#### What are proteins?



•Proteins are important; e.g. for catalyzing and regulating biochemical reactions, transporting molecules, ...

•Linear polymer chain composed of tens (peptides) to thousands (proteins) of monomers

- •Monomers are 20 naturally occurring amino acids
- •Different proteins have different amino acid sequences
- •Structureless, extended unfolded state
- •Compact, 'unique' native folded state (with secondary and tertiary structure) required for biological function
- •Sequence determines protein structure (or lack thereof)
- •Proteins unfold or denature with increasing temperature or chemical denaturants

#### Amino Acids I



Side chains differentiate amino acid repeat units
Peptide bonds link residues into polypeptides

#### Amino Acids II



# The Protein Folding Problem:

What is 'unique' folded 3D structure of a protein based on its amino acidsequence?Sequence  $\rightarrow$  Structure

Lys-Asn-Val-Arg-Ser-Lys-Val-Gly-Ser-Thr-Glu-Asn-Ile-Lys- His-Gln-Pro- Gly-Gly-Gly-...



Why do proteins fold (correctly & rapidly)??

### Levinthal's paradox:

For a protein with N amino acids, number of backbone conformations/minima

 $N_c \sim \mu^{2N}$   $\mu = \#$  allowed dihedral angles

How does a protein find the global optimum w/o global search? Proteins fold much faster.

$$N_c^{~~} 3^{200} \sim 10^{95}$$
  
 $\tau_{fold}^{~~} N_c^{~} \tau_{sample}^{~~} \sim 10^{83} s$  VS  $\tau_{fold}^{~~} \sim 10^{-6} - 10^{-3} s$   
 $\tau_{universe}^{~~} \sim 10^{17} s$  5



#### Roughness of Energy Landscape



#### Critical Assessment of Structure Prediction (CASP)





#### **Driving Forces**

•Folding: hydrophobicity, hydrogen bonding, van der Waals interactions, ...

•Unfolding: increase in conformational entropy, electric charge...

	1	At pH 2 <sup>*</sup>		At pH 7°	
		Very Hydrophobic			
inside		Leu	100	Phe	100
	H (hydrophobic)	lle	100	lle	99
		Phe	92	Trp	97
		Trp	84	Leu	97
		Val	79	Val	76
		Met	74	Met	74
		Hydrophobic			
		Cys	52	Tyr	63
		Tyr	49	Cys	49
		Ala	47	Ala	41
		Neutral			
outside		Thr	13	Thr	13
		Glu	8	His	8
		Gly	0	Gly	0
		Ser	-7	Ser	-5
		Gln	-18	Gln	-10
		Asp	-18		
		Hydrophilic			
	r (polar)	Arg	-26	Arg	-14
		Lys	-37	Lys	-23
		Asn	-41	Asn	-28
		His	-42	Glu	-31
		Pro	-46	Pro	-46 (used pH 2)
				Asp	-55
		^	form for the shall of the second	120.153 (100.0	

Hydrophobicity index

<sup>A</sup>pH 2 values: Normalized from Sereda et al., J. Chrom. 676: 139-153 (1994 <sup>B</sup>pH 7 values: Monera et al., J. Pept. Sci. 1: 319-329 (1995).

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#### Solvent Accessible Surface Area and rSASA





 $rSASA=SASA_{res}/SASA_{dip} = [0,1]$ 



FIGURE 5 Fractions of amino acids with A, rSASA  $\leq 10^{-3}$  and B, rSASA>0.5 for residues in the Dun1.0 (grey), PPI (blue), and TM (red) datasets. The fractions are defined relative to the total number of residues in each rSASA category. C, The fractions of core residues (light bars) and non-core residues (rSASA>0.5, dark bars) among the 11 non-charged residues (Ala, Gly, Ile, Leu, Met, Phe, Ser, Thr, Trp, Tyr, and Val) [Color figure can be viewed at wileyonlinelibrary.com]

#### Secondary Structure: Loops, $\alpha$ -helices, $\beta$ -strands/sheets



•Right-handed; three turns

•Vertical hydrogen bonds between NH<sub>2</sub> (teal/white) backbone group and C=O (grey/red) backbone group four residues earlier in sequence

Side chains (R) on outside; point upwards toward NH<sub>2</sub>
Each amino acid corresponds to 100°, 1.5Å, 3.6 amino acids per turn

•(φ,ψ)=(-60°,-45°)

• $\alpha$ -helix propensities: Met, Ala, Leu, Glu

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•5-10 residues; peptide backbones fully extended
•NH (blue/white) of one strand hydrogen-bonded to C=O (black/red) of another strand

- $^{\bullet}\text{C}_{\alpha}$  ,side chains (yellow) on adjacent strands aligned; side chains along single strand alternate up and down
- •(φ,ψ)=(-135°,135°)

Phe, lle

• $\beta$ -strand propensities: Val, Thr, Tyr, Trp,

#### N<sub>s</sub>=62,938 monomeric xtal structures



#### **Bond Angles**



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**Backbonde Dihedral Angles** 



# 3N-6 DoF -(N-1) Bond lengths -(N-2) Bond angles =N-3 Dihedral angles

 $φ: C'^{i-1}NC_{\alpha}C'$  $\psi$ : NC<sub> $\alpha$ </sub>C'N<sup>i+1</sup>  $\omega_1{:}C^{i\text{-}1}{}_\alpha C'^{i\text{-}1}NC_\alpha$  $\omega_2: C_{\alpha}C'N^{i+1}C^{i+1}{}_{\alpha}$ 



Оc

Он Os

DN DO

#### Ramachandran Plot: Determining Steric Clashes



4 atoms define dihedral angle:

$$C_{-1}NC_{\alpha}C \qquad \phi$$

$$C_{\alpha,-1}C_{-1}NC_{\alpha} \qquad \omega=0,180^{\circ}$$

$$NC_{\alpha}CN_{+1} \qquad \Psi$$



## Backbone dihedral angles from PDB



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#### Dunbrack 1.0



**Figure 5.** Probability distributions  $P(\phi, \psi)$  for the backbone dihedral angles  $\phi$  and  $\psi$  obtained from MD simulations of an Ala dipeptide mimetic using recent versions of the CHARMM and Amber force fields, their associated optimized water models, and with and without the "ILDN-NMR" and "CMAP" dihedral angle potential corrections: (a) Amber99sb+TIP4P-Ew, (b) Amber99sb-ILDN-NMR + TIP4P-Ew, (c) CHARMM27 + TIP3SP, and (d) CHARMM27-CMAP+TIP3SP. Subpanels (e) and (f) correspond to the Ala  $\phi$ - $\psi$  distributions from the Dunbrack Database<sup>38</sup> and the Wu "Coil-3" library,<sup>10</sup> respectively. The Ramachandran hard-sphere<sup>3</sup> normal and outer limits (pink and blue lines, respectively) for  $\tau = 110^{\circ}$  are overlaid on each panel. The Amber and CHARMM MD simulations were thermally equilibrated at 303 K and sampled for 500 ns.



Figure S1: Stick representations of (a) Ile, (b) Phe, (c) Val, (d) Tyr, (e) Trp, (f) Leu, (g) Thr, and (h) Ser dipeptide mimetics. The carbon, nitrogen, oxygen, and hydrogen atoms are shaded green, blue, red, and white, respectively. The side chain dihedral angles  $\chi_1$  and  $\chi_2$  and several key atoms are labeled. The residues before (*i*-1) and after (*i*+1) the *i*th central residue are labeled at the  $C_{\alpha}$  atom.

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Thr



lle



1. Can the structural properties of protein cores be quantitatively modeled using hard-spheres?

2. What is the packing fraction in protein cores?

3. Can simple hard-sphere model improve computational design of protein-protein interactions?