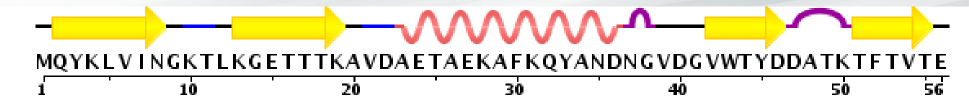
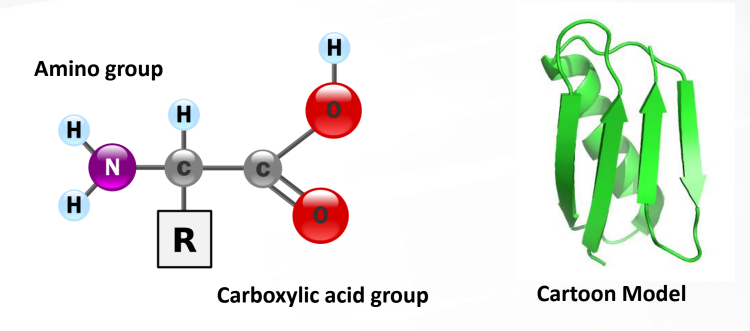
Protein Structure

Biochemistry Boot Camp 2022
Session #1
Matt Thompson
mthompson10@ua.edu

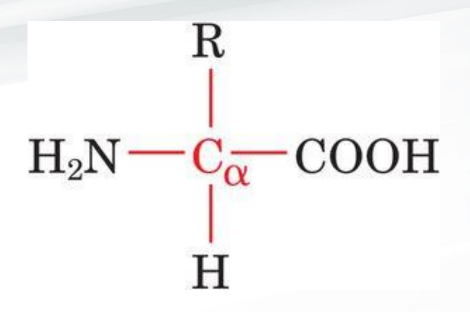
What is a Protein?



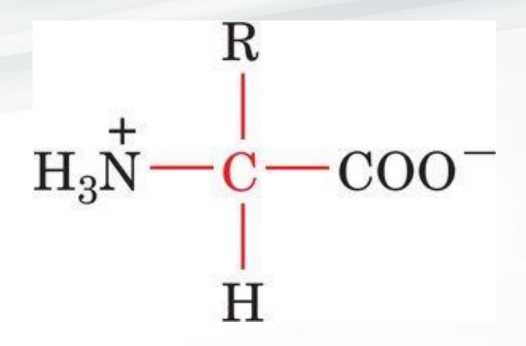


- Answer: Polymer of amino acids connected by peptide bonds
 - Multiple representations, including sequence, chemical structure, simplifying cartoons
- PyMOL Download: Staphylococcal Nuclease (sticks and cartoon)

General Structure of α-Amino Acid



Dipolar Amino Acid: Zwitterion





Structures of Standard Amino Acids

TABLE 4-1 Covalent Structures and Abbreviations of the "Standard" Amino Acids of Proteins, Their Occurrence, and the pK Values of Their Ionizable Groups

Name, Three-letter Syn and One-letter S		Structural Formula ^a	Residue Mass (D) ^b	Average Occurrence in Proteins (%) ^c	pK_1 α -COOH ^d	pK_2 α -NH ₃ ^{+d}	pK _R Side Chain ^d
Amino acids with	nonpolar s	ide chains					
Glycine Gly G	COO- H-C-H NH3		57.0	7.1	2.35	9.78	
Alanine Ala A	COO- H-C-CI NH ⁺ ₃		71.1	8.2	2.35	9.87	
Valine Val V	COO- H-C-CI NH3	,СН ₃ Н `СН ₃	99.1	6.9	2.29	9.74	
Leucine Leu L	COO- H-C-CI NH3	CH ₃ CH ₃	113.2	9.7	2.33	9.74	
Isoleucine Ile I	COO- H-C- I NH3	CH ₃ - C - CH ₂ - CH ₃ H	113.2	6.0	2.32	9.76	
Methionine Met M	COO- H-C-CH NH3	H ₂ -CH ₂ -5-CH ₃	131.2	2.4	2.13	9.28	
Proline Pro P	COO-C C C C C C C C C C C C C C	CH ₂ CH ₂	97.1	4.7	1.95	10.64	
Phenylalanine Phe F	COO- H-C-CI NH3		147.2	3.9	2.20	9.31	
Tryptophan Trp W	COO- H-C-CI NH3		186.2	1.1	2.46	9.41	

[&]quot;The ionic forms shown are those predominating at pH 7.0 (except for that of histidine') although residue mass is given for the neutral compound. The C_{α} atoms, as well as atoms marked with an asterisk, are chiral centers with configurations as indicated according to Fischer projection formulas (Section 4-2). The standard organic numbering system is provided for heterocycles.

^bThe residue masses are given for the neutral residues. For the molecular masses of the parent amino acids, add 18.0 D, the molecular mass of H₂O, to the residue masses. For side chain masses, subtract 56.0 D, the formula mass of a peptide group, from the residue masses.

^cThe average amino acid composition in the complete SWISS-PROT database (http://www.expasy.ch/sprot/relnotes/relstat.html), Release 2013_13. Individual proteins may exhibit large deviations from these quantities.

^dData from Dawson, R.M.C, Elliott, D.C., Elliott, W.H., and Jones, K.M., *Data for Biochemical Research* (3rd ed.), pp. 1–31, Oxford Science Publications (1986).

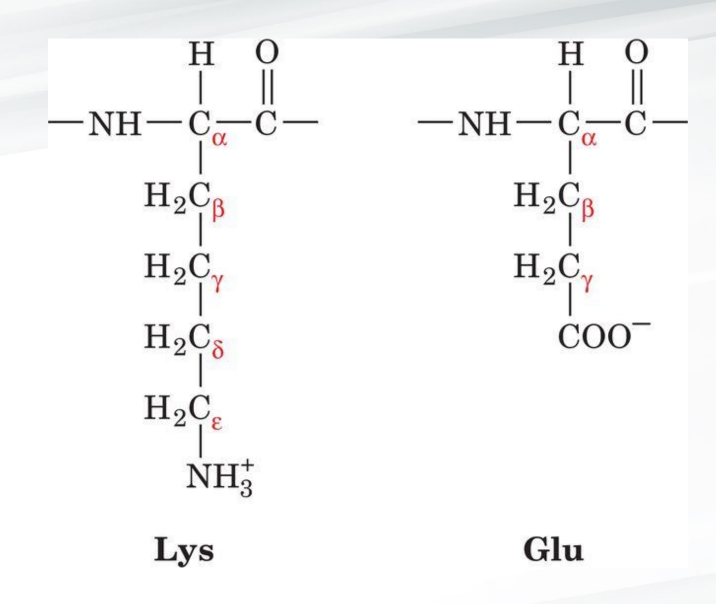
Structures of Standard Amino Acids

Name, Three-letter Syr and One-letter		Structural Formula ^a	Residue Mass (D) ^b	Average Occurrence in Proteins (%) ^c	pK_1 α -COOH ^d	$pK_2 \\ \alpha-NH_3^{+d}$	pK _R Side Chain ^d
Amino acids wit	h uncharged p	oolar side chains					
Serine Ser S	COO- H-C-CH NH3	-ОН	87.1	6.6	2.19	9.21	
Threonine Thr T	H-C	C*-CH ₃	101.1	5.3	2.09	9.10	
Asparagine ^e Asn N	COO- H-C-CH; NH3	2-C NH ₂	114.1	4.1	2.14	8.72	
Glutamine ^e Gln Q	COO- H-C-CH NH3	2-CH ₂ -C NH ₂	128.1	3.9	2.17	9.13	
Tyrosine Tyr Y	COO- H-C-CH I NH‡	2-ОН	163.2	2.9	2.20	9.21	10.46 (phenol)
Cysteine Cys C	COO- H-C-CH NH3	₂ —SH	103.1	1.4	1.92	10.70	8.37 (sulfhydryl)
Amino acids wit	h charged pole	ar side chains					
Lysine Lys K	COO- H-C-CH I NH [‡]	₂ -CH ₂ -CH ₂ -CH ₂ -NH ₃	128.2	5.9	2.16	9.06	10.54 (ε-NH ₃ ⁺)
Arginine Arg R	COO- H-C-CH NH3	2-CH ₂ -CH ₂ -NH-C	156.2	5.5	1.82	8.99	12.48 (guanidino)
Histidine ^f His H	COO- H-C-CH NH3	2—————————————————————————————————————	137.1	2.3	1.80	9.33	6.04 (imidazole)
Aspartic acid ^e Asp D	COO- H-C-CH, NH3	g-c"	115.1	5.4	1.99	9.90	3.90 (β-COOH)
Glutamic acid ^e Glu E	COO- H-C-CH NH‡	-CH ₂ -с	129.1	6.8	2.10	9.47	4.07 (γ-COOH)

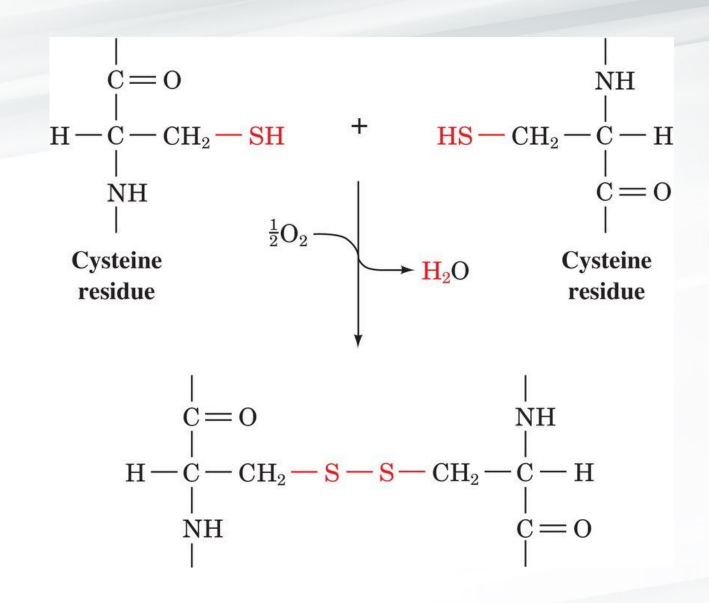
[&]quot;The three- and one-letter symbols for asparagine or aspartic acid are Asx and B, whereas for glutamine or glutamic acid they are Glx and Z. The one-letter symbol for an undetermined or "nonstandard" amino acid is X.

 $[^]f$ Both neutral and protonated forms of histidine are present at pH 7.0, since its p K_R is close to 7.0.

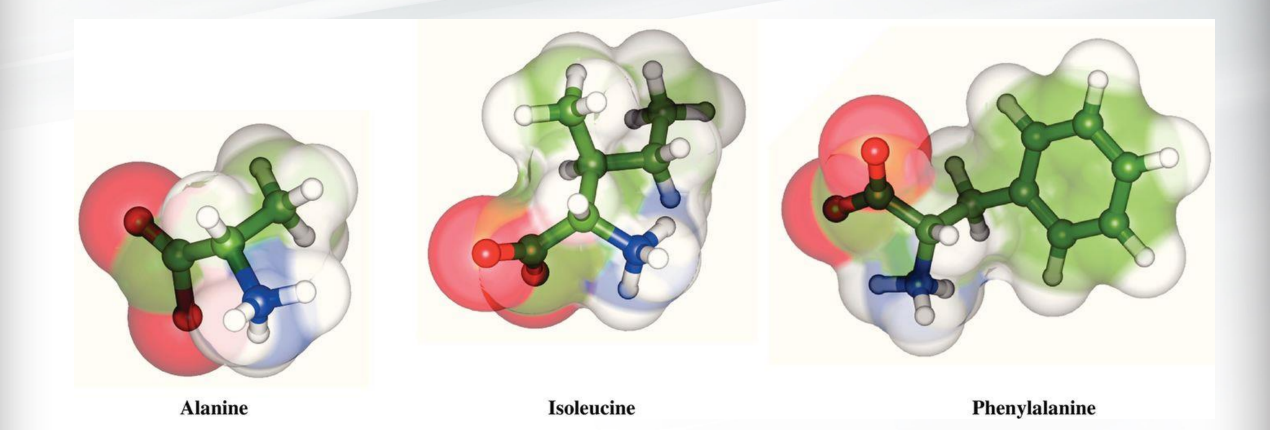
Amino Acid Nomenclature



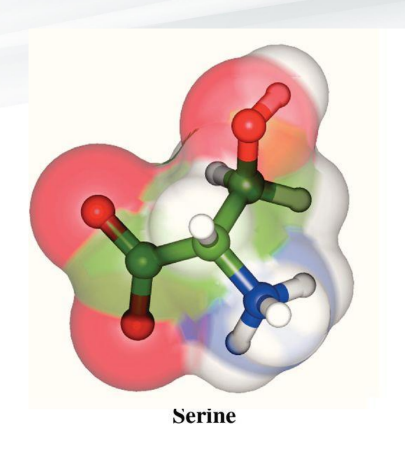
Cysteine Can Form Disulfide Bonds

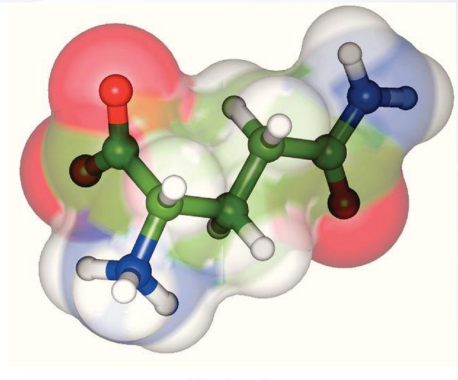


Nonpolar Side Chains



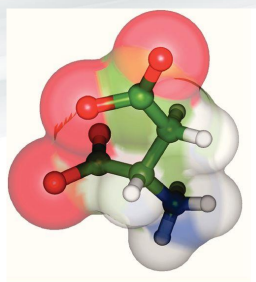
Uncharged Polar Side Chains

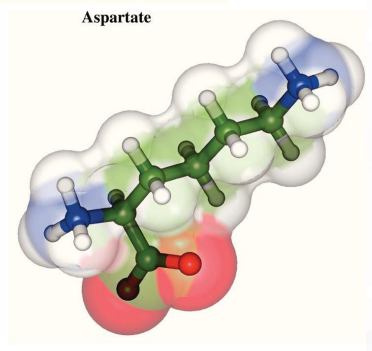




Glutamine

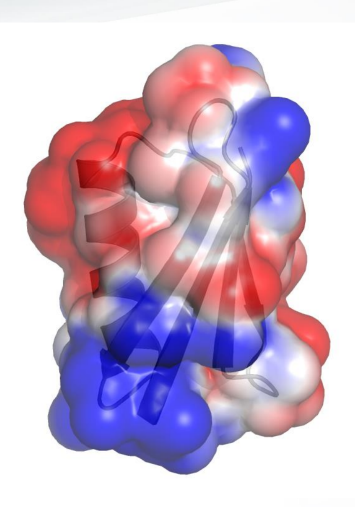
Charged Polar Side Chains



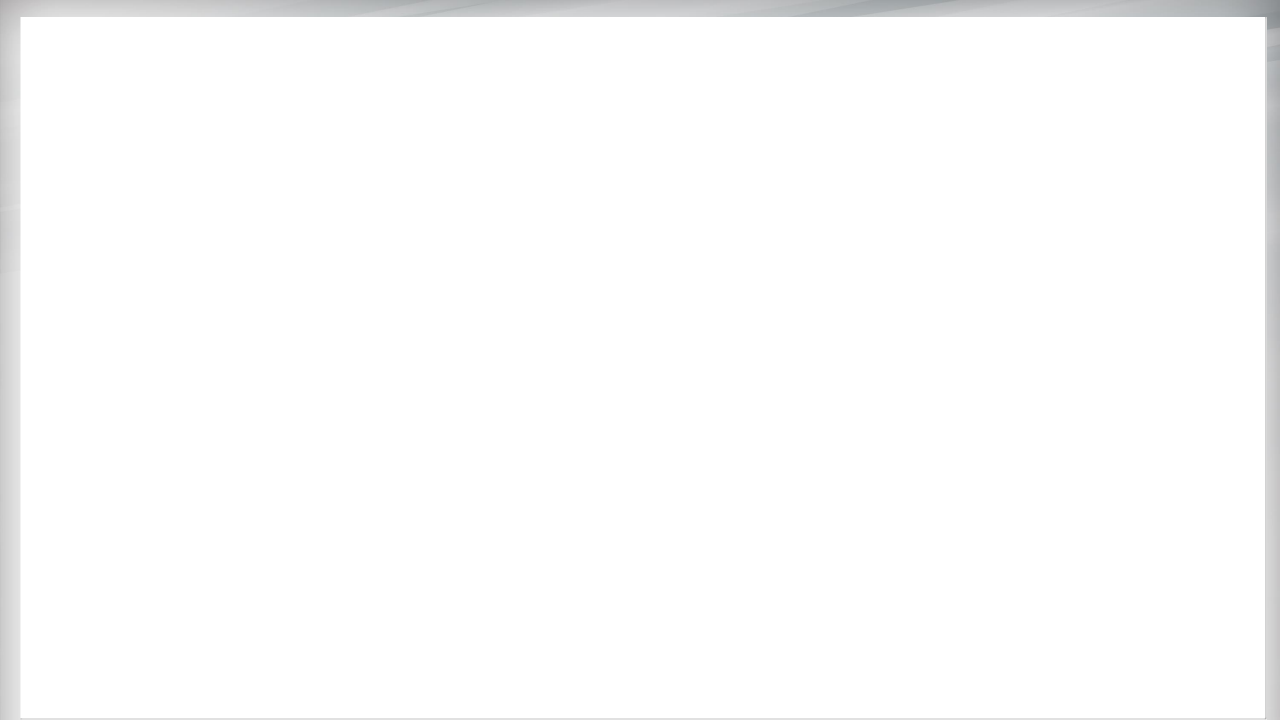


Lysine

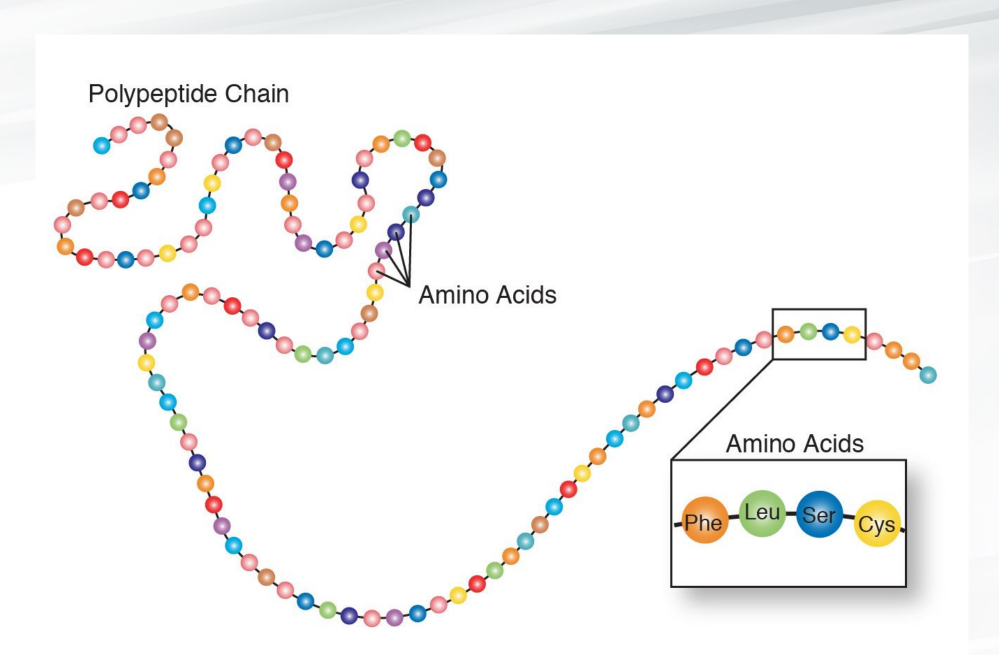
Electrostatic surface



Red – negative charge Blue – positive charge



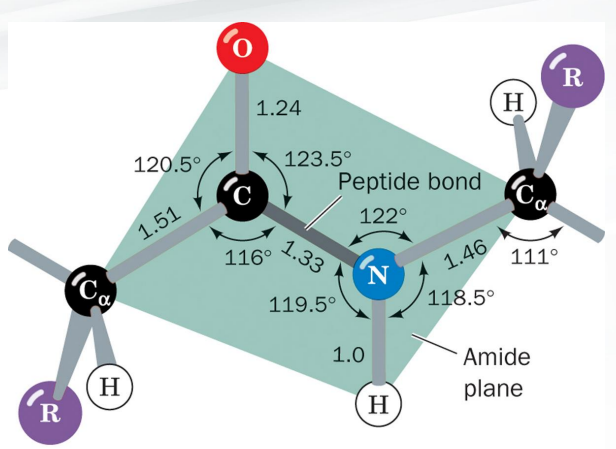
Proteins Are Composed of a "Chain" of Amino Acids



Condensation of Two Amino Acids



Peptide Bonds Assume Trans Conformation



After Marsh, R.E. and Donohue, J., Adv. Protein Chem. 22, 249 (1967).

Resonance of Peptide Bond

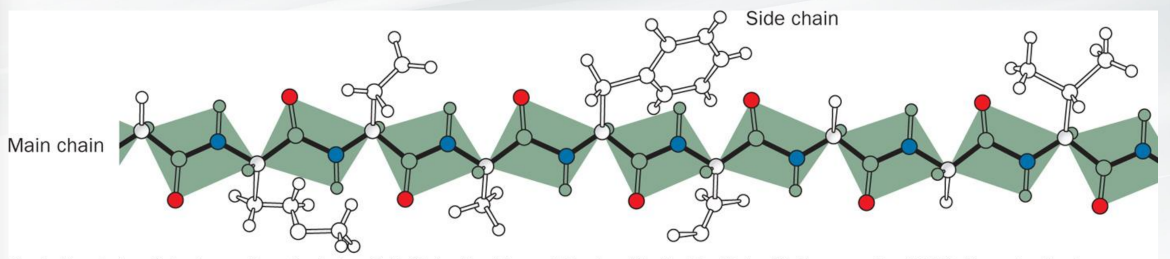
$$\begin{array}{c}
O \\
\parallel \\
C \\
N
\end{array}$$

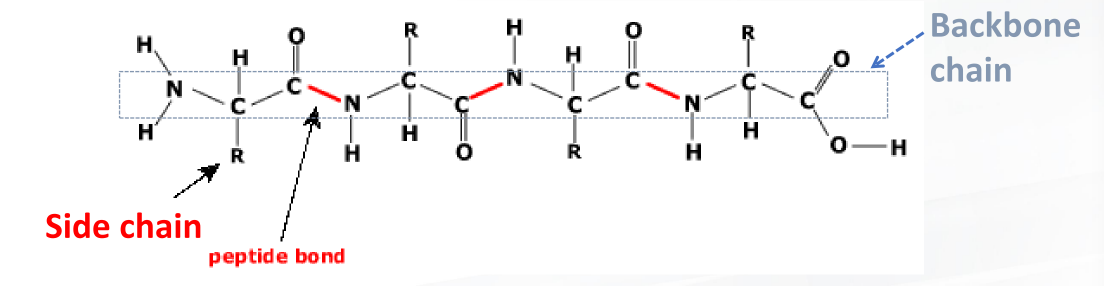
$$\begin{array}{c}
O^{-} \\
\downarrow \\
N
\end{array}$$

$$\begin{array}{c}
C \\
\downarrow \\
H
\end{array}$$

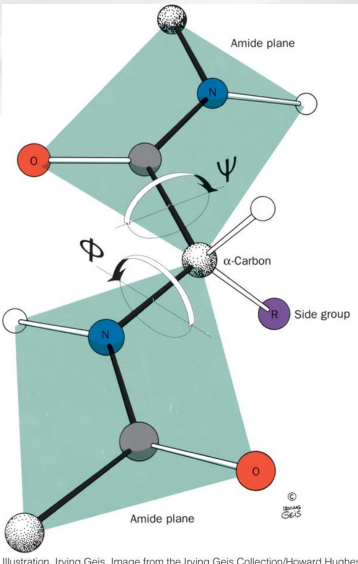
$$\begin{array}{c}
H
\end{array}$$

Extended Conformation of PolypeptideThe Protein "Backbone"



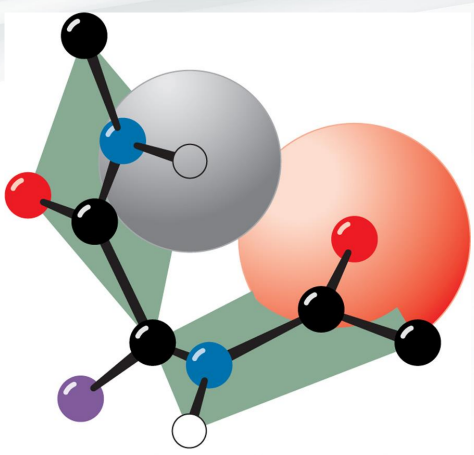


Torsion Angles of Polypeptide Backbone

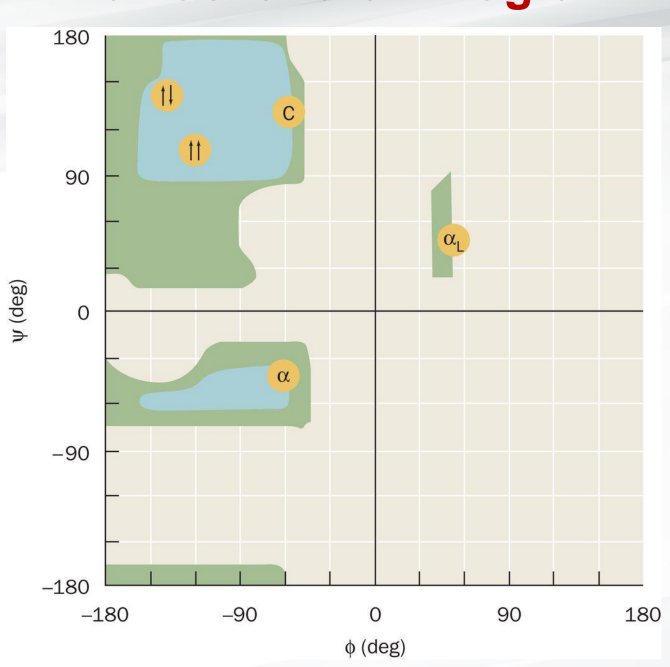


Illustration, Irving Geis. Image from the Irving Geis Collection/Howard Hughes Medical Institute. Rights owned by HHMI. Reproduction by permission only.

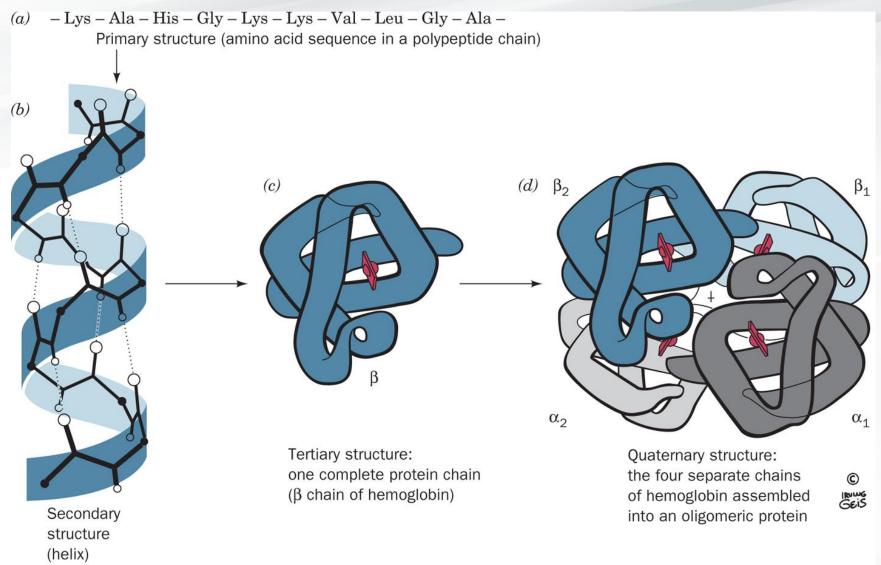
Steric Interference of Adjacent Peptide Groups



Ramachandran Diagram



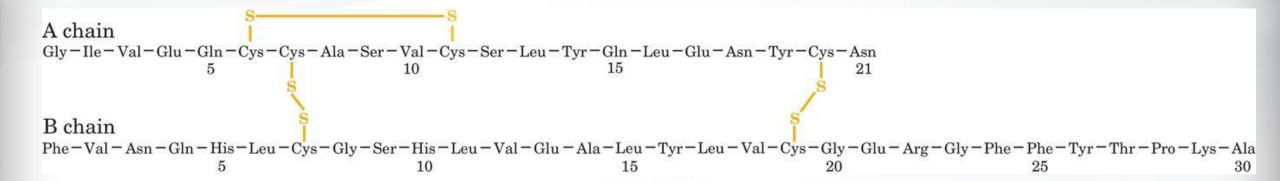
Levels of Protein Structure



1° Structure

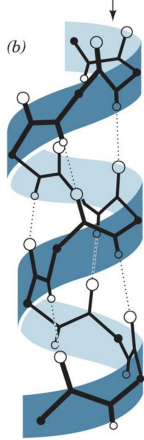
(a) - Lys - Ala - His - Gly - Lys - Lys - Val - Leu - Gly - Ala - Primary structure (amino acid sequence in a polypeptide chain)

Bovine Insulin: 1° Structure



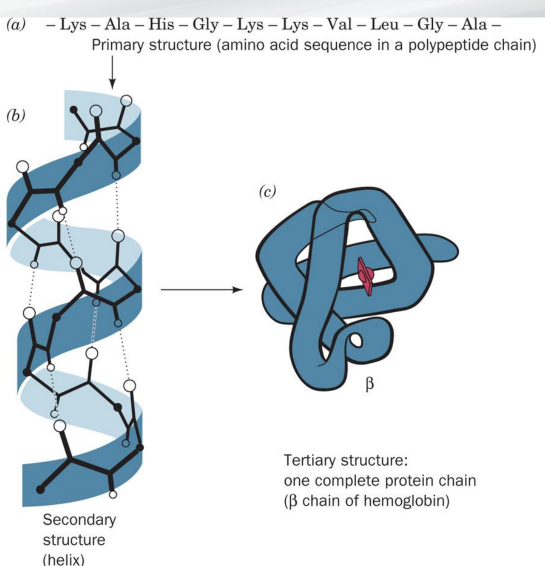
1° & 2° Structure

 $(a) \quad - Lys - Ala - His - Gly - Lys - Lys - Val - Leu - Gly - Ala - \\ \quad \quad \text{Primary structure (amino acid sequence in a polypeptide chain)}$

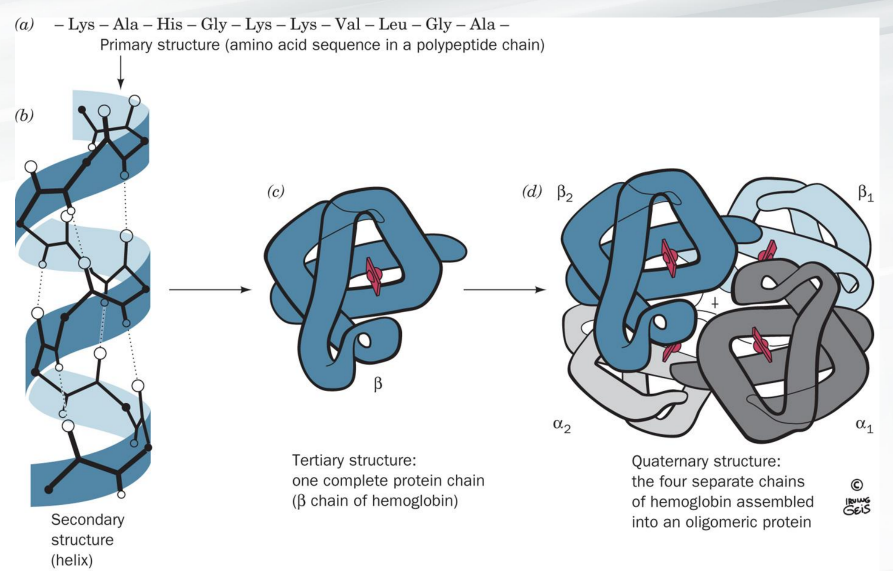


Secondary structure (helix)

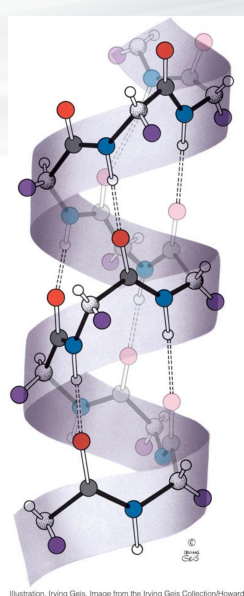
1°, 2°, & 3° Structure



1°, 2°, 3° & 4° Structure

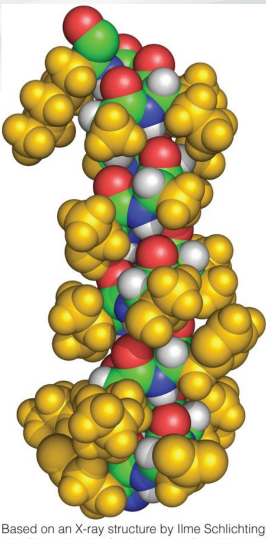


The α Helix



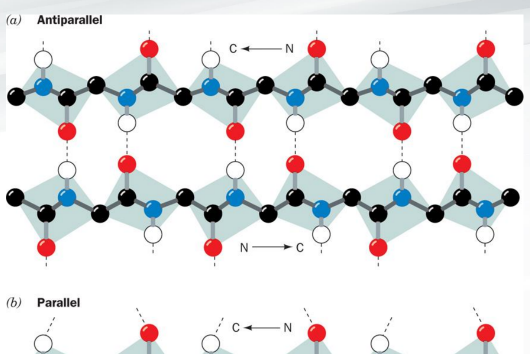
Illustration, Irving Geis. Image from the Irving Geis Collection/Howard Hughes Medical Institute. Rights owned by HHMI. Reproduction by permission only.

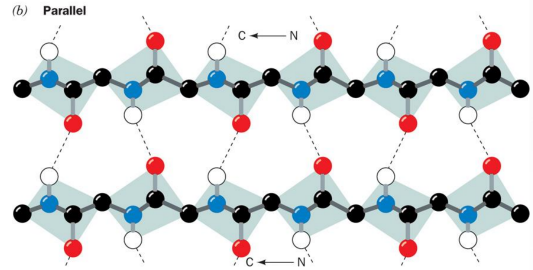
The α Helix: Space Filling Model



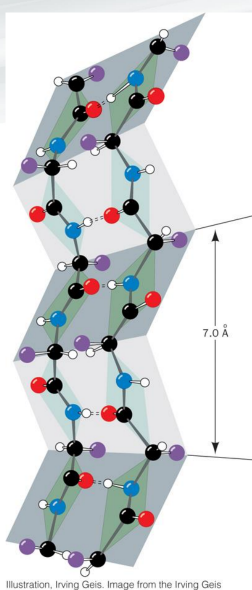
Based on an X-ray structure by Ilme Schlichting, Max Planck Institut für Molekulare Physiologie, Dortmund, Germany. PDBid 1A6M (for the definition of "PDBid" see Section 6-2E).

β Sheets

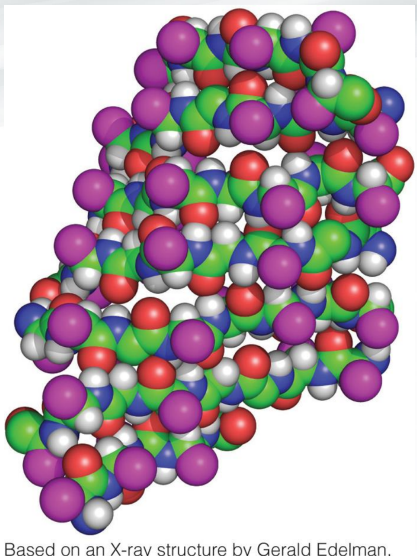




Pleating of β Sheet

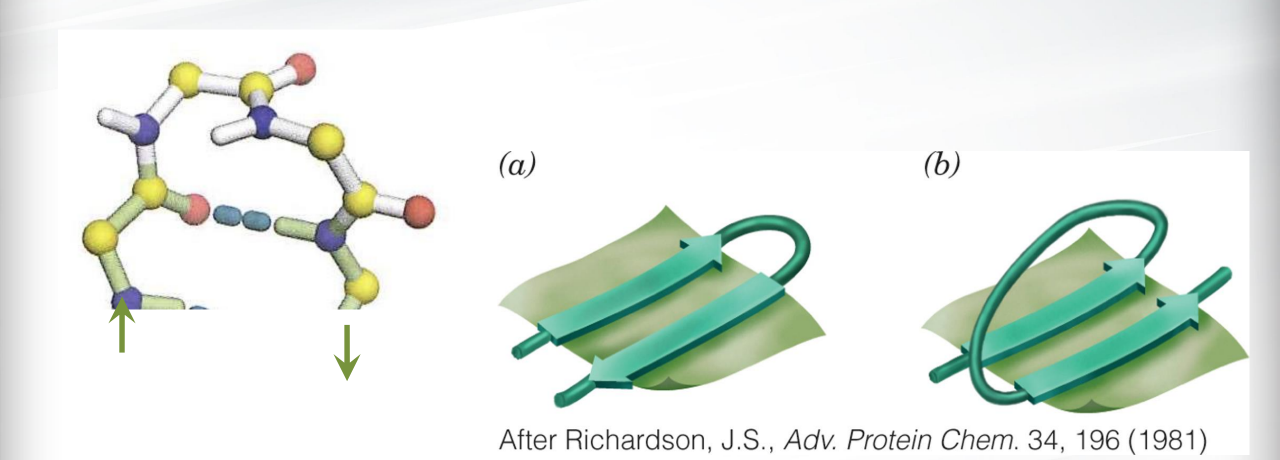


β Sheet: Space Filling Model

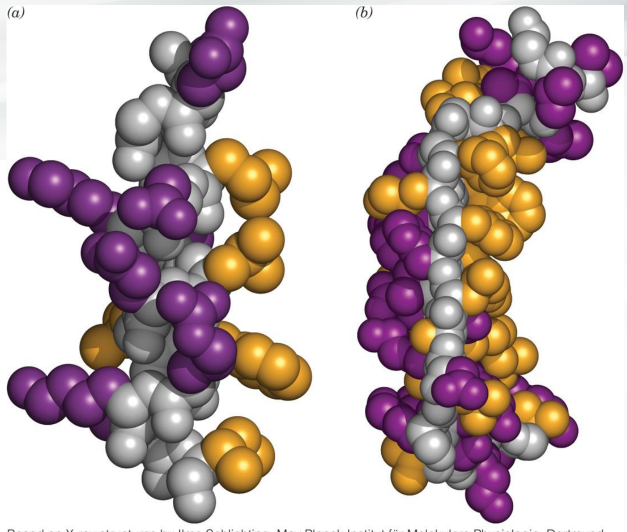


Based on an X-ray structure by Gerald Edelman, The Rockefeller University. PDBid 2CNA.

Connecting Adjacent & StrandsBeta Turns Reverse the Direction of the Backbone



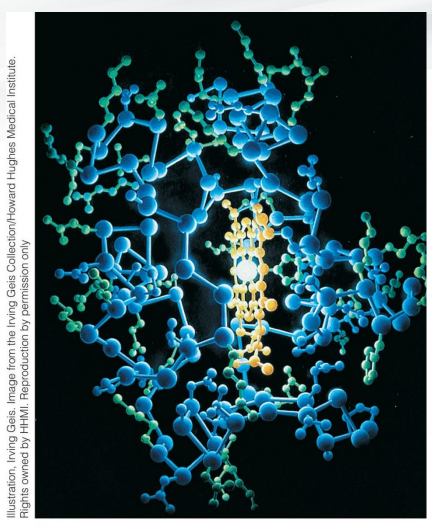
Side Chain Location

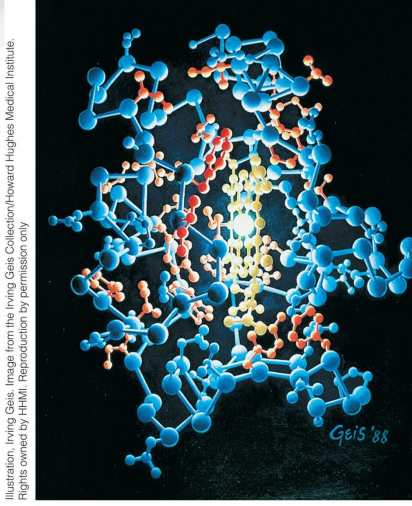


Based on X-ray structures by Ilme Schlichting, Max Planck Institut für Molekulare Physiologie, Dortmund, Germany, and Gerald Edelman, The Rockefeller University. PDBids 1A6M and 2CNA.

Oxy-Myoglobin and Concanavalin A PDBids 1A6M and 2CNA

Side Chain Distribution in Cytochrome

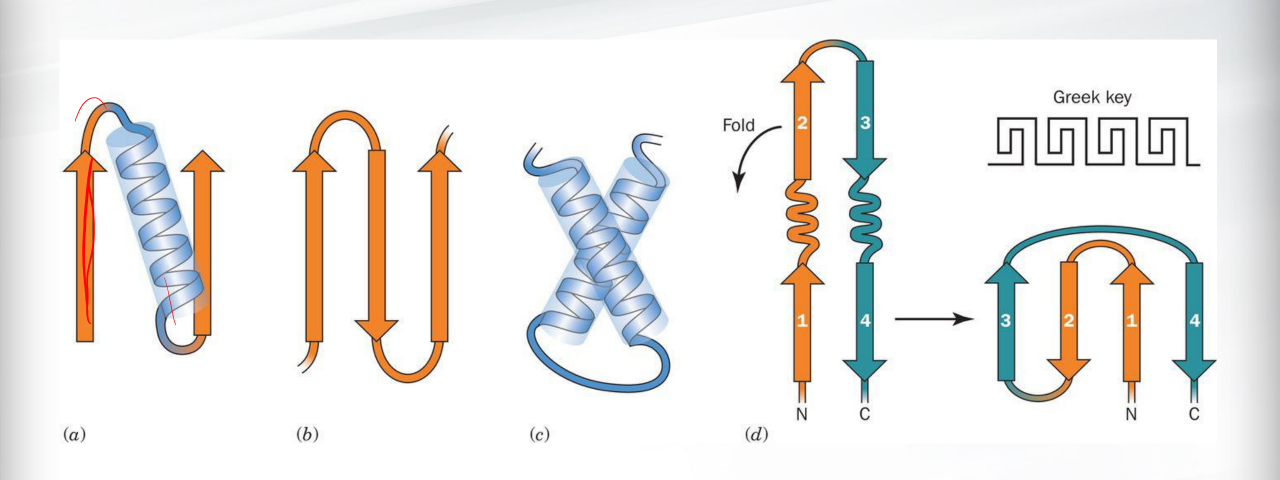




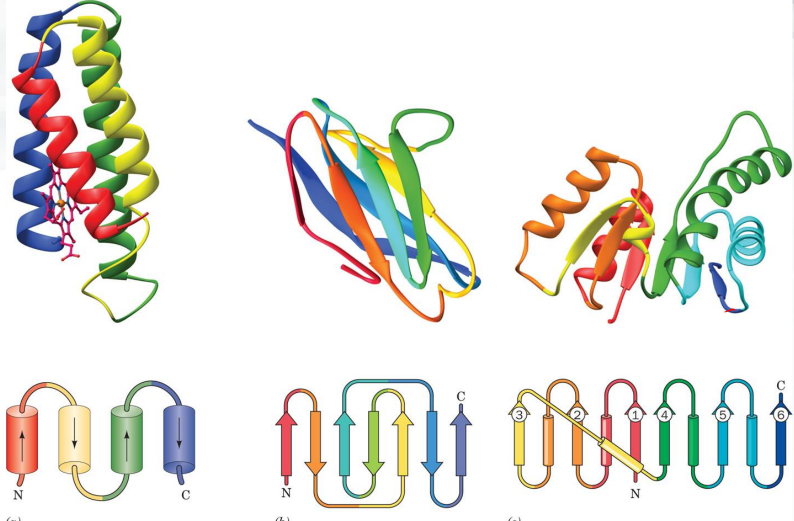
(a)

(b)

Motifs: Supersecondary Structures



Protein Classification: α , β , or α/β



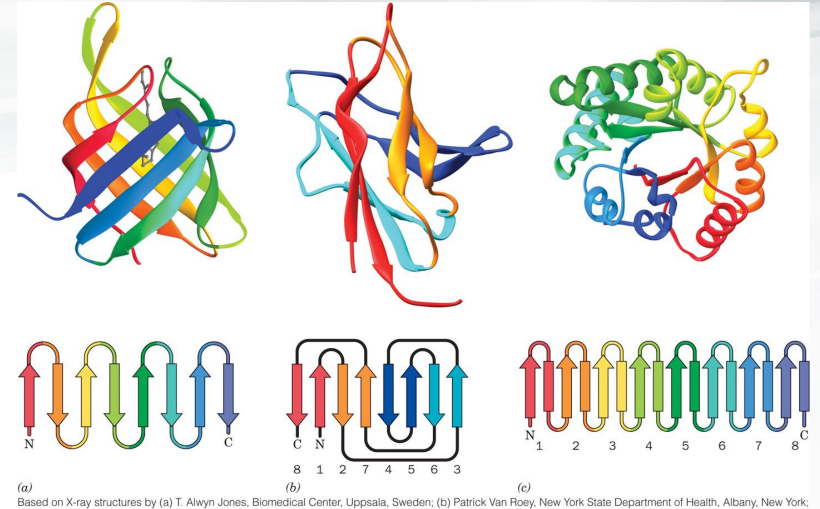
(a) (b) (c) Based on X-ray structures by (a) F. Scott Matthews, Washington University School of Medicine; (b) Roberto Poljak, The Johns Hopkins School of Medicine; and (c) Michael Rossmann, Purdue University. PDBids (a) 256B, (b) 7FAB, and (c) 6LDH.

Cytochrome *b*562 PDBid <u>256B</u>

Human immunoglobulin fragment PDBid <u>7FAB</u>

Dogfish lactate dehydrogenase PDBid <u>6LDH</u>

Protein Topology: 8-Stranded β Barrels



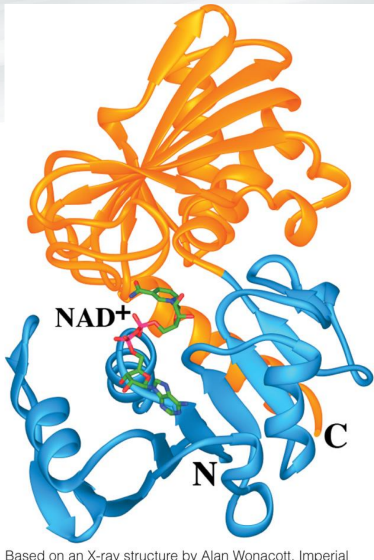
PDBid 1PNG

Human retinol binding protein PDBid 1RBP

Triose phosphate isomerase PDBid 1TIM

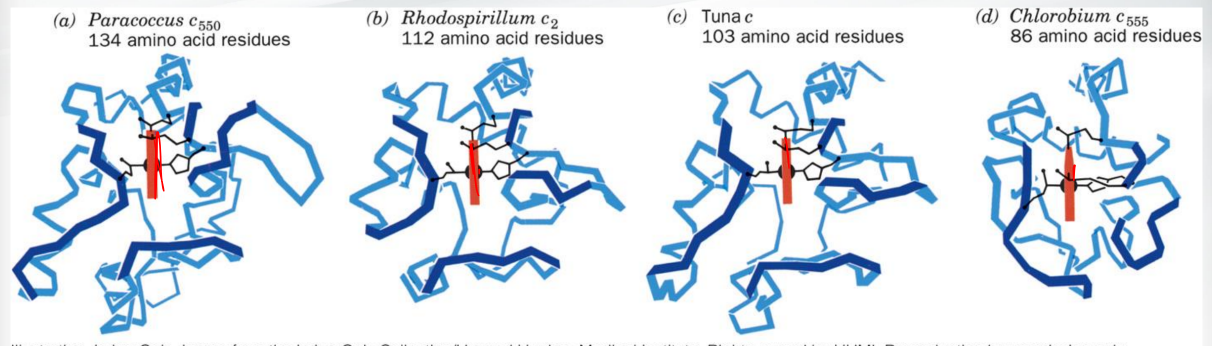


2-Domain Protein: GAPDH

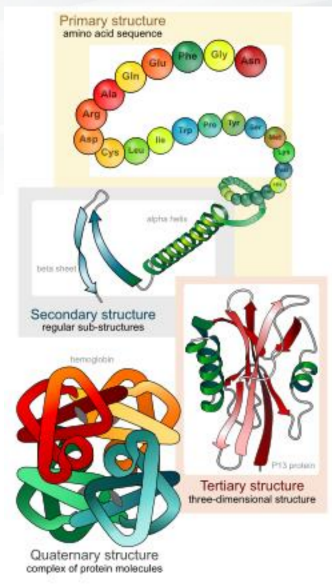


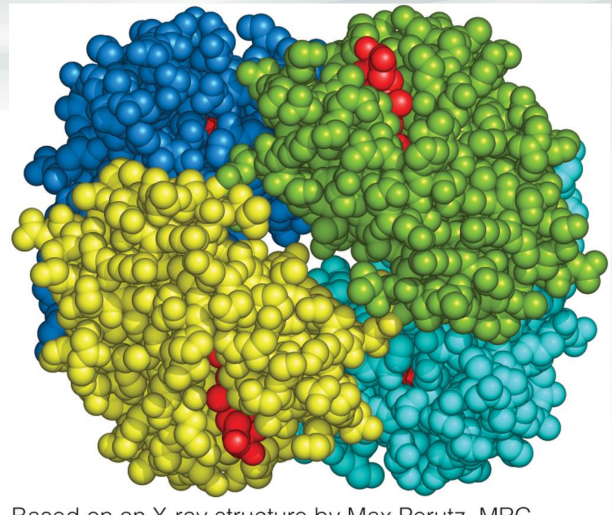
Based on an X-ray structure by Alan Wonacott, Imperial College, London, U.K. PDBid 1GD1.

Structure Conserved More Than Sequence



4º Structure of Hemoglobin

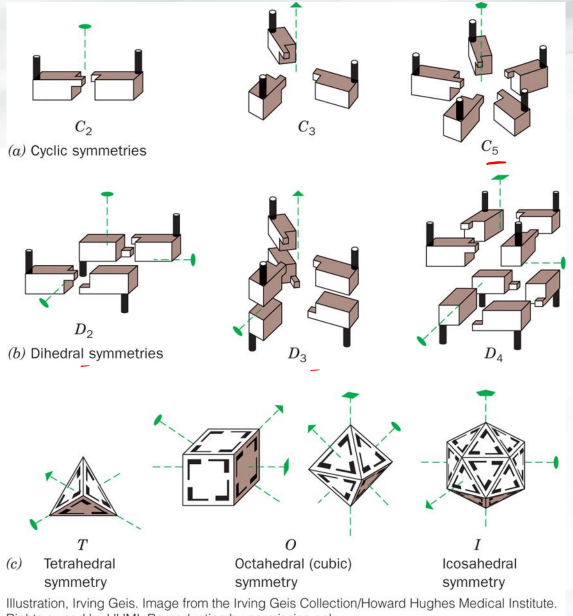




Based on an X-ray structure by Max Perutz, MRC Laboratory of Molecular Biology, Cambridge, U.K. PDBid 2DHB.

Deoxyhemoglobin PDBid 2DHB

Symmetries of Oligomeric Proteins



Rights owned by HHMI. Reproduction by permission only.

Summary

• Proteins are chemical chains made up of 20 different amino acids

 The chemical structure of the backbone has consequences: helices, sheets, and turns

Protein structure is classified into a four-level hierarchy