge not only of the details of this complex process but also of such intriguing topics as the transcriptional and translational control of protein

synthesis. the design of specific drugs to control the growth of harmful bacteria and viruses, and control of the rejection problem in tissue and organ transplants.

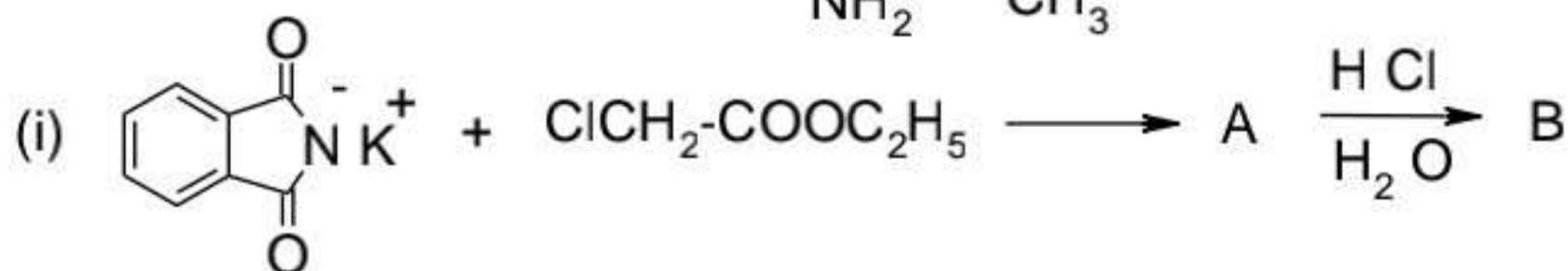
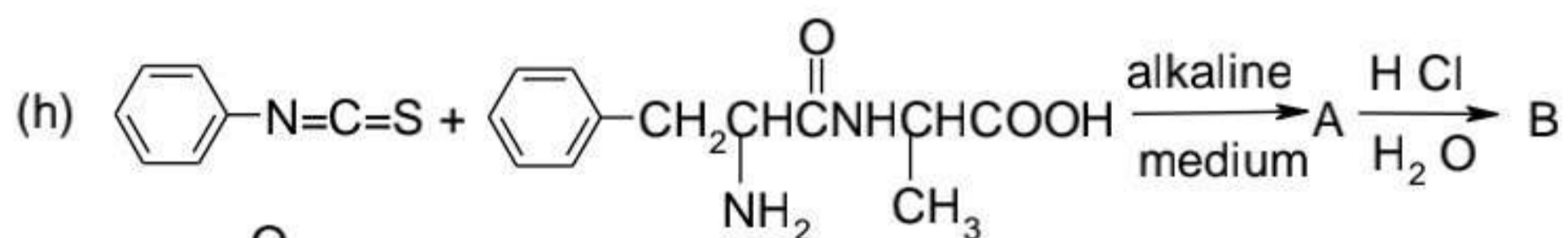
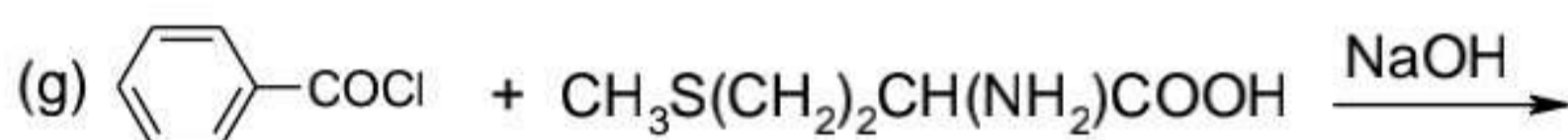
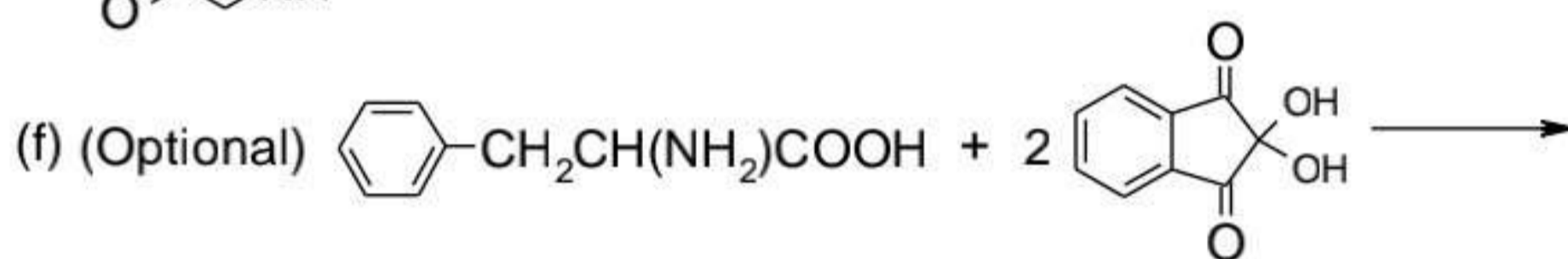
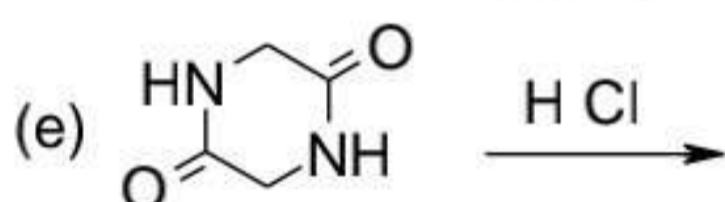
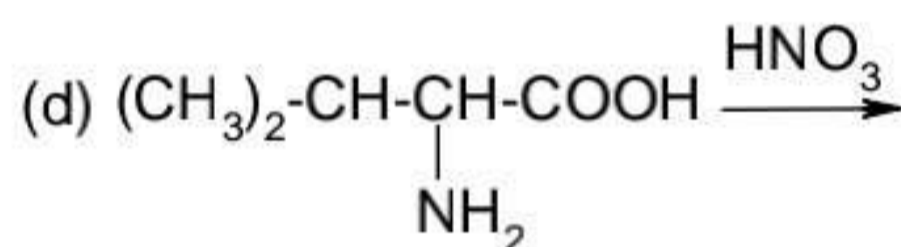
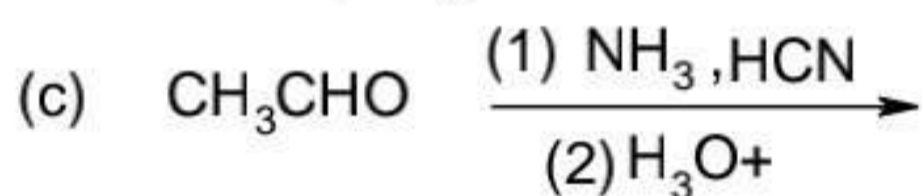
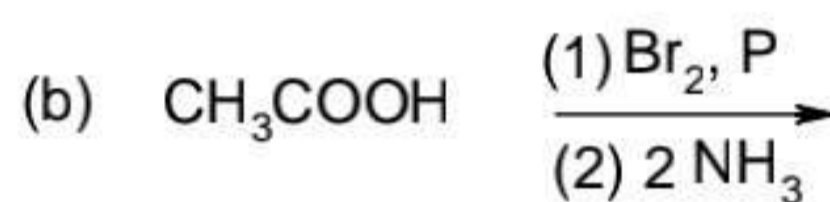
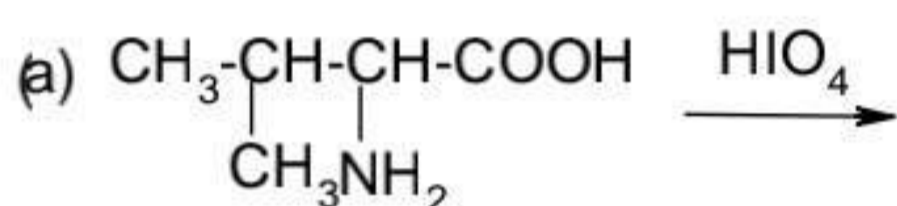
### **PROBLEMS**

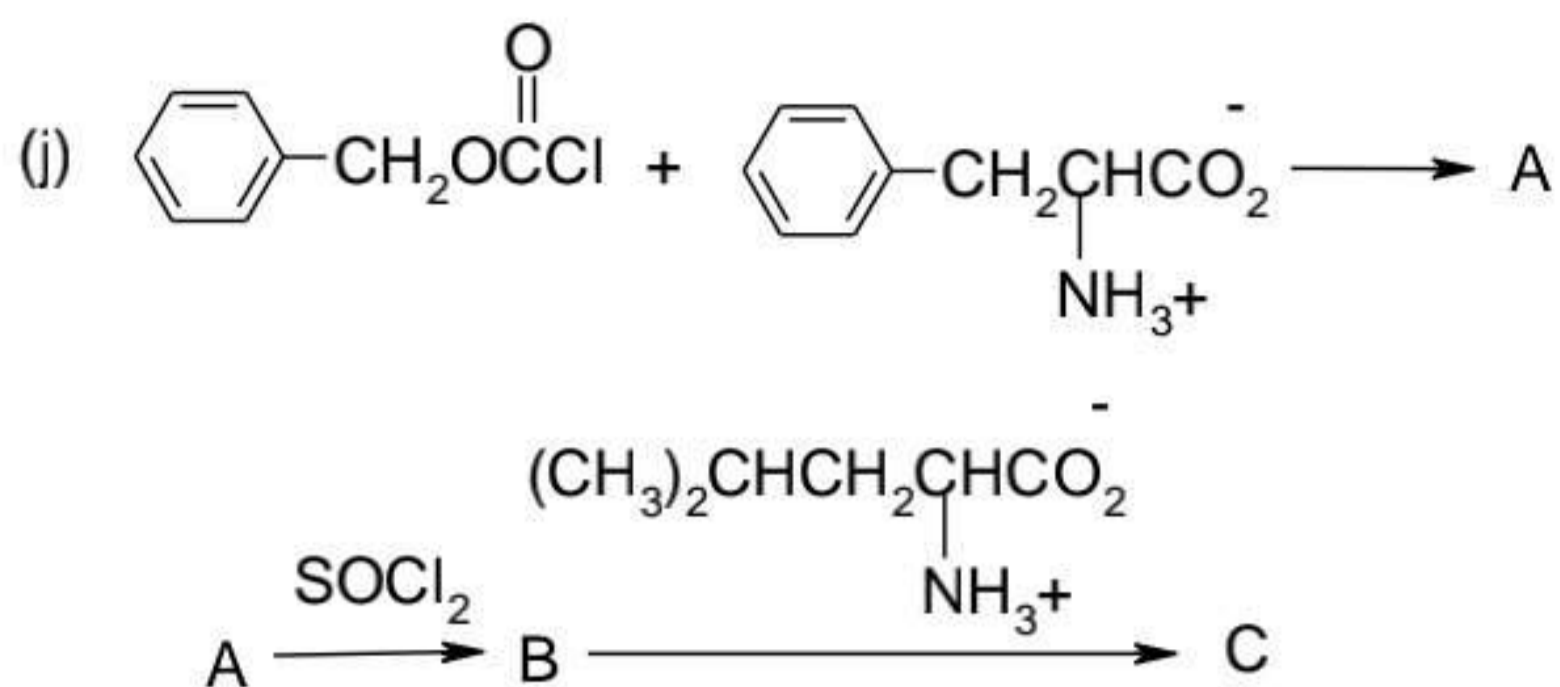
- 1- Name the three types of natural organic polymers that are essential to life processes.
- 2- Give a definition and example of:
  - (a) An essential amino acid
  - (b) The peptide bond
  - (c) Polypeptide
  - (d) Oligopeptide
  - (e) Tripeptide
  - (f) Enzyme
  - (g) Structural protein
- 3- What is meant by the primary, secondary, tertiary, and quaternary structure of polypeptides?
- 4- Define the isoelectric point of an amino acid and explain why arginine has an isoelectric point 5 pH units higher than does glycine, and glutamic acid has one about 3 pH units lower than that of glycine.
- 5- Write the structures that correspond to the following formulas:
  - (a) Val. Trp. Lys. (NH<sub>2</sub>).
  - (b) (Ac) HiS. Gly. CySH. Gln.
- 6- Describe the molecular disease of sickle-cell anemia. Explain why a human being cannot use a blood transfusion from a monkey. but can use insulin from a cow.

- 8- Explain the difference in the chances of the success of a heart transplant between identical twins and a heart transplant between nonidentical twins.
- 9- Describe the molecular basis of immunity.
- 10- Glutathione is a tripeptide that is an important regulator of the oxidation-reduction reactions of cells in animals. From the following experimental results, suggest a structure for glutathione:
- (a) Enzymatic or acid hydrolysis gives glycine, cysteine, and glutamic acid in equimolar amounts.
  - (b) Mild hydrolysis gives two dipeptides: one on further hydrolysis gives cysteine and glutamic acid, and the other gives cysteine and glycine.
  - (c) Carboxypeptidase liberates glycine.
  - (d) 2,4-Dinitrophenylation gives N-(2,4-dinitrophenyl) glutamine.
- 11- Describe by formula two N-terminal amino acid determinations.
- 12- Compare the merits of the chemical C-terminal amino acid determination with the enzymatic determination for the same purpose.
- 13- An octapeptide was found to contain the following amino acids: Ala, Ala, His, Leu, Lys, Pro, Thr, Tyr. Upon partial hydrolysis there were isolated from the resulting mixture four tri-peptides which had the following structures: Leu, Ala, Tyr, Thr, Pro, Leu, Lys, His, and His, Thr, Pro. A C-terminal amino acid determination showed that tyrosine occupied that position. What is the structure of the octapeptide?

14- Another octapeptide upon hydrolysis gave the following amino acids: Met. Asn, CySH. Lys. Pro. Thr, Thr, and Val. Partial hydrolysis gave a mixture from which it was possible to isolate four dipeptides and two tripeptides. which had the following structures: Met, Lys. Val. Thr. Asn. Val. Thr. Met. Pro. Thr. Asn. and Lys. CySH. Pro. What is the structure of the octa-peptide?

15 - Complete:





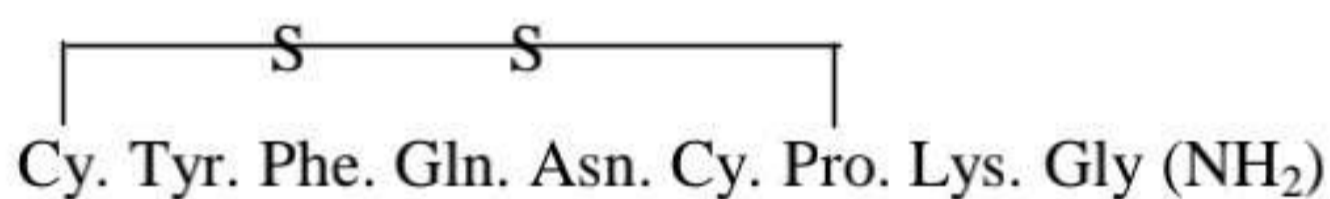
16- Suggest a synthesis for

- Ala. Pro. Val.
- Asp. Ile. Gly.
- Thr. Arg. Lys.

17- Outline a synthesis for lysine from the oxime of cyclohexanone.

18- Discuss the stereochemical implications of obtaining L-Val. L-Ile. L-Lys. by resolution of DL-Val. DL-Ile. DL-LyS.

19- Vasopressin is a posterior pituitary hormone that acts on the kidney to reduce excretion of water and brings about a rise in blood pressure:



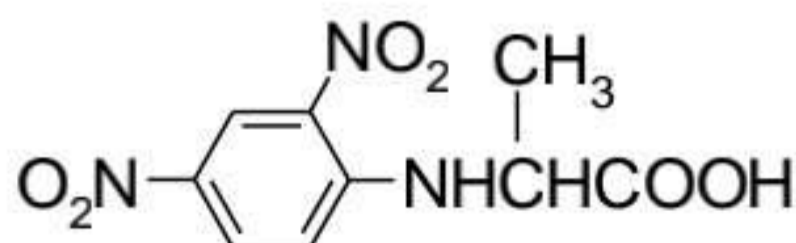
Lysine vasopressin

- Devise a scheme for determining the peptide sequences.
- Devise a synthesis of the following part of the chain: CysH. Pro. Lys. Gly(NH<sub>2</sub>).

20- Cells contain two types of nucleic acids. commonly referred to as RNA and DNA.

- What do the abbreviations RNA and DNA stand for?
- What are the similarities between RNA and DNA?

- (c) What are the differences between RNA and DNA?
- (d) In nucleic acid chemistry, the abbreviations A, C, G, T, and U are commonly employed. Give the structures and names for the compounds represented by these letters.
- (e) What is the chemical basis of the Watson and Crick double-helix structure for DNA?
- 21- Show the mechanism of peptide coupling by dicyclohexylcarbodiimide.
- 22- Describe the role of tRNA in protein biosynthesis.
- 23- Describe the function of the anticodon.
- 24- What is the maximum number of different amino acids possible in human protein?
- 25- A certain tetrapeptide is found to yield on hydrolysis 2 mol of alanine, 1 mol of glycine, and 1 mol of valine. Write the formula for each possible structurally isomeric tetrapeptide that could give this result.
- 26- The careful fractionation of the nonlipid portion of the spleen of a red snurd yielded a pure heptapeptide. Complete hydrolysis of this polypeptide yielded alanine (2 mol), cystine (1 mol), glutamic acid (2 mol), and glycine (2 mol). When the polypeptide was allowed to react with 1-fluoro-2,4-dinitrobenzene and the product was hydrolyzed, the following compound was isolated:



The partial hydrolysis of the polypeptide yielded a mixture from which these three tripeptides could be isolated: CySH. Glu. Glu, Gly. Gly. CySH, and Glu. Ala. What is the structural formula of the heptapeptide?

- 27- Use the Merrifield solid-phase method to outline a synthesis of the tetrapeptide: Phe. Ala. Gly. Gly.

## Enzymes

The word enzyme means “in yeast.” Even without any knowledge of their structures or functions, humans have used enzymes since prehistoric times in the production of wine, vinegar, and cheese. Pasteur thought that living yeast cells were necessary for fermentation processes. We now know that a living cell is not necessary; the proper enzymes, plus reaction conditions that do not cause denaturation, are all that are needed for enzymatic reactions.

An enzyme is a biological catalyst. A higher animal contains thousands of enzymes. Virtually every biochemical reaction is catalyzed by an enzyme. Even the equilibrium  $\text{CO}_2 + \text{H}_2\text{O} \rightleftharpoons \text{H}_2\text{CO}_3$  is enzyme-catalyzed because the rate of the uncatalyzed equilibration does not produce carbonic acid fast enough for an animal's needs.

Enzymes are more efficient catalysts than most laboratory or industrial catalysts (such as Pd in a hydrogenation reaction). Biological reactions in humans occur at  $37^\circ\text{C}$  and in aqueous media. High temperature, high pressure, or very reactive reagents (such as NaOH or  $\text{LiAlH}_4$ ) are not available to an organism. Enzymes also allow a selectivity of reactants and a control over reaction rate that can be obtained with no other class of catalyst.

All enzymes are proteins. Some are relatively simple in structure; however, most are complex. The structures of many enzymes are still unknown. For biological activity, some enzymes require prosthetic groups, or cofactors. These cofactors are nonprotein portions of the enzyme. A cofactor may be a simple metal ion; for example, copper ion is the cofactor for the enzyme ascorbic acid oxidase. Other enzymes contain



nonprotein organic molecules as cofactors. An organic prosthetic group is frequently referred to as a coenzyme.

If an organism cannot synthesize a necessary cofactor, the cofactor must be present in small amounts in the diet. The active units of many cofactors are vitamins. Table 6.5 shows a few cofactors and the corresponding vitamins.

**A. Naming enzymes:**

Most enzymes are named after the reactions that they catalyze. The ending for an enzyme name is usually -ase. The name may be general and refer to a class of enzymes that catalyze a general type of reaction. For example, a polymerase is any enzyme that catalyzes a polymerization reaction, and a reductase is any enzyme that catalyzes a reduction reaction. An enzyme name may also refer to a specific enzyme: ascorbic acid oxidase is the enzyme that catalyzes the oxidation of ascorbic acid, while phosphoglucose isomerase catalyzes the isomerization of glucose 6-phosphate to fructose 6-phosphate.

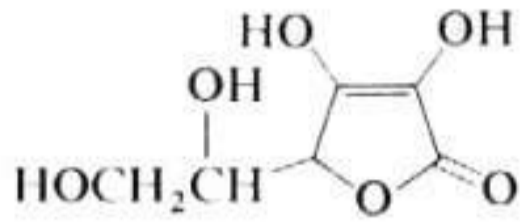
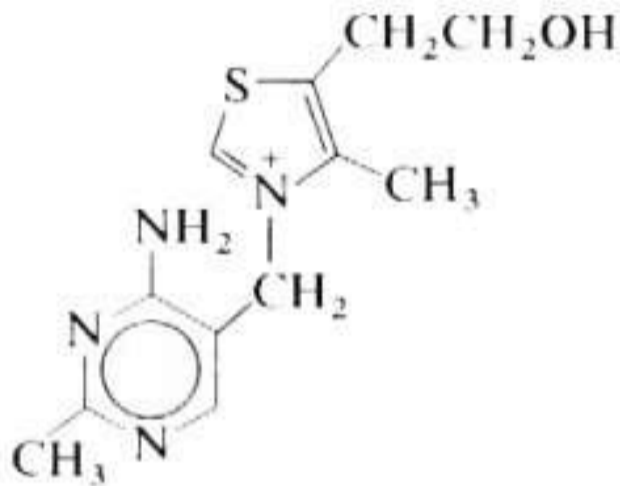
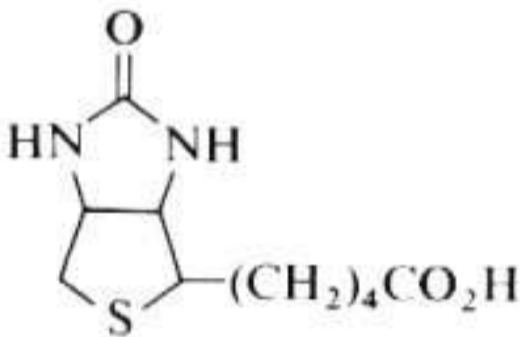
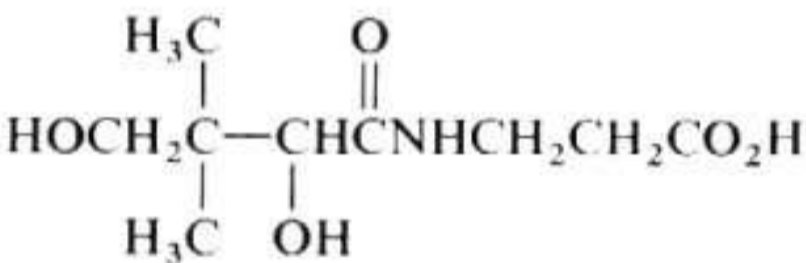
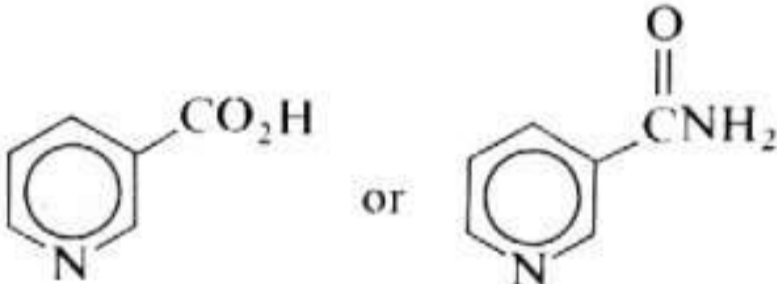
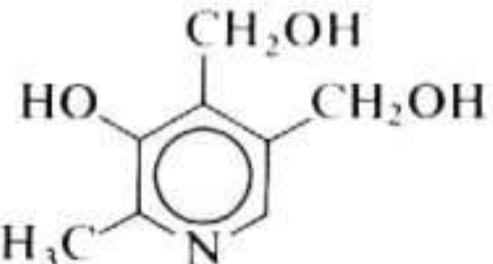
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**STUDY PROBLEM:**

6.9. Suggest the function of each of the following enzymes:

- (a) an acetyltransferase;
  - (b) phenylalanine hydroxylase;
  - (c) pyruvate dehydrogenase.
-

**Table 6.5. Some cofactors that contain vitamins**

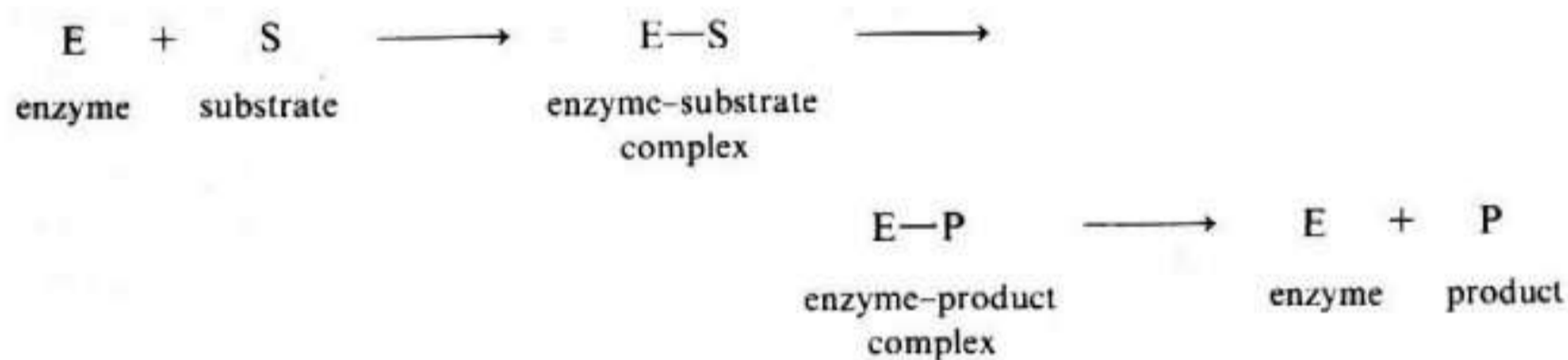
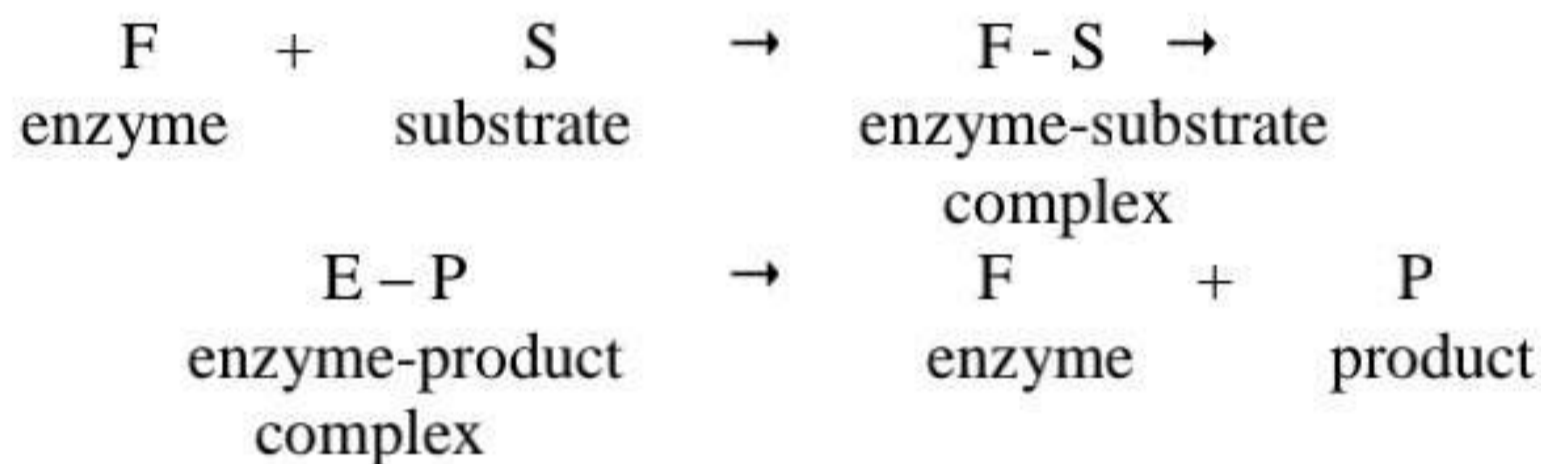
Name of cofactor	Vitamin needed	Structure of vitamin
Vitamin C (ascorbic acid)	Vitamin C	
Vitamin B <sub>1</sub> (Thiamine)	Vitamin B <sub>1</sub>	
biotin	Biotin	
Coenzyme A	Pabtothenic acid	
NAD <sup>+</sup> a	Nicotinic acid (niacin) or nicotinamide (niacinamide)	
Pyridoxyl phosphate	Pyridoxyl	

Nicotinamide adenine dinucleotide, a biological oxidizing agent.

**B. How enzymes work:**

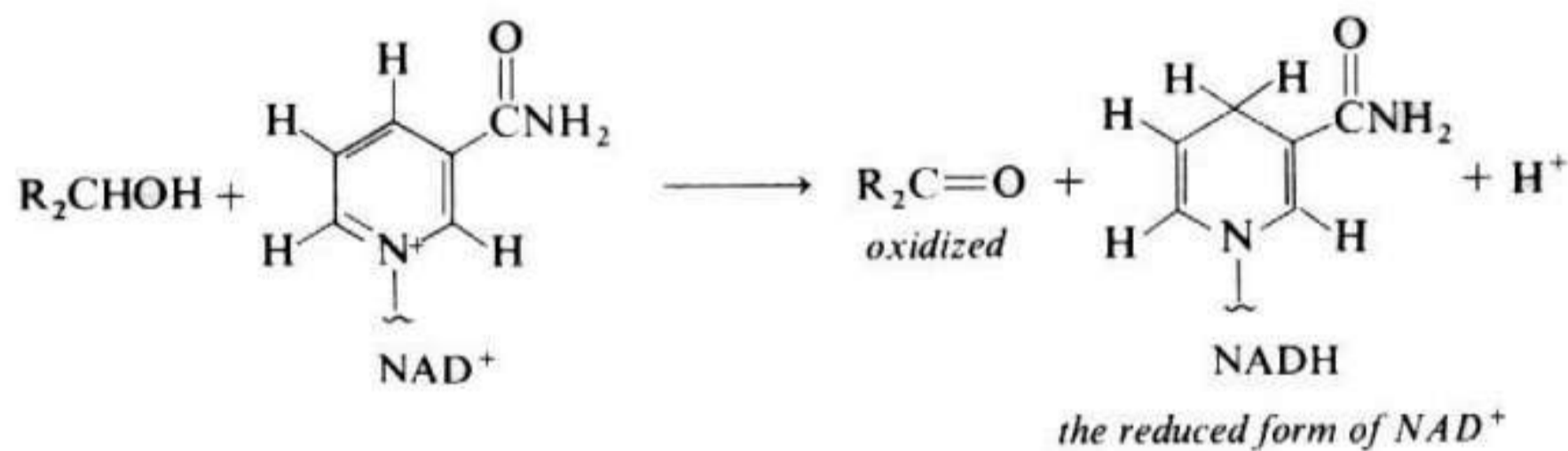
Some enzymes have been studied in detail, yet there is still much to learn about even the well-known enzymes. It is believed that an enzyme fits itself around the substrate (the molecule to be acted upon) to form an enzyme-substrate complex. The bonds of the substrate may be strained by attractions between itself and the enzyme. Strained bonds are of higher energy and are more easily broken; therefore, the desired reaction proceeds easily and yields an enzyme-product complex.

In many cases, the product is not the same shape as the reacting substrate; thus the fit between the product and the enzyme is no longer perfect. The altered shape of the product causes a dissociation of the complex, and the enzyme surface is ready to accept another molecule of substrate. This theory of enzyme activity is called the induced-fit theory.

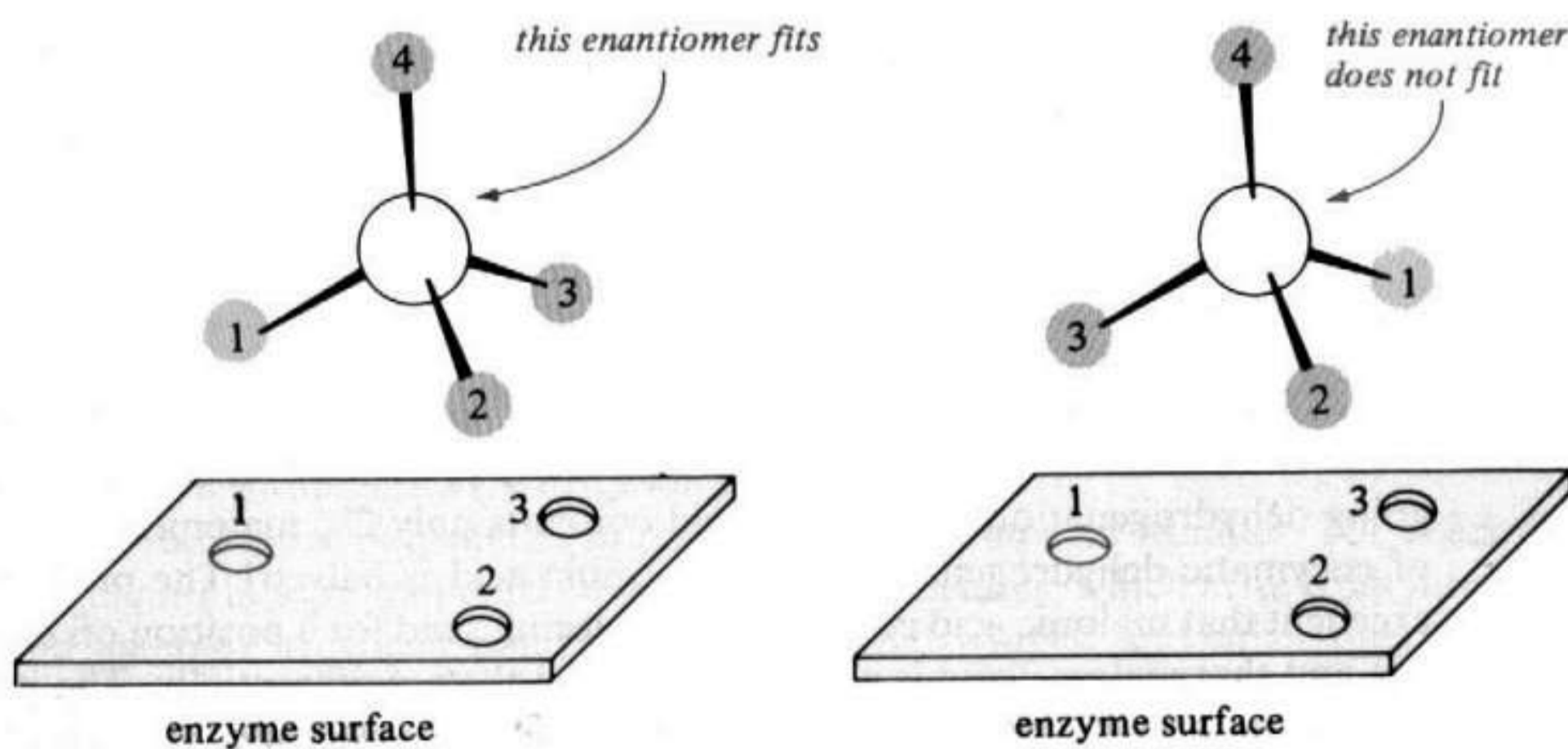


Enzymes have molecular weights of 12,000-120,000 and higher. Most substrates (for example, an amino acid or a unit of glucose) are much smaller molecules. The specific location of the large enzyme structure where reaction occurs is called the active site. This site is where

the prosthetic group (if any) is located. Metallic prosthetic groups are thought to serve as electrophilic agents and, in this way, catalyze the desired reactions. In  $\text{NAD}^+$ , the active site is the nicotinamide end of the cofactor.  $\text{NAD}^+$  is readily reduced and therefore catalyzes oxidation reactions.



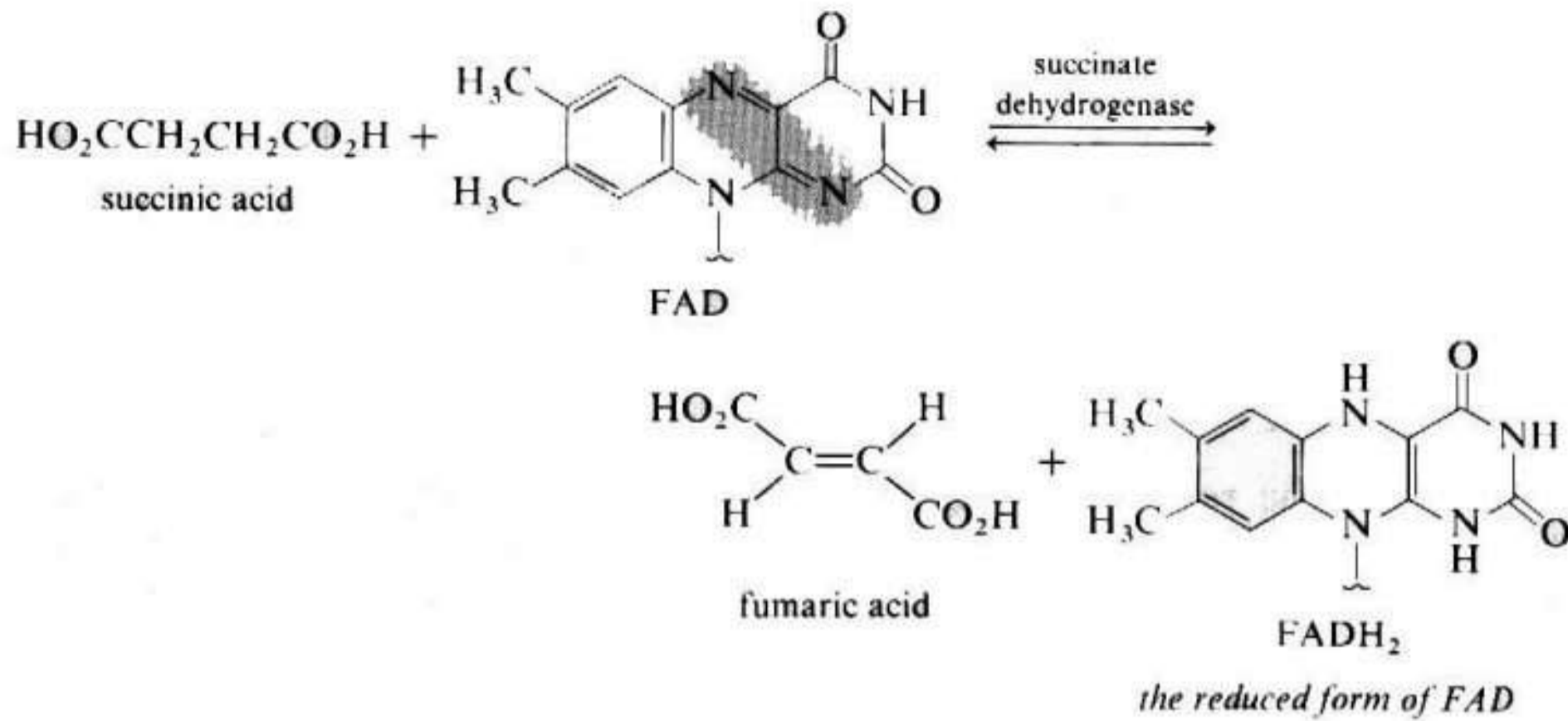
The rest of the enzyme molecule is not simply excess molecular weight! It is believed that this portion of the enzyme recognizes its substrate and holds it in place. It was suggested in the 1890's by Emil Fischer that enzymes are chiral molecules and that reactants must complement this chirality in order to undergo reaction. Fischer compared the fitting together of the substrate structure and the enzyme structure to a key fitting into a lock (see Figure 6.6).



**FIGURE 6.6. One enantiomer fits on the enzyme surface; its mirror image does not.**

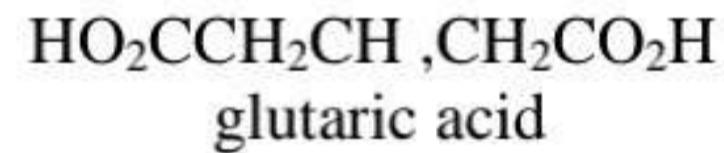
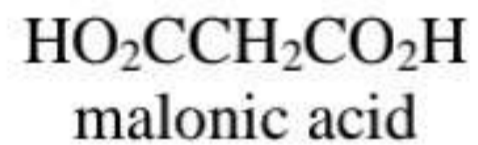
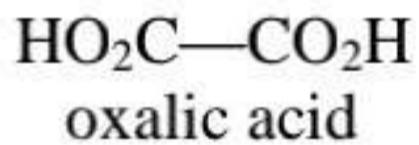
Recognition may occur by a series of dipole-dipole interactions, by hydrogen bonding, or by covalent bonding, in which the stereochemistry must be just right. In some cases, the rest of the enzyme molecule is folded to form a hydrophobic pocket that holds a nonpolar portion of the substrate in place. (We mentioned this type of structure for hemoglobin.) If the nonpolar end of a potential substrate does not fit in the pocket correctly, enzyme catalysis diminishes or is nonexistent. Therefore, the functional group to be acted upon must fit the active site on the enzyme, and the rest of the substrate molecule must fit together with other portions of the enzyme molecule for reaction to proceed. This dual type of recognition is the basis of the unique specificity of most enzymes.

Both the active site and the rest of an enzyme are important in enzyme activity. Let us look at one reaction in which the active site seems to be the more important factor in substrate recognition. The enzyme succinate dehydrogenase catalyzes the dehydrogenation of succinic acid to the trans-diacid fumaric acid. (The cis-isomer, maleic acid, is not produced in this reaction.) The oxidizing agent in this reaction is flavin adenine dinucleotide (FAD), which is reduced by a 1,4-addition of two hydrogen atoms (plus two electrons). (We show only the functional portion of FAD here.)



Other diacids, such as oxalic acid, malonic acid, and glutaric acid, inhibit the dehydrogenation of succinic acid.

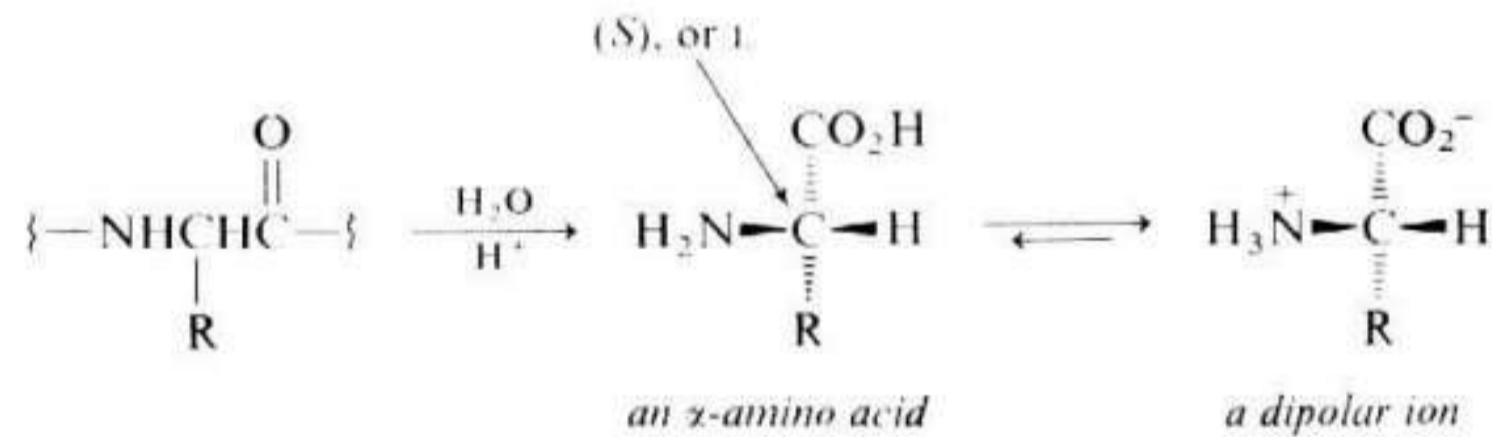
### Inhibitors of succinate dehydrogenase:



Of these diacids, malonic acid has the greatest inhibiting effect. Malonic acid is very similar in structure to succinic acid, but is structurally incapable of undergoing dehydrogenation. If succinic acid contains only 200 malonic acid, the rate of enzymatic dehydrogenation of the succinic acid is halved! The probability is excellent that malonic acid competes with succinic acid for a position on the active site and that malonic acid is attracted and held there preferentially. The presence of malonic acid on the active site thus blocks the approach of succinic acid.

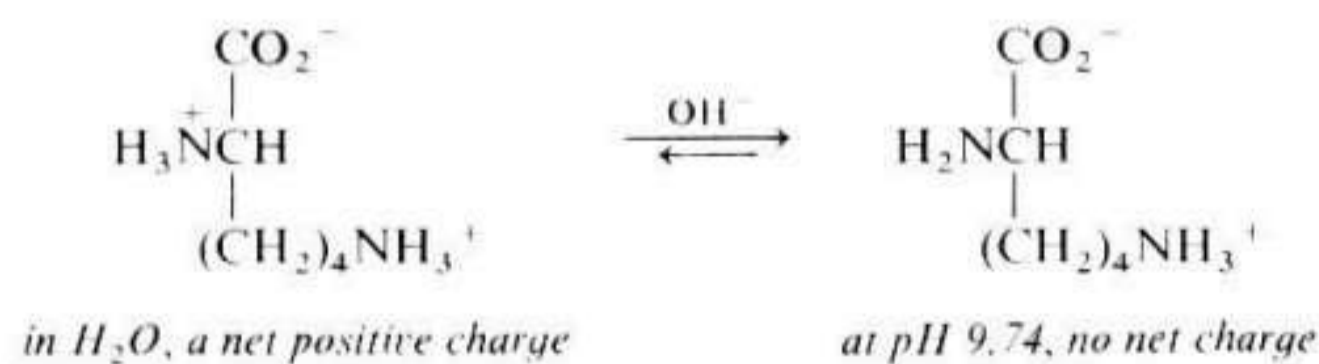
**Summary:**

A protein is a polyamide. Hydrolysis yields  $\alpha$ -amino acids of (S)-configuration at the  $\alpha$  carbon. Amino acids undergo an internal acid-base reaction to yield dipolar ions.



Essential amino acids are those that cannot be synthesized by an organism and must be present in the diet. Acidic amino acids are those with a carboxyl group in the side chain (R in the preceding equation). Basic amino acids contain an amino group in the side chain. Neutral amino acids contain neither  $\text{—CO}_2\text{H}$  nor  $\text{—NH}_2$  in the side chain, but may contain OH, SH, or other polar group. Cross-linking in proteins may be provided by the SH group in cysteine, which can link with another SH in an oxidation reaction:  $2 \text{RSH} \rightarrow \text{RSSR} \pm 2 \text{H}$ .

The isoelectric point of an amino acid is the pH at which the dipolar ion is electrically neutral and does not migrate toward an anode or cathode. The iso-electric point depends on the acidity or basicity of the side chain.



Racemic amino acids may be synthesized by a variety of routes.

A peptide is a polyamide of fewer than 50 amino acid residues. The N-terminal amino acid is the amino acid with a free  $\alpha$ -amino group, while the C-terminal amino acid has a free carboxyl group at carbon 1. End-group analysis to determine the C- and N-terminals and partial hydrolysis to smaller peptides are two techniques for peptide structure-determination.

In the synthesis of a peptide, reactive groups (except for the groups desired to undergo reaction) must be blocked. A carbamate group may be used to protect an amino group. A solid-phase peptide synthesis provides a blocking group for the C-terminal carboxyl group. The biosynthesis of proteins is accomplished by RNA. The order of incorporation of amino acids is determined by the order of attachment of the bases (N-heterocycles) in mRNA.

Proteins are polyamides of more than 50 amino acid residues. The order of side chains in a protein determines its higher structures, which arise from internal and external hydrogen bonding, van der Waals forces, and other interactions between side chains. The higher structures of proteins give them a variety of physical and chemical properties so that they may perform a variety of functions.

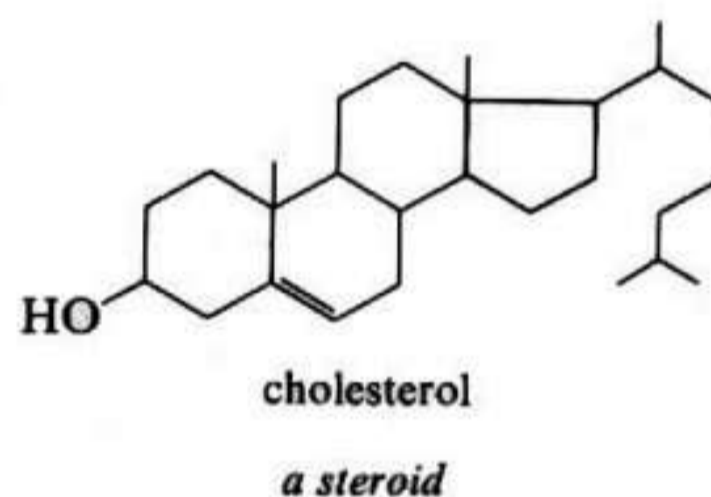
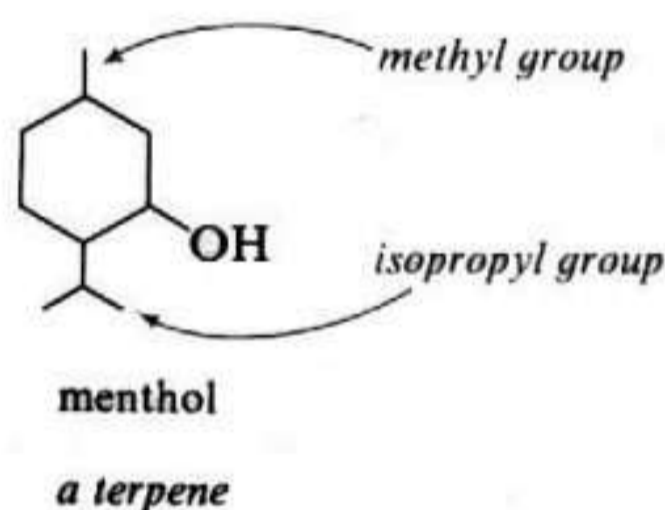
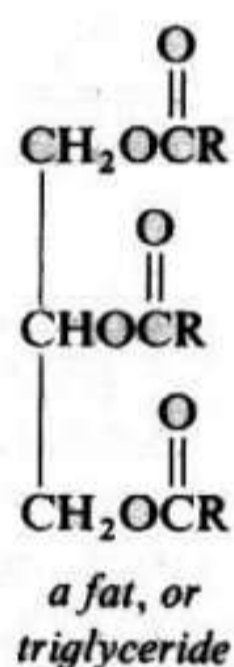
Denaturation is the disruption of hydrogen bonds and thus the disruption of the higher structure of the protein.

Enzymes are proteins that catalyze biochemical reactions. Enzymes are efficient and specific in their catalytic action. The specificity is provided for by the unique shape and by the polar (or nonpolar) groups contained within the enzyme structure. Some enzymes work in conjunction with a nonprotein cofactor, which may be organic or inorganic.



## Lipids and Related Natural Products

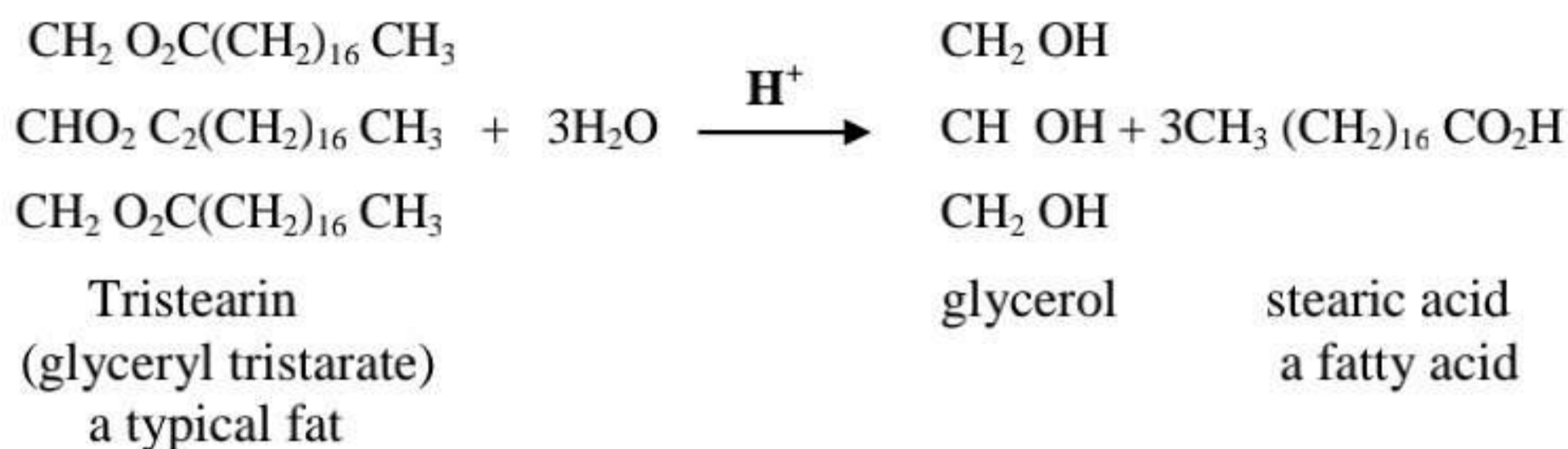
**A lipid:** is defined as a naturally occurring organic compound that is insoluble in water, but soluble in nonpolar organic solvents such as a hydrocarbon or diethyl ether. This definition sounds as if it might include many types of compounds, and indeed it does. The various classes of lipids are related to one another by this shared physical property; but their chemical, functional, and structural relationships, as well as their biological functions, are diverse. We will discuss here the classes usually thought of as lipids: fats and oils, terpenes, steroids, and a few other compounds of interest. (Line formulas are generally used for terpenes and steroids, as the following examples show.



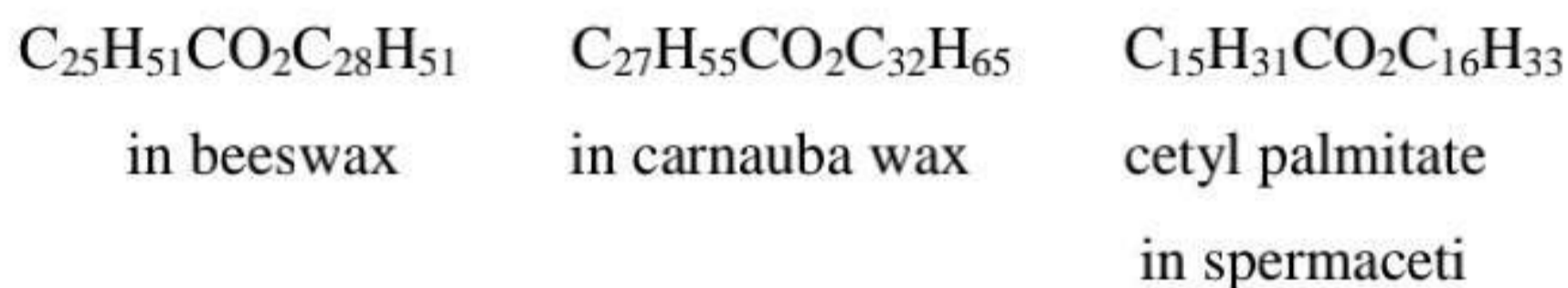
### Fats and Oils:

Fats and oils are triglycerides, or triacylglycerols, both terms meaning “triesters of glycerol.” The distinction between a fat and an oil is arbitrary: at room temperature a fat is solid and an oil is liquid. Most glycerides in animals are fats, while those in plants tend to be oils; hence the terms *animal fats* (bacon fat, beef fat) and *vegetable oils* (corn oil, safflower oil).

The carboxylic acid obtained from the hydrolysis of a fat or oil, called a fatty acid, generally has a long, unbranched hydrocarbon chain. Fats and oils are often named as derivatives of these fatty acids. For example, the tristearate of glycerol is named tristearin, and the tripalmitate of glycerol is named tripalmitin.



Fatty acids can also be obtained from waxes, such as beeswax. In these cases, the fatty acid is esterified with a simple long-chain alcohol.



Most naturally occurring fats and oils are *mixed* triglycerides—that is, the three fatty-acid portions of the glyceride are not the same. Table 7.1 lists some representative fatty acids, and Table 7.2 shows the fatty-acid composition of some plant and animal triglycerides.

Table 7.1. Selected fatty acids and their sources

Name of acid	Structure	Source
Saturated:		
butyric	$\text{CH}_3(\text{CH}_2)_2\text{CO}_2\text{H}$	milk fat
Palmitic	$\text{CH}_3(\text{CH}_2)_{14}\text{CO}_2\text{H}$	animal and plant fat
Stearic	$\text{CH}_3(\text{CH}_2)_{16}\text{CO}_2\text{H}$	animal and plant fat
Unsaturated:		
Palmitoleic	$\text{CH}_3(\text{CH}_2)_5\text{CHCH}(\text{CH}_2)_7\text{CO}_2\text{H}$	animal and plant fat
Oleic	$\text{CH}_3(\text{CH}_2)_7\text{CH}=\text{CH}(\text{CH}_2)_7\text{CO}_2\text{H}$	animal and plant fat
Linoleic	$\text{CH}_3(\text{CH}_2)_4\text{CHCHCH}_2\text{CHCH}(\text{CH}_2)_7\text{CO}_2\text{H}$	plant oils
Linolenic	$\text{CH}_3\text{CH}_2\text{CHCHCH}_2\text{CHCHCH}_2\text{CHCH}(\text{CH}_2)_7\text{CO}_2\text{H}$	Linseed oil

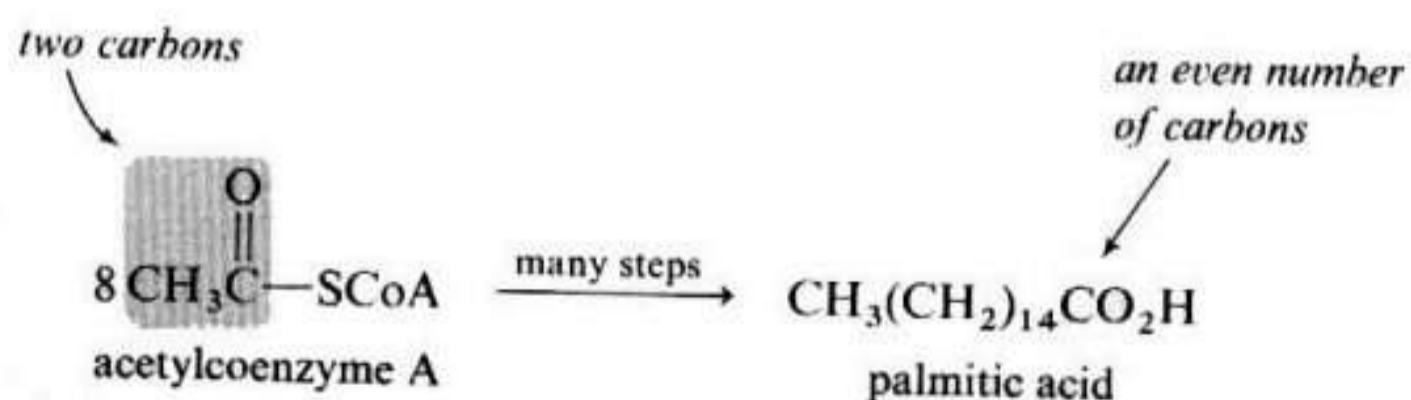
Table 7.2. Approximate fatty-acid composition of some common fats and oils

Composition (%)<sup>a</sup>

Source	Palmitic	Stearic	Oleic	Linoleic
corn oil	10	5	45	38
soybean oil	10	--	25	55
Lard	30	15	45	5
Butter	25	10	35	--
human fat	25	8	46	10

Other fatty acids are also found in lesser amounts.

Almost all naturally occurring fatty acids have an *even* number of carbon atoms because they are biosynthesized from the two-carbon acetyl groups in acetylcoenzyme A.



The hydrocarbon chain in a fatty acid may be saturated or it may contain double bonds. The most widely distributed fatty acid in nature, oleic acid, contains one double bond. Fatty acids with more than one double bond are not uncommon, particularly in vegetable oils; these oils are the so-called *polyunsaturates*.

The configuration around any double bond in a naturally occurring fatty acid is *cis*, a configuration that results in the low melting points of oils. A saturated fatty acid forms zigzag chains that can fit compactly together, resulting in high van der Waals attractions; therefore, saturated fats are solids. If a few *cis* double bonds are present in the chains, the molecules cannot form neat, compact lattices, but tend to coil; polyunsaturated triglycerides tend to be oils. Figure 7.1 shows models of the two types of chains.

Triglycerides are one of the three principal foodstuffs, carbohydrates and proteins being the other two. As an energy source, triglycerides are the most efficient: they provide 9.5 kcal/gram, while the proteins provide 4.4 kcal/gram and the carbohydrates provide 4.2 kcal/gram.

In an organism, ingested fats are hydrolyzed into monoglycerides, di-glycerides, fatty acids, and glycerol, all of which can be absorbed through the intestinal wall. The organism (1) uses these hydrolyzed or partially hydrolyzed fats as raw materials to synthesize its own fats; (2) converts the fatty acids to other compounds such as carbohydrates or cholesterol; or (3) converts the fatty acids to energy.

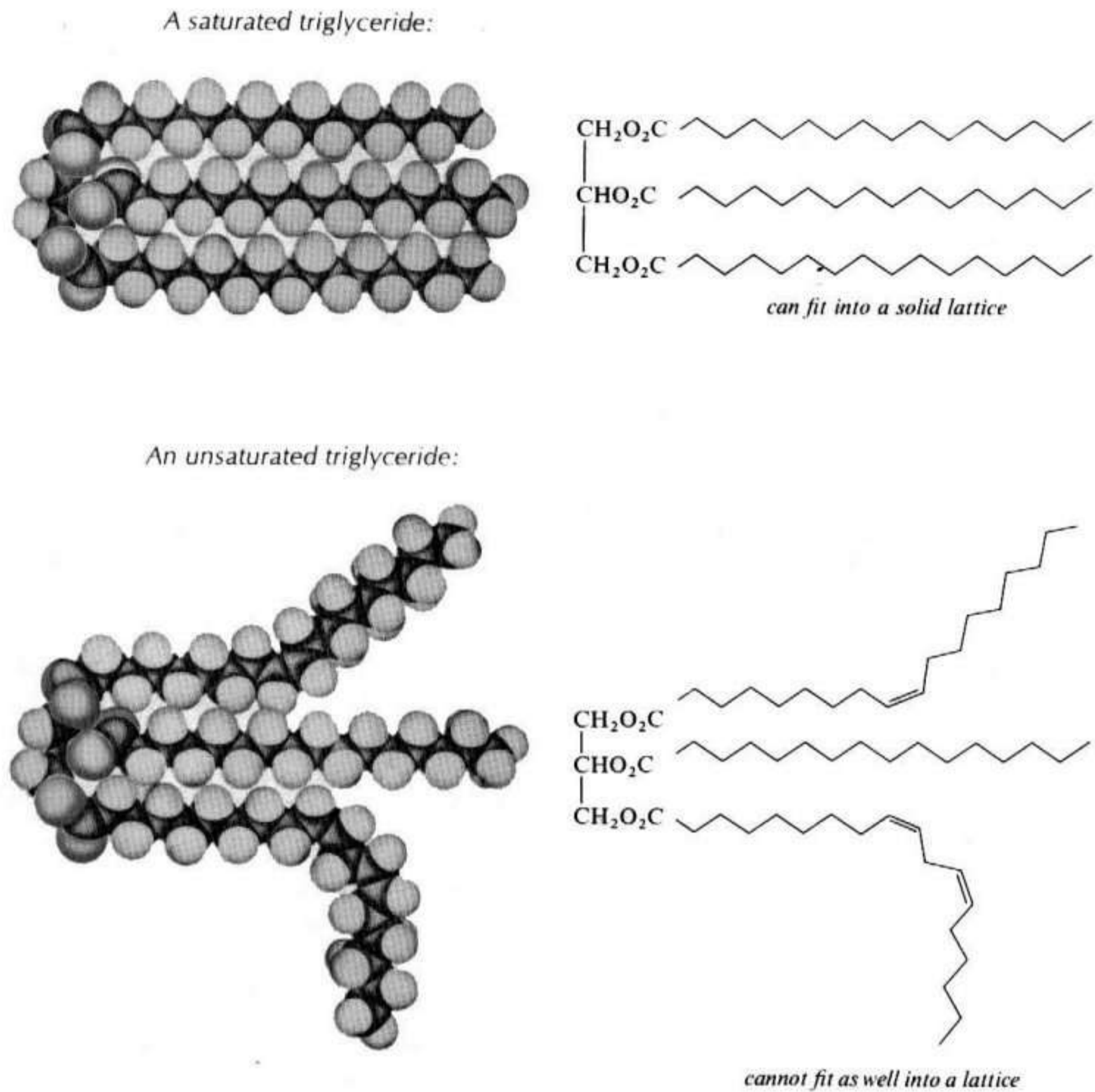
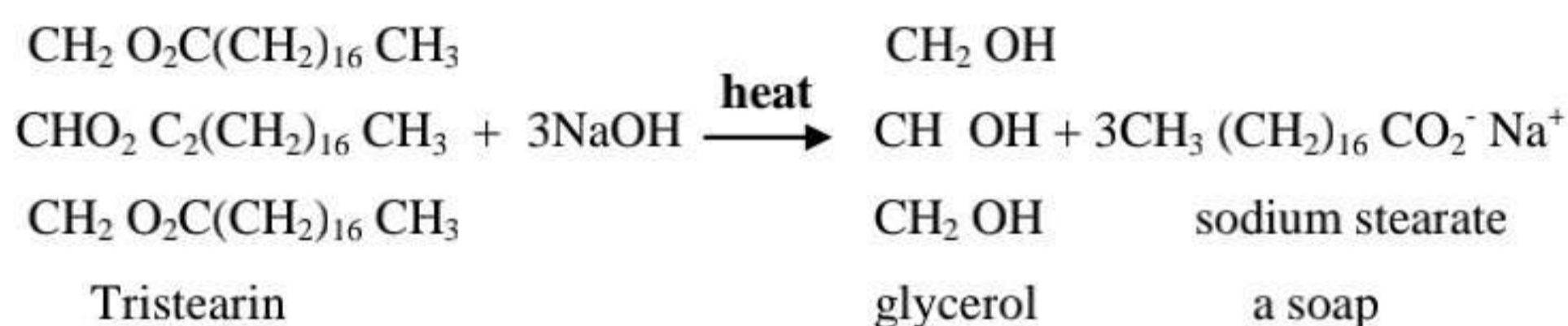


Figure 7.1. The shapes of saturated and unsaturated triglycerides. Adapted from William H. Brown and Judith A. McClarin, *Introduction to Organic and Biochemistry*, 3rd ed. (Willard Grant Press, Boston, 1981).

**SECTION 7.2.****Soaps and Detergents:**

The word *saponify* means “make soap.” Saponification of an ester with NaOH yields the sodium salt of a carboxylic acid. Saponification of a triglyceride yields a salt of a long-chain fatty acid, which is a soap. American pioneers used beef or pork fat and wood ashes (which contain alkaline salts, such as  $K_2CO_3$ ) to make soap. (It was reported by Julius Caesar that Teutonic tribes of his era also made soap this way.)



A molecule of a soap contains a long hydrocarbon chain plus an ionic end. The hydrocarbon portion of the molecule is hydrophobic and soluble in nonpolar substances, while the ionic end is hydrophilic and water-soluble. Because of the hydrocarbon chain, a soap molecule as a whole is not truly soluble in water. However, soap is readily suspended in water because it forms micelles, clusters of hydrocarbon chains with their ionic ends facing the water (see Figure 7.2).

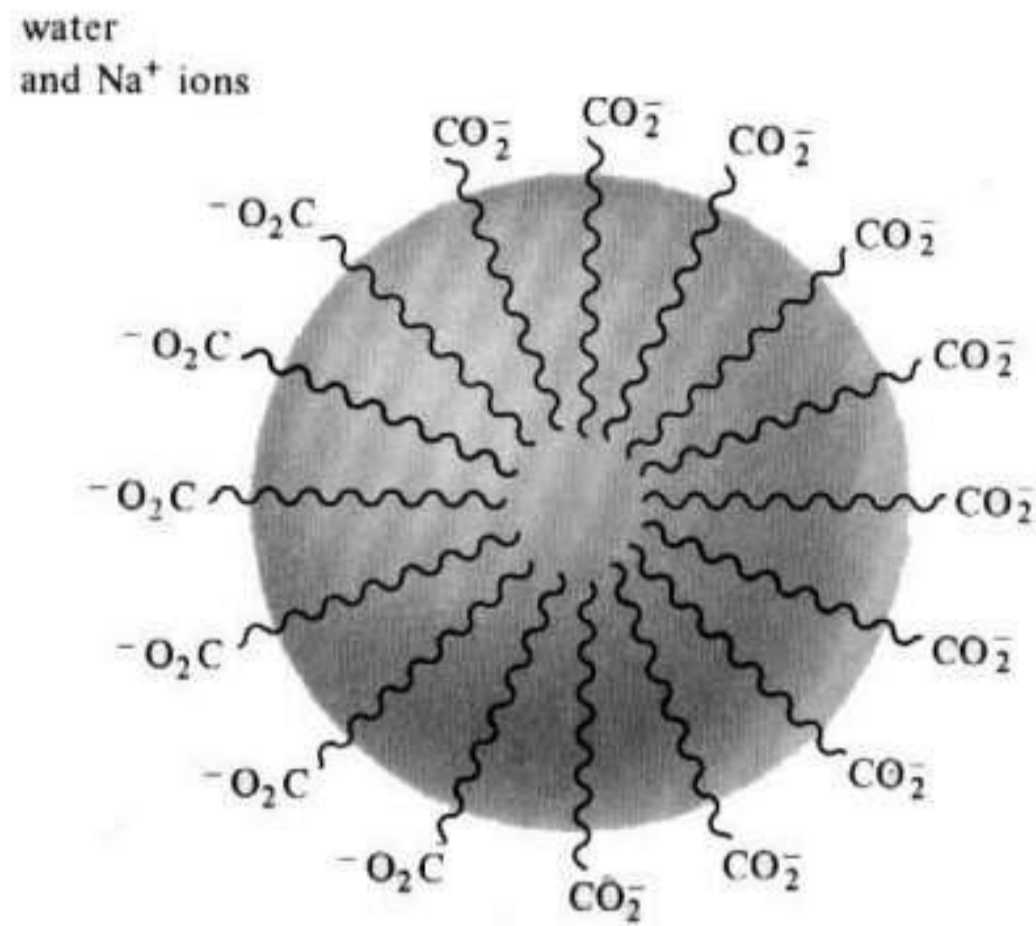
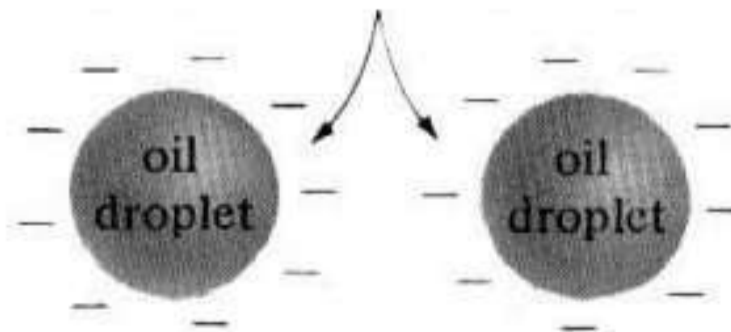


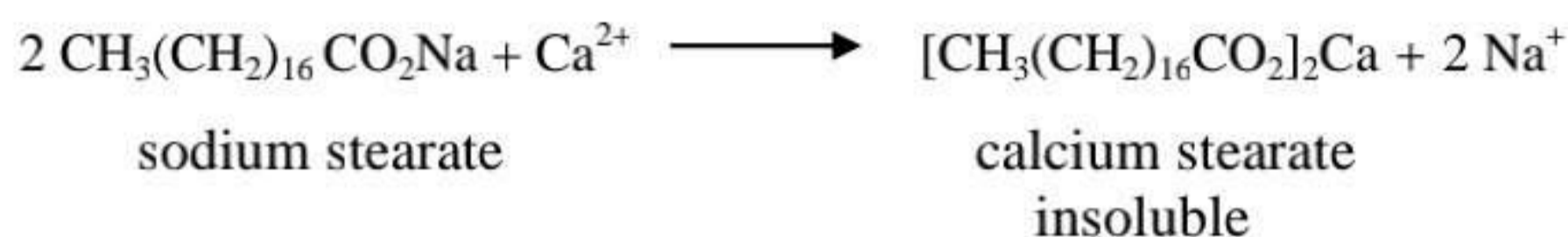
Figure 7.2. A micelle of the alkylcarboxylate ions of a soap.

The value of a soap is that it can emulsify oily dirt so that it can be rinsed away. This ability to act as an emulsifying agent arises from two properties of the soap. First, the hydrocarbon chain of a soap molecule dissolves in nonpolar substances, such as droplets of oil. Second, the anionic end of the soap molecule, which is attracted to water, is repelled by the anionic ends of soap molecules protruding from other drops of oil. Because of these repulsions between the soap-oil droplets, the oil cannot coalesce, but remains suspended.

*in soapy water, oil droplets repel each other because of similar charges of soap's carboxylate groups*

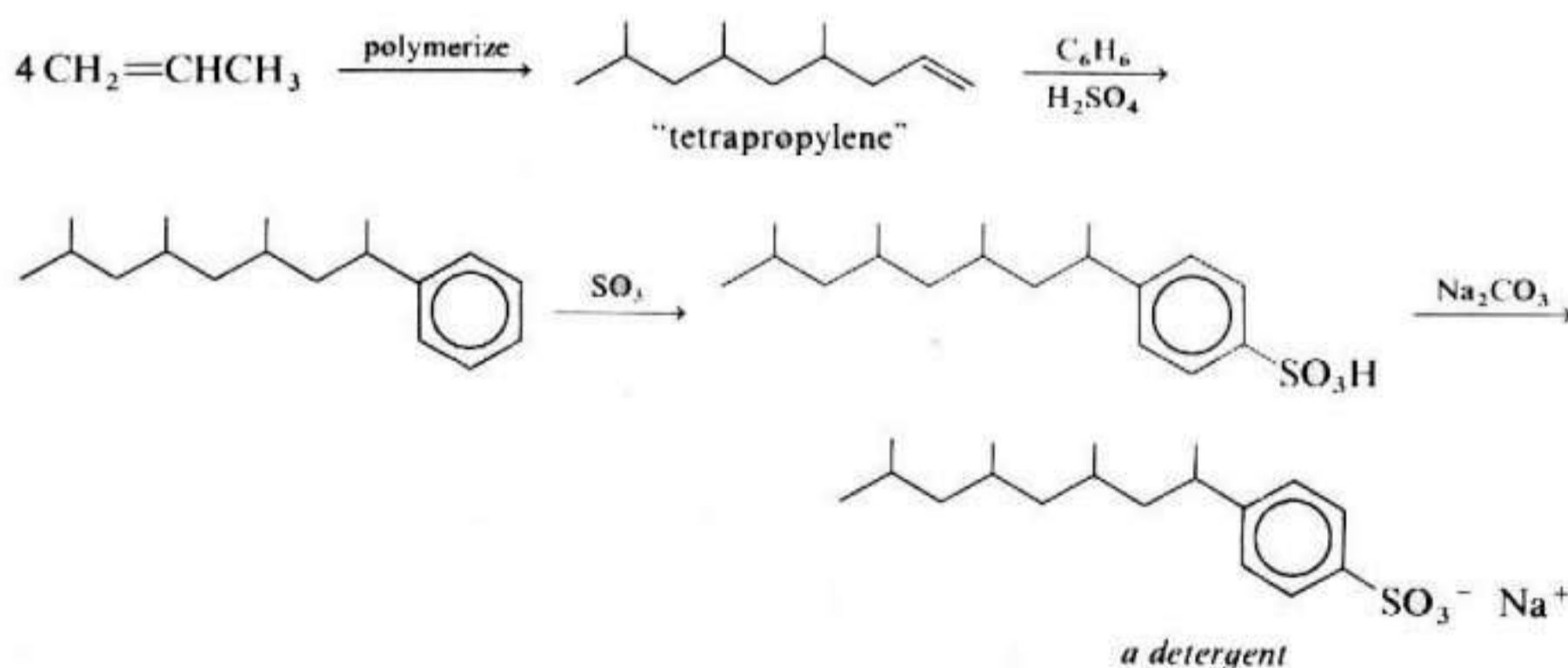


A disadvantage of soaps is that they form insoluble salts (bathtub ring) with  $\text{Ca}^{2+}$ ,  $\text{Mg}^{2+}$  and other ions found in hard water. ("Softening" water involves exchanging these ions for  $\text{Na}^+$ )



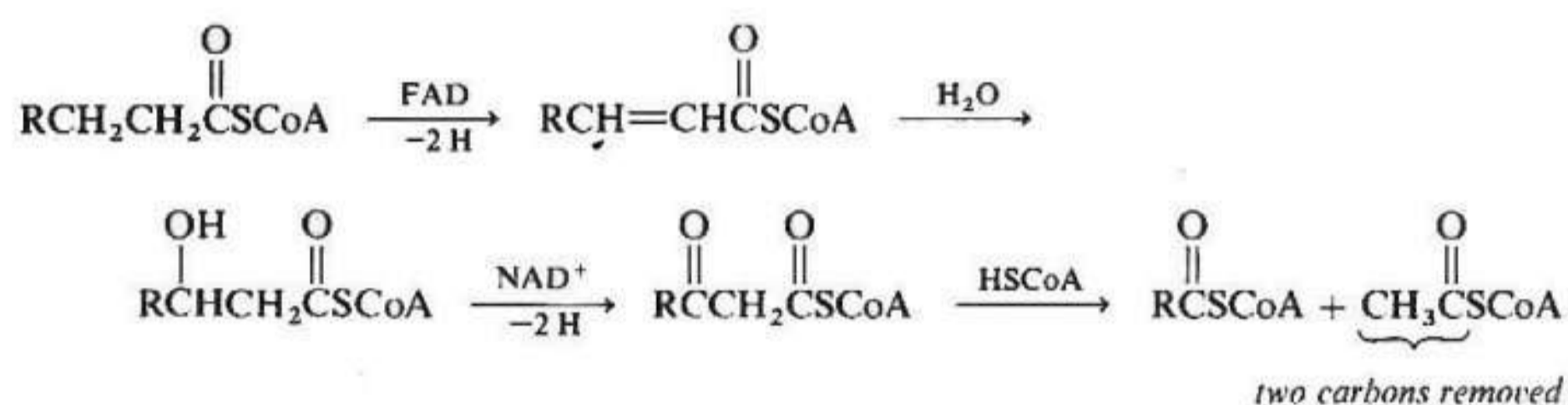
Most laundry products and many toilet "soaps" and shampoos are not soaps, but detergents. A detergent is a compound with a hydrophobic hydrocarbon end plus a sulfonate or sulfate ionic end. Because of this structure, a detergent has the same emulsifying properties as a soap. The advantage of a detergent is that most metal alkylsulfonates and sulfates are water-soluble; detergents do not precipitate with the metal ions found in hard water.

One of the first detergents in common use was a highly branched alkylbenzenesulfonate. The alkyl portion of this compound is synthesized by the polymerization of propylene and is attached to the benzene ring by a Friedel-Crafts alkylation reaction. Sulfonation, followed by treatment with base, yields the detergent.





Although the microorganisms in septic tanks or sewage-treatment plants can break down continuous-chain alkyl groups into smaller organic molecules, they cannot degrade branched chains. The reason for this difference in biodegradability is that long-chain hydrocarbons are degraded two carbons at a time by way of a keto ester. Branching interferes with the formation of the ketone group, and thus blocks the entire sequence. (FAD,  $\text{NAD}^+$  and  $\text{HSCoA}$ , shown in the following equation, are discussed in Sections 13.8 and 19.14B.)



To prevent the build-up of detergents in rivers and lakes, present-day detergents are designed with biodegradability in mind. One type of biodegradable detergent is an alkylbenzenesulfonate with a continuous-chain, rather than a branched-chain, alkyl group. Another type of biodegradable detergent is a continuous-chain alkylsulfate.

**Course Title:**

**Metabolism of Amino Acids ,**  
**Protein and Lipids**

الكلية : العلوم

الشعبة : Biochemisty

الفرقة : الثانية

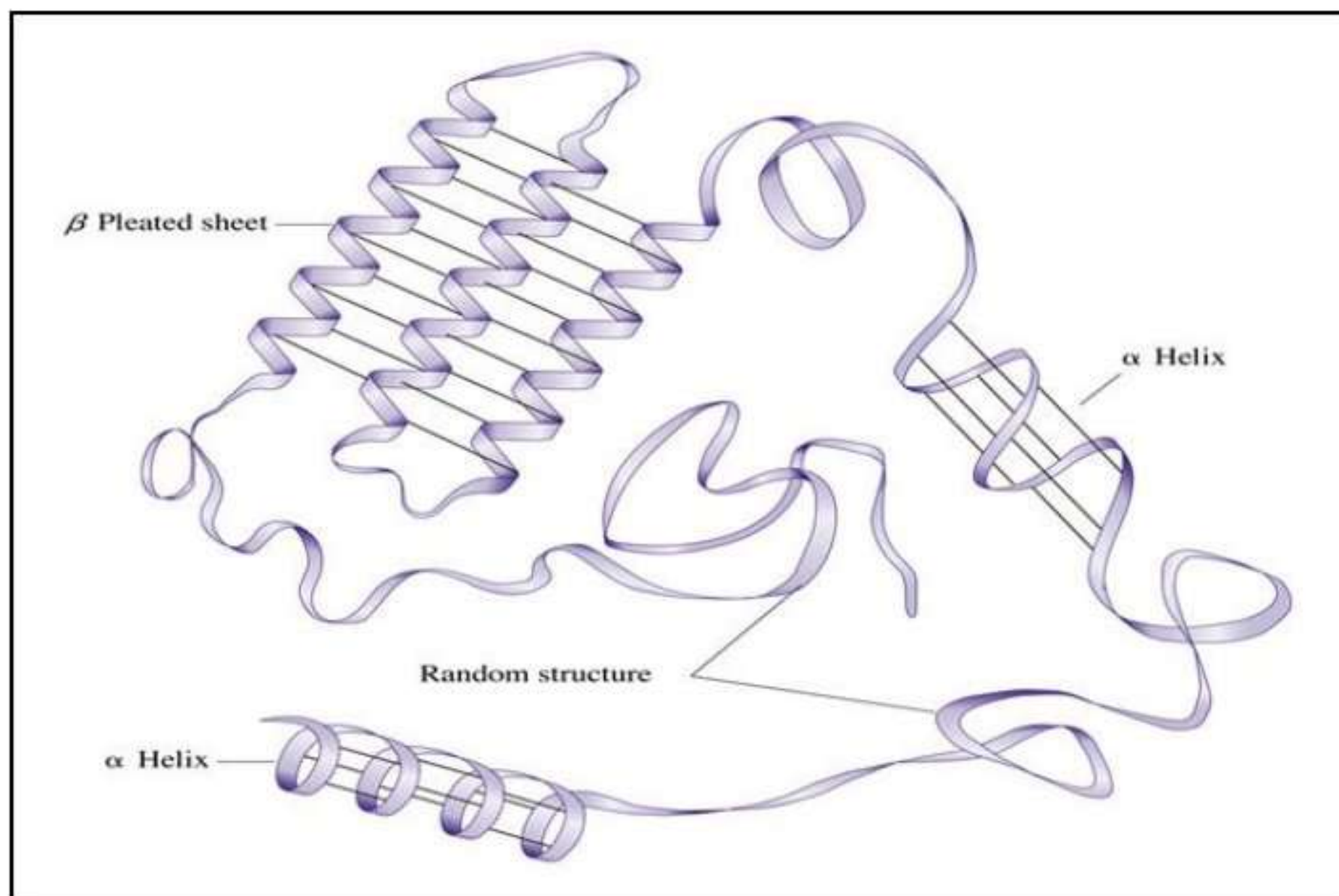
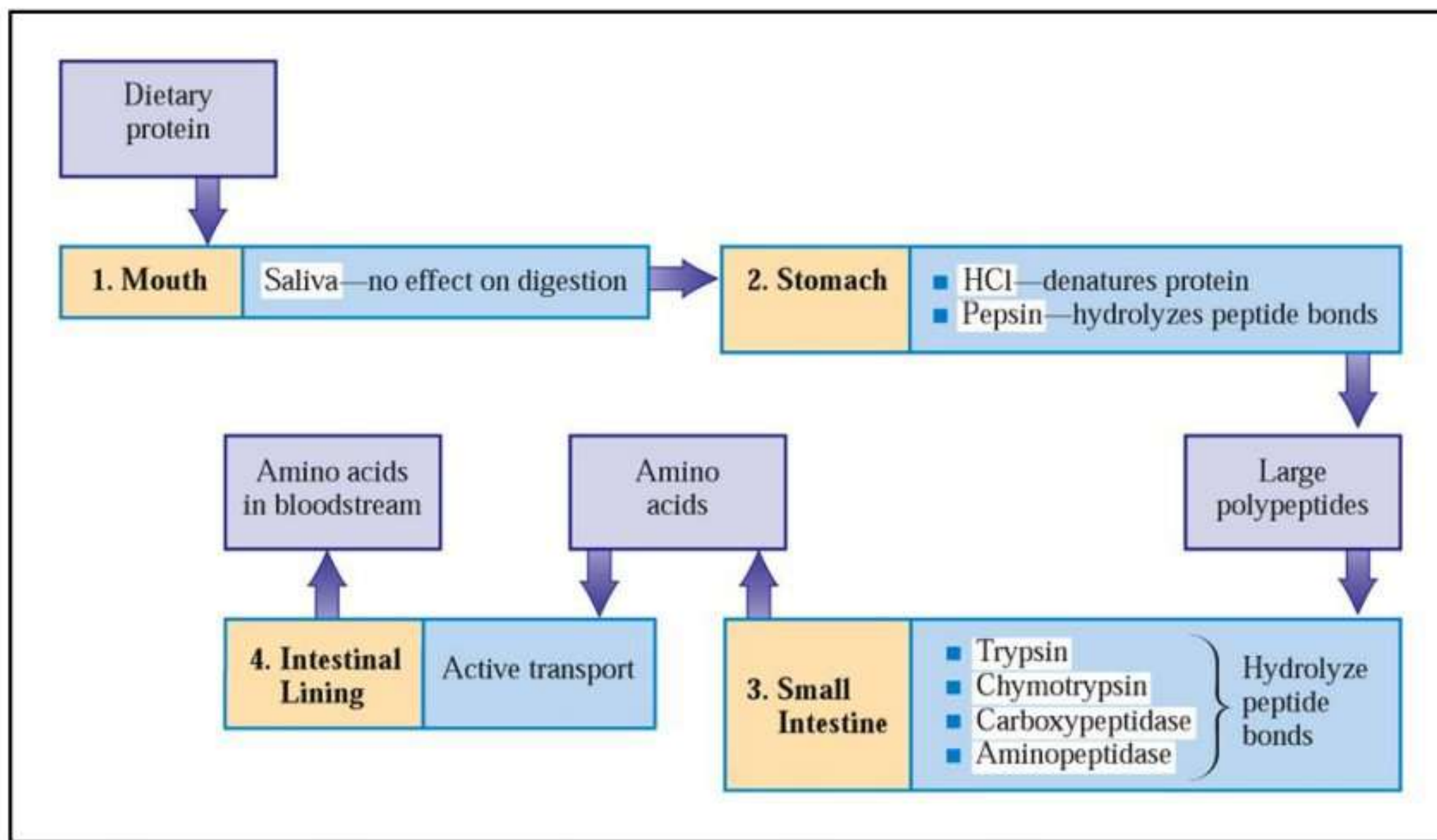
اعداد:

**Ass. Prof. Dr.Hussien Temerik**

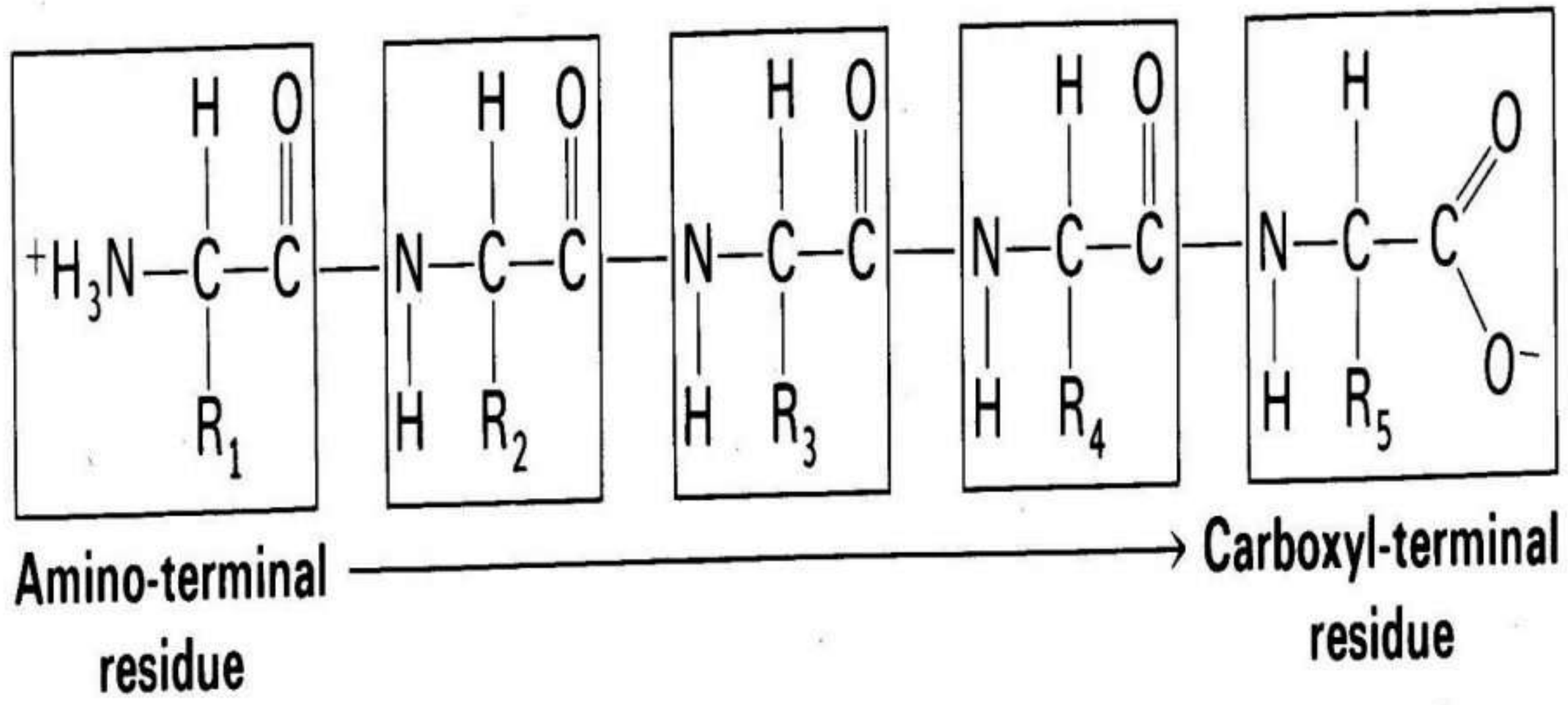
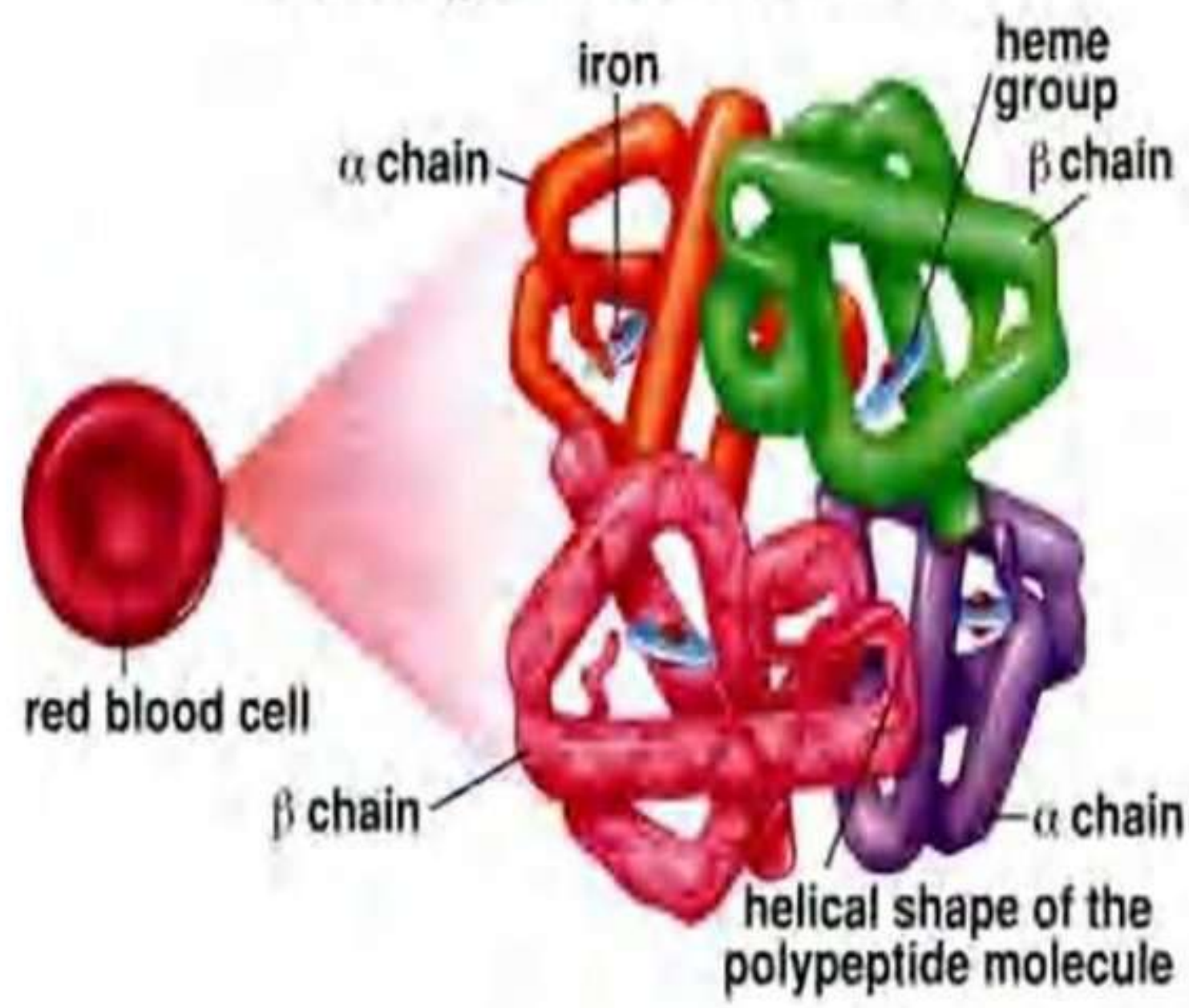
## Protein Digestion

Protein breakdown begins in the **stomach**.

No protein hydrolyzing enzymes are found in saliva.



# Hemoglobin Molecule



**Hydrolysis** (10% of peptide bonds) & **denaturization** by pepsin enzyme & HCl acid produce **short chain polypeptides** in the stomach.

**Trypsin, chymotrypsin, & carboxypeptidase** from Pancreatic juices, and **Aminopeptidase** from cells in the small intestine Brush Zone create “free” **amino acids**.

Free amino acids are absorbed thru intestinal wall via active transport. Enter bloodstream and are brought to cells.

The total supply of free amino acids available is called: the **Amino Acid Pool**.

3 sources of “free” amino acids:

1. Dietary protein breakdown
2. Biosynthesis of amino acids in the Liver
3. Protein turnover (I prefer apple turnovers)

**Protein turnover** is the breakdown & re-synthesis of body protein:

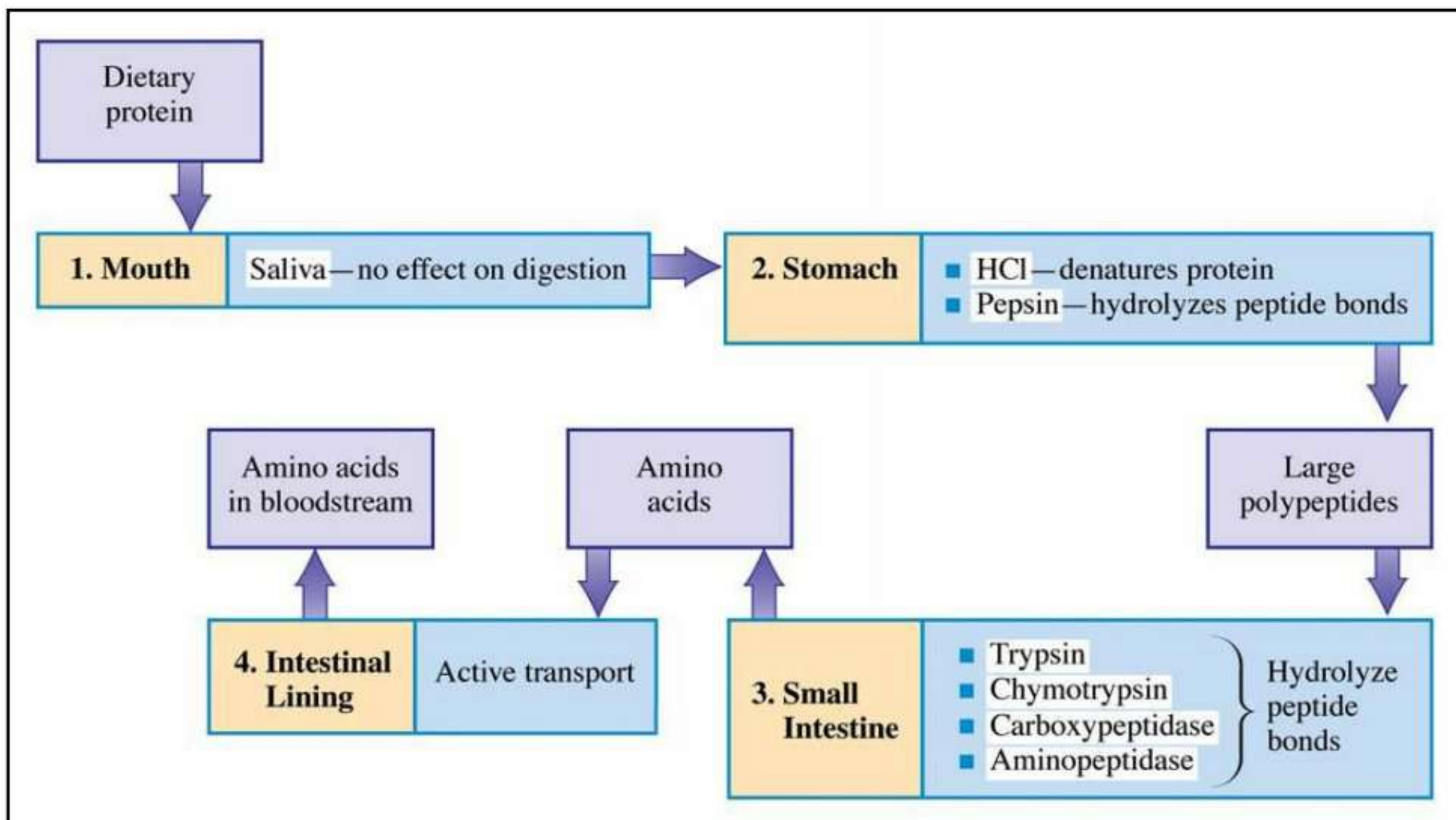
Old tissues

Damage

Recycling enzymes & hormones



Summary of protein digestion in the human body. Possible fates for amino acid degradation products.



### Transamination and Oxidative Deamination:

Two steps in degrading amino acids

- 1) remove  $\alpha$ -amino group
- 2) breakdown & process carbon skeleton

Release of an **amino group** is also two steps:

- 1) **Transamination**
- 2) **Oxidative deamination**

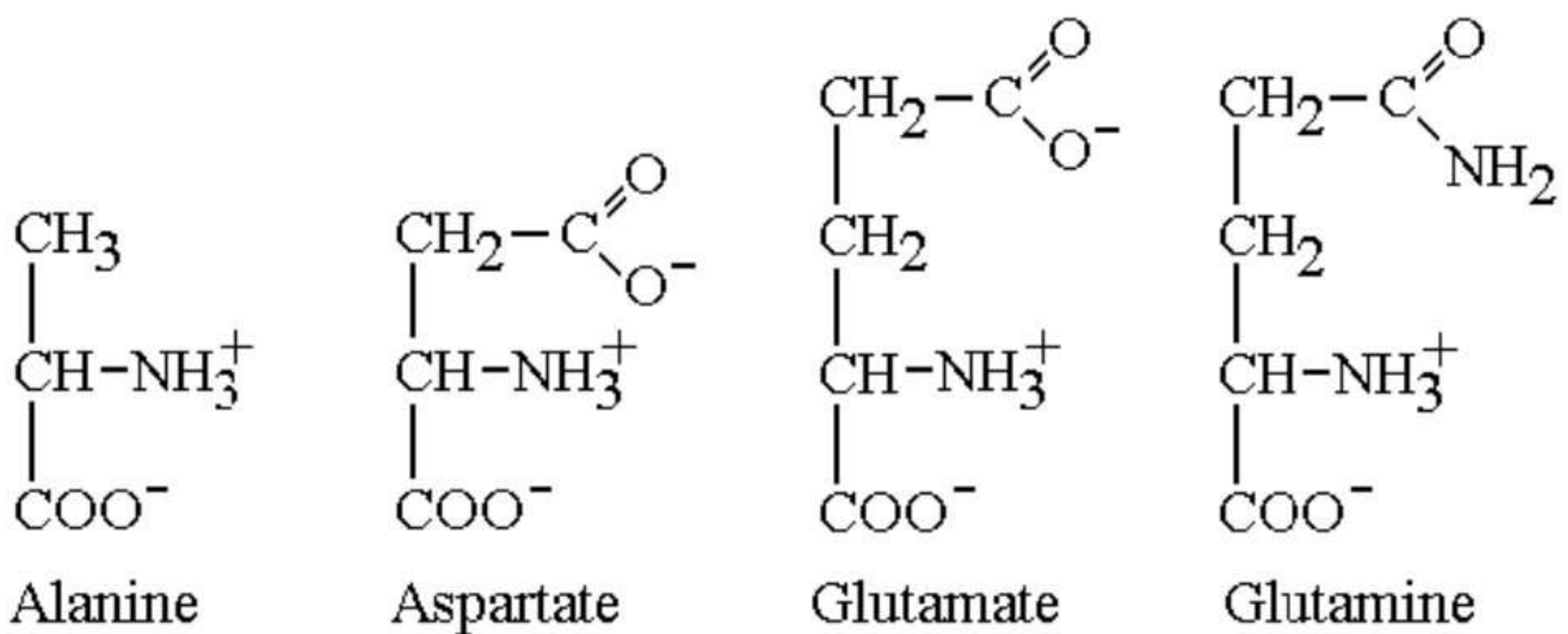
### Central role of glutamate:

Amino acids:

**Glutamate**, **aspartate**, **alanine** & **glutamine**

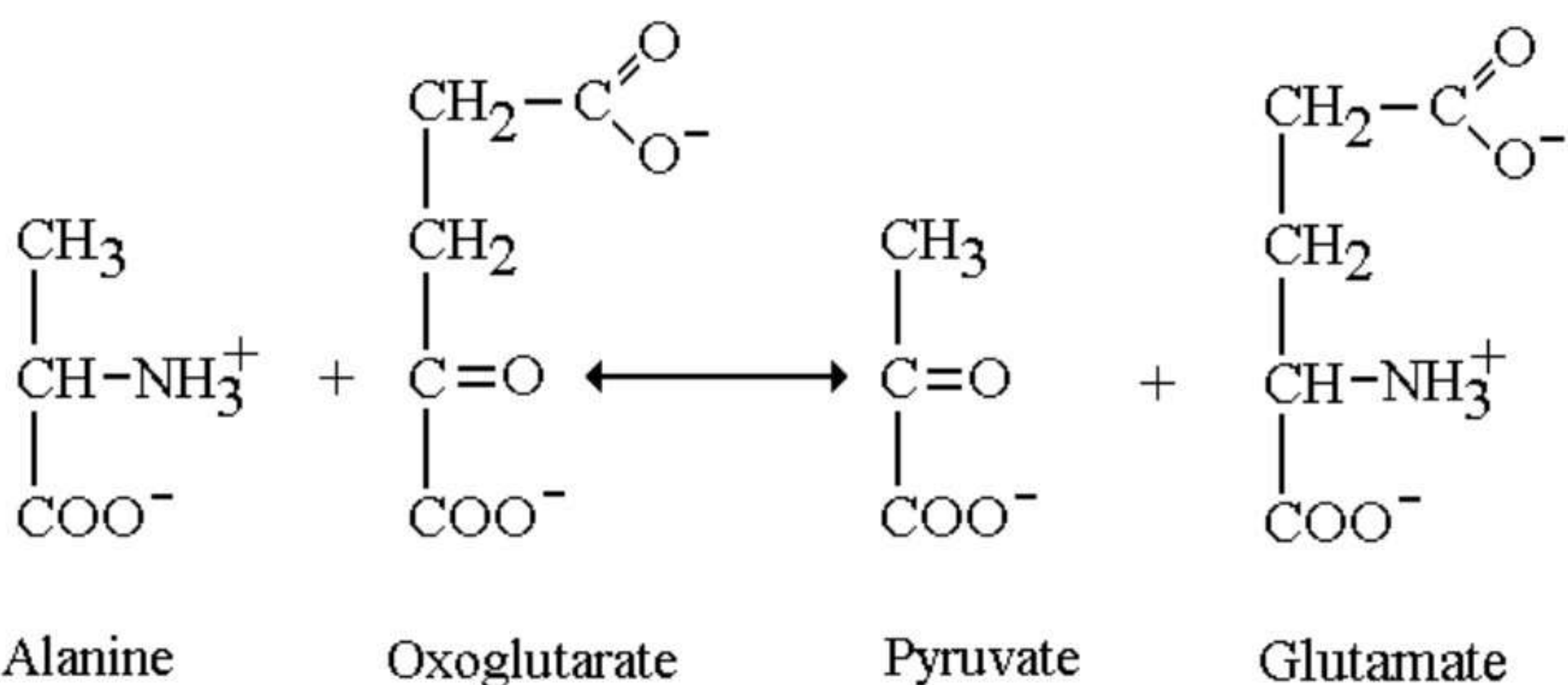
present in higher concentrations in mammalian cells. Have metabolic functions as well as roles in proteins.

**Glutamate** is the most important, metabolically

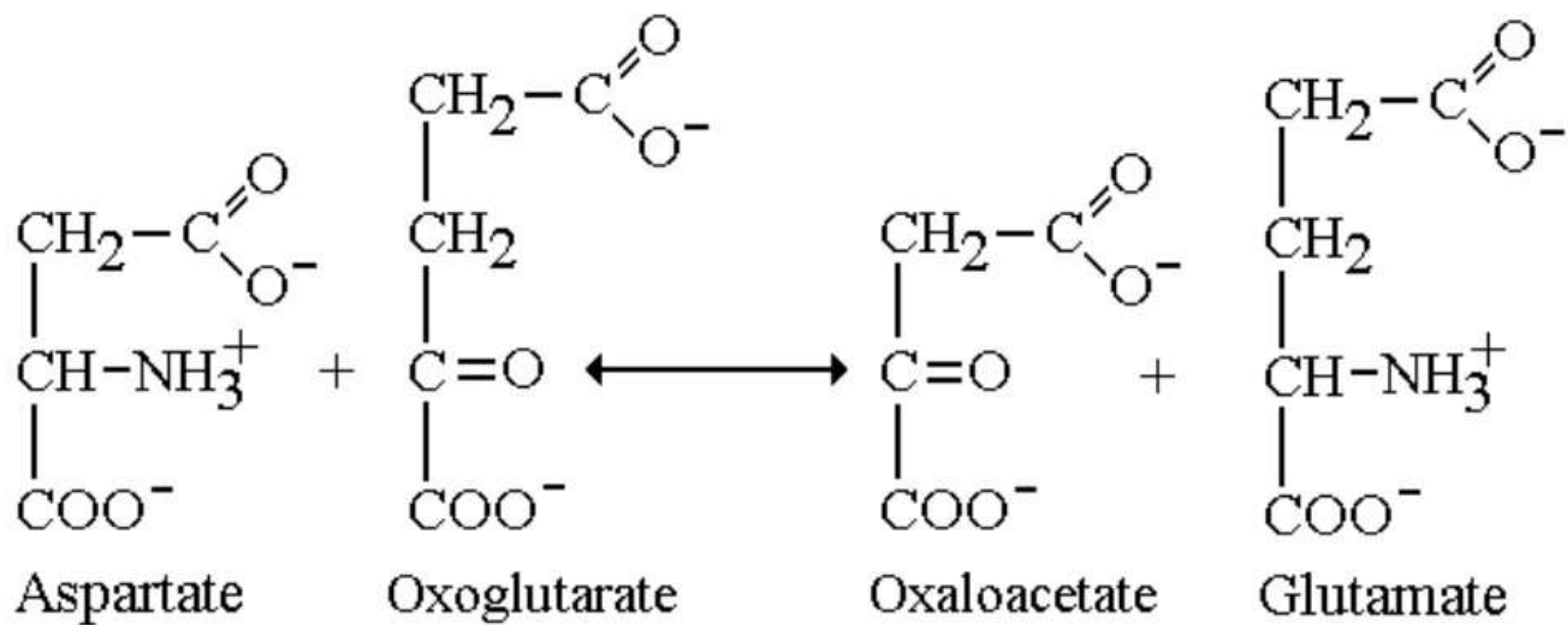


Some **transaminases** are used for diagnosing disorders:

enzyme **alanine aminotransferase**. Escapes in large amounts from dead or dying liver tissue. Measured in blood samples for diagnostic purposes.



Transaminase enzyme **aspartate aminotransferase** very active enzyme inside heart cells. Also escapes in large amounts from dead or dying heart tissues & enters bloodstream. Measured in blood for diagnosing myocardial infarction.

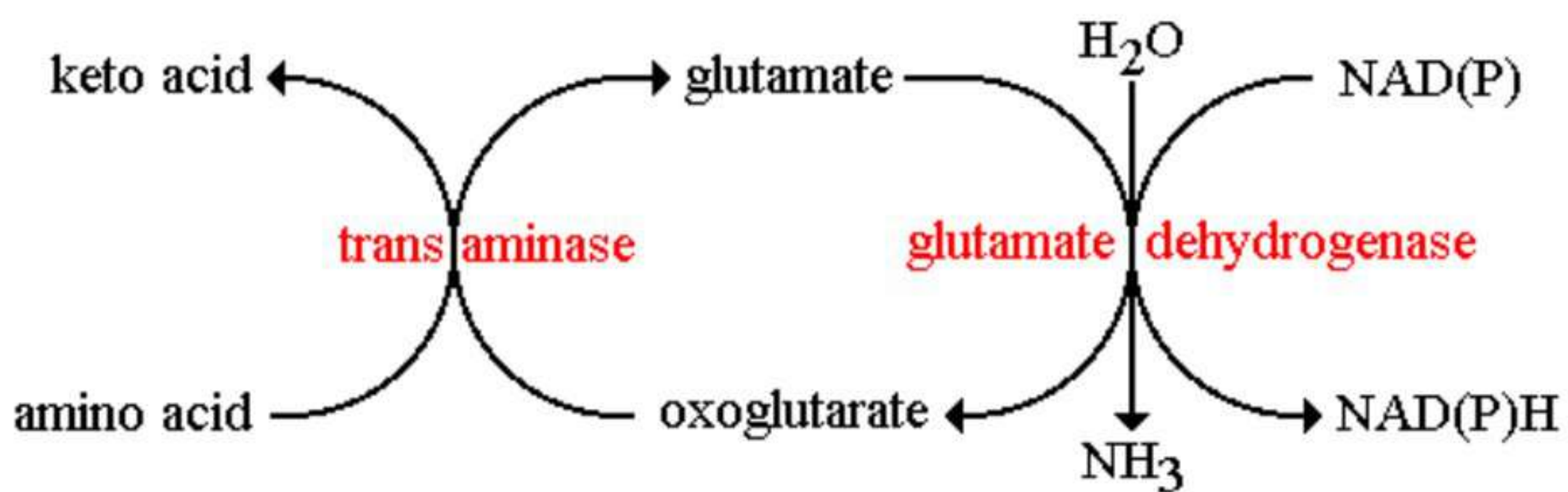


### Trans-deamination (sum it up)

Most **transaminases** share a common substrate and product (oxoglutarate and glutamate) with the enzyme **glutamate dehydrogenase**.

This permits a **combined** N excretion pathway for individual amino acids: "trans-deamination."

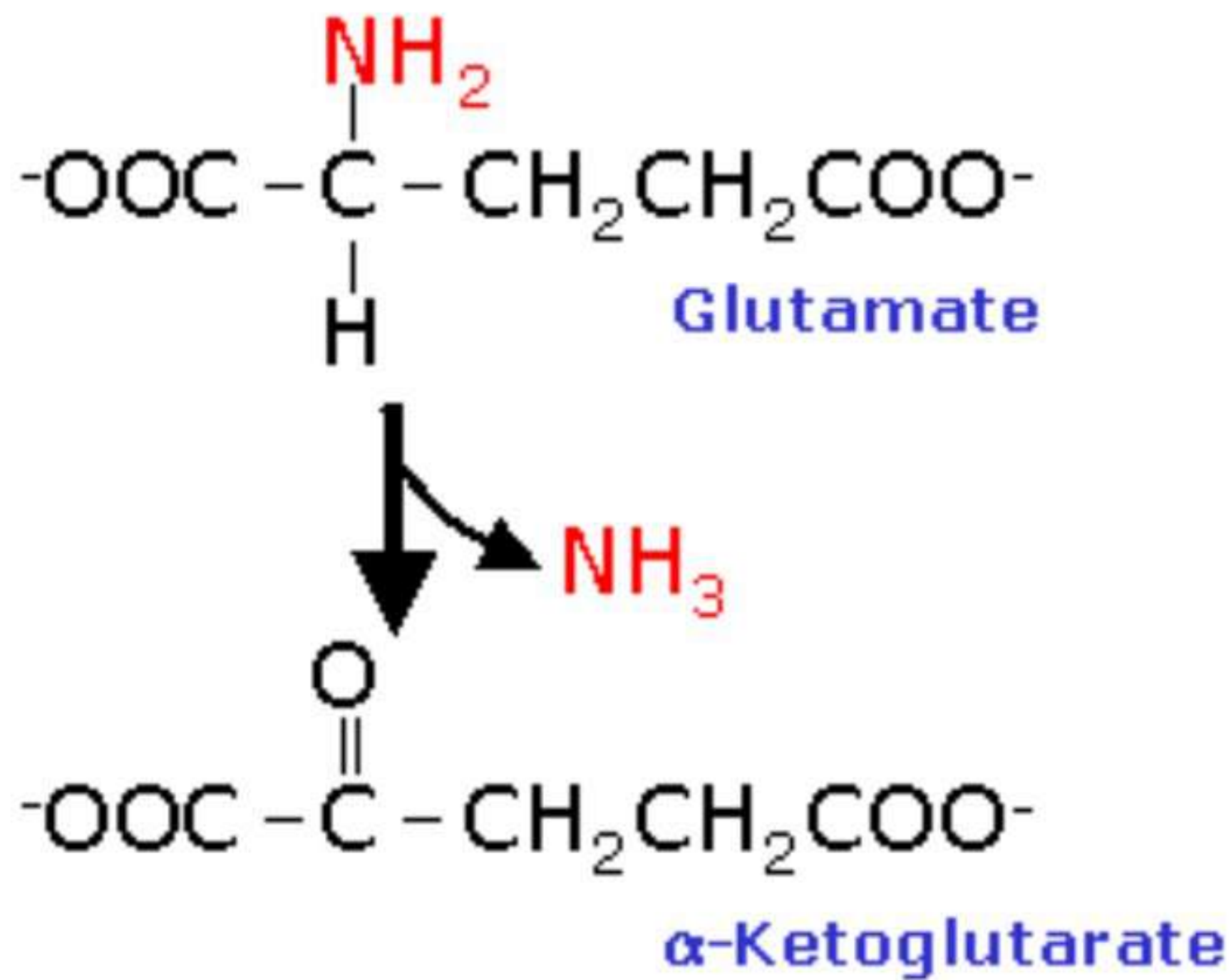
**Glutamate** has a central role in the overall control of nitrogen metabolism.



### Oxidative Deamination

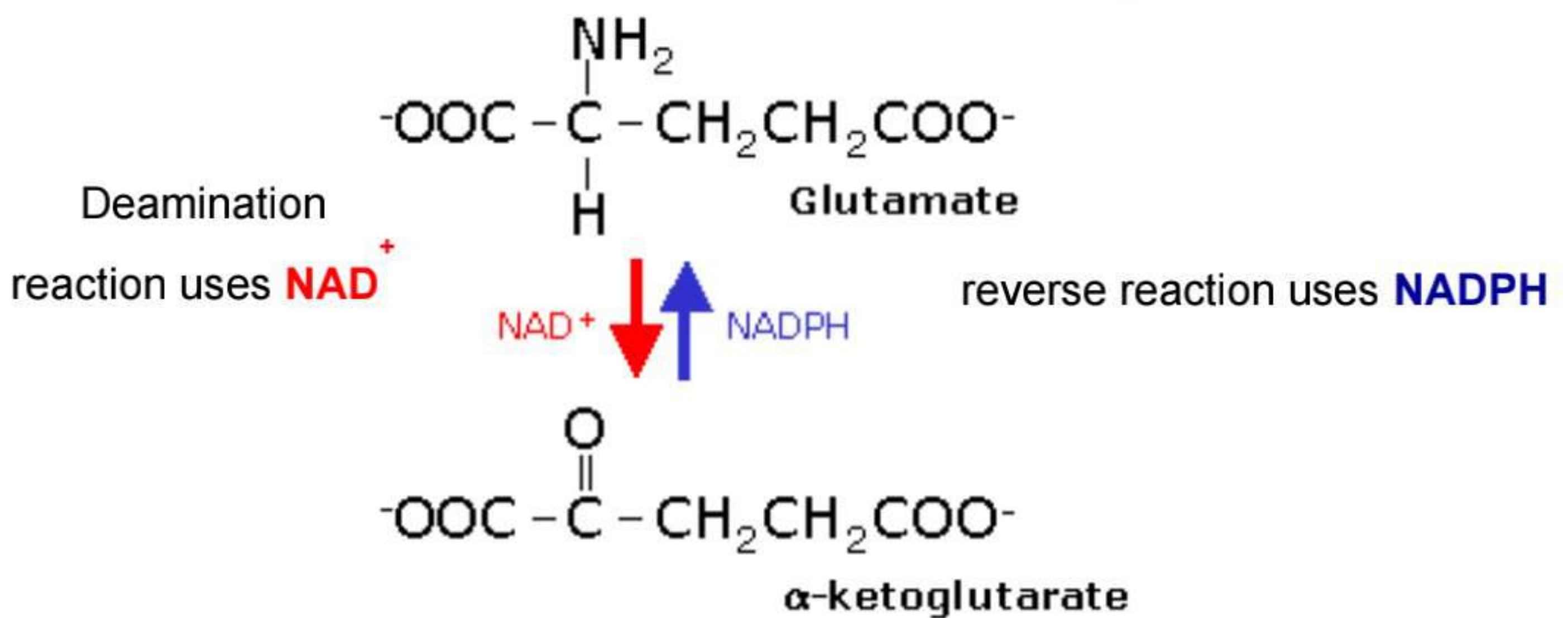
The **glutamate** produced from the transamination step is then deaminated by **oxidative deamination** using the enzyme **glutamate dehydrogenase**.





Recycles back to a ketodiacid & releases ammonia

**Glutamate dehydrogenase [GluDH]** will reversibly convert **glutamate** to **a-ketoglutarate** and **a-ketoglutarate** to **glutamate**.



Uses **both** **NAD<sup>+</sup>** and **NADPH** – how to regulate it?

## Urea cycle:

Ammonium salts ( $\text{NH}_4^+$ ) are toxic compounds.

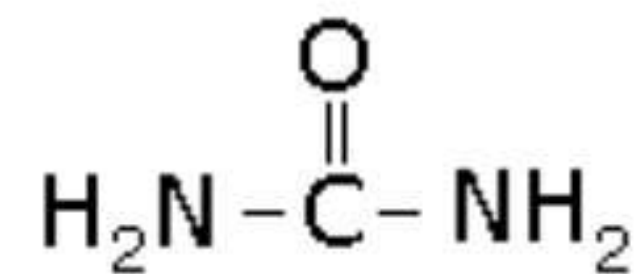
**Oxidative deamination** converting glutamate to  $\alpha$ -ketoglutarate is an easily shifted equilibrium reaction.

**Ammonium ions** building up favors the synthesis of excessive amounts of glutamate, decreasing the Krebs cycle intermediate

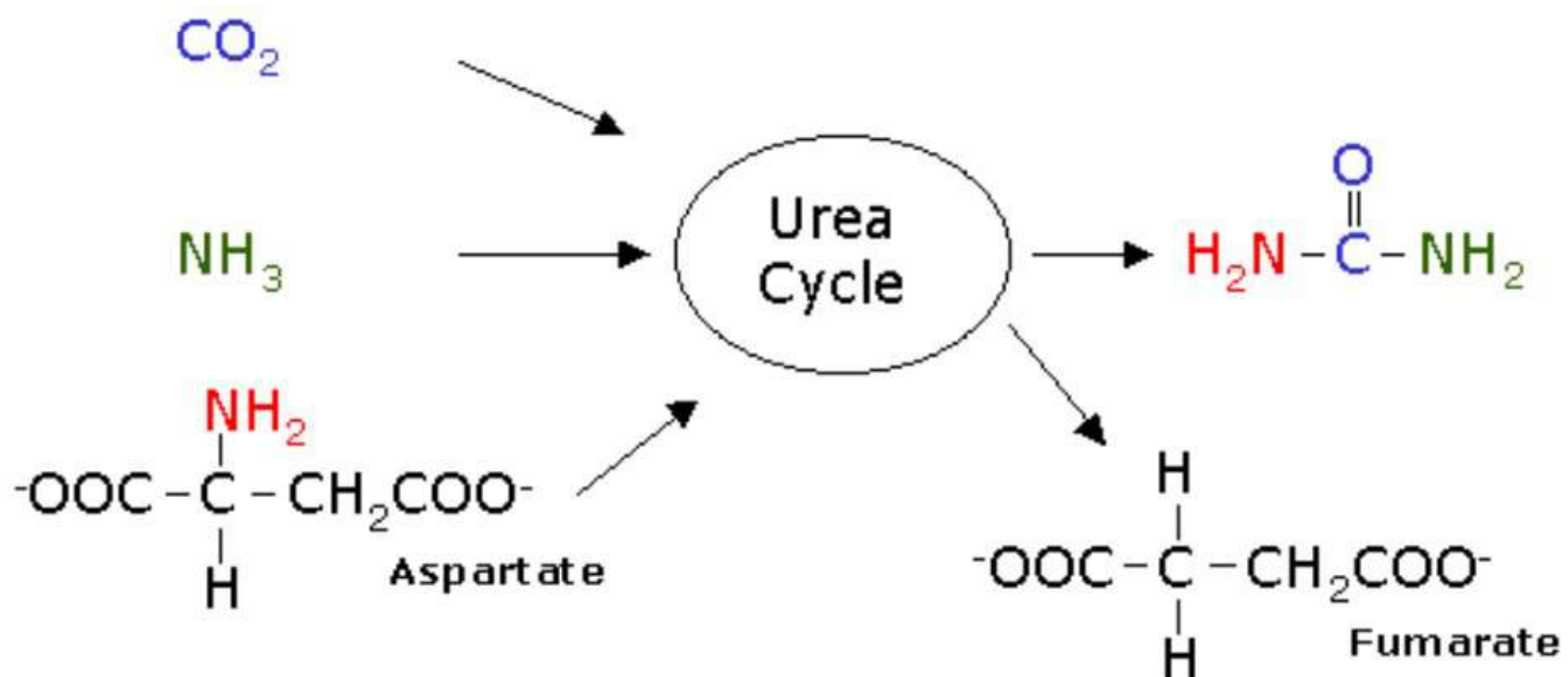
**$\alpha$ -ketoglutarate.**

This in turn decreases **ATP production**, and that affects the nervous system.

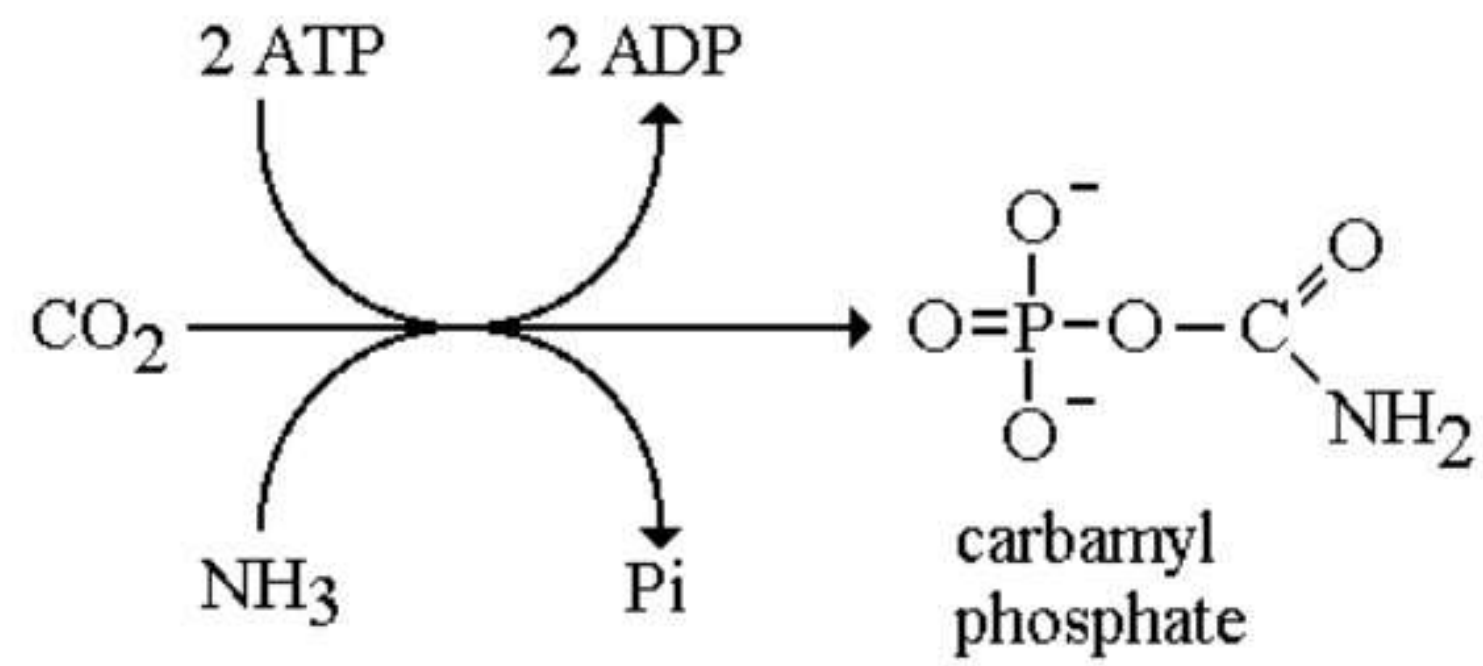
The answer is Urea:



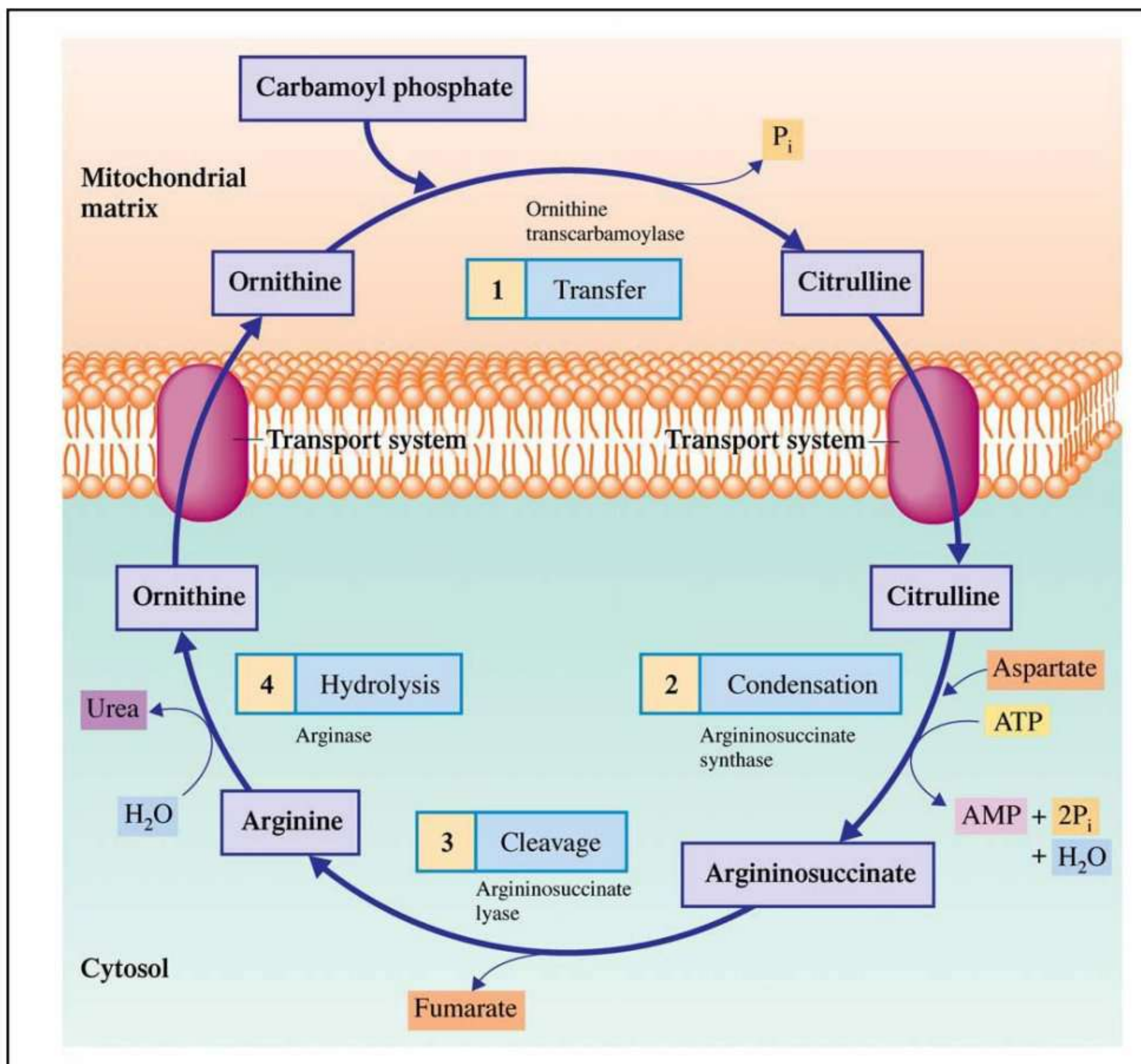
The **inputs** to the urea cycle are  $\text{NH}_3$ ,  $\text{CO}_2$  and aspartic acid and ATP.  
The **outputs** are urea, ADP and fumaric acid.



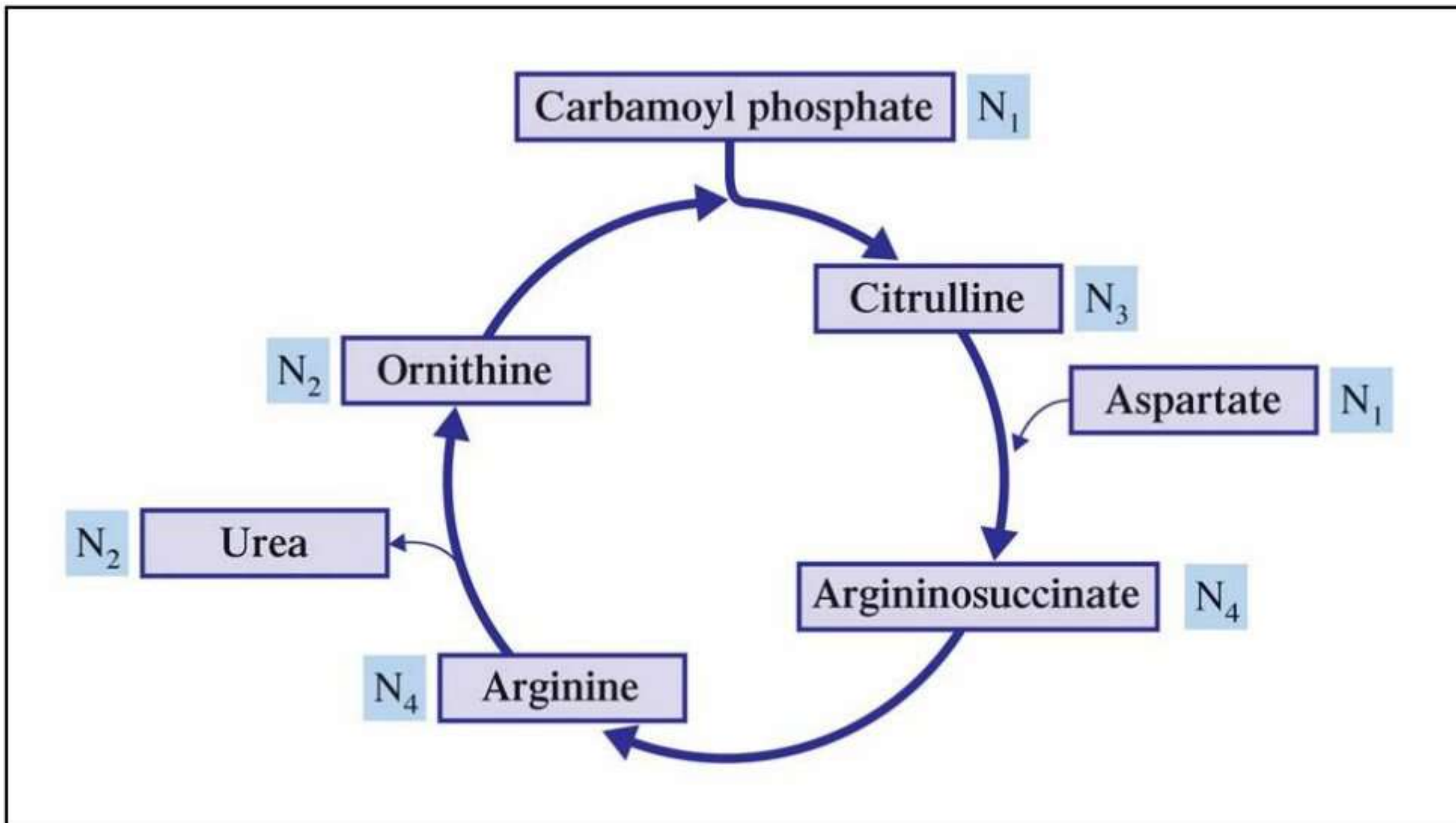
The carbonyl group of urea is derived from  $\text{CO}_2$ , **Ammonia** contributes one of the amine groups on urea



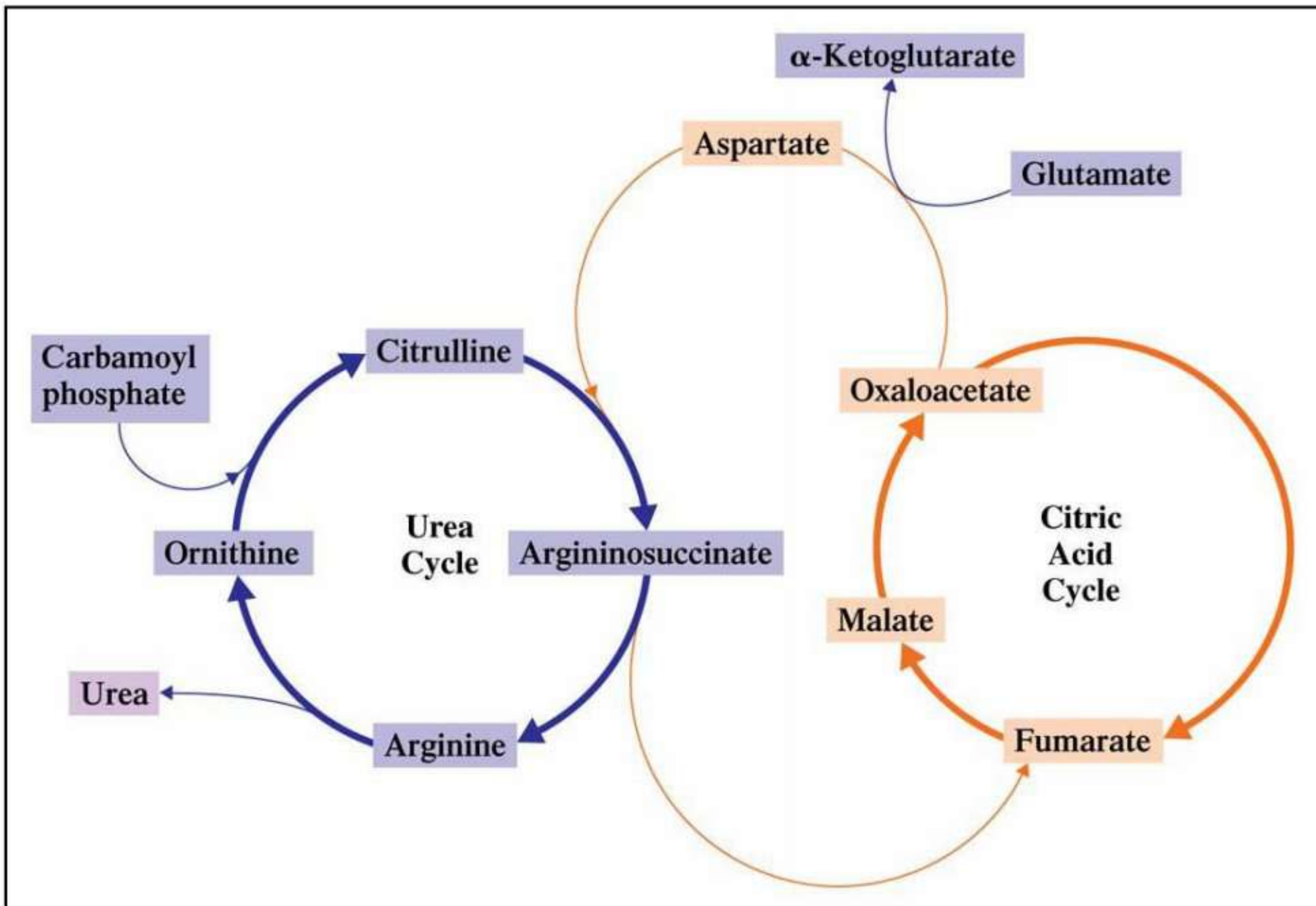
The **four-step urea cycle** in which **carbamoyl phosphate** is converted to **urea**.



The nitrogen content of the various compounds that participate in the urea cycle



**Fumarate** from the urea cycle enters the Krebs cycle. **Aspartate** produced from **oxaloacetate** of the Krebs cycle enters the urea cycle.

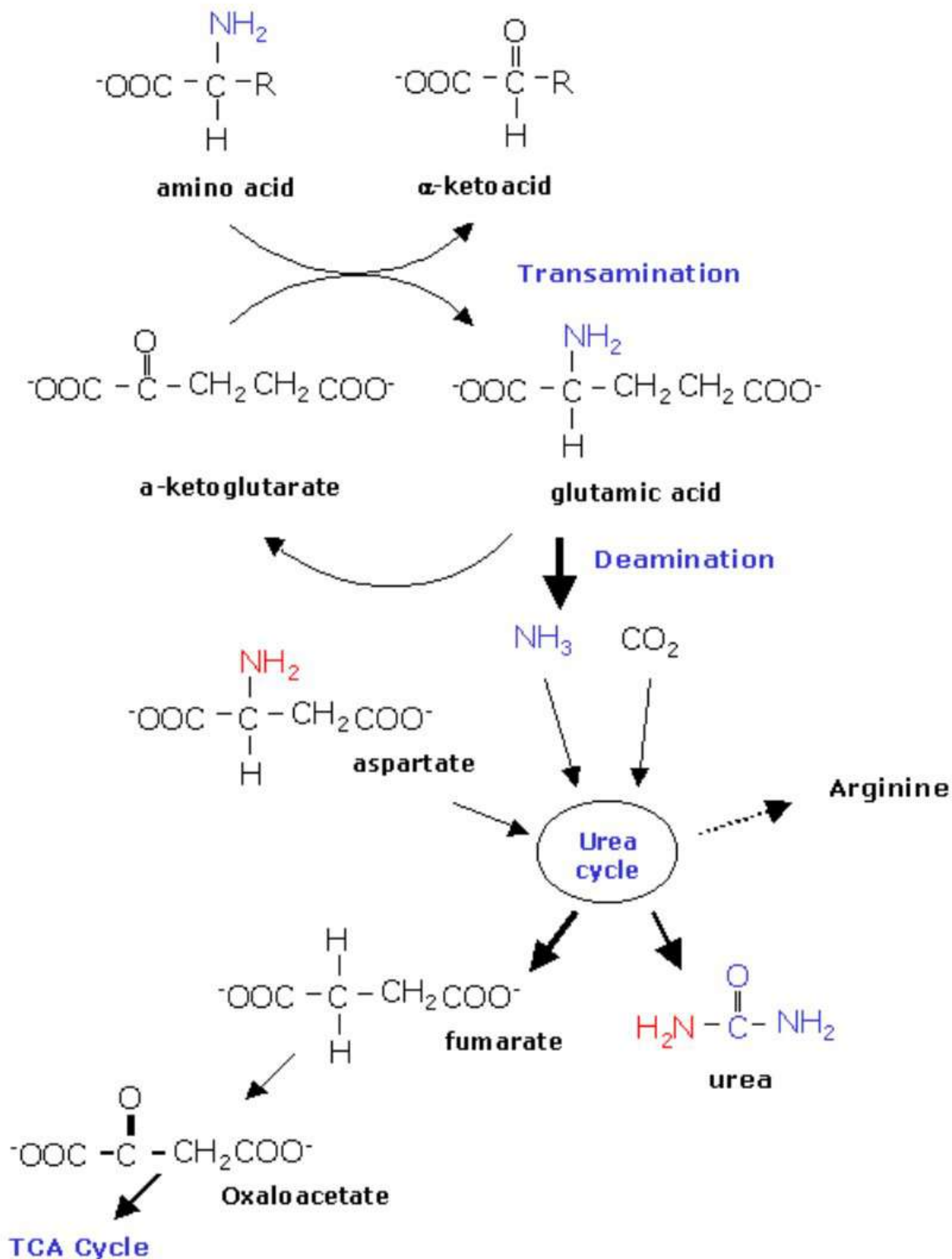


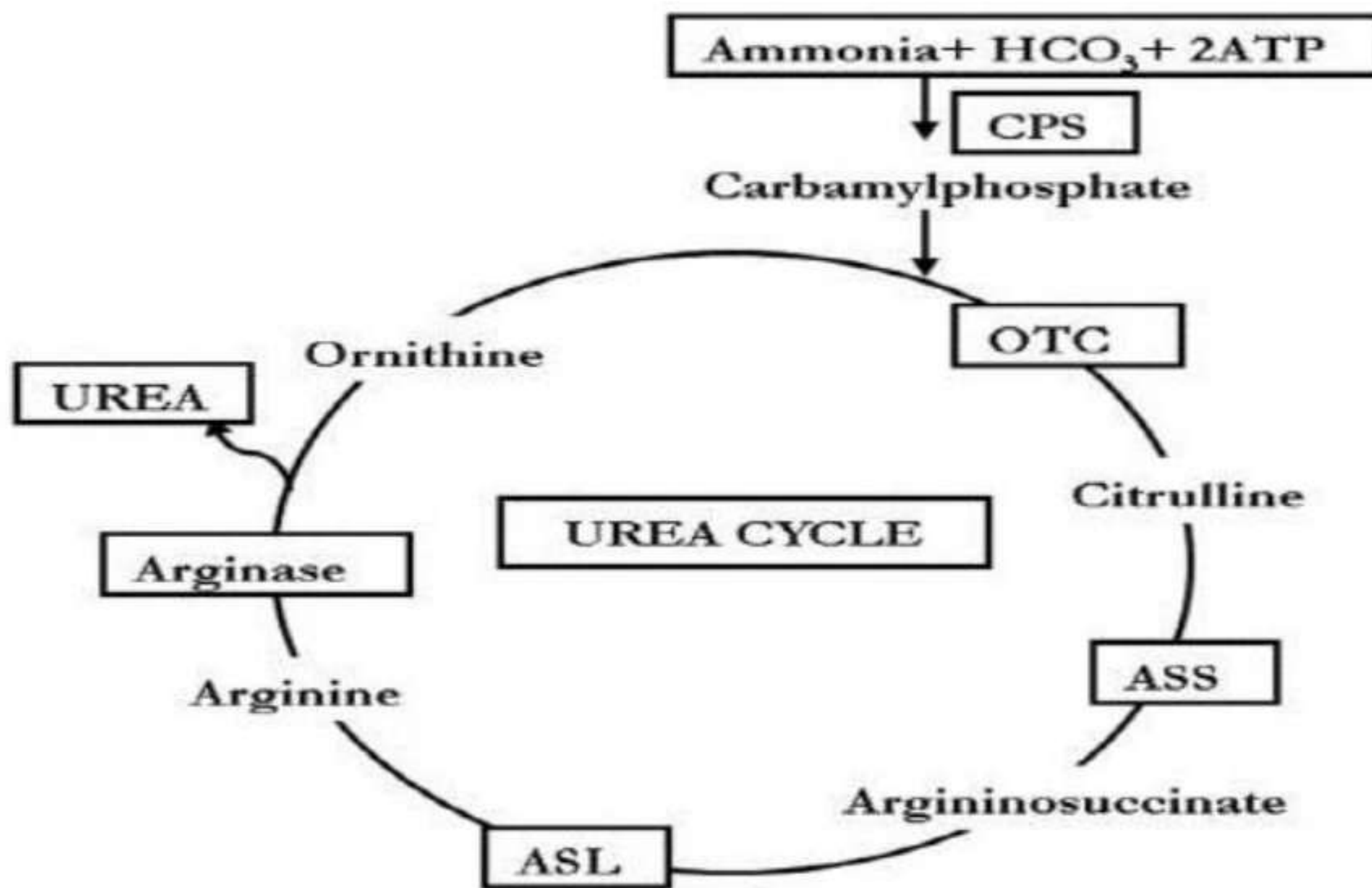
Oxaloacetate has 4 potential fates: transamination; conversion to glucose; formation of citrate; conversion to pyruvate

Summary:

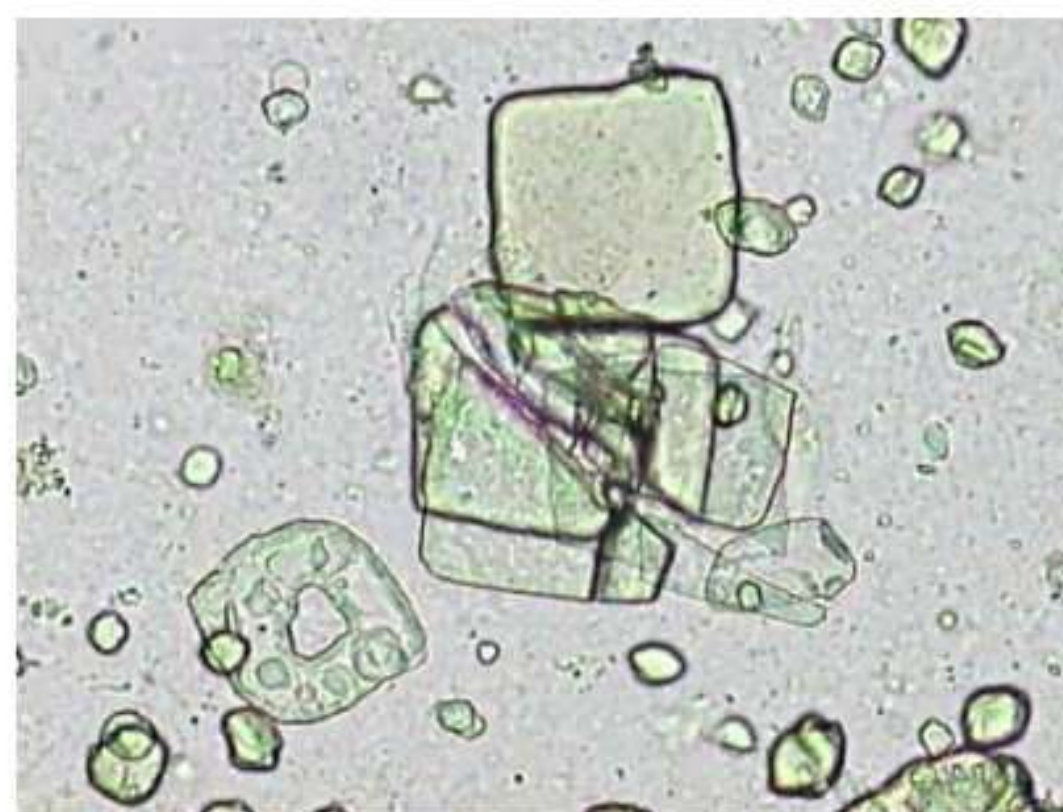
**Transamination** takes off amine groups from amino acids and forms **glutamate** (ionized glutamic acid)

Amine groups form **ammonia** when removed in **deamination**  
This combines with **CO<sub>2</sub>** & **Aspartate**.  
Forms **urea**, **Arginine**,  
& **Fumarate**

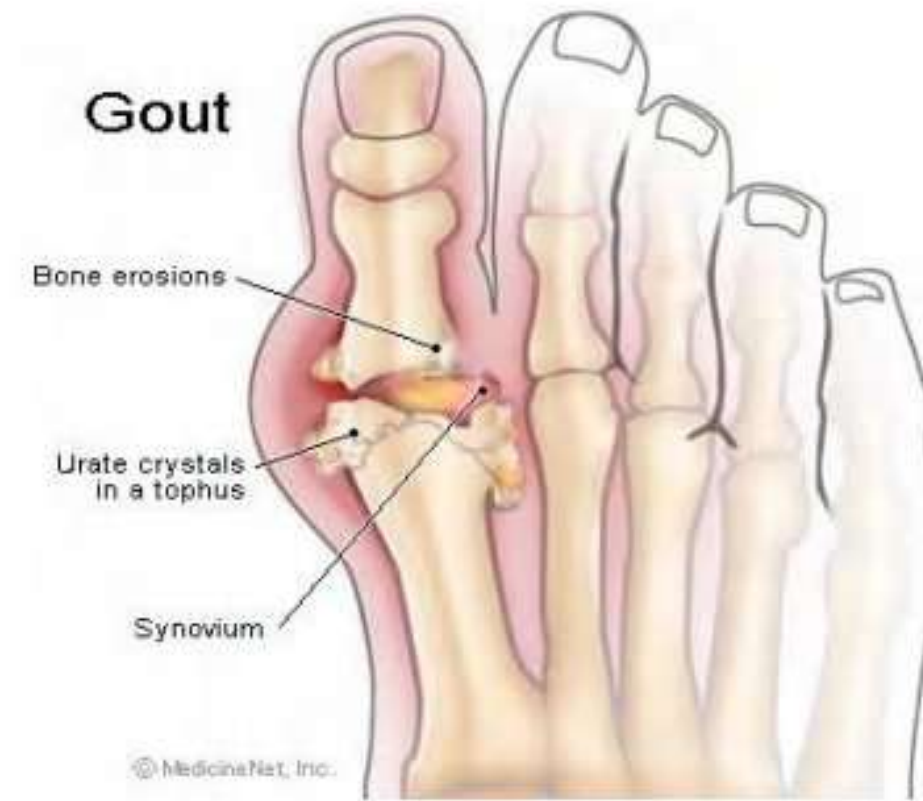




Reptiles & birds excrete **uric acid** – very *insoluble* purine compound – forms supersaturated solutions. Concentrated urine, supersaturated with uric acid, goes from cloaca into hindgut – uric acid crystallizes & water is reabsorbed.



In humans uric acid deposits crystals & causes gout



### Processing Amino Acid Carbon Skeletons

Transamination or Oxidative deamination both produce  $\alpha$ -keto acids  
 Degradation of these carbon skeletons may take several different pathways:

Amino acid C skeletons that degrade to form a Krebs cycle intermediate can then be used to make glucose via **gluconeogenesis**.

These are called **Glucogenic Amino Acids**.

Amino acid C skeletons that degrade to form **acetyl CoA** or **Acetoacetyl CoA** can form fatty acids or ketone bodies.

These are called **Ketogenic Amino Acids**.

### Amino Acid Biosynthesis

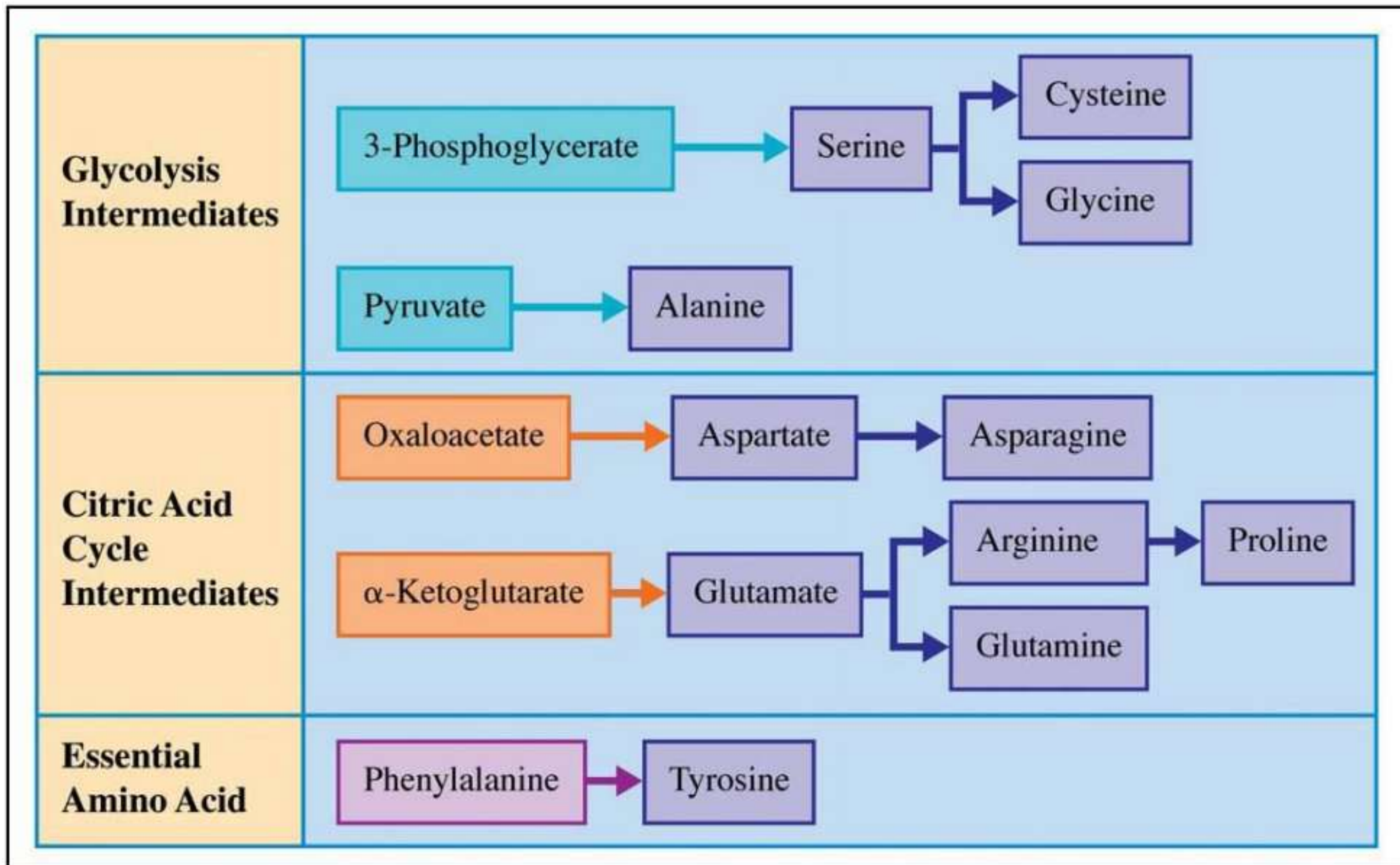
Essential amino acids can be made by plants & bacteria in 7 to 10 steps.

We obtain these amino acids by eating plants. 11 Non-essential amino acids synthesized in 1 to 3 steps. Use glycolysis intermediates:

**3-phosphoglycerate** & **pyruvate** Krebs cycle intermediates:

**Oxaloacetate** &  **$\alpha$ -ketoglutarate**.

Starting materials for biosynthesis of 11 **nonessential** amino acids: 1 step, 2 steps, or 3 steps



Alanine, aspartate, & glutamate use transamination

**Phenylketonuria (PKU):**

Defective phenylalanine hydroxylase – **phenylalanine** accumulates in body. Phenylalanine is transaminated to **phenylpyruvate**.

Accumulation of phenylpyruvate leads to severe mental retardation in infants. Persons suffering from phenylketonuria should not consume foods containing high levels of phenylalanine, such as aspartame.





## Hemoglobin catabolism

Red blood cells contain oxygen carrying pigments of a conjugated protein: Protein part is **Globin** Non-protein prosthetic group is **Heme**. **Heme** contains four pyrrole (**tetrapyrrole**) groups held together by an **iron** atom. Old red blood cells degraded in the spleen. Globin is hydrolyzed into amino acids. Iron atom stored in a protein (**ferritin**) **Tetrapyrrole** degraded to **bile pigments**.

Review: can you...

- Describe the steps in Protein digestion & absorption
- Explain how Amino Acids are utilized in the body
- Explain **Transamination** and **Oxidative De-amination**
- Describe **The Urea Cycle** – purpose and steps
- Describe how a.a. Carbon Skeletons are processed
- Define and explain Amino Acid Biosynthesis.
- Describe the chemical composition of urine.

# Lipid Metabolism



Fatty acids (F.A.s) are taken up by cells.

They may serve as:

- precursors in synthesis of other compounds
- fuels for energy production
- substrates for ketone body synthesis.

Ketone bodies may be exported to other tissues: used for **energy production**. Some cells **synthesize fatty acids** for storage or export.

### Energy

Fats are an important source of calories. Typically 30-40% of calories in American diet are from **fat**. Fat is the major form of **energy storage**.

Typical body fuel reserves are:

**fat:** 100,000 kcal.

**protein:** 25,000 kcal.

**carbohydrate:** 650 kcal.

Provides 60% of energy needs for body at rest TAG reserves would enable someone to survive starvation for ~30 days.

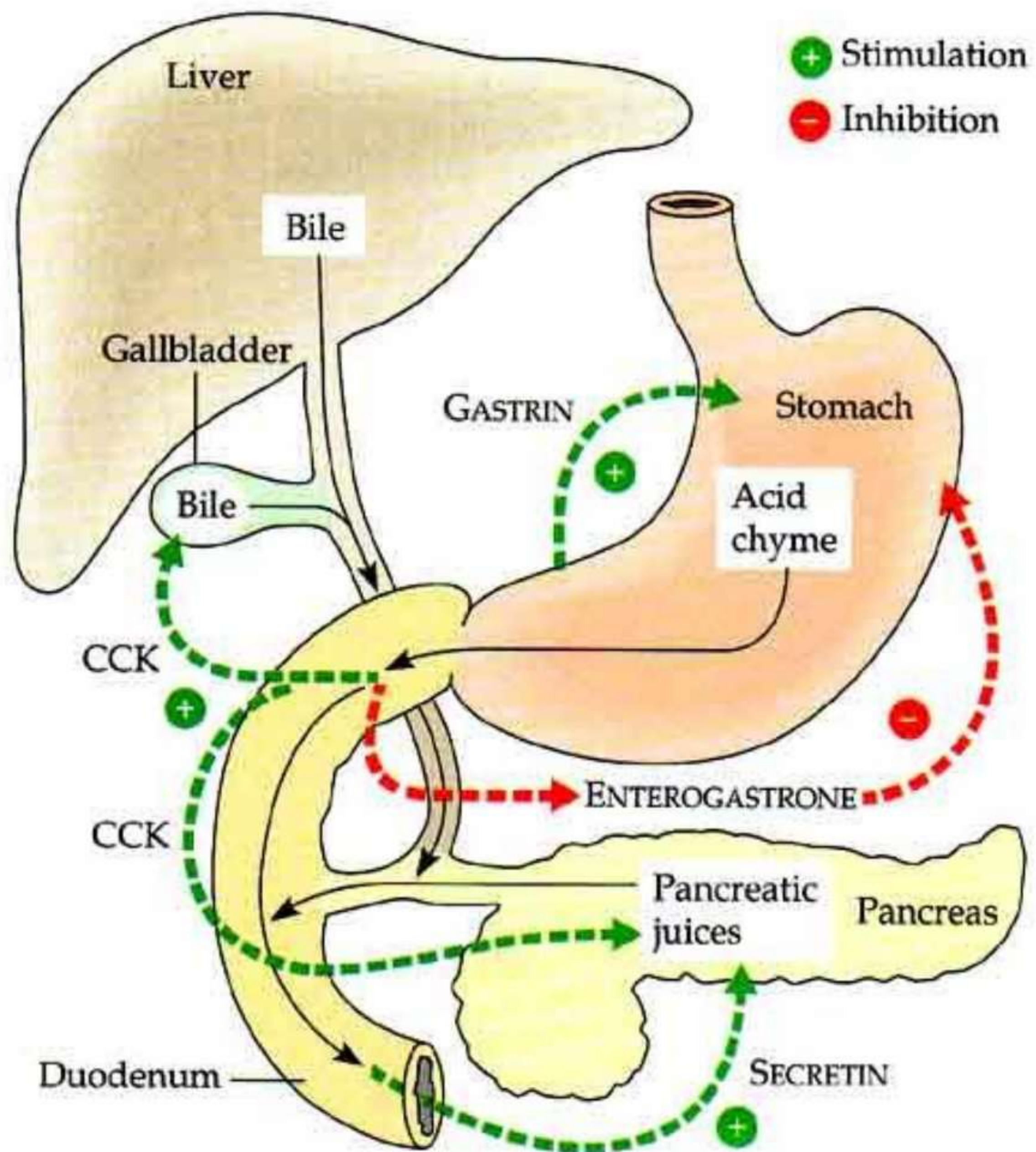
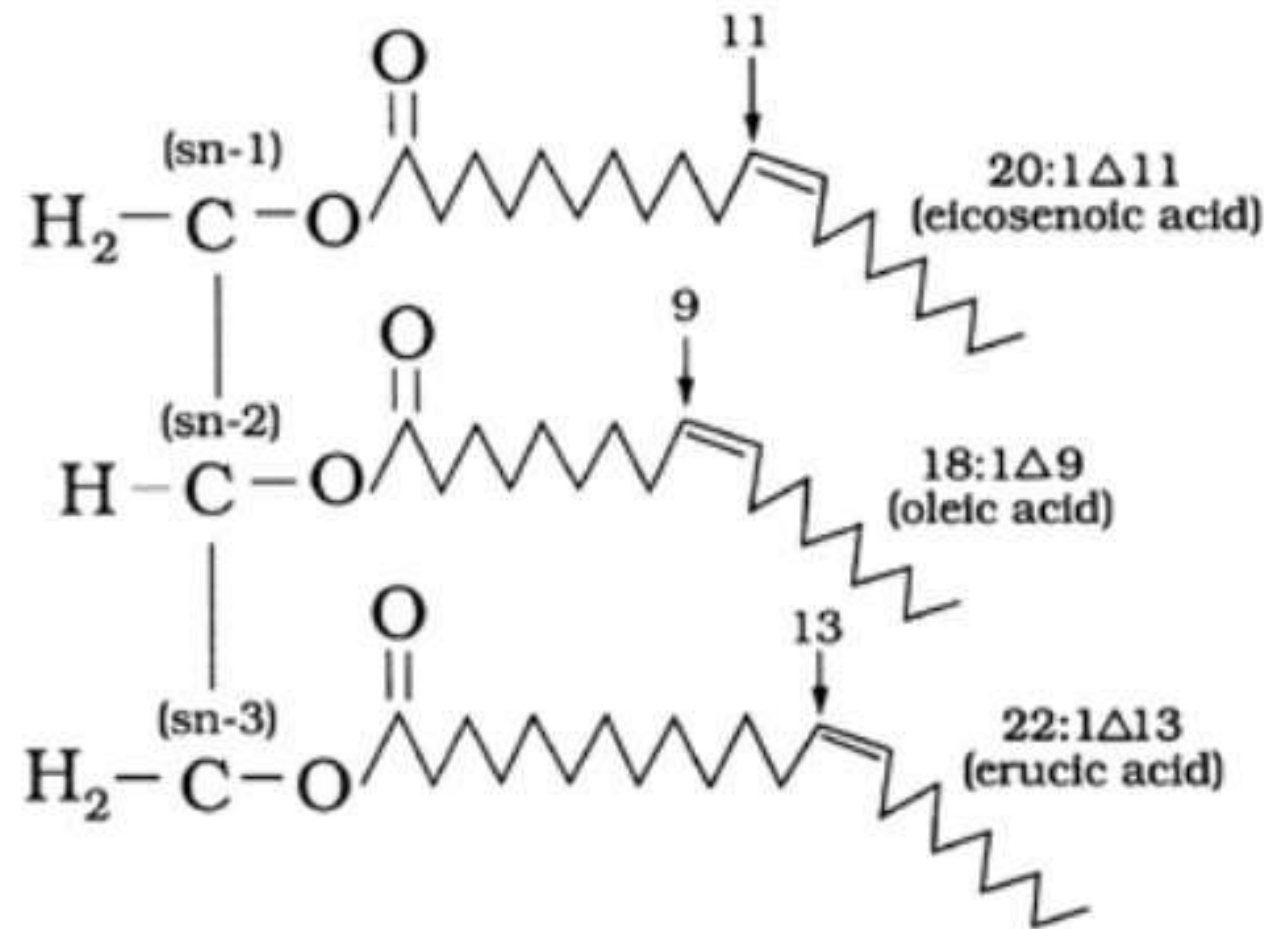
### Digestion and Absorption of Lipids

- 98% of ingested lipids are triacylglycerols (TAGs)
- Digestion in the Mouth: enzymes are **aqueous**-little effect on lipids
- Digestion in the Stomach: causes a large **physical** change-

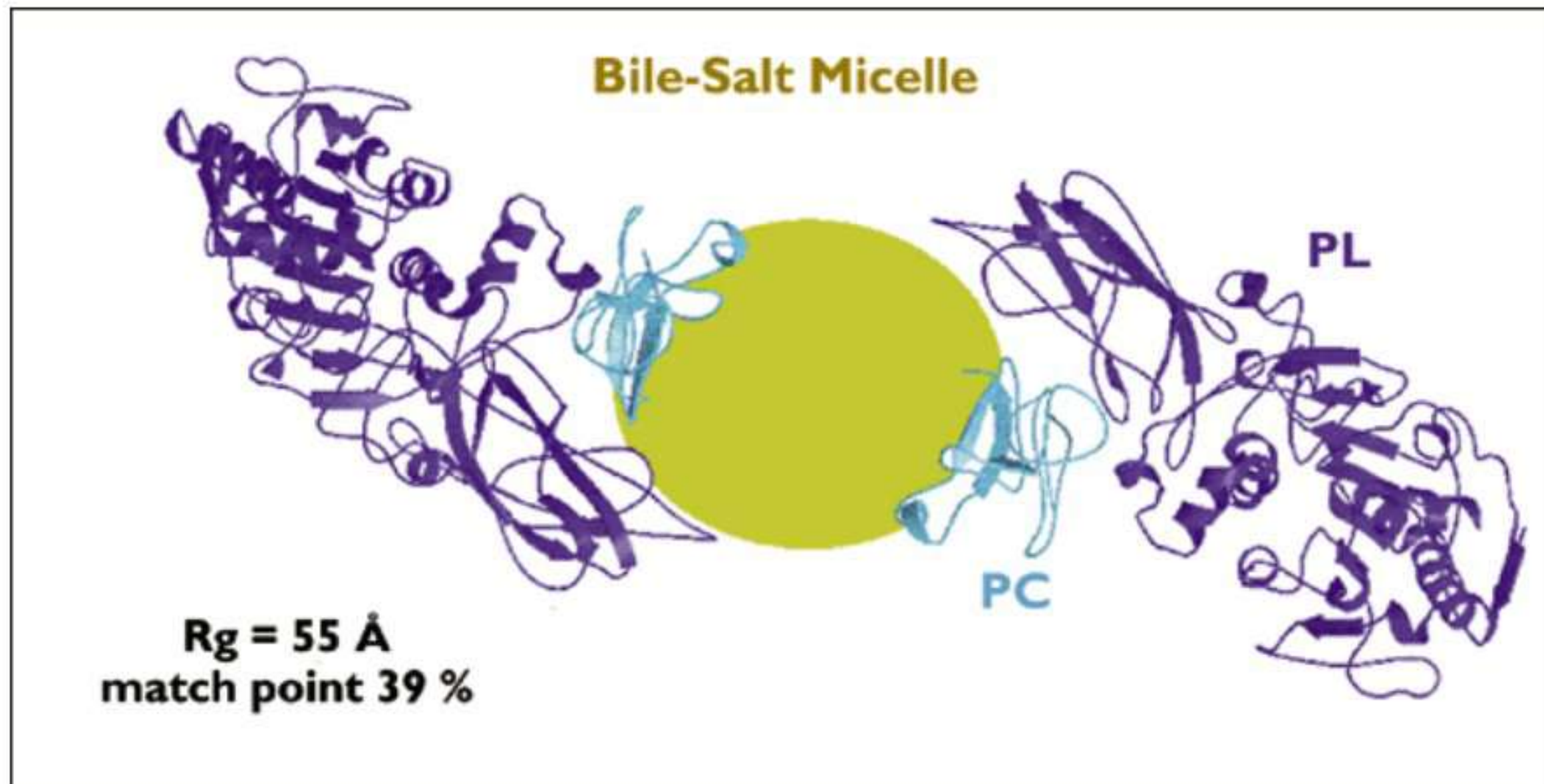
Churned into droplets:

**“Chyme”**

# TRIACYLGLYCEROL

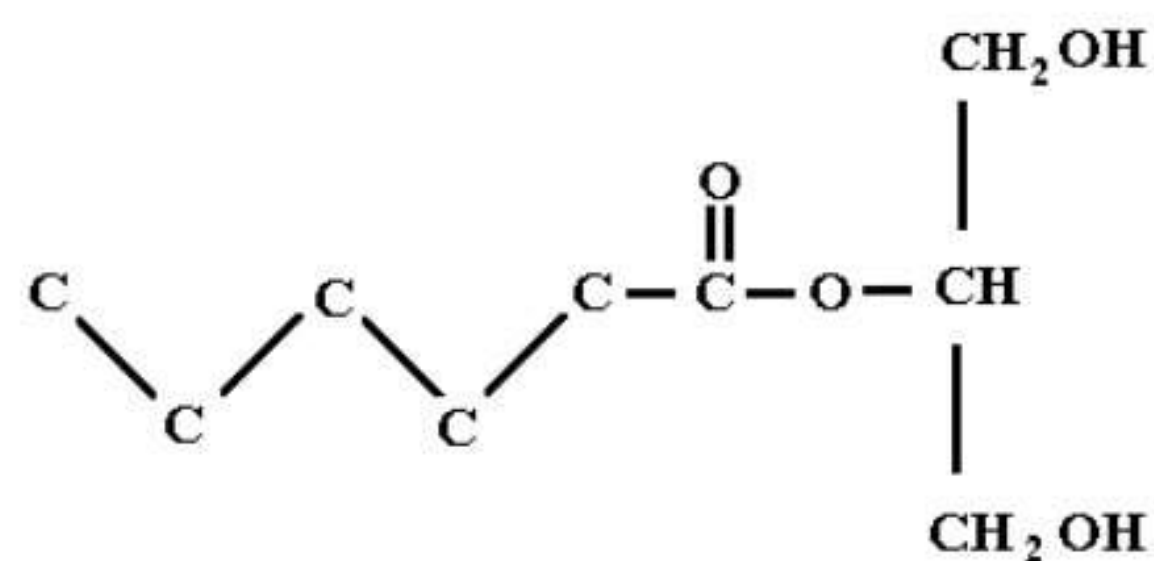


**Gastric Lipase:** Begins actual lipid digestion. ~10% of TAGs are hydrolyzed in the **stomach**. Chyme stimulates **cholecystinin** (CCK) to release **bile** from gallbladder. Bile is an emulsifier

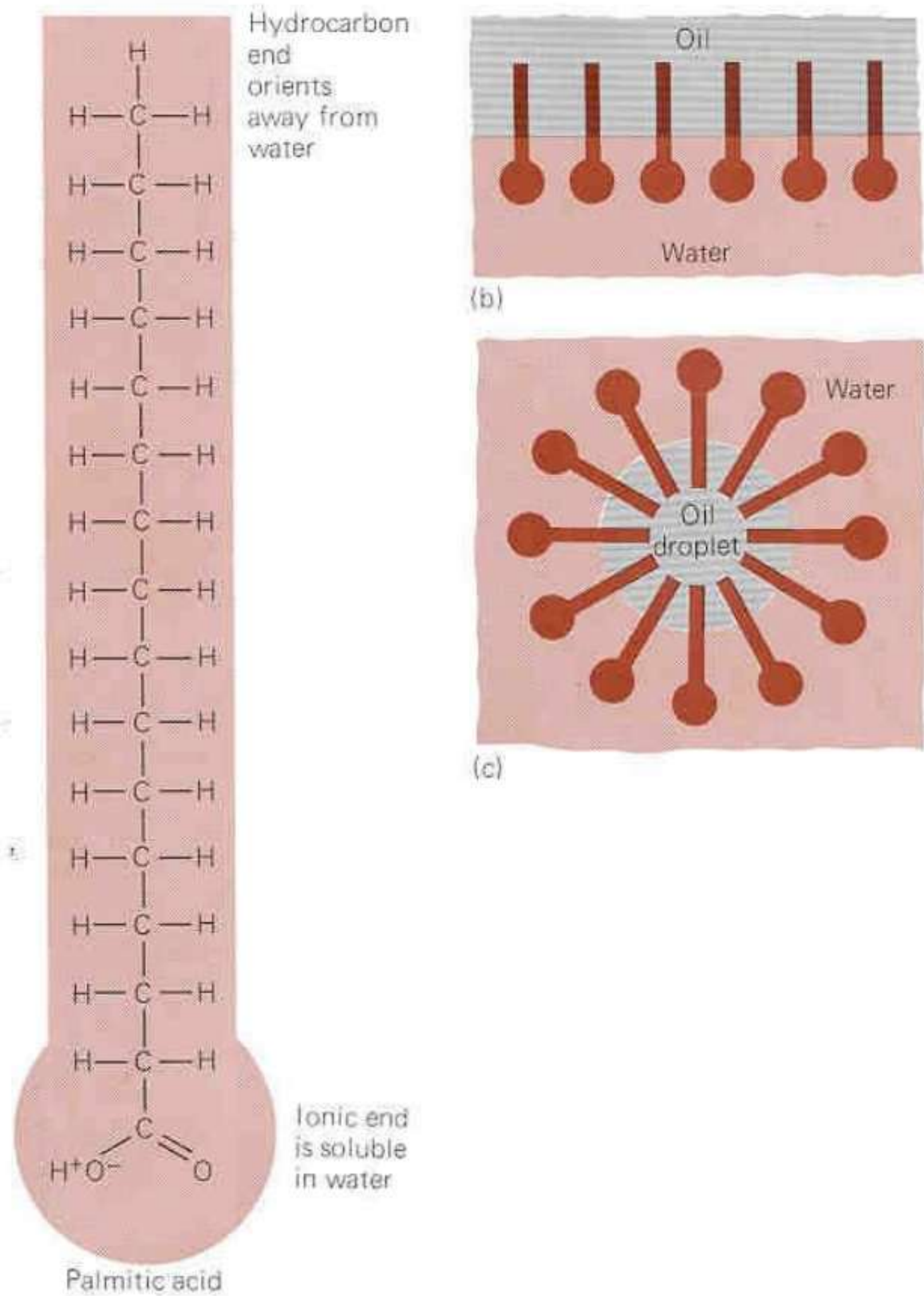
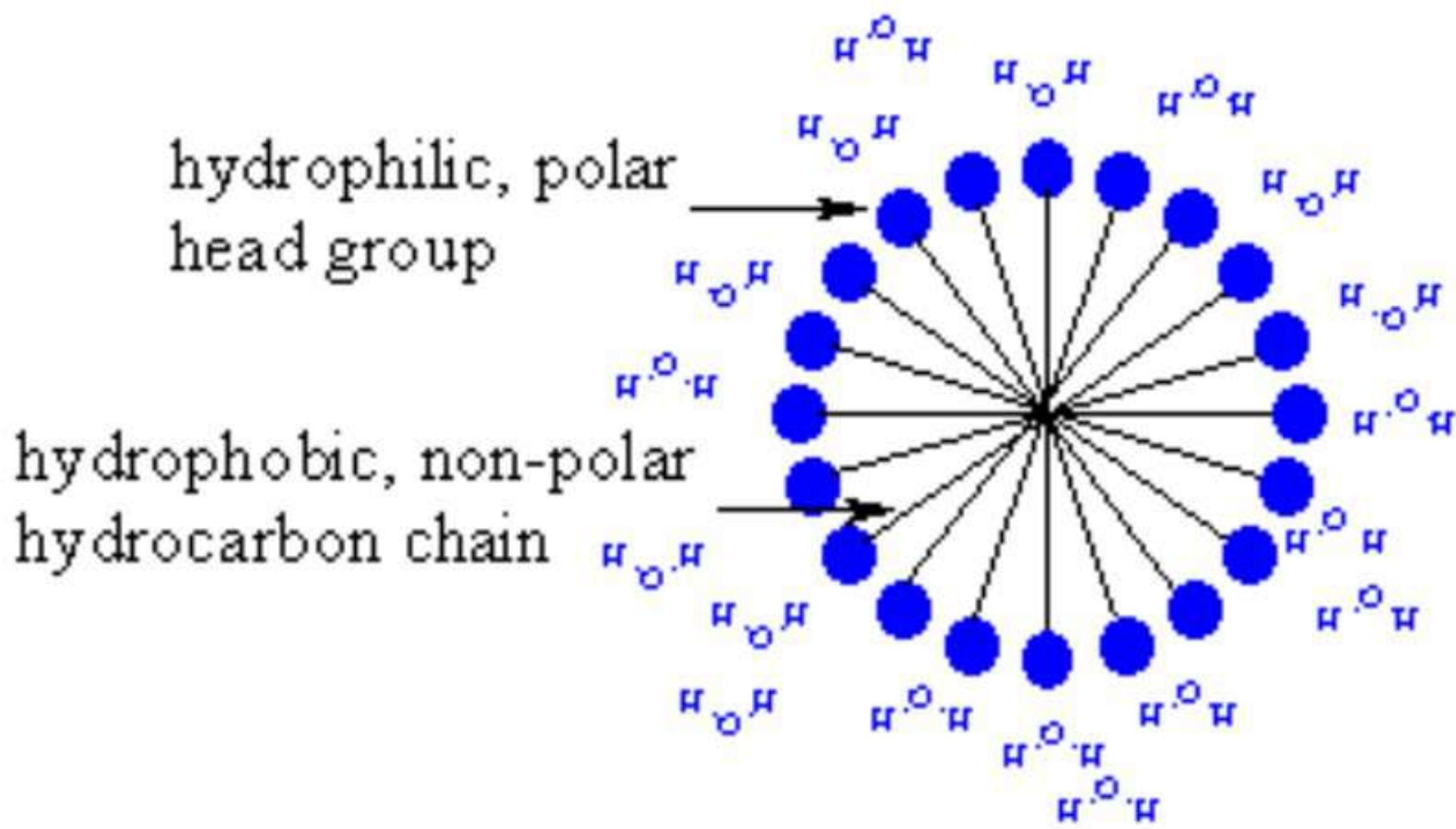


Pancreatic lipase (PL) hydrolyzes insoluble triglyceride by binding to the **bile-salt micelles**. TAGs are **partially** hydrolyzed: 2 of the 3 F.A.s have ester linkages hydrolyzed and are released.

**Monoacylglycerol** remains = glycerol and 1 fatty acid



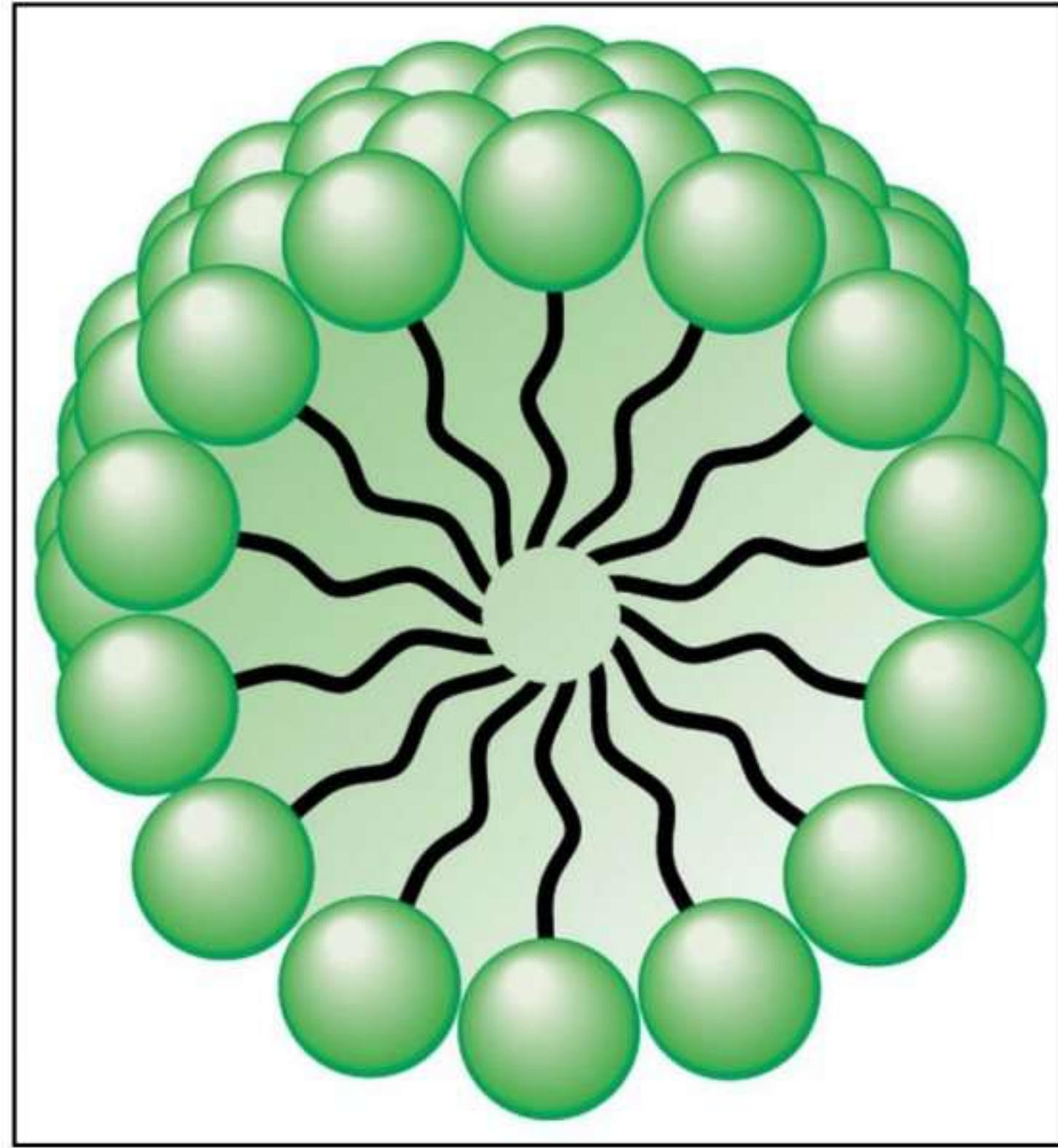
Oil droplets will form spherical **micelle** shapes. Bile salts aid this process clumping fatty acids and monacylglycerols.



Fatty acid micelle: **hydrophobic** fatty acids & monoacylglycerols are in the interior. Bile salts on exterior.

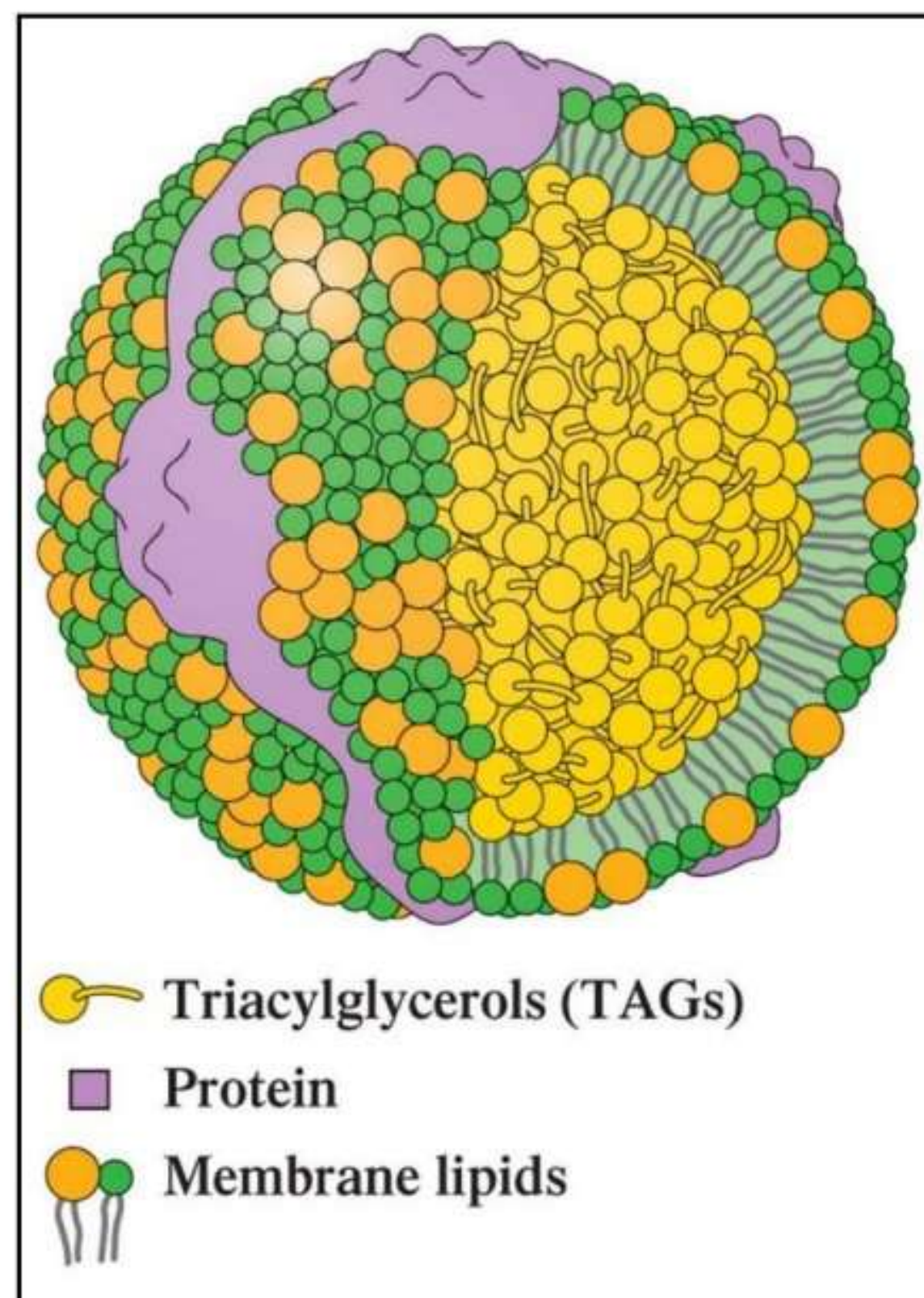
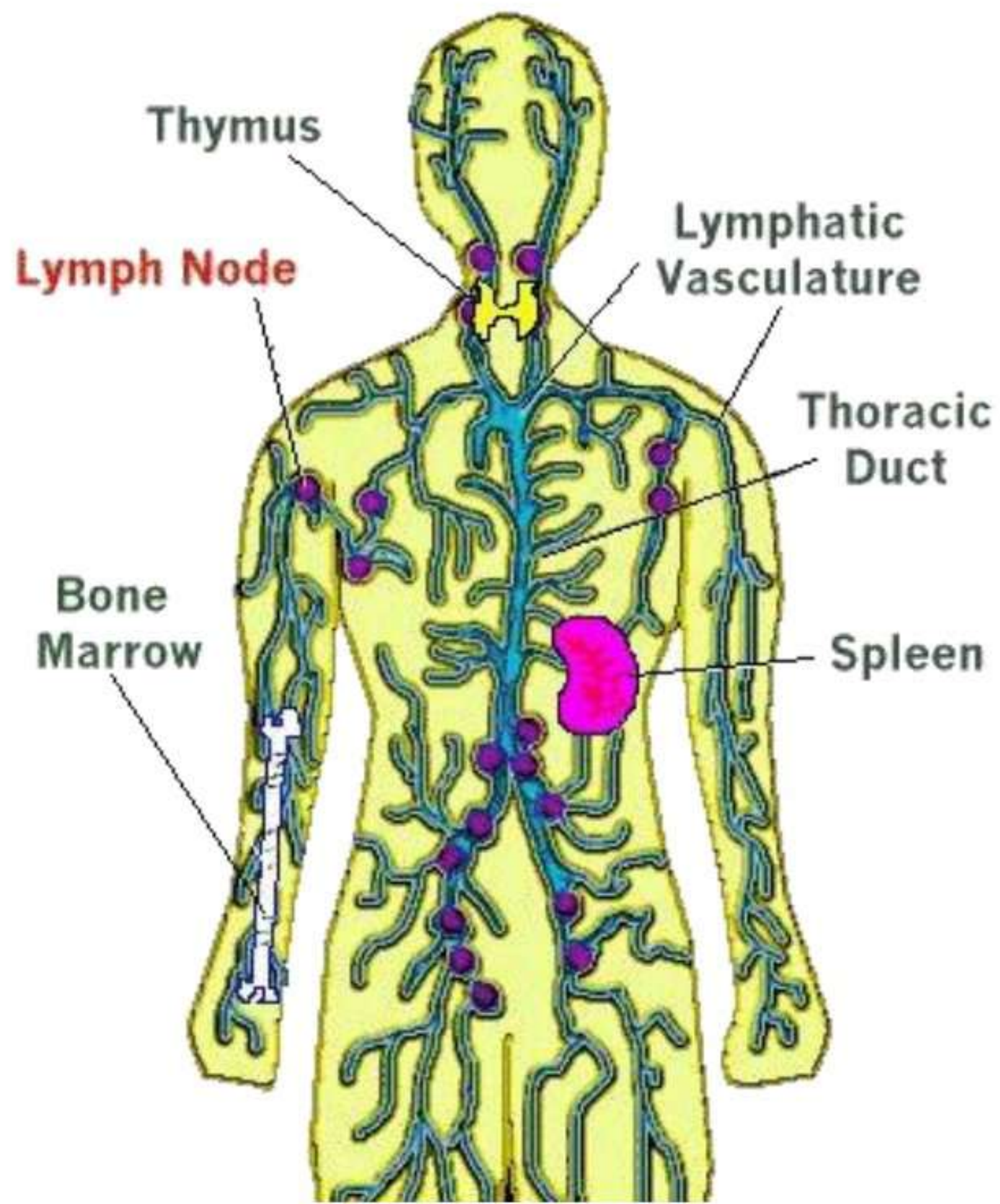
Micelles are small enough to penetrate membrane of intestinal cells. Free fatty acids & monoacylglycerols are reformed into

**triacylglycerols**.



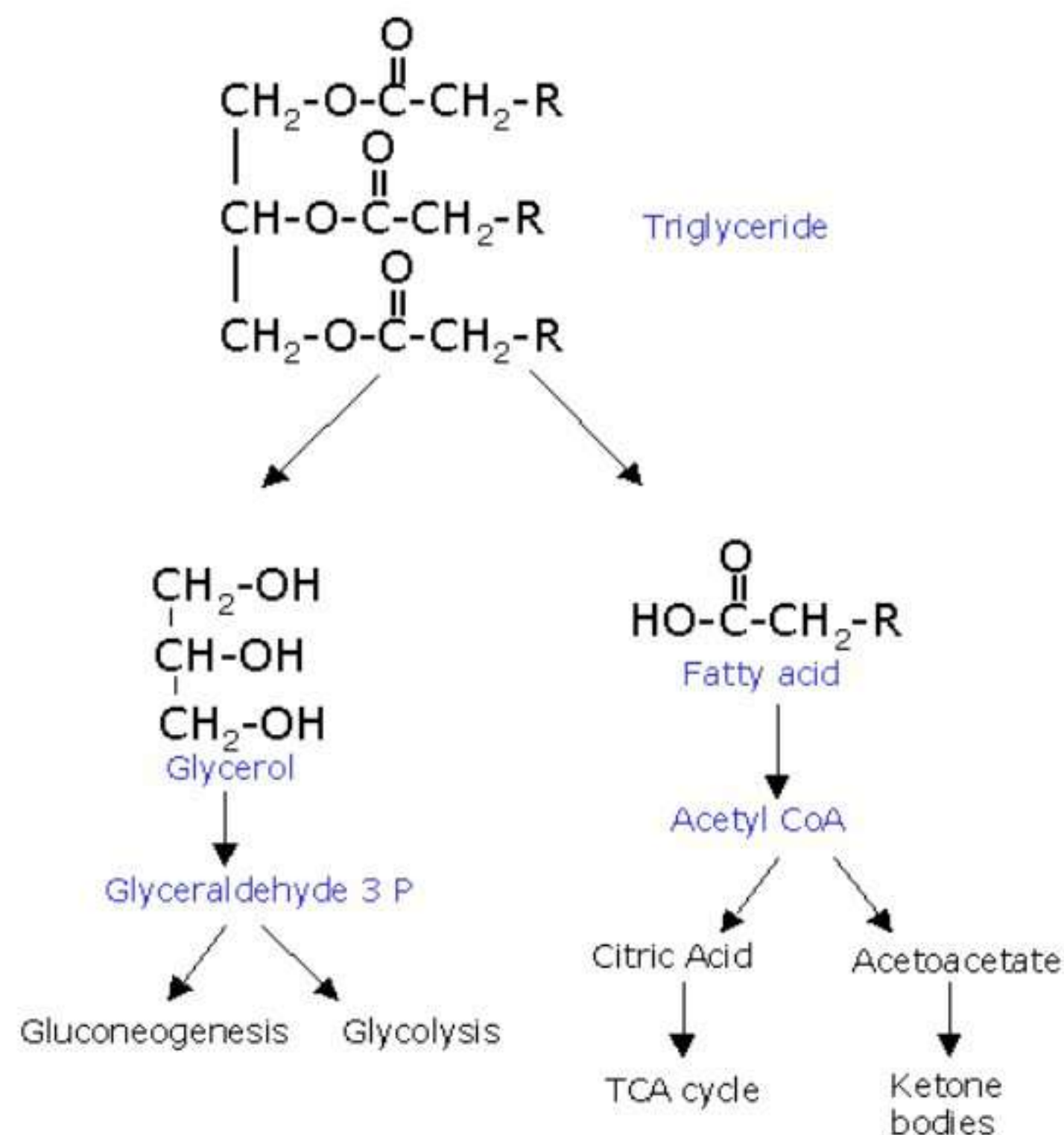
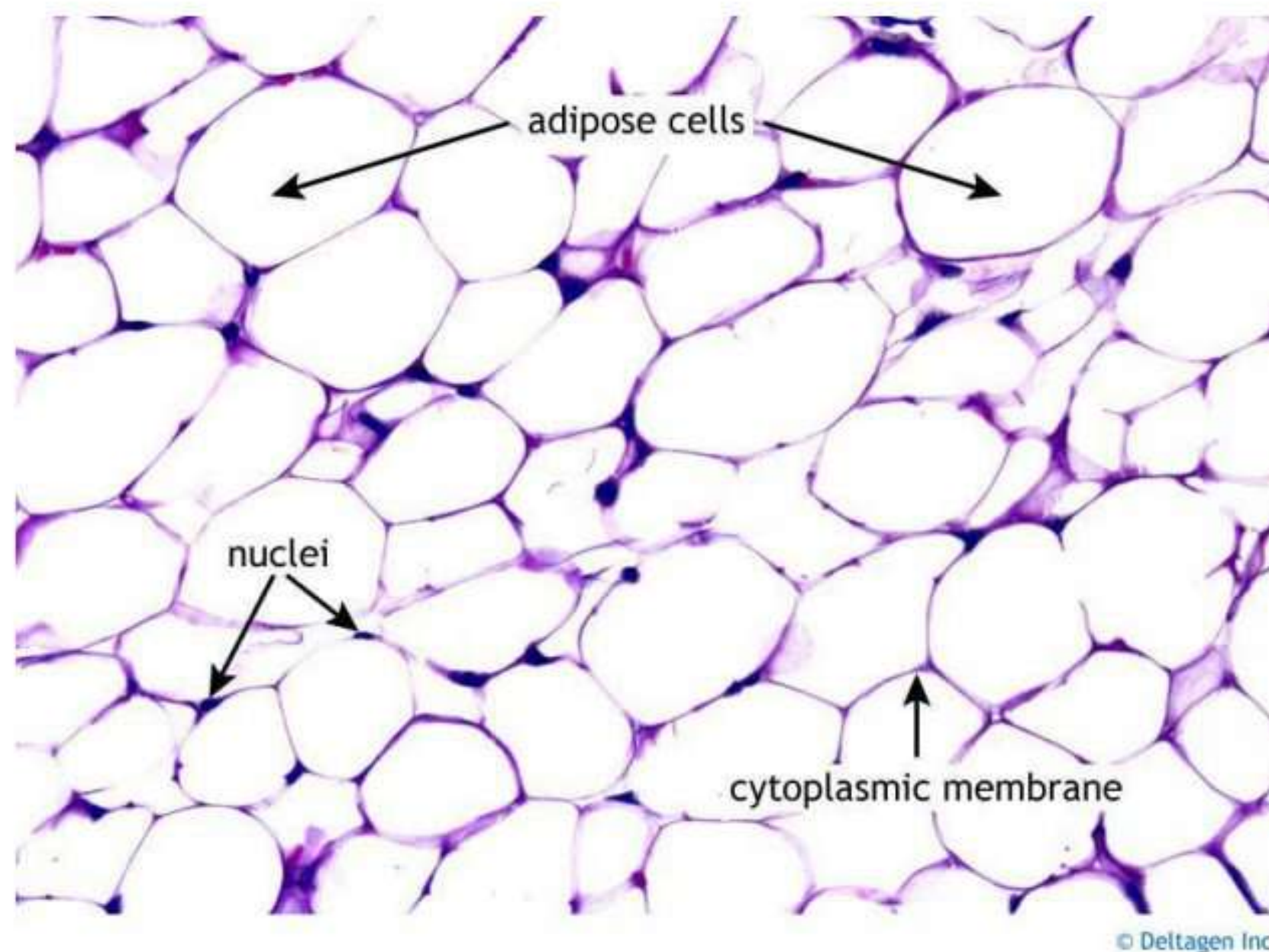
TAGs are combined with membrane & water soluble proteins to form a **chylomicron**, a lipoprotein.

**Chylomicrons** carry TAGs from intestinal cells into bloodstream via the **lymph system**.

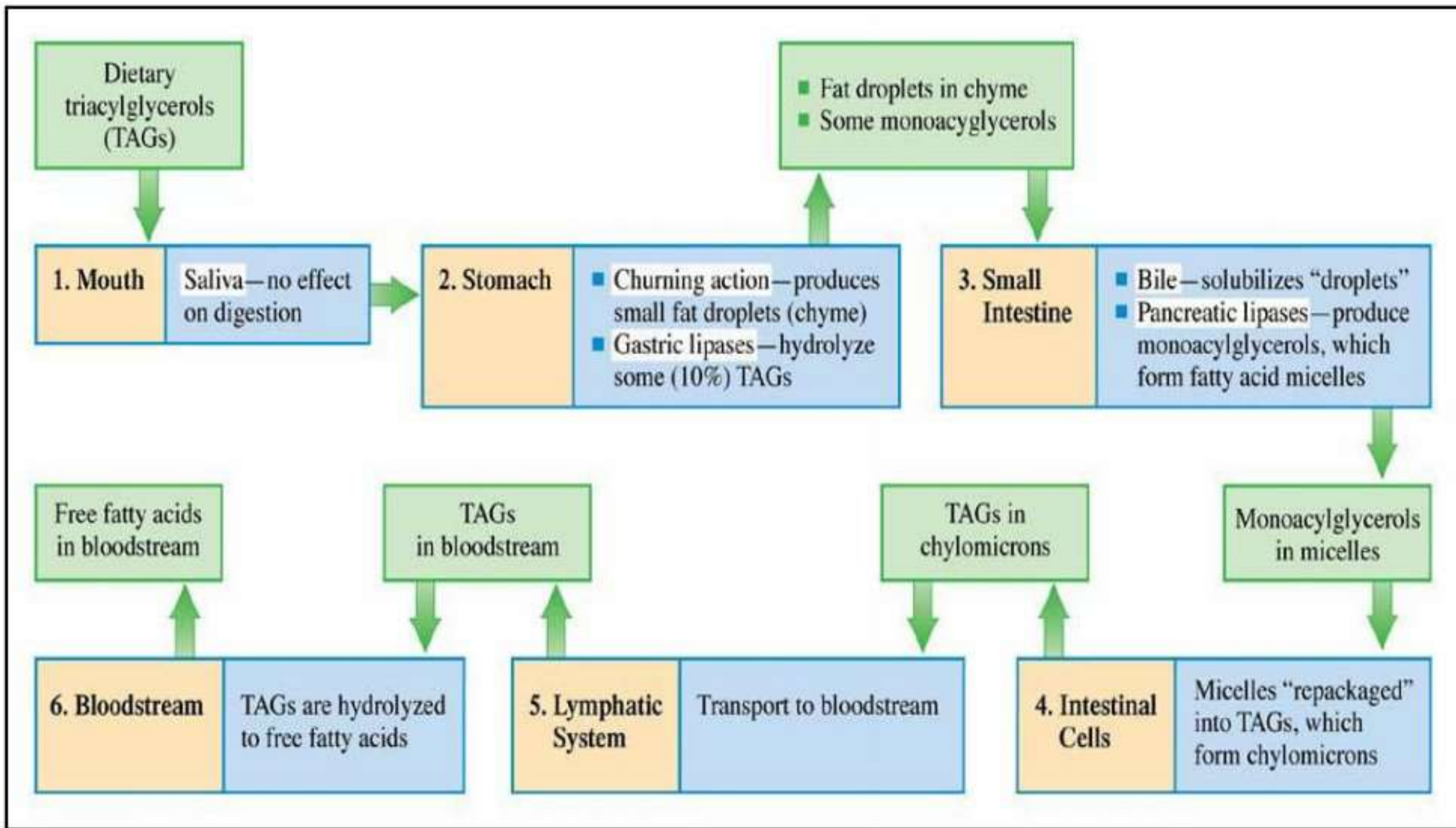




Triacylglycerols reach bloodstream & are hydrolyzed down to **glycerol** and **fatty acids**. These are absorbed by cells and processed further for energy by forming **acetyl CoA**. Or Stored as lipids in fat cells (adipose tissue).

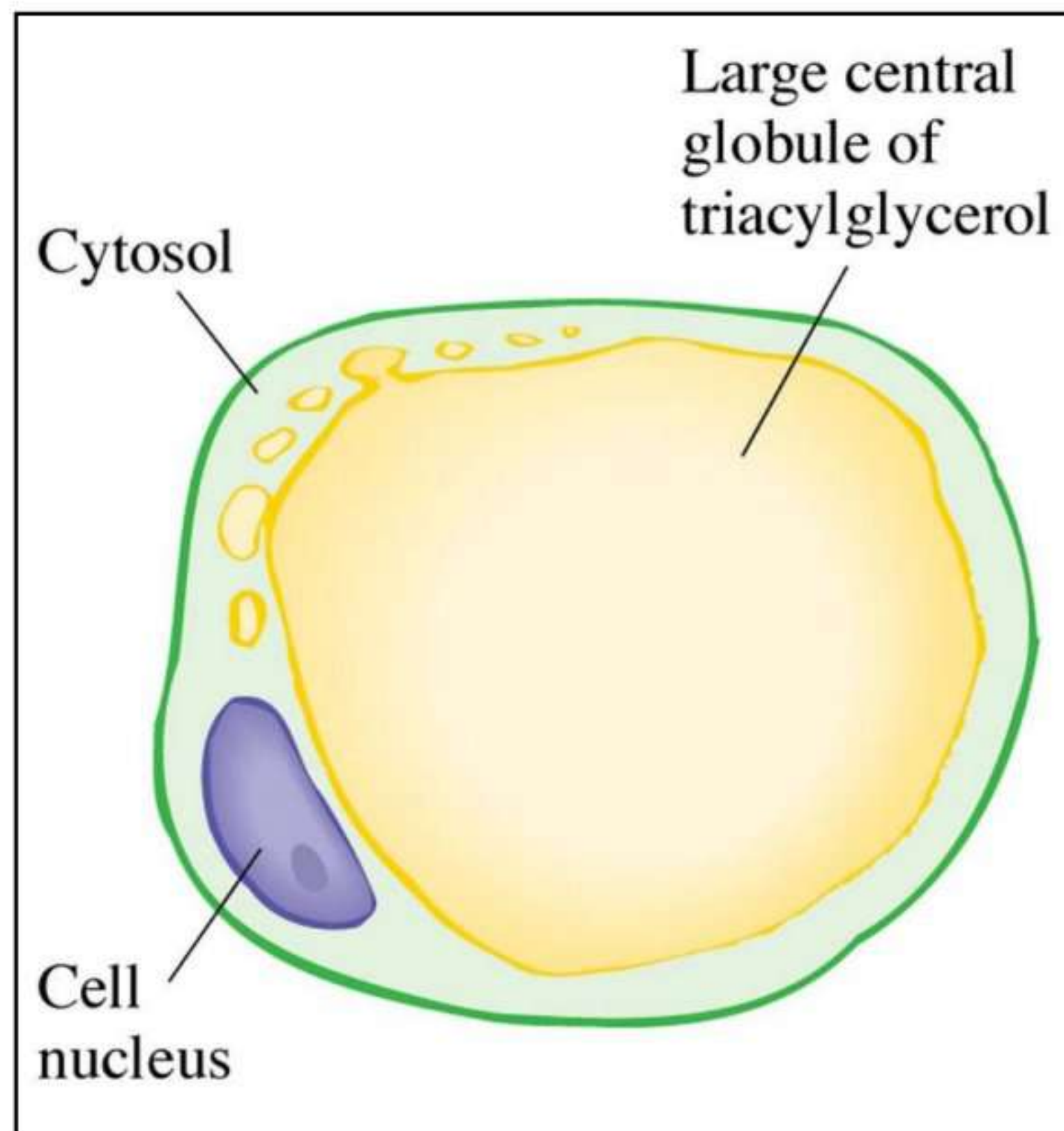


Summary of events that must occur before triacylglycerols (TAGs) can reach the bloodstream through the digestive process.



### Triglyceride Storage & Mobilization

**Storage of triacylglycerol** is in **adipocytes**. Fatty acids stored primarily as triacylglycerol. Triacylglycerol is **hydrolyzed** to release **fatty acids** when needed.



## **Hormonal control of lipolysis**

The breakdown of triglycerides by lipases is under hormonal control.

### **Hormones involved are:**

Epinephrine, glucagon, and insulin.

### **Epinephrine & glucagon:**

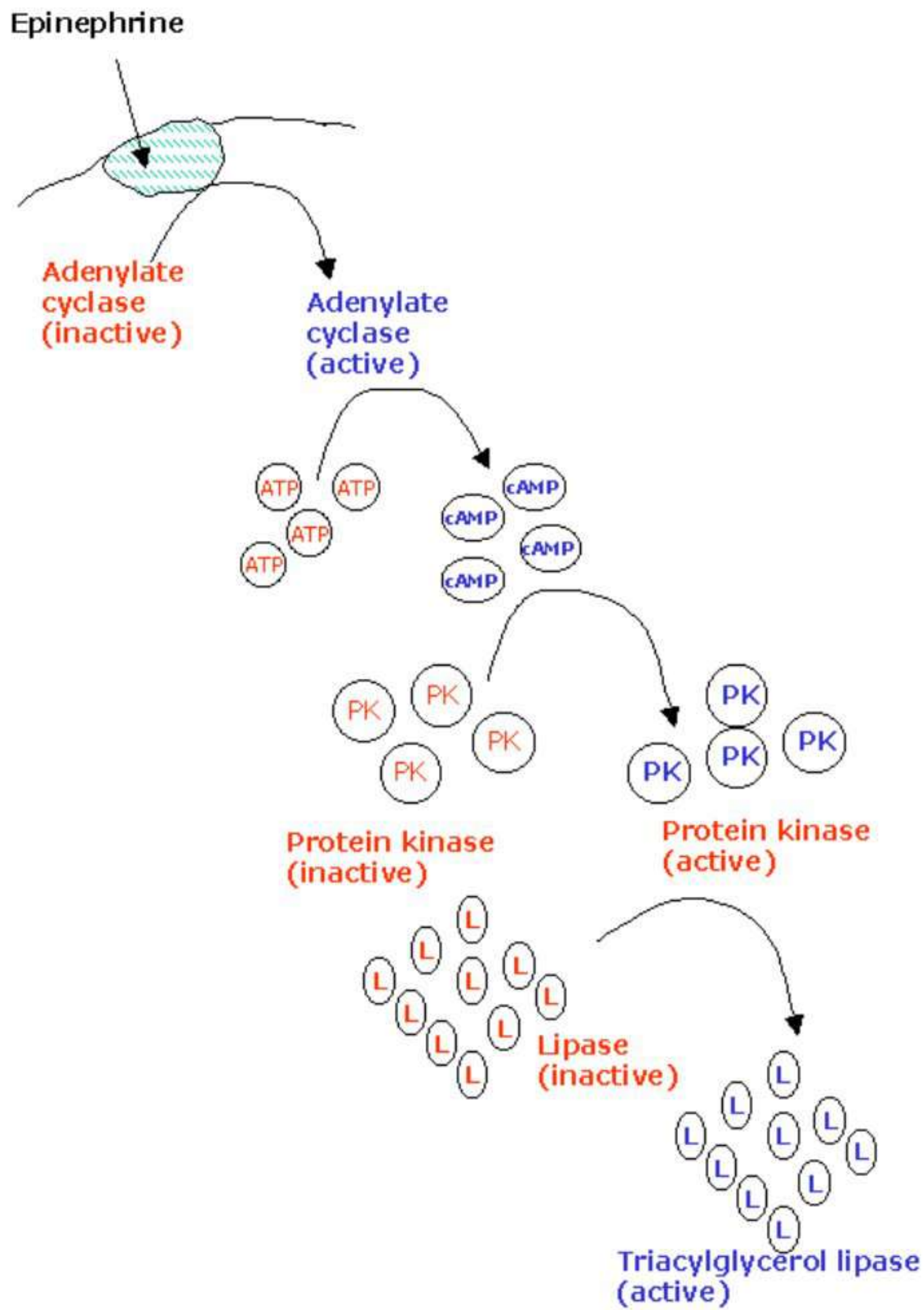
promote breakdown of fat (lipolysis)

### **Insulin:**

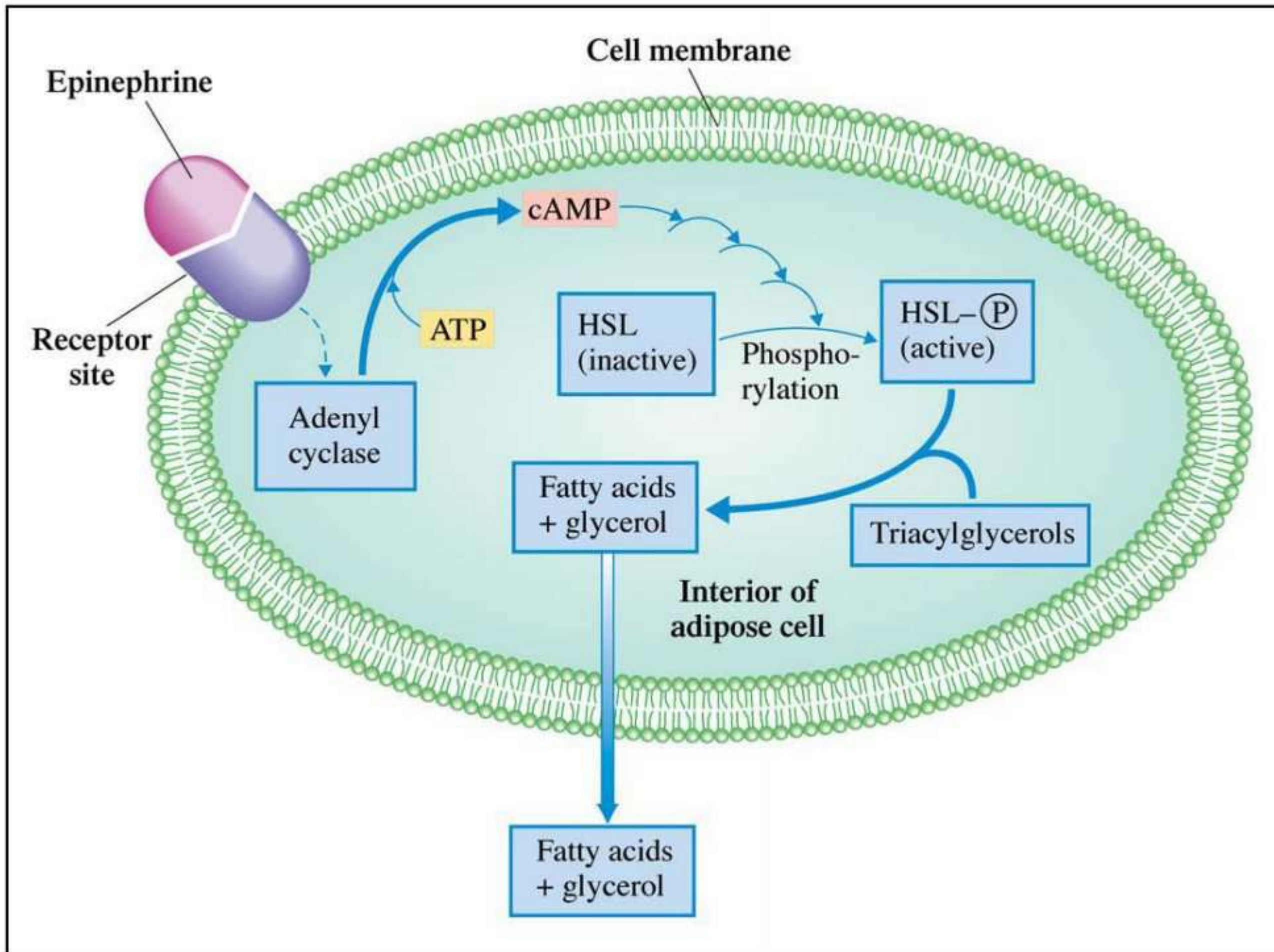
inhibits lipolysis.

### **Triacylglycerol Mobilization:**

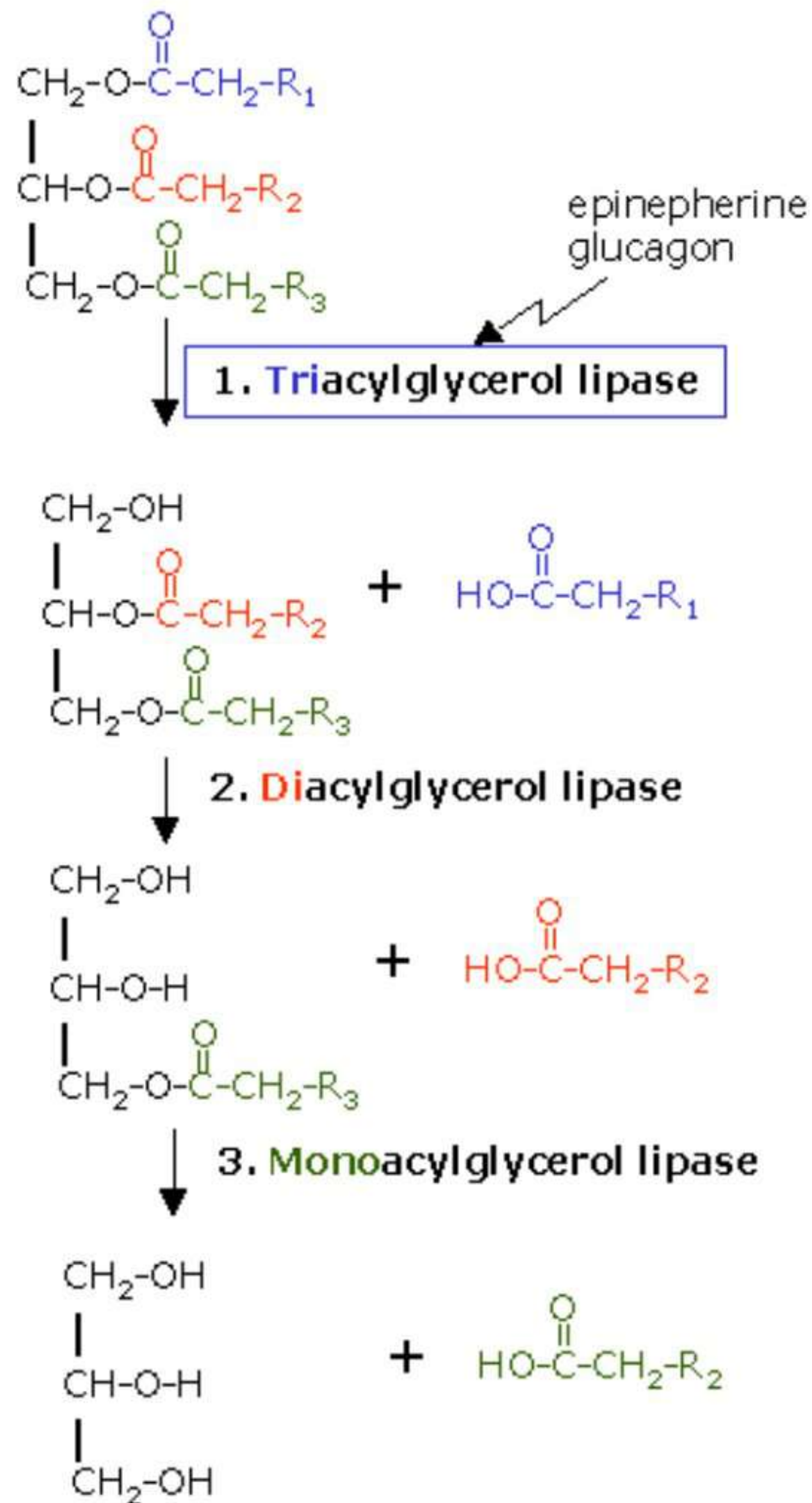
Hydrolyzing lipid reserves in adipose tissue for **energy**. Triggered by hormones~10% TAGs replaced in adipose tissue daily as they get used up for energy.



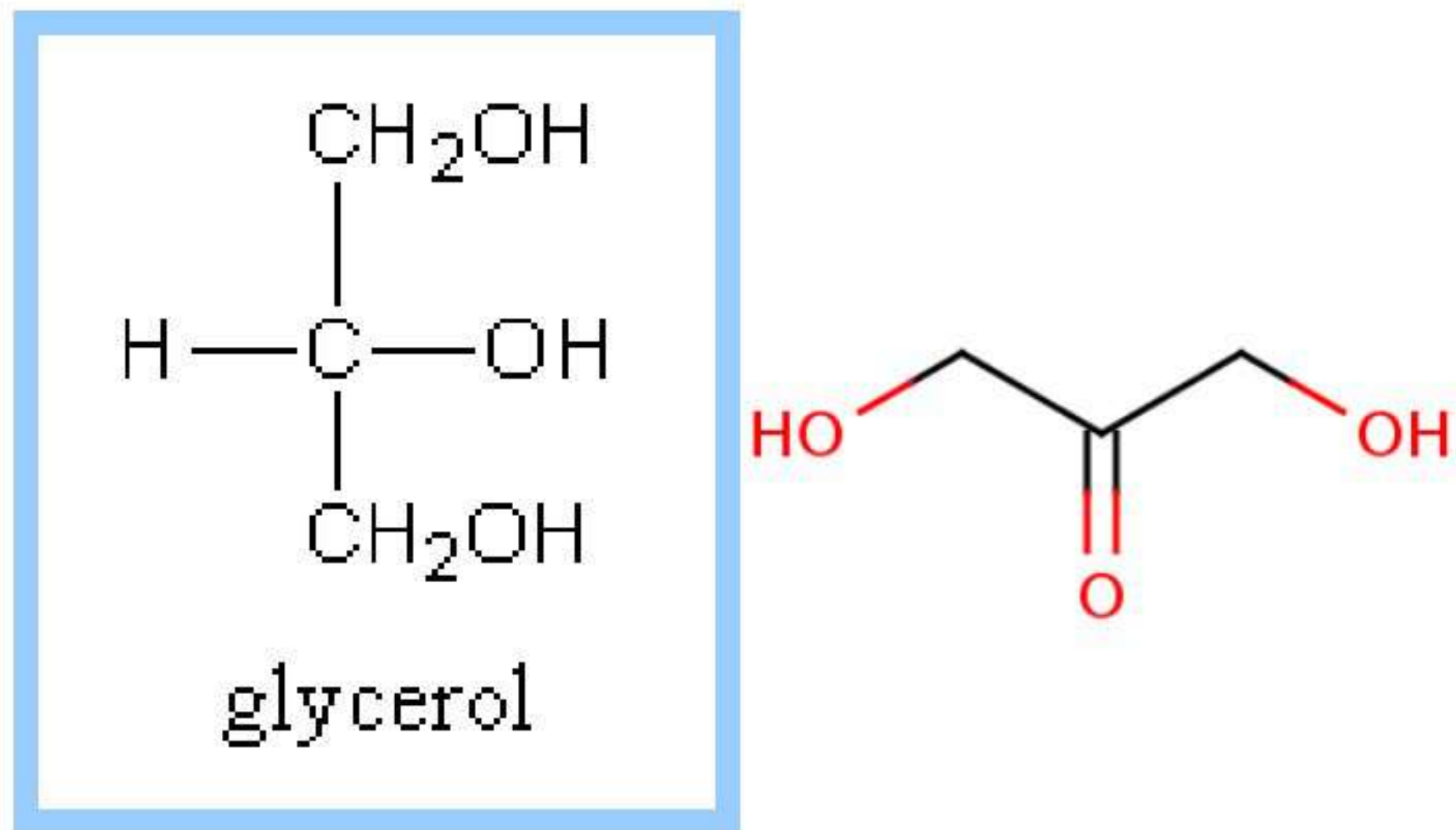
Hydrolysis of stored triacylglycerols in adipose tissue is triggered by hormones that stimulate cAMP production within adipose cells.



Third time is a charm! TAGs hydrolyzed a 3<sup>rd</sup> time to form fatty acids.  
**Tri**acylglycerol lipase **Di**acylglycerol lipase **Mon**oacylglycerol lipase  
 Only triacylglycerol lipase is activated by epinephrine.

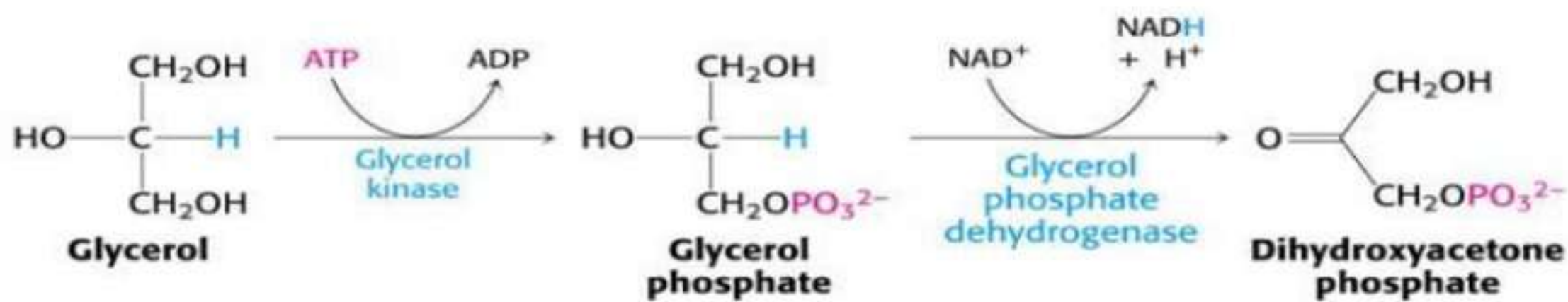


One glycerol formed for each TAG hydrolyzed. Enter bloodstream & go to liver or kidneys for processing. Converted in 2 steps to **Dihydroxyacetone phosphate**



Where will the phosphate be attached?

Uses up one ATP. Reduces one  $\text{NAD}^+$  to  $\text{NADH}$



Primary hydroxyl group is phosphorylated  
**Dihydroxyacetone phosphate**  
 is an intermediate for both

**Glycolysis:**

converted to Pyruvate, then to Acetyl CoA, & eventually to  $\text{CO}_2$ ,  
 releasing its energy.

**Gluconeogenesis:**

creates Glucose from **non-carbohydrate** source

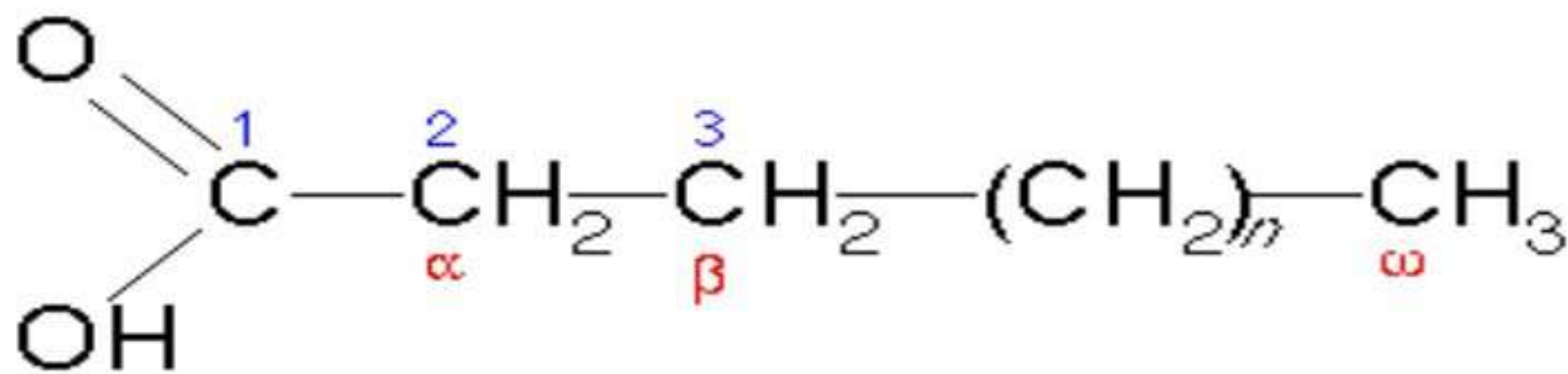
Lipid metabolism & carbohydrate metabolism

are connected.

Fatty acids can also be broken down for energy. What kind of reaction is needed?

**Oxidation!**

Quick review first on fatty acid numbers & letters:



**Fatty acid numbering system**

**Review Important fatty acids:**

<u>Name</u>	# Carbons: (saturation)
Palmitate	16:0
Stearate	18:0
Palmitoleate	16:1 - cis at C9
Oleate	18:1 - cis at C9
Linoleate	18:2 - cis at C9 and C12
Linolenate	18:3 - cis at C9, C12 & C15



## Lipid Metabolism

Lipid nomenclature

- Oxidation of Fatty acids
- β-oxidation
- Ketone Bodies

### Lipid nomenclature

Fatty acids

- triacylglycerols: know structure
- phospholipids
- waxes
- sphingolipids
- Glycosphingolipids
- Isoprenoids
- Steroids
- Nomenclature
- saturated: palmitate, stearate, no double bonds
- unsaturated: palmitoleate, Oleate: double bond at cis9 position
- polyunsaturated
- Melting points: saturated vsunsaturated

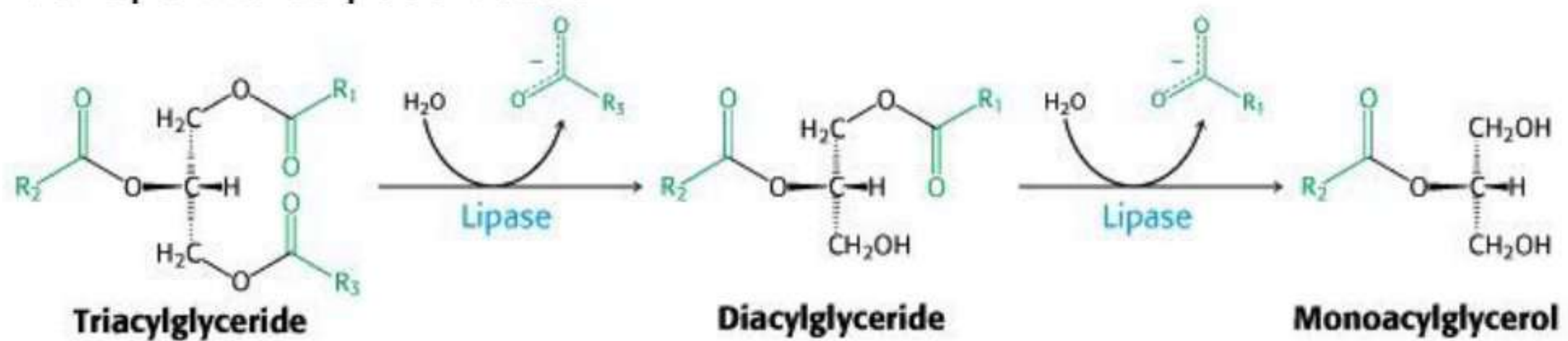
### Oxidation of Fatty acids

- Know equation for palmitate:  $C_{16}H_{32}O + O_2 \rightarrow CO_2 + H_2O$
- Comparison of glucose with palmitatefor ATP production and energy yield
- Mobilization of Triacylglycerols from adipose tissue
- hormonal control: glucagon, epinephrine

- lipases
- transport by lipoproteins
- fate of glycerol
- transport into cytoplasm of cell

## Digestion of lipid in diet

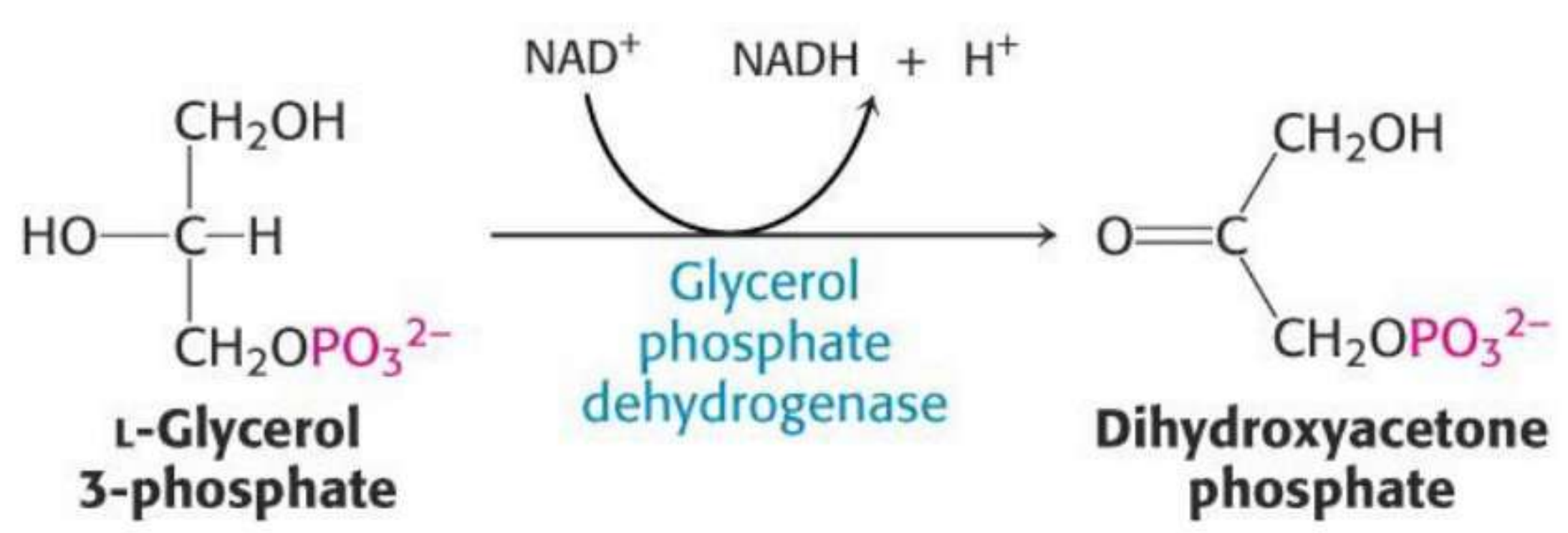
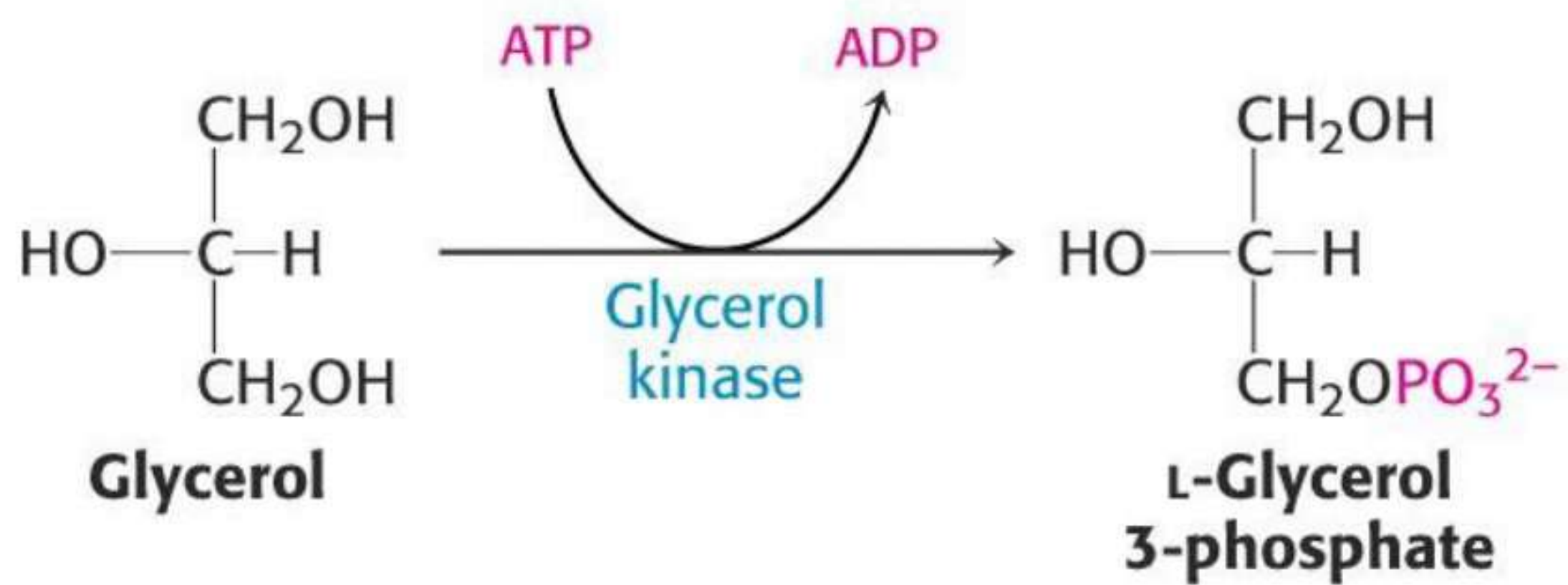
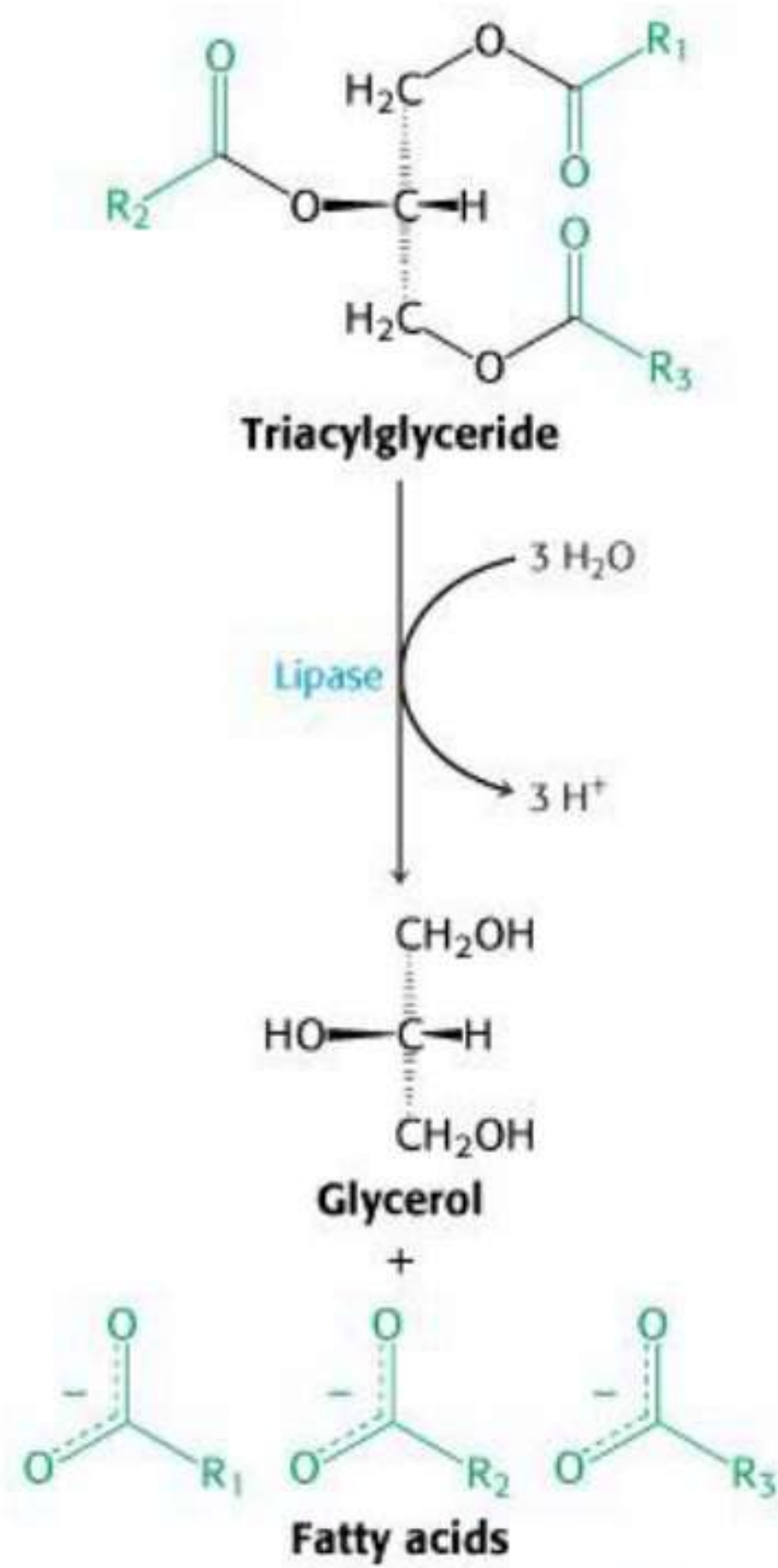
- Triacylglycerols from diet
- broken down in small intestine
- lipases
- bile salts
- transport to adipose tissue



## Mobilization of Triacylglycerols

- hormonal control of lipolysis: glucagon, epinephrine
- lipases
- transport by lipoproteins
- transport into cytoplasm of cell
- Insulin inhibits lipolysis

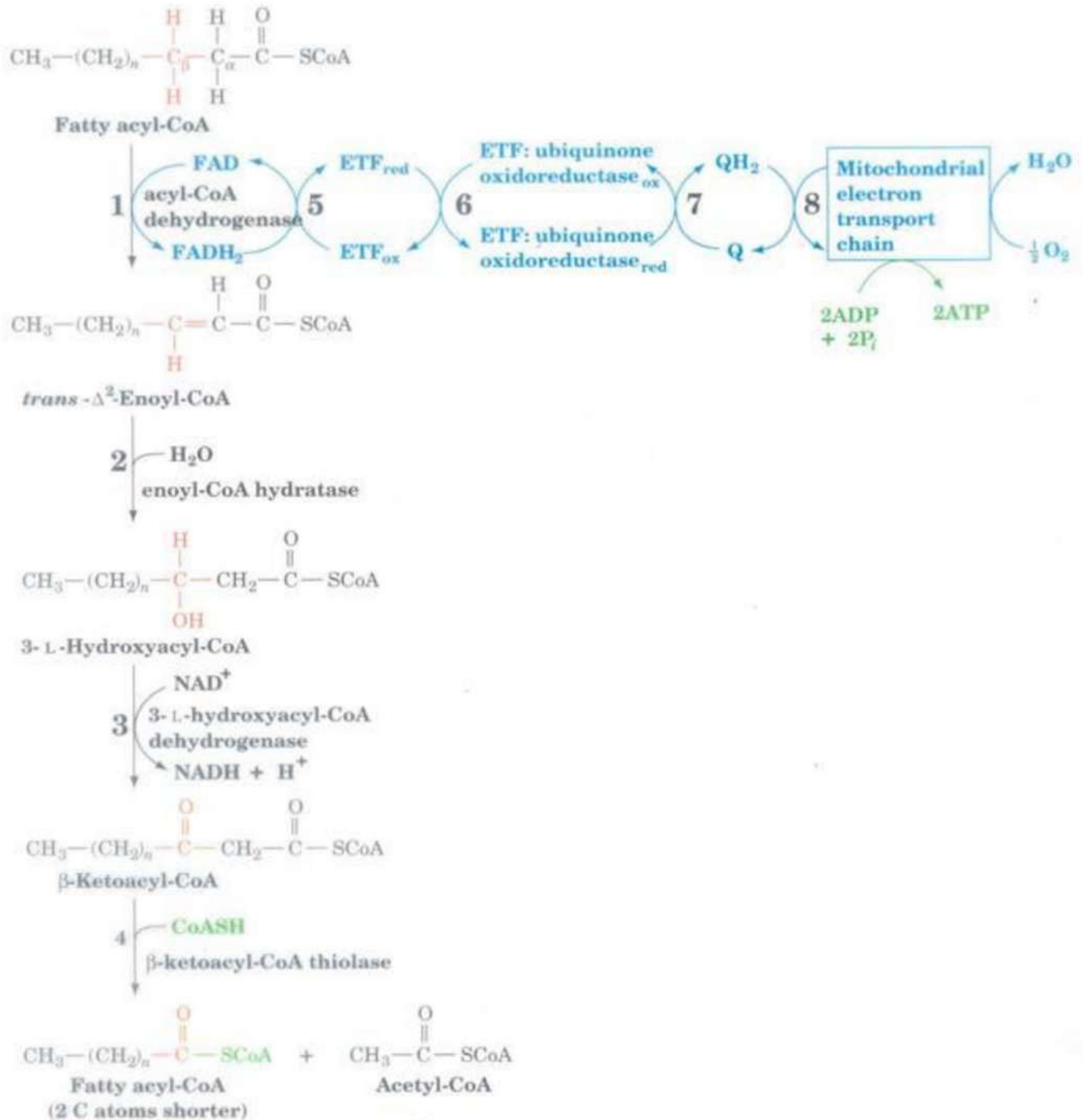
## Breakdown of triacylglycerides



**fate of glycerol**

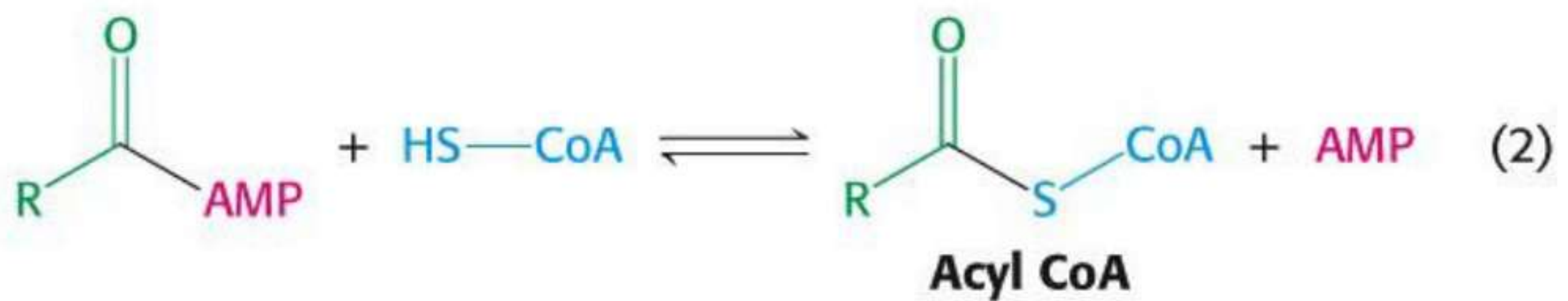
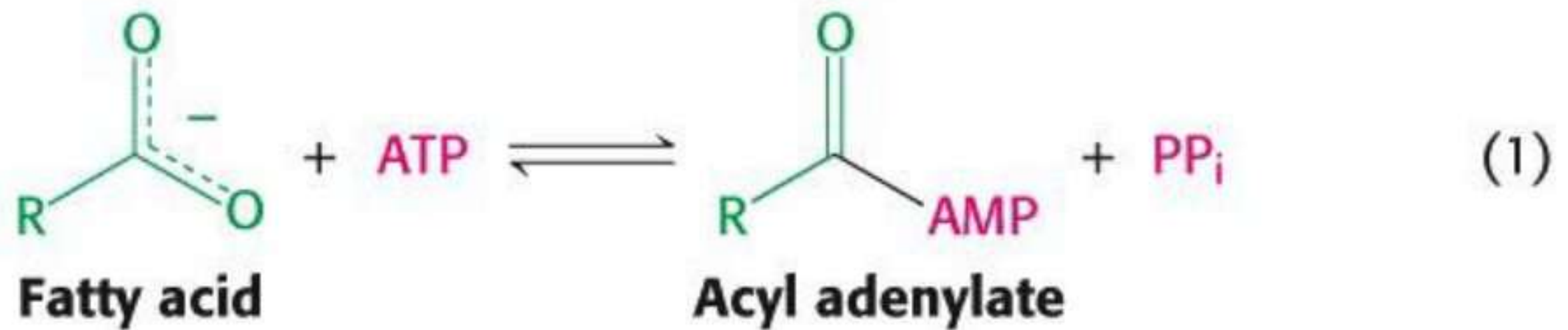
## β-oxidation

- occurs in mitochondria
- uses FAD and NAD
- produces acetyl CoA



## acylCoA synthetase

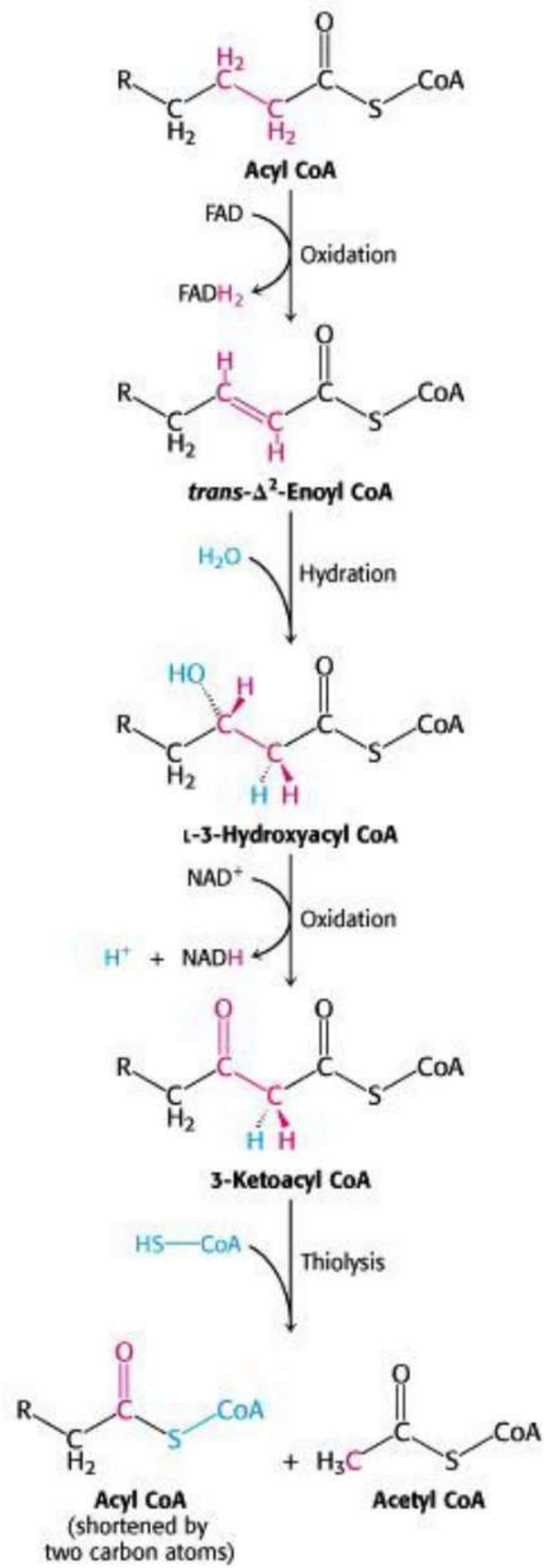
- two step reaction
- ATP + FA  $\rightarrow$  AMP-FA
- AMP-FA + CoASH  $\rightarrow$  FA-CoA + AMP



## **$\beta$ -oxidation**

AcylCoA dehydrogenase

- enoyl-CoA hydratase
- L-hydroxyacyldehydrogenase
- ketoacyl-CoA thiolase
- Repeat steps



## Summary of Reactions

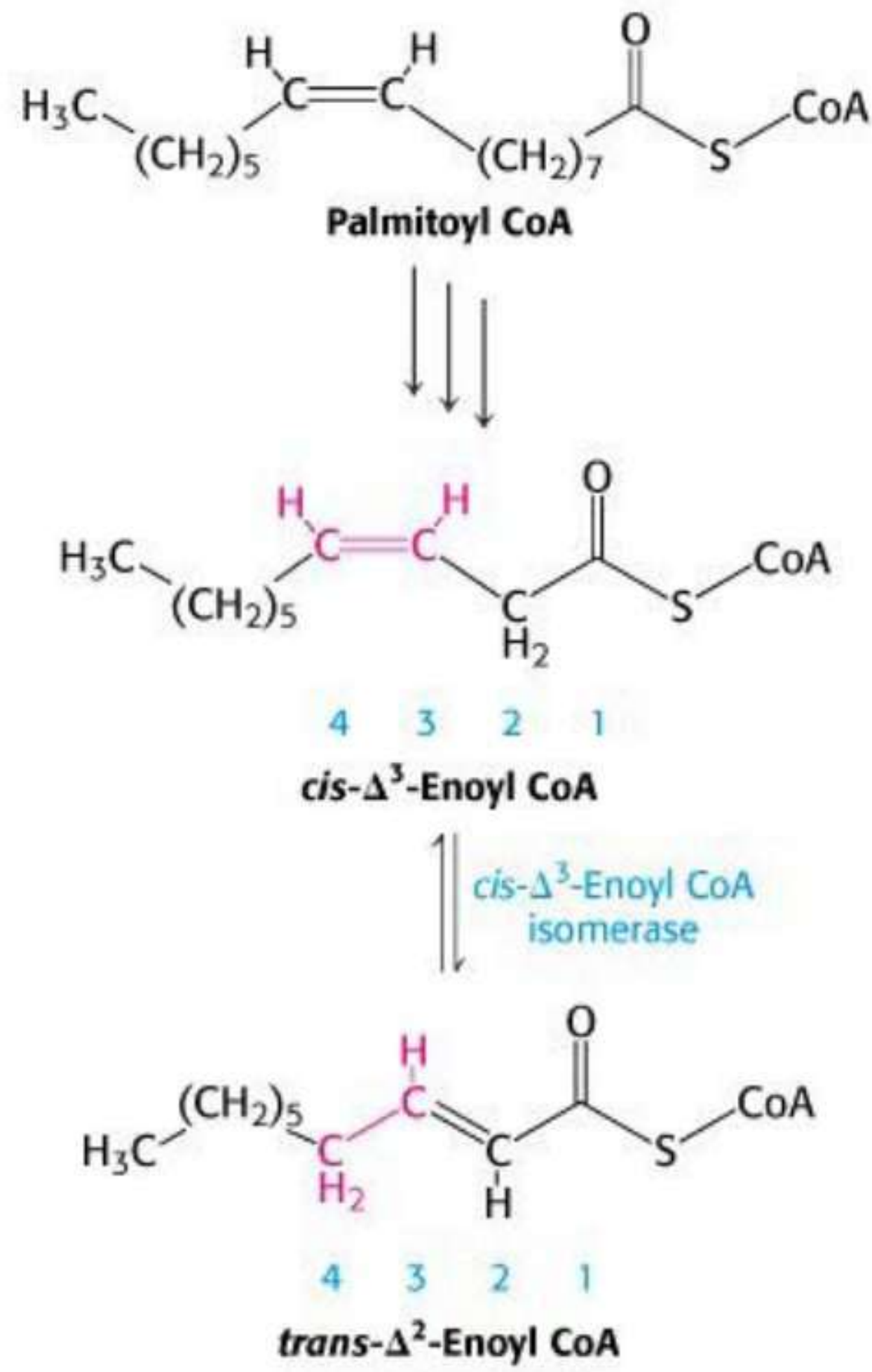
**TABLE 22.1** Principal reactions in fatty acid oxidation

Step	Reaction	Enzyme
1	Fatty acid + CoA + ATP $\rightleftharpoons$ acyl CoA + AMP + PP <sub>i</sub>	Acyl CoA synthetase [also called fatty acid thiokinase and fatty acid:CoA ligase (AMP)]
2	Carnitine + acyl CoA $\rightleftharpoons$ acyl carnitine + CoA	Carnitine acyltransferase (also called carnitine palmitoyl transferase)
3	Acyl CoA + E-FAD $\rightarrow$ <i>trans</i> - $\Delta^2$ -enoyl CoA + E-FADH <sub>2</sub>	Acyl CoA dehydrogenases (several isozymes having different chain-length specificity)
4	<i>trans</i> - $\Delta^2$ -Enoyl CoA + H <sub>2</sub> O $\rightleftharpoons$ L-3-hydroxyacyl CoA	Enoyl CoA hydratase (also called crotonase or 3-hydroxyacyl CoA hydrolyase)
5	L-3-Hydroxyacyl CoA + NAD <sup>+</sup> $\rightleftharpoons$ 3-ketoacyl CoA + NADH + H <sup>+</sup>	L-3-Hydroxyacyl CoA dehydrogenase
6	3-Ketoacyl CoA + CoA $\rightleftharpoons$ acetyl CoA + acyl CoA (shortened by C <sub>2</sub> )	$\beta$ -Ketothiolase (also called thiolase)

## Energy production

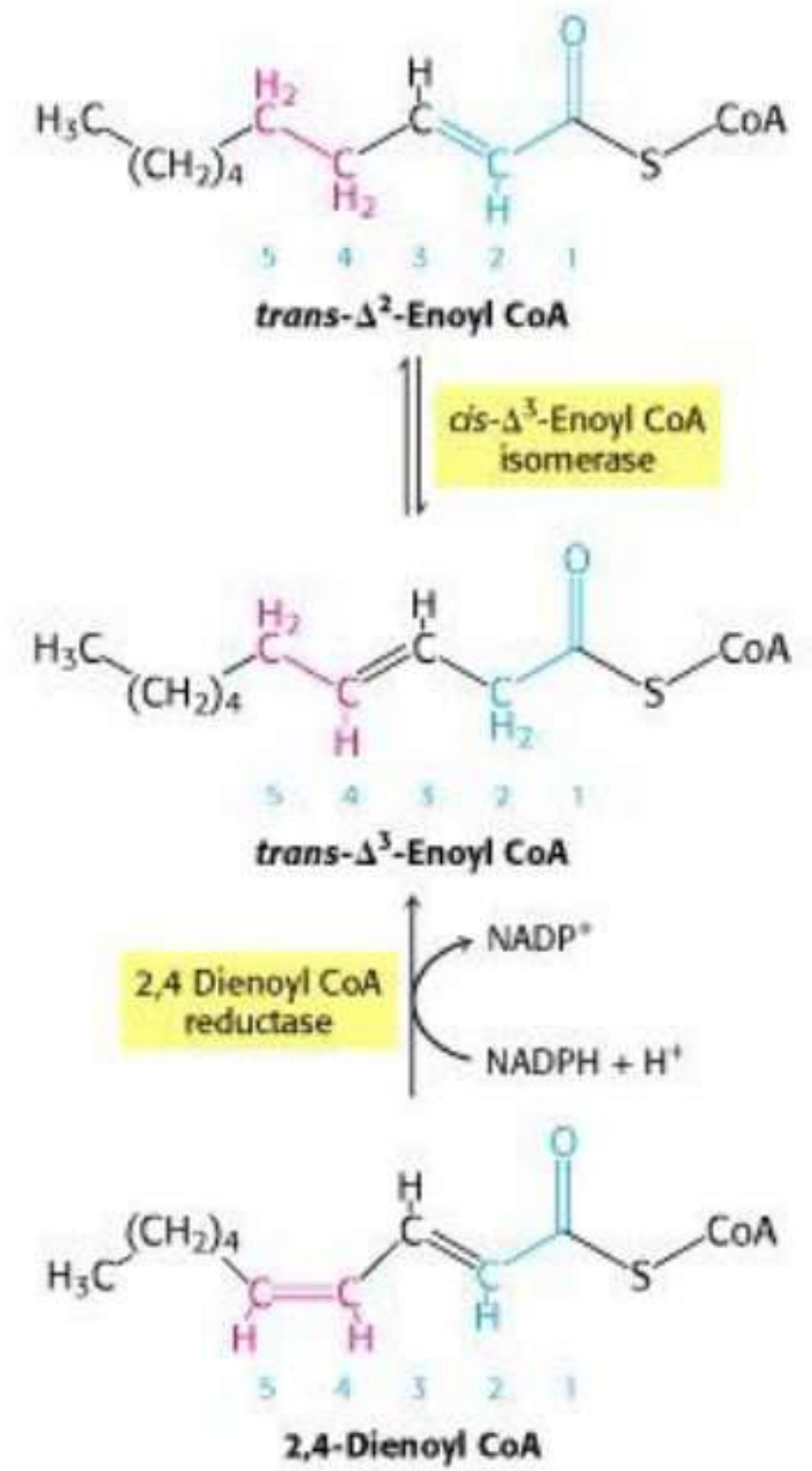
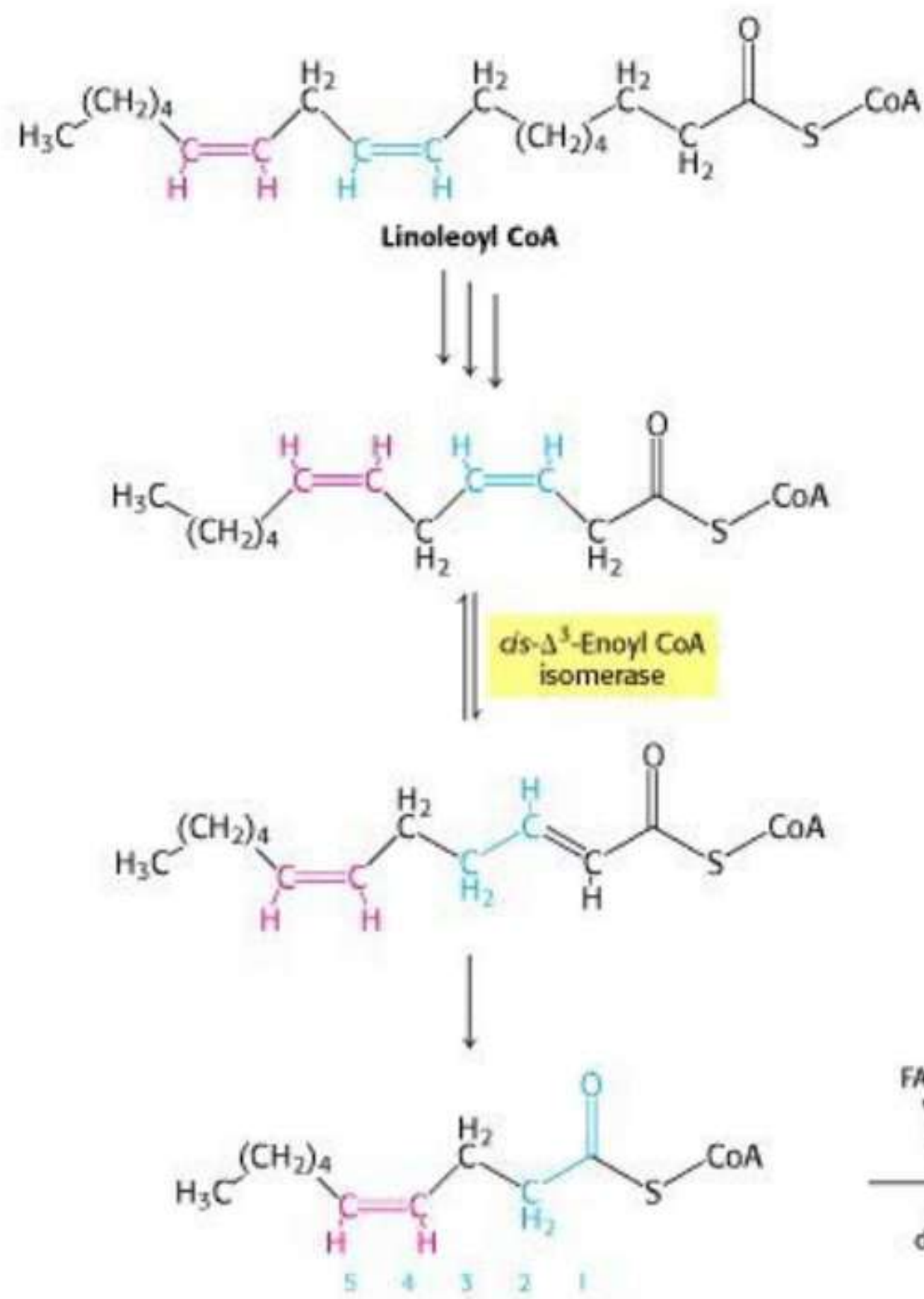
- NADH and FADH from B-oxidation
- TCA cycle from acetyl CoA
- Total net yield is minus 2 ATP from activation

## Oxidation of Unsaturated Fatty acids

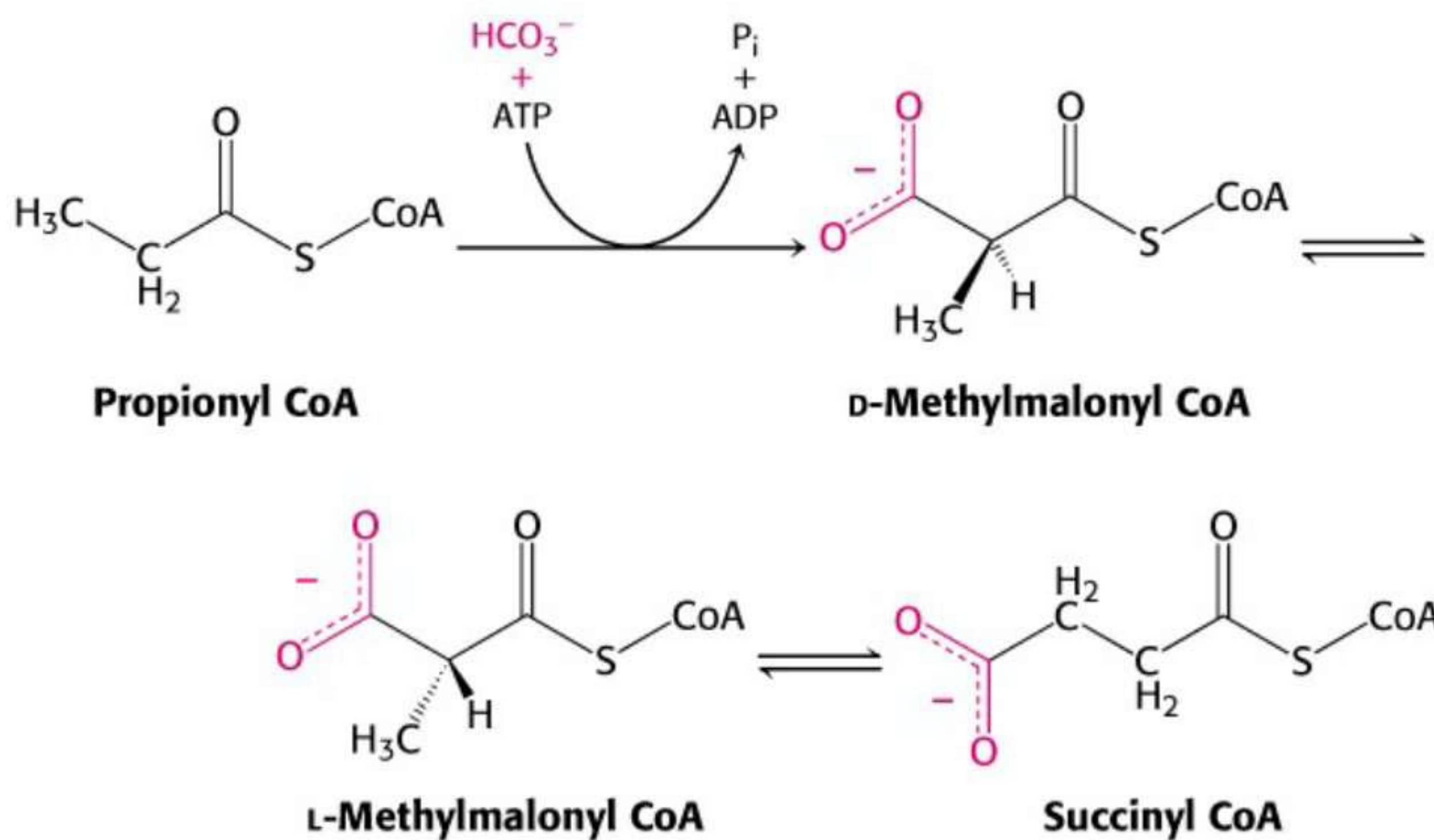


## Unsaturated Fatty acids



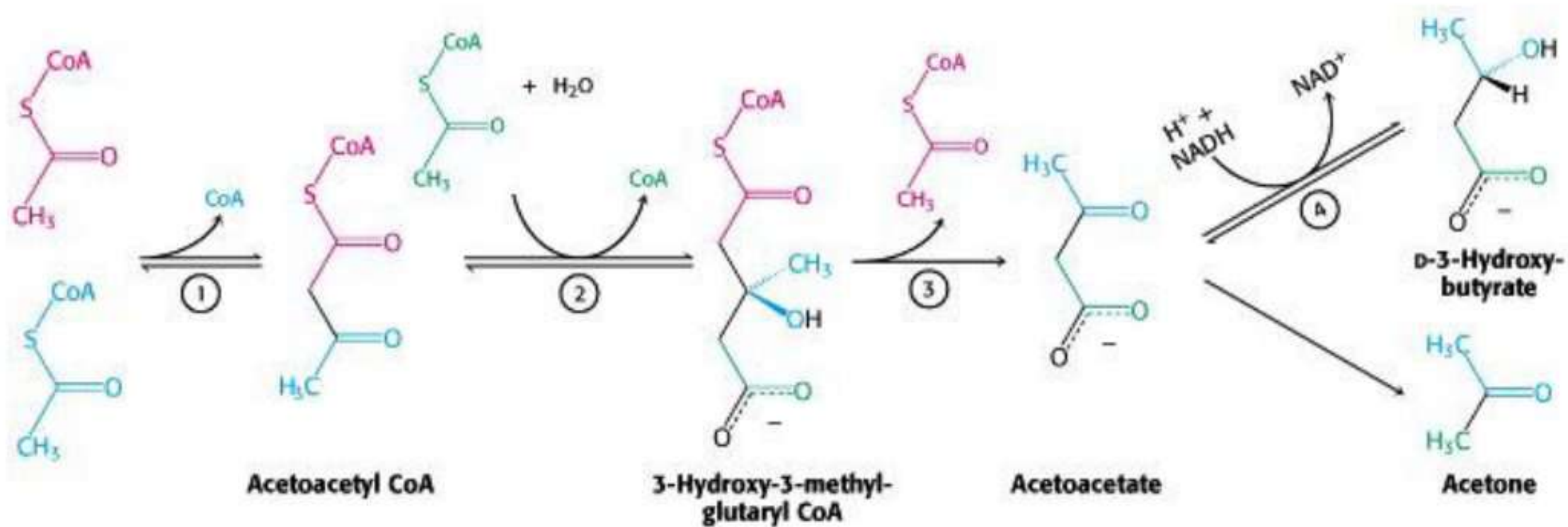


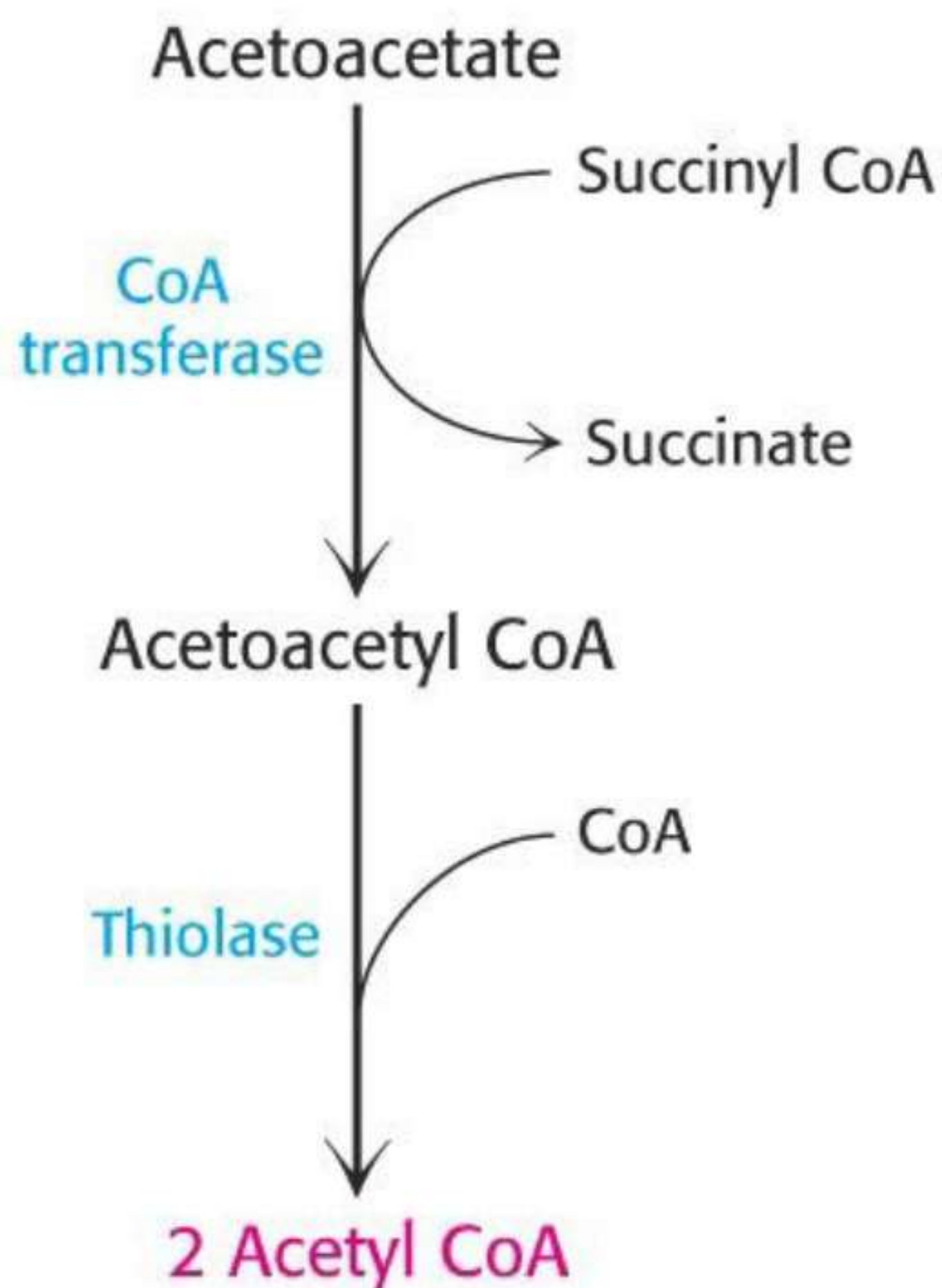
Oxidation of odd chain fatty acids



- form propionylCoA
- produce succinylCoA

## Ketone Bodies





- Acetoacetate
- Acetone
- B-hydroxybutyrate
- HMG CoA synthase

### Referances

Available online

**1-BIOCHEMISTRY IN PERSPECTIVE**

2-METABOLISM OF CARBOHYDRATES, LIPIDS, PROTEINS AND NUCLEIC ACIDS, Course Team

Prof. Anthony, I. O. Ologhobo(Course Writer)-UI

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