

Protein Fold Recognition Weak Similarities

- why do we do sequence alignments ?
 - find related proteins
 - build models
 - guess at function
- For some interesting protein
 - sequence always available
- What should one do with really weak sequence homology ?
- two ideas
 - how to search for very weak similarities
 - can one take advantage of conserved structures ?

Mission

Problem

- for some protein sequence – find as much as possible
 - function
 - build good model
 - build a bad model
- relatively vague information may be useful
 - which residues are near active site ?
 - which residues are near a dimer interface ?
 - which residues are in weakly structured loops ?
 - overall shape (bad model) may be enough for phasing

Approach

- start with most reliable methods
- add more speculative methods as necessary
- Example
 - simple sequence searches
 - searches for more remote homologues
 - searches for possible structures
- methods so far
 - emphasis on speed (in Georgio's lectures)

alignment methods

	slow	fast
methods	Needleman & Wunsch / Smith-Waterman	seeded – blast, fasta, suffix tree methods
time	$O(nm)$ or $O(nm^2)$ (sequence sizes)	$O(nk)$ – database size
guaranteed to find optimal alignment	yes	no
very remote homologues	may work	less likely to work

- does speed matter ?

Slow methods

- Methods for large databases are
 - fast
 - approximate
- Here
 - ultimate use is often a small database (PDB 8×10^4)
 - computer time does not matter
- In lab you have 1 or 10's of proteins
 - each take weeks or months to work on
 - if each search takes hours ? no problem
- remote searches

Remote searches

When to do this ?

- Assume simple (blast / fasta) search returned
 - related sequences
 - unknown function
 - none of related proteins have known structures

Weak sequence similarities

- Your sequence

yours **A B D E F G H I K L M N P Q...**

- finds no helpful proteins. Try searching with a related protein

prot_1 **A B Q E F G R I S L T N P Q...**

- finds a protein whose structure has been solved

prot_2 **Q B Q E Q G R Q S L T N P A...**

- claim
 - yours & prot_2 are related
 - relationship too weak to see directly
 - prot_2 can be used
 - to make a bad model
 - as a guess for function

Weak sequence similarities

- first idea
- take your protein
- collect related proteins
 - foreach (related protein)
 - do a sequence search
 - see if results change
- not practical
- not very systematic
- what else does one get from homologues ?

Information from related sequences

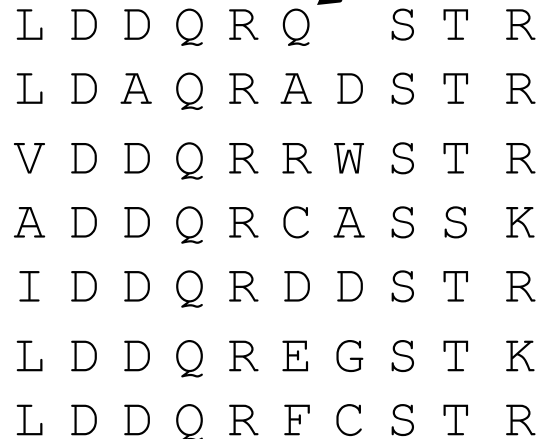
- usually one finds many related sequences.
- consider details...

VLSPADKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
MLSPADKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKTNVKAAWGKVGAGHAGEYGAEALEKMFSLFPTTKTYFPHFDLSHGSAQVKGHG
LSPADKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKTNVKAAWGKVGAGHAGDYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSRDDKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
MLSPADKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKTHVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKTNVKAAWGKVGAGHAGEYGAEAWERMFLSFPTTKTYFPHFDLSHGSAQVKGHG
MLSPADKTNVKAAWGKVGAGHAGEYGAEAWERMFLSFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
MLSPADKTNVKAAYWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKTNVKAHWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSAADKTNVKAAGWSKVGGHAGEYGAEALERMFLGFPTTKTYFPHFDLSHGSAQVKAHG
VLSAADKTNVKAFWSKVGGHAGEYGAEALERMFLGFPTTKTYFPHFDLSHGSAQVKAHG
VLSADDKANIKAEWGKIGGHGAEYGAEALERMFCSFPTTKTYFPHFDVSHGSAQVKGHG
MLSPADKTNVKADWGKVGAGHAGEYGAEAFERMFLSFPTTKTYFPHFDLSHGSAQVKGQG
VLSPADKTNVKACWGKVGAGHAGEYGAEAFERMFLSFPTTKTYFPHFDLSHGSAQVKGQA
VLSAADKSNVKAAWGKVGGNAGAYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
MLSPADKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKSNVKATWDKIGSHAGEYGGEALERTFASFPTTKTYFPHFDLSPGSAQVKAHG
VLSPADKSNVKAWWGKVGGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
MLSPADKTNVKAAWGKVGAGHAGEYGAEALERMFSLFPTTGTYFPHFDLSHGSAQVKGHG
VLSAADKNNVKACWGKIGSHAGEYGAEALERTFCSFPTTKTYFPHFDLSHGSAQVQAHG
VLSAADKSNVKAAWGKVGGNAGAYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSPADKTNVKAQWGKVGAGHAGEYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG
VLSANDKSNVKAAWGKVGNHapeYGAEALERMFSLFPTTKTYFPHFDLSHGSSQVKAHG
VLSPADKSNVKAAWGKVGGHAGDYGAEALERMFSLFPTTKTYFPHFDLSHGSAQVKGHG

... ..

Conservation

- as in secondary structure prediction lectures
- if your sequence has a Q here,
 - may not be helpful to use it in sequence searches

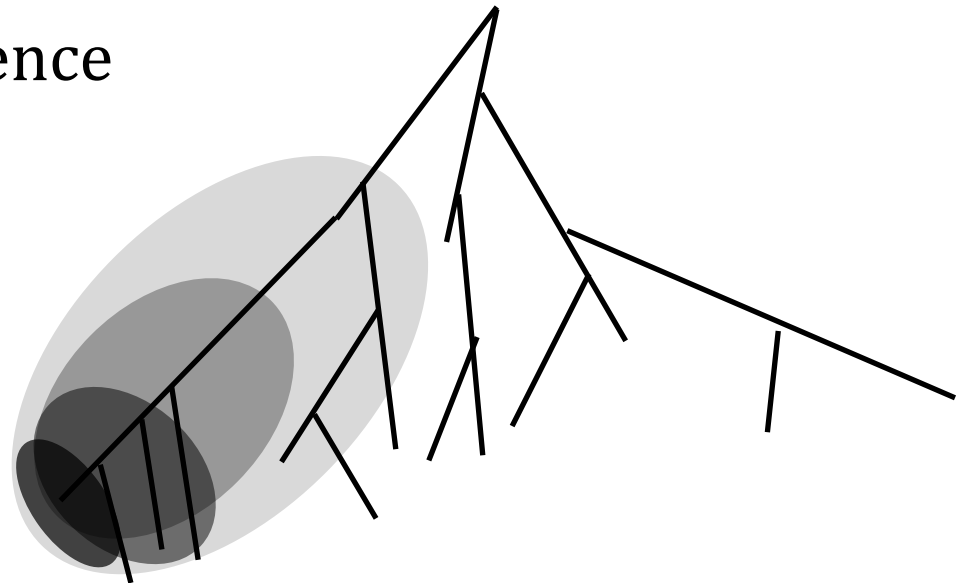


L D D Q R Q S T R
L D A Q R A D S T R
V D D Q R R W S T R
A D D Q R C A S S K
I D D Q R D D S T R
L D D Q R E G S T K
L D D Q R F C S T R

- better to use the "average" residue at this point
- first have to find the "average" residue
- leads to method

Searching with profiles

- initial average_sequence = your_sequence
- while (step < max_steps)
 - search with blast using average_sequence
 - if interesting result (function / structure..)
 - return results
 - else
 - update average_sequence
- basis of "psi-blast"
- does it work ?



Remote sequence searching

- much more sensitive than simple searches, but
- involves weaker sequence similarities, more errors
- alignment not perfect
- statistical significance harder to estimate
- possibility of finding unrelated sequences (rubbish)
- still relies on some significant sequence similarity
- can one move away from sequence similarity ?

Why move away from sequence

- if sequences provide information – use this
- when does it fail ?
 - new functions, not yet

Sequence alignments – implied structures

- From sequence viewpoint

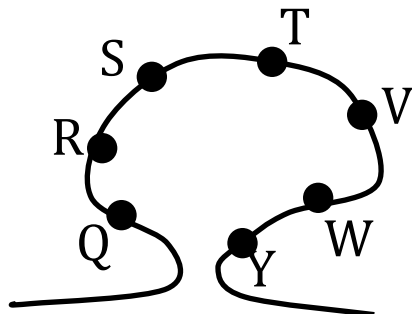
..AC-DEFG..

..QRSTUVWXYZ..

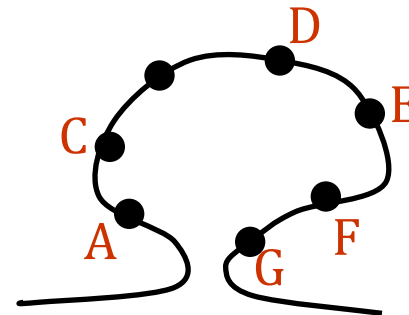
- what if structure of second sequence is known ?

..AC-DEFG.. query sequence

..QRSTUVWXYZ.. known structure



known
structure



model
implied

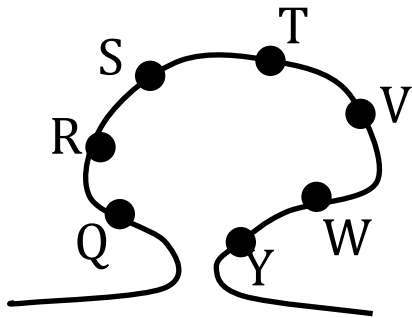
Sequence to structure alignments

- Remember how sequence alignments work
 - similarity / substitution scores
 - fill out score matrix
 - find best path
- Can we use this for sequence to structure alignments ?

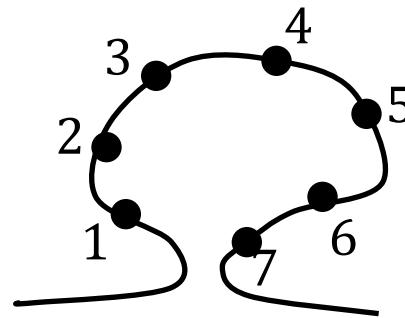
	A	B	C	N	J	R	O	C	L	C	R	P	M
A	4	7	6	6	5	4	4	3	3	2	1	0	0
J	7	7	6	6	6	4	4	3	3	2	1	0	0
C	6	6	7	6	5	4	4	4	3	3	1	0	0
J	6	6	6	5	6	4	4	3	3	2	1	0	0
N	5	5	5	6	5	4	4	3	3	2	1	0	0
R	4	4	4	4	4	5	4	3	3	2	2	0	0
C	3	3	4	3	3	3	3	4	3	3	1	0	0
K	3	3	3	3	3	3	3	3	3	2	1	0	0
C	2	2	3	2	2	2	2	3	2	3	1	0	0
R	2	1	1	1	1	2	1	1	1	1	1	0	0
B	1	2	1	1	1	1	1	1	1	1	1	0	0
P	0	0	0	0	0	0	0	0	0	0	0	1	0

more exotic scoring

- From sequence viewpoint
 - ..AC-DEFG.. my sequence
 - ..QRSTUVWXYZ.. a protein of known structure
- rather than just align sequences, could I use the structure ?



known
structure



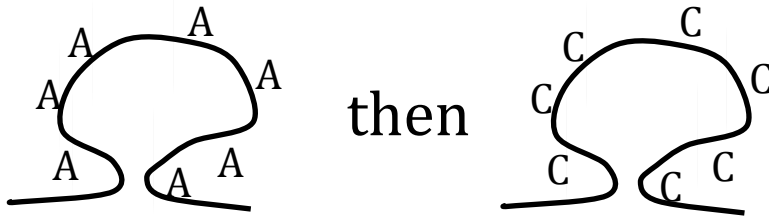
forget
sequence

- score matrix ?

	A	C	D	E	F	G
1	?	...				
2						
3						
4						
5						
6						
7						

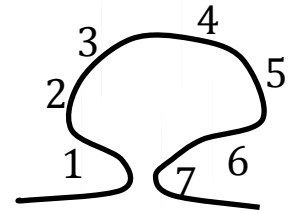
sequence to structure scoring

- I have to be able to place (A, C, D..) at each position and get a suitability score



	A	C	D	E	F	G
1	?	...				
2						
3						
4						
5						
6						
7						

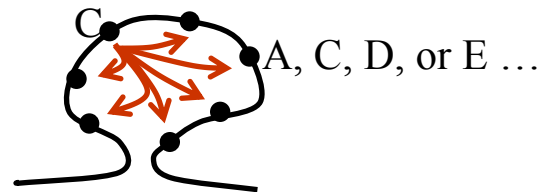
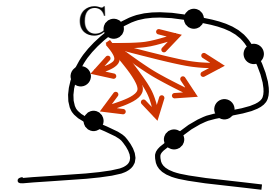
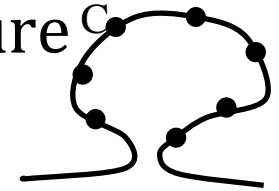
- then it would be easy to do sequence to structure alignments
- advantage:



- we claim that structure is more conserved than sequence
- can find appropriate/fitting/suitable structures for a sequence
- very remote, but homologues
- vorsicht !!!!

sequence to structure scoring

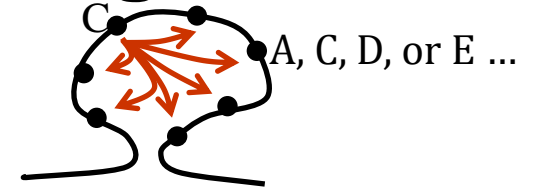
- define an energy function
 - depends on interaction of residue with structure
 - easy
 - depends on interaction with neighbours
 - but who are the neighbours ?



- bad news
 - we cannot even fill out a column in the score matrix
 - to test every combination of neighbours
 - NP-complete
- an excuse to try some approximations

	A	C	D	E	F	G
1	?					
2	?					
3	?					
4	?					
5	?					
6	?					
7	?					

approximations for scoring



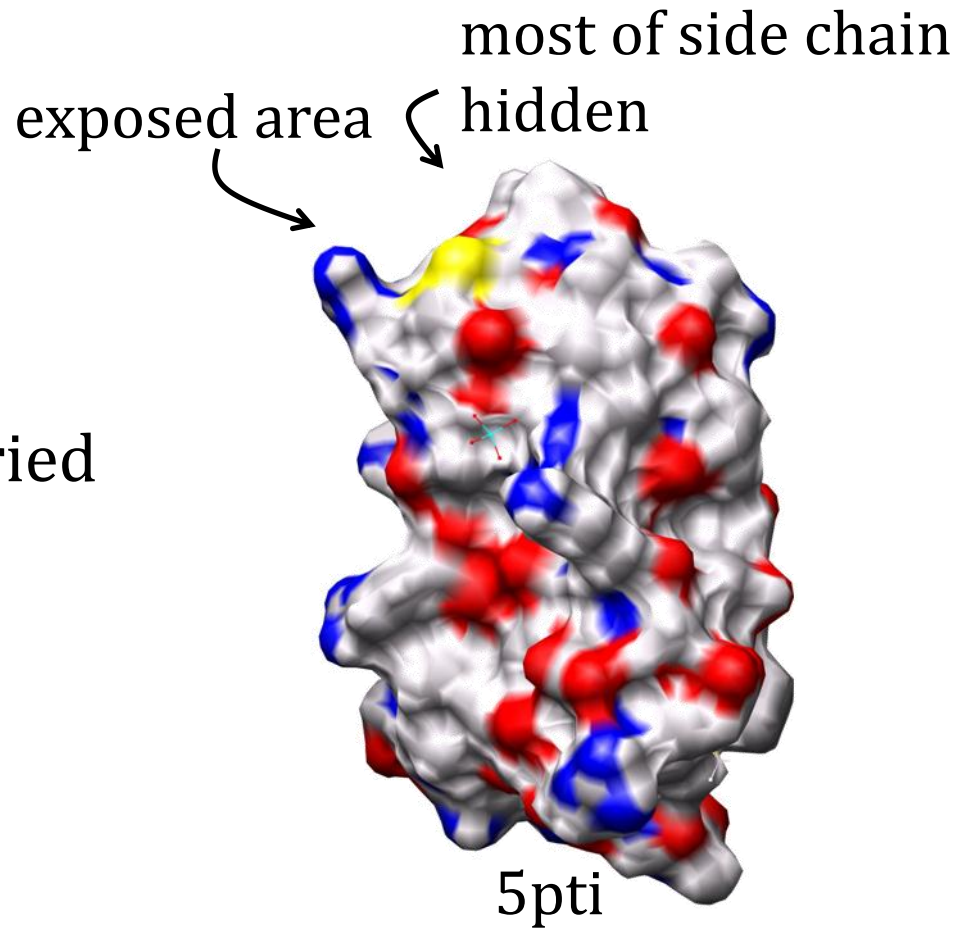
- two problems
 - we do not know where all the atoms are – side chain coordinates
 - to score "C" at each position we need to know neighbours
- side-chains : ignore / average
 - use a score / energy function which averages over all conformations
- neighbour positions : much harder
 - environment description
 - frozen approximation

Environment description

- an example of profiles (case study)
- we know
 - certain sites are hidden from solvent (middle of protein)
 - only compatible with trp, phe, ile, ... (hydrophobic)
 - some sites are involved in "salt bridges"
 - some secondary structures are preferred by certain residues
- can one count the probabilities of residue types ?
- overview
 - collect list (parameterisation set) of proteins
 - classify sites (18 types)
 - collect probability of each residue type in each site type

Environment description

- for each site measure the \AA^2 exposed to solvent
- maybe sometimes one has charges / polar groups touching others
 - measure fraction of buried area covered by polar groups
- define environments...



Environment description

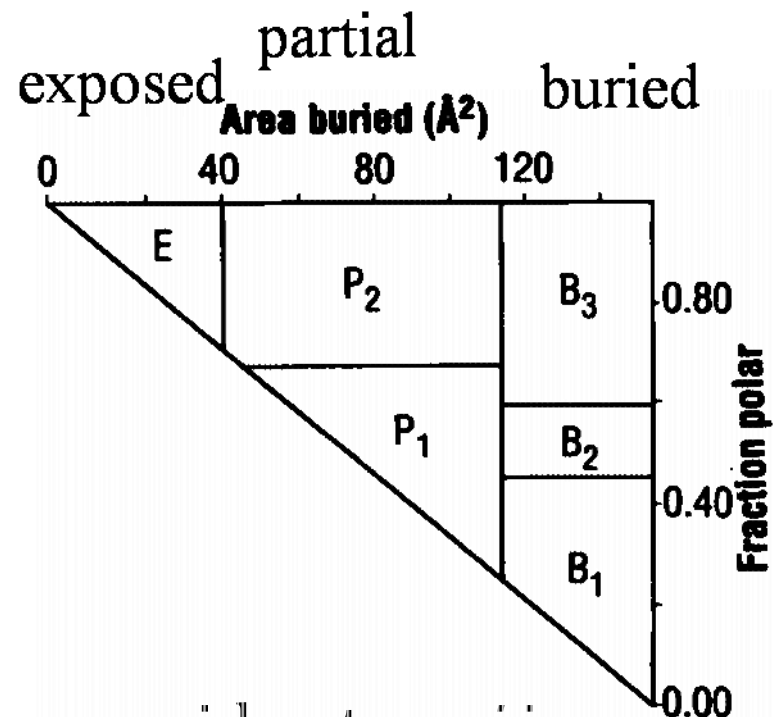
- 6 environment types
- 3 secondary structure types
 - α , β , others
- = 18 environments

- data collection

- 16 proteins
- find environment of each site
- count

- how many times does one see residue type i in environment $j = N(i,j)$

- count – how often does one see residue type $i = N(i)$



Environment description

- how unusual is a residue i in environment j ?

$$score(i, j) = \ln \left(\frac{N(i, j)}{N(i)} \right)$$

- final result ? a big scoring table

likely

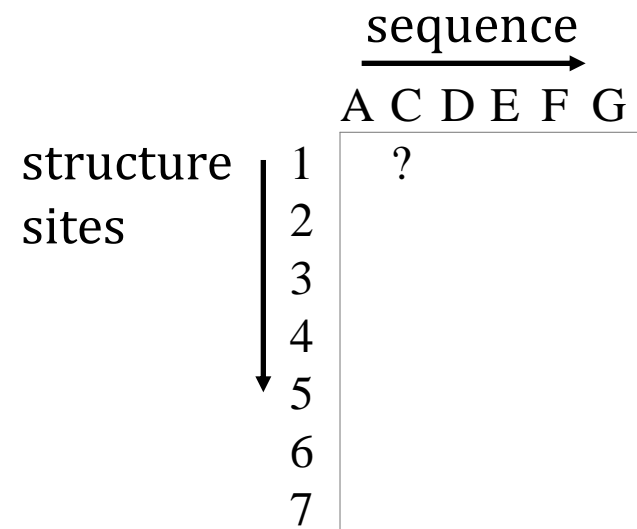
unlikely

what one expects

Environment class	W	F	Y	L	I	V	M	A	G	P	C	T	S	Q	N	E	D	H	K	R
B ₁ α	1.00	1.32	0.18	1.27	1.17	0.66	1.26	-0.66	-2.53	-1.16	-0.73	-1.29	-2.73	-1.08	-1.93	-1.74	-1.97	-0.34	-1.82	-1.67
B ₁ β	1.17	0.85	0.07	1.13	1.47	1.09	0.55	-0.79	-2.02	-0.94	-0.22	-1.12	-2.91	-1.67	-1.42	-1.93	-2.56	-1.91	-2.69	-1.16
B ₁	1.05	1.45	0.17	1.10	1.11	1.02	0.98	-0.91	-1.92	0.26	-1.22	-1.53	-2.81	-1.17	-2.42	-2.52	-1.76	-1.12	-2.59	-2.16
B ₂ α	0.50	0.90	0.85	1.01	0.83	0.68	1.12	-0.89	-1.49	-2.21	-0.10	-1.50	-1.47	-0.23	-0.81	-0.71	-1.62	0.23	-0.78	0.06
B ₂ β	0.01	1.18	1.06	0.76	1.31	1.06	0.64	-1.55	-2.28	-0.49	-0.87	-2.27	-1.77	-1.22	-2.07	-1.07	-1.41	-0.77	-1.14	-0.20
B ₂	1.02	1.05	1.12	0.84	0.81	0.60	0.90	-0.66	-1.66	0.19	-0.05	-0.76	-1.17	-0.76	-0.66	-1.35	-1.28	0.46	-2.34	-0.80
B ₃ α	0.92	-0.03	0.58	0.15	0.04	-0.02	0.89	-0.57	-1.86	-0.68	-1.56	-0.57	-0.96	0.22	-0.06	0.08	-0.50	0.73	0.43	0.96
B ₃ β	0.75	0.81	1.30	0.18	0.54	0.56	-0.57	-0.93	-1.93	-0.34	-0.54	-0.44	-0.74	0.21	-0.24	-0.14	-0.86	0.82	-0.53	0.13
B ₃	1.07	0.70	1.13	0.35	-0.17	-0.03	0.23	-0.96	-0.98	-0.13	-1.20	-0.53	-0.54	0.05	0.04	-0.36	-1.05	1.01	0.10	0.66
P ₁ α	-1.35	-0.82	-0.59	-0.52	-0.24	0.10	-0.03	0.73	-0.49	-0.25	0.95	0.31	0.34	-0.14	-0.54	-0.17	-0.25	-0.52	-0.21	-0.28
P ₁ β	0.36	-0.49	0.17	-1.03	0.20	0.46	-0.27	0.64	-0.82	-0.55	1.49	0.93	0.33	-2.27	-1.32	-0.73	-1.07	-0.42	-1.21	-0.77
P ₁	-1.26	-1.20	-1.31	-0.62	-0.23	-0.01	-1.19	0.46	-0.24	0.66	1.35	0.56	0.49	-0.63	-0.13	-0.61	0.38	-1.12	0.74	-1.29
P ₂ α	-1.14	-1.43	-0.79	-0.35	-0.54	-0.48	-0.45	0.06	-0.50	-0.26	-0.93	-0.05	-0.18	0.55	-0.05	0.56	0.28	0.06	0.61	0.50
P ₂ β	-0.79	-0.54	-0.84	-1.30	-0.33	0.13	-0.72	-0.55	-0.98	-1.29	-0.57	0.84	0.59	-0.08	-0.16	0.32	0.19	-0.87	0.59	0.10
P ₂	-0.82	-0.86	-0.51	-0.70	-1.09	-0.88	-0.89	-0.15	-0.40	0.44	-0.80	0.06	0.26	0.27	0.50	0.27	0.49	0.13	0.44	0.30
E α	-1.35	-2.20	-2.10	-1.58	-2.76	-1.10	-0.72	0.46	0.68	0.04	-0.44	-0.17	0.15	0.36	0.28	0.59	0.44	-0.19	0.13	-0.34
E β	0.64	-0.90	0.30	-1.66	-1.47	-1.74	-0.68	0.06	1.46	-0.96	-0.24	0.14	0.65	-0.19	-0.06	-0.16	-0.78	-0.83	-0.52	-0.49
E	-2.14	-1.90	-0.94	-1.19	-1.61	-0.91	-1.67	0.12	1.13	0.20	-0.46	0.12	0.32	-0.03	0.41	0.03	0.22	-0.25	-0.14	-0.32

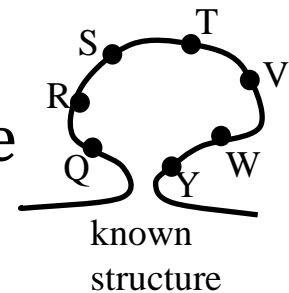
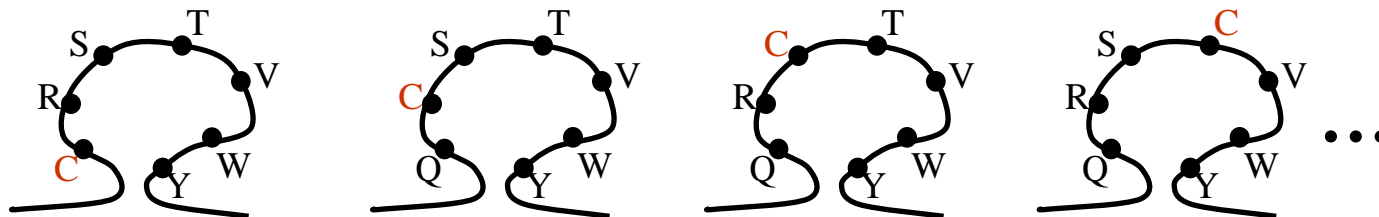
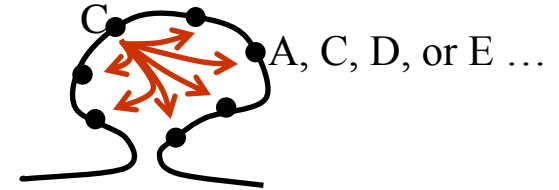
Environment description - application

- given these descriptions – use them
- take a protein structure label each site
- take sequence of interest
- for each residue
 - score at each site of protein
- score matrix
- find best path
 - sequence to structure alignment
- final application
 - take protein databank
 - try to align your sequence to every structure



Frozen approximation

- original problem
 - we want to use a score function which
 - sensitive to sequence
 - sensitive to structure
- remember – original structure did have a sequence
- belief
 - if two proteins are related, the sequences will have similar properties
 - score with the residues of the original sequence



Frozen approximation

- I can score my sequence in the environment of the known structure
- good
 - the environment is well characterised
 - if my structure has polar residues here, they will go into the scoring function
- bad ?
 - we use the sequence of template (known structure)
 - it may only allow very related residues
 - original aim was to move away from close sequences

	sequence						
	A	C	D	E	F	G	
structure	1						
sites	2						
	3						
	4						
	5						
	6						
	7						

Summary so far

- look for closely related templates
- try sequence based methods
- sequence to structure methods are definitely possible
- can I make better scoring schemes ?

Scoring schemes

... S T D G W Y F I L S T ...
polar / charged | small | hydrophobic | polar

- how much structural information is hidden in sequence ?
- look at a sequence
- I already have labels for sites
 - implicit in substitution matrices
- does the structure contain extra information ? ...

Extra information from structures

Residues exist in a protein for different reasons

- gly is easy to substitute – look at diagonal in blosum matrix
- in some turns, gly is essential
 - can only be seen from structure
- cys
 - sometimes a normal hydrophobic residue
 - sometimes the geometry says it must form a disulfide bond
 - structure can say if there is another cys near in space
- ...
- it should be useful to combine sequence and structure information

Extra information from structures

- Claim – hope
 - combination of methods has better signal / noise
- implementation ? easy in principle
 - for each residue i in your query sequence
 - for each site j in template
 - calculate sequence score s_1 based on profile of i
 - calculate structural score s_2 based on fitting residue type i into site j
 - score for alignment matrix = $s_1 + k s_2$
 - for some constant k

In practice

- most fold recognition programs combine sequence terms and structural scores
- results may or may not be better than best pure sequence methods
- problems..

Problems with clever methods

- Simple sequence searches
 - good models for statistical significance
 - (is a related protein really related ?)
- Remote sequence searches (psi-blast)
 - statistics OK, but less reliable
- Structure / Sequence+structure methods ?
 - no good model for scores
 - no good model for statistical significance
- how will score grow with
 - size of query ?
 - size of alignment ?
 - sequence composition ?

Principle

- If you have extra information (structure)
 - must be a good idea to use it

	sequence	structure based
database size	10^6-10^7	10^4
	fast	slow
scores	good models	weaker
statistical significance	good or almost good	weaker

Summarise and stop

- Use sequence information when possible
- use adventurous sequence methods when necessary
- use very speculative methods (sequence to structure) when necessary