#### Final Exam InstructionsHelp Center

Please thoroughly read the instructions below. When you are ready to take the Final Exam, please visit the <u>Final Exam Page</u> and click the blue Attempt Exam button.

Write a Python program that takes as input a file containing DNA sequences in multi-FASTA format, and computes the answers to the following questions. You can choose to write one program with multiple functions to answer these questions, or you can write several programs to address them. We will provide a multi-FASTA file for you, and you will run your program to answer the exam questions.

While developing your program(s), please use the following example file to test your work: <a href="mailto:dna.example.fasta">dna.example.fasta</a>

You'll be given a different input file to launch the exam itself.

Here are the questions your program needs to answer. The quiz itself contains the specific multiplechoice questions you need to answer for the file you will be provided.

- (1) How many records are in the file? A record in a FASTA file is defined as a single-line header, followed by lines of sequence data. The header line is distinguished from the sequence data by a greater-than (">") symbol in the first column. The word following the ">" symbol is the identifier of the sequence, and the rest of the line is an optional description of the entry. There should be no space between the ">" and the first letter of the identifier.
- (2) What are the lengths of the sequences in the file? What is the longest sequence and what is the shortest sequence? Is there more than one longest or shortest sequence? What are their identifiers?
- (3) In molecular biology, a reading frame is a way of dividing the DNA sequence of nucleotides into a set of consecutive, non-overlapping triplets (or codons). Depending on where we start, there are six possible reading frames: three in the forward (5' to 3') direction and three in the reverse (3' to 5'). For instance, the three possible forward reading frames for the sequence AGGTGACACCGCAAGCCTTATATTAGC are:

AGG TGA CAC CGC AAG CCT TAT ATT AGC

A GGT GAC ACC GCA AGC CTT ATA TTA GC

#### AG GTG ACA CCG CAA GCC TTA TAT TAG C

These are called reading frames 1, 2, and 3 respectively. An open reading frame (ORF) is the part of a reading frame that has the potential to encode a protein. It starts with a start codon (ATG), and ends with a stop codon (TAA, TAG or TGA). For instance, ATGAAATAG is an ORF of length 9.

Given an input reading frame on the forward strand (1, 2, or 3) your program should be able to identify all ORFs present in each sequence of the FASTA file, and answer the following questions: what is the length of the longest ORF in the file? What is the identifier of the sequence containing the longest ORF? For a given sequence identifier, what is the longest ORF contained in the sequence represented by that identifier? What is the starting position of the longest ORF in the sequence that contains it? The position should indicate the character number in the sequence. For instance, the following ORF in reading frame 1:

>sequence1

ATGCCCTAG

starts at position 1.

Note that because the following sequence:

>sequence2

ATGAAAAA

does not have any stop codon in reading frame 1, we do not consider it to be an ORF in reading frame 1.

(4) A repeat is a substring of a DNA sequence that occurs in multiple copies (more than one) somewhere in the sequence. Although repeats can occur on both the forward and reverse strands of the DNA sequence, we will only consider repeats on the forward strand here. Also we will allow repeats to overlap themselves. For example, the sequence ACACA contains two copies of the sequence ACA - once at position 1 (index 0 in Python), and once at position 3. Given a length n, your program should be able to identify all repeats of length n in all sequences in the FASTA file. Your program should also determine how many times each repeat occurs in the file, and which is the most frequent repeat of a given length.

Welcome to the final exam.

If you haven't yet read the instructions, you can do so here.

Please run the following data set in the program(s) that you have written: dna1.fasta

If you created your program(s) correctly, you will be able to answer the questions below.

#### **Question 1**

How many records are in the multi-FASTA file?

Your Answer		Score
° 760		
° 740		
° <sub>14</sub>		
© 20	Correct	1.00
Total		1.00 / 1.00
Question 2		

What is the length of the longest sequence in the file?

Your Answer	Score

° 4200

6 4510       Correct       1.00         ○ 6007				
C 10523  Total 1.00 / 1.00  Question 3  What is the length of the shortest sequence in the file?  Your Answer Score  1081  C 475  Correct 1.00	<ul><li>4510</li></ul>	Correct	1.00	
Total 1.00 / 1.00  Question 3  What is the length of the shortest sequence in the file?  Your Answer Score  1081  • 475 Correct 1.00	° 6007			
Question 3 What is the length of the shortest sequence in the file?  Your Answer Score  1081  Correct 1.00	C 10523			
What is the length of the shortest sequence in the file?  Your Answer Score  1081 Correct 1.00 1721	Total		1.00 / 1.00	
<ul> <li>○ 1081</li> <li>○ 475</li> <li>○ Correct</li> <li>○ 834</li> <li>○ 1721</li> </ul>	Question 3 What is the length of the shortest seque	nce in the file?		
© 475 Correct 1.00  © 834  © 1721	Your Answer		Score	
© 834 © 1721	C 1081			
° 1721	<ul><li>475</li></ul>	Correct	1.00	
	© 834			
Total 1.00 / 1.00	° 1721			
	Total		1.00 / 1.00	

# **Question 4**

What is the length of the longest ORF appearing in reading frame 2 of any of the sequences?

Your Answer		Score
° 1271		
• 1134	Correct	1.00
° 1272		
° <sub>1488</sub>		
Total		1.00 / 1.00

# **Question 5**

What is the starting position of the longest ORF in reading frame 3 in any of the sequences? The position should indicate the character number where the ORF begins. For instance, the following ORF:

> sequence1

ATGCCCTAG

starts at position 1.

Your Answer		Score	
	Correct	1.00	
° 2100			

° 237	
° 575	
Total	1.00 / 1.00

## **Question 6**

What is the length of the longest ORF appearing in any sequence in any of the 3 forward reading frames?

Your Answer		Score
C 1488		
<ul><li>1518</li></ul>	Correct	1.00
° 1272		
° 2712		
Total		1.00 / 1.00

# **Question 7**

What is the length of the longest ORF that appears in the sequence with the identifier gi|142022655|gb|EQ086233.1|97?

Your Answer		Score	
C 588			
° 2067			
° 1551			
• 1272	Correct	1.00	
Total		1.00 / 1.00	
Question 8  Find the most frequently occurring rein all?	epeat of length 6 in all sequences. Hov	v many times does it occur	
Your Answer		Score	
° 356			
<ul><li>158</li></ul>	Correct	1.00	
° 98			

° <sub>153</sub>

Total 1.00 / 1.00

## **Question 9**

Find all repeats of length 12 in the input file. Let's use Max to specify the number of copies of the most frequent repeat of length 12. How many different 12-base sequences occur Max times?

Your Answer		Score
° <sub>9</sub>		
° 1		
° 2		
	Correct	1.00
Total		1.00 / 1.00

### **Question 10**

Which one of the following repeats of length 7 has the largest number of occurrences?

Your Answer		Score
• CGGCGGC	Correct	1.00

° CGGCGCT

Total	1.00 / 1.00
GCGGCGC	
° TCGGCGC	