

# Molecular Modeling 2021

## lecture 4 -- Fri Feb 5

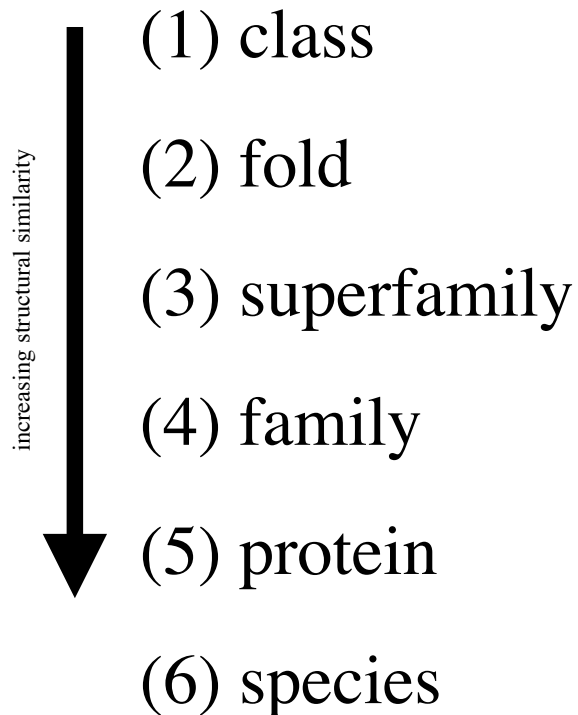
Protein folding

Local structure

Handedness

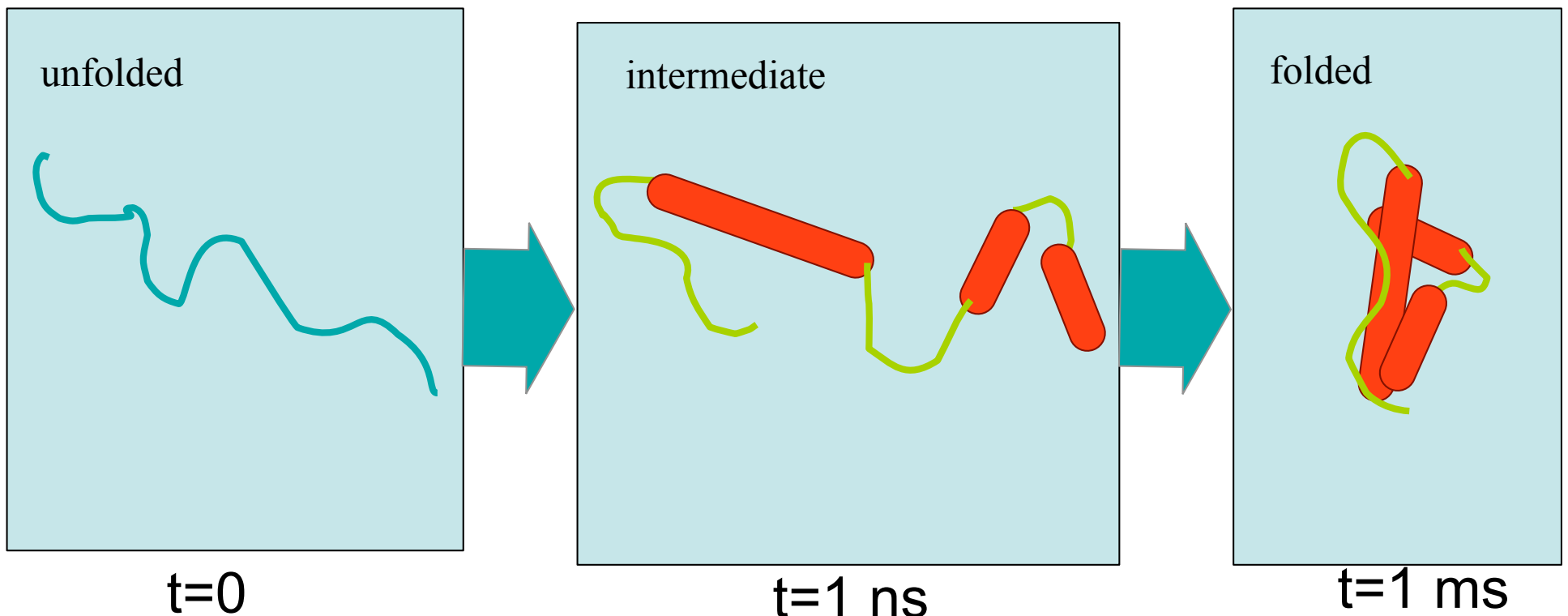
# SCOP -- a hierarchy

■ <http://scop.berkeley.edu>



**SCOP's hierarchy is sequence centered**

# Folding -- also a hierarchy

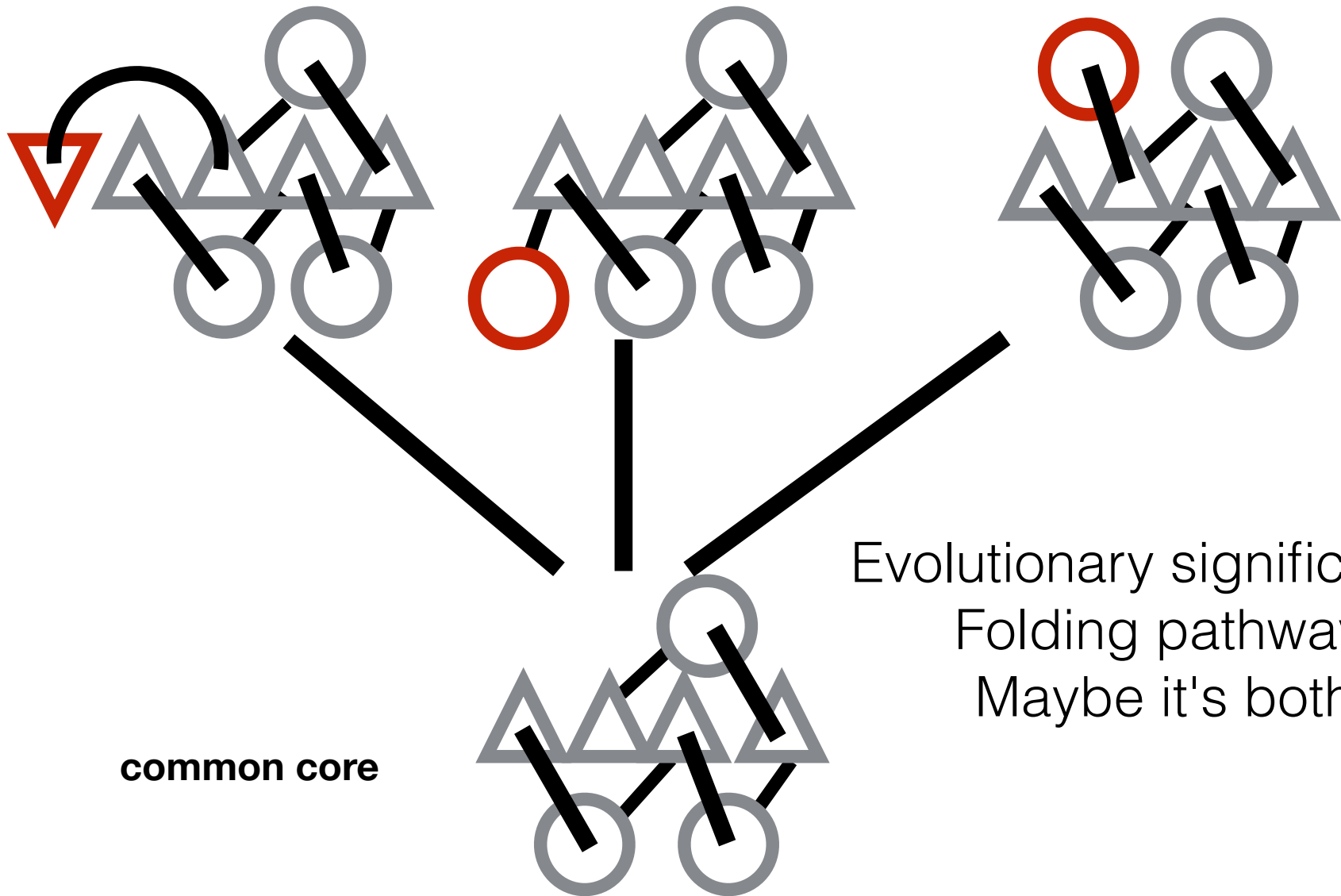


t = time after leaving the ribosome

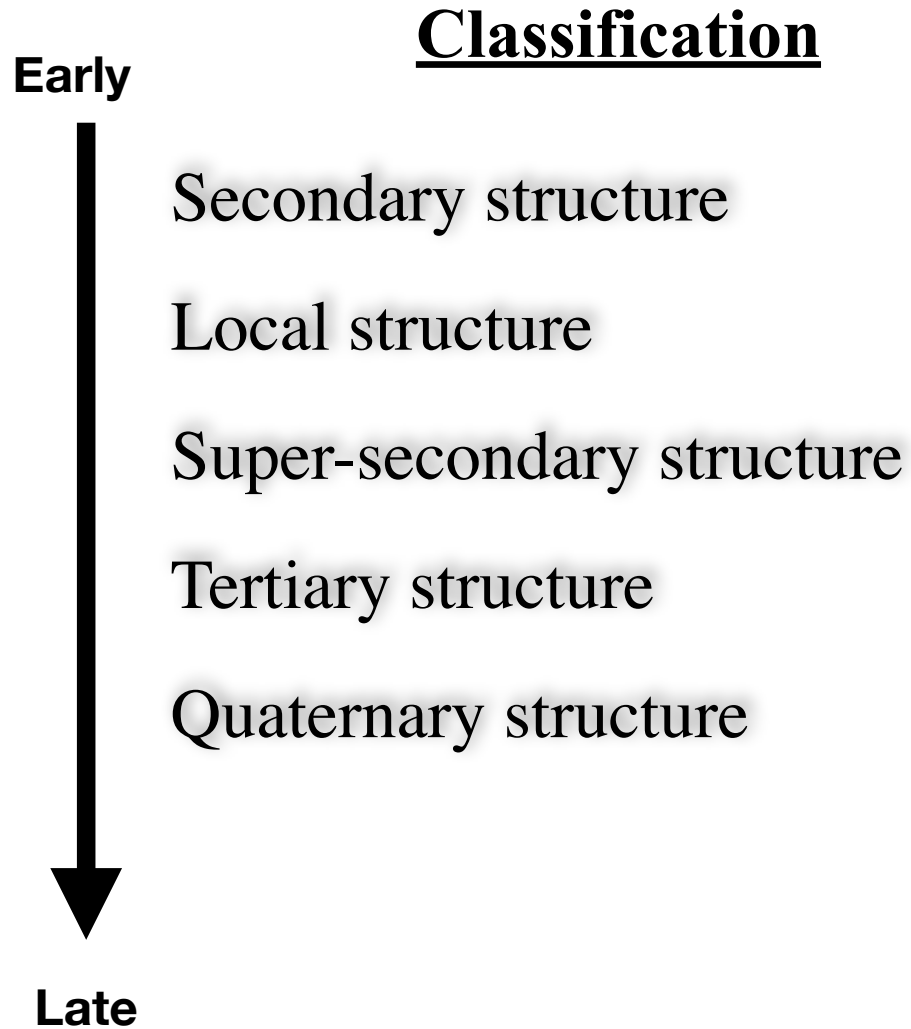
**More about protein folding in later lectures**



# How to grow a Efimov structural tree.



# A classification / folding hierarchy



Folding step 1?

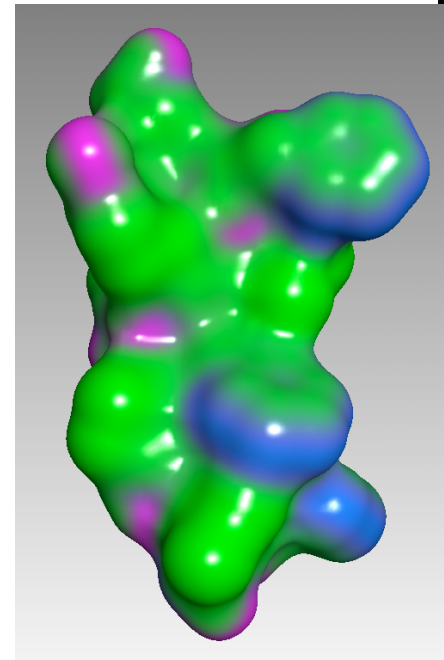
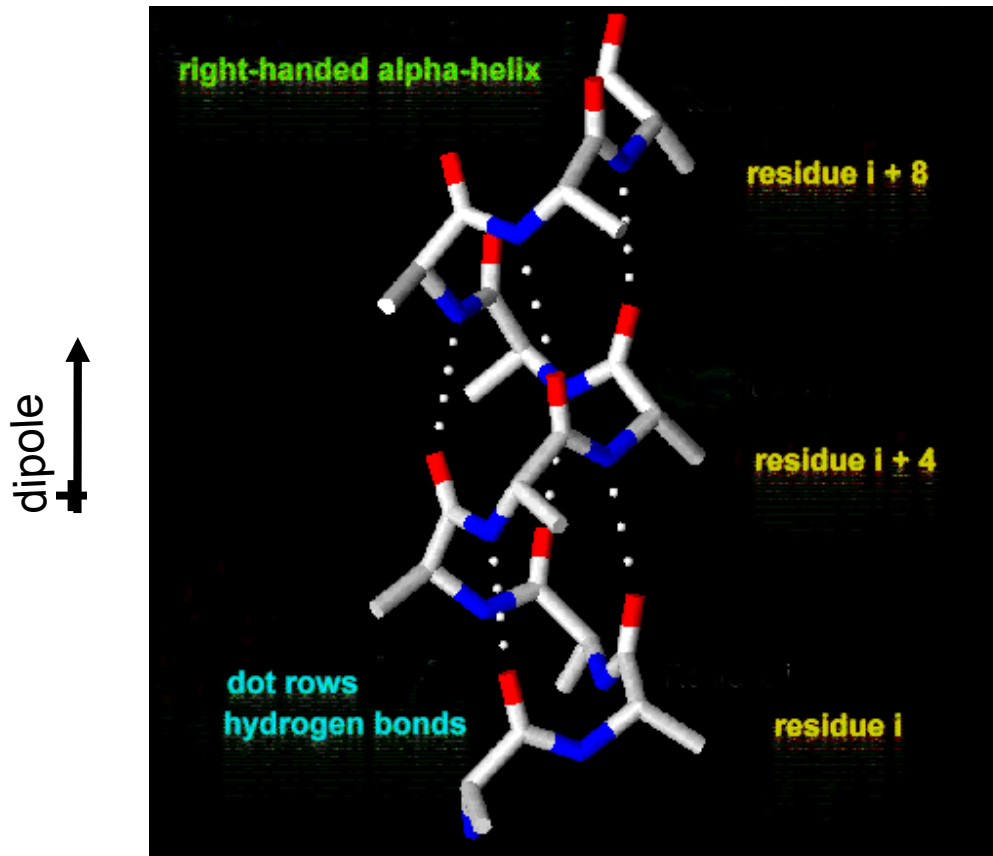
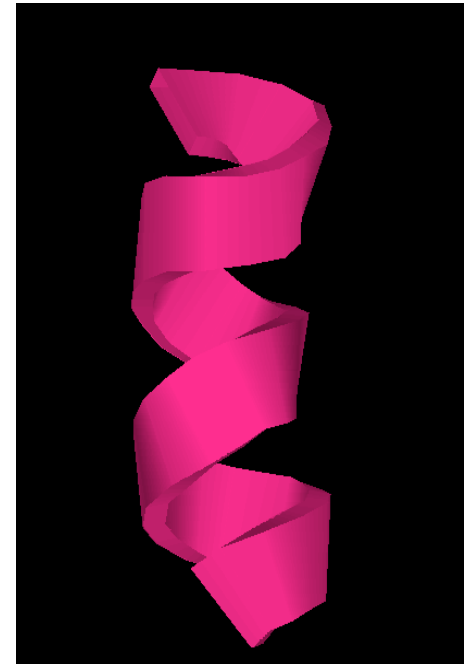
# Secondary structure

# About the Alpha helix

Right-handed helix. H-bond is from the oxygen at  $i$  to the nitrogen at  $i+4$ .  $\alpha$ -helices have an overall dipole because the H-bonds are all in the same direction. Must be  $> 3$  residues.

H-bond rule for donor to acceptor (NH->O):  $i$  to  $i+4$

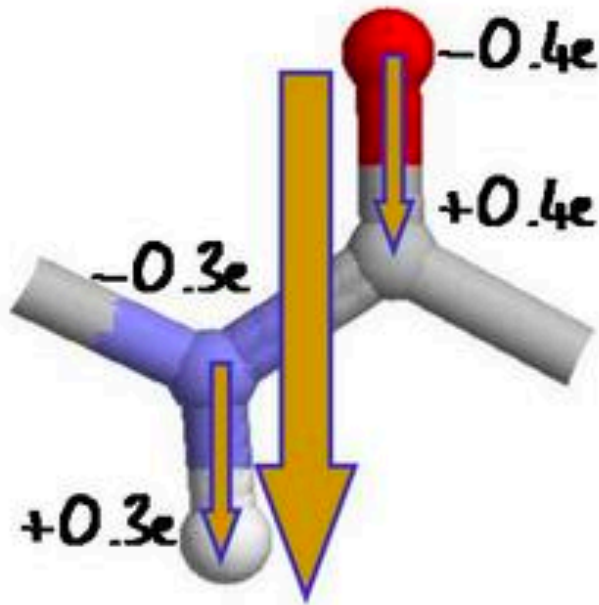
helices  
are right-  
handed



Helices are  
bumpy, not  
"cylindrical".



# The peptide dipole make Hydrogen bonding stronger.



- The peptide group has a strong dipole moment due to partial charges on NH and CO groups.

# Sequence pattern for the amphipathic alpha helix

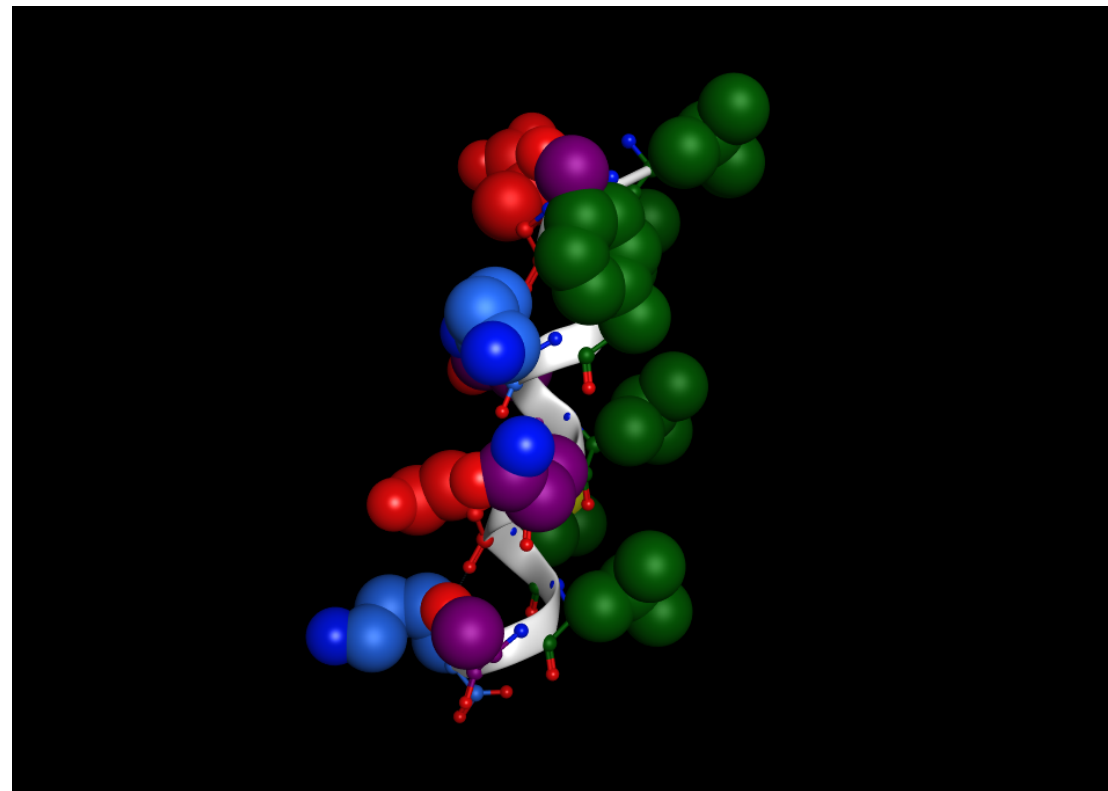
- nppnpp,  
where n = non-polar, p =  
polar
- Example:

**LSELFKNLQDMLSK**

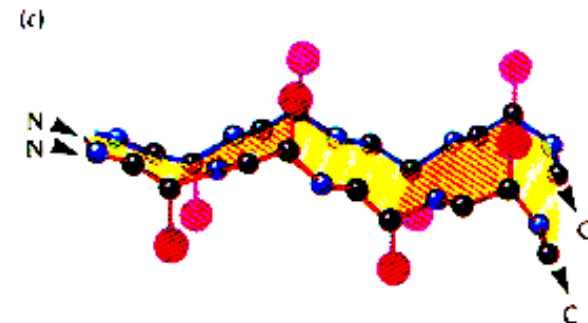
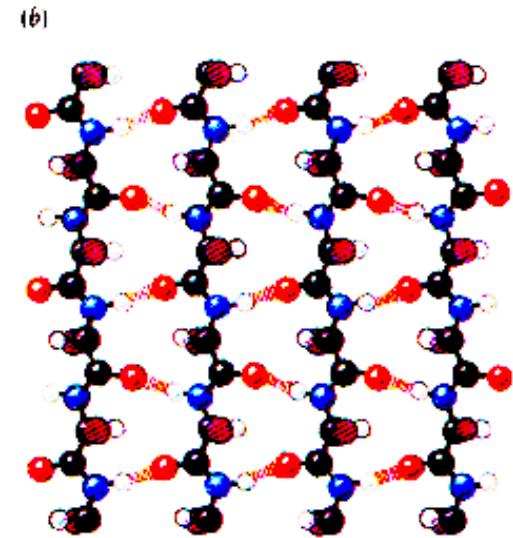
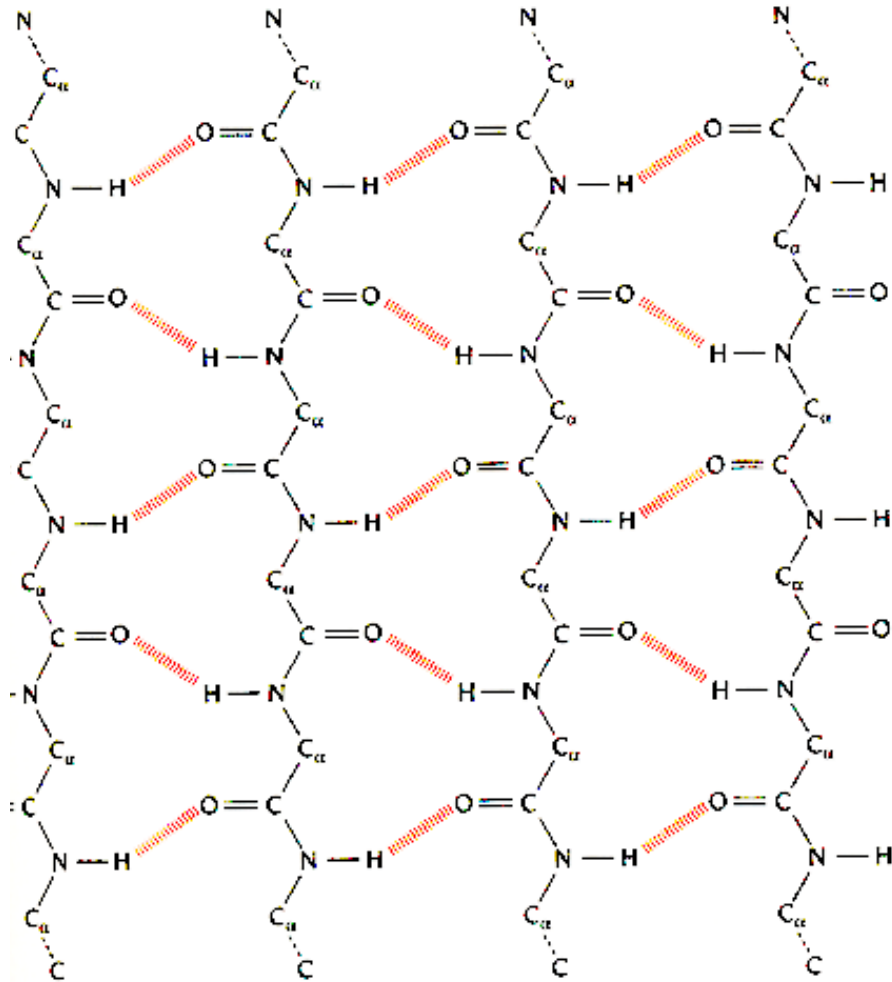
The helix is held together by  
the hydrophobic effect.

Amphipathic helices stick to  
other amphipathic helices.

Hydrophobic all on one side

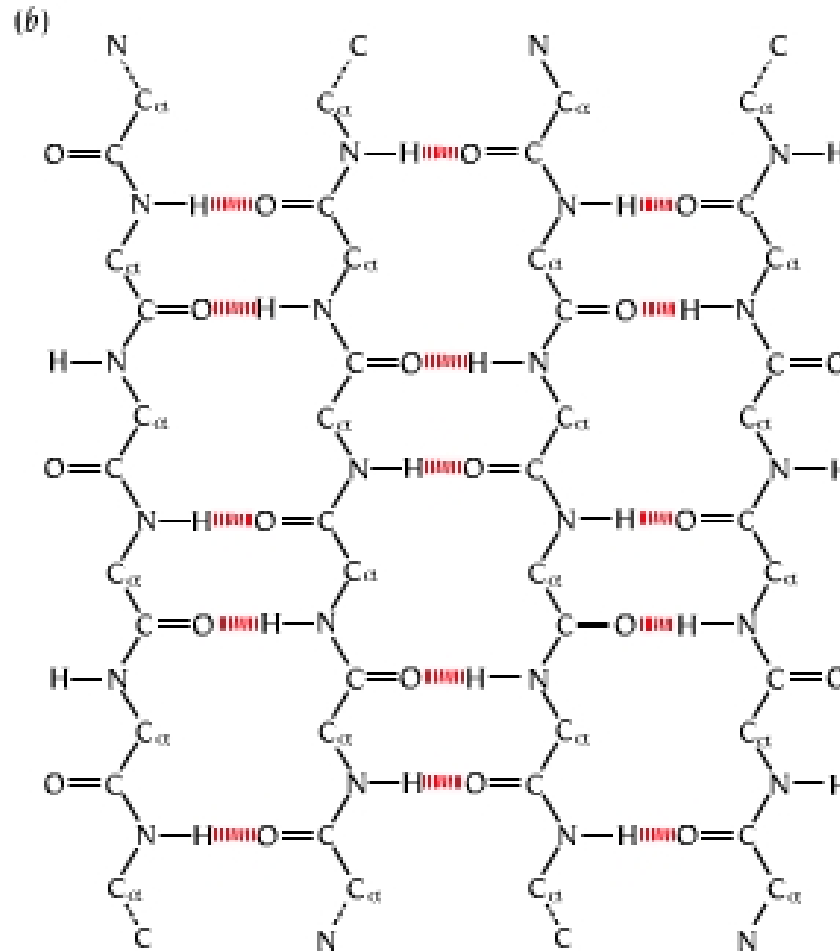
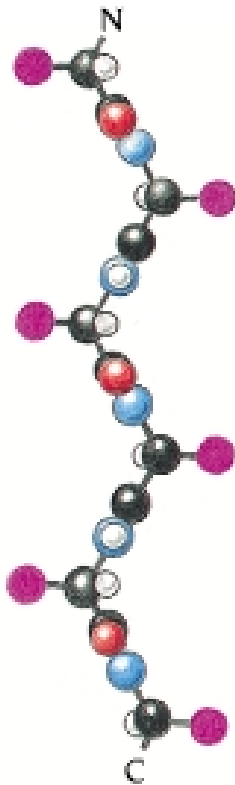


# Parallel beta sheet



H-bonds are evenly spaced.  
H-bonds are not 90° to the chain.

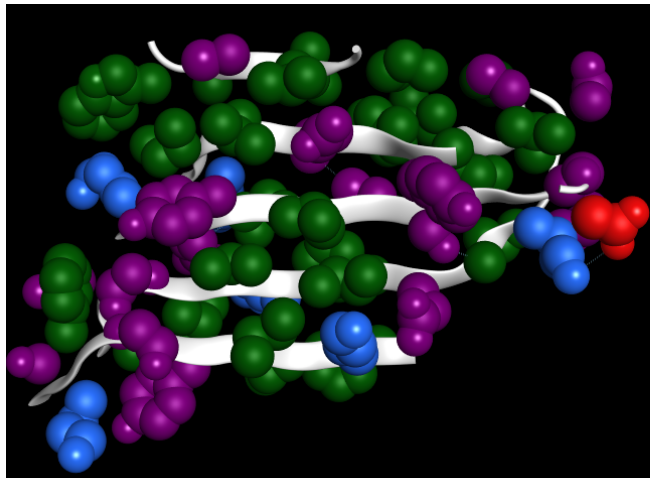
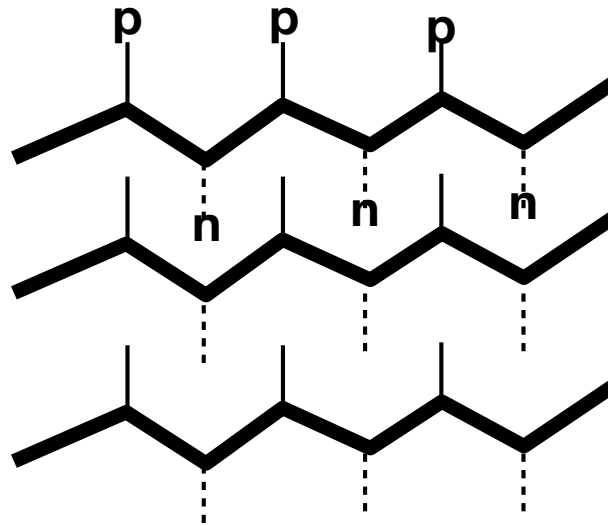
# Anti-parallel beta sheet



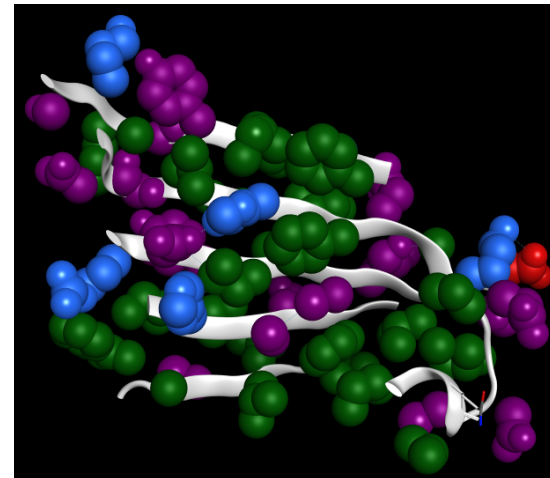
H-bonds are unevenly spaced.  
H-bonds are  $90^\circ$  to the chain.

# Sequence patterns for beta sheet

- npnp, or nnnn , where n=non-polar, p=pola



Non-polar residues  
(green, purple) mostly on  
the face.



Charged residues (blue,  
red) mostly on the ends.

# Mathematical expressions for H-bonding patterns

An H-bonding pattern can be expressed using augmented matrix.

next H-bond donor	=	multiply by donor	multiply by acceptor	add to donor	X	current H-bond donor
next H-bond acceptor		multiply by donor	multiply by acceptor	add to acceptor		current H-bond acceptor

For example, for an alpha helix....

150	=	1	0		1	X	149
146		0	1		1		145

or

150	=	1	0		1	X	149
146		1	0		-3		145

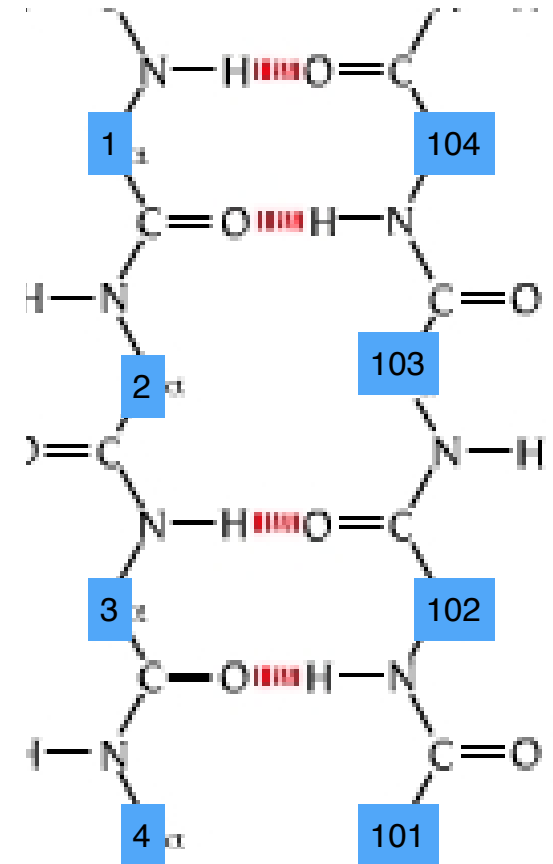
# Secondary structure using matrices: antiparallel sheet

handshake

0	1	0
1	0	0

skip

0	1	2
1	0	-2



Use the augmented matrices to find the next H-bond before/after  
(donor,acceptor)=(102, 3) in a antiparallel sheet

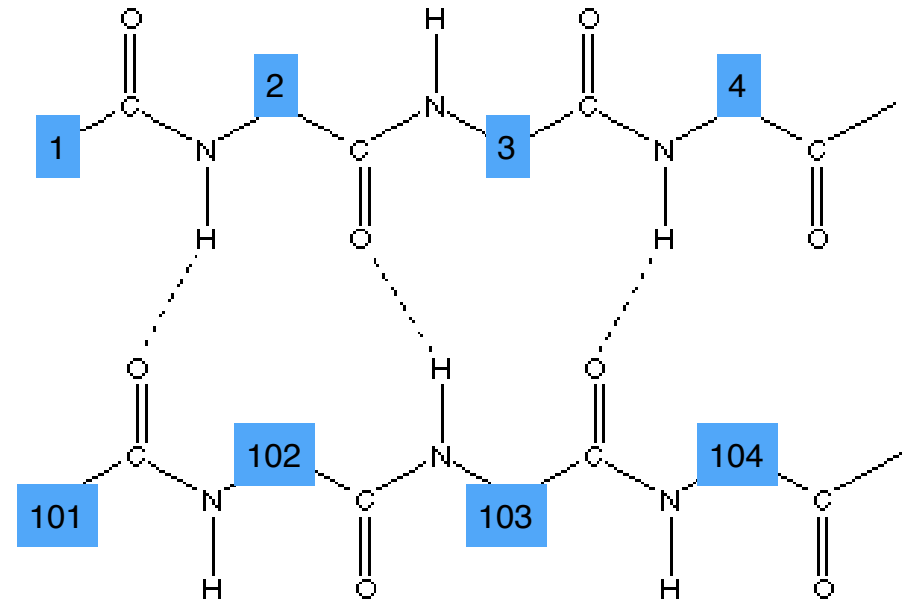
# Secondary structure using matrices: parallel sheet

C

0	1		2
1	0		0

N

0	1		0
1	0		-2

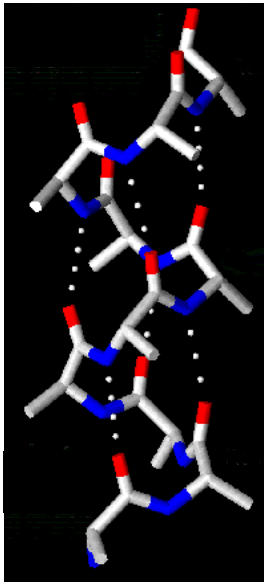


Use the augmented matrix to find the next H-bond before/after  
(donor,acceptor)=(103, 2) in a parallel sheet



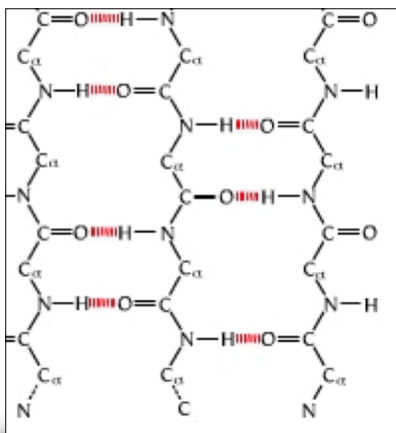
# Are beta sheets really secondary structure? Or tertiary structure?

Early folding is local. Late folding is non-local.



## Alpha helix

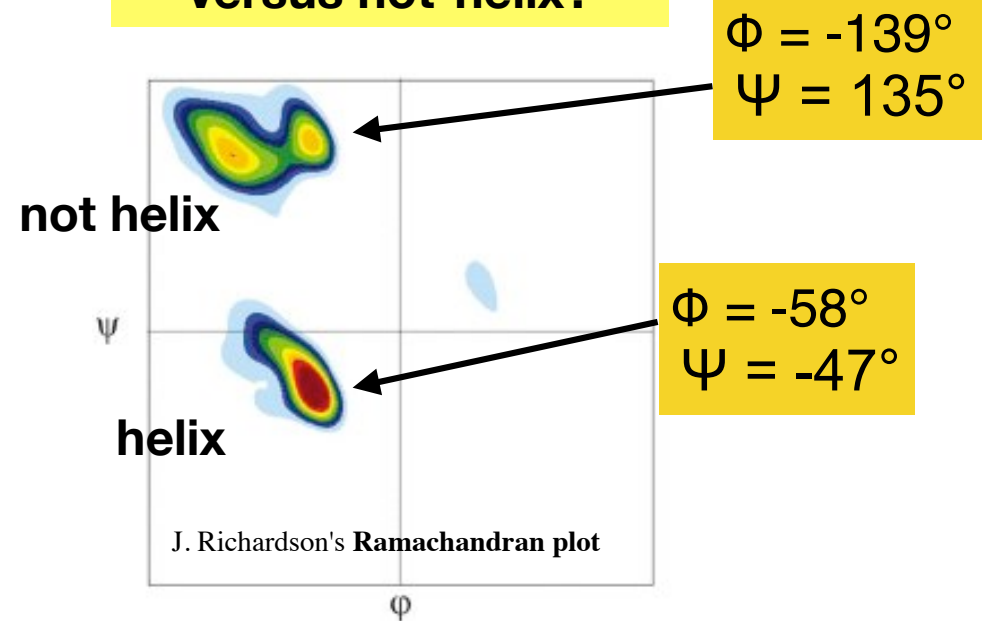
- H-bonds are local
- Side chain contacts are local
- folding is fast (ns)



## Beta sheet

- H-bonds are non-local
- Side chain contacts are non-local
- folding is slow ( $\mu$ s)

Is early folding just a decision of helix versus not-helix?



**Folding step 2?**

# Local structure

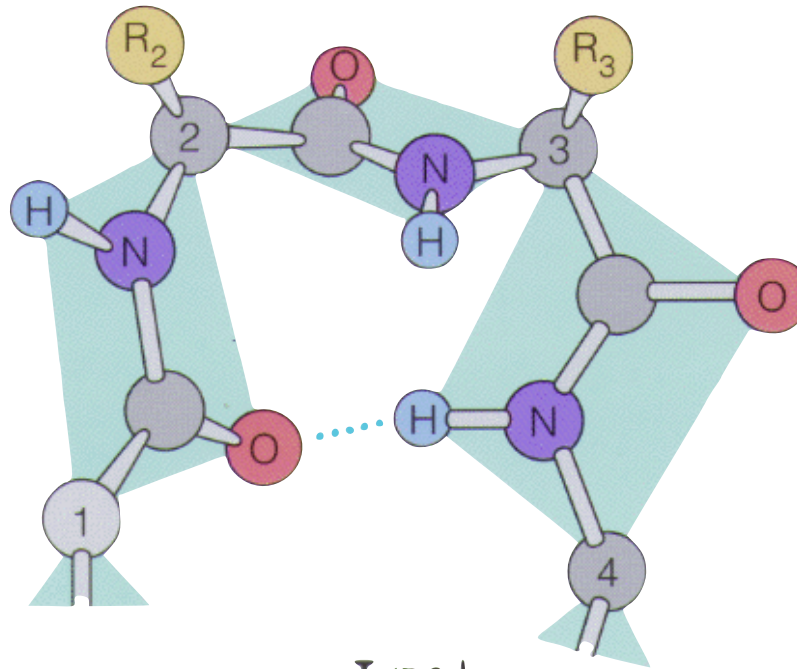
# Turns and caps

- Short pieces of protein chain sample conformational space randomly, driven by energy.
- Sequence determines structure. Non-polar sidechains and glycines are especially important.
- Usually reverse the chain direction.



# beta turns

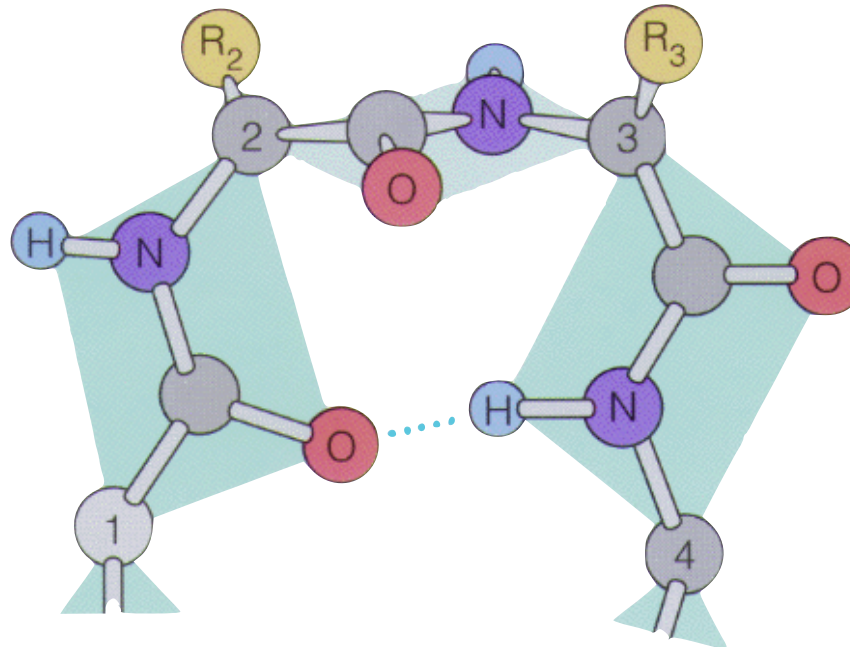
4-residues



Type I

**Type I** (most common). Oxygen points away when viewed clockwise.

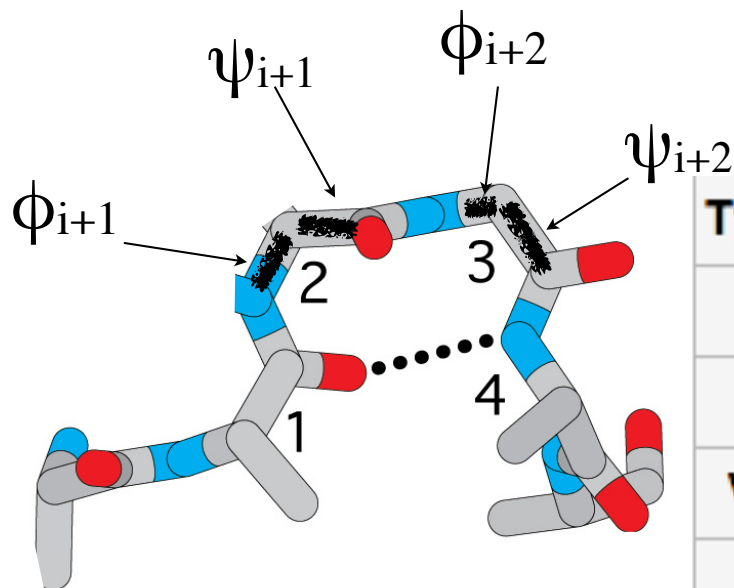
Residue 1 hydrogen bonds to residue 4



Type II

**Type II** (less common). Oxygen points toward when viewed clockwise.

# Backbone angles and sequence of beta turns



Backbone angles  $\pm 30^\circ$

Type	$\phi_{i+1}$	$\psi_{i+1}$	$\phi_{i+2}$	$\psi_{i+2}$
I	-60	-30	-90	0
II	-60	120	80	0
VIII	-60	-30	-120	120
I'	60	30	90	0
II'	60	-120	-80	0
Vla1	-60	120	-90	0*
Vla2	-120	120	-60	0*
Vlb	-135	135	-75	160*
IV	turns excluded from all the above categories			

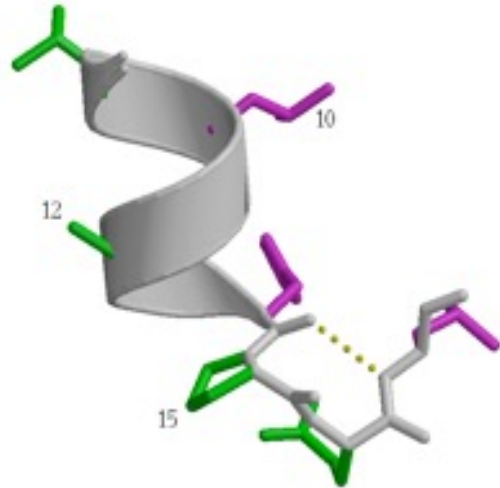
\*have cis-peptide bond at  $i+2$

Glycine rules turn propensity

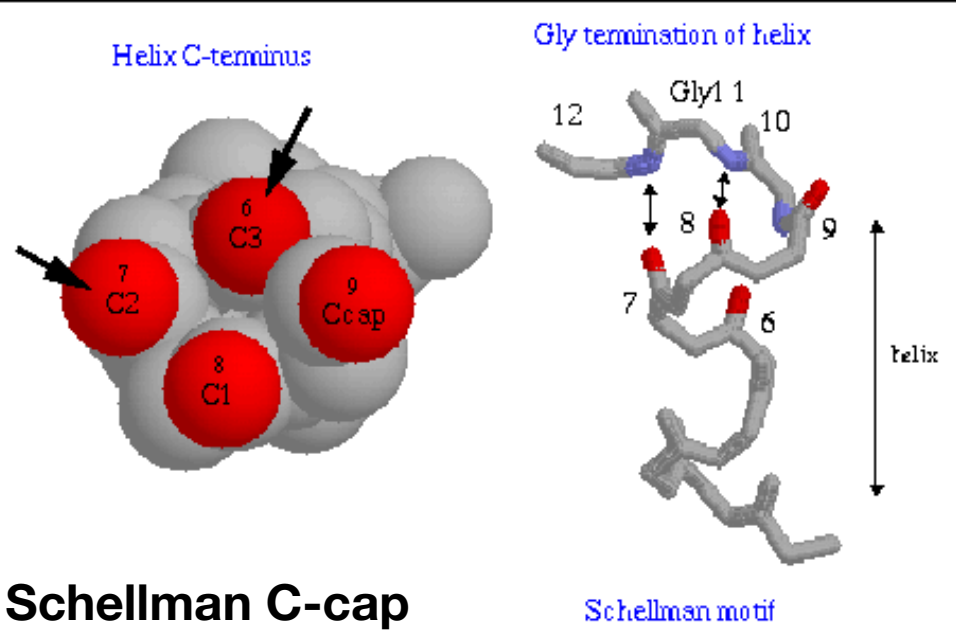
position type \	1	2	3	4
I		P	D/N/S/ T	<b>G</b>
II	P	P	<b>G</b>	
VIII	G/P	P		P
I'		<b>G</b>	<b>G</b>	
II'		<b>G</b>		

<http://www.ebi.ac.uk>

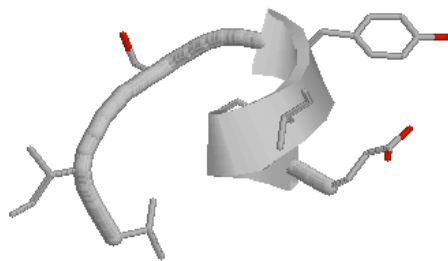
# Helix caps



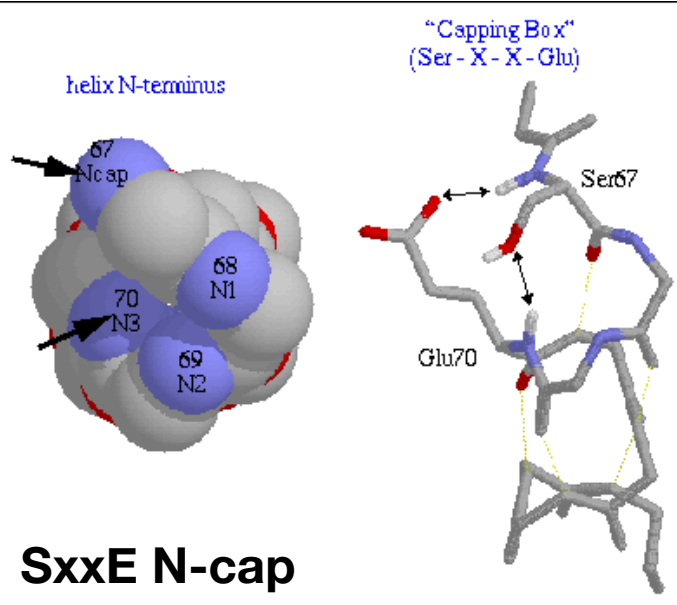
**Proline C-cap**



**Schellman C-cap**



**glycine N-cap**



**SxxE N-cap**

22

# Sequence patterns for turns, caps and secondary structure



Motif	Average boundaries		Average <i>rmsd</i> (len)	Pattern of conserved non- polar residues
	<i>mda</i> (°)	<i>dme</i> (Å)		
1 Amphipathic $\alpha$ -helix	56	0.71	0.78 (15)	1-4-8, 1-5-8
2 Non-polar $\alpha$ -helix	54	0.58	0.40 (11)	1-4-8, 1-5-8
3 Schellman cap type 1	81	1.01	1.02 (15)	1-6-9-11
4 Schellman cap type 2	76	0.94	0.94 (15)	1-6-8-9
5 Proline $\alpha$ -helix C cap	92	1.07	0.89 (13)	1-2-5-8
6 Frayed $\alpha$ -helix	75	0.96	0.69 (15)	1-5-9-13
7 Helix N capping box	99	0.95	0.65 (15)	1-6-9-13
8 Amphipathic $\beta$ -strand	89	0.87	0.87 (6)	1-3, 1-3-5
9 Hydrophobic $\beta$ -strand	101	0.91	0.91 (7)	1-2-3
10 $\beta$ -Bulge	100	0.97	0.78 (7)	1-4-6
11 Serine $\beta$ -hairpin	94	0.76	0.81 (9)	1-8
12 Type-I hairpin	80	0.94	1.23 (13)	1-7-8
13 Diverging type-II turn	87	1.04	1.00 (9)	1-7-9

Bystroff C & Baker D. (1998). Prediction of local structure in proteins using a library of sequence-structure motifs. *J Mol Biol* 281, 565-77.

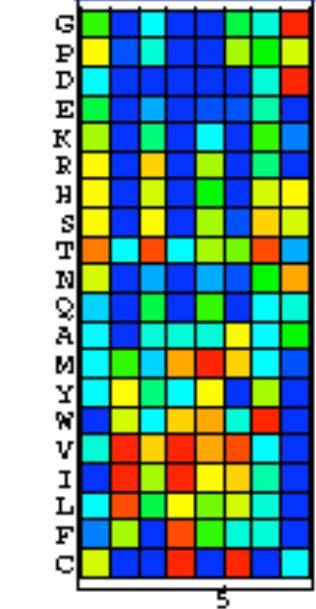
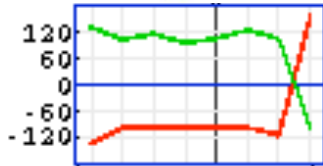
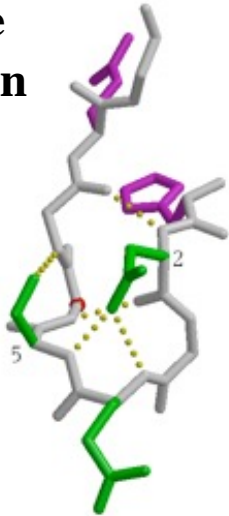
**Folding step 3?**

# Super-secondary structure

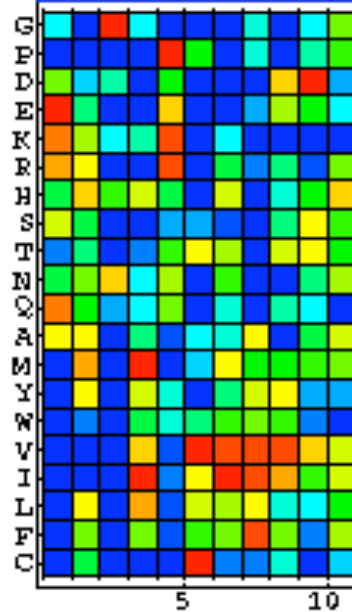
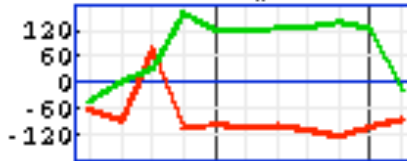
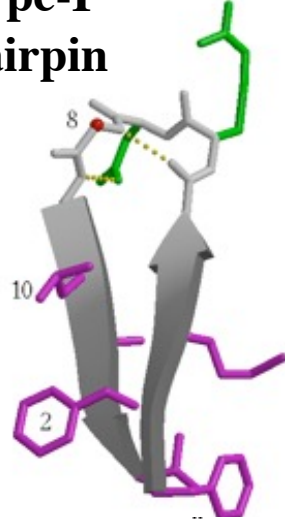


# $\beta$ Hairpins

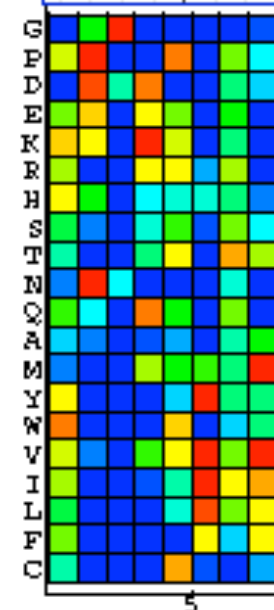
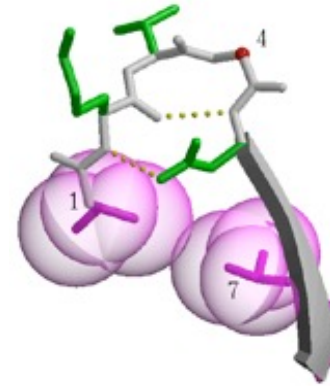
**Serine hairpin**



**Type-I hairpin**



**diverging type-2 turn**

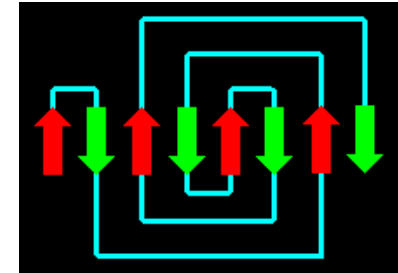
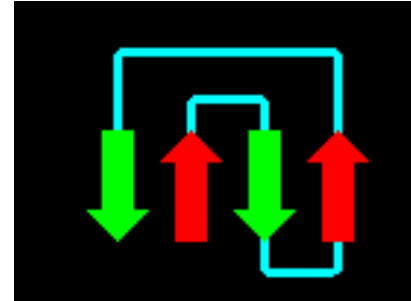


Sequence patterns are expressed as log-likelihood, converted to color scale.

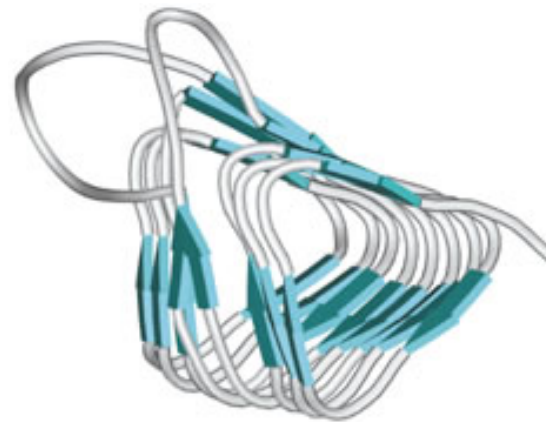
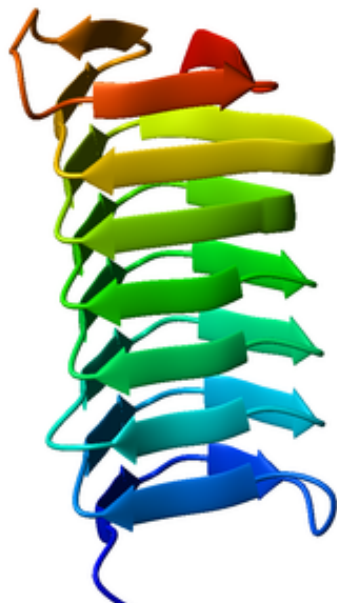
# $\beta$ sheet super-secondary structure.



meander



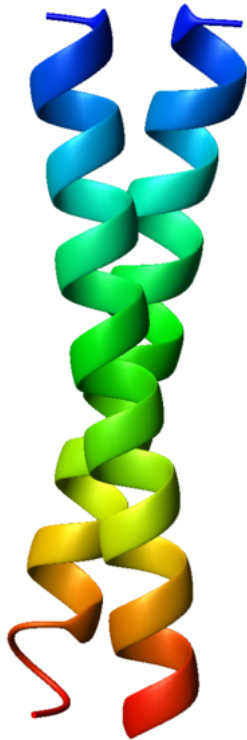
"greek key"



$\beta$  helix (2-sided, 3-sided, left, right)

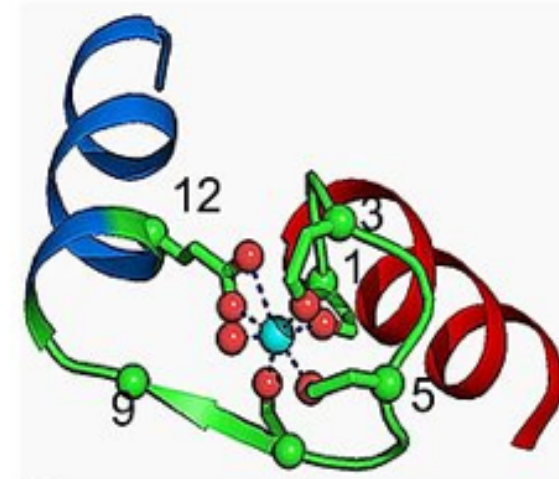
# $\alpha$ Helical Super-Secondary Structures

- SSS contains more than one SSE.
- beta turns and helix caps are usually involved.
- Canonical ones have names.

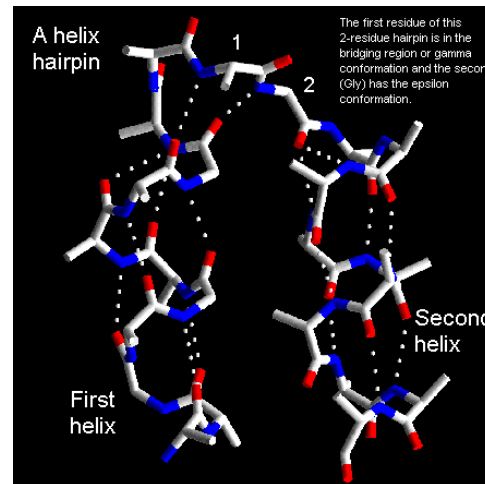


Coiled-coil

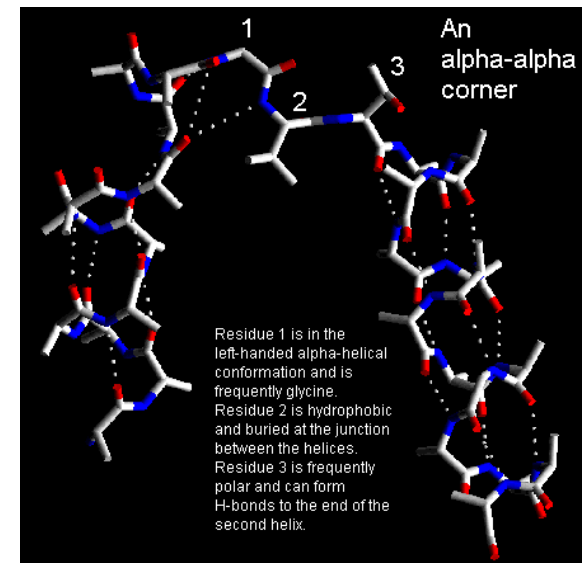
what is the handedness?



EF hand

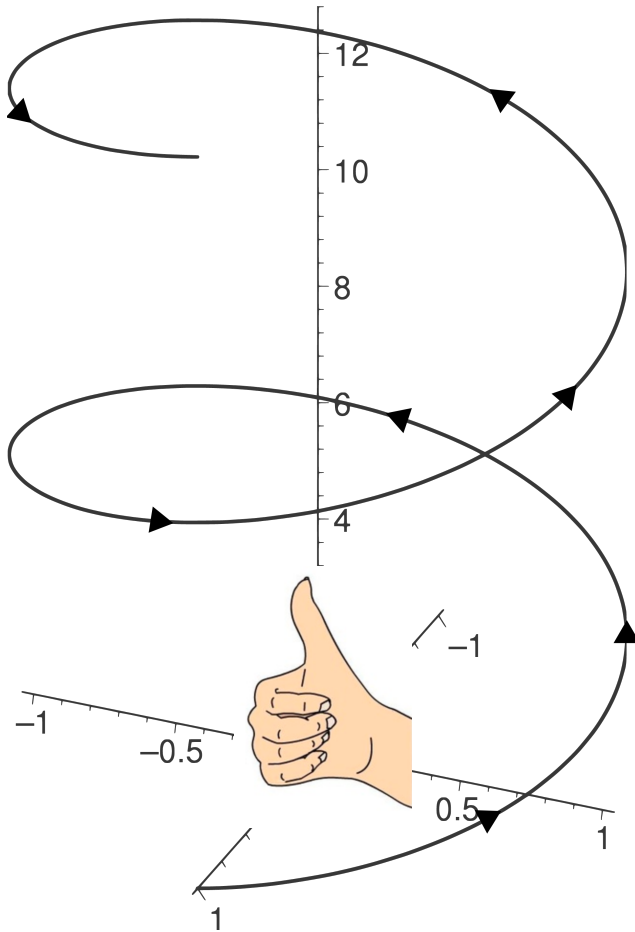


Helix hairpin



alpha-alpha corner

# Handedness



## Right-handed helix.

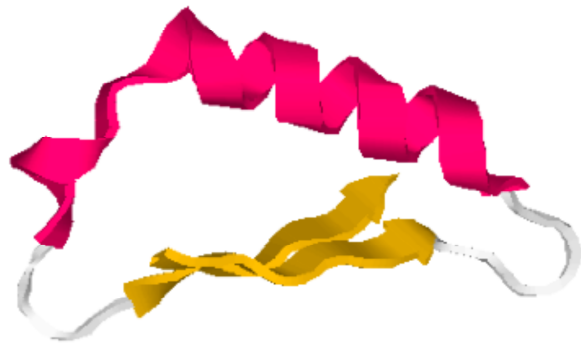
**Put the thumb of the right hand along the axis of rotation.**

**As you travel up the helix (going in the direction of your right thumb) the line curve in the direction of your fingers.**

Yes, that means you are turning left when you walk up a right-handed spiral staircase, and right when you are walking up a left-handed spiral staircase.

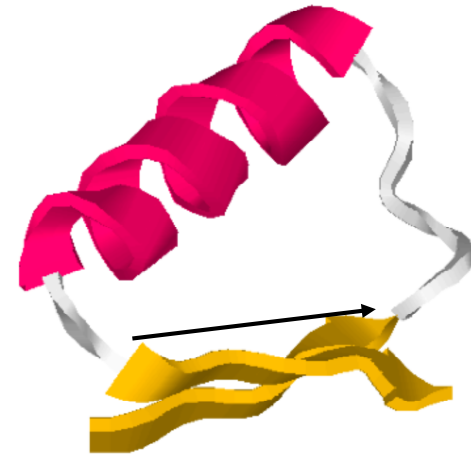
# $\alpha\beta$ Super-secondary structure.

$\beta\alpha\beta$  supersecondary structure units are mostly right-handed



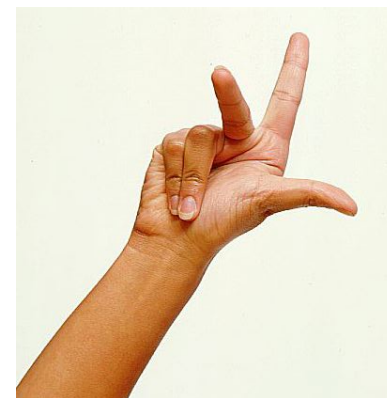
L-handed  $\beta\alpha\beta$

1.5%

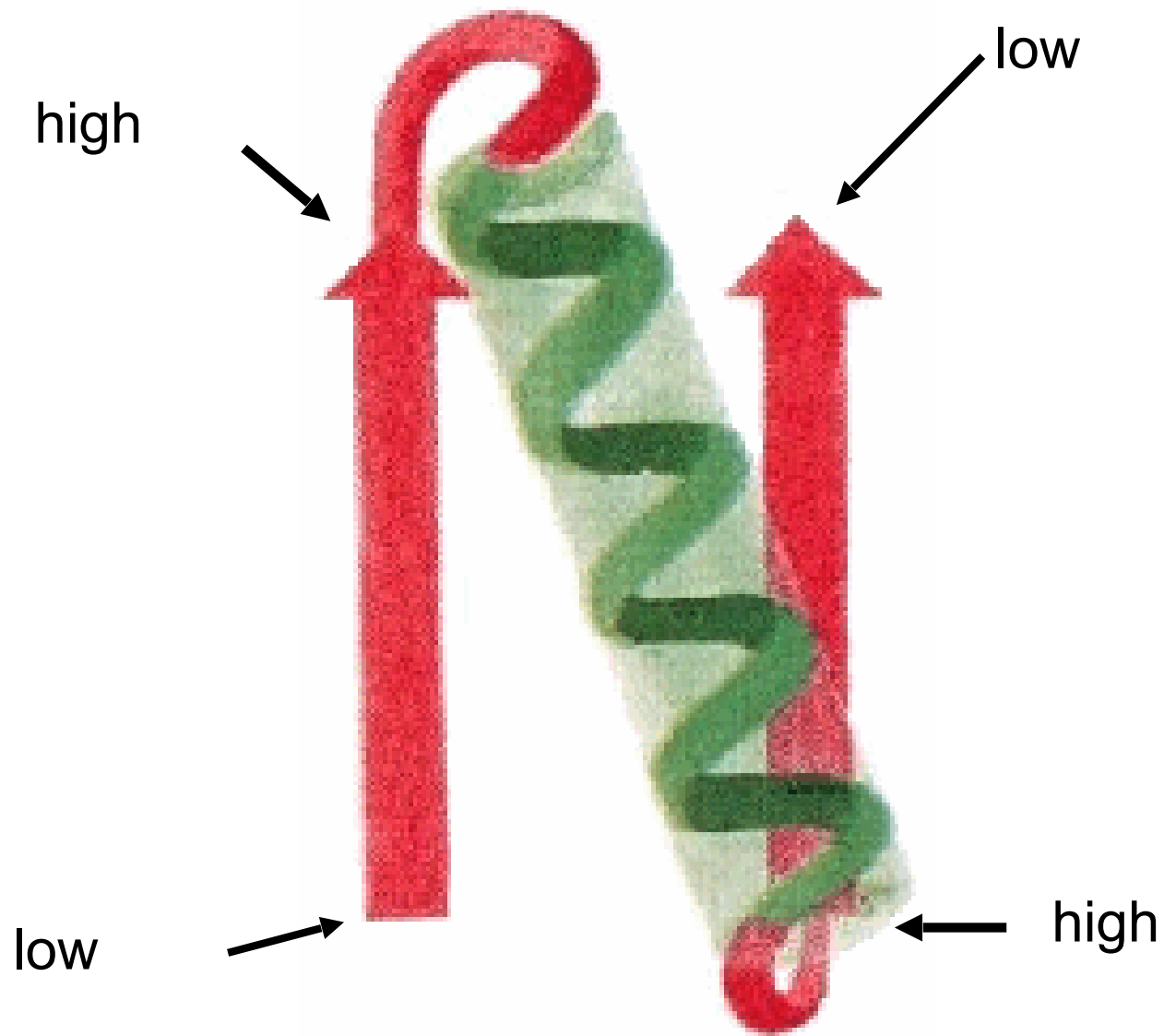


R-handed  $\beta\alpha\beta$

98.5%



Theories for why  $\beta\alpha\beta$  units are right-handed.



Sternberg & Thornton: Twist of beta sheet makes right-handed crossover more of a straight line.

# Theories for why $\beta\alpha\beta$ units are right-handed.

2622 Biochemistry: Richardson

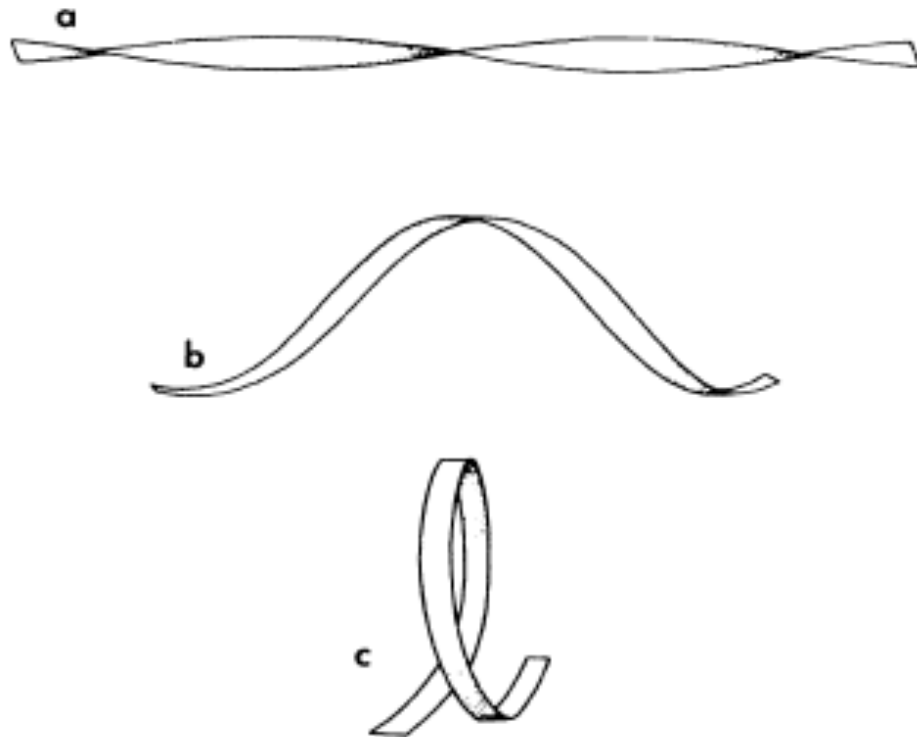


FIG. 4. A possible folding pathway which produces righthanded crossover loops from extended chain. In (a) the section of chain is extended, showing one full turn of the preferred righthanded twist for  $\beta$  strands. In (b) the two ends of this chain segment are moving toward one another, and the ribbon has started to buckle in a righthanded sense constrained by the chain twist. In (c) a complete righthanded loop is formed, with the two ends in position to form parallel  $\beta$  structure.

*Proc. Natl. Acad. Sci. USA 73 (1976)*

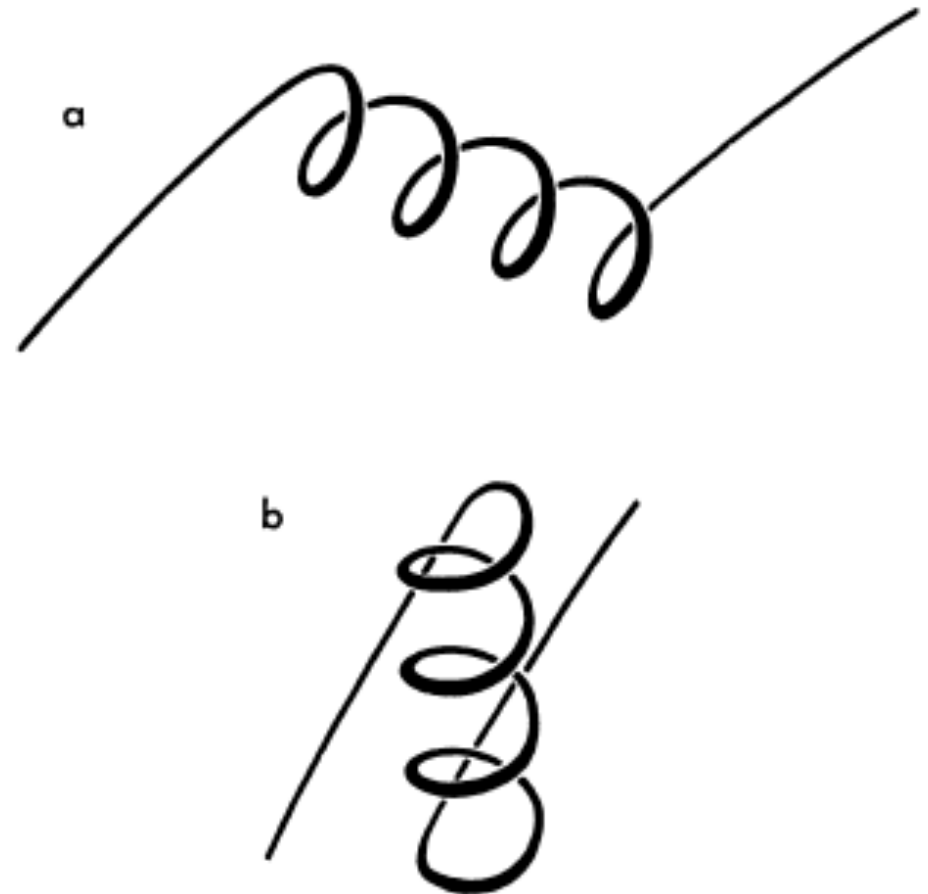


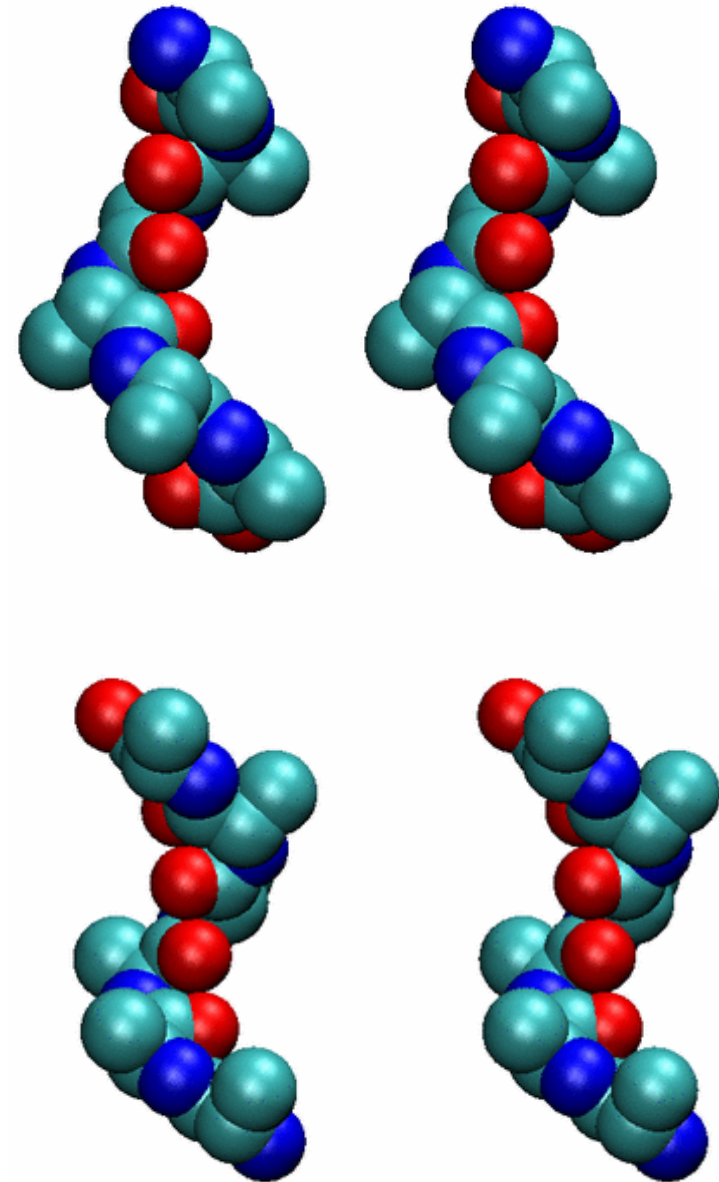
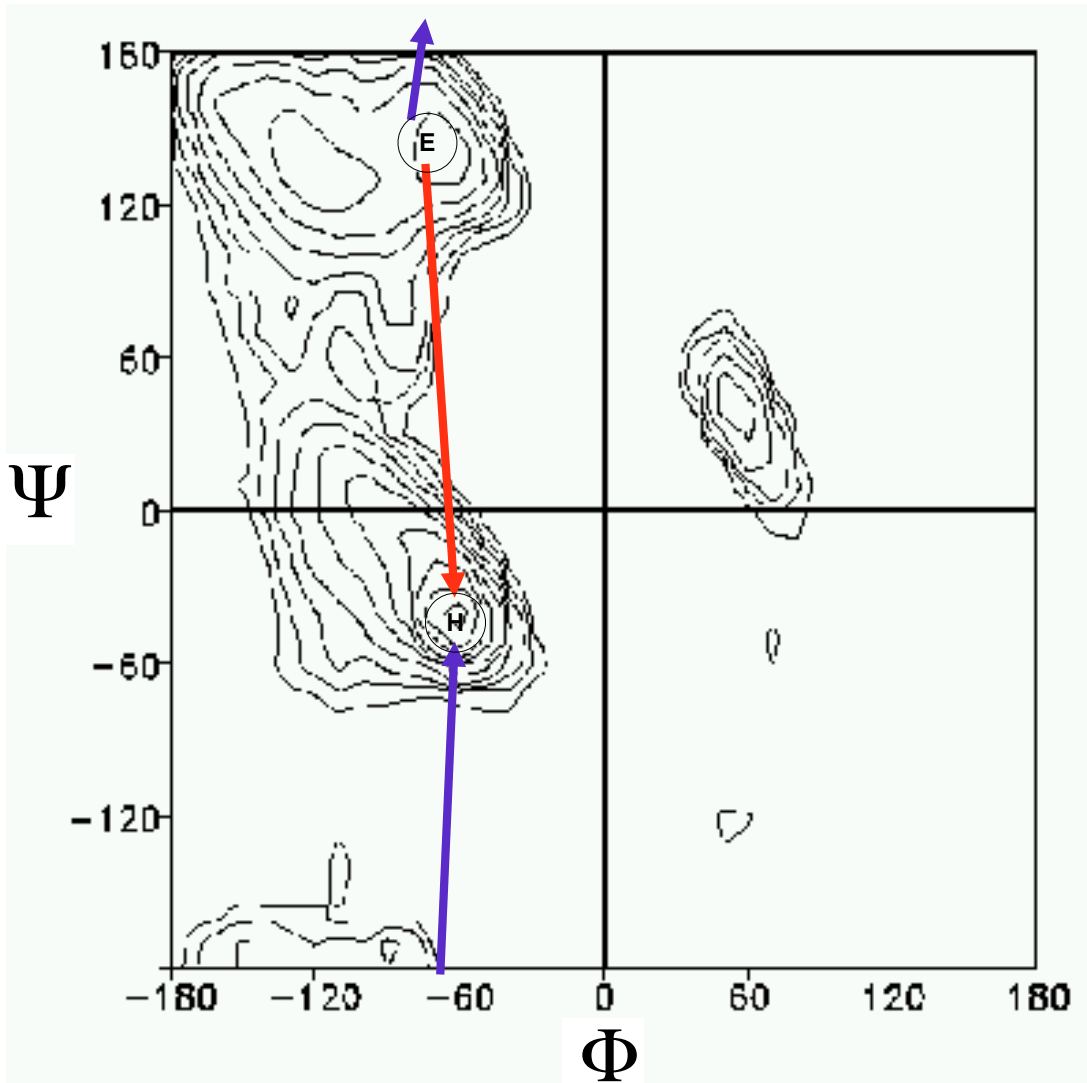
FIG. 5. A possible folding pathway which forms righthanded crossover loops from a righthanded  $\alpha$ -helix with a  $\beta$  strand at each end of it.

Richardson, PNAS, 1976: Right-handed crossovers are trapped early in folding



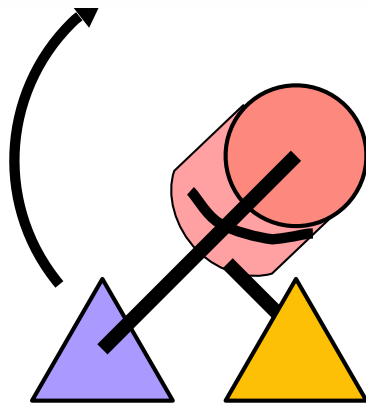
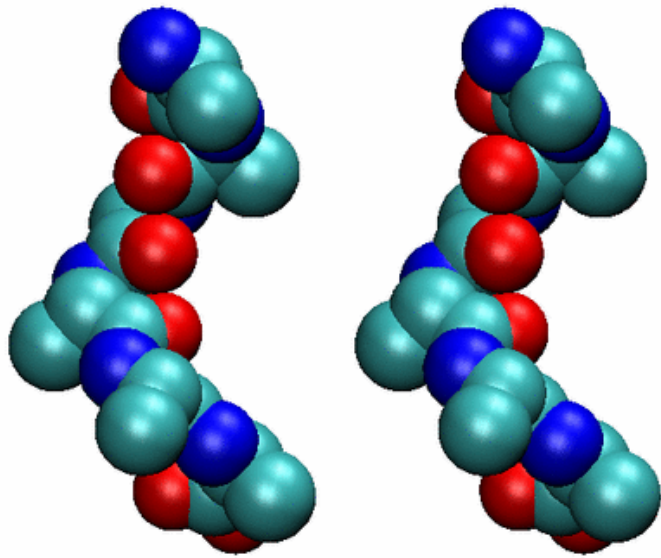
# Theories for why $\beta\alpha\beta$ units are right-handed.

Phone Cord Effect: Northern versus Southern route to helix

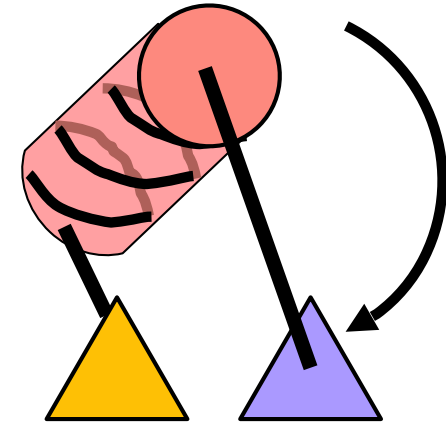
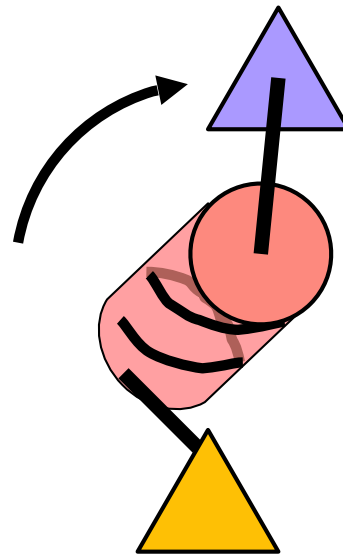




Theories for why  $\beta\alpha\beta$  units are right-handed.



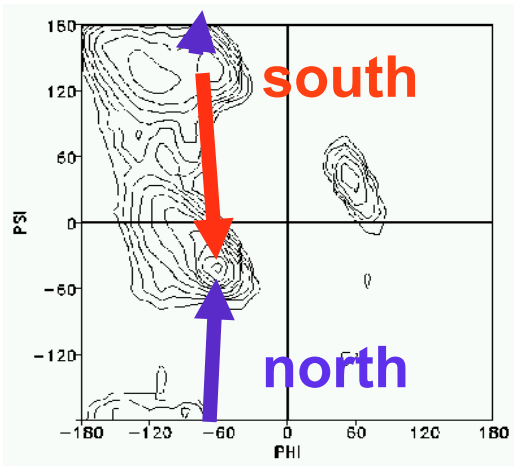
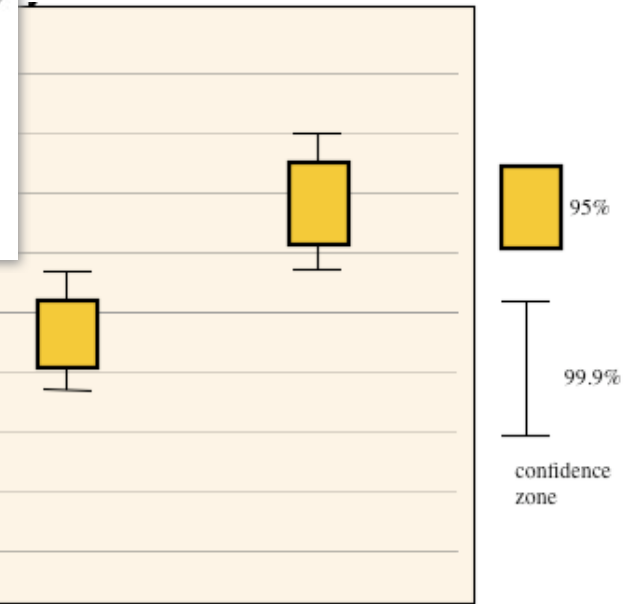
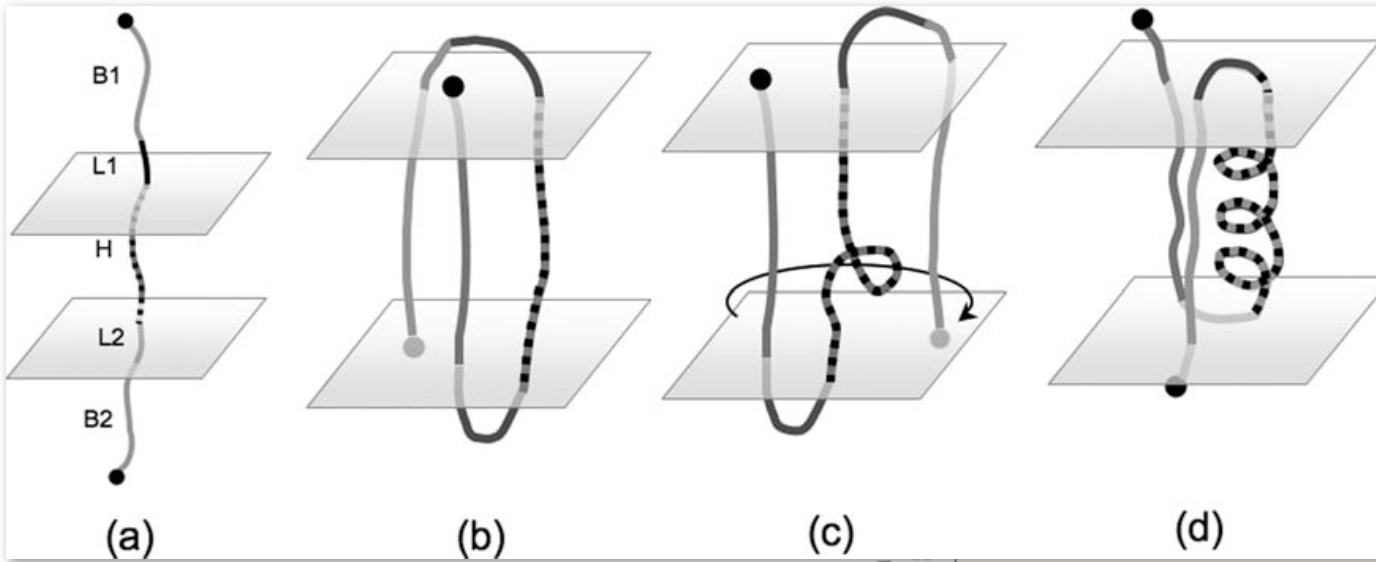
LH



RH

left-handed torque turns left-handed  $\beta\alpha\beta$  to right-handed  $\beta\alpha\beta$

# Phone cord: Brownian Dynamics Simulations



$\Delta\Psi$	0% South	50% South	100% South
Trials	2738	1164	501
Collapsed	2540	1066	418
Helical	851	578	286
Ambiguous	456	299	131
Right-handed	124	130	107
Left-handed	271	149	48

# 3-helix bundles are also right-handed

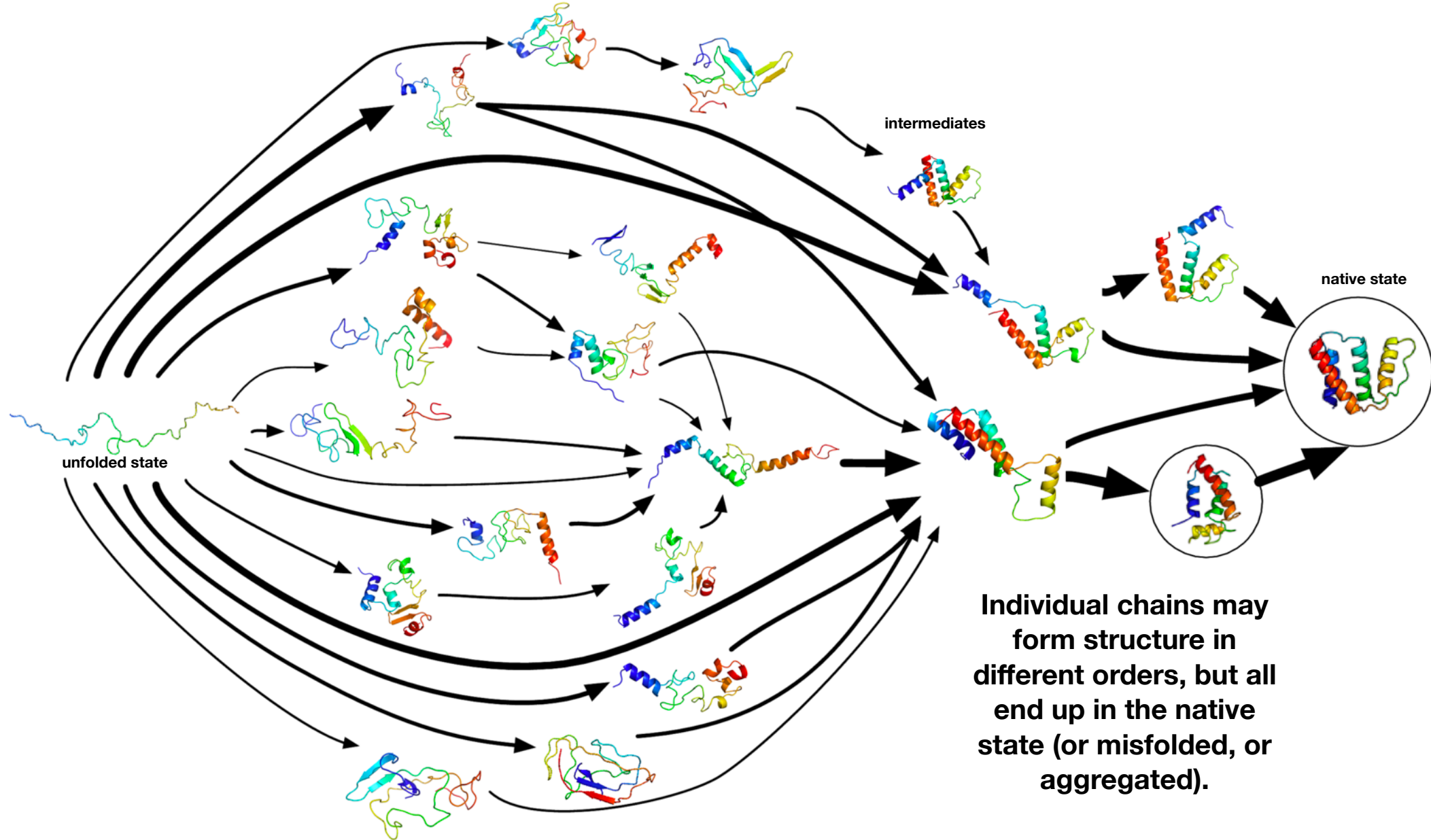
Helix residue ranges										Helix residue ranges										Helix residue ranges										
PDB	R/L	frac R	Contacts	Helix 1			Helix 2			Helix 3			PDB	R/L	frac R	Contacts	Helix 1			Helix 2			PDB	R/L	frac R	Contacts	Helix 1		Helix 2	
code/chain													code/chain										code/chain							
1a26A	R	1.00	14	703	721	726	739	755	778	1j0tA	L	0.06	16	16	31	37	40	49	56	1tm9A	R	0.83	6	57	74	92	106			
1a9xA	R	1.00	6	420	429	433	445	449	456	1jj2O	L	0.00	9	4	14	28	33	37	45	1tx4A	L	0.10	21	192	203	209	214			
1a9xA	R	1.00	7	433	445	449	456	460	479	1jj2O	L	0.00	5	90	111	116	127	134	141	1tx4A	R	0.86	14	64	75	90	102			
1a9xA	R	1.00	8	460	479	486	494	499	506	1jr3A	R	0.78	9	278	297	304	308	310	319	1tx4A	R	1.00	5	165	183	185	188			
1a9xA	R	1.00	1	486	494	499	506	510	519	1jr3A	R	1.00	11	246	258	261	273	278	297	1tx9A	L	0.00	11	88	97	104	109			
1aa7A	R	0.89	28	109	117	121	132	140	157	1jswA	L	0.00	6	47	65	70	83	104	121	1tx9A	R	1.00	2	75	85	88	97			
1aa7A	R	1.00	3	19	33	39	47	54	67	1jswA	L	0.00	18	201	226	246	257	275	302	1tz4A	L	0.00	3	17	21	26	37			
1aa7A	R	1.00	1	39	47	54	67	78	83	1jswA	L	0.00	6	275	302	331	355	365	388	1u84A	R	1.00	2	28	38	44	58			
1aa7A	R	1.00	18	90	105	109	117	121	132	1jswA	R	1.00	1	147	182	201	226	246	257	1ubyA	L	0.00	5	53	67	73	85			
1abvA	L	0.00	3	23	39	41	47	53	64	1k6kA	R	1.00	2	4	20	27	35	38	46	1ubyA	L	0.00	13	167	191	204	214			
1adtA	L	0.00	4	180	194	200	203	212	224	1k6kA	R	1.00	8	27	35	38	46	51	64	1ubyA	L	0.00	6	204	214	216	231			
1aepA	L	0.00	9	34	65	69	86	94	121	1k8kE	R	1.00	8	63	83	88	100	123	148	1ubyA	L	0.00	4	283	291	294	303			
1aepA	R	1.00	2	69	86	94	121	126	129	1kjsA	R	0.76	38	16	26	34	38	45	62	1un8A	R	1.00	24	356	371	373	382			
1aepA	R	1.00	1	94	121	126	129	131	154	1kp8A	L	0.00	4	53	59	65	84	89	109	1un8A	R	1.00	5	388	404	413	427			
1af7A	R	0.80	5	47	61	66	75	80	88	1kp8A	R	0.82	28	10	29	53	59	65	84	1un8A	R	1.00	16	477	490	495	511			
1agrE	R	1.00	2	53	61	63	68	70	82	1l8wA	L	0.00	11	228	240	255	260	277	289	1us7B	L	0.00	8	203	226	234	242			
1ah7A	L	0.00	3	13	27	34	42	44	54	1lbuA	R	1.00	1	17	25	44	56	67	76	1us7B	R	1.00	4	156	164	168	177			
1ah7A	L	0.00	6	187	190	193	204	206	241	1lkpA	R	1.00	1	56	63	83	112	115	144	1us7B	R	1.00	6	294	300	317	321			
1ah7A	L	0.10	10	106	124	141	151	172	185	1llaA	L	0.00	6	266	282	300	309	317	320	1utgA	R	1.00	5	4	14	18	27			
1ailA	R	0.86	29	3	24	30	50	54	69	1llaA	R	1.00	5	300	309	317	320	323	332	1uuja	R	1.00	2	5	21	25	35			
1aorA	L	0.00	1	237	240	243	253	274	280	1llpA	L	0.00	1	166	177	203	209	236	242	1v2zA	R	1.00	7	186	203	211	225			
1aorA	L	0.00	1	274	280	213	217	227	240	1llpA	R	1.00	3	70	73	75	80	87	101	1v2zA	R	1.00	9	211	225	229	246			
1aorA	L	0.00	1	274	280	213	217	227	240	1llpA	R	0.00	9	6	18	24	26	41	54	1v54H	R	1.00	1	26	45	50	63			
1aorA	L	0.00	1	274	280	213	217	227	240	1llpA	R	0.86	9	6	18	24	26	41	54	1v54H	R	1.00	9	9	18	22	38			
1aorA	L	0.14	1	274	280	213	217	227	240	1llpA	R	1.00	1	6	18	24	26	41	54	1v54H	R	1.00	20	22	38	50	63			
1aorA	L	0.20	1	274	280	213	217	227	240	1llpA	R	1.00	11	68	77	87	93	101	111	1v54H	R	1.00	8	135	150	160	178			
1aorA	R	1.00	1	274	280	213	217	227	240	1llpA	R	0.00	8	135	150	160	178	188	198	1v54H	R	1.00	3	67	78	84	97			
1aorA	R	1.00	1	274	280	213	217	227	240	1llpA	R	0.00	17	43	57	60	72	77	91	1v54H	R	1.00	17	43	57	60	72			
1b79A	R	1.00	1	274	280	213	217	227	240	1llpA	R	0.11	10	60	72	77	91	101	111	1v54H	R	1.00	10	60	72	77	91			
1b79A	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	2	102	116	123	132	142	152	1v54H	R	1.00	2	102	116	123	132			
1b79A	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	6	17	23	25	45	55	65	1v54H	R	1.00	6	17	23	25	45			
1bf5A	R	1.00	1	274	280	213	217	227	240	1llpA	R	0.00	14	381	402	412	428	438	450	1v54H	R	1.00	14	381	402	412	428			
1bf5A	R	1.00	1	274	280	213	217	227	240	1llpA	R	0.00	9	412	428	438	450	460	470	1v54H	R	1.00	9	412	428	438	450			
1bvfA	L	0.00	1	274	280	213	217	227	240	1llpA	R	0.00	5	438	450	458	475	485	495	1v54H	R	1.00	5	438	450	458	475			
1bvfA	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	2	458	475	477	485	495	505	1v54H	R	1.00	2	458	475	477	485			
1bmtA	L	0.00	1	274	280	213	217	227	240	1llpA	R	1.00	5	263	297	304	321	331	341	1v54H	R	1.00	5	263	297	304	321			
1bouA	L	0.00	1	274	280	213	217	227	240	1llpA	R	1.00	8	364	377	381	390	399	409	1v54H	R	1.00	8	364	377	381	390			
1bouA	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	2	289	301	305	322	332	342	1v54H	R	1.00	2	289	301	305	322			
1bvp1	R	0.90	1	274	280	213	217	227	240	1llpA	R	1.00	4	11	44	49	71	81	91	1v54H	R	1.00	4	11	44	49	71			
1c1kA	L	0.00	1	274	280	213	217	227	240	1llpA	R	0.00	1	92	128	138	161	171	181	1v54H	R	1.00	1	92	128	138	161			
1c1kA	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	5	6	12	16	20	24	28	1v54H	R	1.00	5	6	12	16	20			
1c75A	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	5	23	41	55	70	85	100	1v54H	R	1.00	5	23	41	55	70			
1cktA	L	0.18	1	274	280	213	217	227	240	1llpA	R	0.79	5	16	20	23	41	55	70	1v54H	R	1.00	5	16	20	23	41			
1crkA	L	0.17	1	274	280	213	217	227	240	1llpA	R	1.00	1	105	120	132	137	147	157	1v54H	R	1.00	1	105	120	132	137			
1cshA	L	0.00	1	274	280	213	217	227	240	1llpA	R	0.95	3	28	36	46	60	74	88	1v54H	R	1.00	3	28	36	46	60			
1cshA	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	2	242	252	260	281	291	301	1v54H	R	1.00	2	242	252	260	281			
1cshA	R	1.00	1	274	280	213	217	227	240	1llpA	R	0.00	14	316	326	332	336	346	356	1v54H	R	1.00	14	316	326	332	336			
1cukA	L	0.08	1	274	280	213	217	227	240	1llpA	R	1.00	4	66	76	80	83	86	92	1y5A	R	0.80	10	260	281	291	309			
1d2tA	R	0.96	1	274	280	213	217	227	240	1llpA	R	1.00	3	120	129	131	145	151	169	1yfsA	R	1.00	3	338	370	380	389			
1d2tA	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	4	2	19	22	39	43	63	1yfsA	R	1.00	14	380	389	394	403			
1dbhA	R	1.00	1	274	280	213	217	227	240	1llpA	R	0.00	15	33	55	62	67	70	83	1yfsA	R	1.00	3	394	403	410	423			
1dbhA	R	1.00	1	274	280	213	217	227	240	1llpA	R	0.00	13	116	142	146	153	156	169	1ygeA	L	0.00	3	636	640	643	656			
1dbhA	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	3	62	67	70	83	91	108	1ygeA	L	0.12	8	255	276	286	292			
1dbhA	R	1.00	1	274	280	213	217	227	240	1llpA	R	1.00	5	146	153	156	169	176	193	1ygeA	R	1.00	1	410	415	417	422			
1dj8A	L	0.14	7	29	39	52	68	74	82	1n93X	L	0.00	3	130	149	152	156	159	172	1ygeA	R	1.00	6	474	516	523	530			
1dj8A	R	1.00	7	18	2																									

Folding step 4?

# Tertiary structure

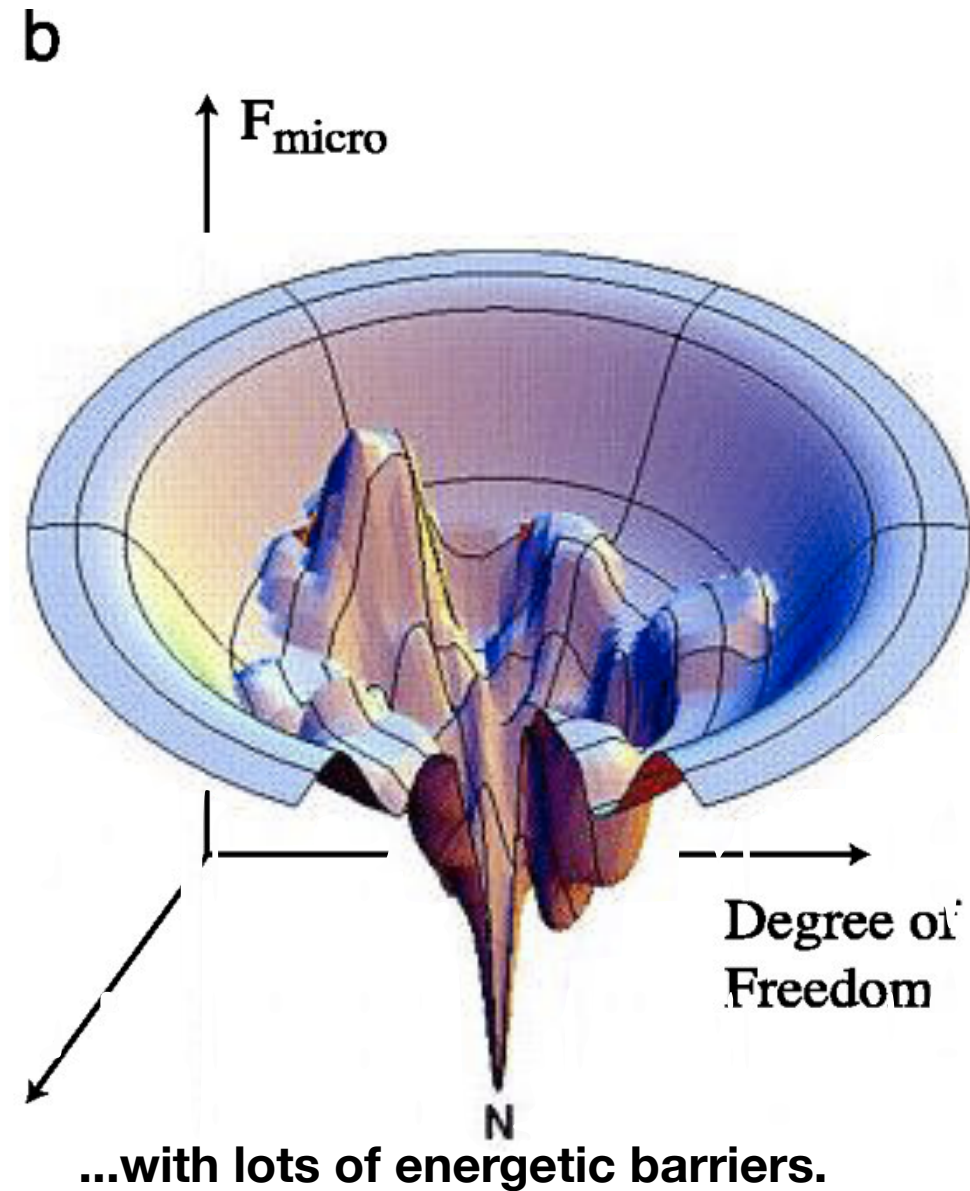
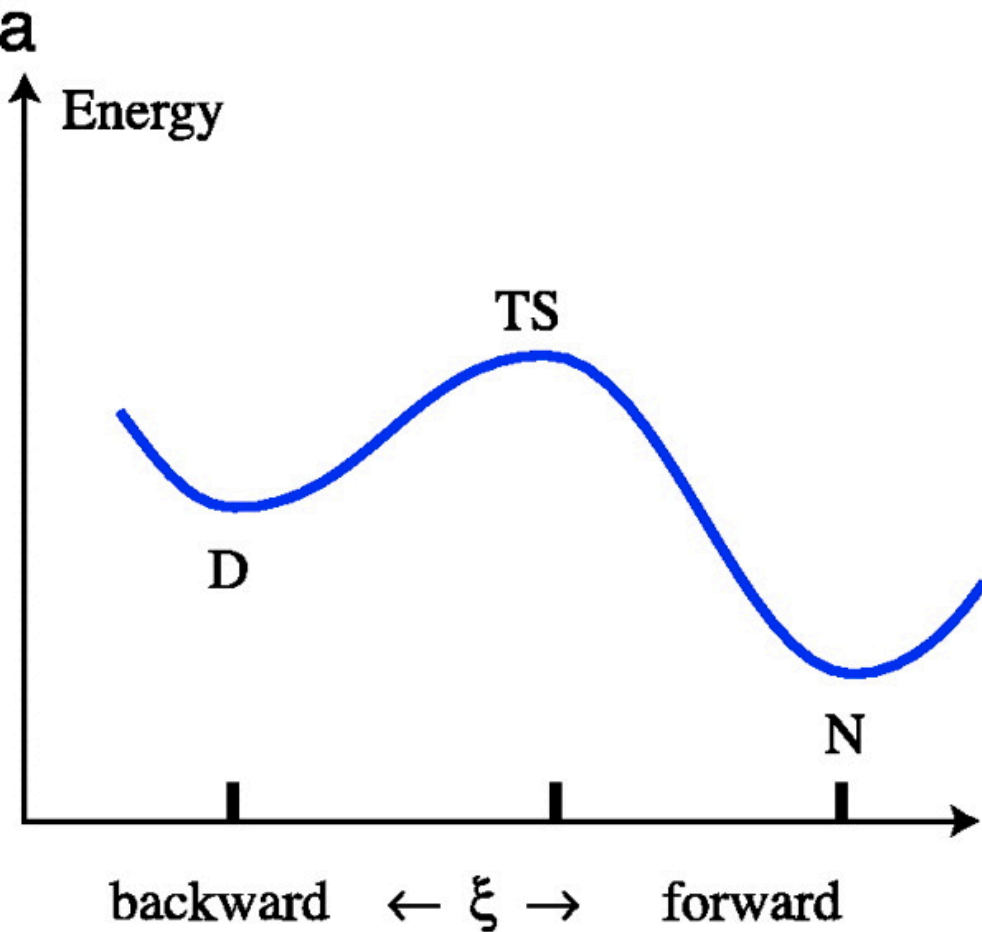
7

# Folding pathway is an ensemble



**Individual chains may form structure in different orders, but all end up in the native state (or misfolded, or aggregated).**

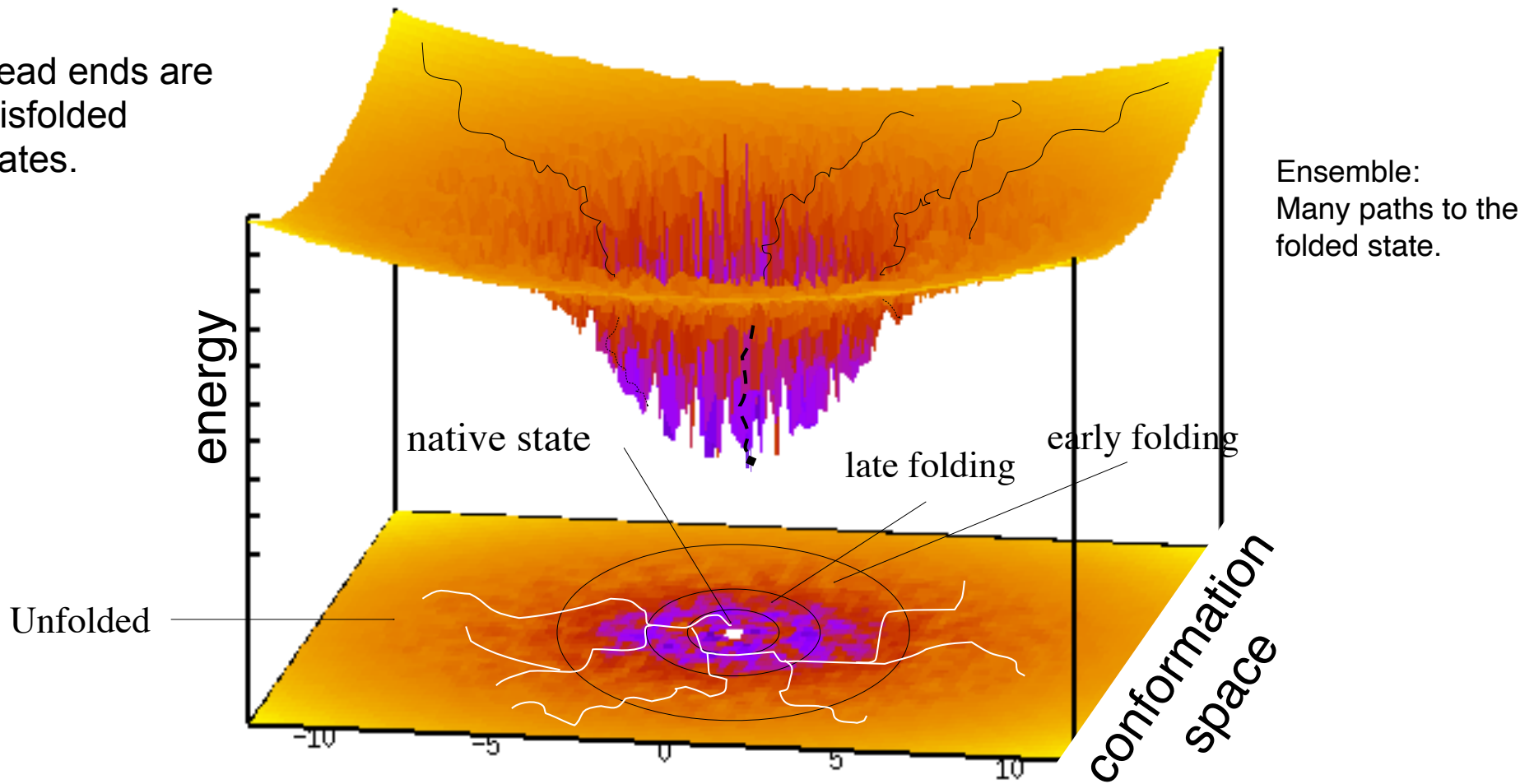
# Folding is a funnel





# Protein folding is an ensemble on a noisy funnel

Dead ends are misfolded states.



Dill, K. A. (1999). Polymer principles and protein folding. *Protein Science*, 8(6), 1166-1180.

Garcia-Manyes, S., Dougan, L., Badilla, C. L., Brujić, J., & Fernández, J. M. (2009). Direct observation of an ensemble of stable collapsed states in the mechanical folding of ubiquitin. *Proceedings of the National Academy of Sciences*

# How to force hydrogen bonds using restraints

- To add a restraint

**Edit | Potential | Restrain**, distance,

Target 1.8, 1.8, Weight 50

Pick amide H and carbonyl O.

Click **Create**.

Cancel | Restrain (or esc) when done

- Energy minimize

**Compute | prepare | Structure preparation**

Checks for missing atoms, assigns energies.

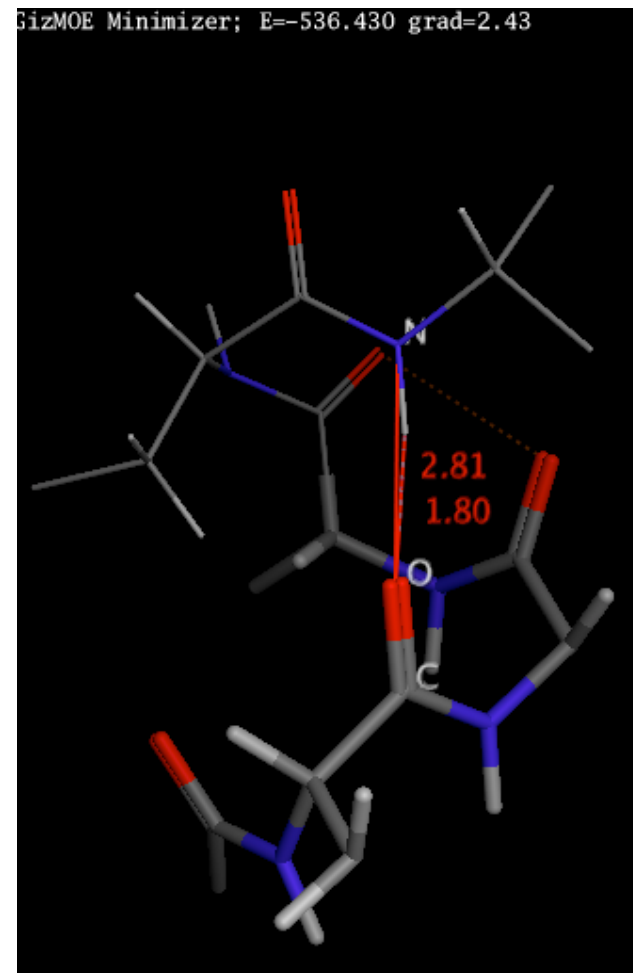
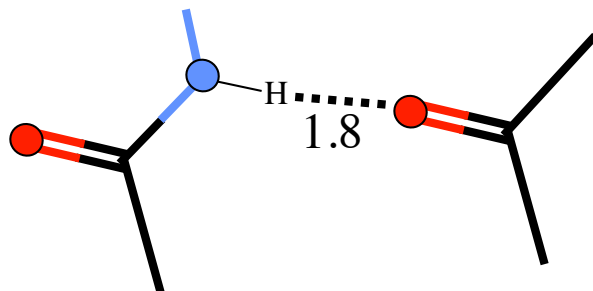
SVL: run 'gizmin.svl'

When finished, be sure to **Cancel | GizMOE\_Minimizer**

- To remove or modify restraints

**Potential setup** (button at far lower left)

Restraints tab





# Exercise 4.1

## Make a beta hairpin

anti-parallel sheet with valine side chains all on the same side of the sheet.

**Edit | Build | Protein**, Geometry: **anti-strand**. Residue: **ADVDVKVSPNGVEVKVRA**

Zoom out.

Select the second half of the chain starting with NG.

Rotate and translate it (**shift-alt-middlemouse**) so that the first three valines (3,5,7) are lined up with other three valines (12,14,16), and the valine backbone H-bonding groups (NH and CO) are close to the H-bonding distance (1.8Å from H to O)

**Hide** side chains to help see the backbone atoms better.

**Edit | Potential | Restrain**.

Set Target 1.8, 1.8, Weight 50. Select H and O atoms. **Create**.

When done you have 2 restraints for each of the three paired valines for a total of 6 restraints.

**Compute | Prepare | Structure preparation**. Hit **Correct** if necessary. **Protonate3D**.

**SVL: run 'gizmin.svl'**.

If there are errors in the restraints, **Cancel/GizMOE**, open **Potential Setup** (extreme lower left of the MOE window). **Restraints**. Click on restraints to delete or modify them.

Restart **SVL: run 'gizmin.svl'**.

Look at out the structure.

It should have beta pleating when viewed from the edge of the sheet. Sidechains should alternate up and down in that view. Residues SPNG form a beta-turn.

**Cancel/Gizmin** . Remove the restraints. Restart **SVL: run 'gizmin.svl'**.

**Does the structure hold together or fall apart?**