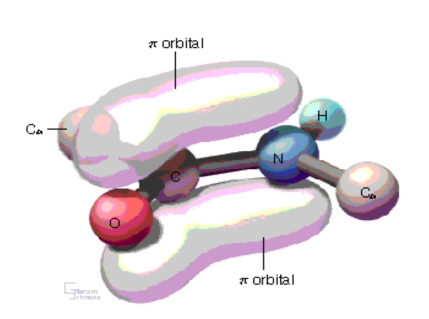
3. Secondary and tertiary structure in proteins

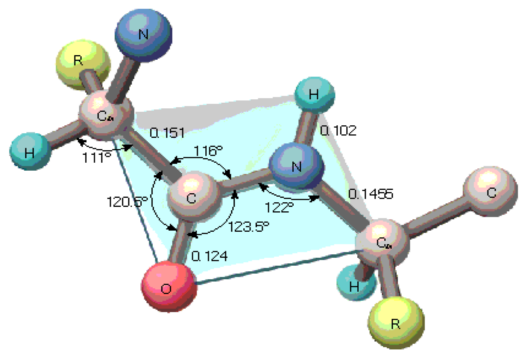
Peptide bond

$$C = N$$

$$C = N$$

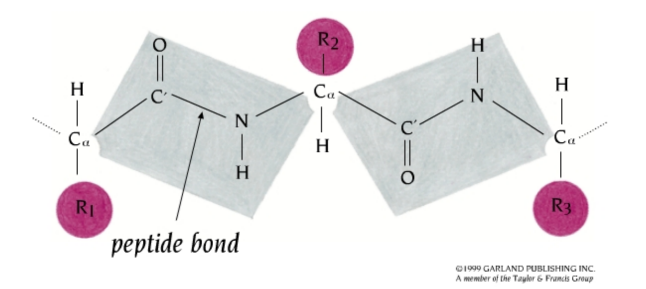
$$C = N$$

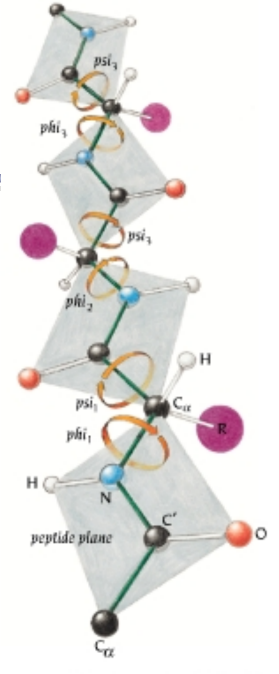




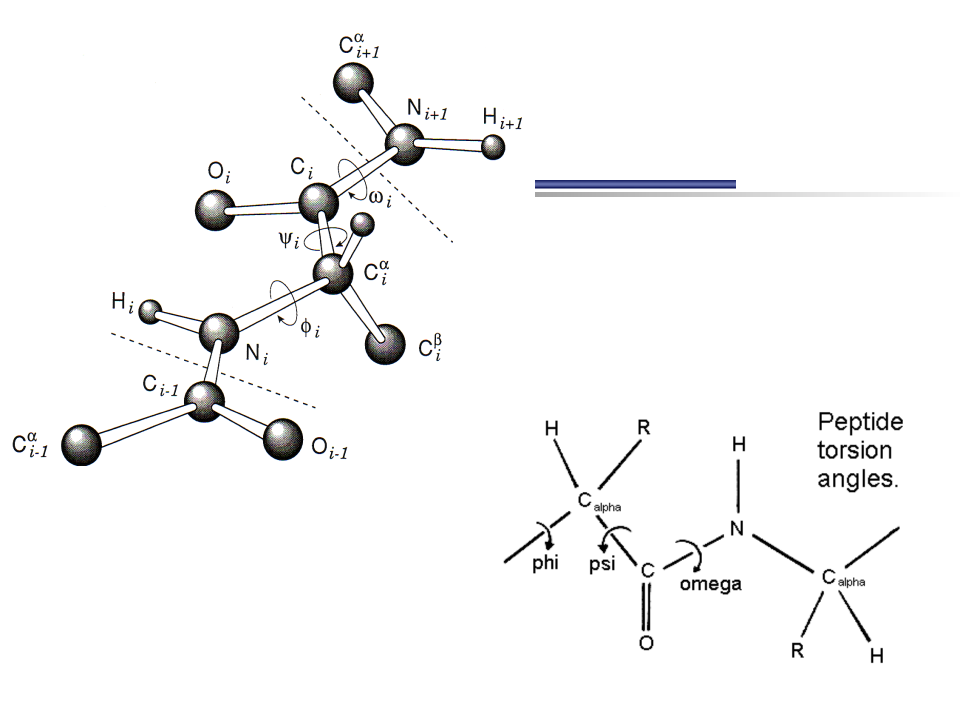
(a) Partial double-bond character of peptide bond

(b) Bond angles and lengths



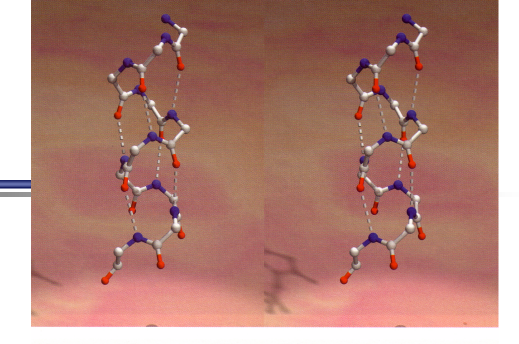


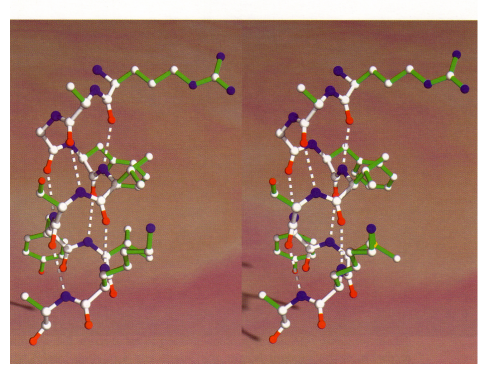
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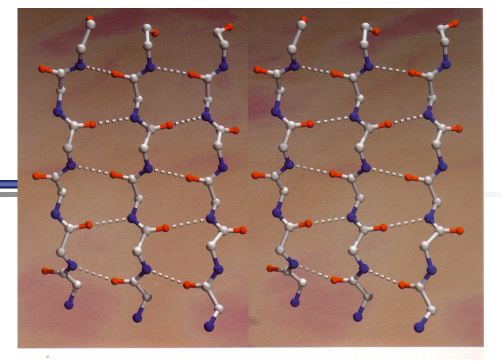
Secondary structure

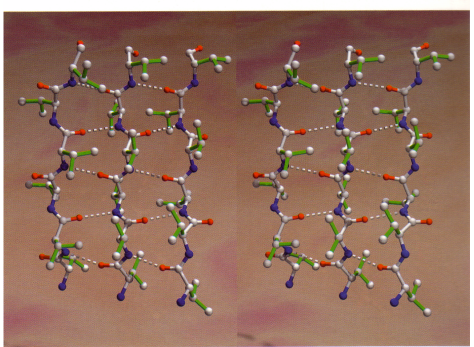
Alpha-helix without/with side chains



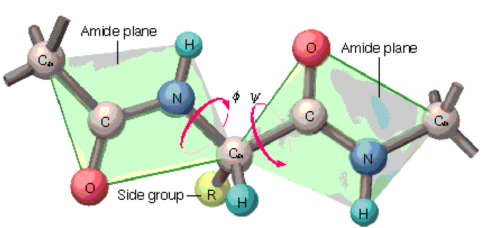


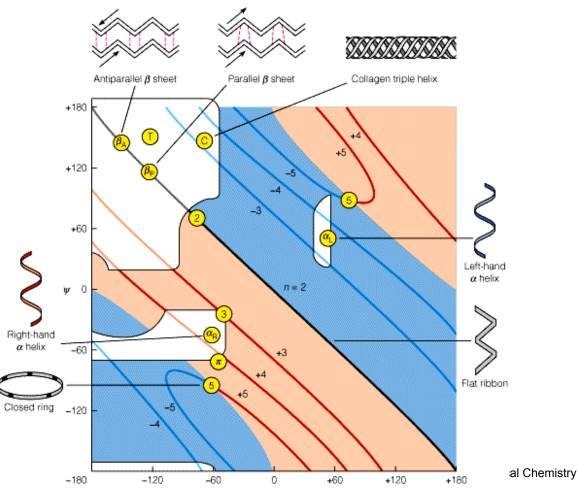
Beta-strands forming a betasheet, without/ with side chains

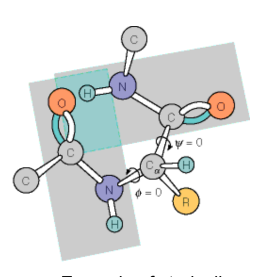




Ramachandran plot:







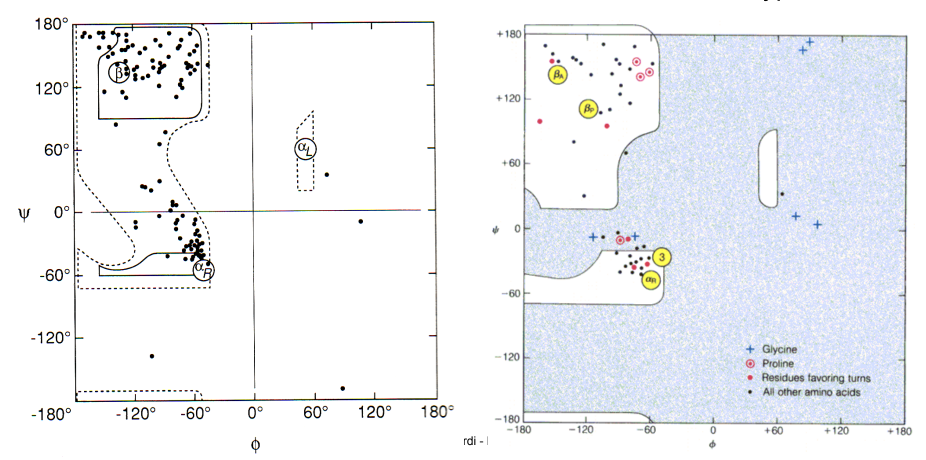
Example of sterically non-allowed conformation for ϕ = 0; ϕ = 0

Sasisekhran-Ramakrishnan-Ramachandran diagram

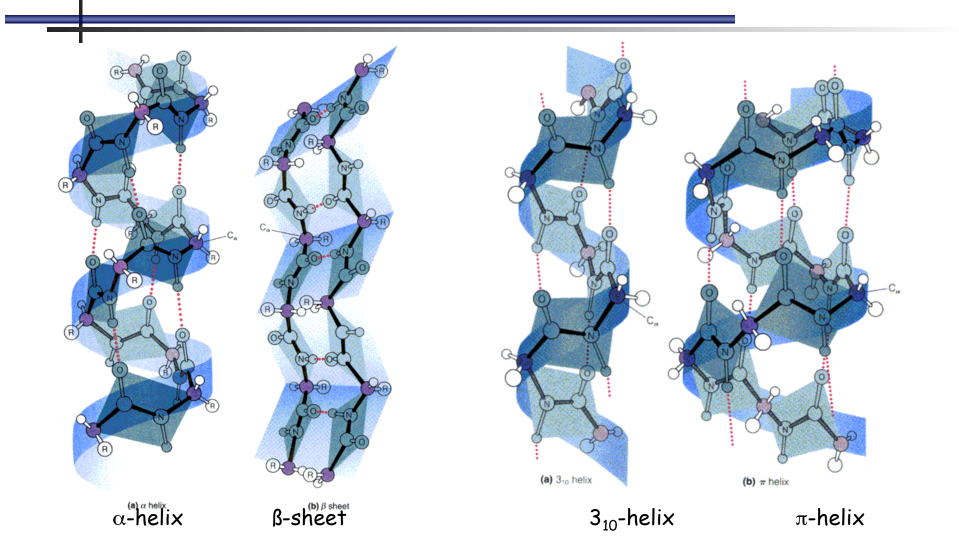




BPTI = Bovin Pancreatic Trypsin Inhibitor



Elements of 2ary structure



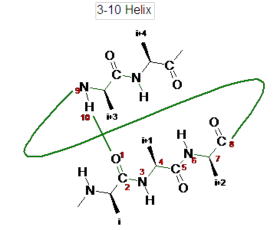
Helices



i+4
o \ _/
H \\ 12\rac{12}{C}
N 10 H13
3 H H C
01 N N 7
H C-N 0 1+2
N Y H
∕ \

Structure Type	Residues/ Turn	Rise (nm)	Number of Atoms in H-Bonded Ring	φ (°)	ψ (°)
Antiparallel β sheet	2.0	0.34	a	-139	+135
Parallel β sheet	2.0	0.32	a	-119	+113
3 ₁₀ helix	3.0	0.20	10	-49	-26
α helix (3.6 ₁₃)	3.6	0.15	13	-57	-47
π helix $(4.4_{16})^b$	4.4	0.12	16	-57	-70

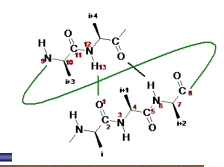
[&]quot;Bonding is between polypeptide chains.



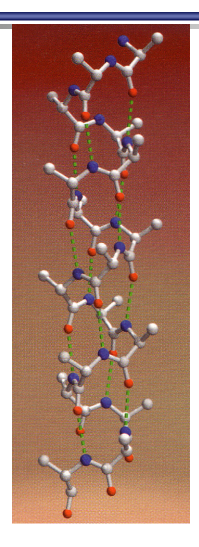
4.4-16 Pi Helix

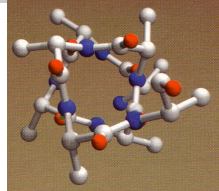
 $^{{}^}b\!\!$ Sterically permitted but not observed in protein.

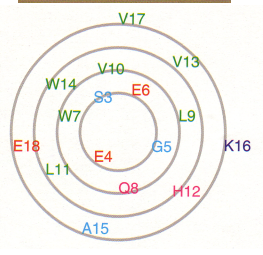
The α -helix



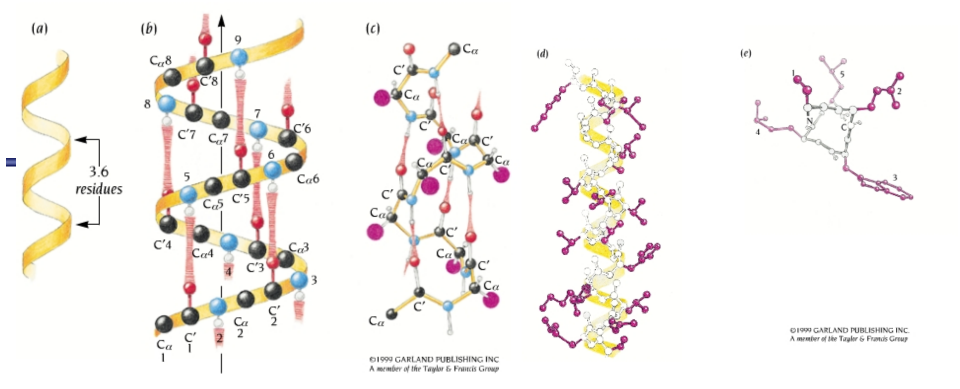
- It is the most common type of helix
- The C=O group of residue i forms a H-bond with the NH group of residue i+4
- One residue is related to the next by a rotation of approximately 100° around the helix axis and with a traslation along the axis by 1.5 Å
- Many α-helices present an asymmetric distribution of hydrophilic/hydrophobic residues







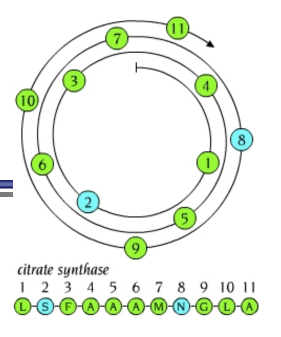
Helical wheel: Helix A from sperm whale myoglobin

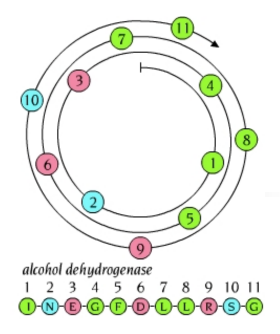


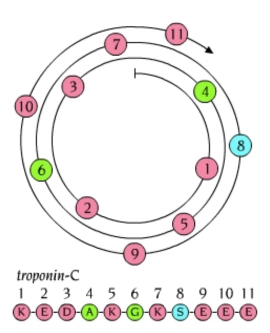
- Length: from 4/5 aa to over 40 aa
- Average length: 10 aa
- 3.6 aa per turn
- 360°: 3.6 = 100° rotation per aa per turn
- Rise: 1.5Å per aa; then an average helix is 15Å long
- Ala, Glu, Leu, Met: good helix formers
- Pro, Gly, Tyr, Ser: bad helix formers

Helices tend to be along the outside of proteins, giving a periodicity of 3.6 with hydrophobic/ hydrophilic aa

Convenient representatio: helical wheel

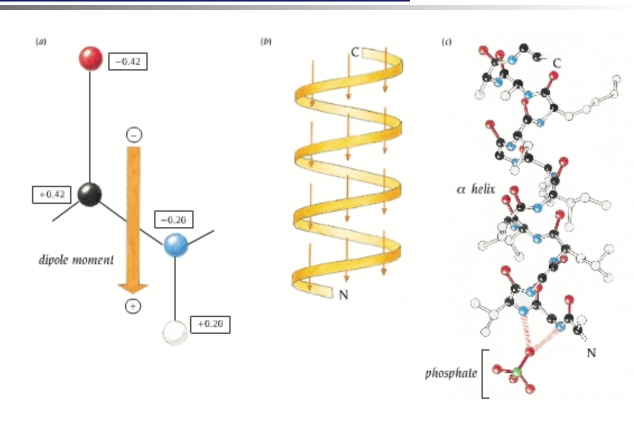




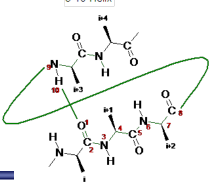


Helices have a dipole moment

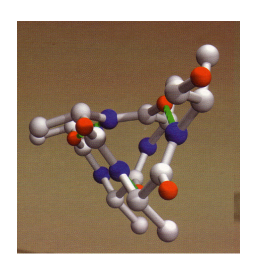
- ✓ A dipole is associated to each peptide unit;
- In an α-helix, these dipoles are aligned along the helix axis;
- ✓ Dipoli associated to the peptide units sum up to gice a helix macrodipole

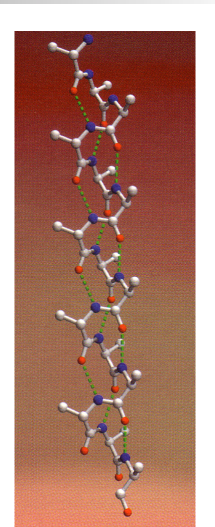


The 3₁₀ helix



- It winds up more tightly than an α -helix
- The C=O group of residue *i* forms a H-bond with the NH group of residue *i*+3

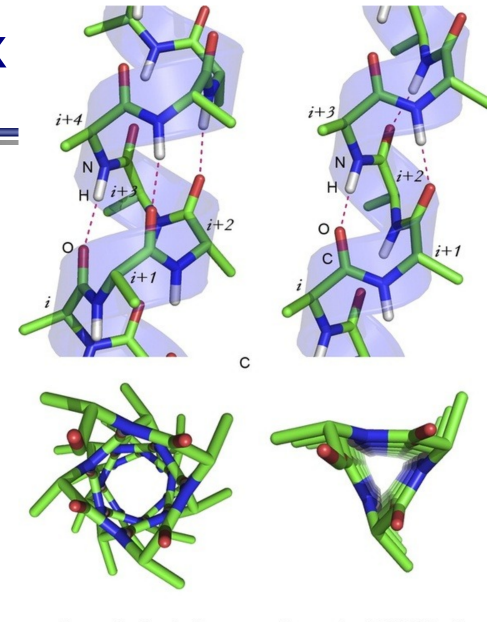




The 3₁₀ helix

The 3_{10} helices are less favored of the α helices:

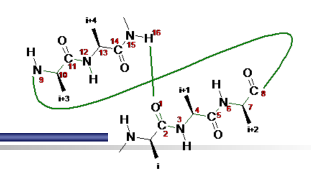
- the atoms of the main chain are too closely packed, leading to interactions of the van der Waals repulsion;
- hydrogen bonds are not linear;
- The dipoles of the peptide deviate of about 30° with respect to the helix axis;
- The position of the side chains (aligned) leads to steric interference.



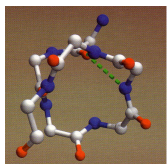
Canonical α -helix

Canonical 3(10) helix

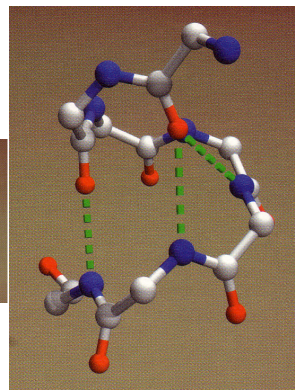
Π-helix



- It winds up less tightly than an α -helix
- The C=O group of residue i forms a H-bond with the NH group of residue i+5
- This type of extended π-helices are rare in proteins
- The π-helix can be outlined as a hollow cylinder.
- The cavity in the cylinder,
- however, is not such as to
- allow the entry of water molecules that may stabilize the helix.

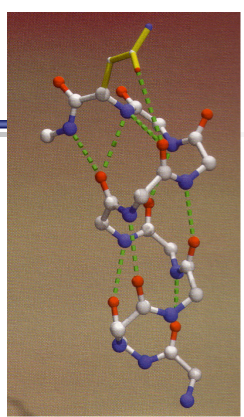


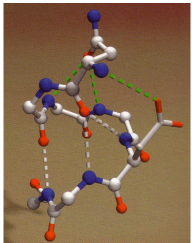
 This makes the π-helix structure a little favored from an energy point of view and extremely uncommon in proteins.



Helix capping

- Often special sequences and conformations appear as caps stabilising helix termini
- Helix-capping motifs are specific patterns of hydrogen bonding and hydrophobic interactions found at or near the ends of helices in both proteins and peptides.
- At the N-terminal helix there are 3 free NH groups, which do not form hydrogen bonds with CO groups, because there is no previous helix turn that provides the necessary partners.
- In this case the necessary partners to form hydrogen bonds are provided to the free -NH groups from the side chains of polar amino acids (Ser, Thr, Asp, Asn) which, in favorable cases, preceding the end dell'α-helix.
- These hydrogen bonds then occur between an element of the main polypeptide chain and one of the side chain.





Structural parameters for protein secondary structures

Structure	ф	Ψ	n	đ	р	
α-helix	-57	-47	3.6	1.5	5.5	
3 ₁₀ helix	-49	-26	3.0	2.0	6.0	
β-helix	-57	-70	4.4	1.1	5.0	
Polyproline II helix	-79	+149	3.0	3.1	9.4	
Parallel β strand	-119	+113	2.0	3.2	6.4	
Antiparallel β strand	-139	+135	2.0	3.4	6.8	

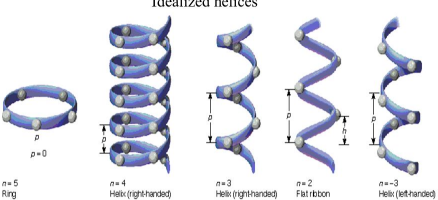
 ϕ and ψ are the conformational angles of the mainchain, with $\omega \sim 180^\circ$ (the trans conformation)

n = the number of residues per turn.

d = the displacement between successive residues along the helix axis.

p = the pitch of the helix, the distance along the helix axis of a complete turn. Note that $p = n \times d$. (The equation is exact; the values of p, n and d in the table have been rounded to two significant figures.)

Idealized helices



The pitch (p) of the helix is the distance parallel to the axis in which the helix makes one turn. There may be an integral number of residues/turn or not.

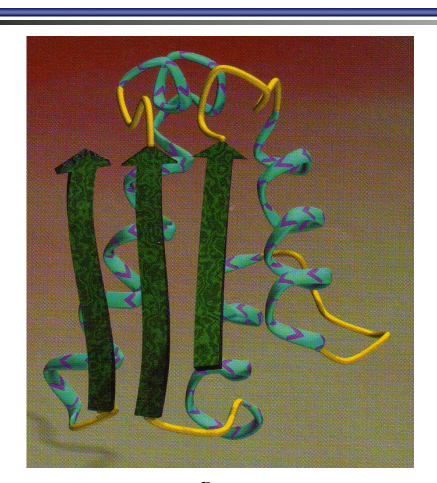
The rise of the helix is the distance parallel to the axis from the level of one residue to the next.

β -sheets

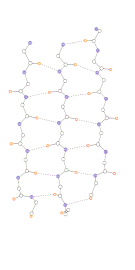


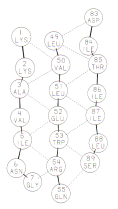
- β-sheets are formed by 2 strands distant in the sequence
- By contrast, helices have H-bonding patterns in the same strand
- They can be
 - Parallel sheets
 - Antiparallel sheets
 - Mixed sheets

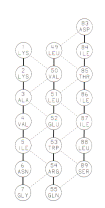
Parallel sheets

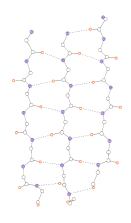


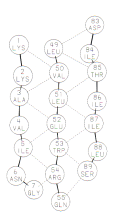
Barstar (inhibitor of Barnase = Bacterial RiboNucleASE), [1BRS]

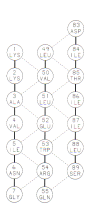


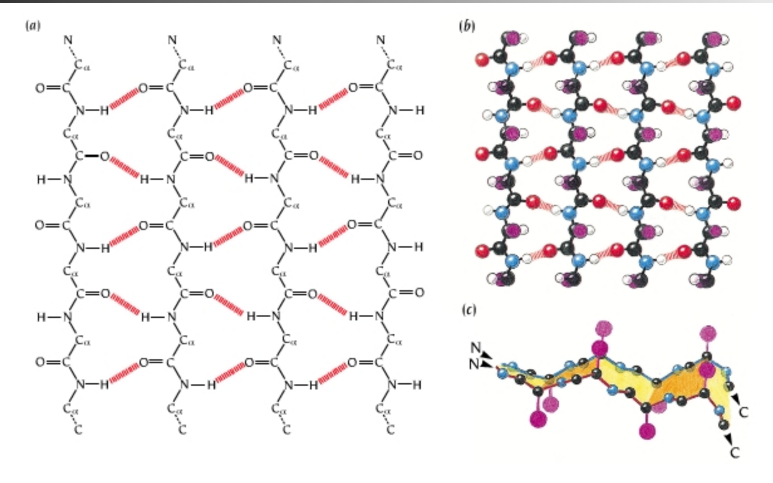




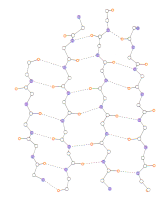


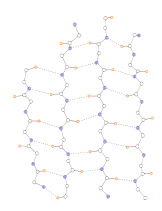




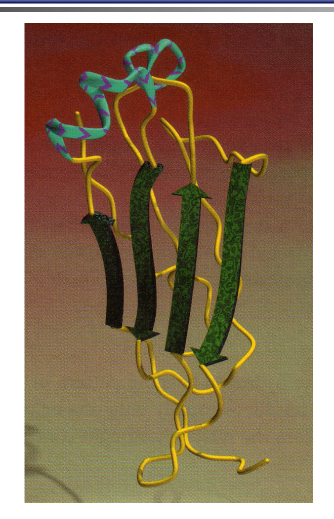


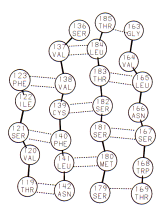
Antiparallel sheets

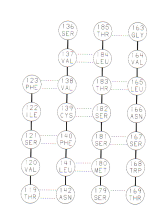


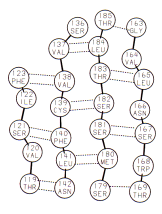


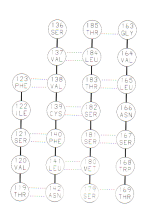
CL domain from Immunoglobulin TE33 [1TET]

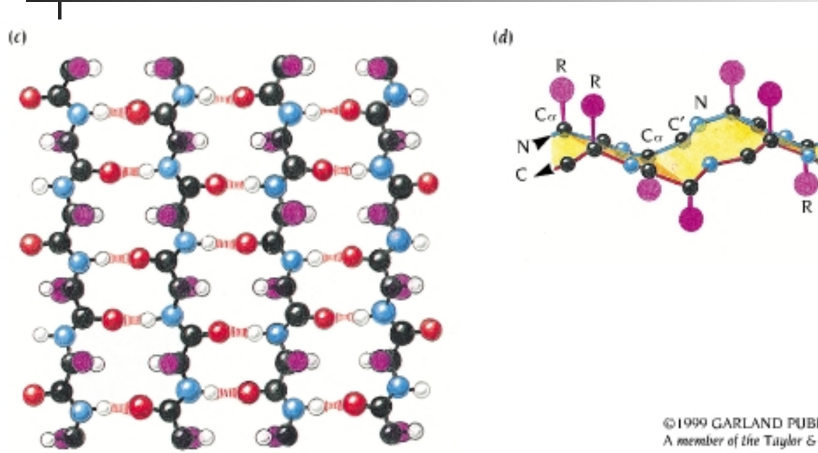








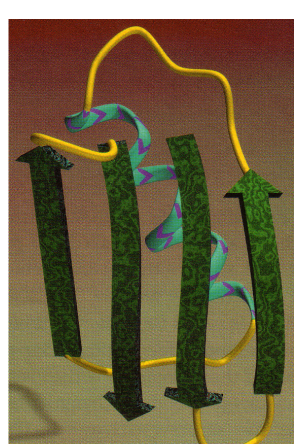


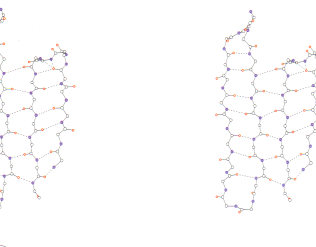


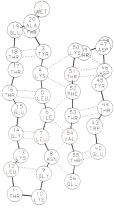
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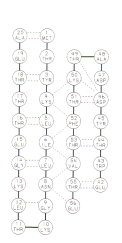
Mixed sheets

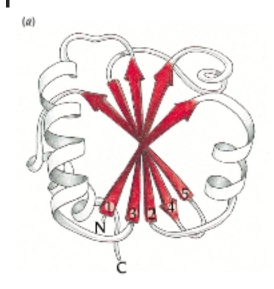
Streptococcal protein G B1 Immunoglobulin binding domain [1PGA]







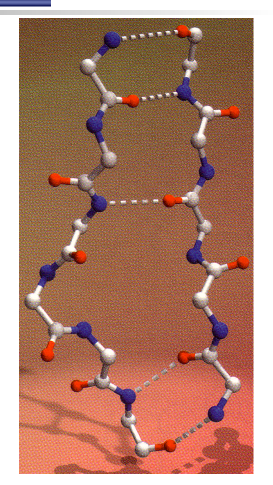




Thioredoxin from *E. coli*

The β-bulge

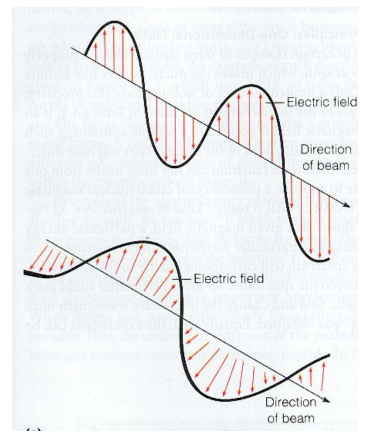
It is an irregularity in the Hbonding pattern of a sheet, where one or two - rarely more residues deviate from the regular patter

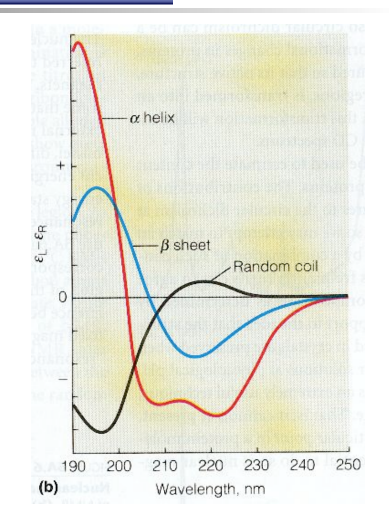


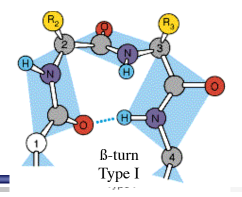
Fab J539 [2FBJ]

Study of protein 2ary structure

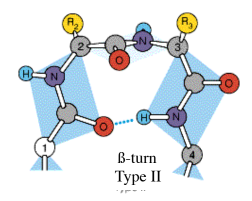
Circular dichroism:

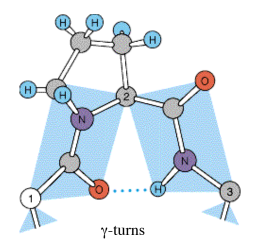






- 2. ß-sheets are usually twisted or wrapped into barrel structures
- 3. There are different types of turns
 - ß-turns:
 - Type I: 4 residues where the C=O of residue i makes H-bond with the N-H of residue (i + 3).
 - Type II: same as type I but residue 2 has a bulky R, so residue 3 is a Gly.
 - γ-turns:
 - Only 3 residues, of which only 1 is out of the Hbonding sequence. Usually Pro is involved.
- 4. Some parts of proteins cannot be classified as helix, ß-sheets or turns: these are 'random coil' regions. Usually are found at the N-ter and C-ter









Levels:

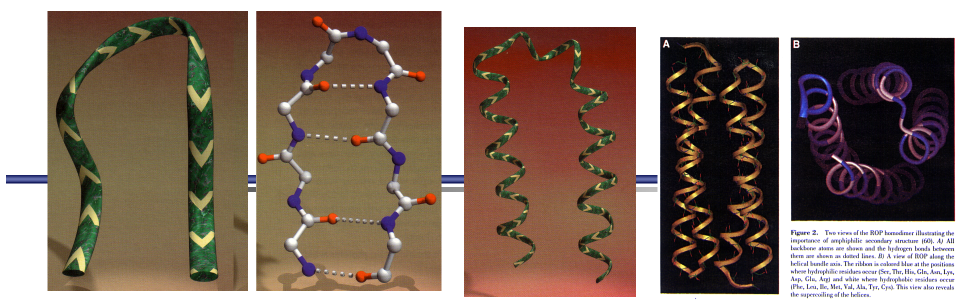
- Linderström-Lang has defined the classical levels of protein structure:
 - I-ary, II-ary, III-ary, IV-ary
- Additional levels have also been introduced:
 - Super-secondary structure:
 - These are recurrent patterns of interaction between helices and sheets close together in sequence. Examples are:
 - α-helical hairpin
 - β-hairpin
 - β -α- β unit
 - Coiled-coil

Domains:

These are compact units that fold independently; they are really boarded-line with III-ary structure

Modular proteins:

These are multi-domains proteins where there are domains repeating themselves. One example is fibronection, an extracellular protein involved in cell-adhesion formed by repeats of 3 identical domains, F1, F2 and F3 in the form of (F1)₆(F2)₂(F1)₃(F3)₁₅(F1)₃



 $\beta\text{-hairpin} \hspace{1cm} \alpha\text{-helical hairpin} \hspace{1cm} \text{coiled coil}$



β– α – β unit

There is a rough analogy between the analysis of protein structures at different levels, and the analysis of text.

The amino acids correspond to letters, the secondary structures to words, supersecondary structures to phrases (or even to clichés), elements of tertiary structure to sentences—this is the level at which true individuality makes its appearance domains to paragraphs, the structure of a full polypeptide chain to a chapter, and the quaternary structure to the assembly of chapters into a complete book.

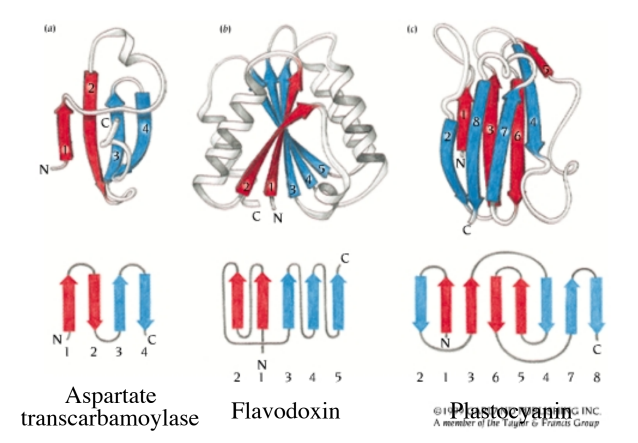


Classification of protein structures

TOPOLOGY

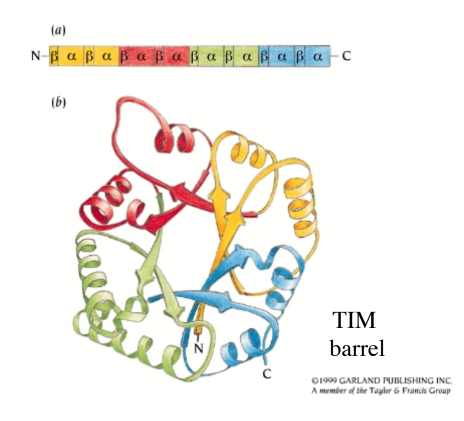
Topology diagrams

Useful for classification of protein structures



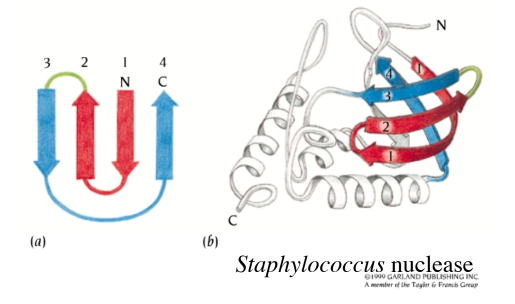
Domains are built from structural motifs

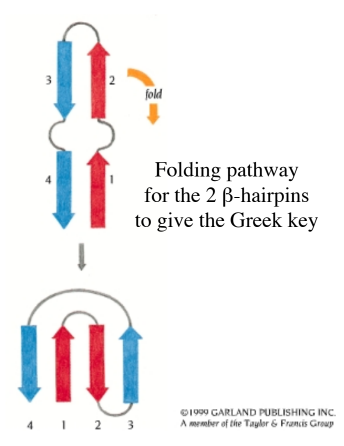
Triose phosphate isomerase is built from 4 β-α-β-α motifs



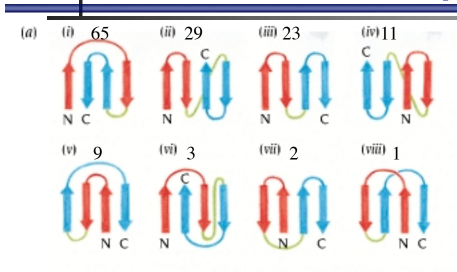
The Greek key motif

4 adjacent antiparallel βstrands (2 β-hairpins) are frequently arranged in the socalled Greek key

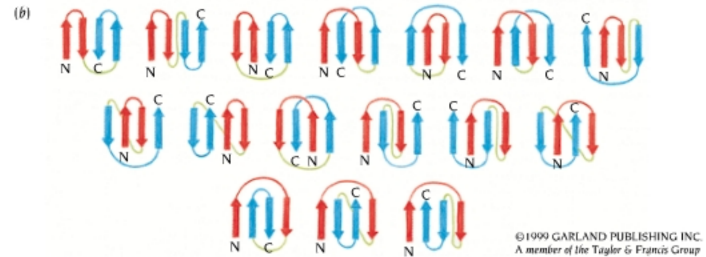




Simple motifs combine to form complex motifs



 2 adjacent β-hairpins can combine in 24 possible ways.



Protein domains can be divided in 3 main classes

- According to Michael Levitt and Cyrus Chothia the taxonomy of protein structures comprehends 3 main groups:
 - α-domains
 - β-domains
 - α/β -domains

Proteins can be broadly classified as:

- Fibrous = structural have only II-ary structure
 - Keratins
 - Fibroin
 - Collagen
 - Elastin
- Globular = functional (enzymes, Ab etc) also have III-ary structure
 - Myoglobin
 - Hemoglobin
 - Cyt c



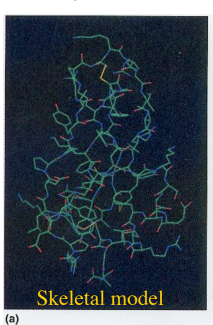
Proteins: Tertiary Structure

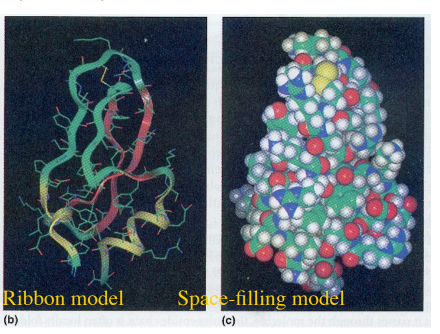
Globular proteins

- Perform different functions:
 - Synthesis
 - Binding and Transport
 - Catalysis and Metabolism
- They are compact structures folded on themselves
- Different ways to show/study/display proteins

Bovin Pancreatic Trypsin Inhibitor BPTI

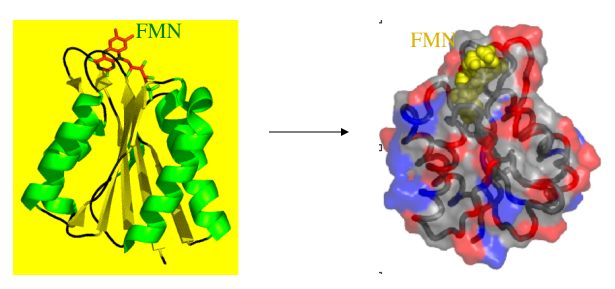
58 aa Prevents trypsin from catalysing peptide hydrolysis





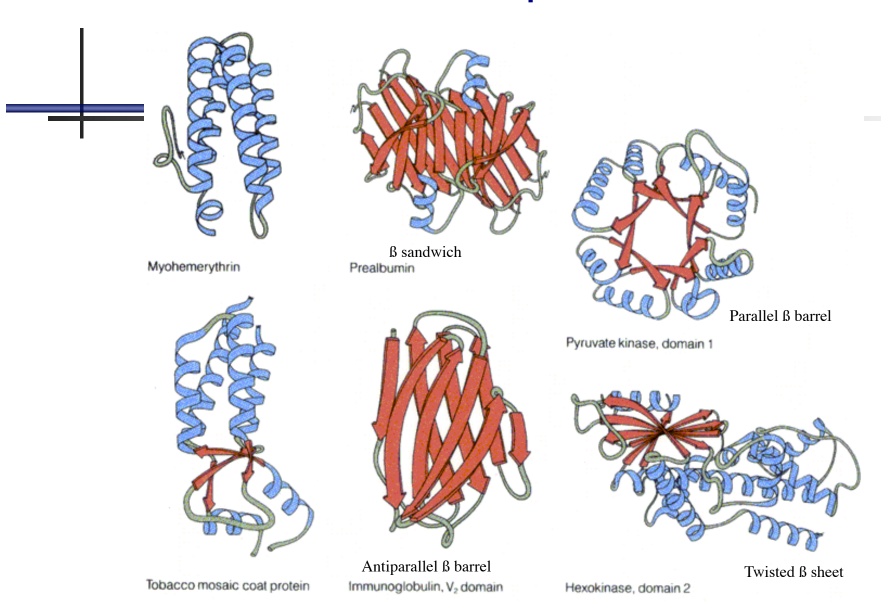
Other examples

- Many proteins have a prosthetic groups
 - These are small molecules non-covalently or covalently bound to the protein to enable function - for example redox centres:



Flavodoxin

Varieties of protein folds



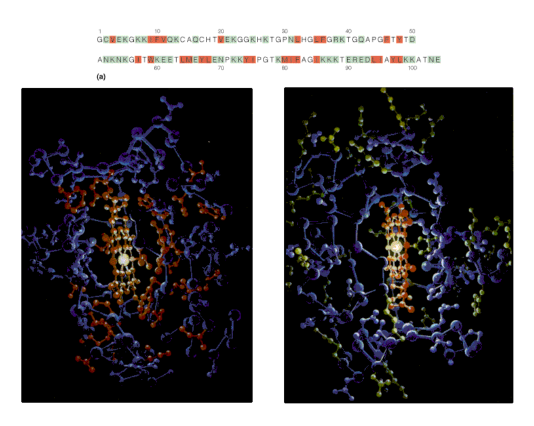
Helix bundles

Predominantly B sheet

Mixed \alpha/\beta

Rules for 3ary folds

- 1. All globular proteins have a define "inside" and "outside"
 - Hydrophilic (green) versus hydrophobic (red)



Horse heart cytochrome c