CSEP 590A Computational Biology Autumn 2008

> Lecture 2 Sequence Alignment

Tonight

Last week's "quiz" & homework Sequence alignment Weekly "bio" interlude - DNA replication More sequence alignment

Week 1 (anonymous) "Quiz"

In your own words, what is DNA? Its main role? What is RNA? What is its main role in the cell?

How many amino acids are there? How many are used in proteins?

Did human beings, as we know them, develop from earlier species of animals?

What are stem cells?

What did Viterbi invent?

What is dynamic programming?

What is a likelihood ratio test?

What is the EM algorithm?

Don't worry, we'll talk about all this stuff before the course ends

How would you find the maximum of f(x) = ax3 + bx2 + cx + d in the interval -10<x<25?

Evolution & Scientific Literacy

"Human beings, as we know them, developed from earlier species of animals"

(avoiding the now politically charged word "evolution")

From 1985 to 2005, the % of Americans

rejecting: declined from 48% to 39% accepting: also declined 45% to 40 uncertain: increased 7% to 21%

In a 2005 survey, the proportion of adults who accept evolution in 34 countries (US, Europe, Japan...), the United States ranked 33rd, just above/below Turkey.

Sequence Alignment

Part I Motivation, dynamic programming, global alignment

Sequence Alignment

What Why A Simple Algorithm Complexity Analysis A better Algorithm: "Dynamic Programming"

Sequence Similarity: What

GGACCA

TACTAAG

TCCAAT

Sequence Similarity: What

GGACCA

T A C T A A G | : | : | | : T C C – A A T

Sequence Similarity: Why

Most widely used comp. tools in biology New sequence always compared to sequence data bases

Similar sequences often have similar origin or function

Recognizable similarity after 10⁸ – 10⁹ yr

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

Try it! pick any protein, e.g. hemoglobin, insulin, exportin,...

Taxonomy Report

root	64 hits	16 orgs
. Eukaryota	62 hits	14 orgs [cellular organisms]
Fungi/Metazoa group	57 hits	11 orgs
Bilateria	38 hits	7 orgs [Metazoa; Eumetazoa]
Coelomata	36 hits	6 orgs
Tetrapoda	26 hits	5 orgs [;;; Vertebrata;;;; Sarcopterygii]
Eutheria	24 hits	4 orgs [Amniota; Mammalia; Theria]
Homo sapiens	20 hits	1 orgs [Primates;; Hominidae; Homo]
Murinae	3 hits	2 orgs [Rodentia; Sciurognathi; Muridae]
Rattus norvegicus	2 hits	1 orgs [Rattus]
Mus musculus	1 hits	1 orgs [Mus]
Sus scrofa	1 hits	1 orgs [Cetartiodactyla; Suina; Suidae; Sus]
Xenopus laevis	2 hits	<pre>1 orgs [Amphibia;;;;;; Xenopodinae; Xenopus]</pre>
Drosophila melanogaster	10 hits	1 orgs [Protostomia;;;; Drosophila;;;]
Caenorhabditis elegans	2 hits	<pre>1 orgs [; Nematoda;;;;;;; Caenorhabditis]</pre>
Ascomycota	19 hits	4 orgs [Fungi]
Schizosaccharomyces pombe	10 hits	<pre>1 orgs [;;;; Schizosaccharomyces]</pre>
Saccharomycetales	9 hits	3 orgs [Saccharomycotina; Saccharomycetes]
Saccharomyces	8 hits	2 orgs [Saccharomycetaceae]
Saccharomyces cerevisiae .	7 hits	1 orgs
Saccharomyces kluyveri	1 hits	1 orgs
Candida albicans	1 hits	<pre>1 orgs [mitosporic Saccharomycetales;]</pre>
Arabidopsis thaliana	2 hits	1 orgs [Viridiplantae;Brassicaceae;]
Apicomplexa	3 hits	2 orgs [Alveolata]
Plasmodium falciparum	2 hits	1 orgs [Haemosporida; Plasmodium]
Toxoplasma gondii	1 hits	1 orgs [Coccidia; Eimeriida; Sarcocystidae;]
. synthetic construct	1 hits	1 orgs [other; artificial sequence]
disease virus	1 hits	1 orgs [Viruses; dsDNA viruses, no RNA …]

Terminology (CS, not necessarily Bio)

String: ordered list of letters TATAAG
Prefix: consecutive letters from front empty, T, TA, TAT, ...
Suffix: ... from end empty, G, AG, AAG, ...
Substring: ... from ends or middle empty, TAT, AA, ...
Subsequence: ordered, nonconsecutive TT, AAA, TAG, ...

Sequence Alignment

acbcdb	acbcdb
/ \	
cadbd	- c a d b - d -

Defn: An alignment of strings S, T is a pair of strings S', T' (with spaces) s.t.
(1) |S'| = |T'|, and (|S| = "length of S")
(2) removing all spaces leaves S, T

Alignment Scoring Alignment Scoring Alignment a c - - b c d b a c b c d b c a d b d - c a d b

-1 2 -1 -1 2 -1 2 -1

Value = 3*2 + 5*(-1) = +1

The *score* of aligning (characters or spaces) x & y is $\sigma(x,y)$. *Value* of an alignment $\sum_{i=1}^{|S'|} \sigma(S'[i],T'[i])$ An *optimal alignment:* one of max value

Optimal Alignment: A Simple Algorithm

for all subseqs A of S, B of T s.t. |A| = |B| doalign A[i] with B[i], $1 \le i \le |A|$ align all other chars to spacescompute its valueretain the maxend

output the retained alignment

Analysis

Assume |S| = |T| = nCost of evaluating one alignment: $\ge n$

How many alignments are there: pick n chars of S,T together say k of them are in S match these k to the k *un*picked chars of T Total time: $\ge n {\binom{2n}{n}} > 2^{2n}$, for n > 3E.g., for n = 20, time is > 2⁴⁰ operations

$$= \begin{pmatrix} 2n \\ n \end{pmatrix}$$

Polynomial vs Exponential Growth



Asymptotic Analysis

How does run time grow as a function of problem size? n^2 or $100 n^2 + 100 n + 100 vs 2^{2n}$ **Defn:** f(n) = O(g(n)) iff there is a constant c s.t. $|f(n)| \le cg(n)$ for all sufficiently large n. $100 n^2 + 100 n + 100 = O(n^2)$ [e.g. c = 101] $n^2 = O(2^{2n})$ 2^{2n} is *not* $O(n^2)$



Utility of Asymptotics

- "All things being equal," smaller asymptotic growth rate is better
- All things are never equal
- Even so, big-O bounds often let you quickly pick most promising candidates among competing algorithms
- Poly time algs often practical; non-poly algs seldom are.
- (Yes, there are exceptions.)

Fibonacci Numbers

```
fib(n) {
    if (n <= 1) {
        return 1;
    } else {
        return fib(n-1) + fib(n-2);
    }
}</pre>
```

Simple recursion, but many repeated subproblems!! => Time = $\Omega(1.61^{n})$

Fibonacci, II

```
int fib[n];
fib[0] = 1;
fib[1] = 1;
for(i=2; i<=n; i++) {
    fin[i] = fib[i-1] + fib[i-2];
}
return fib[n];</pre>
```

Avoid repeated subproblems by tabulating their solutions => Time = O(n) (in this case)

Candidate for Dynamic Programming?

Common Subproblems?

Plausible: probably re-considering alignments of various small substrings unless we're careful.

Optimal Substructure?

Plausible: left and right "halves" of an optimal alignment probably should be optimally aligned (though they obviously interact a bit at the interface).

(Both made rigorous below.)

Optimal Substructure (In More Detail)

Optimal alignment *ends* in 1 of 3 ways: last chars of S & T aligned with each other last char of S aligned with space in T last char of T aligned with space in S (never align space with space; $\sigma(-, -) < 0$) In each case, the rest of S & T should be optimally aligned to each other Optimal Alignment in O(n²) via "Dynamic Programming"

Input: S, T, |S| = n, |T| = m Output: value of optimal alignment

Easier to solve a "harder" problem:

V(i,j) = value of optimal alignment ofS[1], ..., S[i] with T[1], ..., T[j]for all $0 \le i \le n, 0 \le j \le m$.

Base Cases

V(i,0): first i chars of S all match spaces

$$V(i,0) = \sum_{k=1}^{i} \sigma(S[k],-)$$

V(0,j): first j chars of T all match spaces $V(0,j) = \sum_{k=1}^{j} \sigma(-,T[k])$

General Case

Opt align of S[1], ..., S[i] vs T[1], ..., T[j]: $\begin{vmatrix} \sim \sim \sim \sim S[i] \\ \sim \sim \sim \sim T[j] \end{vmatrix}, \quad \begin{vmatrix} \sim \sim \sim \sim \sim S[i] \\ \sim \sim \sim \sim -1 \end{vmatrix}, \text{ or } \begin{vmatrix} \sim \sim \sim \sim -1 \\ \sim \sim \sim \sim T[j] \end{vmatrix}$ Opt align of $V(i,j) = \max \begin{cases} V(i-1,j-1) + \sigma(S[i],T[j]) \\ V(i-1,j) + \sigma(S[i], -) \\ V(i,j-1) + \sigma(-, T[j]) \end{cases},$ for all $1 \le i \le n$, $1 \le j \le m$.

Calculating One Entry

$$V(i,j) = \max \begin{cases} V(i-1,j-1) + \sigma(S[i],T[j]) \\ V(i-1,j) + \sigma(S[i], -) \\ V(i,j-1) + \sigma(-, T[j]) \end{cases}$$



Mismatch = -1 Match = 2

Example

	j	0	1	2	3	4	5	
i			С	а	d	b	d	←T
0		0	-1	-2	-3	-4	-5	
1	а	-1	-1	1				
2	С	-2	1					Time =
3	b	-3						O(mn)
4	С	-4						
5	d	-5						
6	b	-6						

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Mismatch = -1 Match = 2

Example

	j	0	1	2	3	4	5	
i			С	а	d	b	d	←T
0		0	-1	-2	-3	-4	-5	
1	а	-1	-1	1	0	-1	-2	_
2	С	-2	1	0	0	-1	-2	
3	b	-3	0	0	-1	2	1	
4	С	-4	-1	-1	-1	1	1	
5	d	-5	-2	-2	1	0	3	
6	b	-6	-3	-3	0	3	2	

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Finding Alignments: Arrows = **Trace Back**

(ties for) max in V(i,j) def.

<--T



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Complexity Notes

- Time = O(mn), (value and alignment) Space = O(mn)
- Easy to get value in Time = O(mn) and Space = O(min(m,n))
- Possible to get value *and alignment* in Time = O(mn) and Space =O(min(m,n)) but tricky.

Weekly Bio Interlude

DNA Replication

DNA Replication: Basics



Issues & Complications, I

1st ~10 nt's added are called the *primer* In simple model, DNA pol has 2 jobs: prime & extend

Priming is error-prone

So, specialized *primase* does the priming; pol specialized for fast, accurate extension



Still doesn't solve the accuracy problem (hint: primase makes an *RNA* primer)

Issue 2: Rep Forks & Helices

"Replication Fork": DNA double helix is progressively unwound by a DNA helicase, and both resulting single strands are duplicated

- DNA polymerase synthesizes new strand 5' -> 3'(reading its template strand 3' -> 5')
- That means on one (the "leading") strand, DNA pol is chasing/pushing the replication fork
- But on the other "lagging" strand, DNA pol is running away from it.



Issue 3: Fragments

Lagging strand gets a series of "Okazaki fragments" of DNA (~200nt in eukaryotes) following each primer

The RNA primers are later removed by a *nuclease* and *DNA* pol fills gaps (more accurate than primase)

Fragments joined by *ligase*



		T	I	T	T	T						T		T	T I	T L	T	T	T	T	T	T		Т
р	primer Okazaki													р	riı	m	e	r						

Issue 4: Coord Lead/Lag



Alberts et al., Mol. Biol. of the Cell, 3rd ed, p258









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Issue 5: Twirls & Tangles

Unwinding helix (~10 nucleotides per turn) would cause stress. *Topoisomerase I* cuts DNA backbone on *one* strand, allowing it to spin about the remaining bond, relieving stress

Topoisomerase II can cut & rejoin *both* strands, after allowing another double strand to pass through the gap, de-tangling it.



Issue 6: Proofreading

Error rate of pol itself is $\sim 10^{-4}$, but overall rate is 10⁻⁹, due to proofreading & repair, e.g. pol itself can back up & cut off a mismatched base if one happens to be inserted priming the new strand is hard to do accurately, hence RNA primers, later removed & replaced other enzymes scan helix for "bulges" caused by base mismatch, figure out which strand is original, cut away new (faulty) copy; DNA pol fills gap which strand is original? Bacteria: "methylate" some A's, eventually. Euks: strand nicking

Replication Summary

Speed: 50 (eukaryotes) to 500 (prokaryotes) bp/sec Accuracy: 1 error per 10⁹ bp Complex & highly optimized Highly similar across all living cells

More info:

Alberts et al., Mol. Biol. of the Cell

Sequence Alignment

Part II Local alignments & gaps

Variations

Local Alignment

- Preceding gives *global* alignment, i.e. full length of both strings;
- Might well miss strong similarity of part of strings amidst dissimilar flanks
- **Gap Penalties**

10 adjacent spaces cost 10 x one space?

Many others

Local Alignment: Motivations

"Interesting" (evolutionarily conserved, functionally related) segments may be a small part of the whole

"Active site" of a protein

Scattered genes or exons amidst "junk", e.g. retroviral insertions, large deletions

Don't have whole sequence

Global alignment might miss them if flanking junk outweighs similar regions

Local Alignment

Optimal *local alignment* of strings S & T: Find substrings A of S and B of T having max value global alignment

S = abcxdexA = c x d eT = xxxcdeB = c - d evalue = 5

The "Obvious" Local Alignment Algorithm

for all substrings A of S and B of T: Align A & B via dynamic programming Retain pair with max value end;

Output the retained pair

Time: O(n²) choices for A, O(m²) for B, O(nm) for DP, so O(n³m³) total.
[Best possible? Lots of redundant work...] Local Alignment in O(nm) via Dynamic Programming

Input: S, T, |S| = n, |T| = m Output: value of optimal local alignment

Better to solve a "harder" problem for all $0 \le i \le n$, $0 \le j \le m$:

V(i,j) = max value of opt (global) alignment of a suffix of S[1], ..., S[i] with a suffix of T[1], ..., T[j]

Report best i,j

Base Cases

Assume $\sigma(x,-) \le 0$, $\sigma(-,x) \le 0$ V(i,0): some suffix of first i chars of S; all match spaces in T; best suffix is empty V(i,0) = 0 V(0,j): similar

$$V(0,j) = 0$$

General Case Recurrences

Opt suffix align S[1], ..., S[i] vs T[1], ..., T[j]: $\begin{vmatrix} \sim \sim \sim S[i] \\ \sim \sim \sim T[j] \end{vmatrix}, \begin{vmatrix} \sim \sim \sim \sim S[i] \\ \sim \sim \sim - \end{vmatrix}, \begin{vmatrix} \sim \sim \sim \sim - \\ \sim \sim T[j] \end{vmatrix}, \text{or} \end{vmatrix}$ Opt align of suffix of of suffix of $S_{1}...S_{i-1} \&$ $T_{1}...T_{j-1} V(i,j) = \max \begin{cases} V(i-1,j-1) + \sigma(S[i],T[j]) \\ V(i-1,j) + \sigma(S[i], -) \\ V(i,j-1) + \sigma(-,T[j]) \\ 0 \end{cases}, \text{ opt suffix alignment has:} \\ 2, 1, 1, 0 \\ \text{ chars of S/T} \end{cases}$ for all $1 \le i \le n$, $1 \le j \le m$.

Scoring Local Alignments

	j	0	1	2	3	4	5	6	
i			X	Х	Х	С	d	е	←T
0		0	0	0	0	0	0	0	
1	а	0							
2	b	0							
3	С	0							
4	X	0							
5	d	0							
6	е	0							
7	X	0							
	↑ S								

Finding Local Alignments

Again, arrows follow max

	j	0	1	2	3	4	5	6	тах
i			X	X	X	С	d	е	←T
0		0	0	0	0	0	0	0	
1	а	0	0	0	0	0	0	0	
2	b	0	0	0	0	0	0	0	
3	С	0	0	0,	0	2	1	0	
4	X	0	2	2	2		1	0	
5	d	0	1	1	1	1	3	2	
6	е	0	0	0	0	0	2	5	
7	X	0	2	2	2	1	1	4	
	↑ S								Ę

Notes

Time and Space = O(mn) Space O(min(m,n)) possible with time O(mn), but finding alignment is trickier

Local alignment: "Smith-Waterman" Global alignment: "Needleman-Wunsch"

Alignment With Gap Penalties

Gap: maximal run of spaces in S' or T'

abddc-d	2 gaps in S'
addcbd	1 gaps in T'

Motivations, e.g.:

mutation might insert/delete several or even many residues at once

matching cDNA (no introns) to genomic DNA (exons and introns)

Some parts of proteins less critical

A Protein Structure: (Dihydrofolate Reductase)





Sequence Evolution

Nothing in Biology Makes Sense Except in the Light of Evolution Theodosius Dobzhansky, 1973

Changes happen at random

. . .

Deleterious/neutral/advantageous changes unlikely /possibly/likely spread widely in a population Changes are less likely to be tolerated in positions involved in many/close interactions, e.g. enzyme binding pocket protein/protein interaction surface

Gap Penalties

Score = f(gap length) Kinds, & best known alignment time



Global Alignment with Affine Gap Penalties

V(i,j) = value of opt alignment ofS[1], ..., S[i] with T[1], ..., T[j]G(i,j) = ..., s.t. last pair matches S[i] & T[j]F(i,j) = ..., s.t. last pair matches S[i] & -E(i,j) = ..., s.t. last pair matches - & T[j]

Time: O(mn) [calculate all, O(1) each]

Affine Gap Algorithm

Gap penalty = $g + s^*(gap length), g, s \ge 0$ V(i,0) = E(i,0) = V(0,i) = F(0,i) = -g-i*s

$$\begin{split} V(i,j) &= \max(G(i,j), F(i,j), E(i,j)) \\ G(i,j) &= V(i-1,j-1) + \sigma(S[i],T[j]) \\ F(i,j) &= \max(F(i-1,j)-s, V(i-1,j)-g-s) \\ E(i,j) &= \max(E(i,j-1)-s, V(i,j-1)-g-s) \\ Old gap \qquad new gap \end{split}$$

Summary

Functionally similar proteins/DNA often have recognizably similar sequences even after eons of divergent evolution

- Ability to find/compare/experiment with "same" sequence in other organisms is a huge win
- Surprisingly simple scoring works well in practice: score positions separately & add, possibly w/ fancier gap model like affine
- Simple "dynamic programming" algorithms can find *optimal* alignments under these assumptions in poly time (product of sequence lengths)
- This, and heuristic approximations to it like BLAST, are workhorse tools in molecular biology

Significance of Alignments

Is "42" a good score? Compared to what?

Usual approach: compared to a specific "null model", such as "random sequences"

Overall Alignment Significance, II Empirical (via randomization)

Generate N random sequences (say N = $10^3 - 10^6$)

Align x to each & score

If k of them have better score than alignment of x to y, then the (empirical) probability of a chance alignment as good as observed x:y alignment is (k+1)/N

e.g., if 0 of 100 are better, you can say "estimated p < .01" How to generate "random" sequences?

Alignment scores often sensitive to sequence composition So uniform 1/20 or 1/4 is a bad idea Even background p_i can be dangerous Better idea: *permute* y N times

Generating Random Permutations

for (i = n-1; i > 0; i--){
 j = random(0..i);
 swap X[i] <-> X[j];

