Objective

dtype: object

The purpose of this notebook is to employ three unsupervised machine learning techniques to improve our comprehension of the morphological variations present within the Iris species.

This information can be valuable for various stakeholders, including botanists, researchers, and horticulturists, as it enhances our ability to categorize and interpret the diversity present in Iris flowers.

Import libraries and dataset

```
import pandas as pd
In [1]:
         import numpy as np
         import matplotlib.pyplot as plt
         import seaborn as sns
         from sklearn.cluster import KMeans, AgglomerativeClustering
         import scipy.cluster.hierarchy as sch
         from sklearn.mixture import GaussianMixture
         from sklearn.metrics import silhouette_score
In [2]: df = pd.read_csv('./Datasets/IRIS.csv')
In [3]: df.head()
Out[3]:
            sepal_length sepal_width petal_length petal_width
                                                                  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
        df.dtypes
In [4]:
Out[4]: sepal_length
                          float64
         sepal_width
                          float64
         petal_length
                          float64
         petal_width
                          float64
                           object
         species
```

There are 150 observations, with features including sepal length, sepal width, petal length, and petal width. The dataset comprises three types of Iris flowers: Iris-setosa, Iris-versicolor, and Iris-virginica, each with 50 instances.

Check for missing values

```
In [7]: df.isnull().sum()

Out[7]: sepal_length  0
    sepal_width  0
    petal_length  0
    petal_width  0
    species  0
    dtype: int64
```

There are no missing values.

Exploratory data analysis

Check for outliers

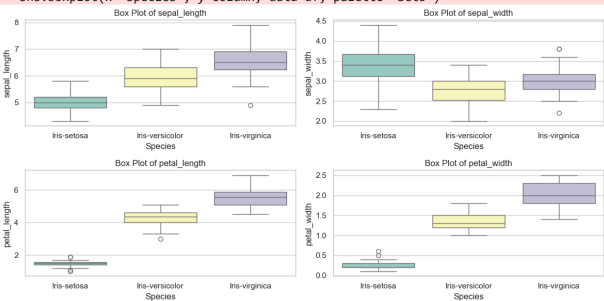
```
In [8]: # Box plots
sns.set(style="whitegrid")
plt.figure(figsize=(12, 6))
attributes = ['sepal_length', 'sepal_width', 'petal_length', 'petal_width']

for i, column in enumerate(attributes):
    plt.subplot(2, 2, i + 1)
    sns.boxplot(x='species', y=column, data=df, palette='Set3')

    plt.title(f'Box Plot of {column}')
    plt.xlabel('Species')
    plt.ylabel(column)
```

```
plt.tight_layout()
plt.show()
```

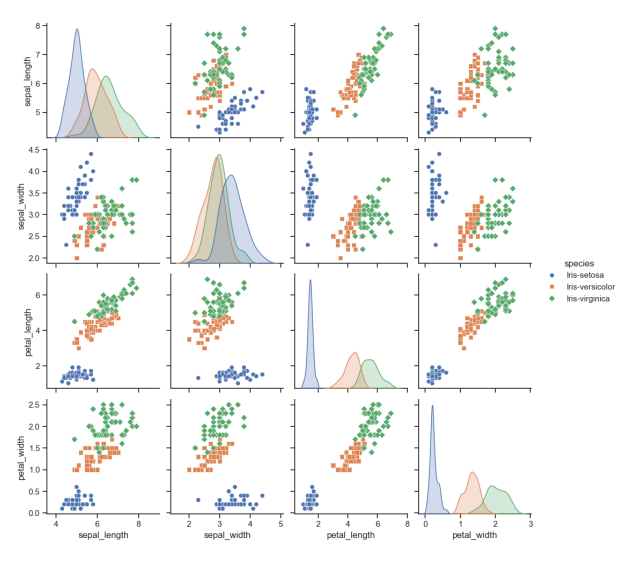
C:\Users\yashk\AppData\Local\Temp\ipykernel_12592\104556284.py:9: FutureWarning: Passing `palette` without assigning `hue` is deprecated and will be removed in v0.1 4.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect. sns.boxplot(x='species', y=column, data=df, palette='Set3') C:\Users\yashk\AppData\Local\Temp\ipykernel_12592\104556284.py:9: FutureWarning: Passing `palette` without assigning `hue` is deprecated and will be removed in v0.1 4.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect. sns.boxplot(x='species', y=column, data=df, palette='Set3') C:\Users\yashk\AppData\Local\Temp\ipykernel_12592\104556284.py:9: FutureWarning: Passing `palette` without assigning `hue` is deprecated and will be removed in v0.1 4.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect. sns.boxplot(x='species', y=column, data=df, palette='Set3') C:\Users\yashk\AppData\Local\Temp\ipykernel 12592\104556284.py:9: FutureWarning: Passing `palette` without assigning `hue` is deprecated and will be removed in v0.1 4.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect. sns.boxplot(x='species', y=column, data=df, palette='Set3')



Outliers are detected in the sepal length and sepal width of Iris-virginica, as well as in the petal length of Iris-setosa and Iris-versicolor, and the petal width of Iris-setosa.

Please note that the box plot for Iris-setosa, particularly in the petal length attribute, exhibits congestion due to the presence of outliers. Before proceeding with the replacement of outliers with the median, let's examine the quantiles.

```
print(setosa_quantiles)
          0.05
                    1.2
          0.95
                    1.7
          Name: petal_length, dtype: float64
In [10]: sns.set(style="ticks")
            sns.pairplot(df,markers=["o", "s", "D"])
            plt.show()
           sepal_length
              6
             5
            4.5
            4.0
         sepal_width 3.0
            2.5
            2.0
             6
           petal_length
            2.5 -
            2.0 -
          petal_width
            1.5
            1.0
            0.5
            0.0
                                                                         4
petal_length
                                      8
                                                                                           ò
                          6
                                                                                                  petal_width
                      sepal_length
                                                sepal_width
In [11]:
           sns.set(style="ticks")
            sns.pairplot(df,hue="species", markers=["o", "s", "D"])
            plt.show()
```



Use a combination of Winsorization and replacing outliers with median values.

```
attributes = ['sepal_length', 'sepal_width', 'petal_length', 'petal_width']
In [12]:
         species_with_outliers = {'Iris-setosa': ['petal_length', 'petal_width'],
                                   'Iris-versicolor': ['petal_length'],
                                   'Iris-virginica': ['sepal_length', 'sepal_width']}
         # Winsorize values for 'Iris-setosa' and 'petal_length'
         df.loc[(df['species'] == 'Iris-setosa') & (df['petal_length'] < 1.3), 'petal_length'</pre>
         df.loc[(df['species'] == 'Iris-setosa') & (df['petal_length'] > 1.7), 'petal_length'
         # Find and replace outliers with median for specific species and attributes
         for species, outlier_columns in species_with_outliers.items():
             for column in outlier_columns:
                 q_low = df[df['species'] == species][column].quantile(0.05)
                 q_high = df[df['species'] == species][column].quantile(0.95)
                 # Replace outliers with median
                 median_value = df[df['species'] == species][column].median()
                 df.loc[(df['species'] == species) & ((df[column] < q_low) | (df[column] > q
         # Box plots for each attribute with 'species' as hue after handling outliers
         sns.set(style="whitegrid")
```

```
plt.figure(figsize=(12, 6))
for i, column in enumerate(attributes):
    plt.subplot(2, 2, i + 1)
    sns.boxplot(x='species', y=column, data=df, palette='Set3')
    plt.title(f'Box Plot of {column} (Outliers replaced )')
    plt.xlabel('Species')
    plt.ylabel(column)
plt.tight_layout()
plt.show()
```

C:\Users\yashk\AppData\Local\Temp\ipykernel_12592\2944058266.py:26: FutureWarning:

Passing `palette` without assigning `hue` is deprecated and will be removed in v0.1 4.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

sns.boxplot(x='species', y=column, data=df, palette='Set3') C:\Users\yashk\AppData\Local\Temp\ipykernel_12592\2944058266.py:26: FutureWarning:

Passing `palette` without assigning `hue` is deprecated and will be removed in v0.1 4.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

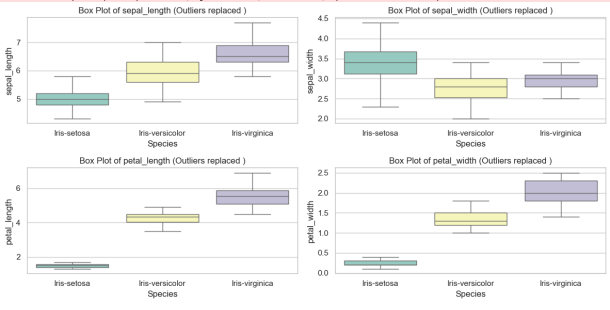
sns.boxplot(x='species', y=column, data=df, palette='Set3') C:\Users\yashk\AppData\Local\Temp\ipykernel_12592\2944058266.py:26: FutureWarning:

Passing `palette` without assigning `hue` is deprecated and will be removed in v0.1 4.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

sns.boxplot(x='species', y=column, data=df, palette='Set3') C:\Users\yashk\AppData\Local\Temp\ipykernel_12592\2944058266.py:26: FutureWarning:

Passing `palette` without assigning `hue` is deprecated and will be removed in v0.1 4.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

sns.boxplot(x='species', y=column, data=df, palette='Set3')



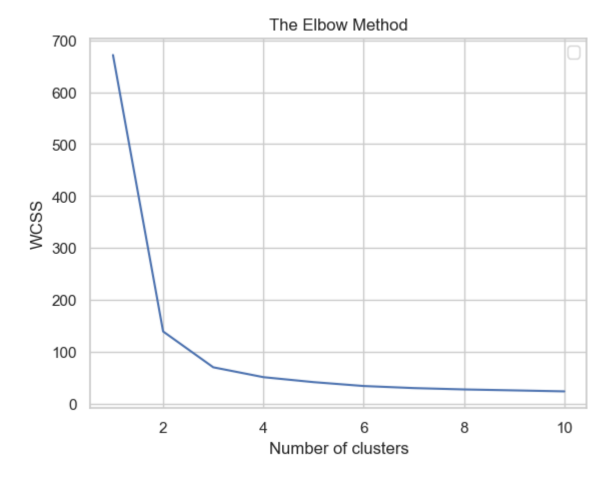
Define X (features)

In this notebook, I'll use three distinct clustering methods: K-means, hierarchical, and Gaussian Mixture Models (GMM).

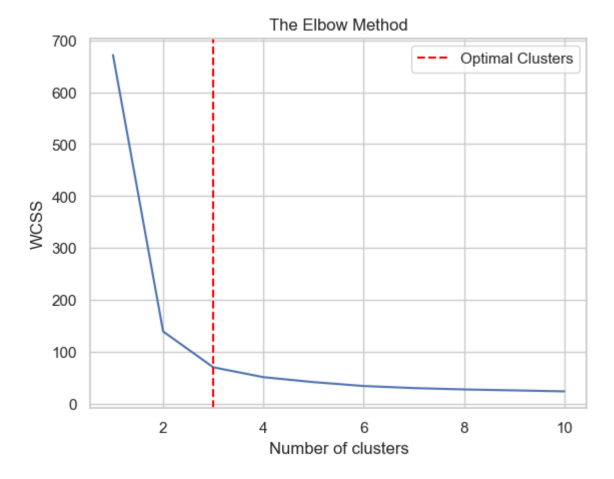
1. K-Means clustering

Define the number of clusters, K

No artists with labels found to put in legend. Note that artists whose label start with an underscore are ignored when legend() is called with no argument.



The elbow bend is evident at the 3rd cluster (as shown below), indicating the presence of three clusters in the dataset.

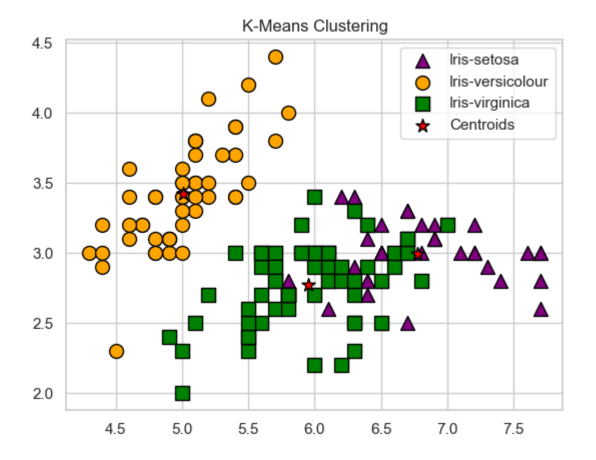


Training the K-Means model on the dataset

```
In [16]: kmeans = KMeans(n_clusters = 3, init = 'k-means++', max_iter = 300, n_init = 10, ra
y_kmeans = kmeans.fit_predict(x)
```

Visualising the clusters

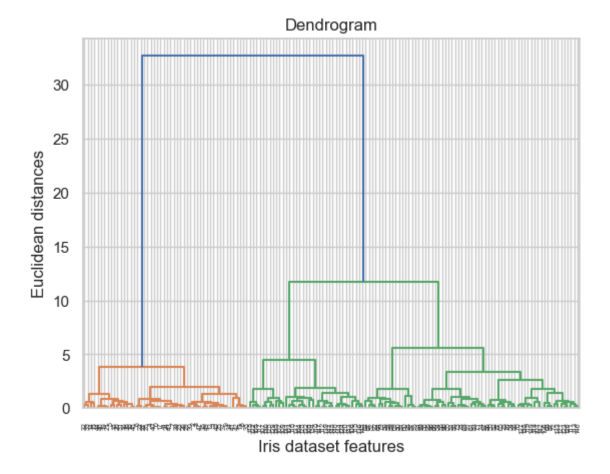
```
In [17]: plt.scatter(x.iloc[y_kmeans == 0, 0], x.iloc[y_kmeans == 0, 1], s=100, c='purple',
    plt.scatter(x.iloc[y_kmeans == 1, 0], x.iloc[y_kmeans == 1, 1], s=100, c='orange',
    plt.scatter(x.iloc[y_kmeans == 2, 0], x.iloc[y_kmeans == 2, 1], s=100, c='green', m
    # Plotting the centroids of the clusters
    plt.scatter(kmeans.cluster_centers_[:, 0], kmeans.cluster_centers_[:, 1], s=100, c=
    plt.title('K-Means Clustering')
    plt.legend()
    plt.show()
```



2. Hierarchical Clustering

Using the dendrogram to find the optimal number of clusters

```
In [18]: dendrogram = sch.dendrogram(sch.linkage(x, method = 'ward'))
    plt.title('Dendrogram')
    plt.xlabel('Iris dataset features')
    plt.ylabel('Euclidean distances')
    plt.show()
```

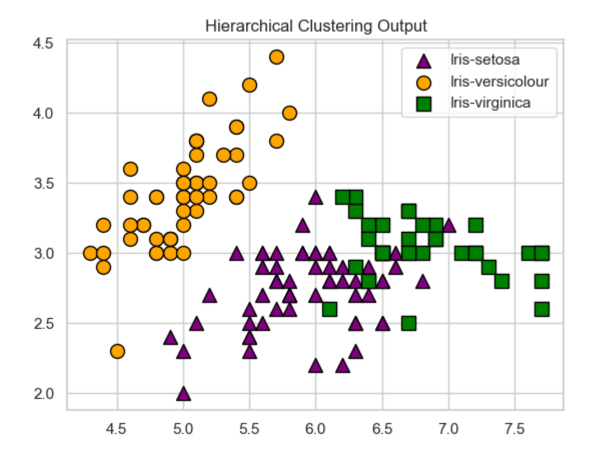


Training the Hierarchical Clustering model on the dataset

```
In [19]: hc = AgglomerativeClustering(n_clusters=3, linkage='ward', metric='euclidean')
y_hc = hc.fit_predict(x)
```

Visualising the clusters

```
In [20]: plt.scatter(x.iloc[y_hc == 0, 0], x.iloc[y_hc == 0, 1], s=100, c='purple', marker='
    plt.scatter(x.iloc[y_hc == 1, 0], x.iloc[y_hc == 1, 1], s=100, c='orange', marker='
    plt.scatter(x.iloc[y_hc == 2, 0], x.iloc[y_hc == 2, 1], s=100, c='green', marker='s
    plt.title('Hierarchical Clustering Output')
    plt.legend()
    plt.show()
```



3. Gaussian Mixture Model (GMM)

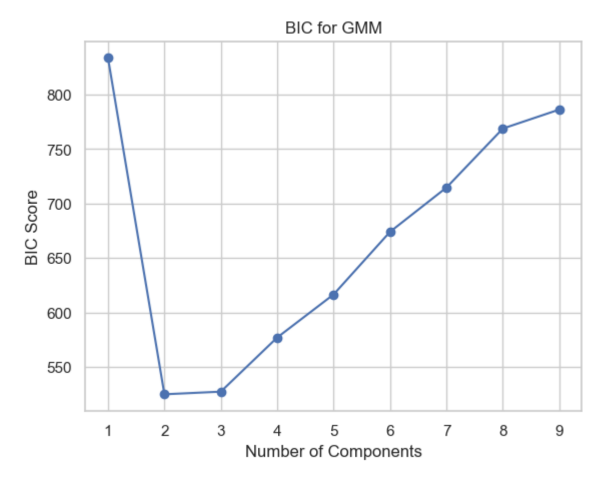
In the context of GMMs, Bayesian Information Criterion (BIC) is often employed to choose the optimal number of components (clusters) for the GMM.

Fit GMM models for different numbers of components and plot BIC

```
In [21]: n_components = range(1, 10)

bic = []
for n in n_components:
        gmm = GaussianMixture(n_components=n, random_state=42)
        gmm.fit(x)
        bic.append(gmm.bic(x))

# Plot the BIC values
plt.plot(n_components, bic, marker='o')
plt.xlabel('Number of Components')
plt.ylabel('BIC Score')
plt.title('BIC for GMM')
plt.show()
```



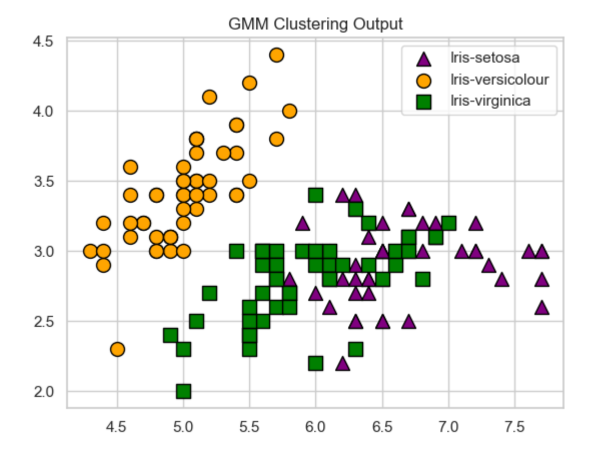
Choose the number of clusters based on the lowest BIC value. While in this example it suggests 2 clusters, remember our dataset has three clusters, so I'm selecting 3.

As a side note, for other datasets, follow the same logic and go with the lowest BIC for optimal clustering.

Training the GMM model on the dataset

Visualising the clusters

```
In [23]: plt.scatter(x.iloc[y_gmm == 0, 0], x.iloc[y_gmm == 0, 1], s=100, c='purple', marker
plt.scatter(x.iloc[y_gmm == 1, 0], x.iloc[y_gmm == 1, 1], s=100, c='orange', marker
plt.scatter(x.iloc[y_gmm == 2, 0], x.iloc[y_gmm == 2, 1], s=100, c='green', marker=
plt.title('GMM Clustering Output')
plt.legend()
plt.show()
```



Summary

In the Iris dataset with features like Sepal length, Sepal width, Petal length, and Petal width, we used three clustering algorithms: K-Means, Hierarchical Clustering, and GMM, to reveal distinct clusters.

```
In [24]: labels = list(df["species"].unique())
labels

Out[24]: ['Iris-setosa', 'Iris-versicolor', 'Iris-virginica']

In [25]: X = df.drop("species", axis=1)
y = df["species"]

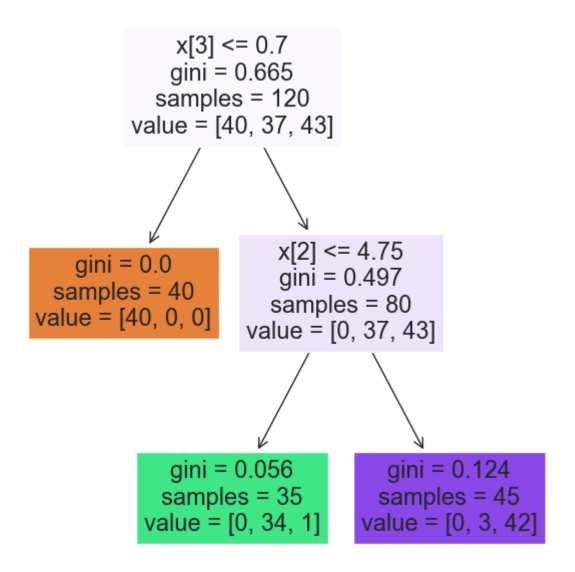
In [26]: from sklearn.model_selection import train_test_split
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2)
```

Post Pruning Decision Tree

```
In [27]: from sklearn.tree import DecisionTreeClassifier
dt = DecisionTreeClassifier()
dt.fit(X_train, y_train)
```

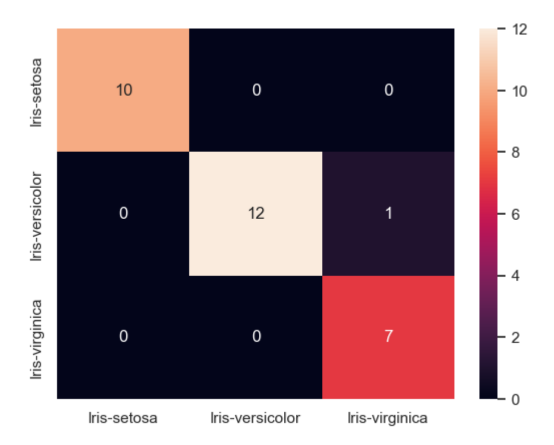
```
Out[27]:
           ▼ DecisionTreeClassifier
          DecisionTreeClassifier()
In [28]: from sklearn import tree
           plt.figure(figsize=(15,10))
           tree.plot_tree(dt, filled=True)
           plt.show()
                                           x[2] \le 2.6
                                           gini = 0.665
                                         samples = 120
                                       value = [40, 37, 43]
                                                    x[2] \le 4.75
                                  gini = 0.0
                                                    gini = 0.497
                                samples = 40
                                                    samples = 80
                               value = [40, 0, 0]
                                                  value = [0, 37, 43]
                                                                                  x[2] <= 4.95
                       x[3] <= 1.65
                       gini = 0.056
                                                                                  gini = 0.124
                                                                                 samples = 45
                      samples = 35
                     value = [0, 34, 1]
                                                                                value = [0, 3, 42]
                                x[2] \le 4.425
                                                                        x[3] \le 1.65
              gini = 0.0
                                                                        gini = 0.469
                                  gini = 0.5
            samples = 33
                                 samples = 2
                                                                        samples = 8
           value = [0, 33, 0]
                                                                                          value = [0, 0, 37]
                               value = [0, 1, 1]
                                                                       value = [0, 3, 5]
                                                                                  x[0] \le 5.95
                                            gini = 0.0
                        gini = 0.0
                                                                gini = 0.0
                                                                                  gini = 0.278
                       samples = 1
                                           samples = 1
                                                              samples = 2
                                                                                  samples = 6
                     value = [0, 1, 0]
                                         value = [0, 0, 1]
                                                             value = [0, 2, 0]
                                                                                 value = [0, 1, 5]
                                                                                  K
                                                                         gini = 0.0
                                                                                            samples = 5
                                                                        samples = 1
                                                                       value = [0, 1, 0]
                                                                                          value = [0, 0, 5]
In [29]: dt = DecisionTreeClassifier(max depth=2)
           dt.fit(X_train, y_train)
Out[29]:
                    DecisionTreeClassifier
          DecisionTreeClassifier(max depth=2)
In [30]:
           plt.figure(figsize=(7, 8))
           tree.plot_tree(dt, filled=True)
```

plt.show()



```
In [31]: from sklearn.metrics import accuracy_score, classification_report, confusion_matrix
y_pred = dt.predict(X_test)
score = accuracy_score(y_test, y_pred)
print("Post Pruning Score: ", score)
cm = confusion_matrix(y_test, y_pred)
sns.heatmap(cm, annot=True, fmt="d", xticklabels=labels, yticklabels=labels)
plt.show()
```

Post Pruning Score: 0.966666666666667



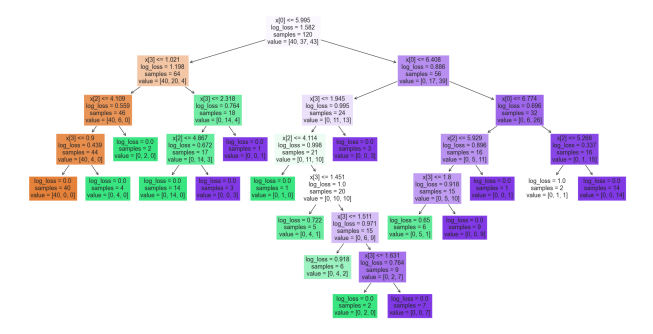
Pre Pruning

```
In [32]: from sklearn.model_selection import GridSearchCV

dt = DecisionTreeClassifier()
param_grid = {
        "criterion": ["gini", "entropy", "log_loss"],
        "splitter": ["best", "random"],
        "max_depth": [2,3,4,5,6,7,8,9,10],
        "min_samples_split": [2,3,4,5,6,7,7,8,9,10],
        "max_features": ["auto", "sqrt", "log2"],
}

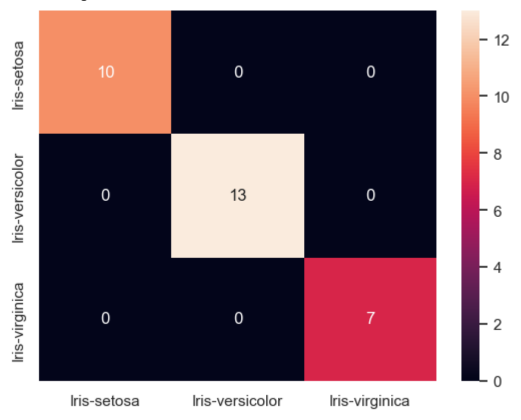
gscv = GridSearchCV(dt, param_grid, cv=5, scoring="accuracy")
gscv.fit(X_train, y_train)
```

```
c:\Users\yashk\AppData\Local\Programs\Python\Python311\Lib\site-packages\sklearn\mod
        el_selection\_validation.py:425: FitFailedWarning:
        2700 fits failed out of a total of 8100.
        The score on these train-test partitions for these parameters will be set to nan.
        If these failures are not expected, you can try to debug them by setting error_score
        ='raise'.
        Below are more details about the failures:
        2700 fits failed with the following error:
        Traceback (most recent call last):
          File "c:\Users\yashk\AppData\Local\Programs\Python\Python311\Lib\site-packages\skl
        earn\model_selection\_validation.py", line 729, in _fit_and_score
            estimator.fit(X_train, y_train, **fit_params)
          File "c:\Users\yashk\AppData\Local\Programs\Python\Python311\Lib\site-packages\skl
        earn\base.py", line 1145, in wrapper
            estimator. validate params()
          File "c:\Users\yashk\AppData\Local\Programs\Python\Python311\Lib\site-packages\skl
        earn\base.py", line 638, in _validate_params
            validate_parameter_constraints(
          File "c:\Users\yashk\AppData\Local\Programs\Python\Python311\Lib\site-packages\skl
        earn\utils\_param_validation.py", line 96, in validate_parameter_constraints
            raise InvalidParameterError(
        sklearn.utils._param_validation.InvalidParameterError: The 'max_features' parameter
        of DecisionTreeClassifier must be an int in the range [1, inf), a float in the range
        (0.0, 1.0], a str among {'sqrt', 'log2'} or None. Got 'auto' instead.
          warnings.warn(some_fits_failed_message, FitFailedWarning)
        c:\Users\yashk\AppData\Local\Programs\Python\Python311\Lib\site-packages\sklearn\mod
        el_selection\_search.py:979: UserWarning: One or more of the test scores are non-fin
                            nan nan ... 0.94166667 0.94166667 0.95
        ite: [
                    nan
         warnings.warn(
                      GridSearchCV
Out[32]: |
          ▶ estimator: DecisionTreeClassifier
                ▶ DecisionTreeClassifier
In [33]: print("Best params: ", gscv.best params )
         print("Best score: ", gscv.best_score_)
        Best params: {'criterion': 'log_loss', 'max_depth': 8, 'max_features': 'log2', 'min
        _samples_split': 9, 'splitter': 'random'}
        Best score: 0.9833333333333333
In [34]: from sklearn import tree
         dt = gscv.best_estimator_
         plt.figure(figsize=(23,12))
         tree.plot_tree(dt, filled=True)
         plt.show()
```



```
In [35]: y_pred = dt.predict(X_test)
    score = accuracy_score(y_test, y_pred)
    print("Pre-Pruning Score: ", score)
    cm = confusion_matrix(y_test, y_pred)
    sns.heatmap(cm, annot=True, fmt="d", xticklabels=labels, yticklabels=labels)
    plt.show()
```

Pre-Pruning Score: 1.0



In []: