

Principles of Protein Structure and Conformational Space

Protein structure

1. Introduction

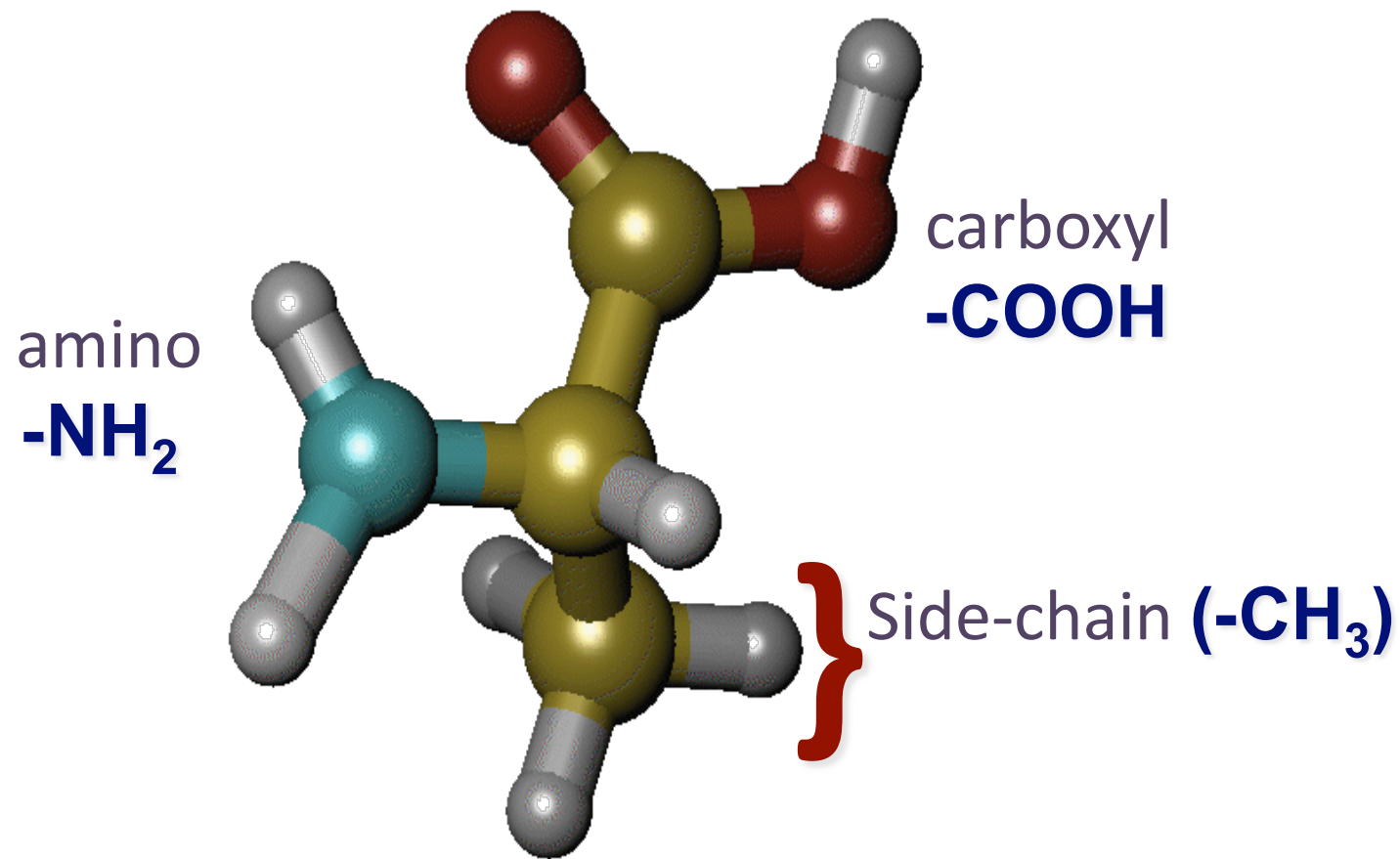
1. Amino acids
2. Peptide bond
3. Native conformation
4. Levels of structure (1,2,3)

2. Secondary Structure

1. Strands
2. Helix
3. Ramachandran Plot
4. Aa propensities

1. Introduction

1. Amino Acids

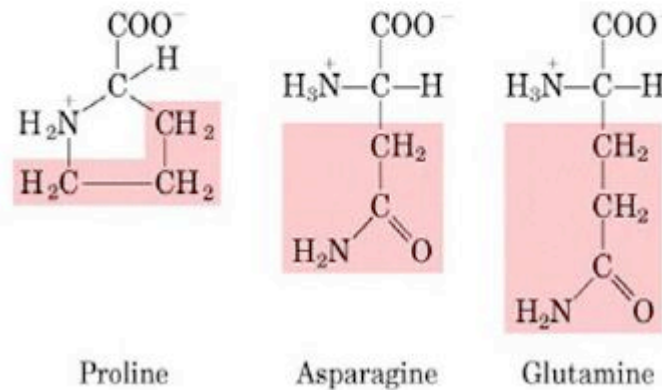
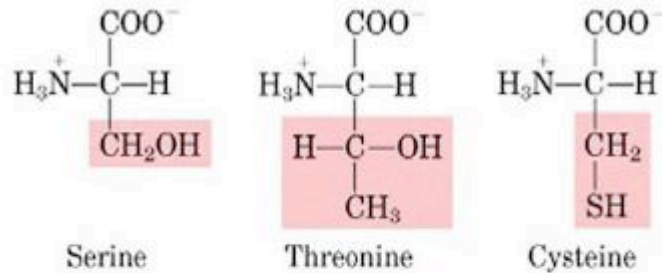


Estereoisomer L

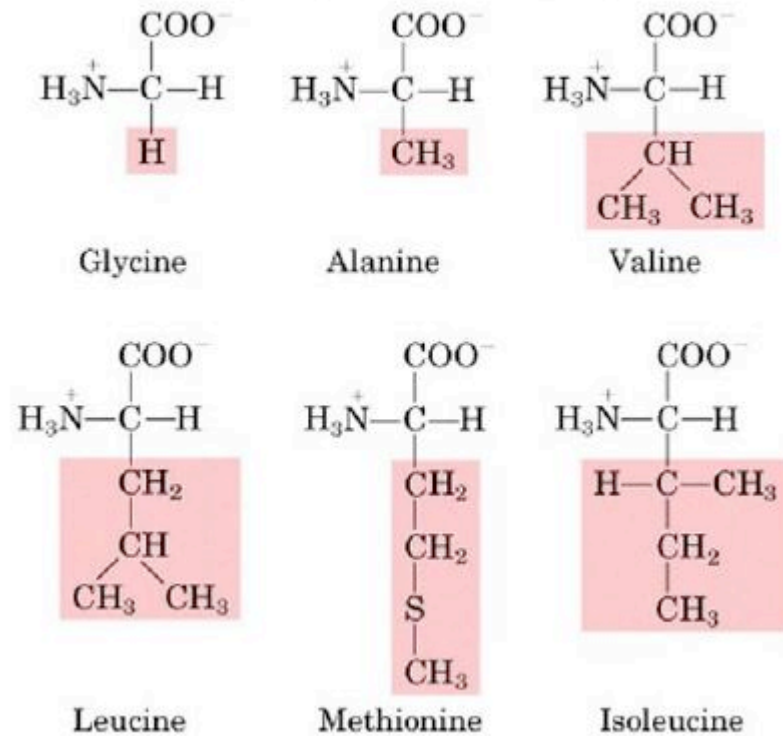
1. Introduction

1. Amino Acids

Polar, uncharged R groups



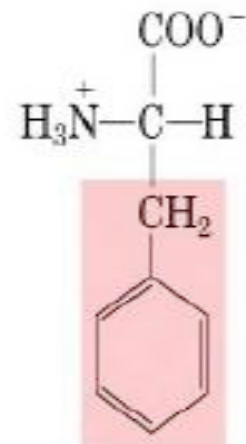
Nonpolar, aliphatic R groups



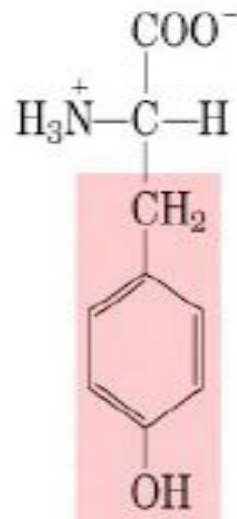
1. Introduction

1. Amino Acids

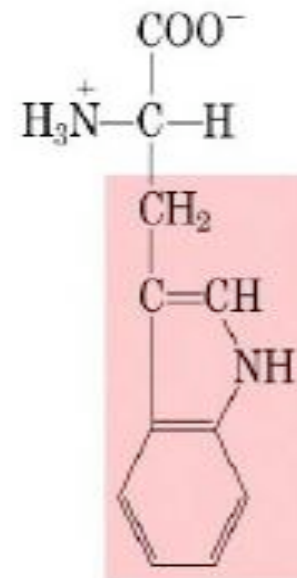
Aromatic R groups



Phenylalanine



Tyrosine

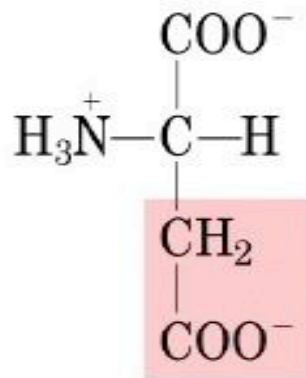


Tryptophan

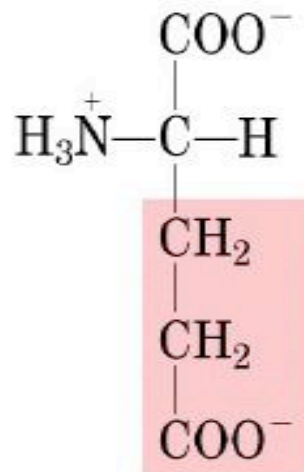
1. Introduction

1. Amino Acids

Negatively charged R groups

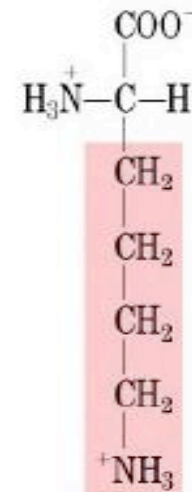


Aspartate

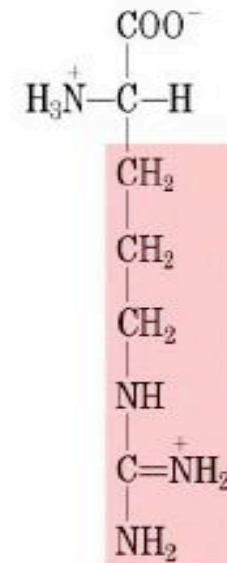


Glutamate

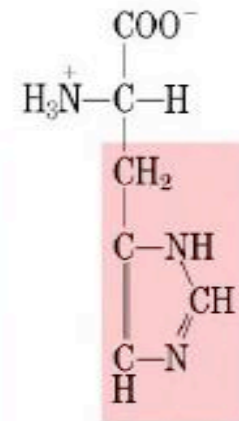
Positively charged R groups



Lysine



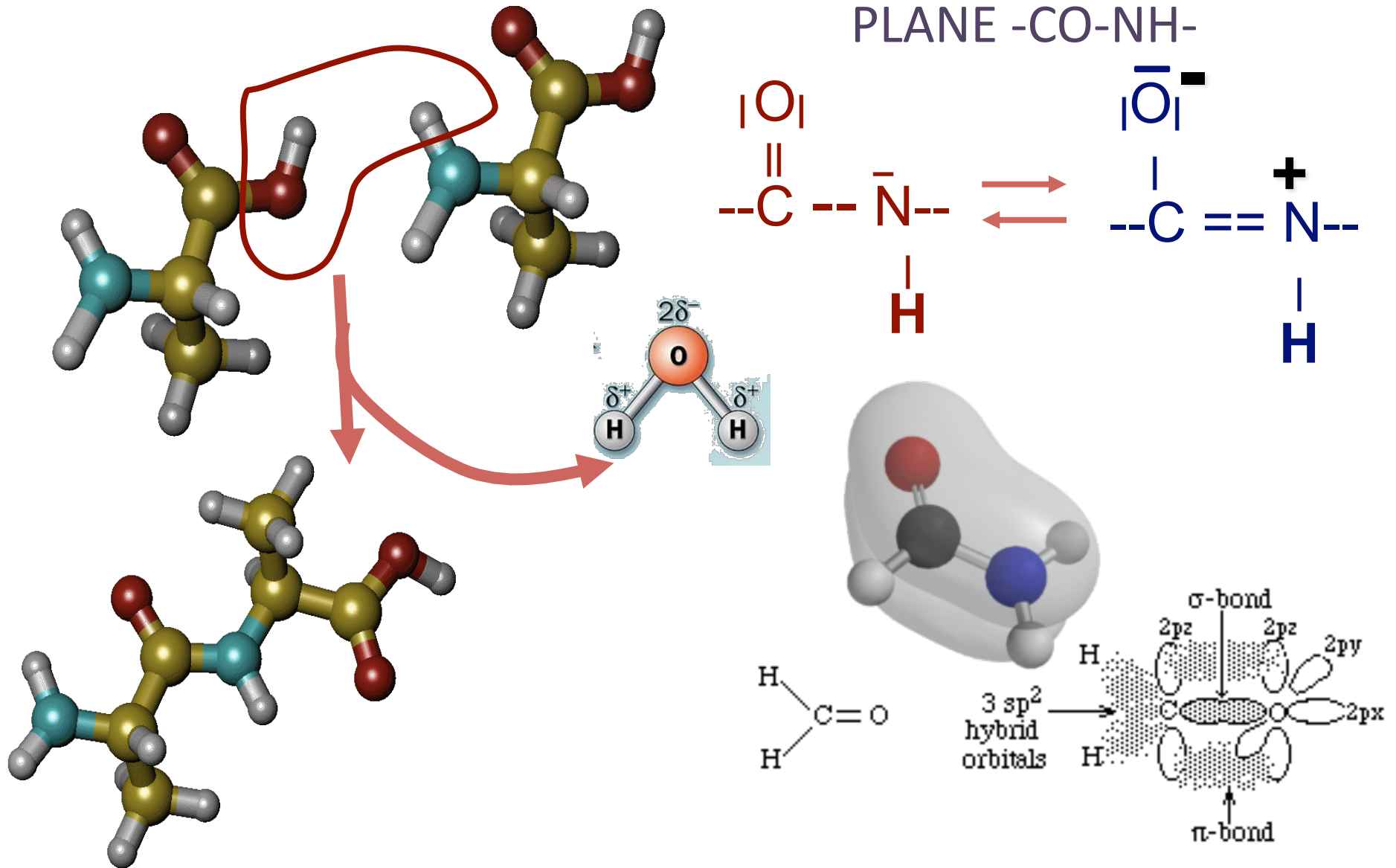
Arginine



Histidine

1. Introduction

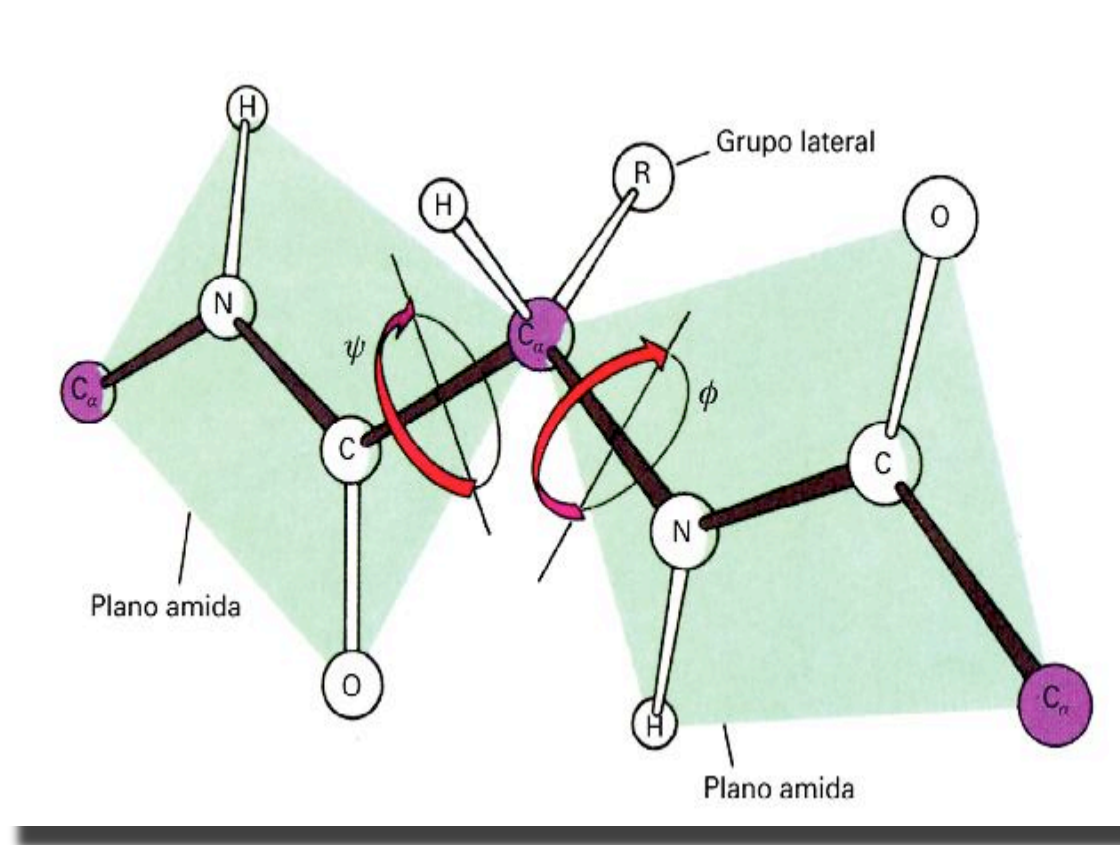
2. Peptide Bond



1. Introduction

2. Peptide Bond

Angles ϕ & ψ



$$\psi = [-180^\circ, +180^\circ]$$

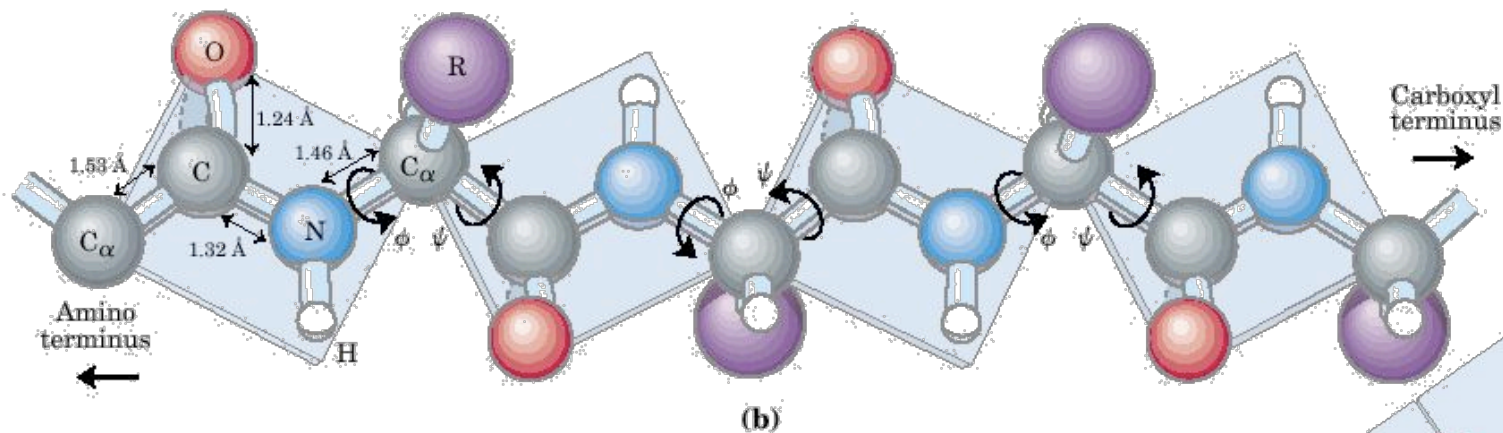
$$\phi = [-180^\circ, +180^\circ]$$

1. Introduction

3. Native Conformation

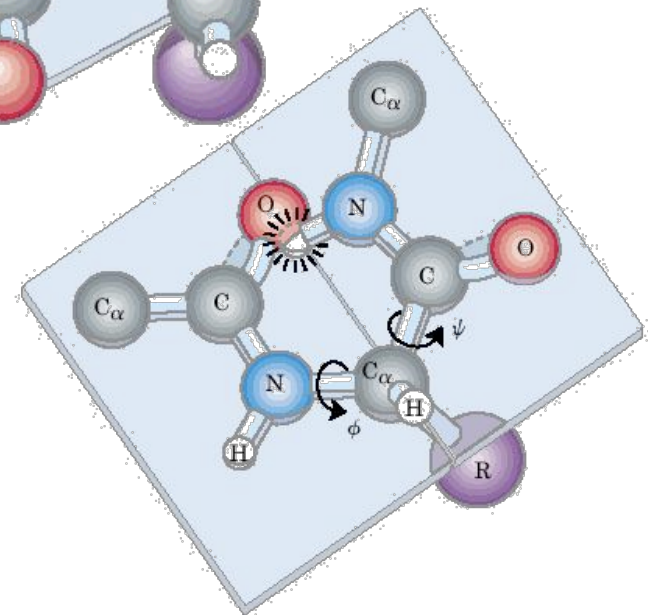
Native conformation:

- 1) Polipeptide + environment
- 2) Function & stability



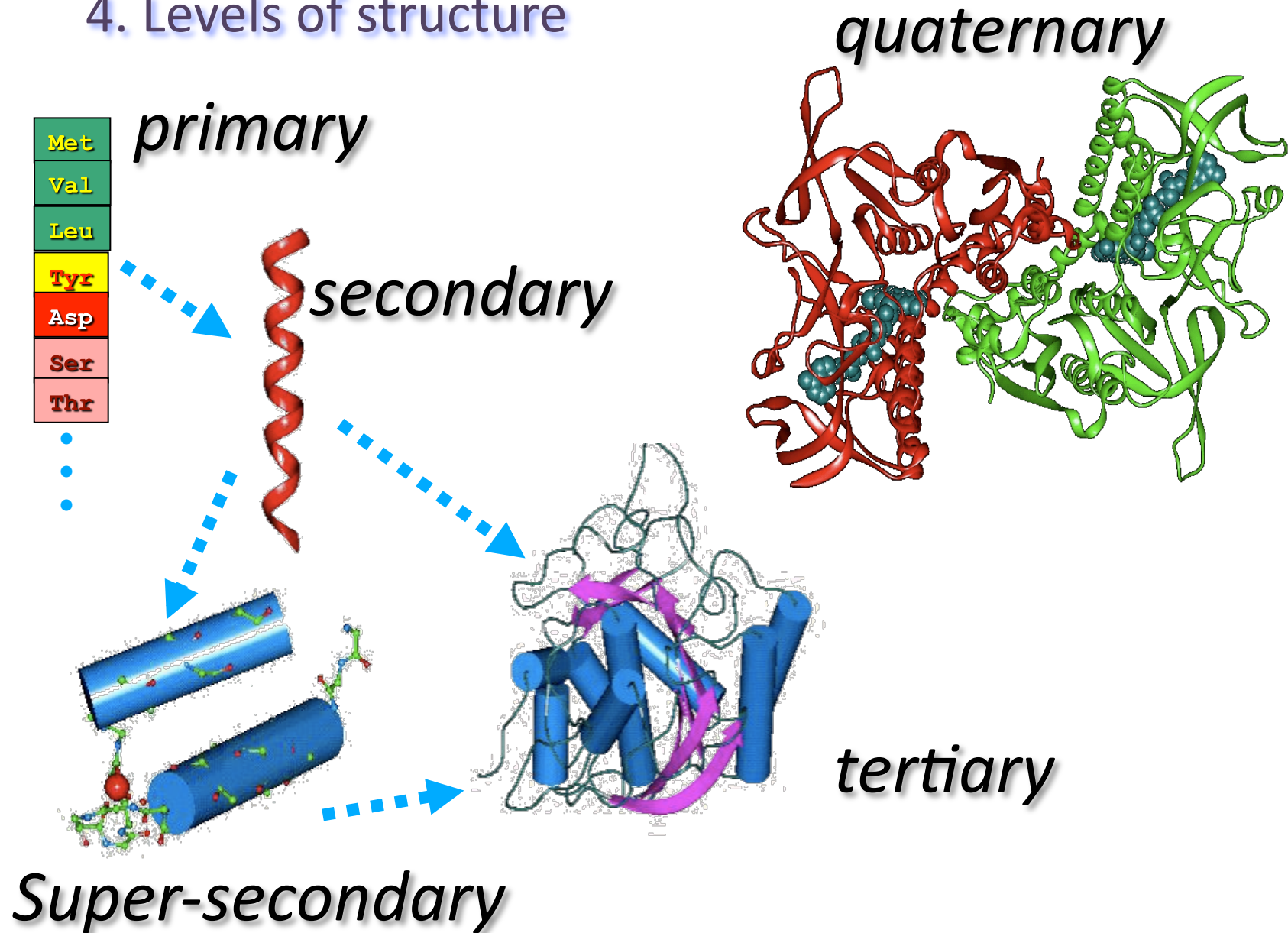
Conformational restraints:

- 1) Peptide bond planarity
- 2) The total space is not 360 x 360



1. Introduction

4. Levels of structure



2. Secondary structure

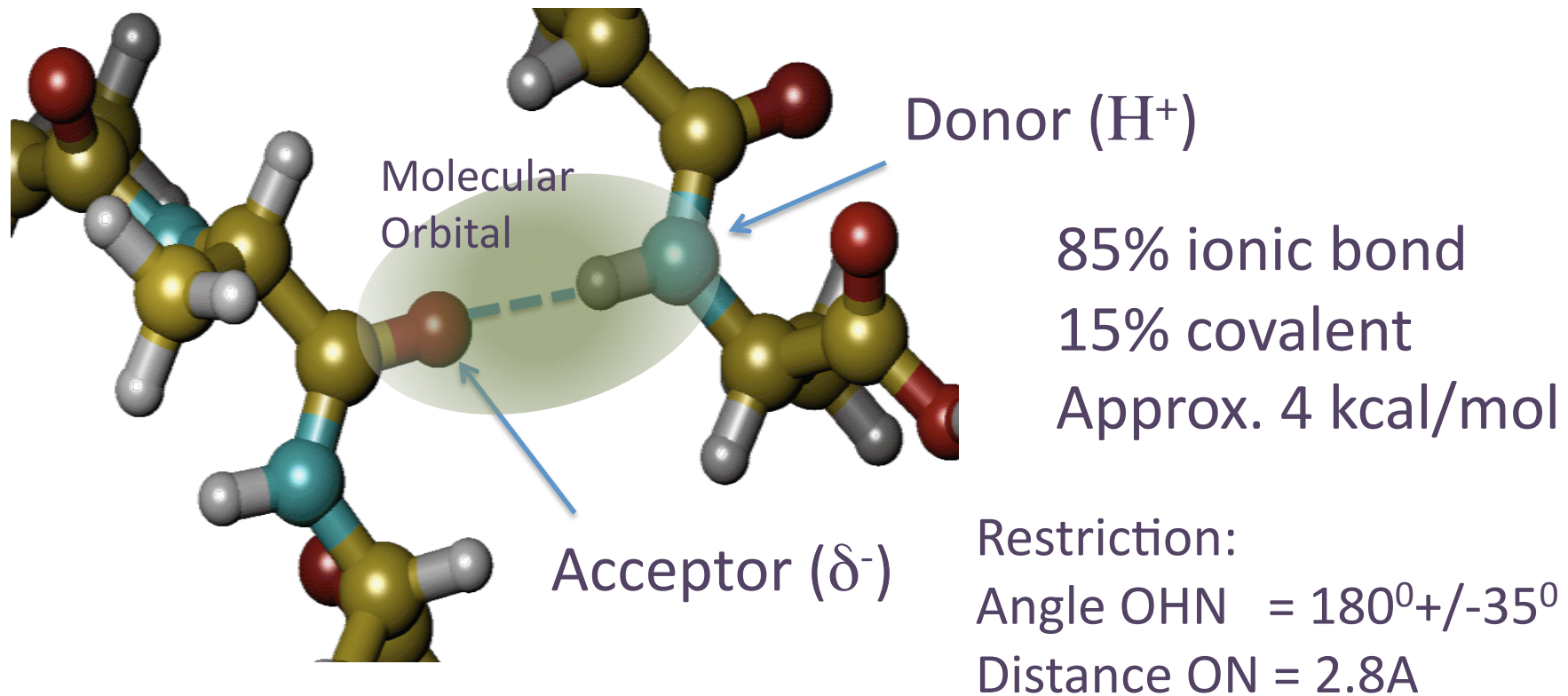
1. Bonding energy terms

Molecular mechanics

$$\begin{aligned} V = & \frac{1}{2} \sum_i K_i^{bond} (d_i - d_{0,i})^2 + \frac{1}{2} \sum_i K_i^{angle} (\alpha_i - \alpha_{0,i})^2 + \frac{1}{2} \sum_i K_i^{dihedral} (\omega_i - \omega_{0,i})^2 \\ & + \frac{1}{2} \sum_i K_i^{torsion} \cos(\lambda_i \phi_i + \delta_i) + \frac{1}{4\pi\epsilon} \sum_i \sum_{j>i} \frac{q_i q_j}{r_{ij}} + \sum_i \sum_{j>i} \left(\frac{C_6(i,j)}{r_{ij}^6} - \frac{C_{12}(i,j)}{r_{ij}^{12}} \right) \\ & + \sum_i (Hydrogen - Bonds) + \sum_i (\pi - \pi) \end{aligned}$$

2. Secondary structure

3. Hydrogen Bonds

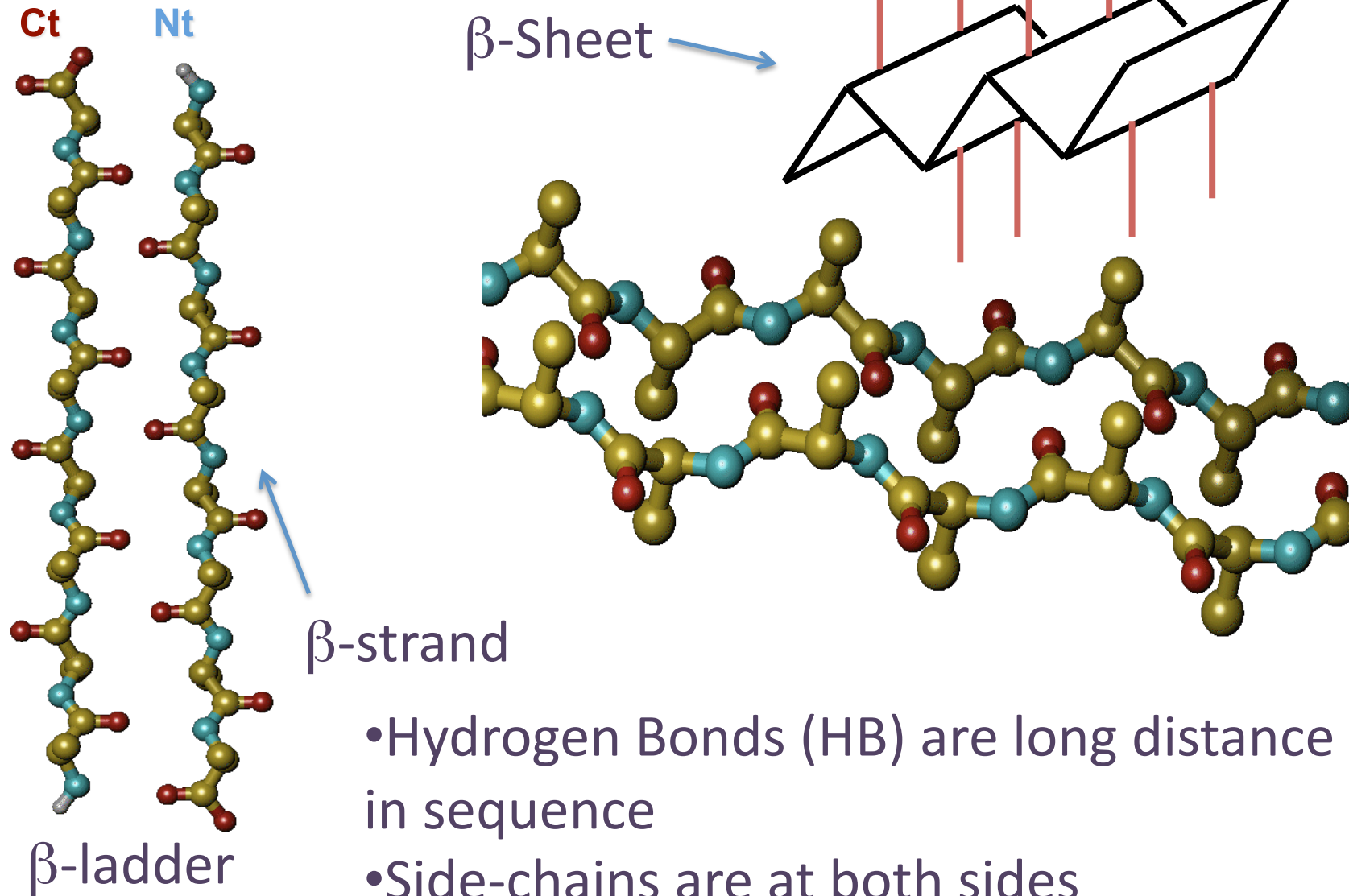


Inter-chain (long distance/short distance in sequence)

Intra-chain

2. Secondary Structure

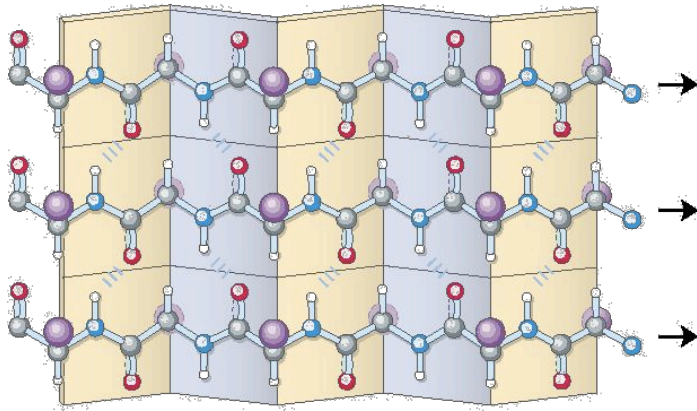
1. Strands



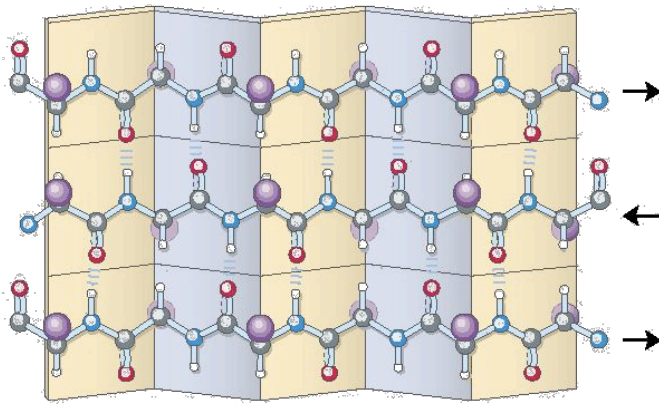
2. Secondary Structure

1. Strands

Parallel β -sheet



Anti-parallel β -sheet



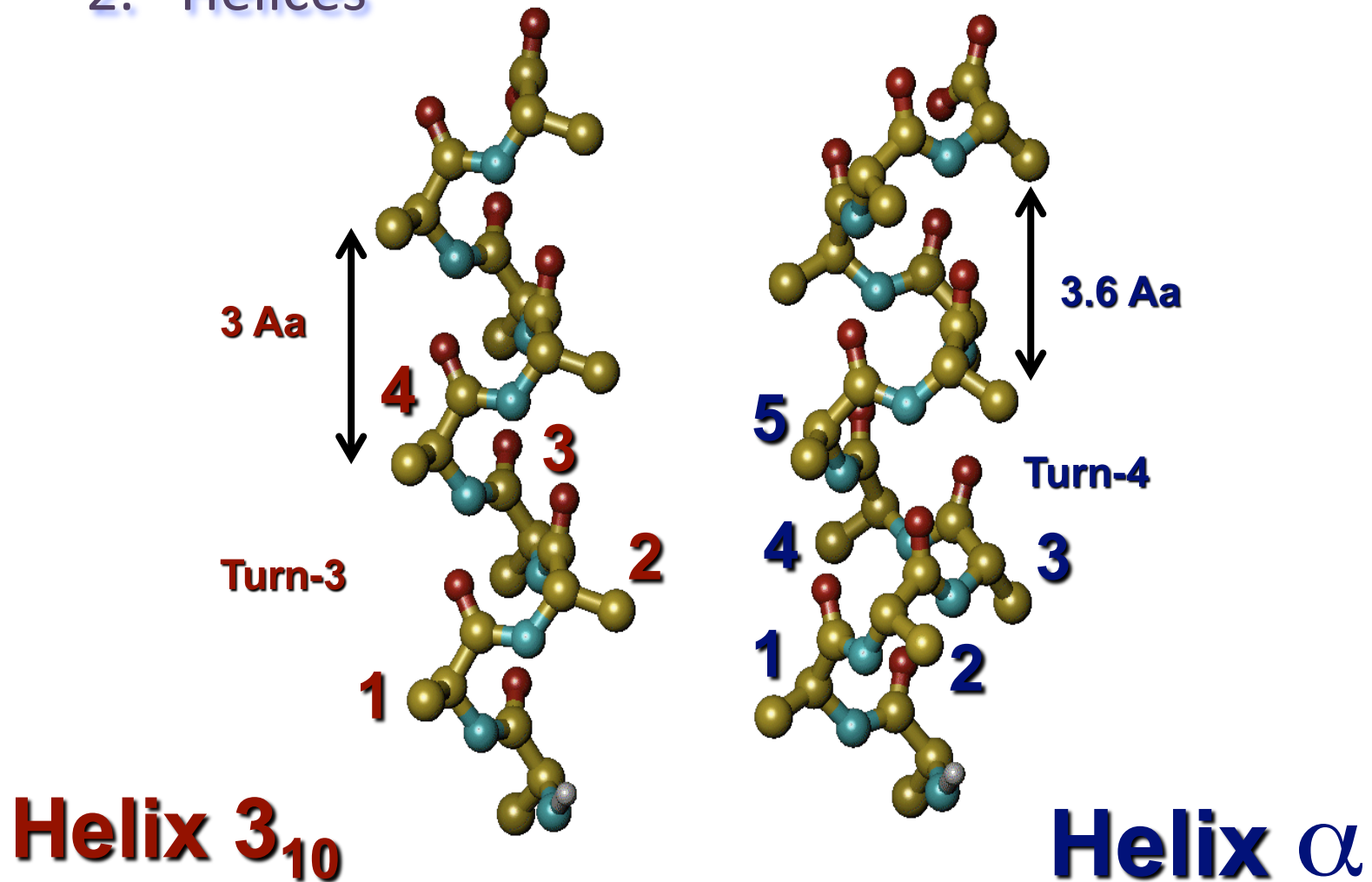
Upper face



Lower face

2. Secondary Structure

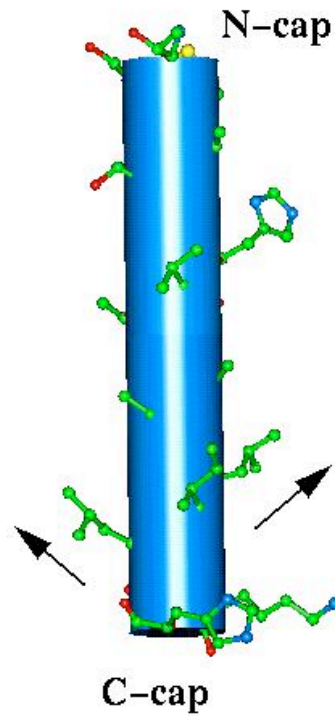
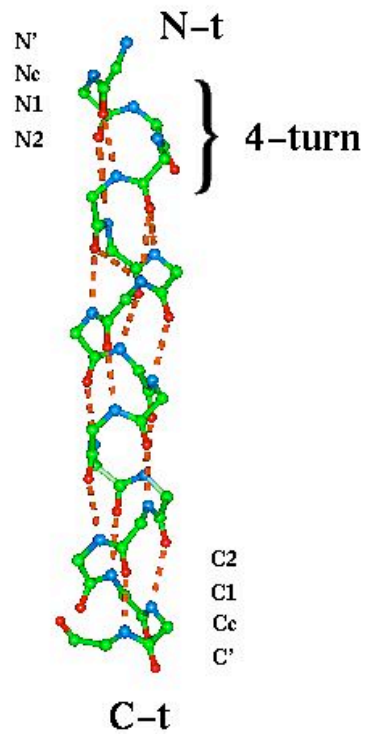
2. Helices



- HB are short distance in sequence
- Side-chains protrude out of the helix

2. Secondary Structure

2. Helices



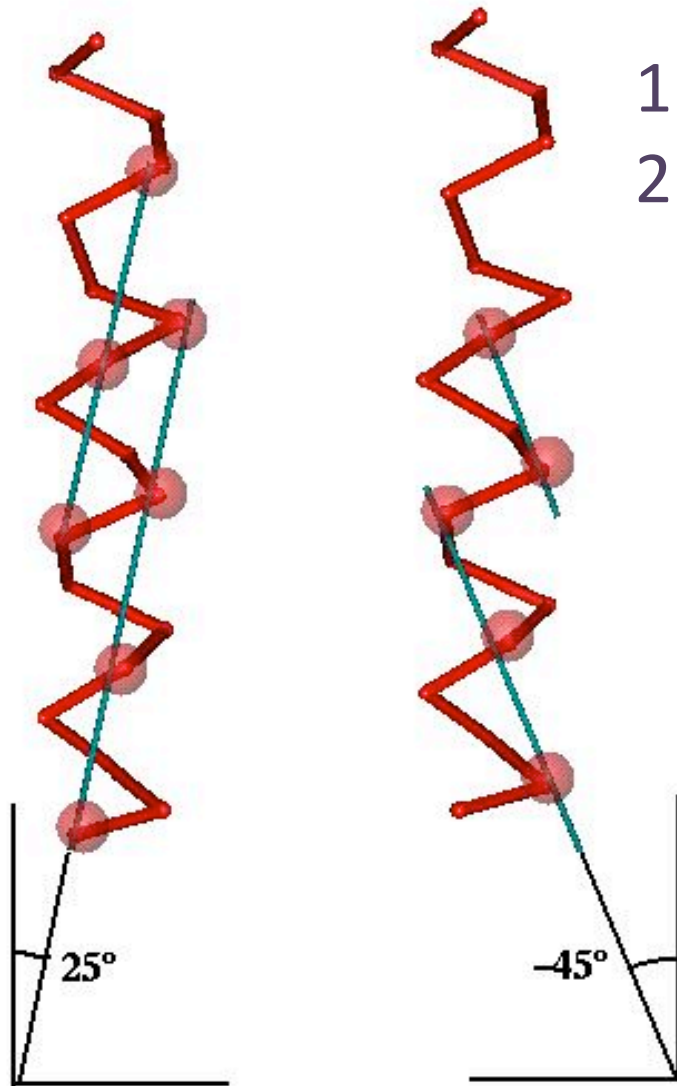
	Hbond	Hydrophobic
N-cap box	N_c-N_3	$N'-N_4$
Big box	N_c-N_3	$N''-N_4$
Shellman	$C''-C_3$ $C'-C_2$	$C''-C_3$
α_L	$C'-C_3$	$C''-C_3$

2. Secondary Structure

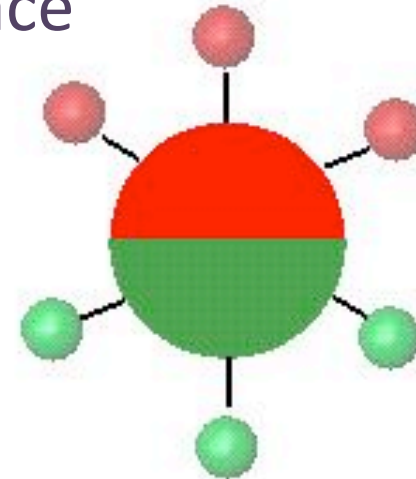
2. Helices

Side-chains location in the helix:

1. Groove formation as in a screw
2. Amphipathic helix: two faces with different solvation properties



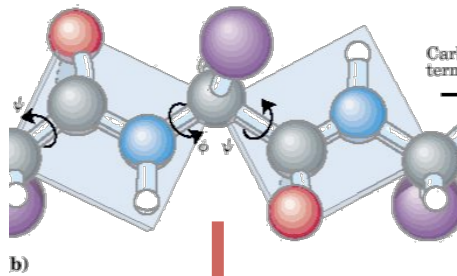
Polar face



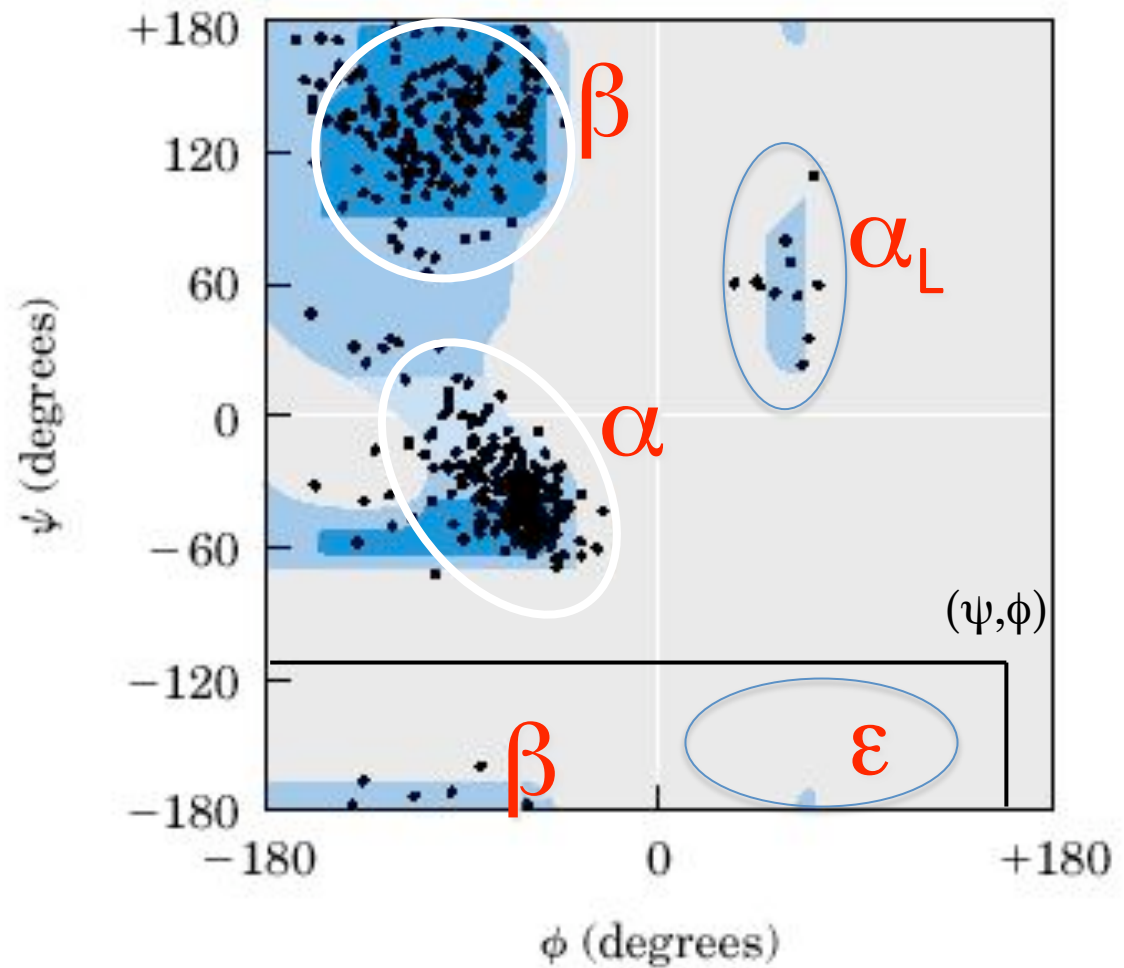
Non-polar face
(hydrophobic)

2. Secondary Structure

3. Ramachandran Plot

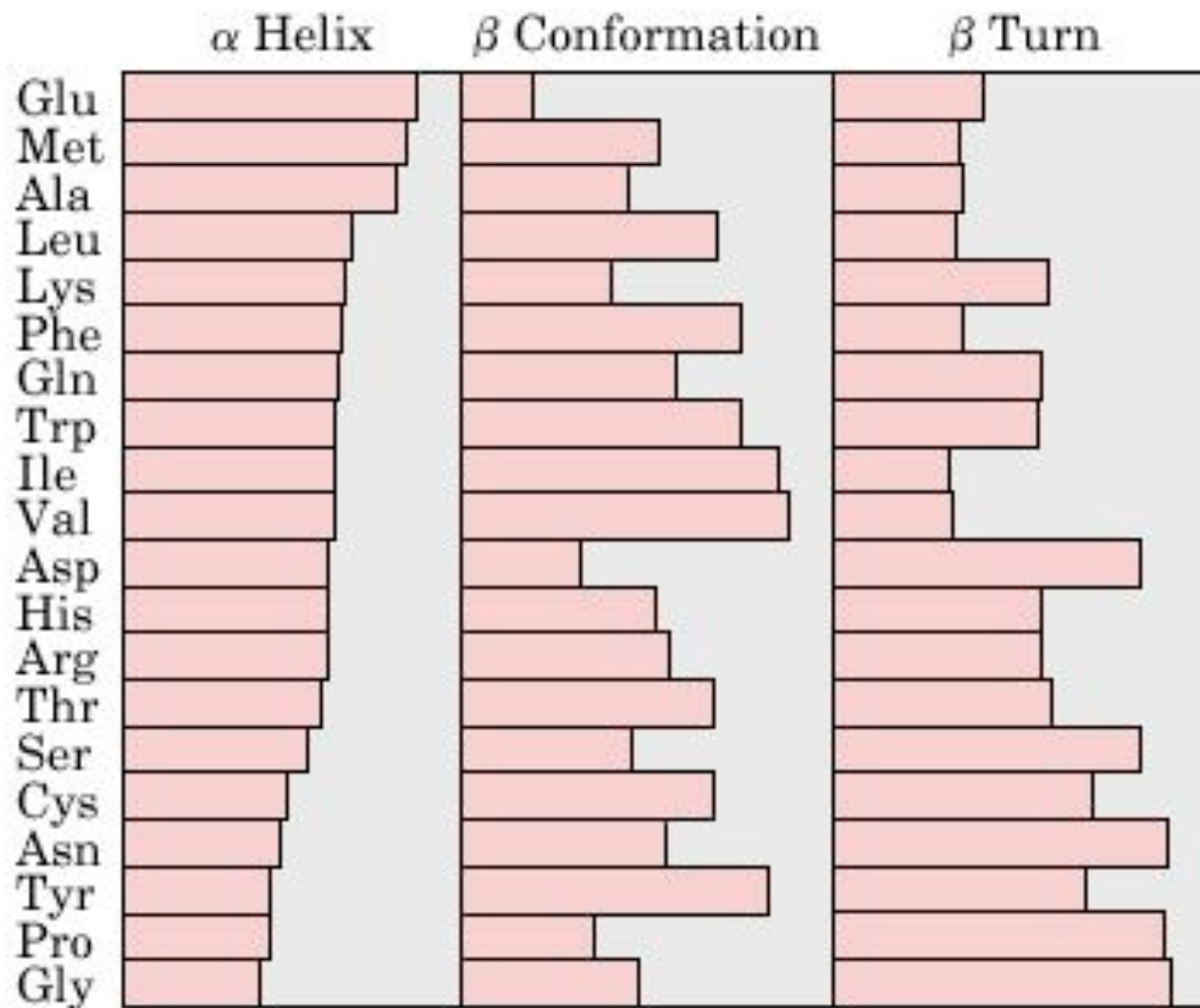


(ψ, ϕ)



2. Secondary Structure

4. Aa propensity



Protein structure

3. Soluble and globular proteins

1. Supersecondary structure

1. Definition

2. α - α

3. β - β

4. β - α & α - β

5. β - α - β

2. Domains

1. Definition

2. Classification

3. Function-Sequence-Structure

4. Databases:

SCOP & CATH

3. Soluble and globular proteins

1. Supersecondary structure

1. Definition

MOTIF or Super-Secondary Structure:

This is defined as a cluster of 2-4 regular secondary structures, usually involved in a particular function.

