

Macromolecules

Bioinformatics for Systems and Synthetic Biology

Emidio Capriotti

<http://biofold.org/>

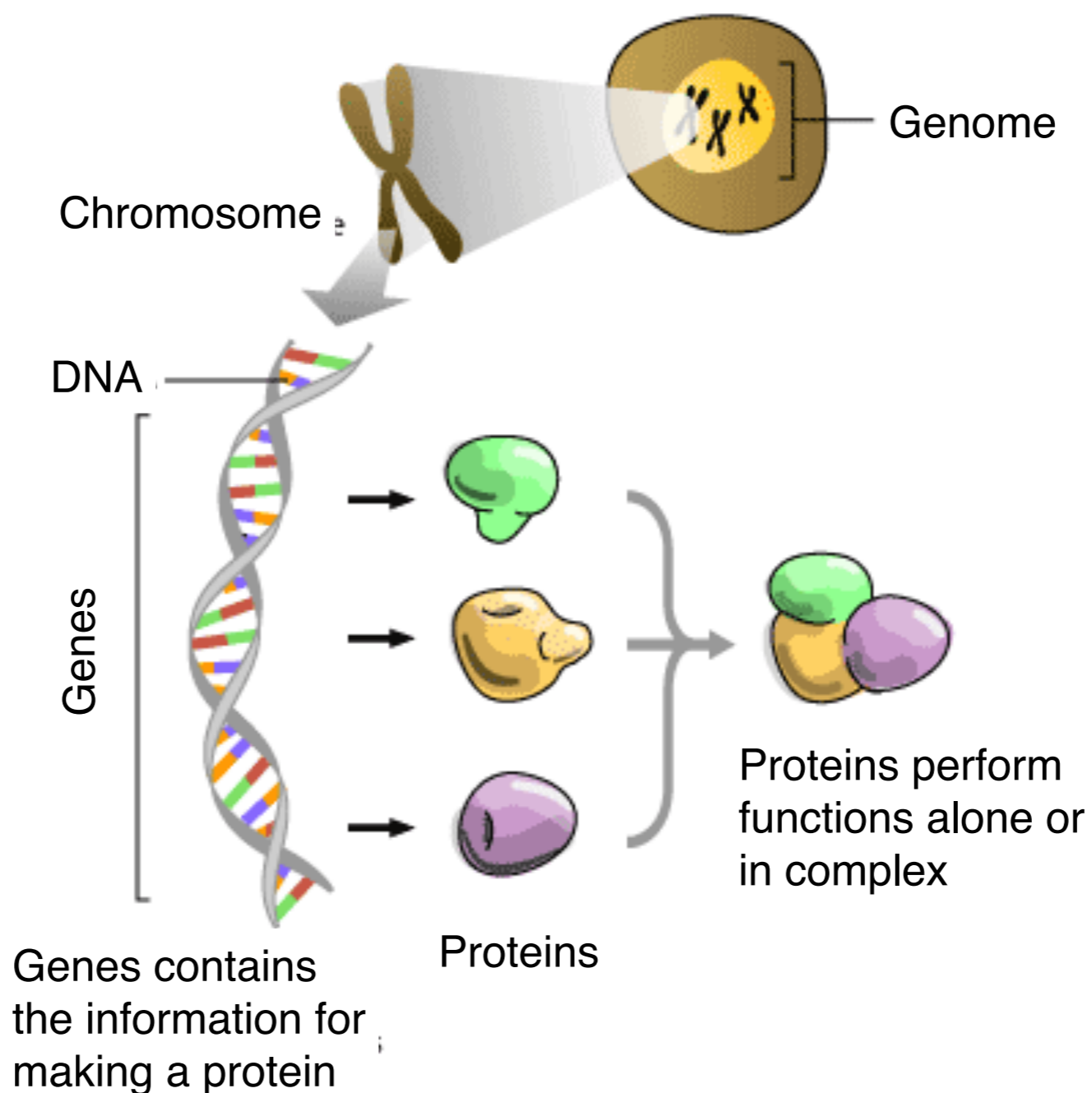


**Biomolecules
Folding and
Disease**

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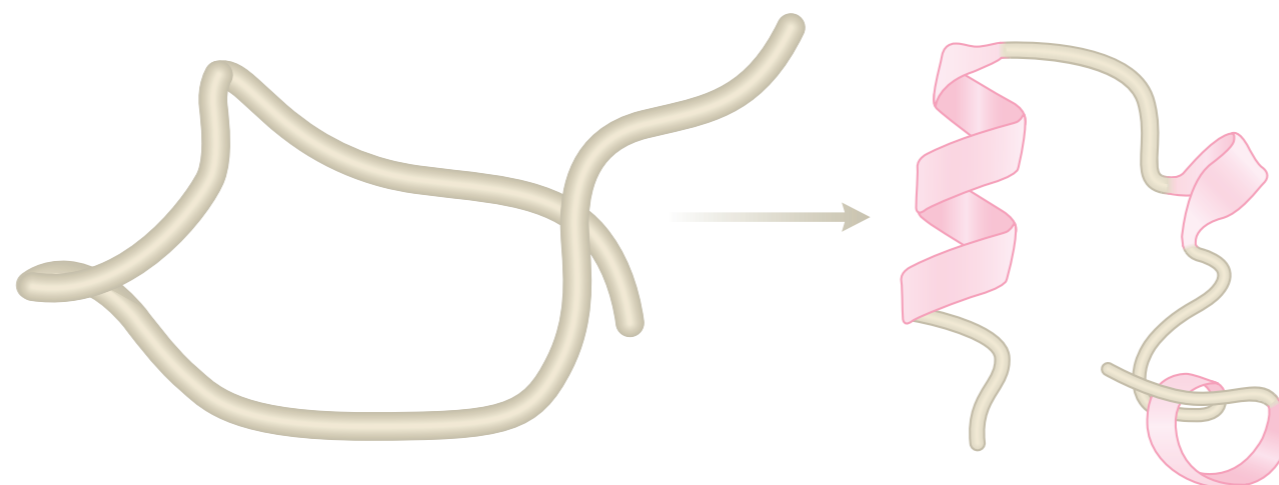


The Central Dogma



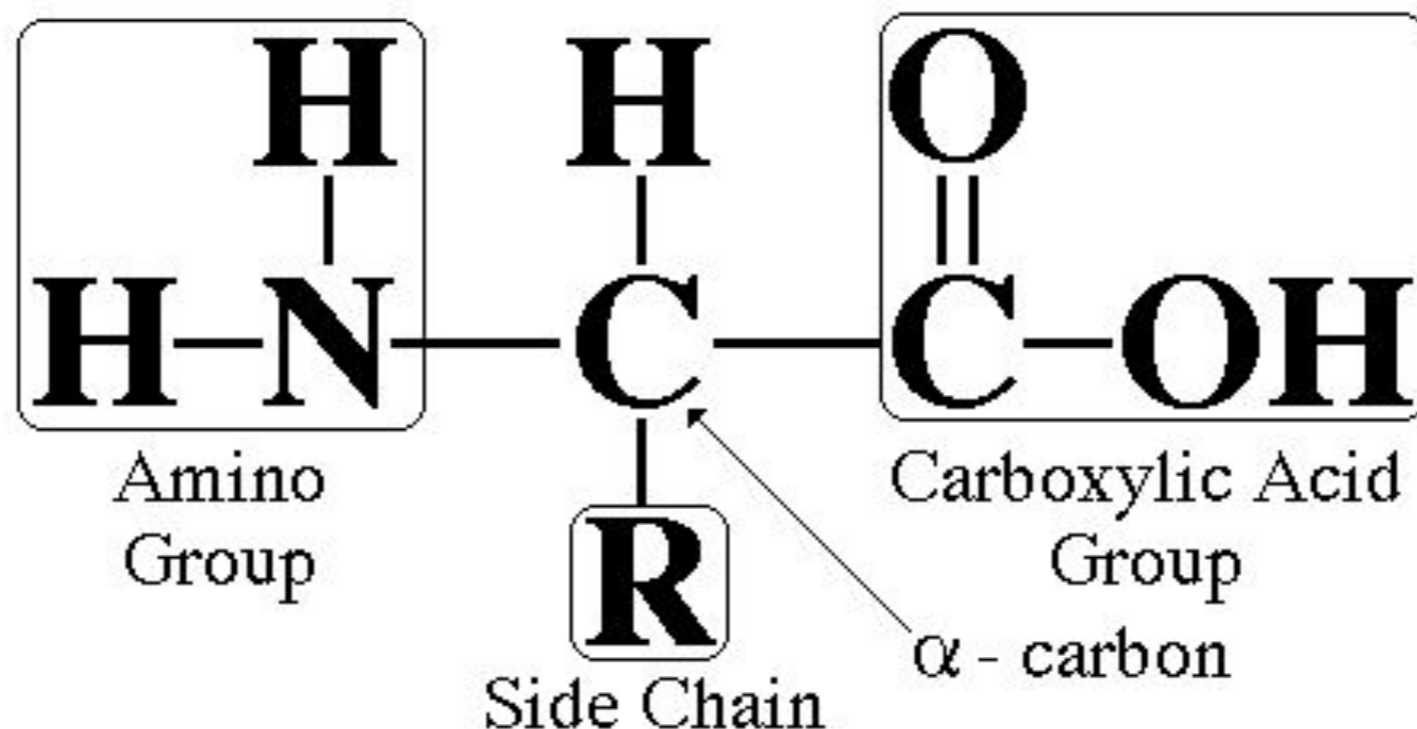
The Protein

- A polypeptide chain **interacts with the solution** adopting a specific three-dimensional structure with a **hydrophobic interior and hydrophilic exterior**.
- The **folded protein** in the proper conformation can **perform a specific function**. In some cases additional polypeptide chains or cofactors must come together before the protein becomes active.
- The **protein structure** can be understood in terms of a conceptual hierarchy, which is **influenced by different interactions**.



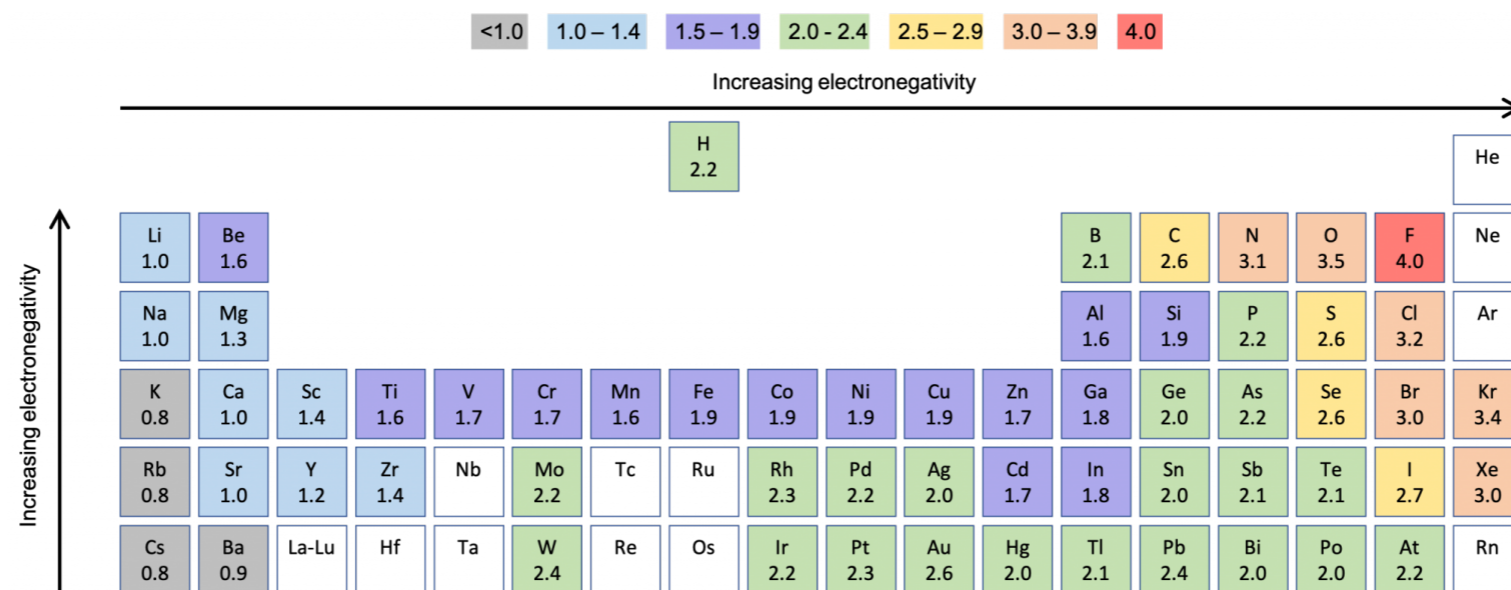
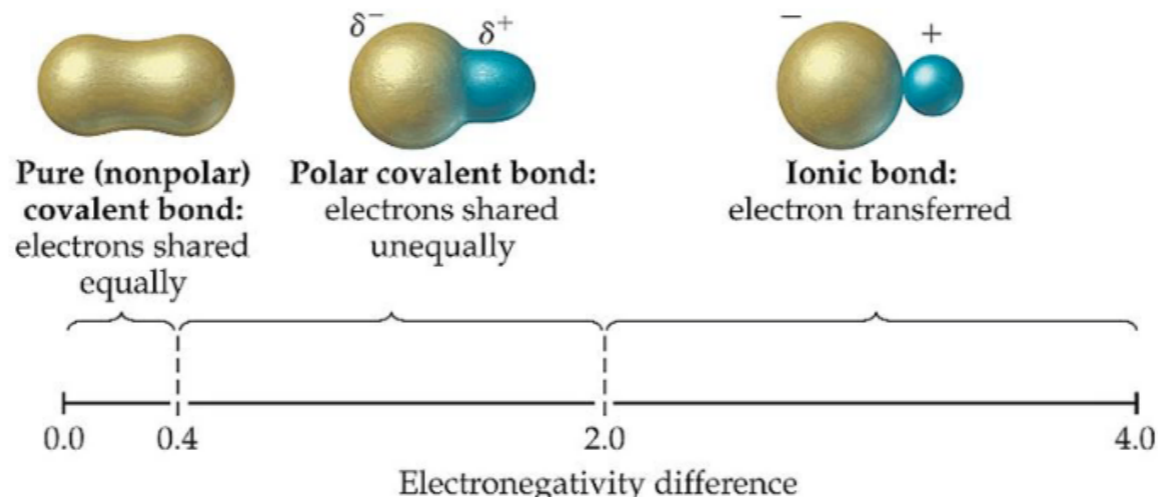
Amino Acid

The side chain (R) determines the type of the amino acid



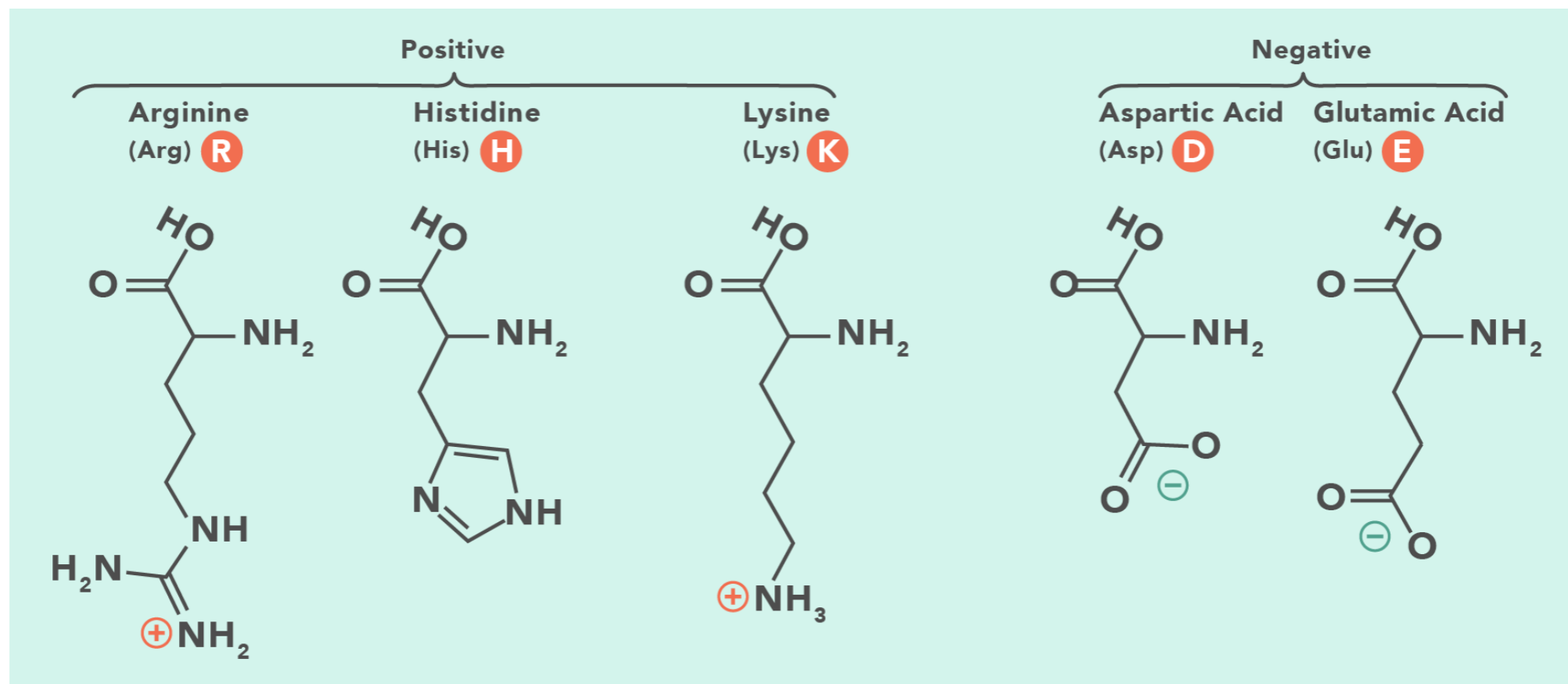
Electronegativity

Tendency for an atom to attract shared electrons when forming a chemical bond.



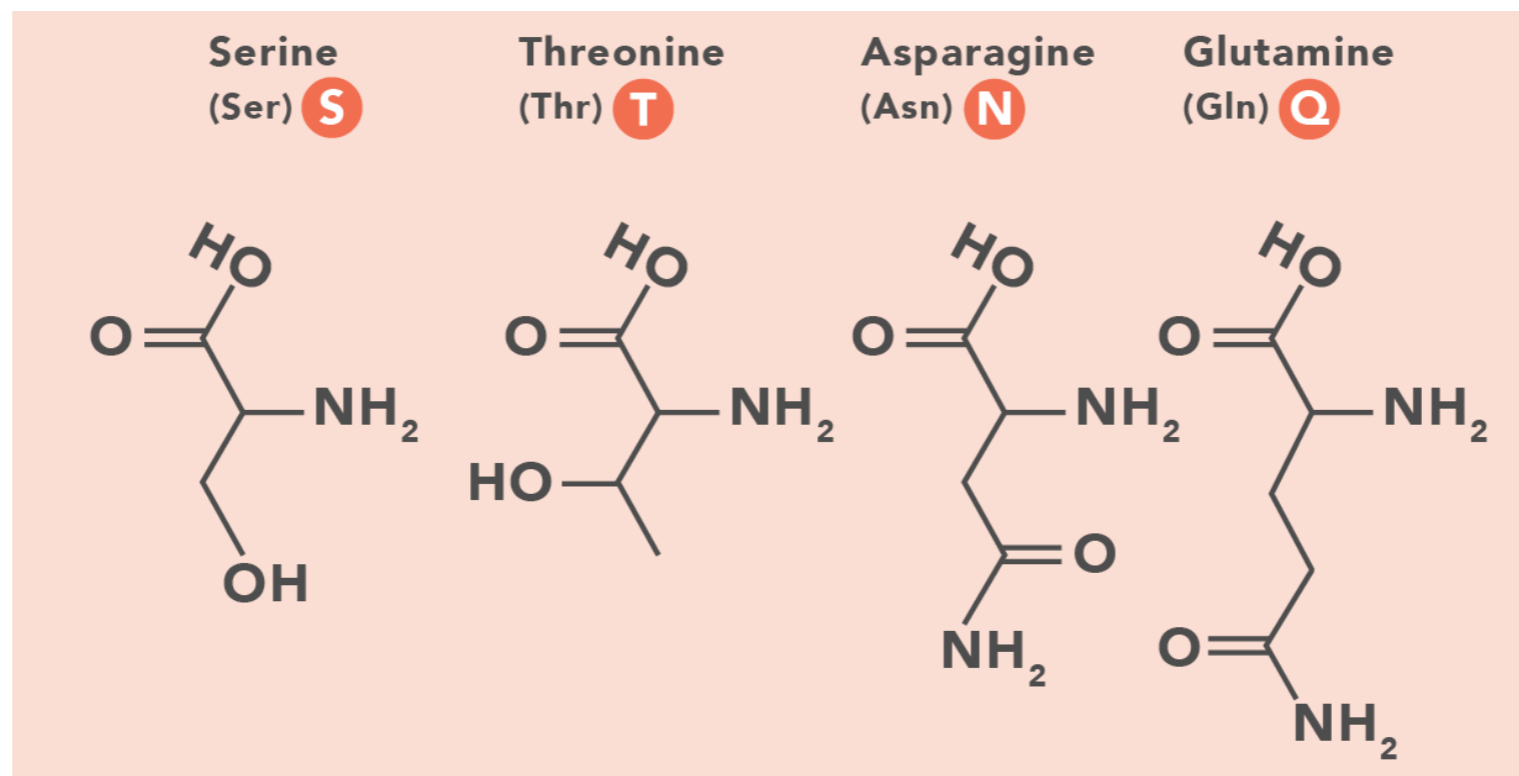
Charged Amino Acid

The side chain end either with amino (-NH₂) or carboxylic (-COOH) groups



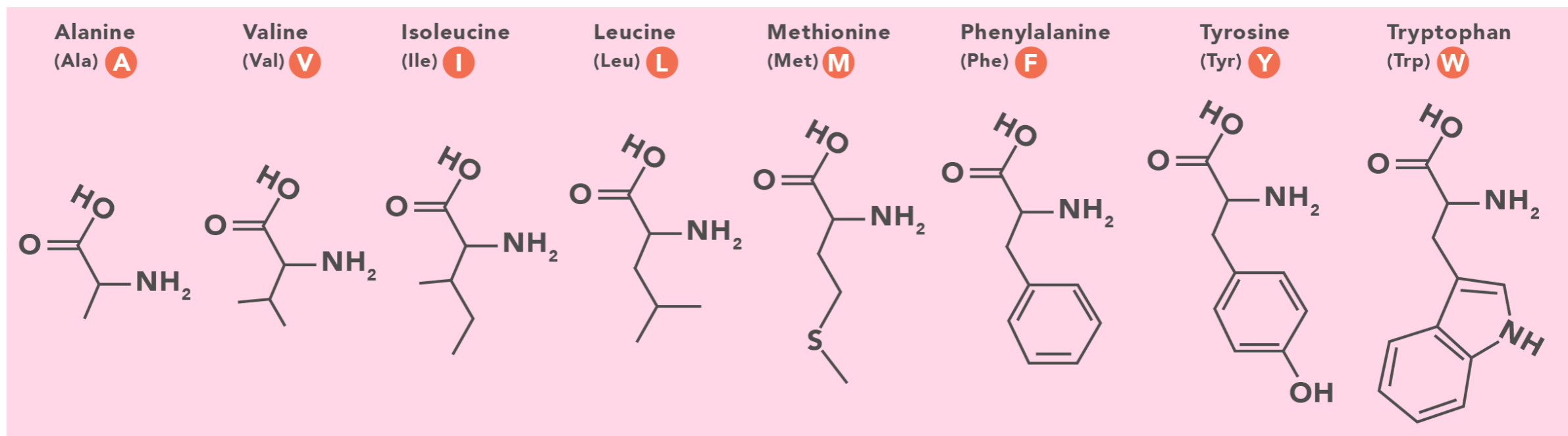
Polar Amino Acid

The side chain end with partially charged groups



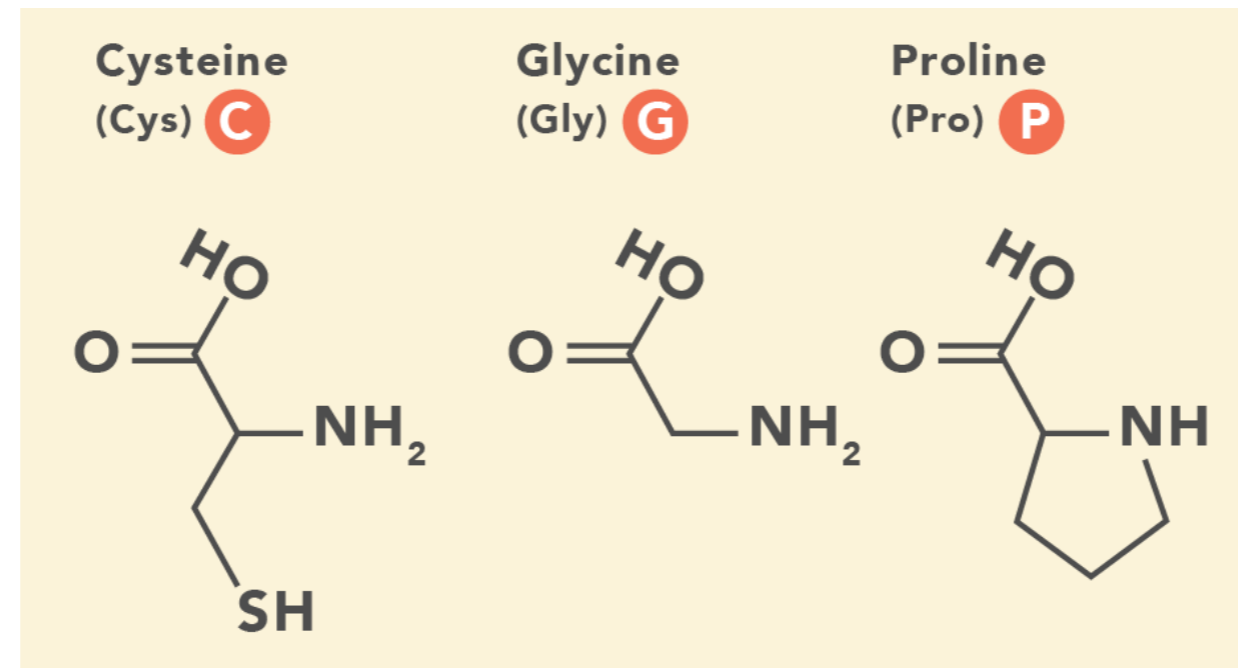
Hydrophobic Amino Acids

The side chain (R) are alkyl or aromatic groups



Special Cases

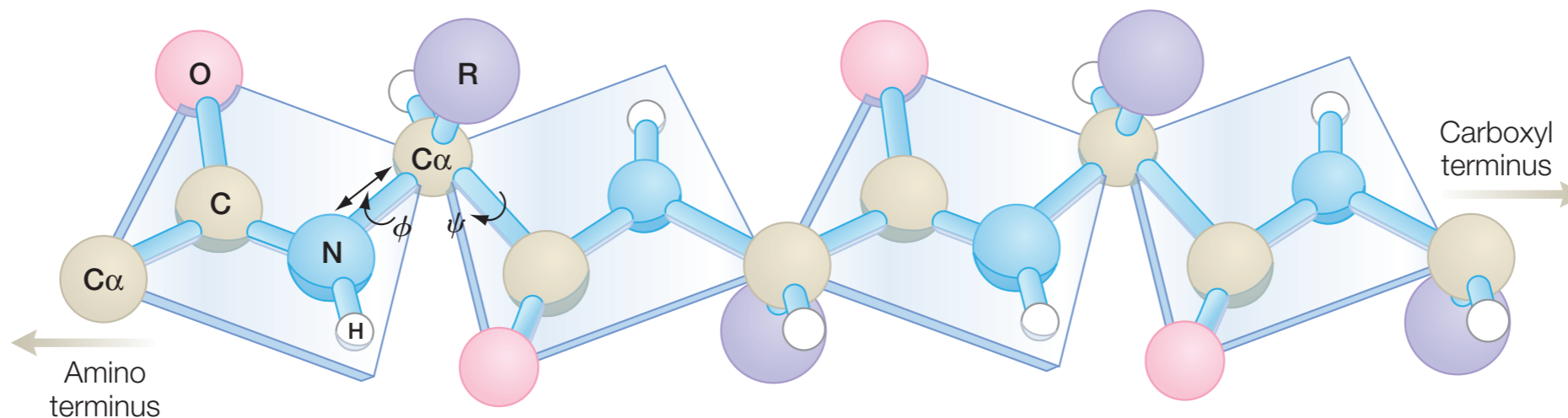
The side chain (R) has particular features



Polypeptide Chain

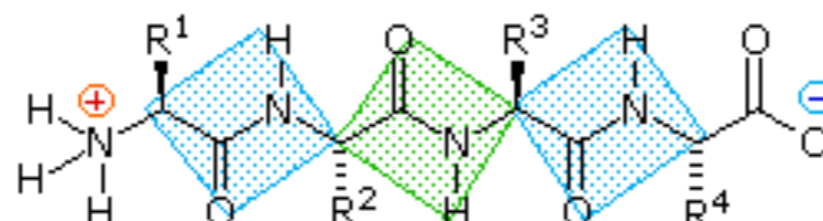
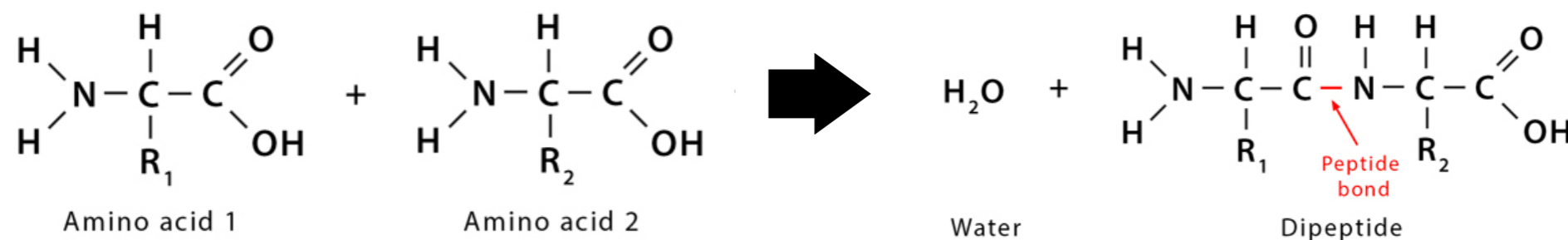
Amino acids form peptide bonds in which the **carboxyl group** of one amino acid is joined to the **amino group** of another amino acid. Many amino acids joined by peptide bonds form a polypeptide chain.

The **protein chain** consists of the regularly repeating main chain or backbone and the side chains.

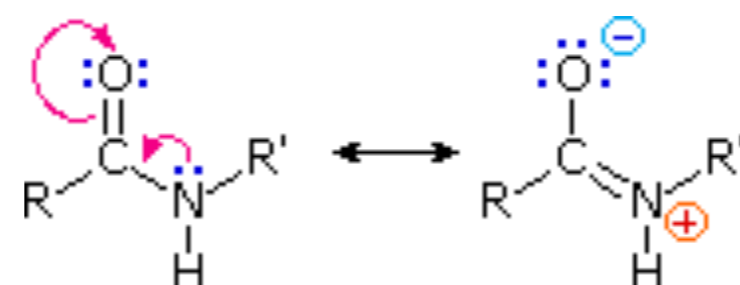


Peptide Bond

The peptide unit is always rigid and planar with the hydrogen of the amino group opposite to the oxygen of the carbonyl group, except for proline.

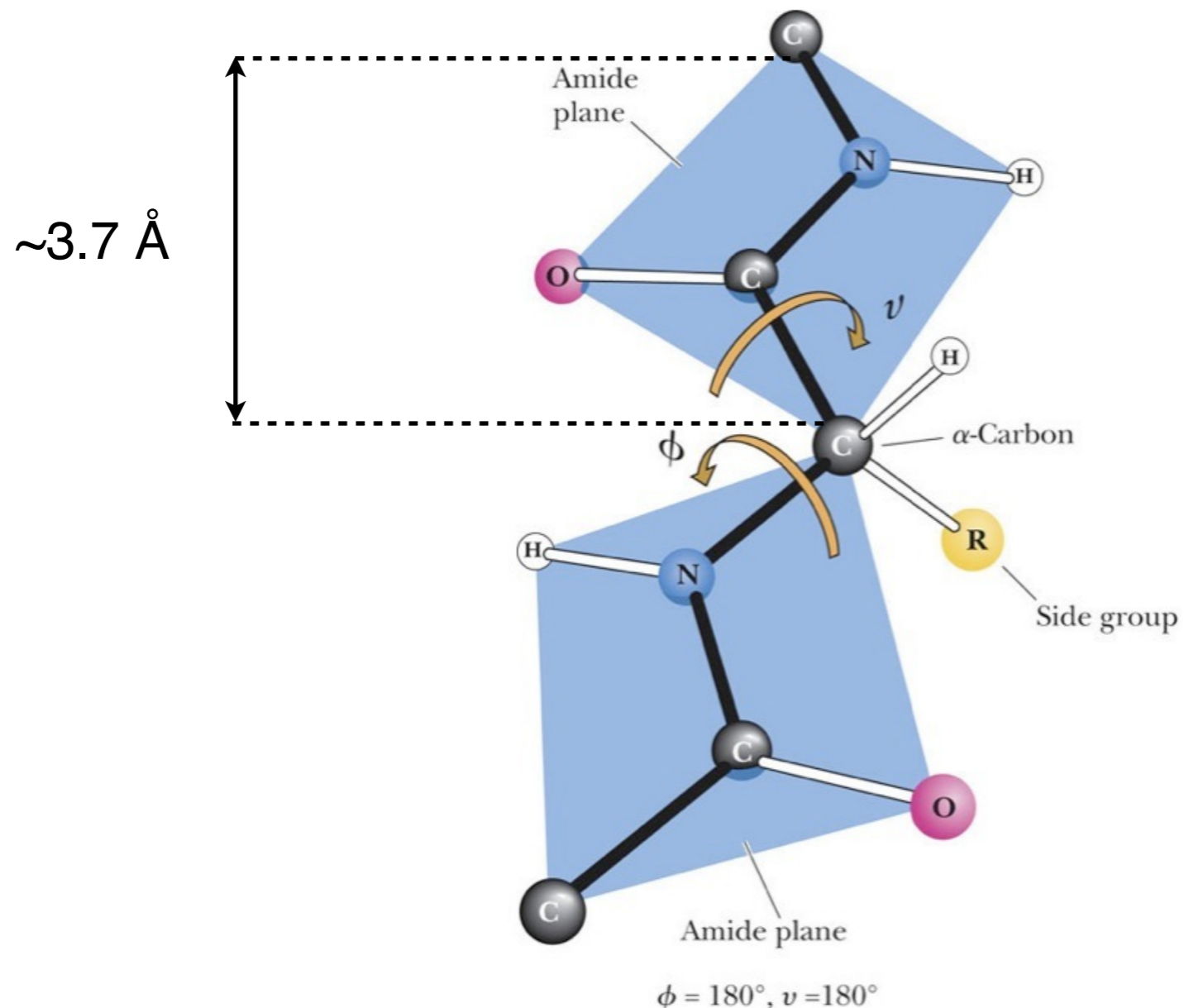


The bond between the carbonyl carbon atom and the nitrogen atom is not free to rotate.



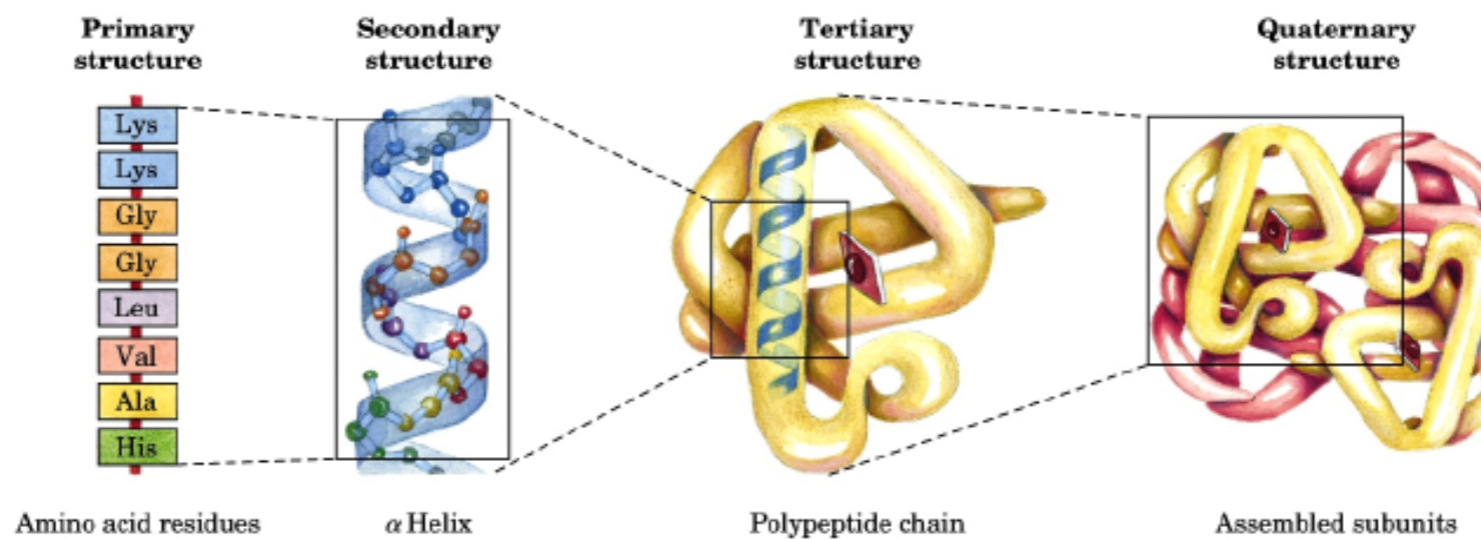
Torsion Angles

The protein consists of a series of planes that can rotate relative to each other. The bond angles centered around at the C_α are identified as ϕ and ψ



Protein Structures

- The **sequence of amino acid residues** linked together by peptide bonds describes the **primary structure** of the protein.
- Local regions of the protein **fold into stable arrangements** of amino acid residues that are recurring in proteins (α helices and β sheets)
- The **overall fold of these secondary-structure elements** describes the **tertiary structure** of the protein.
- The **quaternary structure** is the **arrangement of different polypeptides** of the protein.



Hydrogen Bond

- Hydrogen bond is an electrostatic interaction between the partial negative charge (acceptor) and the partial positive charge (donor).
- The strength of the hydrogen bond will depend upon the relative angles and distances.
- A typical distance is 1.8 Å from the hydrogen to the acceptor, or about 2.8 Å between the nuclei of the donor and acceptor.
- Hydrogen bonds involving the main-chain atoms establish the stability of the secondary structures of proteins.

Electrostatic Interactions

The side chains of the amino acid residues lysine, arginine, glutamate, aspartate, and histidine are ionizable. They can form **electrostatic interactions contribute to protein stability and function.**

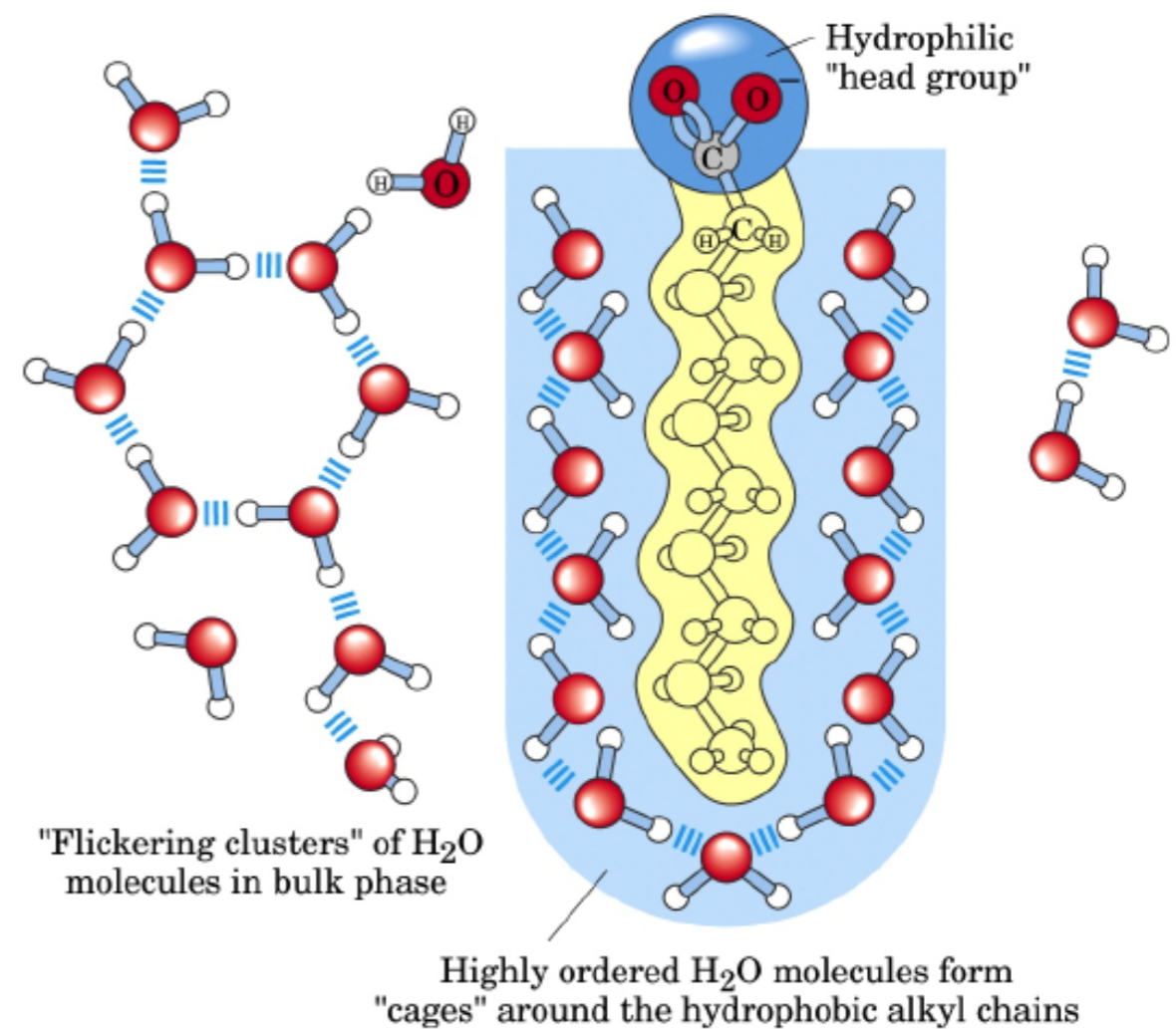
In a typical electrostatic model, the potential between two charges, q_1 and q_2 , that are separated by a distance, r , is given by:

$$V(r) = \frac{q_1 q_2}{4\pi\epsilon r} = (1389 \text{ kJ mol}^{-1}) \frac{q_1(e)q_2(e)}{\epsilon r(\text{\AA})}$$

where ϵ is the **dielectric constant**. The dielectric constant of a vacuum is defined as 1.0 and its value in different solvents **ranges from 80 in a polar solvent such as water to 2 for a nonpolar solvent such as benzene.**

Hydrophobic effect

- Water molecules form a cage-like structure around the nonpolar molecule.
- The positive ΔH is due to the fact that the cage has to be broken to transfer the nonpolar molecule.
- The **positive ΔS** is due to the fact that the water molecules are less ordered (an increase in the degree of disorder) when the cage is broken.



$$\Delta G = \Delta H - T\Delta S$$

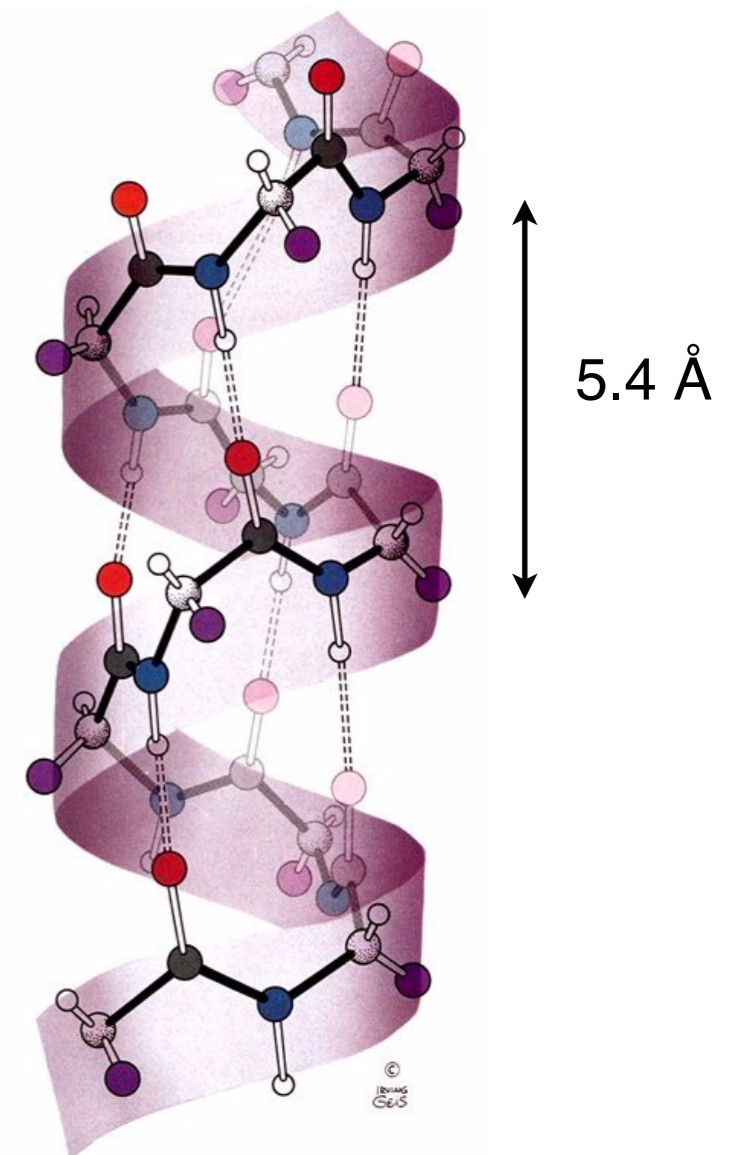
Folding interactions

Several types of **electrostatic interactions** are **contributing** to the **stability** of the native state but they are **not the driving forces** in the folding process

Type	Examples	Binding energy (kcal/mol)	Change of free energy water to ethanol (kcal/mol)
Electrostatic interaction	Salt bridge	$\text{—COO} \cdots \text{N}^+\text{H}_3\text{—}$	-5
	Dipole-dipole		+0.3
Hydrogen bond	Water		-4
	Protein backbone		-3
Dispersion forces	Aliphatic hydrogen		-0.03
Hydrophobic forces	Side chain of Phe		-2.4

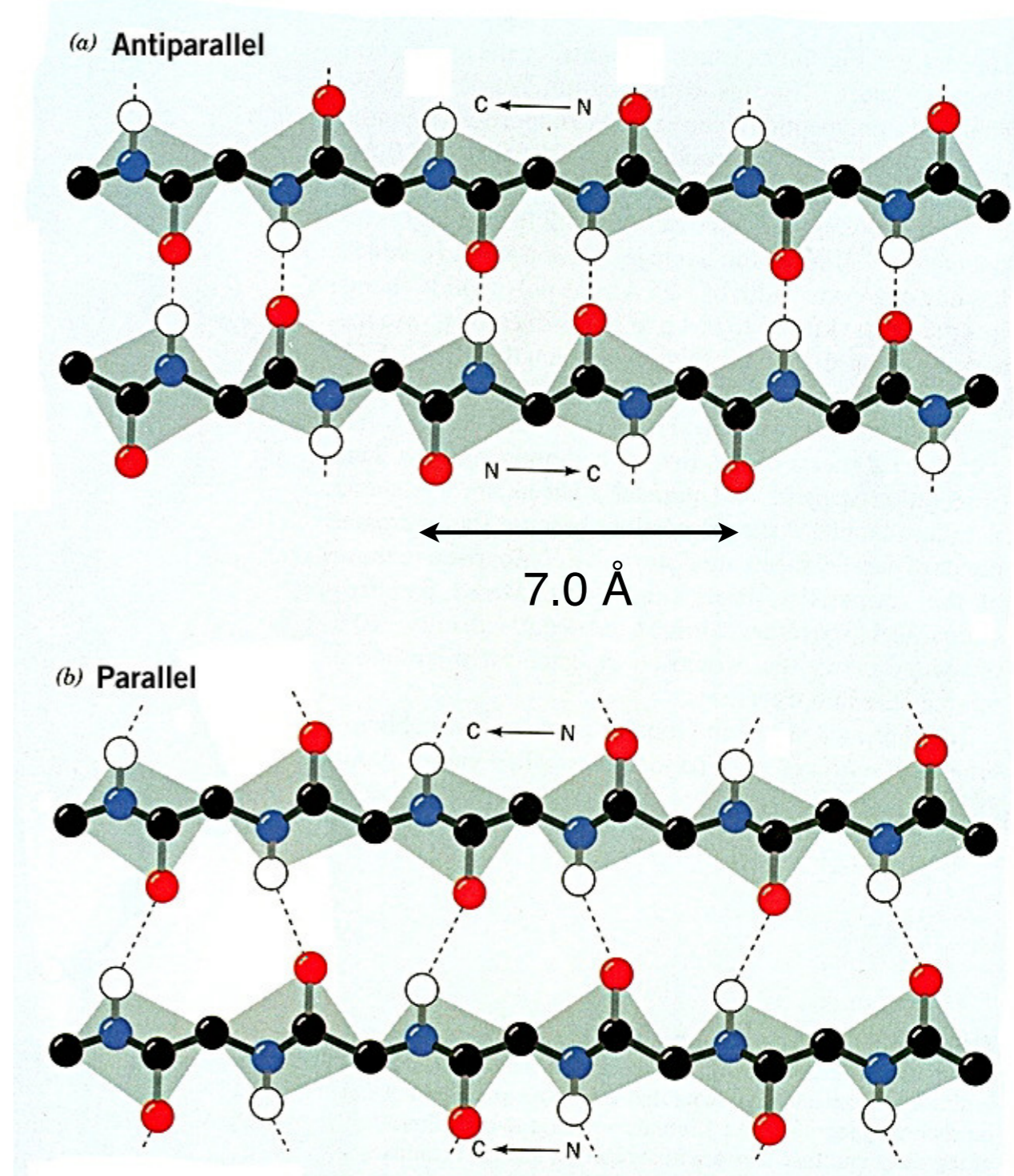
Secondary structure (I)

- Helices observed in proteins are mostly right-handed.
- Typical ϕ , ψ values for residues in α -helix are around -60° ; -50°
- Side chains project backward and outward.
- The core of α -helix is tightly packed.



Secondary structure (II)

- Typical ϕ , ψ values for residues in β -sheet are around 140° , -130°
- Side chains of neighboring residues project in opposite directions.
- The polypeptide is in a more extended conformation.
- Parallel β -sheets are less stable than anti-parallel β -sheets.

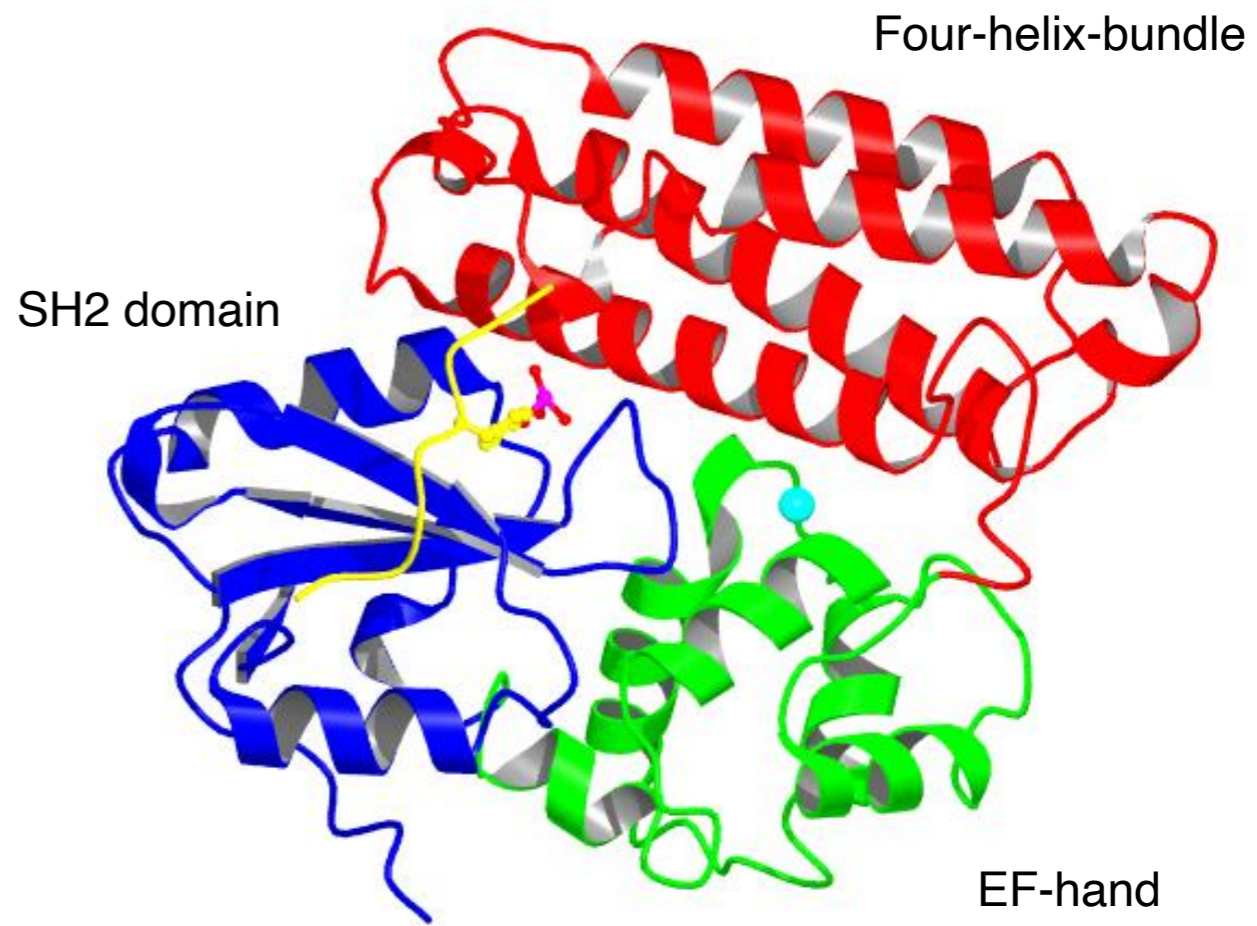
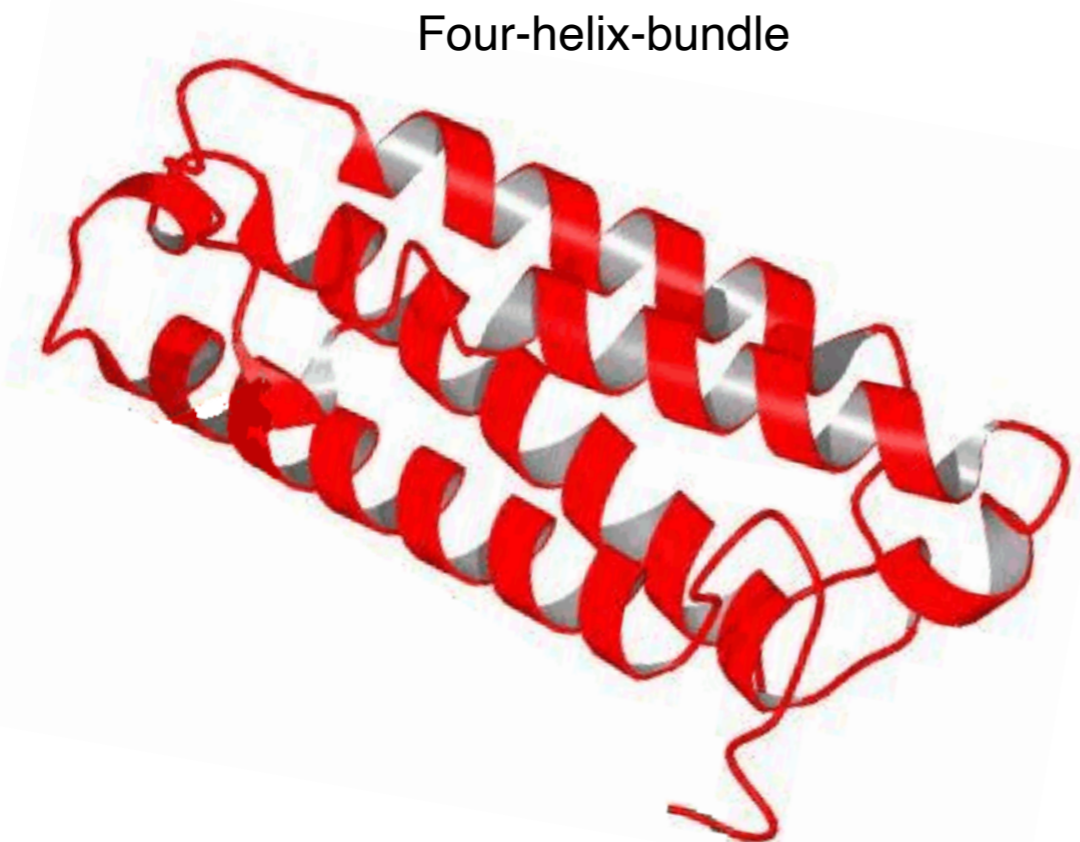


Quaternary structure

The arrangements of secondary structural elements form the Tertiary Structure of the protein.

The complex of **two or more protein domains defines the Quaternary Structure**.

In the example Four-helix-bundle, EF-hand and SH2 domains together form an integrated phosphoprotein that functions as a negative regulator of many signaling pathways from receptors at the cell surface.



The Protein Data Bank

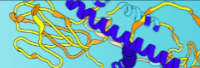
The largest repository of macromolecular structures obtained mainly by X-ray crystallography and NMR

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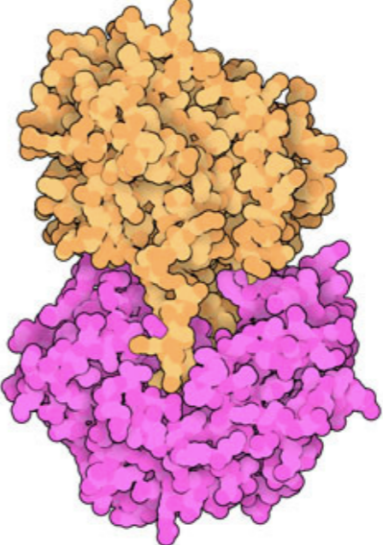
RCSB Protein Data Bank (RCSB PDB) enables breakthroughs in science and education by providing access and tools for exploration, visualization, and analysis of:

- Experimentally-determined 3D structures from the **Protein Data Bank (PDB)** archive
- Computed Structure Models (CSM) from AlphaFold DB and ModelArchive

These data can be explored in context of external annotations providing a structural view of biology.

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October Molecule of the Month



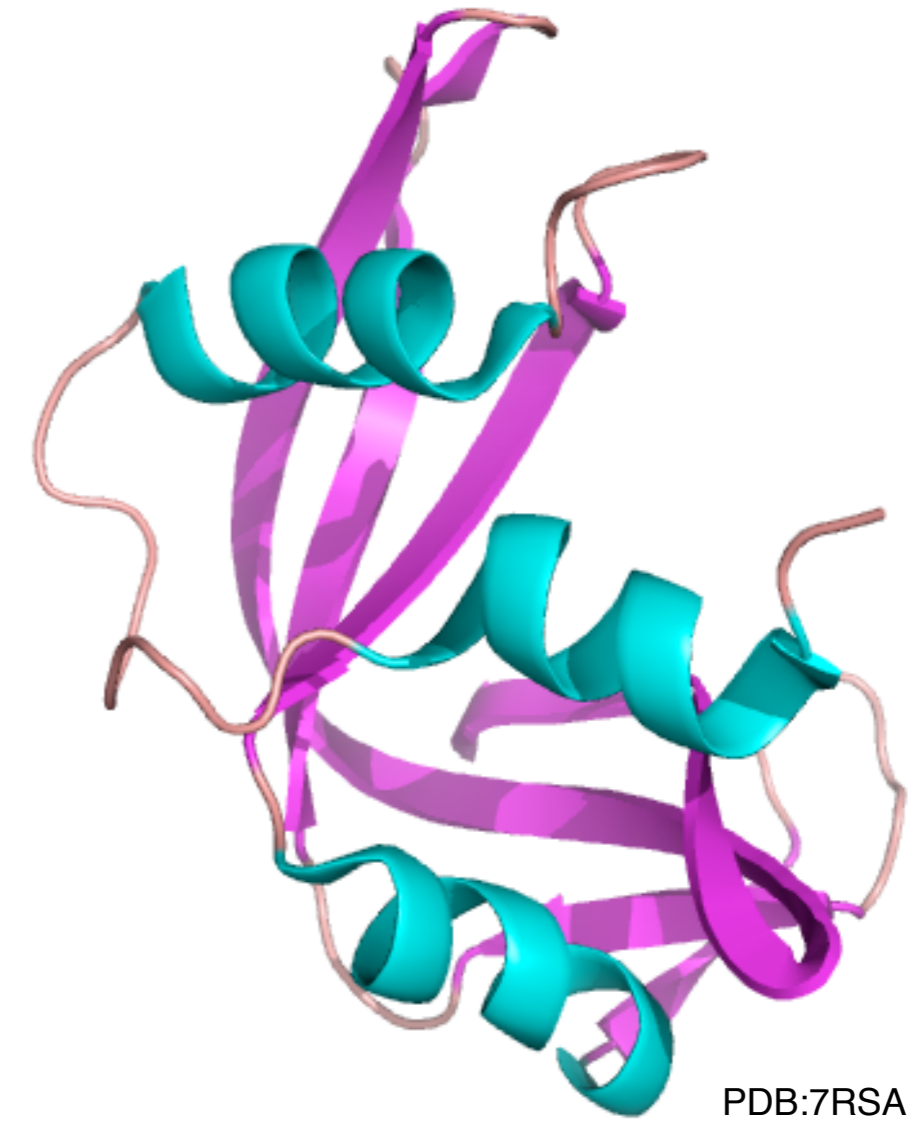
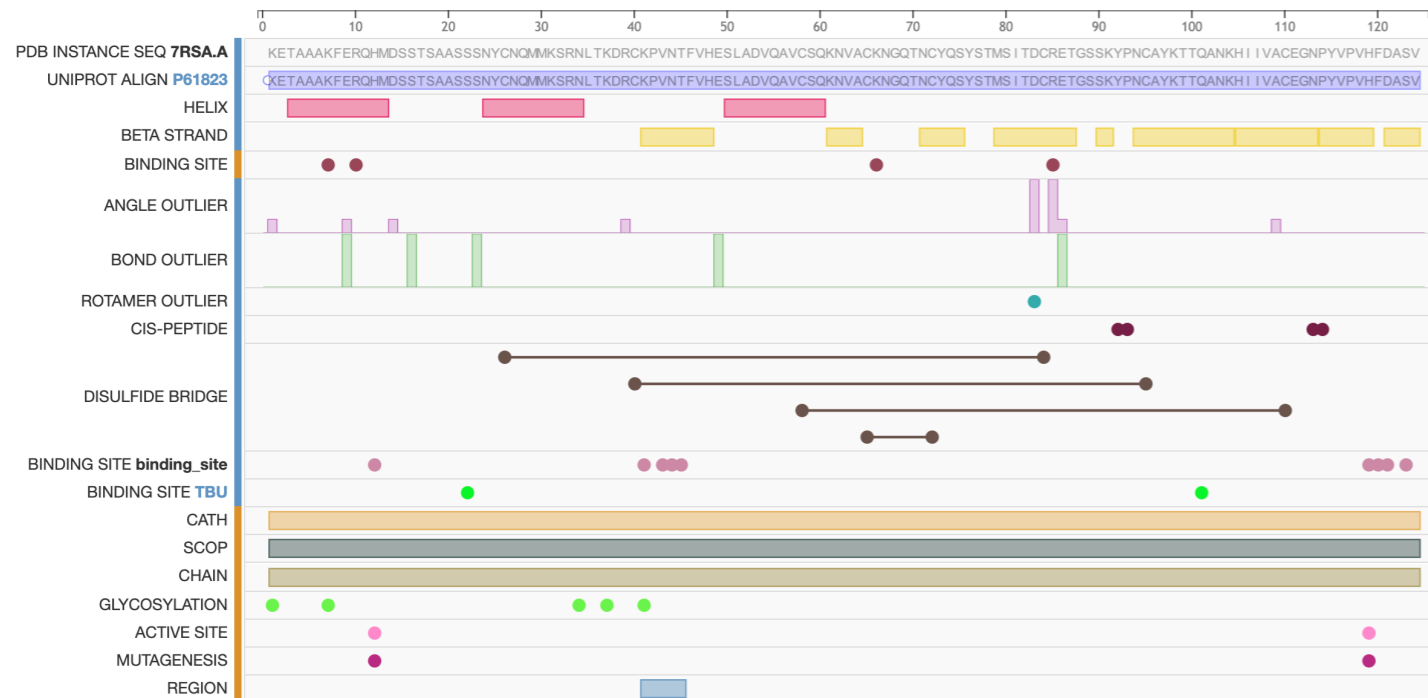
Angiotensin and Blood Pressure

<http://rcsb.org>

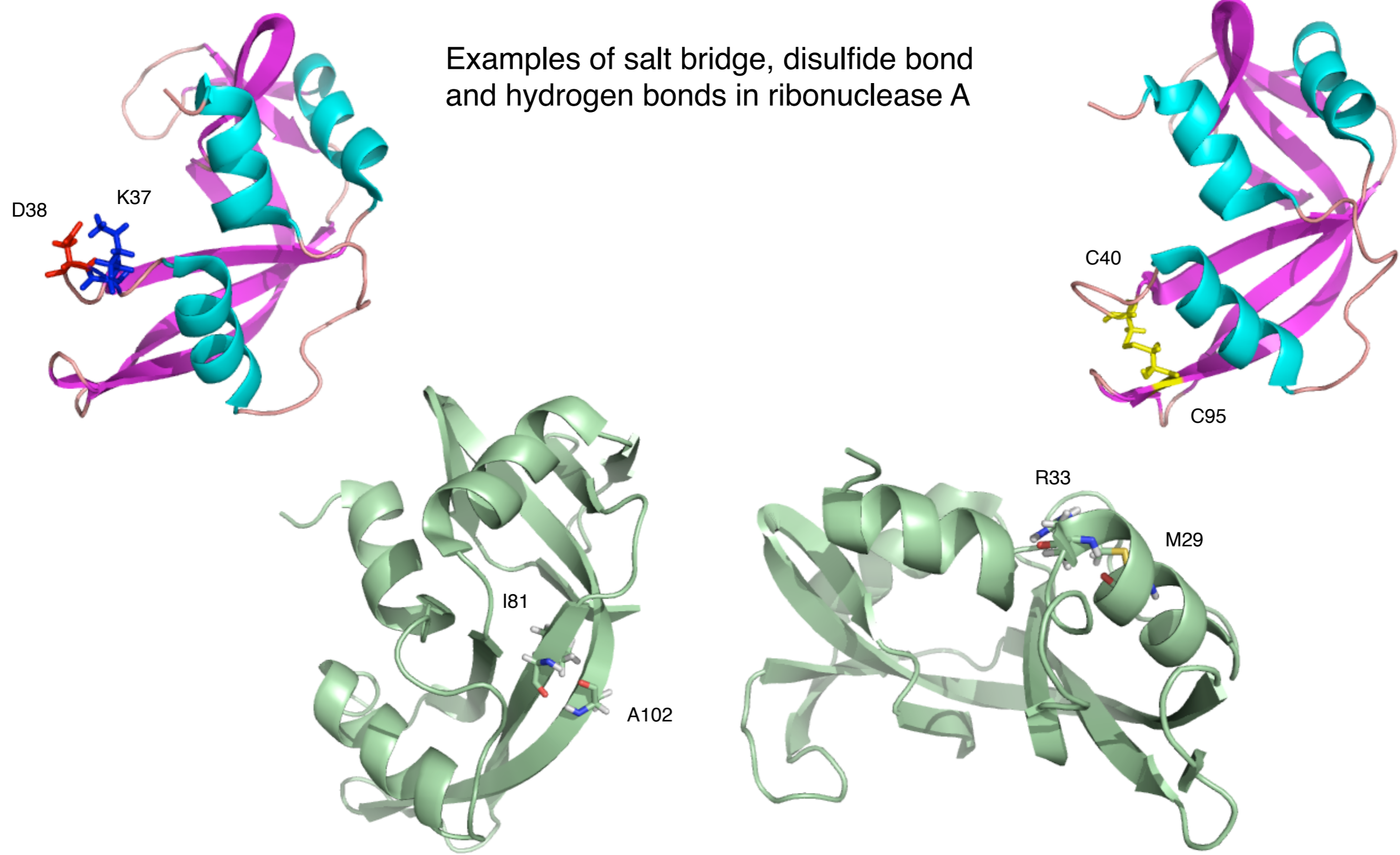
<http://ftp.rcsb.org/pub/pdb/>

The Bovine Ribonuclease A

Ribonuclease A (RNase A) is a [pancreatic ribonuclease](#) which belongs to a class of Lyases. This enzyme cleaves at the 3'-side of pyrimidine (uracil or cytosine) phosphate bonds in RNA.



Bonds and interactions



Protein Binding

Protein binding can be formalized with the following reaction:



where R is the receptor L the ligand and RL receptor-ligand complex. This formalism can **apply both to the study of drug-target and protein-protein interactions**. In the first case the ligand will be a small molecule while in the second case the ligand will be a protein. In general the formation constant (K_f) is:

$$K_f = \frac{[RL]}{[R][L]}$$

To quantify the affinity of the receptor for ligand, the dissociation constant (K_D) is indicated:

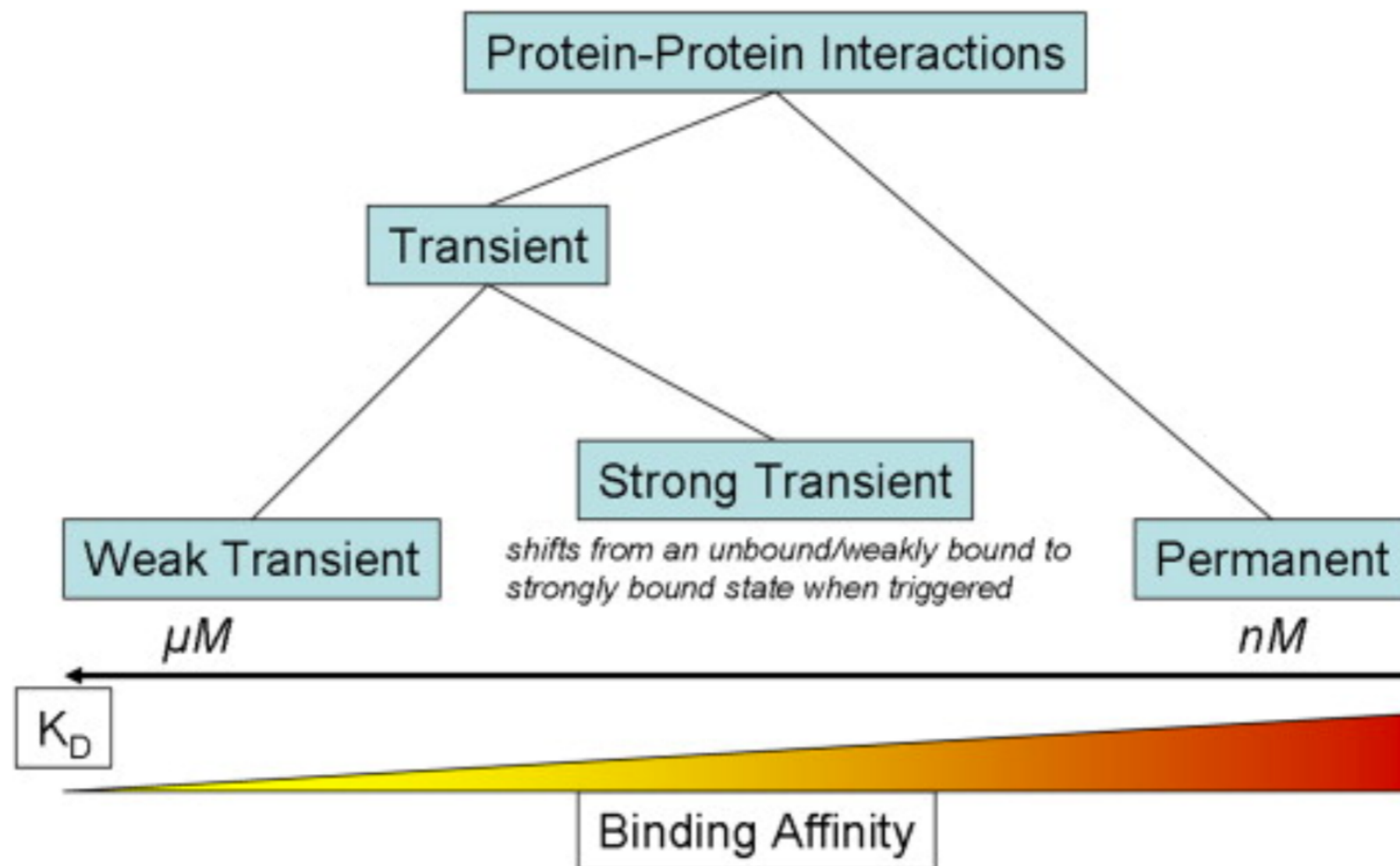
$$K_D = \frac{1}{K_f} = \frac{[R][L]}{[RL]}$$

- A small value for $K_D \Rightarrow$ the equilibrium favors the complex (high affinity)
- A large value for $K_D \Rightarrow$ the equilibrium favors the separation (low affinity)

The Molecular Viewpoint

- The affinity of PPI varies from millimolar to picomolar, depending on the type of interaction and signaling needed (Chen et al. Protein Sci. 2013)
- Despite affinity varies over a wide range, proteins maintain a high degree of specificity for their partners
- Many proteins exhibit specificity for multiple partners (Reichmann et al. Curr. Opin. Struct. Biol. 2007).
- The nature of the interaction surface determines how proteins interact
- A detailed knowledge of the interaction surfaces of proteins and their energetics is necessary to understand the regulatory mechanisms of biochemical pathways (especially to modulate or block these pathways for therapeutic purposes)

Protein-Protein Interactions



Strong transient: This category includes interactions that are triggered/stabilised by an effector molecule or conformational change. An example is given by the Ras proteins, which form tight complexes with their partners when GTP-bound and only weak complexes when GDP-bound.

Surface of Interaction (I)

- The area of PPI interfaces is large (1000 to 4000 Å²)
- Standard-sized interfaces are 1200 to 2000 Å²
- Short-lived and low-stability complexes ⇒ smaller interfaces (1150–1200 Å²)
- large surfaces (2000 to 4600 Å²) ⇒
 - proteases and particular inhibitors
 - G-proteins and other components of the signal transduction system
- Protein-small molecule interaction surfaces have an area of 300 to 1000 Å².

Surface of Interaction (II)

- Surfaces of PPIs are generally **flat** and lack the grooves and pockets that are present at the surfaces of proteins that bind to small molecules.
- PPI **surfaces are generally hydrophobic** in nature.
- Only certain **hydrophobic spots contribute to the free energy** of binding and help to hold the two proteins together.
- Such regions are called **hot spots**.

Hot Spots

- Hot spots account for less than 50% of the contact area of PPI
- A region of protein surface is called a hot spot when replacement of an amino acid residue by alanine in that spot lowers the free energy of binding by at least 2 kcal/mol
- Analysis of the amino acid composition of hot spots shows that some residues are found more frequently in hot spots (Tyr, Trp, and Arg)
- The hot spots are surrounded by energetically less important residues that separate/prevent bulk water from hot spots

Analysis of Protein Complex

- identification of **interface residues/hot spots**
- **details** about the interface
 - solvent accessible surface area, shape, complementarity between surfaces, residue interface propensities, hydrophobicity, segmentation and secondary structure, and conformational changes on complex formation
- assignment of **protein function**
- recognition of **specific residue motifs**

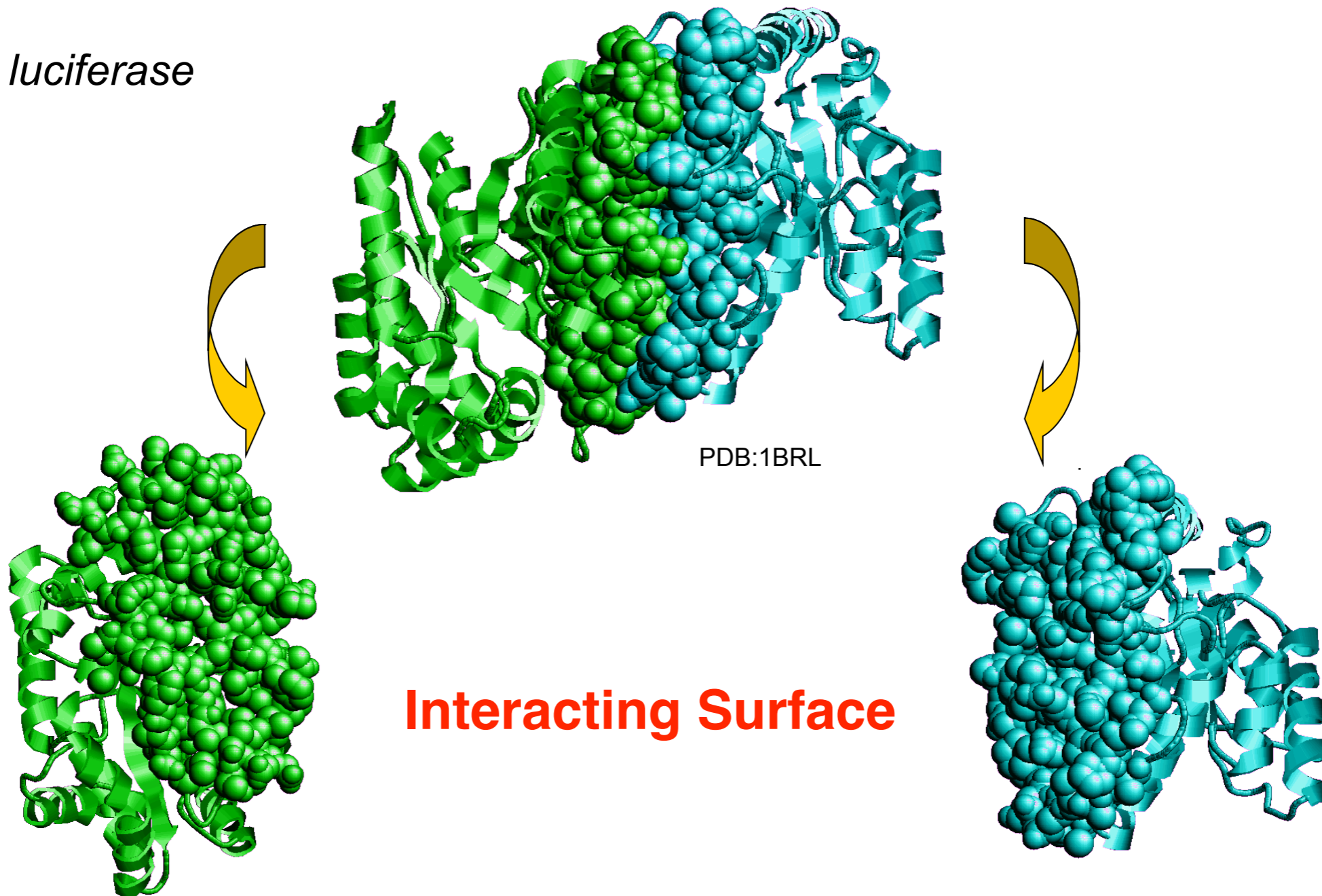
Structure PPI Data

- The most significant contribution to understanding the PPI surface comes from structural biology via **X-ray crystallography** or **NMR** as well as **mutational studies**
- Prediction of interaction/binding sites
- Prediction of protein-protein complexes

Interacting surface

Difference in Accessible Surface Area (ASA) between monomers and complex

Bacterial luciferase



Exercise 1

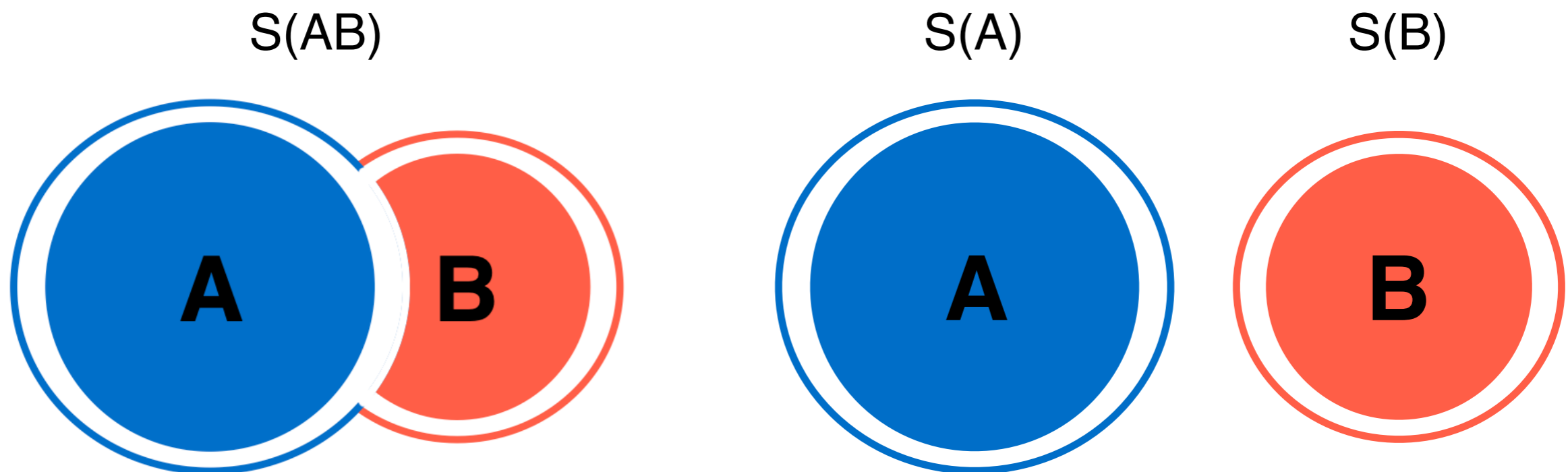
Given the structure of the Bovine Ribonuclease A (PDB: 7RSA) download the structure and measure the distances between the atoms involved in the following interactions:

- LYS37 - ASP38 (salt bridge) d=3.7
- CYS40 - CYS95 (disulphide bond) d=2.0
- MET29 - ARG33 (hydrogen bond) d=3.0 (MET-O ARG-N)
- ILE81 - ALA102 (hydrogen bond) d=2.9 (ILE-O ALA-N)

Suggestion: select the coordinate of the atom involved in the interactions reported above.

Exercise 2

Consider the structure of the Bacterial luciferase (PDB: 1BRL) and determine the size of the surface of interaction between the two monomers A and B. The surface of interaction can be obtained comparing the accessible surface of the complex and the single chains.



Suggestion: Use the [stride](#) server to download the DSSP files of the complex and the monomers. Each file includes the accessible surface of all the residues.