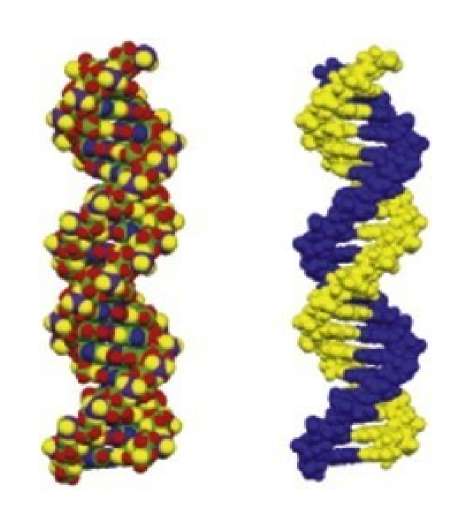
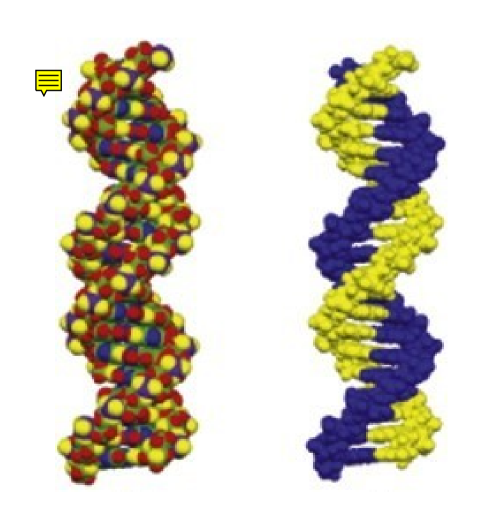
Lab closing Bioinformatics 1 students only



Bioinformatics 1 Lecture 3 Sequencing and Sequence Alignment Basics



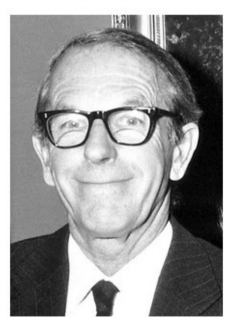
Today's topics

- DNA sequencing
- Sequence alignment, fundamentals
- BLAST

DNA Sequencing



Frederick Sanger - Facts



Frederick Sanger

Born: 13 August 1918, Rendcombe, United Kingdom

Affiliation at the time of the award: MRC Laboratory of Molecular Biology, Cambridge, United Kingdom

Prize motivation: "for their contributions concerning the determination of base sequences in nucleic acids"

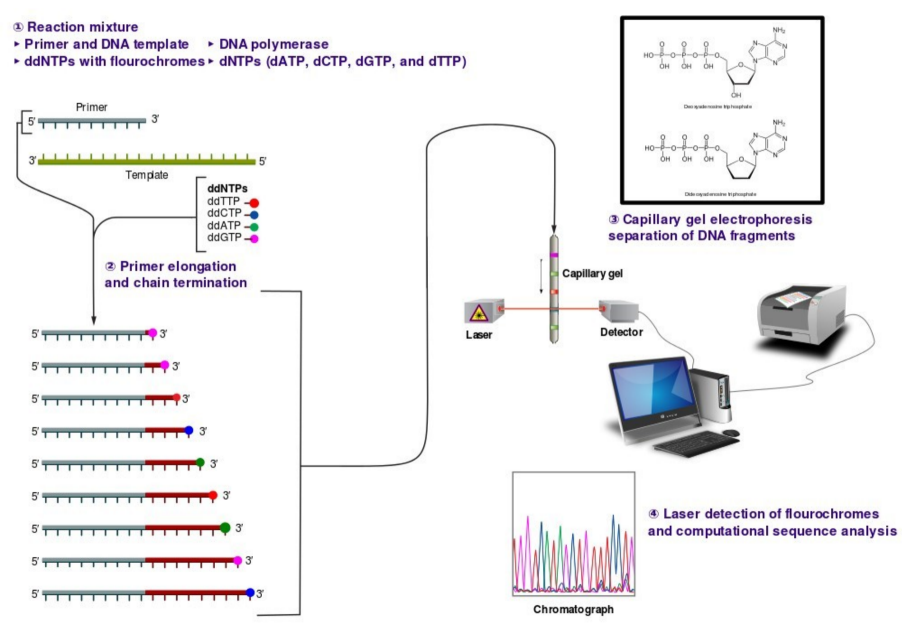
Field: biochemistry

DNA sequencing with chain-terminating inhibitors

F Sanger, S Nicklen... - Proceedings of the ..., 1977 - National Acad Sciences
Abstract A new method for determining nucleotide sequences in DNA is described. It is
similar to the "plus and minus" method [Sanger, F. & Coulson, AR (1975) J. Mol. Biol. 94, 441448] but makes use of the 2', 3'-dideoxy and arabinonucleoside analogues of the ...
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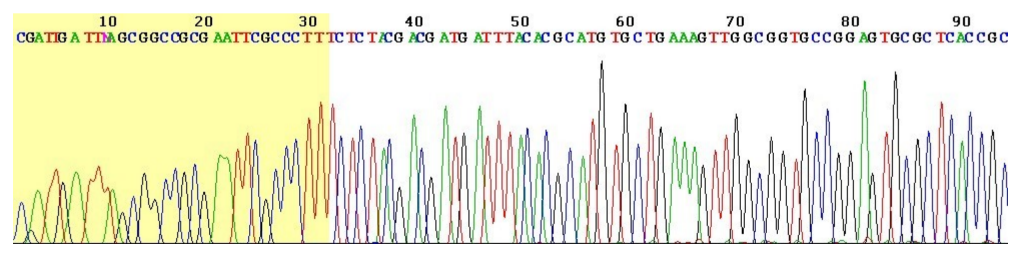
Sanger chain termination method



https://en.wikipedia.org/wiki/Sanger_sequencing

Sanger sequencing





Problem: only short stretches (1000bp) of DNA are sequenced in any one run

The problem of sequence assembly can be compared to taking many copies of a book, passing each of them through a shredder with a different cutter, and piecing the text of the book back together just by looking at the shredded pieces. Besides the obvious difficulty of this task, there are some extra practical issues: the original may have many repeated paragraphs, and some shreds may be modified during shredding to have typos. Excerpts from another book may also be added in, and some shreds may be completely unrecognisable.

Main strategies

• HGP

- create short (100-300kb) Bacterial Artificial Chromosomes (BAC) with overlapping DNA and amplify
- Use repeated shotgun Sequencing: cut DNA up and sequence
 - ++ reliable -- slow -- expensive (\$3bn)

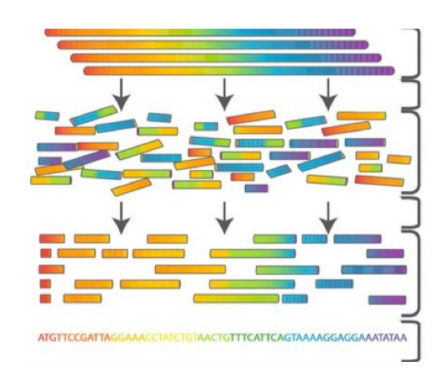


Celerea/Venter



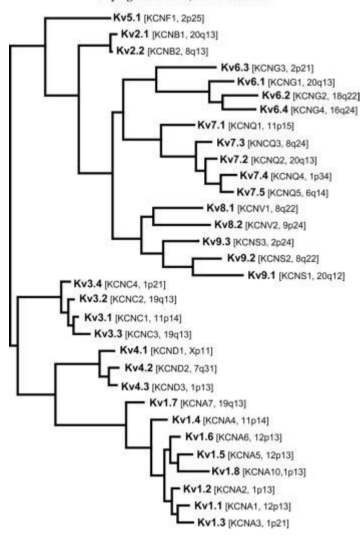
- blast whole genome into 2-10kb fragments and re-assemble in computer
 - - error prone/requires extra data ++ cheap (\$300Mio)

Sequence alignment

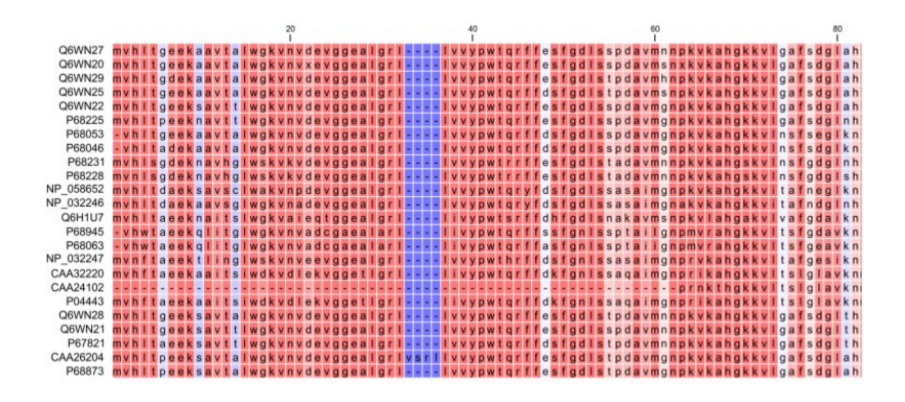


Sequence alignment





Multiple sequence alignments



Interpretation of alignments

- Sequence identity: exactly the same sequence
- Similarity: sufficiently high score, but scoring should have some biological relevance
- Homology: two sequences have a common ancestor

"Homology is the central concept for all of biology. Whenever we say that a mammalian hormone is the 'same' hormone as a fish hormone, that a human gene sequence is the 'same' as a sequence in a chimp or a mouse, that a HOX gene is the 'same' in a mouse, a fruit fly, a frog, and a human - even when we argue that discoveries about a worm, a fruit fly, a frog, a mouse, or a chimp have relevance to the human condition - we have made a bold and direct statement about homology. The aggressive confidence of modern biomedical science implies that we know what we are talking about. But a deeper reflection shows that this confidence is based more on hope than on certainty."

Wake, Comparative terminology. Science 1994, 265:268-269

Graphical sequence comparison (preparation)

Programmatic access to sequences:

```
from Bio import Entrez
gene_id = "some_id"
record = Entrez.efetch(db="nucleotide", id=gene_id1, rettype="gb",
retmode="text")
```

The record object has a number of attributes:

```
    id - Identifier such as a locus tag (string)
    seq - The sequence itself (Seq object or similar)
    name - Sequence name, e.g. gene name (string)
    description - Additional text (string)
    dbxrefs - List of database cross references
    features - Any (sub) features defined
    annotations - Further information about the whole sequence.
    Most entries are strings, or lists of strings.
```

Graphical sequence comparison

- Find a pair of genes to compare. You can choose two species, or two different genes. Get their NCBI IDs.
- Insert these into cell #2 in this Jupyter notebook:

https://www.inf.ed.ac.uk/teaching/courses/bio1/lectures17/Bio1Lecture3GraphAlign.ipynb

HTML version:

https://www.inf.ed.ac.uk/teaching/courses/bio1/lectures17/Bio1Lecture3GraphAlign.html

Alignment scoring

```
(a)
HBA HUMAN
           GSAOVKGHGKKVADALTNAVAHVDDMPNALSALSDLHAHKL
           G+ +VK+HGKKV A+++++AH+D++ +++++LS+LH
           GNPKVKAHGKKVLGAFSDGLAHLDNLKGTFATLSELHCDKL
HBB HUMAN
(b)
HBA HUMAN GSAQVKGHGKKVADALTNAVAHV - - - D - - DMPNALSALSDLHAHKL
           ++ ++++H+ KV
                           + + A + +
                                              +L+L++H+K
LGB2 LUPLU NNPELQAHAGKVFKLVYEAAIQLQVTGVVVTDATLKNLGSVHVSKG
(c)
HBA HUMAN
           GSAOVKGHGKKVADALTNAVAHVDDMPNALSALSD - - - - LHAHKL
           GS+ + G +
                        +D L ++ H+ D+ A +AL D
                                                    ++AH+
F11G11.2
           GSGYLVGDSLTFVDLL - - VAOHTADLLAANAALLDEFPOFKAHOE
```

Figure 2.1 Three sequence alignments to a fragment of human alpha globin. (a) Clear similarity to human beta globin. (b) A structurally plausible alignment to leghaemoglobin from yellow lupin. (c) A spurious high-scoring alignment to a nematode glutathione S-transferase homologue named F11G11.2.

Alignment scoring

- Match: identical residues
- Substitution: different residues
- Insert/deletion (indel, gap): a gap in one of the two sequences
- Sequence alignment attempts to identify sequences originating from a common ancestor
- The functional implications of each sequence difference is important
- Appropriate scoring schemes take this into account (more on this later in the course)

Alignment scoring

- Example scoring:
 - Match +2
 - Substitution -3
 - Gap -4

NW Sco 120	ore	Identities 69/75(92)	%)	Gaps 0/75(0%)	Strand Plus/Plus	
Query	1	ATGGACAATGCAAGAA	TGAACTCCTTCCTG	GAATACCCCATACT	TAGCAGTGGCGACTCG	60
Sbjct	1	ATGGACAATGCAAGAA	tgyyctccttcctg	BAATACCCCATCCT	CAGCGGAGGCGACTCT	60
Query	61	GGGACCTGCTCAGCC				
Sbjct	61	GGGACCTGCTCAGCG	75			

Homo sapiens homeobox A1 (HOXA1), transcript variant 1, mRNA.

VS.

Elephantulus edwardii homeobox A1 (HOXA1), mRNA (Cape elephant shrew)

Optimal Sequence Alignment

- Small sequences can be compared exactly using dynamic programming.
 - → more on this next week
- When searching multiple genomes, exact methods are computationally too expensive.

Solution: heuristic methods

Basic Local Alignment Search Tool

- Compares a query sequence with a full sequence database
- Avoids full alignments through heuristics
- Main assumption: 'true' matches have at least some short highscoring stretches
- Developed by Altschul, Gish, Miller and Myers at NIH in 1980s

Basic local alignment search tool

<u>SF Altschul</u>, W Gish, <u>W Miller</u>, <u>EW Myers</u>... - Journal of molecular ..., 1990 - Elsevier A new approach to rapid sequence comparison, basic local alignment search tool (BLAST), directly approximates alignments that optimize a measure of local similarity, the maximal segment pair (MSP) score. Recent mathmatical results on the stochastic properties of ... Cited by 62090 Related articles All 105 versions Web of Science: 43853 Cite Save

Based on FASTA, which combined DP alignment with heuristics

Rapid and sensitive protein similarity searches

DJ Lipman, <u>WR Pearson</u> - Science, 1985 - sciencemag.org

Abstract An algorithm was developed which facilitates the search for similarities between newly determined amino acid sequences and sequences already available in databases.

Because of the algorithm's efficiency on many microcomputers, sensitive protein database ...

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Strategy

- Remove redundant/low complexity regions from query (SEG or DUST)
- Create set of short words
- Scan database for matching words (seeds)
- Matching words are extended (gap-free), similar to Smith-Waterman
- Extensions below specified threshold score are removed
- Remaining extensions are completed also scoring indels

Step 1: creating short words

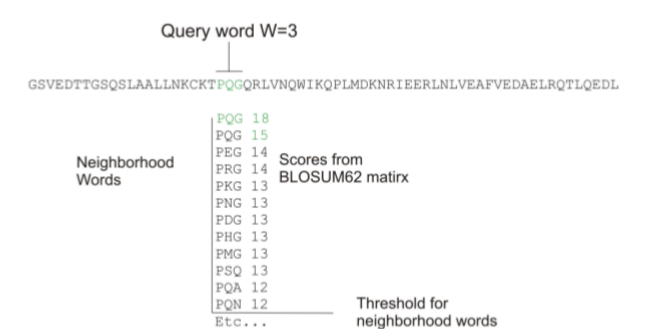
```
Query word W=3

GSVEDTTGSQSLAALLNKCKTPQGQRLVNQWIKQPLMDKNRIEERLNLVEAFVEDAELRQTLQEDL

KCK
CKT
KTP
TPQ
PQG
```

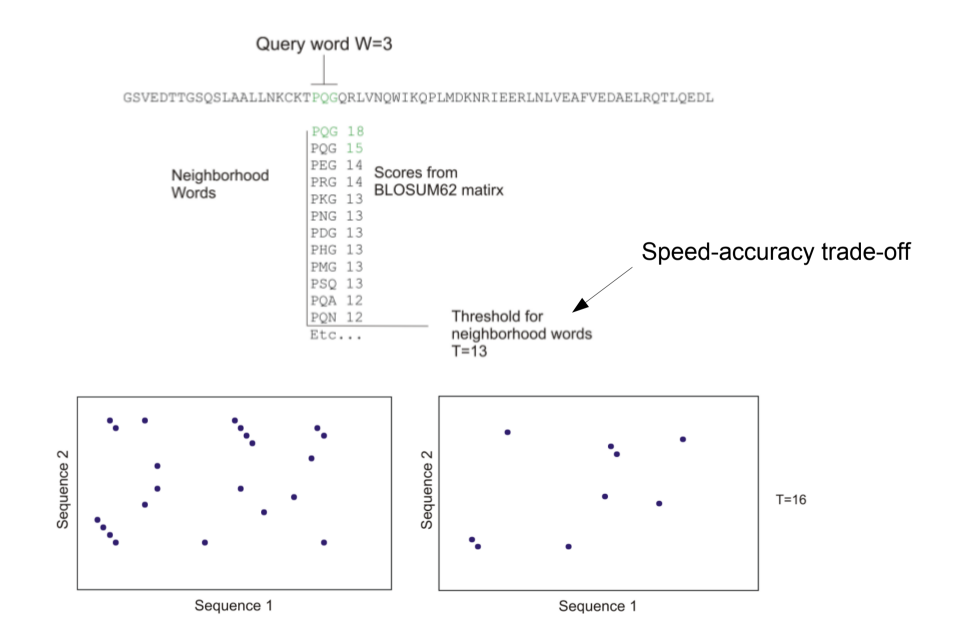
(n=3 for amino acids, n=11 for nucleotides)

Step 1.1: create neighbourhood words

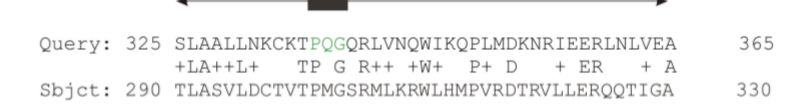


T=13

Step 2: search words in database



Step 3: seed extension



Scoring without indels, extend until threshold reached.

Step 4: thresholding and scoring

- Compare scores with random surrogates of same length, select a significance threshold
- Keep Maximal Scoring Pairs
- Compute significance through comparison with random sequence (using Gumbel distribution that models number of extreme samples in a distribution)
- Output:
 - Raw scores
 - Bit score (sometimes)
 - E value

Scoring

Bit score
 a scaled version of alignment score S

$$S' = \frac{\lambda S - \ln K}{\ln 2}$$

λ and K are parameters of the Grumbel distribution, which describes score probabilities for S-W Algorithm.

E-value

$$E = mn2^{-S'}$$

n query size, m database size

Yields expected number chance alignments with score S' or better = estimate of false positives.

Uses

- Identify species or homologous species in new data
- Search for protein domains
- Phylogeny
- Map chromosome locations
- Map annotations between species

BLAST flavours

- MegaBLAST: similar N-N searches
- BLASTN: more dissimilar N-N
- BLASTP: P-P comparison
- BLASTX: N-P comparison (translation done on the fly)
- BLASTN: P-N comparison (back-translation done on the fly)

NCBI BLAST

Use this code to run a BLAST search:

https://www.inf.ed.ac.uk/teaching/courses/bio1/lectures17/Bio1Lecture3BLAST.ipynbhttps://www.inf.ed.ac.uk/teaching/courses/bio1/lectures17/Bio1Lecture3BLAST.html

- Try manipulating the query sequence, e.g. make indels, reverse the sequence, or insert a piece of different sequence.
- The notebook runs a nucleotide BLAST. Try to modify it to run a protein BLAST.
- Inspect to what extent the E-value reflects biologically plausibility.

Summary

Exact alignment algorithms are too slow to deal with big genomic data search problems.

Algorithms combined with heuristics are highly efficient, but less precise.

The bad

In 1997, the discovery of a new plant adenylyl cyclase gene was published [35]. This was a profound finding because plants were not believed to have adenylyl cyclases. The authors went on to suggest a whole new type of biochemistry for plants. The 'homology' (sequence similarity) they showed was not so weak: there was definitely some similarity, and the homology had a high 'score' (which by itself is not very meaningful) - but when their adenylyl cyclase was aligned to a profile for other known adenylyl cyclases, it was obvious to even first-year graduate students that the characteristics that are common to all other adenylyl cyclases were largely missing.

The ugly

The authors were later forced to retract their paper [36]. What might have saved them from public humiliation was a more careful analysis of their results.

Pertsemlidis, Alexander, John W. Fondon, and W. John. "Having a BLAST with bioinformatics (and avoiding BLASTphemy)." Genome Biol 2.10 (2001).

http://www.ncbi.nlm.nih.gov/pmc/articles/PMC138974/