# PROTEINS: THREE-DIMENSIONAL STRUCTURE AND FUNCTION

Course: Biochemistry I (BIOC 230)

**Textbook:** 

Principles of Biochemistry, 5th Ed., by L. A. Moran and

others. 2014, Pearson. . Chapter 4

## **Terminology**

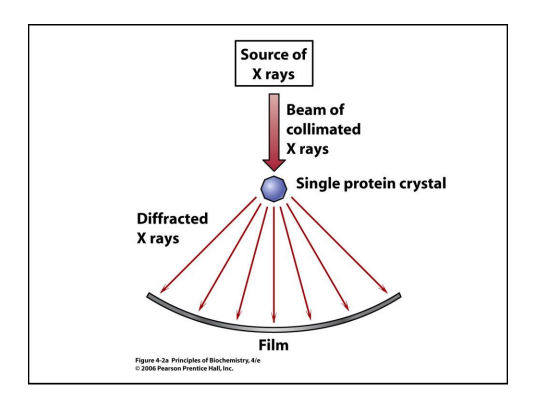
- □ Protein: a chain of amino acids joined by peptide bonds in a specific sequence
- □ **Conformation** spatial arrangement of atoms in a protein, that depends on the rotation of a bond or bonds (?)
- Native conformation conformation of functional protein

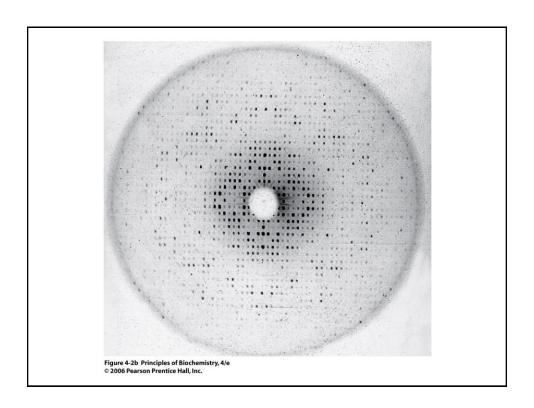
#### From DNA to Protein

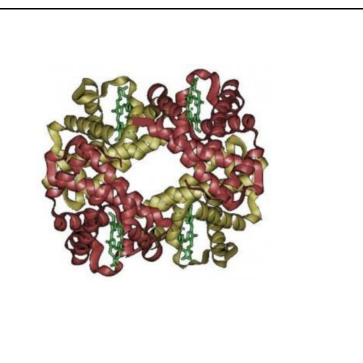
- □ In some species, the size and sequence of every polypeptide can be determined from the sequence of the genome
- □ Genomics: the study of the structure of whole genome
- □ In E. coli, there are 4000 different polypeptides with an average size of about 300 aa
- □ In fruit fly (Drosophila melanogaster) there are 14,000 different polypeptides
- Humans and mammals there are 30,000 different polypeptides
- Proteomics: the science that studies large sets of proteins,
- Proteome: all proteins produced by a cell

#### Methods for determining protein structure

- Primary structure or amino acid sequence of a polypeptide is determined using chemical methods such as Edman degradation or indirectly from the sequence of a gene
- Three-dimensional structure of protein:
  - X-ray crystallography
  - Nuclear magnetic resonance (NMR)

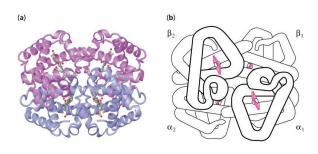






# Protein Classification

- □ **Simple** composed only of amino acid residues
- □ **Conjugated** contain prosthetic groups (metal ions, co-factors, lipids, carbohydrates) Example: Hemoglobin – conjugated to Heme



#### **Protein Classification**

- □ One polypeptide chain: monomeric protein
- □ More than one multimeric protein
- □ Homomultimer one kind of chain
- Heteromultimer two or more different chains
- E.g., Hemoglobin is a heterotetramer. It has two alpha chains and two beta chains:  $\alpha_2\beta_2$

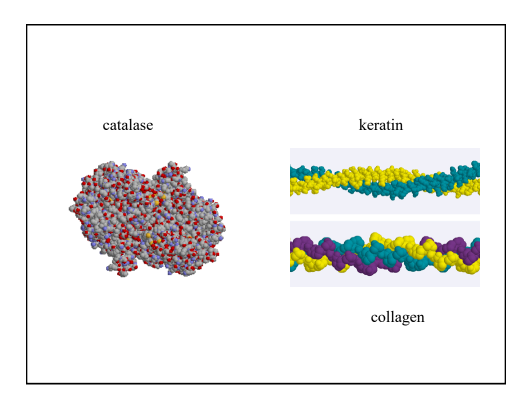
#### **Protein Classification**

#### **Fibrous**

- Polypeptides arranged in long strands or sheets
- Water insoluble (lots of hydrophobic aa's)
- Strong but flexible
- Structural (e.G., Keratin, collagen)

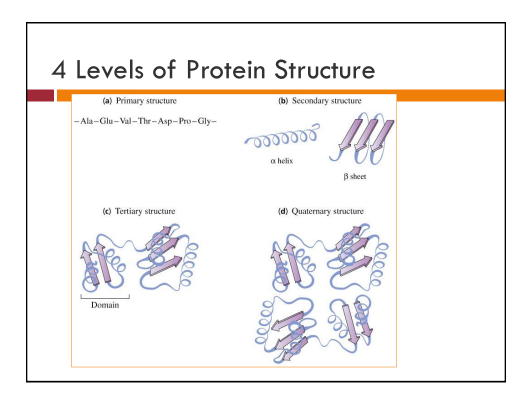
#### Globular

- Polypeptide chains folded into spherical or globular form 1)
- Water soluble 2)
- Contain several types of secondary structure
- Diverse functions (enzymes, regulatory proteins)



#### **Protein Function**

- □ Catalysis enzymes
- □ Structural keratin
- □ Transport hemoglobin
- $\Box$  Trans-membrane transport Na+/K+ ATPases
- □ Toxins rattle snake venom, ricin
- □ Contractile function actin, myosin
- □ Hormones insulin
- □ Storage Proteins seeds and eggs
- □ Defensive proteins antibodies



Non-covalent forces important in determining protein structure

□ van der Waals: 0.4 - 4 kJ/mol

□ hydrogen bonds: 12-30 kJ/mol

□ ionic bonds: 20 kJ/mol

□ hydrophobic interactions: <40 kJ/mol

#### Questions!

- 1. Which of the following level of protein structure determines all other levels?
  - a. Primary structure
  - b. Secondary structure
  - c. Tertiary structure
  - d. Quaternary structure
- The peptide bond that links amino acids in a polypeptide is formed between
  - a. Alpha-carboxyl group of first amino acid and alphaamino group of the next amino acid
  - Alpha-carboxyl group of first amino acid and R-group of the next amino acid

# 1° Structure Determines 2°, 3°, 4° Structure

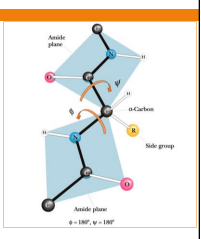
- □ Sickle Cell Anemia single amino acid change in hemoglobin related to disease
- Osteoarthritis single amino acid change in collagen protein causes joint damage

# Classes of 2° Structure

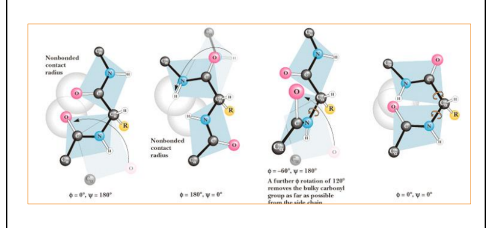
- Alpha helix
- □ B-sheet
- Loops and turns

## 2° Structure Related to Peptide Backbone

- •Double bond nature of peptide bond cause planar geometry
- •Free rotation at N  $\alpha$ C and  $\alpha$ C-carbonyl C bonds
- •Angle about the C(alpha)-N bond is denoted phi (\$\phi\$)
- •Angle about the C(alpha)-C bond is denoted psi  $(\psi)$
- •The entire path of the peptide backbone is known if all phi and psi angles are specified

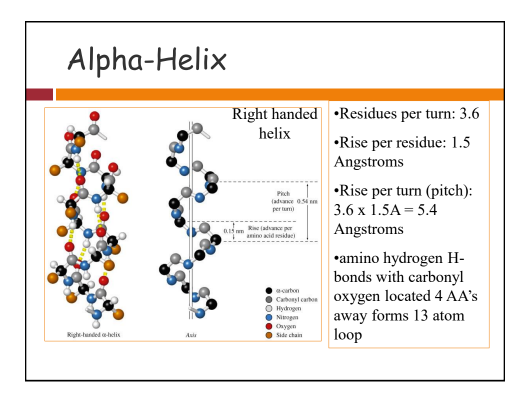


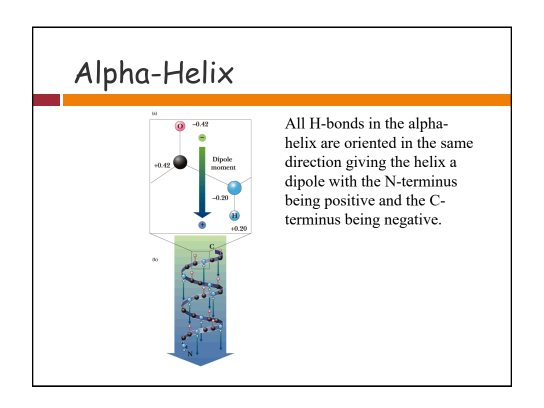
# Not all $\phi/\psi$ angles are possible

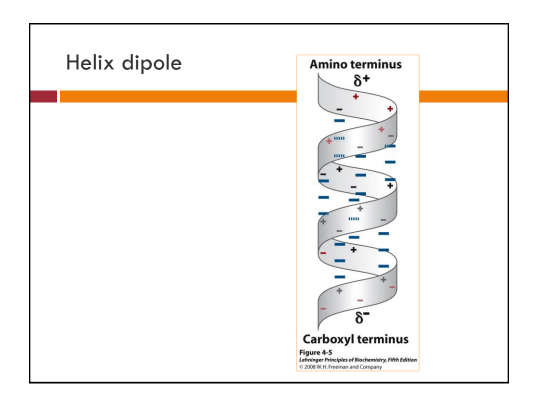


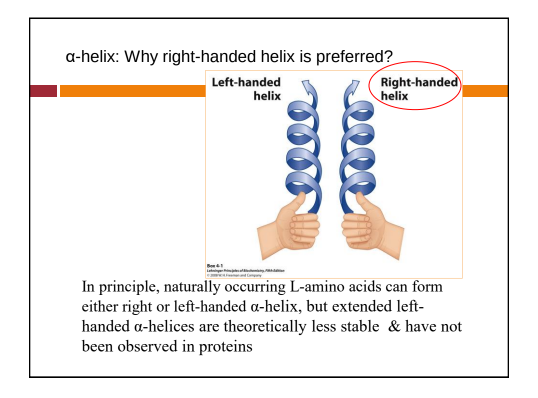
# Alpha-Helix

- First proposed by Linus Pauling and Robert Corey in 1951
- Identified in keratin by Max Perutz
- A ubiquitous component of proteins
- Stabilized by H-bonds

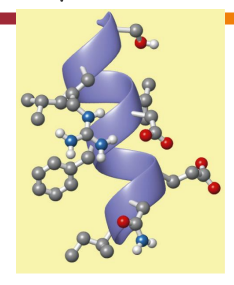






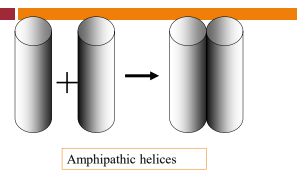


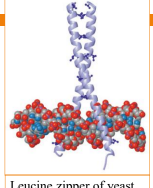
# Alpha-Helix



- •Side chain groups point outwards from the helix
- •AA's with bulky side chains less common in alpha-helix
- •Glycine and proline destabilizes alpha-helix

## **Amphipathic Alpha-Helices**





Leucine zipper of yeast

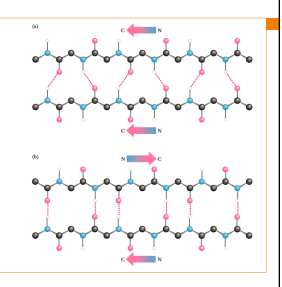
One side of the helix (dark) has mostly hydrophobic AA's Two amphipathic helices can associate through hydrophobic interactions. Such helices often occur on protein surface

#### Beta-Strands and Beta-Sheets

- Also first postulated by Pauling and Corey, 1951
- □ Strands may be parallel or antiparallel
- □ Rise per residue:
  - □ 3.47 Angstroms for antiparallel strands
  - □ 3.25 Angstroms for parallel strands
  - Each strand of a beta sheet may be pictured as a helix with two residues per turn

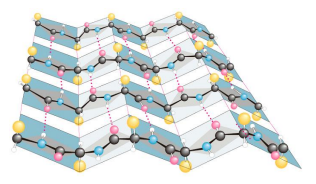
# Beta-Sheets

- Beta-sheets formed from multiple side-by-side beta-strands.
- Can be in parallel or antiparallel configuration
- Anti-parallel beta-sheets more stable
- Typically B-strands twist slightly in a right hand direction or twist clockwise



# Beta-Sheets

- □ Side chains point alternately above and below the plane of the beta-sheet
- □ A typical B-sheet contains 2 to 15 beta-strands/beta-sheet
- $\hfill\Box$  Each strand is made of  $\sim$  6 amino acids on average



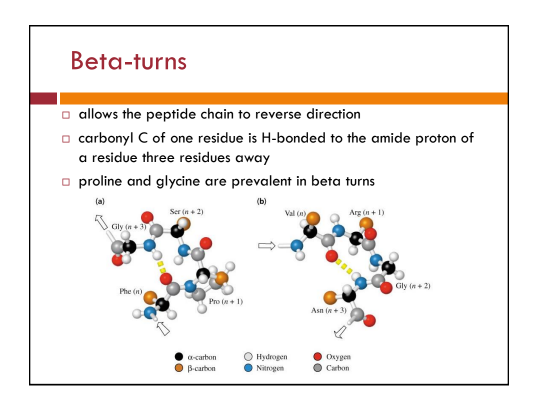
# Loops and turns

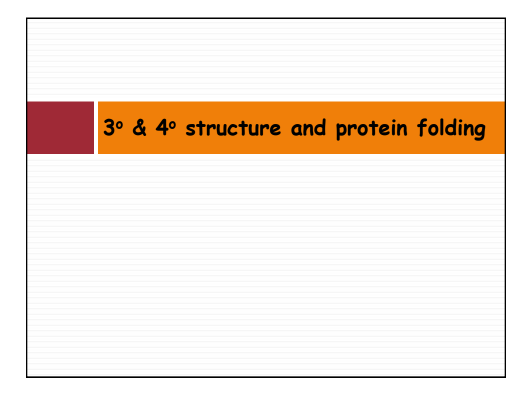
#### **Loops**

- □ Loops usually contain hydrophillic residues.
- □ Found on surfaces of proteins
- □ Connect alpha-helices and beta-sheets
- $\square \sim 10\%$  of residues are found in loops

#### **Turns**

- $\Box$  Loops with < 5 AA's are called turns
- □ Beta-turns are common

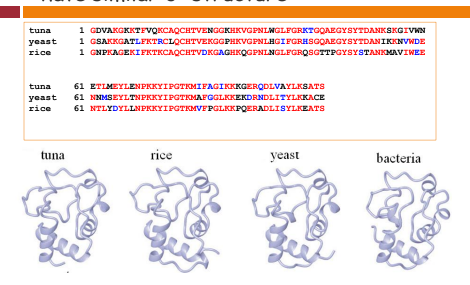




## 3° Structure

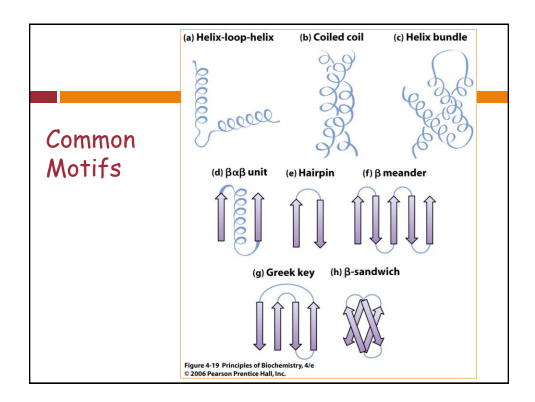
- □ Third level of protein organization
- □ Folding of polypeptide chain causes 2° structures to interact.
- AAs that are far apart in 1 ry structure are brought together, permitting interactions among their side chains.
- 3° structure is stabilized primarily by non-covalent interactions (mostly hydrophobic interactions) between side chains of an residues.
- □ Disulfide bridges are also elements of 3ry structure
- □ Formation of motifs and domains

# Proteins with similar 1° structure also have similar 3° structure



# Super-secondary structures

- □ Super-secondary structures or motifs: are recognizable combinations of  $\alpha$ -helices,  $\beta$ -strands and loops that appear in a different number of proteins.
- Sometimes, motifs are associated with a particular function, but similar motifs may have different functions.

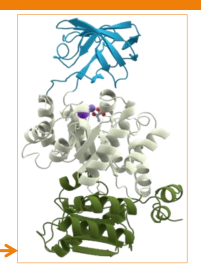


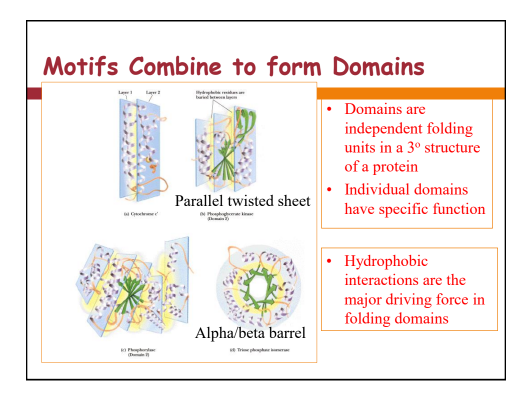
#### Motifs

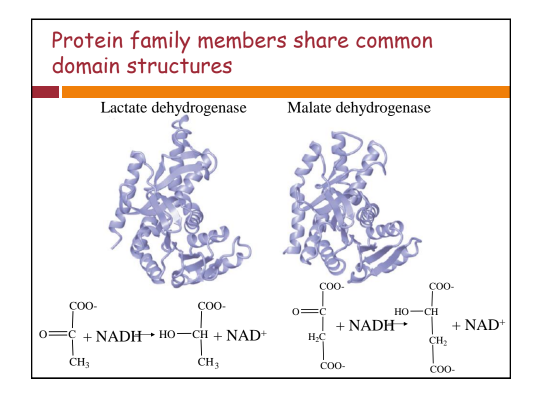
- Helix-loop-helix: occurs in calcium-binding proteins. Glu and Asn forms part of the loop and Ca+2 binding site.
   In some DNA binding proteins it is called, Helix-turnhelix because the loop forms a reverse turn
- □ Coiled –coil motif: two amphipathic a-helices that interact through hydrophobic edges, as in leucine zipper
- □ "βαβ":
- B-meander
- Hairpin
- Greek key
- B-sandwich

#### **Domains**

- Domains: are independently folded compact units
- Domains may consist of combinations of motifs
- Individual domains have specific function
- Size of domain vary from 25 to 30 aa residues to more than 300 aa residues
- Example: Pyruvate dehydrogenase has 3 domains

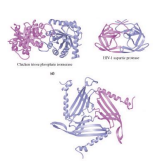




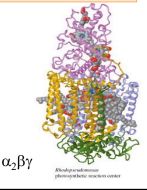


# 4º Structure

- •Quaternary structure describes the organization of subunits in a protein with multiple subunits (oligomeric protein)
- Can have homo-multimers or hetero-multimers
- •Subunits are held together by weak interactions, primarily by hydrophobic interactions







4º Structure

- Determine molecular weight of native protein by gel permeation chromatography
- Determine molecular weight of individual subunits by SDS-PAGE
- $\hfill\Box$  Can use the information to determine subunit composition

If.....

Native protein -160,000 daltons and  $\alpha$ -Subunit -50,000 daltons,  $\beta$ -Subunit -30,000 daltons

Then.....

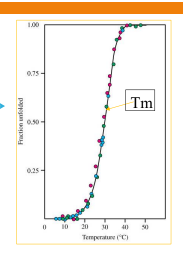
Protein can have  $\alpha_2\beta_2$  structure

# 4º Structure

- Subunits held together by non-covalent interactions
- Oligomeric protein is more stable than disassociated subunits
- Active site often made up of AA residues from different subunits
- 4° and 3° structure is often affected by ligand (substrate or inhibitor) binding. Important in enzyme regulation

#### Protein denaturation

- Denaturation disruption of native conformation
- Heat commonly used to denature proteins
- Tm = temperature where 50% folded/50% unfolded. You can calculate Tm from curve?
- Under physiological conditions, most proteins are stable up to 50-60°C
- □ Typical Tm = 40-60°C
- □ Tm depends on pH and ionic strength
- Tm for thermophiles >100°C (E.g., Taq DNA polymerase)
- Chemical denaturants Chaotrophic agents = Urea, guanidinium salts, KCN detergents = SDS

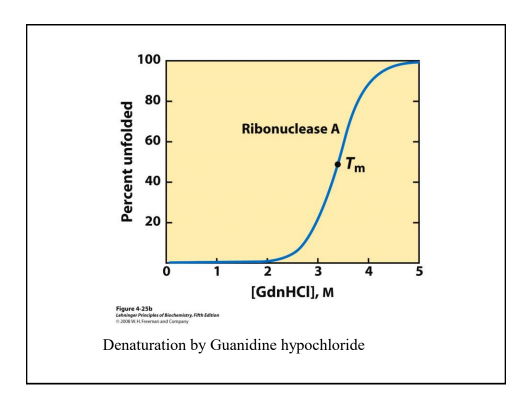


# Protein Folding

- □ Ribonuclease A (RNase A) will refold to native structure spontaneously (1 minute)
- $\Box > 10^{50}$  possible conformations
- $\Box$  If  $10^{-13}$  sec per conformation would take  $10^{30}$  years to sample enough to determine structure
- □ How do proteins fold so quickly?

# Protein Folding

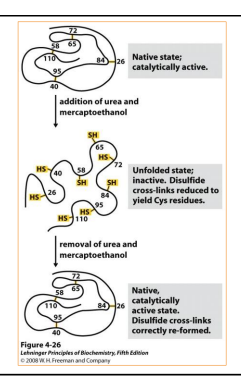
- Structures of globular proteins are not static
- □ Proteins "breathing" between different conformations
- □ Proteins fold towards lowest energy conformation
- Multiple paths to lowest energy form
- All folding paths funnel towards lowest energy form
- Local low energy minimum can slow progress towards lowest energy form



#### **Protein renaturation**

- Protein renaturation: certain globular proteins when denatured can regain their native structure and biological activity
- □ Example: Ribonuclease experiment (see next figure)

Renaturation of unfolded, denatured Ribonuclease

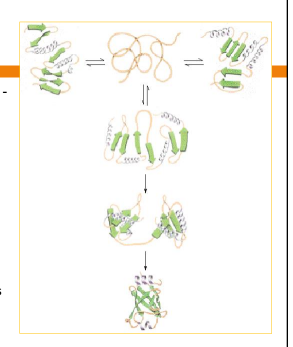


# Ribonuclease Denaturation/Renaturation experiment

- □ Conclusions from the Ribonuclease experiment
- → AA sequence contains all information required to fold the p.p chain into its native, 3D structure.
- > This is true for a minority of proteins, which is small and inherently stable.
- > Even though all proteins have the potential to fold into their native structure, many require some assistance

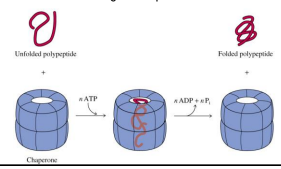
# Pathway of Protein Folding

- 1) Nucleation of folding -Rapid and reversible formation of local 2° structures form
- 2) Formation of domains (Molten Globular intermediates) through aggregation of local 2° structures
- 3) Domain conformations adjust to form native protein



# Chaperonins

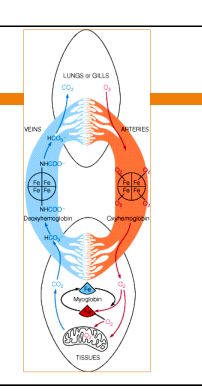
- Protein complexes that promote protein folding
- □ Chaperonins don't determine native structure
- □ Prevent misfolding and aggregation of protein
- Sequesters unfolded protein from other proteins
- Require ATP for protein binding, after ATP hydrolysis native protein released
- □ Thought to bind unfolded regions of protein



# Protein 3-D Structure/ Function

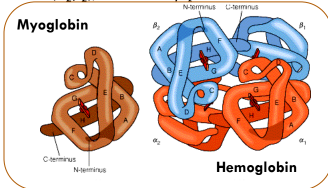
# Myoglobin/ Hemoglobin

- □ First protein structures determined
- Oxygen carriers
- Reversible binding of O2 is called Oxygenation
- $\square$  Hemoglobin transport  $O_2$  from lungs to tissues
- Myoglobin: O<sub>2</sub> storage protein, supply O2 to muscle tissues in reptiles, birds and mammals



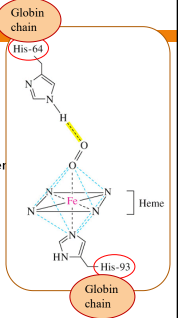
#### Mb and Hb subunits structurally similar

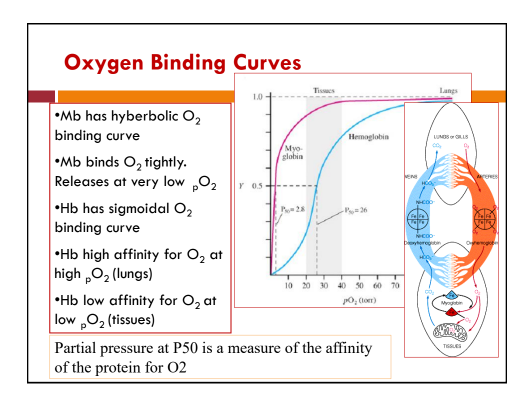
- •8 alpha-helices
- •Contain heme group
- •Mb monomeric protein
- •Hb heterotetramer ( $\alpha_2\beta_2$ ), dimer of  $\alpha\beta$  protomer

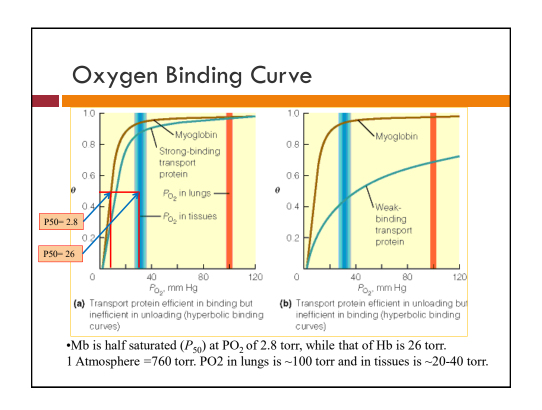


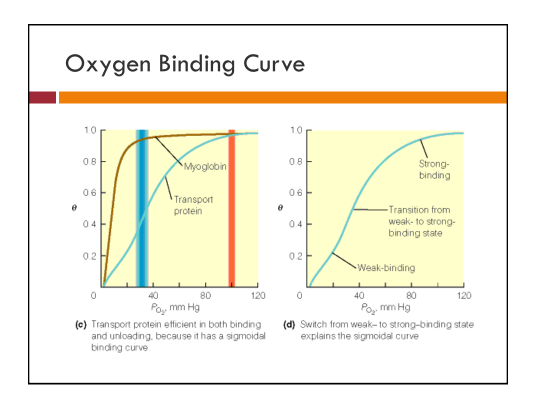
### Heme group

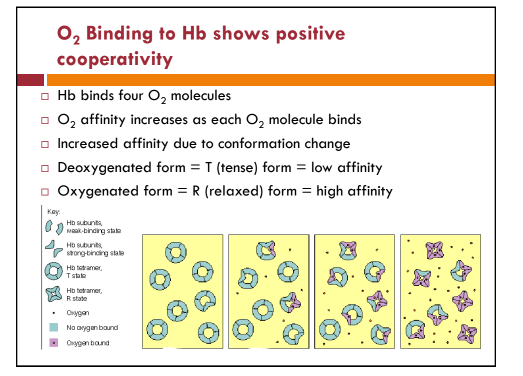
- Heme = Fe<sup>++</sup> bound to tertapyrrole ring (protoporphyrin IX complex)
- Heme non-covalently bound to globin proteins through His residue
- O<sub>2</sub> binds non-covalently to heme Fe<sup>++</sup>, stabilized through H-bonding with another His residue
- □ Heme group in hydrophobic crevice of globin protein

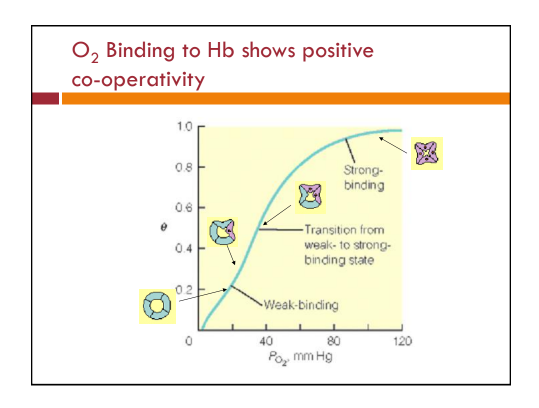


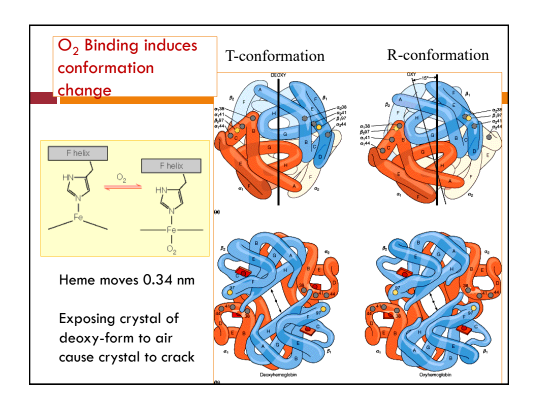










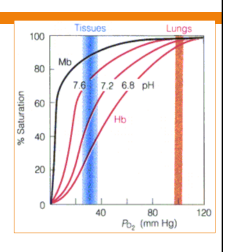


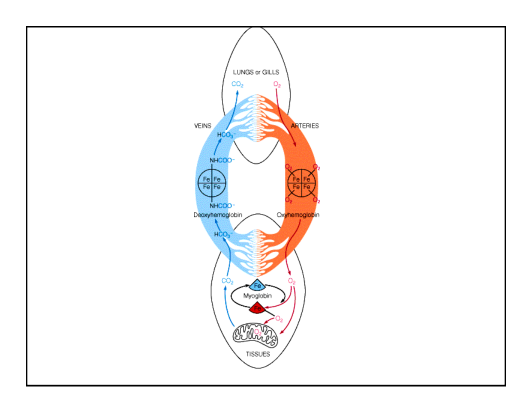
#### **Allosteric Interactions**

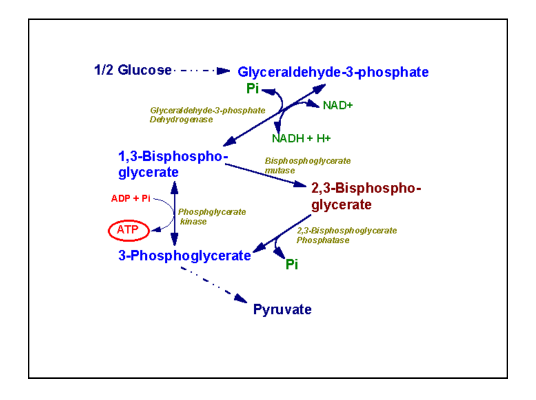
- Allosteric interaction occur when specific molecules bind a protein and modulates activity
- Allosteric modulators or allosteric effectors
- Bind reversibly to site separate from functional binding or active site
- Modulation of activity occurs through change in protein conformation
- 2,3 bisphosphoglycerate (BPG), CO<sub>2</sub> and protons are allosteric effectors of Hb binding of O<sub>2</sub>
- □ The binding of 2,3BPG to Hb raises its P50 to 26 torr, much higher than the P50 for O2 binding to purified Hb in aqueous solution (12 torr).

#### **Bohr Effect**

- □ Increased  $CO_2$  leads to decreased pH  $CO_2 + H_2O < -> HCO_3^- + H^+$
- At decreased pH several key AA's protonated, causes Hb to take on Tconformation (low affinity)
- In R-form same AA's deprotonated, form charge interactions with positive groups, stabilize R-conformation (High affinity)
- HCO<sub>3</sub>- combines with N-terminal alphaamino group to form carbamate group.
   --N<sub>3</sub>H<sup>+</sup> + HCO<sub>3</sub><sup>-</sup> ← → --NHCOO<sup>-</sup>
- Carbamation stabilizes T-conformation

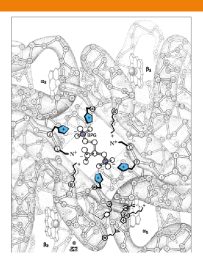






#### Bisphosphoglycerate (BPG)

- BPG involved in adaptation to high altitude
- Binding of BPG to Hb causes low O2 affinity
- BPG binds in the cavity between beta-Hb subunits
- Stabilizes T-conformation
- □ Fetal Hb (HbF,  $\alpha_2\gamma_2$ ) has low affinity for BPG, allows fetus to compete for  $O_2$  with mother's Hb (HbA,  $\alpha_2\beta_2$ ) in placenta.



#### Fetal Hb (HbF) versus adult Hb (HbA)

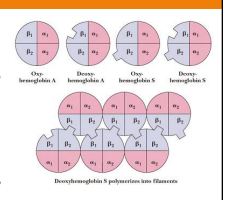
- HbF has higher affinity for O2 than HbA
- □ It lacks two positively charged amino acids that take part in binding 23BPG, i.e., His-143 of each B-globin chain is replaced by Serine. Thus 2,3BPG binds less tightly to HbF than HbA
- □ The P50 for HbF is 18 torr compared to 26 torr for HbA. Thus at PO2 of 20-40 torr in tissues HbF has higher affinity for O2. The difference in affinity allows efficient transfer of O2 from maternal blood to the fetus.

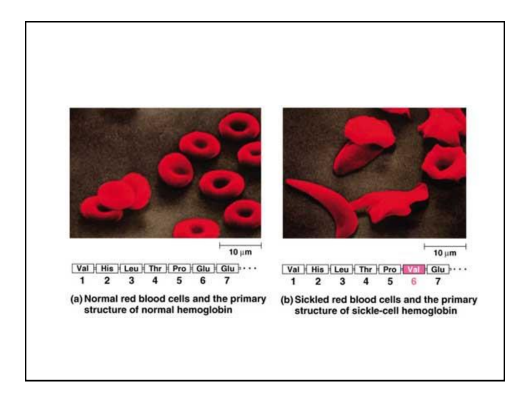
# Class activity!

☐ If HbF and HbA have the same affinity toward oxygen, how this will affect the growth of fetus in utero?

# Mutations in $\alpha$ - or $\beta$ -globin genes can cause disease state

- □ Sickle cell anemia E6 to V6
- Causes V6 to bind to hydrophobic pocket in deoxy-Hb
- Polymerizes to form long filaments
- Cause sickling of cells
- Sickle cell trait offers advantage against malaria
- Fragile sickle cells can not support malaria parasite

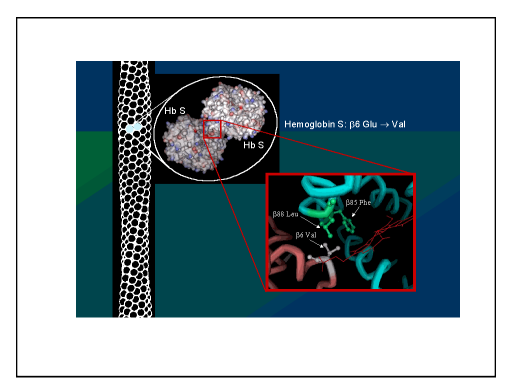




# Sickle cell (HbS) vs. HbC diseases

□ HbS: GAG>GTG, Glu>Val;

□ **HbC**: GAG>AAG, Glu>Lys



# Class activity!

- □ Can you predict the phenotype or severity of phenotype from the amino acid change?
- How sickle cell disease gives an advantage for patients against Malaria?

