BIMM-143: FOUNDATIONS OF BIOINFORMATICS

Lecture 2, "Sequence Alignment Fundamentals" Homework

https://bioboot.github.io/bggn213_S18/lectures/#2

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This unit's homework consists of both (1) an online <u>knowledge assessment quiz</u> and (2) a Needleman-Wunsch dynamic programming assessment exercise. Both components contribute 50% to this unit's grade. For the later we have two sample sequences, and we'd like to use the Needleman-Wunsch algorithm discussed in class to align them.

		Т	А	Т	А	G	С
	0						
G							
Т							
Т							
Α							
Т							
С							

Sequence 1: **TATAGC** Sequence 2: **GTTATC**

Using a match score of 2, a mismatch score of -1, and a gap score of -2. Fill in the table and translate it into a alignment.

Please return your answer to Yuansheng via email: (yuz461@ucsd.edu). This should consist of an attached photo of your completed **alignment matrix** along with your **aligned sequences** and their **optimal score**.

Step	Scoring Rubric/Assessment Criteria	Points	
1	Setup labeled alignment matrix	10	
2	Include initial column and row for GAPs	10	
3	All alignment matrix elements filled in	10	
4	Evidence for correct use of scoring scheme	10	
5	Direction arrows drawn between all cells	10	
6	Evidence of multiple arrows to a given cell if appropriate	10	D
7	Correct optimal score position in matrix used	10	С
8	Correct optimal score obtained for given scoring scheme	10	В
9	Traceback path(s) clearly highlighted	10	А
10	Correct alignment(s) yielding optimal score listed	10	A+

(100 Total points)