Chapter 7: Rapid alignment methods: FASTA and BLAST

- The biological problem
- Search strategies
- FASTA
- BLAST

BLAST: Basic Local Alignment Search Tool

- BLAST (Altschul et al., 1990) and its variants are some of the most common sequence search tools in use
- Roughly, the basic BLAST has three parts:
 - 1. Find local alignments between the query sequence and a database sequence ("seed hits")
 - 2. Extend seed hits into high-scoring local alignments
 - 3. Calculate p-values and a rank ordering of the local alignments
- High-scoring local alignments are called high scoring segment pairs (HSPs)
- Gapped BLAST introduced in 1997 allows for gaps in alignments

Finding seed hits

- First, we generate a set of *neighborhood sequences* for given k, *match score matrix and threshold* T
- Neighborhood sequences of a k-word w include all strings of length k that, when aligned against w, have the alignment score at least T
- For instance, let I = GCATCGGC, J = CCATCGCCATCG and k = 5, match score be 1, mismatch score be 0 and T = 4

Finding seed hits

- I = GCATCGGC, J = CCATCGCCATCG, k = 5, match score 1, mismatch score 0, T = 4
- This allows for one mismatch in each k-word
- The neighborhood of the first k-word of I, GCATC, is GCATC and the 15 sequences

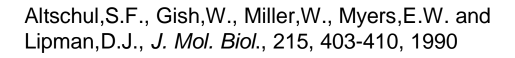
 $\begin{cases} A \\ CCATC, G \\ T \end{cases} \begin{pmatrix} A \\ GATC, GC \\ T \\ T \end{pmatrix} \begin{pmatrix} C \\ GTC, GCA \\ GC \\ CC, GCAT \\ G \\ T \end{pmatrix} \begin{pmatrix} A \\ CC, GCAT \\ G \\ T \end{pmatrix}$

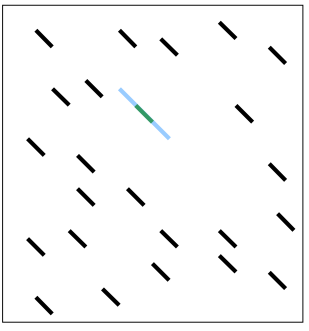
Finding seed hits

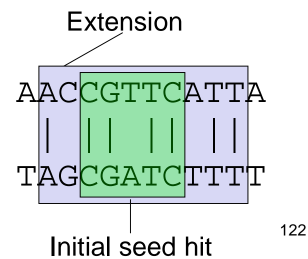
- I = GCATCGGC has 4 k-words and thus 4x16 = 64 5-word patterns (seed hits) to locate in J
- These patterns can be found using exact search in time proportional to the sum of pattern lengths + length of J + number of matches (Aho-Corasick algorithm)
 - Methods for pattern matching are developed on course 58093 String processing algorithms

Extending seed hits: original BLAST

- Initial seed hits are extended
- Extensions do not add gaps to the alignment
- Sequence is extended into a HSP until the alignment score drops below the maximum attained score minus a threshold parameter value
- All statistically significant HSPs
 reported



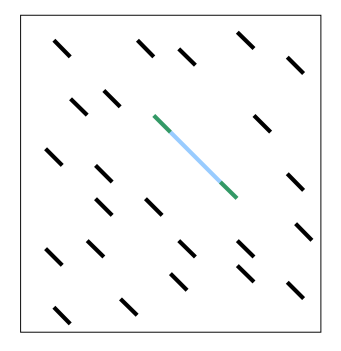




Extending seed hits: gapped BLAST

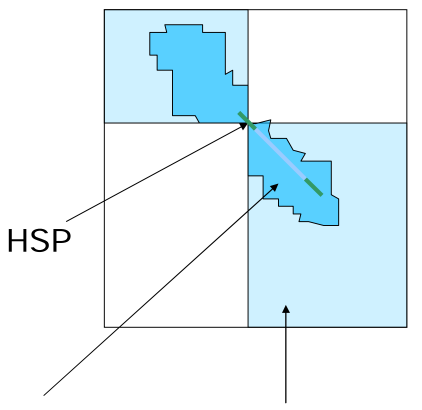
- In later version of BLAST, two seed hits have to be found on the same diagonal
 - Hits have to be non-overlapping
 - If the hits are closer than A (additional parameter), then they are joined into a HSP
- Threshold value T is lowered to achieve comparable sensitivity
- If the resulting HSP achieves a score at least S_g, a *gapped extension* is triggered

Altschul SF, Madden TL, Schäffer AA, Zhang J, Zhang Z, Miller W, and Lipman DJ, *Nucleic Acids Res.* 1;25(17), 3389-402, 1997 Introduction to bioinformatics, Autumn 2007



Gapped extensions of HSPs

- Local alignment is performed starting from the HSP
- Dynamic programming matrix filled in "forward" and "backward" directions (see figure)
- Skip cells where value would be X_g below the best alignment score found so far



above cutoff parameter

Region searched with score Region potentially searched by the alignment algorithm

Estimating the significance of results

- In general, we have a score S(D, X) = s for a sequence X found in database D
- BLAST rank-orders the sequences found by p-values
- The p-value for this hit is P(S(D, Y) ≥ s) where Y is a random sequence
 - Measures the amount of "surprise" of finding sequence X
- A smaller p-value indicates more significant hit
 - A p-value of 0.1 means that one-tenth of random sequences would have as large score as our result

Estimating the significance of results

- In BLAST, p-values are computed roughly as follows
- There are nm places to begin an optimal alignment in the n x m alignment matrix
- Optimal alignment is preceded by a mismatch and has t matching (identical) letters
 - (Assume match score 1 and mismatch score 0)
- Let p = P(two random letters are equal)
- The probability of having a mismatch and then t matches is (1-p)p^t

Estimating the significance of results

- We model this event by a Poisson distribution (why?) with mean $\lambda = nm(1-p)p^t$
- P(there is local alignment t or longer)

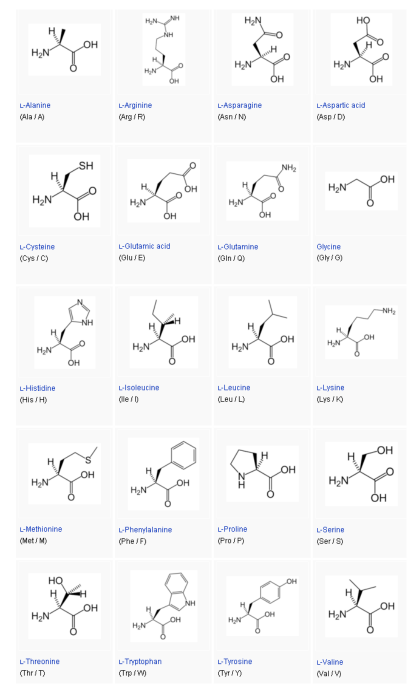
= 1 - P(no such event)

 $= 1 - e^{-\lambda} = 1 - \exp(-nm(1-p)p^{t})$

- An equation of the same form is used in Blast:
- E-value = $P(S(D, Y) \ge s) \approx 1 exp(-nm\gamma\xi^t)$ where $\gamma > 0$ and $0 < \xi < 1$
- Parameters γ and ξ are estimated from data

Scoring amino acid alignments

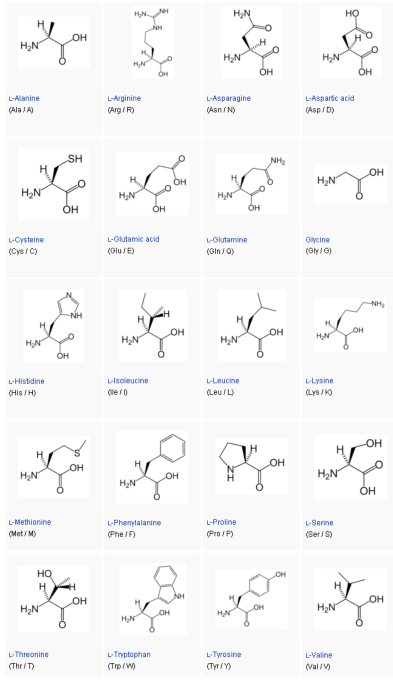
- We need a way to compute the score S(D, X) for aligning the sequence X against database D
- Scoring DNA alignments was discussed previously
- Constructing a scoring model for amino acids is more challenging
 - 20 different amino acids vs. 4 bases
- Figure shows the molecular structures of the 20 amino acids



http://en.wikipedia.org/wiki/List_of_standard_amino_acids

Scoring amino acid alignments

- Substitutions between chemically similar amino acids are more frequent than between dissimilar amino acids
- We can check our scoring model against this



http://en.wikipedia.org/wiki/List_of_standard_amino_acids

Score matrices

- Scores s = S(D, X) are obtained from score matrices
- Let $A = A_1 a_2 \dots a_n$ and $B = b_1 b_2 \dots b_n$ be sequences of equal length (no gaps allowed to simplify things)
- To obtain a score for alignment of A and B, where a_i is aligned against b_i , we take the ratio of two probabilities
 - The probability of having A and B where the characters match (match model M)
 - The probability that A and B were chosen randomly (random model R)

Score matrices: random model

Under the random model, the probability of having X and Y is

$$P(A,B|R) = \prod_i q_{ai} \prod_i q_{bi}$$

- where \boldsymbol{q}_{xi} is the probability of occurence of amino acid type \boldsymbol{x}_i
- Position where an amino acid occurs does not affect its type

Score matrices: match model

- Let p_{ab} be the probability of having amino acids of type a and b aligned against each other given they have evolved from the same ancestor c
- The probability is

$$P(A, B|M) = \prod_i p_{a_i b_i}$$

Score matrices: log-odds ratio score

We obtain the score S by taking the ratio of these two probabilities

$$\frac{P(A,B|M)}{P(A,B|R)} = \frac{\prod_i p_{a_i b_i}}{\prod_i q_{a_i} \prod_i q_{b_i}} = \prod_i \frac{p_{a_i b_i}}{q_{a_i} q_{b_i}}$$

and taking a logarithm of the ratio

$$S = \log_2 rac{P(A,B|M)}{P(A,B|R)} = \sum_{i=1}^n \log_2 rac{p_{a_i b_i}}{q_{a_i} q_{b_i}} = \sum_{i=1}^n s(a_i,b_i)$$

$$S = \log_2 rac{P(A,B|M)}{P(A,B|R)} = \sum_{i=1}^n \log_2 rac{p_{a_i b_i}}{q_{a_i} q_{b_i}} = \sum_{i=1}^n s(a_i,b_i)$$

The score S is obtained by summing over character pair-specific scores:

$$s(a,b) = \log_2 \frac{p_{ab}}{q_a q_b}$$

The probabilities q_a and p_{ab} are extracted from data

Calculating score matrices for amino acids

 $s(a,b) = \log_2 \frac{p_{ab}}{q_a q_b}$

- Probabilities q_a are in principle easy to obtain:
 - Count relative frequencies of every amino acid in a sequence database

Calculating score matrices for amino acids

- To calculate p_{ab} we can use a known pool of aligned sequences
- BLOCKS is a database of highly conserved regions for proteins
- It lists multiply aligned, ungapped and conserved protein segments
- Example from BLOCKS shows genes related to human gene associated with DNA-repair defect xeroderma pigmentosum

 $s(a,b) = \log_2 \frac{p_{ab}}{q_a q_b}$

Block PR00851A			
ID XRODRMPGMNTB; BLOCK			
AC PR00851A; distance from previous block=(52,131)			
DE Xeroderma pigmentosum group B protein signature			
BL adapted; width=21; seqs=8; 99.5%=985; strength=1287			
XPB_HUMAN P19447 (74)	RPLWVAPDGHIFLEAFSPVYK	54	
XPB_MOUSE P49135 (74)	RPLWVAPDGHIFLEAFSPVYK	54	
P91579 (80)	RPLYLAPDGHIFLESFSPVYK	67	
XPB_DROME Q02870 (84)	RPLWVAPNGHVFLESFSPVYK	79	
RA25_YEAST Q00578 (131)	PLWISPSDGRIILESFSPLAE	100	
Q38861 (52)	RPLWACADGRIFLETFSPLYK	71	
013768 (90)	PLWINPIDGRIILEAFSPLAE	100	
000835 (79)	RPIWVCPDGHIFLETFSAIYK	86	

http://blocks.fhcrc.org

BLOSUM matrix

- BLOSUM is a score matrix for amino acid sequences derived from BLOCKS data
- First, count pairwise matches f_{x,y} for every amino acid type pair (x, y)
- For example, for column 3 and amino acids L and W, we find 8 pairwise matches: $f_{L,W} = f_{W,L} = 8$

RF	PLWVAPD	
RF	LWVAPR	
RF	PLWVAPN	
ΡI	.WISPSD	
RF	LWACAD	
ΡI	WINPID	
RF	IWVCPD	

Creating a BLOSUM matrix

Probability p_{ab} is obtained by dividing f_{ab} with the total number of pairs (note difference with course book):

$$p_{ab} = f_{ab} / \sum_{x=1}^{20} \sum_{y=1}^{x} f_{xy}$$

• We get probabilities q_a by

$$q_a = \sum_{b=1}^{20} p_{ab}$$

RPLWVAPD RPLWVAPR RPLWVAPN PLWISPSD RPLWACAD PLWINPID RPIWVCPD

Creating a BLOSUM matrix

The probabilities p_{ab} and q_a can now be plugged into

$$s(a,b) = \log_2 \frac{p_{ab}}{q_a q_b}$$

to get a 20 x 20 matrix of scores s(a, b).

- Next slide presents the BLOSUM62 matrix
 - Values scaled by factor of 2 and rounded to integers
 - Additional step required to take into account expected evolutionary distance
 - Described in the course book in more detail

BLOSUM62

A R N D C O E G H I L K M S Т V B 7 Х F Р W Υ A 4 -1 -2 -2 0 -1 -1 0 -2 -1 -1 -1 -1 -2 -1 1 0 - 3 - 20 - 2 - 10 - 41 0 - 3 - 2R – 1 - 5 0 - 2 - 30 -2 2 -1 -3 -2 -1 -1 -3 -2 -3 -1 0 -1 -4 N - 20 1 - 30 0 0 1 - 3 - 30 - 2 - 3 - 20 -2 -33 0 - 1 - 46 1 D - 2 - 26 -3 1 0 2 -1 -1 -3 -4 -1 -3 -3 -1 0 - 1 - 4 - 3 - 34 1 -1 -4 0 -3 -3 -3 9 -3 -4 -3 -3 -1 -1 -3 -1 -2 -3 -1 -1 -2 -2 -1 -3 -3 -2 -4 C 5 2 -2 0 -3 -2 0 -3 -1 0 -1 -2 -1 -23 -1 -4 0 -1 1 0 0 -3 1 0 E -1 0 2 - 42 5 - 20 - 3 - 31 - 2 - 3 - 10 - 1 - 3 - 2 - 24 -1 -4 0 1 6 -2 -4 -4 -2 -3 -3 -2 0 -2 -2 -3 -3 -1 -2 -1 -4 G 0 - 20 -1 -3 -2 -2 н – 2 0 1 -1 -3 0 0 -2 8 -3 -3 -1 -2 -1 -2 -1 -2 -2 2 - 3 0 0 -1 -4 2 - 3 I -1 -3 -3 -3 -1 -3 -3 -4 -3 4 1 0 -3 -2 -1 -3 -1 3 -3 -3 -1 -4 2 4 -2 L -1 -2 -3 -4 -1 -2 -3 -4 -3 2 0 -3 -2 -1 -2 -11 -4 -3 -1 -4 1 1 -2 -1 -3 -2 5 -1 -3 -1 0 -1 -3 -2 -2 к -1 2 0 - 1 - 30 1 -1 -4 1 2 -1 M - 1 - 1 - 2 - 3 - 10 - 2 - 3 - 25 0 -2 -1 -1 -1 -1 1 -3 -1 -1 -4 F -2 -3 -3 -3 -2 -3 -3 -1 0 0 -3 0 6 -4 -2 -2 1 3 -1 -3 -3 -1 -4 P -1 -2 -2 -1 -3 -1 -1 -2 -2 -3 -3 -1 -2 -4 7 -1 -1 -4 -3 -2 -2 -1 -2 -4 0 - 1 - 2 - 1S 1 –1 1 0 -1 0 0 0 -1 -2 -24 - 3 -2 -2 0 0 0 1 -4 5 - 2 - 2 Т 0 -1 0 -1 -1 -1 -1 -2 -2 -1 -1 -1 -1 -2 -1 1 0 -1 -1 0 - 4W -3 -3 -4 -4 -2 -2 -3 -2 -2 -3 -2 -3 -1 1 -4 -3 -2 11 2 -3 -4 -3 -2 -4 Y -2 -2 -2 -3 -2 -1 -2 -3 2 -1 -1 -2 -1 3 - 3 - 2 - 2 2 7 -1 -3 -2 -1 -4 1 -2 1 -1 -2 -2 0 -3 -1 4 -3 -2 -1 -4 0 -3 -3 -3 -1 -2 -2 -3 -3 3 V 3 4 -3 0 1 -1 0 -3 -4 0 -3 -3 -2 0 -1 -4 -3 -3 B -2 -1 4 1 -1 -4 7. – 1 0 0 1 - 33 4 -2 0 -3 -3 1 -1 -3 -1 0 -1 -3 -2 -21 4 -1 -4 Х 0 -1 -1 -1 -2 -1 -1 -1 -1 -1 -1 -1 -1 -1 -2 0 0 -2 -1 -1 -1 -1 -1 -4

Using BLOSUM62 matrix

MQLEANADTSV

LQEQAEAQGEM

$$s = \sum_{i=1}^{11} s(a_i, b_i)$$

$$= 2 + 5 - 3 - 4 + 4 + 0 + 4 + 0 - 2 + 0 + 1$$
$$= 7$$

Demonstration of BLAST at NCBI

http://www.ncbi.nlm.nih.gov/BLAST/