

Bioinformatics for Biologists

Comparative Protein Analysis: Part II. Sequence Pattern and Profile Database Searching

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Knowledge Exploration

- **Phylogenetic Trees** and **Multiple Sequence Alignments** are important tools to understand relationships between known sequences.
- How do you apply what you know about a group of sequences to finding additional, related sequences?
- What can the relationship between your sequences and newly discovered tell you about their function?
- Discovering sequence **Families**

Syllabus

(Finding Family Members)

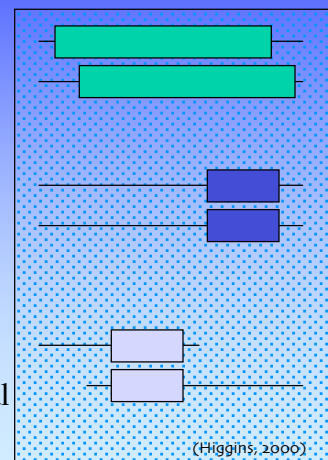
- **Protein Families**
 - Protein Domains
 - Family Databases & Searches
- Searching for Homologous Sequences Using Patterns/Profiles
 - Pattern Searches
 - Patscan
 - Profile Searches
 - PSI-BLAST/HMMER2

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Proteins As Modules

- Proteins are derived from a limited number of basic building blocks (**Domains**)
- Evolution has shuffled these modules giving rise to a diverse repertoire of protein sequences
- As a result, proteins can share a global or local relationship

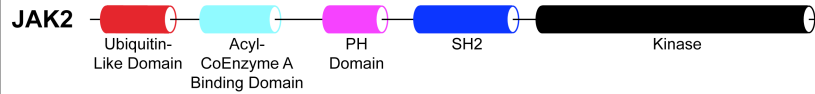


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Protein Domains

Janus Kinase 2 Modular Sequence Architecture



SH2 Motif

```

BLK_MOUSE 117-198 WFFRTLEKDAERQQLAPKAGSFLIREENKGAFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
LCK_MOUSE 126-208 WFFKRLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
LYN_MOUSE 128-210 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
FSR_HUMAN 144-224 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
SRC_RSVF 148-230 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
NCK1_HUMAN 282-356 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
VAV_MOUSE 671-745 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
ARI2_HUMAN 173-248 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
P85A_HUMAN 624-698 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
SIC_HUMAN 488-559 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
ITK_HUMAN 239-323 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
BTX_HUMAN 281-362 WFFKDLERKDAERQQLAPGTHGFLIREENKAGSFLRVKDTLTCQEV--VHYKIRSLDNG--GFYISPRIT--FFLQAVQVHY
    
```

Motifs describe the domain

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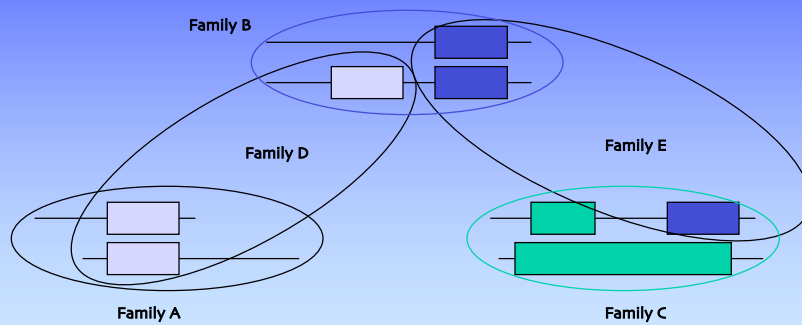
Protein Families

- **Protein Family** - a group of proteins that share a common function and/or structure, that are potentially derived from a common ancestor (set of homologous proteins)
- **Characterizing a Family** - Compare the sequence and structure patterns of the family members to reveal shared characteristics that potentially describe common biological properties
- **Motif/Domain** - sequence and/or structure patterns common to protein family members

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Protein Families



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Family Database Resources

- **Curated Databases***
 - Proteins are placed into families with which they share a specific sequence pattern
- **Clustering Databases***
 - Sequence similarity-based without the prior knowledge of a specific patterns
- **Derived Databases***
 - Pool other databases into one central resource
- **Search and Browse**

*(Higgins, 2000)

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Curated Family Databases

- **Pfam** (<http://pfam.wustl.edu/hmmsearch.shtml/>) **
 - Uses manually constructed seed alignments and PSSM to automatically extract domains
 - db of protein families and corresponding profile-HMMs
 - Searches report e-value and bits score
- **Prosite** (<http://www.expasy.ch/tools/scanprosite/>)
 - Hit or Miss -> no stats
- **PRINTS** (<http://www.bioinf.man.ac.uk/fingerPRINTScan/>)

Pfam HMM search results, glocal+local alignments merged (Pfam_Is+Pfam_fs)
[\[Go here for an explanation of the format of the results\]](#)

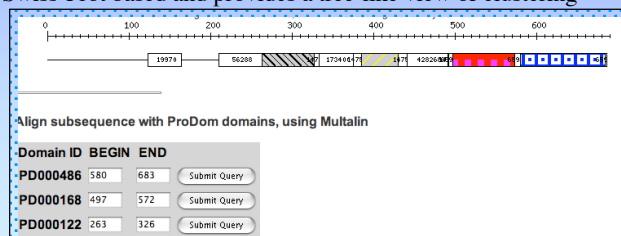
Model	Seq-from	Seq-to	HMM-from	HMM-to	Score	E-value	Alignment	Description
!! GTP_EFTU	258	483	1	298	315.7	5.5e-92	glocal	Elongation factor Tu GTP binding domain
!! GTP_EFTU_D2	502	570	1	75	46.1	8e-11	glocal	Elongation factor Tu domain 2
!! GTP_EFTU_D3	576	684	1	112	142.9	6.1e-40	glocal	Elongation factor Tu C-terminal domain

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Clustering Family Databases

- Search a database against itself and cluster similar sequences into families
- **ProDom** (<http://prodes.toulouse.inra.fr/prodom/doc/prodom.html>)
 - Searchable against MSAs and consensus sequences
- **Protomap** (<http://protomap.cornell.edu/>)
 - Swiss-Prot based and provides a tree-like view of clustering

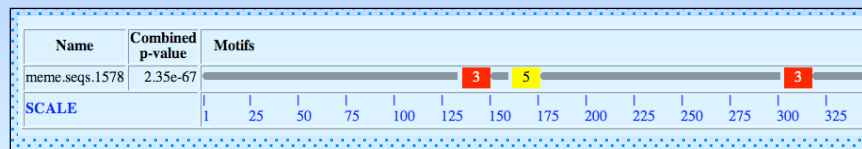


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Derived Family Databases

- Databases that utilize protein family groupings provided by other resources
- **Blocks** - Search and Make (<http://blocks.fhrc.org/blocks/>)
 - Uses Protomap system for finding blocks that are indicative of a protein family (GIBBS/MOTIF)
- **Proclass** (<http://pir.georgetown.edu/gfserver/proclass.html>)
 - Combines families from several resources using a neural network-based system (relationships)
- **MEME** (<http://meme.sdsc.edu/meme/website/intro.html>)



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Searching Family Databases

- BLAST searches provide a great deal of information, but it is difficult to select out the important sequences (listed by score, not family)
- Family searches can give an immediate indication of a protein's classification/function
- Use Family Database search tools to identify domains and family members

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Syllabus

(Finding Family Members)

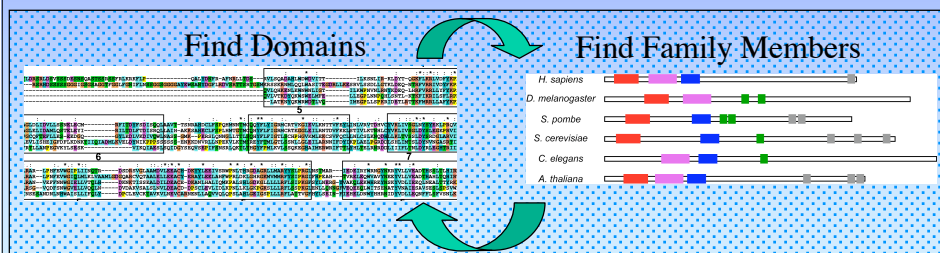
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Creating Protein Families

- Use domains to identify family members
 - Use a sequence to search a database and characterize a pattern/profile
 - Use a specific pattern/profile to identify homologous sequences (family members)



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Patterns & Profiles

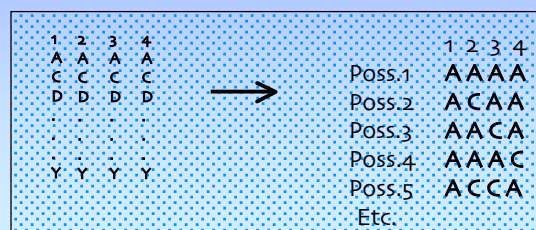
- Techniques for searching sequence databases to uncover common domains/motifs of biological significance that categorize a protein into a family
- **Pattern** - a deterministic syntax that describes multiple combinations of possible residues within a protein string
- **Profile** - probabilistic generalizations that assign to every segment position, a probability that each of the 20 aa will occur

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Discovery Algorithms

- Pattern Driven Methods
 - Enumerate all possible patterns in solution space and try matching them to a set of sequences

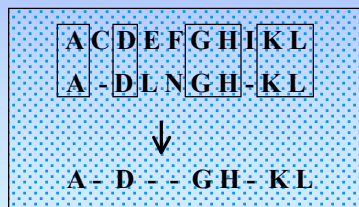


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Discovery Algorithms

- Sequence Driven Methods
 - Build up a pattern by pair-wise comparisons of input sequences, storing positions in common, removing positions that are different



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Pattern Building

- Find patterns like “aa1 xx aa2 xxxx aa3”
 - Definition of a non-contiguous motif

1.	C	Y	D	C	A	F	T	L	R	Q	S	A	V	M	H	K	H	A	R	E	H		
2.	C	A	T	Y	C	R	T	A	I	D	T	V	K	N	S	L	K	H	H	S	A	H	
3.	C	W	D	G	G	C	G	I	S	V	E	R	M	D	T	V	H	K	H	D	T	V	H
4.	C	Y	C	C	S	D	H	M	K	K	D	A	V	E	R	M	H	K	K	D	H		
5.	C	N	M	F	C	M	P	I	F	R	Q	N	S	L	A	R	E	H	E	R	M	H	
6.	C	L	N	N	T	C	T	A	F	W	R	Q	K	K	D	D	T	V	H	N	S	L	H
	C	xxxx	C	xxxx	[LIVMFW]	xxxxxxxx	H	xxxxx	H														

Define/Search A Motif <http://us.expasy.org/tools/scanprosite/>

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Pattern Properties

- **Specification**
 - a single residue K, set of residues (KPR), exclusion {KPR}, wildcards X, varying lengths x(3,6) -> variable gap lengths
- **General Syntax**
 - C-x(2,4)-C-x(3)-[LIVMFYWC]-x(8)-H-x(3,5)-H
- **Patscan Syntax**
 - C 2...4 C 3...3 any(LIVMFYWC) 8...8 H 3...5 H
- **Pattern Database Searching**
 - `%scan_for_matches -p pattern_file < /db0/Data/nr > output_file`

Sequence Pattern Concerns

- Pattern descriptors must allow for approximate matching by defining an acceptable distance between a pattern and a potential hit
 - Weigh the sensitivity and specificity of a pattern
- What is the likelihood that a pattern would randomly occur?

Sequence Profiles

- **Consensus** - mathematical probability that an aa will be located at a given position
- **Probabilistic** pattern constructed from a MSA
- Opportunity to assign penalties for insertions and deletions, but not well suited for variable gap lengths
- **PSSM** - (Position Specific Scoring Matrix)
 - Represents the sequence profile in tabular form
 - Columns of weights for every aa corresponding to each column of a MSA

PSSM Example

1. I T I S
 2. T D L S
 3. V T M G
 4. I T I G
 5. V G F S
 6. I E L T
 7. T T T S
 8. I T L S

(i.e. Distribution of aa in an MSA column)

← Target sequences

Resulting Consensus: I T L S

PSSM



P O S	A	C	D	E	F	G	H	I	K	L	M	N	P	Q	R	S	T	V	W	Y
1	8	-2	5	-4	5	5	-4	24	0	15	13	1	1	1	-7	2	22	21	-18	-6
2	13	-5	24	18	-18	19	7	1	7	-7	-4	14	11	-10	-1	9	29	3	-28	-14
3	5	-5	3	4	13	4	2	8	-4	14	12	8	-5	0	-10	0	10	10	-1	5
4	17	17	13	10	-12	29	-5	-5	6	-14	-9	12	10	0	-2	34	19	1	-8	-15

PSSM Properties

- Score-based sequence representations for searching databases
 - Calculations determined by Log odds score
- Goal
 - Limit the diversity in each column to improve reliability
- Problems
 - Differing length gaps between conserved positions (unlike patterns)

PSSM Weighting

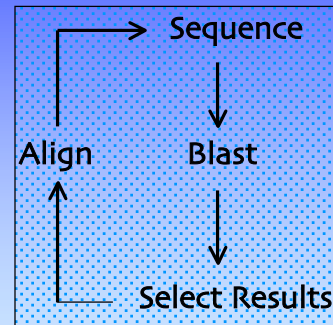
- Differentially weigh sequences to reduce redundancy from non-representative sampling
 - Similar sequences get low weights, diverged sequences get higher weights

PSI-BLAST Implementation

- **PSI-BLAST**

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

- Start with a sequence, BLAST it, align select results to query sequence, estimate a profile with the MSA, search DB with the profile - constructs PSSM
- Iterate until process stabilizes
- Focus on domains, not entire sequences
- Greatly improves sensitivity



PSI-BLAST Sample Output

Sequences with E-value WORSE than threshold

<input type="checkbox"/>	gi19629055 ref NP_044074.1	(NC_001731) MC123R [Molluscum contag...	37	0.16
<input type="checkbox"/>	gi18176554 gb AA35488.2	(S79774) bile salt-dependent lipase; B...	36	0.25
<input type="checkbox"/>	gi14502771 ref NP_001798.1	(NM_001807) carboxyl ester lipase (b...	35	0.86
<input type="checkbox"/>	gi12316291 sp P18835 BAL_HUMAN	Bile-salt-activated lipase precurs...	35	0.89
<input type="checkbox"/>	gi115242929 ref NP_200612.1	(NM_125189) putative protein [Arabi...	34	1.1
<input type="checkbox"/>	gi197595291 db BAB10995.1	(AB024029) gene_id:K21L19.3-unknown p...	34	1.3
<input type="checkbox"/>	gi11804821 gb AAA52014.1	(M85201) cholesterol esterase [Homo sap...	33	1.8
<input type="checkbox"/>	gi1187061 sp P21173 DNAA_MICLU	Chromosomal replication initiator...	32	4.6
<input type="checkbox"/>	gi11266791 sp P161101 LEG3_MOUSE	GALECTIN-3 (GALACTOSE-SPECIFIC LE...	32	4.9
<input type="checkbox"/>	gi152851 emb CAA34206.1	(X16074) L-34 protein (AA 1-264) [Mus sp.]	32	5.0
<input type="checkbox"/>	gi15399071 pir IAA45983	lactose-binding lectin Mac-2 - mouse	32	5.0
<input type="checkbox"/>	gi1387111 gb AAA37311.1	(J03723) carbohydrate binding protein 3...	32	5.4
<input type="checkbox"/>	gi195064271 ref NP_062019.1	(NM_019146) bassoon [Rattus norvegic...	32	5.5

HMM Building

- **Hidden Markov Models** are Statistical methods that considers all the possible combinations of matches, mismatches, and gaps to generate a consensus (Higgins, 2000)
- Sequence ordering and alignments are not necessary at the onset (but in many cases alignments are recommended)
- Ideally use at least 20 sequences in the training set to build a model
- Calibration prevents over-fitting training set (i.e. Ala scan)
- Generate a model (profile/PSSM), then search a database with it

HMM Implementation

- **HMMER2** (<http://hmmer.wustl.edu/>)
 - Determine which sequences to include/exclude
 - Perform alignment, select domain, excise ends, manually refine MSA (pre-aligned sequences better)
 - Build profile
 - `%hmmbuild [-options] <hmmfile output> <alignment file>`
 - Calibrate profile (re-calc. Parameters by making a random db)
 - `%hmmcalibrate [-options] <hmmfile>`
 - Search database
 - `%hmmsearch [-options] <hmmfile> <database file> > out`

HMMER2 Output

- Hmsearch returns e-values and bits scores
- Repeat process with selected results
 - Unfortunately need to extract sequences from the results and manually perform MSA before beginning next round of iteration

```
HMMER 2.2g (August 2001)
Copyright (C) 1992-2001 HHMI/Washington University School of Medicine
Freely distributed under the GNU General Public License (GPL)
-----
HMM file:          pfam_had.hmm [Hydrolase]
Sequence database: /cluster/db0/Data/nr
per-sequence score cutoff: [none]
per-domain score cutoff: [none]
per-sequence Eval cutoff: <= 10
per-domain Eval cutoff: [none]
-----
Query HMM: Hydrolase
Accession: PF00702
Description: haloacid dehalogenase-like hydrolase
           [HMM has been calibrated; E-values are empirical estimates]

Scores for complete sequences (score includes all domains):
Sequence      Description      Score  E-value  N
-----
gi|16131263|ref|NP_417844.1|  phosphoglycolat  168.4  2.9e-45  1
gi|24114648|ref|NP_709158.1|  phosphoglycolat  167.8  4.2e-45  1
gi|15803888|ref|NP_289924.1|  phosphoglycolat  167.8  4.2e-45  1
gi|26249979|ref|NP_756019.1|  Phosphoglycolat  166.4  1.1e-44  1
```

Patterns vs. Profiles

- **Patterns**
 - Easy to understand (human-readable)
 - Account for different length gaps
- **Profiles**
 - Sensitivity, better signal to noise ratio
 - Teachable

Demonstration

- Family/Domain Search
- Pattern Search
 - scan_for_matches (Patscan)
- Profile Search
 - PSI-BLAST
 - HMMER2

References

- Bioinformatics: Sequence and genome Analysis. David W. Mount. CSHL Press, 2001.
- Bioinformatics: A Practical Guide to the Analysis of Genes and Proteins. Andreas D. Baxevanis and B.F. Francis Ouellete. Wiley Interscience, 2001.
- Bioinformatics: Sequence, structure, and databanks. Des Higgins and Willie Taylor. Oxford University Press, 2000.