- Regular secondary structures are connected by loops.
- Loops are highly flexible and produce change of the polypeptide chain orientation.
- Length of loops varies from 1 to more than 40 residues
- Some super-secondary structures are stabilized with an hydrophobic core

3. Soluble and globular proteins

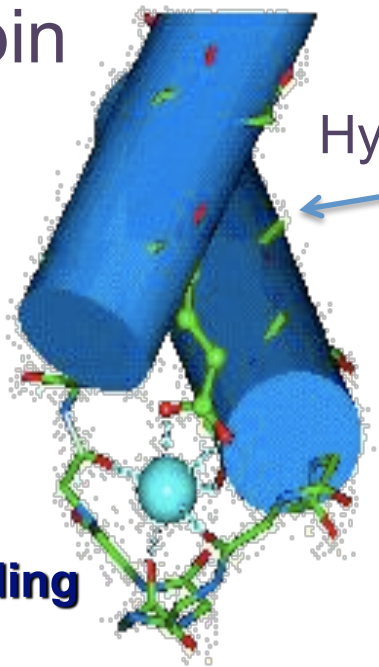
1. Supersecondary structure

2. α - α

Ca^{+2} EF hand

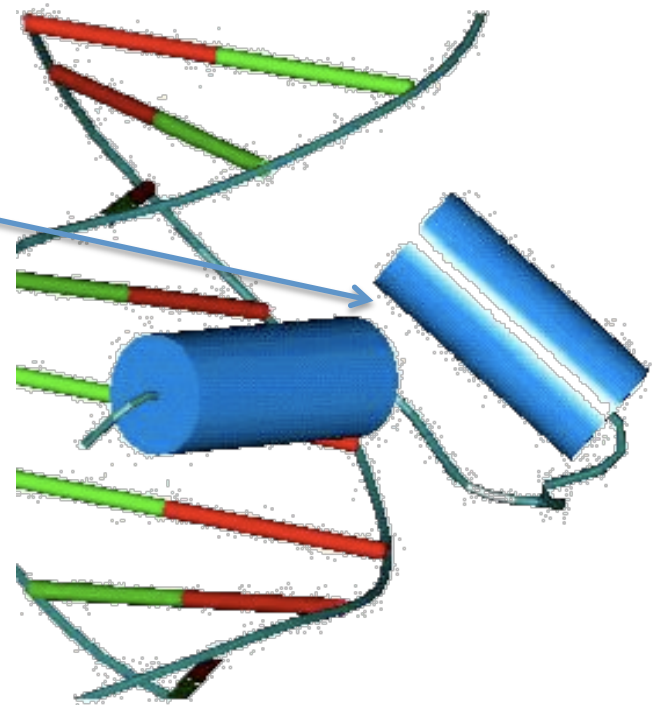
α hairpin

Ca^{+2} binding



Hydrophobic core

DNA binding motif
(Cro repressor)

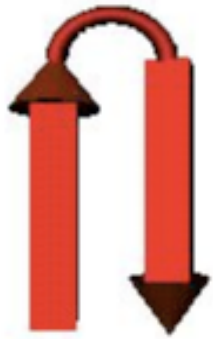


3. Soluble and globular proteins

1. Supersecondary structure

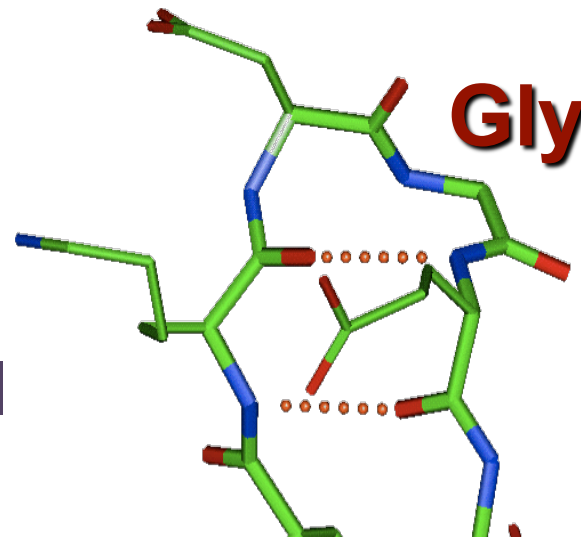
3. β - β

β - β hairpin



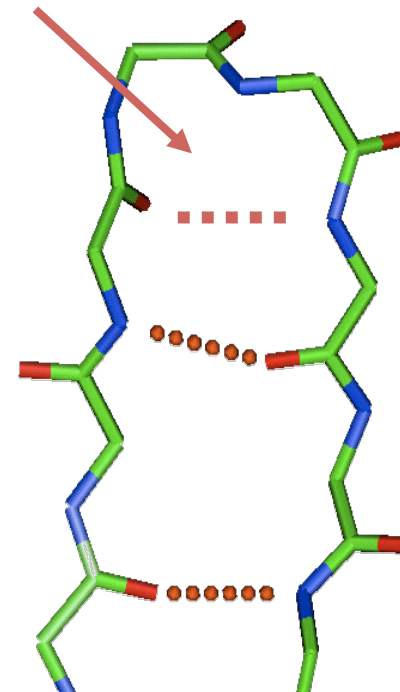
Anti-parallel
 β -ladder

Turn $\beta = 2$ Aa



Type I'

H.B. lost

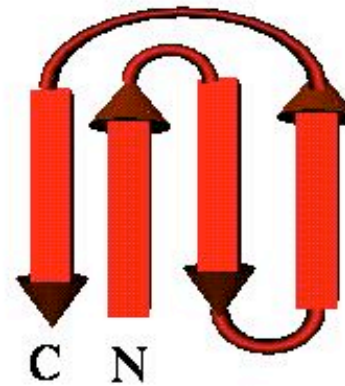
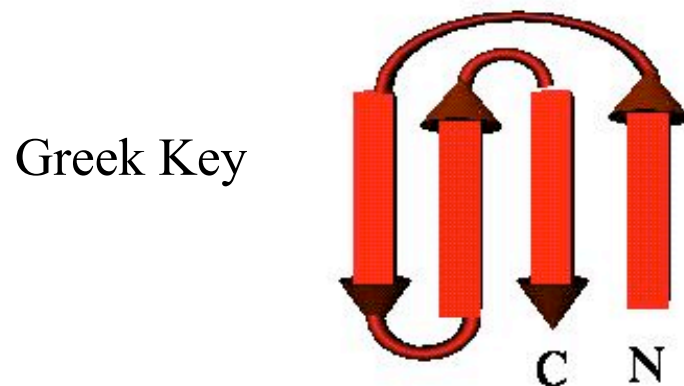
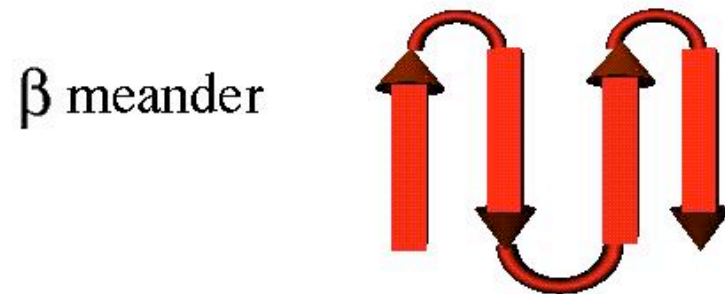


Type II

3. Soluble and globular proteins

1. Supersecondary structure

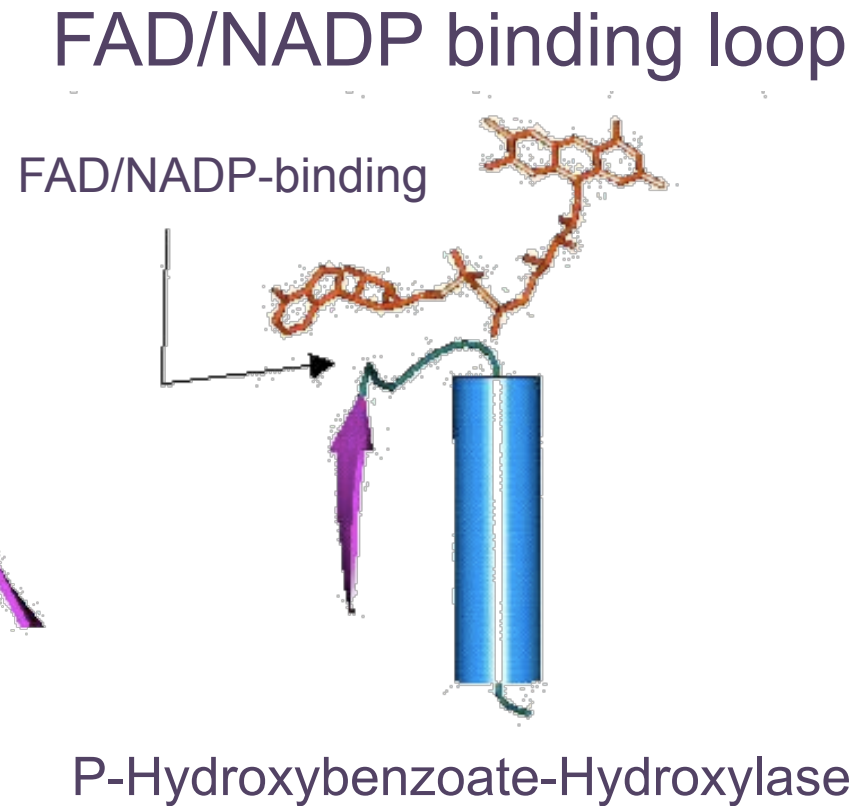
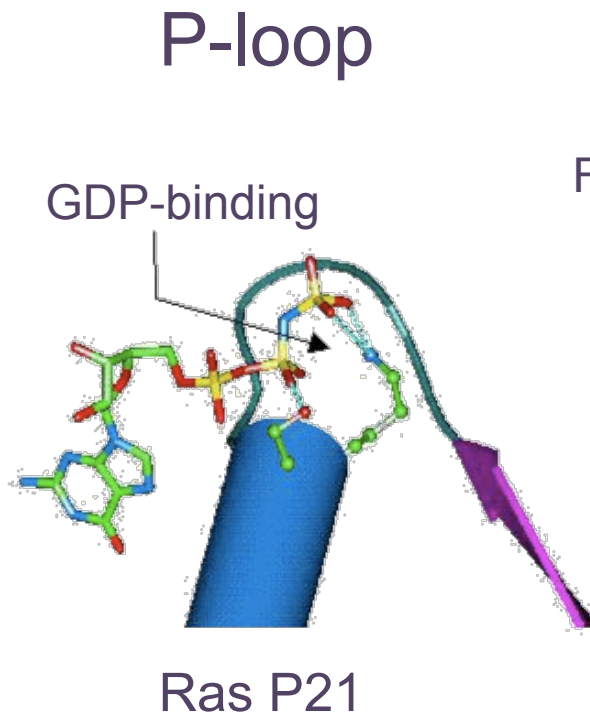
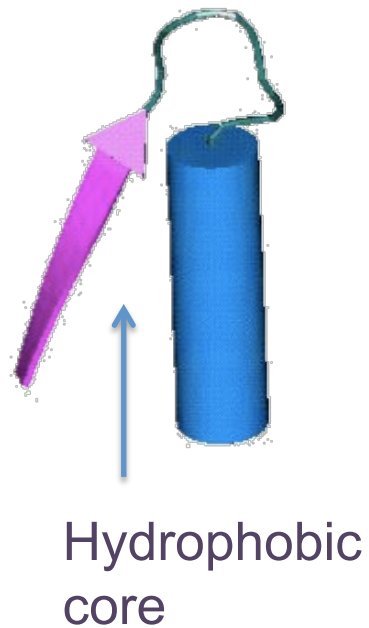
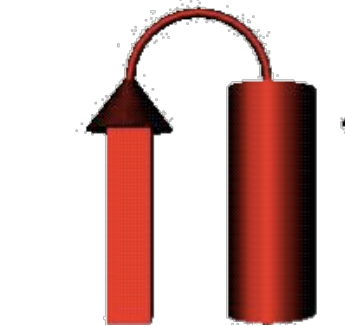
3. β - β



3. Soluble and globular proteins

1. Supersecondary structure

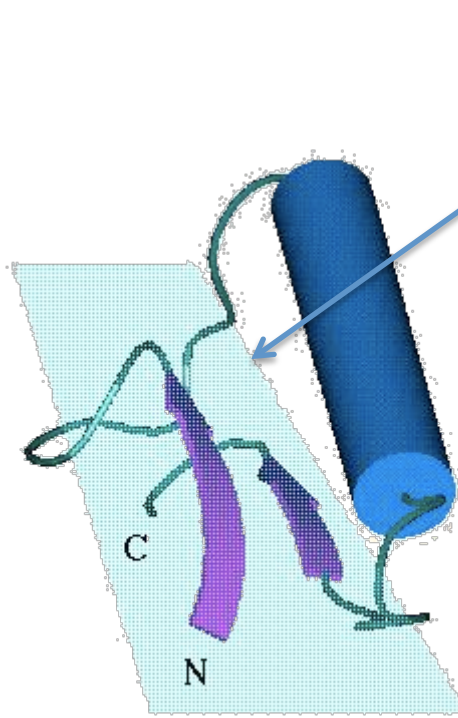
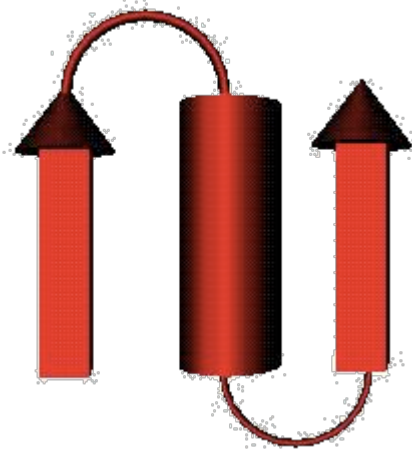
4. β - α & α - β



3. Soluble and globular proteins

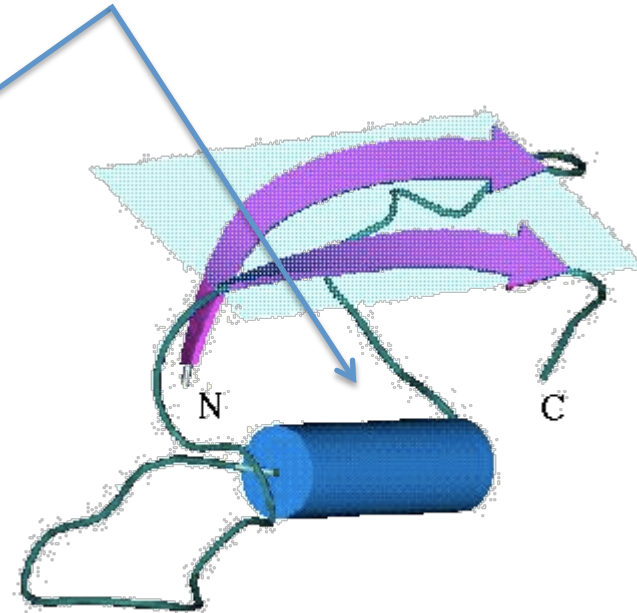
1. Supersecondary structure

5. β - α - β



Right handed

Hydrophobic core

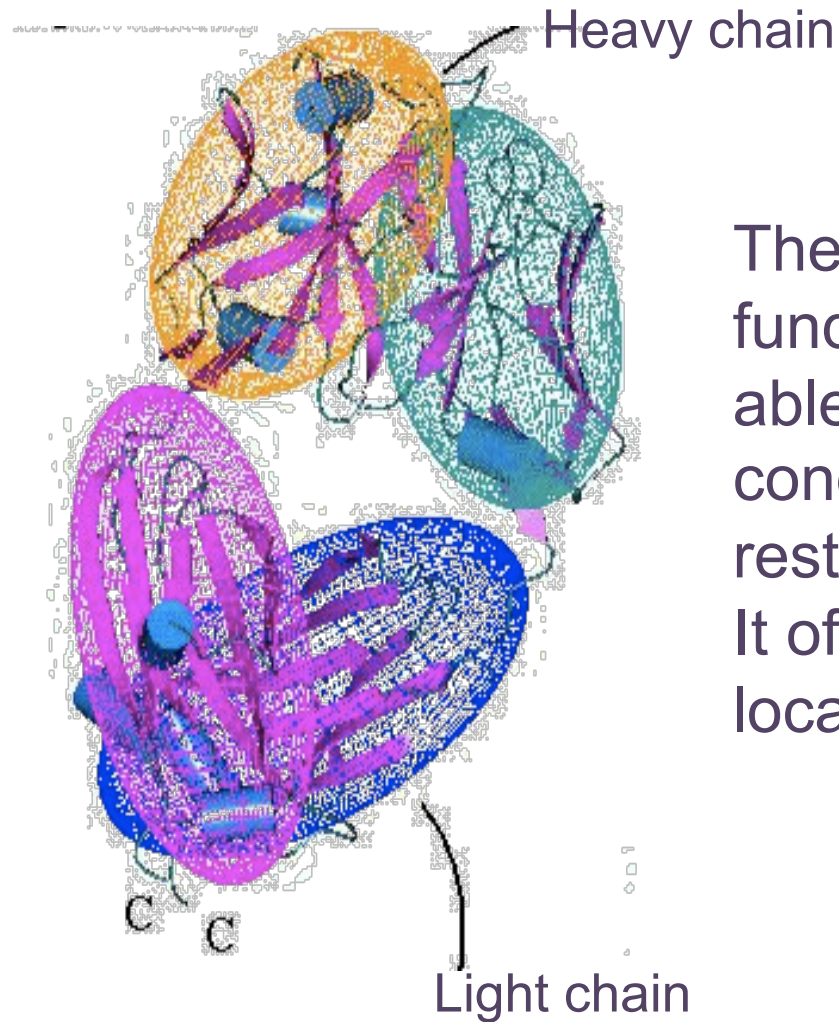


Left handed

3. Soluble and globular proteins

2. Domains

1. Definition



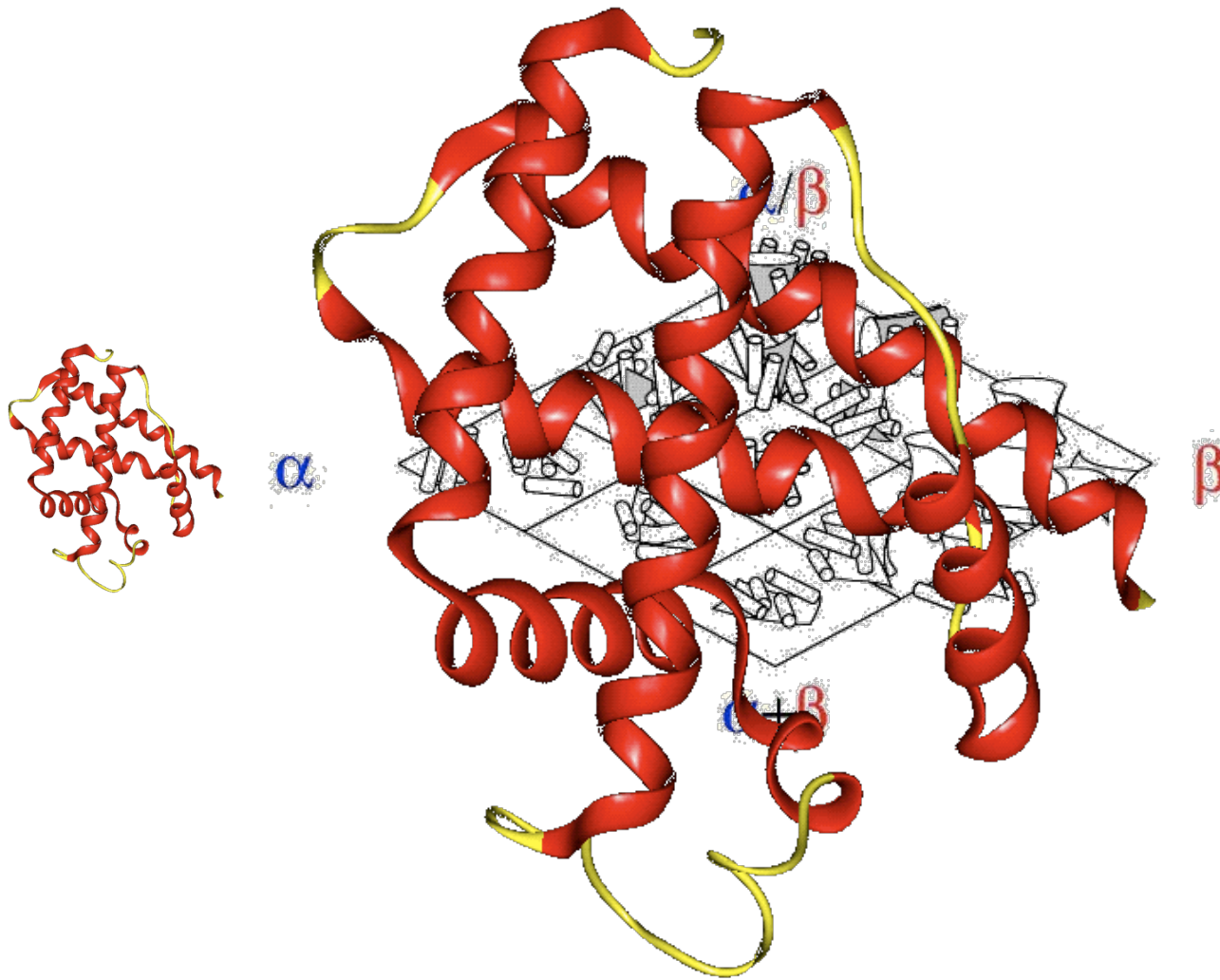
The **protein domain** is defined as the fundamental unit of 3D structure, able to fold by itself in the right conditions with independence of the rest of the protein.

