Assignment 1: Counting nucleotides and determining if a sequence is i.i.d.

Due date: Friday, 1/29

The goal of this homework assignment is to get comfortable with handling genomic sequences using simple scripts to test the hypothesis that genomic sequences are "i.i.d." (independent and identically distributed) an assumption that most statistical models make. We will obtain sequences from human chromosome 22 to calculate the frequencies of nucleotides and dinucleotides and compare these frequencies to those generated with a first-order random background model.

Part 1

Obtain the human chr22 sequence.

Preparatory Steps

ssh to the class server using your username and password

You should see username@genomic:c:c(or something similar) where username is replaced with your username. This is the command prompt.

Setup your assignment 1 directories

Make your assignment1 directory

Note: do not type the '\$', it's only included to tell you this is a command prompt.

```
$ mkdir assignment1
```

Change directories to your assignment1 directory

```
$ cd assignment1
```

Make your work and submission directories

```
$ mkdir work
```

```
$ mkdir submission
```

Change directories to your work directory

```
$ cd work
```

Download human chromosome 22 from NCBI using <u>FTP</u> Connect to NCBI

```
$ ftp ftp.ncbi.nih.gov
```

Follow the instructions to log into ftp anonymously

type "anonymous" as your username and "email" as your password.

Download human chromosome 22. The "ftp>" is the FTP prompt (which will show up on the command prompt/terminal). Do not type "ftp>".

```
ftp> cd genomes/Homo_sapiens/CHR_22
ftp> get hs_ref_GRCh38.p2_chr22.fa.gz
```

Terminate the FTP session

```
ftp> bye
Unzip the file
$ gunzip hs ref GRCh38.p2 chr22.fa.gz
```

There should be a file called hs_ref_GRCh38.p2_chr22.fa now in your directory.

At the end of this process, you will have a sequence file in <u>fasta format</u>. You can look at the first 10 lines of this file by typing

```
$ head hs_ref_GRCh38.p2_chr22.fa
```

We have created a template README.txt file for you to edit and turn in. Please replace '{}' and everything in between with your answers, but keep everything else the same. Copy the template to your working directory

```
$ cp /home/assignments/assignment1/README.txt .
```

Part 2

The script nuc_count.py counts the number of As, Cs, Gs, and Ts, and Ns in a fasta sequence file and prints the results. This script only counts one strand and is case-insensitive, e.g., both a and A bases are used to count the number of As in the sequence.

```
The usage of nuc_count.py is:

$ python3 nuc_count.py <fasta>
```

Before using this script, copy it to your work directory

\$ cp /home/assignments/assignment1/nuc_count.py .

Question 1

Run nuc_count.py on hs_ref_GRCh38_chr22.fa. How many times do each of the 4 nucleotides occur in chr22?

Part 3

Modify nuc_count.py, so that it also outputs frequencies of A, C, G, T. <u>Ignore N (and any other nonACGT nucleotides) from this point forward.</u>

Question 2

Run your modified nuc_count.py on hs_ref_GRCh38_chr22.fa. What are the frequencies of the 4 nucleotides on chr22?

Part 4

In this section, you will finish writing a script, make_seq.py, that generates a random sequence given a sequence length and nucleotide frequencies.

The usage of make_seq.py is:

```
$ python3 make_seq.py <sequence_length> <A_freq> <C_freq> <G_freq> <T_freq>
```

The script prints a random sequence of length <sequence length> to the terminal (stdout). The random sequence should have the same nucleotide frequencies as the input nucleotide frequencies.

```
First, copy make_seq.py to your work directory.

$ cp /home/assignments/assignment1/make seq.py . ← note the period.
```

Finish the script by writing code where it says "TODO". Refer to the assignment 1 presentation for tips on random number generators. Test to see if your code is working by generating files with different nucleotide frequency inputs and then checking the nucleotide frequencies within those files using your modified nuc_count.py script.

Using make_seq.py, generate a random sequence with length 1,000,000 using the nucleotide frequencies calculated in part 3 and save it to random_seq_1M.txt. Please keep at least two decimal places from the original calculated frequencies when generating your new random sequence file.

To save the sequence to a file, redirect the standard output using ">". Here's an example of this: python3 make_seq.py 1000 0.25 0.25 0.25 0.25 > random_seq_1k.txt

Part 5

Modify nuc_count.py to also output frequencies of all dinucleotides using an overlapping window method. By an overlapping method we mean if the sequence is 'ACGC', then there are 3 di-nucleotides, 'AC', 'CG', and 'GC' instead of 'AC' and 'GC'.

When you run nuc_count.py the output should look similar to this:

```
$ python3 nuc_count.py <fasta>
```

```
Dinucleotide Frequencies
AA:0.058
AC:0.128
AG:0.02
AT:0.06
CA:0.135
CC:0.14
CG:0.023
CT:0.083
GA:0.018
GC:0.038
GG:0.005
GT:0.013
TA:0.055
TC:0.075
TG:0.025
```

TT:0.04

Question 3

Run the modified nuc_count.py for both human chr22 and your generated 'random_seq_1M.txt' from part 4. Compare the two lists of frequencies. What are the differences? Can you provide a biological explanation for these differences?

What to turn in

Two modified scripts nuc_count.py and make_seq.py.
A completed README.txt.
The sequence file random_seq_1M.txt

These four files should be in your assignment1/submission folder.

Note: to copy your work files to your submission folder, type \$ cp <file_name> ~/assignment1/submission/ where <file_name> is the name of the file you want to copy.

Changes

V2 (1/22 12pm): Minor: updated typos in path names in part 2 and part 4 V3 (1/22 4pm): Minor: updated typo in fasta filename in part 1