Lab 04 Files and Modules

Deliverables:

- sequenceAnalysis.py module 35 total points classes included:

 - NucParams,
 - FastAreader

ProteinParams,

- genomeAnalyzer.py 15 total points possible extra credit - 10 additional points
- Due: Monday April 27, 2020 11:55pm Congratulations. You have started to build an inventory of some pretty useful functions. Because

now, you may decide to keep it somewhere else later.

deposit into your very own sequenceAnalysis toolkit. In Python, these toolkits are called modules.

We are also going to start using real sequence files. The fastA format, described here: en.wikipedia.org/wiki/FASTA_format is very convenient to use and fully capable of storing sequences of many types. You will be reading these from an input file for this assignment.

these are written as classes, you can easily reuse them. Your ProteinParam class is your first

There are a few things that we can do that mirror and extend the analyses that we did previously on protein sequences. We can calculate composition statistics on the genome (gc content for example), we can calculate relative codon usage in the genome, and we can calculate amino acid

Genomic analysis

composition by translating those codons used the genome. For this lab, I have provided a NucParams class, with the required methods that it implements (see below). You will need to design and write those methods, and these are to be placed in a file called sequenceAnalysis.py This is a *module* that you can use from now on.

You will also need to place the ProteinParams class (Lab 3) into this module also. This class will not be used for this assignment, but place it into your toolbox. I have written the FastAreader class. It is included (below). Keep it as is part of your module for

The input file for this assignment will be named testGenome.fa, and will be available in Canvas soon. You will not need to submit testGenome.fa, but it will be necessary for your testing. For

development and testing, create a new directory (Lab04) and place the data file (testGenome.fa),

your program (genomeAnalyzer.py) and your new module (sequenceAnalysis.py) into your Lab04

Hints • Python modules have the .py extension as files, but when they are imported, the name without the extension is used in the import statement in your program. • File placement: Make sure to place your program, your sequenceAnalysis module and the

required data files in the same folder. This will allow Python to find them. Read over the FastAreader usage to see how to specify file names that you can use for your data.

directory.

- Codon frequency calculations
- Notice that NucParams does all of the counting you need. It is responsible for counts of codons and their translated amino acids. Your genomeAnalyzer.py program has the task of determining which codons are preferred for

each of the amino acids and calculating the relative percentage. For any given amino acid, the

relative codon usage (percentages) should sum to 100.0%. Notice that Methionine and

Tryptophan only have 1 codon that codes for those, so these will have relative codon usages of 100%. For example: Lysine is coded by both AAA (607) and AAG (917) (example counts in parentheses).

From our aaComposition() method, we are given the aaComposition dictionary and we can

607/1524 for AAA and 917/1524 for AAG. The associated percentages are thus: 39.8 for AAA and 60.2 for AAG. AAA = 607/1524 * 100 = 39.8%AAG = 917/1524 * 100 = 60.2%

best structure? This class (NucParams) is intended to be very similar to ProteinParam. Make sure

to read addSequence() before making this decision, and remember that objects of this class may

need to handle an arbitrarily large number of sequences (hint: dictionaries are good). As a second

lookup 'K' to find 1524 counts (these came from those 607+917 codons). We can then calculate

The constructor of the class has one optional parameter, a sequence of type string. It may include upper or lower case letters of the set {ACGTUN} or whitespace. These will be gene sequences and they begin in frame 1. In other words the first 3 letters of the sequence encode the first AA of the sequence. Carefully consider in what form this class should maintain its data. Is a string the

addSequence() - 5 pts

_init__

hint, notice that init and addSequence are doing VERY similar things - you could just make one of them do most of the work.

Design specification - sequenceAnalysis.py

(if any). aaComposition() - 10 pts This method will return a dictionary of counts over the 20 amino acids and stop codons. This dictionary is VERY similar to the lab 3 aaComposition, though you must decode the codon first. The translation table from codon to AA is provided. You are counting amino acids by translating from the proper codon table.

This method returns a dictionary of counts of valid nucleotides found in the analysis. (ACGTNU).

DNA nucleotides, they should be counted as DNA nucleotides. Any N bases found should be

This dictionary returns counts of codons. Presume that sequences start in frame 1, accept the

alphabet {ACGTUN} and store codons in RNA format, along with their counts. Any codons found

with invalid bases should be discarded. Discard codons that contain N bases. This means that

If you were given RNA nucleotides, they should be counted as RNA nucleotides. If you were given

This method must accept new sequences, from the {ACGTUN} alphabet, and can be presumed

to start in frame 1. This data must be added to the data that you were given with the **init** method

all codon counts are stored as RNA codons, even if the input happens to be DNA. If you discard a codon, make sure to not alter the frame of subsequent codons. nucCount()

Input must be from STDIN Your FastaReader object will read data from sys.stdin if it is not given a filename. You can specify

using standard files.

summaries and final display of the data.

myReader = FastAreader('testGenome.fa') # make sure to change this to use stdin with:

a filename for your testing in jupyter and then remove that filename argument when you move to

• lines 5 - 68 are the output statistics on relative codon usage for each codon ordered by codon within each amino acid group as follows:

Contain in-line comments

Nuc Params

rnaCodonTable = {

RNA codon table

In [33]: class NucParams:

U

A

G

onTable.items()}

pass

pass

Output format - 15 pts

second line is blank

fourth line is blank

XXX : A F (D)

32.8 (2270) ...

The function to output the results of your analysis has specific formatting rules that you must follow to get full credit. These rules are as follows: • First line: sequence length = X.XX Mb with two digits after the decimal and labeled Mb (you

• third line: GC content = XX.X% as a percentage with one digit after the decimal

sequence length = 3.14 Mb GC content = 60.2% UAA: - 32.6 (1041) UAG: - 38.6 (1230) UGA: - 28.8 (918) GCA: A 14.1 (10605) GCC: A 40.5 (30524) GCG: A 30.5 (22991) GCU: A 14.9 (11238) UGC: C 67.2 (4653) UGU: C

• Include any assumptions or design decisions you made in writing your code

• contain proper docstrings for the program, classes, modules and any public functions.

where XXX is the three letters for an RNA codon, A is the 1-letter amino acid code, F is relative

codon frequency, use {:5.1f} for the format, and D is for codon count, use the format {:6d}. There

is a single space between each of these fields. For example (this is not representative of any real

- Extra credit 10 pts possible You now have a very powerful set of classes for evaluating genomes. Write a compareGenomes.py program that compares GC content, aaComposition and relative codon
 - # C 'CUU': 'L', 'CCU': 'P', 'CAU': 'H', 'CGU': 'R', # CXU 'CUC': 'L', 'CCC': 'P', 'CAC': 'H', 'CGC': 'R', # CXC 'CUA': 'L', 'CCA': 'P', 'CAA': 'Q', 'CGA': 'R', # CXA 'CUG': 'L', 'CCG': 'P', 'CAG': 'Q', 'CGG': 'R', # CXG

codonComposition() - 10 pts

counted also. Invalid bases are to be ignored in this dictionary.

nucComposition() - 10 pts

This returns an integer value, summing every valid nucleotide (ACGTUN) found. This value should exactly equal the sum over the nucleotide composition dictionary. Design specification - genomeAnalyzer.py

This program must import your sequenceAnalysis module. It is responsible for preparing the

You would replace:

need to calculate the number of bases in Mb).

myReader = FastAreader() # make sure to change this to use stdin

genome):

To get full credit on this assignment, your code needs to:

Run properly (execute and produce the correct output).

Congratulations, you have finished your fourth lab assignment!

bias of 2 genomes. You will have a halophile genome and a hyperthermophile genome to compare. Submit your code using canvas

'UUU': 'F', 'UCU': 'S', 'UAU': 'Y', 'UGU': 'C', # UXU

'UUC': 'F', 'UCC': 'S', 'UAC': 'Y', 'UGC': 'C', # UxC

'UUA': 'L', 'UCA': 'S', 'UAA': '-', 'UGA': '-', # UXA

'UUG': 'L', 'UCG': 'S', 'UAG': '-', 'UGG': 'W', # UxG

'GUU': 'V', 'GCU': 'A', 'GAU': 'D', 'GGU': 'G', # GXU

'GUC': 'V', 'GCC': 'A', 'GAC': 'D', 'GGC': 'G', # GXC

'GUA': 'V', 'GCA': 'A', 'GAA': 'E', 'GGA': 'G', # GXA

'GUG': 'V', 'GCG': 'A', 'GAG': 'E', 'GGG': 'G' # GXG

dnaCodonTable = {key.replace('U', 'T'):value for key, value in rnaCod

'AUU': 'I', 'ACU': 'T', 'AAU': 'N', 'AGU': 'S', # AxU 'AUC': 'I', 'ACC': 'T', 'AAC': 'N', 'AGC': 'S', # AxC 'AUA': 'I', 'ACA': 'T', 'AAA': 'K', 'AGA': 'R', # AXA 'AUG': 'M', 'ACG': 'T', 'AAG': 'K', 'AGG': 'R', # AXG

def init (self, inString=''):

def addSequence (self, inSeq):

header = ''

sequence = ''

skip to first fasta header

while not line.startswith('>') :

if line.startswith ('>'):

sequence = ''

yield header, sequence

header = line[1:].rstrip()

sequence += ''.join(line.rstrip().split()).upper()

line = fileH.readline()

line = fileH.readline()

for line in fileH:

else:

yield header, sequence

header = line[1:].rstrip()

```
def aaComposition(self):
                 return self.aaComp
             def nucComposition(self):
                 return self.nucComp
             def codonComposition(self):
                 return self.codonComp
             def nucCount(self):
                 return sum(self.nucComp)
         FastAreader
In [34]:
         import sys
         class FastAreader :
              , , ,
             Define objects to read FastA files.
             instantiation:
             thisReader = FastAreader ('testTiny.fa')
             usage:
             for head, seq in thisReader.readFasta():
                 print (head, seq)
             def init (self, fname=''):
                 '''contructor: saves attribute fname '''
                 self.fname = fname
             def doOpen (self):
                  ''' Handle file opens, allowing STDIN.'''
                 if self.fname is '':
                     return sys.stdin
                 else:
                     return open(self.fname)
             def readFasta (self):
                  ''' Read an entire FastA record and return the sequence header/s
         equence'''
                 header = ''
                 sequence = ''
                 with self.doOpen() as fileH:
```

```
Main
        Here is a jupyter framework that may come in handy
In [ ]: def main ():
            myReader = FastAreader('testGenome.fa') # make sure to change this t
        o use stdin
            myNuc = NucParams()
            for head, seq in myReader.readFasta() :
                myNuc.addSequence(seq)
            # sort codons in alpha order, by Amino Acid
            # calculate relative codon usage for each codon and print
            for nucl in nucs:
                print ('{:s}: {:s} {:5.1f} ({:6d})'.format(nuc, aa, val*100, th
        isCodonComp[nuc]))
        if name == " main ":
            main()
```