Lab 2: Importing and plotting data

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Table of Contents

- 1. Review of Numpy arrays
- 2. Importing data from a file into a Numpy array
- 3. Examining and plotting data in a Numpy array
- 4. Bonus exercise

Helpful Resources

- <u>Python Data Science Handbook (http://shop.oreilly.com/product/0636920034919.do)</u> by Jake VanderPlas
- <u>Python Basics Cheat Sheet (https://datacamp-community-prod.s3.amazonaws.com/e30fbcd9-f595-4a9f-803d-05ca5bf84612)</u> by Python for Data Science
- <u>Jupyter Notebook Cheat Sheet (https://datacamp-community-prod.s3.amazonaws.com/48093c40-5303-45f4-bbf9-0c96c0133c40)</u> by Python for Data Science
- <u>Matplotlib Cheat Sheet (https://datacamp-community-prod.s3.amazonaws.com/28b8210c-60cc-4f13-b0b4-5b4f2ad4790b)</u> by Python for Data Science
- Numpy Cheat Sheet (https://datacamp-community-prod.s3.amazonaws.com/e9f83f72a81b-42c7-af44-4e35b48b20b7) by Python for Data Science

Data

The data in this lab was downloaded from <u>Kaggle (https://www.kaggle.com/uciml/iris)</u> (originally from <u>Fisher 1936 (http://rcs.chemometrics.ru/Tutorials/classification/Fisher.pdf)</u>) and was edited for teaching purposes.

Lab 2 Part 1: Review of Numpy arrays

In lecture this week we used Numpy arrays to generate random numbers, look at data, and make patterns. In this first lab section we'll review how to create, access, and edit parts of a Numpy array.

To use the Numpy library we need to first import it using the command <code>import numpy</code> as <code>np</code>. We'll also import Matplotlib in this same code block, since we'll use this library later in the lab. It's good practice to import all of your libraries at the very beginning of your code file, so that anyone can quickly see what external libraries are necessary to run your code.

```
In [1]: import numpy as np
   import matplotlib.pyplot as plt

# Magic command to turn on in-line plotting
# (show plots within the Jupyter Notebook)
%matplotlib inline
```

Creating a Numpy array from existing data

To review some important concepts about Numpy arrays, let's make a small 3x3 array called alphabet data, filled with different letters of the alphabet:

```
In [2]: row_A = ["A", "B", "C"]
    row_D = ["D", "E", "F"]
    row_G = ["G", "H", "I"]

    alphabet_data = np.array([row_A, row_D, row_G])
    print(alphabet_data)

[['A' 'B' 'C']
    ['D' 'E' 'F']
    ['G' 'H' 'I']]
```

We can use the print command to look at the entire alphabet_data Numpy array. But often we'll work with very large arrays full of data, and we'll want to pick small subsets of the data to look at. Therefore, it's useful to know how to ask Python to give you just a section of any Numpy array.

Selecting subsets of Numpy arrays

In lab 1, we talked about how index values describe where to find a specific item within a Python list or array. For example, the variable <code>example_list</code> is a list with one row, containing three items. To print the first item in the list we would print <code>example_list[0]</code>, or the value in the variable <code>example_list</code> at index 0. Remember that the first item in a Python list corresponds to the index 0.

```
In [3]: example_list = ["avocado", "tomato", "onion"]
    print("example_list is:", example_list)
    print("example_list[0] is:", example_list[0])

example_list is: ['avocado', 'tomato', 'onion']
    example_list[0] is: avocado
```

Selecting a single value in a Numpy array

alphabet_data is a little more complicated since it has rows and columns, but the general principle of indexing is still the same. Each value in a Numpy array has a unique index value for its row location, and a separate unique index value for its column location. We can ask Numpy to give us just the value we want by using the syntax alphabet_data[row index, column index].

Exercise 1: Use indexing to print the second item in the first row of alphabet data:

```
In [4]: print(alphabet_data[0, 1])
```

В

Selecting a range of values in a Numpy array

In addition to selecting just one value, we can use the syntax lower index range: upper index range to select a range of values. Remember that ranges in Python are *exclusive* - the last index in the range is not included. Below is an example of range indexing syntax used on example list:

```
In [5]: example_list = ["avocado", "tomato", "onion"]
    print("example_list is:", example_list)
    print("example_list[0:2] is:", example_list[0:2])

example_list is: ['avocado', 'tomato', 'onion']
    example_list[0:2] is: ['avocado', 'tomato']
```

We can use exactly the same notation in a Numpy array. However, since we have both row and column indices, we can declare one range for the rows and one range for the columns. For example, the following code prints all rows from index 0 to index 3, and all columns from index 0 to index 2. Note that index 3 doesn't actually exist - but since the upper index range is not included in a Python range, we need to use an index of 3 to print everything up to index 2.

Exercise 2: Print the first two rows of the first two columns in alphabet data.

Exercise 3: Print the last two rows of the last two columns in alphabet_data.

```
In [8]: print(alphabet_data[1:, 1:])
        [['E' 'F']
        ['H' 'I']]
```

Once we know how to select subsets of arrays, we can use this knowledge to *change* the items in these selections. For example, in a list we can assign a value found at a specific index to be something else. In this example we use indexing to reference the first item in <code>example_list</code>, and then change it.

```
In [9]: example_list = ["avocado", "tomato", "onion"]
    print("before assignment, the example_list is:", example_list)
    example_list[0] = "banana"
    print("after assignment, the example_list is:", example_list)
```

before assignment, the example_list is: ['avocado', 'tomato', 'onion'] after assignment, the example_list is: ['banana', 'tomato', 'onion']

Similarly, we can change items in a Numpy array using indexing:

```
In [10]: print("before assignment, alphabet_data is:")
    print(alphabet_data)

    alphabet_data[0, 0] = "Z"
    print("after assignment, alphabet_data is:")
    print(alphabet_data)

before assignment, alphabet_data is:
    [['A' 'B' 'C']
    ['D' 'E' 'F']
    ['G' 'H' 'I']]
    after assignment, alphabet_data is:
    [['Z' 'B' 'C']
    ['D' 'E' 'F']
    ['G' 'H' 'I']]
```

Exercise 4: Replace the item in the third row and second column of $alphabet_data$ with "V".

```
In [11]: alphabet_data[2, 1] = "V"
    print(alphabet_data)

[['Z' 'B' 'C']
        ['D' 'E' 'F']
        ['G' 'V' 'I']]
```

Exercise 5: Replace the entire second row of alphabet_data with a new row: ["X", "Y", "X"]

```
In [12]: alphabet_data[1] = ["X", "Y", "X"]
    print(alphabet_data)

[['Z' 'B' 'C']
       ['X' 'Y' 'X']
       ['G' 'V' 'I']]
```

Lab 2 Part 2: Importing data from a file into a Numpy array

Let's apply these principles of Numpy arrays to some real biological data. In the Lab_02 data folder there are three data files:

```
• ./data/Lab 02/Iris setosa data.csv
```

- ./data/Lab 02/Iris versicolor data.csv
- ./data/Lab_02/Iris_virginica_data.csv

These files contain data collected by botanist Edgar Anderson in the Gaspe Peninsula for three species of irises: *Iris setosa*, *Iris versicolor*, and *Iris virginica*. Anderson was interested in investigating the morphologic variation in these three related species. He collected 50 individual flowers for each species, and recorded four different measurements for each flower: the length and width of the sepals and petals.

The data is formatted as a large table, with one file for each species. The files contain 50 rows, each representing one individual, and four columns, which represent sepal length, sepal width, petal length, and petal width. For example, <code>Iris_setosa_data.csv</code> corresponds to the column and row labels shown below:

Sample ID	Sepal length (cm)	Sepal width (cm)	Petal length (cm)	Petal width (cm)
Individual 1	5.1	3.5	1.4	0.2
Individual 2	4.9	3	1.4	0.2
Individual 3	4.7	3.2	1.3	0.2
Individual 50	5	3.3	1.4	0.2

We'll use the Numpy command loadtxt to read in our first file, Iris_setosa_data.csv. We will save this data in a Numpy array called iris_data.

```
In [13]: # Load our file data from "filename" into a variable called iris_data
    filename = "./data/Lab_02/Iris_setosa_data.csv"
    iris_data = np.loadtxt(fname=filename, delimiter=",")
```

The data description above tells us that <code>iris_data</code> should contain 50 rows and 4 columns, so let's use the Numpy <code>shape</code> command to double check that's the case. Numpy <code>shape</code> will print two numbers in the format (number of rows, number of columns).

Exercise 6: Right now, the code below prints a warning if we don't have the expected 50 rows. Edit the code so that the warning is also printed if the number of columns is not 4.

```
In [14]: # Print the shape of the loaded dataset
    data_shape = iris_data.shape
    print("Iris data shape is:", data_shape)

# Print a warning if the data shape is not what we expect
    if (data_shape[0] != 50) or (data_shape[1] != 4):
        print("Unexpected data shape!")
    else:
        print("Correct data shape of 50 rows, 4 columns!")
```

```
Iris data shape is: (50, 4)
Correct data shape of 50 rows, 4 columns!
```

It looks like our <code>iris_data</code> Numpy array is the shape we expect. Now let's look at a subset of data to see what kind of data we're working with.

