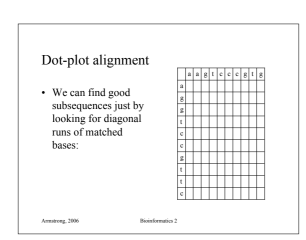


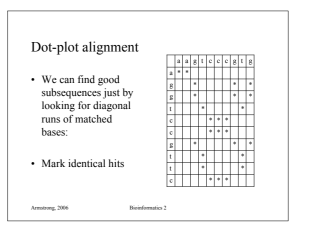
Bioinformatics 2

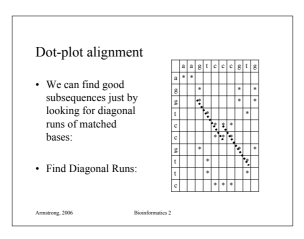
Armstrong, 2006

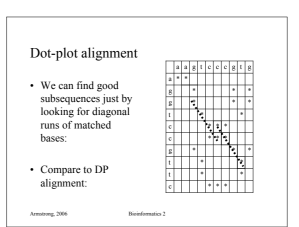


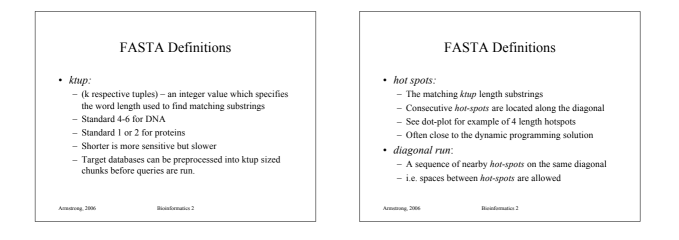
Bioinformatics 2

Armstrong, 2006











• init₁:

- The best scoring run
- *init_n*:
 The best local alignment
 - Combination of good diagonal runs and indels/gaps between them.

Armstrong, 2006

Bioinformatics 2



1. Look for hot-spots:

Armstrong, 2006

- The stage can be done by using a look-up table or a hash.
- Pre-process the database and store the location of each possible *ktup* (AA=20², DNA=4⁶)

Bioinformatics 2

• Move a *ktup* sized window along the query sequence and record the position of matching locations in the database.

2

FASTA Process

- 2. Find best diagonal runs:
- Each hot spot gets a positive score.
- Distance between *hot spots* is negative and length dependant
- Score of the diagonal run
- Fasta finds and stores the 10 best diagonal runs

Bioinformatics 2

Armstrong, 2006

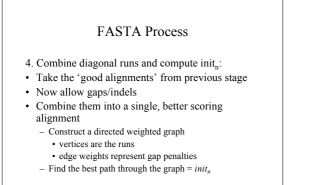
FASTA Process

Bioinformatics 2

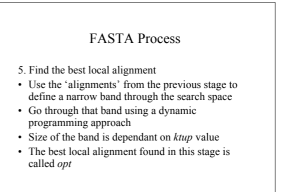
3. Compute $init_1$ & filter:

- · Diagonal runs specify a potential alignment
- Evaluate properly using a substitution matrix
- Define the best scoring run as init₁
- · Discard any much lower scoring runs

Armstrong, 2006







Armstrong, 2006

FASTA Process

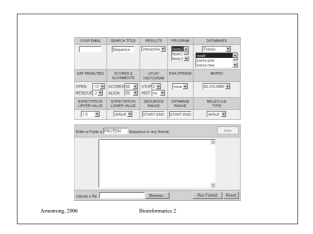
- 6. Compare the alignments
- Take the *opt* or $init_n$ scores for each sequence in the database
- · Rank according to score
- Use a full dynamic programming algorithm to align the query sequence with the highest ranking result sequences

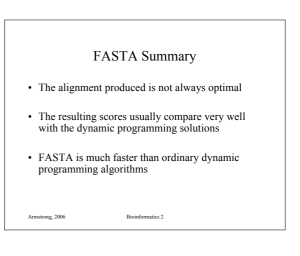
Armstrong, 2006 Bioir

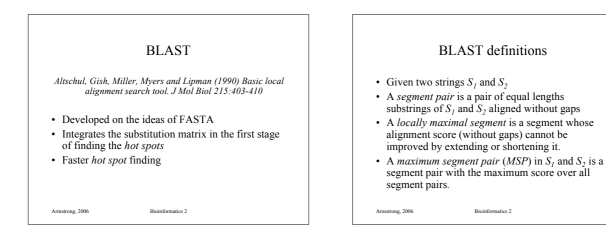
Bioinformatics 2

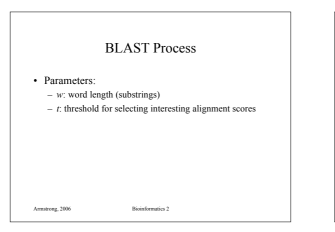
FASTA Programs

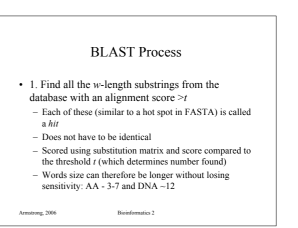
• fasta3	scan a protein or DNA sequence library for similar sequences
• fastax/y3	compare a DNA sequence to a protein sequence database, comparing the translated DNA sequence in forward and reverse frames
• tfastax/y3	compares a protein to a translated DNA data bank
fasts3fastf3	compares linked peptides to a protein databank compares mixed peptides to a protein databank
Armstrong, 2006	Bioinformatics 2









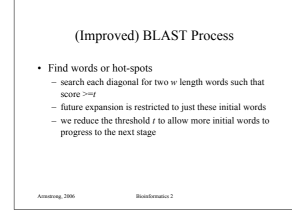


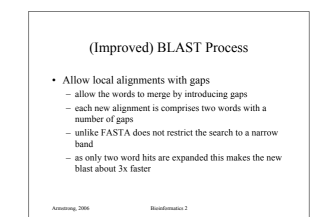


- 2. Extend hits:
 - extend each hit to a local maximal segment
 - extension of initial w size hit may increase or decrease the score
 - terminate extension when a threshold is exceeded
 - find the best ones (HSP)
- This first version of Blast did not allow gaps....

Armstrong, 2006 Bioinformatics 2

(Improved) BLAST Altshul, Madden, Schaffer, Zhang, Zhang, Miller & Lipman (1997) Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Research 25:3389-3402 inster database search 25:3389-3402 inste





PSI-BLAST

- Iterative version of BLAST for searching for protein domains
 - Uses a dynamic substitution matrix
 - Start with a normal blast

Armstrong, 2006

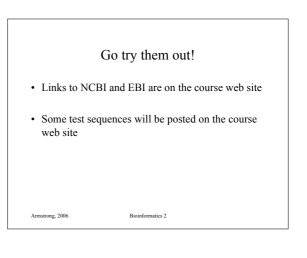
- Take the results and use these to 'tweak' the matrix
- Re-run the blast search until no new matches occur
- Good for finding distantly related sequences but high frequency of false-positive hits

Bioinformatics 2

BLAST Programs

blastp compares an amino acid query sequence against a protein sequence database.
 blastn compares a nucleotide query sequence against a nucleotide sequence database.
 blastx compares a nucleotide query sequence translated in all reading frames against a protein sequence database.
 tblastn compares a protein query sequence against a nucleotide sequence database.
 tblastn compares a protein query sequence against a nucleotide sequence database.
 tblastn compares a protein query sequence against a nucleotide sequence database.
 tblastx compares the six-frame translations of a nucleotide query sequence against the six-frame translations of a nucleotide sequence database. (SLOW)
 Armstrong, 2006 Bioinformatics 2

YOUR EMAIL	SEARCH TITLE	RESULTS	PROGRAM	DAT	ABASE
	Sequenc	interactive 🛟	WU-blastp 🛟	Protein UniProt	:
MATRIX	DNA STRAND	EXP.THR	FILTER	VIEW	FILTER
blosum62 🛟	none 🛟	default 🗘	none 🕄	nc	;
SENSITIVITY	STATS	SORT	topcomboN	SCORES	ALIGNMENTS
normal ‡	sump 🛟	pvalue \$	default \$	default \$	default :
Enter or Paste a	PROTEIN : S	Sequence in any fe	ormat:		Help
Enter or Paste a	PROTEIN : S	Sequence in any fr	ormat:		Help

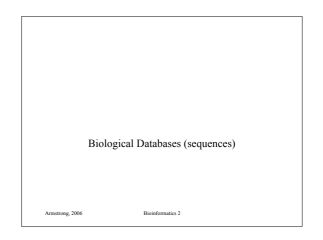


Alignment Heuristics

- Dynamic Programming is better but too slow
- FASTA and BLAST based on several assumptions about good alignments

 substitutions more likely than gaps
- good alignments have runs of identical matches
- FASTA good for DNA sequences but slower
- BLAST better for amino acid sequences and pretty good for DNA, fastest.

Armstrong, 2006 Bioinformatics 2



Biological Databases

- Introduction to Sequence Databases
- Overview of primary query tools and the databases they use (e.g. databases used by BLAST and FASTA)
- Demonstration of common queries
- Interpreting the results
- Overview of annotated 'meta' or 'curated' databases

Armstrong, 2006

Bioinformatics 2

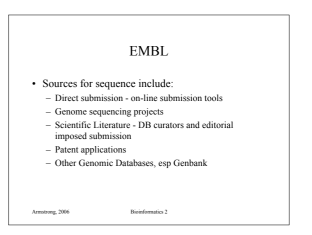
DNA Sequence Databases

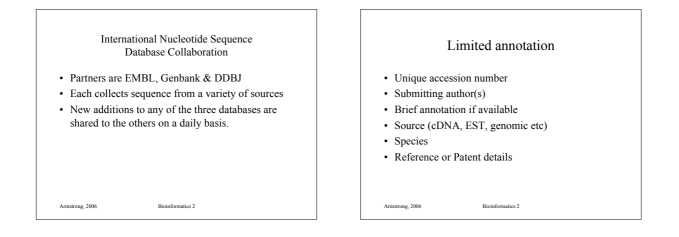
- Raw DNA (and RNA) sequence
- · Submitted by Authors
- Patent, EST, Gemomic sequences
- Large degree of redundancy
- Little annotation

Armstrong, 2006

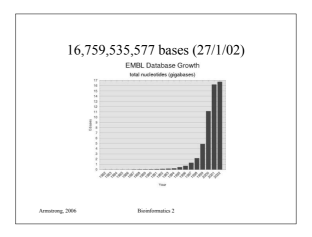
· Annotation and Sequence errors!

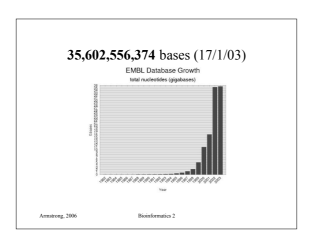
Main DNA DBs				
• Genbank	US			
• EMBL	EU			
• DDBJ	Japan			
Celera genomics	Commercial DB			
Armstrong, 2006	Bioinformatics 2			

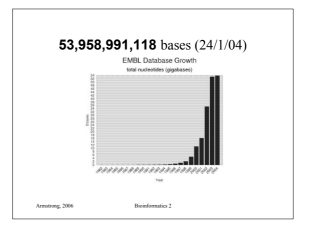


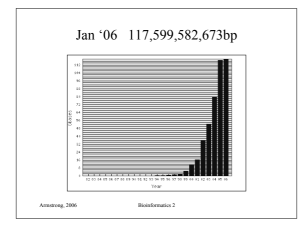


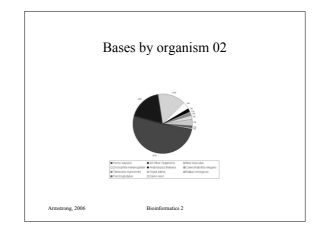


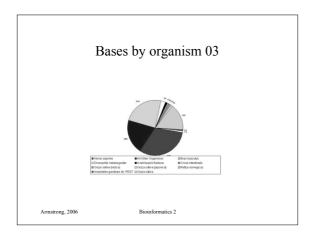


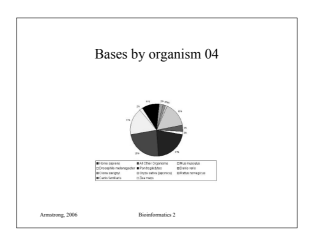


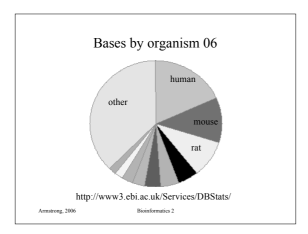


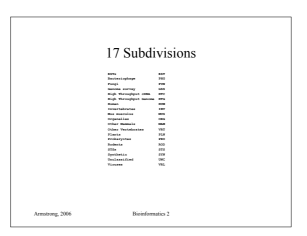


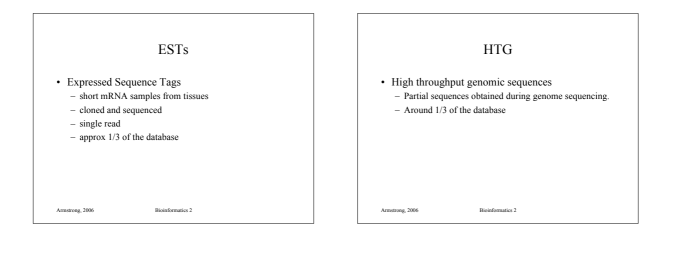












Specialist DNA Databases

- Usually focus on a single organism or small related group
- Much higher degree of annotation
- · Linked more extensively to accessory data
 - Species specific:

Armstrong, 2006

- Drosophila: FlyBase,
- C. elegans: AceDB
- Other examples include Mitochondrial DNA, Parasite Genome DB

Bioinformatics 2

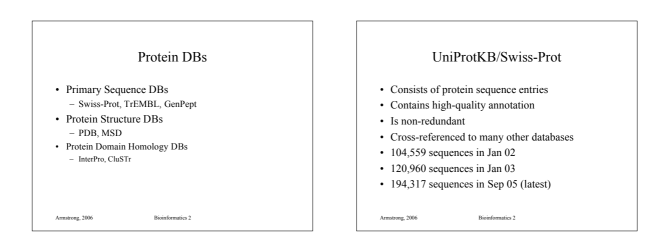


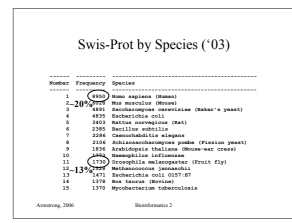
• Includes the entire annotated genome searchable by BLAST or by text queries

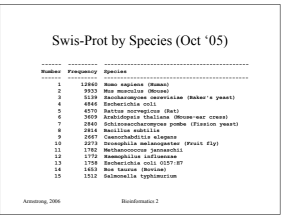
Bioinformatics 2

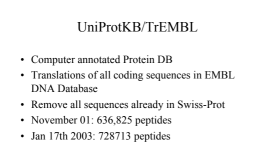
- Also includes a detailed ontology or standard nomenclature for *Drosophila*
- Also provides information on all literature, researchers, mutations, genetic stocks and technical resources.
- Full mirror at EBI

Armstrong, 2006



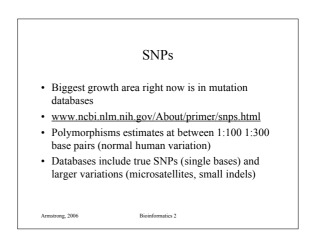


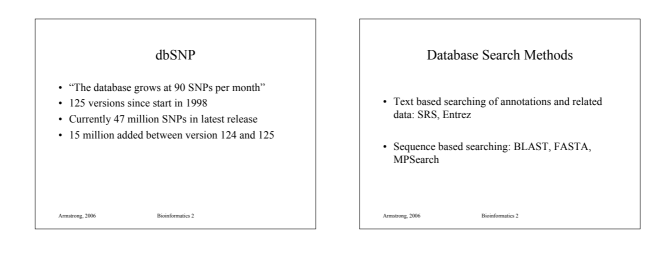


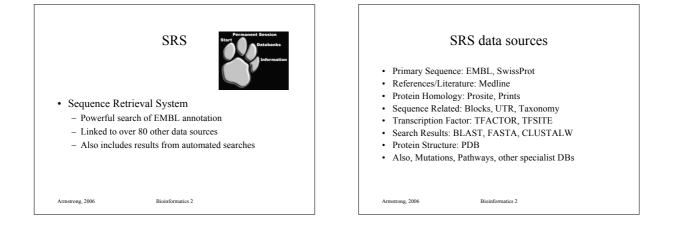


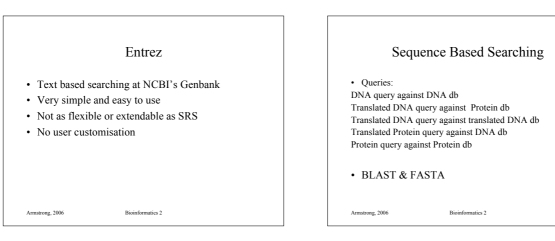
- TrEMBL new is a weekly update
- · GenPept is the Genbank equivalent

Armstrong, 2006 Bioinformatics 2

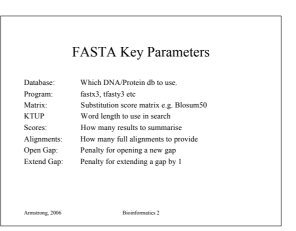


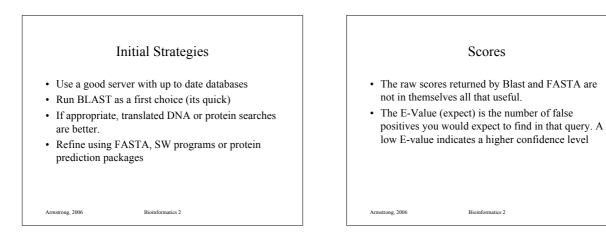


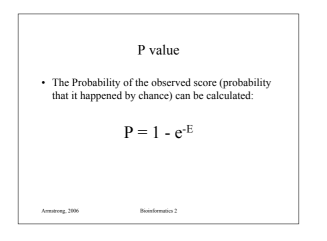


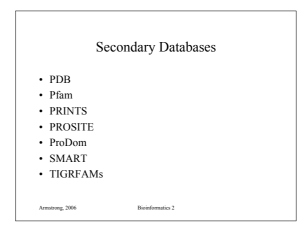


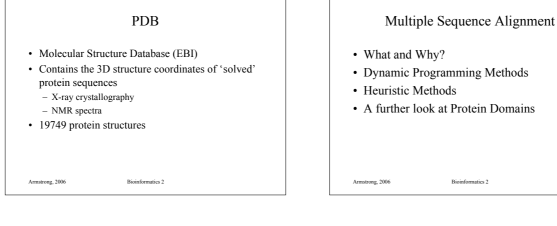
	BLAS	Т
Version	Query	<u>DB</u>
Blastn	DNA	DNA
Blastp	Peptide	Peptide
Blastx	DNA	Peptide
tBlastn	Peptide	DNA
tBlastx	DNA	DNA
Armstrong, 2006	Bioinformatic	s ₂ Iranslated

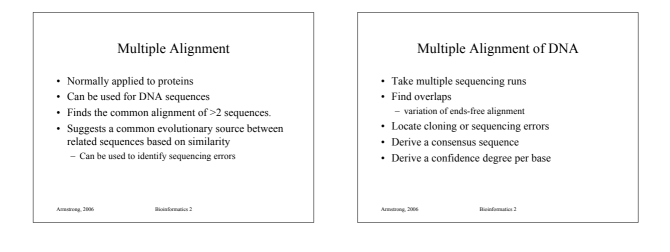


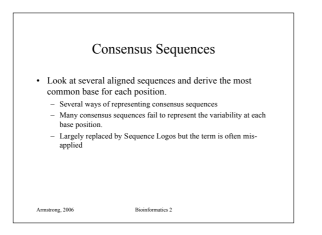


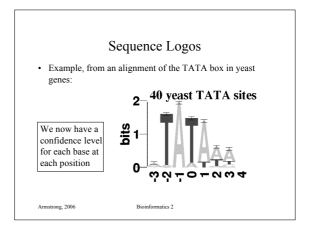


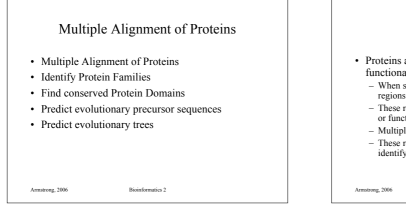


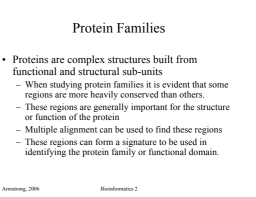


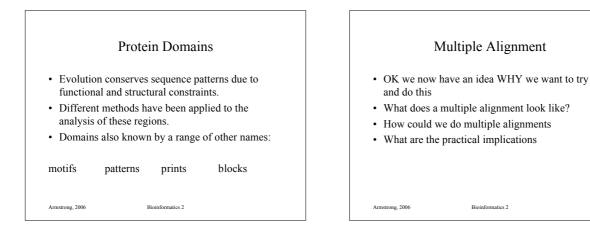


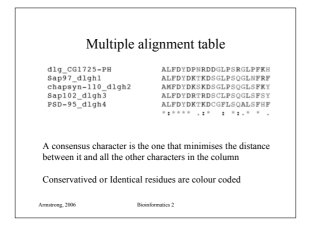


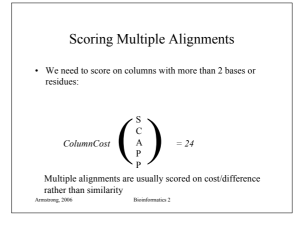


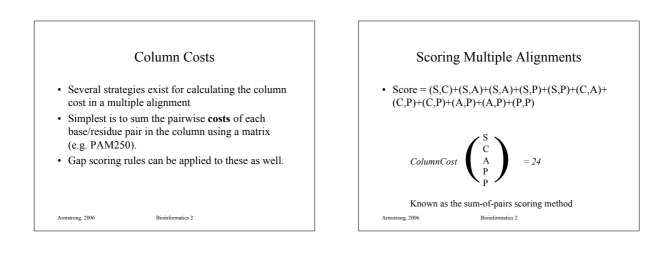


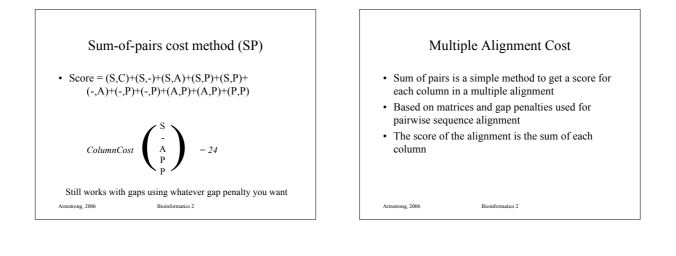


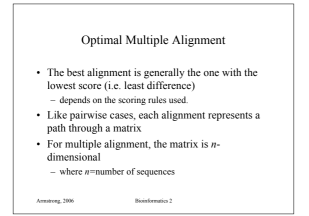


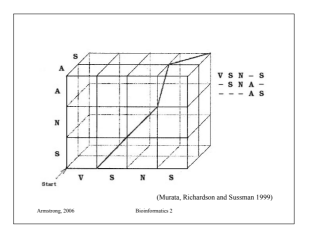


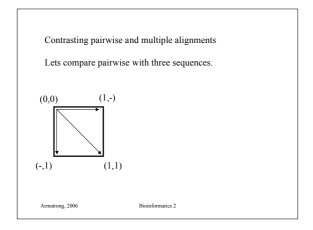


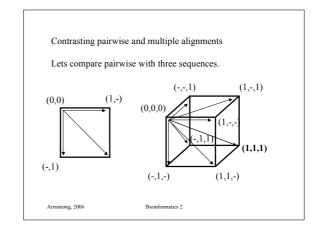


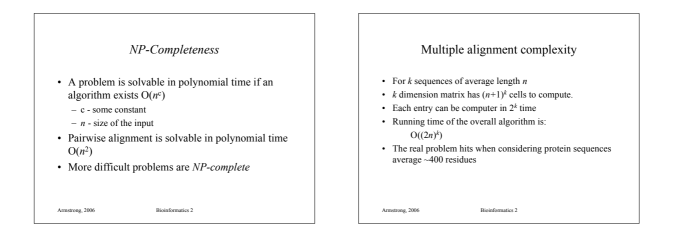


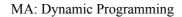








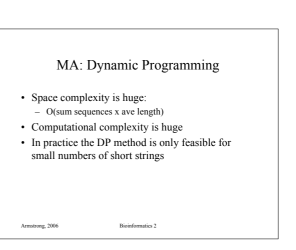


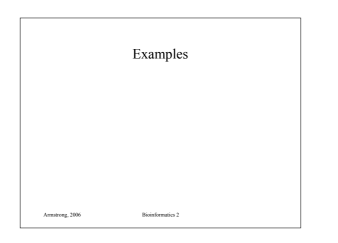


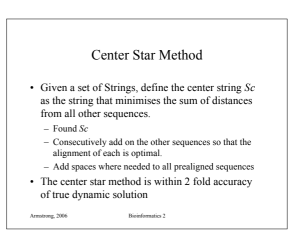
- We can use dynamic programming in some small cases.
- For *x* sequences, build an *x* dimensional hypercube.

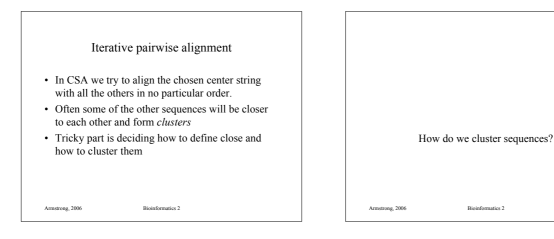
Armstrong, 2006

• Solve as before using gap and substitution penalties but remembering that there are more routes to each cell in the hypercube









Building trees

• Need to define how the sequences are related to

• Most use the distances between pairs in the set of

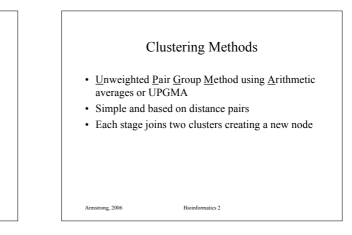
• Key parameter is in defining the distance score.

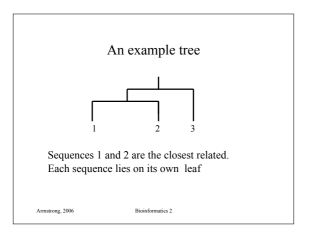
Bioinformatics 2

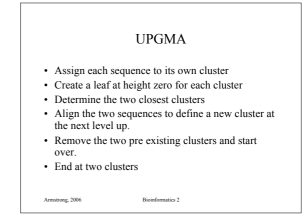
one another.

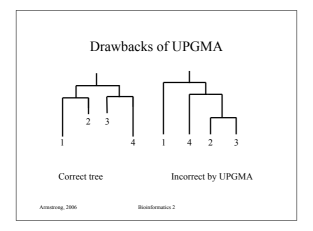
sequences.

Armstrong, 2006

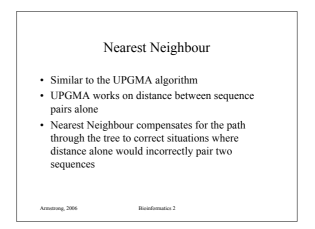


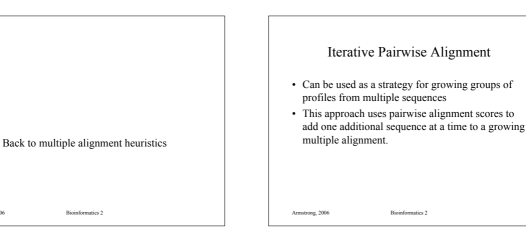


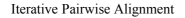




Armstrong, 2006







- · First align all pairs of strings where one is already in a multiple alignment and one is aligned.
- · Find the closest matches.
- · Align the unassigned sequence with the family profile of the closest group
- Realign the group and get a new profile.

Armstrong, 2006

Bioinformatics 2

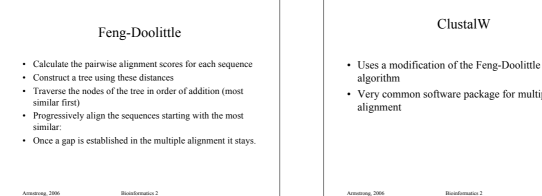
Feng-Doolittle

- · Feng-Doolittle 1987 Journal of Molecular Evolution 25:351-360
- · The key principal is that the two most similar sequences in a multiple alignment are the most recently diverged.
- Therefore the pairwise alignment of these two sequences is . the most reliable of the entire group

Bioinformatics 2

· Gaps present in the alignment should therefore be preserved in the multiple alignment.

Armstrong, 2006



Very common software package for multiple

ClustalW

- · Starts by calculating pairwise alignments and converting scores to distances
- Uses a neighbour joining algorithm to build a tree from the distances
- · Aligns sequences to each other
- · Aligns sequences to profiles
- · Aligns profiles to profiles
- Can output multiple alignment as well a predicted evolutionary tree

Armstrong, 2006

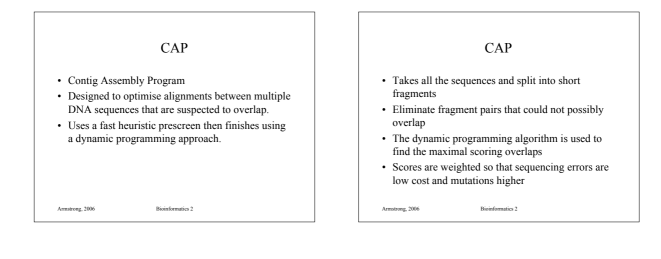
Bioinformatics 2

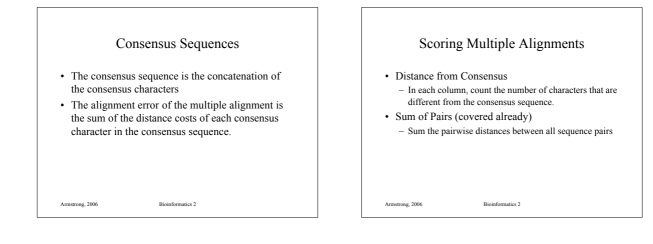
MSA

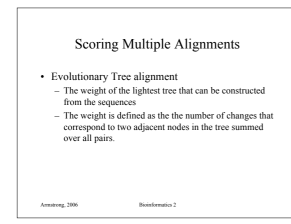
- · Exploits the fact that closely aligned sequence paths will be close to the main diagonal on a DP table.
- · Estimates a good solution, removes cells from the hypercube where the score could not feasibly pass through them.

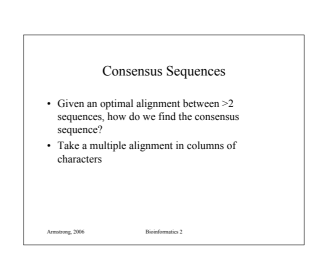
Bioinformatics 2

trong, 2006









Multiple alignment table

 dlg_CG1725-PH
 ALFDYDPNRDDGLPSRGLPFKH

 Sap97_dlgh1
 ALFDYDKTKDSGLPSQGLNFRF

 chapsyn-110_dlgh2
 AMFDYDKSKDSGLPSQGLSFKY

 sap102_dlgh3
 ALFDYDRTRSCLPSQGLSFKY

 PSD-95_dlgh4
 ALFDYDKTKDCGFLSQALSFHF

The consensus character is the one that minimises the distance between it and all the other characters in the column

Bioinformatics 2

Armstrong, 2006

Finally some examples We are interested in the protein DLG DLG is a molecular scaffold 1 gene in Drosophila 4 human genes (DLG1-4 with synonyms) Tarpey et al 2004 found mutations linking DLG3/Sap102 to Mental Retardation Obtained sequences for all 5 proteins Run through ClustalW (results on-line)

Another example

- We are also interested in PDE4B
 - PDE4B is a phosphodiesterase
 - 1 gene in Drosophila (dunce) linked to memory
 multiple human genes closest PDE4B
- Millar et al 2005 found a link between PDE4B and schizophrenia
- A database search funds many possible PDE4B proteins, need to make sense of it all...

Armstrong, 2006 Bioinformatics 2