Biological Command Line Interpreter

Project Instructions: - Please read very well

- 1. The team should be composed of 4-5 students (min 4 and max 5) no more no less. You can form your team from different groups.
- 2. The deadline is **Friday 31**st of **December.** This deadline is sharp and won't be changed, so manage your time well.
- The submissions will be via blackboard.
- 4. You should submit all your code as ".py" python file named as follows:

ID1_ID2_ID3_ID4_Group.py

No txt files, no zip files, just the python file named properly.

5. **Cheating** cases will take -3 without discussions. (BE CAREFUL, you will not just get ZERO, you will take -3 if you cheated any part of the code)

Project Description

You shall use what you have learned in this course to develop a Biological Command Line Interpreter Simulator in python. It will be similar to the cmd in windows, but will take commands that apply some biological operations on biological data. You can use any built in libraries explained in the lectures.

The Input:

The input to your project will be a **command** to execute along with any **parameters** the command demands (this will be more clarified in the following section). You **MUST** use the **getopt** functionality that was explained in lectures to parse the input because you will run your python file as follows:

You can see that the command can is divided into two sections the first one is your program itself as an argument to python to be parsed and interpreted. The second part is the command your program shall consume and parse to be executed.

The parsing step should be done using **getopt** functionality to identify the command to be execute, get each argument and each option to be execute the command.

Your program should call the correct function in your code, that correspond to the given command, and pass it the given parameters needed to execute this command. E.g. calc_gc(args[1])

```
D:\FCI\BioPython>python project_trial.py gc agct
GC=50.0
```

For the above example, the command consists of two parts: the first one is "python project_trial.py" is the mandatory step for running your program. The second part is "gc agat", that is the command "gc"

and its argument seq "agat" that your program will parse. Your program should recognize which function to call corresponding to the "gc" command and pass it the sequence "agat" and the result of this function is your output which is in this case is the gc percentage of the given sequence.

Note: you should handle any errors that may occur like "wrong command name" or "missing parameters".

The required commands details are illustrated in the following section.

Commands Description

gc	
Usage	gc seq
Parameters	seq: a string represents the sequence
Description	This command takes a seq as a string and returns the gc percentage of it.
Sample run	gc AGCAT

transcribe		
Usage	transcribe seq	
Options and arguments	seq: a string represents the sequence	
Description	This command takes a seq as a string and returns its transcription.	
Sample run	transcribe AGCAT	

reverse_complement		
Usage	reverse_complement seq	
Options and arguments	seq: a string represents the sequence	
Description	This command takes a seq as a string and returns its reverse complement.	
Sample run	reverse_complement AGCAT	

calc_nbases		
Usage	calc_nbases seq	
Options	seq: a string represents the sequence	
and		
arguments		
Description	This command takes a seq and calculates its nbases	
Sample run	calc_nbases NGCTN	

	is_valid		
Usage	is_valid seq type		
Options and arguments	seq : a string represents the sequence type: a string that represents the type of the sequence. It can be one of these keywords [protein, dna, rna]		
Description	This command takes a seq and a type (protein, dna, rna) and returns a Boolean value of whether it's a valid type or not		
Sample run	is_valid NGCTN protein		

filter_nbases		
Usage	filter_nbases seq	
Options and arguments	seq: a string represents the sequence	
Description	This command takes a seq and returns the Seq after removing n bases	
Sample run	filter_nbases NGCTN	

seq_alignment		
Usage	seq_alignment seq1 seq2 [-o file]	
Options	seq1, seq2 : strings represents the sequence	
and	-o file : specifies the path of the output file	
arguments		
Description	This command takes 2 sequences as input and get all their alignments along with the score. The -o is an optional parameter if we need the output to be written on a file instead of the screen.	
Sample run	seq_alignment AGCCT AGC -o output.txt	

seq_alignment_files		
Usage	seq_alignment_files file1 file2 [-o file3]	
Options	file1, file2: specifies the paths of the files to be aligned that contain the	
and	sequences	
arguments	-o file3 : specifies the path of the output file	
Description	This command takes 2 fasta files as input, each file contains a single sequence. It reads the 2 sequences from files and get all their alignments along with the score. The -o is an optional parameter if we need the output to be written on a file instead of the screen.	
Sample run	seq_alignment s1.fasta s1.fasta -o output.txt	

online_alignment		
Usage	online_align seq [-o file]	
Options	seq : a string represents the sequence	
and	-o file : specifies the path of the output file	
arguments		
Description	This command takes a sequence and uses BLAST to search the internet for its alignments. The output should be all the information in the resultant BLAST record. The -o is an optional parameter if we need the output to be written on a file instead of the screen.	
Sample run	seq_alignment ACTGCCGTCAAGTCAG -o output.txt	

	merge_fasta		
Usage	merge_fasta file1 file2 [file3] [-o output_file]		
Options and arguments	file1, file2, etc.: specifies the paths of the files to be merged -o output_file: specifies the path of the output file		
Description	This command takes any number of fasta files (at least two) and merge their contents into one fasta output file. There is an option to write the merge result in a file using -o option, otherwise the merge result will be displayed on the console. Hint: use variadic parameters in functions to handle the unknown number of parameters. Don't use lists.		
Sample run	merge_fasta f1.fasta f2.fasta f3.fasta f4.fasta		

convert_to_fasta		
Usage	convert_to_fasta file	
Options and arguments	file: specifies the path of a genbank file	
Description	This command converts the input genbank file with multiple records onto a fasta formatted file. The output is to be written in a different output fasta file.	
Sample run	convert_to_fasta "ls-orchid.gbk"	

Grading Criteria

Using Getopt	1
Error Handling	0.5
Gc	0.5
Transcribe	0.5
Reverse_complement	0.5
Calc_n_bases	0.5
Is_valid	0.5
Filter_n_bases	0.5
Seq_alignment S1 S2 [-o output_file]	1.5
seq_alignment file1 file2 [-o file name]	1.5
online_align s1 [-o file name]	1.5
merge file1 file2 (variadic parameters)	1.5
convert_to_fasta genbank_file	1.5
Total	12