Programming for Biololgy Similarity Searching II -

Practical search strategies

Bill Pearson wrp@virginia.edu

CSHL Programming for Biology

1

Effective Similarity Searching

- Always search protein databases (possibly with DNA blastx, fastx)
- 2. Use E()-values, not percent identity, to infer homology
 - E() < 0.001 is significant in a single search
- 1. Search smaller (comprehensive) databases
 - Less redundancy; higher sensitivity
- 2. Change the scoring matrix for:
 - short sequences (exons, reads)
 - short evolutionary distances (mammals, vertebrates, aproteobacteria)
 - high identity (>50% alignments) to reduce over-extension

CSHL Programming for Biology

Review – Sequence Similarity - Conclusions

- <u>Homologous</u> sequences share a common ancestor, but most sequences are <u>non-</u> <u>homologous</u>
- · Always compare Protein Sequences
- Sequence Homology can be reliably inferred from statistically significant similarity (non-homology cannot from non-similarity)
- Homologous proteins share common structures, but not necessarily common functions
- Sequence statistical significance estimates are accurate (verify this yourself)10⁻⁶ < E() < 10⁻³ is statistically significant

CSHL Programming for Biology

3

3

Similarity Searching II

- 1. What question to ask?
- 2. What program to use?
- 3. What database to search?
- 4. When to do something different (changing scoring matrices)
- 5. Is every aligned domain homologous?

CSHL Programming for Biology

1. What question to ask?

- Is there an homologous protein (a protein with a similar structure)?
- Does that homologous protein have a similar function?
- Does XXX genome have YYY (kinase, GPCR, ...)?

Questions not to ask:

- Does this DNA sequence have a similar regulatory element (too short – never significant)?
- Does (non-significant) protein have a similar function/modification/antigenic site?

CSHL Programming for Biology

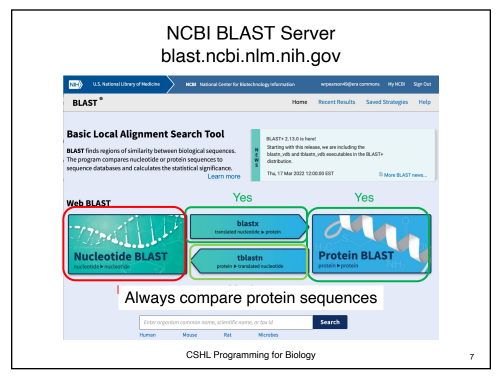
5

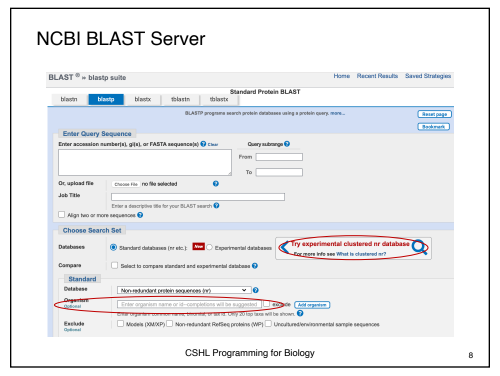
5

2. What program to run?

- What is your query sequence?
 - protein BLASTP (NCBI), SSEARCH (EBI)
 - protein coding DNA (EST) –
 BLASTX (NCBI), FASTX (EBI)
 - DNA (structural RNA, repeat family) –
 BLASTN (NCBI), FASTA (EBI)
- Does XXX genome have YYY (protein)?
 - TBLASTN YYY vs XXX genome
 - TFASTX YYY vs XXX genome
- Does my protein contain repeated domains?
 - LALIGN (UVa http://fasta.bioch.virginia.edu, EBI)

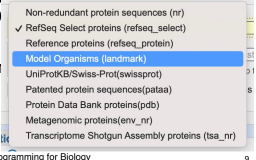
CSHL Programming for Biology





3. What database to search?

- Search the smallest comprehensive database likely to contain your protein
 - vertebrates human proteins (40,000)
 - NCBI Landmark sequences (human, mouse, no rat)
 - Quest for Orthologs reference proteomes (1,000,000)
- Search a richly annotated protein set (SwissProt: 500,00
- Always search NR
- Never Search "Gen



CSHL Programming for Biology

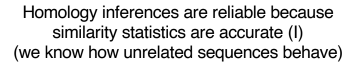
Effective Similarity Searching

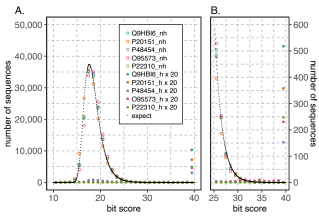
- 1. Always search protein databases (possibly with translated DNA)
- 2. Use E()-values, not percent identity, to infer homology
 - E() < 0.001 is significant in a single search

1. Search smaller (comprehensive) databases

- 2. Change the scoring matrix for:
 - short sequences (exons, reads)
 - short evolutionary distances (mammals, vertebrates, aproteobacteria)
 - high identity (>50% alignments) to reduce over-extension
- 3. Is every aligned residue homologous?
 - alignment overextension
- 4. (Tomorrow) All methods (pairwise, HMM, PSSM) miss homologs, and find homologs the other methods miss

CSHL Programming for Biology



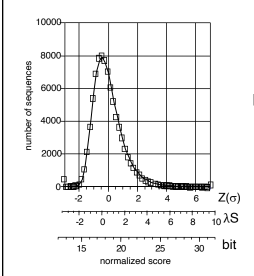


Distributions of similarity scores in searches with 5 human enzymes. Open circles (_nh) show scores for non-homologs. Closed circles show homolog (_h) scores.

CSHL Programming for Biology

11

Why smaller databases are better - statistics



 $S' = \lambda S_{raw}$ - In K m n $S_{bits} = (\lambda S_{raw} - In K)/In(2)$ $P(S'>x) = 1 - exp(-e^{-x})$ $P(S_{bits} > x) = 1 - exp(-mn2^{-x})$

E(S'>x ID) = P D

Bonferroni correction

P(B bits) = m n 2^{-B} P(40 bits)= 1.5×10^{-7} E(40 | D=4000) = 6×10^{-4}

E(40 | D=500E6) = 75

CSHL Programming for Biology

12

When do you trust search results?

- BLAST(P)
 - Statistically significant similarity
 - Expectation E()-value
 - Why should you trust the statistics??
 - What about false negatives??

CSHL - Programming for Biology

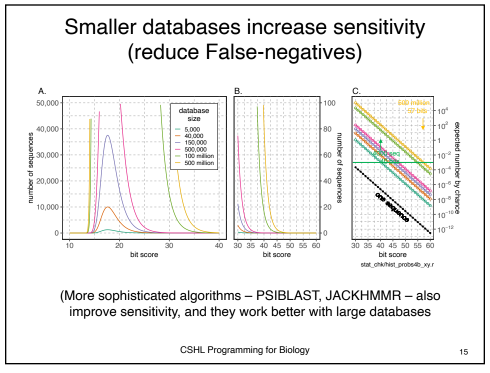
13

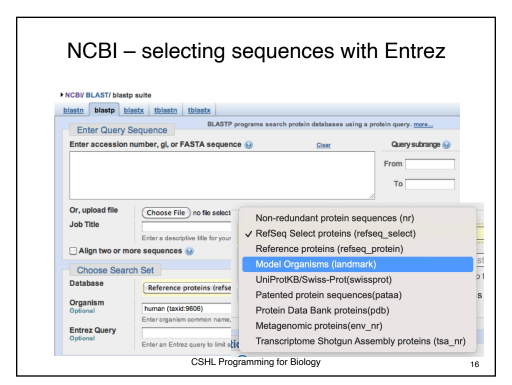
13

Local similarity statistics

```
\begin{split} S' &= \lambda S_{raw} \text{ - In K m n } \quad \text{m: query length, n: subj length} \\ S_{bit} &= (\lambda S_{raw} \text{ - In K)/In(2)} \\ P(S'>x) &= 1 \text{ - exp(-e^{-x})} \\ P(S'>x) &= e^{-x} \quad \text{(for P < 0.1)} \\ \\ P(S_{bits} > \text{bits)} &= 1 \text{ -exp(-mn2^{-x})} \\ P(S_{bits} > \text{bits)} &= \text{mn2-bits} \quad \text{(for P < 0.1)} \\ \\ E(S', S_{bits} \text{ ID)} &= \text{PD} \\ E(S_{bits} \text{ ID)} &= \text{D mn2-bits} \quad \text{Bonferroni correction} \\ \\ \text{dblength} &= \text{D n} \\ E(S_{bit}) &= \text{m dblength 2-bits} \quad \text{(BLAST)} \end{split}
```

CSHL Programming for Biology





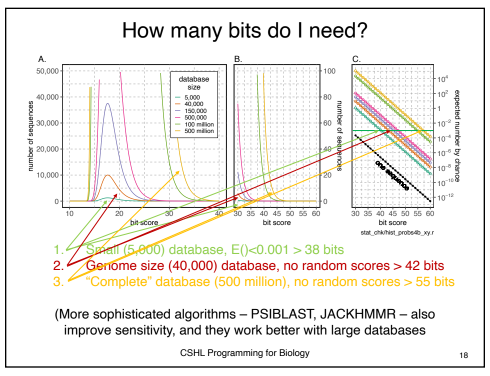
3. What database to search?

Database	Size	Bits (0.001)
Landmark	441 thousand	47
SwissProt	480 thousand	47
Refseq_Select	64 million	53
Refseq_Protein	234 million	55
NR (clustered)	242 million	55
NR	510 million	56

CSHL Programming for Biology

17

17



Effective Similarity Searching

- Always search protein databases (possibly with translated DNA)
- 2. Use E()-values, not percent identity, to infer homology
 E() < 0.001 is significant in a single search
- 1. Search smaller (comprehensive) databases
- 2. Change the scoring matrix for:
 - short sequences (exons, reads)
 - short evolutionary distances (mammals, vertebrates, aproteobacteria)
 - high identity (>50% alignments) to reduce over-extension
- 3. Is every aligned residue homologous?
 - alignment overextension

CSHL Programming for Biology

19

19

Scoring matrices – shifting lookback (where do those bits come from?)

- Scoring matrices can set the evolutionary lookback time for a search
 - Lower PAM (PAM10/VT10 ... PAM/VT40) for closer (10% ... 50% identity)
 - Higher BLOSUM for higher conservation (BLOSUM50 distant, BLOSUM80 conserved)
- Shallow scoring matrices for short domains/short queries (metagenomics)
 - Matrices have "bits/position" (score/position), 40 aa at 0.45 bits/position (BLOSUM62) means 18 bit ave. score (50 bits significant)
- Deep scoring matrices allow alignments to continue, possibly outside the homologous region

CSHL Programming for Biology

Scoring matrices and alignment length

$$\lambda S_{i,j} = \log_b(\frac{q_{i,j}}{p_i p_j})$$

 $\begin{array}{ll} q_{ij}: \mbox{homolog frequency wat PAM40, 250} \\ q_{R:N\,(\ 40)} = 0.000435 & p_R = 0.051 \\ q_{R:N\,(250)} = 0.002193 & p_N = 0.043 \\ \lambda_2 \ S_{ij} = \lg_2 \ (q_{ij}/p_ip_j) \ \lambda_e \ S_{ij} = \ln(q_{ij}/p_ip_j) & p_Rp_N = 0.002193 \\ \lambda_2 \ S_{R:N(\ 40)} = \lg_2 \ (0.000435/0.00219) = -2.333 \\ \lambda_2 = 1/3; \ S_{R:N(\ 40)} = -2.333/l_2 = -7 \\ \lambda \ S_{R:N(250)} = \lg 2 \ (0.002193/0.002193) = 0 \end{array}$

CSHL Programming for Biology

21

21

Empirical matrix performance (median results from random alignments)

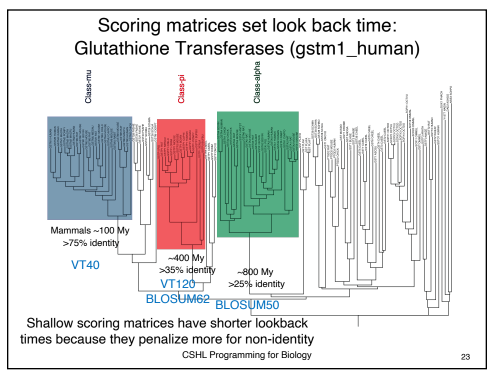
	target % ident	bits/position	aln len (50 bits)
VT160 -12/-2	23.8	0.26	192
BLOSUM50 -10/-2	25.3	0.23	217
BLOSUM62* -11/-1	28.9	0.45	111
VT120 -11/-1	27.4	1.03	48
VT80 -11/-1	51.9	1.55	32
PAM70* -10/-1	33.8	0.64	78
PAM30* -9/-1	45.5	1.06	47
VT40 -12/-1	72.7	2.76	18
VT20 -15/-2	84.6	3.62	13
VT10 /16/-2	90.9	4.32	12

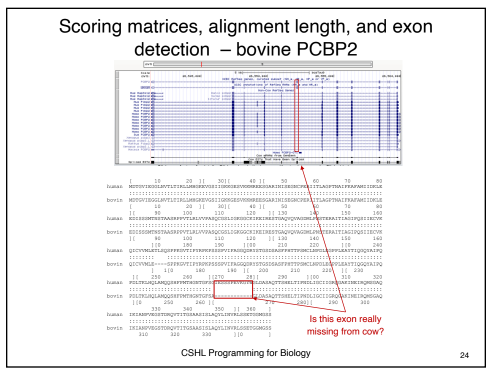
HMMs can be very "deep"

Pearson (2013) Curr. Prot. Bioinformatics 3.5.1

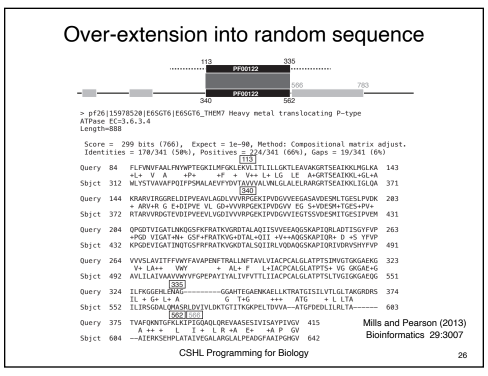
CSHL Programming for Biology

22





S	corin	g m	atri	ces,	aligr	nmei	nt le	ngth	n, and	d ex	on
Scoring matrices, alignment length, and exon detection – bovine PCBP2											
	name	start	end	len	VT10	bits	BP62	Bits	PAM30	Bits	
	ex_1	1	23	23	=	58	+5	45	+5	57	
	ex_2	24	31	8	=	30	_	<25	=	19	
	ex_3	32	42	11	=	36	+1	27	=	26	
	ex_4	43	81	39	=	88	=	61	+5	95	
	ex_5	82	125	44	=	96	=	66	=	104	
	ex_6	126	168	43	=	96	+5	65	+7	106	
	ex_7	169	197	29	=	69	+2	50	+2	70	
	ex_8	198	228	31	=	76	+43	53	=	80	
	ex_9	229	242	14	+4	40	+4	32	+2	37	
	ex_10	243	266	24	+5	60	+87	45	+5	68	
	ex_11	267	280	14	=	42	+38	32	=	43	
	ex_12	281	297	17	=	49	+2	34	_		
	ex_13	298	354	57	=	120	=	78	+9	135	
	ex_14	355	365	11	=	37	=	32	_		
VT10 BP62 qregion: bits=71.5; id=1.00; exon_8-8 qregion: 181-197 : bits=1.6; id=0.500: exon_7-7 qregion: bits=72.7; id=1.000: exon_8-8 qregion: 229-242 : bits=52.7; id=1.000: exon_8-8 qregion: 23-255 : bits=5.4; id=0.286 exon_10-10 properties of the properties of											
Shallow matrix (MD10) finds exon and exon boundaries Deep (sensitive) matrix (BP62) finds exon, but overextends exon boundaries The exon is present in cow, but not detected because it is sho											
CSHL Programming for Biology											



Scoring Matrices - Summary

- PAM and BLOSUM matrices greatly improve the sensitivity of protein sequence comparison – low identity with significant similarity
- PAM matrices have an evolutionary model lower number, less divergence – lower=closer; higher=more distant
- BLOSUM matrices are sampled from conserved regions at different average identity – higher=more conservation
- Shallow matrices set maximum look-back time
- Short alignments (domains, exons, reads) require shallow (higher information content) matrices

CSHL Programming for Biology

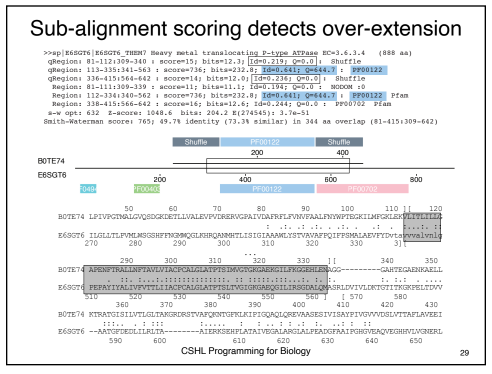
27

27

Effective Similarity Searching

- Always search protein databases (possibly with DNA blastx, fastx)
- 2. Use E()-values, not percent identity, to infer homology
 - E() < 0.001 is significant in a single search
- Search smaller (comprehensive) databases
 - Less redundancy; better sensitivity
- 2. Change the scoring matrix for:
 - short evolutionary distances (mammals, vertebrates, aproteobacteria)
 - short sequences (exons, reads)
 - high identity (>50% alignments) to reduce over-extension

CSHL Programming for Biology



Homology, non-homology, and over-extension

- Sequences that share statistically significant sequence similarity are homologous (simplest explanation)
- But not all regions of the alignment contribute uniformly to the score
 - lower identity/Q-value because of non-homology (overextension) ?
 - lower identity/Q-value because more distant relationship (domains have different ages) ?
- Test by searching with isolated region
 - can the <u>distant domain (?)</u> find closer (significant) homologs?
- Similar (homology) or distinct (non-homology) structure is the gold standard
- Multiple sequence alignment can obscure over-extension
 - if the alignment is over-extended, part of the alignment is NOT homologous

CSHL Programming for Biology

Effective Similarity Searching

- Always search protein databases (possibly with translated DNA)
- 2. Use E()-values, not percent identity, to infer homology
 - E() < 0.001 is significant in a single search
 - Control for statistical accuracy highest scoring unrelated sequence
- 1. Search smaller (comprehensive) databases
- 2. Change the scoring matrix for:
 - short sequences (exons, reads)
 - short evolutionary distances (mammals, vertebrates, aproteobacteria)
 - high identity (>50% alignments) to reduce over-extension
- 3. Is every aligned residue homologous?
 - alignment overextension

CSHL Programming for Biology

31

31

workshop II – parsing blast results

Goto:

fasta.bioch.virginia.edu/mol_evol/pfb_python_matrices.html

Your goal is to reproduce a version of this table:

Matrix		align_len	evalue
VT160	29.7	67	2.1
BLOSUM50	34.0	121	1.2
BLOSUM62* -11/-1	31.2	90	0.37
VT80	66.7	50	1.8
VT40	72.7	11	1.3

CSHL Programming for Biology