Genome 559 Introduction to Statistical and Computational Genomics Winter 2010

> Lecture 14a: BLAST Larry Ruzzo

### 1 minute responses

Pacing was: (a) A little slow (1), (b) great (3) [maybe we don't need semesters after all!], or (c) I was lost/equation-dense (4) (but,I'll try harder to keep up with reading)

Paper slides for note-taking *really* help. *Agreed* More time for problems helped. *Hopefully again today*. Is revised hw schedule on web? *Some*.

Liked it, but need some practice problems for it to sink in. See hw5!

Fuzzy on purpose of relative entropy; why does it matter. *If motif distribution is like background (low entropy), WMM prediction will be error-prone. Similarly, columns of low entropy may only add noise; at edges, especially, maybe delete them.* 

Didn't explain substring matches/match objects (2) Today

# **BLAST**:

### **Basic Local Alignment Search Tool**

Altschul, Gish, Miller, Myers, Lipman, J Mol Biol 1990

#### The most widely used comp bio tool

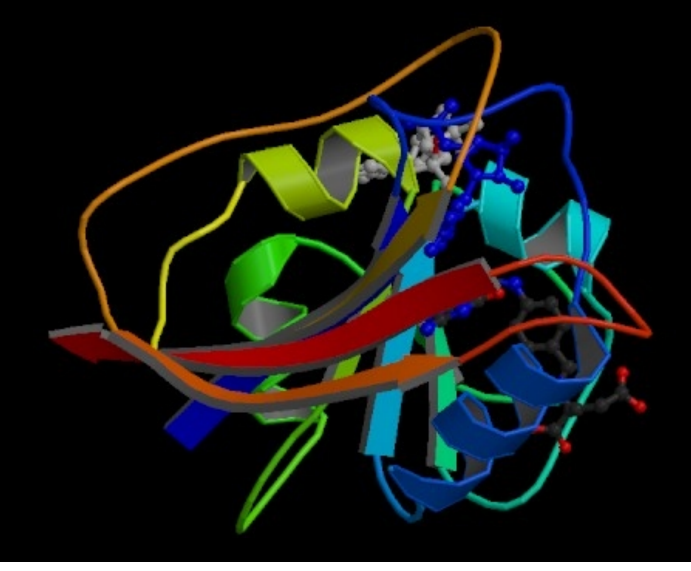
- Which is better: long mediocre match or a few nearby, short, strong matches with the same total score?
  - score-wise, exactly equivalent
  - biologically, later may be more interesting
  - if must miss some, rather miss the former (?)

#### BLAST is a heuristic emphasizing the later

speed/sensitivity tradeoff: BLAST may miss weak matches, but gains greatly in speed

*Heuristic*: A method proceeding towards a solution by trial and error, intuition or loosely defined rules. Cf. Algorithm; Smith-Waterman, etc.

# A Protein Structure: (Dihydrofolate Reductase)



# **BLAST: What**

Input:

a query sequence (say, 50-300 residues)

a data base to search for other sequences similar to the query (say, 10<sup>6</sup> - 10<sup>9</sup> residues)

a score matrix  $\sigma(r,s)$ , giving cost of substituting r for s (& perhaps gap costs)

various score thresholds & tuning parameters

Output:

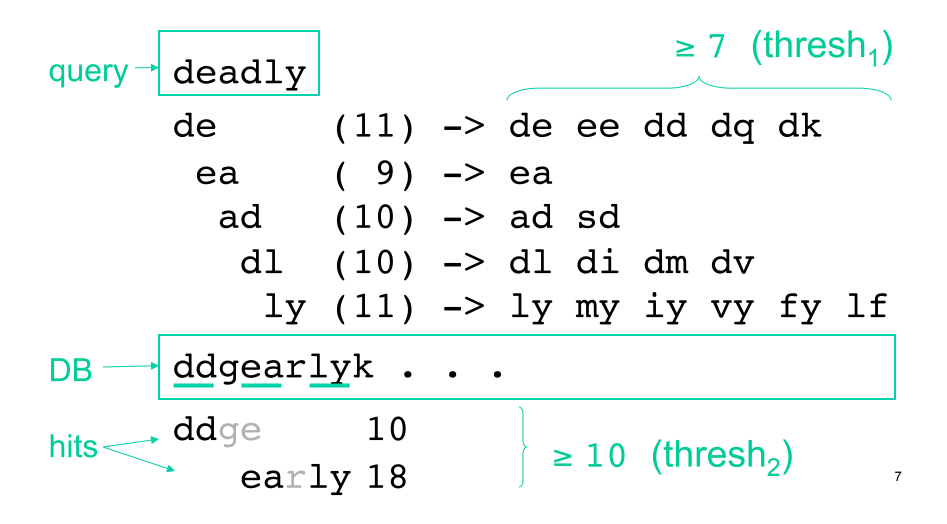
"all" matches in data base above threshold

"E-value" of each

## **BLAST: How**

- Idea: emphasize parts of data base near a good match to some short subword of the query
- Break query into overlapping words w<sub>i</sub> of small fixed length (e.g. 3 aa or 11 nt)
- For each w<sub>i</sub>, find (empirically, ~50) "neighboring" words v<sub>ij</sub> with score  $\sigma(w_i, v_{ij}) > \text{thresh}_1$
- Look up each v<sub>ii</sub> in database (via prebuilt index) --
- i.e., exact match to short, high-scoring word Extend each such "seed match" (bidirectional)
- Report those scoring > thresh<sub>2</sub>, calculate E-values

### **BLAST: Example**



# BLOSUM 62

	Α	R	Ν	D	С	Q	Ε	G	Н	Ι	L	Κ	Μ	F	Ρ	S	Т	W	Υ	V
Α	4	-1	-2	-2	0	-1	-1	0	-2	-1	-1	-1	-1	-2	-1	1	0	-3	-2	0
R	-1	5	0	-2	-3	1	0	-2	0	-3	-2	2	-1	-3	-2	-1	-1	-3	-2	-3
Ν	-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
С	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
Е	-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
н	-2	0	1	-1	-3	0	0	-2	8	-3	-3	-1	-2	-1	-2	-1	-2	-2	2	-3
Ι	-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
К	-1	2	0	-1	-3	1	1	-2	-1	-3	-2	5	-1	-3	-1	0	-1	-3	-2	-2
Μ	-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
Ρ	-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
Т	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
Υ	-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

## **BLAST Refinements**

"Two hit heuristic" – need 2 nearby, nonoverlapping, gapless hits before trying to extend either

"Gapped BLAST" – run heuristic version of Smith-Waterman, bi-directional from hit, until score drops by fixed amount below max

PSI-BLAST – For proteins, iterated search, using "weight matrix" pattern from initial pass to find weaker matches in subsequent passes (PSI=pos specific iter) Many others

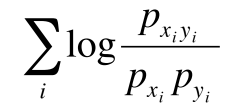
## A Likelihood Ratio

Defn: two proteins are *homologous* if they are alike because of shared ancestry; similarity by descent

Suppose among proteins overall, residue x occurs with frequency p<sub>x</sub> Then in a random ungapped alignment of 2 random proteins, you would expect to find x aligned to y with prob p<sub>x</sub>p<sub>y</sub>

Suppose among *homologs*, x & y align with prob p<sub>xy</sub>

Are seqs X & Y homologous? Which is more likely, that the alignment reflects chance or homology? Use a *likelihood ratio test.* 



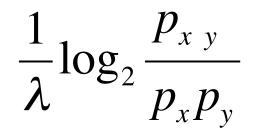
E.g., BLOSUM62: trusted "homologues" = BLOCKS w/ ≥ 62% identity.

## Non-ad hoc Alignment Scores

Take alignments of homologs and look at frequency of x-y alignments vs freq of x, y overall

BLOSUM approach

large collection of trusted alignments (the BLOCKS DB) subsetted by similarity, e.g. BLOSUM62 => 62% identity



http://blocks.fhcrc.org/blocks-bin/getblock.pl?IPB013598

# ad hoc Alignment Scores?

Make up any scoring matrix you like

Somewhat surprisingly, under pretty general assumptions<sup>\*\*</sup>, it is *equivalent* to the scores constructed as above from some set of probabilities p<sub>xy</sub>, so you might as well understand what they are

NCBI-BLASTN: +1/-2  $\leftrightarrow$  95% identity WU-BLASTN: +5/-4  $\leftrightarrow$  66% identity

<sup>\*\*</sup> e.g., average scores should be negative, but you probably want that anyway, otherwise local alignments turn into global ones, and some score must be > 0, else best match is empty

# Summary

BLAST is a highly successful search/alignment heuristic. It looks for alignments anchored by short, strong, ungapped "seed" alignments

Strengths:

Speed, E-values, well-supported implementation & web server

Weaknesses:

Heuristic search can miss weaker matches