

15-853: Algorithms in the Real World

Computational Biology II

- Introduction to Comp. Bio.
- Longest Common Subsequence and Minimum Edit Distance

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DNA

DNA: sequence of **base-pairs** (bp):
{A, C, T, G}

Human Genome

$\approx 3 \cdot 10^9$ bps divided into
46 chromosomes with between
 $5 \cdot 10^7$ and $25 \cdot 10^7$ bps each

Each chromosome is a **sequence** of base-pairs

DNA is used to generate proteins:

DNA $\xrightarrow{\text{transcription}}$ mRNA $\xrightarrow{\text{translation}}$ Protein

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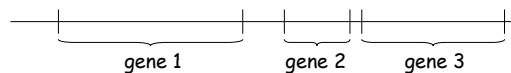
Proteins

Proteins: sequence of **Amino Acids**

{gly, trp, cys, ...} (20 of them)

Each DNA bp triple (a "**codon**") encodes 1 amino acid
Since there are 64 possible codons, this is a many
to one mapping. Some triples have special
meanings, e.g. EOF.

Chromosomes are partitioned into **genes** each of which
codes a protein. Some regions of the chromosome
do not code anything (**intergene DNA**).



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Form and Function

The Amino Acid sequence determines the protein's 3d **structure**. The structure is also affected by the environment.

- The **primary structure** refers to the amino acid sequence.
- The **secondary structure** refers to general configuration into **alpha helixes** and **beta sheets**
- The **tertiary structure** refers to the full 3d structure

Protein's 3d structure determines its **function**.

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Some Goals in Molecular Biology

1. Extract and compare genome sequence for various organisms.
 2. Determine what proteins they code.
 3. Determine structure and purpose of coded proteins.
- Goals 2. and 3. can often be aided by **matching** genome or protein sequences to previously studied sequences

Use to:

- study and cure genetic diseases
- design drugs
- study evolution
- understand molecular processes

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Example of MS

Multiple Sclerosis is a disease in which the immune system attacks the myelin sheaths of nerve cells

Conjecture: The immune system T-cells incorrectly identify the myelin sheaths as a virus or bacteria from an earlier infection.

This was tested by comparing the proteins sequences of myelin sheaths with a database of viral and bacterial proteins.

The ones that matched were tested in the laboratory to see if they were attacked by the T-cells. **Some were.**

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Approximate Matching and Sequence Alignment

How similar are:

actagtctac
cgacgtcgata ?

Allow for:

mutations: $x \rightarrow y$

insertions: $_ \rightarrow y$

deletions: $x \rightarrow _$

} "indels"

e.g.

acta_gtc__tac
| | | | |
cgacgtcgata

1 mutation
3 insertions
2 deletions

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Applications of matching and alignment

Used in many ways in computational biology

- Sequencing (finding the sequence of DNA or Proteins for a particular individual or species)
- Physical mapping (locating unique tags in DNA)
- Database searches (does this DNA match any other DNA?)
- Evolutionary trees (how similar are two species?)

Before talking about the general matching problem in computational Biology, we will talk about a closely related, but simpler problem: **longest common subsequence (LCS)**

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Longest Common Subsequence

Subsequence (C): Any subset of the elements of a sequence that maintain relative order

e.g. $A = a_1 a_2 a_3 a_4 a_5 a_6$

$A' = a_2 a_4 a_5$ (a subsequence of A)

$A' = a_2 a_1 a_5$ (not a subsequence of A)

Longest Common Subsequence (LCS):

$LCS(A,B) = C$, where $C \subset A$, $C \subset B$, $|C|$ is maximized

e.g. $A = a \ b \ a \ c \ d \ a \ c$
 $B = c \ a \ d \ c \ d \ d \ c$
 $C = a \ c \ d \ c \quad |C| = 4$

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Minimum Edit Distance

Minimum Edit Distance: $D(A,B) =$ minimum # of insertions and deletions needed to change A to B .

e.g. $A = a \ \boxed{b} \ \boxed{a} \ c \ d \ \boxed{a} \ c$ \square delete
 $B = \boxed{c} \ a \ \boxed{d} \ c \ d \ \boxed{d} \ c$ \square insert

Claim: $D(A,B) = |A| + |B| - 2|LCS(A,B)|$

Proof outline:

$C = LCS(A,B)$ (some common subsequence)

$A - C =$ deletions, $B - C =$ insertions

#deletions = $|A| - |C|$ #insertions = $|B| - |C|$

This reduction works both ways, hence equality.

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Applications

Unix diff:

Find Minimum Edit Distance and print edits

GNU diff based on algorithm by Eugene Myers, who was VP of Informatics at Celera

Screen redisplay:

Find minimum number of changes that need to be sent to the display along a "skinny" wire.

Used, for example, by Emacs.

Computational biology:

This will use generalizations of LCS, but the algorithms will work with slight modifications.

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Outline

Will work our way up to the GNU diff algorithm

- Recursive solution
- Memoized solution
- Dynamic programming
- Memory efficient solution
- Myers/Ukkonon algorithm

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Recursive Solution

```
D(A, empty) = |A|   delete(A)
D(empty, B)  = |B|   insert(B)
D(A:x, B:x)  = D(A, B)
D(A:a, B:b)  = min(1 + D(A:a, B), 1 + D(A, B:b))
                insert(b)   delete(a)
```

Note that this just returns the edit distance, but it is easy to include the edits.

Can work from start or end (here from end).

Why does it work?

What is the running time?

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Recursive Solution

Convert to use indices

```
int ED(int i, int j) {
    if (i==0) r = j;
    if (j==0) r = i;
    if (A[i] == B[j])
        r = ED(i-1, j-1);
    else
        r = min(1 + ED(i-1, j), 1 + ED(i, j-1));
    return r;
}
ED(n, m);
```

What is the running time?

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Memoized Solution

```
int ED(int i, int j) {
    if (M[i, j] != -1) return M[i, j];
    if (i==0) r = j;
    if (j==0) r = i;
    if (A[i] == B[j])
        r = ED(i-1, j-1);
    else
        r = min(1 + ED(i-1, j), 1 + ED(i, j-1));
    return M[i, j] = r;
}
M[1..n, 1..m] = -1;
ED(n, m);
```

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Dynamic programming

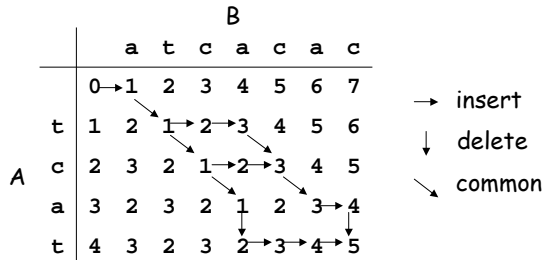
```
for i = 1 to n
    M[i, 1] = i;
for j = 1 to m
    M[1, j] = j;

for i = 1 to n
    for j = 1 to m
        if (A[i] == B[j])
            M[i, j] = M[i-1, j-1];
        else
            M[i, j] = 1 + min(M[i-1, j], M[i, j-1]);
```

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Example



Note: can be filled in any order as long as the cells to the left and above are filled.
 Can follow path back through matrix to construct edits.

Space Efficient Solution

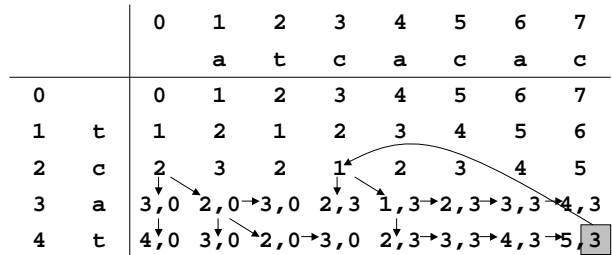
```

for i = 1 to n
  for j = 1 to m
    if (A[i] == B[j])
      R[j] = Rprev[j-1];
    else
      R[j] = 1 + min(Rprev[j],R[j-1]);
  Rprev[0..m] = R[0..m]
  
```

Requires only $O(m)$ space.
 What is the problem?

Space Efficient Solution

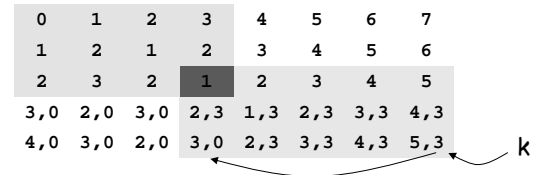
For each entry in a row past $n/2$ keep track of which column it comes from in row $n/2$.



The function $ED'(A,B)$ returns column: 3

Space efficient solution

Now solve recursively:



```

function RecED(A,B) =
  k = ED'(A,B)
  RecED(A[1..n/2],B[1..k]) ++ RecED(A[n/2..n],B[k..m])
  
```


Dijkstra's algorithm (review)

Single source (v_s) single destination (v_d) shortest path

$d(v) = \infty$ for all $v \neq v_s$

$d(v_s) = 0$

insert(v_s , EmptyQ)

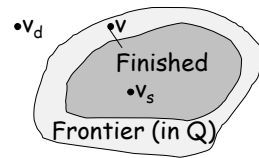
while Min(Q) $\neq v_d$

$v = \text{deleteMin}(Q)$

for each neighbor v' of v

$d(v') = \min(d(v'), d(v) + \text{weight}(v,v'))$

insert(v', Q), or replace if already in there



Takes $|E|$ inner iterations, each taking $O(\log |V|)$ time.
Can be improved to $O(|E| + |V|\log|V|)$ time.

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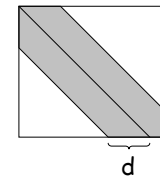
Bounding Visited Vertices

Theorem: $D_{ij} \geq |j-i|$

Proof: every step away from the diagonal is along a horizontal or vertical edge. Each such edge contributes 1, so the total D , must be at least the distance from the diagonal

Corollary: Dijkstra's algorithm visits at most $\min(n,m) \cdot 2d$ vertices.

■ searched vertices



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Bounding time

Priority Queue can be kept in constant time per operation (basically a modified breadth-first search), so **total time** is $O(\min(n,m) \cdot d)$.

In practice this can be much less than $O(nm)$.

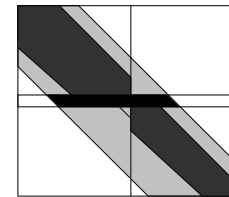
What about space?

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Optimizing Space

Use recursive trick from before



Problem: We do not know d ahead of time. Scanning row by row will not work without d .

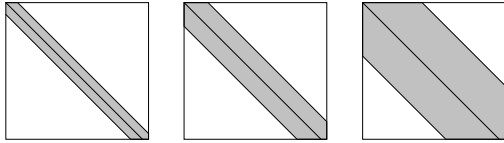
Need to bound size of frontier.

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Increasing Band-Widths

Start with band of $|m-n|$ and double on each step
until large enough



Visits at most twice as many vertices as it should.