Heuristic methods for alignment Sequence databases Multiple alignment Gene and protein prediction

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Assumptions for Heuristic Approaches

- Even linear time complexity is a problem for large genomes
- Databases can often be pre-processed to a degree
- Substitutions more likely than gaps
- Homologous sequences contain a lot of substitutions without gaps which can be used to help find start points in alignments

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BLAST

Heuristic Methods

• FASTA

• BLAST

· Gapped BLAST

• PSI-BLAST

Basic Local Alignment Search Tool

Altschul, Gish, Miller, Myers and Lipman (1990) Basic local alignment search tool. J Mol Biol 215:403-410

- Developed on the ideas of FASTA
 - uses short identical matches to reduce search = hotspot
- Integrates the substitution matrix in the first stage of finding the hot spots
- Faster hot spot finding

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BLAST definitions

- Given two strings S_1 and S_2
- A segment pair is a pair of equal lengths substrings of S_1 and S_2 aligned without gaps
- A locally maximal segment is a segment whose alignment score (without gaps) cannot be improved by extending or shortening it.
- A maximum segment pair (MSP) in S₁ and S₂ is a segment pair with the maximum score over all segment pairs.

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BLAST Process

- Parameters:
 - − w: word length (substrings)
 - *t*: threshold for selecting interesting alignment scores

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BLAST Process

- 1. Find all the *w*-length substrings from the database with an alignment score >*t*
 - Each of these (similar to a hot spot in FASTA) is called a hit
 - Does not have to be identical
 - Scored using substitution matrix and score compared to the threshold t (which determines number found)
 - Words size can therefore be longer without losing sensitivity: AA - 3-7 and DNA ~12

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BLAST Process

- · 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....

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(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

- Improved algorithms allowing gaps
 - these have superceded the older version of BLAST
 - two versions: Gapped and PSI BLAST

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(Improved) BLAST Process

- Find words or hot-spots
 - search each diagonal for two w length words such that score >=t
 - future expansion is restricted to just these initial words
 - we reduce the threshold t to allow more initial words to progress to the next stage

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(Improved) BLAST Process

- Allow local alignments with gaps
 - allow the words to merge by introducing gaps
 - each new alignment comprises two words with a number of gaps
 - unlike FASTA does not restrict the search to a narrow band
 - as only two word hits are expanded this makes the new blast about 3x faster

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PSI-BLAST

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

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BLAST Programs

 blastp compares an amino acid query sequence against a protein sequence database.

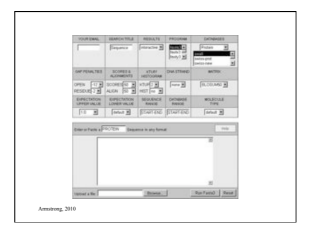
 blastn compares a nucleotide query sequence against a nucleotide sequence database.

• 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 dynamically translated in all reading

 tblastx compares the six-frame translations of a nucleotide query sequence against the six-frame translations of a nucleotide sequence database. (SLOW)

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Alignment Heuristics

- · Dynamic Programming is better but too slow
- BLAST (and FASTA) 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, pretty good for DNA, fastest, now dominant.

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Biological Databases (sequences)

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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

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DNA Sequence Databases

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

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Main DNA DBs

Genbank USEMBL EUDDBJ Japan

• Celera genomics Commercial DB

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EMBL

- Sources for sequence include:
 - Direct submission on-line submission tools
 - Genome sequencing projects
 - Scientific Literature DB curators and editorial imposed submission
 - Patent applications
 - Other Genomic Databases, esp Genbank

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International Nucleotide Sequence Database Collaboration

- Partners are EMBL, Genbank & DDBJ
- Each collects sequence from a variety of sources
- New additions to any of the three databases are shared to the others on a daily basis.

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Limited annotation

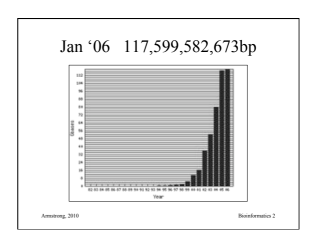
- Unique accession number
- Submitting author(s)
- Brief annotation if available
- Source (cDNA, EST, genomic etc)
- · Species
- · Reference or Patent details

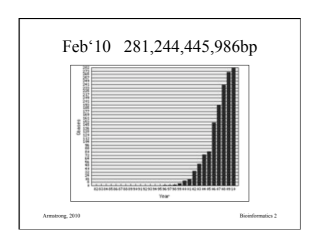
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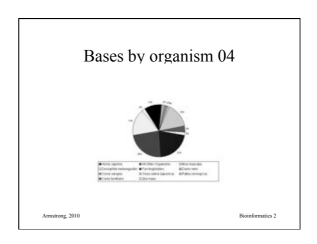
EMBL file tags

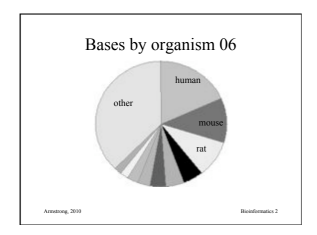


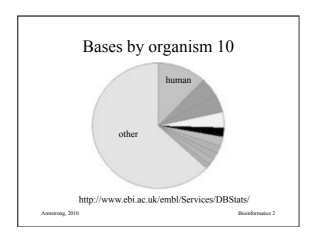
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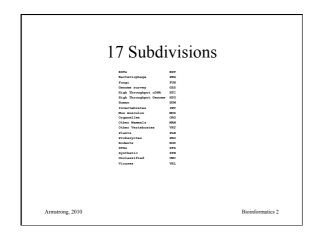












Specialist DNA Databases

- Usually focus on a single organism or small related group
- Much higher degree of annotation
- Linked more extensively to accessory data
 - Species specific:
 - Drosophila: FlyBase,
 - C. elegans: AceDB
 - Other examples include Mitochondrial DNA, Parasite Genome DB

FlyBase

flybase.bio.indiana.edu

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

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Protein DBs

- Primary Sequence DBs
 - UniProt, TrEMBL, GenPept
- Protein Structure DBs
 - PDB, MSD
- Protein Domain Homology DBs
 - InterPro, CluSTr

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UniProtKB/Swiss-Prot

- Consists of protein sequence entries
- · Contains high-quality annotation
- · Is non-redundant
- Cross-referenced to many other databases
- 104,559 sequences in Jan 02
- 120,960 sequences in Jan 03
- 514,789 sequences in Feb 10 (latest)

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Swis-Prot by Species ('03)

Swis-Prot by Species (Oct '05)

```
Number Frequency Species

1 12860 Homo sapiens (Human)
2 9933 Mus musculus (Mouse)
3 5139 Saccharomyces cerevisiae (Baker's yeast)
4 4846 Escherichia coll
5 4570 Rattus norvegicus (Rat)
6 3609 Arabidopsis thaliana (Mouse-ear cress)
7 2840 Schizosaccharomyces pombe (Fission yeast)
8 2212 Ratillus subrillas
9 2867 Caenorhabditis elegans
10 2257 Caenorhabditis elegans
11 277 Decembrilla malanogaster (Fruit fly)
12 1772 Mesemphila malanogaster (Fruit fly)
13 1758 Escherichia coll ol57:H7
14 1653 Bos taurus (Bovine)
15 1512 Salmonella typhimurium
```

Swis-Prot by Species (Oct '05)

UniProtKB/TrEMBL

- Computer annotated Protein DB
- Translations of all coding sequences in EMBL DNA Database
- Remove all sequences already in Swiss-Prot
- November 01: 636,825 peptides
- Feb 10: 10,376,872 peptides
- TrEMBL is a weekly update
- ArmsGenPept is the Genbank equivalent Bioinformatics 2

SNPs

- Biggest growth area right now is in mutation databases
- www.ncbi.nlm.nih.gov/About/primer/ snps.html
- Polymorphisms estimates at between 1:100 1:300 base pairs (normal human variation)
- Databases include true SNPs (single bases) and larger variations (microsatellites, small

 Amagnetics)

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dbSNP

- "The database grows at 90 SNPs per month"
- 130 versions since start in 1998
- Currently 156 million SNPs in v130
- 23 million added between version 129 and 130!

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Database Search Methods

- Text based searching of annotations and related data: SRS, Entrez
- Sequence based searching: BLAST, FASTA, MPSearch

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SRS



- Sequence Retrieval System
 - Powerful search of EMBL annotation
 - Linked to over 80 other data sources
 - Also includes results from automated searches

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SRS data sources

- Primary Sequence: EMBL, SwissProt
- References/Literature: Medline
- Protein Homology: Prosite, Prints
- Sequence Related: Blocks, UTR, Taxonomy
- Transcription Factor: TFACTOR, TFSITE
- Search Results: BLAST, FASTA, CLUSTALW
- Protein Structure: PDB
- Also, Mutations, Pathways, other specialist DBs

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Entrez

- Text based searching at NCBI's Genbank
- Very simple and easy to use
- Not as flexible or extendable as SRS
- No user customisation

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Sequence Based Searching

· Queries:

DNA query against DNA db Translated DNA query against Protein db Translated DNA query against translated DNA db Translated Protein query against DNA db Protein query against Protein db

· BLAST & FASTA

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Secondary Databases

- PDB
- Pfam
- PRINTS
- PROSITE
- ProDom
- SMART
- TIGRFAMs

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PDB

- Molecular Structure Database (EBI)
- Contains the 3D structure coordinates of 'solved' protein sequences
 - X-ray crystallography
 - NMR spectra
- 19749 protein structures

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Multiple Sequence Alignment

- What and Why?
- Dynamic Programming Methods
- Heuristic Methods
- A further look at Protein Domains

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Multiple Alignment

- · Normally applied to proteins
- Can be used for DNA sequences
- Finds the common alignment of >2 sequences.
- Suggests a common evolutionary source between related sequences based on similarity
- Can be used to identify sequencing errors
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Multiple Alignment of DNA

- Take multiple sequencing runs
- · Find overlaps
 - variation of ends-free alignment
- Locate cloning or sequencing errors
- Derive a consensus sequence
- Derive a confidence degree per base

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Consensus Sequences

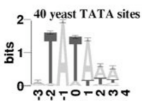
- Look at several aligned sequences and derive the most common base for each position.
 - Several ways of representing consensus sequences
 - Many consensus sequences fail to represent the variability at each base position.
 - Largely replaced by Sequence Logos but the term is often misapplied.

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Sequence Logos

• Example, from an alignment of the TATA box in yeast genes:

We now have a confidence level for each base at each position



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Multiple Alignment of Proteins

- Multiple Alignment of Proteins
- Identify Protein Families
- · Find conserved Protein Domains
- Predict evolutionary precursor sequences
- · Predict evolutionary trees

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Protein Families

- Proteins are complex structures built from functional and structural sub-units
 - When studying protein families it is evident that some regions are more heavily conserved than others.
 - These regions are generally important for the structure or function of the protein
 - Multiple alignment can be used to find these regions
- These regions can form a signature to be used Armstrong 1701 Identifying the protein family or functional

Protein Domains

- Evolution conserves sequence patterns due to functional and structural constraints.
- Different methods have been applied to the analysis of these regions.
- Domains also known by a range of other names:

motifs patterns prints blocks

Multiple alignment

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Multiple Alignment

- OK we now have an idea WHY we want to try and do this
- What does a multiple alignment look like?
- · How could we do multiple alignments
- What are the practical implications

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Multiple alignment table

dlg_CG1725-PH Sap97_dlgh1 chapsyn-110_dlgh2 Sap102_dlgh3 PSD-95_dlgh4

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

Conservatived or Identical residues are colour coded

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Scoring Multiple Alignments

 We need to score on columns with more than 2 bases or residues:

ColumnCost
$$\begin{pmatrix} S \\ C \\ A \\ P \\ P \end{pmatrix} = 24$$

Multiple alignments are usually scored on cost/difference rather than similarity

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Column Costs

- Several strategies exist for calculating the column cost in a multiple alignment
- Simplest is to sum the pairwise **costs** of each base/residue pair in the column using a matrix (e.g. PAM250).
- Gap scoring rules can be applied to these as well.

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Scoring Multiple Alignments

• Score = (S,C)+(S,A)+(S,A)+(S,P)+(S,P)+ (C,A)+(C,P)+(C,P)+(A,P)+(A,P)+(P,P)



Known as the sum-of-pairs scoring method

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Sum-of-pairs cost method (SP)

• Score = (S,C)+(S,-)+(S,A)+(S,P)+(S,P)+(S,P)+(S,A)+(S,P)+

ColumnCost
$$\begin{pmatrix} S \\ -A \\ P \\ P \end{pmatrix} = 24$$

Still works with gaps using whatever gap penalty you want

Multiple Alignment Cost

- Sum of pairs is a simple method to get a score for each column in a multiple alignment
- Based on matrices and gap penalties used for pairwise sequence alignment
- The score of the alignment is the sum of each column

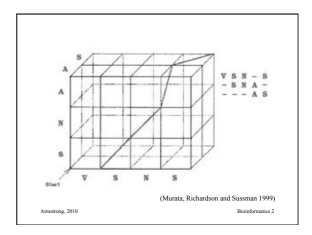
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Optimal Multiple Alignment

- The best alignment is generally the one with the lowest score (i.e. least difference)
 - depends on the scoring rules used.
- Like pairwise cases, each alignment represents a path through a matrix
- For multiple alignment, the matrix is *n*-dimensional
 - where *n*=number of sequences

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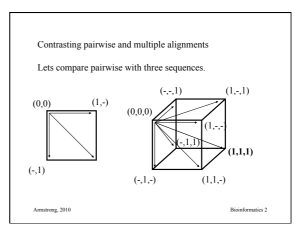


Contrasting pairwise and multiple alignments

Lets compare pairwise with three sequences.



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Multiple alignment table

dlg_CG1725-PH Sap97_dlgh1 chapsyn-110_dlgh2 Sap102_dlgh3 PSD-95_dlgh4

ALFDYDPNRDDGLPSRGLPFKH ALFDYDKTKDSGLPSQGLNFRF AMFDYDKSKDSGLPSQGLSFKY ALFDYDRTRDSCLPSQGLSFSY ALFDYDKTKDCGFLSQALSFHF *:**** : * : *: : *:* :

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

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Gene and Protein Prediction

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Gene prediction

- What is a gene?
 - Simple definition: A stretch of DNA that encodes a protein and includes the regulatory sequences required for temporal and spatial control of gene transcription.
- Characteristics of genes.
 - What genetic features can we use to recognise a gene?

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DNA structure



Bases: A,C,G and T

Chemically, A can only pair with T and G with C

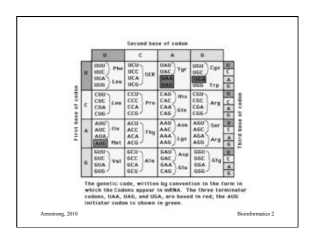
Two strands, 5' and 3' Genes are encoded along one side of the DNA molecule. The 5' end being at the left hand side of the gene.

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Codons and ORFs

- Three bases that encode an amino acid or stop site.
- A run of valid codons is an Open Reading Frame.
- An ORF usually starts with a Met
- Ends with a nonsense or stop codon.

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12

Predicting ORFs

- 64 total codons
- 3 stop codons, 61 codons for amino acids
- Random sequence 1:21 ratio for stop:coding.
- = 1 stop codon every 63 base pairs
- Gene lengths average around 1000 base pairs.

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Finding ORFs

- One algorithm slides along the sequence looking stop codons.
- · Scans back until it finds a start codon.
- Fails to find very short genes since it it looking for long ones
- · Also fails to find overlaping ORFs
- There are many more ORFs than genes

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Amino Acid Bias

- The amino acids in proteins are not random
 - leucine has 6 codons
 - alanine has 4 codons
 - tryptophan has 1 codon
- The random the ratio would be 6:4:1
- In proteins it is 6.9:6.5:1
 - i.e. it is not random

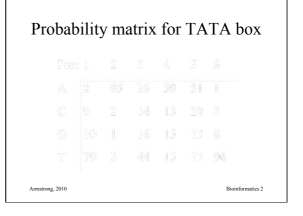
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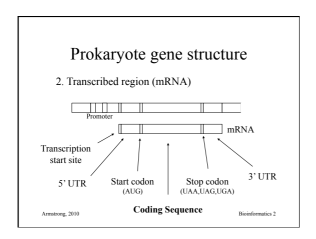
Gene Prediction

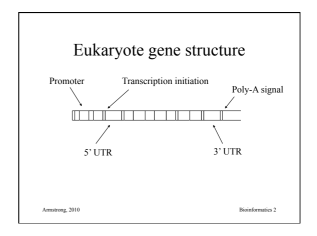
- · Take all factors into consideration
- · Prokaryotes
 - No Nucleus
 - 70% of the genome encodes protein
 - No introns

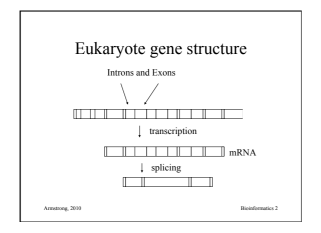
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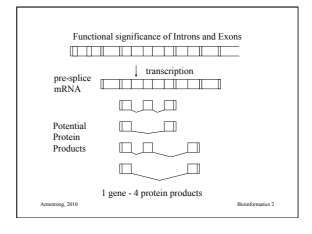
Prokaryote gene structure 1. Promoter region nnn<u>TTGACA</u>nnnnnnnnnnnnnnnnn<u>TATAAT</u>nnnnnnS (consensus sequence for *E.coli*.) Armstrong, 2010 Bioinformatics 2

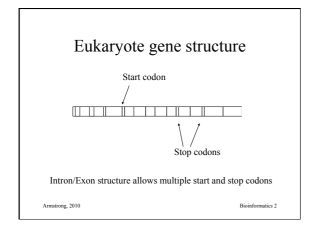












HMMs for codons

- Model based on examining 6 consecutive bases (i.e. all three reading frames).
- Based on statistical differences between coding and non coding regions
- 5th order Markov Model.
- Given 5 preceding bases, what is the probability of the 6th?
- Homogenous model (ignores reading frame)

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HMMs for codons

- Homogenous models have two tables, one for coding, one non coding.
- Each table is has 4096 entries for the potential 6 base pair sequences
- Non-homogenous models have three tables for possible reading frames
- Short exons cause these models problems
- Hard to detect splice sites

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Glimmer

- Uses non-homogenous HMMs to predict prokaryote gene sequences
- · Identifies ORFs
- Trains itself on a prokaryote genome using ORFs over 500 bp
- http://www.cs.jhu.edu/labs/compbio/glimmer.html

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Predicting Splice Sites

- There are some DNA features that allow splice sites to be predicted
- These are often species specific
- They are not very accurate.

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NetGene2

- Neural network based splice site prediction
- Trained on known genes
- Claims to be 95% accurate
- Human, C. elegans & Arabidopsis thaliana
- http://www.cbs.dtu.dk/services/NetGene2/

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HMMgene

- Based on an HMM model of gene structure
- Predicts intron/exon boundries
- Predicts start and stop codons
- Known information can be added (e.g. from ESTS etc)
- · Outputs in GFF format

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GFF Format

- Exchange format for gene finding packages
- Fields are:
 - < seqname > name, genbank accession number
 - <source> program used
 - <feature> various inc splice sites
 - <start> start of feature

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GFF Format

- <end> end of feature
- <score> floating point value
- − <strand> +, (or .. for n/a)
- < frame > 0,1 or 2

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GenScan

- Probabilistic model for gene structure based on a general HMM
- Can model intron/exon boundries, UTRs, Promoters, polyA tails etc
- http://genes.mit.edu/GENSCAN.html

ng 2010

Given a new protein sequence...

- What is the function?
- Where is the protein localised?
- What is the structure?
- What might it interact with?

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Given a new protein sequence...

- What is the function?
- Have we seen this protein or a very similar one before?
 - If yes then we can infer function, structure, localisation and interactions from homologous sequence.
- Are there features of this protein similar to

 Armothers?

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Protein Families

- Proteins are complex structures built from functional and structural sub-units
 - When studying protein families it is evident that some regions are more heavily conserved than others.
 - These regions are generally important for the structure or function of the protein
 - Multiple alignment can be used to find these regions
- These regions can form a signature to be used Armstrong 170 of dentifying the protein family or function are

Protein Domains

- Evolution conserves sequence patterns due to functional and structural constraints.
- Different methods have been applied to the analysis of these regions.
- Domains also known by a range of other names:

motifs patterns prints

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patterns prints

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16

Profiles

- Given a sequence, we often want to assign the sequence to a family of known sequences
- We often also want to assign a subsequence to a family of subsequences.

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Profiles

- Examples include assigning a gene/protein to a known gene/protein family, e.g.
 - G coupled receptors
 - actins
 - globins

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Profiles

- Also we may wish to find known protein domains or motifs that give us clues about structure and function
 - Phosphorylation sites (regulated site)
 - Leucine zipper (dna binding)
 - EGF hand (calcium binding)

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Creating Profiles

- Aligning a sequence to a single member of the family is not optimal
- Create profiles of the family members and test how similar the sequence is to the profile.
- A profile of a multiply aligned protein family gives us letter frequencies per column.

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Matching sequences to profiles

- We can define a distance/similarity cost for a base in each sequence being present at any location based on the probabilities in the profile.
- We define define costs for opening and extending gaps in the sequence or profile.
- Therefore we can essentially treat the alignment of a sequence to a profile as a Ampairwise alignment and use dynamic Bioinformatics 2

Protein profiles

- Multiple alignments can be used to give a consensus sequence.
- The columns of characters above each entry in the consensus sequence can be used to derive a table of probabilities for any amino acid or base at that position.

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Protein profiles

- The table of percentages forms a profile of the protein or protein subsequence.
- With a gap scoring approach sequence similarity to a profile can be calculated.
- The alignment and similarity of a sequence / profile pair can be calculated using a dynamic programming algorithm.

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Protein profiles

- Alternative approaches use statistical techniques to assess the probability that the sequence belongs to a family of related sequences.
- This is calculated by multiplying the probabilities for amino acid *x* occurring at position *y* along the sequence/profile.

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Tools for HMM profile searches

- Meme and Mast at UCSD (SDSC)
- http://meme.sdsc.edu/
- MEME
 - input: a group of sequences
 - output: profiles found in those sequences
- MAST
 - input: a profile and sequence database
- output: locations of the profile in the database

Summary

- Multiple alignment is used to define and find conserved features within DNA and protein sequences
- Profiles of multiply aligned sequences are a better description and can be searched using pairwise sequence alignment.
- Many different programs and databases available.

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Secondary Databases

- PDB
- Pfam
- PRINTS
- PROSITE
- ProDom
- SMART
- TIGRFAMs

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PDB



- Molecular Structure Database
- Contains the 3D structure coordinates of 'solved' protein sequences
 - X-ray crystallography
 - NMR spectra
- 29429 protein structures

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SUPERFAMILY is a library of profile hidden Markov models that represent all proteins of known structure, based on SCOP.

The SCOP database aims to provide a detailed and comprehensive description of the structural and evolutionary relationships between all proteins whose structure is known (based on PDB)

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Pfam



- Database of protein domains
- Multiple sequence alignments and profile HMMs
- · Entries also annotated
- · Swiss-Prot DB all pre-searched
- New sequences can be searched as well.
 - 7973 entries in Pfam last update

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- Database of 'protein fingerprints'
- Group of motifs that combined can be used to characterise a protein family
- ~11,000 motifs in PRINTS DB
- Provide more info than motifs alone

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'linear' motifs

- · Not all protein motifs are easy to find
- Linear motifs involved in protein-protein interactions
 - Very degenerate
 - Found in specific regions of proteins
 - Require special treatment
 - Neduva et al, PLOS 2005

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Linking it all together...

- Database Searches
 - Multiple Alignments
 - Find known motifs and domains
 - Find possible similar folds
- · Prediction algorithms
 - Properties of amino acids
 - Predicting folding
 - Finding cysteine bonds

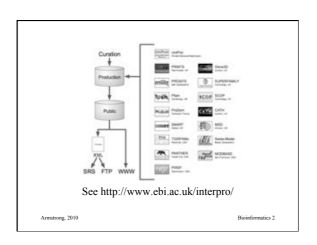
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InterPro

- EBI managed DB
- Incorporates most protein structure DBs
- Unified query interface and a single results output.

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InterPro

DATABASE	VERSION	ENTRIES
SWISS-PROT	48	197228
PRINTS	38	1900
TREMBL	31.1	2342938
PFAM	18	7973
PROSITE	19.10	1882

Currently 15 databases, plans to add 3 new ones this

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PredictProtein



http://www.embl-heidelberg.de/predictprotein/

Database searches:

- generation of multiple sequence alignments (MaxHom)
- detection of functional motifs (PROSITE)
- detection of composition-bias (SEG)
- detection of protein domains (PRODOM)
- fold recognition by prediction-based threading (TOPITS)

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PredictProtein

Predictions of

- secondary structure (PHDsec, and PROFsec)
- residue solvent accessibility (PHDacc, and PROFacc)
- transmembrane helix location and topology (PHDhtm PHDtopology)
- protein globularity (GLOBE)
- coiled-coil regions (COILS)
- cysteine bonds (CYSPRED)
- structural switching regions (ASP)

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Data and methods in Data and methods in Data and methods in Data and methods in Add data and programs run at centrar sine and updated on a regular basis Armstrong, 2010 Bioinformatics 2

Too many programs/databases

- How do we keep track of our own queries?
 - Repeat an old query
 - Run the same tests on a new sequence
 - Run 100s of sequences..
 - Document the process for a paper or client or for quality assurance

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Workflow managers

- Locate and manage connections to software and databases
- · Record actions
- Replay a workflow at a later date or against multiple sequences
- Manages redundant external sources (e.g. multiple blast servers)
- Can connect to specialist local sources

 Armstrong, 2010



- http://taverna.sourceforge.net/
- Open source and free to download
- Runs on PC/linux/mac
- Drag-n-Drop interface to bioinformatics Am**analy**sis

