Multiple sequence alignments similarity without sequence similarity

Andrew Torda, Bioinformatics, Sommersemester 2015

Bis jetzt

- Man hat eine Sequenz (Protein oder Nukleotid)
- Man will so viel wie möglich finden, um
 - Struktur vorherzusagen
 - Funktion vorherzusagen
- Jetzt Alignments, Evolution & Funktion

Multiple alignments

...

...

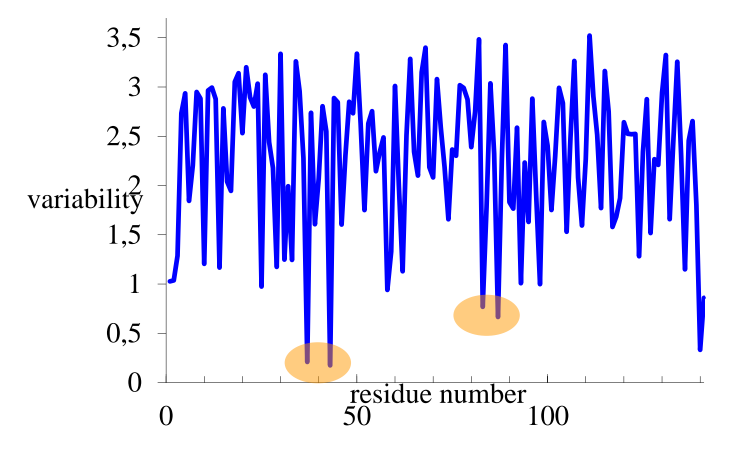
... ...

- what does a set of sequences look like?
- data for a haemoglobin
- summarise this data

VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG $mostly \ for \ proteins^{\text{vlspadktnvkaawgkvgahageygaealermflsfpttktyfphfdlshgsaqvkghg}}_{\text{wlspadktnvkaawgkvgahageygaealermflsfpttktyfphfdlshgsaqvkghg}}$ VLSPADKTNVKAAWGKVGAHAGEYGAEALEKMFLSFPTTKTYFPHFDLSHGSAQVKGHG LSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTNVKAAWGKVGAHAGDYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPDDKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTHVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEAWERMFLSFPTTKTYFPHFDLSHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEAWERMFLSFPTTKTYFPHFDLSHGSAOVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSAADKTNVKAAWSKVGGHAGEYGAEALERMFLGFPTTKTYFPHFDLSHGSAQVKAHG VLSAADKTNVKAAWSKVGGHAGEYGAEALERMFLGFPTTKTYFPHFDLSHGSAOVKAHG VLSADDKANIKAAWGKIGGHGAEYGAEALERMFCSFPTTKTYFPHFDVSHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEAFERMFLSFPTTKTYFPHFDLSHGSAQVKGQG VLSPADKTNVKAAWGKVGAHAGEYGAEAFERMFLSFPTTKTYFPHFDLSHGSAQVKGQA VLSAADKSNVKAAWGKVGGNAGAYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG VLSPADKSNVKATWDKIGSHAGEYGGEALERTFASFPTTKTYFPHFDLSPGSAQVKAHG VLSPADKSNVKAAWGKVGGHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTGTYFPHFDLSHGSAQVKGHG VLSSADKNNVKACWGKIGSHAGEYGAEALERTFCSFPTTKTYFPHFDLSHGSAQVQAHG VLSAADKSNVKAAWGKVGGNAGAYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSANDKSNVKAAWGKVGNHAPEYGAEALERMFLSFPTTKTYFPHFDLSHGSSQVKAHG VLSPADKSNVKAAWGKVGGHAGDYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG

Conservation / variability

Look at residues 37, 43, 83 and 87



- how do we get these and what does it mean?
- what does it mean for this protein ?

26/05/2015 [3]

Conserved residues

Proximity to haem group

• green residues

• more on pro 37 later

Beliefs in multiple sequence alignments

Similar proteins found in many organisms

- where they are conserved connected with function
- variation reflects evolution (phylogeny)

How many homologues might you have ?

- many
 - some DNA replication proteins almost every form of life
 - profilin cell mobility bacteria, mammals, plants
 - .
- few
 - exotic viral proteins
 - messengers exclusively in human biochemistry

Trees / Phylogeny

Multiple sequence alignments are fun

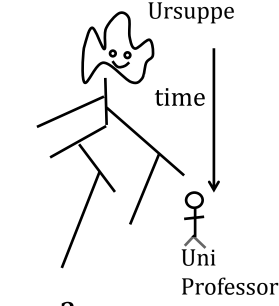
• conservation, function...

What next? Phylogeny - making trees

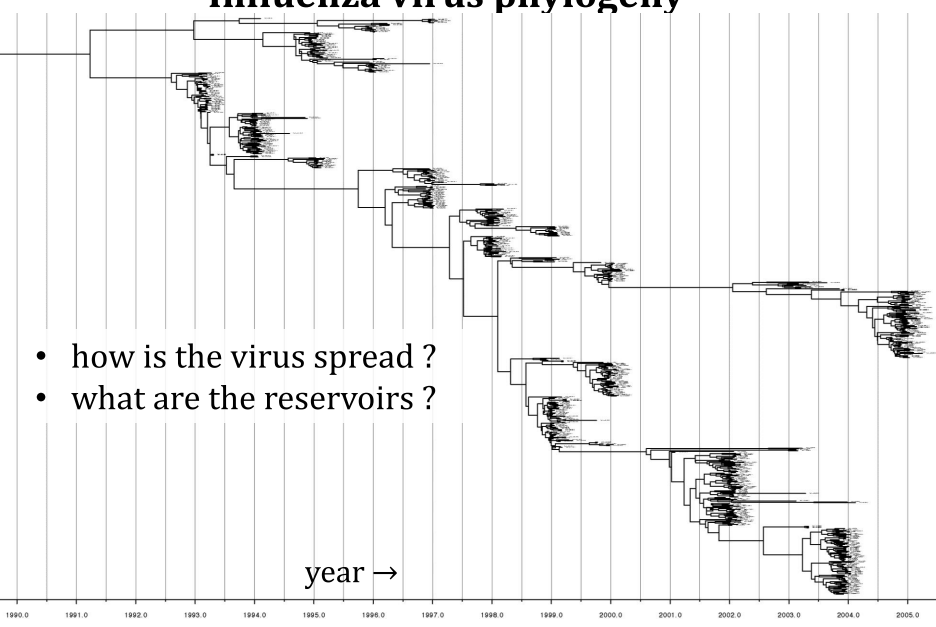
Need multiple sequence alignments to make trees

Do you just want the tree of life ?