It often corresponds to functional local and compact units of a protein

3. Soluble and globular proteins

4. Domains

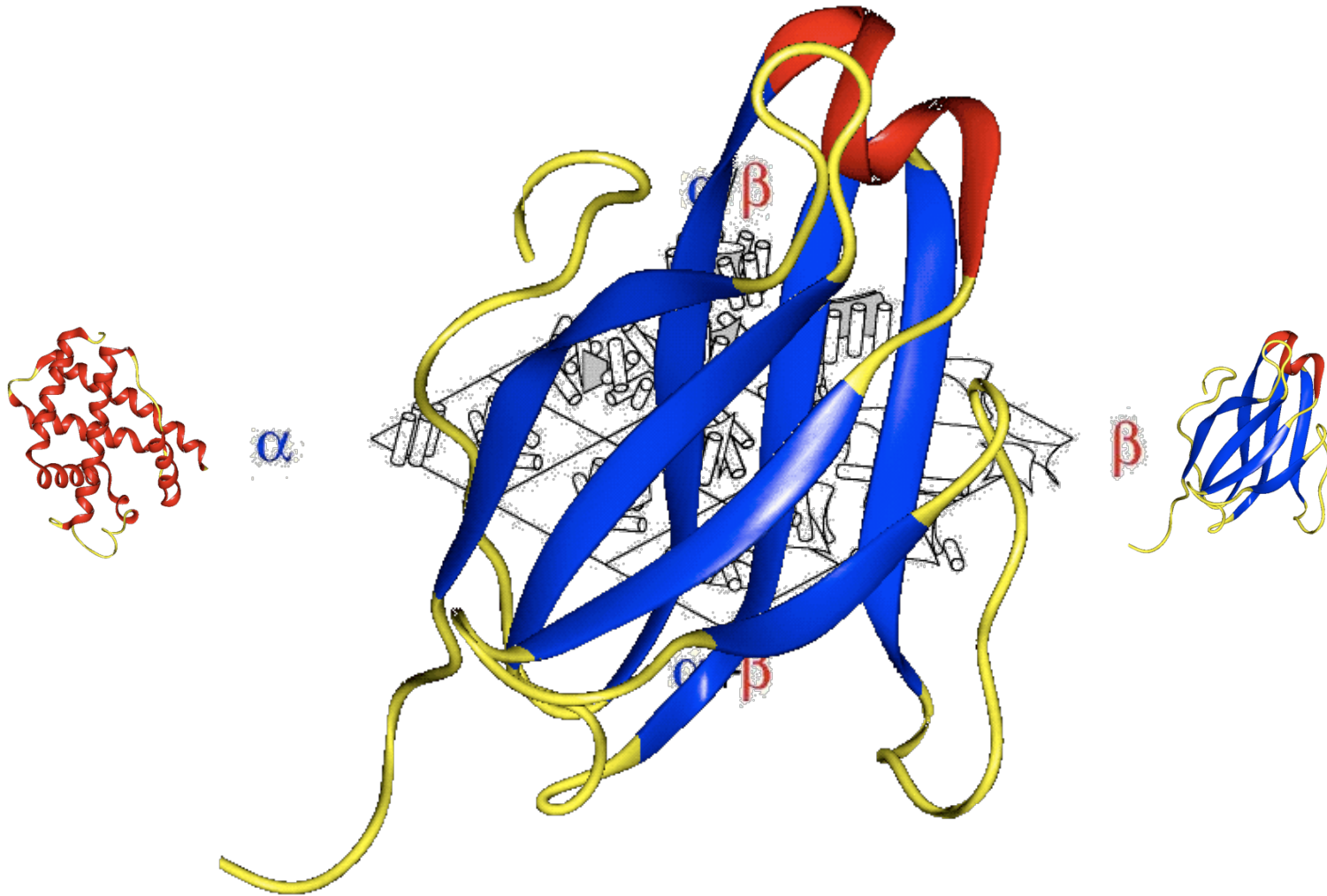
2. Classification



3. Soluble and globular proteins

4. Domains

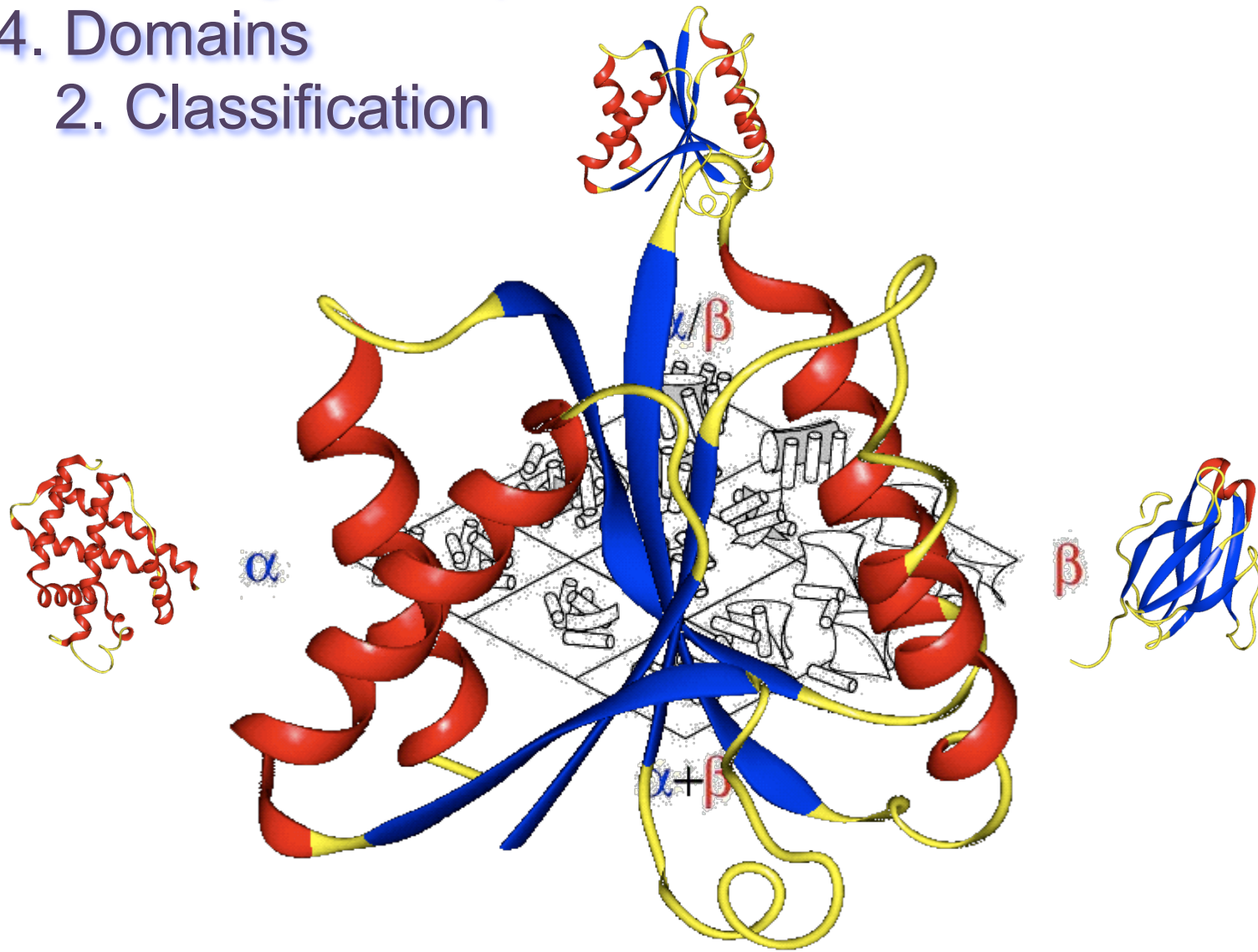
2. Classification



3. Soluble and globular proteins

4. Domains

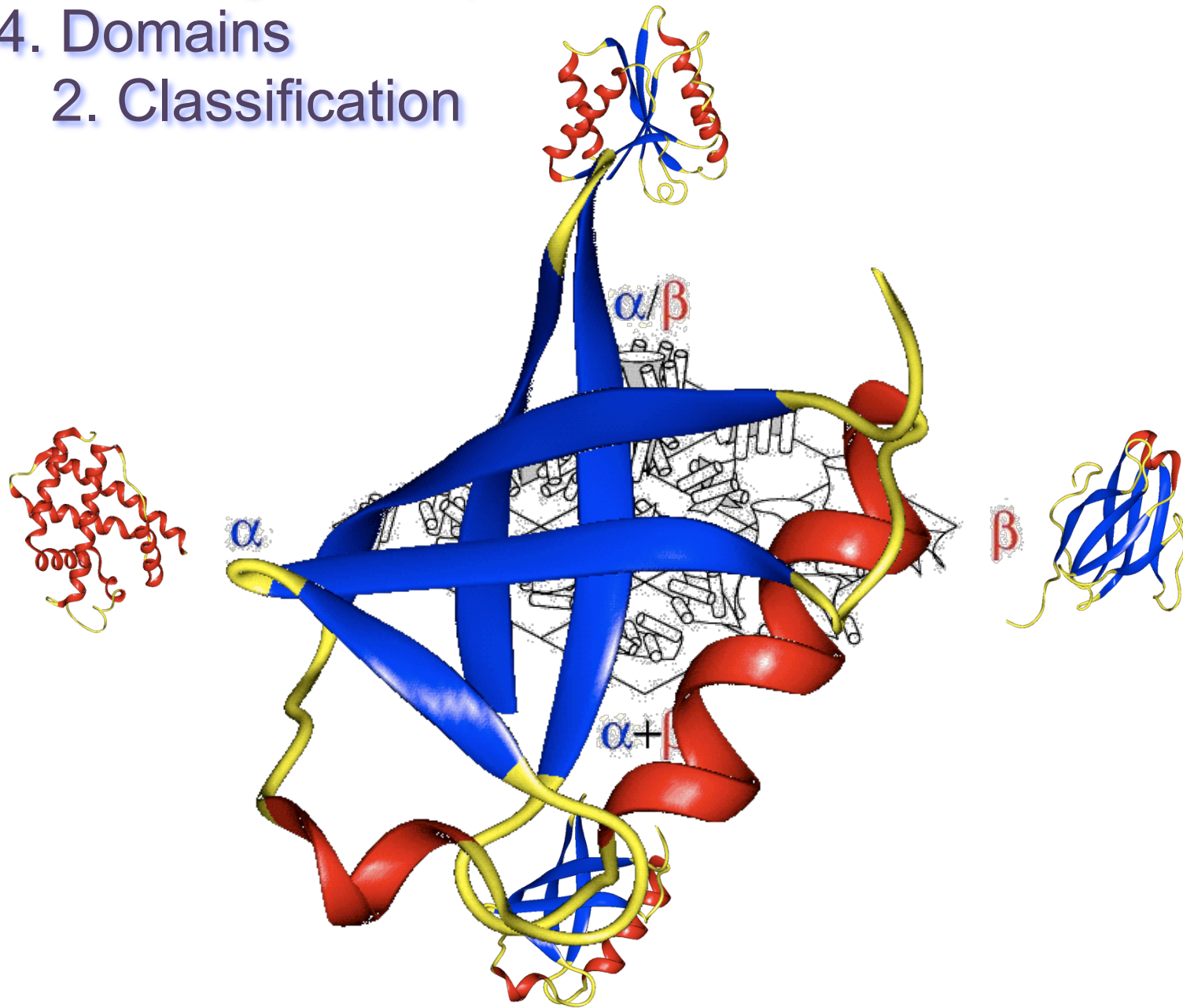
2. Classification



3. Soluble and globular proteins

4. Domains

2. Classification



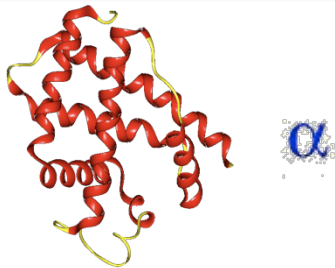
3. Soluble and globular proteins

4. Domains

2. Classification

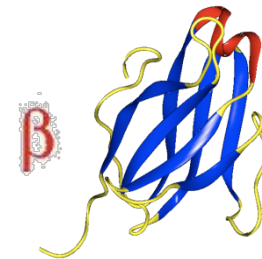
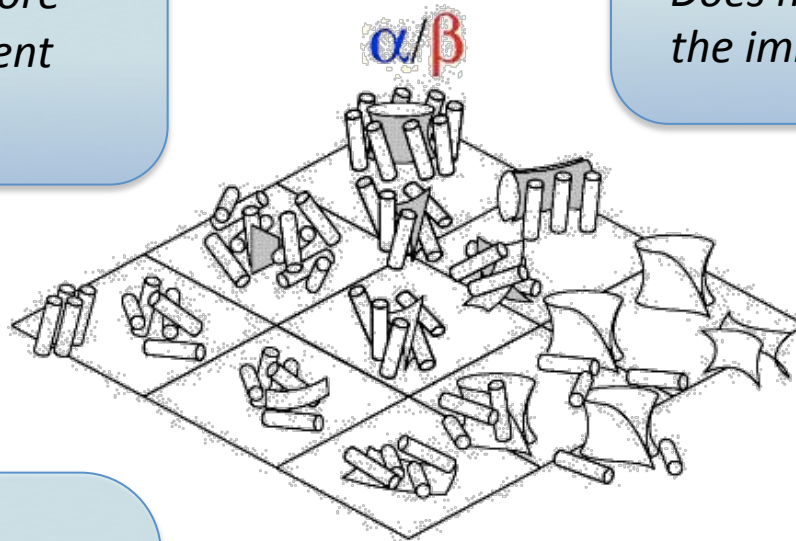
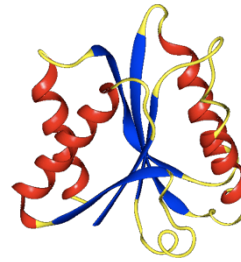
5th class

Multi-domain proteins α and β
Folds consisting of two or more domains belonging to different classes



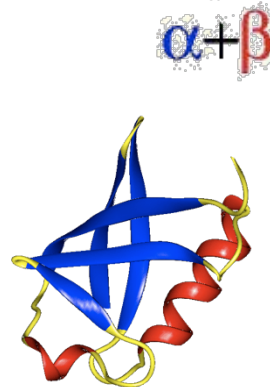
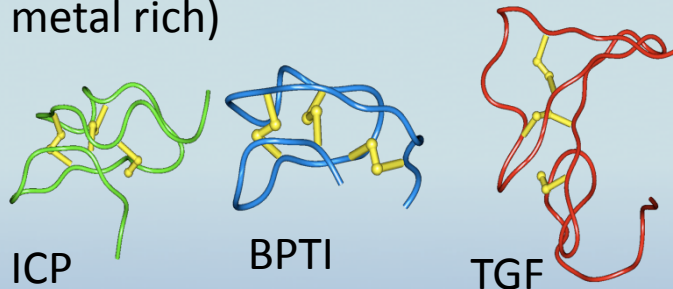
6th class

Membrane and cell surface proteins and peptides.
Does not include proteins in the immune system



7th class

Small proteins (disulphide-rich, metal rich)

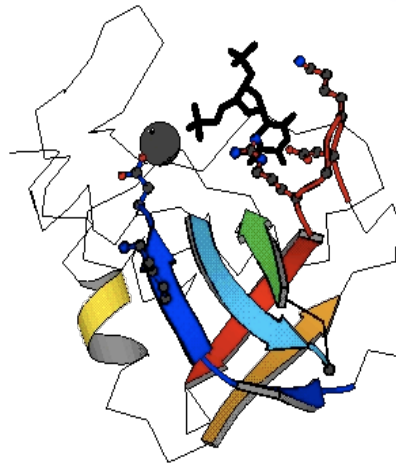


3. Soluble and globular proteins

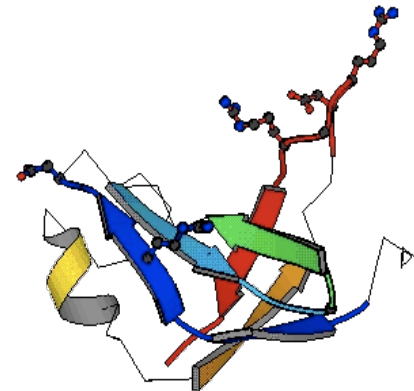
4. Domains

3. Function-Sequence-Structure

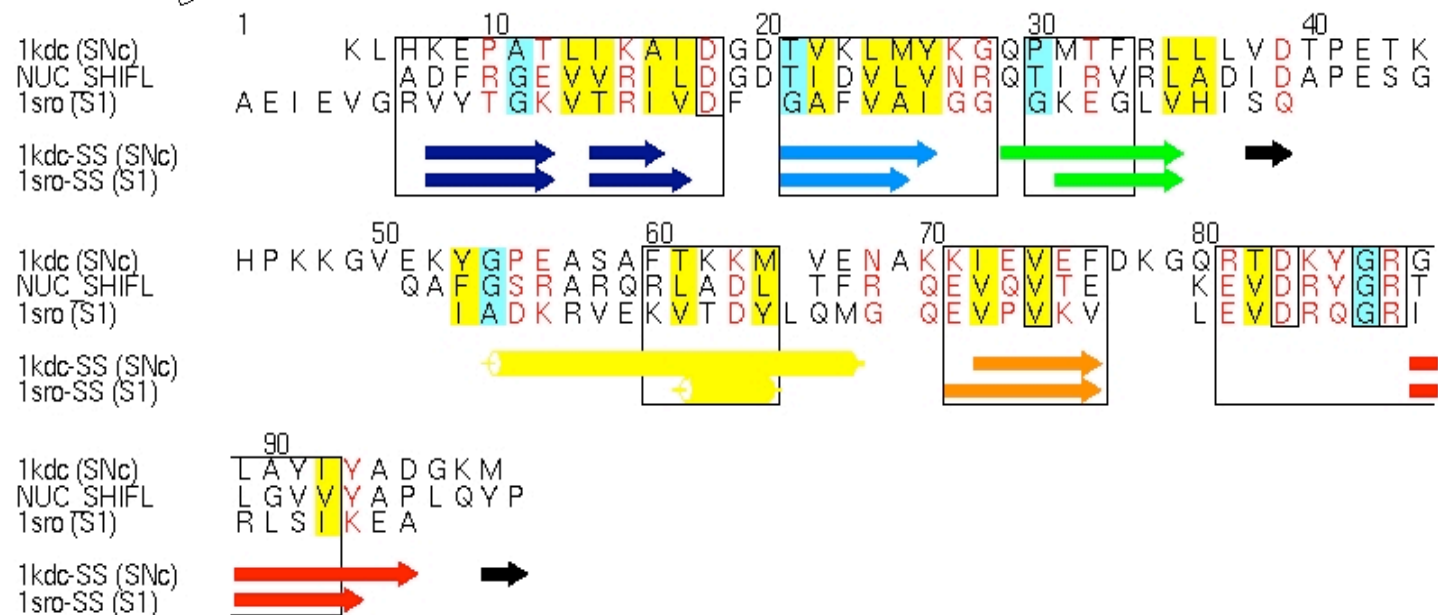
Staphylococcal nuclease



S1 RNA binding domain

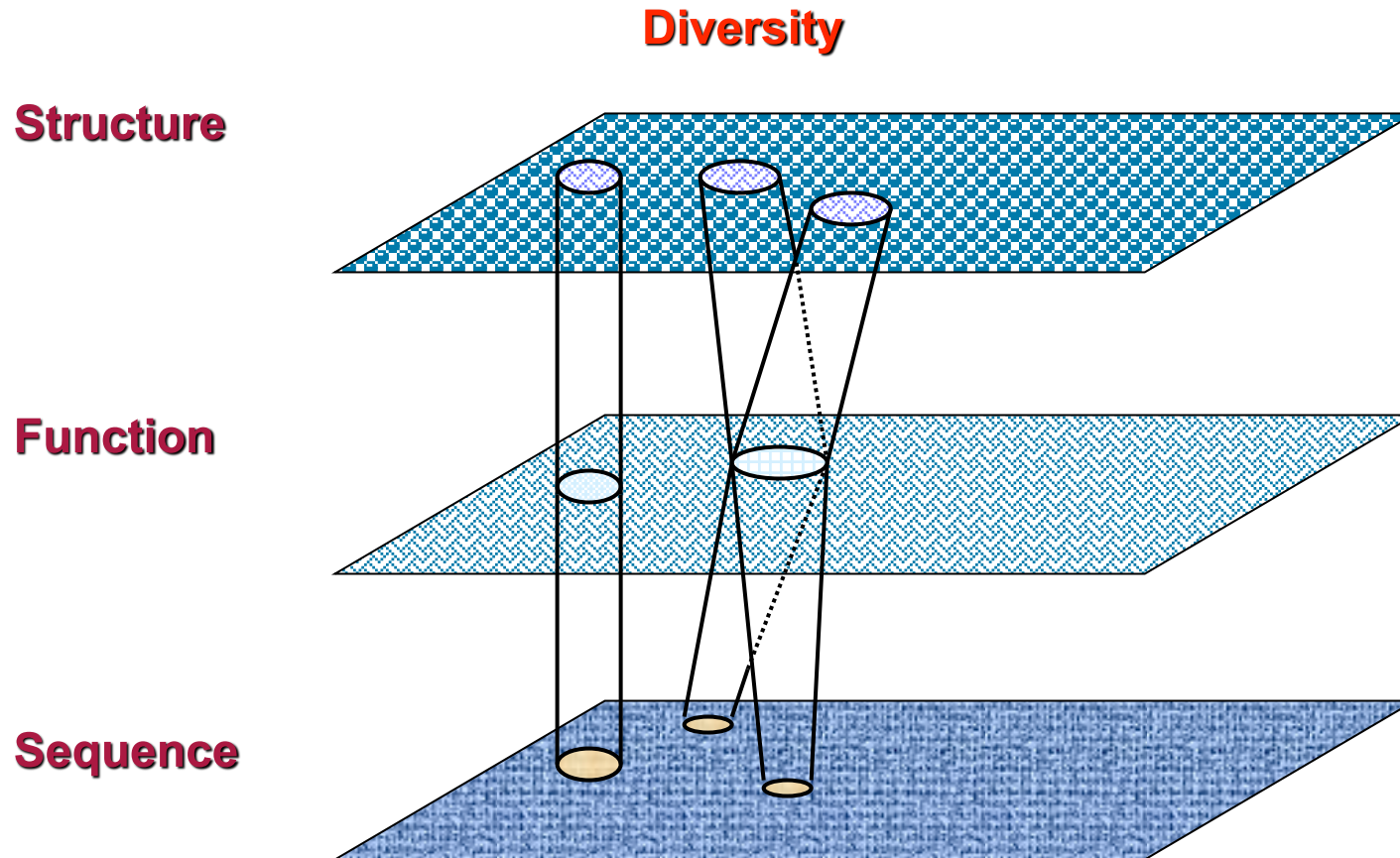


The structural alignment shows the best conservation of residues and a possible relationship in evolution



3. Soluble and globular proteins

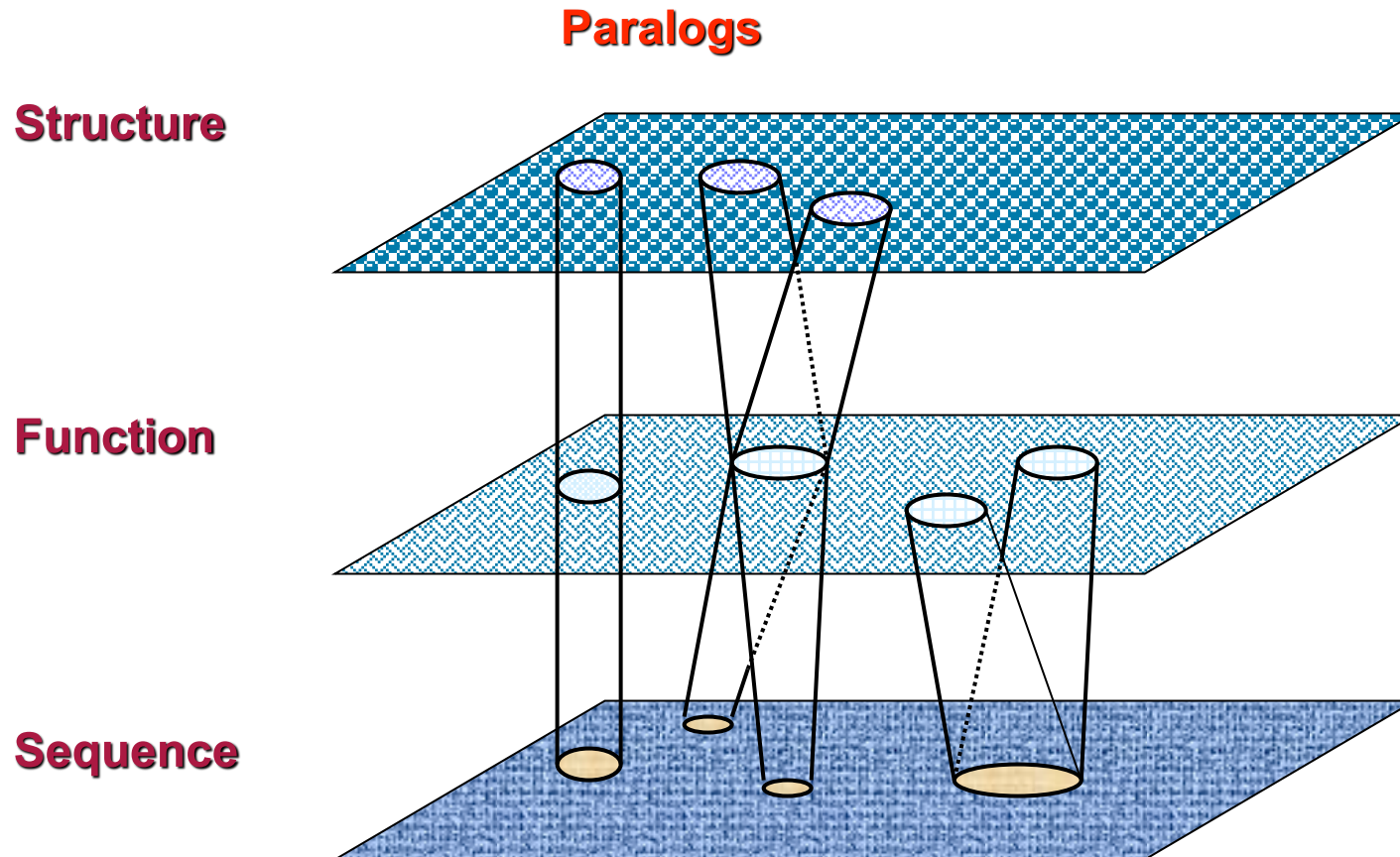
4.3. Function-Sequence-Structure



Homology: two proteins are homologous if they are the products of genes that evolved from the same ancestor

3. Soluble and globular proteins

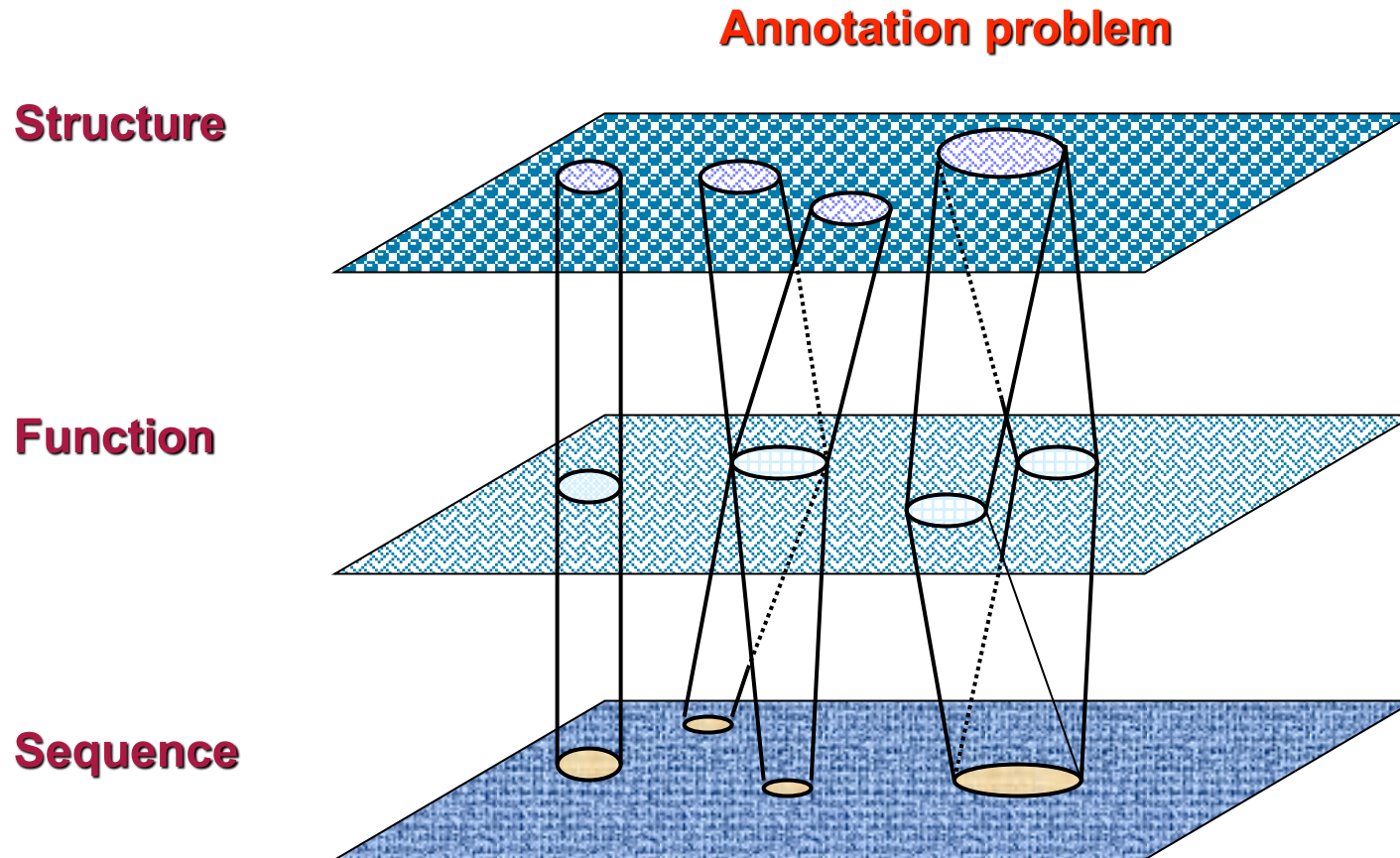
4.3. Function-Sequence-Structure



Homology: two proteins are homologous if they are the products of genes that evolved from the same ancestor

3. Soluble and globular proteins

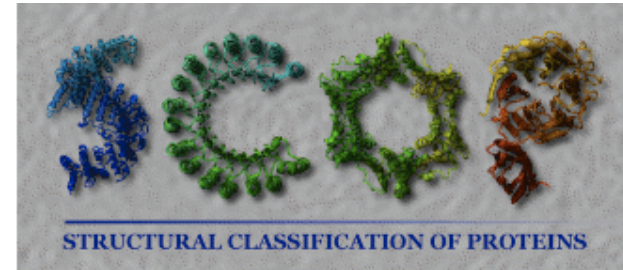
4.3. Function-Sequence-Structure



Homology: two proteins are homologous if they are the products of genes that evolved from the same ancestor

3. Soluble and globular proteins

4. database SCOP



Structural Classification Of Proteins (SCOP)

Class: It groups the folds according to the percentage and 3D disposition of the regular secondary structures.

Family: It groups proteins clearly related by homology. In general we assume they are homologs if the alignment has $>30\%$ ID, they have the similar structure and similar function

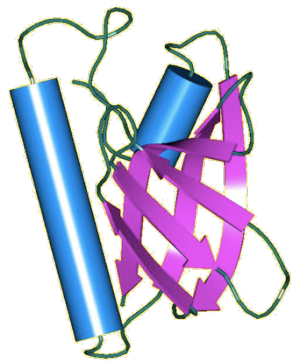
Superfamily: Proteins which sequences align with very few %ID but showing similar structural patterns and similar function.

Fold: Proteins with very similar disposition of the regular secondary structures

3. Soluble and globular proteins

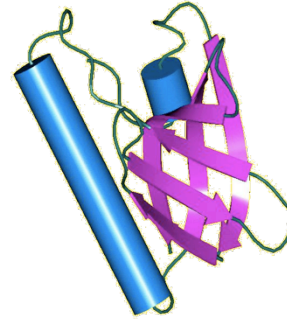
SCOP

- Family
- Superfamily
- Fold
- Class



Enterotoxin

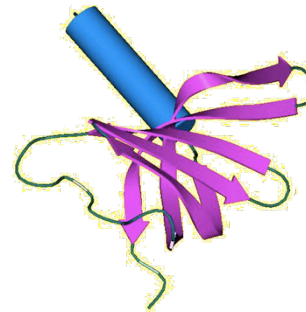
Homology



Cholera toxin

80% Id

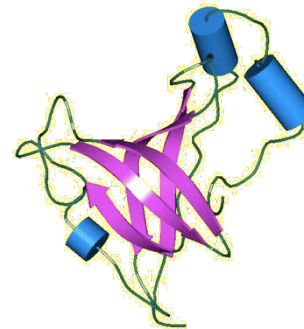
Remote
Homology



TSS toxin

8.8% Id

Analogs



Aminoacyl
tRNA synthetase

4.4% Id

3. Soluble and globular proteins

4. database CATH



- **Class(C)**

Class is determined according to the secondary structure composition and packing within the structure. It is **assigned automatically**

- **Architecture(A)**

This describes the overall shape of the domain structure as determined by the orientations of the secondary structures but ignores the connectivity between the secondary structures. It is currently **assigned manually**

- **Topology(T)**

Structures are grouped into fold groups at this level depending on both the overall shape and connectivity of the secondary structures. This is done using **an automated structure comparison algorithm**.

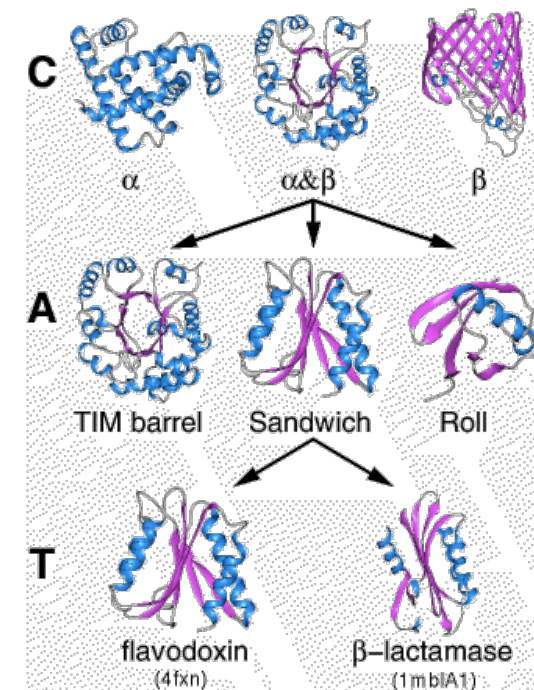
- **Homologous superfamily (H).**

This level groups together protein domains which are thought to share a common ancestor and can therefore be described as homologous. Similarities are **automatically** identified either by high sequence identity or **structure comparison**.

- **Sequence Family Levels: (S,O,L,I, D)**

Domains within each H-level are subclustered into sequence families using multi-linkage clustering S(35%), O(60%), L (90%), I (100%)

Class 1 Class 3 Class 2
Mainly α Mixed Mainly β
 α & β

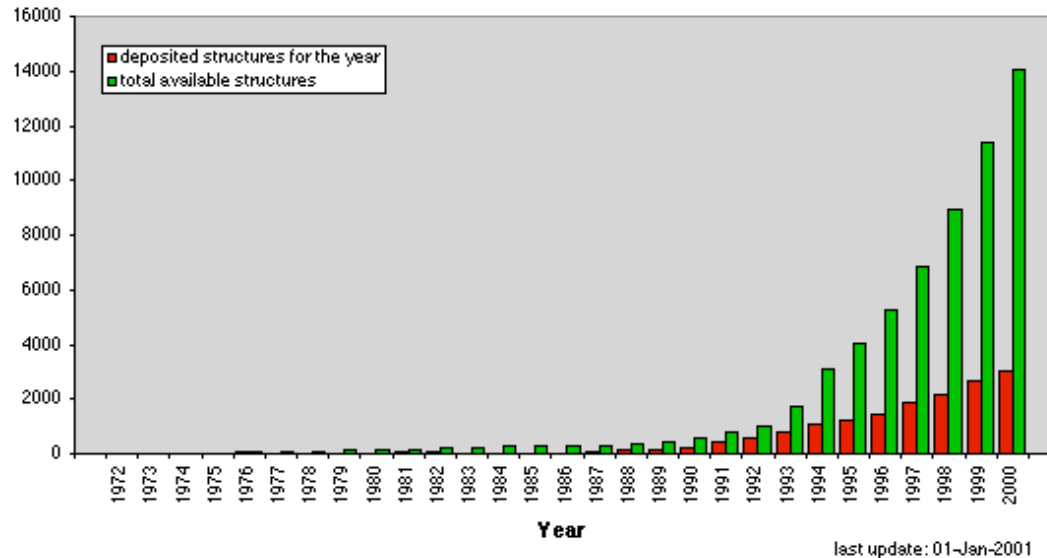


Protein structure

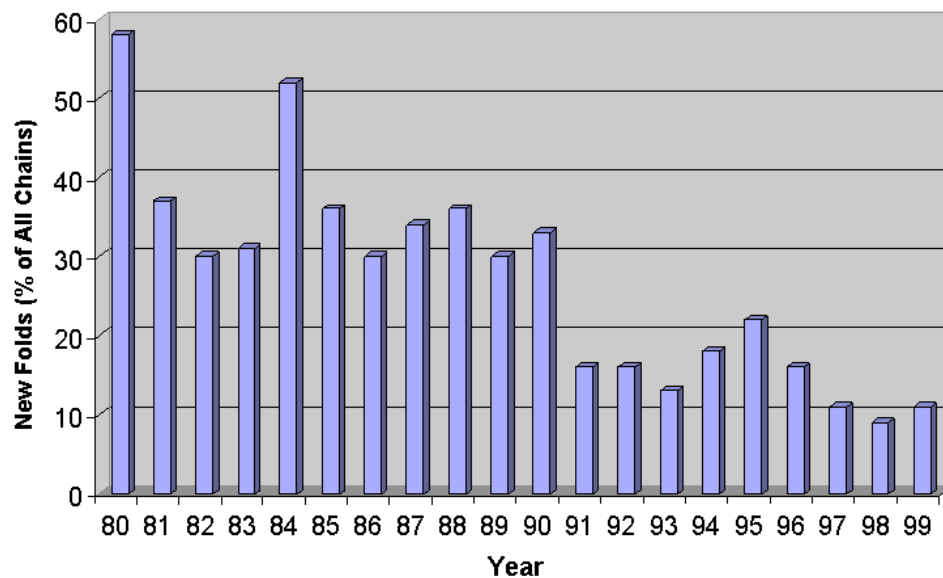
3. Soluble and globular proteins
 5. Principles observed in folds
 6. All α
 7. All β
 8. α/β
 9. $\alpha+\beta$
4. Globular membrane proteins
 1. Definition
 2. Classification
 1. Integral
 2. Periferal

3. Soluble and globular proteins

5. Principles observed in folds



Geometric increase of the number of structures in PDB



The % of new folds has reached a plateau and the tendency will be to decrease until no more new folds are found.

3. Soluble and globular proteins

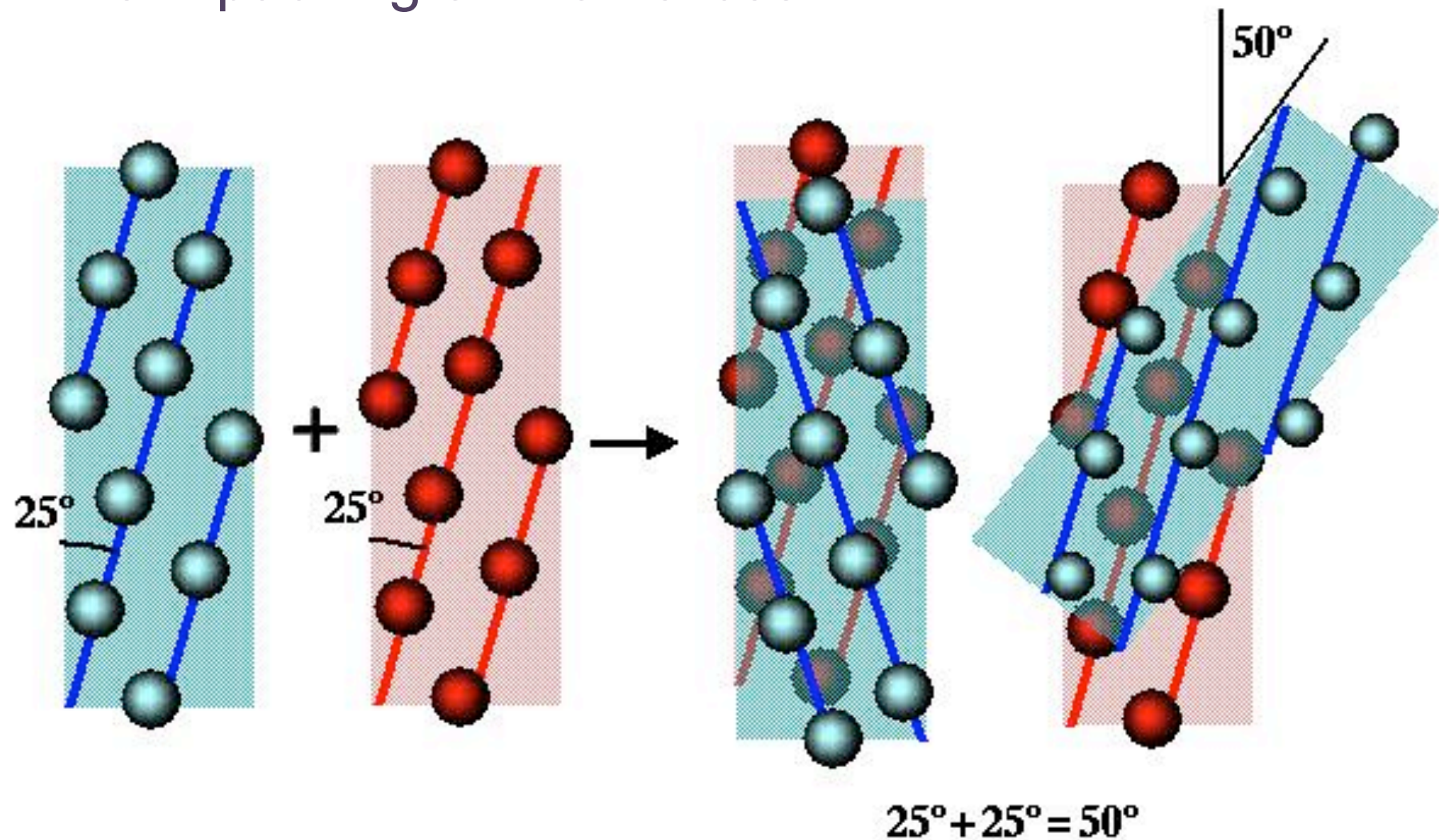
5. Principles observed in folds

- Folds have a compact structure formed by the organization of regular secondary structures
- Folds have an hydrophobic core that stabilizes its conformation
- The hydrophobic core is formed by non-polar residues, while polar residues are mostly in the surface and active sites
- Structure is conserved among proteins with similar sequence
- The total number of folds is finite
- The structure of a protein can be built by joining several folds (this can be produced by gene fusion).

3. Soluble and globular proteins

6. All α

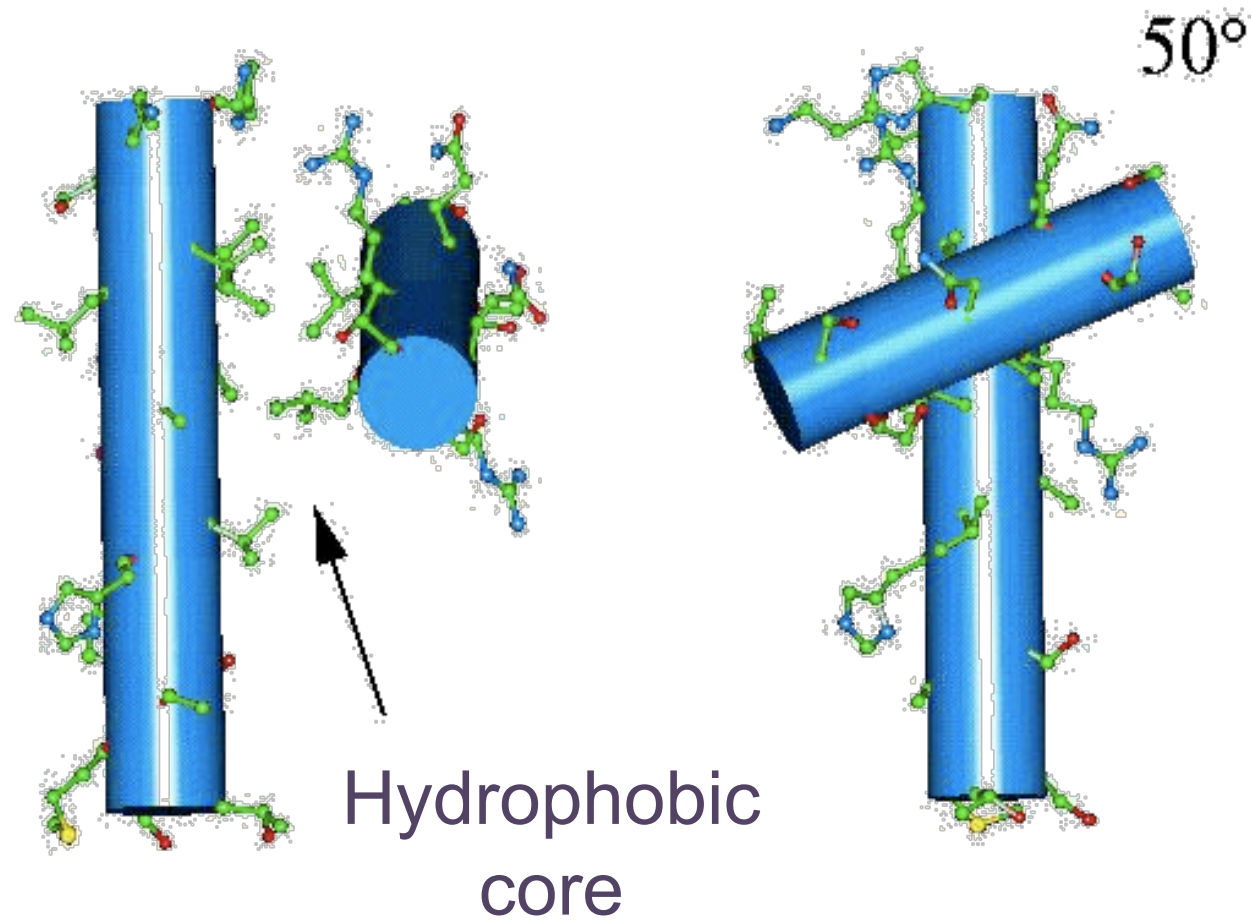
α -helix packing of two helices



3. Soluble and globular proteins

6. All α

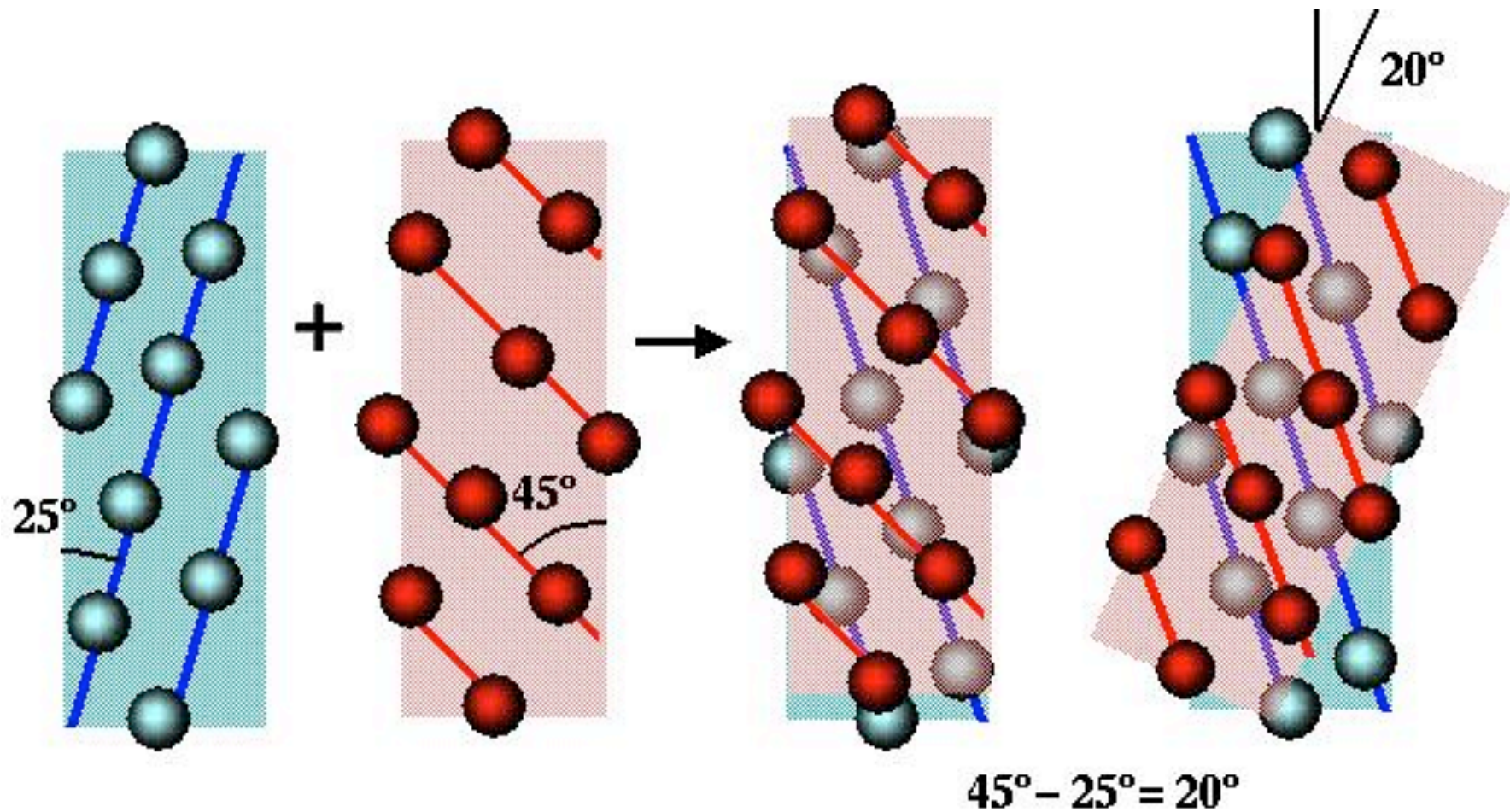
α -helix packing of two helices



3. Soluble and globular proteins

6. All α

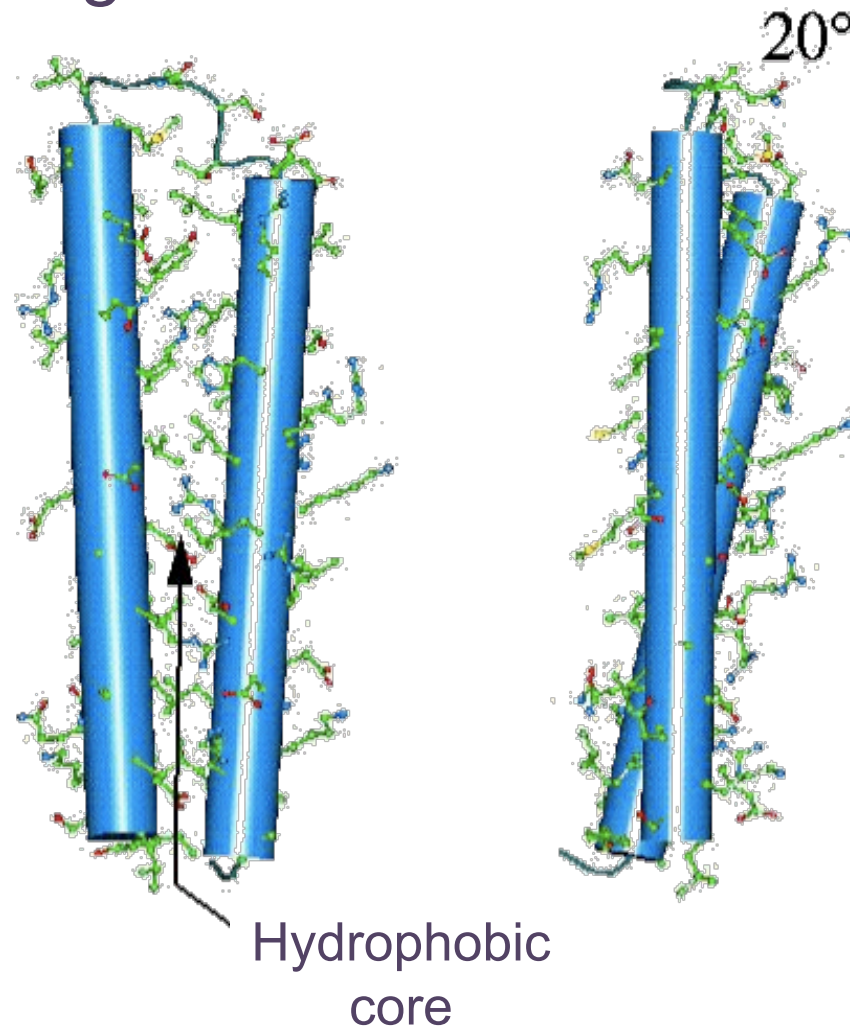
α -helix packing of two helices



3. Soluble and globular proteins

6. All α

α -helix packing of two helices



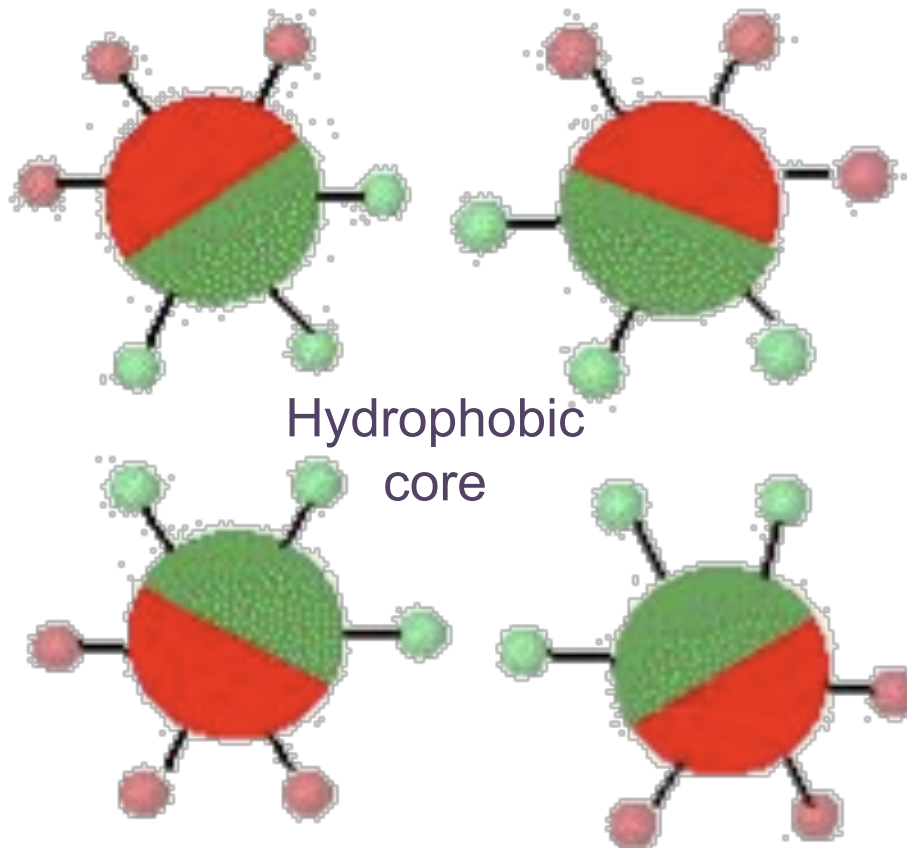
3. Soluble and globular proteins

6. All α

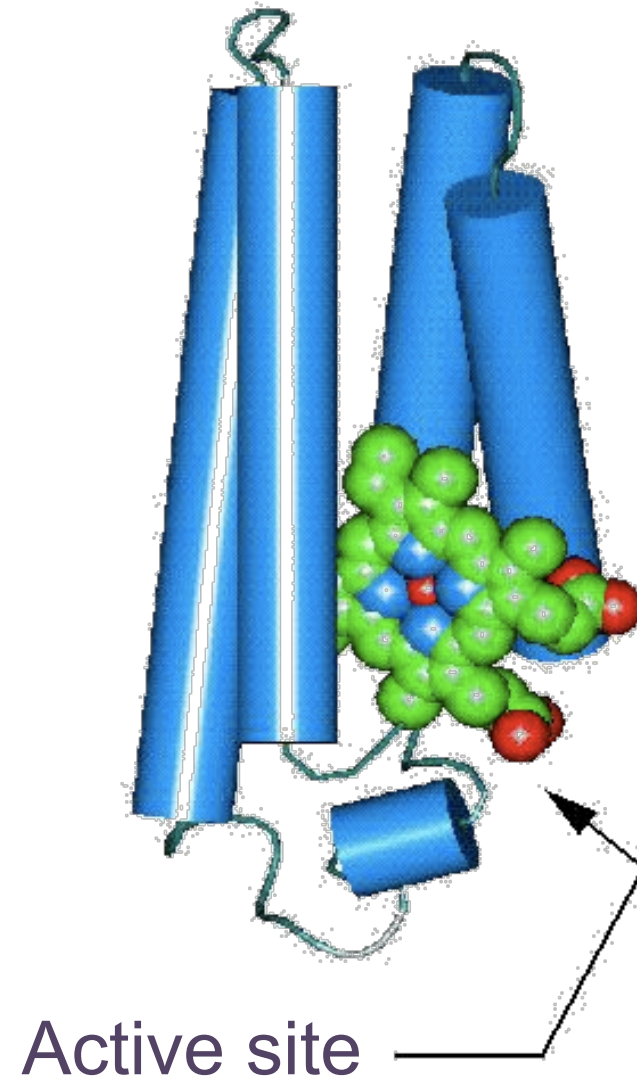
4 helix bundle

Polar residues

Non-polar residues



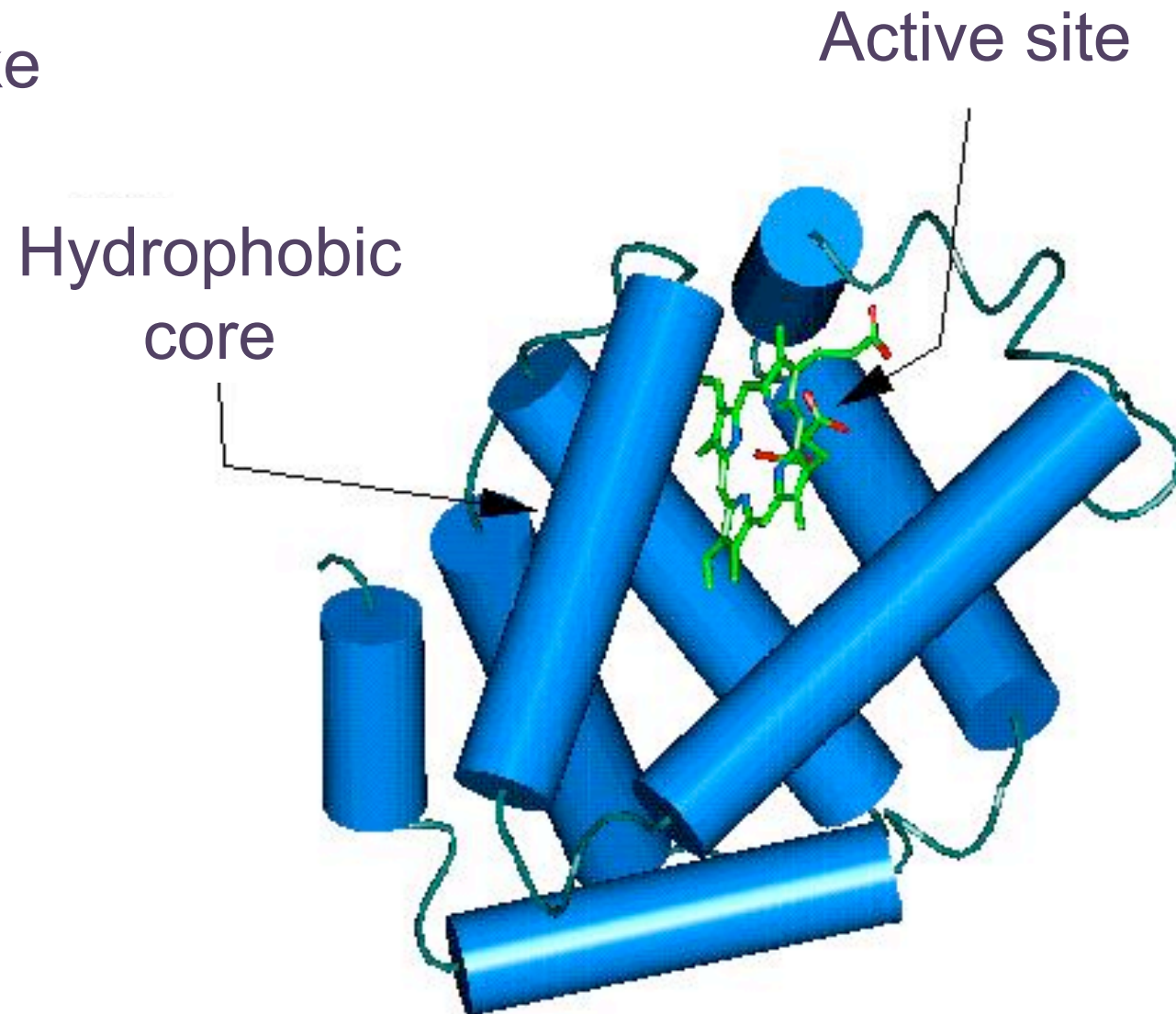
Cytochrome b_{562}



3. Soluble and globular proteins

6. All α

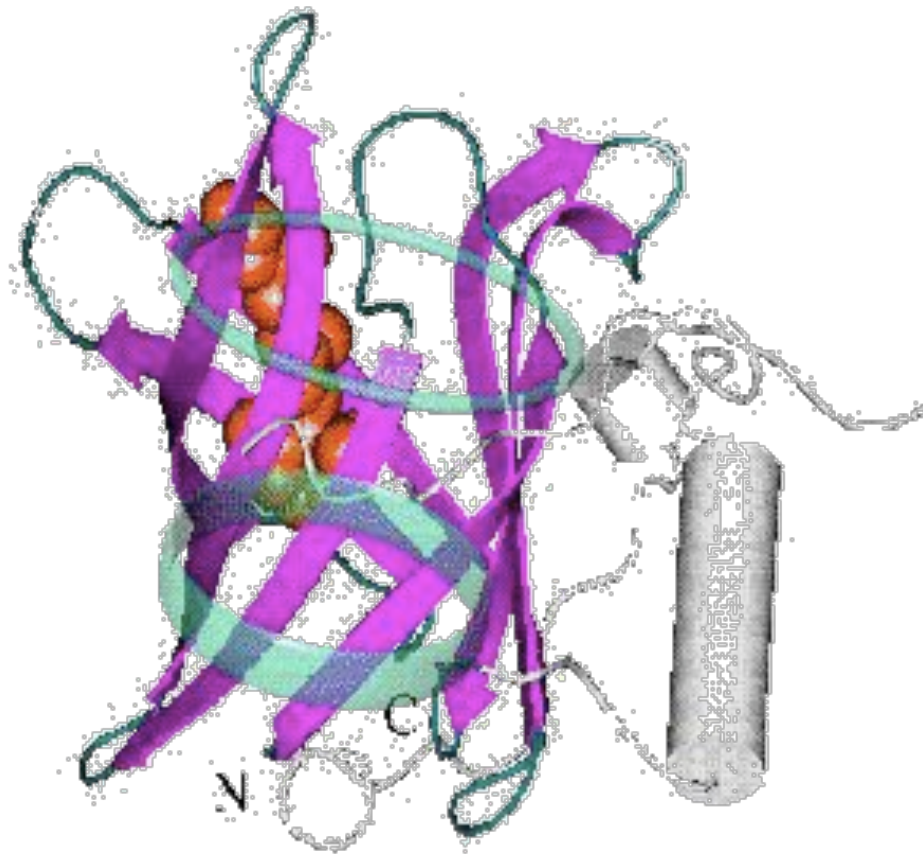
Globin-like



3. Soluble and globular proteins

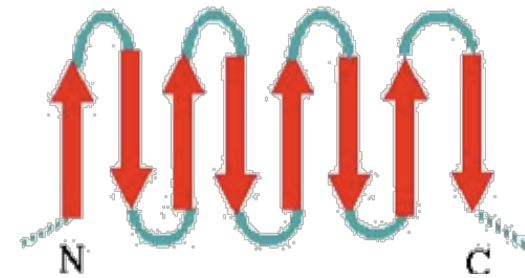
7. All β

Up & Down β -barrel

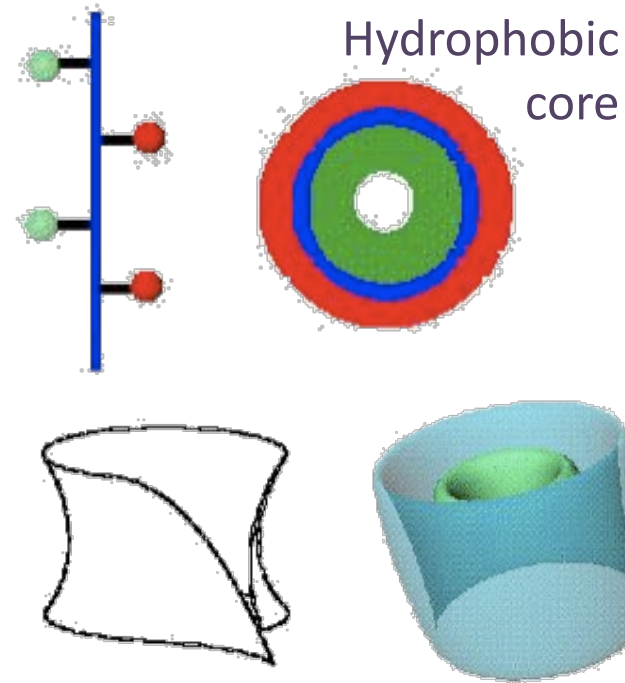


Retinol binding protein is a carrier of retinol (hydrophobic)

Topologic diagram



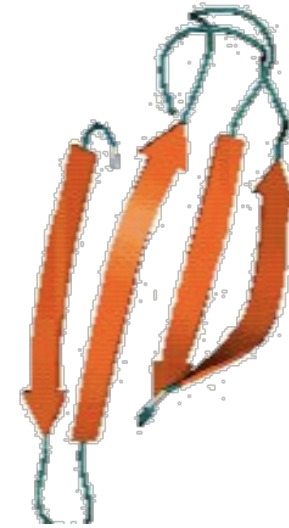
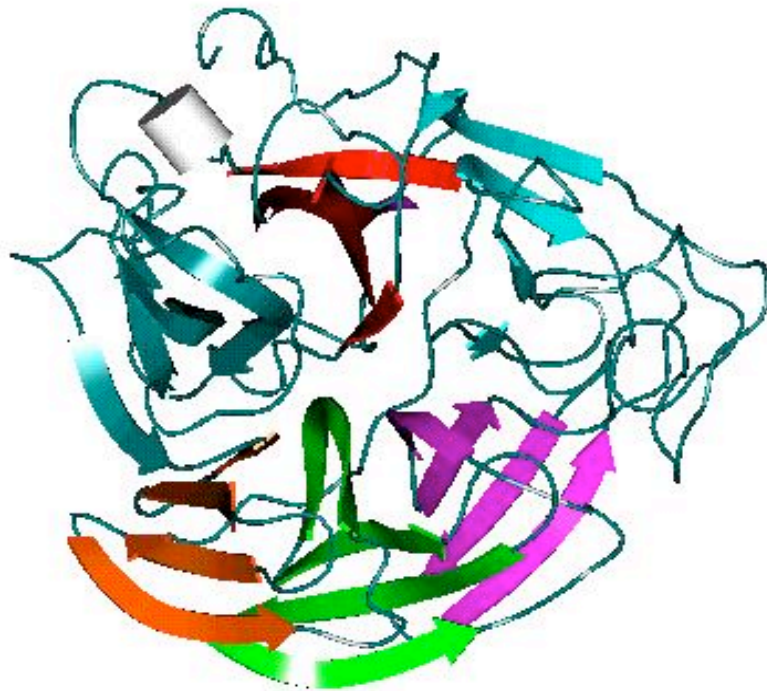
β -meanders are the main super-secondary structure



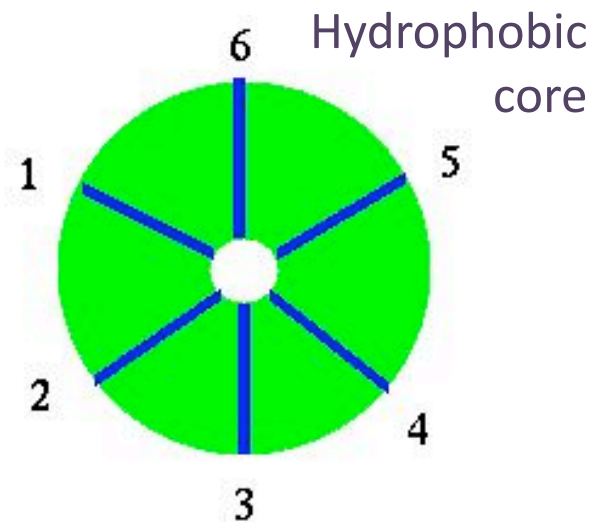
3. Soluble and globular proteins

7. All β

β -propeller (super barrel)



formed by N
 β -meanders
placed as
blades

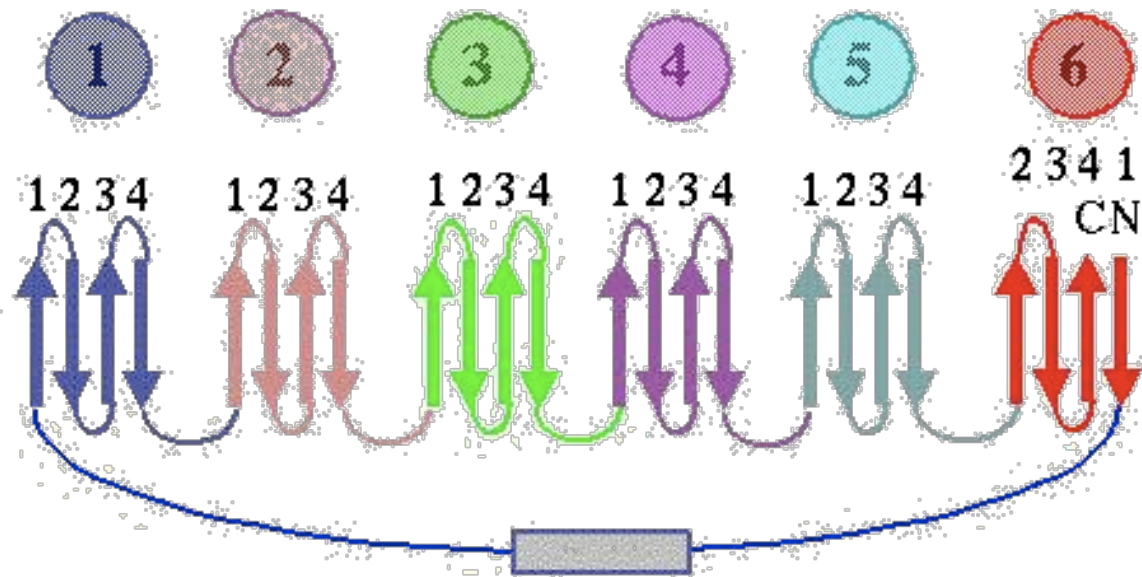


3. Soluble and globular proteins

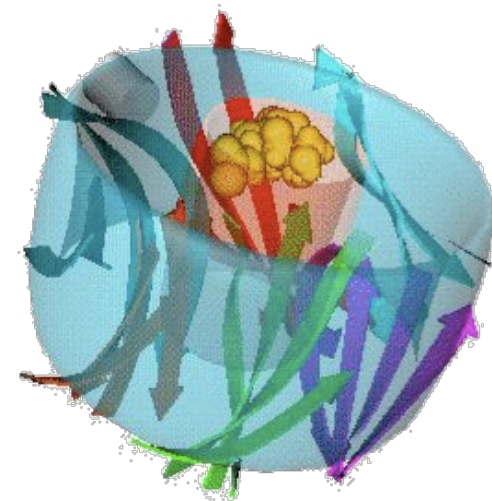
7. All β

β -propeller (super barrel)

Neuraminidase is a 6 β -meander β -propeller



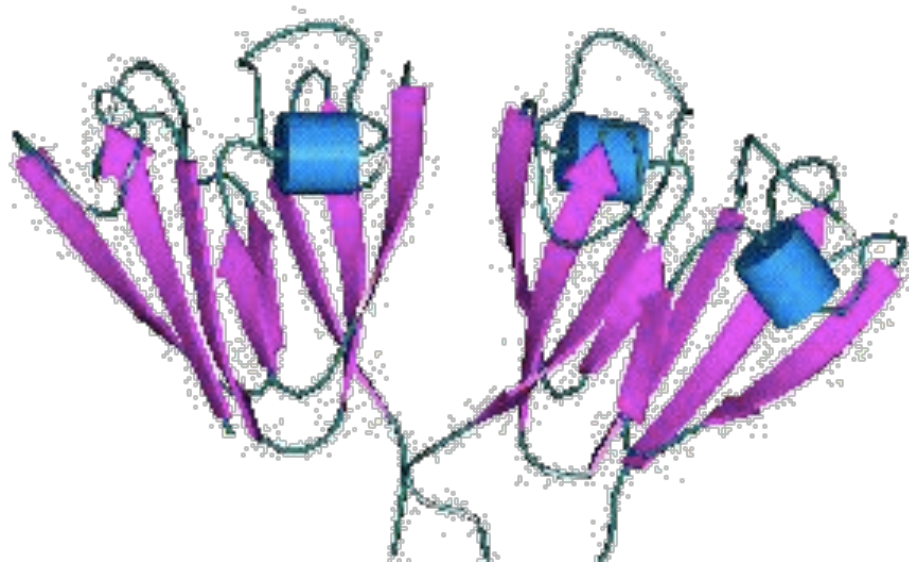
Active site



3. Soluble and globular proteins

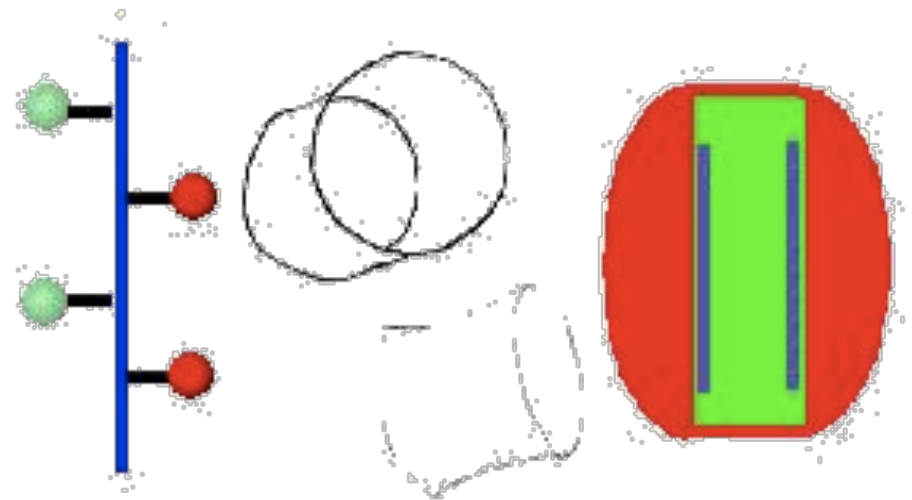
7. All β

Greek key β -sandwich



γ -crystallin

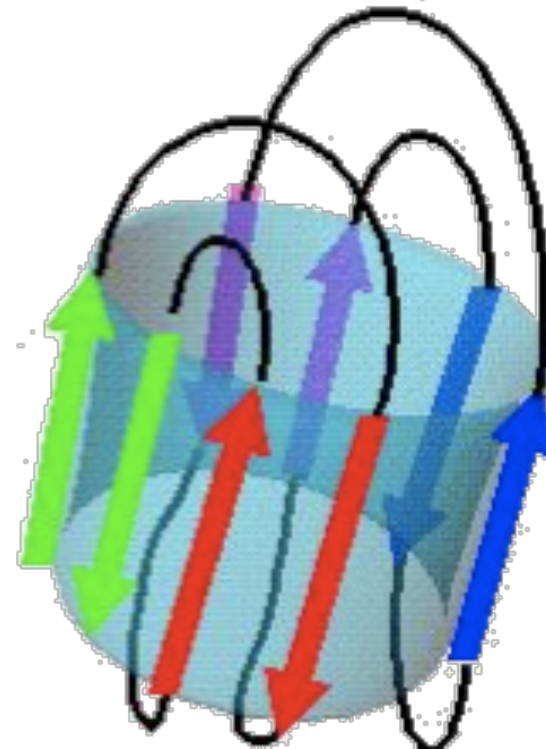
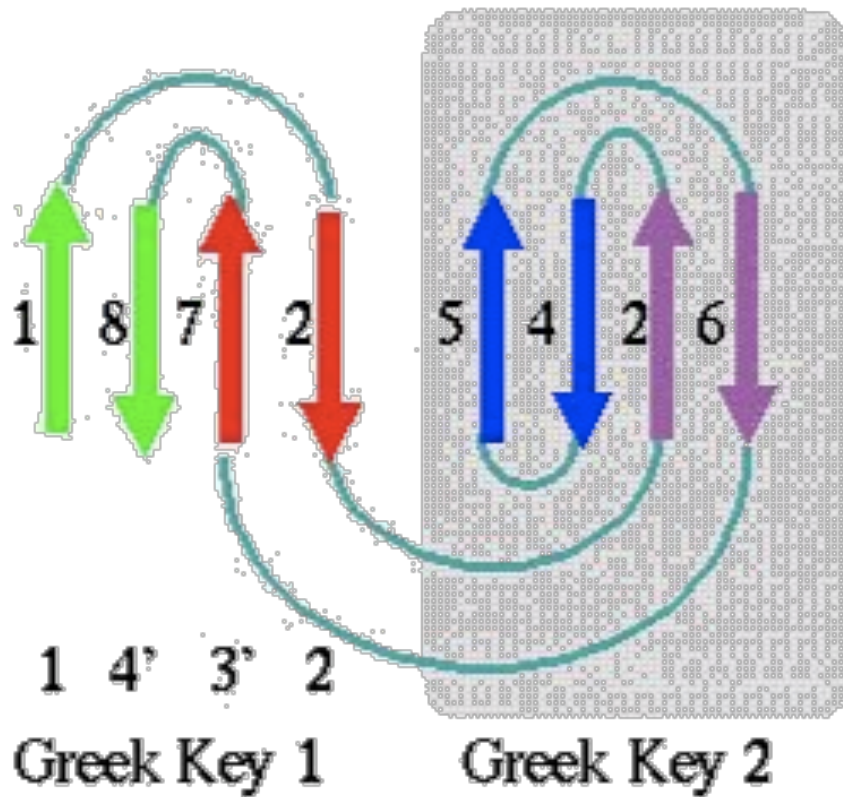
Hydrophobic core



3. Soluble and globular proteins

7. All β

Greek key Jelly-Roll

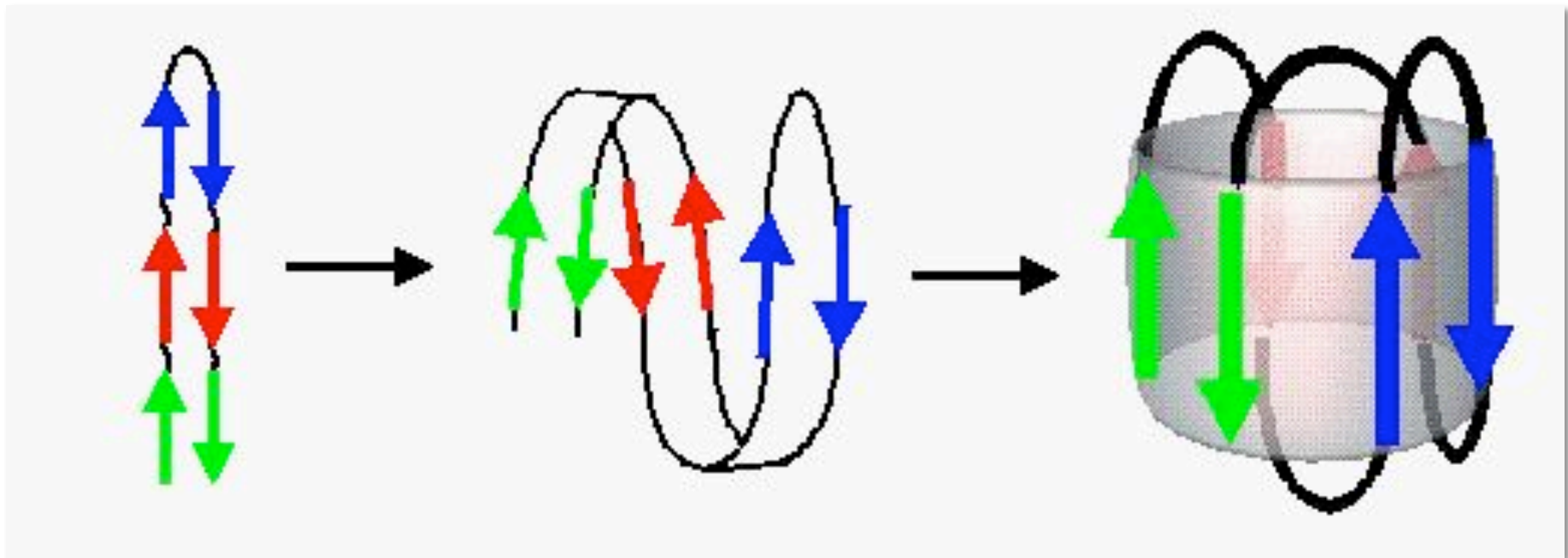


3. Soluble and globular proteins

7. All β

Greek key Jelly-Roll

Proposed mechanism of folding



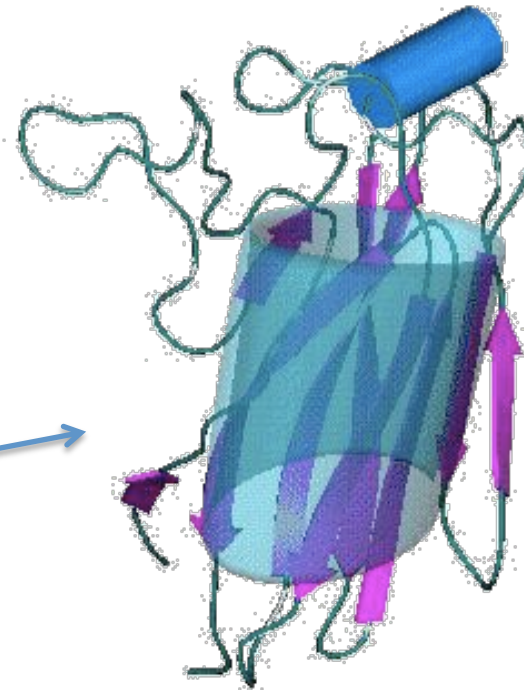
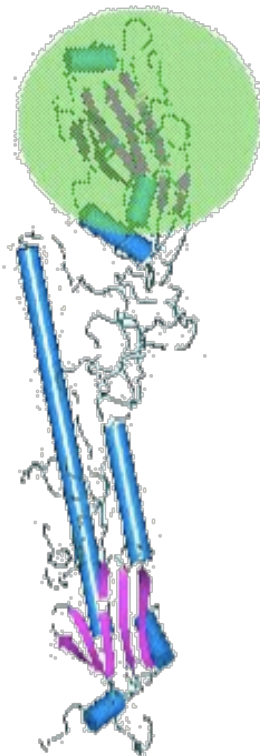
3. Soluble and globular proteins

7. All β

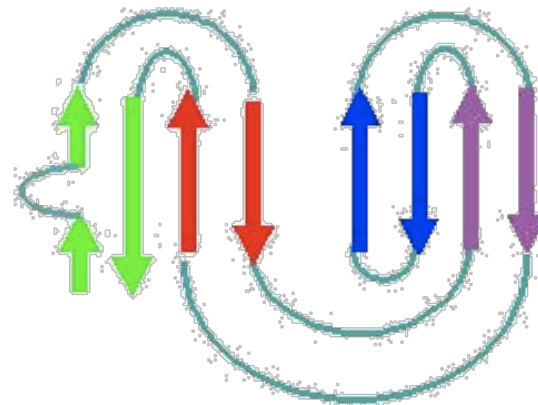
Greek key Jelly-Roll

Hemagglutinin

Trimer of the capsid of the virus of flu. Membrane fusion



Jelly-roll
barrel



Topologic
diagram

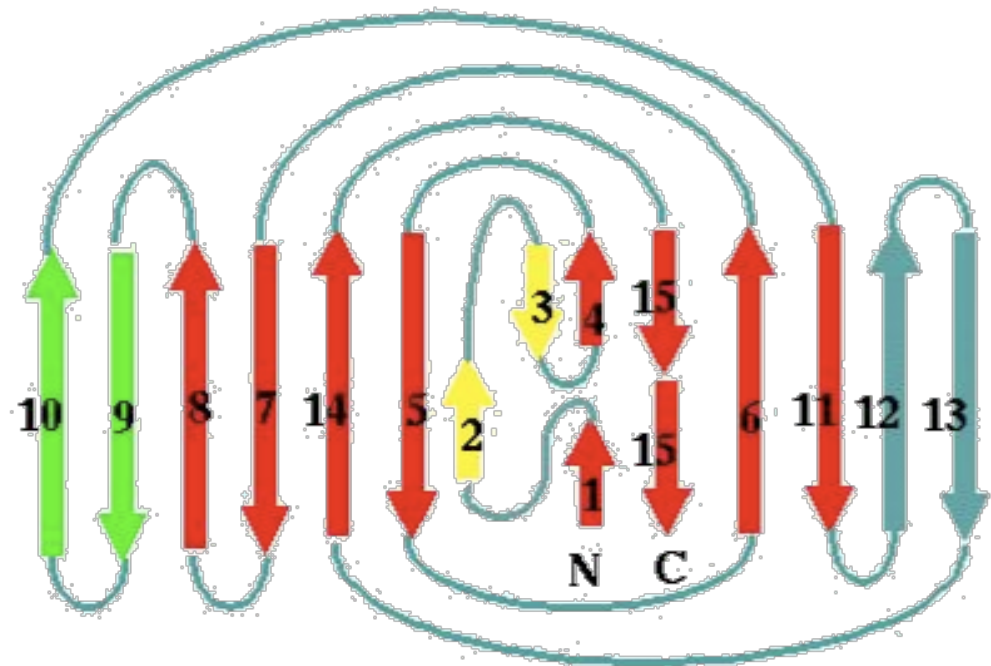
3. Soluble and globular proteins

7. All β

Greek key Jelly-Roll



Glucanase

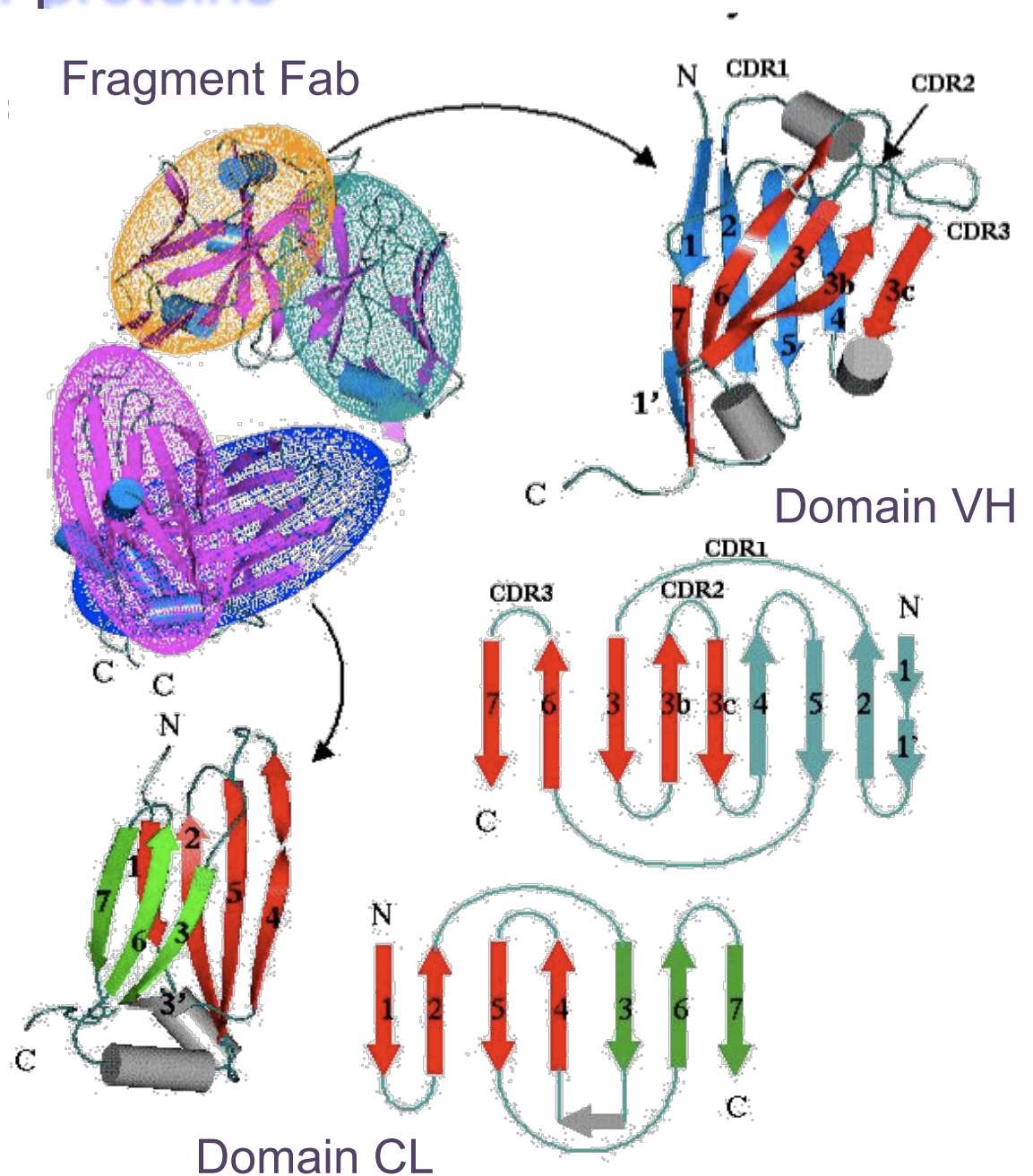
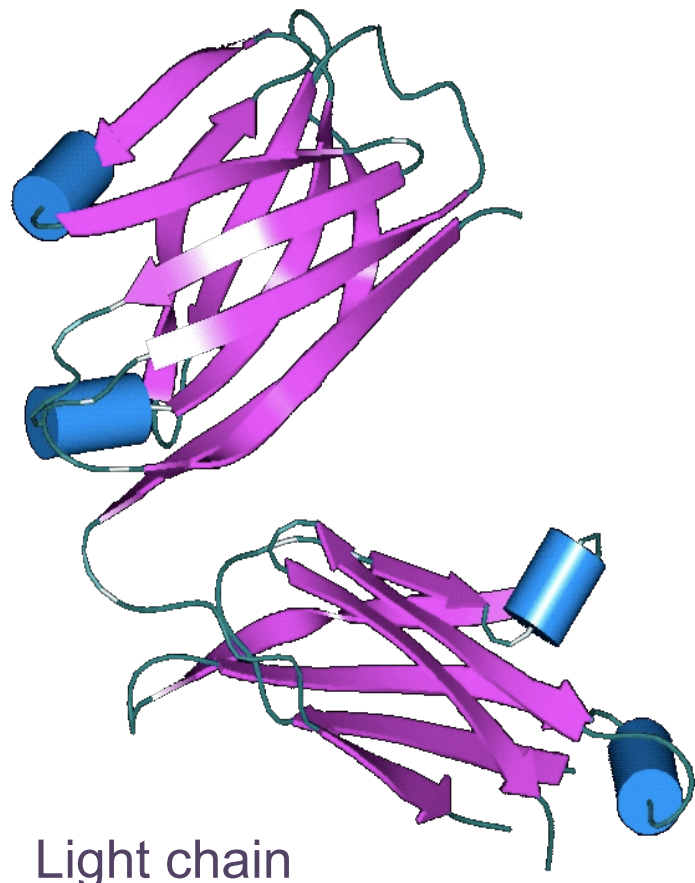


Topologic diagram

3. Soluble and globular proteins

7. All β

β -sandwich
Immunoglobulin-like

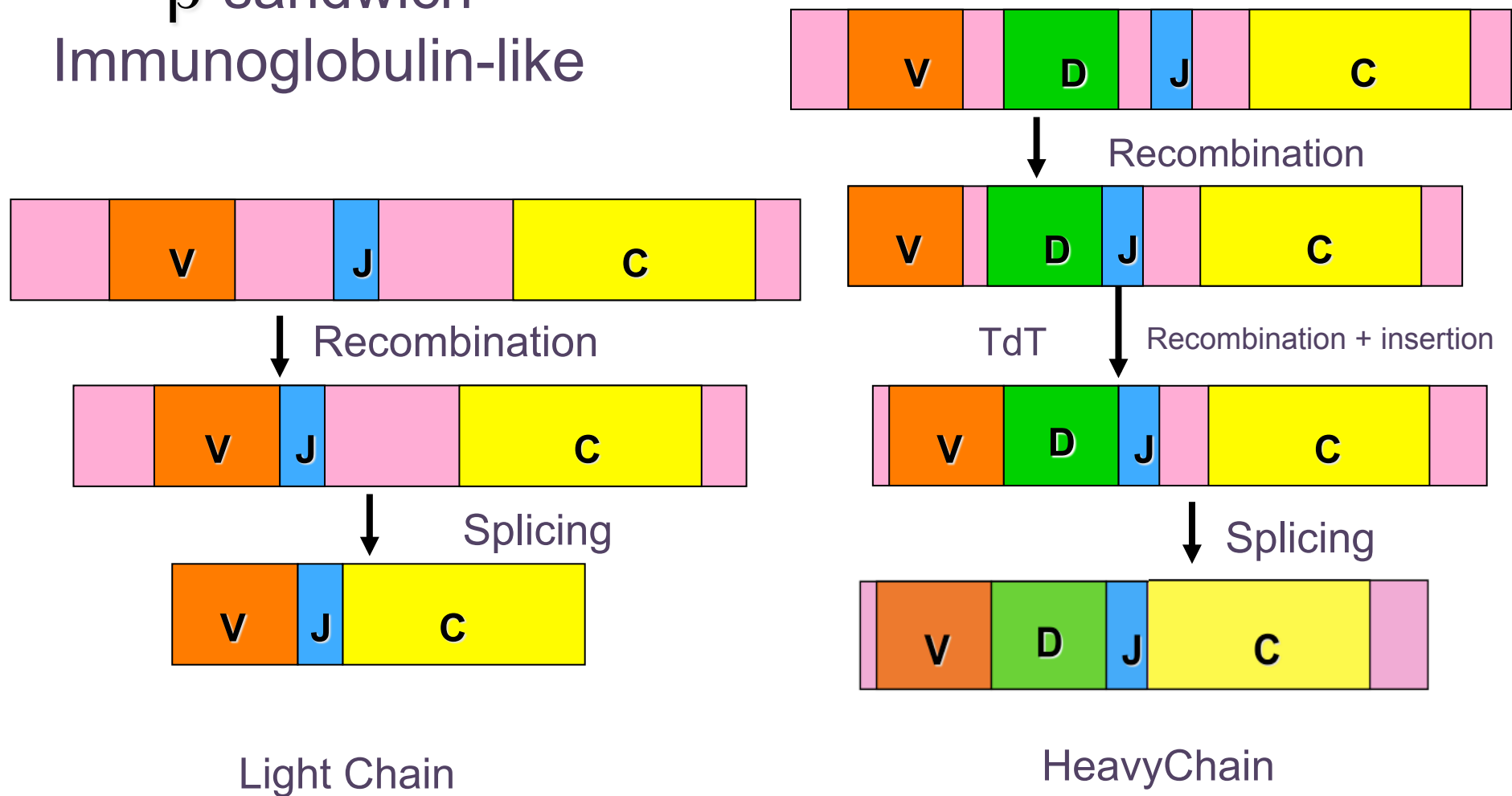


3. Soluble and globular proteins

7. All β

β -sandwich

Immunoglobulin-like

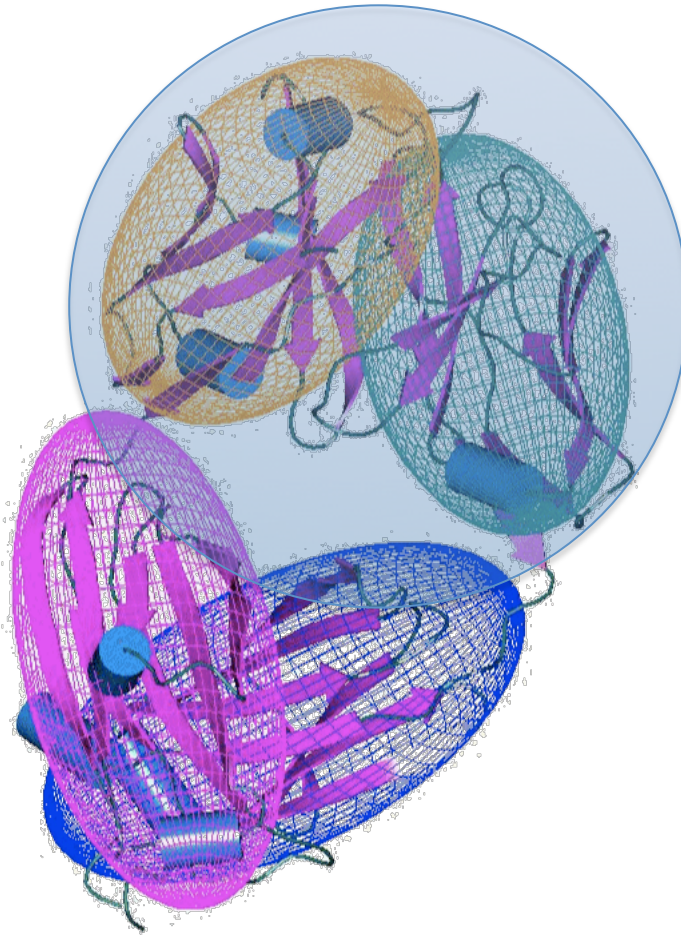


3. Soluble and globular proteins

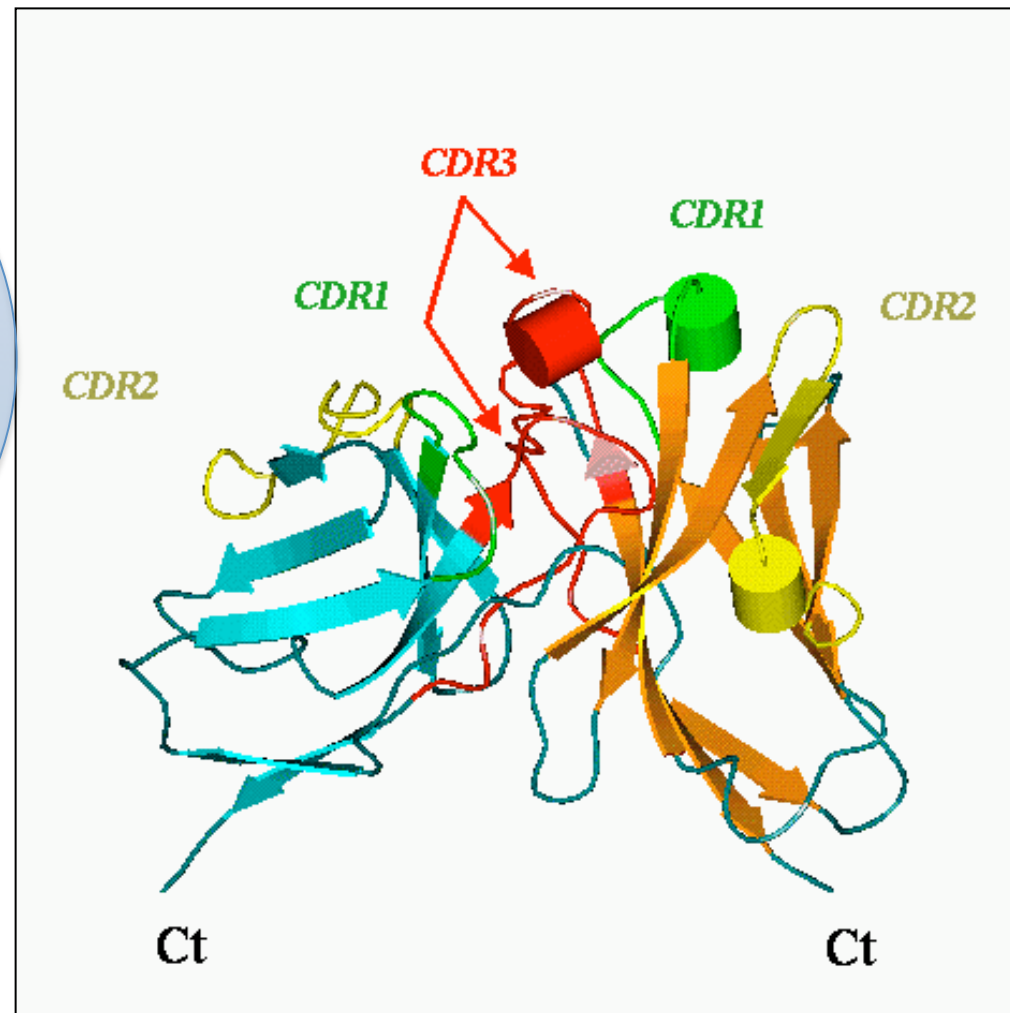
7. All β

β -sandwich

Immunoglobulin-like



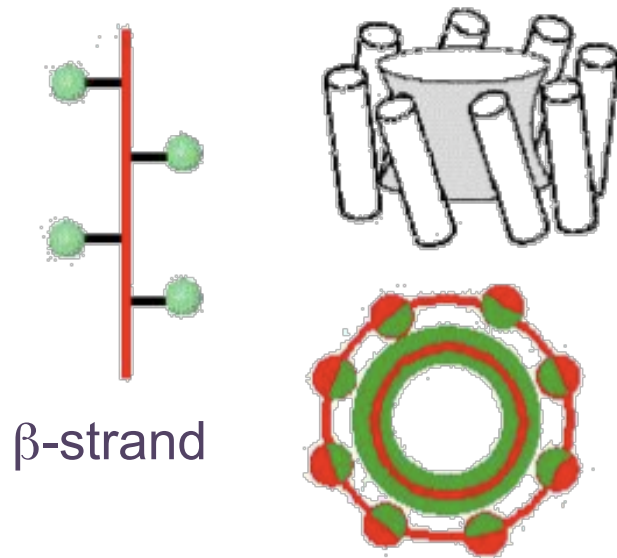
CDR loops



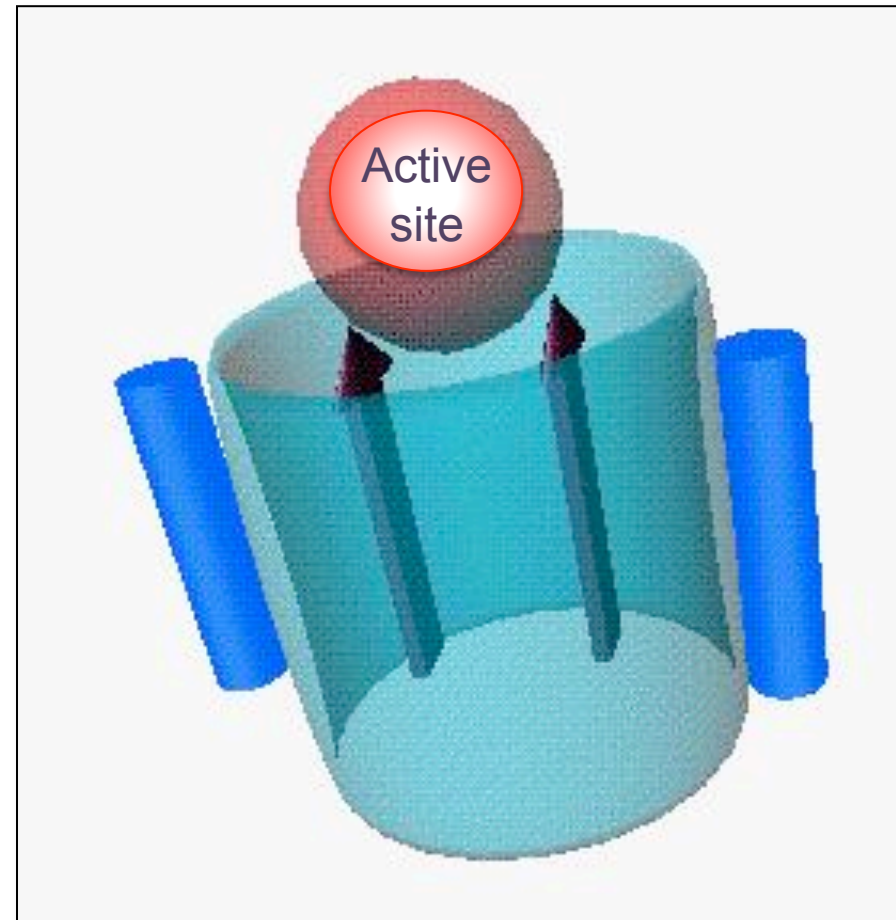
3. Soluble and globular proteins

8. α/β

TIM Barrel



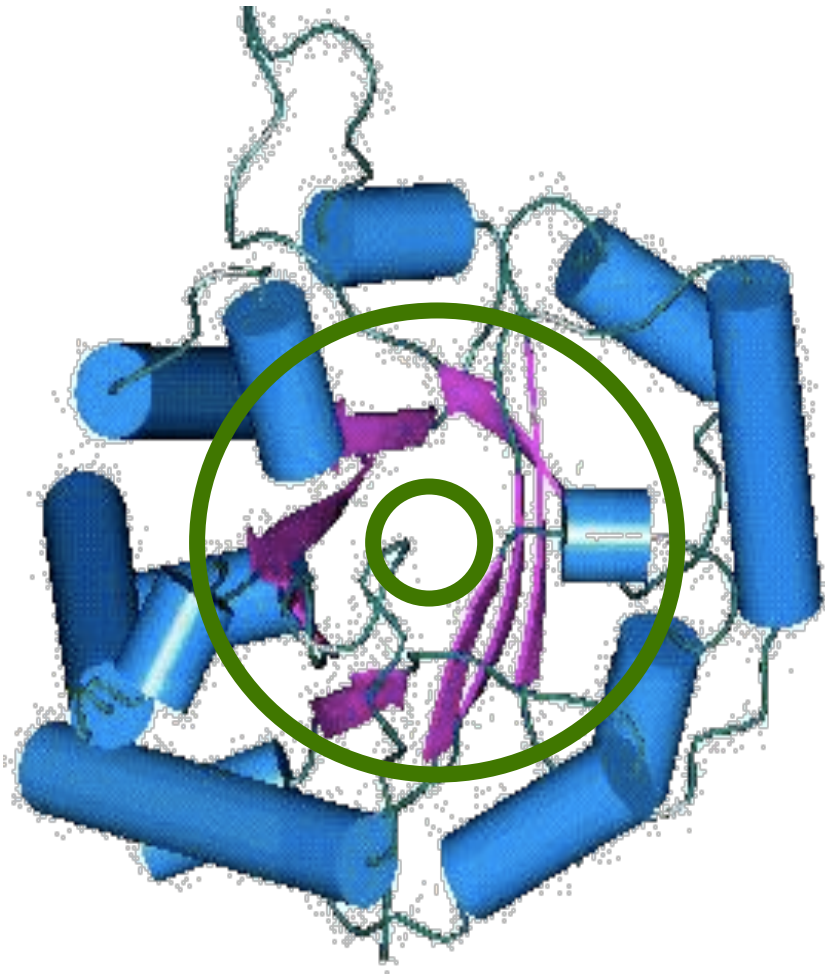
Highly stable structure due to the double hydrophobic core



3. Soluble and globular proteins

8. α/β

Triose Phosphate Isomerase (TIM Barrel)

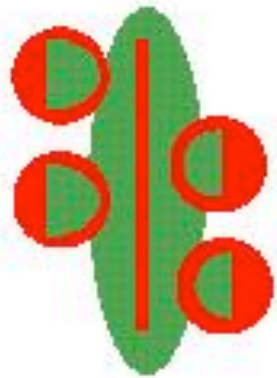


- Double hydrophobic core
- Highly stable
- Found in many proteins, mostly enzymes.
- It was recently proved that all of them have the same ancestor

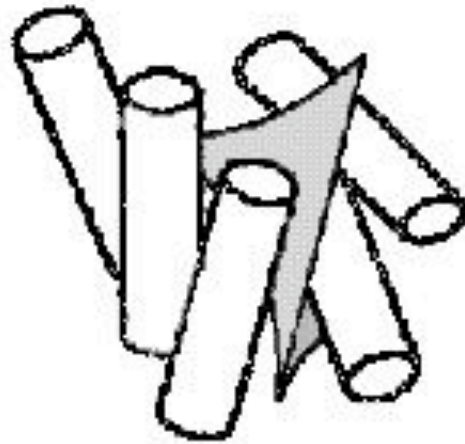
3. Soluble and globular proteins

8. α/β

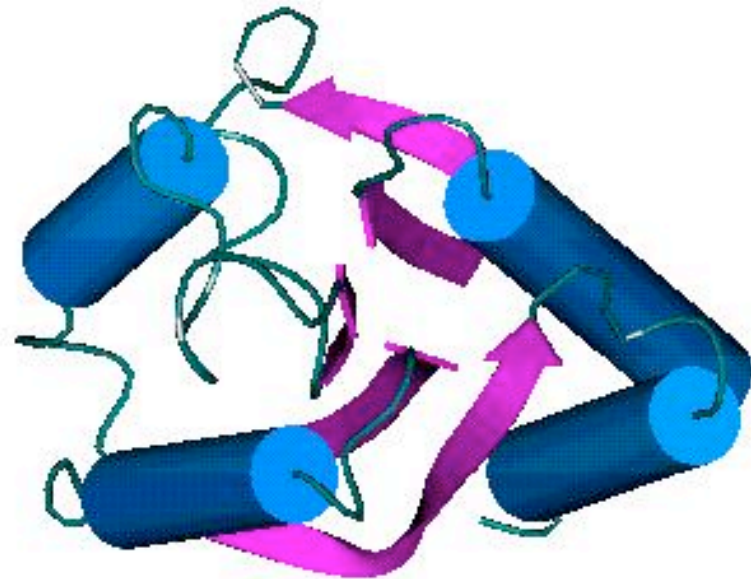
Rossmann Fold



Open sheet

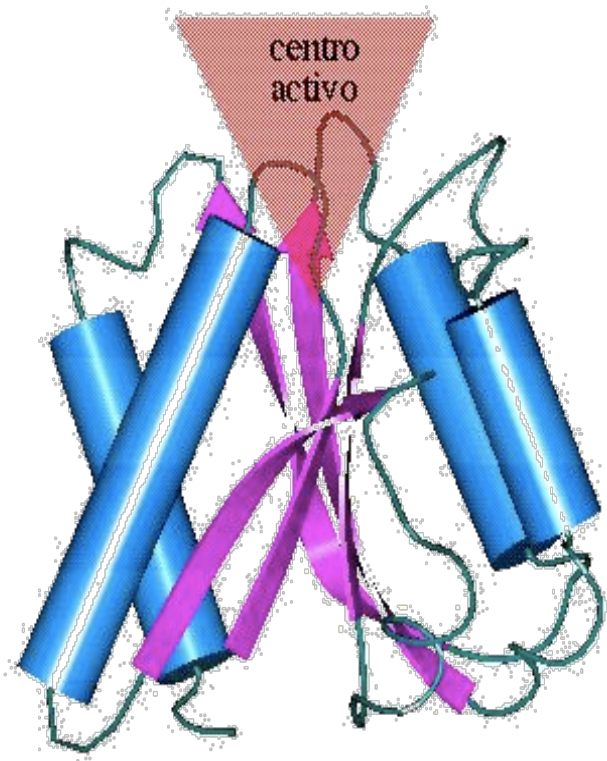


Flavodoxin



3. Soluble and globular proteins

Rossmann Fold



Often found as binding domain of mono-nucleotides (NAD, FAD, FMN, etc.).

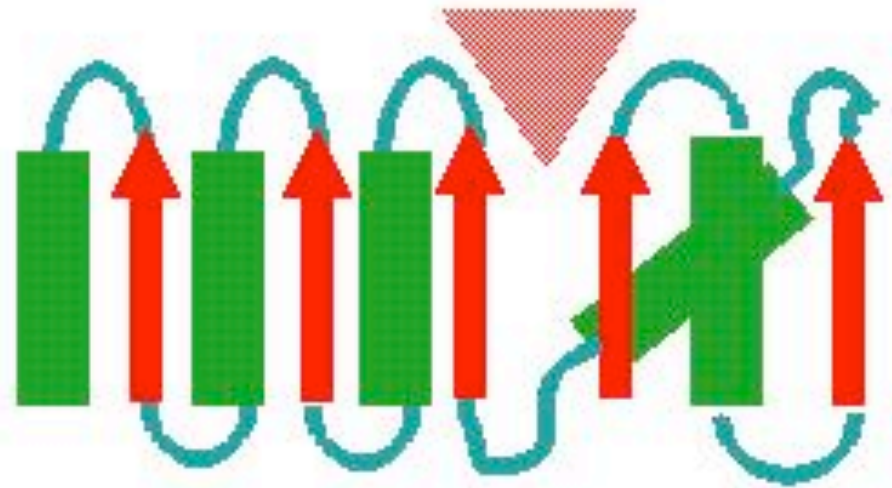
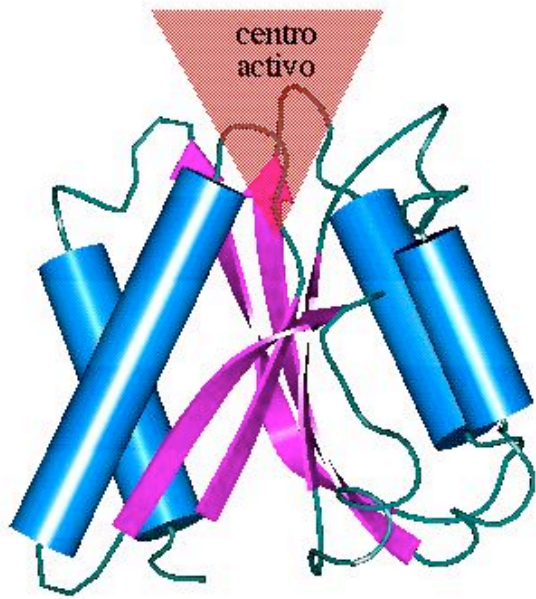
- Found in 1974 by Rossmann
- This domain appears in many proteins
- **Hypothesis:** many proteins have incorporated this domain by gene fusion, changing the activity by its interaction with mononucleotides

Rossmann Fold of Flavodoxin

3. Soluble and globular proteins

Rossmann Fold

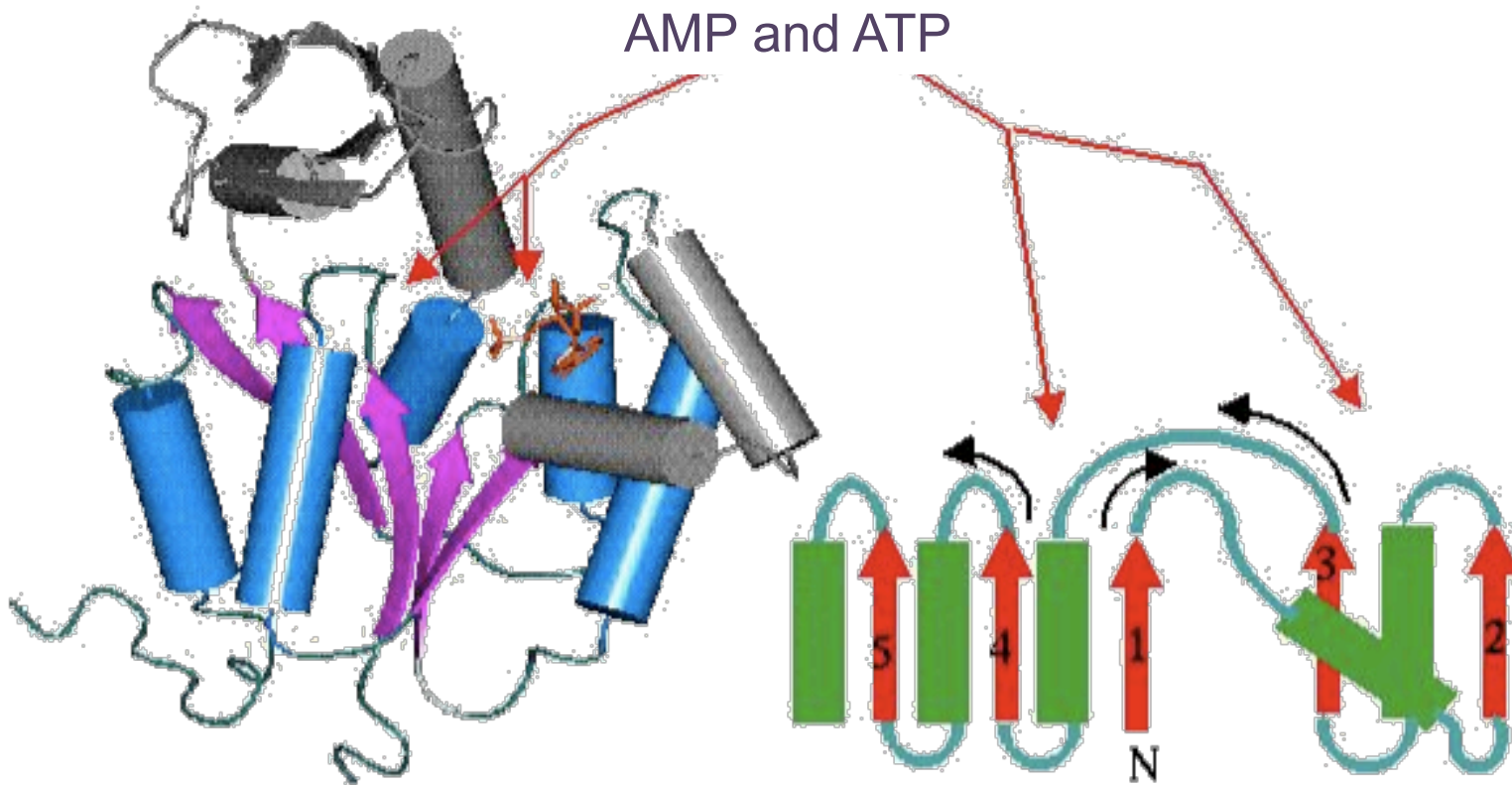
- The active site is located in a $\beta\alpha$ motif of the type NADP/FAD binding loop
- The location of the loop was often found in the change of orientation of the **topological diagram**. This is named **topological switch point**



3. Soluble and globular proteins

Rossmann Fold

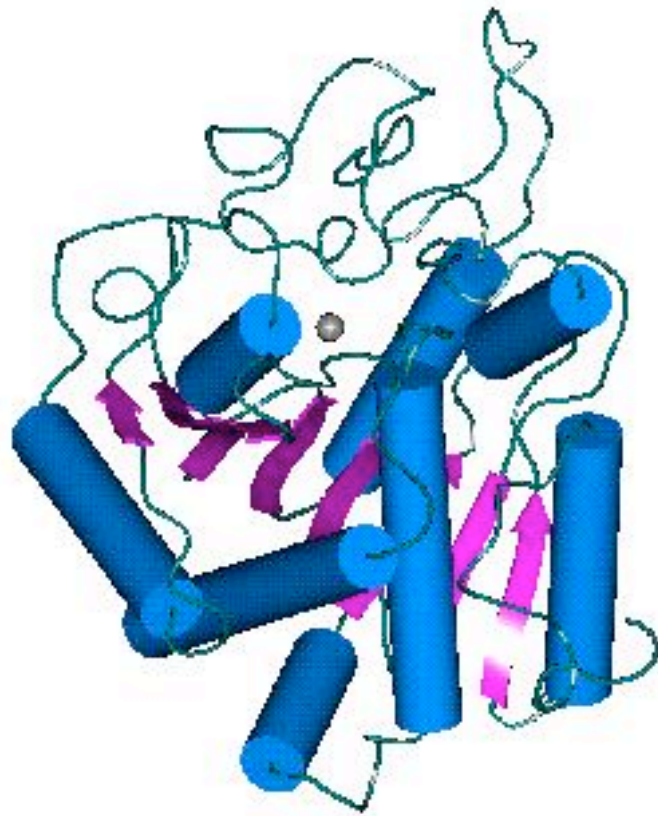
Active site has two
binding sites for
AMP and ATP



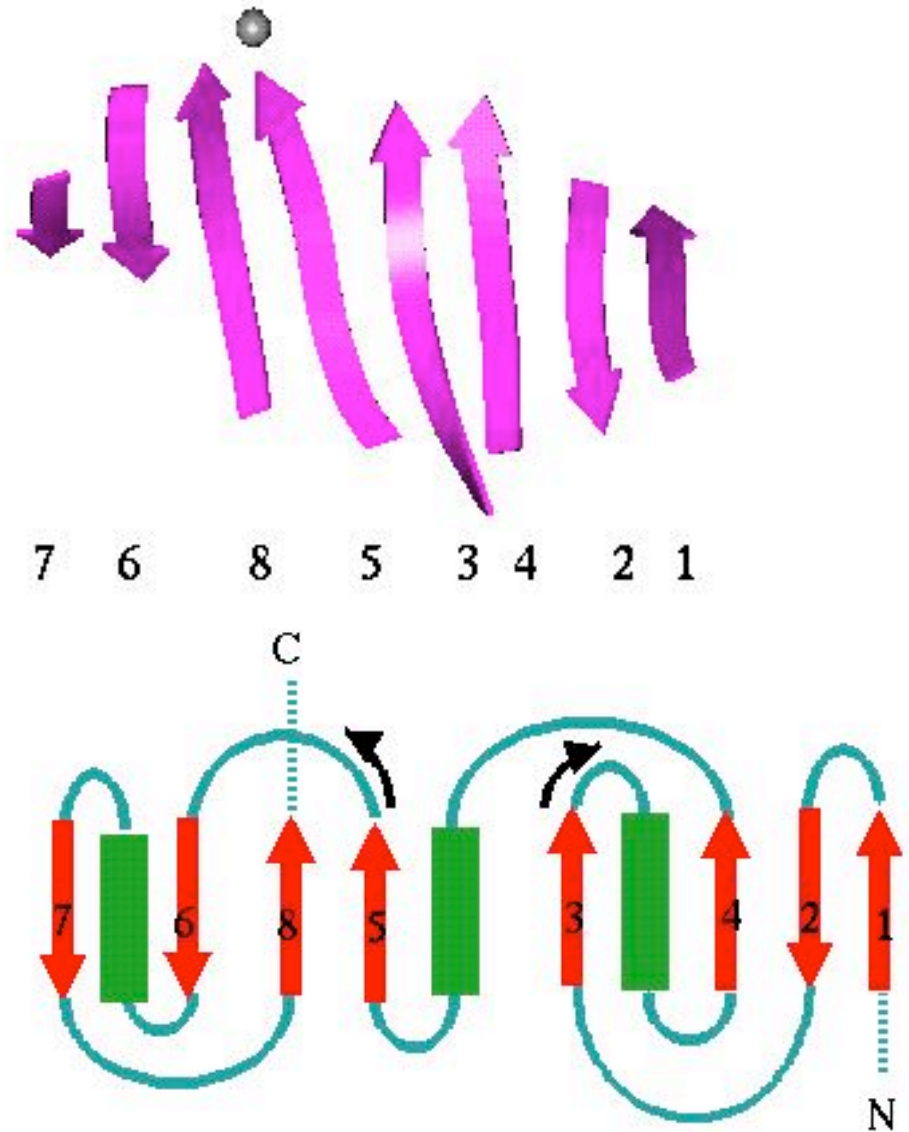
Adenylate Kinase

3. Soluble and globular proteins

Rossmann Fold



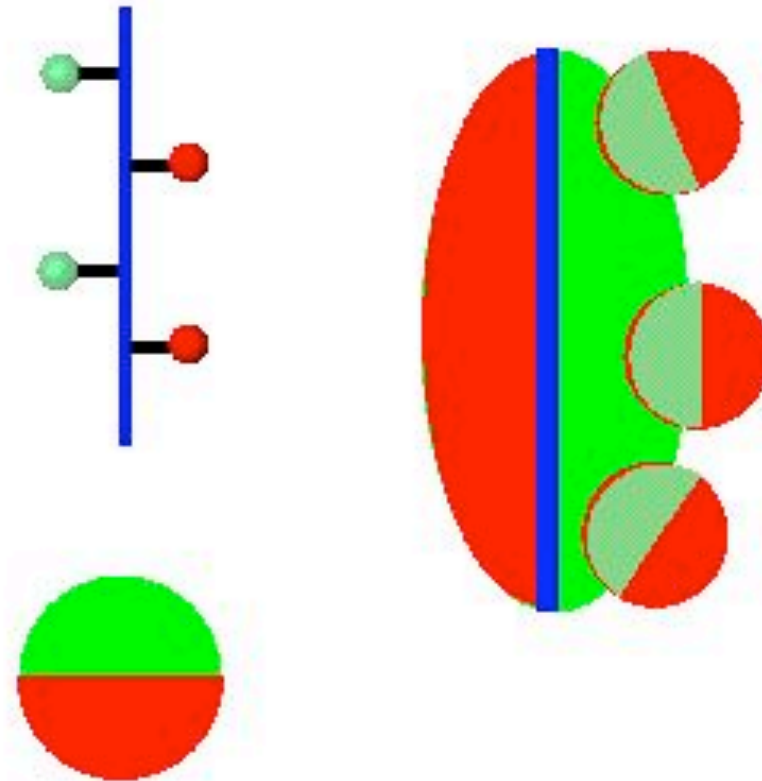
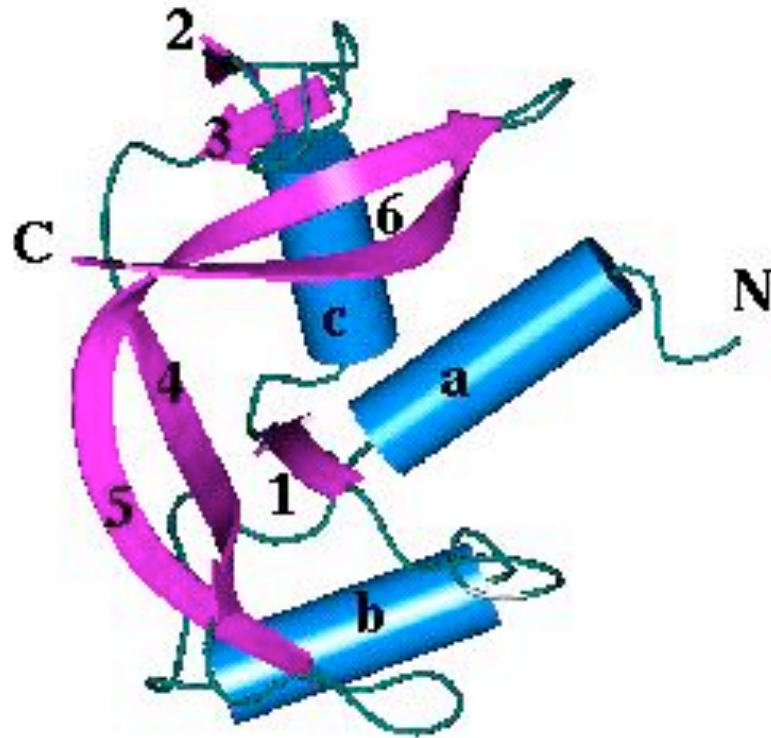
Carboxypeptidase



3. Soluble and globular proteins

9. $\alpha+\beta$

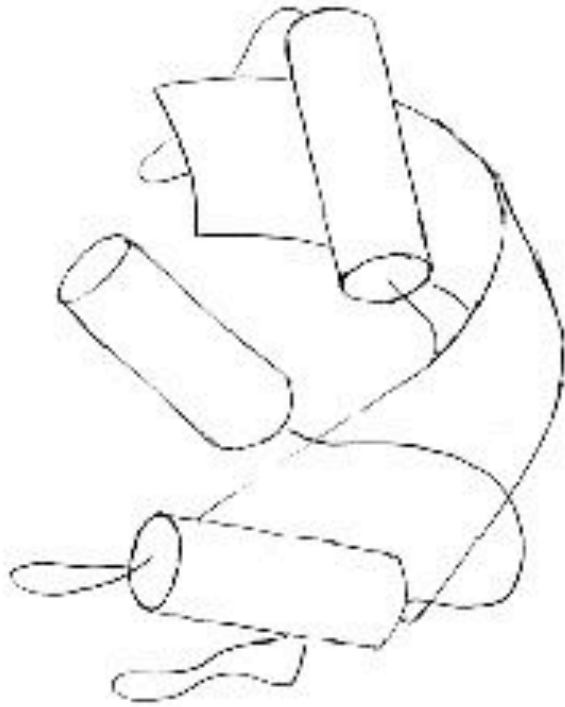
Ribonuclease



3. Soluble and globular proteins

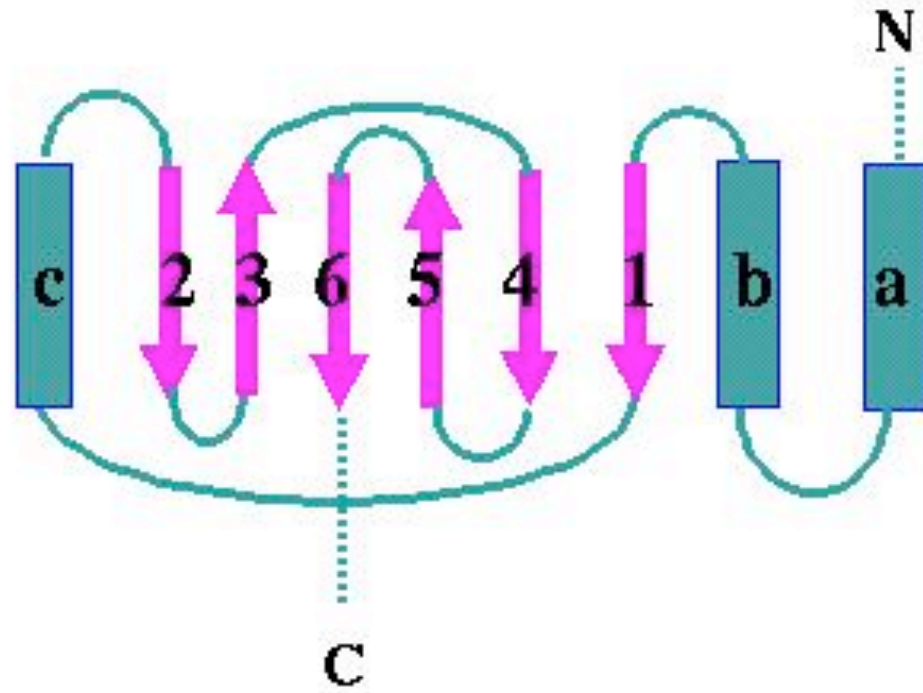
9. $\alpha+\beta$

Ribonuclease



RNase A-like fold

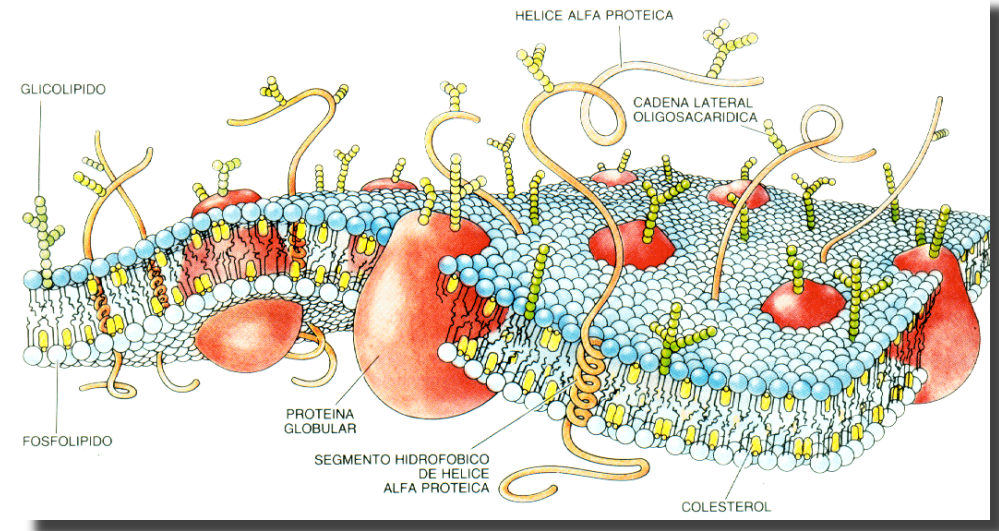
$\alpha+\beta$ class segregates α and β regular secondary structures in 3D and in the topologic diagram



4. Globular membrane proteins

1. Definition

- Membrane proteins are proteins bound to the membrane
- Membrane Proteins can be of two types: **peripheral and integral**
- Integral membrane proteins are inserted in the membrane, where the main bulk of the domain is buried inside the membrane
- Peripheral membrane proteins are only linked by few residues to the membrane
- The residues of the integral proteins are mostly non-polar, changing the rules of hydrophobicity patterns described for soluble domains.



4. Globular membrane proteins

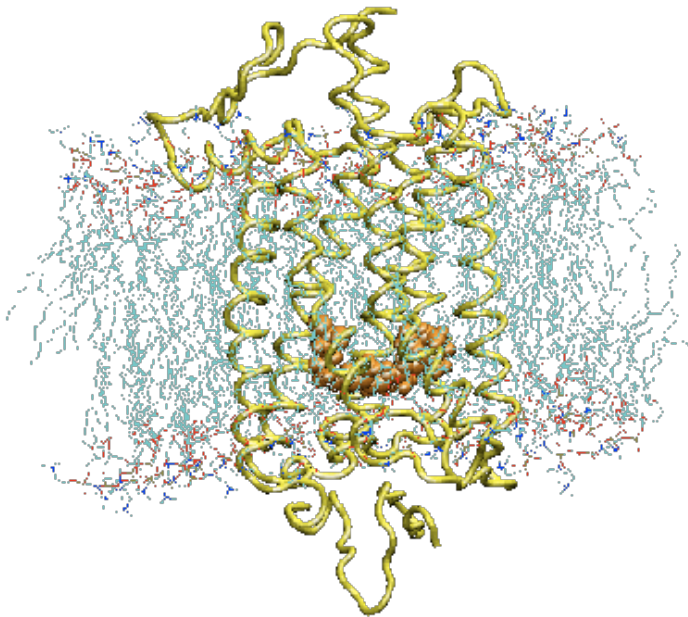
2. Classification

1. Integral

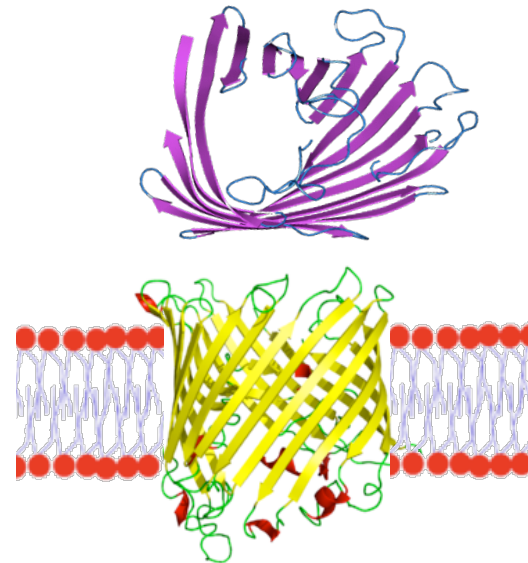
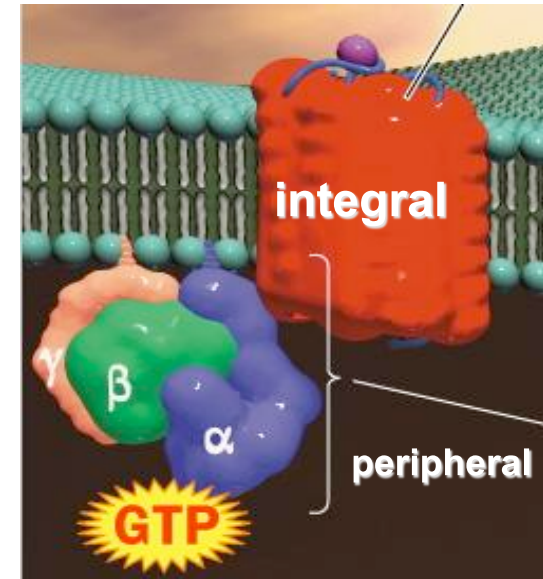
Two types of integral proteins:

α -helix transmembrane

β -sheet transmembrane



α -helix transmembrane
Rhodopsin



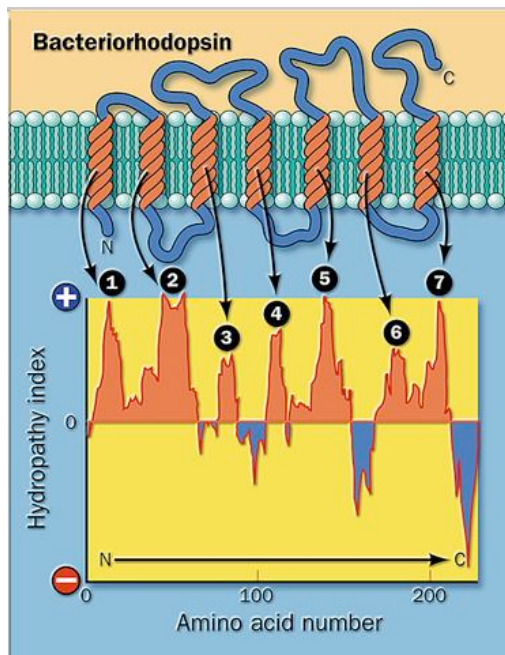
β -sheet transmembrane
Porin

4. Globular membrane proteins

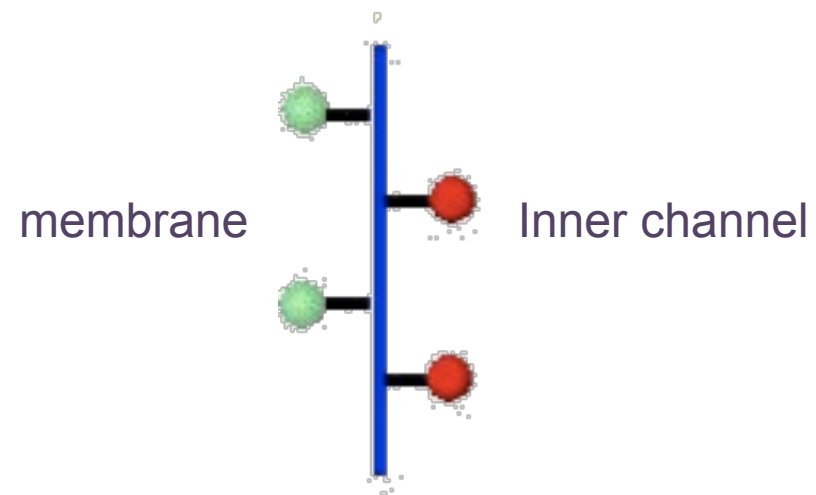
2. Classification

1. Integral

Hydrophobic properties



α -helix transmembrane
Rhodopsine



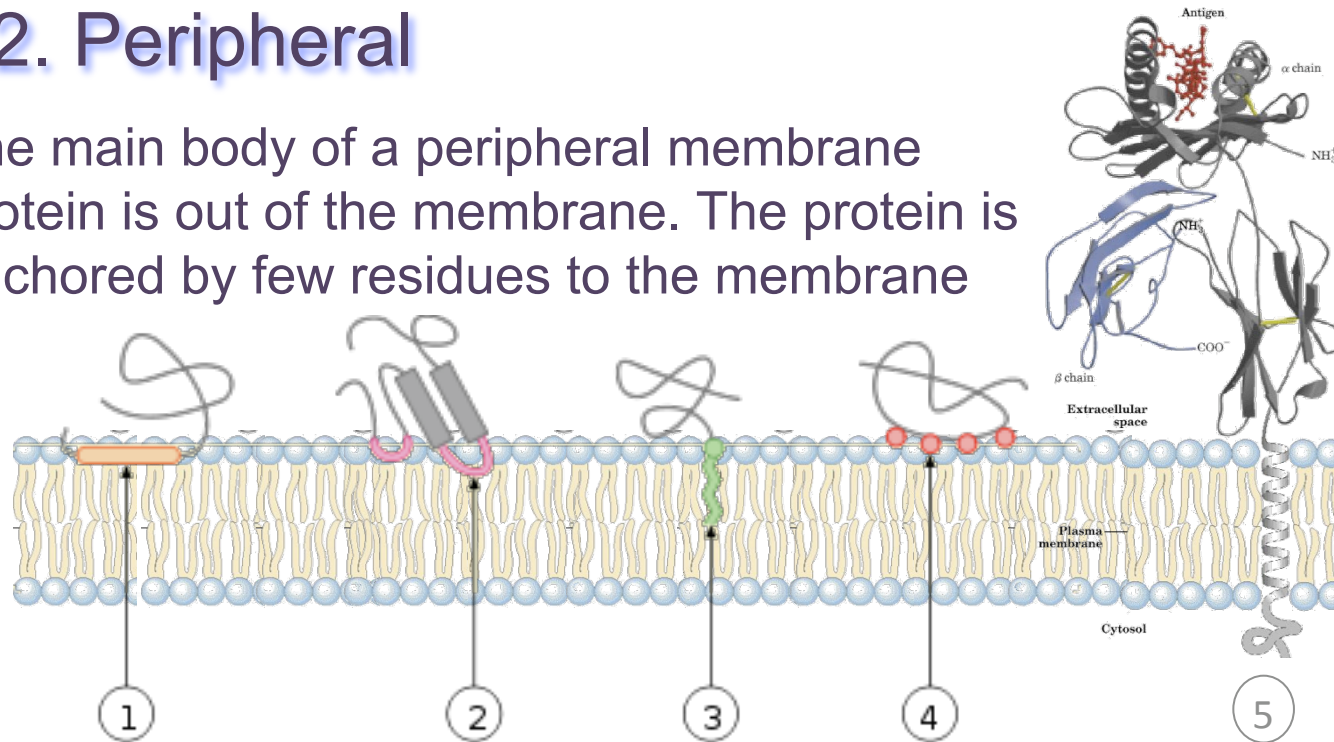
β -sheet transmembrane
Porin

4. Globular membrane proteins

2. Classification

2. Peripheral

The main body of a peripheral membrane protein is out of the membrane. The protein is anchored by few residues to the membrane



Example of different types of interaction between membrane proteins and the cell membrane: 1. interaction by an **amphipathic helix** parallel to the membrane plane 2. interaction by a **hydrophobic loop** 3. interaction by a covalently bound membrane lipid (**lipidation**, i.e. myristilation) 4. **electrostatic** or ionic interaction with membrane lipids (e.g. through a calcium ion) 5. **hydrophobic helix** transmembrane.

4. Globular membrane proteins

2. Classification

2. Peripheral

Some proteins have two main bodies one at each side of the membrane, and in general its function is to transfer a biochemical signal from one side to the other

