CSE 427 Computational Biology Winter 2008

> Sequence Alignment; DNA Replication

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/

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	1	orgs	[Amphibia;;;;;; Xenopodinae; Xenopus]
Drosophila melanogaster	10	hits	1	orgs	[Protostomia;;;; Drosophila;;;]
Caenorhabditis elegans	2	hits	1	orgs	<pre>[; Nematoda;;;;;; Caenorhabditis]</pre>
Ascomycota	19	hits	4	orgs	[Fungi]
Schizosaccharomyces pombe	10	hits	1	orgs	[;;;; Schizosaccharomyces]
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	1	orgs	[mitosporic Saccharomycetales;]
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]
lymphocystis disease virus	1	hits	1	orgs	[Viruses; dsDNA viruses, no RNA]

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

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

Mismatch = -1 Match = 2

Alignment Scoring

- a c b c d b c a d b d - c a d b - d - $-1 2 -1 -1 2 -1 2 -1 \leftarrow$ Value = 3*2 + 5*(-1) = +1
- The score of aligning (characters or spaces) x & y is σ(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

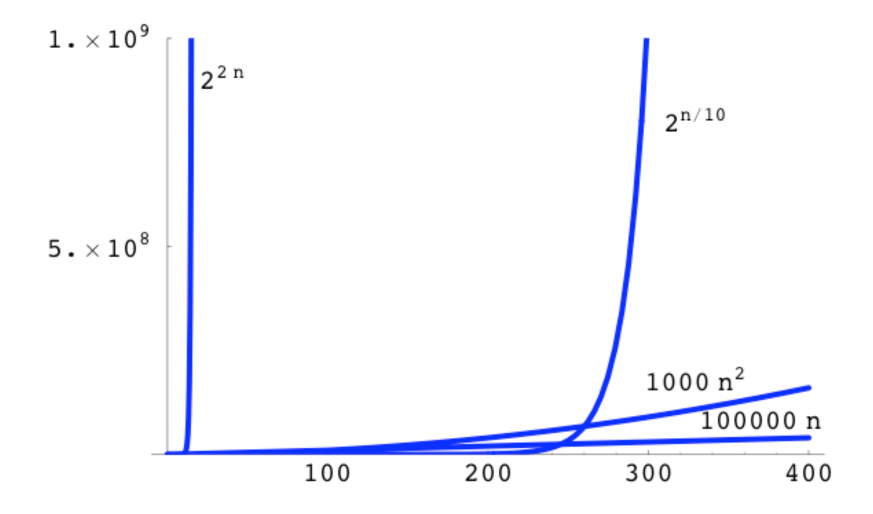
for all subseqs A of S, B of T s.t. |A| = |B| do
 align A[i] with B[i], 1 ≤ i ≤ |A|
 align all other chars to spaces
 compute its value
 retain the max
end
S = abcd A = cd
T = wxyz B = xz
-abc-d a-bc-d
w--xyz -w-xyz

output the retained alignment

Analysis

- Assume |S| = |T| = n
- Cost of evaluating one alignment: ≥ n
- How many alignments are there: $\geq \binom{2n}{n}$ • pick n chars of S,T together say k of them are in S match these k to the k unpicked chars of T
- Total time: $\ge n \binom{2n}{n} > 2^{2n}$, for n > 3• E.g., for n = 20, time is > 2⁴⁰ operations

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 = 300, or 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 algorithms often practical; non-poly algorithms 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++) {
    fib[i] = fib[i-1] + fib[i-2];
}
return fib[n];</pre>
```

```
"Dynamic
Programming"
Avoid repeated work by
tabulating solutions to
repeated subproblems
=>
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

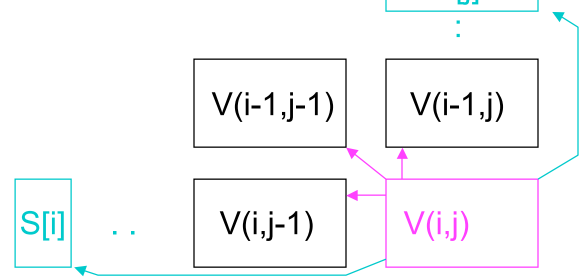
$$V(i,0) = \sum_{k=1}^{i} \sigma(S[k],-)$$

• 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]: $\begin{vmatrix} \sim \sim \sim & S[i] \\ \sim \sim \sim & T[j] \end{vmatrix}, \quad \begin{vmatrix} \sim \sim \sim & \sim & S[i] \\ \sim \sim \sim & - \end{vmatrix}, \text{ or } \begin{vmatrix} \sim \sim \sim & - & - \\ \sim \sim & T[j] \end{vmatrix}$ Opt align of $S_{1}...S_{i-1} \& T_{1}...T_{j-1}$ $V(i,j) = \max \begin{cases} V(i-1,j-1) + \sigma(S[i],T[j]) \\ V(i-1,j) + \sigma(S[i], -) \\ V(i,j-1) + \sigma(-, T[j]) \end{cases}$ Opt align of 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 2 3 5 j 0 1 4 i b d d ←T а С 0 -3 -5 -2 0 -1 -4 1 -1 1 -1 а 2 1 -2 С Time = 3 -3 b O(mn) 4 -4 С 5 d -5 6 -6 b

Ś

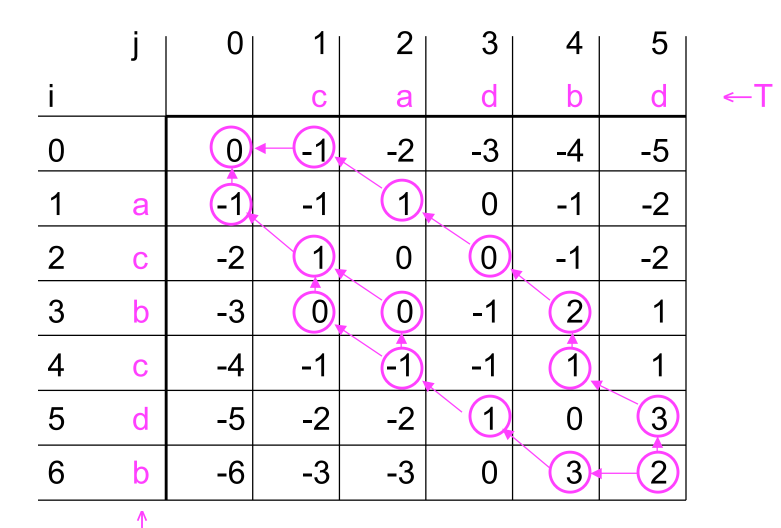
Mismatch = -1 Match = 2

	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	
				1	1	1	1	l i i i i i i i i i i i i i i i i i i i

Example

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Finding Alignments: Trace Back



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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.

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

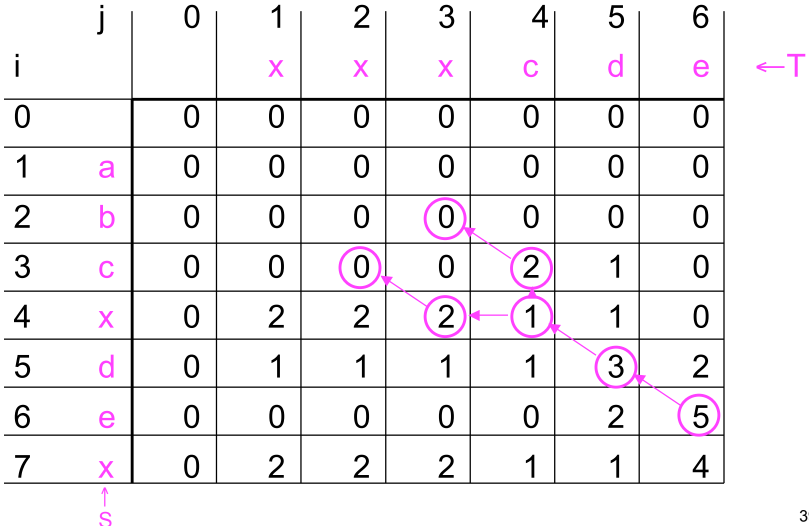
General Case Recurrences

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 $\sum_{\substack{S_{1},..,S_{i-1} \\ T_{1},..,T_{j-1}}}^{\text{SUIIX OT}} V(i,j) = \max \begin{cases} V(i-1,j-1) + \sigma(S[i],T[j]) \\ V(i-1,j) + \sigma(S[i], -) \\ V(i,j-1) + \sigma(-,T[j]) \\ 0 \end{cases}, \begin{array}{c} \text{opt suffix alignment has:} \\ 2, 1, 1, 0 \\ \text{chars of S/T} \end{cases}$ suffix of 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	←Ţ
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								3

Finding Local Alignments



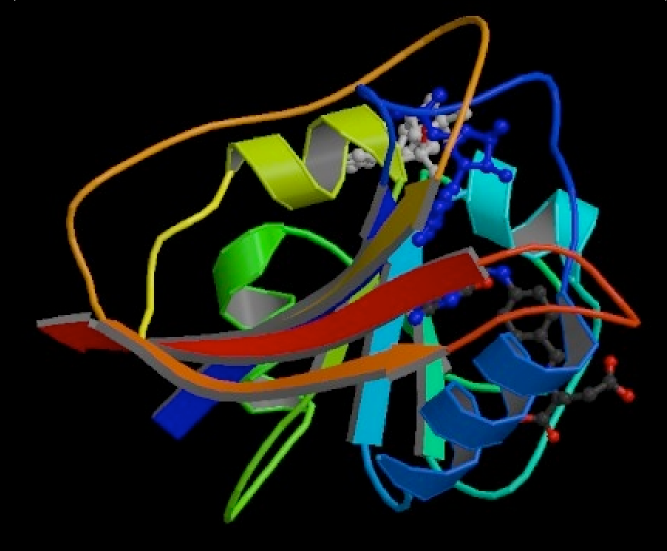
Notes

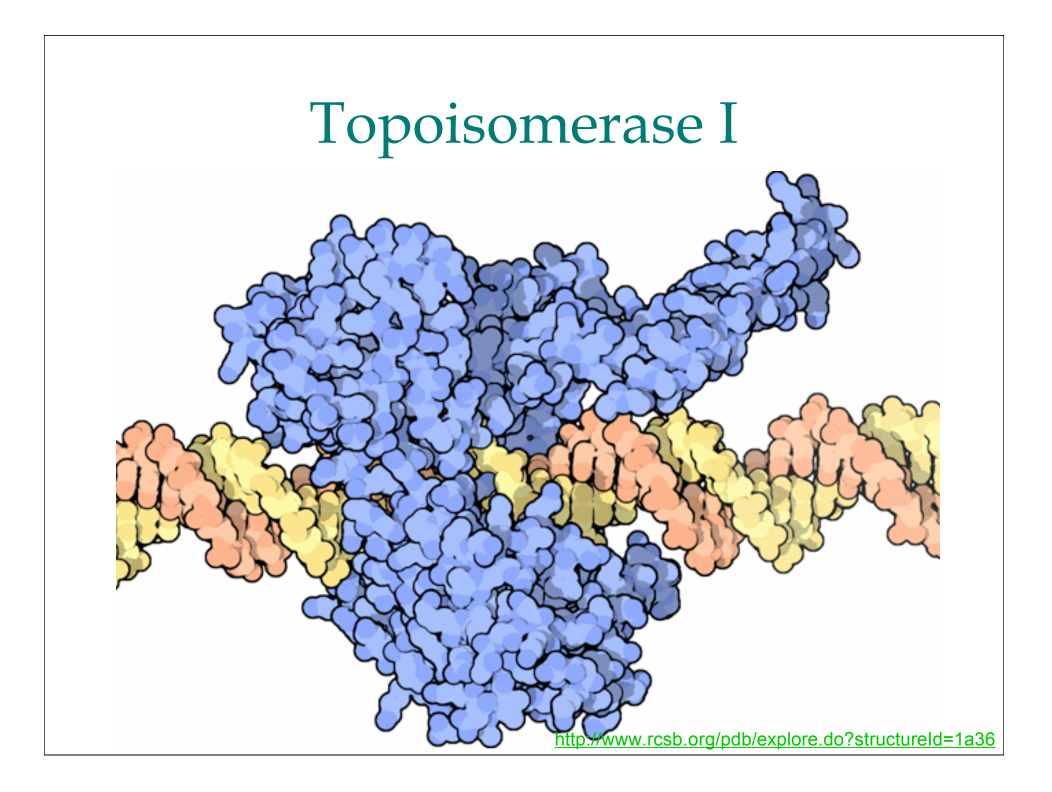
- Time and Space = O(mn)
- Space O(min(m,n)) possible with time O(mn), but finding alignment is trickier
- Local alignment: "Smith-Waterman"
- Global alignment: "Needleman-Wunsch"

Alignment With Gap Penalties

- Gap: maximal run of spaces in S' or T' ab----c-d a-ddddcbd
 2 gaps in S', 1 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

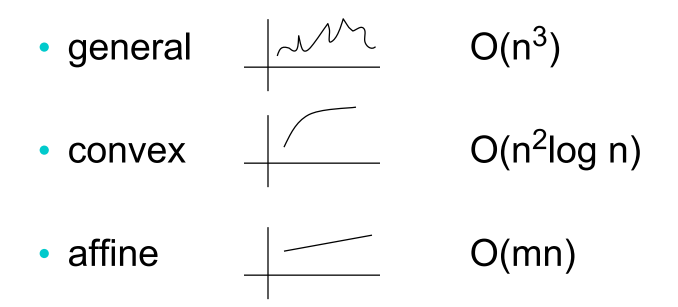
Changes happen at random

. . .

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 ≥ 0 V(i,0) = E(i,0) = V(0,i) = F(0,i) = -g-i*s 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

new gap

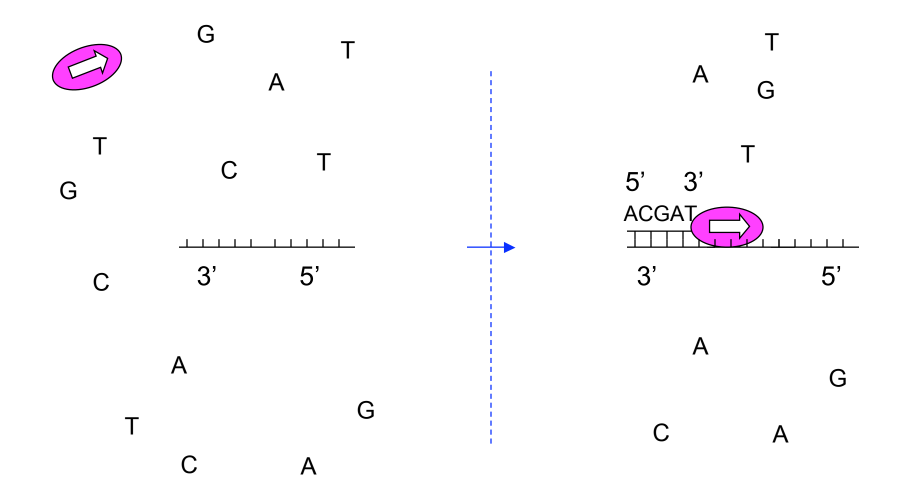
Summary

- 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 model works well in practice: score each position 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

Weekly Bio Interlude

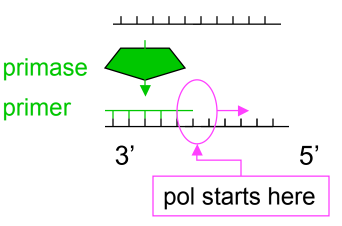
DNA Replication

DNA Replication: Basics



Issues & Complications, I

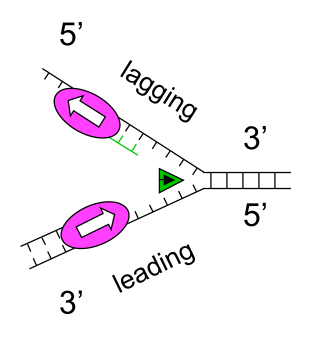
- 1st ~10 nt's added are called the primer
- In simple model, DNA pol has 2 jobs: prime & extend
- Priming is error-prone
- So, specialized *primase* does the priming; pol specialized for fast, accurate extension



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

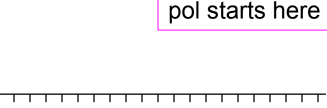
Issue 2: Rep Forks & Helices

- "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.



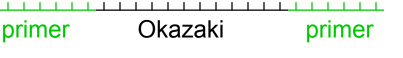
Issue 3: Fragments

- 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)



3'

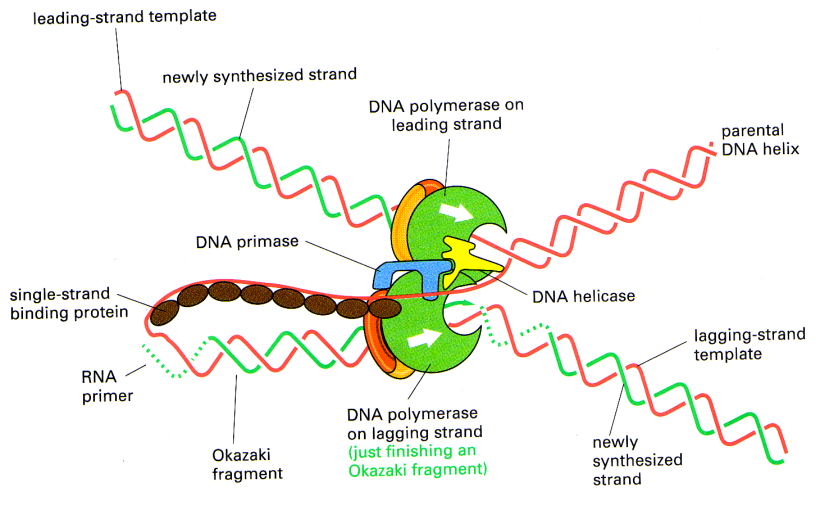
Fragments joined by *ligase*



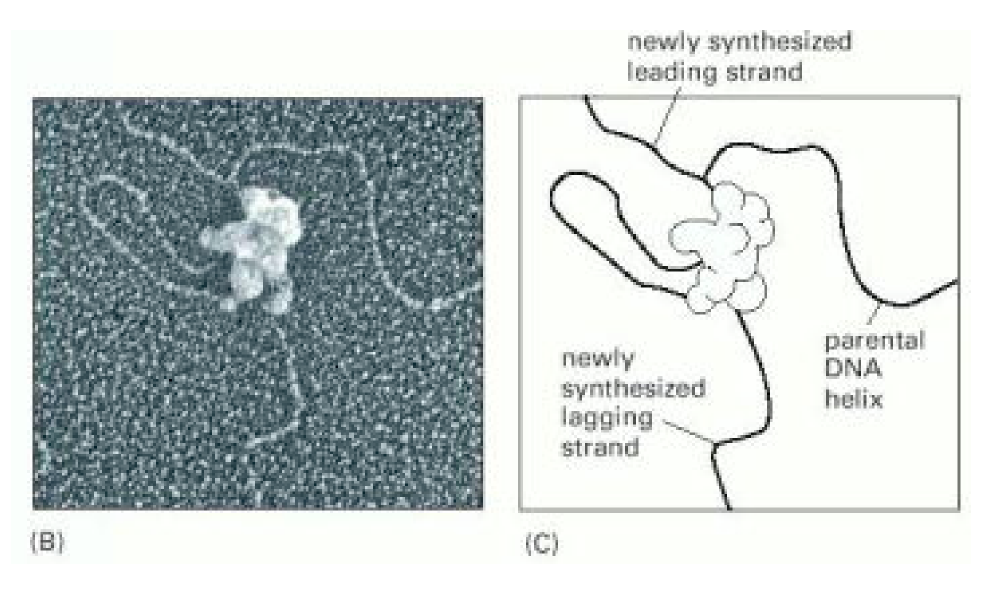
5

primer

Issue 4: Coord Lead/Lag

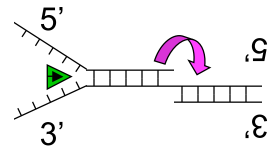


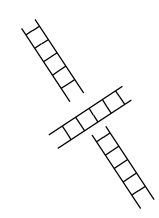
Alberts et al., Mol. Biol. of the Cell, 3rd ed, p258



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 ~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? In bacteria, some A's are "methylated", but not immediately after replication

Replication Summary

- Speed: 50 (eukaryotes) 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