- who killed the bananas ?
- who killed the ladies ?
- where does influenza come from ?
- where did the American Indians come from ?



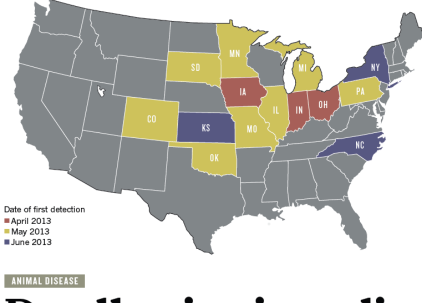
Influenza virus phylogeny



Rambaut, A., .. Holmes, C. The genomic.. influenza A virus, Nature 452, 1-6, 2008

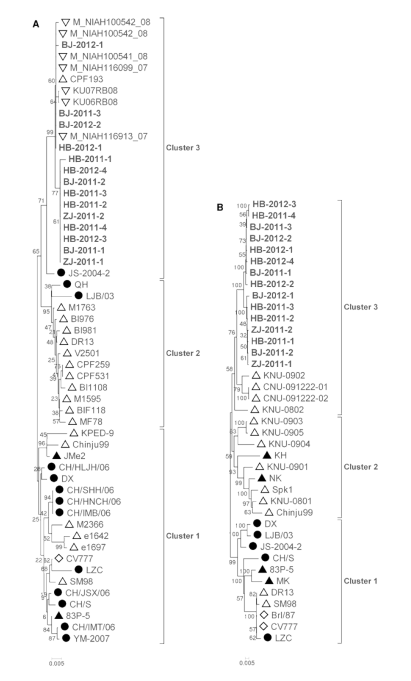
PIG VIRUS ON THE WING

Porcine epidemic diarrhoea virus, a type of coronavirus that can kill piglets, has been detected in 14 US states.



Deadly pig virus slips through US borders

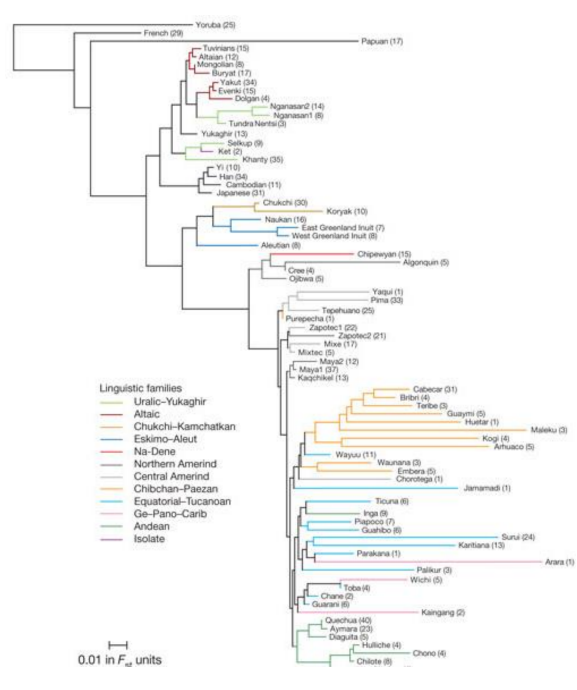
Researchers race to track spread of coronavirus.



Mole, B., Nature, 499, 388 (2013)

Gao, Y. Kou, Q., Ge, X, Zhou, L., Guo, Yang, H., Arch Virol, 158, 711-715 (2012)

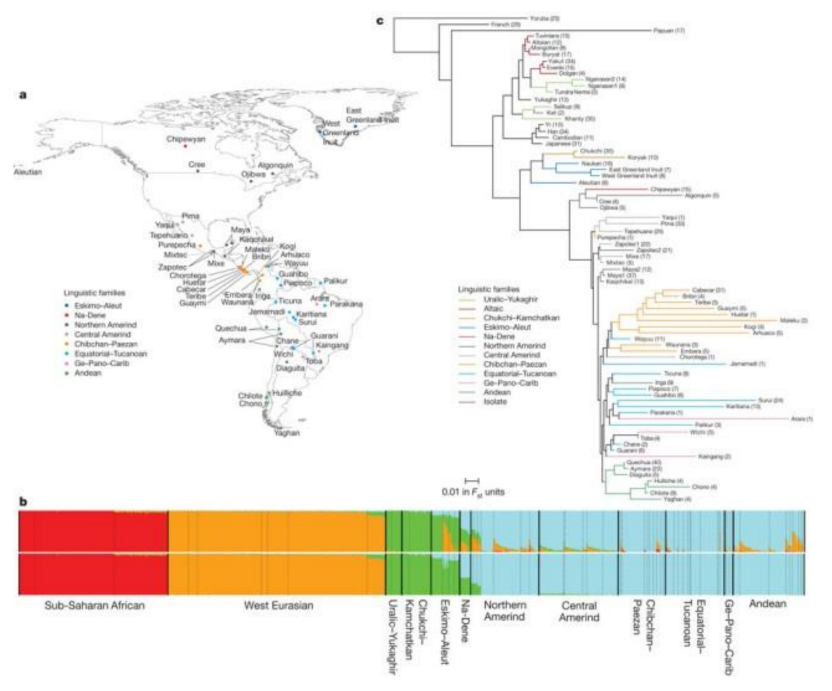
26/05/2015 [8]



52 Native American groups

Reich, D., ...Ruiz-Linares, A., Nature, 488, 370 (2012), Reconstructing Native American .. History

26/05/2015 [9]

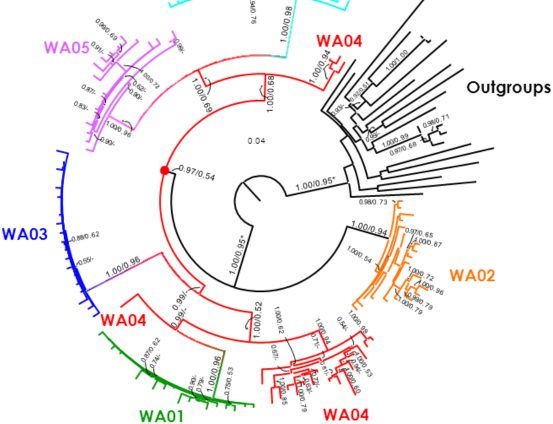


Reich, D., ...Ruiz-Linares, A., Nature, 488, 370 (2012), Reconstructing Native American .. History

1000 acts of sex

"the defendant intended to inflict "great bodily harm"... Between 1999 and 2004, he engaged in more than 1000 oral, vaginal, and anal acts of unprotected sex with his female partners"

Colour changes where WA04 infected ladies



Scaduto, D.I., Brown, JM, Haaland, WC, Zwicki, DJ, Hillis, DM, Metzker, ML, (2010) Proc Natl Acad Sci USA, 107, 21242 26/05/2015 [11]

The plan

• optimise and alignment and tree simultaneously

Many sequences - rigorous alignment

- two sequence alignment
 - optimal path through $n \times m$ matrix
- three sequence alignment
 - optimal path through $n \times m \times p$ matrix
- four sequence alignment
 - ...
- *m* sequence alignment of *n* residues.... $O(n^m)$

Excuse to use lots of approximations

• no guarantee of perfect answer

Reasonable starting point

• begin with pairs of proteins

Scoring schemes

$$S_{a,b} = \sum_{i=1}^{N_{res}} \operatorname{match}(s_{a,i}, s_{b,i})$$

In pairwise problem

VLSPADKSNVKAGWGQVGAHAGDYGAEAIERMYLSFPSTKTYFPHTDISHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG

- Sum over match()
 N_{res} is sequence length
- match(s_{a,i}, s_{b,i}) is the match/mismatch score of sequence a and b at position i
- invent a distance between two sequences like

$$d_{a,b} = \frac{1}{S_{a,b}}$$

distance measure..
 which sequences are most dissimilar to each other

Scoring schemes for a multiple alignment

In the best alignment

- 1 is aligned to 2, 3, ..
- 2 to 3,4, ...

VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG
 VITP-EQSNVKAAWGKVGAHAGEYGAEALEQMFLSYPTTKTYFP-FDLSHGSAQIKGHG
 MLSPGDKTQVQAGFGRVGAHAG--GAEALDRMFLSFPTTKSFFPYFELTHGSAQVKGHG
 VLSPAEKTNIKAAWGKVGAHAGEYGAEALEKMF-SYPSTKTYFPHFDISHATAQ-KGHG
 -VTPGDKTNLQAGW-KIGAHAGEYGAEALDRMFLSFPTTK-YFPHYNLSHGSAQVKGHG
 VLSPAEKTNVKAAWGRVGAHAGDYGAEALERMFLSFPSTQTYFPHFDLS-GSAQVQAHA
 VLSPDDKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG

• then I move 5 and 2 & 5 and 3 – messes up 2 and 3

Mission: for N_{seq} sequences

• S_{ab} : alignment score sequences *a* and *b*

$$score = \sum_{b \neq a}^{N_{seq}} \sum_{a=1}^{N_{seq}} S_{a,b}$$

- not quite possible
 - this method is just an approximation

Aligning average sequences

VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VITPAEKTNVKAAWGKVGAHAGEYGAEALEQMFLSYPTTKTYFPHFDLSHGSAQIKGHG

and

IITPGDKTNVKAAFGKVGAHGGEYGAEALDRMFISFPSTKTYYPHFDLSHASAQVKAHG VITPAEQTNIKGAWGQIGAHAGDYAADALEQMFLSYPTSKTYFPYFDLTHGSAQIKGHG VITPAEKTQVKAAWGKVGGHAGEYGAEAIEQMFLTYPTTQTYFPHFELSHGTAQIKGHG

- At each position
- use some kind of average in scoring
- if a column has $2 \times D$ and $1 \times E$ score
 - score as 2/3 D + 1/3 E
- later.. call the average of S1 and S2: av(S1, S2)

Summarise ingredients

- pairwise scores + distances
- ability to align little groups of sequences

Progressive alignments

Guide tree / progressive / neighbour joining method

Steps

- build a distance matrix
- build a guide tree
- build up overall alignment in pieces

Progressive alignment - tree

Compute pairwise

- S1 ATCTCGAGA
- S2 ATCCGAGA
- S3 ATGTCGACGA
- S4 ATGTCGACAGA
- S5 ATTCAACGA

alignments, S1 calculate the S2 .11 distance matrix S3 .20 .30 .27 S4.36 .09 .30 .33 S5 .23 .27 S2 S1 S3 S4 S5 S1 calculate guide tree S2 S3 S4 S5

Multiple alignment from guide tree

- gaps at early stages remain
- problems..
- S1/S2 and S3/S4 good
 - no guarantee of S1/S4 or S2/S3

 av(S1,S2) is average of S1 and S2

align S1 with S2

S1	ATCTCGAGA
S2	ATC-CGAGA

align S3 with S4 S3 ATGTCGAC-GA S4 ATGTCGACAGA

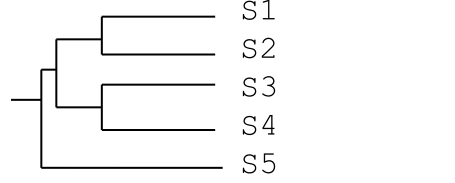
align av(S1,S2) with av(S3,S4)

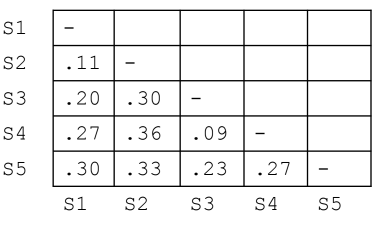
S1	ATCTCGAGA
S2	ATC-CGAGA
S3	ATGTCGAC-GA
S4	ATGTCGACAGA

align av(S1,S2,S3,S4) with S5

- S1 ATCTCGA--GA
- S2 ATC-CGA--GA
- S3 ATGTCGAC-GA
- S4 ATGTCGACAGA
- S5 AT-TCAAC-GA

Problems and variations

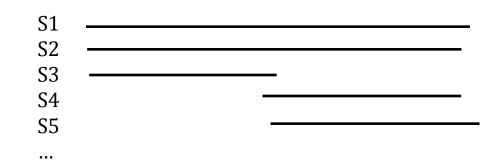




What order should we join ?

- pairs are easy (S1+S2) and (S3+S4)
- which next?

Real breakdown



- S1 and S2 are multi-domain proteins
 - S3 is not really related to S4 or S5
 - distance matrix elements are rubbish

Given an alignment

How reliable / believable ?

- set of very related proteins (an enzyme from 100 mammals)
 - no problem
- diverse proteins (an enzyme from bacteria to man)
 - lots of little errors
- can break completely (domain example)

Is the tree a "phylogeny" ? A reflection of evolution ?

• more later

Measuring conservation / entropy

Entropy

- how much disorder do I have ? $S = -k \sum_{i=1}^{N_{states}} p_i \ln p_i$
- in how many states may I find the system? Our question
- look at a column how much disorder is there ?

VLSPADKTNVKAAWGKVC AHAGEYGAEALERMFLSFPTTKTYFPHE DI.SHGSAQVKGHG VITP-EQSNVKAAWGKVC AHAGEYGAEAIEQMFLSYPTTKTYFP-E DI.SHGSAQIKGHG MLSPGDKTQVQAGFGRVC AHAG--GAEAVDRMFLSFPTTKSFFPYE EI.THGSAQVKGHG VLSPAEKTNIKAAWGKVC AHAGEYGAEAAEKMF-SYPSTKTYFPHE DI.SHATAQ-KGHG -VTPGDKTNLQAGW-KIC AHAGEYGAEALDRMFLSFPTTK-YFPHYNI.SHGSAQVKGHG VLSPAEKTNVKAAWGRVC AHAGDYGAEAGERMFLSFPSTQTYFPHE DI.S-GSAQVQAHA VLSPDDKTNVKAAWGKVC AHAGEYGAEALERMFLSFPTTKTYFPHE DI.SHGSAQVKGHG

> no disorder

much disorder

Calculate an "entropy" for each column

Entropy

- forget k (Boltzmann just scaling) $S = -\sum_{i=1}^{N_{states}} p_i \ln p_i$ We have a protein
- 20 possible states

What if a residue is always conserved ? $p_i = 1$ or $p_i = 0$

 $S = \ln 1 = 0$ (no entropy)

What if all residues are equally likely ? $p_i = \frac{1}{20}$

$$S = -\sum_{i=1}^{20} \frac{1}{20} \ln \frac{1}{20} = -20 \cdot \frac{1}{20} \ln \frac{1}{20}$$

≈ 3

• my toy alignment...

Entropy

- First column is boring

- Second

$$p_{\rm D} = \frac{5}{7}$$

 $p_{\rm E} = \frac{1}{7}$

$$p_{\rm N} = {}^1\!/_7$$

VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDISHGSAQVKGHG VITP-EQSNVKAAWGKVGAHAGEYGAEAIEQMFLSYPTTKTYFP-FDISHGSAQIKGHG MLSPGDKTQVQAGFGRVGAHAG--GAEAVDRMFLSFPTTKSFFPYFEITHGSAQVKGHG VLSPAEKTNIKAAWGKVGAHAGEYGAEAAEKMF-SYPSTKTYFPHFDISHATAQ-KGHG -VTPGDKTNLQAGW-KIGAHAGEYGAEALDRMFLSFPTTK-YFPHYNISHGSAQVKGHG VLSPAEKTNVKAAWGRVGAHAGDYGAEAGERMFLSFPSTQTYFPHFDIS-GSAQVQAHA VLSPDDKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDISHGSAQVKGHG

$$S = -\left(\frac{5}{7}\ln\frac{5}{7} + \frac{1}{7}\ln\frac{1}{7} + \frac{1}{7}\ln\frac{1}{7}\right)$$

\$\approx 0.8\$

Entropy from DNA

Exactly as for proteins Will numbers be larger or smaller ?

max possible entropy

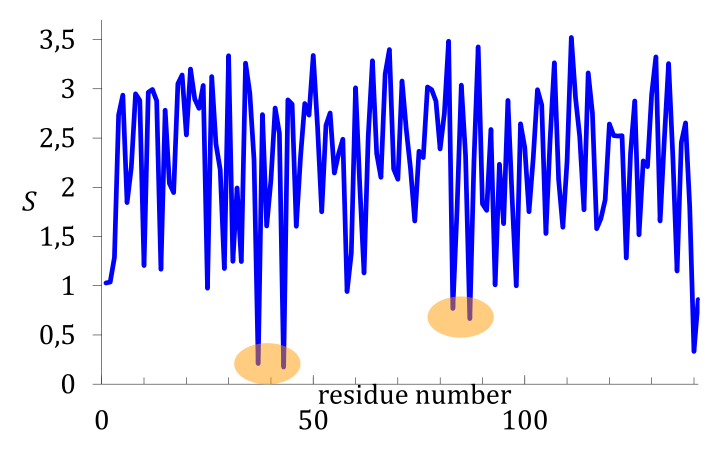
$$S = -4\left(\frac{1}{4}\ln\frac{1}{4}\right)$$
$$= -\ln\frac{1}{4}$$

≈ 1.4

example from start of this topic

Haemoglobin conservation

Look at residues 37, 43, 83 and 87



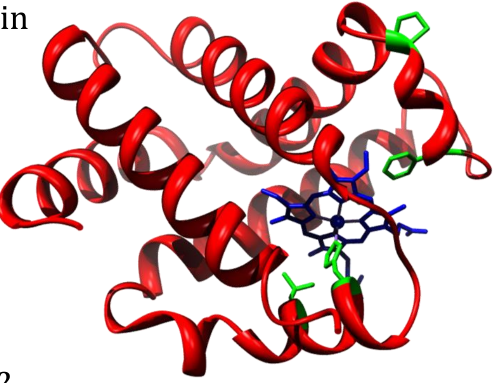
4 residues (maybe more) stand out as conserved

• why ?

26/05/2015 [26]

Conserved residues in haemoglobin

- 3 of the sites are easy to explain
- interact with haem group
- Look at fourth site
- proline
- end of a helix



What is special about proline ?

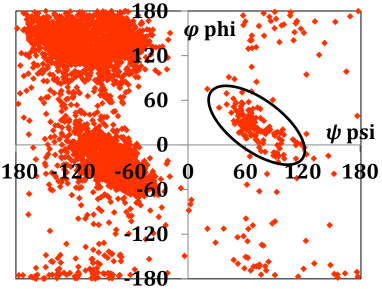
- no H-bond donor
- here if it mutates, maybe haemoglobin does not fold

Conservation for structure

Some residues have very special structural roles

- proline not an H-bond donor
 - often end of a helix
- glycine can visit part of $\varphi \psi$ plot
 - found in some turns
- Are all gly residues so important?
- NO they occur in many places sometimes in turns

Are all pro residues very conserved ? No



Conservation for function

In a serine protease

- always a "catalytic serine"
- can it mutate ? Not often

In haemoglobin – residues necessary for binding haem

- can they mutate ? rarely
- changes properties of haemoglobin (bad news)

Dogma

residues in active site will be more conserved than other sites

Important summary

Conservation may reflect

- important function
- structural role
- Mutagenesis / chemistry
- what residue may I change to allow binding to a solid substrate ? (for biosensor/immobilized enzyme ?)
- try error prone PCR to select for new enzyme activity which sites might I start with (active site) ?
- Drug design example
- target is an essential protein (basic metabolism, DNA synthesis, protein synthesis..)
- is there some set of sequence features common to pathogen, different to mammalian protein ?

Evolution – do not trust conservation

Imagine: two possible systems for some important enzyme

- 1. active site fits to essential biochemistry

 - any mutation you lose active site residues are conserved in a conservation plot
- 2. maybe enzyme is not absolutely perfect
 - some mutations kill you
 - some mutations OK lacksquare
 - site does not appear perfectly conserved •

Where would you evolve to?

- 1. very fragile
- 2. likely to survive mutations

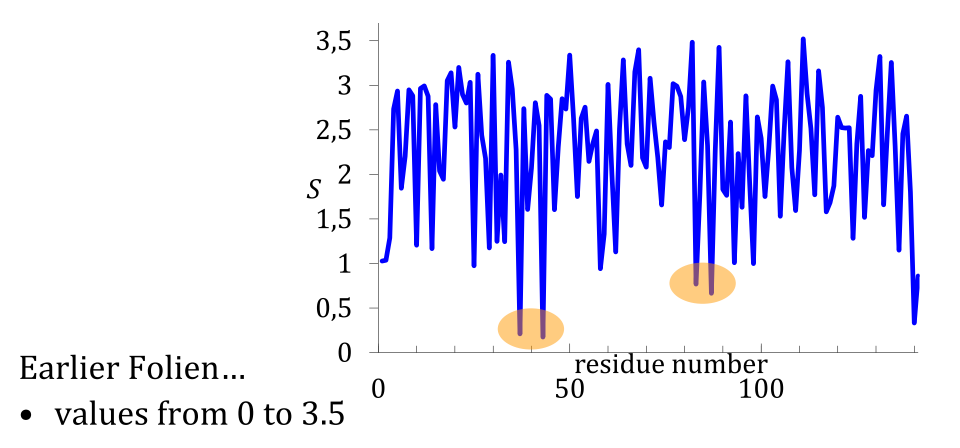
Resistance to mutations...

Resistance to Mutations

Why might one tolerate mutations?

- boring answer
 - some amino acids are similar to each other
- better answer
 - it will be selected for
 - you genes have a better chance of being passed on
 - it is a Darwinian trait

Conservation – how meaningful ?



What if I used more homologues ?

Conservation – how meaningful ?

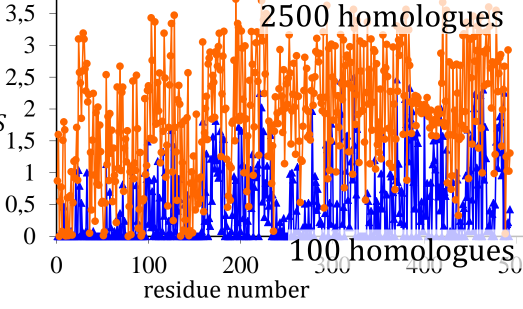
Example sequence (1ab4, DNA gyrase)

- find 100 close homologues (mostly > 80% similarity)
 calculate conservation
- find 2500 close homologues (mostly > 50 % similarity) calculate conservation

S

Fewer sequences

- lots of conserved sites
- you can get the answer $\begin{bmatrix} 1\\0,5\\0\end{bmatrix}$ you want 0



Consequences - summarise

Significance of conservation

You read in a paper – residue 37 is conserved

- how many sequences did they look at ?
 - 10 ? bad 100 better 1000 better
- choosing the number of sequences lets you manipulate results
- statistically
 - have you sampled over enough sequences ?

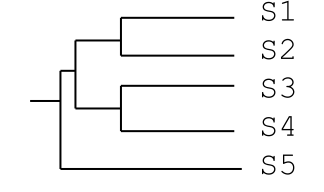
Phylogeny / Evolution

The trees in text books are almost never perfect One rarely knows the correct history

Problems..

Previously we had a "guide tree"

did (S1,S2) and (S3,S4) share an ancestor but not S5 ?



- branch lengths do not reflect evolutionary time
- there may be other similar trees which could be evolutionary paths

Evolutionary time

Compare two DNA sequences see 1 mutation (represents time *t*) 2 mutations (time 2*t*) 3 mutations (time 3*t*)... No !

After some evolution

 $A \rightarrow C \rightarrow G$ two events (although looks like $A \rightarrow G$) $A \rightarrow C \rightarrow G \rightarrow C \rightarrow A$ looks like zero mutations

If I have infinite time

- all bases / residues equally likely
- $p_{mut} = 3/4 = 0.75$ (DNA) or $p_{mut} = 19/20$ (protein)

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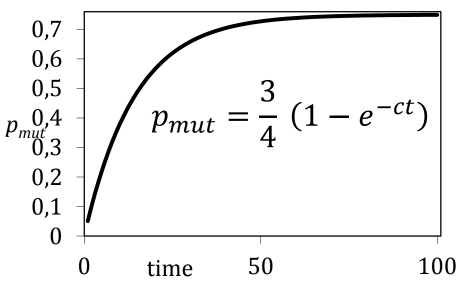
Mutation probability

- time units are arbitrary
- how would I estimate time ? (for DNA)

•
$$t \propto -\ln\left(1-\frac{4}{3}p_{mut}\right)$$

• p_{mut} ? count $\frac{n_{mut}}{n_{res}}$



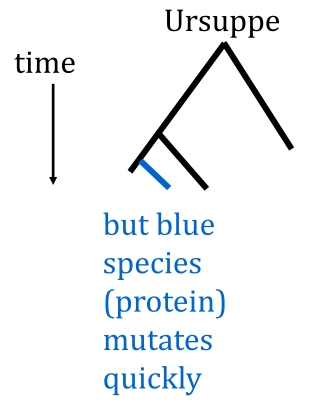


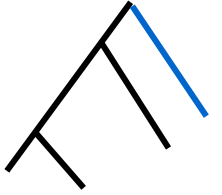
For short times, p_{mut} changes fast

- for small *t*, distances will be more reliable
 - as will be alignments
- Is this enough for phylogeny ?
- what about reliability ?

Problems in phylogeny

- not all sites mutate equally quickly
- not all species mutate equally quickly





blue appears to have branched off earlier

- Backwards mutations?
- not really a problem (Klausurerfahrung)

Problems estimating time

- 1. mutation rates vary wildly
 - changing environments pH, temperature,..
- 2. imagine time *t* is such that p_{mut} = 0.25
 - we have random events
 - sometimes you see 23% mutation, sometimes 28%
- time estimates will never be accurate
- maybe we cannot find the correct tree
 - can we roughly estimate reliability?

Reliability

Think of first alignment

VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VITP-EQSNVKAAWGKVGAHAGEYGAEAIEQMFLSYPTTKTYFP-FDLSHGSAQIKGHG MLSPGDKTQVQAGFGRVGAHAG--GAEAVDRMFLSFPTTKSFFPYFELTHGSAQVKGHG VLSPAEKTNIKAAWGKVGAHAGEYGAEAAEKMF-SYPSTKTYFPHFDISHATAQ-KGHG -VTPGDKTNLQAGW-KIGAHAGEYGAEALDRMFLSFPTTK-YFPHYNLSHGSAQVKGHG VLSPAEKTNVKAAWGRVGAHAGDYGAEAGERMFLSFPSTQTYFPHFDLS-GSAQVQAHA VLSPDDKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG

What would happen if you deleted a column ?

- if the data is robust /reliable
 - not much
- if the tree is very fragile /sensitive
 - tree will change

better...

Reliability

Repeat 10^2 to 10^3 times

- delete 5 to 10 % of columns
- copy random columns so as to have original size
- recalculate tree

How often did you see each branch?



Monster example

- generate 1000 trees
- for each sub-tree
 - see how often it is present
- example from cover of nature

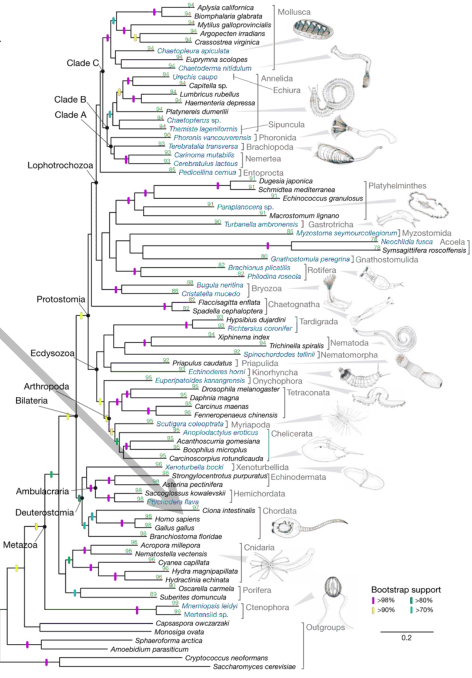
Monster calculation

How long did this take?

months on 120 processors

We are

- usually placed near Hühne
- not so reliably placed with little worms



DNA or protein sequences ?

The issues

- regulatory regions, RNA genes
- synonymous mutations (common only seen in DNA)
- non-synonymous mutations (amino acid changes)
 - more information $D \rightleftharpoons E$, $I \rightleftharpoons L \rightleftharpoons V$, ...

Alignment reliability

- proteins
 - uses codon structure (implicitly)
 - better, amino acid similarity, $I \rightleftharpoons L \rightleftharpoons V$ is not bad
- DNA
 - less information

DNA or protein sequences ?

	protein	DNA	time
synonymous changes	no	yes	short
a.a. changes	yes	no	longer
frame shifts	no	yes	
non-coding regions	no	yes	

Very short time

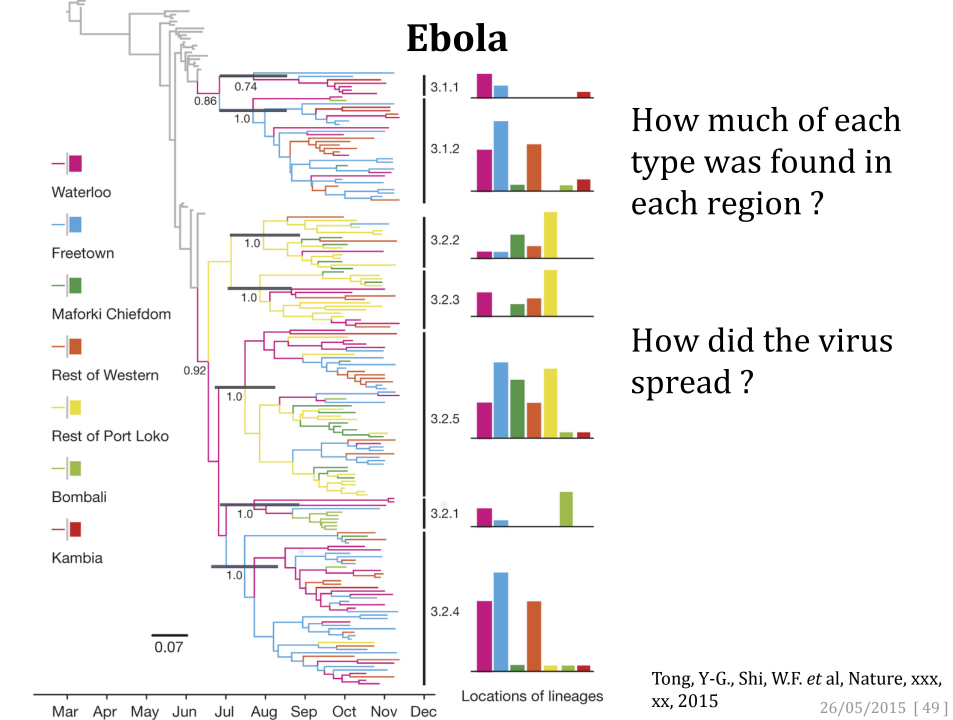
• use DNA

Longer time

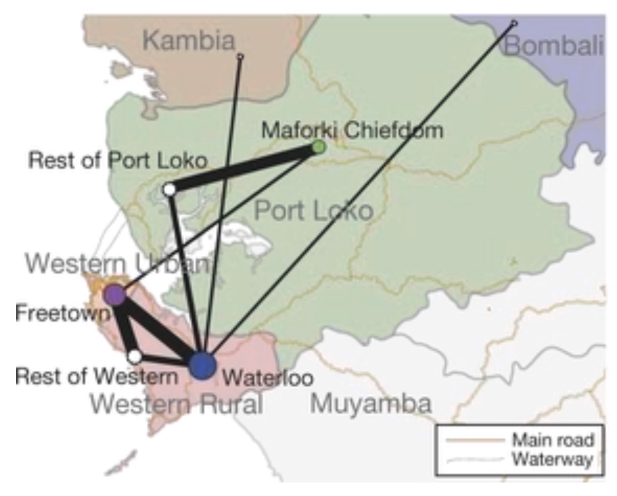
• use proteins

Summary

- multiple sequence alignment conservation
 - find important residues (function or structure)
 - can quantify conservation
- relations between most similar proteins are most reliable
- best tree is never found
 - too difficult algorithmically
 - lots of errors evolution is a random process
- rough idea of reliability
- quick tree possible for 1000s of sequences
- more complicated methods Frau Dobler's Vorlesungen



How did the virus spread?



Thickness of lines – closeness of sequences

Protein structures and comparisons

Ultimate aim

- how to find out the most about a protein
- what you can get from sequence and structure information

On the way..

- remote similarities between proteins
- sequence versus structural similarity
- Detour
 - protein coordinates representation, accuracy
- measures for similarity of coordinates

Sequence and structure similarity

Claim from before

- if two sequences are similar
 - they are related structures are similar

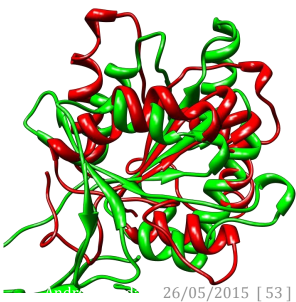
Question

- if two sequences are different
 - are their structures different ?

Remote similarities

1cbl & 1eca (haemoglobin & erythrocruorin) 14 % sequence id

> 1fyv & 1udx, TLR receptor and nucleotide binder, 9 % sequence id



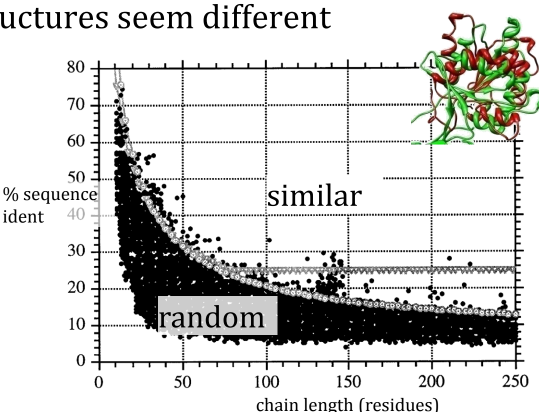
No sequence similarity – similar structures

Are these rare ?

- easy to find 100s of examples Does this agree with previous claims ?
- dot in diagram two structures seem different

If sequences are similar

- structures will be similar
- If sequences are different
- one does not know



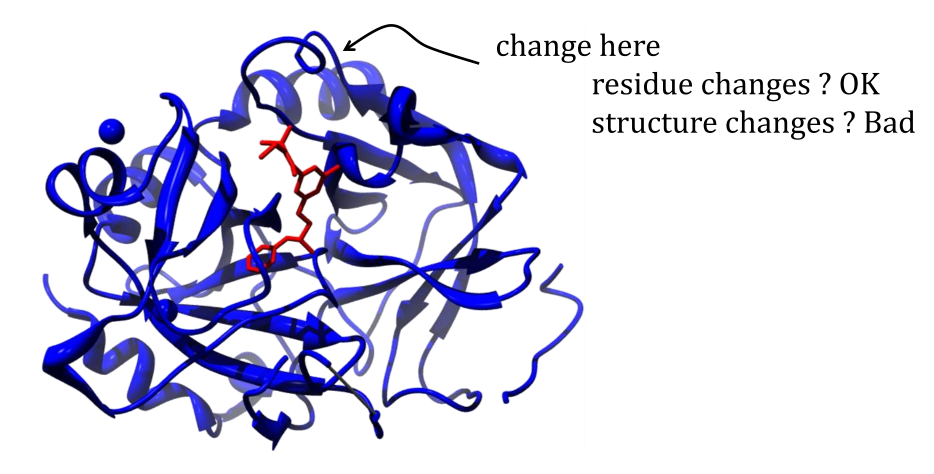
Structure versus sequence similarity

Clear statement

- sequence changes faster than structure
 Reason ? Unclear
- possibility..
- protein function depends on having groups in orientation in space

Why can sequence change

View of molecular evolution...



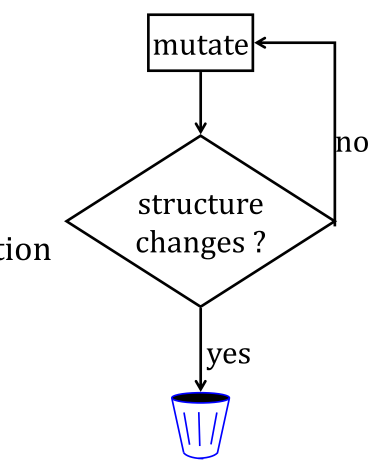
Simple view of molecular evolution

mutate continuously

- mutations which are not lethal
 - may be passed on (fixed)
- if structure changes
 - protein probably will not function
 - not passed on

Result

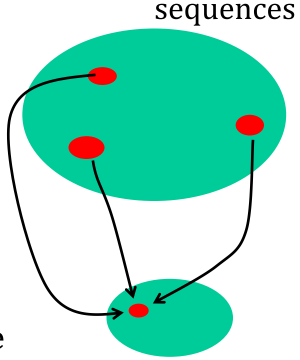
- evolution will find many sequences
 - compatible with structure
 - compatible with function
- how else would we see this ?



Sequence vs structure evolution

Sequence and structure space

- sequence space is larger
 - many different sequences map to similar structure
- sequence evolves faster than structure



structures

Practical Consequences

Sequences of proteins are nearly always known

Similar sequence

• usually similar structure, similar function

Sequences not (obviously) related

- maybe similar structure
- maybe similar function

Sequence vs structure similarity

When comparing proteins

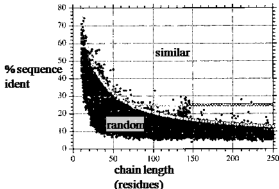
Similar sequences

- structure and function will be similar
 - remember threshold graphs from earlier

Similar structures, different sequences

- evolutionary relationship implied but
 - bigger evolutionary distance
- not enough to be confident about function
- what do we mean by similar structures ?
 - coming soon

practical consequences ...



Little summary

Multiple sequence alignments

- for conservation
- first step to phylogenies

Phylogenies

• not as reliable as the pictures imply

Structure vs sequence evolution

- sequence changes faster
- sequence similarity means a closer evolutionary relationship
 - functional similarity

Comparing structures

- what are protein coordinates ?
- comparison

Representation

- Proteins are not as smooth as we draw them
 - very discrete set of atoms

Protein coordinate files

Detour - Protein data bank (www.rcsb.org)

- only significant database of protein coordinates
- deposition of coordinates often requirement of publication
- $\approx 10^5$ structures
 - huge redundancy (> 500 T4 lysozyme)
- X-ray crystallography $\approx 85 \%$
- NMR ≈ 14 % (more in smaller proteins)

Protein Databank – biased ?

10⁵ structures – a good summary of the world ? Maybe not

- 1. Chemistry we see proteins
- that could be crystallised
- that can be expressed
- protein 2 is easier if protein 1 has a known structure
- 2. Sociology
- human proteins
- "model organisms"
- disease-causing proteins / therapeutic targets

the bias

Lots of proteins that are

- smaller
- soluble / globular (few membrane bound)
- stable / can be handled in laboratory
- not toxic can be expressed
- similar to ones in the databank
- from humans and special projects

Comparing coordinates

These are very similar



we want to put numbers on this property

First some notation

- We have spoken of *x*, *y*, *z* coordinates. Easier..
 - vector \vec{r} or $\vec{r_i}$ for atom *i*,
 - for two proteins let us have position *i* in protein *a* and *b*
 - \vec{r}_i^a and \vec{r}_i^b

Comparing two proteins

- take one atom (C^{α}) from residue *i*
- what do I know from the picture ?
- if my two proteins are similar $\vec{r}_i^a \vec{r}_i^b$ will be a short vector
- for each residue *i*
- define $|\vec{r}_i^a \vec{r}_i^b|$ distance between \vec{r}_i^a and \vec{r}_i^b

I want a single number that tells me

- how close is a residue in *a* to the corresponding residue in *b*
- think of the set of distances $|\vec{r}_i^a \vec{r}_i^b|$
- how spread out is this population of distances ?
 - like a standard deviation (standard Abweichung)

Root mean square (rms)

Normal formula for standard deviation

$$\sigma_{\chi} = \left(\frac{1}{N}\sum_{i=1}^{N}(x_i - \bar{x})^2\right)^{\frac{1}{2}}$$

Something similar for coordinates

$$r_{rmsd} = \left(\frac{1}{N_{res}} \sum_{i=1}^{N_{res}} |\vec{r}_i^a - \vec{r}_i^b|^2\right)^{\frac{1}{2}}$$

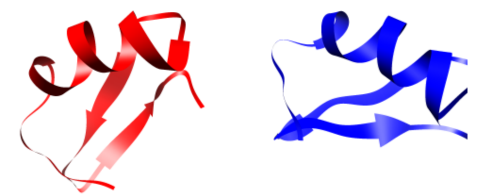
1

- where proteins *a* and *b* have *N*_{res} residues
- rmsd is "root mean square difference"
- complications

Before calculating rmsd

Two very similar proteins

- coordinates are in different orientations
- not on top of each other



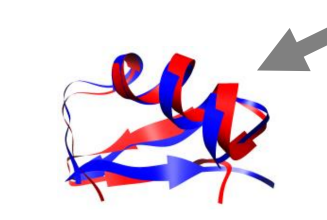
What are the orientations of files in PDB?

• totally arbitrary

Consequence

• put the proteins on top of each other

Superposition of coordinates



now use formula for *rmsd*

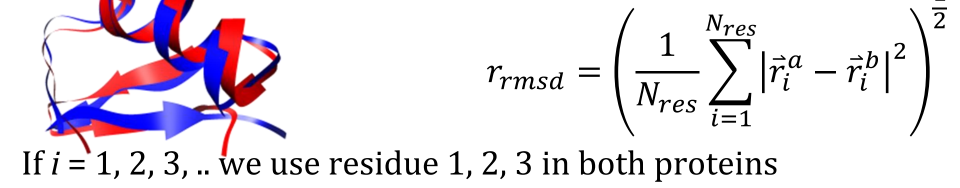
Meaning of *rmsd*

Before calculating *rmsd*

- units Å
- rmsd is size dependent
 - 5 Å in a small protein (50 residues) will not look similar
 - 5 Å in a big protein (250 residues) will look similar

Difficulty with *rmsd*

These two proteins have the same number of residues

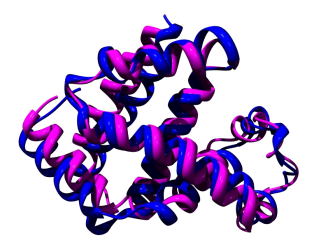


These two proteins have different numbers of residues

we cannot compare residue 1 to 1, 2 to 2..

rmsd different sized proteins

- make a list of residues in each protein
- just work with corresponding residues (amazingly difficult)



rmsd summary

- formula bit like standard deviation
- needs translation and rotation
- size dependent (just remember)
- difficult for proteins of different sizes

Summary of comparisons

Closely related proteins

- sequence alignments
- multiple sequence alignments
 - conservation, phylogeny

Structures

- see similarity even when sequences are very different
- necessary for drug design / Wirkstoffentwurf
- explaining patterns of conservation
- predictions of which residues are near/far from active site