# ML LAB: ASSIGNMENT7

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#### CSE-A

#### A7: Predicting Diabetes using decision tree

**Aim**: Develop a python program to predict diabetics using Decision Tree Model. Visualize the features from the dataset and interpret the results obtained by the model using Matplotlib library.

The Diabetes prediction dataset is a collection of medical and demographic data from patients, along with their diabetes status (positive or negative). The data includes features such as age, gender, body mass index (BMI), hypertension, heart disease, smoking history, HbA1c level, and blood glucose level. This dataset can be used to build machine learning models to predict diabetes in patients based on their medical history and demographic information. This can be useful for healthcare professionals in identifying patients who may be at risk of developing diabetes and in developing personalized treatment plans. Additionally, the dataset can be used by researchers to explore the relationships between various medical and demographic factors and the likelihood of developing diabetes.

import pandas as pd
import numpy as np
import seaborn as sns
from scipy.stats import ZSCOre
import matplotlib.pyplot as plt
from sklearn import tree
from sklearn.preprocessing import MinMaxScaler
from sklearn.metrics import classification\_report, roc\_auc\_score,
\_\_\_\_\_\_confusion\_matrix, ConfusionMatrixDisplay, RocCurveDisplay, roc\_auc\_score
from sklearn.model\_selection import train\_test\_split
from sklearn.neighbors import KNeighborsClassifier
from sklearn.naive\_bayes import MultinomialNB
from sklearn.tree import DecisionTreeClassifier
from sklearn import preprocessing
from sklearn.decomposition import PCA

#### import scipy.stats

from sklearn.metrics import PrecisionRecallDisplay from sklearn.metrics import confusion\_matrix, ConfusionMatrixDisplay from sklearn.neighbors import KNeighborsClassifier

```
pd.set_option("display.float_format", lambda x: "%.2f" \% x) %matplotlib inline
```

C:\Users\Nithi\AppData\Local\Temp\ipykernel\_13632\1512000984.py:1: DeprecationWarning: Pyarrow will become a required dependency of pandas in the next major release of pandas (pandas 3.0),

(to allow more performant data types, such as the Arrow string type, and better interoperability with other libraries) but was not found to be installed on your system. If this would cause problems for you,

import pandas as pd

count 100000.00

### 3 Loading dataset

[2]: data = pd\_read\_csv("C:/Users/ashwi/Downloads/ML Lab/A7/ ¬diabetes\_prediction\_dataset.csv
) [3]: data.head() gender age Female 80.00 hypertension heart disease [3]: age smoking\_history bmi \0 never 25.19 1 1 Female 54.00 0 0 No Info 27.32 2 Male 28.00 0 0 never 27.32 3 Female 36.00 0 0 current 23.45 4 Male 76.00 1 current 20.14 HbA1c level blood\_glucose\_level diabetes 0 140 6.60 0 1 6.60 80 0 2 5.70 0 158 3 5.00 0 155 4 4.80 155 0 [4]: data.describe() age bmi HbA1c level \ [4]: hypertension heart disease

100000.00

100000.00 100000.00

100000.00

mean	41.89	0.07	0.04	27.32	5.53
std	22.52	0.26	0.19	6.64	1.07
min	0.08	0.00	0.00	10.01	3.50
25%	24.00	0.00	0.00	23.63	4.80
50%	43.00	0.00	0.00	27.32	5.80
75%	60.00	0.00	0.00	29.58	6.20
max	80.00	1.00	1.00	95.69	9.00

```
blood_glucose_level
                              diabetes
                  100000.00 100000.00
count
                     138.06
                                   0.09
mean
std
                      40.71
                                   0.28
                      80.00
                                   0.00
min
25%
                     100.00
                                   0.00
```

[5]: #Find number of rows and columns

```
num_rows, num_columns = data.shape
print("Number of rows:", num_rows)
print("Number of columns:", num_columns)
```

Number of rows: 100000 Number of columns: 9

#### 4 Pre-Processing the data

[6]:

#Distribution

data["diabetes"].value\_counts()

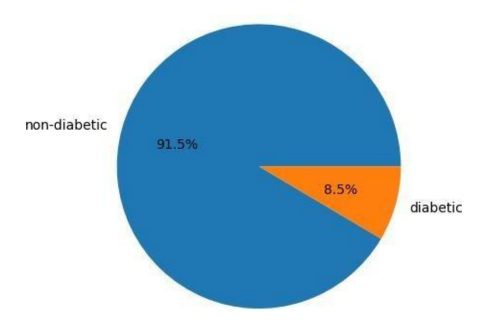
[6]: diabetes

915008500

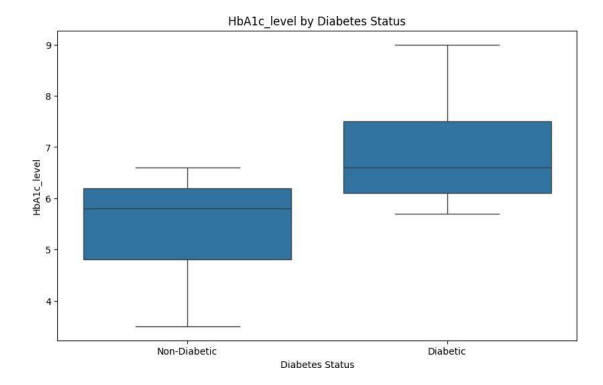
Name: count, dtype: int64

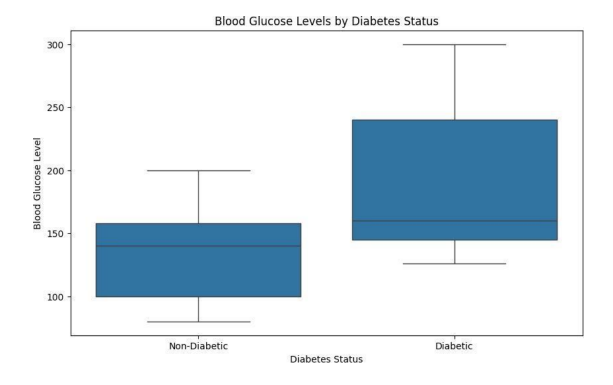
[7]: plt\_pie(data["diabetes"].value\_counts(), labels = ["non-diabetic", "diabetic"], autopct = "%1.1f%")
plt.title("Distribution of diabetics in dataset")
plt.show()

#### Distribution of diabetics in dataset



```
[8]: plt_figure(figsize=(10, 6))
sns_boxplot(x="diabetes", y="HbA1c_level", data=data)
plt_title("HbA1c_level by Diabetes Status")
plt_xlabel("Diabetes Status")
plt_ylabel("HbA1c_level")
plt_xticks([0, 1], ["Non-Diabetic", "Diabetic"]) # Adjust based on your_
a'diabetes' column encoding
plt.show()
```





```
[10]: #Find Missing Values
       missing_values = data.isnull().sum().sum()
       percentage_missing = (missing_values / data.shape[0]) * 100
print("Percentage of missing values:", percentage_missing)
[11]:
       #Convert Categorical variables into binary and numeric
       label_encoder = preprocessing.LabelEncoder()
       data["smoking_history"] = label_encoder_fit_transform(data["smoking_history"])
       data["gender"] = label_encoder_fit_transform(data["gender"])
       data.head()
[11]:
          0 80.00
                     0
                                                             4 25.19
                                                          0
                                                                             0 27.32
       1
                0 54.00
                                        0
       2
                                                                             4 27.32
                 1 28.00
                                        0
                                                          0
       3
                                        0
                                                          0
                                                                             1 23.45
                0 36.00
       4
                                        1
                                                          1
                1 76.00
                                                                             1 20.14
          HbA1c level blood glucose level
                                                   diabetes
       0
                   6.60
                   6.60
                                              80
                                                           0
       1
```

```
3
                  5.00
                                                         0
                                           155
       4
                  4.80
                                           155
                                                         0
       numeric_data = data_select_dtypes(include="number")
print(numeric_data)
[12]:
                    er age
0 80.00
                              hypertensionheart_disease
                                                           smoking_history
                                                                               bmi \0
              gender
                                                                               4 25.19
      1
                    0 54.00
                                          0
                                                           0
                                                                               0 27.32
      2
                                                           0
                    1 28.00
                                          0
                                                                               4 27.32
      3
                    0 36.00
                                          0
                                                           0
                                                                               1 23.45
      4
                    1 76.00
                                          1
                                                           1
                                                                               1 20.14
      99995
                    0 80.00
                                          0
                                                           0
                                                                               0 27.32
      99996
                    0 2.00
                                          0
                                                           0
                                                                               0 17.37
                                                           0
                                                                               3 27.83
      99997
                    1 66.00
                                          0
      99998
                    0 24.00
                                          0
                                                           0
                                                                               4 35.42
      99999
                    0 57.00
                                          0
                                                           0
                                                                               1 22.43
              HbA1c_level blood_glucose_level
                                                     diabetes
      0
                      6.60
                                               140
      1
                      6.60
                                                80
                                                             0
      2
                      5.70
                                               158
                                                             0
      3
                      5.00
                                               155
                                                             0
      4
                      4.80
                                               155
                                                             0
      99995
                                                90
                                                             0
                      6.20
      99996
                      6.50
                                                             0
                                               100
[13]: 99997
                                                             0
                      5.70
                                               155
      99998
                      4.00
                                               100
                                                             0
                                                             0
      99999
                      6.60
                                                90
[14]:
      # Calculate Z-scores for numeric columns
       # Selecting the numerical columns (excluding binary columns for hypertension,
        →heart_disease, diabetes)
       numerical_columns = ["age", "bmi", "HbA1c_level", "smoking_history",_
        ⇔"blood_glucose_level"]
```

158

0

2

5.70

age bmi HbA1c\_level smoking\_history blood\_glucose\_level0 1.69 -0.32 1.00 0.96 0.05

1	0.54 -0.00	1.00	-1.15	-1.43
2	-0.62 -0.00	0.16	0.96	0.49
3	-0.26 -0.58	-0.49	-0.62	0.42
4	1.52 -1.08	-0.68	-0.62	0.42
 Q:	 9995 1.69 -0.00	0.63	 -1.15	 -1.18
	9996 -1.77 -1.50	0.91	-1.15	-0.93
9	9997 1.07 0.08	0.16	0.43	0.42
9:	9998 -0.79 1.22	-1.43	0.96	-0.93
[15]: 9	9999 0.67 -0.74	1.00	-0.62	-1.18

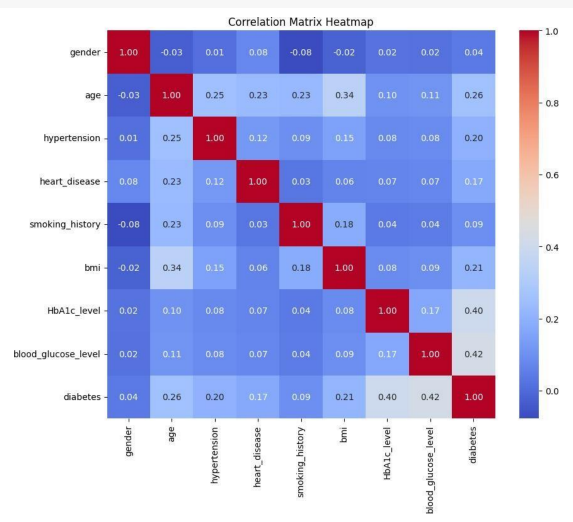
[100000 rows x 5 columns]

```
# Define threshold for outlier detection (e.g., Z-Score > 3)
threshold = 3
# Find outliers
outliers = data[z_scores > threshold]
print(outliers)
          INAIN INAIN
                             เงสเง
                                             IVAIN
                                                               ivaiv ivaiv
2
          NaN NaN
                             NaN
                                             NaN
                                                               NaN NaN
3
                                                               NaN NaN
          NaN NaN
                             NaN
                                             NaN
4
          NaN NaN
                             NaN
                                             NaN
                                                               NaN NaN
99995
                             NaN
                                                               NaN NaN
          NaN NaN
                                             NaN
99996
          NaN NaN
                             NaN
                                             NaN
                                                               NaN NaN
99997
          NaN NaN
                             NaN
                                             NaN
                                                               NaN NaN
99998
          NaN NaN
                             NaN
                                             NaN
                                                               NaN NaN
99999
          NaN NaN
                             NaN
                                             NaN
                                                               NaN NaN
       HbA1c_level blood_glucose_level
                                          diabetes
0
               NaN
                                     NaN
                                               NaN
1
               NaN
                                     NaN
                                               NaN
2
               NaN
                                     NaN
                                               NaN
3
               NaN
                                     NaN
                                               NaN
4
               NaN
                                     NaN
                                               NaN
99995
               NaN
                                               NaN
                                     NaN
                                               NaN
99996
               NaN
                                     NaN
99997
               NaN
                                     NaN
                                               NaN
99998
               NaN
                                     NaN
                                               NaN
99999
               NaN
                                     NaN
                                               NaN
[100000 rows x 9 columns]
```

```
[16]: # Count outliers
       outliers_count = (z_scores.abs() > threshold).sum().sum()
       print("Number of outliers:", outliers_count)
      Number of outliers. To Le
[17]: # normalization
       # Initialize the MinMaxScaler
       scaler = MinMaxScaler()
       # Fit the scaler to the data and transform it
       data[numerical_columns] = scaler.fit_transform(data[numerical_columns])
       # Display the first few rows to verify the normalization
       print(data.head())
                                                                   0.00 0.20
               1 0.35
      2
                                   0
                                                    0
                                                                   0.80 0.20
      3
               0 0.45
                                   0
                                                    0
                                                                   0.20 0.16
      4
               1 0.95
                                   1
                                                    1
                                                                   0.20 0.12
         HbA1c_level blood_glucose_level
                                              diabetes
      0
                 0.56
                                        0.27
                 0.56
                                        0.00
                                                      0
      1
                 0.40
                                        0.35
                                                      0
      2
[18]: 3
                 0.27
                                        0.34
                                                      0
                 0.24
                                        0.34
                                                      0
       data.head()
[18]:
          U 1.UU
                                    U
                                                        U.8U U.18
               0 0.67
                                                                    0.00 0.20
                                    0
                                                     0
       2
                                                                    0.80 0.20
               1 0.35
                                    0
                                                     0
       3
                                    0
                                                     0
               0 0.45
                                                                    0.20 0.16
       4
               1 0.95
                                    1
                                                     1
                                                                    0.20 0.12
          HbA1c_level
                       blood_glucose_level
                                               diabetes
       0
                 0.56
                                        0.27
                                                       0
       1
                 0.56
                                        0.00
                                                       0
       2
                 0.40
                                        0.35
                                                       0
       3
                 0.27
                                        0.34
                                                       0
```

4 0.24 0.34 0

### 5 Exploratory Data Analysis

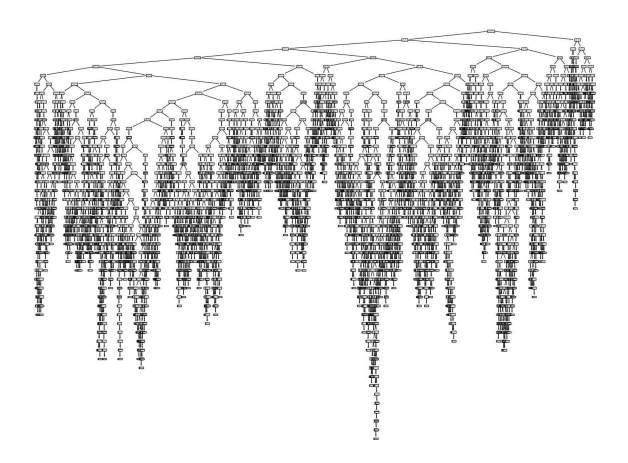


### 6 Split the data into training, testing and validation sets

```
[20]:
      X = data_drop(columns=['diabetes'], axis=1)
      y = data["diabetes"]
[21]:
     X_train_raw, X_test_raw, y_train, y_test = train_test_split(X, y, test_size=0.
        →30, random_state=42)
[22]:
      X_train = pd_DataFrame((X_train_raw), columns=X_train_raw_columns)
      X_test = pd_DataFrame((X_test_raw), columns=X_test_raw_columns)
      X_test.head()
             у<del>с</del>пист аус
0 0.16
[22]:
                            hypertensionneart_uisease_sinoking_nistory
      75721
                                                                      0.00 0.13
                                                       0
                                        0
      80184
                   0 0.04
                                       0
                                                       0
                                                                      0.00 0.13
      19864
                   1 0.79
                                       0
                                                       0
                                                                      0.60 0.18
      76699
                   0 0.02
                                       0
                                                       0
                                                                      0.80 0.09
                                                       0
      92991
                   0.0.41
                                       0
                                                                      1.00 0.35
              HbA1c level
                            blood alucose level
      75721
                     0.42
                            0.21
      80184
                     0.27
                                            0.30
      19864
                     0.00
                                            0.55
      76699
                     0.47
                                            0.21
      92991
                     0.49
                                            0.55
          Train the model
[23]:
      pca = PCA(random_state=42)
      pca.fit(X_train)
[23]: PCA(random state=42)
[24]: # Calculate the cumulative explained variance ratio
      cumulative_variance = pca.explained_variance_ratio_.cumsum()
      # Find the number of components needed for 90% variance
      n_components_90 = (cumulative_variance <= 0.90).sum()
      print("Number of components to explain 90% variance:", n_components_90)
[25]:
      pca = PCA(n_components=5, random_state=42)
      pca.fit(X_train)
```

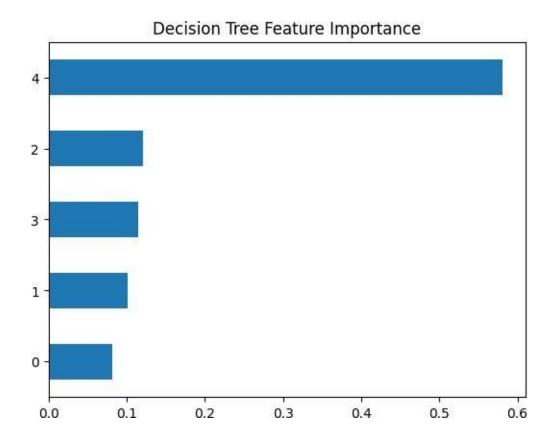
```
X_{train\_PCA} = pd.DataFrame(pca.transform(X_{train}))

X_{test\_PCA} = pd.DataFrame(pca.transform(X_{test}))
[26]:
        X_{train\_PCA.columns} = [str(column\_name) for column\_name in X_train\_PCA.columns] X_test\_PCA.columns = [str(column\_name) for column\_name in X_test\_PCA.columns]
[27]:
        X_train_PCA.head()
[27]:
                 0
                                  2
                                           3
                                                   4
               0 0.58 0.06 -0.01 -0.11 -0.06
               1 0.51 0.47 -0.09 -0.19 -0.33
                2 -0.37 -0.39 -0.13 0.07 -0.26
         3 -0.38 -0.22
                              0.20 - 0.17
                                               0.17
         4 0.66 -0.46 -0.13 0.12
                                               0.02
       8
               Test the model
[28]:
        model = tree_DecisionTreeClassifier()
[29]:
        model.fit(X_train_PCA, y_train)
[29]: DecisionTreeClassifier()
[30]:
        fig = plt_figure(figsize=(20,15))
tree.plot_tree(model);
```



```
[31]: feature_importances = pd.Series(model.feature_importances_, model.

_feature_names_in_).sort_values()
feature_importances.plot.barh();
plt.title("becision Tree Feature Importance");
```



y\_pred\_test = model.predict(X\_test\_PCA)
y\_pred\_train= model.predict(X\_train\_PCA)

## 9 Measure the performance of the trained model

[33]: train\_decision\_tree\_report = classification\_report(y\_train, y\_pred\_train) print(train\_decision\_tree\_report)

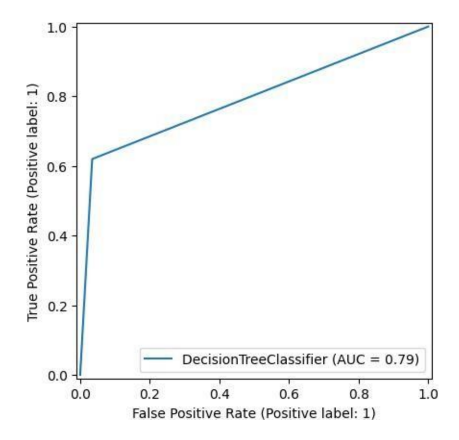
	precision	recall	f1-score	support
0	1.00	1.00	1.00	64047
1	1.00	0.99	1.00	5953
accuracy			1.00	70000
macro avg	1.00	1.00	1.00	70000
weighted avg	1.00	1.00	1.00	70000

[34]: test\_decision\_tree\_report = classification\_report(y\_test, y\_pred\_test) print(test\_decision\_tree\_report)

	precision	recall	f1-score	support
0	0.96	0.97	0.97	27453
1	0.62	0.62	0.62	2547
accuracy			0.94	30000
macro avg	0.79	0.79	0.79	30000
weighted avg	0.94	0.94	0.94	30000

## 10 Represent the results using graphs

[35]: RocCurveDisplay.from\_estimator(model, X\_test\_PCA, y\_test) plt.show()



[36]: from sklearn.tree import DecisionTreeClassifier from sklearn.metrics import accuracy\_score from sklearn.decomposition import PCA from sklearn.model\_selection import train\_test\_split import numpy as np

```
X_train, X_test, y_train, y_test = train_test_split(X_train_PCA, y_train,_

stest_size=0.2, random_state=42)

       dt_entropy = DecisionTreeClassifier(criterion="entropy", random_state=42)
       dt_entropy.fit(X_train, y_train)
       y_pred_entropy = dt_entropy.predict(X_test)
       accuracy_entropy = accuracy_score(y_test, y_pred_entropy)
       print("Accuracy using Entropy impurity measure with PCA:", accuracy_entropy)
       dt_gini = DecisionTreeClassifier(criterion="gini", random_state=42)
       dt_gini.fit(X_train, y_train)
       y_pred_gini = dt_gini.predict(X_test)
       accuracy_gini = accuracy_score(y_test, y_pred_gini)
       print("Accuracy using Gini-index impurity measure with PCA:", accuracy_gini)
      Accuracy using Entropy impurity measure with PCA: 0.9377857142857143 Accuracy using
      Gini-index impurity measure with PCA: 0.9399285714285714
[37]: from sklearn.tree import DecisionTreeClassifier
       from sklearn.metrics import accuracy_score
       dt_classifier = DecisionTreeClassifier(random_state=42)
       dt_classifier.fit(X_train, y_train)
      y_pred = dt_classifier.predict(X_test)
      accuracy_without_pca = accuracy_score(y_test, y_pred)
print("Accuracy without PCA:", accuracy_without_pca)
[44]:
      from sklearn.metrics import accuracy_score
       from sklearn.model_selection import train_test_split
       import pandas as pd
       dt_entropy = DecisionTreeClassifier(criterion="entropy", random_state=42)
       dt_entropy.fit(X_train_raw, y_train)
```

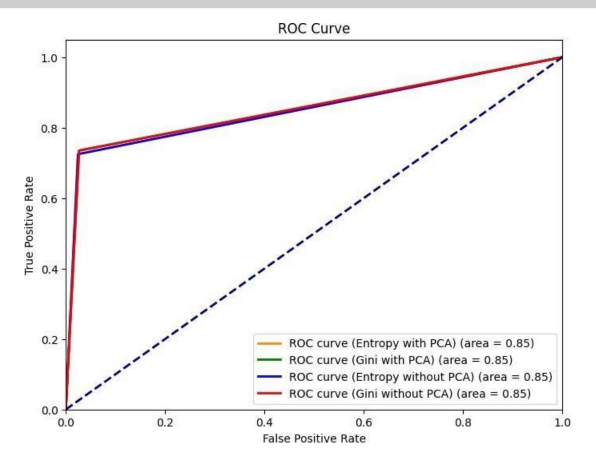
```
y_pred_entropy = dt_entropy.predict(X_test_raw)
       accuracy_entropy = accuracy_score(y_test, y_pred_entropy)
       print("Accuracy using Entropy impurity measure without PCA:", accuracy_entropy)
       dt_gini = DecisionTreeClassifier(criterion="gini", random_state=42)
       dt_gini.fit(X_train_raw, y_train)
       y_pred_gini = dt_gini.predict(X_test_raw)
       accuracy_gini = accuracy_score(y_test, y_pred_gini)
       print("Accuracy using Gini-index impurity measure without PCA:", accuracy_gini)
      Accuracy using Entropy impurity measure without PCA: 0.9542666666666667 Accuracy using
      Gini-index impurity measure without PCA: 0.953033333333333333
[43]: X = data_drop(columns=['diabetes'], axis=1) y =
       data['diabetes']
       X_train_raw, X_test_raw, y_train, y_test = train_test_split(X, y, test_size=0.
        →3, random state=42)
       print("X_train_raw shape:", X_train_raw.shape)
       print("y_train_shape:", y_train.shape)
print("X_test_raw_shape:", X_test_raw.shape)
       print("y_test_shape:", y_test.shape)
       dt entropy = DecisionTreeClassifier(criterion='entropy', random state=42)
       dt_entropy.fit(X_train_raw, y_train)
       y pred entropy = dt entropy.predict(X test raw) accuracy entropy =
       accuracy_score(y_test, y_pred_entropy)
       print("Accuracy using Entropy impurity measure without PCA:", accuracy entropy)
       dt_gini = DecisionTreeClassifier(criterion='gini', random_state=42) dt_gini.fit(X_train_raw,
       y_train)
       y_pred_gini = dt_gini.predict(X_test_raw) accuracy_gini =
       accuracy_score(y_test, y_pred_gini)
       print("Accuracy using Gini-index impurity measure without PCA:", accuracy_gini)
      X train raw shape: (70000, 8)
      y_train shape: (70000,)
      X test raw shape: (30000, 8)
```

```
[45]: from sklearn.metrics import roc_curve, auc
       import matplotlib.pyplot as plt
       y_pred_entropy_no_pca = dt_entropy.predict(X_test_raw)y_pred_gini_no_pca =
       dt gini.predict(X test raw)
       fpr_entropy_pca, tpr_entropy_pca, _ = roc_curve(y_test, y_pred_entropy)roc_auc_entropy_pca =
       auc(fpr_entropy_pca, tpr_entropy_pca)
       fpr_gini_pca, tpr_gini_pca, _ = roc_curve(y_test, y_pred_gini)roc_auc_gini_pca
       = auc(fpr_gini_pca, tpr_gini_pca)
       fpr_entropy_no_pca, tpr_entropy_no_pca, _ = roc_curve(y_test, _

    y_pred_entropy_no_pca)

       roc_auc_entropy_no_pca = auc(fpr_entropy_no_pca, tpr_entropy_no_pca)
       fpr_gini_no_pca, tpr_gini_no_pca, _ = roc_curve(y_test, y_pred_gini_no_pca)
       roc_auc_gini_no_pca = auc(fpr_gini_no_pca, tpr_gini_no_pca)
       # Plot ROC curve
       plt_figure(figsize=(8, 6))
       plt_plot(fpr_entropy_pca, tpr_entropy_pca, color='darkorange', lw=2, label='ROC ___
         curve (Entropy with PCA) (area = %0.2f)' % roc_auc_entropy_pca) plt.plot(fpr_gini_pca,
       tpr_gini_pca, color='green', lw=2, label='ROC curve__
         (Gini with PCA) (area = %0.2f)' % roc auc gini pca) plt.plot(fpr entropy no pca,
       tpr_entropy_no_pca, color='blue', lw=2, label='ROC ___
         curve (Entropy without PCA) (area = %0.2f)' % roc_auc_entropy_no_pca)
       plt.plot(fpr_gini_no_pca, tpr_gini_no_pca, color='red', lw=2, label='ROC curve__
         Gini without PCA) (area = %0.2f)' % roc_auc_gini_no_pca) plt.plot([0,
       1], [0, 1], color='navy', lw=2, linestyle='--')
       plt.xlim([0.0, 1.0])
       plt.xliff([0.0, 1.05])
plt.ylim([0.0, 1.05])
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title('ROC Curve')
plt.legend(loc="lower right")
```

#### plt.show()



[46]: print("X\_train\_raw shape:", X\_train\_raw.shape)
print("y\_train shape:", y\_train.shape)
print("X\_test\_raw shape:", X\_test\_raw.shape)
print("y\_test\_shape:", y\_test.shape)

λ\_train\_raw snape: (τυυυυ, δ)

y\_train shape: (70000,)

X\_test\_raw shape: (30000, 8)

y\_test shape: (30000,)

#### 11 Inference

- 1.Accuracy using Entropy impurity measure without PCA: 0.954266666666667 Accuracy using Gini-index impurity measure without PCA: 0.9530333333333333333
- 2. Accuracy using Entropy impurity measure with PCA: 0.9377857142857143

Accuracy using Gini-index impurity measure with PCA: 0.9399285714285714

As we can clearly see that the accuracy of the decision tree classifier using Gini-index and Entropyusing PCA gives less accuracy than without using PCA.

#### 12 Learning Outcomes

- 1. Applying standardization and other pre-processing techniques.
- 2. Splitting the data according to PCA.
- 3. Understanding Decision Tree classifier Algorithm.
- 4. Implementing decision tree classifier model.
- 5. Implementing the decision tree model with using PCA and

without using PCA.GITHUB LINK:

https://github.com/Anandh-007/Machine-learning-lab/tree/main/ML\_A7