Bioinformatics Algorithms

(Fundamental Algorithms, module 2)

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Database search with BLAST (summary)

Database search

- Until now: compare two sequences
 - how similar/different are they? (score/value)
 - where are the similarities/differences? (alignment)

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Database search

- Until now: compare two sequences
 - how similar/different are they? (score/value)
 - where are the similarities/differences? (alignment)
- Now: compare one sequence to a database (i.e. to many sequences)

Database search

Goal

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Identifying sequences in the DB which have high local similarity with the query.

- We know how to do this: Smith-Waterman DP-algorithm.
- But: too slow

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Say all sequences have length n (query t and all DB seq's), and there are r sequences in the DB.

• exact solution (Smith-Waterman): $O(r \cdot n^2)$

Example

- UniProt/SwissProt (protein database): 548 454 sequences, 195 409 447 aa's (avg. length 350 aa's)

 version 29/04/15
- NCBI Genbank (nucleotide database): 182 188 746 sequences, 189 739 230 107 nucleotides (avg. length 1041 nucl.) April 2015, no WGS

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So we would get something like $350\cdot 350\cdot 548454=67\,185\,615\,000=$ about 67 billion $(67\cdot 10^9)$ steps, which takes 18 hours on a computer that performs 1 million operations per second (for UniProt), and $197\,434\,482\,454\,026~(\approx 1.9\cdot 10^{12}),$ about 6 years, for Genbank. And still about 1 hour on a computer performing 1 billion operations per second.

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And this is for one query only!

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Basic idea

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If there is a good local alignment between two sequences, then this local alignment is likely to contain two short substrings with high score when aligned without gaps.

Basic steps of BLAST

- 1. create list of high-scoring words with query
- 2. scan DB for these words (called seeds)
- extend seeds in both directions to form good local alignment (these are called MSPs = maximum segment pairs)

BLAST then gives a significance score to the MSPs and only retains them if above a certain threshold.

(for an example, see class notes)

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The NCBI BLAST website

- Different versions of BLAST, depending on the task (nucl-nucl: blastn, megablast, ..., prot-prot: blastp, psi-blast, nucl-prot: blastx, prot-nucl: tblastn, ...)
- Different databases (nucl vs. prot, different organisms, different types of db, different levels of assembly, . . .)
- Very good explanations and help pages!

BLAST: Basic Local Alignment Search Tool

- Altschul et al. 1990, 1997
- looks for sequences in a database with high local similarity to query
- heuristic algorithm
- solid mathematical foundations (Karlin-Altschul statistics)
- extremely successful, now the database search tool ("to blast a sequence against a database")
- NCBI¹ Blast at: http://blast.ncbi.nlm.nih.gov/Blast.cgi

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BLAST2

Some of the main changes in BLAST2 (Altschul el al. 1997)

- start with two seeds instead of one, not too far apart
- gapped alignments
- extension of statistical theory to HSPs (high-scoring segment pairs)

Note: All versions of BLAST include many complex pre- and postprocessing steps, optimizations, ...These are explained in the cited papers, and followup publications. Here we are looking only at the basic ideas underlying the algorithm.

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 $^{^{1}}$ NCBI = National Center for Biotechnology Information