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# Multiple alignments

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... ...

- mostly for proteins
- what does a set of sequences look like ?

- data for a haemoglobin
- summarise this data

VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALEKMFLSFPTTKTYFPHFDLSHGSAQVKGHG LSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTNVKAAWGKVGAHAGDYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG VLSPDDKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTHVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEAWERMFLSFPTTKTYFPHFDLSHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEAWERMFLSFPTTKTYFPHFDLSHGSAOVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSAADKTNVKAAWSKVGGHAGEYGAEALERMFLGFPTTKTYFPHFDLSHGSAQVKAHG VLSAADKTNVKAAWSKVGGHAGEYGAEALERMFLGFPTTKTYFPHFDLSHGSAOVKAHG VLSADDKANIKAAWGKIGGHGAEYGAEALERMFCSFPTTKTYFPHFDVSHGSAOVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEAFERMFLSFPTTKTYFPHFDLSHGSAQVKGOG VLSPADKTNVKAAWGKVGAHAGEYGAEAFERMFLSFPTTKTYFPHFDLSHGSAOVKGOA VLSAADKSNVKAAWGKVGGNAGAYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKSNVKATWDKIGSHAGEYGGEALERTFASFPTTKTYFPHFDLSPGSAOVKAHG VLSPADKSNVKAAWGKVGGHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTGTYFPHFDLSHGSAOVKGHG VLSSADKNNVKACWGKIGSHAGEYGAEALERTFCSFPTTKTYFPHFDLSHGSAOVOAHG VLSAADKSNVKAAWGKVGGNAGAYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAOVKGHG VLSANDKSNVKAAWGKVGNHAPEYGAEALERMFLSFPTTKTYFPHFDLSHGSSOVKAHG 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 ?

#### conserved residues

- proximity to haem group •
  - green residues



# **Beliefs in multiple sequence alignments**

- Most proteins found in many organisms
- rarely identical
- where they vary will be connected with function
- how much they vary will reflect evolution (phylogeny)
- How many homologues might you have ?
- many
  - some DNA replication proteins almost every form of life
  - some glycolysis proteins from bacteria to man
  - ...
- few
  - some exotic viral proteins
  - some messenger exclusively in human biochemistry

### 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

. . .

- excuse to use lots of approximations
  - no guarantee of perfect answer
- reasonable starting point
  - begin with pairs of proteins

## **Scoring schemes**

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

• In pairwise problem

VLSPADKSNVKAGWGQVGAHAGDYGAEAIERMYLSFPSTKTYFPHTDISHGSAQVKGHG MLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG

- Sum over where  $N_{res}$  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

$$d_{a,b} = 1 - \frac{S_{a,b}}{100 \times N_{res}}$$
 or  $d_{a,b} = \frac{1}{S_{a,b}}$ 

• distance measure – mainly to see which sequences are most similar 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

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

 $b \neq a a = 1$ 

- Mission: for  $N_{seq}$  sequences
- $S_{ab}$ : alignment score sequences *a* and *b*
- not quite possible
  - if I move sequences 4 and 5, may make a mess of 5 and 2

# 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 D (cheating but fast)
    - 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**

- known as guide tree / progressive 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

S1 S2 .11 \_ S3 .20 .30 S4 .27 .36 .09 ----.33 S5 .30 .23 .27 S2 S3 S1 S4 S5

calculate guide tree

alignments,

calculate the

distance matrix



#### multiple alignment from guide tree

alıgn S1	with S2
<b>S</b> 1	ATCTCGAGA
S2	ATC-CGAGA

align Sa	3 with S4
<b>S</b> 3	ATGTCGAC-GA
S4	ATGTCGACAGA

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

<b>S</b> 2	ATC-CGA-	-GA

- 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

• av(S1,S2) is average of S1 and S2

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

### **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 100 organisms, bacteria to man)
  - maybe lots of little errors
- can break completely (domain example)

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

• more later

## **Measuring conservation / entropy**

• Gibbs entropy

$$S = -k \sum_{i=1}^{N_{states}} p_i \ln p_i$$

- how much disorder do I have ?
- in how many states may I find the system ?
- Our question
  - look at a column how much disorder is there ?

VLSPADKTNVKAAWGKVGAFAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VITP-EQSNVKAAWGKVGAFAGEYGAEAIEQMFLSYPTTKTYFPHFDLSHGSAQIKGHG MLSPGDKTQVQAGFGRVGAFAGEYGAEAAEXMF-SYPSTKTYFPHFDLSHGSAQVKGHG VLSPAEKTNIKAAWGKVGAFAGEYGAEAAEXMF-SYPSTKTYFPHFDLSHGSAQVKGHG VLSPAEKTNVKAAWGRVGAFAGEYGAEAAEXMFLSFPTTK-YFPHFDLSHGSAQVKGHG VLSPAEKTNVKAAWGRVGAFAGEYGAEAAEXMFLSFPSTQTYFPHFDLSGSAQVKGHG

> no disorder

much disorder

• Calculate and "entropy" for each column

# **Entropy**

- We can 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 ?
  - $S = \ln(1) = 0$  (no entropy)
- What if all residues are equally likely?

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

• my toy alignment..

# Entropy

- first column is boring
- second
  - $p_{\rm D} = 5/7$
  - $p_{\rm E} = 1/7$
  - $p_{\rm N} = 1/7$

VLSPADKTNVKAAWGKVGAFAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG VITP-EQSNVKAAWGKVGAFAGEYGAEAIEQMFLSYPTTKTYFP-FDLSHGSAQIKGHG MLSPGDKTQVQAGFGRVGAFAGE-GAEAVDRMFLSFPTTKSFFPYFELTHGSAQVKGHG VLSPAEKTNIKAAWGKVGAFAGEYGAEAAEKMF-SYPSTKTYFPHFDLSHGSAQVKGHG VLSPAEKTNVKAAWGRVGAFAGEYGAEALDRMFLSFPTTK-YFPHYDLSHGSAQVKGHG VLSPAEKTNVKAAWGRVGAFAGEYGAEAAERMFLSFPSTQTYFPHFDLS-GSAQVQAHA

$$S = -\left(\frac{5}{7}\ln\frac{5}{7} + \frac{1}{7}\ln\frac{1}{7} + \frac{1}{7}\ln\frac{1}{7}\right)$$
  
\$\approx 0.8\$

• 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 ?

## **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 Hbond 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  $\phi \psi$  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



φ phi

### 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 ? sometimes
  - 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 ?)
  - I want to 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
  - you see active site residues as conserved in a conservation plot
- 2. maybe enzyme is not absolutely perfect
  - some mutations kill you
  - some mutations OK
  - site does not appear perfectly conserved

If you have the choice, where would you evolve to ?

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

### 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
- 2500 homologues 3.5 fewer sequences 3 • lots of conserved sites 2.5 2 S 1.5 you can get the answer you want 0.5 00 homologues 100 200 residue number 300 400500 0

# **Phylogeny / Evolution**

Purely academic ? For fun ? Not always

- possibly useful in explaining disease propagation
  - where did HIV come from ?
  - where did the flu pandemics come from ?
  - virus infects banana crop where did it come from ?
- previously we had a "guide tree"
  - did (S1,S2) and (S3,S4) share an ancestor but not S5 ?
  - not so good



- 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  $p_{mut} = 19/20$

## mutation probability

- time units are rather arbitrary
- how would I estimate time?

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



- p<sub>mut</sub> ? count n<sub>mut</sub> / n<sub>res</sub>
   scaling of t not so important (relative time)
- 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
- less drastic..

# problems in trees

- blue evolves a bit faster
- when we make average sequences
  - $av(S1, S_{blue})$  and sub-tree seems further from other sequences
  - all nearby nodes will be distorted



# **Problems estimating time**

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

# Reliability

- think of first alignment
- what would happen if you deleted a column ?

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

- if the data is robust /reliable
  - not much
- if the tree is very fragile /sensitive
  - tree will change
- better
  - repeat  $10^2$  to  $10^3$  times
    - delete 5 to 10 % of columns
    - copy random columns so as to have original size
    - recalculate tree

#### Monster example

- generate lots of trees
- for each subtree
  - see how often it is is present
- example from cover of nature

# **Monster calculation**

- we are usually placed near Hühne
- we are not so reliably placed with little worms
- how long does this take?
  - months on 120 processors
- a more applied example..





#### influenza virus phylogeny



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

# 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 hundreds of sequences
- more complicated methods only practical for smaller numbers of sequences