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Deliverables: 36 total points

    seqCleaner 9 points

    fastqParse 9 points

    coordinateMathSoln 9 points

    converter 9 points

    Optional: Triad class rewrite (5 points extra credit)

              These must be submitted as 4(or 5) .py files,
              Due: Monday April 13, 2020 11:55pm
         Lab 2 – Manipulating Data Types (36 points)
         Overview
         For this assignment, create a Lab02 folder inside your BME160 folder. Download the Lab02 ipynb file to that new folder and we will work on the notebook in
         that folder.
         Last week we introduced the Python function input(), which is used to take in string data. This week we will use input() to take in string and numeric (float and
         integer) data.
          Remember that all data returned from input() is a string object, so this will mean that you need to convert any numeric data to their respective numeric objects.
         The exercises in this lab will give you practice manipulating various types of data that commonly arise in computational biology problems.
         We are again using jupyter for this assignment, and as a final step we will create distinct command line programs from each of the cells that make up these
         programs. Each of those program files are text files and will have names like: seqCleaner.py, or fastqParse.py. Please save each of the four (or five) python
         programs as .py files into your LAB02 folder. You can copy the text from the code cell, then paste it into an editor like notepad or textedit. Save that new file
         with the appropriate name (eg. seqCleaner.py) into your LAB02 folder. Using terminal or cmd or conda, you can now do a final test on that new program with
         the command:
         python3 seqCleaner.py
          All together, you will submit:

    seqCleaner.py,

           fastqParse,py,

    coordinateMathSoln.py,

           converter.py, and
           (optionally) Triad.py
         Sequence cleanup
         In this exercise, you will create a program to "clean up" a sequence of DNA by removing ambiguous bases (denoted by "N") output from a sequencer. Your task
         is to create a Python program called segCleaner that

    asks for and collects a sequence of DNA using input()

           • removes the ambiguous parts of the sequence, outputs the "cleaned" sequence, replacing the ambiguous parts with a count in {}'s. For example, if I enter
              the sequence of DNA "AaNNNNNNGTC" (without quotes), the program will output:
             AA{6}GTC
         Hints:
           • The input sequence is not guaranteed to be uppercase, but should be interpreted as though it is all uppercase.
           • Only the letters (A,C,G,T,N) will be included in the input.

    Only the DNA characters (A,C,G,T) should remain after the cleanup.

    The input will include, at most, one block of 'N' characters.

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

    Run properly (execute and produce the correct output)

           • Contain docstrings and specific line or block comments that explain the semantics of your implementation.

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

           • Include an overview describing what your program does with expected inputs and outputs. This should be in the form of a program level dosctring.
         seqCleaner
In [ ]: #!/usr/bin/env python3
          # Name: Your full name (CATS account username)
          # Group Members: List full names (CATS usernames) or "None"
          Read a DNA string from user input and return a collapsed substring of embedded Ns to: {count}.
          Example:
           input: AaNNNNNNGTC
          output: AA{6}GTC
          Any lower case letters are converted to uppercase
          class DNAstring (str):
               def length (self):
                   return (length(self))
               def purify(self):
                    ''' Return an upcased version of the string, collapsing a single run of Ns.'''
                   pass # this is just a placeholder, so dont leave "pass" here
          def main():
               ''' Get user DNA data and clean it up.'''
              data = input('DNA data?')
              thisDNA = DNAstring (data)
              pureData = thisDNA.purify()
              print (pureData)
          main()
         Sequence information parsing
          In this exercise, you will create a program to "parse" sequence name information from a single line of a FASTQ formatted file. Your task is to create a Python
          script called fastqParse that:

    asks for and collects the seqname line of a FASTQ file using input()

           • parses out each field of the run information from the string and displays each of them on a new line For example, if I enter the FASTQ seqname line:
              @EAS139:136:FC706VJ:2:2104:15343:197393 then the program will output:
             Instrument = EAS139
              Run ID = 136
              Flow Cell ID = FC706VJ
              Flow Cell Lane = 2
              Tile Number = 2104
             X-coord = 15343
              Y-coord = 197393 ### Hints:

    The input string is guaranteed to have 7 fields.

           • The first character of the FASTQ segname line is "@" and each field of the run information is separated by a colon":".
           • A reasonable solution would be around 16 lines of code excluding comments.
          To get full credit on this assignment, your code needs to:

    Run properly (execute and produce the correct output)

    Contain documentation/comments

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

    Include an overview describing what your program does with expected inputs and outputs

         fastqParse
In [ ]: #!/usr/bin/env python3
          # Name: Your full name (CATS account username)
          # Group Members: List full names (CATS usernames) or "None"
          Program docstring goes here.
          class FastqString (str):
               ''' Class docstring goes here.'''
               def parse(self):
                    ''' Method docstring goes here.'''
          def main():
               ''' Function docstring goes here.'''
              pass
          main()
         Protein coordinates
         In this exercise, you will create a program that takes three sets of atomic coordinates, all provided on a single line. The program then calculates the bond
          lengths and angles. For this program, you can start with the Triad class (provided). Your task is to create a Python program called coordinateMathSoln.py that:

    asks for and collects three sets of coordinates using input(), only use 1 line for this data!!

           • outputs the N-C and N-Ca bond lengths and the C-N-Ca bond angle with correct number of significant digits (see below)
         For example, if I enter the following coordinates (notice.. they are all on one line !!!):
         C = (39.447, 94.657, 11.824) N = (39.292, 95.716, 11.027) Ca = (39.462, 97.101, 11.465)
         then the program will output the following three lines:
         N-C bond length = 1.33
         N-Ca bond length = 1.46
         C-N-Ca bond angle = 124.0
         (Note: make sure that the angle returned is in degrees !!)
         Hints:
           • Each coordinate will contain only 3 numeric values.
           • Bond lengths are the distance between two points, which for points P and Q in 3-space, (P_x, P_y, P_z) and (Q_x, Q_y, Q_z) respectively, the distance
              between them is:
                                                       ||PQ|| = \sqrt{(P_x - Q_x)^2 + (P_y - Q_y)^2 + (P_z - Q_z)^2}= \sqrt{\sum_{i \in x, y, z} (P_i - Q_i)^2}

    Bond angles can be calculated from the dot product.

         Let's say that we have three points in space labeled P, Q and R. We are interested in the angle at point Q that is made from line segments QP and QR. The dot
          product tells us (for standard vectors P and R) that:
                                                                      P \cdot R = ||P|| ||R|| \cos \theta
         in this notation, ||P|| refers to the length of vector P as a standard vector (assumed to begin at the origin (0,0,0)). We can then see that the angle between
         vectors P and R can be found by:
                                                                        \cos \theta = \frac{P \cdot R}{R}
                                                                                  ||P|||R||
          We can calculate the dot product using the sum of products of the vector components:
                                                                       P \cdot R = \sum_{i \in x, y, z} P_i R_i
         Now, to find vector P in standard form, we need to remember that QP starts at Q, so we need to place the origin at Q and find out where P is in that new space.
         We do that by subtracting the components of Q from P. Putting all of this together, we get:
                                                              \theta = \cos^{-1} \frac{\sum_{i \in x, y, z} (P_i - Q_i)(R_i - Q_i)}{\|QP\| \|QR\|}
          Remember, \theta is in radians.
          Below I have given you a class (Triad) with methods to calculate dot products (dot), dot products of translated vectors (ndot), distances (dPQ, dPR, dPR) and
         angles in radians (angleP, angleQ and angleP) for each of the three points in a Triad object. A reasonable solution for this exercise involves around 12 lines of
         additional code excluding comments.
          To get full credit on this assignment, your code needs to:

    Run properly (execute and produce the correct output)

    Contain docstrings and line comments (using #)

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

    Include an overview describing what your program does with expected inputs and outputs as a program level docstring

         coordinateMathSoln
In [ ]: #!/usr/bin/env python3
          # Name: Your full name (CATS account username)
          # Group Members: List full names (CATS usernames) or "None"
          Program docstring goes here
          import math
          class Triad:
               Calculate angles and distances among a triad of points.
               Author: David Bernick
               Date: March 21, 2013
               Points can be supplied in any dimensional space as long as they are consistent.
               Points are supplied as tupels in n-dimensions, and there should be three
               of those to make the triad. Each point is positionally named as p,q,r
               and the corresponding angles are then angleP, angleQ and angleR.
               Distances are given by dPQ(), dPR() and dQR()
               Required Modules: math
               initialized: 3 positional tuples representing Points in n-space
                         p1 = Triad(p=(1,0,0), q=(0,0,0), r=(0,1,0))
               attributes: p,q,r the 3 tuples representing points in N-space
               methods: angleP(), angleR(), angleQ() angles measured in radians
                      dPQ(), dPR(), dQR() distances in the same units of p,q,r
               11 11 11
               def __init__(self,p,q,r) :
                    """ Construct a Triad.
                    Example construction:
                       p1 = Triad(p=(1.,0.,0.), q=(0.,0.,0.), r=(0.,0.,0.)).
                   self.p = p
                   self.q = q
                   self.r = r
          # private helper methods
               def d2 (self,a,b) : # calculate squared distance of point a to b
                    return float(sum((ia-ib)*(ia-ib) for ia,ib in zip (a,b)))
               def dot (self,a,b) : # dotProd of standard vectors a,b
                    return float(sum(ia*ib for ia,ib in zip(a,b)))
               def ndot (self,a,b,c) : # dotProd of vec. a,c standardized to b
                    return float(sum((ia-ib)*(ic-ib) for ia,ib,ic in zip (a,b,c)))
          # calculate lengths(distances) of segments PQ, PR and QR
               def dPQ (self):
                    """ Provides the distance between point p and point q """
                    return math.sqrt(self.d2(self.p,self.q))
               def dPR (self):
                    """ Provides the distance between point p and point r """
                   return math.sqrt(self.d2(self.p,self.r))
               def dQR (self):
                    """ Provides the distance between point q and point r """
                    return math.sqrt(self.d2(self.q,self.r))
               def angleP (self) :
                    """ Provides the angle made at point p by segments pq and pr (radians). """
                    return math.acos(self.ndot(self.q,self.p,self.r) / math.sqrt(self.d2(self.q,self.p)*self.d2(self.r,self.p)))
               def angleQ (self) :
                    """ Provides the angle made at point q by segments qp and qr (radians). """
                   return math.acos(self.ndot(self.p,self.q,self.r) / math.sqrt(self.d2(self.p,self.q)*self.d2(self.r,self.q)))
               def angleR (self) :
                    """ Provides the angle made at point r by segments rp and rq (radians). """
                    return math.acos(self.ndot(self.p,self.r,self.q) / math.sqrt(self.d2(self.p,self.r)*self.d2(self.q,self.r)))
          def main():
               ''' Function docstring goes here'''
          main()
         Extra credit (5 points): Rewrite the Triad class.
         For extra-credit, provide a direct replacement for the Triad class. The external methods that calculate angles, distances, and points (tuples) p,q and r must be
         maintained such that either version of the Triad class can be used.
          You could use the cosine law to calculate angles instead of the dot product. You might make use of the numpy module. You might recode each of the methods
         to avoid using zip. You might consider using list iterations. Your Triad replacement must reimplement all of Triad public function, without using zip and without
         being a trivial rewrite. Your implementation need not be as compact as the current implementation, and it needs to be correct and fully documented to receive
         full credit.
          Triad rewrite
```

## self.r = r def dPQ (self): """ Provides the distance between point p and point q """ pass def dPR (self):

p1 = Triad(p=(1.,0.,0.), q=(0.,0.,0.), r=(0.,0.,0.)).

""" Provides the distance between point p and point r """

""" Provides the distance between point q and point r """

def init (self,p,q,r):

self.p = pself.q = q

pass

def dQR (self):

""" Construct a Triad.

Example object construction:

In [ ]: class Triad:

```
pass
     def angleP (self) :
          """ Provides the angle made at point p by segments pq and pr (radians). """
          pass
     def angleQ (self) :
          """ Provides the angle made at point q by segments qp and qr (radians). """
          pass
          """ Provides the angle made at point r by segments rp and rq (radians). """
Codon tables and amino acid letter converters
In this exercise, you will create a program that uses mappings to convert sequence information between different amino acid representations. This includes the
3-letter codon code (RNA and DNA), the one letter amino acid code and the 3-letter amino acid code. The program will use different dictionaries that represent
: codon tables – one for DNA and the other for RNA, and amino acid letter representation converters.
Your task is to create a Python program called converter that asks for and collects a single input string using input() parses the string, looks up the information
in the appropriate dictionary, and outputs the correct conversion For example:
if I enter "ATG" (without quotes), then the program will output:
ATG = MET
if I enter "UAG" (without quotes), then the program will output:
UAG = ---
if I enter "E" (without quotes), then the program will output:
E = GLU
if I enter "Asp" (without quotes), then the program will output:
ASP = D
Hints:
  • The program might not get a valid codon. In that case, it should output 'unknown'. You can use the dictionary method 'get' and include a default_value to
    handle the 'unknown' case. See Model p. 72 for an example.
To get full credit on this assignment, your program needs to:

    Run properly (execute and produce the correct output)

    Contain docstrings and line comments (using #)

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

  • Include an overview describing what your program does with expected inputs and outputs as a program level docstring
```

## converter In [ ]: #!/usr/bin/env python3

# Name: Your full name (CATS account username)

Program docstring goes here

Email your code files to yourself

coordinateMathSoln.py 9 points

• triad.py Triad rewrite (optional) 5 points

• Run properly (execute and produce the correct output)

Congratulations, you finished your second lab assignment!

• Contain proper docstrings and appropriate line style comments (#)

seqCleaner.py 9 pointsfastqParse.py 9 points

• converter.py 9 points

files:

Copy your code files to a pen drive

Save your work on your own computer if you are using Anaconda (or equiv)

Important: to get full credit on this lab assignment, each of the code files you submit needs to:

short\_AA = {

# Group Members: List full names (CATS usernames) or "None"

'CYS': 'C', 'ASP': 'D', 'SER': 'S', 'GLN': 'Q', 'LYS': 'K',

'ILE': 'I', 'PRO': 'P', 'THR': 'T', 'PHE': 'F', 'ASN': 'N',

'GLY': 'G', 'HIS': 'H', 'LEU': 'L', 'ARG': 'R', 'TRP': 'W',

'ALA': 'A', 'VAL': 'V', 'GLU': 'E', 'TYR': 'Y', 'MET': 'M'

}

```
long AA = {value:key for key,value in short AA.items()}
RNA_codon_table = {
# Second Base
\# U C A
'UUU': 'Phe', 'UCU': 'Ser', 'UAU': 'Tyr', 'UGU': 'Cys',
'UUC': 'Phe', 'UCC': 'Ser', 'UAC': 'Tyr', 'UGC': 'Cys',
'UUA': 'Leu', 'UCA': 'Ser', 'UAA': '---', 'UGA': '---',
'UUG': 'Leu', 'UCG': 'Ser', 'UAG': '---', 'UGG': 'Trp',
'CUU': 'Leu', 'CCU': 'Pro', 'CAU': 'His', 'CGU': 'Arg',
'CUC': 'Leu', 'CCC': 'Pro', 'CAC': 'His', 'CGC': 'Arg',
'CUA': 'Leu', 'CCA': 'Pro', 'CAA': 'Gln', 'CGA': 'Arg',
'CUG': 'Leu', 'CCG': 'Pro', 'CAG': 'Gln', 'CGG': 'Arg',
'AUU': 'Ile', 'ACU': 'Thr', 'AAU': 'Asn', 'AGU': 'Ser',
'AUC': 'Ile', 'ACC': 'Thr', 'AAC': 'Asn', 'AGC': 'Ser',
'AUA': 'Ile', 'ACA': 'Thr', 'AAA': 'Lys', 'AGA': 'Arg',
'AUG': 'Met', 'ACG': 'Thr', 'AAG': 'Lys', 'AGG': 'Arg',
'GUU': 'Val', 'GCU': 'Ala', 'GAU': 'Asp', 'GGU': 'Gly',
'GUC': 'Val', 'GCC': 'Ala', 'GAC': 'Asp', 'GGC': 'Gly',
'GUA': 'Val', 'GCA': 'Ala', 'GAA': 'Glu', 'GGA': 'Gly',
'GUG': 'Val', 'GCG': 'Ala', 'GAG': 'Glu', 'GGG': 'Gly'
dnaCodonTable = {key.replace('U','T'):value for key, value in rnaCodonTable.items()}
def main():
    ''' Function docstring goes here'''
    pass
main()
Submit your code and answers
Important: please save your work before logging out of the computer. This will ensure that you have a copy of your work and you will avoid having to redo
everything in the event that something happens with the lab computers. The two recommended solutions (pick one) at this point are to:
```

The class will be using canvas to submit the source code files created as part of this lab assignment. For this lab, you should upload the following programs as