

# CSE 584A Class 20

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## 1 Eliminating Redundant Work in Tree Alignment

When can we stop downward DP in a tree?

- Neither local nor semi-local alignments are *a priori* anchored at the first row of the DP matrix (i.e. top of tree).
- Hence, it seems that we need to continue DP all the way to the bottom of the tree on every branch to find all alignments.
- (This would cost  $\Theta(|P||T|^2)$  – blech!)
- But wait – a full computation repeats a lot of work!
- Example:

- An alignment starting at  $T[i]$  will be found  $i$  times, one for each suffix of  $T$  starting at or before position  $i$ .

The following argument is key to removing redundancy.

- Suppose we are aligning  $P$  to the path in  $\tau$  labeled with text suffix  $T[h..n]$ .
- Now suppose that an optimal alignment skips over some nonempty prefix  $T[h..j]$  of this suffix before using any characters in  $P$ .
- The same alignment, at the same location in  $T$ , will also be discovered when we perform DP on the path in  $\tau$  labeled with  $T[j + 1..n]$ .
- Hence, WLOG, *we can always anchor* our alignments to the top of the tree, since paths for all suffixes of  $T$  begin there.

- Above argument applies to either local or semi-local alignment w/r to  $P$ .

How do we apply this observation in practice?

- *First step*: prevent skipping of any nonempty prefix of  $T$  before starting to align  $P$ .
- Initialize first column of DP matrix with  $-\infty$  after first (initialization) row.
- This prevents us from finding the same alignment on several branches, but it doesn't seem to help us stop DP before the bottom of the tree.
- To win big, we need a stronger rule, as follows.
- Suppose we have found an DP alignment path  $\Pi$  that uses at least one character of  $T$ .
- Suppose further that  $\Pi$  can be replaced by another path  $\Pi'$ , ending at the same cell as  $\Pi$ , that does *not* use any of  $T$  and has at least the same score as  $\Pi$ .
- Surely, any alignment path  $\Pi \cdot \Phi$  can be replaced by  $\Pi' \cdot \Phi$  with at least the same score, so  $\Pi'$  dominates  $\Pi$ .
- (something trivial to check for affine gap case)
- Moreover,  $\Pi'$  will be found by alignment on another branch of the tree that starts later in  $T$ .
- Conclude that we may *cut off extension of*  $\Pi$  because some other branch will find a better or equal alignment path through the same part of  $P$  and a later suffix of  $T$ .

We now have an X-drop-like rule for cutting off DP.

- Let's first consider fully local case.
- Suppose that, after the initialization row, we set a cell  $(i, j)$  in the DP matrix to 0.
- By our Smith-Waterman recurrence, any nonempty path ending at  $(i, j)$  (which must use some of  $T$ ) is no better than the empty path (which does not use any of  $T$ ).
- Hence, by our reasoning above, we can cut off extension from  $(i, j)$ .
- As with X-dropping, we cut off extension by replacing the 0 in cell  $(i, j)$  with  $-\infty$ .
- The same hacks as for cutting off BLAST extension now tell us when we can safely stop aligning!
- A detailed version of the affine Smith-Waterman recurrence with this form of cutting off is used in the BWT-SW program, described in "Compressed indexing and local alignment of DNA," by Lam et al. (*Bioinformatics* 2008).

But what about the semi-local case?

- In semi-local alignment, it is not free to skip a prefix of  $P$ .

- In particular, an alignment that starts in column  $j$  of the DP matrix incurs a gap cost  $c(j) = -g_o - j \cdot g_e$  (for the affine case).
- Now suppose that the score of a cell  $(i, j)$ , after the initialization row, is  $\sigma \leq c(j)$ .
- We can replace any alignment path ending at this cell by a path of equal or better score that skips the first  $j$  characters of  $P$ , at cost  $c(j)$ , and uses none of  $T$ .
- By our argument above, some other path in the tree will find this better alignment.
- Hence, *in column  $j$ , we can replace any score  $\leq c(j)$  after the initialization row by  $-\infty$ .*
- Again, usual cutoff tricks apply.

## 2 Further Pruning of Tree Alignment

So far, we have one provably correct criterion for cutting off alignments. Can we do even better?

- Suppose we seek alignments to a pattern  $P[1..m]$  above some score threshold  $\theta$ .
- Now suppose that, after using up characters  $P[1..j]$  (or some suffix thereof for local alignment), we have a partial alignment path ending at cell  $(i, j)$  with score  $\sigma < \theta$ .
- Is it possible to reach score  $\theta$  by aligning the rest of  $P$  to the subsequent chars of  $T$ ?
- If not, we might as well cut off this path by setting  $(i, j)$  to  $-\infty$ .
- But how can we prove that we cannot reach  $\theta$  without actually doing the rest of the alignment?
- *Idea:* if we can prove that no alignment of  $P[j + 1..m]$  to the subsequent characters of  $T$  can exceed some score  $\psi$ , and  $\sigma + \psi < \theta$ , then we can never reach score  $\theta$ .
- So, any heuristic that can overestimate (but *not* underestimate) the true remaining alignment score can be used for cutting off.
- (like the “admissible heuristic” criterion in state-space search)

Can we come up with some heuristics that don’t require as much work as just completing the alignment?

- Here’s a trivial one: compute the score  $\rho(j)$  of aligning each character of  $P[j..m]$  against itself.
- If the scoring system always scores matches higher than mismatches or gaps, then no alignment of  $P[j..m]$  to  $T$  can score better than  $\rho(j)$ .
- This seems like a pretty weak heuristic.
- BWA uses a better approximation for the *special case* that the scoring function
  - assigns all matches the same bonus  $a$ ,

- assigns all edits (mismatches or gaps) the same linear penalty  $b$
- *Idea*: compute the *matching statistics* of  $P$  vs  $T$ , and use them to upper-bound cost of matching  $P$  against an arbitrary substring of  $T$ .
- **Defn**: the *matching statistic*  $\mu(j)$  is the length of the longest prefix match between  $P[j..m]$  and  $T$ .
- We can compute  $\mu(j)$  in time proportional to its value using a top-down tree walk to match  $P[j..m]$  vs  $T$ .
- Now consider trying to match  $P[j..j + \mu(j)]$  against  $T$ .
- By definition, this substring, which is one longer than  $\mu(j)$ , does not occur anywhere in  $T$ , so we must incur at least one edit in *any* alignment of it to  $T$ .
- Can recursively compute this cost for the rest of  $P$ .
- Here's BWA's edit-distance bounding algorithm.

```

EDITBOUND( $P, T$ )
   $j \leftarrow 1$ 
   $d \leftarrow 0$ 
  while  $j \leq m$  do
    if  $j + \mu(j) \leq m$ 
       $d \leftarrow d + 1$ 
       $j \leftarrow j + \mu(j) + 1$ 
  return  $d$ 

```

- To convert  $d$  to a cost bound, just compute  $(m - d) \cdot a - d \cdot b$ .
- We can precompute the cost bound for every suffix of  $P$ . For each suffix, we may do  $O(m)$  work to compute matching stats.
- Hence, total cost is  $O(m^2)$ .
- However, if we do the equivalent computation starting from the *end* of  $P$ , we still get valid bounds (not necessarily the same ones), but we can reuse the matching stats computed for  $P[k..m]$ ,  $k > j$ , when computing  $d$  for  $P[j - 1..m]$ .
- Total cost for all suffixes of  $P$  is then only  $O(m)$ .
- BWA actually does the computation the other way around, i.e. bounding the number of edits to align each *prefix* of  $P$ , because it computes alignments of  $P$  to  $T$  starting from the end of  $P$ , using the BWT to simulate traversal backwards in  $T$ .

Coming up with better-than-trivial heuristics for cutting off DP with arbitrary match, substitution, and gap costs (as for protein alignment) is (AFAIK) an open problem.