CSC 314**, Bioinformatics Lab #11: Name:\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_**

**Dynamic Programming / Pairwise Alignments**

***Note: For all alignments, we will use a linear gap penalty of 4 points, a match score of +5 points, and a mismatch score of -1 point.***

1. Consider the dynamic programming matrix for the optimal global alignment of *ttcag* with *taca*, which has been partially completed for you.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  |  | t | a | c | a |
|  | 0 | -4 | -8 | -12 | -16 |
| t | -4 | 5 | 1 |  |  |
| t | -8 | 1 | 4 |  |  |
| c | -12 | **-3** | **0** |  |  |
| a | -16 | **-7** |  |  |  |
| g | -20 |  |  |  |  |

* 1. The highlighted squares with bold scores correspond to the scores that should be used when calculating the optimal global alignment score between *ttca* and *ta*. The optimal global alignment score is the maximum of what 3 values?
  2. What is the score of this optimal alignment (between *ttca* and *ta*)?
  3. What is the optimal global alignment (between *ttca* and *ta*). Note: if multiple optimal alignments exist, you can arbitrarily select one.
  4. Fill in the rest of the matrix to find the score of the optimal global alignment between the two sequences *ttcag* and *taca*.
     1. What is the optimal global alignment score
     2. What is the optimal global alignment?

1. Find the optimal global alignment and optimal global alignment score between the sequences *east* and *hat*, using the scoring system at the top of this assignment, by constructing the appropriate dynamic programming matrix. You must show your dynamic programming matrix to receive credit.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  |  | h | a | t |
|  | 0 | -4 | -8 | -12 |
| e | -4 |  |  |  |
| a | -8 |  |  |  |
| s | -12 |  |  |  |
| t | -16 |  |  |  |

1. What is the optimal global alignment score?
2. What is the optimal global alignment?
3. Complete the dynamic programming matrix below to find the score of the optimal *semiglobal* alignment, and the optimal semiglobal alignment between *acac* and *tacc*. The semiglobal alignment is found in the same way as the global alignment except that (1) the 1st row and 1st column of the matrix are initialized to all 0s (completed for you) and (2) traceback begins from the highest scoring cell in the bottom row or last column, with opening and closing gaps added as necessary.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  |  | t | a | c | c |
|  | 0 | 0 | 0 | 0 | 0 |
| a | 0 |  |  |  |  |
| c | 0 |  |  |  |  |
| a | 0 |  |  |  |  |
| c |  |  |  |  |  |

1. What is the optimal global alignment score?
2. What is the optimal global alignment?
3. Find the optimal semiglobal alignment between *tata* and *caag* by completing the dynamic programming matrix below.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  |  | c | a | a | g |
|  | 0 | 0 | 0 | 0 | 0 |
| t | 0 |  |  |  |  |
| a | 0 |  |  |  |  |
| t | 0 |  |  |  |  |
| a | 0 |  |  |  |  |

1. What is the optimal global alignment score?
2. What is the optimal global alignment?
3. Find the optimal *local* alignment between *steps* and *toe*, by constructing the appropriate dynamic programming matrix. Refer to your notes for the steps for finding the optimal local alignment. Note that the optimal local alignment is the following which has a score of 9.

t e p

t o e

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  |  | t | o | e |
|  | 0 | 0 | 0 | 0 |
| s | 0 |  |  |  |
| t | 0 |  |  |  |
| e | 0 |  |  |  |
| p | 0 |  |  |  |
| s | 0 |  |  |  |