

# Inspection Introduction

For this assignment, provide a description of your MyBio code. Describe where you put the string constants and why you think they go there. Notice, in my example, that I put all of the string constants in the main part of the program, not in the class. Why might that be a useful way to design this simple program?

Deliverables: 30 total points

- helloMe 4 points
- helloAgain 5 points
- myBio 6 points
- countAT-bugsSoln 9 points
- countNucleotides 6 points

Due: Monday April 10, 2023 11:55pm

## Lab 1 – Introduction to Python (30 points)

Work in small groups. Find yourself a team who you will hopefully work with throughout the quarter. Although you are required to turn in independent code and answers yourself, you are encouraged to ask questions and work with each other to solve these exercises. Programming has a lot of area-specific vocabulary and concepts so it is often extremely helpful to ask for help when you run up against hurdles. Make sure to include your partners name(s) as a comment in the first few lines of your program, like this:

```
In [1]:#!/usr/bin/env python3
# Name: Your full name
# Group Members: List full names or "None"
```

### Setup your workspace

#### Install Anaconda on your machine

From the machine that you intend to do all of your work for BME 160, go to: <https://www.anaconda.com/products/individual/> From that page, select the Python 3.10 version by using the download button. This may take awhile, depending on your internet connection speed.

#### Downloading your first notebook

The document you are now reading is a jupyter notebook. It is available for download using our class website service Canvas. First we will need to create a folder that you'll use for this lab. To do this create a folder called bme160 on the desktop. From Canvas :: files :: jupyter, download Lab01.ipynb, and place it into your bme160 folder.

There are five programs you will need to write. They are all embedded in Lab01.ipynb as jupyter "cells". Make sure to place Lab01.ipynb into your Lab01 folder.

#### Accessing Jupyter

##### Open jupyter

There are a number of ways to startup jupyter. Here are a few:

- LocalHost Open *terminal* ( located in the utilities folder on MacOS ), then change to the Lab01 directory and startup: jupyter notebook
- <https://try.jupyter.org> you will then need to upload Lab 01.ipynb to that remote server. This server enforces a 10-minute inactivity limit, so you will need to save your work often. Note that this remote server sometimes doesnt run.
- Anaconda Navigator. Startup Anaconda, then select: jupyter notebook. From Jupyter, you can then navigate to your Desktop, then to bme160, and then open Lab01.ipynb. That should bring you to a page that looks just like the Lab01 assignment.

Try out **jupyter** by selecting the following cell, then push the "run" button in the toolbar above, or pull down the Cell/ menu above and select Run Cells. Notice that cells that contain runnable code start with In [ ]:

```
In [ ]:print ('hello Sally')
```

make sure that "hello" is returned.

From here on, you will be writing programs.

#### First python program

The hello.py program is one of the first python programs that people write.

Documenting code is an important and often overlooked process in software engineering. Don't fall into the trap of not adding comments to your code! Note, you will not receive full credit on your assignment if your code is not clearly documented. To add comments you use the # character for single line comments, or you can use triple-quotes (""") to begin a block of comments and then add another set of triple-quotes to end that block comment.

To get an idea of what documenting and comments look like, here is an old-school version of the 'hello.py' program, documented:

```
In [3]:#!/usr/bin/env python3
# Name: David Bernick (dbernick)
# Group Members: John Cleeze (jcleeze) and Eric Idle (eidle)
```

```
"""
Print out the string Hello World when run.
"""
# remember that strings in python use either single (') or double (") quotes.
print ("Hello World")

Hello World

Add your own similar-looking documentation and comments to the the following code, customizing it for yourself, and lets make it print: Hello your name instead. In my case, it would print "Hello David"
```

#### helloMe

```
In [5]:#!/usr/bin/env python3
# Name: David Bernick(dbernick)
# Group Members: none
```

```
"""
Print out the most common statement in computer programming, Hello World.
"""
```

```
''' this is a single line docstring '''
class Announcer (str): # Announcer is now defined as a kind of python string.
    def printMe (self):
        print (self)
```

```
student = Announcer ('Hello David')
dog = Announcer ('bark bark')
student.printMe()
dog.printMe()
```

Hello David  
bark bark

There are a few things to mention about this style of coding. Classes are templates for *objects*. The *announcer* class is a kind of *string* class, and we tell python that by including (str) after the name of the class. We have added a method ( printme ) to the announcer class, and that method allows objects of class announcer to *print* themselves. In python, we access data that is saved in an object using the word *self*. Since objects of this class are strings, *self* refers to the data ('Hello David' in this case). You will notice that python methods are defined using the keyword: `def name ( self, parameters)`

The line: `student = announcer ('Hello David')` creates a new name (student), of class *announcer* and it is given a parameter '*Hello David*'. That parameter is given to announcer as its data. Objects of class announcer refer to that data using the name *self*.

The line: `student.printme()` sends a printme() message to the student object, and the student object then prints out its data.

#### Return of "Hello World"

We will now take a look at a more sophisticated, second person version of *helloMe*. As currently written, it should ask you to enter a name; you type in the name and then press the *enter* key (maybe labelled return).

The output you should see is:

Hi there, I am name

Look at *helloAgain* below. You'll notice that it's more complex and longer than *helloMe*. Now modify *helloAgain* so that, in addition to asking for that person's name, the program also prompts the user for their favorite pet like this:

"What is your favorite pet? :"

To make this clearer, here are a few lines that belong in your revised *helloAgain*.

```
In [ ]:name = input( "What is your name? : " )
pet = input( "what is your favorite kind of pet? : " )
newPerson = Person (name, pet)
newPerson.introduce()
```

#### helloAgain

```
In [ ]:#!/usr/bin/env python3
# Name: David Bernick(dbernick)
# Group Members: none
```

```
"""
Build a Person object and have it introduce itself.
"""
```

```
input: a string of arbitrary length, which is used to name the new person object
output: greeting printed to screen
"""
```

```
class Person:
    def __init__(self,name,pet):
        self.myName = name
        self.myPet = pet

    def introduce (self):
        print ("Hi there, I am {}}, and i like {}s".format(self.myName,self.myPet))
```

# put your new code here.

Your new version of *helloAgain* should now print something that looks like this:

"Hi there, I am name and I like pets!"

replacing the italicized text with what the user entered at the command line. Note the statement uses a plural form by appending the letter s, your program should do this too.

So, if you decided that your new person is named Fred, and Fred is fond of his pet aardvark, then your program would output:

Hi there, I am Fred and I like aardvarks!

Remember to save and add documentation/comments to *helloAgain* .

#### Brief Biography Program

Now that you've gotten a little experience programming and seen a few python programs, next you'll write a program that will do the following:

- Print your full name (first and last)
- Print your CATS/CruzMail username
- Print whether you are an undergraduate or graduate student
- Print your major
- Print why you are taking this class and what you hope to get out of it
- What problem in molecular biology most interests you.
- Print your prior computer programming experience

This program should NOT ask the user for this data. This data should be constants in your program. The constants should NOT be part of the person class -- use parameters that are given to your person object when it is instantiated. The name of the program is "myBio" and a sample execution of the program should look something like this:

My name is **David Bernick**.

My username is **dbernick**.

I am a **professor**.

My major is **Bioinformatics and Biomolecular Engineering**.

**I'm not taking this class, but I teach it.**

**I am a synthetic bioengineer and I'm interested in extreme life, especially hyperthermophilic and hypersaliphilic, with a focus on small RNA and novel proteins from extreme viruses.**

I have prior programming experience using **Python, Perl, Java, Fortran, Pascal, PL/1, ADA, Basic,TPL, APL, Assembly, and C++** (I'm not very fond of this though).

Each piece of info should occupy only a single line.

In order to get full credit, the program must use attributes of the person class for each of the **emphasized strings** above and contain appropriate documentation/comments.

Note that in Python, *names* that are part of a class are called *attributes*. Functions that are part of a class are called *methods*. In other languages, *names* are sometimes called *variables*.

#### myBio

```
In [ ]:#!/usr/bin/env python3
# Name: David Bernick(dbernick)
# Group Members: none
```

```
"""
Build a Person object and have it introduce itself.
"""
```

```
input: a string of arbitrary length, which is used to name the new person object
output: greeting printed to screen
"""
```

```
class Person:
    """ Build a new Person objects with all of hte attributes given at instantiation time. """
    def __init__(self,name, major, studentType, height):
        """ Save all of the instance arguments. """
        self.myName = name
        self.myMajor = major

    def introduce (self):
        """ Print the introductionusing all of the object attributes """
        print ("Hi there, I am {}},format(self.myName))
```

```
name = 'David Bernick'
username = 'dbernick'
studentType = 'professor'
major = ' '
madison = Person ('Madison',
                 'BME',
                 'student',
                 '5-5')
```

```
tara = Person ('Tara', 'BioC', 'student', '5-5')
madison.introduce()
tara.introduce()
```

#### Debug buggy code

Practice makes us better; usually debugging makes code closer to perfect. This program is supposed to ask the user to enter a DNA sequence and then output the AT content of the sequence— total number of As and Ts divided by the sequence length—as a fraction of the total sequence length (a number between 0 and 1).

For example, given the string:

AAATGAATGGCTAACTTTTGAA

the program should output: (make sure that you pay attention to the number of significant digits in the output!)

AT content = 0.739

Unfortunately there are a few bugs in the program. Your task is to identify and correct those bugs. This part of the assignment is to help familiarize you with the debugging process. It is possible to identify the bugs without using any of the following techniques just by looking over the program, but this will be good practice and will be essential when your code gets more complex.

To debug countAT-bugs.py, run the program, which will produce an error. Figure out what the error is and correct the code. It's likely that after you correct the first error you will notice another error. Repeat this process until you fix all the bugs and the program runs properly.

Save a corrected copy of the program called countAT-bugsSoln.py that includes:

- Documentation and comments
- A program overview that includes how many bugs you found and what lines from the original file they were found at.

#### countAT-bugs

```
In [6]:#!/usr/bin/env python3
# Name: David Bernick(dbernick)
# Group Members: none
```

```
class dnaString (str):
    def length (self):
        return (len(self))

    def getAT (self):
        num_A = self.count(A)
        num_T = self.count("T")
        return ((num_A + num_T)/ self.len() )
```

```
dna = input("Enter a dna sequence: ")
upperDNA = dna.upper()
coolString = dnaString(upperDNA)
```

```
print ("AT content = {:.1f}".format(coolString.getAT() ) )
```

Enter a dna sequence: ACGT

```
NameError                                Traceback (most recent call last)
<ipython-input-6-49107681beca> in <module>
    16 coolString = dnaString(upperDNA)
    17
--> 18 print ("AT content = {:.1f}".format(coolString.getAT() ) )
```

```
<ipython-input-6-49107681beca> in getAT(self)
     8
-->  9     def getAT (self):
    10         num_A = self.count(A)
    11         num_T = self.count("T")
    12         return ((num_A + num_T)/ self.len() )
```

```
NameError: name 'A' is not defined
```

#### Counting DNA Letters

Now that you are inspired by your ability to debug the countAT-bugs code, let's create a new program that will ask the user to enter a DNA sequence and then output the total sequence length and the number of each base {total As, Cs, Gs, Ts} found in the sequence. For example, given the string:

AAATGAATGGCTAACTTTTGAA

the program should output:

Your sequence is 23 nucleotides long with the following breakdown of bases:

number As = 10 number Cs = 2 number Gs = 4 number Ts = 7

Write a program called countNucleotides (put it below) that will take in a DNA sequence entered by a user and produce the outputs specified above. Note that "specified" means ... "required"

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

- Run properly
- Contain documentation/comments
- Include an overview about what your program is designed to do with expected inputs and outputs. Use a Markdown Cell to write your overview.

#### countNucleotides

```
In [ ]:#!/usr/bin/env python3
# Name: David Bernick(dbernick)
# Group Members: none
```

```
class dnaString (str):
    def length (self):
        return (len(self))

    def getAT (self):
        num_A = self.count(A)
        num_T = self.count("T")
        return ((num_A + num_T)/ self.len() )
```

```
def countNucleotideA (self):
    num_A = self.count('A')
    return num_A

def countNucleotideC (self):
    num_C = self.count('C')
    return num_C
```

```
dna = input("Enter a dna sequence: ")
upperDNA = dna.upper()
coolString = dnaString(upperDNA)
```

```
print ("Number As = {} number C = {}".format(
    coolString.countNucleotideA(),
    coolString.countNucleotideC() ) )
```

#### 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. The two recommended solutions (pick one) at this point are to:

- Email your jupyter notebook to yourself
- Copy your notebook to a pen drive

The class will be using canvas to submit the notebook created as part of this lab assignment.

For this lab, you should upload your notebook with the following cells filled in

- helloMe 4 points
- helloAgain 5 points
- myBio 6 points
- countAT-bugsSoln 9 points
- countNucleotides 6 points

Important: to get full credit on this lab assignment, everything needs to

- Run properly
- Contain informative documentation/comment, including your name and your partner
- be your own work

Congratulations, you finished your first lab assignment!

#### Inspection results ( Who and When )

What did your inspection team have to say about the work and who are they? Did you discuss the many coding choices that are available, even in a beginning assignment like this? What did you decide to do as a result of the inspection?