Molecular Cell Biology A

"Protein structure and pathways"

BIOX24ZL

Tuesdays 9-10:30

Ray LC

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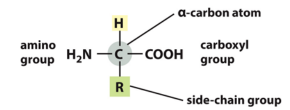
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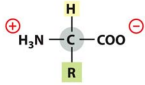
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THE AMINO ACID

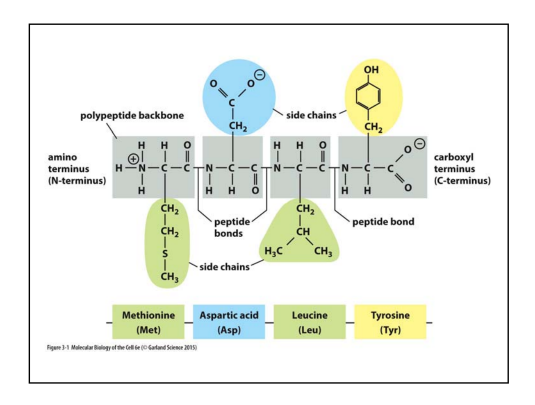
The general formula of an amino acid is

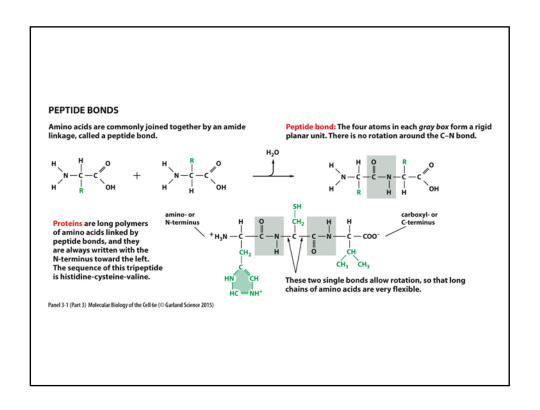


R is commonly one of 20 different side chains. At pH 7 both the amino and carboxyl groups are ionized.



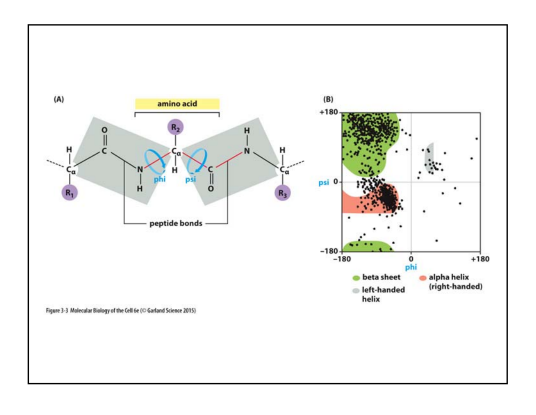
Panel 3-1 (Part 1) Molecular Biology of the Cell 6e (© Garland Science 2015)

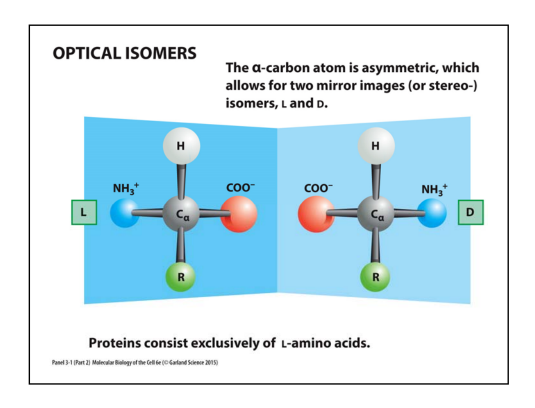


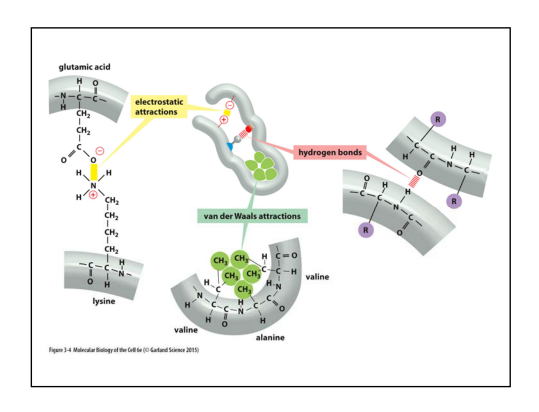


Proteins are specified by their amino acid sequence.

- Polypeptide backbone with AA side chains
- Bond angle restricted, weak noncovalent bond: H+, electrostatic, van der Waals determine final 3D shape
- Nonpolar (hydrophobic) side chains (Leu Phe Trp Val) cluster together interior to H+ water
- Polar (hydrophilic) side chains (Asp- Arg+ His+ Gln polar) H+ bond outside, backbone interact







	D	negative	Alanine	Ala		_
Glu	_		/1101111110	Ala	Α	nonpolar
	E	negative	Glycine	Gly	G	nonpolar
Arg	R	positive	Valine	Val	V	nonpolar
Lys	K	positive	Leucine	Leu	L	nonpolar
His	н	positive	Isoleucine	lle	1	nonpolar
Asn	N	uncharged polar	Proline	Pro	Р	nonpolar
Gln	Q	uncharged polar	Phenylalanine	Phe	F	nonpolar
Ser	S	uncharged polar	Methionine	Met	М	nonpolar
Γhr	т	uncharged polar	Tryptophan	Trp	w	nonpolar
Гуr	Υ	uncharged polar	Cysteine	Cys	c	nonpolar
5	lis Asn Gln Ger Thr	lis H Asn N Gln Q Ger S Thr T	ys K positive His H positive Asn N uncharged polar Gln Q uncharged polar For S uncharged polar Thr T uncharged polar	Leucine Isoleucine Isoleucine Isoleucine Isoleucine Proline Isoleucine Proline Proline Phenylalanine Isoleucine Proline Proline Phenylalanine Tryptophan	Leucine Leu Leucine Ile Leucine Leu Leucine Leucine Leucine Leucine	Leucine Leu L Isoleucine Ile I Isoleucine Pro P Isoleucine Pro P Isoleucine When Pro P Isoleucine Ile I Isoleucine Pro P Isoleucine Ile I Isoleucine When Pro P Isoleucine Ile I Isole

FAMILIES OF AMINO ACIDS

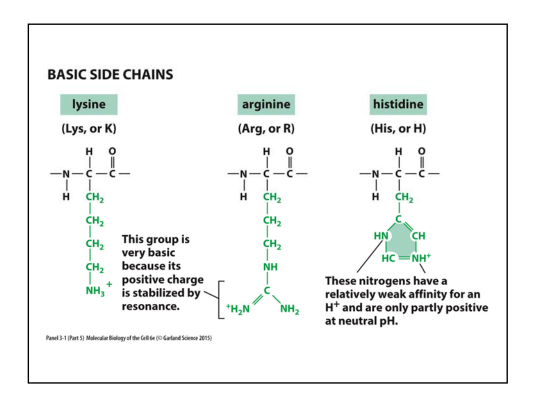
The common amino acids are grouped according to whether their side chains are

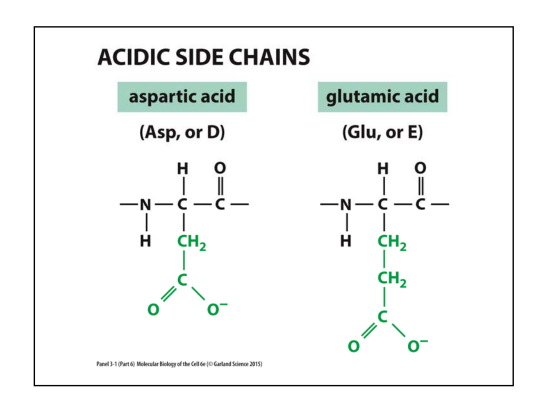
acidic basic uncharged polar nonpolar

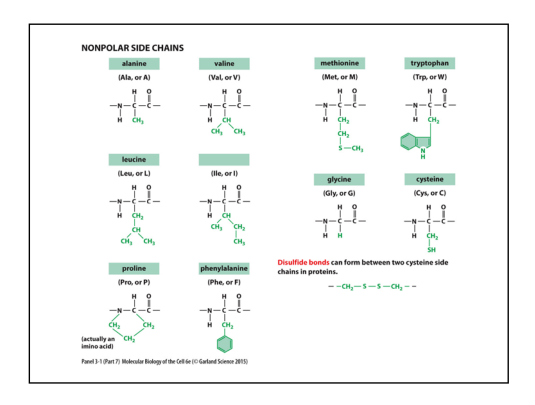
These 20 amino acids are given both three-letter and one-letter abbreviations.

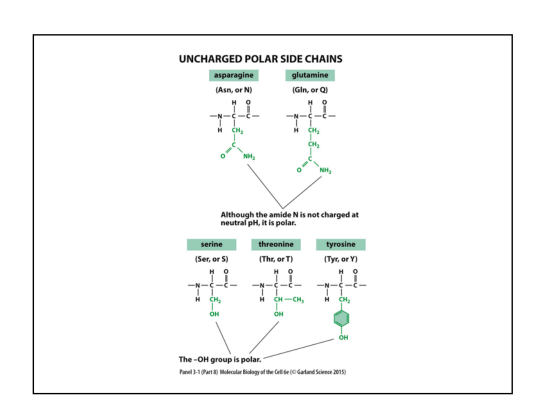
Thus: alanine = Ala = A

Panel 3-1 (Part 4) Molecular Biology of the Cell 6e (© Garland Science 2015)



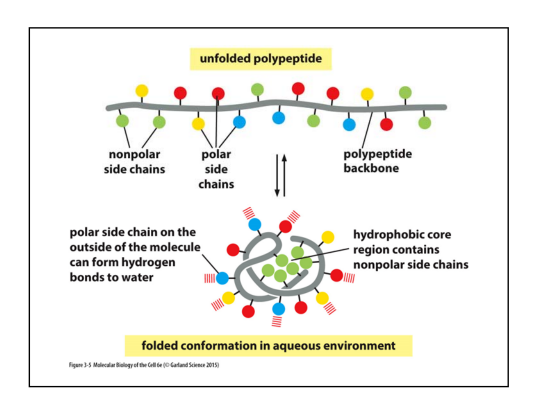


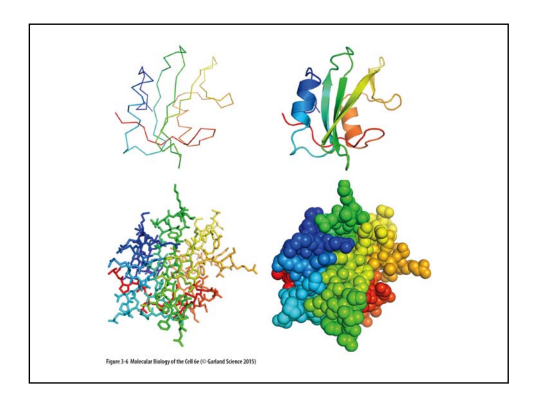




Proteins fold into unique 3D conformations of lowest energy.

- Study by denaturing noncovalent bonds (urea), allow to renature from flexible chain
- Conformation altered if interact with cell mol
- Chaperones: prevent hydrophobic aggregates making folding more reliable, assist folding
- SH2 (GTGA domains) 100AA backbone, ribbon, wire, space-filling models





Team work.

Which of the following pairs of amino acid residues would you expect to form ionic bonds?

- A. Glutamic acid and glutamine
- B. Arginine and lysine
- C. Tryptophan and tyrosine
- D. Tyrosine and glutamine
- E. Lysine and glutamic acid

Which of the following stretches of amino acid residues would you expect to find in the interior of protein molecules?

- A. Ala-Val-Leu-Ile-Trp
- B. Ala-Asp-Asp-Tyr-Arg
- C. Phe-Glu-Gln-Glu-Asn
- D. Gly-Tyr-His-Arg-His
- E. Gly-Lys-Ser-Pro-Thr

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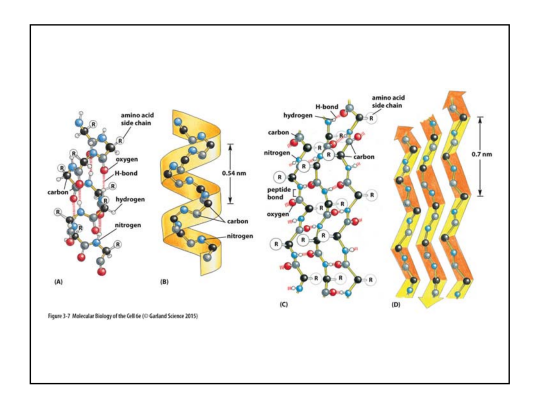
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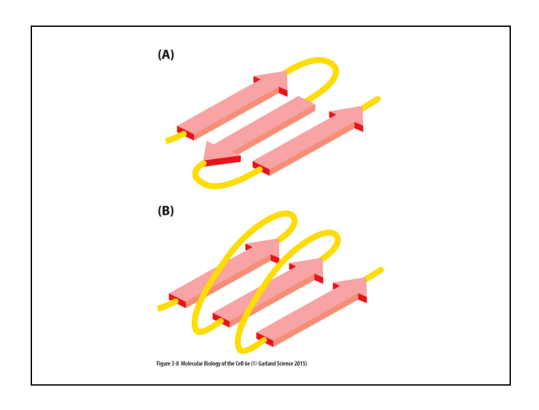
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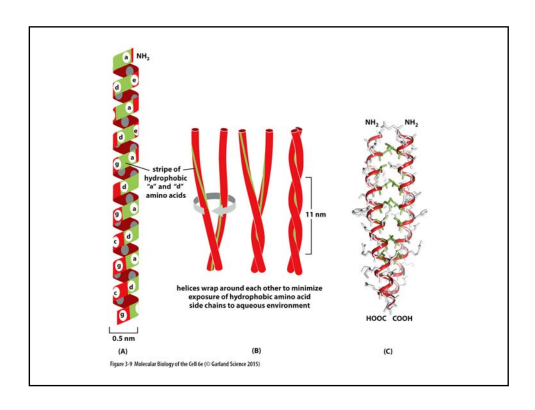


Proteins form common patterns (alpha helix, beta sheet) by backbone H bond.

- Alpha helix: keratin in skin hair, N-H (up) H bond to C=O (down) 4 away, turn every 3.6, C end part neg, N part pos, membrane proteins inside H bond each other, outside nonpolar
- Beta sheet: fibroin in silk, H bond between diff chains, R alternate up down, protein core rigid
- Beta sheet run in parallel (same orientation) or anti-parallel (fold back on itself) directions
- Coiled coil alpha helix 2-3 chains, nonpolar on one side inward so twist around each other

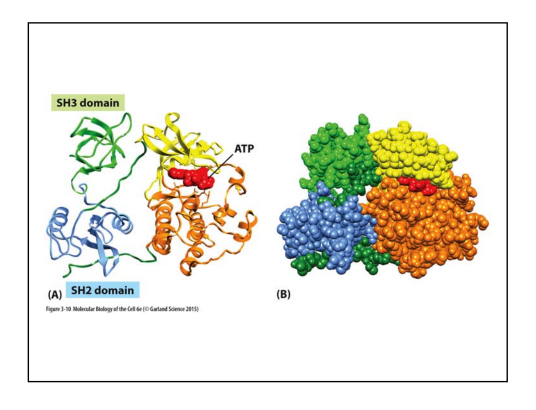


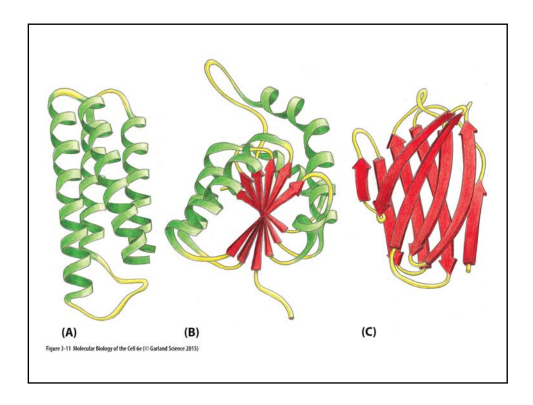




Proteins are characterized by four (plus one) levels of structure.

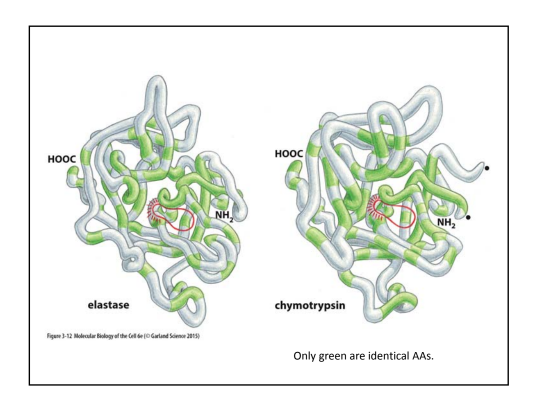
- Primary structure: AA sequence
- Secondary structure: alpha helix, beta sheet
- Tertiary structure: 3D peptide organization
- Quaternary structure: multiple peptide chains
- Domains: part of chain into stable modules, e.g. Src Kinase SH2 SH3 regulate C-term kinase
- 20ⁿ possible seq but evolution selects for only 1 in a billion for its stable structure

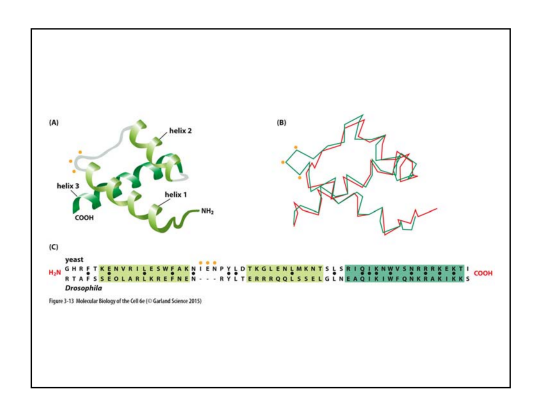


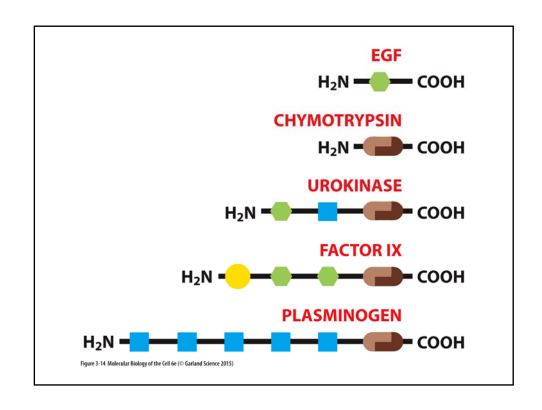


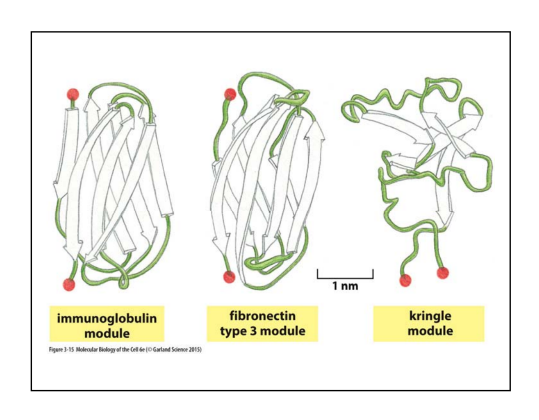
Protein diversity result from reuse of same modules as homologs.

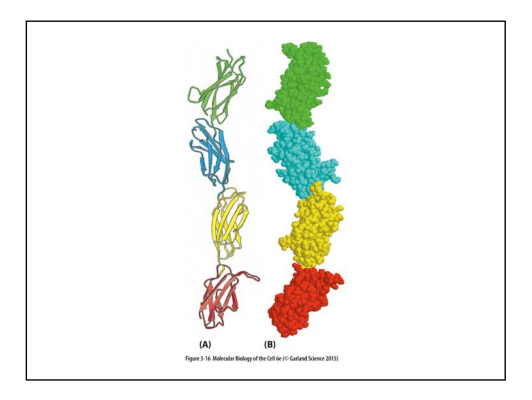
- Families of protein from duplicated mutations
- Diff AA seq (25% same) still similar structure due to small number of characteristic shapes
- Pattern matches in signature seq allow identification with proteins of known function
- Domain shuffling forms new proteins from existing motifs, binding sites mutated for diff ligands, N-C terms at opposite or same ends





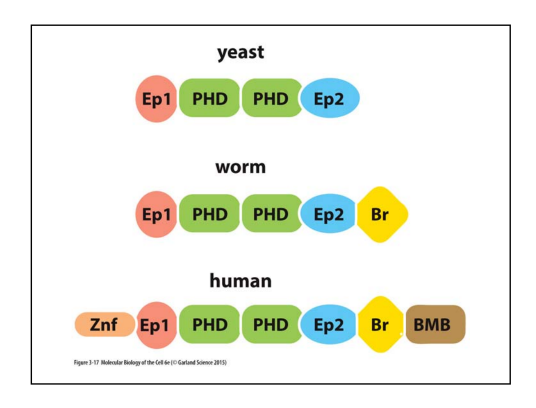


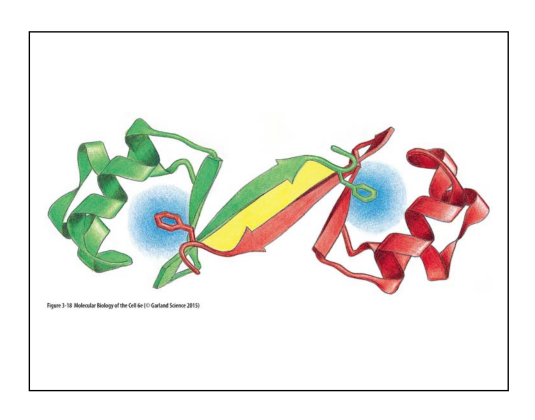


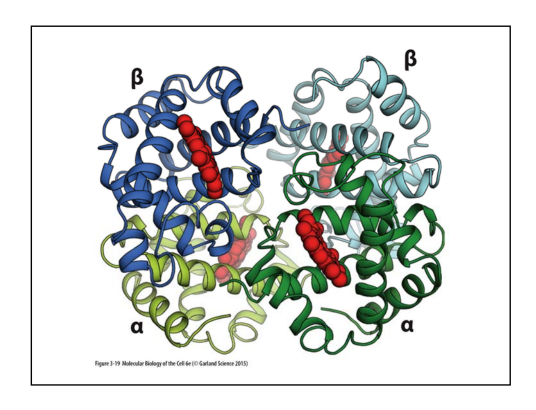


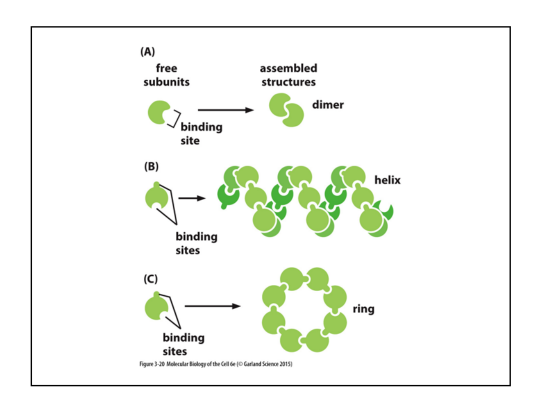
Diversity of protein structures allow for adaptability.

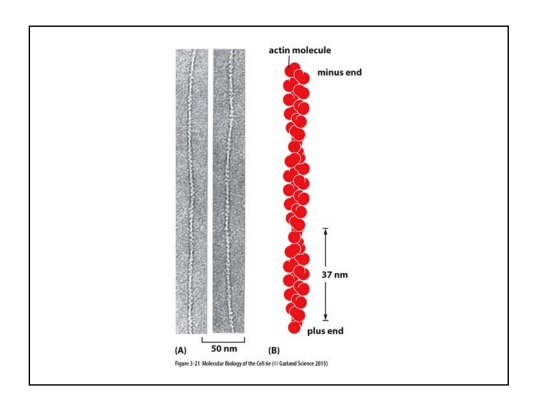
- Half of domains shared between all organisms only 5% of 2-domain combos shared ev recent
- · Humans: similar genes vastly shuffled domains
- Binding sites noncovalent interactions with other polypeptides lead to multi-chain proteins
- Repeat helical filaments common, e.g. actin
- Fibrous proteins span large distance in extracellular matrix, e.g. collagen triple helix
- Unstructured chains allow stretch, e.g. elastin

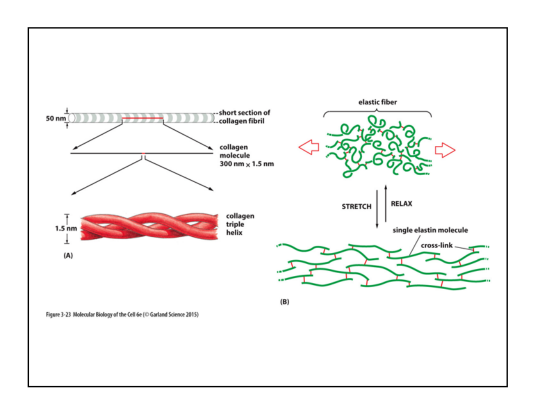






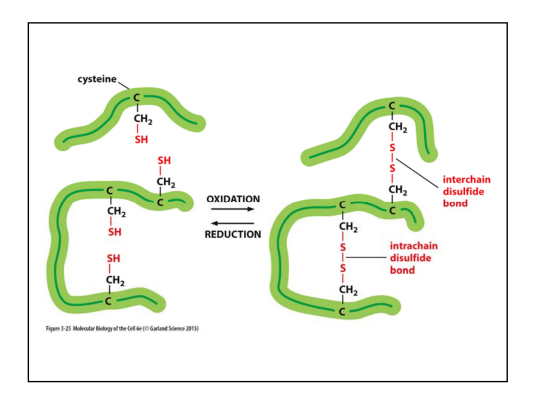


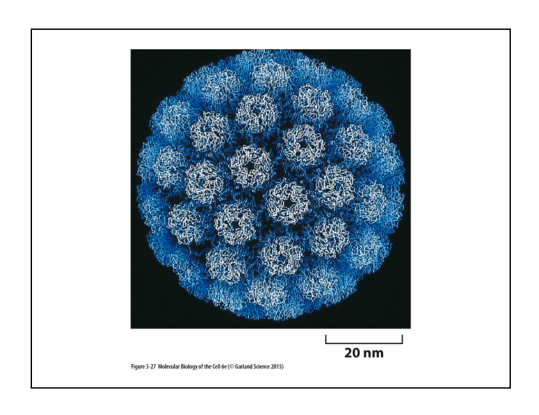


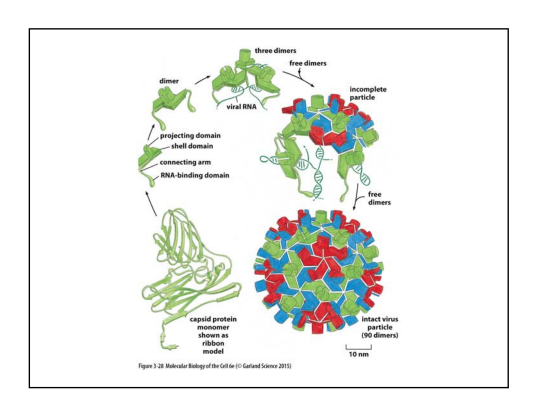


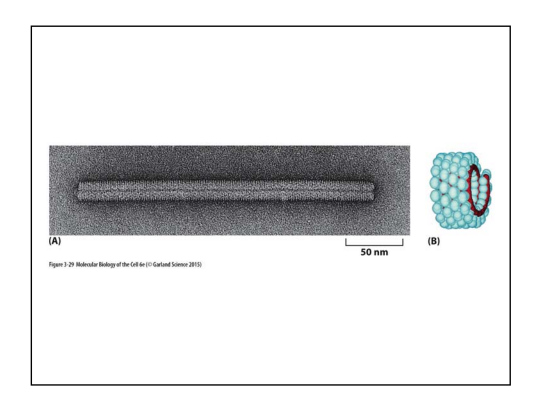
Protein assemblies are formed by various means.

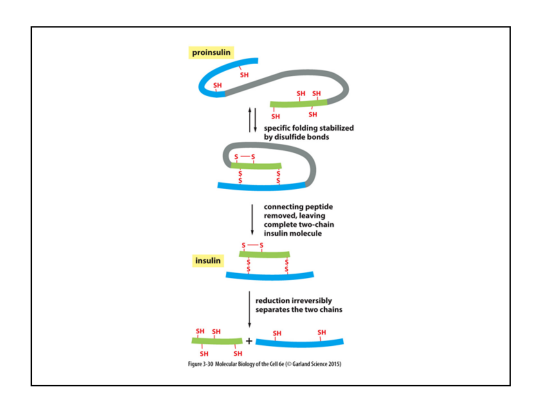
- Disulfide bonds (SH cysteine) cross-links proteins together in extracellular environ only, reducing (provide electrons) to separate
- Protein subunits assemblies, e.g. icosohedral viral capsid
- Self assembly: tobacco mosaic virus, ribosome
- Irreversibility: cannot self assemble after proteolytic cleavage, mitochondria











Team work.

Protein secondary structure elements such as α helices and β sheets constitute the major regular folding patterns in proteins. With regard to these elements, \dots

- the folding patterns result from hydrogen-bonding between the N–H and C=O groups in the polypeptide backbone.
- В. a certain short amino acid sequence always adopts the same secondary structure.
- hydrogen-bonding between the amino acid side chains defines the type of secondary structure.
- only a few specific amino acid sequences can adopt these repetitive structures. D.
- All of the above.

You have purified a multisubunit extracellular protein that has several interchain disulfide bonds. Which of the following chemicals would you add to your purified protein mixture if you wanted to eliminate the disulfide bonds?

- NaCl, a salt
- DTT, a reducing agent
- C. $\rm H_2O_2$, an oxidizing reagent
- SDS, an ionic detergent and denaturing agent
- Tris, a buffering agent

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