Exercise 7: Use Python array indexing to print the first three rows, first four columns of iris_data. Check to make sure that the printed data matches what is given to you in the data description above.

```
In [15]: print(iris_data[0:3, 0:5])

[[ 5.1     3.5     1.4     0.2]
       [ 4.9     3.     1.4     0.2]
       [ 4.7     3.2     1.3     0.2]]
```

Lab 2 Part 3: Examining and plotting data in a Numpy array

Calculate interesting characteristics of a Numpy array

Now that we have loaded our Iris data into a Numpy array, there are several interesting commands we can use to find out more about our data. First let's look at the sepal length column for *Iris setosa* (the first column in the dataset). Using array indexing, we will put this entire first column into a new variable called <code>sepal_lengths</code>. When indexing between a range of values, leaving the upper range bound blank causes Python to include everything until the end of the array:

```
In [16]: # put the sepal lengths for this dataset in a variable called sepal length
         sepal lengths = iris data[0:, 0]
        print("The sepal lengths in this dataset is:")
        print(sepal lengths)
        The sepal lengths in this dataset is:
                                                4.4
               4.9 4.7 4.6
                             5.
                                  5.4 4.6 5.
                                                     4.9 5.4
                                                               4.8
                                                                    4.8
                                                                        4.3
        [ 5.1
        5.8
          5.7
               5.4 5.1 5.7 5.1 5.4 5.1 4.6 5.1
                                                     4.8
                                                          5.
                                                               5.
                                                                    5.2
                                                                        5.2
        4.7
               5.4 5.2 5.5 4.9 5. 5.5 4.9 4.4 5.1 5.
          4.8
                                                               4.5
                                                                        5.
        5.1
          4.8
               5.1 4.6 5.3 5. ]
```

Numpy contains many useful functions for finding out different characteristics of a dataset. The code below shows some examples:

```
In [17]: # Print some interesting characteristics of the data
    print("Mean:", np.mean(sepal_lengths))
    print("Standard deviation:", np.std(sepal_lengths))
    print("Median:", np.median(sepal_lengths))
    print("Minimum:", np.min(sepal_lengths))
    print("Maximum:", np.max(sepal_lengths))
Mean: 5.006
```

Standard deviation: 0.348946987378
Median: 5.0
Minimum: 4.3
Maximum: 5.8

We can use our sepal_lengths variable and the useful characteristics we found above to make a histogram of our data. In the below code we've created a histogram, and added a line that shows where the mean of the dataset is.

Exercise 8: Edit the code block below to plot the maximum and minimum data values as two additional vertical lines.

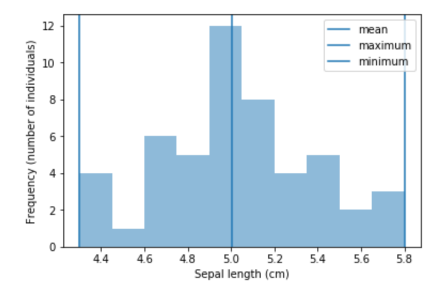
```
In [18]: # Create a histogram with an opacity of 50% (alpha=0.5)
plt.hist(sepal_lengths, alpha=0.5)

# Add a vertical line to the plot showing the mean.
plt.axvline(np.mean(sepal_lengths), label="mean")
# Your code here!
plt.axvline(np.max(sepal_lengths), label="maximum")
plt.axvline(np.min(sepal_lengths), label="minimum")

# Don't forget to label the axes!
plt.xlabel("Sepal length (cm)")
plt.ylabel("Frequency (number of individuals)")

# Add a legend to the plot
plt.legend()

# Show the plot in our jupyter notebook
plt.show()
```



Review of for loops using indexing

Last week in lab we went over an example of a for loop that uses indices to loop through a list. Let's pretend that in this *Iris setosa* dataset, we have marked in our lab notebook that the first, 12th, 26th, and 44th irises we sampled seemed suspiciously small. Let's use a for loop to print out the sepal length of each of these irises.

Exercise 9: Instead of using a for loop to look at just the indices in

interesting_indices, use a for loop to look at *all* indices in the sepal_lengths dataset. Remember that you can use the command len(sepal_lengths) to find out how many values are in the data. Print the sepal length and index if the sepal length is larger than the mean sepal length.

```
all indices = np.arange(0, len(iris data))
In [20]:
         for index in all indices:
             sepal = sepal lengths[index]
             if sepal > sepal lengths.mean():
                 print("The sepal length at index", index, ":", sepal)
         The sepal length at index 0:5.1
         The sepal length at index 5: 5.4
         The sepal length at index 10: 5.4
         The sepal length at index 14: 5.8
         The sepal length at index 15: 5.7
         The sepal length at index 16: 5.4
         The sepal length at index 17: 5.1
         The sepal length at index 18: 5.7
         The sepal length at index 19: 5.1
         The sepal length at index 20: 5.4
         The sepal length at index 21: 5.1
         The sepal length at index 23: 5.1
         The sepal length at index 27: 5.2
         The sepal length at index 28: 5.2
         The sepal length at index 31: 5.4
         The sepal length at index 32: 5.2
         The sepal length at index 33: 5.5
         The sepal length at index 36: 5.5
         The sepal length at index 39 : 5.1
         The sepal length at index 44: 5.1
         The sepal length at index 46: 5.1
         The sepal length at index 48: 5.3
```

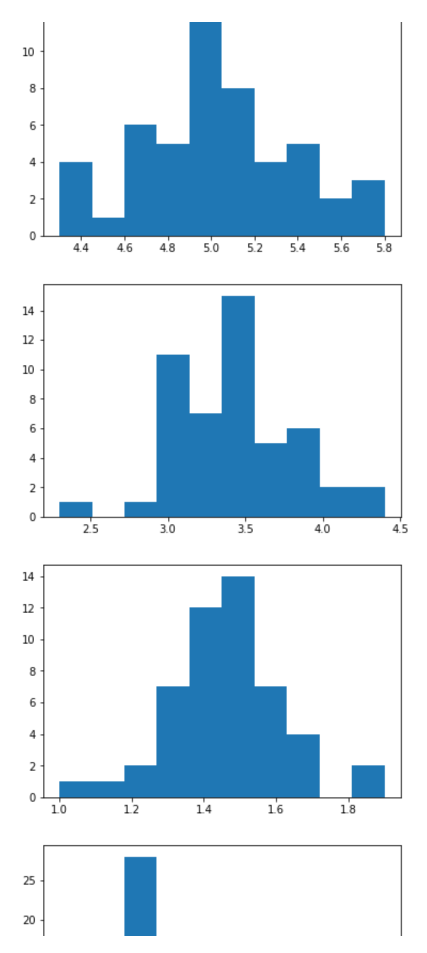
So far we've only looked at the sepal lengths in this dataset. Let's use a for loop to also look at the sepal widths, petal lengths, and petal widths. Remember that the columns in this dataset stand for:

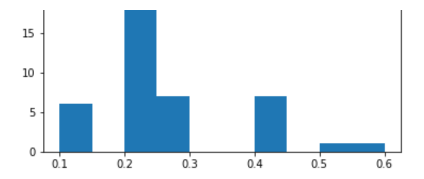
```
In [21]: sepal_lengths = iris_data[0:, 0]
    sepal_widths = iris_data[0:, 1]
    petal_lengths = iris_data[0:, 2]
    petal_widths = iris_data[0:, 3]

morphologies = [sepal_lengths, sepal_widths, petal_lengths, petal_widths

for morphology in morphologies:
    # Create a histogram
    plt.hist(morphology)
    # Show the plot in our jupyter notebook
    plt.show()
```

12 -





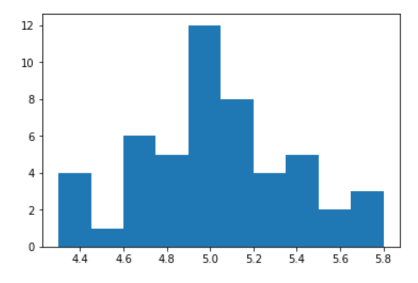
Notice that the code in the above box is doing the same action for every column in the array. So instead of re-assigning every column in the array to a new variable called <code>sepal_lengths</code>, <code>sepal_widths</code>, etc, let's use array indexing to loop through the data instead. Notice that the only thing changing when looking at different columns is the *column index*.

Exercise 10: Change the following code so that it creates a histogram for all columns in *Iris* setosa, like in the previous block. However, instead of making a new variable for each column called sepal lengths, sepal widths, etc, use indexing instead.

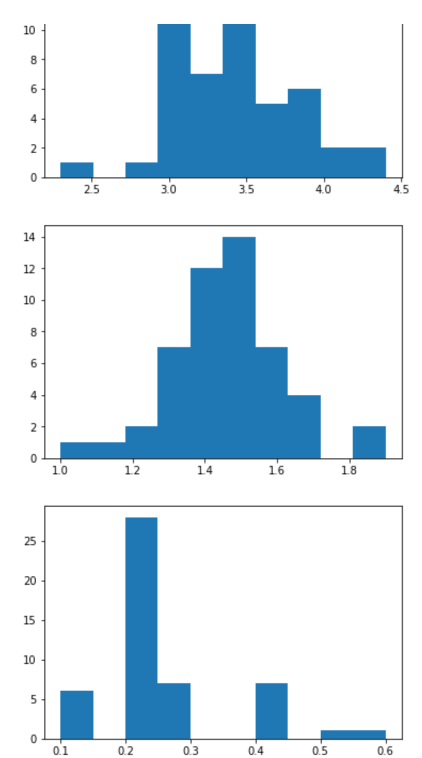
```
In [22]:
```

```
column_indices = [0, 1, 2, 3]

for index in column_indices:
    data_subset = iris_data[0:, index]
    # Create a histogram
    plt.hist(data_subset)
    # Show the plot in our jupyter notebook
    plt.show()
```







Putting it all together: Using a for loop to load, analyze, and plot multiple data files

We've now found some interesting things about *Iris setosa*. But our original dataset included three different species - *Iris setosa*, *Iris versicolor*, and *Iris virginica*. We probably want to run these exact same analyses for each species, and this is a great opportunity to use a for loop to make our lives easier. Because all three of our datasets are exactly the same shape and format, we can reuse all of our code that we've already written.

Now that we have a list of filenames to analyze, we can turn this into a for loop that loads each file and then runs analyses on the file. The code block below has started the process - for each filename, we load in the file data as a variable called <code>iris_data</code>. Note that we're not actually doing anything with the data yet, so we don't see many interesting things being printed.

```
In [24]: for filename in filenames:
    # Load our file data from "filename" into a variable called iris_dat
    iris_data = np.loadtxt(fname=filename, delimiter=",")
    print("NOW ANALYZING DATASET: ", filename)

NOW ANALYZING DATASET: ./data/Lab_02/Iris_setosa_data.csv
NOW ANALYZING DATASET: ./data/Lab_02/Iris_versicolor_data.csv
NOW ANALYZING DATASET: ./data/Lab_02/Iris_virginica_data.csv
```

The data loading doesn't seem to have caused any errors, so we'll continue to copy and paste the code we've already written to work with the data. Note that everything we've copied and pasted is code we've already written - but now we're asking Python to run this same code on *all* the data files, instead of just *Iris setosa*. For the purposes of this exercise, we'll analyze just the sepal lengths of the dataset, so that we end up with a manageable number of output plots.

```
In [25]: for filename in filenames:
    # Load our file data from "filename" into a variable called iris_dat
    iris_data = np.loadtxt(fname=filename, delimiter=",")
    print("----")
    print("NOW ANALYZING DATASET: ", filename)

# Print the shape of the loaded dataset
    data_shape = iris_data.shape
    print("Iris_data_shape is:", data_shape)
```

```
# Print a warning if the data shape is not what we expect
    if (data shape[0] != 50) or (data shape[1] != 4):
       print("Unexpected data shape!")
    else:
       print("Correct data shape of 50 rows, 4 columns!")
    # put the sepal lengths for this dataset in a variable called sepal
    sepal lengths = iris data[0:, 0]
    print("The sepal lengths in this dataset is:")
    print(sepal lengths)
NOW ANALYZING DATASET: ./data/Lab_02/Iris_setosa_data.csv
Iris data shape is: (50, 4)
Correct data shape of 50 rows, 4 columns!
The sepal lengths in this dataset is:
[ 5.1
     4.9 4.7 4.6
                    5.
                          5.4 4.6 5.
                                        4.4 4.9 5.4
                                                       4.8
                                                            4.8
                                                                 4.3
5.8
  5.7
      5.4 5.1
               5.7
                    5.1
                          5.4 5.1 4.6 5.1
                                             4.8
                                                                 5.2
4.7
      5.4 5.2 5.5 4.9 5.
  4.8
                              5.5 4.9 4.4
                                             5.1
                                                  5.
                                                       4.5
                                                            4.4
                                                                5.
5.1
  4.8 5.1 4.6 5.3 5. ]
NOW ANALYZING DATASET: ./data/Lab 02/Iris versicolor data.csv
Iris data shape is: (50, 4)
Correct data shape of 50 rows, 4 columns!
The sepal lengths in this dataset is:
<sup>7</sup>.
      6.4 6.9 5.5
                     6.5
                         5.7 6.3 4.9 6.6
                                             5.2 5.
                                                                 6.1
                                                       5.9 6.
5.6
  6.7
      5.6
          5.8
               6.2 5.6
                         5.9 6.1 6.3 6.1
                                             6.4
                                                  6.6
                                                       6.8
                                                            6.7
                                                                 6.
5.7
                              6.7 6.3 5.6
  5.5
      5.5 5.8
                6.
                     5.4 6.
                                             5.5 5.5 6.1
                                                            5.8
5.6
 5.7
      5.7 6.2 5.1 5.71
NOW ANALYZING DATASET: ./data/Lab 02/Iris virginica data.csv
Iris data shape is: (50, 4)
Correct data shape of 50 rows, 4 columns!
The sepal lengths in this dataset is:
      5.8 7.1 6.3 6.5
                         7.6 4.9 7.3 6.7 7.2 6.5 6.4
[ 6.3
                                                            6.8
                                                                 5.7
5.8
               7.7
  6.4
      6.5 7.7
                     6.
                          6.9 5.6 7.7 6.3
                                             6.7 7.2
                                                       6.2
                                                            6.1
                                                                 6.4
7.2
      7.9 6.4 6.3 6.1 7.7 6.3 6.4 6.
  7.4
                                             6.9 6.7 6.9
                                                            5.8
                                                                6.8
6.7
```

6.7 6.3 6.5 6.2 5.91

Exercise 11: Similarly, add in the code you've already written to print the interesting characteristics of the data (mean, median, max, etc.) and create a histogram for each data file that includes the mean and median. Run your final for loop. Which iris species has the longest mean sepal length? Smallest minimum sepal length?

```
In [26]: for filename in filenames:
             # Load our file data from "filename" into a variable called iris_dat
             iris data = np.loadtxt(fname=filename, delimiter=",")
             print("---")
             print("NOW ANALYZING DATASET: ", filename)
             # Print the shape of the loaded dataset
             data shape = iris data.shape
             print("Iris data shape is:", data shape)
             # Print a warning if the data shape is not what we expect
             if (data shape[0] != 50) or (data shape[1] != 4):
                 print("Unexpected data shape!")
             else:
                 print("Correct data shape of 50 rows, 4 columns!")
             # put the sepal lengths for this dataset in a variable called sepal
             sepal lengths = iris data[0:, 0]
             print("The sepal lengths in this dataset is:")
             print(sepal lengths)
             # Print some interesting characteristics of the data
             print("Mean:", np.mean(sepal lengths))
             print("Standard deviation:", np.std(sepal lengths))
             print("Median:", np.median(sepal lengths))
             print("Minimum:", np.min(sepal lengths))
             print("Maximum:", np.max(sepal lengths))
             # Create a histogram with an opacity of 50% (alpha=0.5)
             plt.hist(sepal lengths, alpha=0.5)
             # Add a vertical line to the plot showing the mean.
             plt.axvline(np.mean(sepal lengths), label="mean")
             plt.axvline(np.median(sepal lengths), label="median")
             # Your code here!
             # Don't forget to label the axes!
             plt.xlabel("Sepal length (cm)")
             plt.ylabel("Frequency (number of individuals)")
             # Add a legend to the plot
             plt.legend()
```

```
# snow tne piot in our jupyter notebook
plt.show()
```

