## **Classifying and comparing proteins**

#### Plan

- why ?
- domains
- classifications
  - hierarchical vs pragmatic / empirical
  - continuous or clustered ?
- sequence similarity vs structure similarity
- example classifications
- comparison measures

Andrew Torda, Wintersemester 2016 / 2017, GST...

# Why?

Background – details later

- evolutionarily close proteins similar structures
- evolutionarily remote proteins may have similar structures
- function prediction / annotation
- interpretation
- structure prediction can I predict this sequence fits to that structural class ?

Examples..

## **Transfer of properties**

Arguments as with homology

- Homology modelling
  - can I find a related protein ?
  - can I say my protein has similar function / structure ?
- Classifications of proteins
  - I have classes of proteins some members are well characterised
  - If you can say your protein is in class X,
    - probably has similar function to other members

## **Structure prediction**

How many possible protein structures are there?

• astronomical

How many interesting / different protein structures actually occur on earth ?
2×10<sup>3</sup> to 5×10<sup>3</sup>

*de novo / ab initio* prediction ?

• search in giant space

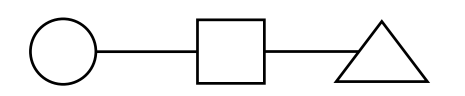
Find most likely protein fold?

- search amongst  $10^3$  to  $10^4$  structures
- find the class of your protein crude structure prediction

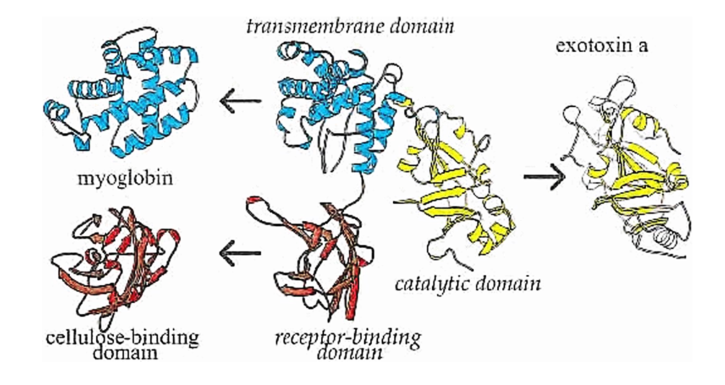
### Domains

We will usually talk about protein domains (not whole proteins)

• association of domains with function and evolution..



• most literature classifications work with domains



#### **Domains for these lectures**

- Usually structure based
- compact units
- In these lectures
- no functional domains
- no sequence-based

Should we classify by structure or sequence?

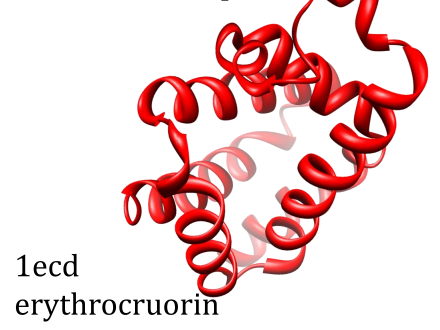
### **Structure vs Sequence similarities**

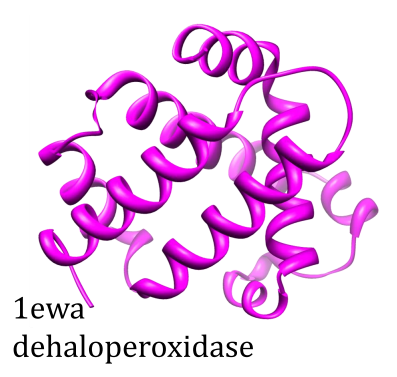
More different than you might expect

Similar sequences

- have not evolved for too long
- expect similar structures

Other way round ? Examples





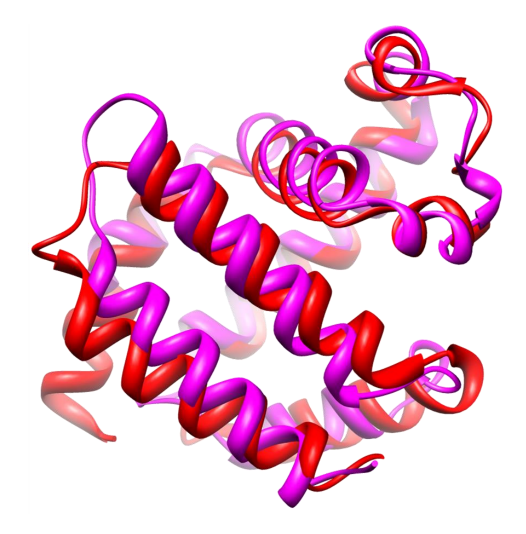
## very different sequences

1ecd & 1ewa

- 17% sequence identity (very low)
- structures almost identical

Is this an exception ?

- 100's of examples
- totally normal
- play with our server



http://flensburg.zbh.uni-hamburg.de/~wurst/salami/

### **Example family**

Example, neighbours of 1cun chain A

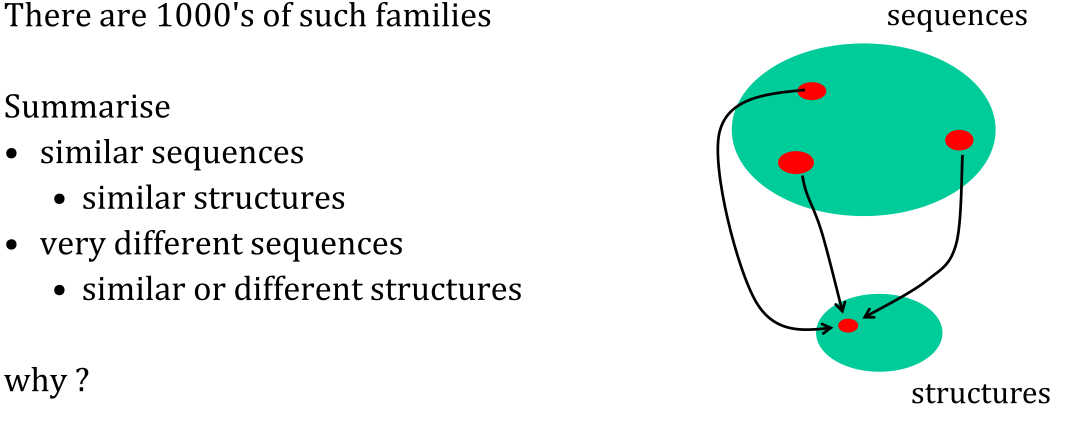
• look at sequence identity (% id)

root mean square diff in Å

	No	Chain	%id	lali	rmsd	Description
		1cunA				ALPHA SPECTRIN
	2	<b>1hciA</b>	24	111	1.6	ALPHA-ACTININ 2
	3	lek8A	12	106	4.4	RIBOSOME RECYCLING FACTOR
	4	10xzA	9	91	2.5	ADP-RIBOSYLATION FACTOR BINDING PROTEIN GGA1
	5	leh1A	8	102	4.6	RIBOSOME RECYCLING FACTOR
	6	1hx1B	5	105	3.1	HEAT SHOCK COGNATE 71 KDA
	7	1dd5A	8	103	4.7	RIBOSOME RECYCLING FACTOR
	8	<b>1lvfA</b>	9	98	2.6	SYNTAXIN 6
		1bg1A		99	2.3	STAT3B
	10	1hg5A	5	98	3.0	CLATHRIN ASSEMBLY PROTEIN SHORT FORM
		1hs7A		92	2.5	SYNTAXIN VAM3
	12	1dn1B	10	101	2.7	SYNTAXIN BINDING PROTEIN 1
	13	1ge9A	6	108	4.6	RIBOSOME RECYCLING FACTOR
	14	<b>lfewA</b>	8	125	3.5	SECOND MITOCHONDRIA-DERIVED ACTIVATOR OF
	15	1qsdA	4	90	2.4	BETA-TUBULIN BINDING POST-CHAPERONIN COFACTOR
-	16	1e2aA	6	95	2.8	ENZYME IIA
th	17	1i1iP	7	95	3.3	NEUROLYSIN
	18	1fioA	8	100	2.6	SSO1 PROTEIN
	19	1m62A	8	81	2.8	BAG-FAMILY MOLECULAR CHAPERONE REGULATOR-4
	20	1k4tA	6	147	25.8	DNA TOPOISOMERASE I

alignment length

#### **Structure vs Sequence**



Summarise

- similar sequences
  - similar structures
- very different sequences
  - similar or different structures

why?

## **Structures < Sequences... Why ?**

Evolution

- many small changes
- if structure changes, function breaks, you die
- sequences change as much as possible within this constraint

Chemistry

- sequence determines structure
  - many sequences could fit structure (more next semester)

Surprising?

- consider near universal proteins
  - 100's millions years evolution, function largely preserved
  - sequence has changed radically

# **Classifying by sequence**

#### Forget hierarchy (for now)

- tools any alignment program (blast, fasta, clustal, ...)
- method
  - survey all proteins in the protein databank
  - collect all pairs > x %
- result (Nov 2016)
- how many structure classes ?  $2 \times 10^3$  to  $5 \times 10^3$  ?
- some sequence classes are not really different from each other

Now.. examples of structure based classifications

similarity	num clusters
90 %	41 097
70%	34 547
50%	27 787

## **Clusters and hierarchies**

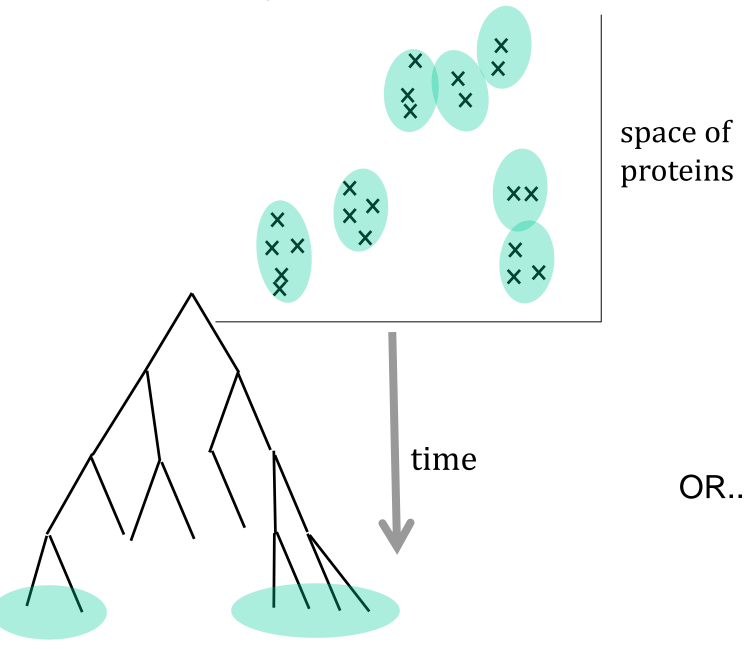
Are there clusters ? Yes

- Sequence-based ? Do a sequence search for a haemoglobin or profilin
  - find 10<sup>3</sup> to 10<sup>4</sup> homologues this is some kind of cluster
- Structure-based ?
  - search for haemoglobins (or your favourite protein)
  - find  $10^2 10^3$  similar structures a cluster

Are they hierarchical ? No idea

• what is the question ? (reminder from last lecture)

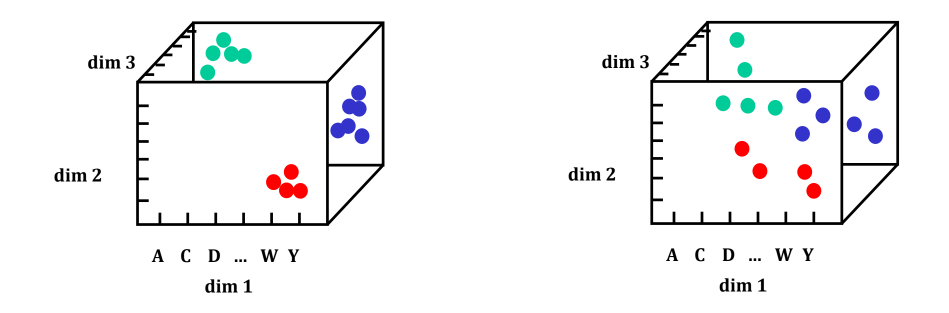
#### Maybe there should be protein clusters



12/5/2016 [14]

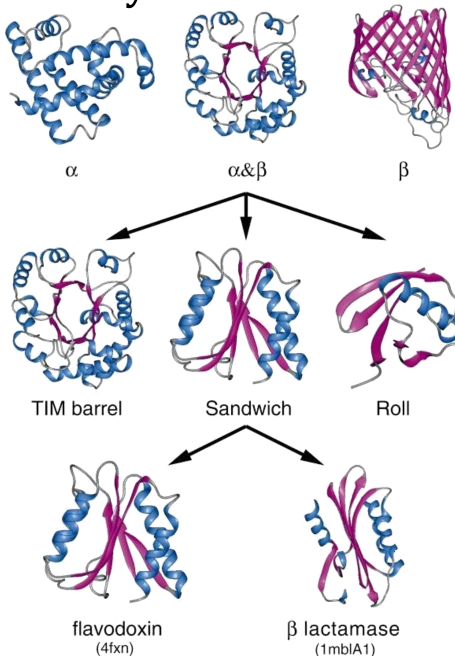
If we knew every protein that every existed anywhere

• would we be able to connect the clusters ?



• An example of a hierarchical classification

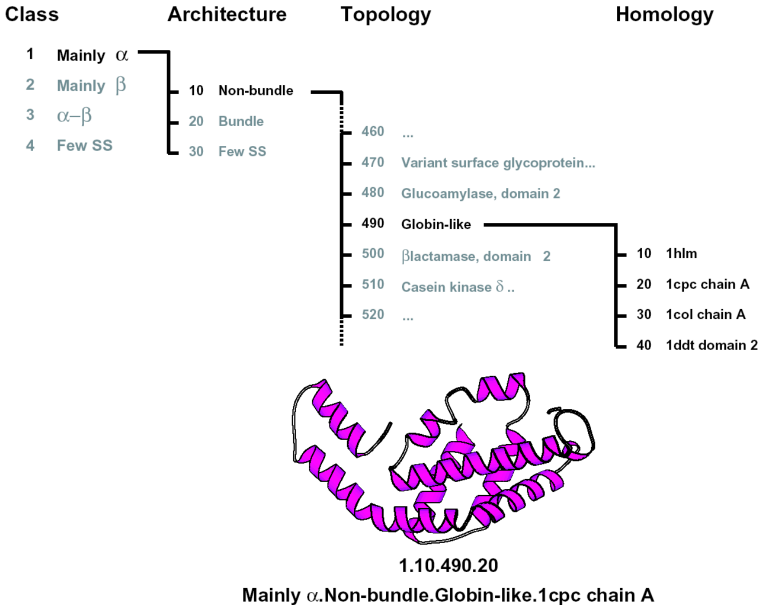
#### **Imposing a Hierarchy on Proteins**



- parts may correspond to evolution
- top level ?
- How useful and applicable ?
- examples

12/5/2016 [16]

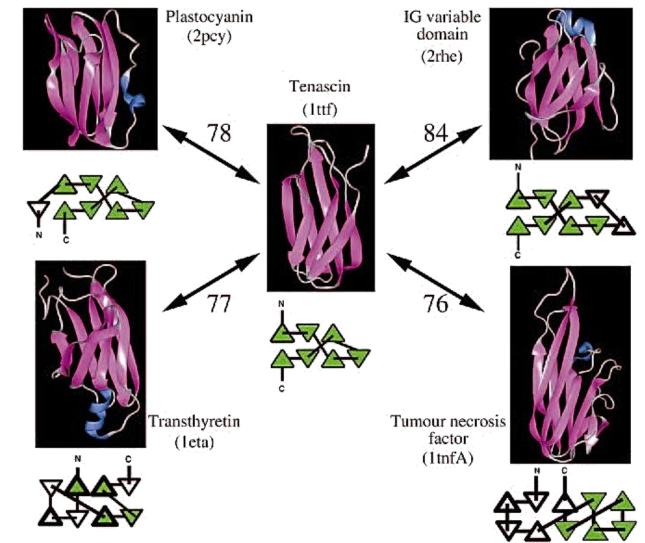
#### **Example from "CATH"**

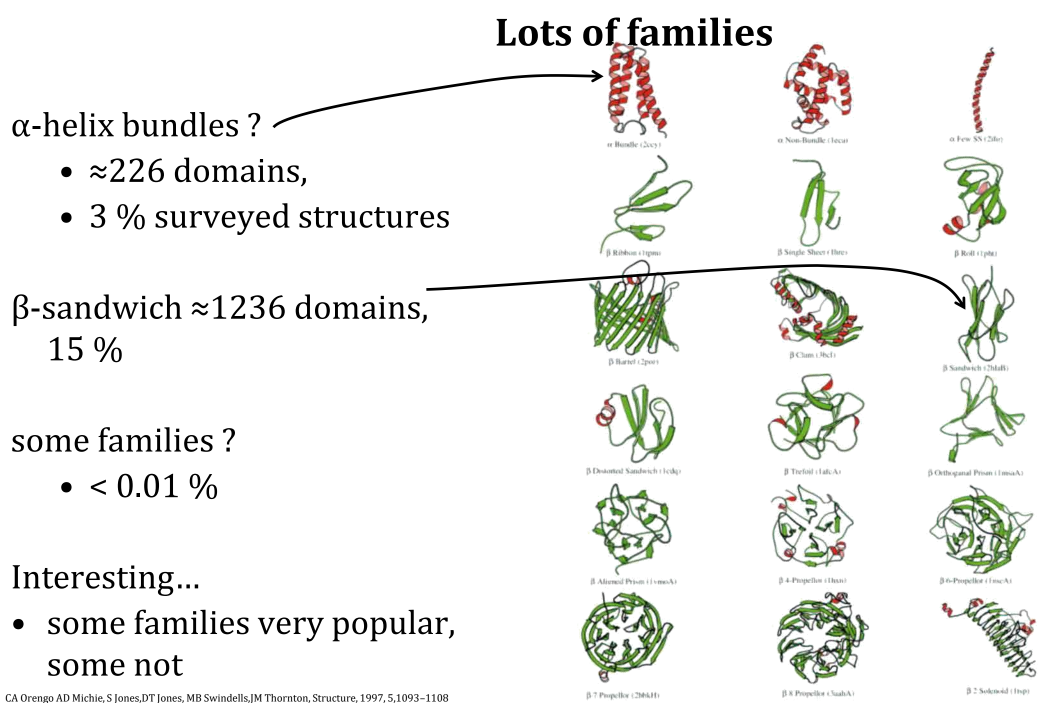


## **Evolution and Classification**

Can we interpret structures in evolutionary terms?

- sometimes
- for more remote proteins
   not really possible
- some typical figures from a literature classification





12/5/2016 [19]

### Some families populated more than others ?

Are some structures more stable ? physics ?

Can some "accommodate" more sequences / tolerate more mutations?

• next semester

Are some older in evolutionary terms?

Biases? PDB has

- mainly soluble, globular proteins which crystallised
- few membrane-bound proteins

# **Hierarchy**?

- Is the hierarchy really justified ?
- at low levels maybe
- at higher levels ?  $(\alpha, \alpha/\beta, ...)$

Better to discover relationships automatically

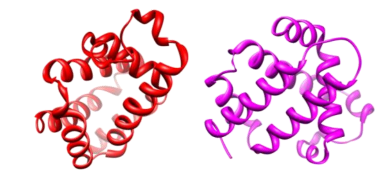
Imagine I can compare arbitrary proteins

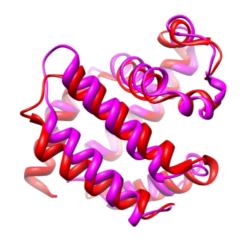
- have some measure of similarity
- use this to classify

Huge problem

- proteins are different sizes and shapes
- how to compare ?







## **Summary**

- Classification would be useful
- Given a distance (dissimilarity) one can invent a space for sequences or structures
- not known if it
  - exists
  - is hierarchical
- sequence vs structure similarity
  - different sequences can fold to same structure
- imposing a hierarchy on protein structures very *ad hoc*
- one can forget hierarchy simply use a clustering method
  - one will need a measure of similarities
  - big topic...

#### **FORGET HIERARCHIES**

- forget evolution
- forget hierarchies
- just look for similarities

### **Protein Structure Comparison / Numerical**

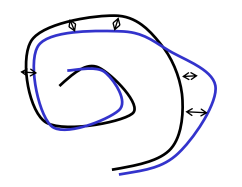
Most common protein structural question

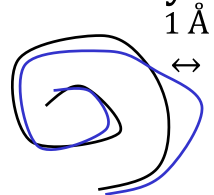
- how much has my protein moved over a simulation?
- how similar are these NMR models for a structure ?
- how close is my model to the correct answer ?
- more difficult
  - how similar is rat to human haemoglobin?
- two cases
  - 1. same protein, same number of atoms
  - 2. different proteins
- first
  - measures for easy cases

#### **Numerical Comparison of Structures - Easy**

What units would we like?

- scale of similarity (0 to 1.0)?
- comparison of angles
- distance / Å ? most common / easy to interpret
- looks a bit like the average difference between coordinates
- consider analogy with standard deviation / variance





#### **From Standard Deviation to RMSD**

Analogy with comparing a set of numbers

• get average (mean)  $\bar{x} = N^{-1} \sum_{i=1}^{N} x_i$ 

• standard deviation 
$$\sigma = \left(N^{-1}\sum_{i=1}^{N}(x_i - \bar{x})^2\right)^{1/2}$$

• apply this to coordinates of *r* and *r*'

$$rmsd = \left(\frac{1}{N}\sum_{i=1}^{N} |\vec{r_i} - \vec{r'_i}|^2\right)^{1/2}$$

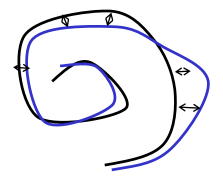
• rms / rmsd / RMSD = root mean square difference

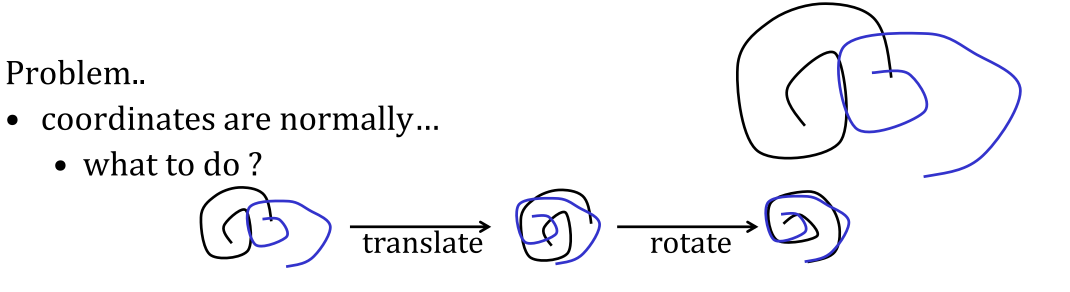
## **Calculating rmsd**

$$rmsd = \left(\frac{1}{N}\sum_{i=1}^{N}|\vec{r_{i}} - \vec{r_{i}'}|^{2}\right)^{1/2}$$

start at one end

- difference between pairs of atoms
- $|\vec{r}_i \vec{r}'_i|^2 = (x_i x'_i)^2 + (y_i y'_i)^2 + (z_i z'_i)^2$





#### **Translation and Rotation**

translation

- c.o.m. = centre of mass  $\vec{r}_{com} = (\sum_{i=1}^{N} m_i)^{-1} \sum_{i=1}^{N} \vec{r}_i m_i$
- subtract difference vector  $\vec{r}_{diff} = \vec{r}_{com} \vec{r}'_{com}$ rotation
- rotation matrix to minimise

$$rmsd = \left(\frac{1}{N}\sum_{i=1}^{N}|\vec{r_i} - \vec{r'_i}|^2\right)^{1/2}$$

summary

- translate
- rotate
- apply formula
- Still not finished

## Which Atoms ?

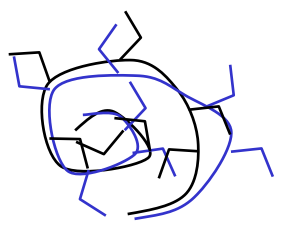
What tells me the shape of a protein ?

• backbone trace

What happens if you include all atoms ?

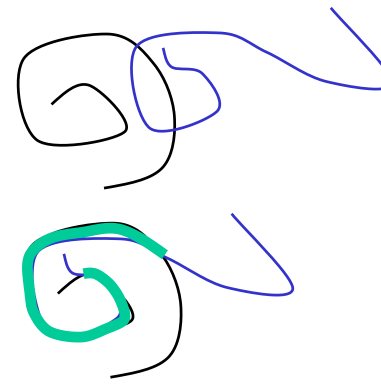
- bigger *rmsd*
- normal choice
  - C<sup>α</sup>
- sometimes
  - N, C<sup>*α*</sup>, C
- all atoms ?
  - when a model is very close

Still not finished with simple *rmsd* 



### **Parts Of Proteins**

- Two models of a molecule
- mostly very similar
- is *rmsd* a good measure ?
  Identify similar parts (method used in chimera)



#### define

```
superimpose ({r}, {r'}, {d}) {
    translate ({r,}, {r'}, {d})
    rotate ({r}, {r'}, {d})
}
where {d} is some subset of sites
```

#### **Selection of Interesting Atoms**

Define a threshold like thresh =2 Å

```
{d}={|r<sub>i</sub>-r'<sub>i</sub>|} i=1..N
sort {d}
```

```
diff= rmsd ({r_i}, {r_i'})
while (diff > thresh) {
  remove largest d
  superimpose ({r}, {r'}, {d})
  recalculate distances
 diff = rmsd (\{r\}, \{r'\}, \{d\})
}
if (diff < thresh)
  return {d}, diff
else
  return broken
```

Result ? a subset of interesting atoms

### **Subsets of Atoms**

Originally, quantify structural differences as Å *rmsd* 

Alternative quantity implied

• number of residues used for *rmsd* below threshold

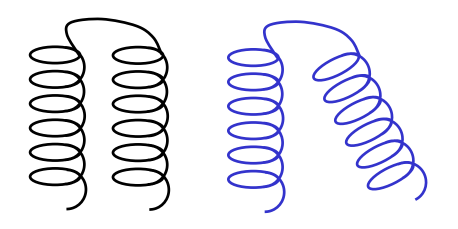
Implicit rule

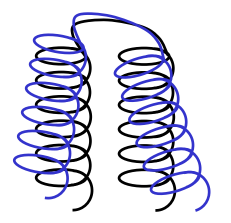
• as number of atoms  $\downarrow$  calculated *rmsd*  $\downarrow$ 

### Why not to use *rmsd*

#### Helices identical, fold identical

• *rmsd* ?





• superposition requires rotation, affects all atoms

- big *rmsd*, but structure has hardly changed
- do not see that helices are identical
- more problems

### Size dependence

Two proteins with 5 Å *rmsd* – similar or not ?

Consider proteins of different sizes

- maximum difference with  $N_{res} = 50$  or  $N_{res} = 100$ ?
- consider random structures with  $N_{res}$  = 50 or  $N_{res}$  = 100
- for small proteins 5 Å *rmsd* may be bad
- for large proteins 5 Å *rmsd* may be almost identical

extends to comparisons of small molecules

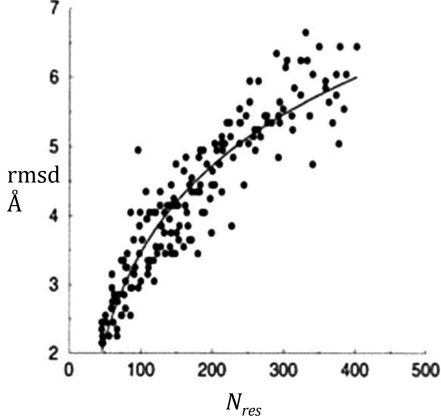
• ligands / medikamente...

What would one expect for random structures ?...

## Size dependence

Empirical

 survey of random protein comparisons



#### Theoretical

• can find result from compact polymer theory (Florey) not in these lectures

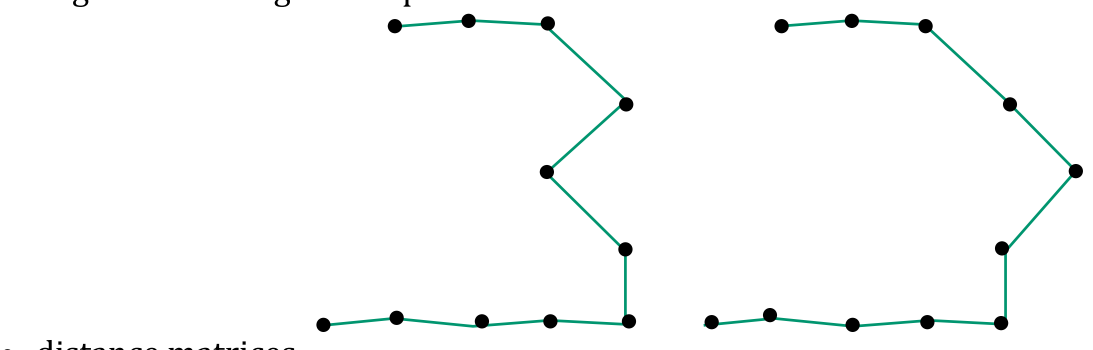
## rmsd size dependence

good rule

•  $rmsd_{interesting} = a + b(N_{res})^{1/3}$  for some constants *a*, *b* 

problems with *rmsd* measure – alternatives

• angles ? OK – angles compensate for another



distance matrices ...

#### **Distance Matrices With Numbers**

Another characteristic of structures

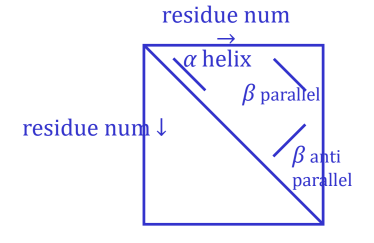
- C<sup>α</sup> distance matrices
- measure the distance between  $C^{\alpha}$  atoms

	1	2	3	4	5	6	7			Ν
1	0	3.8	6	7						
2		0	3.8	5						
3			0	3.8	4.5					
4				0	3.8					
5					0	3.8				
6						0	3.8			
7							0	3.8		
								0	3.8	
									0	3.8
N										0

#### **Distance Matrix for Recognizing Structure**

One way to summarise a structure

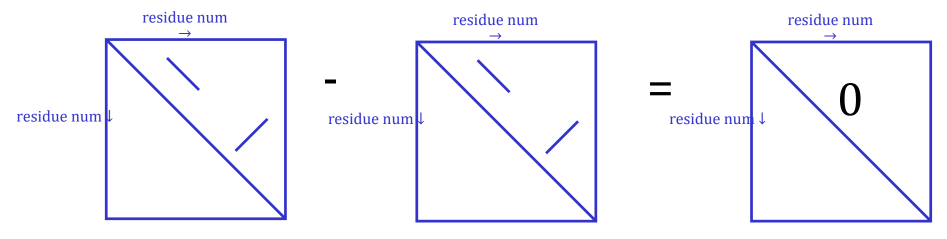
- plot  $C^{\alpha}$  distance matrix, points below 4 Å
- can make  $\alpha$ -helices and  $\beta$ -sheets clear



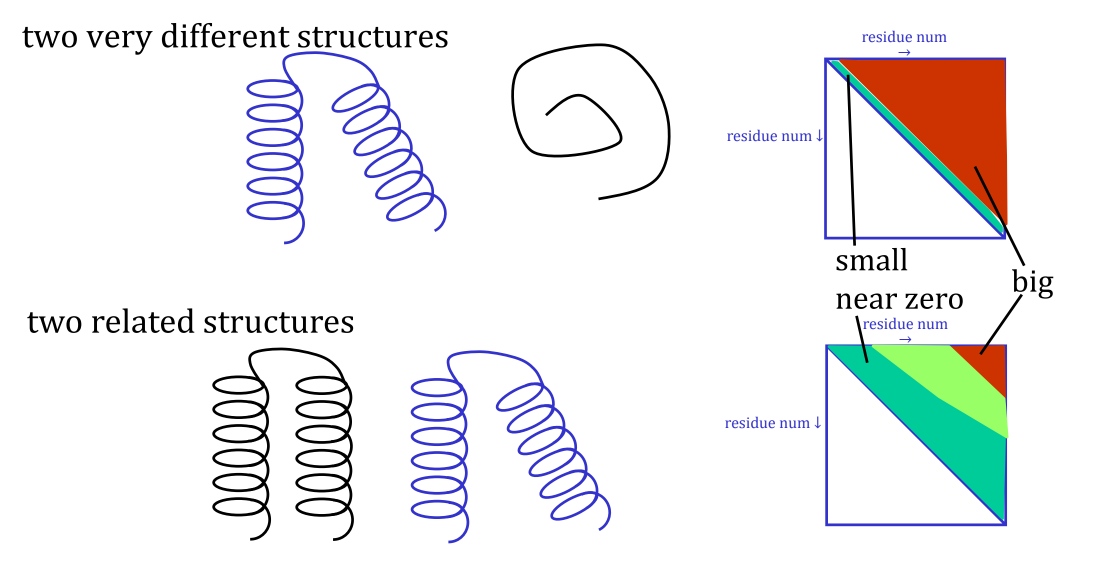
## **Distance matrix for comparing structures**

#### Take two similar proteins

#### look at the difference of distance matrices



## **Comparing Distance Matrices**



pictures are better than any single measure, but...

#### **From Distance Matrices to Single Number**

For lots of comparisons, single number is more convenient

Root mean square (*rms*) difference of distance matrices

• distance between  $C^{\alpha}$  atoms *i* and *j*  $d_{ij} = |\vec{r}_i - \vec{r}_j|$ 

*rms* of distance matrices measure is

$$rms = \left(\frac{2}{N(N-1)}\sum_{i=1}^{N}\sum_{j>i}^{N} (d'_{ij} - d_{ij})^{2}\right)^{1/2}$$

Like all other *rms* quantities

• normalised over top half of matrix

#### **Summary – Comparing Models / Structures**

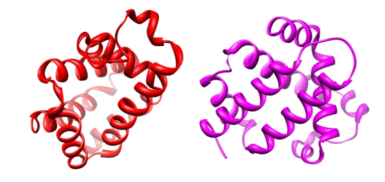
- rmsd
  - most popular
  - requires superposition (translate + rotate)
  - can be fooled by "hinge" movements
  - size dependent
- to look at the shape of a molecule use  $C^{\alpha}$  or backbone atoms
- numbers in Å have a physical meaning
- to look for the common core of a structure, find a subset of backbone
- other measures may be better than *rmsd*
- weakness of all measures
  - a single number can never capture all information

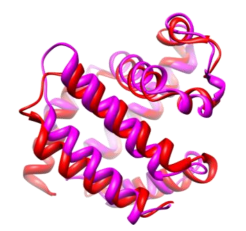
## **Comparing Proteins – different sizes**

- compare red and blue proteins
- if we know which residues match
  - easy (use any *rms* formula)
- which residues match ?
  - sequence alignment?

protein 1A C D W Y T RP K L H G FD S A C V Nprotein 2A C D W V T -P K V H G YD S A C V N

- green residues mismatches (no problem)
- blue residues ignore
- is this useful for similar proteins ? very (rat vs human haemoglobin)
- for very different proteins ? no



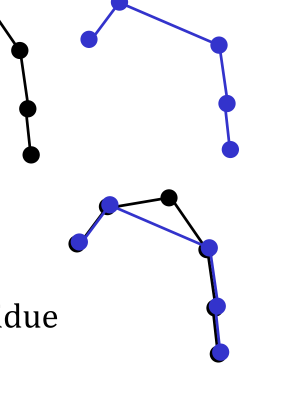


## **Comparing Very Different Proteins**

- Sequence alignment vs identity
- as identity ↓, errors ↑
- Consequence
- methods needed
  - operate on  $C^{\alpha}$
  - do not require sequence
- How difficult?
- superposition requires recognising the deleted residue
- can we use standard dynamic programming?

• no

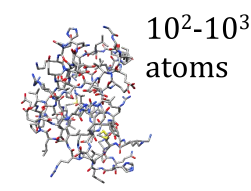
- gap/insertion at any position, any length
  - combinatorial explosion

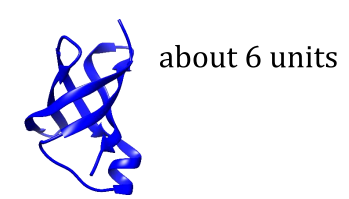


# Strategies For Comparing Different Structures 1. use secondary structure

Combinatorial explosion is the problem

- reduce size of problem
- use elements of secondary structure





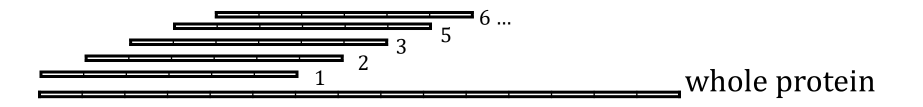
- define secondary structure
- search for superposition
- for each residue
  - find closest  $C^{\alpha}$  in partner structure
- use the set of matching residues to calculate *rmsd*

# 2. Peptide fragment strategy

- more general version of idea on previous page
- basis of most popular methods

Ingredients

- break protein into overlapping fragments of structure (length 6 or 8)
- protein is no longer a string of residues nor a whole structure



• each fragment is a little distance matrix

## **Fragment Based Comparison**

- any two distance matrices can be compared
- two proteins length *N* and *M* can now be compared...

		1	2	3	4	5			<i>N</i> -7	protein 1
unatain 2	1	1.3	1.0	2.0	0.9					fragments $\rightarrow$
protein 2 fragments J	2	2.7	2.3	0.5						
maginentis	3	5.5	4.4							
	4	0.1	0.5	0.3	3.3	4.2				
	5	1.9	4.4	5.5	0.3	3.3				
	6	4.4	1.6	1.7	5.0	2.3				
		4.1	3.1	3.3	4.4	0.2	3.3	•••		
	<i>M</i> -7	5.2	1.1	0.1	5.5	4.4	0.1	3.3	0.1	

• imagine *rmsd* 

• this is now like a sequence comparison problem

## **Finding Equivalent Fragments**

- find optimal path through matrix
- classic dynamic programming method like sequence comparison

	1	2	3	4	5			N-7
1	1.3	1.0	2.0	0.9				
2	2.7	2.3	0.5					
3	5.5	4.4						
4	0.1	0.5	<u>5.</u> %	3.3	4.2			
5	1.9	4.4	5.5	0.8	3.3			
6	4.4	1.6	1.7	5.0	23			
	4.1	3.1	3.3	4.4	0.8	3.3		
N-7	5.2	1.1	0.1	5.5	4.4	0.1	3.3	0.1

Like sequence comparison

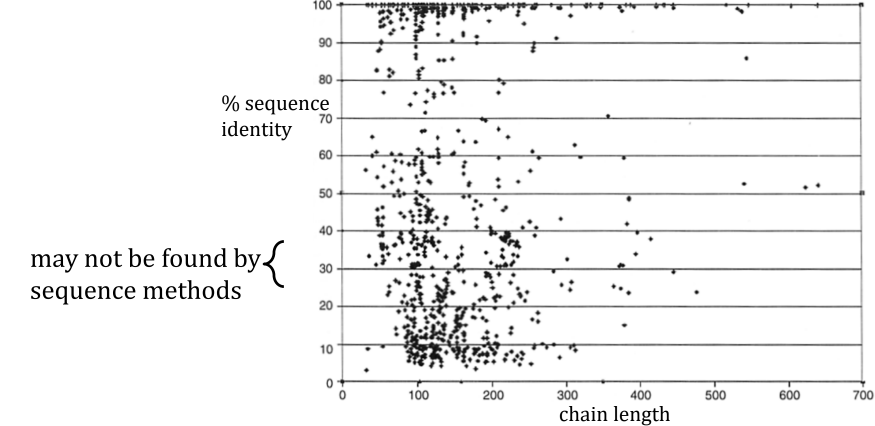
- find optimal path through matrix
- classic dynamic programming method (N & W, S & W)
- uses gap penalties

#### **Comparing Different Size Protein Structures**

- Break protein into overlapping fragments
- fragments can be compared to each other via distance matrices
- align like sequences
- from aligned fragments, get list of aligned residues
- using aligned residues, calculate *rmsd*, *rms* of overall distance matrices

#### **How Important Are These Similarities ?**

- survey 1 000 proteins
- find structurally similar pairs
- plot sequence identity



pictures from "Structural Bioinformatics", ed Bourne, PE and Weissig, H.

#### **Summary of All Protein Comparisons**

Classification of proteins

• could be done by sequence, better by structure

Structure comparison

- for one protein
  - selection of atoms
- for different proteins
  - requires list of matching atoms
- for similar proteins
  - can use pairs from sequence alignment
- for often dissimilar proteins
  - pure structure based method

## **Summary of everything**

- classification is appealing
- very different answers using sequence or structure
- even if we believe in evolution
  - complete hierarchical scheme may be artificial