IRIS DATASET VISUALIZATION(SEABORN, MATPLOTLIB)

Importing libraries

```
In [29]: import numpy as np
   import pandas as pd
   import seaborn as sns
   import matplotlib.pyplot as plt
   import warnings
   warnings.filterwarnings('ignore')
```

Iris Flower

importing iris data set

In [34]: iris=pd.read_csv(r"C:/Users/user/Documents/Iris.csv")
 iris

Out[34]:		ld	SepalLengthCm	SepalWidthCm	PetalLengthCm	PetalWidthCm	Species
	0	1	5.1	3.5	1.4	0.2	lris- 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
	•••						
	145	146	6.7	3.0	5.2	2.3	lris- virginica
	146	147	6.3	2.5	5.0	1.9	lris- virginica
	147	148	6.5	3.0	5.2	2.0	lris- virginica
	148	149	6.2	3.4	5.4	2.3	lris- virginica
	149	150	5.9	3.0	5.1	1.8	lris- virginica

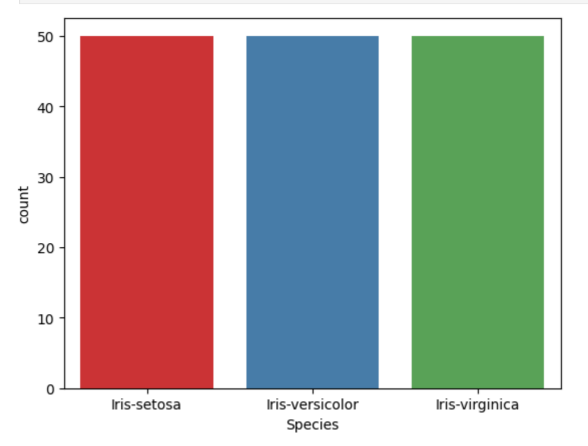
150 rows × 6 columns

^{***} Displaying Data ***

```
In [17]:
          iris.head()
Out[17]:
                 SepalLengthCm SepalWidthCm PetalLengthCm PetalWidthCm
             ld
                                                                                 Species
          0
              1
                            5.1
                                            3.5
                                                           1.4
                                                                          0.2 Iris-setosa
              2
                            4.9
                                            3.0
                                                           1.4
                                                                          0.2 Iris-setosa
          1
          2
              3
                            4.7
                                            3.2
                                                           1.3
                                                                          0.2 Iris-setosa
              4
                            4.6
                                            3.1
                                                           1.5
                                                                          0.2 Iris-setosa
              5
                            5.0
                                            3.6
                                                                          0.2 Iris-setosa
          4
                                                           1.4
         iris.drop('Id',axis=1,inplace=True) # axis=0 refers to rows. axis=1 refers to co
In [19]:
In [21]: iris.head()
Out[21]:
             SepalLengthCm SepalWidthCm PetalLengthCm PetalWidthCm
                                                                             Species
          0
                        5.1
                                        3.5
                                                        1.4
                                                                      0.2 Iris-setosa
          1
                         4.9
                                        3.0
                                                        1.4
                                                                      0.2 Iris-setosa
          2
                                        3.2
                        4.7
                                                        1.3
                                                                      0.2 Iris-setosa
          3
                         4.6
                                        3.1
                                                        1.5
                                                                      0.2 Iris-setosa
          4
                         5.0
                                        3.6
                                                        1.4
                                                                      0.2 Iris-setosa
          Checking if there are any missing values
In [24]: iris.info()
         <class 'pandas.core.frame.DataFrame'>
        RangeIndex: 150 entries, 0 to 149
        Data columns (total 5 columns):
         #
             Column
                             Non-Null Count Dtype
             ____
                             -----
         0
            SepalLengthCm 150 non-null
                                              float64
             SepalWidthCm 150 non-null
                                              float64
         1
                                              float64
         2
             PetalLengthCm 150 non-null
             PetalWidthCm
                                              float64
         3
                             150 non-null
             Species
                             150 non-null
                                              object
        dtypes: float64(4), object(1)
        memory usage: 6.0+ KB
In [26]: iris['Species'].value counts()
Out[26]: Species
          Iris-setosa
                              50
                              50
          Iris-versicolor
          Iris-virginica
                              50
          Name: count, dtype: int64
```

2.Bar Plot : Here the frequency of the observation is plotted. In this case we are plotting the frequency of the three species in the Iris Dataset





We can see that there are 50 samples each of all the Iris Species in the data set.

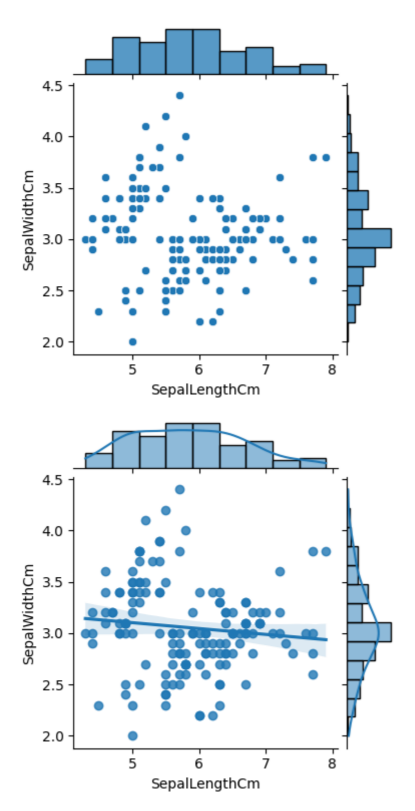
3. Joint Plot:

Jointplot is seaborn library specific and can be used to quickly visualize and analyze the relationship between two variables and describe their individual distributions on the same plot.

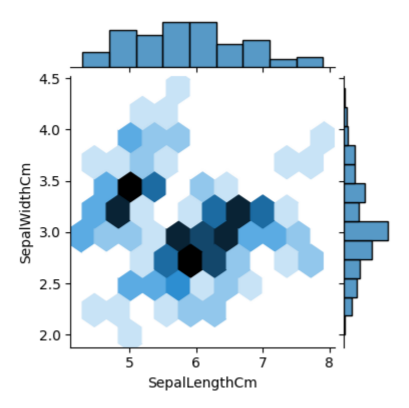
In [33]:	iris.head()	
----------	-------------	--

Out[33]:		SepalLengthCm	SepalWidthCm	PetalLengthCm	PetalWidthCm	Species
	0	5.1	3.5	1.4	0.2	Iris-setosa
	1	4.9	3.0	1.4	0.2	Iris-setosa
	2	4.7	3.2	1.3	0.2	Iris-setosa
	3	4.6	3.1	1.5	0.2	Iris-setosa
	4	5.0	3.6	1.4	0.2	Iris-setosa

```
In [35]: fig=sns.jointplot(x='SepalLengthCm',y='SepalWidthCm',data=iris,height=4)
In [36]: sns.jointplot(x="SepalLengthCm", y="SepalWidthCm", data=iris, kind="reg",height=plt.show()
```



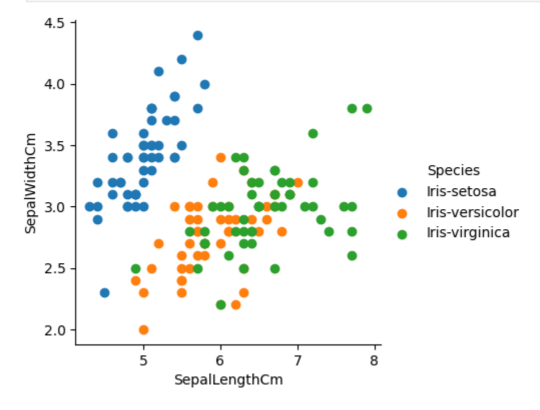
In [38]: fig=sns.jointplot(x='SepalLengthCm',y='SepalWidthCm',kind='hex',data=iris,height
plt.show()



4. FacetGrid Plot

```
import matplotlib.pyplot as plt
%matplotlib inline

sns.FacetGrid(iris,hue='Species',height=4)\
.map(plt.scatter,'SepalLengthCm','SepalWidthCm')\
.add_legend()
plt.show()
```



5. Boxplot or Whisker plot

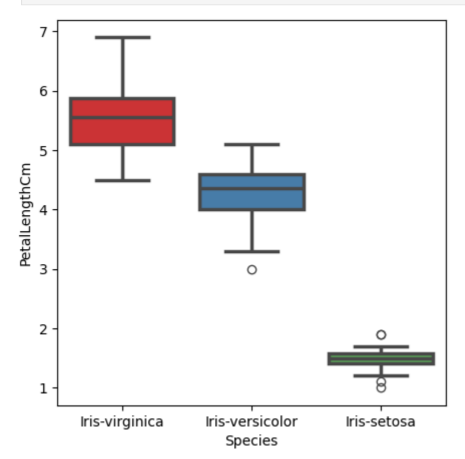
Box plot was was first introduced in year 1969 by Mathematician John Tukey.Box plot give a statical summary of the features being plotted.Top line represent the max value,top edge of box is third Quartile, middle edge represents the median,bottom edge represents the first quartile value.The bottom most line respresent the minimum value of the feature.The height of the box is called as Interquartile range.The black dots on the plot represent the outlier values in the data.

In [89]: iris.head()

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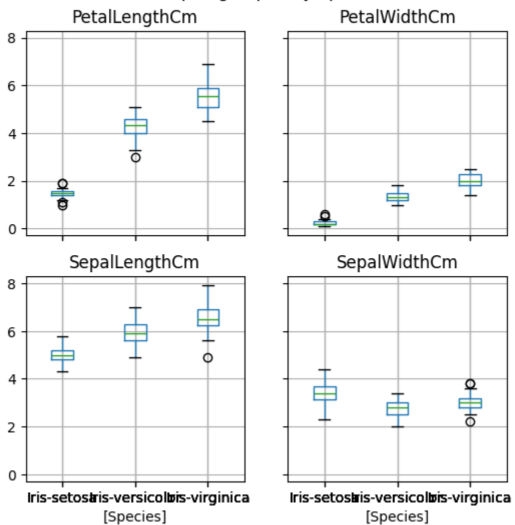
	SepalLengthCm	SepalWidthCm	PetalLengthCm	PetalWidthCm	Species
0	5.1	3.5	1.4	0.2	Iris-setosa
1	4.9	3.0	1.4	0.2	Iris-setosa
2	4.7	3.2	1.3	0.2	Iris-setosa
3	4.6	3.1	1.5	0.2	Iris-setosa
4	5.0	3.6	1.4	0.2	Iris-setosa

```
In [91]: fig=plt.gcf()
    fig.set_size_inches(5,5)
    fig=sns.boxplot(x='Species',y='PetalLengthCm',data=iris,order=['Iris-virginica',
    plt.show()
```



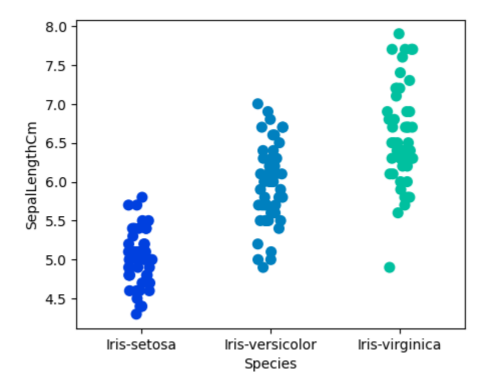
```
In [96]: iris.boxplot(by="Species", figsize=(6, 6))
plt.show()
```

Boxplot grouped by Species



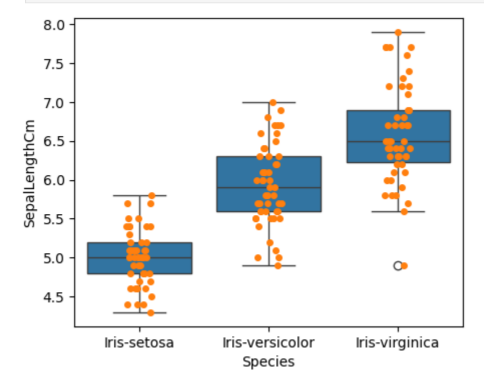
6. Strip plot

```
In [103... fig=plt.gcf()
    fig.set_size_inches(5,4)
    fig=sns.stripplot(x='Species',y='SepalLengthCm',data=iris,jitter=True,edgecolor=
    plt.show()
```



7. Combining Box and Strip Plots

```
In [45]: fig=plt.gcf()
    fig.set_size_inches(5,4)
    fig=sns.boxplot(x='Species',y='SepalLengthCm',data=iris)
    fig=sns.stripplot(x='Species',y='SepalLengthCm',data=iris,jitter=True,edgecolor=
    plt.show()
```



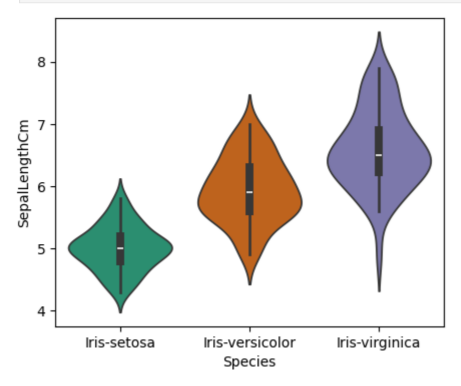
```
In []: ax= sns.boxplot(x="Species", y="PetalLengthCm", data=iris)
    ax= sns.stripplot(x="Species", y="PetalLengthCm", data=iris, jitter=True, edgeco
    boxtwo = ax.artists[2]
    boxtwo.set_facecolor('yellow')
    boxtwo.set_edgecolor('black')
```

```
boxthree=ax.artists[1]
boxthree.set_facecolor('red')
boxthree.set_edgecolor('black')
boxthree=ax.artists[0]
boxthree.set_facecolor('green')
boxthree.set_edgecolor('black')
```

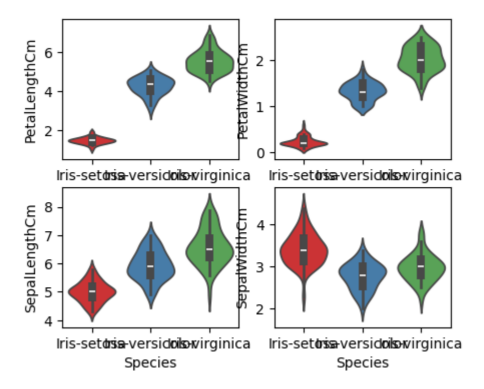
8. Violin Plot

It is used to visualize the distribution of data and its probability distribution. This chart is a combination of a Box Plot and a Density Plot that is rotated and placed on each side, to show the distribution shape of the data. The thick black bar in the centre represents the interquartile range, the thin black line extended from it represents the 95% confidence intervals, and the white dot is the median. Box Plots are limited in their display of the data, as their visual simplicity tends to hide significant details about how values in the data are distributed

```
fig=plt.gcf()
    fig.set_size_inches(5,4)
    fig=sns.violinplot(x='Species',y='SepalLengthCm',data=iris,palette="Dark2")
    plt.show()
```



```
In [133... plt.figure(figsize=(5,4))
    plt.subplot(2,2,1)# subplot(nrows, ncols, index, **kwargs)
    sns.violinplot(x='Species',y='PetalLengthCm',data=iris,palette="Set1")
    plt.subplot(2,2,2)
    sns.violinplot(x='Species',y='PetalWidthCm',data=iris,palette="Set1")
    plt.subplot(2,2,3)
    sns.violinplot(x='Species',y='SepalLengthCm',data=iris,palette="Set1")
    plt.subplot(2,2,4)
    sns.violinplot(x='Species',y='SepalWidthCm',data=iris,palette="Set1")
    plt.show()
```

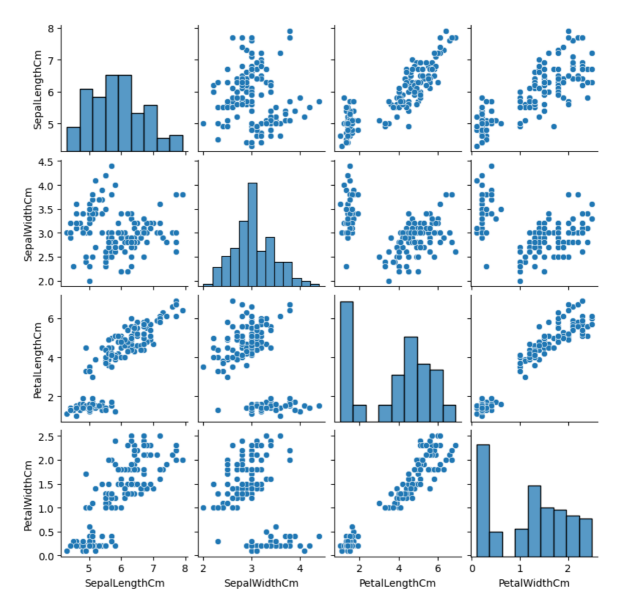


9. Pair Plot:

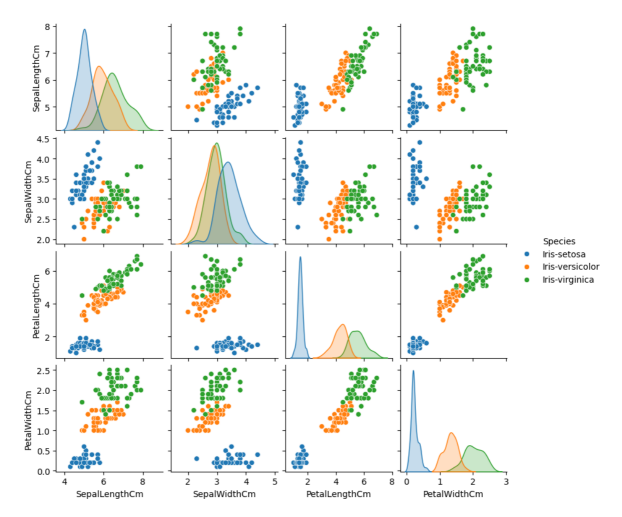
A "pairs plot" is also known as a scatterplot, in which one variable in the same data row is matched with another variable's value, like this: Pairs plots are just elaborations on this, showing all variables paired with all the other variables.

```
In [146... plt.figure(figsize=(5,4))
    sns.pairplot(data=iris,kind='scatter',height=2)
    plt.show()
```

<Figure size 500x400 with 0 Axes>



In [158... sns.pairplot(iris,hue='Species',kind='scatter',height=2)
 plt.show()



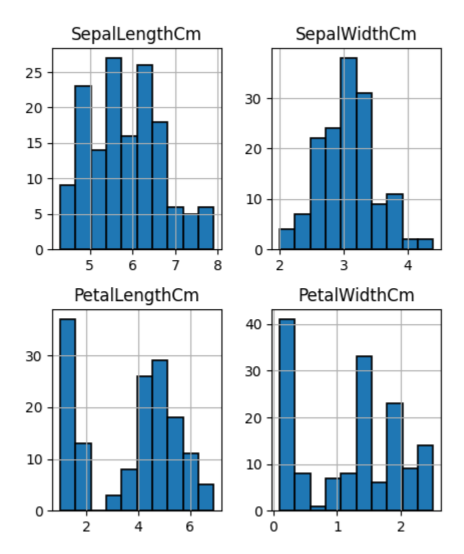
10. Heat map Heat map is used to find out the correlation between different features in the dataset. High positive or negative value shows that the features have high correlation. This helps us to select the parmeters for machine learning.

```
In [ ]: fig=plt.gcf()
    fig.set_size_inches(10,7)
    fig=sns.heatmap(iris.corr(),annot=True,cmap='cubehelix',linewidths=1,linecolor='
    plt.show()
```

12. Distribution plot:

The distribution plot is suitable for comparing range and distribution for groups of numerical data. Data is plotted as value points along an axis. You can choose to display only the value points to see the distribution of values, a bounding box to see the range of values, or a combination of both as shown here. The distribution plot is not relevant for detailed analysis of the data as it deals with a summary of the data distribution.

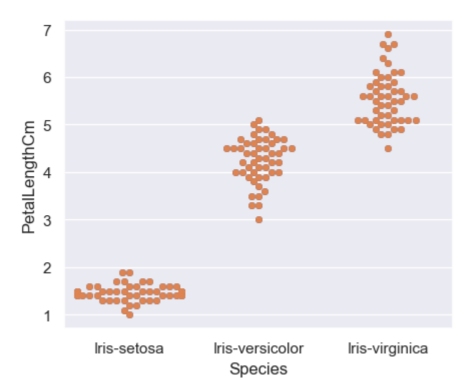
```
In [175... iris.hist(edgecolor='black', linewidth=1.2)
    fig=plt.gcf()
    fig.set_size_inches(5,6)
    plt.show()
```



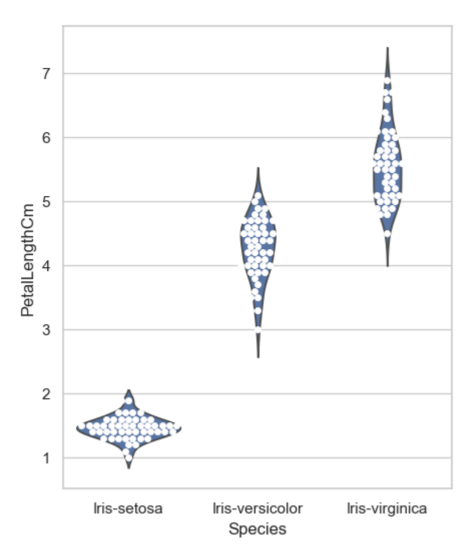
12. Swarm plot

It looks a bit like a friendly swarm of bees buzzing about their hive. More importantly, each data point is clearly visible and no data are obscured by overplotting. A beeswarm plot improves upon the random jittering approach to move data points the minimum distance away from one another to avoid overlays. The result is a plot where you can see each distinct data point, like shown in below plot

```
In [180... sns.set(style="darkgrid")
    fig=plt.gcf()
    fig.set_size_inches(5,4)
    fig = sns.swarmplot(x="Species", y="PetalLengthCm", data=iris)
    plt.show()
```

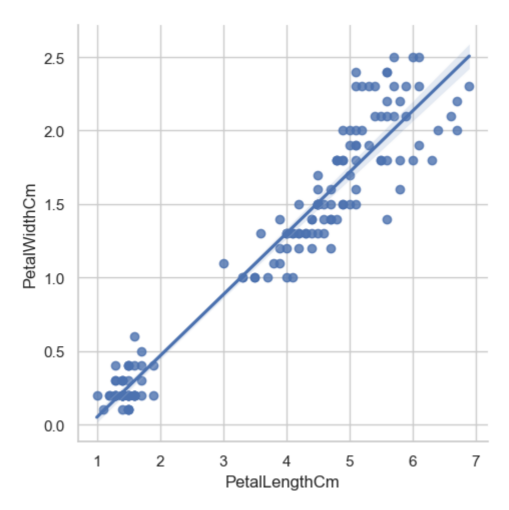


```
In [184...
sns.set(style="whitegrid")
fig=plt.gcf()
fig.set_size_inches(5,6)
ax = sns.violinplot(x="Species", y="PetalLengthCm", data=iris, inner=None)
ax = sns.swarmplot(x="Species", y="PetalLengthCm", data=iris,color="white", edge
plt.show()
```



13. LM PLot

```
In [192... fig=sns.lmplot(x="PetalLengthCm", y="PetalWidthCm",data=iris)
    plt.show()
```



14. FacetGrid

```
sns.FacetGrid(iris, hue="Species") \
In [202...
              .map(sns.kdeplot, "PetalLengthCm") \
              .add_legend()
           plt.ioff()
           plt.show()
             2.5
             2.0
             1.5
                                                     Species
                                                      Iris-setosa
             1.0
                                                      Iris-versicolor
                                                      Iris-virginica
             0.5
             0.0
                                       6
                                              8
                        2
                        PetalLengthCm
           ** 15. Factor Plot **
```

sns.factorplot('Species','SepalLengthCm',data=iris)

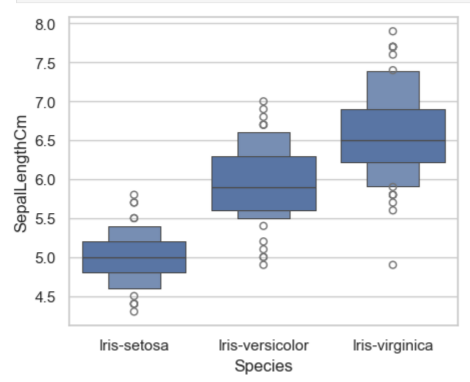
plt.ioff()

In []:

```
plt.show()
```

** 16. Boxen Plot**

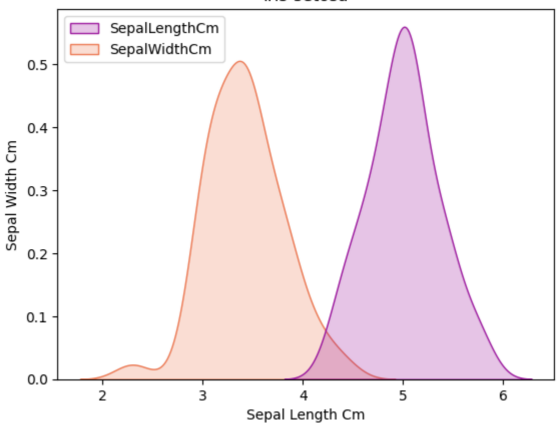
```
In [215... fig=plt.gcf()
    fig.set_size_inches(5,4)
    fig=sns.boxenplot(x='Species',y='SepalLengthCm',data=iris)
    plt.show()
```



17.KDE Plot

```
In [49]: sub=iris[iris['Species']=='Iris-setosa']
    sns.kdeplot(data=sub[['SepalLengthCm','SepalWidthCm']],palette="plasma", shade=T
    plt.title('Iris-setosa')
    plt.xlabel('Sepal Length Cm')
    plt.ylabel('Sepal Width Cm')
    plt.show()
```

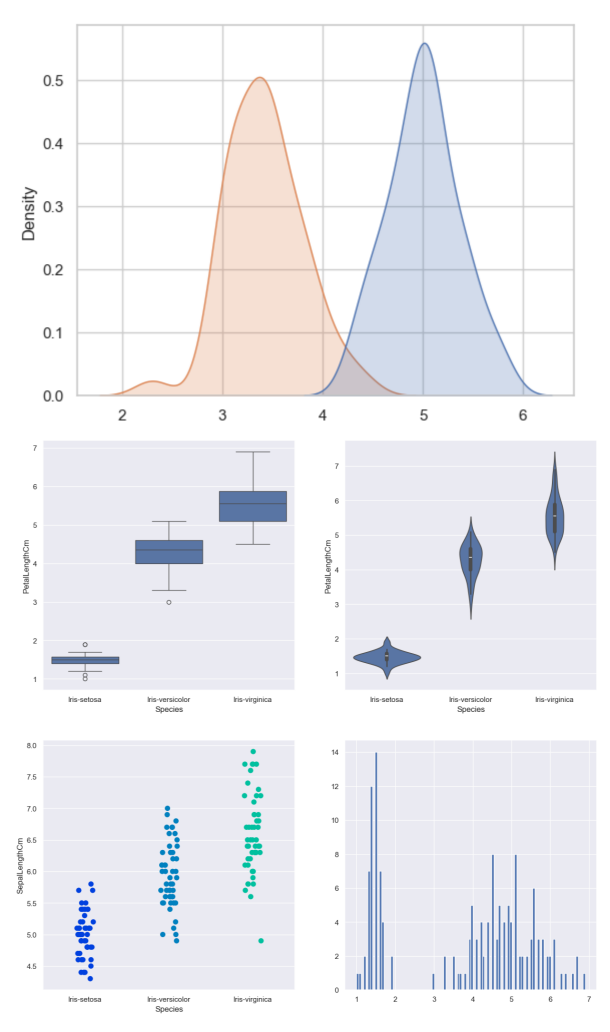
Iris-setosa



18.Dashboard

```
In [221...
sns.set_style('darkgrid')
f,axes=plt.subplots(2,2,figsize=(15,15))

k1=sns.boxplot(x="Species", y="PetalLengthCm", data=iris,ax=axes[0,0])
k2=sns.violinplot(x='Species',y='PetalLengthCm',data=iris,ax=axes[0,1])
k3=sns.stripplot(x='Species',y='SepalLengthCm',data=iris,jitter=True,edgecolor='#axes[1,1].hist(iris.hist,bin=10)
axes[1,1].hist(iris.PetalLengthCm,bins=100)
#k2.set(xlim=(-1,0.8))
plt.show()
```

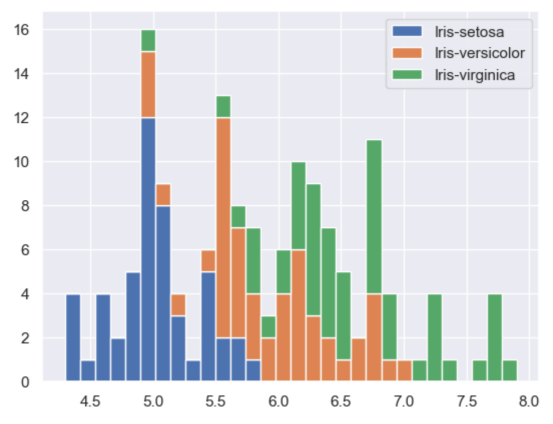


19.Stacked Histogram

```
In [224... iris['Species'] = iris['Species'].astype('category')

In [226... list1=list()
    mylabels=list()
    for gen in iris.Species.cat.categories:
        list1.append(iris[iris.Species==gen].SepalLengthCm)
        mylabels.append(gen)

    h=plt.hist(list1,bins=30,stacked=True,rwidth=1,label=mylabels)
    plt.legend()
    plt.show()
```



With Stacked Histogram we can see the distribution of Sepal Length of Different Species together. This shows us the range of Sepan Length for the three different Species of Iris Flower.

20.Area Plot: Area Plot gives us a visual representation of Various dimensions of Iris flower and their range in dataset.

```
iris.plot.area(y=['SepalLengthCm','SepalWidthCm','PetalLengthCm','PetalWidthCm']
plt.show()
```



In []: