

Lehninger

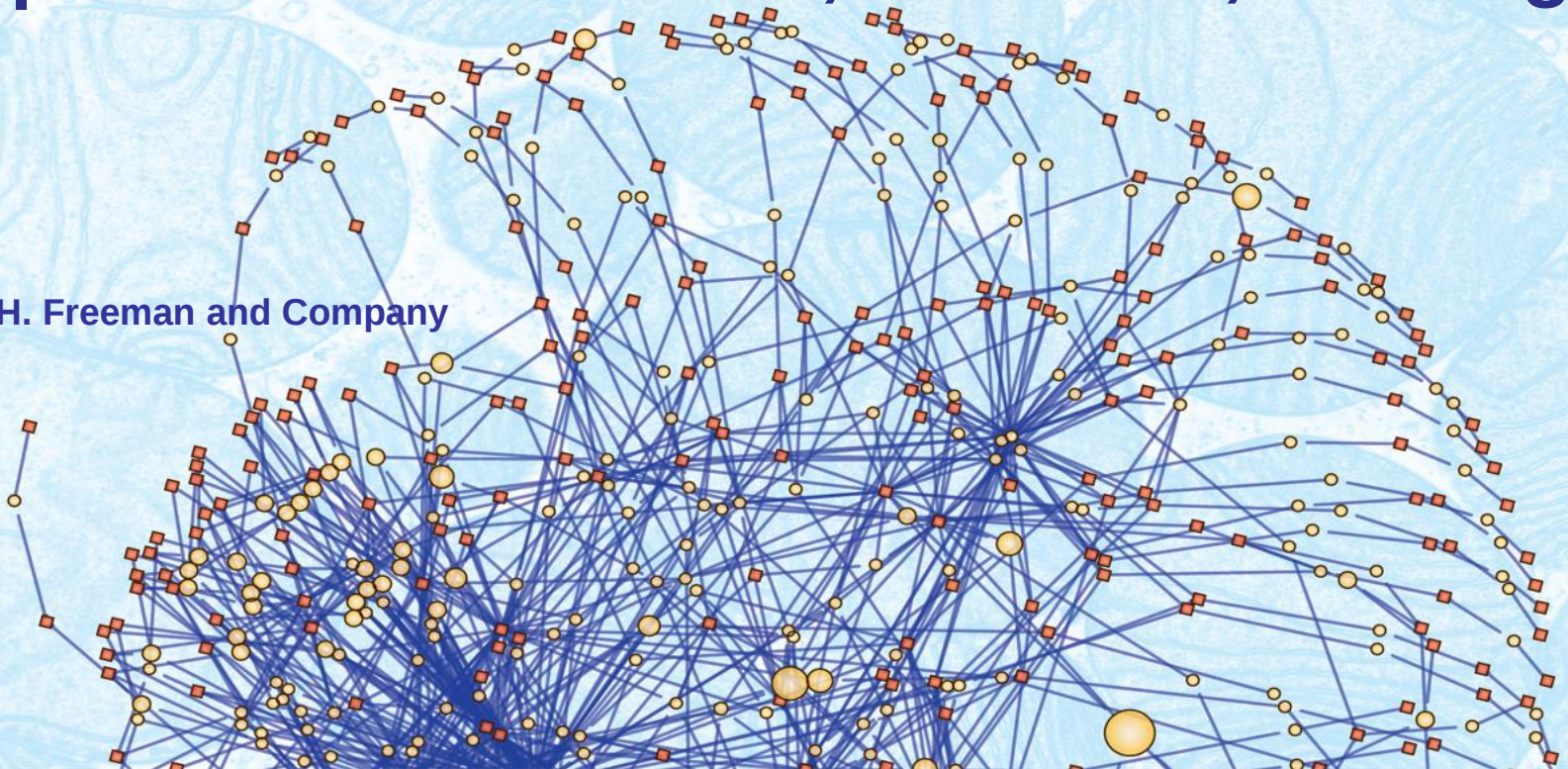
SIXTH EDITION

Principles of Biochemistry

David L. Nelson | Michael M. Cox

4 | Proteins: Structure, Function, Folding

© 2013 W. H. Freeman and Company



CHAPTER 4

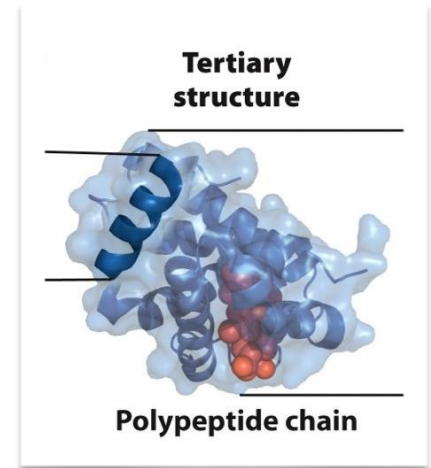
Proteins: Structure, Function, Folding

Learning goals:

- Structure and properties of the **peptide bond**
- Structural **hierarchy** in proteins
- Structure and function of **fibrous** proteins
- Structure analysis of **globular** proteins
- Protein **folding** and **denaturation**

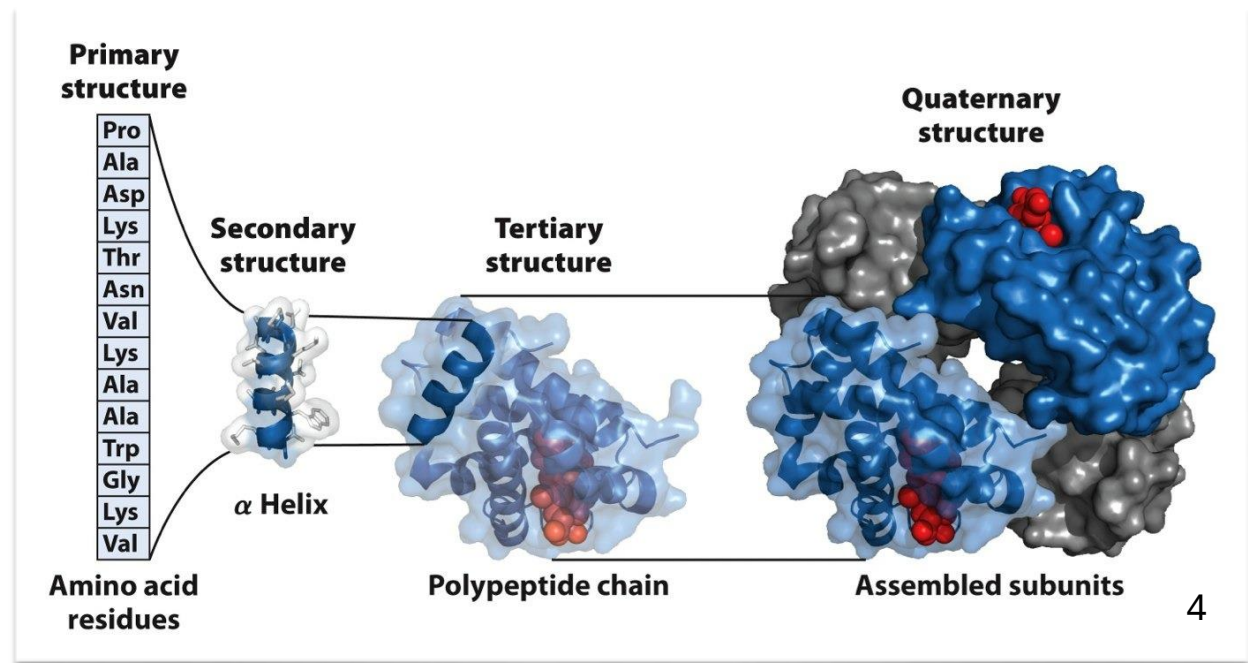
Structure of Proteins

- Protein adopts a specific **3D conformation**.
 - In principle, an uncountable number of conformations.
 - In reality, one or a few unique structures.
 - Structure fulfills a specific biological function.
- This structure is called the **native fold**.
 - A large number of **favorable interactions** within the protein.
 - A **cost** in conformational **entropy** of folding into native fold.



Six Themes About Protein Structure

1. 3D structure determined by amino acid sequence.
2. Function depends on structure.
3. One or a few **stable** structures.
4. Most important force is **noncovalent** bond.
5. Structural patterns can be organized.
6. Not static.



Three-Dimensional Structure of Proteins

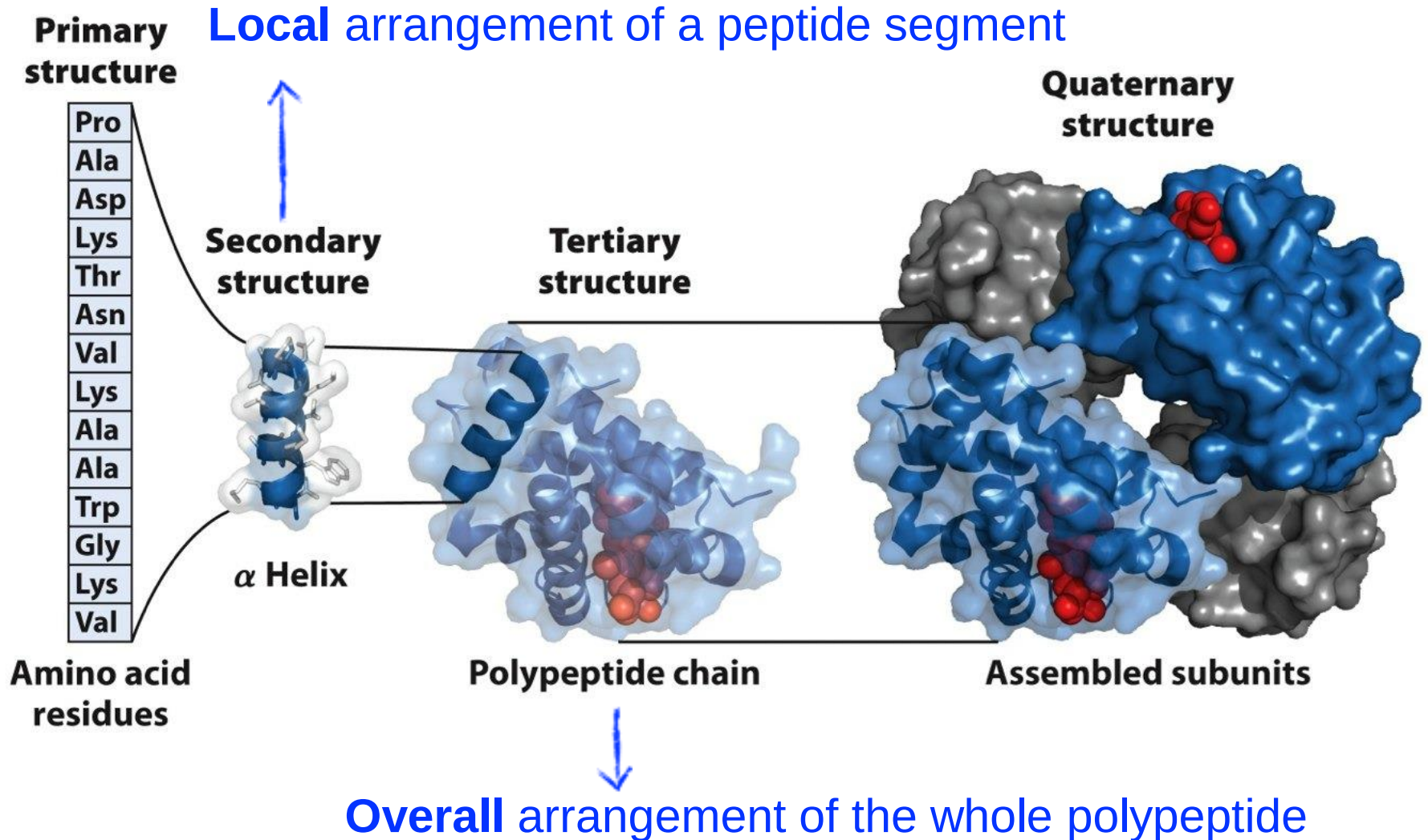
4.1 Overview of Protein Structure

4.2 Protein Secondary Structure

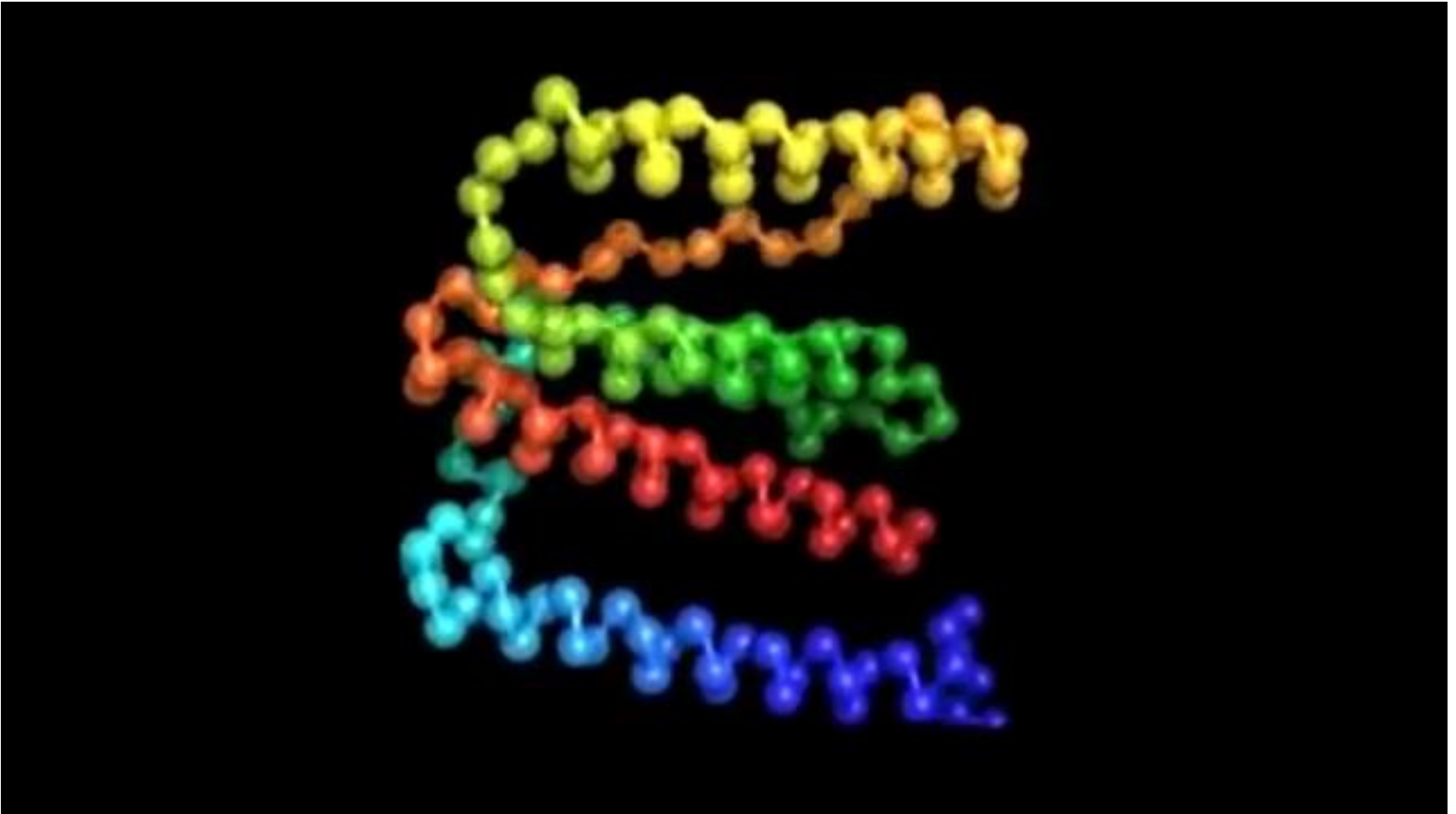
4.3 Protein Tertiary and Quaternary Structure

4.4 Protein Denaturation and Folding

4 Levels of Protein Structure

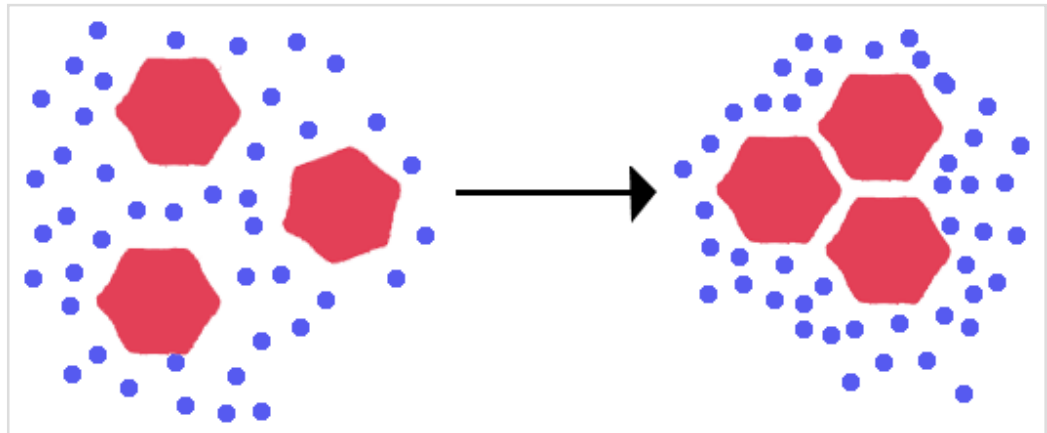


From Primary to Tertiary Structure



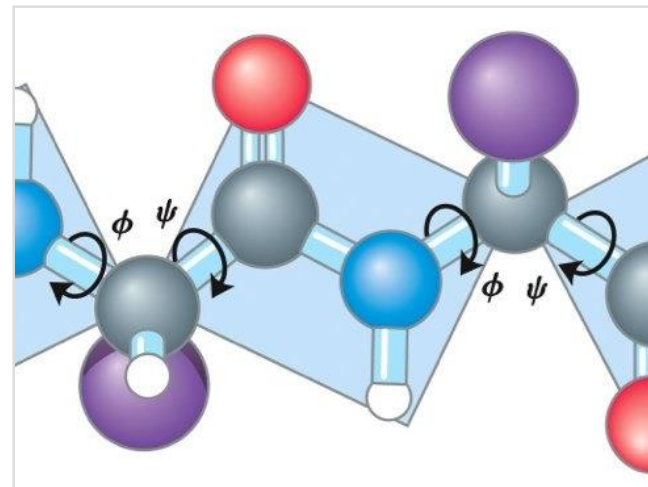
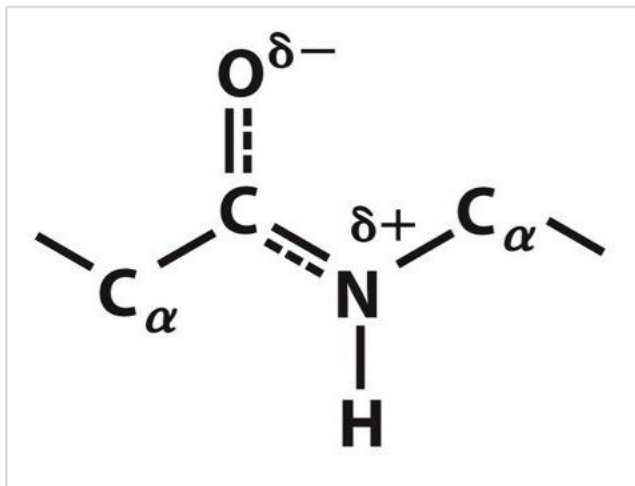
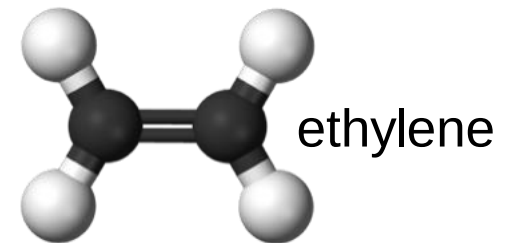
Structure Stabilized by **Weak Interactions**

- Covalent bond (disulfide)
- Noncovalent bond
 - **hydrophobic interaction**
 - hydrogen bond
 - ionic interaction

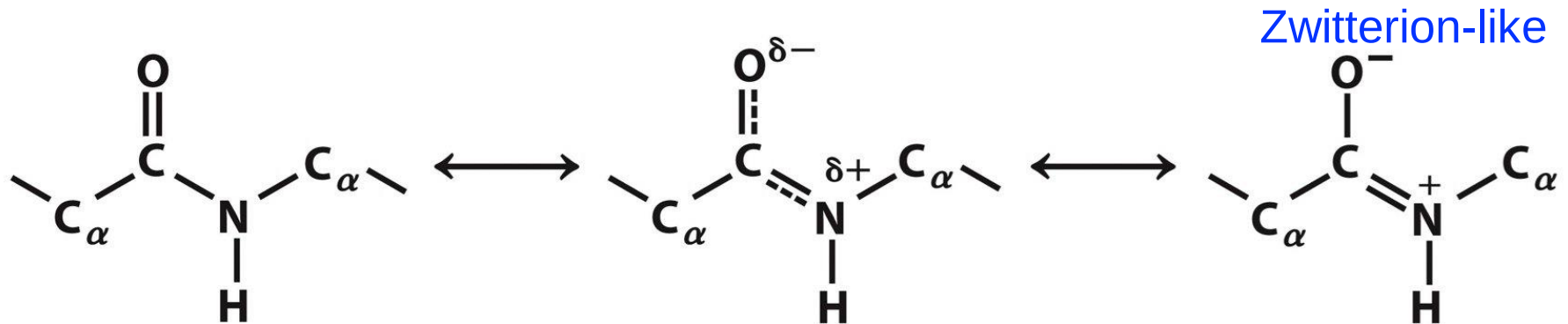


Structure of Peptide Bond

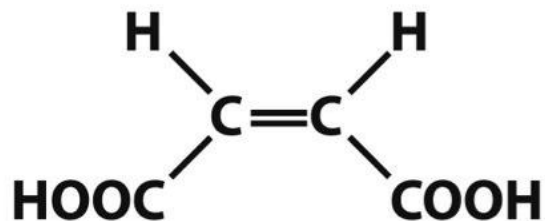
- Structure partially dictated by peptide bond.
- A resonance hybrid of two canonical structures.
- The resonance causes the peptide bonds:
 - to be less reactive.
 - to be quite **rigid** and nearly **planar**.



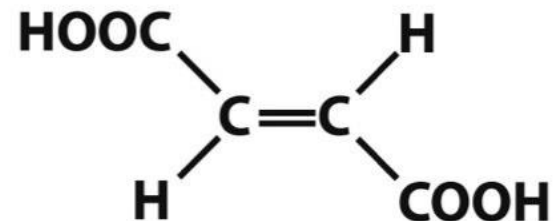
Resonance in Peptide Bond



The carbonyl oxygen has a partial negative charge and the amide nitrogen a partial positive charge, setting up a small electric dipole. Virtually all peptide bonds in proteins occur in this trans configuration; an exception is noted in



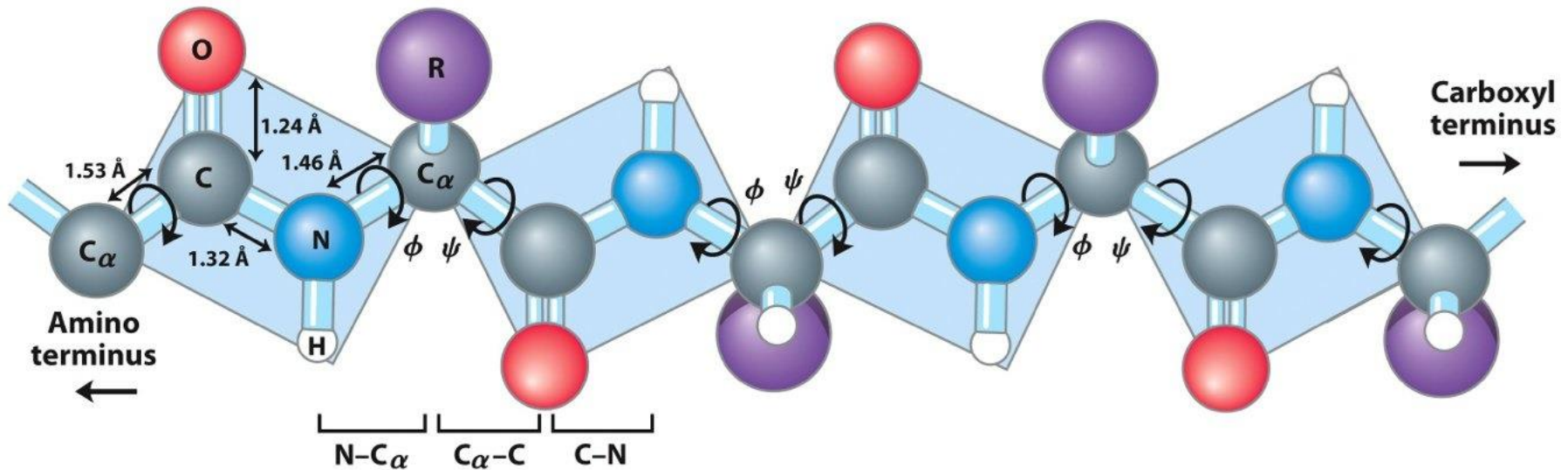
Maleic acid (cis)



Fumaric acid (trans)

Polypeptide Made up of A Series of Planes Linked at α Carbons

Two C-N bond lengths are different?



on the
SAME
plane

α -carbon

carbonyl group (C and O)

amino group (N and H)

α -carbon

Summary 4.1 Overview of Protein Structure

- A typical protein has **one or more** stable three-dimensional structures or conformations.
- Protein structure is stabilized largely by multiple **weak** interactions, with **hydrophobic** interactions being the major contributors.
- The peptide bond has a partial double-bond character that keeps it in a rigid **planar** configuration.

Example Question

Any given protein is characterized by a unique amino acid sequence (primary structure) and three-dimensional (tertiary) structure. How are these related?

Example Question

All of the following are considered “weak” interactions in proteins *except*:

- A) hydrogen bonds.
- B) hydrophobic interactions.
- C) ionic bonds.
- D) peptide bonds.
- E) all above interactions are weak interactions.

Example Question

Which statement about the peptide bond is correct?

- A) peptide bonds have many different conformations that are equally probable.
- B) peptide bonds are essentially planar, with no rotation about the C-N axis.
- C) peptide bonds in proteins are uncommon.
- D) peptide bond structure is extraordinarily complex and poorly understood.
- E) primary structure of all proteins is similar, although the secondary and tertiary structure may differ greatly.

Three-Dimensional Structure of Proteins

4.1 Overview of Protein Structure

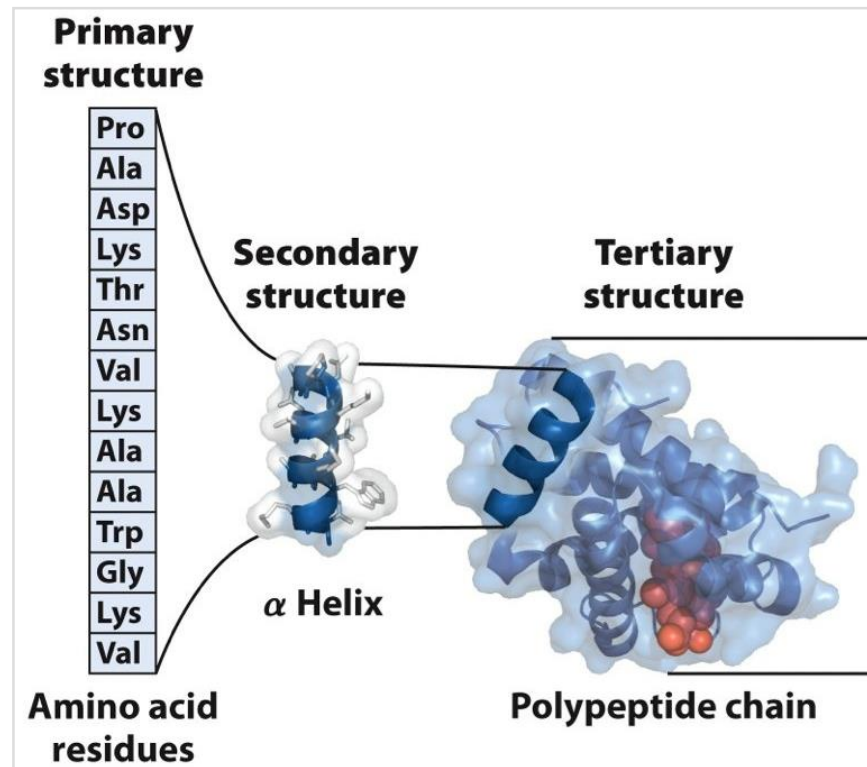
4.2 Protein Secondary Structure

4.3 Protein Tertiary and Quaternary Structure

4.4 Protein Denaturation and Folding

Secondary Structures

- Any chosen segment of a polypeptide chain.
- Local spatial arrangement of **main-chain atoms**.
- NOT positioning of **side chains**.
- NOT **relationship to other** segments.



Secondary Structures

Three common secondary structures:

➤ α helix

- stabilized by hydrogen bonds between **nearby** residues.

➤ β sheet

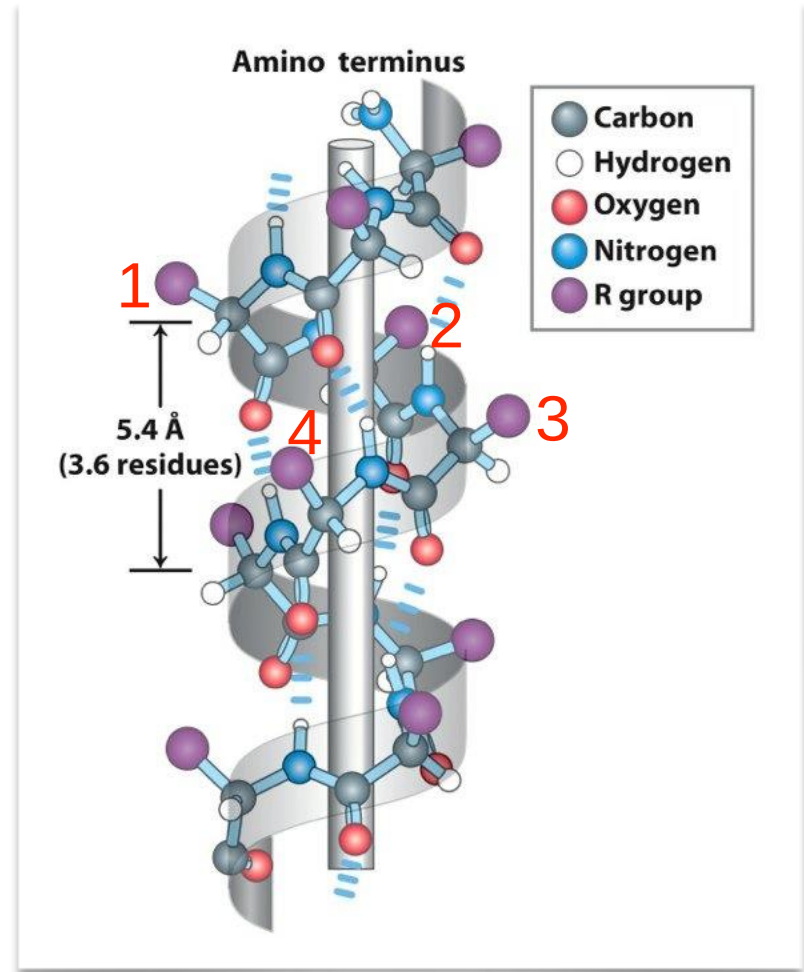
- stabilized by hydrogen bonds between adjacent segments that **may not be nearby** in primary sequence.

➤ β turn

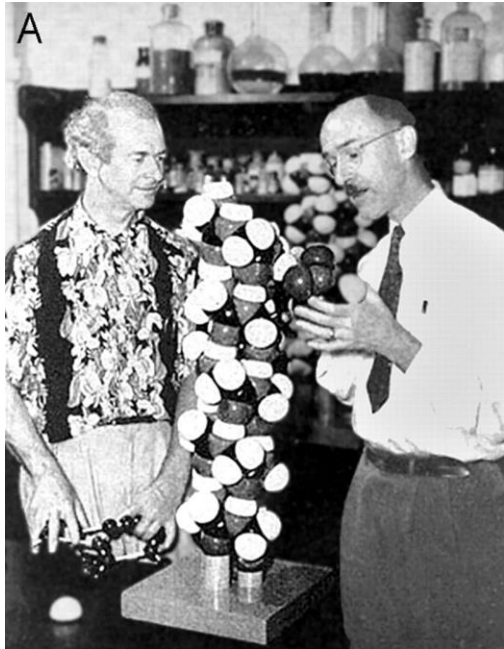
- used to connect two adjacent β sheets.

α Helix

- Side chains point outward and are roughly perpendicular with the axis.
- 3.6 residues per turn.
 - 5.4 Å along the axis
 - 1 Å (angstrom) = 10^{-10} m = 0.1 nm.
- Hydrogen bonds between the backbone amides of 1st and 4th peptide bonds.



Discovery of α Helix



- Paper discovery in 1948
 - Pauling bedridden with a cold
 - Paper drawing and folding
- Experimental proof in 1951
 - Theory before observation
- DNA double helix in 1953
 - Pauling-Corey model-building

A

Linus Pauling (left), deep understanding of chemical structure and bonding
Robert Corey (right), x-ray crystallographer with know-how and patience
Wooden helix, scale of 1 inch per Å, an enlargement of _____ times

B

Herman Branson, physicist capable of calculating helical structures

What is a Right-handed Helix?

**Left-handed
helix**

**Not
Observed in
Proteins**

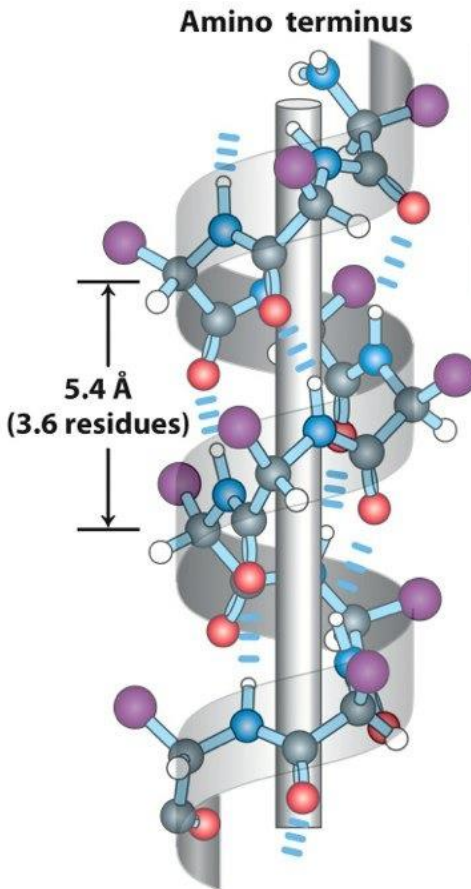


**Right-handed
helix**

Common Form



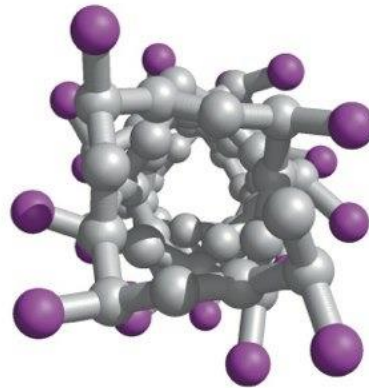
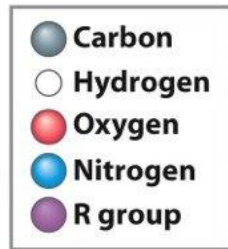
α Helix



(a)

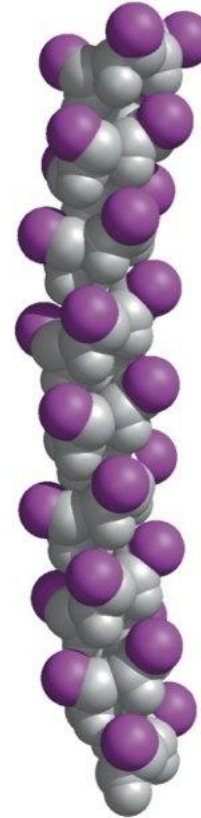
Carboxyl terminus

Side view



(b)

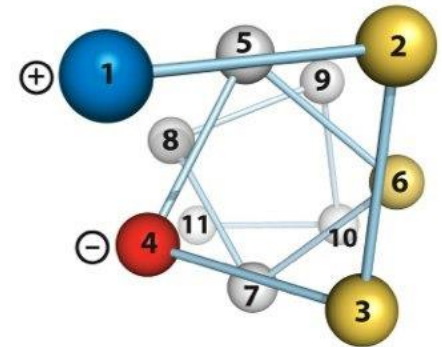
Top view



(c)

Space-filling

- Hydrophobic interface
- Ionic interaction



(d)

Interactions

(a and b) ball-and-stick model
gives a false impression that helix interior is hollow

Sequence Affects Helix Stability

- Not all polypeptide sequences adopt α -helical structures.
- **Ala** is the strongest helix former.
 - Small hydrophobic residues.
- **Pro** acts as a helix breaker.
 - Rotation around the N-C $_{\alpha}$ bond is impossible.
- **Gly** acts as a helix breaker.
 - the tiny R-group supports other conformations.
- Attractive or repulsive interactions between side chains 3-4 amino acids apart will affect formation.

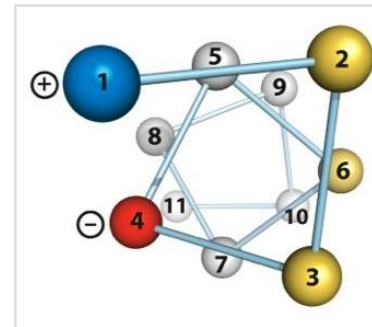
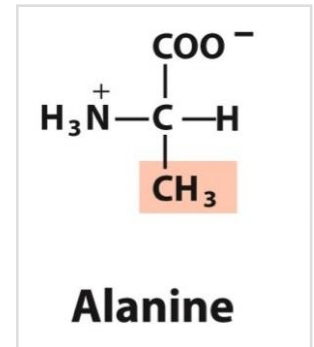
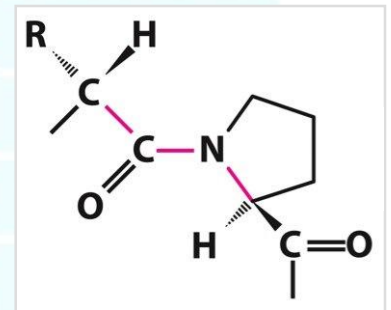


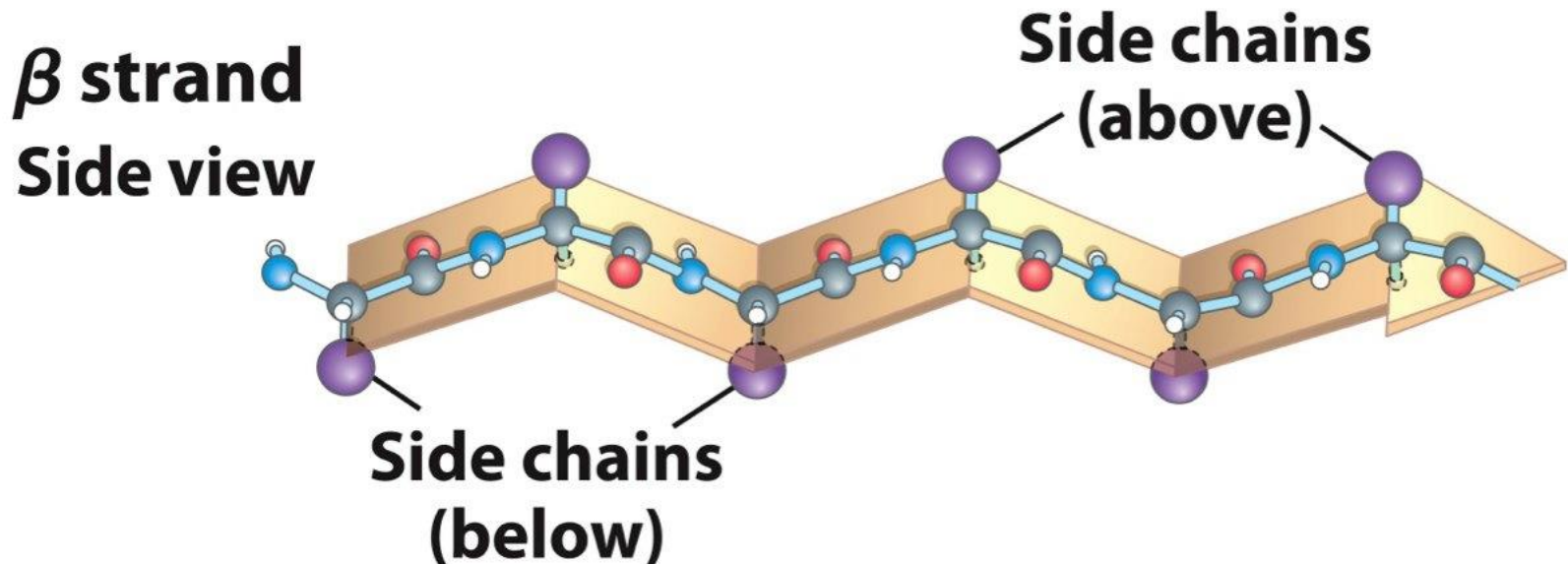
TABLE 4-2**Propensity of Amino Acid Residues to Take Up an α -Helical Conformation**

Amino acid	$\Delta\Delta G^\circ$ (kJ/mol)*	Amino acid	$\Delta\Delta G^\circ$ (kJ/mol)*
Ala	0	Leu	0.79
Arg	0.3	Lys	0.63
Asn	3	Met	0.88
Asp	2.5	Phe	2.0
Cys	3	Pro	>4
Gln	1.3	Ser	2.2
Glu	1.4	Thr	2.4
Gly	4.6	Tyr	2.0
His	2.6	Trp	2.0
Ile	1.4	Val	2.1



β Sheet

- A **pleated sheet-like** structure.
- Held together by **hydrogen bonds** between the backbone amides in different segments.
- Side chains protrude from the sheet **alternating** in up and down direction.

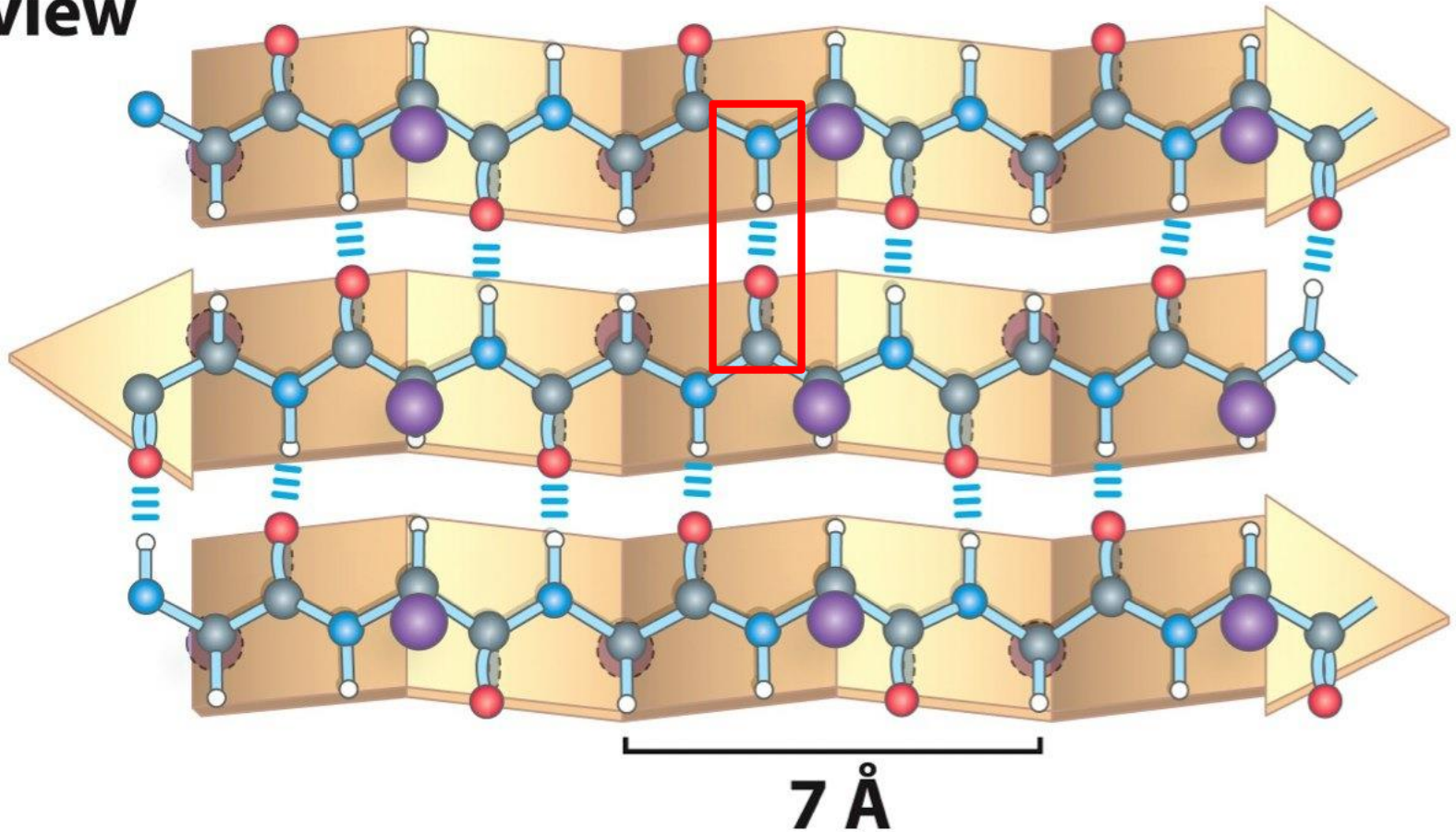


Parallel and Antiparallel β Sheets

- **Parallel or antiparallel** orientation of two chains within a sheet are possible
- In parallel β sheets the H-bonded strands run in the **same direction**
 - Resulting in bent H-bonds (weaker)
- In antiparallel β sheets the H-bonded strands run in **opposite directions**
 - Resulting in linear H-bonds (stronger)

Antiparallel β sheet

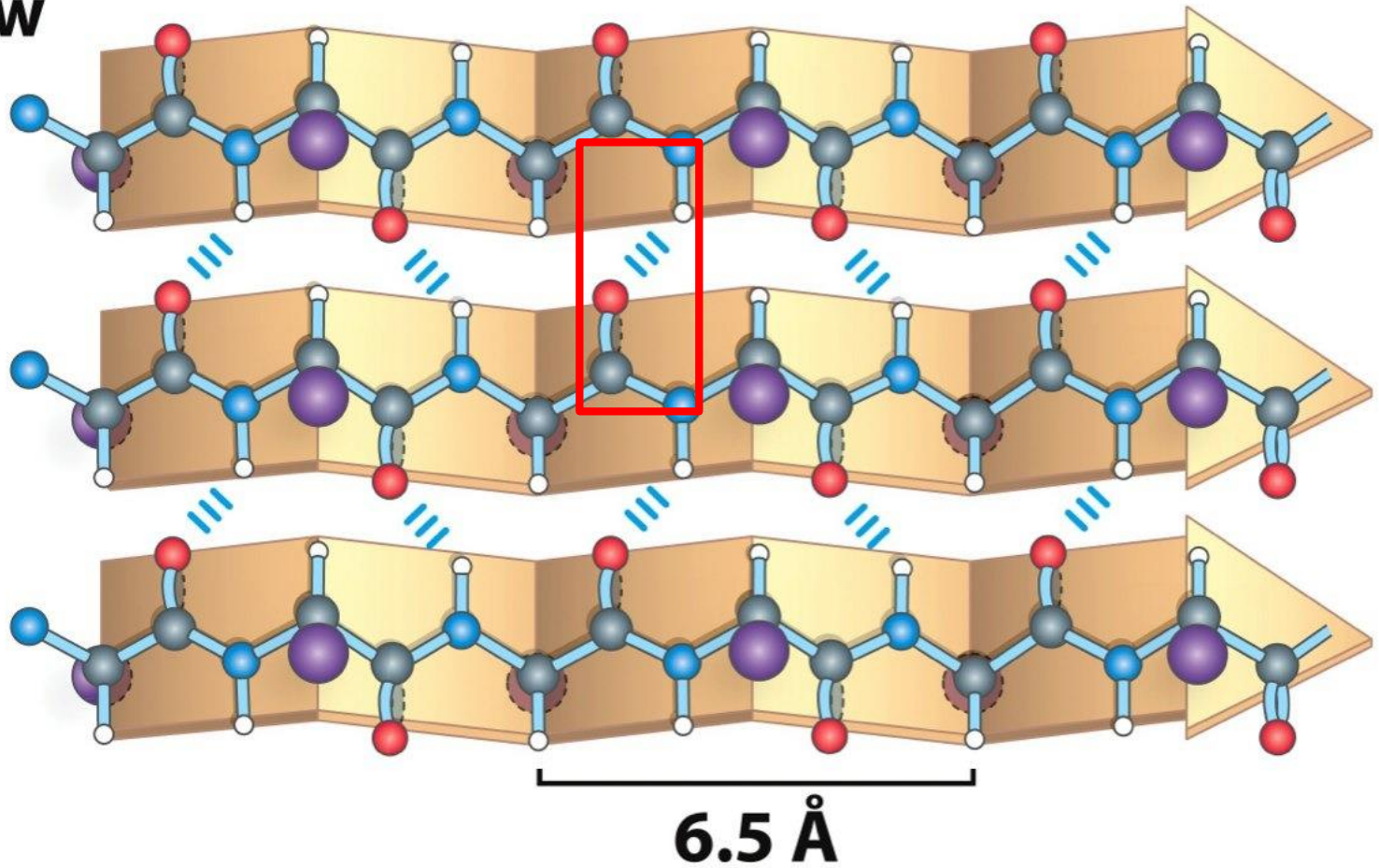
Top view



hydrogen bonds: **In-line.**

Parallel β sheet

Top view

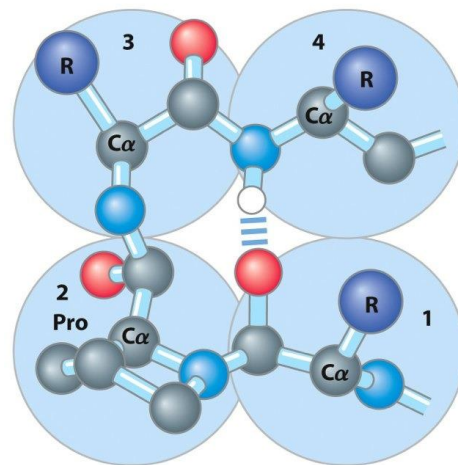


hydrogen bonds: **Not in-line, or distorted.**

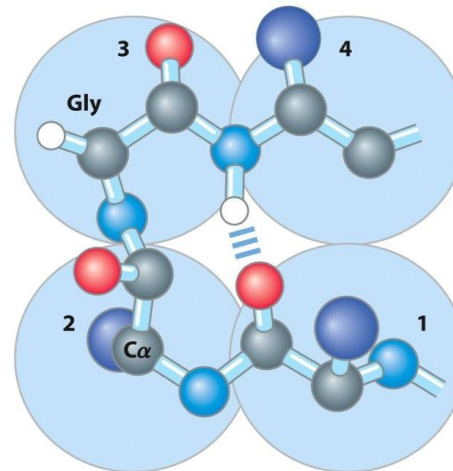
β Turn

β turns occur frequently whenever strands in β sheets change the direction

- The 180° turn is accomplished over four amino acids
- The turn is stabilized by a hydrogen bond.
- **Proline** in position 2 or **glycine** in position 3 are common in β turns

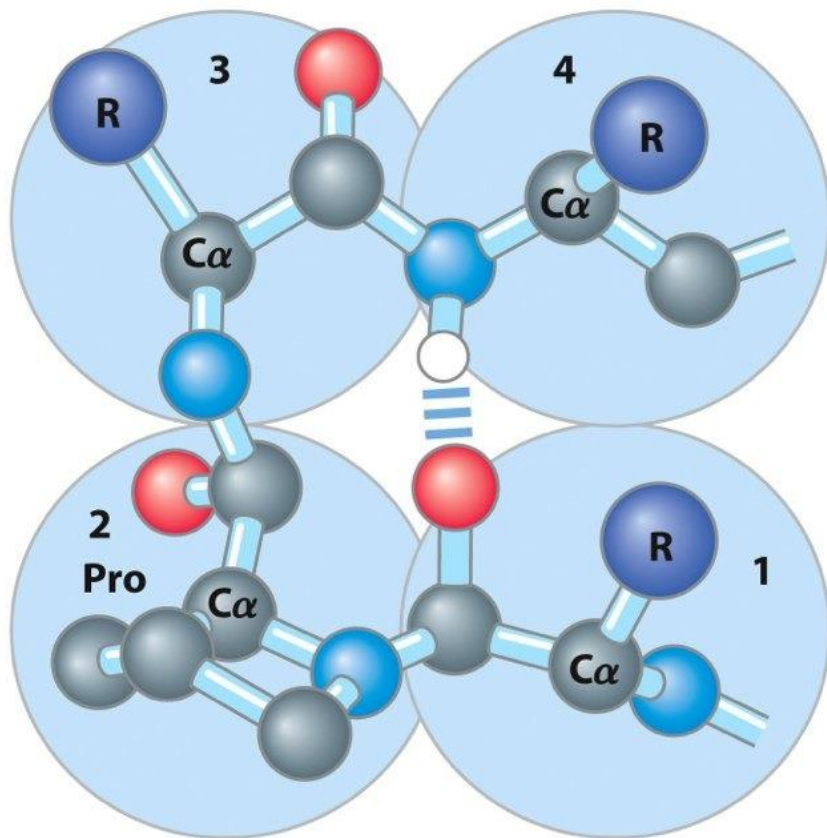


Type I β turn

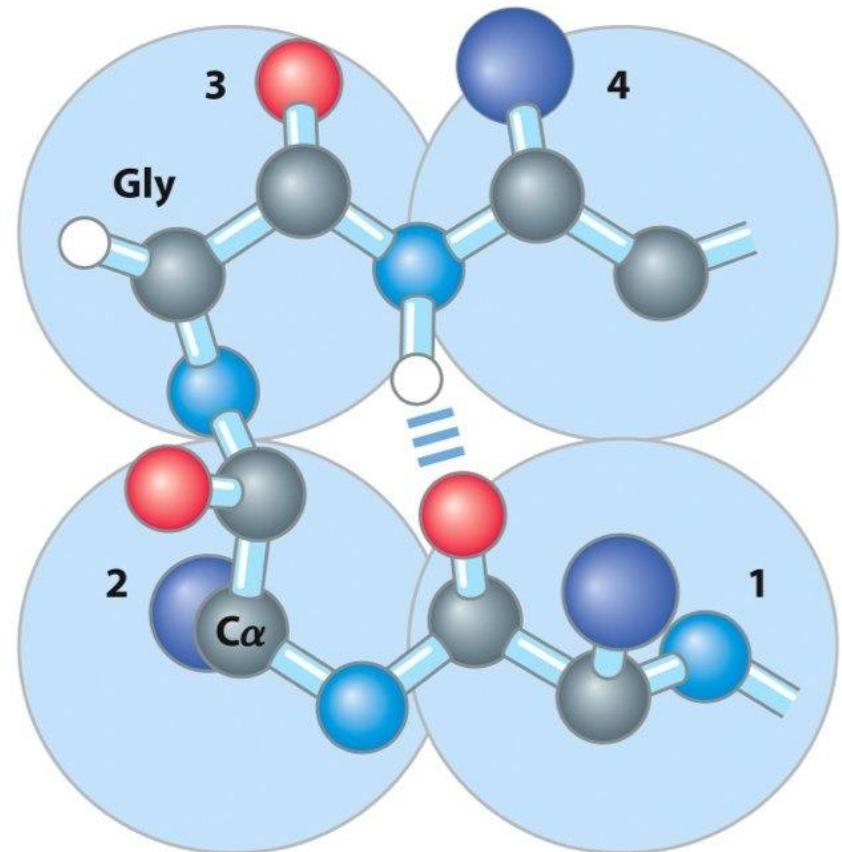


Type II β turn

β Turn



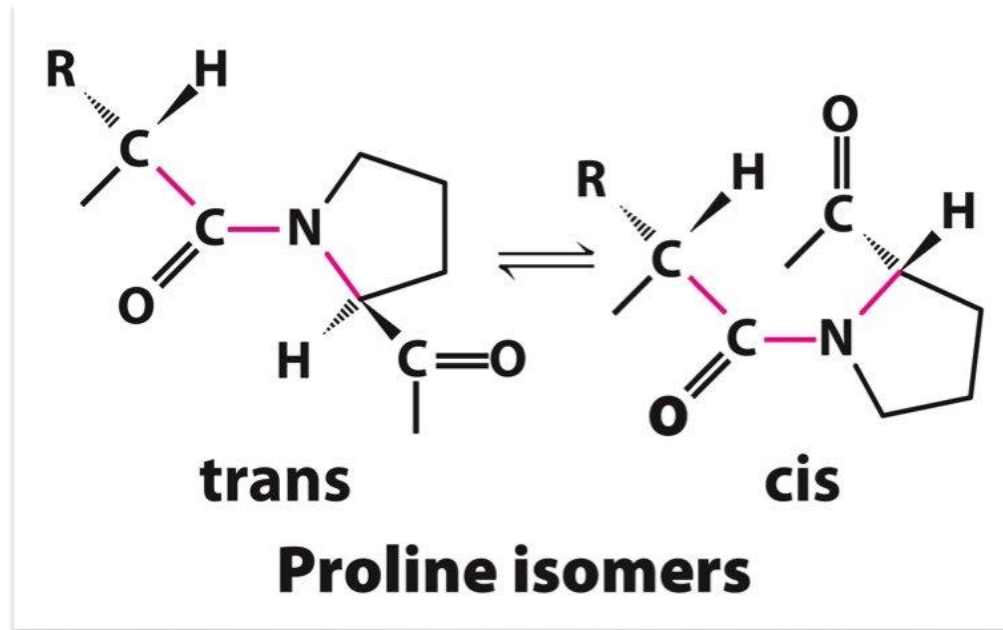
Type I β turn



Type II β turn

Proline Isomers

- Most peptide bonds **not involving proline** are in the *trans* configuration (>99.95%).
- For peptide bonds **involving proline**, about 6% are in the *cis* configuration. Most of this 6% involve β -turns.
- Proline isomerization is catalyzed by proline isomerases.



Summary 4.2 Secondary Structure

- Secondary structure is the **local** spatial arrangement of main-chain atoms in a selected segment of polypeptide chain.
- The α helix (3.6 residues, 5.4 Å, right-handed, Ala).
- The β sheet (parallel vs antiparallel).
- The β turn (180° , four residues, Pro).

Example Question

Secondary structure describes the relationship and interaction of amino acid residues that are:

- A) always side by side.
- B) generally near each other in sequence.
- C) restricted to about 7 of the 20 standard amino acids.
- D) often on different polypeptide strands.
- E) usually near the polypeptide chain's amino terminus or carboxyl terminus.

Example Question

Roughly how many amino acids are there in one turn of an α -helix?

A) 1

B) 2.8

C) 3.6

D) 4.2

E) 10

Example Question

What is the length of a polypeptide with 80 amino acid residues in a single contiguous α helix?

Example Question

A D-amino acid would interrupt an α -helix made of L-amino acids. Another naturally occurring hindrance to the formation of an α -helix is the presence of:

- A) a negatively charged Arg residue.
- B) a nonpolar residue near the carboxyl terminus.
- C) a positively charged Lys residue.
- D) a Pro residue.**
- E) two Ala residues side by side.

Example Question

A sequence of amino acids in a certain protein is found to be -Ser-Gly-Pro-Gly-. The sequence is most probably part of a(n):

- A) antiparallel β sheet.
- B) parallel β sheet.
- C) alpha helix.
- D) beta sheet.
- E) β turn.

Three-Dimensional Structure of Proteins

4.1 Overview of Protein Structure

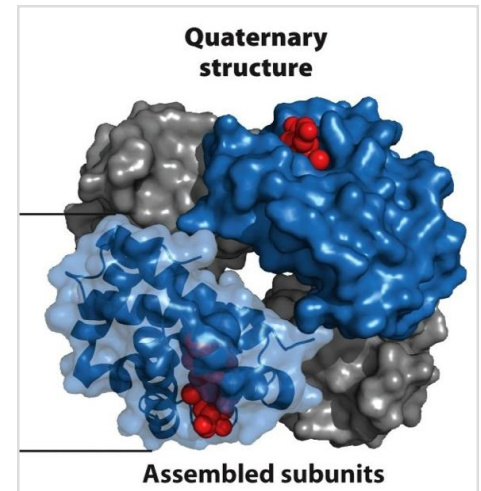
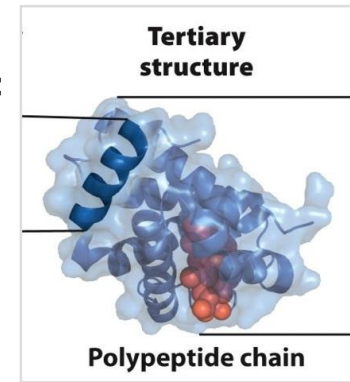
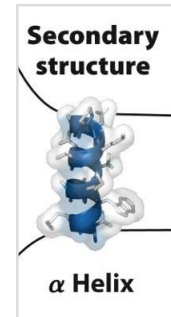
4.2 Protein Secondary Structure

4.3 Protein Tertiary and Quaternary Structure

4.4 Protein Denaturation and Folding

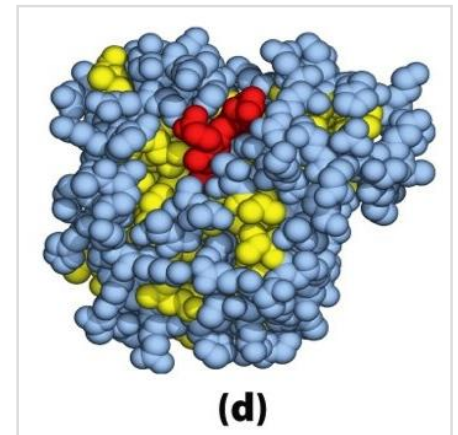
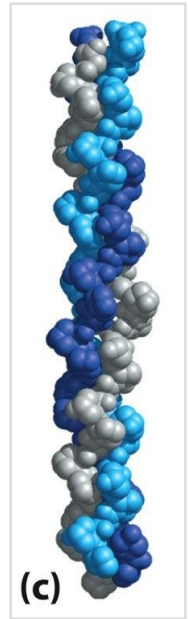
Protein Tertiary and Quaternary Structure

- 3° structure refers to the overall spatial arrangement of atoms in a protein
 - Secondary structure: adjacent residues
 - Tertiary structure: **longer-range** interactions
- 4° structure refers to the arrangement of subunits in a **multisubunit** protein in three-dimensional complex.
- Stabilized by numerous **weak** interactions between amino acid side chains.
 - Largely **hydrophobic** and polar interactions
 - Can be stabilized by disulfide bonds



Fibrous Proteins and Globular Proteins

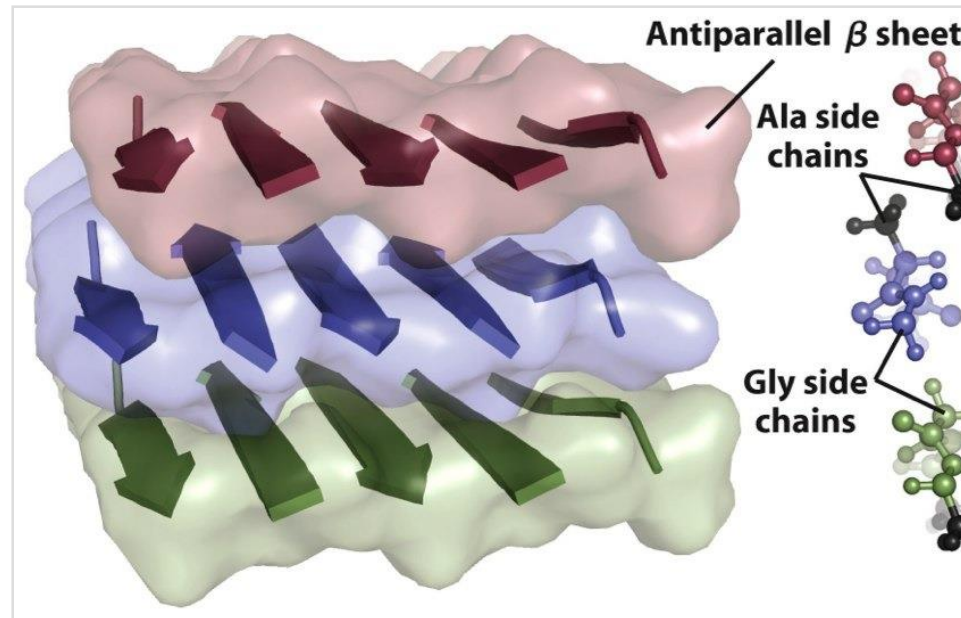
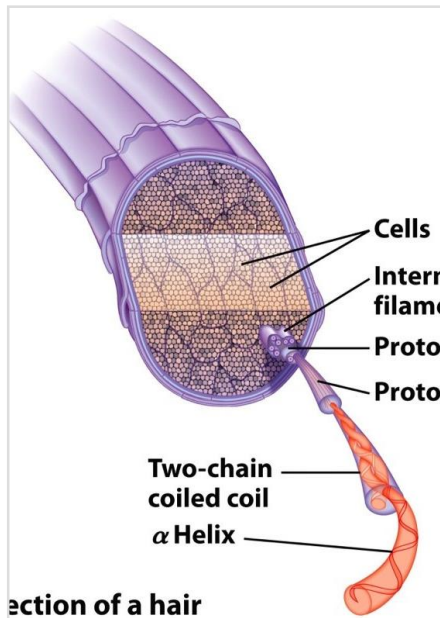
- Fibrous Proteins
 - Long strands or sheets
 - A **single** type of secondary structure
 - Provide support, shape, and external protection
- Globular Proteins
 - Spherical or globular shape
 - **Several** types of secondary structure
 - Enzymes and regulatory protein



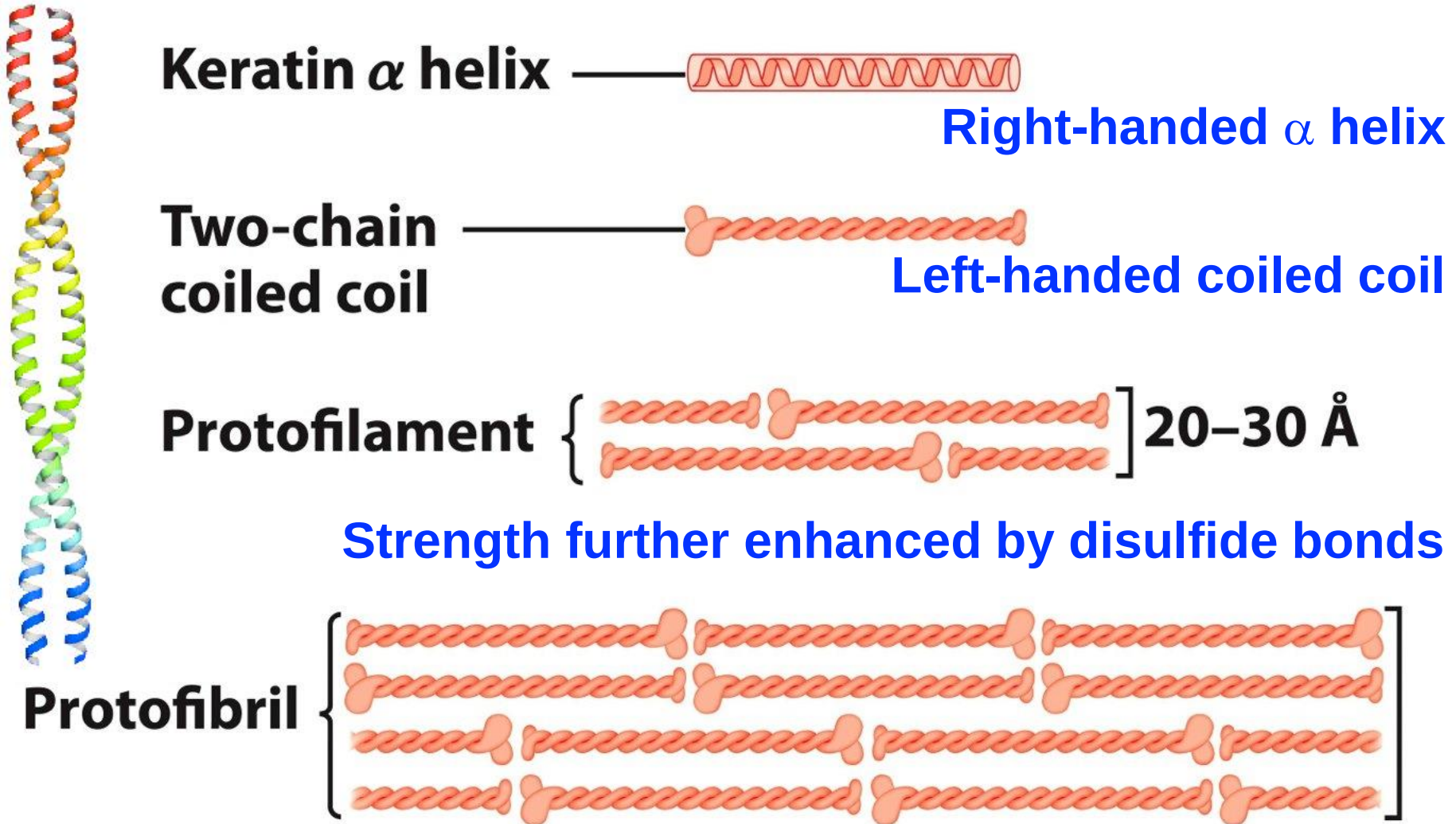
Fibrous Proteins: From Structure to Function

TABLE 4-3 Secondary Structures and Properties of Some Fibrous Proteins

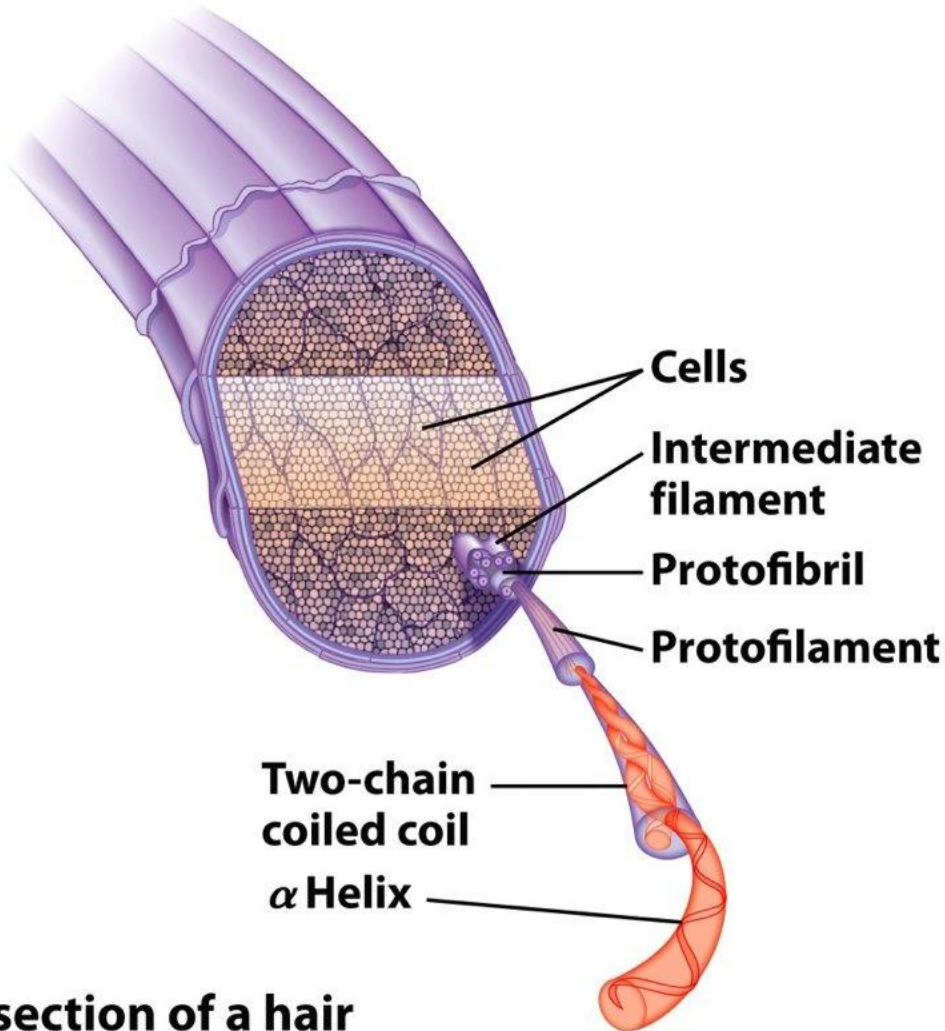
Structure	Characteristics	Examples of occurrence
α Helix, cross-linked by disulfide bonds	Tough, insoluble protective structures of varying hardness and flexibility	α -Keratin of hair, feathers, nails
β Conformation	Soft, flexible filaments	Silk fibroin
Collagen triple helix	High tensile strength, without stretch	Collagen of tendons, bone matrix



Structure of α -Keratin in Hair



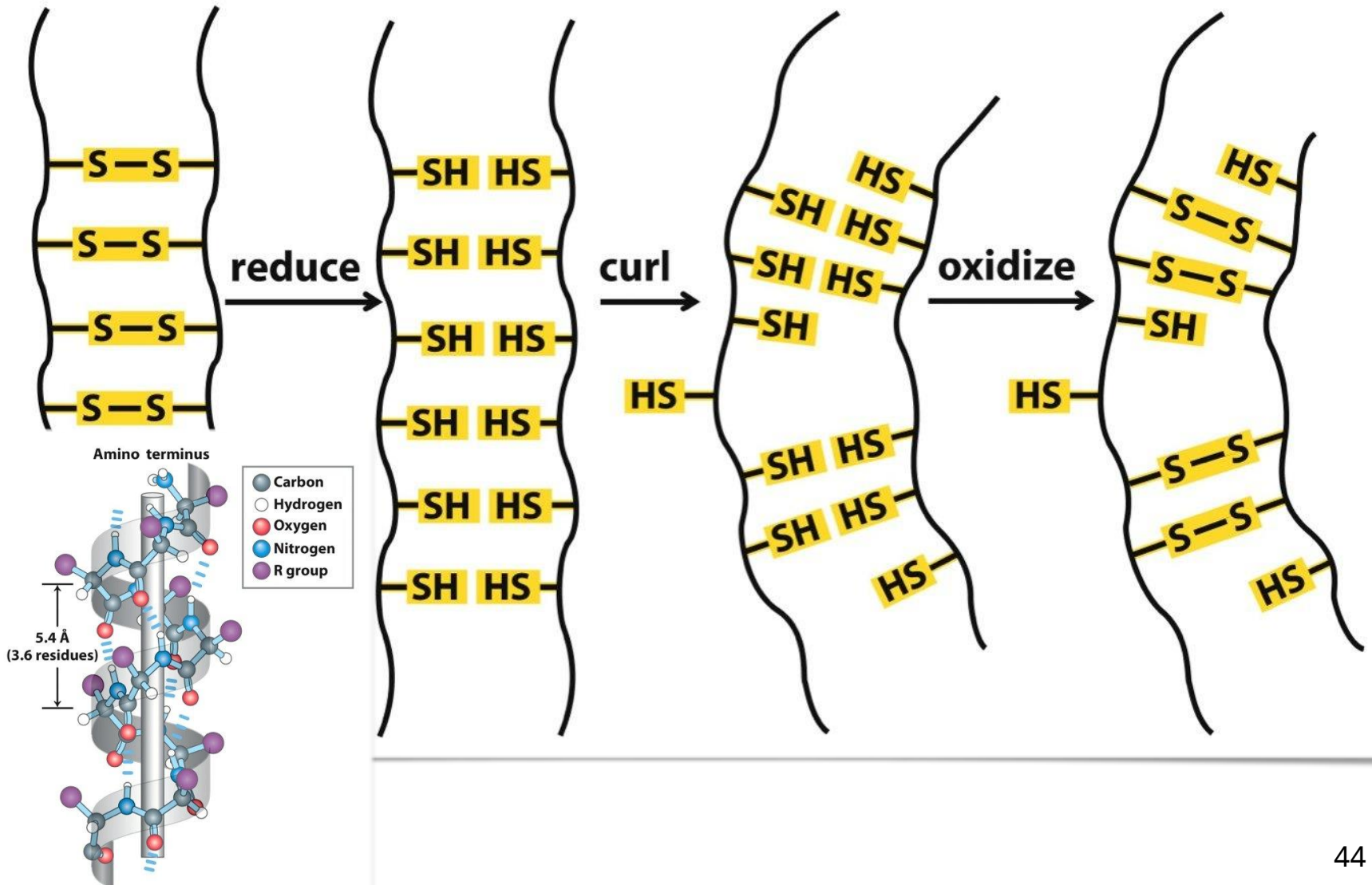
Structure of Hair



- Right-handed α helix
- Left-handed coiled coil
- Cross-linked by disulfide bonds

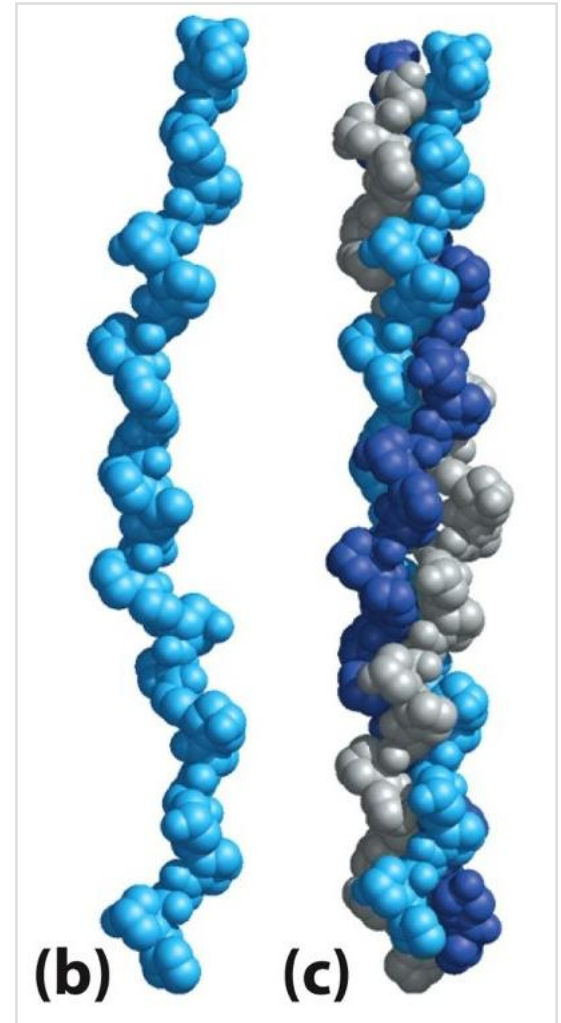
Cross section of a hair

Chemistry of Permanent Waving



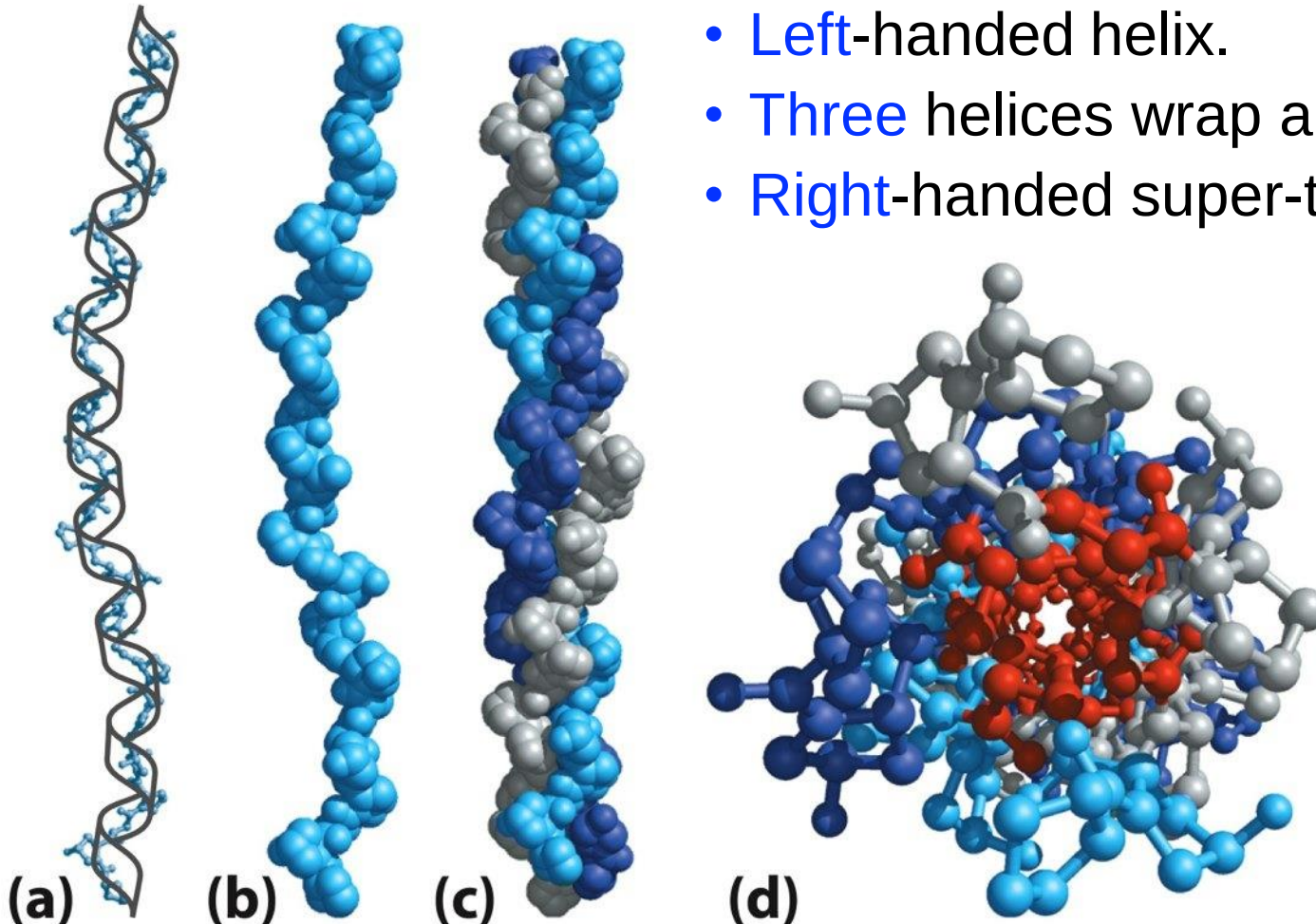
Structure of Collagen

- Each collagen chain is a long **Gly-rich** and **Pro-rich** **left-handed** helix.
- Three collagen chains intertwine into a **right-handed** superhelical **triple** helix.
- Many triple-helices assemble into a collagen fibril.



Structure of Collagen

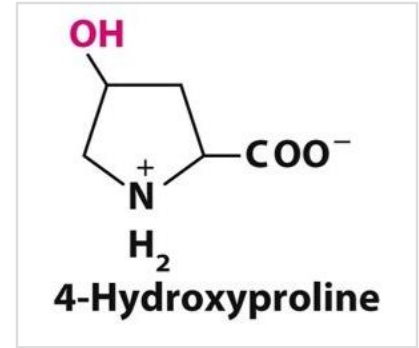
- **NOT** α -helix.
- **Left**-handed helix.
- **Three** helices wrap around.
- **Right**-handed super-twisting.



Glycine shown in red

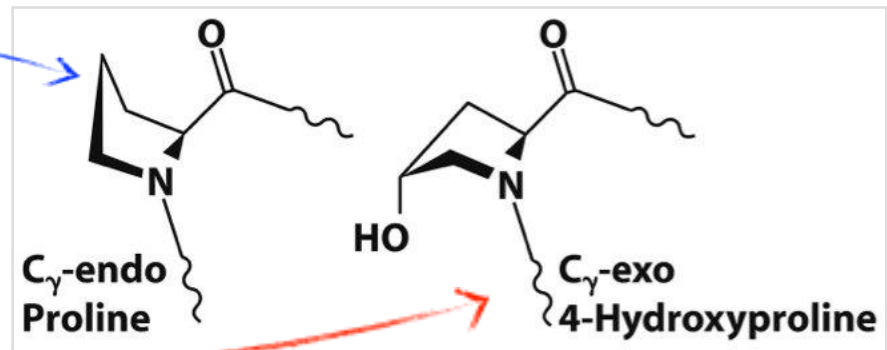
Proline & 4-Hydroxyproline in Collagen

- Repeating unit in collagen: Gly-X-Y (X is often Pro, Y is often 4-Hyp).
 - **Three** amino acid residues per turn.
 - Only **Gly** can be accommodated at very tight junctions.
 - Pro and 4-Hyp permit **sharp** twisting of collagen helix.
- 4-Hyp forces proline ring into a favorable pucker.
- 4-Hyp offers more **hydrogen bonds** between three strands of collagen.
- **Post-translational processing** is catalyzed by prolyl hydroxylase and requires **ascorbate (vitamin C)**.



C_γ-endo required in X position

C_γ-exo required in Y position



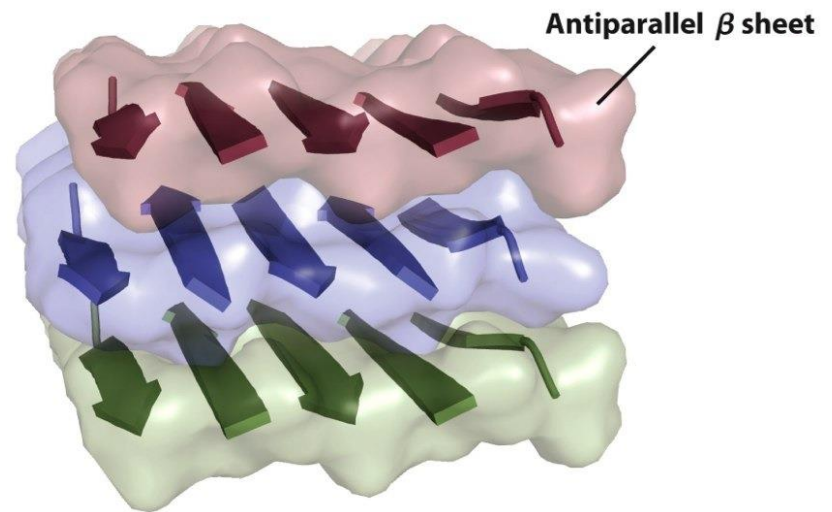
Collagen



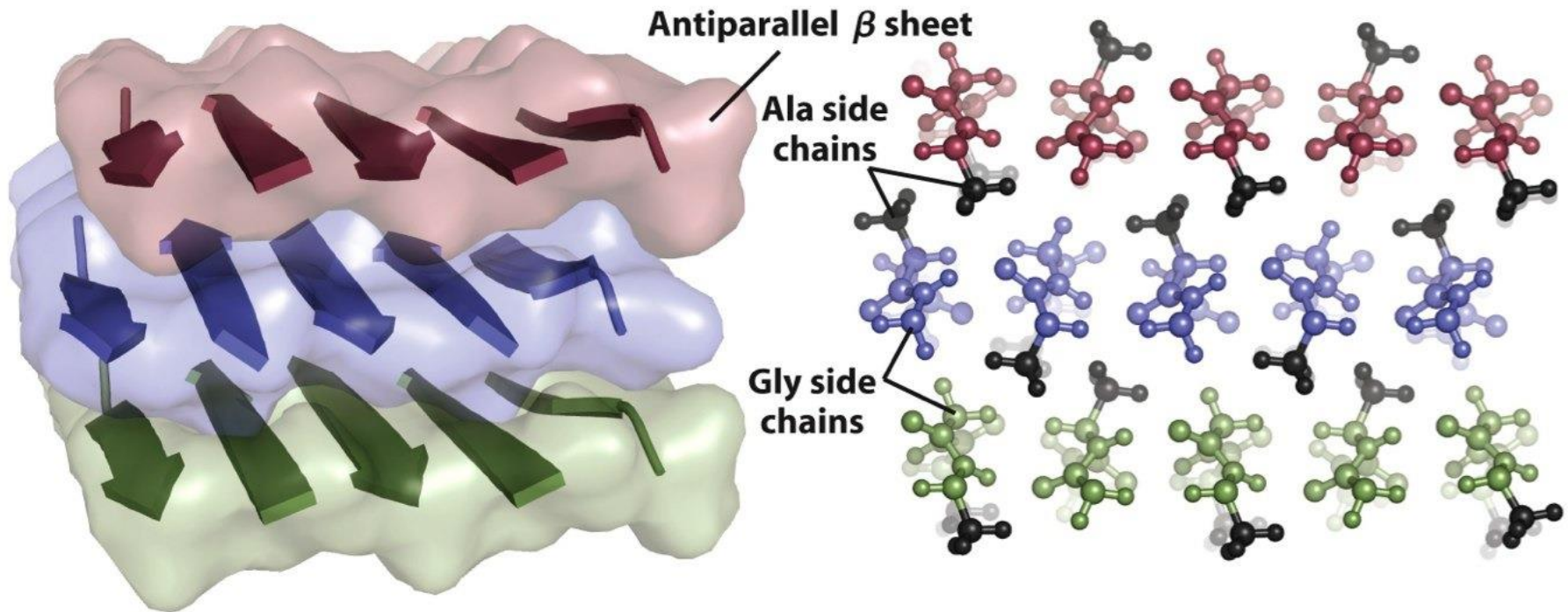
blood vessel 血管 These filaments are NOT individual protein strands. Each collagen monomer consists of **three left-handed** helical protein strands, wound around each other to form a **right-handed triple helix**.

Silk Fibroin

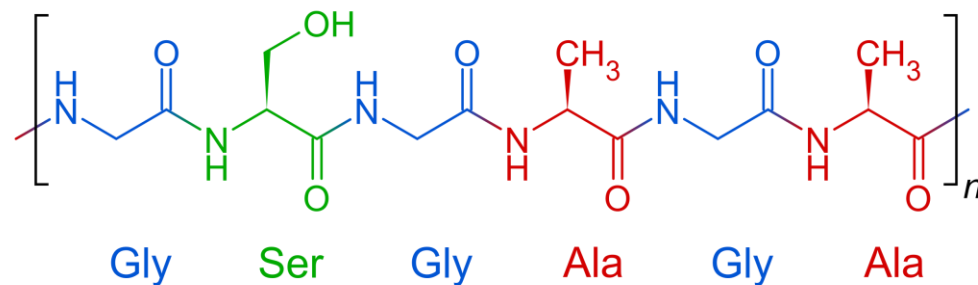
- Fibroin is the main protein in silk from insects and spiders.
- Antiparallel β sheet structure.
- Small side chains (**Ala** and **Gly**) allow the close packing of sheets.
- Structure is stabilized by
 - **Hydrogen bonding**.
 - **NOT covalent** bonds.



Structure of Silk



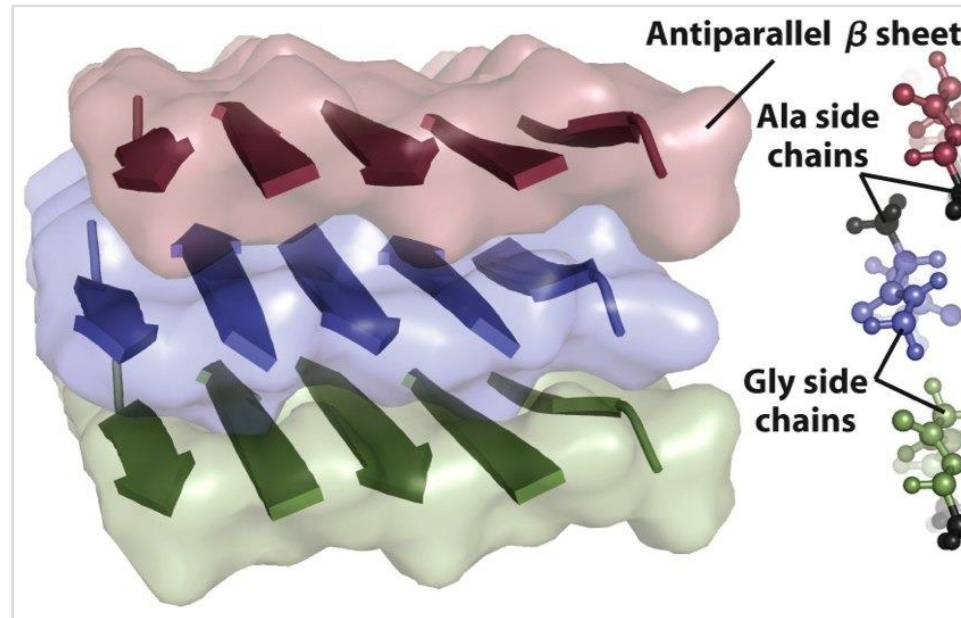
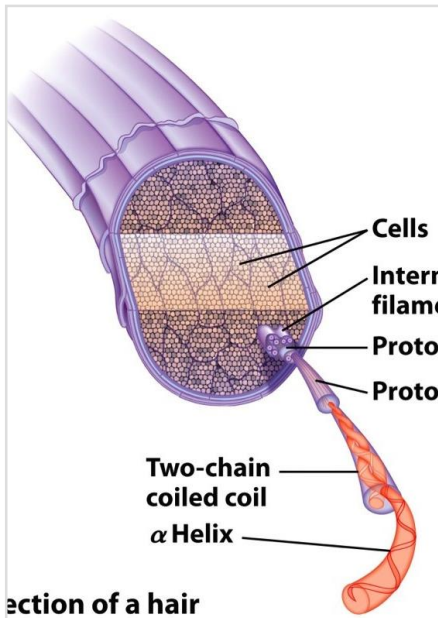
recurrent amino acid sequence Gly-Ser-Gly-Ala-Gly-Ala



Fibrous Proteins Review

TABLE 4-3 Secondary Structures and Properties of Some Fibrous Proteins

Structure	Characteristics	Examples of occurrence
α Helix, cross-linked by disulfide bonds	Tough, insoluble protective structures of varying hardness and flexibility	α -Keratin of hair, feathers, nails
β Conformation	Soft, flexible filaments	Silk fibroin
Collagen triple helix	High tensile strength, without stretch	Collagen of tendons, bone matrix



Structural Diversity -> Functional Diversity

- Globular proteins more **compact** than fibrous proteins.
- A wide array of biological functions.
 - Enzymes, regulatory proteins, transport proteins, etc.

β Conformation

$2,000 \times 5 \text{ \AA}$

Human serum albumin has 585 residues in a single chain

α Helix

$900 \times 11 \text{ \AA}$

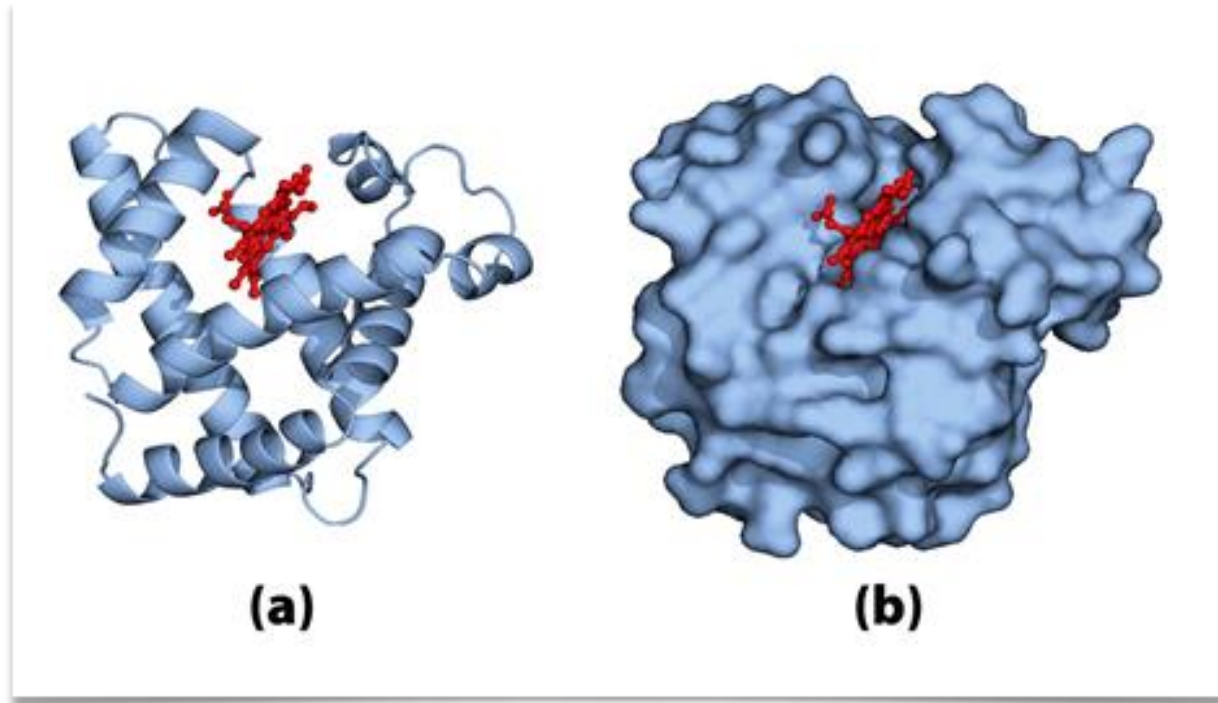


Native globular form

$100 \times 60 \text{ \AA}$

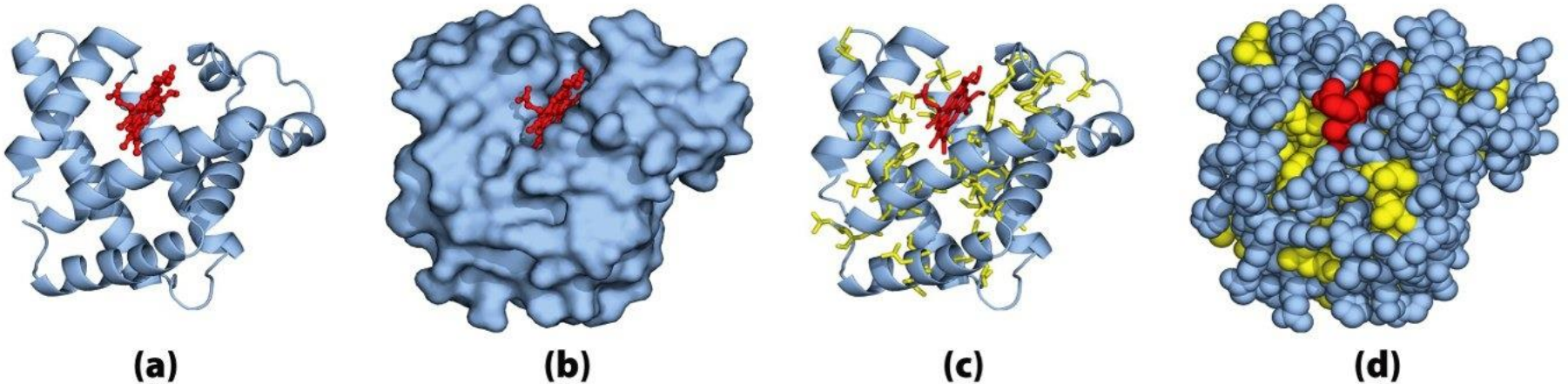
Myoglobin

- A single peptide chain with 153 residues.
- Oxygen binding.
- Contains a single heme group (iron protoporphyrin).



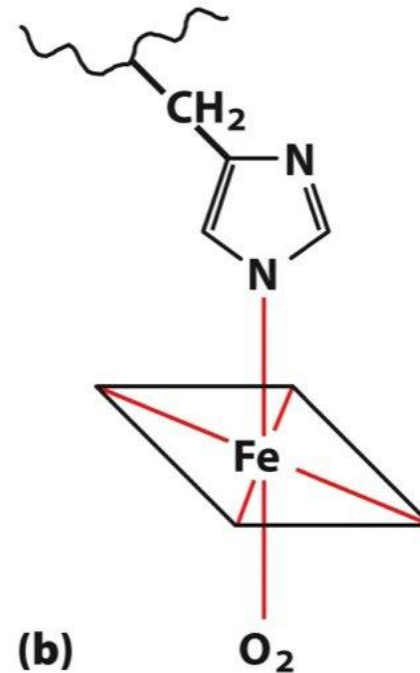
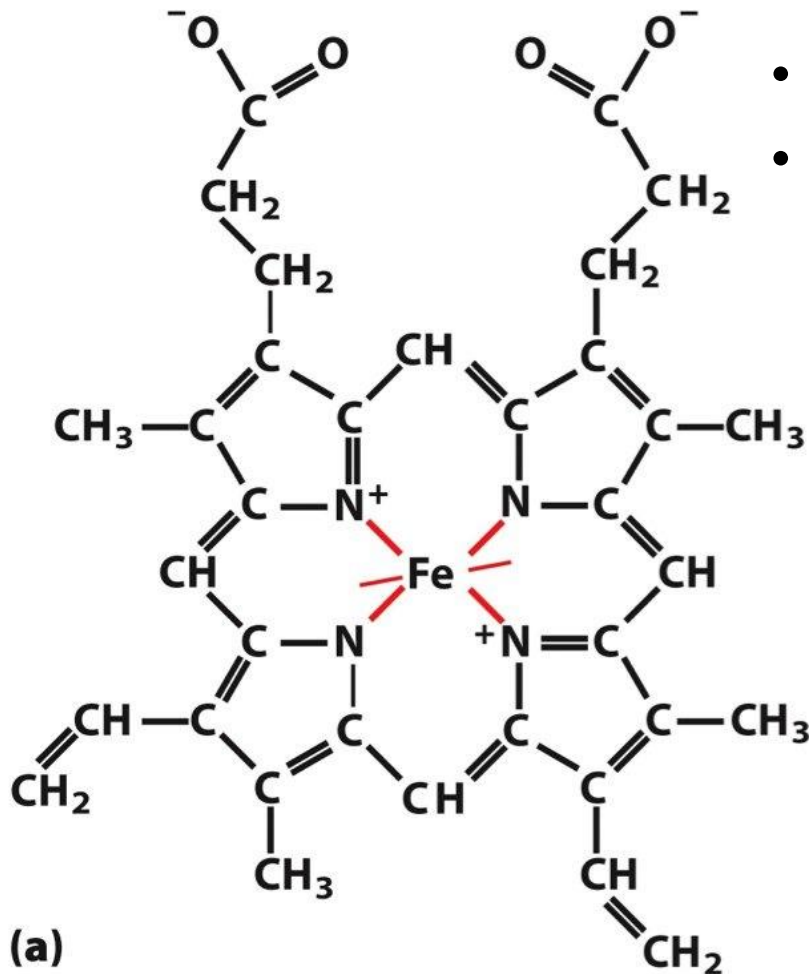
Structure of Myoglobin

- (a) Ribbon representation
- (b) Surface representation
- (c) Ribbon representation with several side chains shown as sticks
- (d) Space-filling model



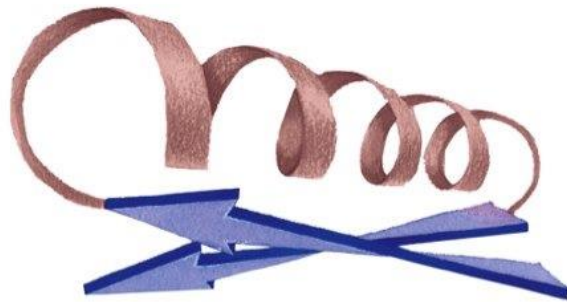
Structure of Heme

- Center: Fe^{2+} (ferrous) ion.
- Heterocyclic macrocycle porphyrin.
- Histidine side chain.



Motif or Fold

- A recognizable folding pattern
- Involves two or more elements of secondary structure
- Can be very simple or complex



β - α - β Loop

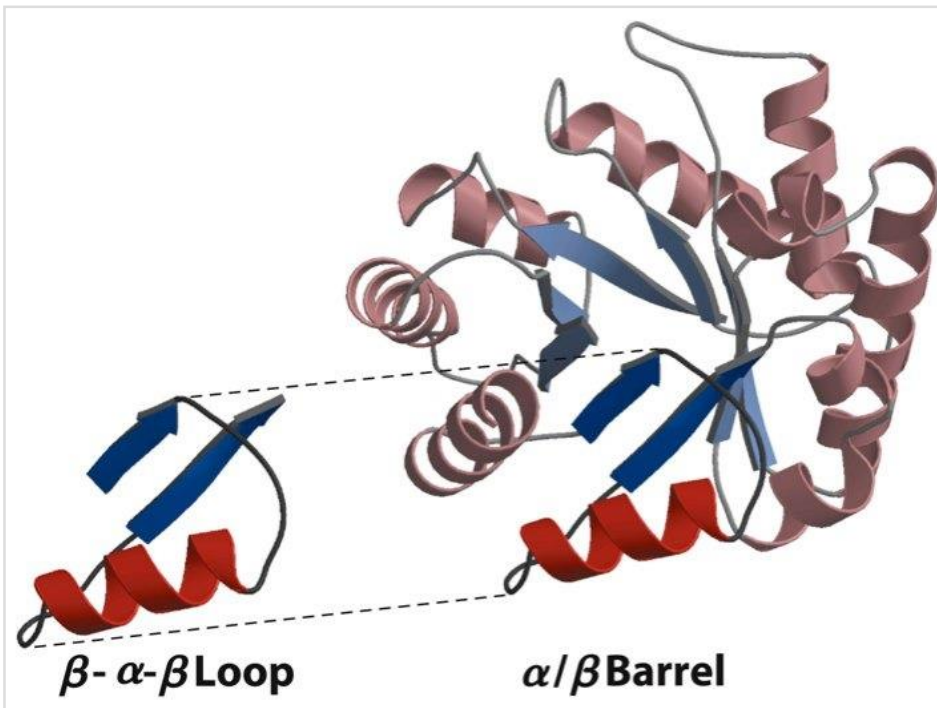


β Barrel

Examples of Motifs

Keratin α helix — 

Two-chain coiled coil — 

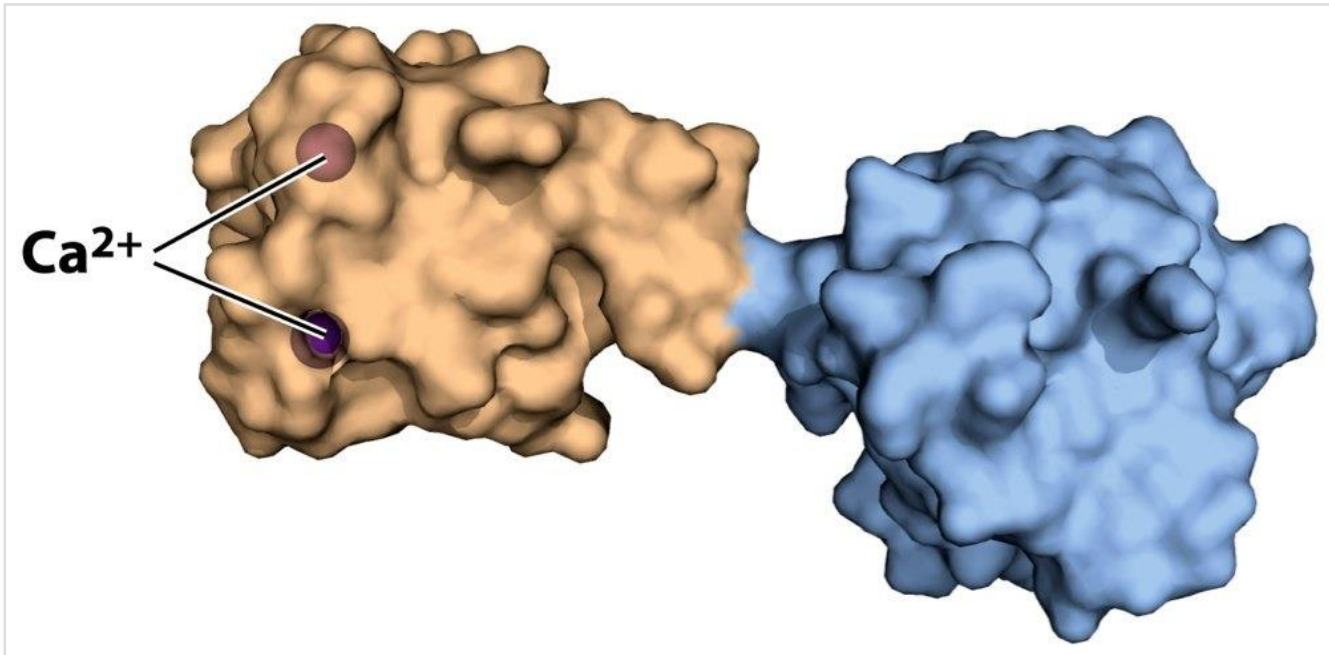


Globin Fold

- Reoccurring in many proteins.
- A single large motif may be the entire protein.
 - NOT a hierarchical structural element.
 - A folding pattern.

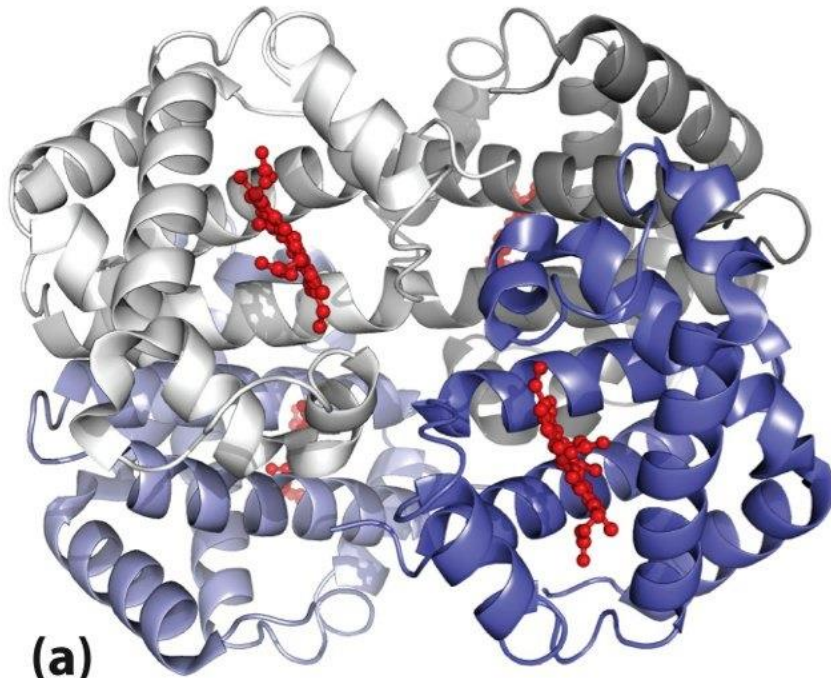
Domain

- A part of a polypeptide chain.
- Independently stable or move as a single entity.
- **Maintain** structure and function when separated.
- For small proteins, **the domain is the protein**.

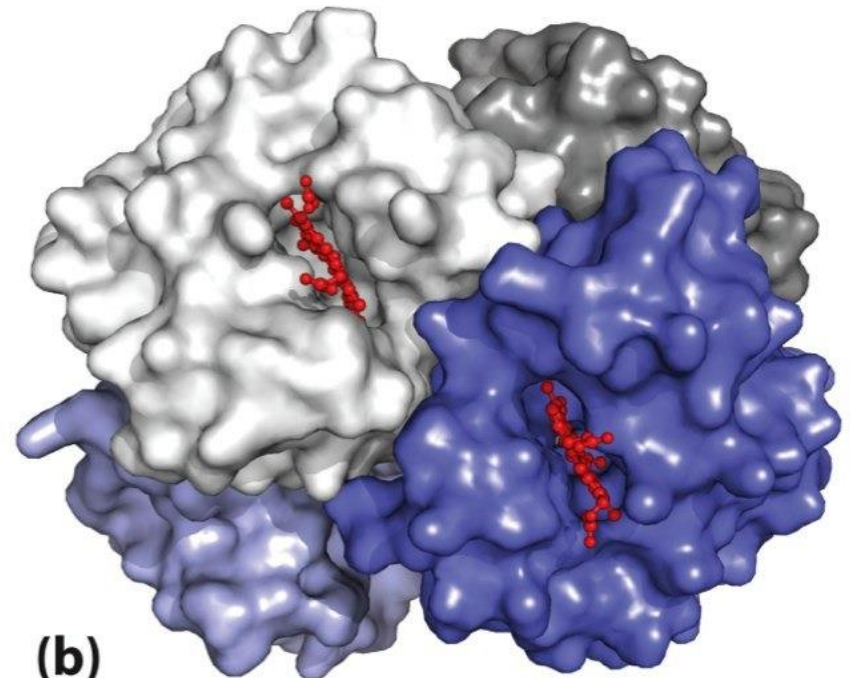


Quaternary Structure

Quaternary structure is formed by the assembly of **individual polypeptides** into a larger functional cluster.



Ribbon representation

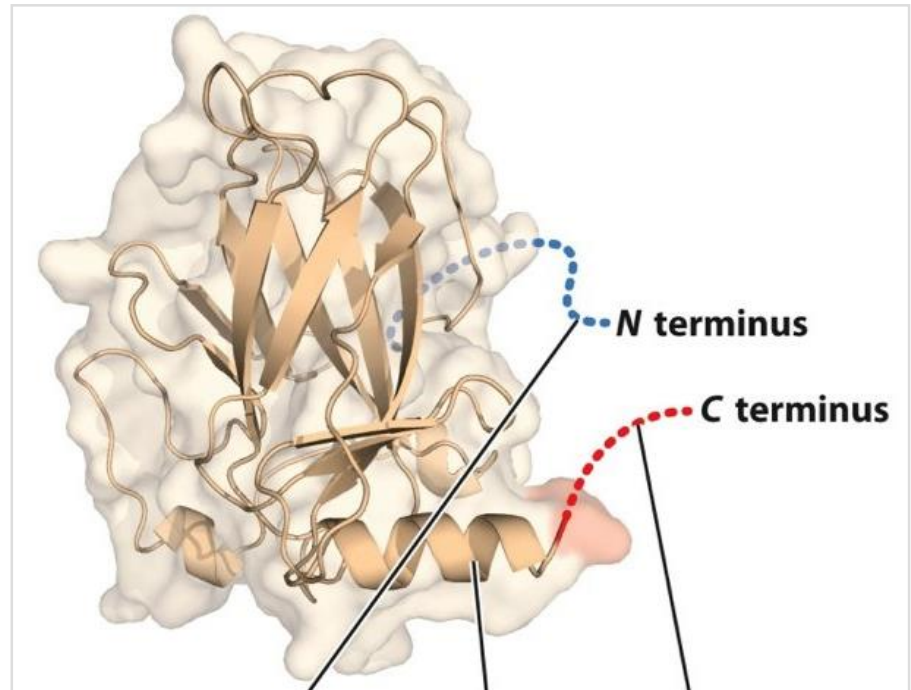


Surface model

Hemoglobin

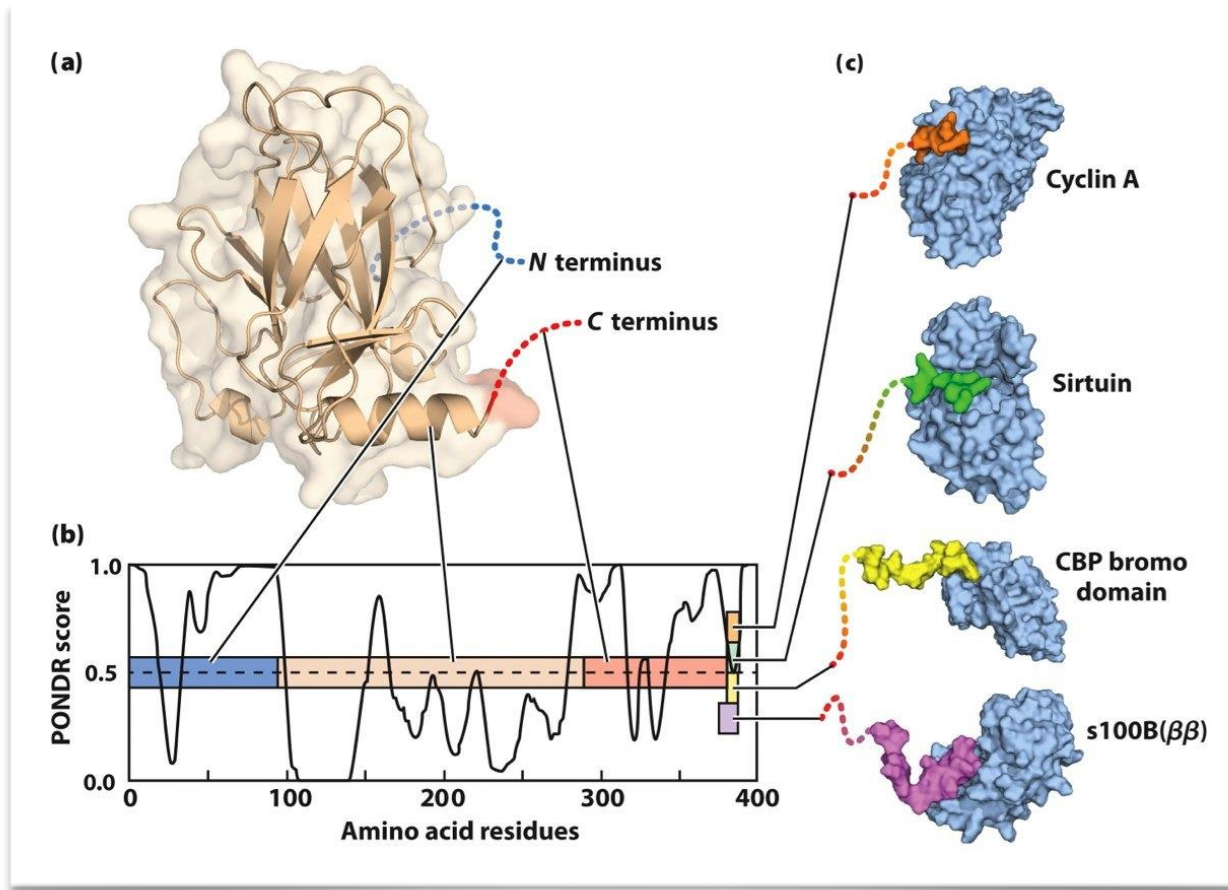
Intrinsically Disordered Proteins

- Contain protein segments **lack** definable structure.
- Composed of amino acids whose higher concentration **forces less-defined structure**.
 - Lys, Arg, Glu, and Pro
- Disordered regions can interact with **many different partner** proteins

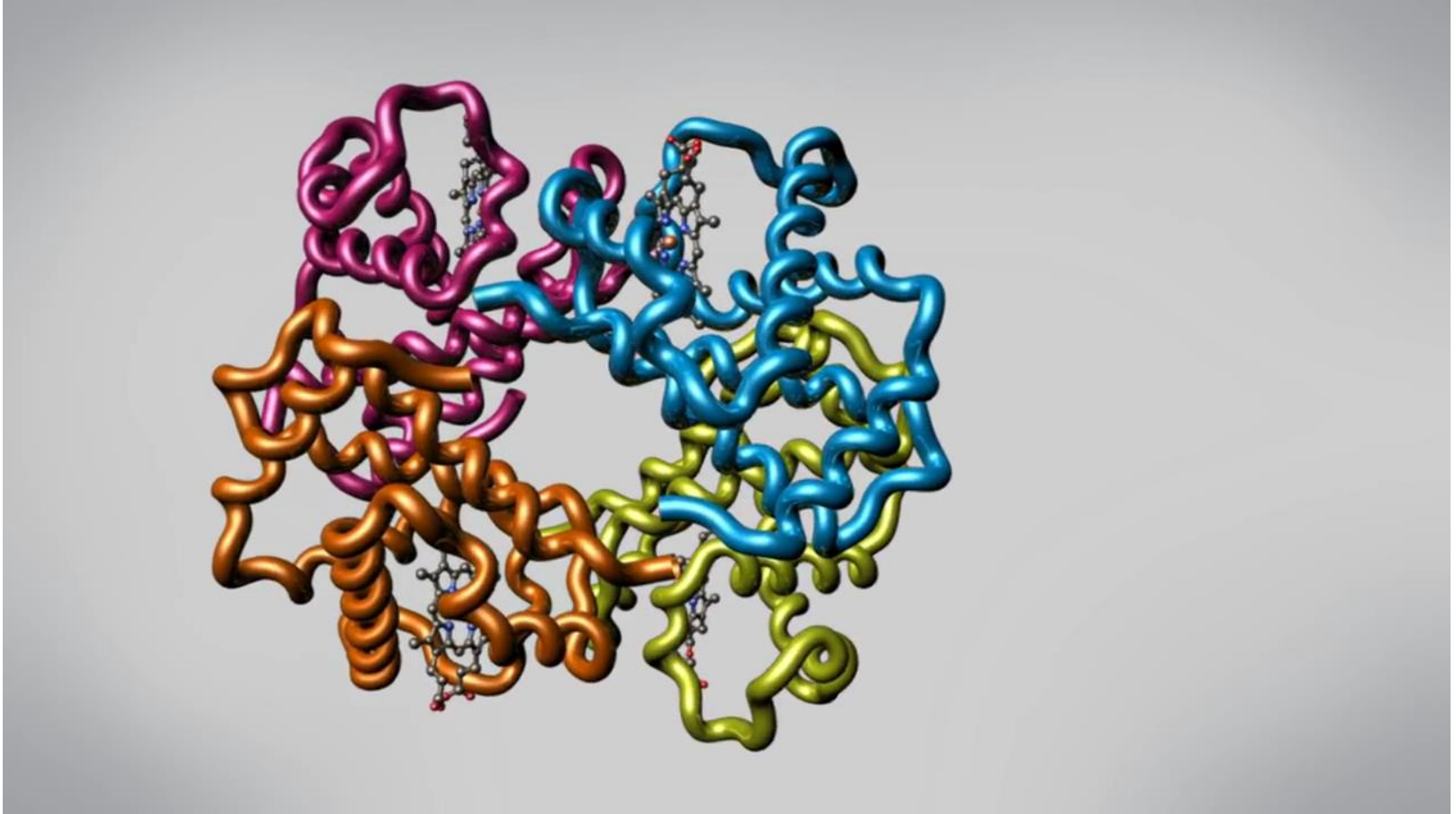


Intrinsically Disordered p53 Protein

- Inhibit by wrapping around target proteins.
- Multiple targets.
- Bind different targets in different ways.



What is A Protein?

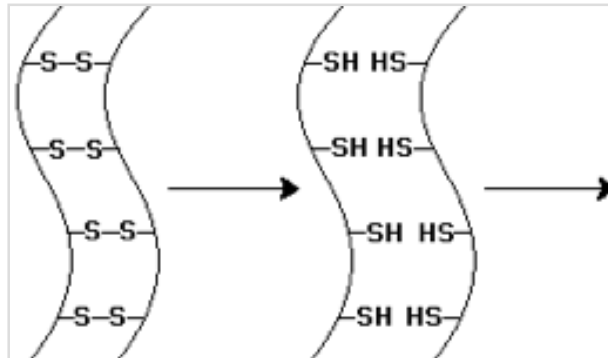


Summary 4.3 Tertiary & Quaternary Structure

- Tertiary structure is the **complete** three-dimensional structure of a polypeptide chain. Quaternary structure describes interactions between subunits of **multisubunit** proteins.
- **Fibrous** proteins mainly serve structural roles. **Globular** proteins have more complicated tertiary structures.
- Some proteins are intrinsically disordered. Some **function** as versatile inhibitors.

Example Question

The α -keratin chains indicated by the diagram below have undergone one chemical step. To alter the shape of the α -keratin chains—as in hair waving—what subsequent steps are required?



- A) Chemical oxidation and then shape remodeling
- B) Chemical reduction and then chemical oxidation
- C) Chemical reduction and then shape remodeling
- D) Shape remodeling and then chemical oxidation
- E) Shape remodeling and then chemical reduction

Example Question

Which of the following statements concerning protein domains is *true*?

- A) They are a form of secondary structure.
- B) They are examples of structural motifs.
- C) They consist of separate polypeptide chains (subunits).
- D) They have been found only in prokaryotic proteins.
- E) They may retain their correct shape even when separated from the rest of the protein.

Example Question

Which statement about intrinsically disordered proteins is *true*?

- A) They contain small hydrophobic cores.
- B) They represent misfolded conformations of cellular proteins.
- C) They have no stable three-dimensional structure and therefore have no cellular function.
- D) They can interact with multiple protein-binding partners and are central to protein interaction networks.

Three-Dimensional Structure of Proteins

4.1 Overview of Protein Structure

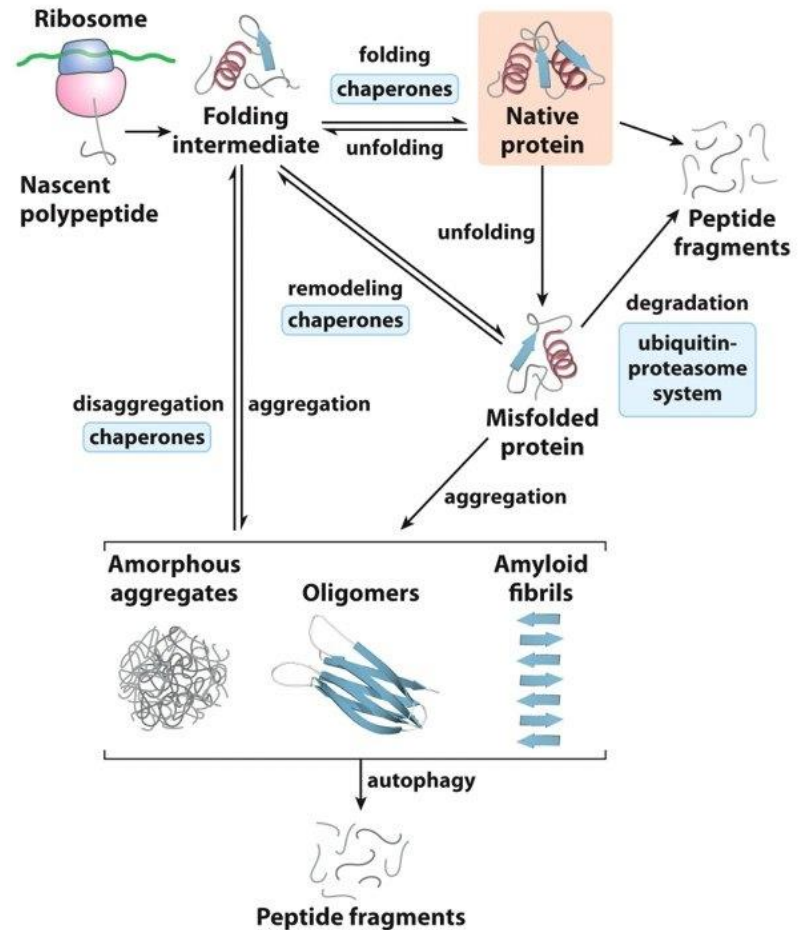
4.2 Protein Secondary Structure

4.3 Protein Tertiary and Quaternary Structure

4.4 Protein Denaturation and Folding

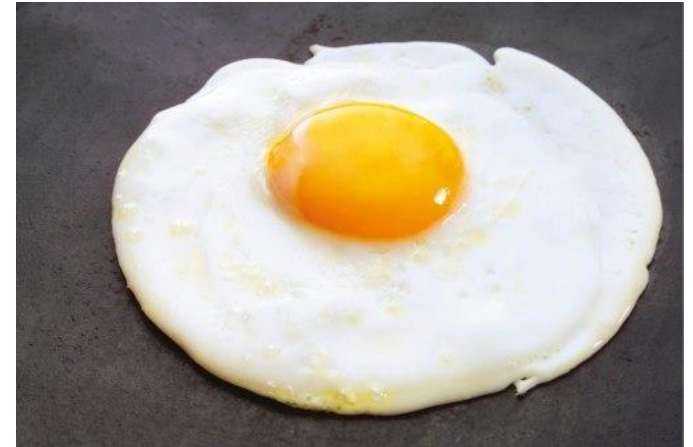
Proteostasis

- Continual maintenance of an **active** set of cellular proteins under a given set of conditions.
- Three processes:
 - Synthesis
 - **Folding**
 - Degradation

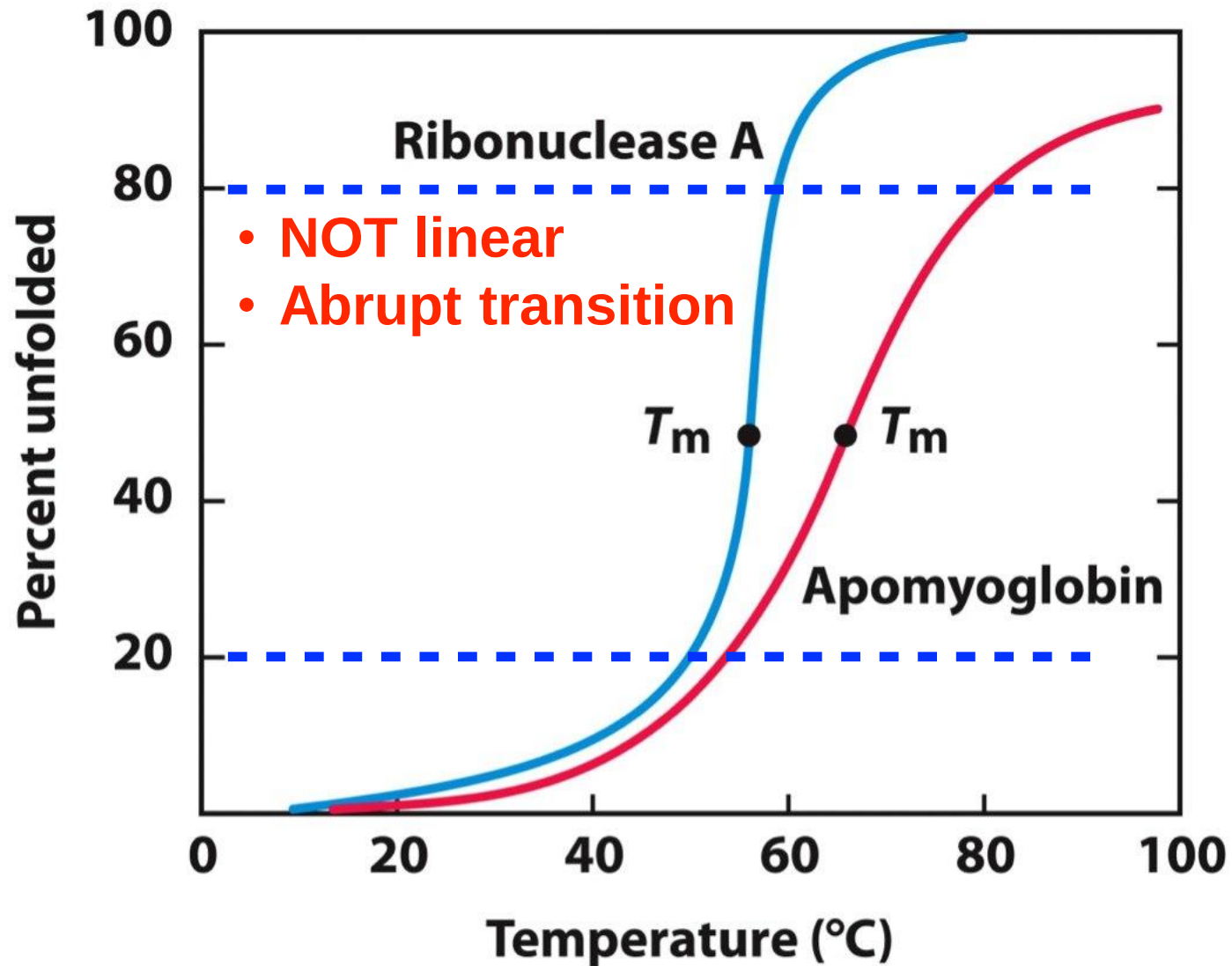


Loss of Structure -> Loss of Function

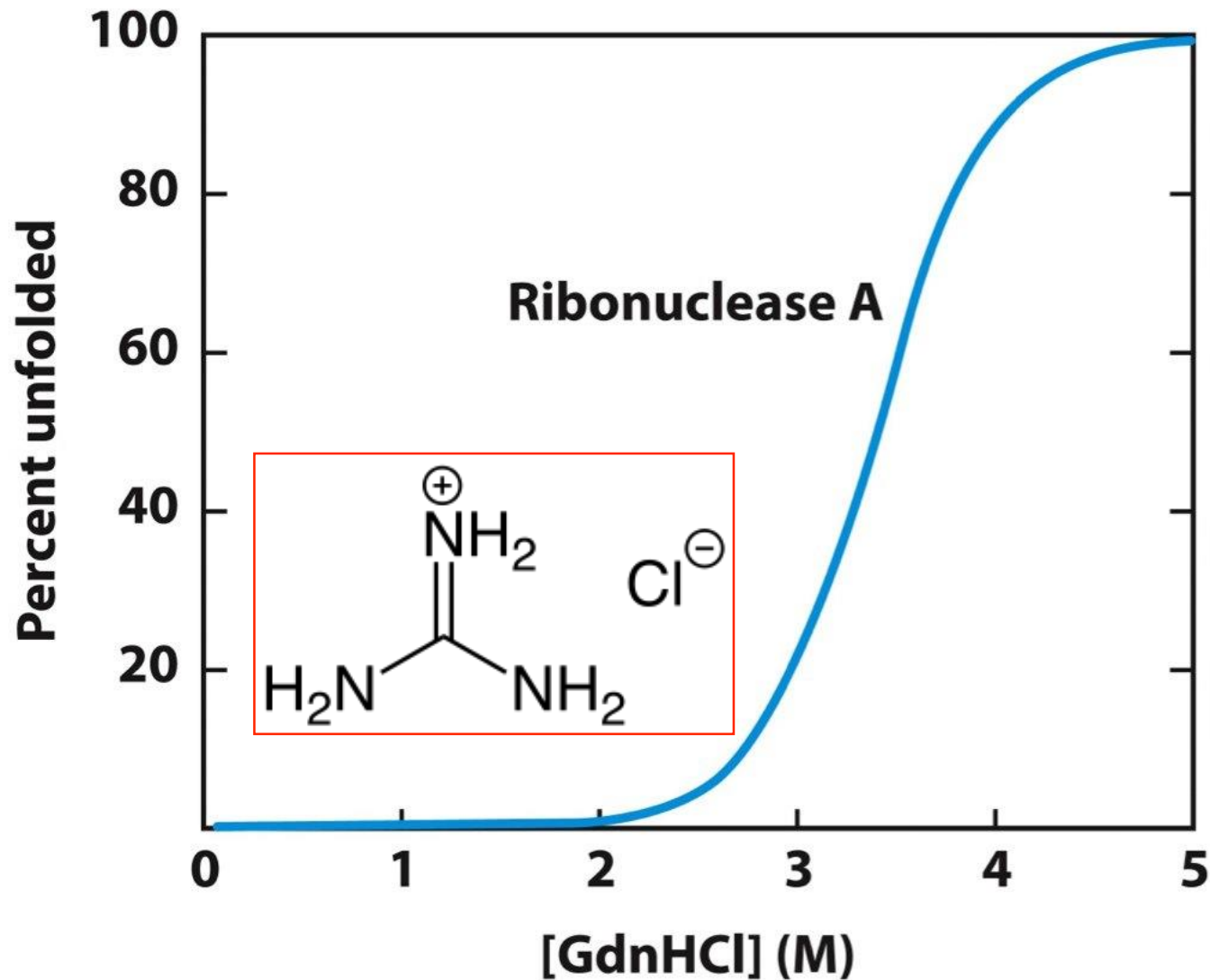
- A protein's function depends on its 3D-structure.
- Loss of structural integrity with accompanying loss of activity is called **denaturation**.
- Proteins can be denatured by:
 - Heat
 - pH extremes
 - Organic solvents
 - Urea, detergent and guanidinium hydrochloride



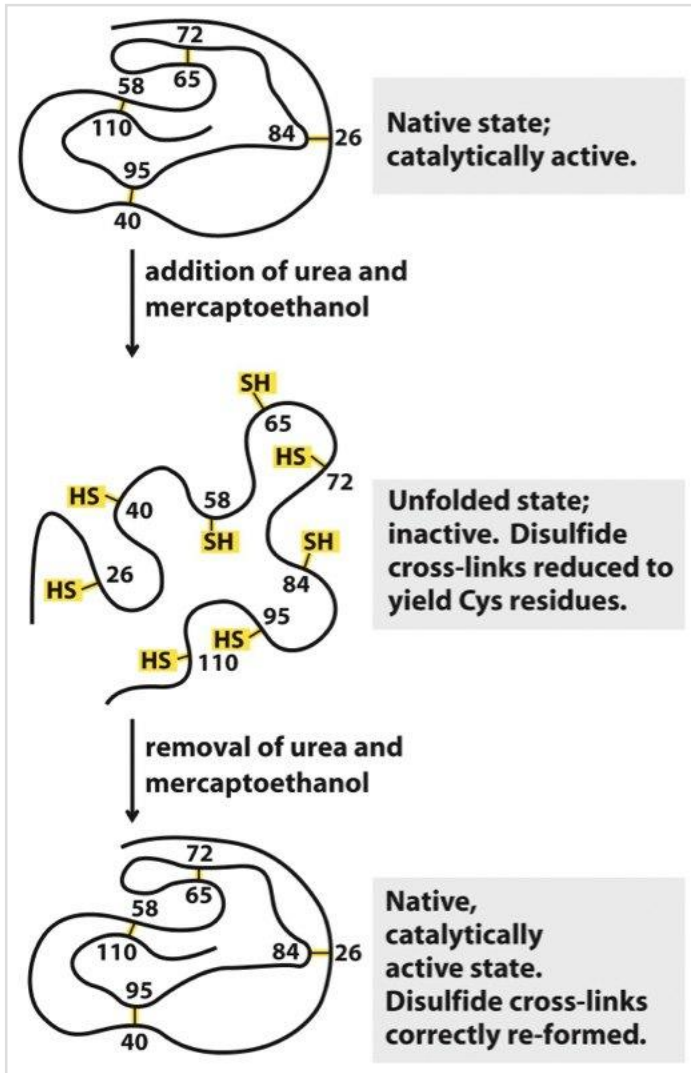
Protein Denaturation by Heat



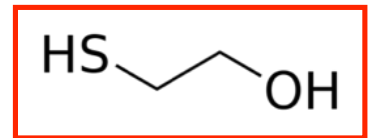
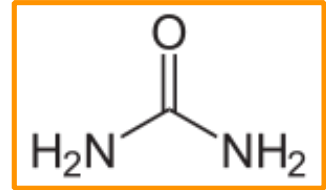
Denaturation by Guanidine Hydrochloride



Ribonuclease Refolding Experiment



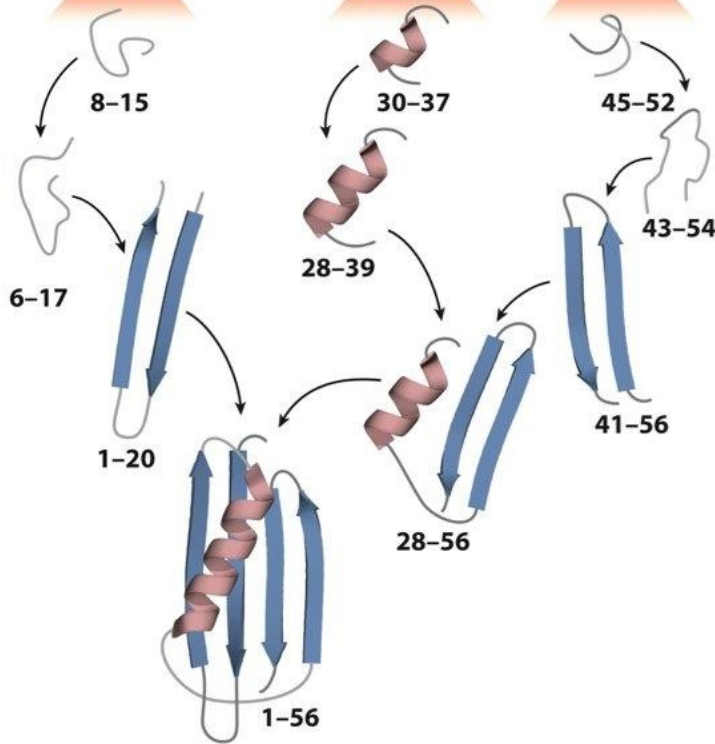
- A small protein
- 8 cysteines
- 4 disulfide bonds
- **Urea** and **BME** fully denatures
- When urea and BME are removed, the protein spontaneously **refolds**, and the correct disulfide bonds are reformed
- **Sequence alone determines the native conformation.**



Proteins Fold by a Stepwise Process

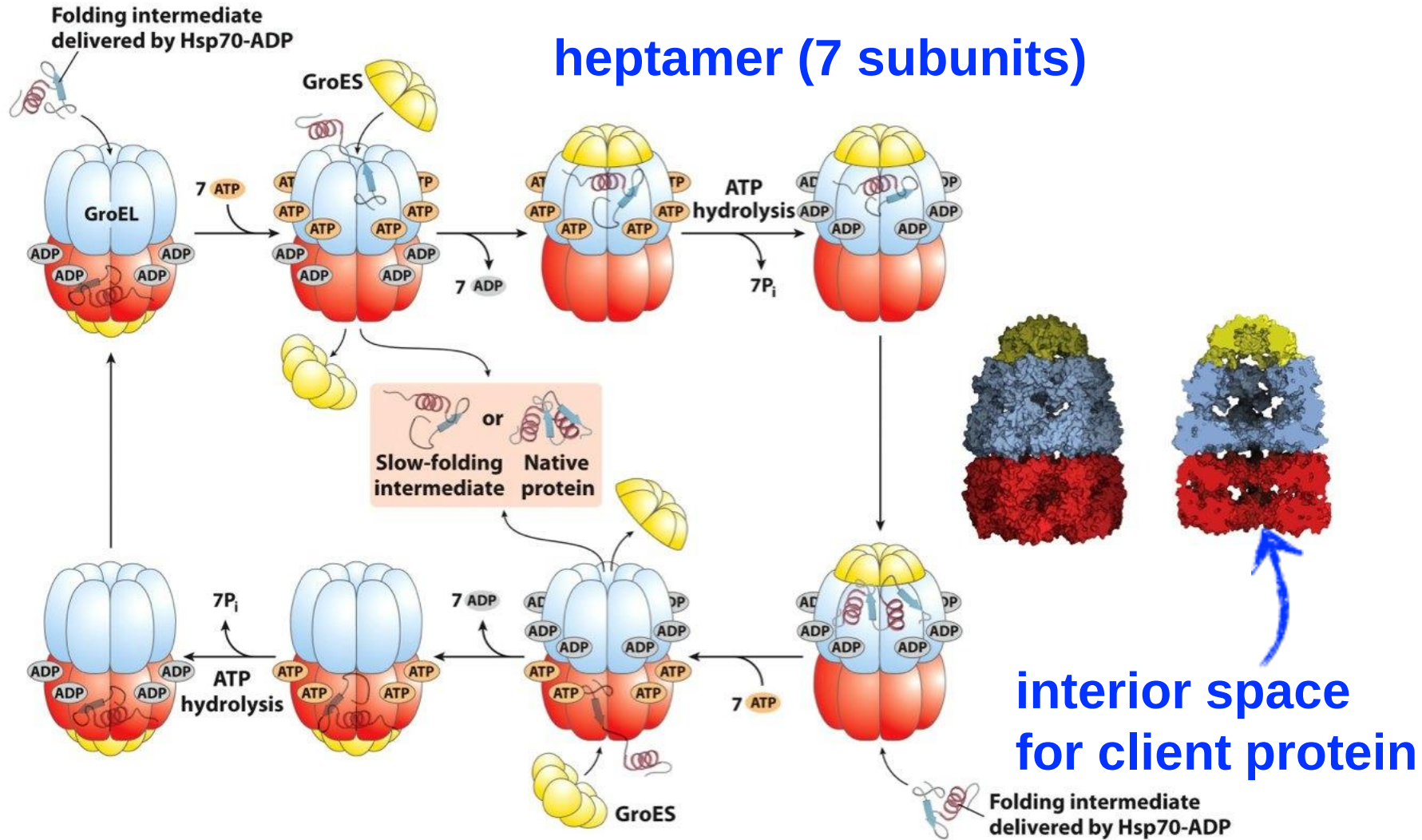
Amino acid sequence of a 56-residue peptide

MTYKLIL**NGKTLKG**ETTTEAVDAATAEKV **FKQYANDN**GVDGEWT **YDDATKTF**TVTE

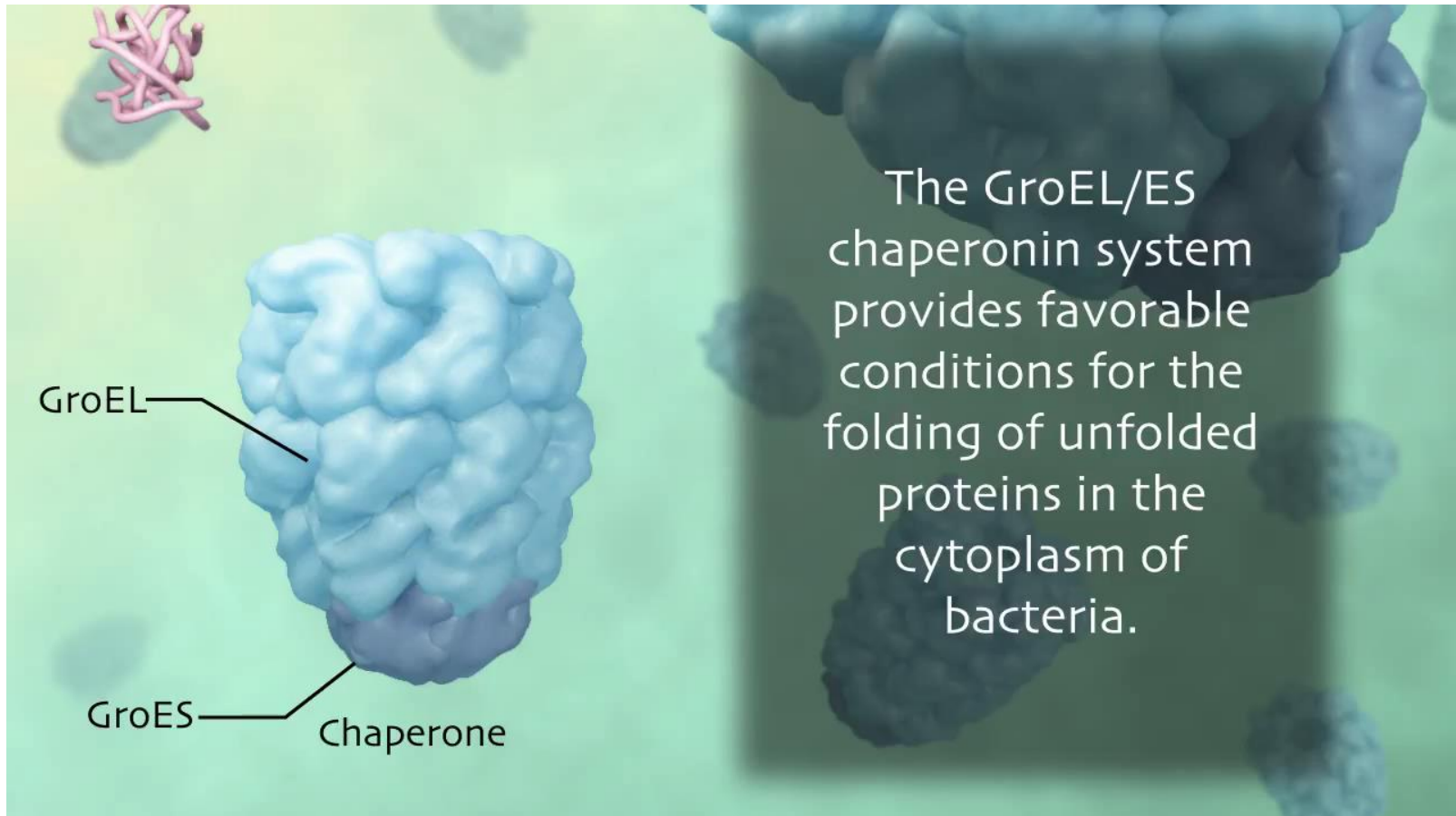


- **Local** structures first.
2° structures assembled.
- Then **long-range** interactions.
 - 2° structures further interact to form 3° structure.
- **Hydrophobic** interactions play a significant role throughout the process.

Assisted Folding by Chaperones



Assisted Folding by Chaperones



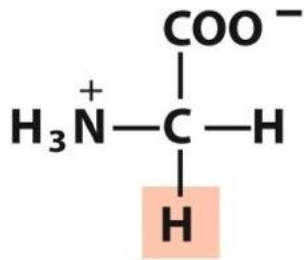
Summary 4.4 Protein Folding & Denaturation

- Maintenance of a collection of active cellular proteins is called **proteostasis**. It involves three processes: synthesis, folding, and degradation.
- Structure and function can be destroyed by **denaturation**. Some denatured proteins can **renature** spontaneously. 3° structure is determined by 1° sequence.
- Protein folding is **hierarchical** and often assisted by **chaperones**.

Example Question

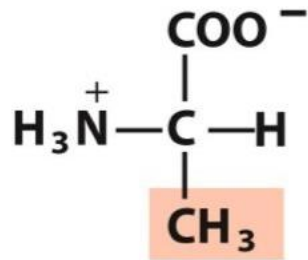
Experiments on denaturation and renaturation after the reduction and re-oxidation of the —S—S— bonds in the enzyme ribonuclease (RNase) have shown that:

- A) folding of denatured RNase into the native, active conformation, requires input of energy in the form of heat.
- B) native ribonuclease does not have a unique secondary and tertiary structure.
- C) the completely unfolded enzyme, with all —S—S— bonds broken, is still enzymatically active.
- D) the enzyme, dissolved in water, is thermodynamically stable relative to the mixture of amino acids whose residues are contained in RNase.
- E) the primary sequence of RNase is sufficient to determine its specific secondary and tertiary structure.



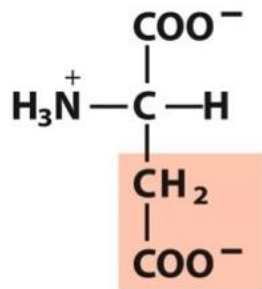
Glycine

Gly, G



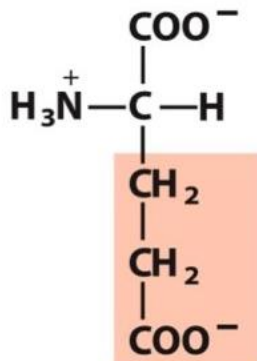
Alanine

Ala, A



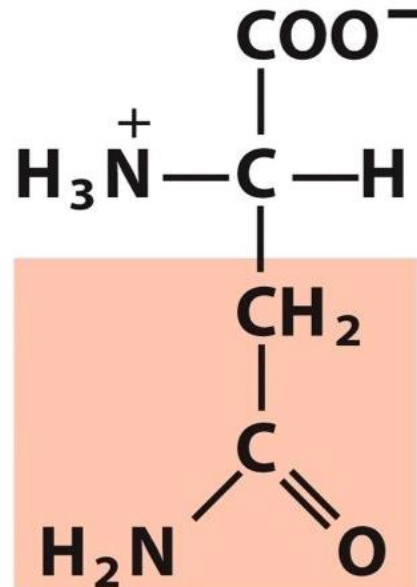
Aspartate

Asp, D



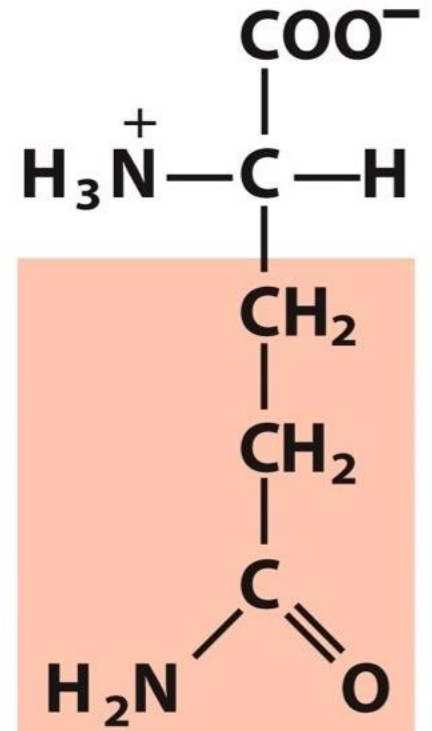
Glutamate

Glu, E



Asparagine

Asn, N



Glutamine

Gln, Q

Amino acid for 3rd week

Chapter 4: Summary

In this chapter, we learned about:

- Two most important secondary structures.
 - α helices
 - β sheets
- Three common fibrous proteins and one globular protein.
 - α keratin
 - collagen
 - silk fibroin
 - myoglobin
- Protein denaturation & folding.