

Recap From Last Time:

- Sequence alignment is a fundamental operation underlying much of bioinformatics.
- Introduced dot matrices, dynamic programing and the BLAST heuristic approaches.
- Key point: Even when optimal solutions can be obtained they are not necessarily unique or reflective of the biologically correct alignment.
- Introduced classic global and local alignment algorithms (Needleman–Wunsch and Smith–Waterman) and their major application areas.
- Heuristic approaches are necessary for large database searches and many genomic applications.

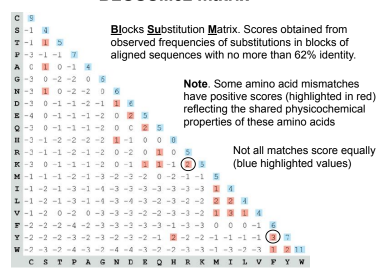
Todays Menu

- Sequence motifs and patterns: Simple approaches for finding functional cues from conservation patterns
- Sequence profiles and position specific scoring matrices (PSSMs): Building and searching with profiles, Their advantages and limitations
- PSI-BLAST algorithm: Application of iterative PSSM searching to improve BLAST sensitivity
- Hidden Markov models (HMMs): More versatile probabilistic model for detection of remote similarities

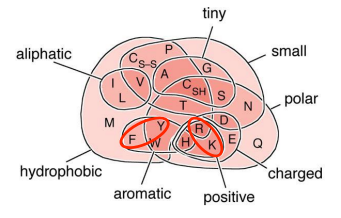
Side Note:

Q. Where do our alignment match and mis-match scores typically come from?

By default BLASTp match scores come from the BLOSUM62 matrix

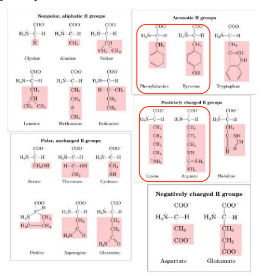


Protein scoring matrices reflect the properties of amino acids



Key Trend: High scores for amino acids in the same biochemical group and low scores for amino acids from different groups.

Protein scoring matrices reflect the properties of amino acids



N.B. BLOUSM62 does not take the local context of a particular position into account

(i.e. all like substitutions are scored the same regardless of their location in the molecules).

We will revisit this later...

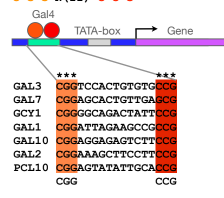
Todays Menu

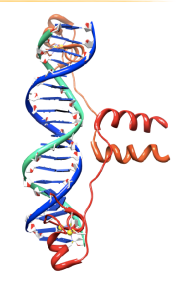
- Sequence motifs and patterns: Simple approaches for finding functional cues from conservation patterns
- Sequence profiles and position specific scoring matrices (PSSMs): Building and searching with profiles, Their advantages and limitations
- PSI-BLAST algorithm: Application of iterative PSSM searching to improve BLAST sensitivity
- Hidden Markov models (HMMs): More versatile probabilistic model for detection of remote similarities

Functional cues from conservation patterns...

Many DNA patterns are binding sites for Transcription Factors.

• E.g., The Gal4 binding sequence C-G-G-N(11)-C-C-G





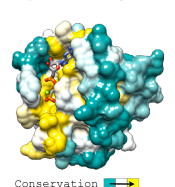
Functional cues from conservation patterns

Within a protein or nucleic acid sequence there may be a small number of characteristic residues that occur consistently. These conserved "sequence fingerprints" (or **motifs**) usually contain functionally important elements

 E.g., the amino acids that are consistently found at enzyme active sites or the nucleotides that are associated with transcription factor binding sites.

ATP/GTP-binding proteins: G-x(4)-G-K-T





Representing recurrent sequence patterns

Beyond knowledge of invariant residues we can define **position-based** representations that highlight the range of permissible residues per position.

 Pattern: Describes a motif using a qualitative consensus sequence (e.g., IUPAC or regular expression). N.B. Mismatches are not tolerated!

- Profile: Describes a motif using quantitative information captured in a position specific scoring matrix (weight matrix).
 Profiles quantify similarity and often span larger stretches of sequence.
- Logos: A useful visual representation of sequence motifs.



Image generated by: weblogo.berkeley.edu

PROSITE is a protein pattern and profile database

Currently contains > 1790 patterns and profiles: http://prosite.expasy.org/
Example PROSITE patterns:

PS00087; SOD_CU_ZN_1
[GA]-[IMFAT]-H-[LIVF]-H-{S}-x-[GP]-[SDG]-x-[STAGDE]
The two Histidines are copper ligands

- Each position in the pattern is separated with a hyphen
- · x can match any residue
- [] are used to indicate ambiguous positions in the pattern e.g., [SDG] means the pattern can match S, D, or G at this position
- {} are used to indicate residues that are not allowed at this position e.g., {S} means NOT S (not Serine)
- () surround repeated residues, e.g., A(3) means AAA

Information from http://ca.expasy.org/prosite/prosuser.html

Pattern advantages and disadvantages

Advantages:

- Relatively straightforward to identify (exact pattern matching is fast)
- · Patterns are intuitive to read and understand
- Databases with large numbers of protein (e.g., PROSITE) and DNA sequence (e.g., JASPER and TRANSFAC) patterns are available.

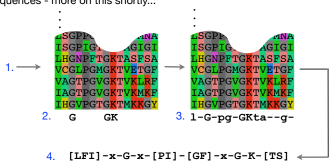
Disadvantages:

- Patterns are qualitative and deterministic (i.e., either matching or not!)
- We lose information about relative frequency of each residue at a position E.g., [GAC] vs 0.6 G, 0.28 A, and 0.12 C
- Can be difficult to write complex motifs using regular expression notation
- · Cannot represent subtle sequence motifs

Defining sequence patterns

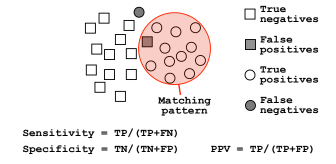
There are four basic steps involved in defining a new PROSITE style pattern:

- 1. Construct a multiple sequence alignment (MSA)
- 2. Identify conserved residues
- 3. Create a core sequence pattern (i.e. consensus sequence)
- 4. Expand the pattern to improve **sensitivity** and **specificity** for detecting desired sequences more on this shortly...



Side note: pattern sensitivity, specificity, and PPV

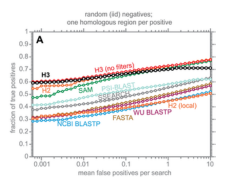
In practice it is not always possible to define one single regular expression type pattern which matches all family sequences (*true positives*) while avoiding matches in unrelated sequences (*true negatives*).



The positive predictive value (or PPV) assesses how big a proportion of the sequences matching the pattern are actually in the family of interest. (i.e., the probability that a positive result is truly positive!)

ROC plot example

ROC plot of sequence searching performance...



H3 (HMMER3) has a much higher search sensitivity and specificity than BLASTp

In each benchmark, true positive subsequences have been selected to be no more than 25% identical to any sequence in the query alignment ... (see paper for details).

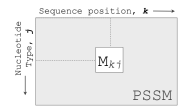
See: Eddy (2011) PLoS Comp Biol 7(10): e1002195

Sequence profiles

A sequence profile is a **position-specific scoring matrix** (or **PSSM**, often pronounced 'possum') that gives a *quantitative* description of a sequence motif.

Unlike deterministic patterns, profiles assign a score to a query sequence and are widely used for database searching.

A simple PSSM has as many columns as there are positions in the alignment, and either 4 rows (one for each DNA nucleotide) or 20 rows (one for each amino acid).



$$M_{kj} = \log\left(\frac{p_{kj}}{p_j}\right)$$

 \mathbf{M}_{kj} score for the *j*th nucleotide at position k \mathbf{p}_{kj} probability of nucleotide *j* at position k

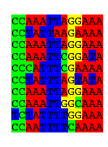
 p_j "background" probability of nucleotide j

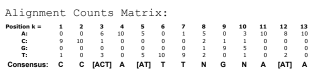
See Gibskov et al. (1987) PNAS 84, 4355

Todays Menu

- Sequence motifs and patterns: Simple approaches for finding functional cues from conservation patterns
- Sequence profiles and position specific scoring matrices (PSSMs): Building and searching with profiles, Their advantages and limitations
- PSI-BLAST algorithm: Application of iterative PSSM searching to improve BLAST sensitivity
- Hidden Markov models (HMMs): More versatile probabilistic model for detection of remote similarities

Computing a transcription factor bind site PSSM





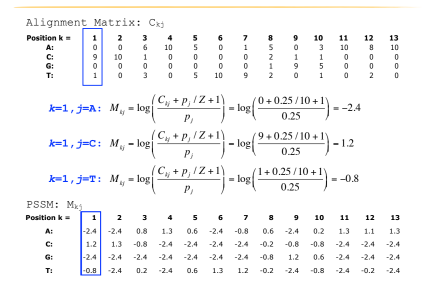
$$M_{kj} = \log\left(\frac{p_{kj}}{p_j}\right) \qquad p_{kj} = \frac{C_{kj} + p_j}{Z + 1}$$

$$M_{kj} = \log\left(\frac{C_{kj} + p_j / Z + 1}{p_j}\right)$$

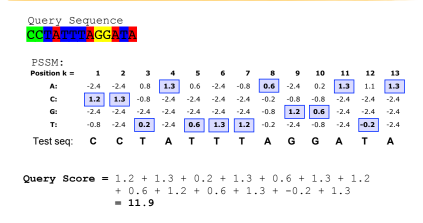
- C_{kj} Number of jth type nucleotide at position k
- Z Total number of aligned sequences
- pi "background" probability of nucleotide i
- \mathbf{p}_{kj} probability of nucleotide j at position k

Adapted from Hertz and Stormo, Bioinformatics 15:563-577

Computing a transcription factor bind site PSSM...

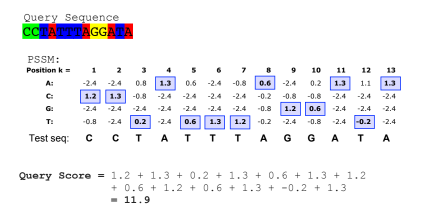


Scoring a test sequence

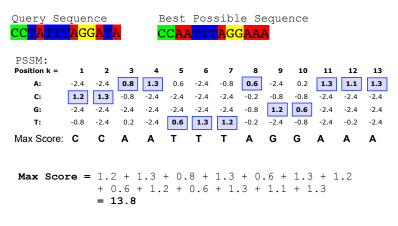


Q. Does the guery sequence match the DNA sequence profile?

Scoring a test sequence



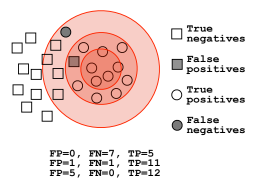
Scoring a test sequence...



A. Following method in Harbison *et al.* (2004) Nature 431:99-104 Heuristic threshold for match = 60% x Max Score = (0.6 x 13.8 = 8.28); 11.9 > 8.28; Therefore our query is a potential TFBS!

Picking a threshold for PSSM matching

Again, you want to select a threshold that **minimizes FPs** (e.g., how many shuffled or random sequences does the PSSM match with that score) and **minimizes FNs** (e.g., how many of the 'real' sequences are missed with that score).



Q. Which threshold has the best PPV (TP/(TP+FP))?

Side note: Building PSSMs from unaligned sequences

Patterns and profiles are most often built on the basis of known site equivalences (i.e. from a pre-calculated MSA).

However, a number of programs have been developed that employ local multiple alignments to search for common sequence elements in unaligned sequences.



Gibbs sampling methods:

Motif Sampler - http://bayesweb.wadsworth.org/gibbs/gibbs.html AlignAce - http://atlas.med.harvard.edu/cgi-bin/alignace.pl

Expectation maximization method: MEME - http://meme.sdsc.edu/

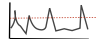
See: Lawrence et al. (1993) Science. 262, 208-14

Searching for PSSM matches

If we do not allow gaps (i.e., no insertions or deletions):

 Perform a linear scan, scoring the match to the PSSM at each position in the sequence - the "sliding window" method





If we allow gaps:

 Can use dynamic programming to align the profile to the protein sequence(s) (with gap penalties)

We will discuss PSI-BLAST shortly... see Mount, Bioinformatics: sequence and genome analysis (2004)

Can use hidden Markov Model-based methods

We will cover HMMs in the next lecture... see Durbin et al., Biological Sequence Analysis (1998)

Profiles software and databases

Pftools is a package to build and search with profiles,

http://www.isrec.isb-sib.ch/ftp-server/pftools/

The package contains (among other programs):

- pfmake for building a profile starting from multiple alignments
- pfsearch to search a protein database with a profile
- pfscan to search a profile database with a protein

PRINTS database of PSSMs

http://bioinf.man.ac.uk/dbbrowser/PRINTS

Collection of conserved motifs used to characterize a protein

- Uses fingerprints (conserved motif groups).
- Very good to describe sub-families.

BLOCKS is another PSSMs database similar to prints

http://www.blocks.fhcrc.org

ProDom is collection of protein motifs obtained automatically using PSI-BLAST http://prodes.toulouse.inra.fr/prodom/doc/prodom.html

Profiles software and databases...

InterPro is an attempt to group a number of protein domain databases. http://www.ebi.ac.uk/interpro

It currently includes:

- ▶ Pfam
- ▶ PROSITE
- ▶ PRINTS
- ▶ ProDom
- SMART
- ▶ TIGRFAMs
- InterPro tries to have and maintain a high quality of annotation
- The database and a stand-alone package (iprscan) are available for UNIX platforms, see:

ftp://ftp.ebi.ac.uk/pub/databases/interpro

Your Turn!

Hands-on sections 1 & 2:

Comparing methods and the trade-off between sensitivity, selectivity and performance

~50 mins

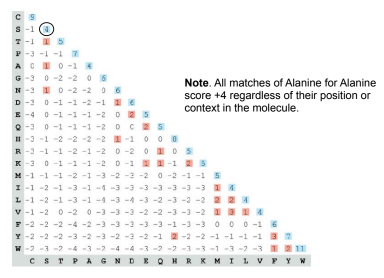
Todays Menu

- Sequence motifs and patterns: Simple approaches for finding functional cues from conservation patterns
- Sequence profiles and position specific scoring matrices (PSSMs): Building and searching with profiles, Their advantages and limitations
- PSI-BLAST algorithm: Application of iterative PSSM searching to improve BLAST sensitivity
- Hidden Markov models (HMMs): More versatile probabilistic model for detection of remote similarities

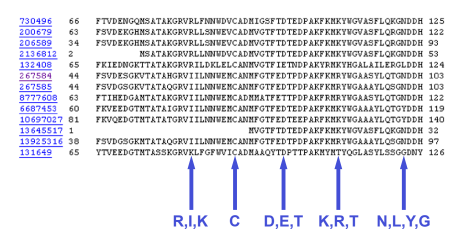
Recall: BLOUSM62 does not take the local context of a particular position into account

(i.e. all like substitutions are scored the same regardless of their location in the molecules).

By default BLASTp match scores come from the BLOSUM62 matrix



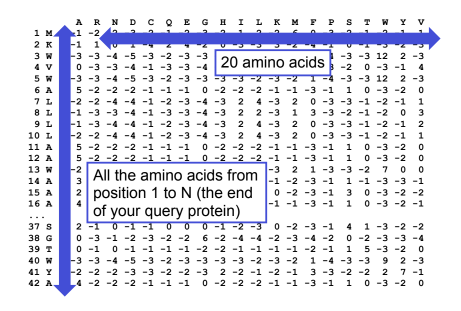
Inspect the blastp output to identify empirical "rules" regarding amino acids tolerated at each position

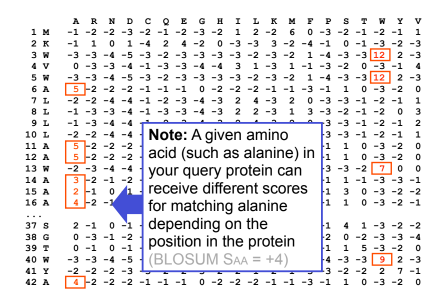


PSI-BLAST: Position specific iterated BLAST

- The purpose of PSI-BLAST is to look deeper into the database for matches to your query protein sequence by employing a scoring matrix that is customized to your query
 - PSI-BLAST constructs a multiple sequence alignment from the results of a first round BLAST search and then creates a "profile" or specialized position-specific scoring matrix (PSSM) for subsequent search rounds

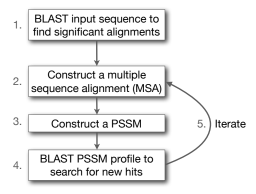
34



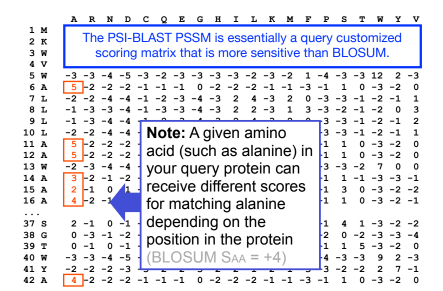


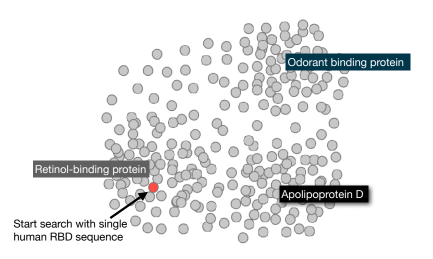
PSI-BLAST: Position-Specific Iterated BLAST

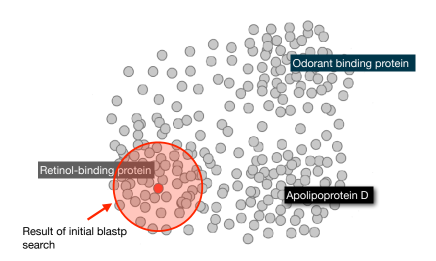
Many proteins in a database are too distantly related to a query to be detected using standard BLAST. In many other cases matches are detected but are so distant that the inference of homology is unclear. Enter the more sensitive PSI-BLAST

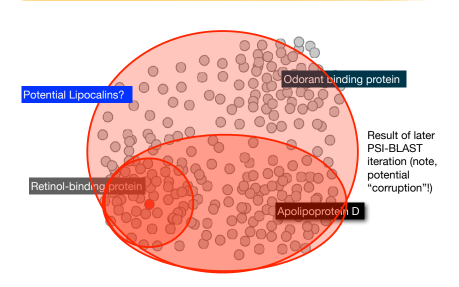


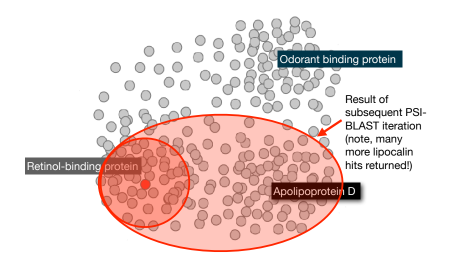
(see Altschul et al., Nuc. Acids Res. (1997) 25:3389-3402)











PSI-BLAST returns dramatically more hits

PSI-BLAST frequently returns many more hits with significant E-values than blastp

The search process is continued iteratively, typically about five times, and at each step a new PSSM is built.

 You must decide how many iterations to perform and which sequences to include!

You can stop the search process at any point - typically whenever few new results are returned or when no new "sensible" results are found.

Iteration	Hits with E < 0.005	Hits with E > 0.005
1	34	61
2	314	79
3	416	57
4	432	50
5	432	50

Human retinol-binding protein 4 (RBP4; P02753) was used as a query in a PSI-BLAST search of the RefSeq database.

```
(a) Iteration 1
        >ref|NP 001638.1| apolipoprotein D precursor [Homo sapiens] Length=T89
         Score = 57.4 bits (137), Expect = 3e-07, Method: Composition-based stats. Identities = 47/151 (31%), Positives = 78/151 (51%), Gaps = 39/151 (25%)
       Query 29 VKENFDKARFSGTWYAMAKKDPEGLFLQDNIVAEFSVDETGQMSATAKGRVRLLNNWDVC 88
       V+ENFD ++ G WY + +K P I A +S+ E G ++++LN ++
Sbjct 33 VQENFDVNKYLGRWYEI-EKIPTTFENGRCIQANYSLMENG------KIKVLNQ-ELR 82

        Query
        89
        ADMVGTFTDTE------DPAKFKMKY-WGVASFLQKGNDDHWIVDTDYDTYAVQYSC
        138

        AD GT
        E
        +PAK
        +K+ W
        S
        +WI+TDY+YALYYSC
        34

        Sbjct
        83
        AD--GTVNQISGEATPVALTPAKLEVKFSWFMPS-----PAPWILLATDYSNYALYYSC
        134
        134

       Query 139 ----RLINLDGTCADSYSFVFSRDPNGLPPE 165
        +L ++D ++++ +R+PN LPPE
Sbjct 135 TCIIQLFHVD-----FAWILARNPN-LPPE 158
(b) Iteration 2
       >ref|NP 001638.1| apolipoprotein D precursor [Homo sapiens] Length=T89  
         Score = 175 bits (443), Expect = 1e-42, Method: Composition-based stats. Identities = 45/163 (27%), Positives = 77/163 (47%), Gaps = 31/163 (19%)
      Query 14 GSGRAERDCRVSSFRVKENFDKARFSGTWYAMAKKDPEGLFLQDNIVAEFSVDETGQMSA 73
GYA + Y-VENED + GWY + K F I A 184 E G++
Sbjct 18 AEGQAFHGKCPNPFVQBNFDVNKYLGMNUEI-EKEPTFTENARCIQANYSLMENMSKKV 76
       Query 74 TAK----GRVRLLNNWDVCADMVGTFTDTEDPAKFKMKY-WGVASFLQKGNDDHWIVDT 127
       + G V + T + +PAK ++K+ W + S +WI+ T
Sbjct 77 LNQELRADGTVNQIEG-----EATPVNLTEPAKLEVKFSWFMPS-----APYWILAT 123
      Query 128 DYDTYAVQYSCR----LINLDGTCADSYSFVFSRDPNGLPPEA 166
DY+ YA+ YSC L ++D ++++ +R+PN LPPE
Sbjct 124 DYENYALVYSCTCIIQIFHVD-----FAWILARNPN-LPPET 159
(c) Iteration 3
       >ref[NP_000597.1| complement component 8, gamma polypeptide [Homo sapiens] Length=702
         Score = 104 bits (260), Expect = 2e-21, Method: Composition-based stats. Identities = 40/186 (21%), Positives = 74/186 (39%), Gaps = 29/186 (15%)
       Query 24 VSSFRVKENFDKARFSGTWYAMAKKDPEGLFLQDNIVAEFSVDETG-QMSATAKGRVRLL 82
       +S+ + K NFD +F+GTW +A + AE + Q +A A R L
Sbjct 33 ISTIQPKANFDAQQFAGTWLLVAVGSACRFLQEQGHRAEATTLHVAPQGTAMAVSTFRKL 92
      Query 83 NNWDVCADMVGTFTDTEDPAKFKMKYWGVASFLQKGNDDHWIVDTDYDTYAVQY----- 136
+ 1C + 1DT + F + G + HTDY + HAV Y
Sbjct 93 DG-ICWQVRQLVGDTGVLGRFLLQARGA----RGAVHVVVAETDYGSFAULYLERAGQ 145
      Query 137 -SCRLLNLDGTCADSYSFVFSRDPNGLPPEAQKIVRQRQEELCLARQYRLIVHNGYCDGR 195 S +L F +DS F + EA ++++ +Y G+C+ Sbjct 146 LSVKLVARSLPVSDSVLSGFEDRVQ---EA---HLTEDQTFYFFXY-----GFCEAA 191
       Query 196 SERNLL 201
```

blastp E-value for this hit was 0.27

Profile advantages and disadvantages

Advantages:

· Quantitate with a good scoring system

+ ++L Sbjct 192 DQFHVL 197

- Weights sequences according to observed diversity Profile is specific to input sequence set
- Very sensitive
 Can detect weak similarity
- Relatively easy to compute
 Automatic profile building tools available

Disadvantages:

- If a mistake enters the profile, you may end up with irrelevant data The corruption problem!
- Ignores higher order dependencies between positions
 i.e., correlations between the residue found at a given position and those found
 at other positions (e.g. salt-bridges, structural constraints on RNA etc...)
- · Requires some expertise and oversight to use proficiently

PSI-BLAST errors: the corruption problem

The main source of error in PSI-BLAST searches is the spurious amplification of sequences that are unrelated to the query.

There are three main approaches to stopping corruption of PSI-BLAST queries:

- Perform multi-domain splitting of your query sequence
 If a query protein has several different domains PSI-BLAST may find database matches related to both individually. One should not conclude that these hits with different domains are related.
 - Often best to search using just one domain of interest.
- Inspect each PSI-BLAST iteration removing suspicious hits.
 E.g., your query protein may have a generic coiled-coil domain, and this may cause other proteins sharing this motif (such as myosin) to score better than the inclusion threshold even though they are not related.
 - Use your biological knowledge!
- Lower the default expect level (e.g., E = 0.005 to E = 0.0001).
 This may suppress appearance of FPs (but also TPs)

Todays Menu

- Sequence motifs and patterns: Simple approaches for finding functional cues from conservation patterns
- Sequence profiles and position specific scoring matrices (PSSMs): Building and searching with profiles, Their advantages and limitations
- PSI-BLAST algorithm: Application of iterative PSSM searching to improve BLAST sensitivity
- Hidden Markov models (HMMs): More versatile probabilistic model for detection of remote similarities



Problems with PSSMs: Positional dependencies

Do not capture positional dependencies

WEIRD
WEIRD
WEIRD
WEIRD
WEIOH

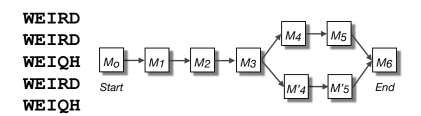
D					0.6
ш		—			
Η					0.4
I			I		
Q				0.4	
R				0.6	
W	I				

Note: We <u>never</u> see **QD** or **RH**, we only see **RD** and **QH**. However, P(RH)=0.24, P(QD)=0.24, while P(QH)=0.16

Markov chains: Positional dependencies



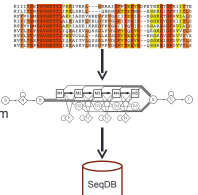
The connectivity or **topology** of a Markov chain can easily be designed to capture dependencies and variable length motifs.



Recall that a PSSM for this motif would give the sequences **WEIRD** and **WEIRH** equally good scores even though the **RH** and **QR** combinations were not observed

Use of HMMER

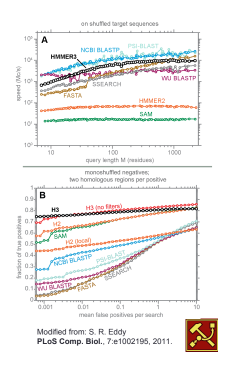
- Widely used by protein family databases
 - Use 'seed' alignments
- Until 2010
 - Computationally expensive
 - Restricted to HMMs constructed from multiple sequence alignments
- Command line application





HMMER vs BLAST

	HMMER	BLAST	
Progra m	P HMM E R	BLASTP	
Quer y	Single sequence		
Target Databas e	Sequenc e databas e		
Progra m	HMM SCA N	RP SB LA S T	
Quer y	Single sequence		
Target Database	Profil e HM M database, e.g. Pfa m	PSSM database, e.g. CD D	
Progra m	HM M SE ARCH	PSI-BLAS T	
Quer y	Profil e HM M	PSS M	
Targe t Databas e	Sequenc e databas e		
Progra m	JACKHMMER	PSI-BLAS T	
Quer y	Single sequence		
Targe t Databas e	Sequenc e databas e		



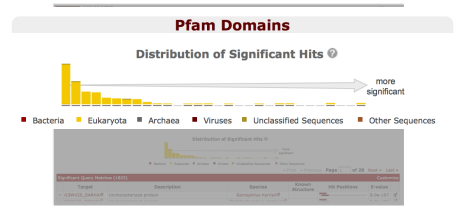
Fast Web Searches

- · Parallelized searches across compute farm
 - Average query returns ~1 sec
- Range of sequence databases
 - Large Comprehensive
 - Curated / Structure
 - Metagenomics
 - Representative Proteomes
- Family Annotations
 - Pfam
- Batch and RESTful API
 - Automatic and Human interface

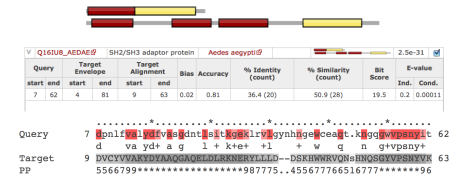




Visualization of Results – By Score



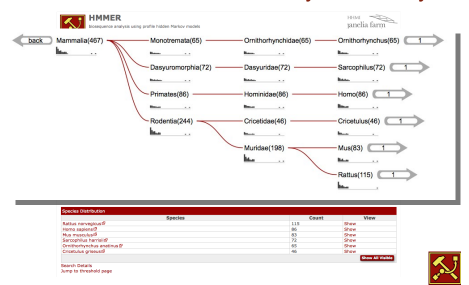
Visualization of Results – By Score







Visualization of Results – By Taxonomy



PFAM: Protein Family Database of Profile HMMs

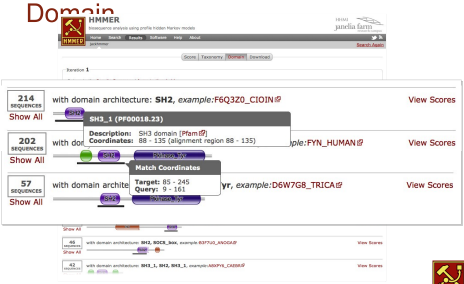
Comprehensive compilation of both multiple sequence alignments and profile HMMs of protein families.

http://pfam.sanger.ac.uk/

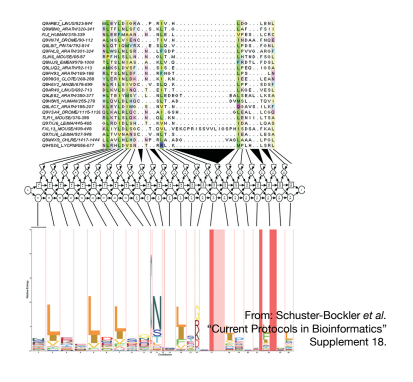
PFAM consists of two databases:

- Pfam-A is a manually curated collection of protein families in the form of multiple sequence alignments and profile HMMs. HMMER software is used to perform searches.
- **Pfam-B** contains additional protein sequences that are automatically aligned. Pfam-B serves as a useful supplement that makes the database more comprehensive.
- Pfam-A also contains higher-level groupings of related families, known as clans

Visualization of Results – By







HMM limitations

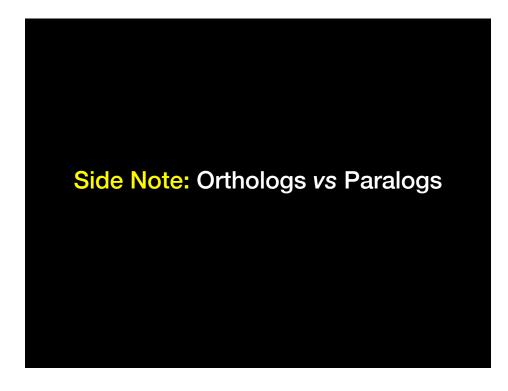
HMMs are linear models and are thus **unable to capture higher order correlations** among positions (e.g. distant cysteins in a disulfide bridge, RNA secondary structure pairs, etc).

Another flaw of HMMs lies at the very heart of the mathematical theory behind these models. Namely, that the probability of a sequence can be found from the product of the probabilities of its individual residues.

This claim is only valid if the probability of a residue is independent of the probabilities of its neighbors. In biology, there are frequently **strong dependencies between these probabilities** (e.g. hydrophobic residues clustering at the core of protein domains).

These biological realities have motivated research into new kinds of statistical models. These include hybrids of HMMs and neural nets, dynamic Bayesian nets, factorial HMMs, Boltzmann trees and stochastic context-free grammars.

See: Durbin et al. "Biological Sequence Analysis"



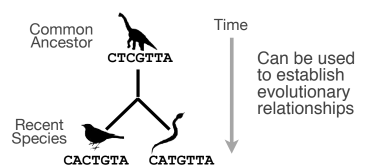


That's it!

Sequence comparison is most informative when it detects **homologs**

Homologs are sequences that have common origins *i.e.* they share a **common ancestor**

They may or may not have common activity



Key terms

When we talk about related sequences we use specific terminology.

Homologous sequences may be either:

Orthologs or Paralogs(Note. these are all or nothing relationships!)

Any pair of sequences may share a certain level of:

Identity and/or Similarity
 (Note. if these metrics are above a certain level we often infer homology)

Paralogs tend to have slightly different functions

Paralogs: are homologs produced by **gene duplication**. They represent genes derived from a common ancestral gene that <u>duplicated within an organism</u> and then subsequently <u>diverged by accumulated mutation</u>.

- Para = [greek: along side of]

Single Species CTCGTTA Duplication

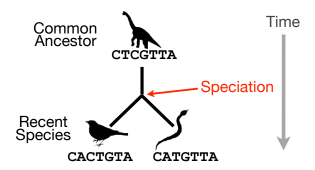
CACGTTA Divergence

CACTGTA CATGTTA

Orthologs tend to have similar function

Orthologs: are homologs produced by <u>speciation</u> that have diverged due to divergence of the organisms they are associated with.

- Ortho = [greek: straight] ... implies direct descent



Orthologs vs Paralogs

- In practice, determining ortholog *vs* paralog can be a complex problem:
 - gene loss after duplication,
 - lack of knowledge of evolutionary history,
 - weak similarity because of evolutionary distance
- Homology does not necessarily imply exact same function
 - may have similar function at very crude level but play a different physiological role

66

67