EDA using Data Visualization

Data visualization is the most effective tool in exploratory data analysis as it gives a birds eye view of the entire data in the most efficient way and amplifies the cognition. Python library package matplotlib, provides a very user firiently approach in data visualization. There are two approached namely stateless and stateful.

Stateful and Stateless Approaches

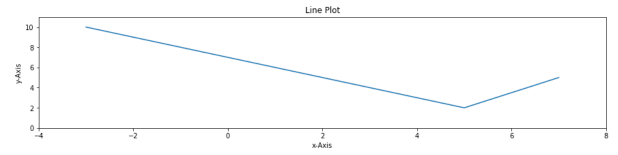
In stateful approach, the program refer the pyplot directly and any changes made to the state reflect across all plots. Where as in stateless approach is an object oriented approach, where an object with proprties of figure and axes is created from pyplot and that object is refrenced in creation of subsequent plots instead of directly calling pyplot function.

Pros and cons of each approach

Pros and cons of each approach is dependent on the usage. For simple and fast ploting stateful is preferred but will become difficult to manage as the number of plots increases. On the other hand stateless approach has the overhead of creating an object but is easier to keep track of each plots and properties.

Before diving into the EDA, providing an example of each of these approaches for better understanding

```
In [1]: # import necessary packages
        import pandas as pd
        import numpy as np
        import matplotlib.pyplot as plt
        %matplotlib inline
        #Create two lists
        x = [-3, 5, 7]
        y = [10, 2, 5]
        # Plot the above lists elements pairs on x-y axis using stateful approac
        plt.figure(figsize=(15,3))
        plt.plot(x,y)
        plt.xlim(-4,8)
        plt.ylim(0,11)
        plt.xlabel("x-Axis")
        plt.ylabel("y-Axis")
        plt.title('Line Plot')
        plt.show()
```



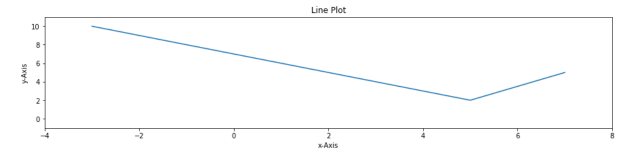
```
In [2]: %matplotlib inline
# Plot the above lists elements pairs on x-y axis using stateless approa
ch

#create a pyplot object for figure and axis
fig,ax = plt.subplots(figsize=(15,3))

ax.plot(x,y)
ax.set_xlim(-4,8)
ax.set_ylim(-1,11)
ax.set_ylim(-1,11)
ax.set_ylabel("x-Axis")
ax.set_ylabel("y-Axis")
ax.set_title('Line Plot')
fig.show()
```

/Users/jacobdenny/anaconda3/lib/python3.7/site-packages/ipykernel_launc her.py:13: UserWarning: Matplotlib is currently using module://ipykernel.pylab.backend_inline, which is a non-GUI backend, so cannot show the figure.

del sys.path[0]



The above two figures are plotted using the two approaches and the difference is that in stateless approach an object is created of type pyplot and the object properties and functions are used to plot data whereas in stateful the ploting directly refer to pyplot

Import and load data set

A dataset from a csv file needs to be loaded into the python environment and is stored in a dataframe which is analogous to a data table, in that the data is seen as rows and columns. Two packages DataFrame and read csv are used for this.

```
In [3]: #Import additional packages necessary for loading data from csv source a
    nd manipulating it in python
    from pandas.plotting import scatter_matrix
    from pandas import DataFrame, read_csv
```

Out[4]:

	ld	SepalLengthCm	SepalWidthCm	PetalLengthCm	PetalWidthCm	Species
0	1	5.1	3.5	1.4	0.2	Iris-setosa
1	2	4.9	3.0	1.4	0.2	Iris-setosa
2	3	4.7	3.2	1.3	0.2	Iris-setosa
3	4	4.6	3.1	1.5	0.2	Iris-setosa
4	5	5.0	3.6	1.4	0.2	Iris-setosa

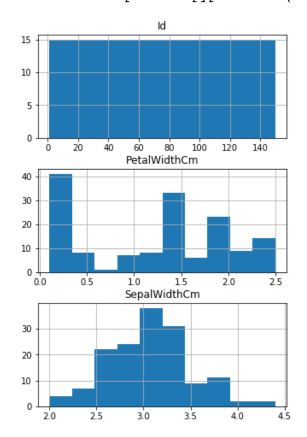
Univariate Plots

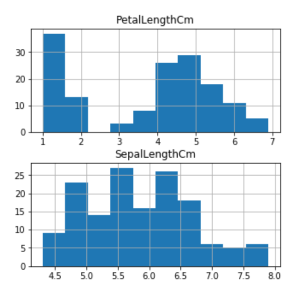
Plot each column to get an idea of the distribution. We will use the following

- Histogram
- Densityplot
- Boxplot

```
In [5]: #Histogram
    dfIris.hist(figsize=(12,8))
    plt.show
```

Out[5]: <function matplotlib.pyplot.show(*args, **kw)>

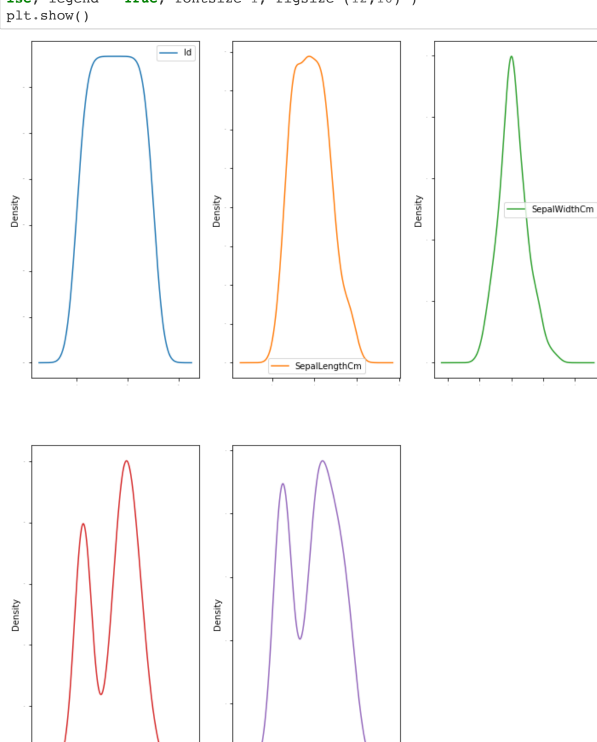




What Histogram Tells About Data

The histograms are displayed for each column of the data, which means the id column also gets plotted, which has no value in the data analysis and is ignored. Sepallength and sepalwidth shows almost normal distribution, petal length and width seems to be bimodal

In [6]: #Density plot
 dfIris.plot(kind="density", subplots = True, layout = (2,3), sharex = Fa
 lse, legend = True, fontsize=1, figsize=(12,16))
 plt.show()

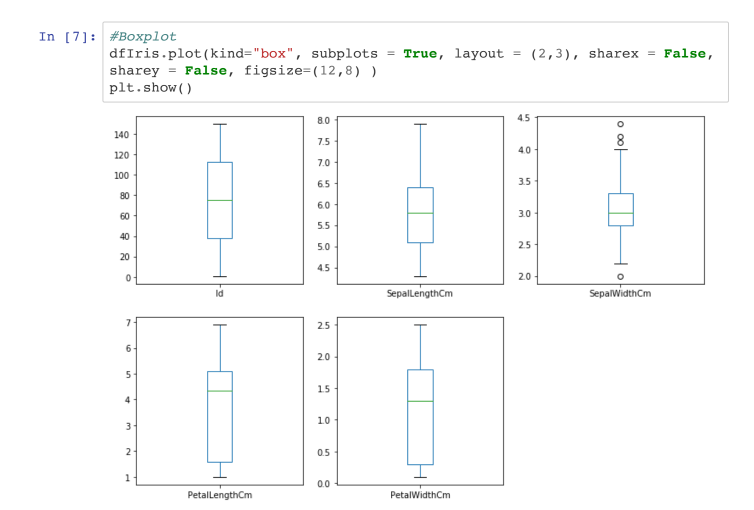


PetalWidthCm

PetalLengthCm

What Densityplots Tells About Data

The density plot is consistant with histogram and are displayed for each column of the data, which means the id column also gets plotted, which has no value in the data analysis and is ignored. Sepallength and sepalwidth shows almost normal distribution, petal length and width seems to be bimodal



What Boxplots Tells About Data

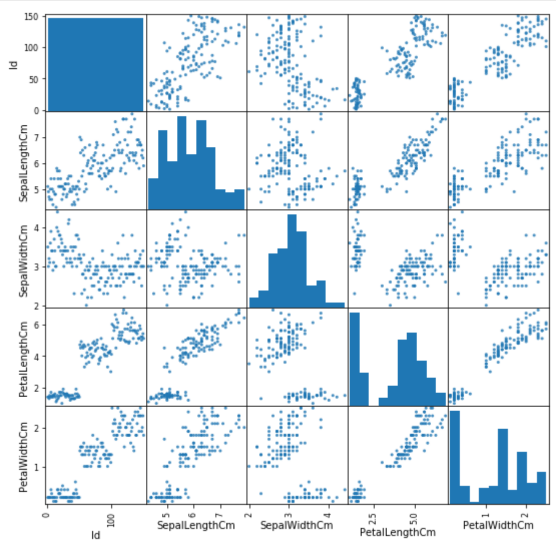
- · Ignore the plot for id
- For sepal length the values range from 4.5cm to 8cm
- For sepal width the values range from 2cm to 4.5cm
- For petal length the values range from 1cm to 7cm
- For petal widht the values range from 0 to 2.5cm
- · Sepalwidth has evident outliers

Multivariate Analysis

The above plots shows chrecteristics of each column like distribution of data, range, dispersion or independent features of each column. Now we would like to see how each feature is affected by other features and we use multivariate analysis for this. Scatter matrix is the best technique for this analysis.

```
In [8]: #Scatter matrix

scatter_matrix(dfIris, alpha=0.8, figsize=(9,9))
plt.show()
```



What scatter matrix tells us

- As sepal length increase petal length nd width increase
- Sepal width exhibits almost null corelation with other features and hence has data points scattered all over
- Petal lenght and petal width also shows a positive linear corelation

```
In [ ]:

In [ ]:

In [ ]:
```

```
In [ ]:
```

Supervised Linear Regression

Creating a model defining the outcome variable as a linear combination of a set of independent features. Steps involve:

- · Preprocess data
- EDA
- · Split data into test and train set
- · Build and train model
- Test/evaluate model

```
In [9]: # import scikit module for linear regression
    from sklearn.linear_model import LinearRegression
    # import scikit module for spliting data into training and testing datas
    ets
    from sklearn.model_selection import train_test_split
    # import scikit module for kfold cross validation for evaluation and cro
    ss validation
    from sklearn.model_selection import KFold
    from sklearn.model_selection import cross_val_score
```

Out[10]:

	CRIM	ZN	INDUS	CHAS	NOX	RM	AGE	DIS	RAD	TAX	PTRATIO	В	LSTAT
0	0.00632	18.0	2.31	0	0.538	6.575	65.2	4.0900	1	296	15.3	396.90	4.98
1	0.02731	0.0	7.07	0	0.469	6.421	78.9	4.9671	2	242	17.8	396.90	9.14
2	0.02729	0.0	7.07	0	0.469	7.185	61.1	4.9671	2	242	17.8	392.83	4.03
3	0.03237	0.0	2.18	0	0.458	6.998	45.8	6.0622	3	222	18.7	394.63	2.94
4	0.06905	0.0	2.18	0	0.458	7.147	54.2	6.0622	3	222	18.7	396.90	5.33

Preprocessing Data

We are trying to predict the median value of owner occupied houses in 1000 dollors(MEDV) using linear regression with 'RM', 'AGE', 'DIS', 'RAD', 'PTRATIO' as predictors. Hence the subset is extracted from dataframe and is stored into another data frame. Also, from domain knowledge we know that the average value of a home, the pupil teacher ratio and average number of rooms per dwelling cannot be zero. If any row cell contain zero for these three columns that needs to be marked as mising. So preprocessing includes:

- · Extract the columns under consideration
- · Data cleaning
- · Check for invalid Values

```
In [11]: # Extract a sub-dataset from the original one -- > dataframe: df2
         df2 = df[['RM', 'AGE', 'DIS', 'RAD', 'PTRATIO', 'MEDV']]
         #Data cleaning
         #Replace 0 value with NaN for columns 'RM', 'PTRATIO', 'MEDV'
         df[['RM', 'PTRATIO', 'MEDV']] = df[['RM', 'PTRATIO', 'MEDV']].replace(0,
         np.NaN)
          #Verify there are no missing values in the dataset
         print(df.isnull().sum())
                     0
         CRIM
         7N
                     0
         INDUS
                     0
         CHAS
         NOX
                     0
                     0
         RM
                     0
         AGE
         DIS
                     0
         RAD
                     0
         TAX
                     0
         PTRATIO
                     0
         LSTAT
                     0
         MEDV
         dtype: int64
```

There are no missing values in the dataset. Now we are ready to move to Exploratory data analysis

Exploratory Data Analysis

In this step we are getting to know our data. The number of columns and rows, what each column represent etc. The following are explored in this section:

- · Shape of data set
- Type of each value in the dataset
- · View a snippet of the dataset
- · Descriptive statistics of each column
- · Univariate plots
- · Multivariate plots

```
In [12]: #SHAPE
# Get the dimensions or Shape of the dataset
# i.e. number of records/rows x number of variables/columns
print(df2.shape)
(452, 6)
```

We have 6 variables and 452 rows.

```
In [13]:
         #TYPE
         # Get the data types of all variabLes/attributes of the data set
         # The results show
         print(df2.dtypes)
                     float64
         RM
         AGE
                     float64
                     float64
         DIS
         RAD
                       int64
                     float64
         PTRATIO
                     float64
         MEDV
         dtype: object
```

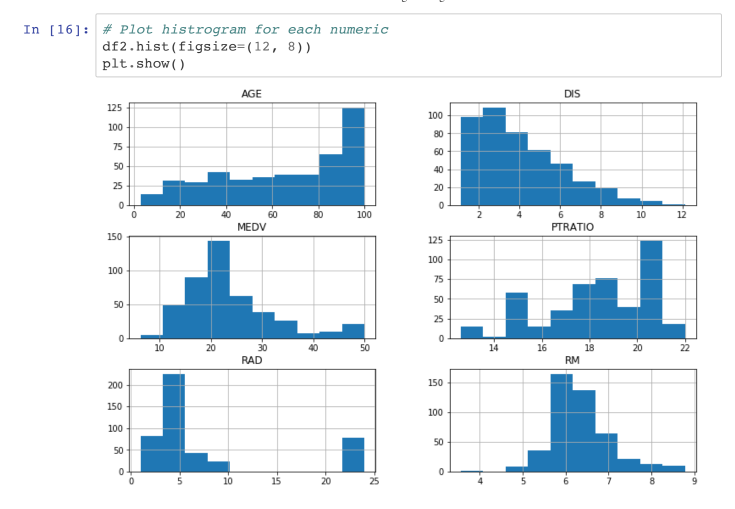
All the features and the dependent variable(MEDV) are of type floating point except RAD which is integer type.

```
In [14]: #View of a snippet of Dataset
         # Get the first five records
         print(df2.head(5))
              RM
                   AGE
                           DIS
                               RAD PTRATIO MEDV
           6.575 65.2 4.0900
                                       15.3 24.0
         0
                                1
         1
           6.421 78.9
                       4.9671
                                 2
                                       17.8
                                            21.6
         2
                                 2
                                       17.8 34.7
           7.185 61.1
                        4.9671
         3
           6.998 45.8
                                       18.7
                        6.0622
                                 3
                                             33.4
           7.147 54.2
                        6.0622
                                       18.7
                                             36.2
```

	RM	AGE	DIS	RAD	PTRATIO	
MEDV						
count	452.000000	452.000000	452.000000	452.000000	452.000000	452.
000000						
mean	6.343538	65.557965	4.043570	7.823009	18.247124	23.
750442						
std	0.666808	28.127025	2.090492	7.543494	2.200064	8.
808602						
min	3.561000	2.900000	1.129600	1.000000	12.600000	6.
300000						
25%	5.926750	40.950000	2.354750	4.000000	16.800000	18.
500000						
50%	6.229000	71.800000	3.550400	5.000000	18.600000	21.
950000						
75%	6.635000	91.625000	5.401100	7.000000	20.200000	26.
600000						
max	8.780000	100.000000	12.126500	24.000000	22.000000	50.
000000						

The above table gives the summary statistics for each of the columns. This includes number of records, mean, standard deviation, minimum 25 percentile 50 percentile 75 percentile and the maximum values of each column namely RM, AGE, DIS, RAD, PTRATIIO and MEDV

Histograms

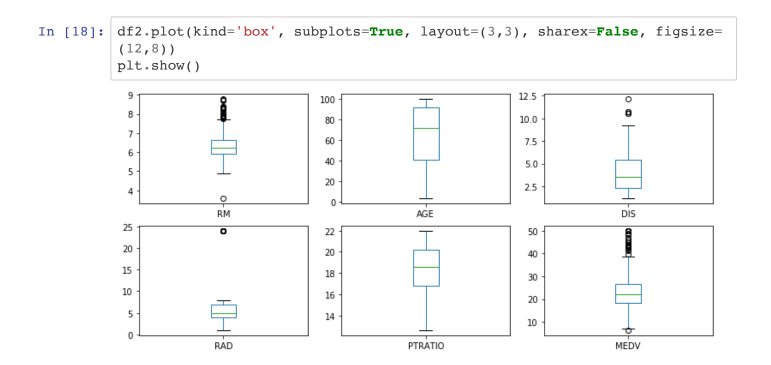


MEDV,PTRATIO and RM has a distribution that is almost normal but the rest are not.

Density Plots

```
In [17]: # Density plots
          # IMPORTANT NOTES: 5 numeric variables -> at Least 5 plots -> Layout (2,
          3): 2 rows, each row with 3 plots
          df2.plot(kind='density', subplots=True, layout=(2, 3), sharex=False, leg
          end=True, fontsize=1,
          figsize=(12, 16))
          plt.show()
                                - RM
                                             AGE
                                                                                      - DIS
           Density
                                                                 Density
                               - RAD
                                             PTRATIO
                                                                                  — MEDV
                                      Density
```

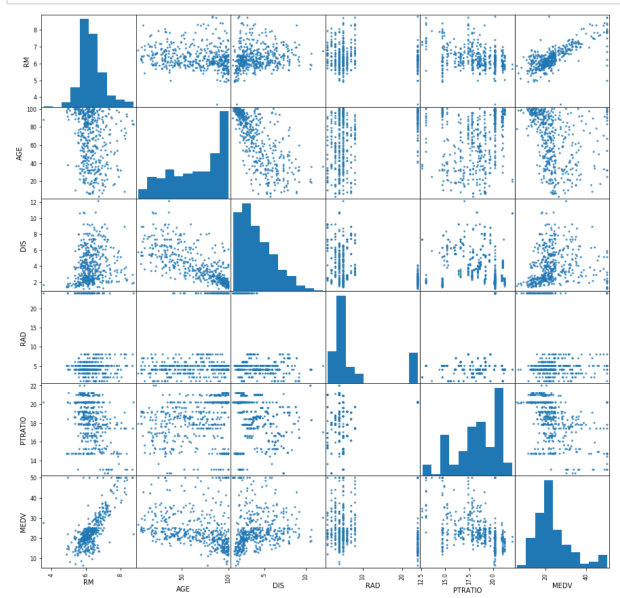
Consistent with the histogram, density plots also gives an almost normal distribution for MEDV,PTRATIO and RM. but DIS is also fairly normal in density plot. But RAD and AGE are bimodal.



- · RAD has a single outlier
- · MEDV, RM and DIS has outliers at both ends
- Min value of RM is 1 and max at 9
- Min value od AGE is 4 and max just below 100
- Min value of DIS is 1 and max just below 12.5
- Min value of RAD is 1 and max just below 10 with an outlier at 25
- Min value of PTRATIO is 5 and max just below 22
- Min value of MEDV is 41 and max just above 50

Multivariate Analysis : Scatter Matrix

```
In [19]: # scatter plot matrix
    scatter_matrix(df2, alpha=0.8, figsize=(15, 15))
    plt.show()
```



- · There is an evident linear relation between MEDV and RM
- MEDV and AGE is showing an inverse linear relation
- DIS and MEDV on the other hand has a direct linear relation

Separate Dataset into Input & Output NumPy Arrays

```
In [20]: # Store dataframe values into a numpy array
array = df2.values
# separate array into input and output components by slicing
# For X (input)[:, 5] --> all the rows, columns from 0 - 4 (5 - 1)
X = array[:,0:5]
# For Y (output)[:, 5] --> all the rows, column index 5 (Last column)
Y = array[:,5]
```

- InDependent variables are 'RM', 'AGE', 'DIS', 'RAD' and 'PTRATIO' columns 0 to 4
- Dependent variable is 'MEDV' column 5

Split Input/Output Arrays into Training/Testing Datasets

```
In [21]: # Split the dataset --> training sub-dataset: 67%; test sub-dataset:
    test_size = 0.33
    # Selection of records to include in which sub-dataset must be done rand
    omly
    # use this seed for randomization
    seed = 7
    # Split the dataset (both input & outout) into training/testing datasets
    X_train, X_test, Y_train, Y_test = train_test_split(X, Y, test_size=test
    _size,
    random_state=seed)
```

- 33% of the data is test data
- 67% of the data is train data
- · Split is based on random sampling

Build and Train the Model

```
In [22]: # Build the model
         model = LinearRegression()
         # Train the model using the training sub-dataset
         model.fit(X_train, Y_train)
         # Print out the coefficients and the intercept
         # print intercept and coefficients
         print ("Intercept", model.intercept )
         #print (model.coef )
         # If we want to print out the list of the coefficients with their corres
         pondent variable name
         # pair the feature names with the coefficients
         names 2 = ['RM', 'AGE', 'DIS', 'RAD', 'PTRATIO']
         coeffs_zip = zip(names_2, model.coef )
         # Convert iterator into set
         coeffs = set(coeffs zip)
         #print (coeffs)
         for coef in coeffs:
          print(coef, "\n")
         Intercept -4.53602172360959
         ('DIS', -0.800386625412262)
```

```
Intercept -4.53602172360959
('DIS', -0.800386625412262)
('RAD', -0.14269109757375334)
('AGE', -0.08160589380401152)
('PTRATIO', -0.8641387261012642)
('RM', 8.457012650599342)
```

The model build by the algorithm is

- MEDV = -4.53602172360959 +(8.45701265 RM) -(0.08160589 AGE) -(0.80038663 DIS) -(0.1426911 RAD) -(0.86413873 * PTRATIO)
 - As per the model for unit increase in RM, with all other features held constant MEDV increases 8.45701265
 - As per the model for unit increase in AGE, with all other features held constant MEDV decreases 0.08160589
 - As per the model for unit increase in DIS, with all other features held constant MEDV decreases
 0.80038663
 - As per the model for unit increase in RAD, with all other features held constant MEDV decreases
 0.1426911
 - As per the model for unit increase in PTRATIO, with all other features held constant MEDV decreases 0.86413873

Calculate R-Squared

```
In [23]: R_squared = model.score(X_test, Y_test)
    print(R_squared)
    0.4921278466441376
```

The model has successfully explained 49 % of variance in MEDV using variables 'RM', 'AGE', 'DIS', 'RAD' and 'PTRATIO'

Prediction

We have trained the model. Let's use the trained model to predict the "Median value of owner-occupied homes in 1000 dollars" (MEDV)_

The suburb area has the following predictors:

- RM: average number of rooms per dwelling = 6.0
- AGE: proportion of owner-occupied units built prior to 1940 = 55
- DIS: weighted distances to five Boston employment centers = 5
- RAD: index of accessibility to radial highways = 2
- PTRATIO: pupil-teacher ratio by town = 16

```
In [24]: Predicted_Value = model.predict([[6.0, 55, 5, 2, 16]])
    print("Predicted Median value of owner-occupied homes in 1000 dollars fo
    r the above house is :",Predicted_Value[0],"Thousand" )
```

Predicted Median value of owner-occupied homes in 1000 dollars for the above house is: 23.60419508093679 Thousand

Evaluate/Validate Algorithm/Model Using K-Fold Cross-Validation

```
In [25]: # Evaluate the algorithm
         # Specify the K-size
         num folds = 10
         # Fix the random seed
         # must use the same seed value so that the same subsets can be obtained
         # for each time the process is repeated
         seed = 7
         # Split the whole data set into folds
         kfold = KFold(n splits=num folds, random state=seed)
         # For Linear regression, we can use MSE (mean squared error) value
         # to evaluate the model/algorithm
         scoring = 'neg mean squared error'
         # Train the model and run K-foLd cross-validation to validate/evaluate t
         he model
         results = cross val score(model, X, Y, cv=kfold, scoring=scoring)
         # Print out the evaluation results
         # Result: the average of all the results obtained from the k-foLd cross-
         validation
         print(results.mean())
```

-31.17777676938244

After we train we evaluate Use K-Fold to determine if the model is acceptable We pass the whole set because the system will divide for us -31 avg of all error (mean of square errors) this value would traditionally be positive value, but scikit reports as neg Square root would be between 5 and 6

With the test train split we calculated an R square of 49 but that is applicable to only that sample. Where as in K-fold validation with number of folds set to 10, we have 10 different accuracies which averages to 31. 17777. K-fold validation describes the population better than a predicting on single sample from population and that deceases the accuracy to 31.2.

Logistic Regression

Logistic regrssion is a classification technique.

Dataset

Predicted attribute: class of iris plant. Number of Instances: 150 (50 in each of three classes) Number of predictors: 4 numeric, predictive attributes and the class Attribute Information:

- 1. sepal length in cm
- 2. sepal width in cm
- 3. petal length in cm
- 4. petal width in cm
- 5. class: Iris Setosa, Iris Versicolour, Iris Virginica

Import necessary packages

```
In [26]: # Import scikit-Learn module for the algorithm/modeL: Logistic Regressio
         from sklearn.linear model import LogisticRegression
         # Import scikit-Learn module classification report to later use for info
         rmation about how the system
         # try to classify / lable each record
         from sklearn.metrics import classification report
In [27]: #Load data into a pandas dataframe
         # Already loaded into dfIris during visualization exercise
         print(dfIris.head(5))
         dfIris=dfIris.drop(['Id'], axis=1)
         print("After droping column Id \n", dfIris.head(5))
                SepalLengthCm SepalWidthCm PetalLengthCm PetalWidthCm
                                                                                Sp
         ecies
             1
                           5.1
                                         3.5
                                                         1.4
                                                                       0.2 Iris-s
         etosa
         1
             2
                           4.9
                                         3.0
                                                         1.4
                                                                       0.2 Iris-s
         etosa
                           4.7
                                         3.2
                                                        1.3
                                                                       0.2 Iris-s
             3
         etosa
                                                                       0.2 Iris-s
         3
                           4.6
                                         3.1
                                                        1.5
         etosa
             5
                           5.0
                                         3.6
                                                        1.4
                                                                       0.2 Iris-s
         etosa
         After droping column Id
             SepalLengthCm SepalWidthCm PetalLengthCm PetalWidthCm
                                                                             Speci
         es
         0
                       5.1
                                     3.5
                                                                   0.2 Iris-setos
                                                    1.4
         a
                                                                   0.2 Iris-setos
         1
                       4.9
                                     3.0
                                                    1.4
         a
         2
                       4.7
                                     3.2
                                                    1.3
                                                                   0.2 Iris-setos
         а
         3
                       4.6
                                     3.1
                                                    1.5
                                                                   0.2 Iris-setos
         а
```

Preprocess Dataset

4

a

Clean Data: Find & Mark Missing Values

5.0

- Zero values are invalid in these columns as petal and sepal width cannot have 0 as a valid value. It they exist, we need to mark them as missing value or numpy. NaN

3.6

1.4

0.2 Iris-setos

```
In [28]: # mark zero values as missing or NaN
         dfIris[[ 'SepalLengthCm' , 'SepalWidthCm' , 'PetalLengthCm' ,'PetalWidth
         Cm' ]] = dfIris[['SepalLengthCm' , 'SepalWidthCm' , 'PetalLengthCm' , 'Pe
         talWidthCm' ]].replace(0,np.NaN)
         # count the number of NaN values in each column
         print (dfIris.isnull().sum())
         SepalLengthCm
                          0
         SepalWidthCm
                          0
         PetalLengthCm
                          0
         PetalWidthCm
                          0
         Species
         dtype: int64
```

There are no missing values in the dataset. Now we are ready to move to Exploratory data analysis

Exploratory Data Analysis

In this step we are getting to know our data. The number of columns and rows, what each column represent etc. The following are explored in this section:

- · Shape of data set
- Type of each value in the dataset
- · View a snippet of the dataset
- · Descriptive statistics of each column
- Univariate plots
- · Multivariate plots

```
In [29]: # get the dimensions or shape of the dataset
# i.e. number of records / rows X number of variables / columns
print(dfIris.shape)
(150, 5)
```

We have 5 variables and 150 rows as we have deleted the column labeled ld.

```
In [30]: #get the data types of all the variables / attributes in the data set
         print(dfIris.dtypes)
          #return the first five records / rows of the data set
         print(dfIris.head(5))
         SepalLengthCm
                           float64
         SepalWidthCm
                           float64
         PetalLengthCm
                           float64
         PetalWidthCm
                           float64
         Species
                            object
         dtype: object
            SepalLengthCm SepalWidthCm
                                          PetalLengthCm PetalWidthCm
                                                                             Specie
         s
         0
                       5.1
                                     3.5
                                                     1.4
                                                                   0.2
                                                                        Iris-setos
         a
                       4.9
                                     3.0
                                                     1.4
                                                                   0.2
                                                                        Iris-setos
         1
         а
         2
                       4.7
                                     3.2
                                                     1.3
                                                                   0.2 Iris-setos
         а
         3
                       4.6
                                     3.1
                                                     1.5
                                                                   0.2 Iris-setos
         a
```

All the features are of type floating point except species which is object type.

5.0

4

а

1.4

0.2 Iris-setos

3.6

	SepalLengthCm	SepalWidthCm	PetalLengthCm	PetalWidthCm
count	150.000000	150.000000	150.000000	150.000000
mean	5.843333	3.054000	3.758667	1.198667
std	0.828066	0.433594	1.764420	0.763161
min	4.300000	2.000000	1.000000	0.100000
25%	5.100000	2.800000	1.600000	0.300000
50%	5.800000	3.000000	4.350000	1.300000
75%	6.400000	3.300000	5.100000	1.800000
max	7.900000	4.400000	6.900000	2.500000

The above table gives the summary statistics for each of the numeric columns.

This includes number of records, mean, standard deviation, minimum 25 percentile 50 percentile 75 percentile and the maximum values of each column namely SepalLengthCm, SepalWidthCm, PetalLengthCm, PetalWidthCm

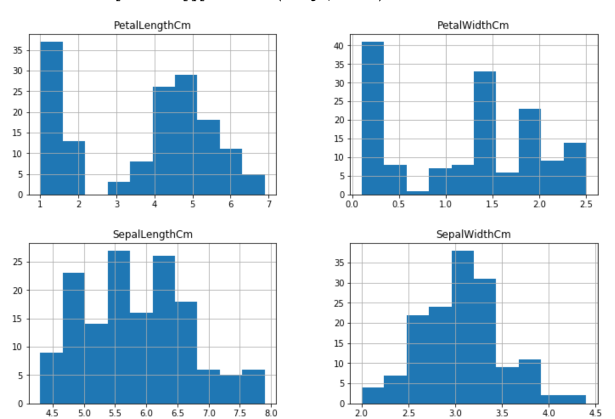
Univariate Plots

Plot each column to get an idea of the distribution. We will use the following

- Histogram
- Densityplot
- Boxplot

```
In [32]: #Histogram
dfIris.hist(figsize=(12,8))
plt.show
```

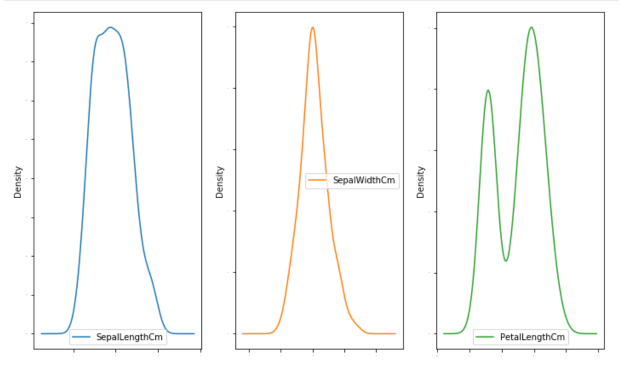
Out[32]: <function matplotlib.pyplot.show(*args, **kw)>

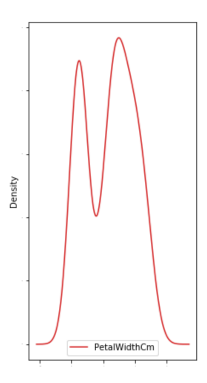


What Histogram Tells About Data

The histograms are displayed for each column of the data. Sepallength and sepalwidth shows almost normal distribution, petal lenght and width seems to be bimodal

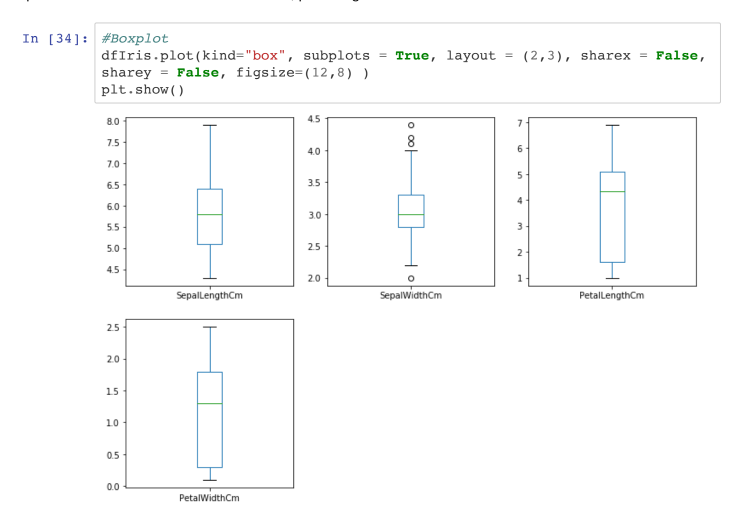
```
In [33]: #Density plot
    dfIris.plot(kind="density", subplots = True, layout = (2,3), sharex = Fa
    lse, legend = True, fontsize=1, figsize=(12,16) )
    plt.show()
```





What Densityplots Tells About Data

The density plot is consistant with histogram and are displayed for each column of the data. Sepallength and sepalwidth shows almost normal distribution, petal length and width seems to be bimodal



What Boxplots Tells About Data

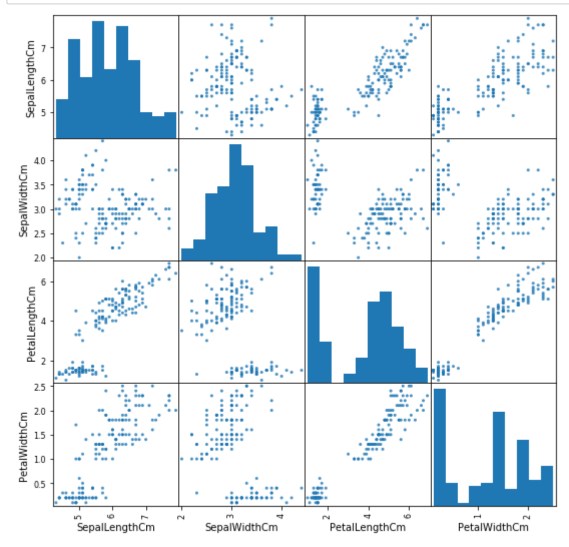
- For sepal length the values range from 4.5cm to 8cm
- For sepal width the values range from 2cm to 4.5cm
- For petal length the values range from 1cm to 7cm
- For petal widht the values range from 0 to 2.5cm
- · Sepalwidth has evident outliers

Multivariate Analysis

The above plots shows chrecteristics of each column like distribution of data, range, dispersion or independent features of each column. Now we would like to see how each feature is affected by other features and we use multivariate analysis for this. Scatter matrix is the best technique for this analysis.

```
In [35]: #Scatter matrix

scatter_matrix(dfIris, alpha=0.8, figsize=(9,9))
plt.show()
```



What scatter matrix tells us

As sepal length increase petal length and width increase Sepal width exhibits almost null corelation with other features and hence has data points scattered all over Petal lenght and petal width also shows a positive linear corelation

Each of the groups are equally represented in the data.

Separate Dataset into Input & Output NumPy arrays

Training data Set (X) consist of the independent variables or predictors Desired Output (Y) consist of the dependent variable or that which we are trying to predict

Split Input/Output Arrays into Training/Testing Datasets

```
In [38]: # split the dataset --> training sub-dataset: 67%; test sub-dataset: 33%
    test_size = 0.33
    #selection of records to include in each data sub-dataset must be done r
    andomly
    seed = 7
    #split the dataset (input and output) into training / test datasets
    X_train, X_test, Y_train, Y_test = train_test_split(X, Y, test_size=test
    _size,
    random_state=seed)
```

- 33% of the data is test data
- · 67% of the data is train data
- Split is based on random sampling

Build and Train the Model

```
In [39]: #build the model
    model = LogisticRegression()
    # train the model using the training sub-dataset
    model.fit(X_train, Y_train)
    #print the classification report
    predicted = model.predict(X_test)
    report = classification_report(Y_test, predicted)
    print(report)
```

	precision	recall	f1-score	support
Iris-setosa	1.00	1.00	1.00	14
Iris-versicolor	0.93	0.78	0.85	18
Iris-virginica	0.81	0.94	0.87	18
accuracy			0.90	50
macro avg	0.91	0.91	0.91	50
weighted avg	0.91	0.90	0.90	50

/Users/jacobdenny/anaconda3/lib/python3.7/site-packages/sklearn/linear_model/logistic.py:432: FutureWarning: Default solver will be changed to 'lbfgs' in 0.22. Specify a solver to silence this warning.

FutureWarning)

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"this warning.", FutureWarning)

A note on classification report - precision, recall, F1score and support

Precision

The precision is the ratio tp / (tp + fp) --> where tp is the number of true positives and fp the number of false positives. The precision represents the ability of the classifier not to label a positive sample as negative

Recall

The recall is the ratio tp / (tp + fn) --> where tp is the number of true positives and fn the number of false negatives. The recall represents the ability of the classifier to find all the positive samples

F-beta

The F-beta score can be interpreted as a weighted harmonic mean of the precision and recall --> where an F-beta score reaches its best value at 1 and worst score at 0 The F-beta score weights recall more than precision by a factor of beta beta == 1.0 means recall and precision are equally important

Support

The support is the number of occurrences of each class in y_true

We have perfect prediction of the Iris-setosa

For Iris-versicolor, 93% Iris-versicolor were classified correctly but 7% were classified as Iris-versicolor wrongly

For Iris-virginica, 81% Iris-virginica were classified correctly but 19% were classified as Iris-virginica wrongly 94% Iris-virginica were classified correctly and 6% were misclassified

```
In [40]: #score the accuracy leve
    result = model.score(X_test, Y_test)
    #print out the results
    print(("Accuracy: %.3f%%") % (result*100.0))
```

Accuracy: 90.000%

The model correctly classifies 90% of the times

Classify/Predict Model

Now use the model to predict the group of a given flower with given measurents for different features The new record has the following predictors:

- sepal length in cm = 5.3
- sepal Width in cm = 3.0
- petal length in cm = 4.5
- petal width in cm 1.5

```
In [41]: predicted=model.predict([[5.3, 3.0, 4.5, 1.5]])
   print("The model prediction for the given feaures is :",predicted[0])
```

The model prediction for the given feaures is : Iris-virginica

Based on the model's accuracy score: there is 90% chance that this new record is a Iris-virginica.

Evaluate the model using the 10-fold cross-validation technique.

- Specify the number of time of repeated splitting,
- Fix the random seed, Must use the same seed value so that the same subset can be obtained for each time the process is repeated,
- · Split the whole data set into folds, for logistic regression
- Use accuracy level to evaluate the model/algorithm, train the model
- Run k-fold cross-validation to validate/evaluate the model.
- · Print out the evaluation results

```
In [42]: # evaluate the algorythm
         # specify the number of time of repeated splitting, in this case 10 fold
         S
         n \text{ splits} = 10
         # fix the random seed
         # must use the same seed value so that the same subsets can be obtained
         # for each time the process is repeated
         seed = 7
         # split the whole dataset into folds
         '''In k-fold cross-validation, the original sample is randomly partition
         ed into k equal sized
         subsamples. Of the k subsamples, a single subsample is retained as the v
         alidation data for
         testing the model, and the remaining k-1 subsamples are used as traini
         ng data. The crossvalidation
         process is then repeated k times, with each of the k subsamples used exa
         ctly once as
         the validation data. The k results can then be averaged to produce a sin
         gle estimation. The
         advantage of this method over repeated random sub-sampling is that all o
         bservations are used for
         both training and validation, and each observation is used for validatio
         n exactly once
         kfold = KFold(n splits, random state=seed)
         # for logistic regression, we can use the accuracy level to evaluate the
         model / algorithm
         scoring = 'accuracy'
         # train the model and run K-fold cross validation to validate / evaluate
         results = cross val score(model, X, Y, cv=kfold, scoring=scoring)
         # print the evaluationm results
         # result: the average of all the results obtained from the K-fold cross
          validation
         print("Accuracy: %.3f (%.3f)" % (results.mean(), results.std()))
```

Accuracy: 0.880 (0.148)

/Users/jacobdenny/anaconda3/lib/python3.7/site-packages/sklearn/linear_model/logistic.py:432: FutureWarning: Default solver will be changed to 'lbfgs' in 0.22. Specify a solver to silence this warning.

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 using the 10-fold cross-validation to evaluate the model / algorithm, the accuracy of this logistic regression model is 88%

Above, the model predicted the flower type of the new record as Iris-virginica. According to the model's accuracy score obtained from the model evaluation using 10-told cross-validation there is only 88% chance that this new record is an Iris-virginica as opposed to 90% obtained by single sampling done earlier.

In []: