

CS425

ASSIGNMENT 1 (DUE SEP 24, 2019)

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September 17, 2019

1. Global alignment with a limited number of gaps [30 pts].

Suppose $k \in \mathbb{Z}$ is a non-negative third input (in addition to input sequences v and w) to the alignment algorithm. Devise a dynamic programming algorithm that finds an optimal global alignment between v and w subject to the constraint that the alignment contains at most k blocks of consecutive indels. You may denote the scoring matrix by δ which is pursuant to our notation in class.

2. Semi-global alignment [35 pts].

In this question, we consider a variant of the global alignment problem, where we impose no penalty on gaps at the end of one of the input sequences. Consider the following two alternative alignments:

```
v: CAGCA-CTTGGATTCTCGG
w: ---CAGCGTGG-----
```

```
v: CAGCACTTGGATTCTCGG
w: CAGC-----G-T----GG
```

When doing global alignment under the scoring scheme of +1 for a match, -1 for a mismatch, and -1 for a gap, the second alignment is preferred despite our intuition that the first alignment is more biologically relevant. If the gaps on the ends of w are not penalized, then the first alignment scores higher. This approach is called *semi-global alignment*. Note that in this approach, the end gaps in one of the sequences (v) are penalized as in the standard global alignment. Show how to modify the Needleman-Wunsch algorithm to compute a semi-global alignment (including the initialization of the matrix, and the backtrace operations). Modify `align.py` to perform semi-global alignment and illustrate that your code is indeed performing semi-global alignment by running it on some examples and providing the output.

3. Local alignment [35 pts].

Modify the program `align.py` to perform local sequence alignment of proteins using a substitution matrix (here's a link to a BLOSUM 62 matrix: <http://www.ncbi.nlm.nih.gov/Class/FieldGuide/BLOSUM62.txt>). Use your program to align the *Aniridia* protein from *H. sapiens* to the eyeless protein from *D. melanogaster* using the BLOSUM 62 substitution matrix. The

sequences of these proteins can be obtained from the Uniprot protein database using the following links: <http://www.uniprot.org/uniprot/P26367> (aniridia) and <http://www.uniprot.org/uniprot/O18381> (eyeless). Note that Uniprot has an option for saving an entry in FASTA format. Answer the following questions:

- What is the alignment and score you obtained?
- Align 100 randomly generated protein sequences with the same length and amino acid composition as the *Aniridia* protein with the eyeless protein. Compare the scores with that obtained above. Does that suggest a statistically significant similarity event?
- * Optional: Quantify that statistical significance by providing a p -value. Any idea about the underlying null probability distribution?

Upload your answer on Canvas in one zip file or tarball. Include all the code/scripts you have written in your submission.