Example questions for Bioinformatics, first semester half

Sommersemester 2011

Note

- Die schriftliche Klausur wird in deutsch geschrieben.
- The questions will be based on material from the Übungen and the Lectures.
- These are typical questions. It is not a "Fragenkatalog". It is a sample of possibilities.

Example questions

1. You are given two DNA sequences to align

ACGTCCTTCATT and GTCTCATG

You have a scoring scheme where a

- match gives you +1
- a mismatch gives you 0
- gap opening costs -10

Write down the best alignment of the two sequences

- 2. You have a scoring scheme where
 - A match gives you +1
 - a mismatch gives you -1
 - opening a gap costs you -1

Write down the best alignment for the same two DNA sequences.

3. You are aligning protein sequences using a substitution matrix :

A 4 -1 -2 -2 0 -1 -1 0 -2 -1 -1 -1 -1 -2 -1 1 0 -3 - R -1 5 0 -2 -3 1 0 -2 0 -3 -2 2 -1 -3 -2 -1 -1 -3 -	
N -2 0 6 1 -3 0 0 0 1 -3 -3 0 -2 -3 -2 1 0 -4 -	2 -3
D -2 -2 1 6 -3 0 2 -1 -1 -3 -4 -1 -3 -3 -1 0 -1 -4 -	3 - 3
C 0 -3 -3 -3 9 -3 -4 -3 -3 -1 -1 -3 -1 -2 -3 -1 -1 -2 -	2 -1
Q -1 1 0 0 -3 5 2 -2 0 -3 -2 1 0 -3 -1 0 -1 -2 -	1 -2
E -1 0 0 2 -4 2 5 -2 0 -3 -3 1 -2 -3 -1 0 -1 -3 -	2 -2
G 0 -2 0 -1 -3 -2 -2 6 -2 -4 -4 -2 -3 -3 -2 0 -2 -2 -	3 - 3
H -2 0 1 -1 -3 0 0 -2 8 -3 -3 -1 -2 -1 -2 -1 -2 -2	2 - 3
I -1 -3 -3 -3 -1 -3 -3 -4 -3 4 2 -3 1 0 -3 -2 -1 -3 -	1 3
L -1 -2 -3 -4 -1 -2 -3 -4 -3 2 4 -2 2 0 -3 -2 -1 -2 -	1 1
K -1 2 0 -1 -3 1 1 -2 -1 -3 -2 5 -1 -3 -1 0 -1 -3 -	2 -2
M -1 -1 -2 -3 -1 0 -2 -3 -2 1 2 -1 5 0 -2 -1 -1 -1 -	1 1
F -2 -3 -3 -3 -2 -3 -3 -3 -1 0 0 -3 0 6 -4 -2 -2 1	3 -1
P -1 -2 -2 -1 -3 -1 -1 -2 -2 -3 -3 -1 -2 -4 7 -1 -1 -4 -	3 -2
S 1 -1 1 0 -1 0 0 0 -1 -2 -2 0 -1 -2 -1 4 1 -3 -	2 -2
T 0 -1 0 -1 -1 -1 -1 -2 -2 -1 -1 -1 -1 -2 -1 1 5 -2 -	2 0
W -3 -3 -4 -4 -2 -2 -3 -2 -2 -3 -2 -3 -1 1 -4 -3 -2 11	2 -3
Y -2 -2 -2 -3 -2 -1 -2 -3 2 -1 -1 -2 -1 3 -3 -2 -2 2	7 -1
V 0 -3 -3 -3 -1 -2 -2 -3 -3 3 1 -2 1 -1 -2 -2 0 -3 -	1 4

Gap opening costs -8. Gap widening (extension) costs -1.

You are given an alignment

AACDQRST A-CD-RST What is the score of this alignment ?

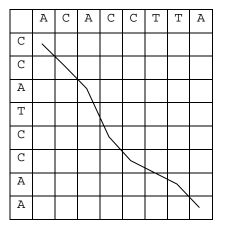
4. Given

AACDQRST A-CD--ST What is the score of this alignment ?

5. AACDQRST A-CD-SST

What is the score of this alignment?

6. You have calculated a score matrix for a pair of DNA sequences. You have performed the traceback calculation and found a result like:



Write down the corresponding sequence alignment with gaps in the correct positions.

7. Outline the steps used to find values for a BLOSUM amino acid similarity matrix.

- 8. What is the advantage of a Needleman-Wunsch alignment compared to a seeded alignment ?
- 9. What is the advantage of a seeded method like BLAST compared to a Needleman-Wunsch alignment ?
- 10. Name an application where you would use a method like BLAST and not a Needleman-Wunsch alignment.
- Name an application where you would use a slow method like Needleman and Wunsch, rather than a BLAST-like method.
- 12. You have two proteins with weak, remote similarity. You know their sequences. You also know the sequences of the original DNA. Why would you expect a better alignment using the protein sequences ?
- 13. In protein alignments, we do not just look at match/mismatches. We look at the similarities between amino acids. How are these represented ?
- 14. There are several different substitution matrices used in protein alignments. Why would you prefer one over another ?
- 15. What is the difference between an iterated blast (psi-blast) search and a simple blast search ?
- 16. What is the advantage of an iterated blast search compared to a simple blast search ?
- 17. I have written a program that generates random DNA sequences. I expect to see 25 % sequence identity between pairs of random sequences in gapped alignments. When I try the calculation, I usually see more than 25 % sequence identity. Why ?
- 18. If I take random biological sequences from a data bank, I see even more sequence similarity. Why ?
- 19. I have two proteins with 20 % sequence identity. I ask you if this is likely to be significant. What other simple piece of information do you need to answer this question properly ?
- 20. If you use the program "chimera", what representation would you pick in order to see the secondary structure ?
- 21. I want to calculate a multiple sequence alignment for N_{seq} sequences. How many pair-wise alignments will I have to calculate ?
- 22.

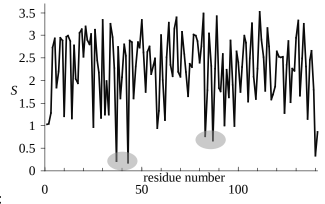
In a multiple sequence alignment, you want to maximise a score, $score = \sum_{b \neq a}^{N_{seq}} \sum_{a=1}^{N_{seq}} S_{a,b}$ What is this score in

words?

- 23. In a multiple sequence alignment, I want to build a "guide tree". What determines the order in which I join the nodes together ?
- 24. I have 3 sequences, A, B, C. The sequences B and C are both related to A, but I cannot get a good alignment score when I align B and C. What could be a reason ? Draw a diagram if it is easier to explain.
- 25. I have calculated a multiple sequence alignment. I want to find which sites in the alignment are conserved and which vary. I would like to make a plot of variability/conservation as a function of sequence position.

You remember a formula $S = \sum_{i=1}^{N_{states}} p_i \ln p_i$ What is the meaning of p_i ?

26. From a multiple sequence alignment, I have calculated variability/conservation as a function of sequence





position:

conserved. Why might they be important residues ?

- 27. In the picture above, some sites are not very conserved. I say these residues cannot be important to the function of the protein. Why may I be wrong ?
- 28. I have calculated a sequence alignment of 400 tyrosine kinases and I find that very few sites seem to be conserved in evolution. How could I change my results, so that more sites seem to be conserved ?
- 29. We have a family of sequences and all pair-wise alignments. I can count the number of differences (mutations) between any two sequences and calculate the fraction of residues that have changed :

$$p_{mut} = \frac{N_{diff}}{N_{length}}$$
. I would like to estimate evolutionary time by saying $t = k p_{mut}$ for some constant k.

This is not a good measure. Why not?

- 30. I want to use aligned DNA sequences to build a phylogenetic tree. Name two reasons that the branches in the tree may not be reliable.
- 31. I have built a phylogenetic tree using a neighbour joining method. Describe a general approach I could use to see how reliable the tree is.
- 32. It is believed that protein sequence evolves and changes faster than protein structure. What could be an evolutionary explanation for this.
- 33. Which graphical representation would you use in order to emphasize the secondary structure content of protein? all-atom, chaintrace, ribbon, ...?
- 34. You have a protein of unknown function from a bacterium. You have made a knock-out mutant, but the bacteria die immediately without the corresponding gene. You have sequenced the protein. What steps would you take to guess the function of the protein ? What kind of information would you look for ?
- 35. Wie würden Sie vorgehen um in einem Multiplen Sequenzalignment potentiell katalytisch wichtige Seitenketten im aktiven Zentrum zu identifizieren ?