## **Lecture 3 Homework**

http://thegrantlab.org/

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To complete this homework you must be working on your own copy in **Google Docs**. Once you have filled in your answers to **Q1-Q4** in the space provided click **File > Download > PDF document** and upload to gradescope (link can be found on the class website).

**Q1**. [4pts] Consider the following multiple alignment of Transcription Factor Binding site DNA sequences

	1	2	3	4	5
Sequence 1	-	G	Α	G	С
Sequence 2	С	Т	А	G	А
Sequence 3	С	G	А	-	Α
Sequence 4	Α	G	С	G	А

Give the average profile (frequency matrix) of the above alignment by filling out the table below. The first position of the first column (i.e. position in the alignment) has been done for your, now complete the rest. You will use this table for answering questions 2 and 3 below.

	1	2	3	4	5
A	0.25	0	0.75	0	0.75
С	0.5	0	0.25	0	0.25
Т	0	0.25	0	0	0
G	0	0.75	0	0.75	0
-	0.25	0	0	0.25	0

what is it's so	;ore?
Sequence:	CGAGA
Score:	3.5
Q3. [2pts] Usequences (\$	Using your completed profile table above (from question 1) score the following two
\$1. CTGGC	
<b>31.</b> 01000	1.75
S2. AGCTA	2
<b>Q4</b> . [2pts] F	following the heuristic threshold for a positive match proposed in Harbison et al.
-	4) 431:99-104.] namely using the threshold for a positive match = 60% x Max Sco the two sequences in question 3 potential transcription factor binding sites? If so,
	shold is 60%x3.5=2.1, which is bigger than the scores in question 3. Therefore, nces are not potential transcription factor binding sites.