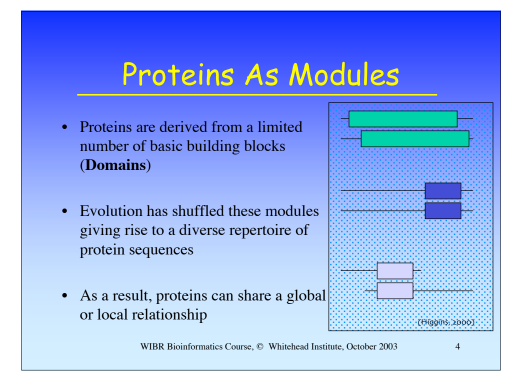


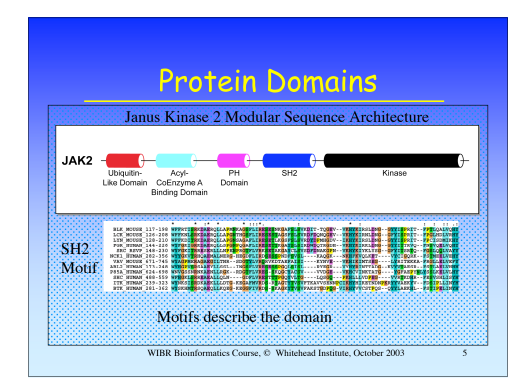
Syllabus

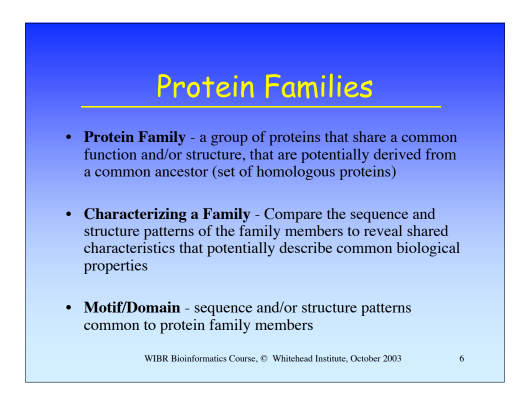
(Finding Family Members)

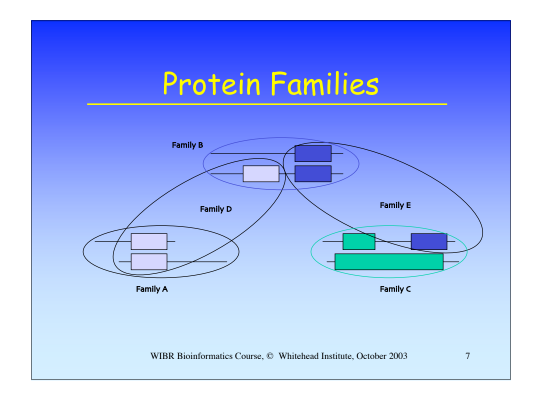
- Protein Families
 - Protein Domains
 - Family Databases & Searches
- Searching for Homologous Sequences Using Patterns/Profiles
 - Pattern Searches
 - Patscan
 - Profile Searches
 - PSI-BLAST/HMMER2

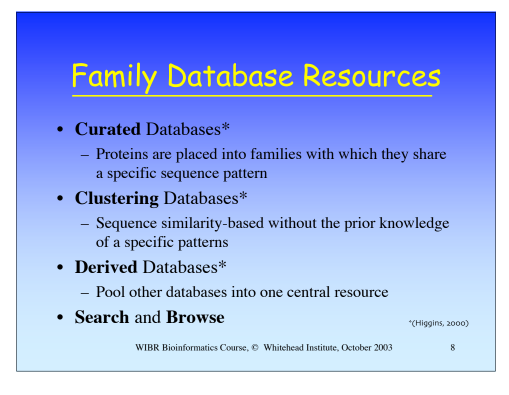
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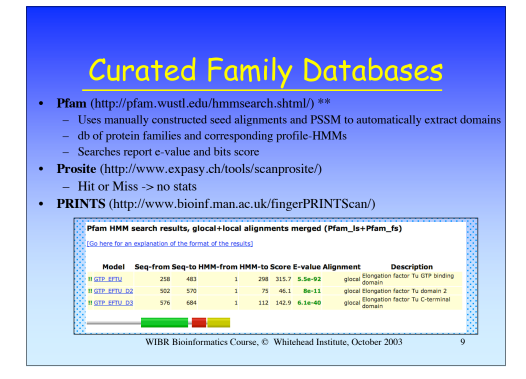


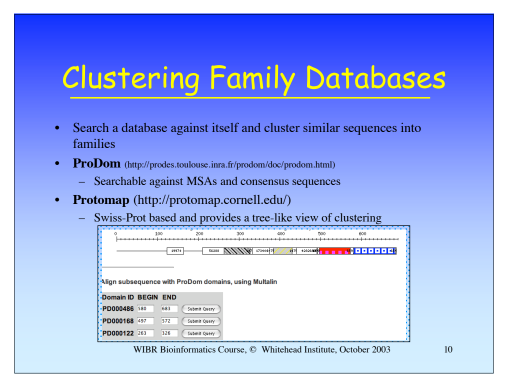


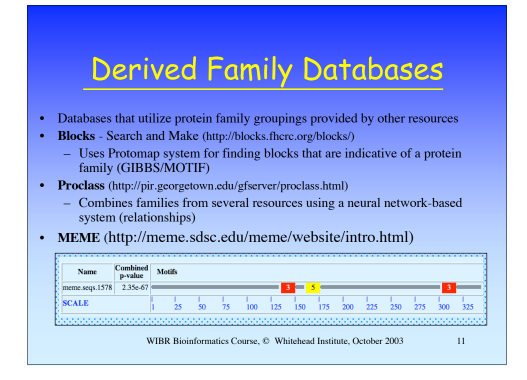


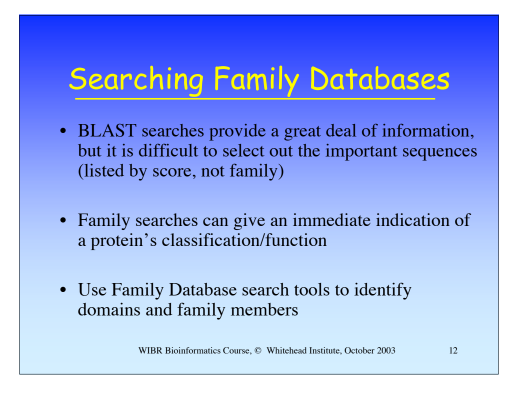










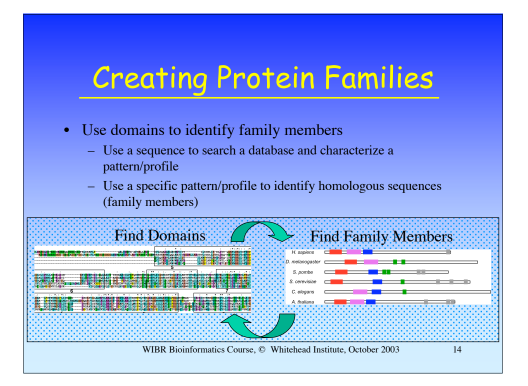


Syllabus

(Finding Family Members)

- Protein Families
 - Protein Domains
 - Family Databases & Searches
- Searching for Homologous Sequences (Finding Family Members)
 - Pattern Searches
 - Patscan
 - Profile Searches
 - PSI-BLAST/HMMER2

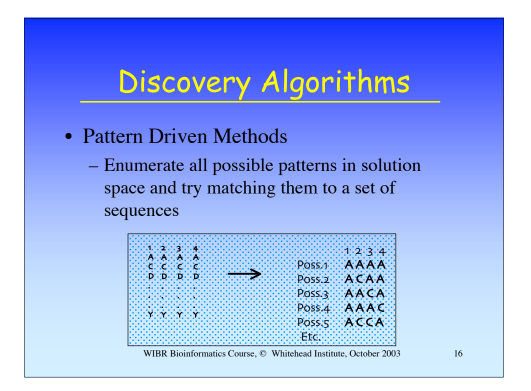
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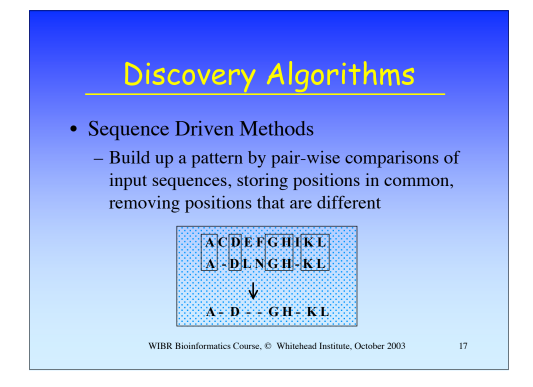


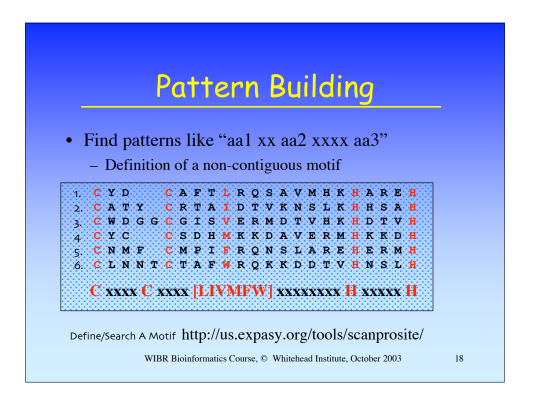
Patterns & Profiles

- 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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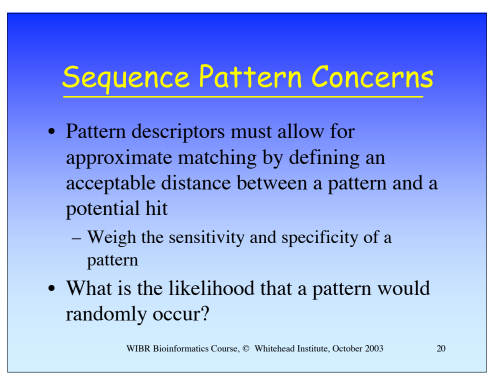
Pattern Properties

• 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

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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

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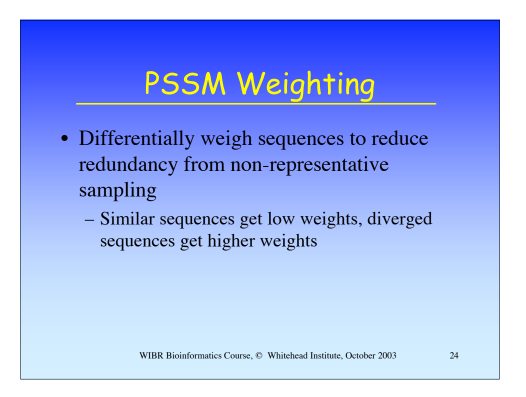
21

PSSM Example ITIS DLS (i.e. Distribution of aa in an MSA column) MG TIG ← Target sequences Resulting Consensus: ITLS G F ELT **PSSM** TTS ŦĽ. 0 s A С D Е F G н Ť K L м N Р Q R s т v w Y 1 8 -2 5 4 5 5 -4 <u>24</u> 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 <u>29</u> 3 -28 -14 5 -5 3 13 12 8 -5 Ö -10 0 10 10 -1 5 3 4 4 2 8 -4 <u>14</u> 17 17 13 10 -12 29 -5 -5 6 -14 -9 12 10 0 -2 <u>34</u> 19 1 -8 -15 4 WIBR Bioinformatics Course, © Whitehead Institute, October 2003 22

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)

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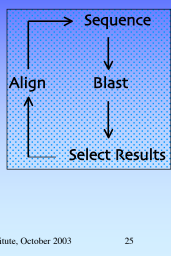




• PSI-BLAST

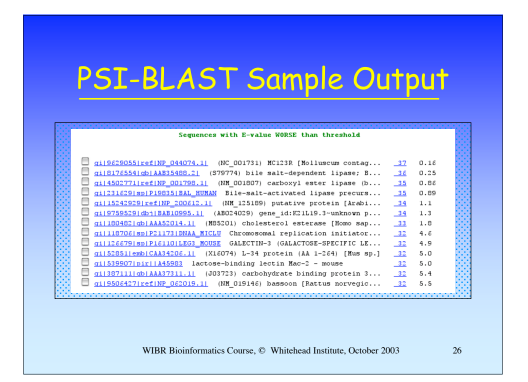
http://www.ncbi.nlm.nih.gov/BLAST/

 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

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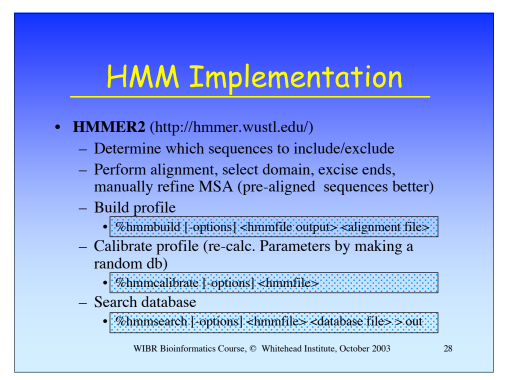


HMM Building

- **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

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HMMER2 Output

- Hmmsearch returns evalues 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: <a 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]

Sequence	Description		Score	E-value	Ν
 gil16131263lreflNP	417844.11	phosphoglycolat	168.4	2.9e-45	
gil24114648lreflNP		phosphoglycolat			
gil15803888lreflNP	289924.11	phosphoglycolat	167.8	4.2e-45	1
gil26249979lreflNP	756019.11	Phosphoglycolat	166.4	1.1e-44	- 1

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