

University of Puerto Rico



Puerto Rico - IDeA Networks of Biomedical Research Excellence



Python3 Part 3 – Crash Course in Python3 for Future STEM Coders



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- The following material is the result of PR-INBRE research and curriculum development effort to provide a set of educational materials for research training and curriculum changes in biology programs across the island to support PR-INBRE efforts to establish a Community of Practice in Bioinformatics that offers a fruitful environment to increase computational and bioinformatics skills among traditional researchers and students (undergraduate and graduate) in the island. They have been developed as a part of the NIH funded project "Puerto Rico IDeA Network Biomedical Research Excellence (PRINBRE)" (Award Number 5P20GM103475).
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Objectives

- Open and Read data of different file formats using either a for loop, while loop, or a Python Library
 - TXT File
 - CSV File
- Write to different file formats
 - TXT File
 - CSV File
- Import a Python Library to your code
- Using Panda Library, create a **Pandas DataFrame** with columns and **export the DataFrame as a CSV File**





Competencies

June 5, 2023 - Record and write simple and common Python scripts to deal with Bioinformatic needs for biological data analyses using a Jupyter Notebook

June 7, 2023 — Exploit datatypes that can record objects, as well as, reason logical statements

Today — Open, read, write, manipulate, analyze files using Python





Objectives

- Continue to use Google Colab as an environment to practice and learn common Python lines of code.
- Create a new file using the function open()
- Write information to a file using the write() function
- Using for Loops analyze lines of a file

• import python libraries





Target Audience

• This training is addressed to beginners, highly motivated (eager to learn what is needed in order to be competitive without the tendency to self-limit when learning computational skills by saying that is difficult) wanting to become familiar with the **Python** programming language and become the Script Master of their bioinformatic analysis.







The importance of Python

- R is great however... eventually, you'll have to use Python language
- No matter your field, python will be a necessity (chemistry, biology, engineering, ecology, mathematics, etc)
- Knowing Python gives you an edge when job searching and navigating graduate research
- The majority bioinformatic tools are python-based and will lead you to understand under the hood code in order to successfully execute said program's purpose.





Lesson 3.0.0 Create Files





Indicate with 'w' that you want to write to this file



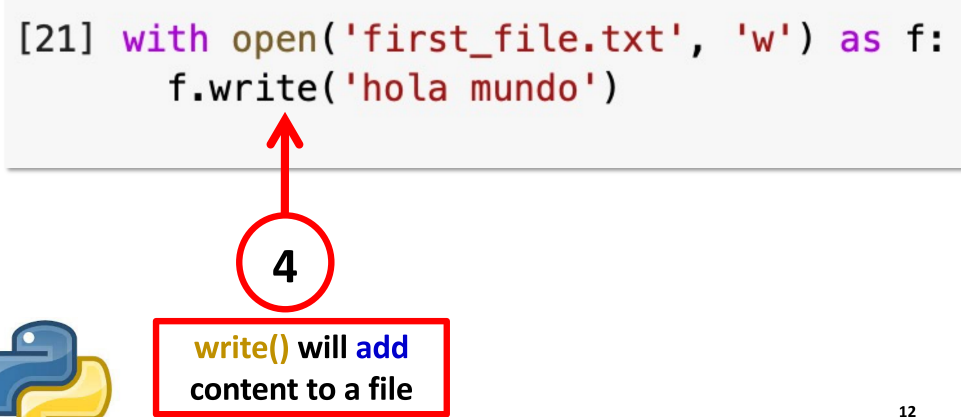
[21] with open('first_file.txt', 'w') as f:



Assign a variable to this new file

[21] with open('first_file.txt', 'w') as f:





```
[21] with open('first_file.txt', 'w') as f:
    f.write('hola mundo')
```

Use write() to write 'hola

mundo' to your new file called f

Success! Check out your first file!









Exercise 3.0.0 Create a file with gene names found in a List



input is a list

```
[28] crtam_genes_list = ['CRTAM_HUMAN', 'CRTAM_MOUSE', 'CADM1_HUMAN']
    print(crtam_genes_list)
```



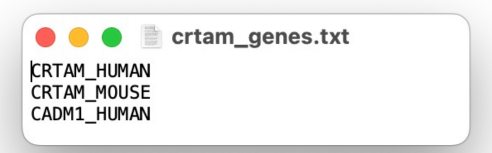
input is a list

```
[28] crtam_genes_list = ['CRTAM_HUMAN', 'CRTAM_MOUSE', 'CADM1_HUMAN']
    print(crtam_genes_list)

['CRTAM_HUMAN', 'CRTAM_MOUSE', 'CADM1_HUMAN']
```



output the following txt file





input is a list

```
[31] crtam_genes_list = ['CRTAM_HUMAN', 'CRTAM_MOUSE', 'CADM1_HUMAN']
    print(crtam_genes_list)

['CRTAM_HUMAN', 'CRTAM_MOUSE', 'CADM1_HUMAN']
```

Answer





input is a list

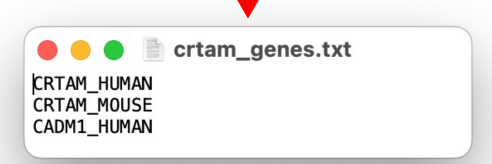
```
[31] crtam_genes_list = ['CRTAM_HUMAN', 'CRTAM_MOUSE', 'CADM1_HUMAN']
    print(crtam_genes_list)

['CRTAM_HUMAN', 'CRTAM_MOUSE', 'CADM1_HUMAN']
```

Answer

Tip: Use '\n' to indicate a new line





input is a list

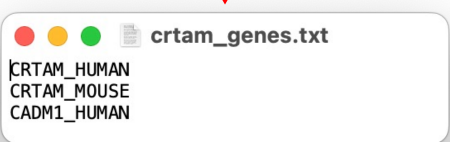
```
[31] crtam_genes_list = ['CRTAM_HUMAN', 'CRTAM_MOUSE', 'CADM1_HUMAN']
    print(crtam_genes_list)

['CRTAM_HUMAN', 'CRTAM_MOUSE', 'CADM1_HUMAN']

[32] with open('crtam_genes.txt', 'w') as f:
    for gene in crtam_genes_list:
        f.write(gene)
        f.write('\n')
```











Lesson 3.1.0 Reading in your newly created files







Lesson 3.1.1 Read in the lines of a file using a For Loop



crtam_genes_file = open('crtam_genes.txt','r')

Use open()
function



crtam_genes_file = open('crtam_genes.txt','r')

Identify your
file of
interest



crtam_genes_file = open('crtam_genes.txt','r')

Indicate that you want to read the file with 'r'

with 'r'



crtam_genes_file = open('crtam_genes.txt','r')
for crtam_genes_line in crtam_genes_file:



Use a for loop to loop through the lines in a file



```
crtam_genes_file = open('crtam_genes.txt','r')
for crtam_genes_line in crtam_genes_file:
    print(crtam_genes_line)
```

print() each lines



```
crtam_genes_file = open('crtam_genes.txt','r')
for crtam_genes_line in crtam_genes_file:
    print(crtam_genes_line)
    crtam_genes_file.close()
```





```
crtam_genes_file = open('crtam_genes.txt','r')
for crtam_genes_line in crtam_genes_file:
  print(crtam_genes_line)
crtam_genes_file.close()
CRTAM_HUMAN
CRTAM_MOUSE
CADM1_HUMAN
```



How many lines are being printed out?

```
crtam_genes_file = open('crtam_genes.txt','r')
for crtam_genes_line in crtam_genes_file:
  print(crtam_genes_line)
crtam_genes_file.close()
CRTAM_HUMAN
CRTAM_MOUSE
CADM1_HUMAN
```



How many lines are being printed out?

```
crtam_genes_file = open('crtam_genes.txt','r')
 for crtam_genes_line in crtam_genes_file:
   print(crtam_genes_line)
 crtam_genes_file.close()
OCRTAM_HUMAN
2CRTAM_MOUSE
3
4CADM1_HUMAN
```



Remove these blank lines using strip()

```
crtam_genes_file = open('crtam_genes.txt','r')
 for crtam_genes_line in crtam_genes_file:
   print(crtam_genes_line)
 crtam_genes_file.close()
OCRTAM_HUMAN
2CRTAM_MOUSE
3
4CADM1_HUMAN
```



Remove these blank lines using strip()







Lesson 3.1.2 Read in the lines of a file using a While Loop



Read in your file using a while loop

crtam_genes_file = open('crtam_genes.txt','r')

Use open()
function



Read in your file using a while loop

crtam_genes_file = open('crtam_genes.txt','r')
crtam_genes_lines = crtam_genes_file.readline()

Using readline() function will allow you to while loop the lines of a file



crtam_genes_file = open('crtam_genes.txt','r')
crtam_genes_lines = crtam_genes_file.readline()
while crtam_genes_lines:

Perform while loop through lines of this file



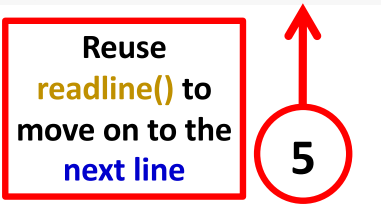


```
crtam_genes_file = open('crtam_genes.txt','r')
crtam_genes_lines = crtam_genes_file.readline()
while crtam_genes_lines:
    print(crtam_genes_lines.strip('\n'))
```

Strip and print each line



crtam_genes_file = open('crtam_genes.txt','r')
crtam_genes_lines = crtam_genes_file.readline()
while crtam_genes_lines:
 print(crtam_genes_lines.strip('\n'))
 crtam_genes_lines = crtam_genes_file.readline()





```
crtam_genes_file = open('crtam_genes.txt','r')
crtam_genes_lines = crtam_genes_file.readline()
while crtam_genes_lines:
    print(crtam_genes_lines.strip('\n'))
    crtam_genes_lines = crtam_genes_file.readline()
crtam_genes_file.close()
```





```
crtam_genes_file = open('crtam_genes.txt','r')
crtam_genes_lines = crtam_genes_file.readline()
while crtam_genes_lines:
    print(crtam_genes_lines.strip('\n'))
    crtam_genes_lines = crtam_genes_file.readline()
    crtam_genes_file.close()
OCRTAM_HUMAN
1CRTAM_MOUSE
```



2CADM1_HUMAN





Lesson 3.2.0 Manipulate files the Panda way





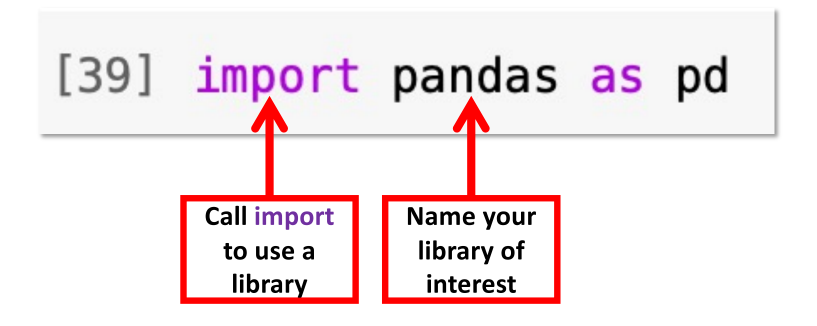


Lesson 3.2.1 Import the Panda Library

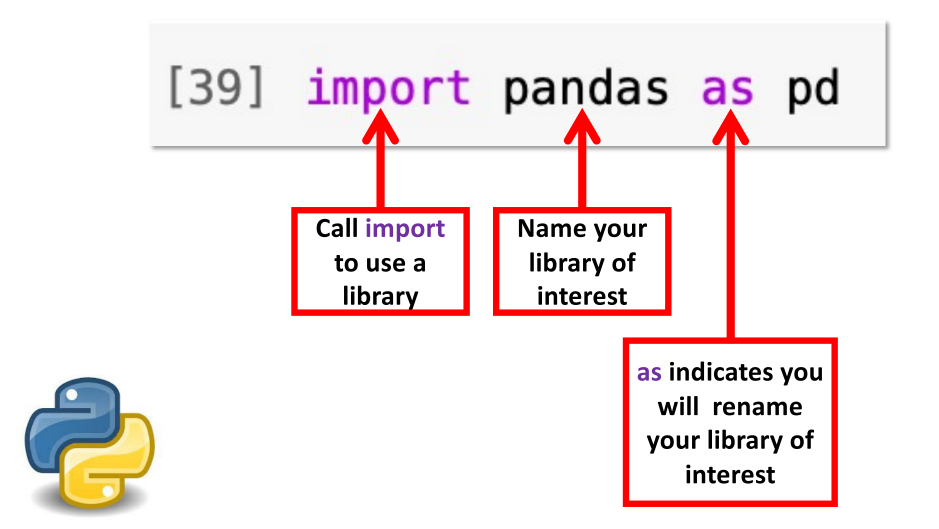


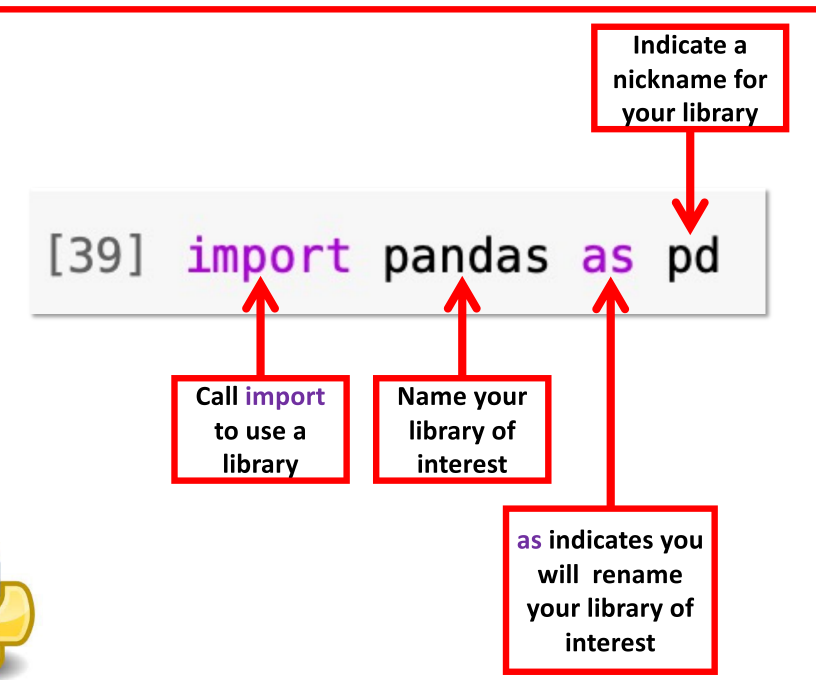
















Lesson 3.2.2 Read in a CSV file using Panda Library







Lesson 3.2.2a Reading in a CSV file with a header using Panda Library





gene, length CRTAM_HUMAN, 393 CRTAM_MOUSE, 781 CADM1_HUMAN, 442



```
crtam_with_header.csv

gene, length
CRTAM_HUMAN, 393
CRTAM_MOUSE, 781
CADM1_HUMAN, 442
```

```
[66] pd.read_csv('crtam_with_header.csv',header=0,delimiter=',')
```



```
crtam_with_header.csv

gene, length
CRTAM_HUMAN, 393
CRTAM_MOUSE, 781
CADM1_HUMAN, 442
```

```
[66] pd.read_csv('crtam_with_header.csv',header=0,delimiter=',')

Call pandas
library by its
nickname
```



```
crtam_with_header.csv

gene, length
CRTAM_HUMAN, 393
CRTAM_MOUSE, 781
CADM1_HUMAN, 442
```



```
gene, length
CRTAM_HUMAN, 393
CRTAM_MOUSE, 781
CADM1_HUMAN, 442
```

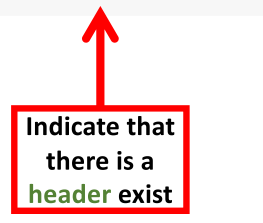
```
[66] pd.read_csv('crtam_with_header.csv',header=0,delimiter=',')
```

Name your file of interest



```
gene, length
CRTAM_HUMAN, 393
CRTAM_MOUSE, 781
CADM1_HUMAN, 442
```

```
[66] pd.read_csv('crtam_with_header.csv',header=0,delimiter=',')
```



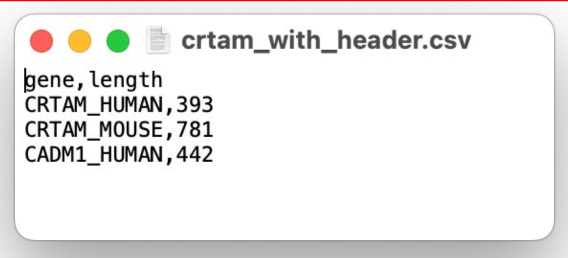


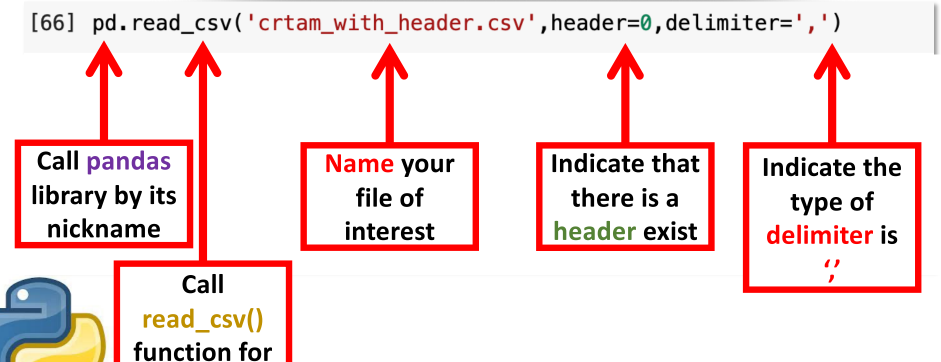
```
gene, length
CRTAM_HUMAN, 893
CRTAM_MOUSE, 781
CADM1_HUMAN, 142
```

```
[66] pd.read_csv('crtam_with_header.csv',header=0,delimiter=',')
```

Indicate the type of delimiter is







a csv file

```
crtam_with_header.csv

gene, length
CRTAM_HUMAN, 393
CRTAM_MOUSE, 781
CADM1_HUMAN, 442
```

```
gene length header

O CRTAM_HUMAN 393

1 CRTAM_MOUSE 781

2 CADM1_HUMAN 442
```

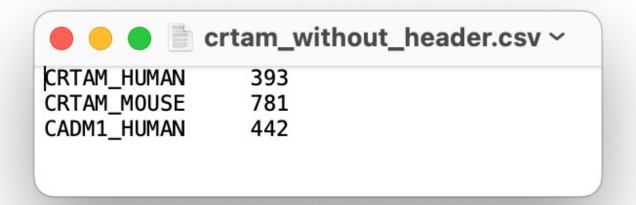






Lesson 3.2.2b Reading in a CSV file with no header using Panda Library



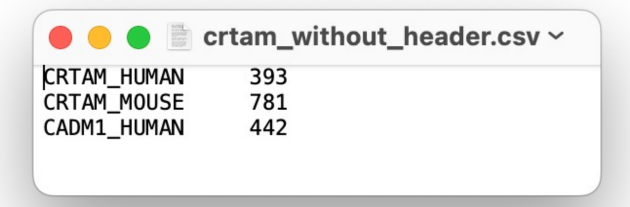


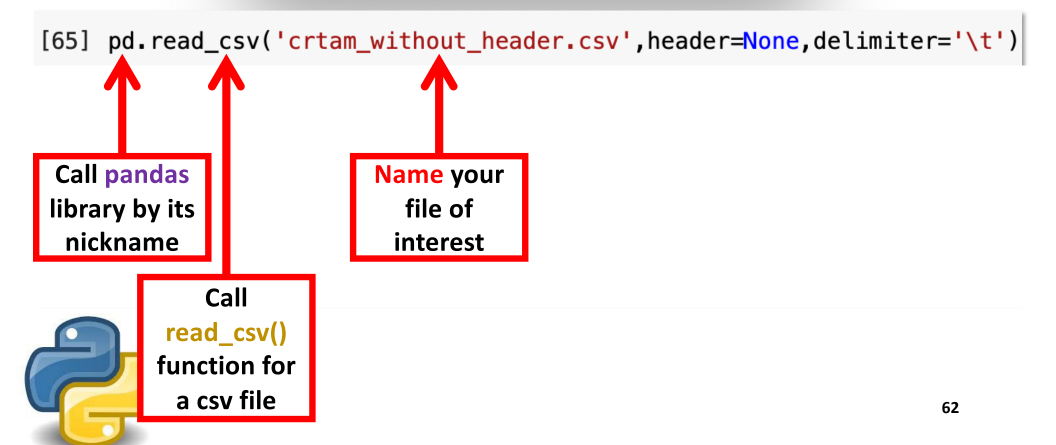


```
CRTAM_HUMAN 393
CRTAM_MOUSE 781
CADM1_HUMAN 442
```

```
[65] pd.read_csv('crtam_without_header.csv',header=None,delimiter='\t')
```







```
CRTAM_HUMAN 393
CRTAM_MOUSE 781
CADM1_HUMAN 442
```

```
[65] pd.read_csv('crtam_without_header.csv',header=None,delimiter='\t')
```

Indicate that there is no header exist

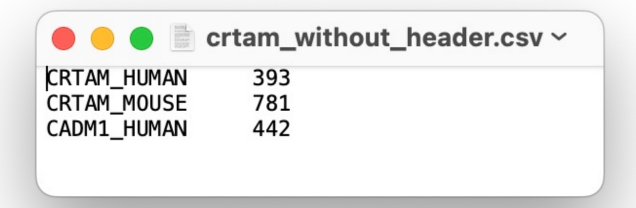


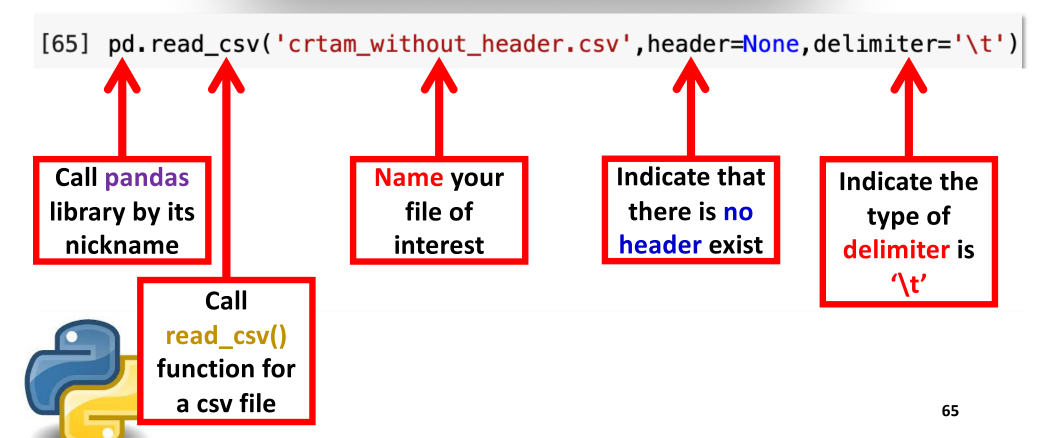
```
CRTAM_HUMAN 393
CRTAM_MOUSE 781
CADM1_HUMAN 442
```

```
[65] pd.read_csv('crtam_without_header.csv',header=None,delimiter='\t')
```

```
Indicate the type of delimiter is '\t'
```







```
CRTAM_HUMAN 393
CRTAM_MOUSE 781
CADM1_HUMAN 442
```

```
[65] pd.read_csv('crtam_without_header.csv',header=None,delimiter='\t')
```

```
0 1 header
```

- O CRTAM_HUMAN 393
- 1 CRTAM_MOUSE 781
- 2 CADM1_HUMAN 442







Lesson 3.2.3 Create a DataFrame and export a new CSV file using Panda Library



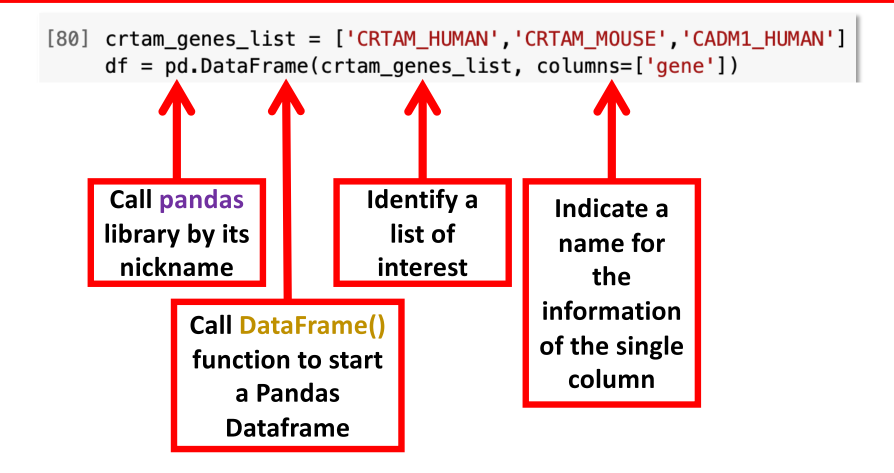


Lesson 3.2.3a Create a Panda DataFrame from a single List



[80] crtam_genes_list = ['CRTAM_HUMAN','CRTAM_MOUSE','CADM1_HUMAN']









```
[80] crtam_genes_list = ['CRTAM_HUMAN','CRTAM_MOUSE','CADM1_HUMAN']
    df = pd.DataFrame(crtam_genes_list, columns=['gene'])
    df
                gene
        CRTAM_HUMAN
        CRTAM_MOUSE
        CADM1_HUMAN
   index
```







Lesson 3.2.3b Create a Panda DataFrame from multiple Lists



```
[81] crtam_human_list = ['CRTAM_HUMAN',393,'human']
    crtam_mouse_list = ['CRTAM_MOUSE',781,'mouse']
    cadm1_human_list = ['CADM1_HUMAN',442,'human']
```



Call pandas library by its nickname

Call DataFrame()
function to start
a Pandas
Dataframe





List of Lists





```
crtam human list = ['CRTAM HUMAN',393,'human']
crtam_mouse_list = ['CRTAM_MOUSE',781,'mouse']
cadm1 human list = ['CADM1 HUMAN',442,'human']
df = pd.DataFrame([crtam_human_list,crtam_mouse_list,cadm1_human_list],
                  columns=['gene','length','organisms'])
df
            gene length organisms
   CRTAM_HUMAN
                     393
                             human
   CRTAM MOUSE
                     781
                              mouse
   CADM1 HUMAN
                     442
                              human
```





```
crtam human list = ['CRTAM HUMAN',393,'human']
crtam_mouse_list = ['CRTAM_MOUSE',781,'mouse']
cadm1 human list = ['CADM1 HUMAN',442,'human']
df = pd.DataFrame([crtam_human_list,crtam_mouse_list,cadm1_human_list],
                  columns=['gene','length','organisms'])
df
            gene length organisms
                                             column names
   CRTAM HUMAN
                     393
                             human
   CRTAM_MOUSE
                     781
                             mouse
   CADM1 HUMAN
                     442
                             human
```



```
crtam human list = ['CRTAM HUMAN',393,'human']
crtam_mouse_list = ['CRTAM_MOUSE',781,'mouse']
cadm1 human list = ['CADM1 HUMAN',442,'human']
df = pd.DataFrame([crtam_human_list,crtam_mouse_list,cadm1_human_list],
                  columns=['gene','length','organisms'])
df
            gene length organisms
                                             crtam_human_list
   CRTAM_HUMAN
                     393
                             human
   CRTAM_MOUSE
                     781
                             mouse
   CADM1 HUMAN
                     442
                             human
```



```
crtam human list = ['CRTAM HUMAN',393,'human']
crtam_mouse_list = ['CRTAM_MOUSE',781,'mouse']
cadm1 human list = ['CADM1 HUMAN',442,'human']
df = pd.DataFrame([crtam_human_list,crtam_mouse_list,cadm1_human_list],
                  columns=['gene','length','organisms'])
df
            gene length organisms
   CRTAM HUMAN
                     393
                             human
                                             crtam_mouse_list
   CRTAM_MOUSE
                     781
                             mouse
   CADM1 HUMAN
                     442
                             human
```



```
crtam human list = ['CRTAM HUMAN',393,'human']
crtam_mouse_list = ['CRTAM_MOUSE',781,'mouse']
cadm1 human list = ['CADM1 HUMAN',442,'human']
df = pd.DataFrame([crtam_human_list,crtam_mouse_list,cadm1_human_list],
                  columns=['gene','length','organisms'])
df
            gene length organisms
   CRTAM HUMAN
                     393
                             human
   CRTAM_MOUSE
                     781
                             mouse
                                             cadm1_human_list
   CADM1_HUMAN
                     442
                             human
```







Lesson 3.2.3c Create a Pandas DataFrame from a Nested Dictionary

























Lesson 3.2.4 Export a Panda DataFrame as a CSV File



	gene	length	organism
0	CRTAM_HUMAN	393	human
1	CRTAM_MOUSE	781	mouse
2	CADM1_HUMAN	442	human



crtam_information.csv





```
,gene,length,organism
0,CRTAM_HUMAN,393,human
1,CRTAM_MOUSE,781,mouse
2,CADM1_HUMAN,442,human
```

	gene	length	organism
0	CRTAM_HUMAN	393	human
1	CRTAM_MOUSE	781	mouse
2	CADM1_HUMAN	442	human

[90] df.to_csv('crtam_information.csv', sep=',')

Call variable of your dataframe



	gene	rength	organism
0	CRTAM_HUMAN	393	human
1	CRTAM_MOUSE	781	mouse
2	CADM1_HUMAN	442	human





Call to_csv()
function to start
a Pandas
Dataframe

	gene	length	organism
0	CRTAM_HUMAN	393	human
1	CRTAM_MOUSE	781	mouse
2	CADM1_HUMAN	442	human

[90] df.to_csv('crtam_information.csv', sep=',')

Name your file of interest



	gene	length	organism
0	CRTAM_HUMAN	393	human
1	CRTAM_MOUSE	781	mouse
2	CADM1_HUMAN	442	human



[90] df.to_csv('crtam_information.csv', sep=',')

Indicate the type of separator or delimiter is



			gene	length	organism	n 🎢
	0 (CRTAM_H	HUMAN	393	human	n
	1 (CRTAM_I	MOUSE	781	mouse	
	2 (CADM1_l	HUMAN	442	human	ı
90] d			'crtam_	informat	ion.csv'	, sep=',')
of y data	your fram			Name yo		Indicate the type of
		Call to_cs	()v	interes	st	separator or
	(, v ()			delimiterie
		nction to a Panda	start			delimiter is

Check out your new CSV file!



,gene,length,organism
0,CRTAM_HUMAN,393,human
1,CRTAM_MOUSE,781,mouse
2,CADM1_HUMAN,442,human









Proficiency assessment Show off your new skills!

- 1. Read in the Fasta File called crtam.faa
- 2. Choose at least 3 of the following options below to create a dataframe with your columns of interest.
 - Gene_names
 - Length in amino acid
 - Length in it's DNA sequence
 - Total RSNNEETS
 - Total KAH
 - Total PGLRLRLLL
- 3. Export a file to import into Excel