NOW ANALYZING DATASET: ./data/Lab_02/Iris_setosa_data.csv Iris data shape is: (50, 4)

Correct data shape of 50 rows, 4 columns!

The sepal lengths in this dataset is:

[5.1 4.9 4.7 4.6 5. 5.4 4.6 5. 4.4 4.9 5.4 4.8 4.8 4.3 5.8

5.7 5.4 5.1 5.7 5.1 5.4 5.1 4.6 5.1 4.8 5. 5. 5.2 5.2 4.7

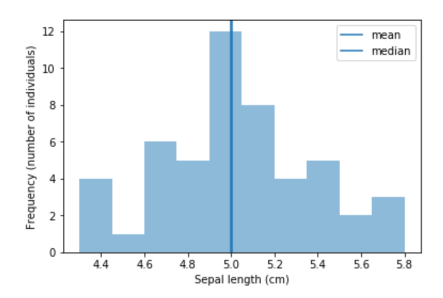
4.8 5.4 5.2 5.5 4.9 5. 5.5 4.9 4.4 5.1 5. 4.5 4.4 5. 5.1

4.8 5.1 4.6 5.3 5.]

Mean: 5.006

Standard deviation: 0.348946987378

Median: 5.0 Minimum: 4.3 Maximum: 5.8



NOW ANALYZING DATASET: ./data/Lab 02/Iris versicolor data.csv

Iris data shape is: (50, 4)

Correct data shape of 50 rows, 4 columns!

The sepal lengths in this dataset is:

[7. 6.4 6.9 5.5 6.5 5.7 6.3 4.9 6.6 5.2 5. 5.9 6. 6.1

5.6

6.7 5.6 5.8 6.2 5.6 5.9 6.1 6.3 6.1 6.4 6.6 6.8 6.7 6.

5.7

5.5 5.5 5.8 6. 5.4 6. 6.7 6.3 5.6 5.5 5.5 6.1 5.8 5.

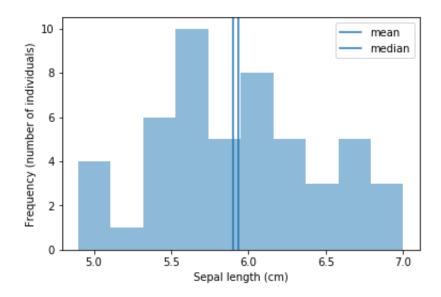
5.6

5.7 5.7 6.2 5.1 5.7]

Mean: 5.936

Standard deviation: 0.510983365678

Median: 5.9
Minimum: 4.9
Maximum: 7.0



NOW ANALYZING DATASET: ./data/Lab_02/Iris_virginica_data.csv

Iris data shape is: (50, 4)

Correct data shape of 50 rows, 4 columns!

The sepal lengths in this dataset is:

[6.3 5.8 7.1 6.3 6.5 7.6 4.9 7.3 6.7 7.2 6.5 6.4 6.8 5.7

5.8

6.4 6.5 7.7 7.7 6. 6.9 5.6 7.7 6.3 6.7 7.2 6.2 6.1 6.4

7.2

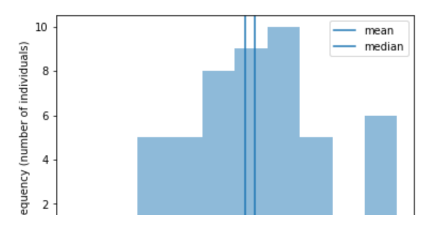
7.4 7.9 6.4 6.3 6.1 7.7 6.3 6.4 6. 6.9 6.7 6.9 5.8 6.8 6.7

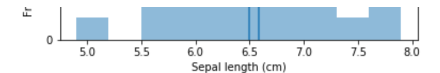
6.7 6.3 6.5 6.2 5.9]

Mean: 6.588

Standard deviation: 0.629488681391

Median: 6.5
Minimum: 4.9
Maximum: 7.9





Lab 2 Bonus exercise

Bonus Exercise 1: Now take the above code and edit it so that we analyze all of the 4 flower morphology variables, for all of the species of plants. Label the plot axis and title with the appropriate information (flower species for title, and the morphological variable on the x axis).

```
for filename in filenames:
In [27]:
             # Load our file data from "filename" into a variable called iris dat
             iris data = np.loadtxt(fname=filename, delimiter=",")
             print("----")
             print("NOW ANALYZING DATASET: ", filename)
             # Print the shape of the loaded dataset
             data shape = iris data.shape
             print("Iris data shape is:", data shape)
             # Print a warning if the data shape is not what we expect
             if (data shape[0] != 50) or (data shape[1] != 4):
                 print("Unexpected data shape!")
             else:
                 print("Correct data shape of 50 rows, 4 columns!")
             # the number of columns is the same as the number of items in the fi
             num columns = len(iris data[0])
             axis labels = ["Sepal length (cm)",
                             "Sepal width (cm)",
                             "Petal length (cm)",
                             "Petal width (cm)"]
             # THIS IS CALLED A NESTED FOR LOOP!
             # A NESTED FOR LOOP HAS A FOR LOOP INSIDE OF ANOTHER FOR LOOP.
             for index in np.arange(0, num columns):
                 data subset = iris data[0:, index]
                 # Print some interesting characteristics of the data
                 print("Mean:", np.mean(data subset))
                 print("Standard deviation:", np.std(data subset))
                 print("Median:", np.median(data subset))
                 print("Minimum:", np.min(data subset))
                 print("Maximum:", np.max(data subset))
                 # Create a histogram with an opacity of 50% (alpha=0.5)
                 plt.hist(data subset, alpha=0.5)
```

```
# Add a vertical line to the plot showing the mean.
plt.axvline(np.mean(data_subset), label="mean")
plt.axvline(np.median(data_subset), label="median")
# Your code here!

# Don't forget to label the axes!
plt.xlabel(axis_labels[index])
plt.ylabel("Frequency (number of individuals)")

species_name = filename.strip('.csv')
plt.title(species_name)

# Add a legend to the plot
plt.legend()

# Show the plot in our jupyter notebook
plt.show()
```

NOW ANALYZING DATASET: ./data/Lab_02/Iris_setosa_data.csv

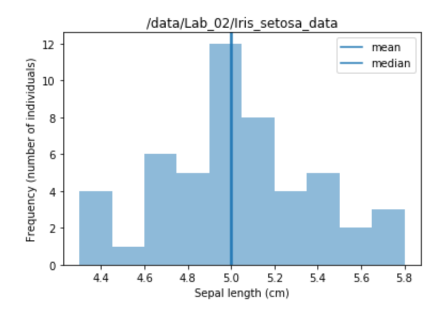
Iris data shape is: (50, 4)

Correct data shape of 50 rows, 4 columns!

Mean: 5.006

Standard deviation: 0.348946987378

Median: 5.0
Minimum: 4.3
Maximum: 5.8

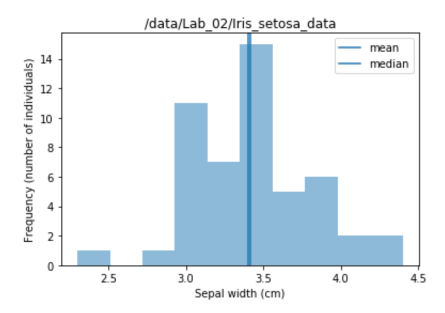


Mean: 3.418

Standard deviation: 0.377194909828

Median: 3.4 Minimum: 2.3

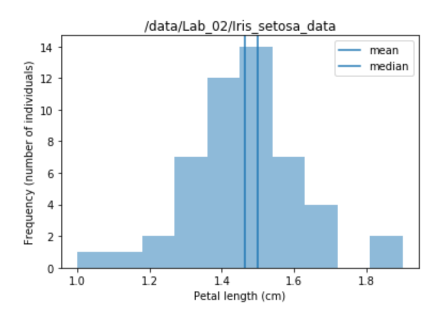
Maximum: 4.4



Mean: 1.464

Standard deviation: 0.171767284429

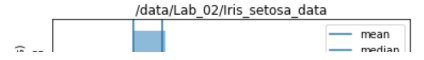
Median: 1.5
Minimum: 1.0
Maximum: 1.9

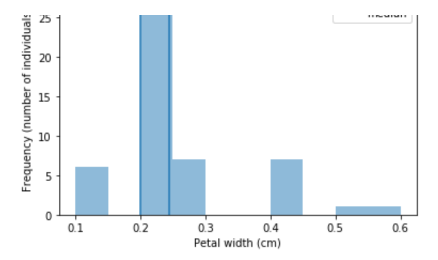


Mean: 0.244

Standard deviation: 0.106131993291

Median: 0.2 Minimum: 0.1 Maximum: 0.6





NOW ANALYZING DATASET: ./data/Lab_02/Iris_versicolor_data.csv

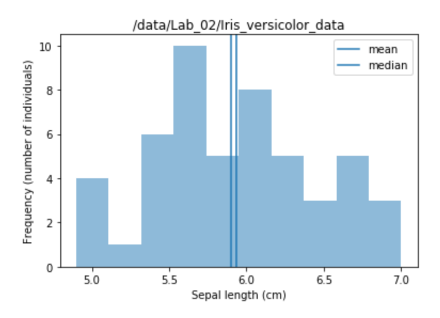
Iris data shape is: (50, 4)

Correct data shape of 50 rows, 4 columns!

Mean: 5.936

Standard deviation: 0.510983365678

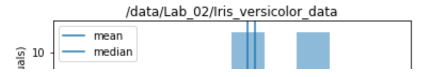
Median: 5.9
Minimum: 4.9
Maximum: 7.0

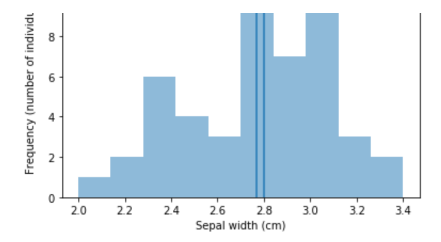


Mean: 2.77

Standard deviation: 0.31064449134

Median: 2.8
Minimum: 2.0
Maximum: 3.4

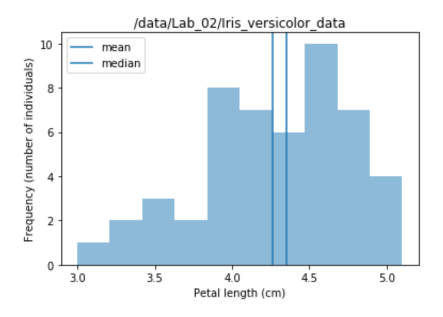




Mean: 4.26

Standard deviation: 0.465188133985

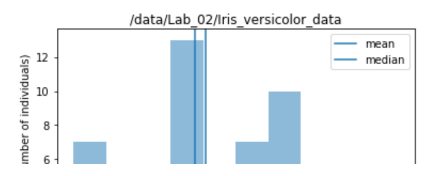
Median: 4.35
Minimum: 3.0
Maximum: 5.1

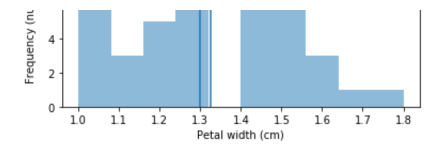


Mean: 1.326

Standard deviation: 0.195765165441

Median: 1.3 Minimum: 1.0 Maximum: 1.8





NOW ANALYZING DATASET: ./data/Lab_02/Iris_virginica_data.csv

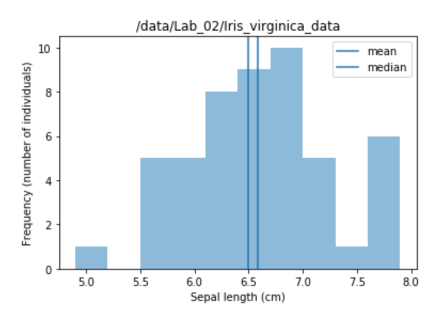
Iris data shape is: (50, 4)

Correct data shape of 50 rows, 4 columns!

Mean: 6.588

Standard deviation: 0.629488681391

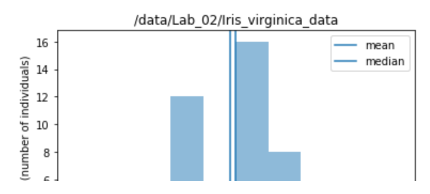
Median: 6.5
Minimum: 4.9
Maximum: 7.9

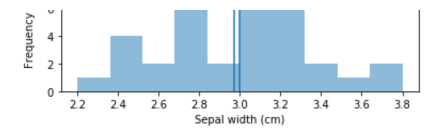


Mean: 2.974

Standard deviation: 0.319255383666

Median: 3.0 Minimum: 2.2 Maximum: 3.8

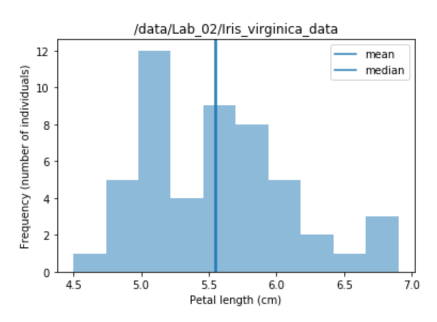




Mean: 5.552

Standard deviation: 0.546347874527

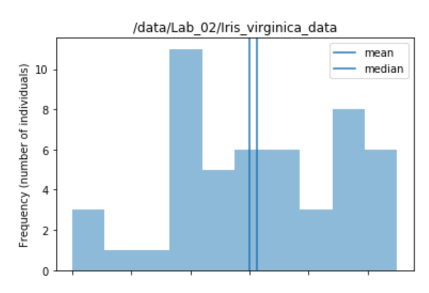
Median: 5.55
Minimum: 4.5
Maximum: 6.9



Mean: 2.026

Standard deviation: 0.271889683512

Median: 2.0
Minimum: 1.4
Maximum: 2.5



1.4 1.6 1.8 2.0 2.2 2.4 Petal width (cm)

In []: