# Programming Bootcamp 2015: Lab 7

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#### Name:

If you would like to receive points for your work, copy and paste your code into this document, save it, and send it to **sarahmid@mail.med.upenn.edu** with "Bootcamp Lab7" as the subject line **before \*10PM\* on Thursday**, **\*5/25\*** (NOTE DIFFERENT TIME/DAY!). This will be the final graded problem set! The winner(s) will be announced at the meeting on 5/26.

## 1 os and glob practice (6pts)

For this problem, use horrible.fasta and the fasta-reading function we created in lab6 (use the my solutions if you did not get to this yet).

(a) (2 pt) Write a script that takes two command line arguments: an input file name (assumed to be a fasta file) and an output folder name. Check if the input file exists; if it doesn't, print an error message. Then check if the output folder exists; if it doesn't, create it.

(b) (2 pt) Change the script so that it also reads in the fasta file (if it exists) and prints each individual sequence to a separate file. The files should be named <SEQID>.fasta, where <SEQID> is the name of the sequence, and all files should be output to the specified output folder.

(c) (2 pt) Now use glob to get a list of all files in the output folder from part (b) that have a .fasta extension. For each file, print just the file name (not the file path) to the screen.

## 2 optparse practice (6pts)

Here you will create a program that generates a random dataset of sequences. This program will accept the following args/options:

```
OUTFILE (required) - file the sequences will be printed to NUM_SEQS (optional; default = 1) - number of sequences to create MIN_LEN (optional; default = 100) - minimum sequence length MAX_LEN (optional; default = 100) - maximum sequence length RNA (optional; a flag (true or false); default = False) - whether U or T is used ID_PREFIX (optional; default = "seq") - Prefix to be added to beginning of each seq ID
```

The program should print the indicated number of randomly generated sequences, each with a length randomly chosen from the indicated range, to the indicated output file. If the –rna flag is used, use U's instead of T's. The output file should be in fasta format, and the sequence ID of each sequence should be the indicated prefix followed by a unique number (make sure each ID is unique; you can just use a counter for the number).

Run your script to create a large file of 10,000 random sequences of random length 500-5,000 nt. Call this file fake.fasta – it will be used in the next problem.

## 3 time practice (6pts)

(a) (2 pt) Is it faster to use the built-in function str.replace() or to loop through a string and replace characters manually? Compare the two by replacing all the T's in fake.fasta with U's using each method separately and comparing how long they take to run. Put both your code and your answer below.

(b) (2 pt) Is it faster to use the built-in function str.count() or to loop through a string and count characters manually? Compare the two by counting all the A's in fake.fasta using each method separately and comparing how long they take to run. Put both your code and answer below.

(c) (2 pt) Is it faster to use a list or a dictionary as a lookup table? Read in fake.fasta, ignoring everything but the header lines. Count the number of unique IDs (headers) using a dictionary and then a list, and compare how long each method takes to run. Put both your code and answer below.

## 4 subprocess practice (6pts)

(a) (2 pt) Write a script that runs the command 1s -1. Print the captured output to the screen.

(b) (4 pt) Write a script that (1) creates a random dataset (by calling the script you created in question 2), (2) gets statistics about the sequences in the newly generated file (by calling the script you wrote in lab6, question 4b) and then (3) (optionally) launches an R script to graph the output (using the supplied graph\_gc.r script). You will need R installed with ggplot to make the script work; if you don't have it then you can skip that last step for now. All scripts should be called using subprocess.Popen() (in other words, DO NOT copy/paste all the code from those other scripts into one script; the point is to practice using subprocess to launch existing scripts!).

The R script can be called as follows:

"Rscript graph\_gc.r seq\_info.txt"

where seq\_info.txt can be replaced with the name of the output file of statistics (Length/GC) that you create in step(2).