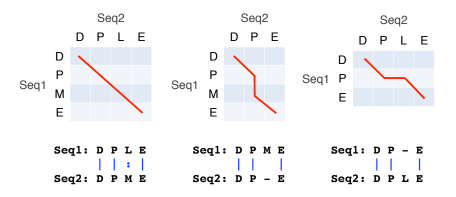


# Muddy Point: Different paths represent different alignments



(Mis)matches are represented by <u>diagonal paths</u> & Indels with <u>horizontal or vertical path</u> segments

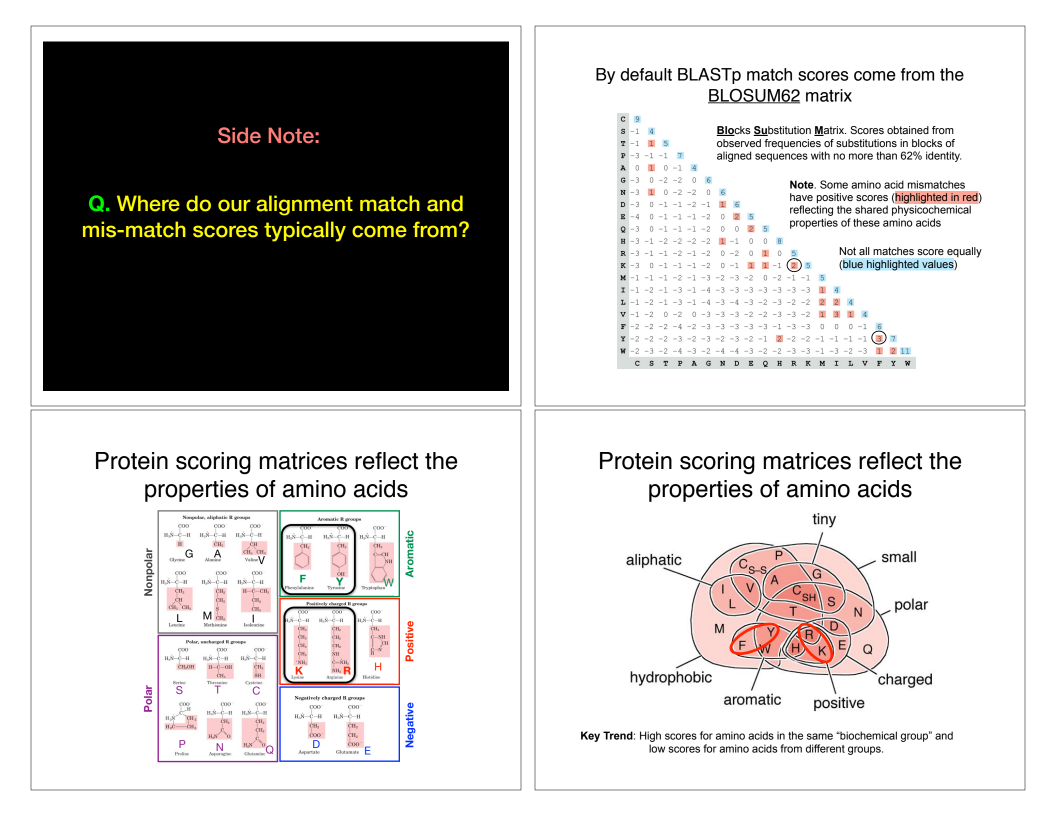
### **Recap From Last Time:**

- 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.

**Feedback** 

## 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



**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

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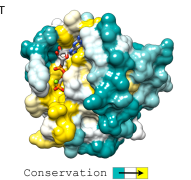
#### Functional cues from conservation patterns

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

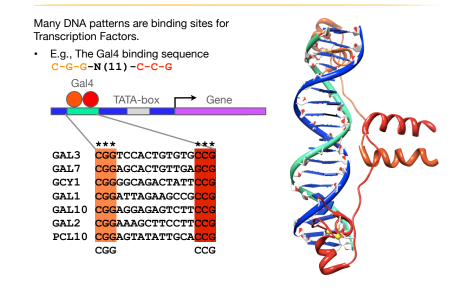
• E.g., the amino acids that are consistently found at enzyme active sites or the nucleotides that are associated with transcription factor binding sites.

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





Functional cues from conservation patterns...



#### 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!

[LFI]-x-G-[PT]-P-G-x-G-K-[TS]-[AGSI]

- 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.

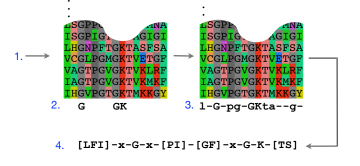


Image generated by: weblogo.berkeley.edu

#### Defining sequence patterns

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

- 1. Construct a <u>multiple sequence alignment</u> (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...



#### PROSITE is a protein pattern and profile database

Currently contains > 1790 patterns and profiles: <u>http://prosite.expasy.org/</u> 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:

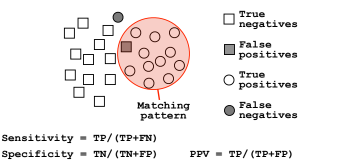
- Relatively straightforward to identify (exact pattern matching is fast)
- · Patterns are intuitive to read and understand
- Databases with large numbers of protein (e.g., PROSITE) and DNA sequence (e.g., JASPER and TRANSFAC) patterns are available.

#### **Disadvantages:**

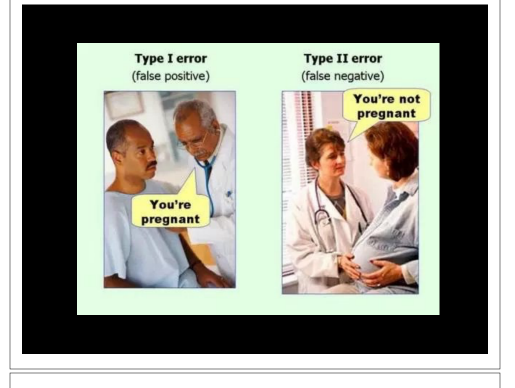
- Patterns are qualitative and *deterministic* (i.e., either matching or not!)
- We lose information about relative frequency of each residue at a position E.g.,  $\,$  [GAC]  $\,vs\,$  0.6 G, 0.28 A, and 0.12 C  $\,$
- · Can be difficult to write complex motifs using regular expression notation
- ·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

#### Side note: pattern sensitivity, specificity, and PPV

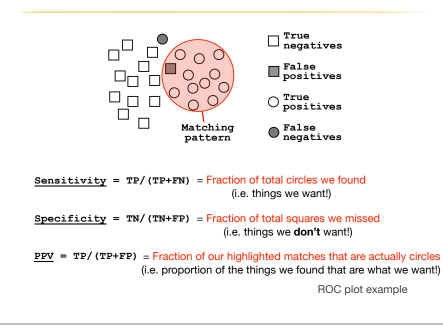
In practice it is not always possible to define one single regular expression type pattern which matches all family sequences (*true positives*) while avoiding matches in unrelated sequences (*true negatives*).



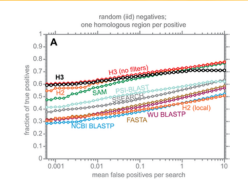
The positive predictive value (or PPV) assesses how big a proportion of the sequences matching the pattern are actually in the family of interest. (i.e., the probability that a positive result is truly positive!)



#### Side note: pattern sensitivity, specificity, and PPV



#### 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

## Todays Menu

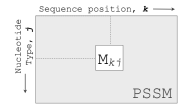
- Sequence motifs and patterns: Simple approaches for finding functional cues from conservation patterns
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#### Sequence profiles

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).





 $M_{kj}$  score for the *j*th nucleotide at position *k*   $p_{kj}$  probability of nucleotide *j* at position *k*  $p_1$  "background" probability of nucleotide *j* 

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

#### Computing a transcription factor bind site PSSM

AAA <mark>TT</mark> AGGAAA														
TA <mark>TTAA</mark> GAAAA														
AAA <mark>TT</mark> AGGAAA	Alignme	nt	Со	unts	M	atri	x:							
AAA <mark>TT<mark>C</mark>GGATA</mark>	Position k =	1	2	3	4	5	6	7	8	9	10	11	12	13
CATTTCGAAAA	A:	0	0	6	10	5	0	1	5	0	3	10	8	10
ATTTAGTATA	C:	9	10	1	0	0	0	0	2	1	1	0	0	0
ATTIAGIAIA	G:	0	0	0	0	0	0	0	1	9	5	0	0	0
AAATTAGGAAA	T:	1	0	3	0	5	10	9	2	0	1	0	2	0
AAA <mark>TT<mark>GGC</mark>AAA</mark>	Consensus:	с	С	[ACT]	Α	[AT]	т	т	Ν	G	Ν	Α	[AT]	Α

$M_{kj} = \log\left(\frac{p_{kj}}{r}\right)$	$p = \frac{C_{kj} + p_j}{2}$
$M_{kj} = \log\left(\frac{1}{p_i}\right)$	$P_{kj} = \overline{Z+1}$

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

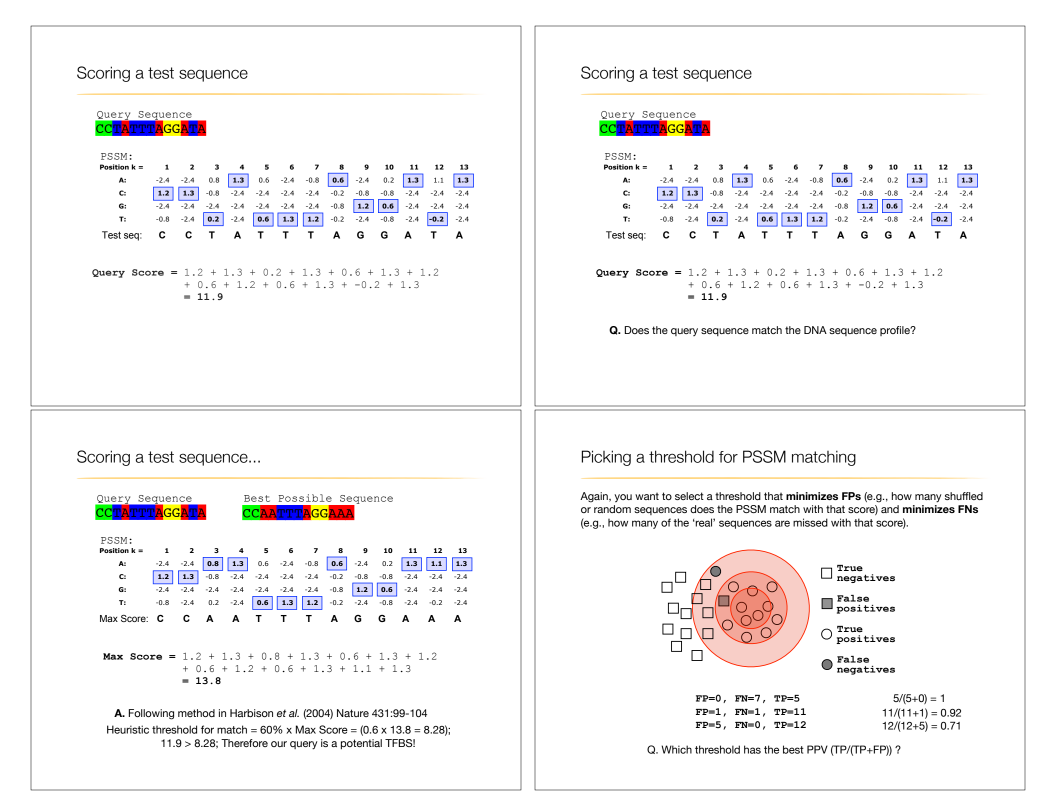
Ckj	Number of <i>j</i> th type nucleotide at position $k$

- **Z** Total number of aligned sequences
- **p**<sub>j</sub> "background" probability of nucleotide *j*
- $\mathbf{p}_{kj}$  probability of nucleotide *j* at position *k*

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

#### Computing a transcription factor bind site PSSM...

Position k =	1	2	3	4	5	6	7	8	9	10	11	12	13
A:	0	0	6	10	5	0	1	5	0	3	10	8	10
C:	9	10	1	0	0	0	0	2	1	1	0	0	(
G:	0	2 0 10 0 0	0	0	0	0	0	1	9	5	0	0	
т:	1	0	3	0	5	10	9	2	0	1	0	2	
<b>k=1</b> ,				`	- )								
<b>k=1</b> ,	j=C:	$M_{_{kj}}$	$= \log \left( \frac{1}{2} \right)$	$C_{kj}$ +	$p_j / Z$	$\frac{2+1}{2}$	= log	<u>9+0</u>	.25/1	$\frac{0+1}{}$	= 1.2		
			,	(	1 )	/		`	0.25				
<b>k=1</b> ,													
<b>k=1</b> ,	<b>j=T :</b>	$M_{kj}$	= log	$\frac{C_{kj}}{C_{kj}}$ +	$\frac{p_j}{p_j}$	$\left(\frac{2+1}{2}\right)$	= log	(1+0	.25 / 10 0.25				
<b>k=1</b> ,	<b>j=T :</b>	$M_{kj}$	= log	$\frac{C_{kj}}{C_{kj}}$ +	$\frac{p_j}{p_j}$	$\left(\frac{2+1}{2}\right)$	= log	(1+0	.25 / 10 0.25				1
<b>k=1</b> ,	<b>j=T :</b>	$M_{kj}$	= log	$\frac{C_{kj}}{C_{kj}}$ +	$\frac{p_j}{p_j}$	$\left(\frac{2+1}{2}\right)$	= log	(1+0	.25 / 10 0.25	$\left(\frac{D+1}{2}\right)$	= -0.3	8	_
<b>k=1</b> ,	<b>j=T :</b>	$M_{kj}$	= log	$\frac{C_{kj}}{C_{kj}}$ +	$\frac{p_j}{p_j}$	$\left(\frac{2+1}{2}\right)$	= log	(1+0	.25 / 10 0.25	<u>0 + 1</u> ) 10	= -0.3 11	8 12	1.
<b>k=1</b> ,	<b>j=T :</b>		= log	$\frac{C_{kj}}{C_{kj}}$ +	$\frac{p_j}{p_j}$	$\left(\frac{2+1}{2}\right)$	= log	(1+0	.25 / 10 0.25	$\frac{10}{10}$	= -0.3 11 1.3	8 12 1.1	<b>1</b> 1. -2.



#### 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) Side note: 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:

- **b** Pfam
- PROSITE
- PRINTS
- ProDom SMART
- TIGREAMs
- 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

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Hands-on sections 1 & 2:

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

~50 mins

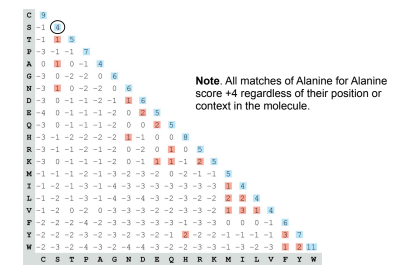
**Recall:** BLOUSM62 does not take the local context of a particular position into account

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

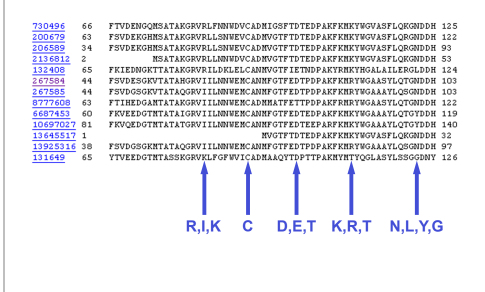
### PSI-BLAST: Position specific iterated BLAST

- 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

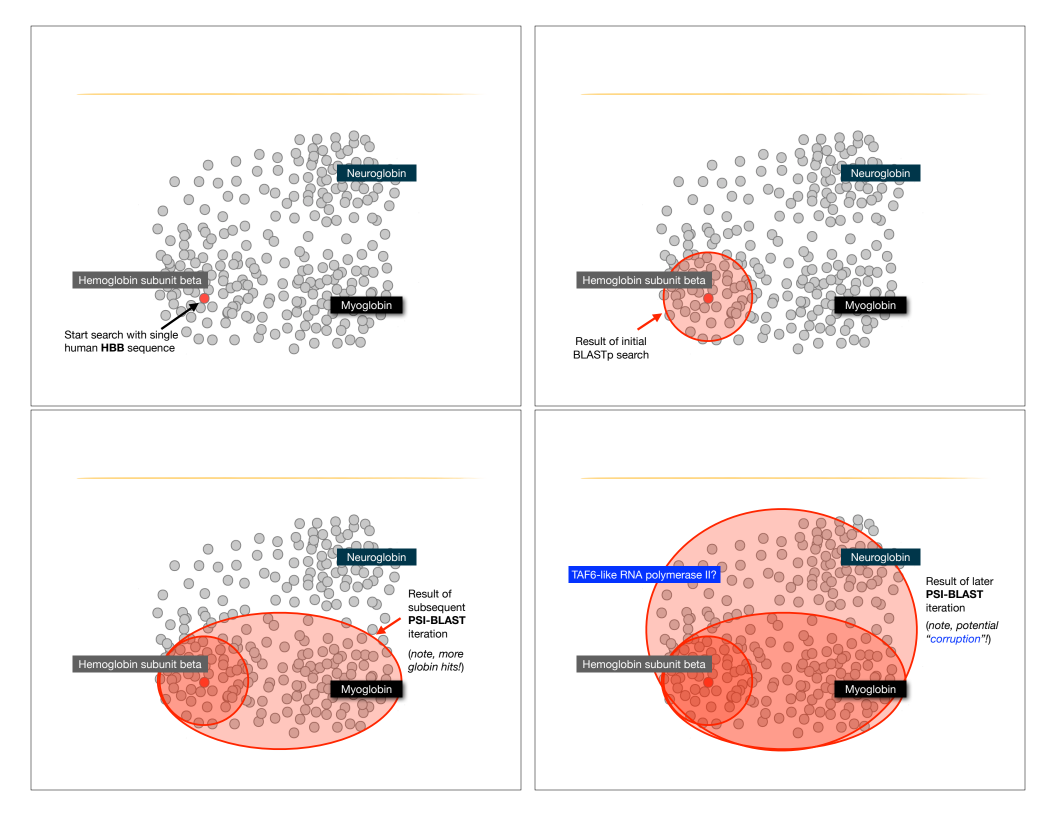
## By default BLASTp match scores come from the BLOSUM62 matrix



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



A R N D C Q E G H I L K M F P S T W Y V 1 M 1 -2 2 2 3 2 1 2 2 3 3 3 2 4 1 0 1 2 3 2 1 0 1 0 1 -3 2 -3 3 W -3 -3 -4 -5 -3 -2 -3 -3 4 V 0 -3 -3 -4 -1 3 -3 -4 5 W -3 -3 -4 -5 -3 -2 -3 -3 6 A 5 -2 -2 -2 -1 -1 -1 0 -2 -2 -2 -2 -1 -4 -3 -3 12 2 -3 6 A 5 -2 -2 -2 -1 -1 -1 0 -2 -2 -2 -2 -1 -4 -3 -3 12 2 -3 6 A 5 -2 -2 -2 -1 -1 -1 0 -2 -2 -2 -2 -1 -4 -3 -3 12 2 -3 6 A 5 -2 -2 -2 -1 -1 -1 0 -2 -2 -2 -2 -1 -1 -3 -1 1 0 -3 -2 0 7 L -2 -2 4 -4 -1 -2 -3 -4 -3 2 4 -3 2 2 4 -3 2 0 -3 -3 -1 -2 -1 1 8 L -1 -3 -3 -4 -1 -2 -3 -4 -3 2 2 4 -3 2 2 0 -3 -3 -1 -2 -1 2 10 L -2 -2 4 -4 -1 -2 -3 -4 -3 2 2 4 -3 2 0 -3 -3 -1 -2 -1 2 10 L -2 -2 -4 -4 -1 -2 -3 -4 -3 2 4 -3 2 2 0 -3 -3 -1 -2 -1 1 11 A 5 -2 -2 -2 -1 -1 -1 0 -2 -2 -2 -2 -1 -1 -3 -1 1 0 -3 -2 0 12 A 5 -2 -2 -2 -1 -1 -1 0 -2 -2 -2 -2 -1 -1 -3 -1 1 0 -3 -2 0 13 W -2 All the amino acids from 15 A 2 16 G 0 -3 -1 -2 -3 -2 -2 -3 -3 -3 -3 -2 -3 -1 4 1 -3 -3 -1 17 -3 -3 -1 2 -1 -1 0 -2 -2 -2 -2 -1 -1 -3 -1 1 0 -3 -2 -2 38 G 0 -3 -1 -2 -3 -2 -2 6 -2 -4 -4 -2 -3 -4 -2 0 -2 -3 -1 4 1 -3 -2 -2 38 G 0 -3 -1 -2 -3 -2 -2 6 -2 -4 -4 -2 -3 -4 -2 0 -2 -3 -1 4 1 -3 -2 -2 39 T 0 -1 0 -1 -1 -1 0 0 0 -1 -2 -3 -3 0 -2 -3 -1 4 1 -3 -2 -2 38 G 0 -3 -1 -2 -3 -2 -2 6 -2 -4 -4 -2 -3 -2 -3 -1 4 1 -3 -3 -2 0 40 W -3 -3 -4 -5 -3 -2 -3 -3 -3 -3 -2 -3 -2 -1 -1 -1 1 0 -3 -2 0 40 W -3 -3 -4 -5 -3 -2 -3 -3 -3 -3 -2 -2 -3 -2 -1 -1 -1 1 0 -3 -2 0 41 Y -2 -2 -2 -2 -1 -1 -1 0 0 -2 -2 -2 -2 -1 -1 -1 1 0 -3 -2 0 41 Y -2 -2 -2 -3 -3 -2 -3 -3 -3 -3 -3 -2 -3 -2 -3 -2 -1 -2 -3 -1 -2 -3 -2 -3 -2 -2 -3 -1 -2 -1 -1 -1 -3 -3 -2 0 41 Y -2 -2 -2 -3 -3 -2 -3 -3 -3 -3 -2 -3 -2 -3 -2 -3 -2 -2 -3 -2 -2 -3 -2 -2 -3 -2 -3 -2 -2 -3 -2 -2 -2 -3 -2 -2 -3 -2 -2 -3 -2 -2 -2 -3 -2 -2 -2 -1 -1 -1 -1 0 -2 -2 -2 -2 -2 -1 -1 -1 -1 0 -3 -2 0	A R N D C Q E G H I L K M F P S T W Y V 1 M $-1 -2 -2 -3 -2 -1 -2 -3 -2 1 2 -2 6 0 -3 -2 -1 -2 -1 1$ 2 K $-1 1 0 1 -4 2 4 -2 0 -3 -3 3 -2 -4 -1 0 -1 -3 -2 -3$ 3 W $-3 -3 -4 -5 -3 -2 -3 -3 -3 -3 -2 -3 -2 1 -4 -3 -3 12 2 -3$ 4 V $0 -3 -3 -4 -5 -3 -2 -3 -3 -3 -3 -2 -3 -2 1 -4 -3 -3 12 2 -3$ 6 A $5 -2 -2 -2 -1 -1 -1 0 -2 -2 -2 -1 -1 -3 -1 1 0 -3 -2 0$ 7 L $-2 -2 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -3 -2 -3 -2 1 -4 -3 -3 12 2 -3$ 8 L $-1 -3 -3 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -1 -2 -1 1$ 8 L $-1 -3 -3 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -2 -2 -1 -2 0 3$ 9 L $-1 -3 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -2 -1 -2 0 3$ 9 L $-1 -3 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -2 -1 -2 0 3$ 9 L $-1 -3 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -2 -1 -2 0 3$ 9 L $-1 -3 -4 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -2 -1 -2 0 3$ 9 L $-1 -3 -4 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -3 -2 -1 -2 0 3$ 9 L $-1 -3 -4 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -2 -1 -2 0 3$ 9 L $-1 -3 -4 -4 -4 -1 -2 -3 -4 -3 2 2 -3 -3 -3 -1 -2 -1 1 -3 -1 -1 0 -3 -2 0$ 10 L $-2 -2 -4 -4 -4 -1 -2 -3 -1 -1 0 -3 -2 0$ 11 U $-3 -2 -2 -2 -2 -1 -1 -1 0 -2 -2 -2 -2 -1 -1 -3 -1 1 0 -3 -2 0$ 12 A $5 -2 -2 -2 -2 -1 -1 -1 0 -2 -2 -2 -2 -1 -1 -3 -1 1 0 -3 -2 0$ 13 W $-2 -3 -3 -4 -4 -4 -2 -1 -1 -1 0 -2 -2 -2 -2 -1 -1 -3 -1 1 0 -3 -2 0$ 14 L $-3 -3 -2 -2 -2 -3 -3 -4 -4 -3 -2 -2 -3 -3 -4 -3 -3 -3 -1 -2 -1 -3 -1 -1 0 -3 -2 0$ 14 L $-3 -3 -2 -2 -2 -3 -3 -4 -4 -3 -3 -3 -1 -2 -2 -2 -2 -3 -3 -4 -3 -3 -3 -1 -2 -2 -2 -2 -3 -3 -4 -3 -3 -3 -1 -2 -3 -3 -4 -3 -3 -3 -3 -3 -3 -3 -3 -3 -3 -3 -3 -3 $
A R N D C Q E G H I L K M F P S T W Y V The PSI-BLAST PSSM is essentially a query customized scoring matrix that is more sensitive than BLOSUM. 4 V 5 W $-3 -3 -4 -5 -3 -2 -3 -3 -3 -3 -2 -3 -2 -1 -4 -3 -3 12 -2 -3 -3 -3 -4 -5 -3 -2 -2 -2 -1 -1 -3 -1 -1 -0 -3 -2 0 -7 -1 -2 -2 -4 -4 -1 -2 -3 -4 -3 -2 -2 -2 -1 -1 -3 -1 -1 -0 -3 -2 0 -7 -1 -2 -2 -4 -4 -1 -2 -3 -4 -3 -2 -2 -3 -3 -3 -1 -2 -1 -1 -3 -1 -1 -3 -3 -4 -1 -3 -3 -4 -3 -2 -2 -2 -3 -3 -3 -1 -2 -1 -2 -1 -3 -3 -4 -1 -3 -3 -4 -3 -2 -2 -3 -3 -3 -1 -2 -1 -2 -1 -3 -3 -4 -1 -3 -3 -4 -4 -1 -3 -3 -4 -3 -2 -2 -3 -3 -3 -3 -1 -2 -1 -2 -1 -3 -3 -1 -2 -1 -2 -1 -3 -1 -3 -1 -2 -2 -2 -2 -2 -1 -1 -3 -1 -2 -2 -2 -2 -2 -1 -1 -3 -1 -2 -2 -2 -2 -2 -2 -1 -1 -3 -1 -1 -2 -2 -2 -2 -2 -2 -1 -1 -3 -1 -1 -0 -3 -2 -0 -2 -2 -2 -2 -1 -1 -3 -1 -1 -0 -3 -2 -0 -2 -2 -2 -2 -1 -1 -3 -1 -1 -0 -3 -2 -0 -2 -2 -2 -2 -1 -1 -3 -1 -1 -0 -3 -2 -0 -3 -2 -2 -2 -2 -1 -1 -3 -1 -1 -3 -2 -0 -3 -2 -1 -2 -2 -2 -2 -2 -2 -1 -1 -3 -1 -1 -2 -2 -2 -2 -2 -2 -2 -2 -2 -2 -2 -2 -2 $	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 1. BLAST input sequence to find significant alignments 2. Construct a multiple sequence alignment (MSA) 3. Construct a PSSM 4. BLAST PSSM profile to search for new hits (see Altschul <i>et al.</i> , Nuc. Acids Res. (1997) 25:3389-3402)



Description	Max score	Total score	Query cover	E value	Ident	Accession
hemoglobin subunit beta [Homo sapiens]	301	301	100%	2e-106	100%	NP_000509.1
hemoglobin subunit delta [Homo sapiens]	284	284	100%	7e-100	93%	NP_000510.1
hemoglobin subunit epsilon [Homo sapiens]	240	240	100%	2e-82	76%	NP_005321.1
hemoglobin subunit gamma-2 [Homo sapiens]	235	235	100%	2e-80	73%	NP_000175.1
hemoglobin subunit gamma-1 [Homo sapiens]	232	232	100%	3e-79	73%	NP_000550.2
hemoglobin subunit alpha [Homo sapiens]	114	114	97%	7e-33	43%	NP_000508.1
hemoglobin subunit zeta [Homo sapiens]	100	100	97%	3e-27	36%	NP_005323.1

Description	Max score	Total score	Query cover	E value	Ident	Accession
hemoglobin subunit beta [Homo sapiens]	301	301	100%	2e-106	100%	NP_000509.1
hemoglobin subunit delta [Homo sapiens]	284	284	100%	7e-100	93%	NP_000510.1
hemoglobin subunit epsilon [Homo sapiens]	240	240	100%	2e-82	76%	NP_005321.1
hemoglobin subunit gamma-2 [Homo sapiens]	235	235	100%	2e-80	73%	<u>NP_000175.1</u>
hemoglobin subunit gamma-1 [Homo sapiens]	232	232	100%	3e-79	73%	NP_000550.2
hemoglobin subunit alpha [Homo sapiens]	114	114	97%	7e-33	43%	NP_000508.1
hemoglobin subunit zeta [Homo sapiens]	100	100	97%	3e-27	36%	NP_005323.1
myoglobin [Homo sapiens]	80.5	5 80.5	5 97%	2e-19	26%	ND 005250 1
<u>myogroun (nomo sapiens)</u>	80.5	80.5	5 97%	28-19	20%	NP_005359.1
neuroglobin [Homo sapiens]	54.7	<b>7</b> 54.7	92%	2e-09	23%	NP_067080.1

New relevant globins found only by PSI-BLAST

Description	Max score	Total score	Query cover	E value	Ident	Accession
hemoglobin subunit beta [Homo sapiens]	301	301	100%	2e-106	100%	NP_000509.1
hemoglobin subunit delta [Homo sapiens]	284	284	100%	7e-100	93%	NP_000510.1
hemoglobin subunit epsilon [Homo sapiens]	240	240	100%	2e-82	76%	NP_005321.1
hemoglobin subunit gamma-2 [Homo sapiens]	235	235	100%	2e-80	73%	NP_000175.1
hemoglobin subunit gamma-1 [Homo sapiens]	232	232	100%	3e-79	73%	NP_000550.2
hemoglobin subunit alpha [Homo sapiens]	114	114	97%	7e-33	43%	NP_000508.1
hemoglobin subunit zeta [Homo sapiens]	100	100	97%	3e-27	36%	NP_005323.1
myoglobin [Homo sapiens]	80.5	80.5	97%	2e-19	26%	NP_005359.1
neuroglobin [Homo sapiens]	54.7	54.7	92%	2e-09	23%	NP_067080.1
					<u> </u>	
myoglobin [Homo sapiens]	159	159	97%			NP_005359.1
hemoglobin subunit alpha [Homo sapiens]	151	151	97%	3e-47	42%	NP_000508.1
hemoglobin subunit mu [Homo sapiens]	147	147	97%	6e-46	35%	NP_001003938.1
hemoglobin subunit theta-1 [Homo sapiens]	147	147	97%	2e-45	37%	NP_005322.1
neuroglobin [Homo sapiens]	134	134	92%	3e-40	23%	NP_067080.1
PREDICTED: cytoglobin isoform X2 [Homo sapiens]	115	115	66%	3e-33	25%	XP_016879605.1
PREDICTED: microtubule cross-linking factor 1 isoform X1 [Homo sapie	46.3	46.3	27%	7e-06	39%	XP_011523942.1
PREDICTED: microtubule cross-linking factor 1 isoform X4 [Homo sapie	46.3	46.3	27%	7e-06	39%	XP_005258156.1
Score and E value dep	bend	s on	PSS	Μ		

Description	Max score	Total score	Query cover	E value	Ident	Accession	
hemoglobin subunit beta [Homo sapiens]	301	301	100%	2e-106	100%	NP_000509.1	
hemoglobin subunit delta [Homo sapiens]	284	284	100%	7e-100	93%	NP_000510.1	
hemoglobin subunit epsilon [Homo sapiens]	240	240	100%	2e-82	76%	NP_005321.1	
hemoglobin subunit gamma-2 [Homo sapiens]	235	235	100%	2e-80	73%	NP_000175.1	1
hemoglobin subunit gamma-1 [Homo sapiens]	232	232	100%	3e-79	73%	NP_000550.2	
hemoglobin subunit alpha [Homo sapiens]	114	114	97%	7e-33	43%	NP_000508.1	
hemoglobin subunit zeta [Homo sapiens]	100	100	97%	3e-27	36%	NP_005323.1	
myoglobin [Homo sapiens]	80.5	80.5	97%	2e-19	26%	NP_005359.1	2
neuroglobin [Homo sapiens]	54.7	54.7	92%	2e-09	23%	NP_067080.1	2
mvoglobin [Homo sapiens]	159	159	97%	3e-50	26%	NP 005359.1	
hemoglobin subunit alpha [Homo sapiens]	151	151	97%			NP 000508.1	
hemoglobin subunit mu [Homo sapiens]	147	147	97%			NP 001003938.1	
hemoglobin subunit theta-1 [Homo sapiens]	147	147	97%			NP 005322.1	
neuroglobin [Homo sapiens]	134	134	92%			NP 067080.1	3
PREDICTED: cvtoglobin isoform X2 [Homo sapiens]	115	115	66%			XP 016879605.1	
						_	
PREDICTED: microtubule cross-linking factor 1 isoform X1 [Homo sapie	46.3	46.3	27%	7e-06	39%	XP_011523942.1	4
PREDICTED: microtubule cross-linking factor 1 isoform X4 [Homo sapie	46.3	46.3	27%	7e-06	39%	XP_005258156.1	

nclusion of irrelevant hits can lead to PSSM corruption

#### PSI-BLAST is performed in five steps

- A normal blastp search uses a scoring matrix (e.g., BLOSUM62) to perform pairwise alignments of your query sequence (such as RBP) against the database. PSI-BLAST also begins with a protein query that is searched against a database of choice.
- PSI-BLAST constructs a multiple sequence alignment (MSA) from an initial blastp-like search. It then creates a **PSSM** based on that multiple alignment.
- This **PSSM** is then used as a query to search the database again.
- PSI-BLAST estimates the statistical significance of the database matches, essentially using the parameters we described for gapped alignments.
- The search process is continued iteratively, typically 3 to 5 times. At each step a new PSSM is built.

#### PSI-BLAST returns dramatically more hits

#### 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.

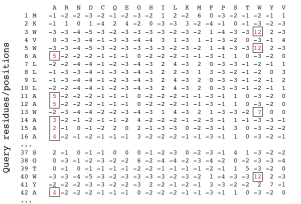
Iteration	Hits with E < 0.005	Hits with E > 0.005
1	34	61
2	314	79
3	416	57
4	432	50
5	432	50

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

#### Example PSI-BLAST PSSM at iteration 3

The PSI-BLAST PSSM is essentially a query customized scoring matrix that is more sensitive than BLOSUM (e.g. BLOSUM  $S_{AA} = +4$ )

#### 20 amino acids types



#### 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)

#### Profile advantages and disadvantages

#### Advantages:

- · Quantitate with a good scoring system
- 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

## **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



#### Hands-on sections 3 & 4:

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

~30 mins

Problems with PSSMs: Positional dependencies

Do not capture positional dependencies

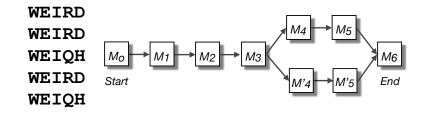
WEIRD WEIRD WEIQH WEIRD WEIRD

D					0.6
Е		Ι			
Н					0.4
Ι			Ι		
Q				0.4	
Q R				0.6	
W	Ι				

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

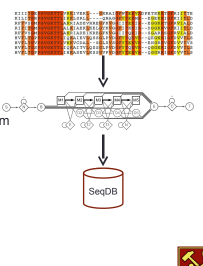
The connectivity or **topology** of a Markov chain can easily be designed to capture dependencies and variable length motifs.



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

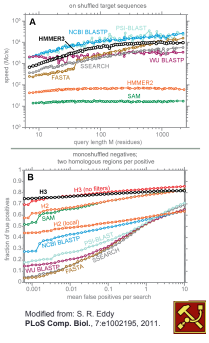
### Use of HMMER

- 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	BLAS <b>T</b>			
Progra <b>m</b>	PHMMER	B LA STP			
Quer <b>y</b>	Single sec	luence			
Targe <b>t</b> Databas <b>e</b>	Sequenc <b>e</b> d	atabas <b>e</b>			
Progra <b>m</b>	HMM SCAN	RP SB LA S <b>T</b>			
Quer <b>y</b>	Single sec	luence			
Targe <b>t</b> Databas <b>e</b>	Profil <b>e</b> HM <b>M</b> database, e.g. Pfa <b>m</b>	PSSM database e.g. CDD			
Progra <b>m</b>	HM M SE ARC <b>H</b>	PSI-BLAS <b>T</b>			
Quer <b>y</b>	Profil <b>e</b> HM <b>M</b>	PSSM			
Targe <b>t</b> Databas <b>e</b>	Sequenc <b>e</b> d	atabas <b>e</b>			
Progra <b>m</b>	JACKHMME <b>R</b>	PSI-BLAS <b>T</b>			
Quer <b>y</b>	Single sec	luence			
Targe <b>t</b> Databas <b>e</b>	Sequenc <b>e</b> d	atabas <b>e</b>			



## **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



Signi	ficant Query Matches (12)	in swissprot (v.2018_11)		Custo	omise Customia
	Target	Description	Species	Cross-references	E-value
>	HBB_HUMAN ₽	Hemoglobin subunit beta	Homo sapiens@	🗰 🕃 😸 🤮 🚍 📾	6.8e-99
>	HBD_HUMAN @	Hemoglobin subunit delta	Homo sapiens@	🚾 🕃 🌐 🕸 🛟 🖪	1.6e-91
>	HBE_HUMAN ₽	Hemoglobin subunit epsilon	Homo sapiens@	m 🔅 🧱 🕴 😽 🕃 🖿	1.5e-74
>	HBG2_HUMAN	Hemoglobin subunit gamma-2	Homo sapiens@	🚾 🕃 🏭 🕴 🔁 🖪	8.8e-73
>	HBG1_HUMAN⊮	Hemoglobin subunit gamma-1	Homo sapiens@	<b>(1)</b>	6.2e-72
>	HBA_HUMAN ₪	Hemoglobin subunit alpha	Homo sapiens@	m 😠 🗮 🕴 😽 🕞 🖪	3.8e-29
>	HBAZ_HUMAN@	Hemoglobin subunit zeta	Homo sapiens@	m 🗟 🗮 k 😣 🕞 🖪	4.5e-23
>	HBAT_HUMAN₽	Hemoglobin subunit theta-1	Homo sapiens@	m 9 🔳 😽 🗘 🖪	5.2e-22
>	HBM_HUMAN®	Hemoglobin subunit mu	Homo sapiens@	x & = * C E	3.4e-19
>	CYGB_HUMAN⊯	Cytoglobin	Homo sapiens@	m 😨 🗮 🖡 😵 🔁 🖪	3.1e-14
>	MYG_HUMAN	Myoglobin	Homo sapiens@		2.3e-06
>	NGB_HUMAN⊯	Neuroglobin	Homo sapiensø		0.0017
(shov	v all) alignments	Your search took: 0.	06 secs	show	ing rows 1 - 12 of

## Visualization of Results – By Score



	Q16IU8_AEDAE샵 응		SH2/SH3 ac	laptor p	rotein	Aedes	aegypti岱		2.5e-31				
Qu	ery			% Similarity	Bit	E-value							
start	end	start	end	d start	end			(count)	,	(count)	Score	Ind.	Cond.
7	62	4	81	9	63	0.02	0.81	36.4 (20	))	50.9 (28)	19.5	0.2	0.0001

Query	7														* q <mark>gwvpsny</mark> it	62
															g+vpsny+	
Target	9	DVCY	VVAI	KYDY	AA	QGA	QELI	DLF	RKNER	YLLLD-	DSK	HWW	RVQN	SHNQ	SGYVPSNYVK	63
PP		5566	799	****	**	***	* * *	* * *	****9	87775.	455	677	7665	1677	7******96	



### Visualization of Results – By Taxonomy

litere	Monotremata(65)	Ornithorhynchidae(65)	Ornithorhynchus(65)
	listen	litetatus	litite
	Primates(86)	Hominidae(86)	— Homo(86) 1
	limas	lina	lm.a
	Rodentia(244)	Cricetidae(46)	Cricetulus(46)
	ht.a	h	<u>h</u>
		Muridae(198)	— Mus(83)
		htsa	liten
			Rattus(115)
			httan .
Species Distribution			
Rattus norvegicus	Species	Count 115	View Show
Homo saplens @		86	Show
Mus musculus @ Sarcophilus harrisil@		83	Show
Ornithorhynchus anatinus		65	Show
Cricetulus griseust		46	Show
			Show All Visible

## Visualization of Results – By



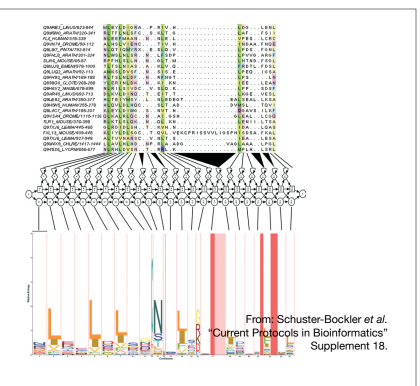
#### **PFAM:** Protein Family Database of Profile HMMs

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



## Summary

- 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

### Homework: DataCamp!

Check your email for an DataCamp invite and sign up with your UCSD username (i.e. first part of your email address) please.

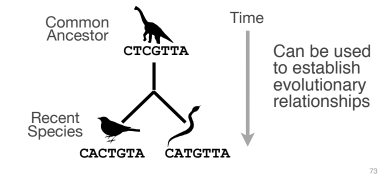
Let me know <u>NOW</u> if you don't see the invite email!



## Sequence comparison is most informative when it detects homologs Homologs are sequences that have common origins When

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)

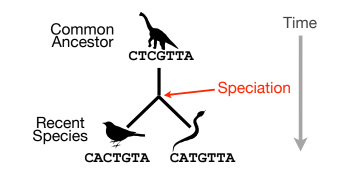
74

76

## 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

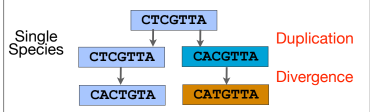


75

# 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]



### 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

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