## What is Python?

Python is a high-level, interpreted, and general-purpose programming language known for its simple and readable syntax. It was created by Guido van Rossum and first released in 1991.

Python is very popular among beginners and professionals because it is:

- Easy to learn and write
- Versatile (used in many fields)
- Open-source and has a large community
- Rich in libraries and frameworks

## **Uses of Python**

Python is used in a wide range of areas:

#### 1. Artificial Intelligence & Machine Learning

- Libraries: TensorFlow, PyTorch, Scikit-learn
- Used to build models that can learn and make predictions

#### 2. Web Development

- Frameworks: Django, Flask, FastAPI
- Used to create dynamic websites and REST APIs

#### 3. Data Science & Data Analysis

- Libraries: Pandas, NumPy, Matplotlib, Seaborn
- Used to analyze, visualize, and manipulate data

#### 4. Automation & Scripting

- Automate boring or repetitive tasks
- Example: Web scraping, file management, email sending

#### 5. Desktop Applications

- Libraries: Tkinter, PyQt
- Used to create GUI-based applications

#### 6. Game Development

- Libraries: Pygame
- Used to build simple 2D games

# What is NumPy?

NumPy (short for Numerical Python) is a powerful Python library used for numerical and scientific computing. It provides support for:

- Multidimensional arrays (ndarrays)
- · Mathematical functions
- Linear algebra
- Random number generation

NumPy is fast and efficient, and it's the foundation of many other libraries like Pandas, TensorFlow, and Scikit-learn.

#### **Key Features of NumPy**

- Powerful N-dimensional arrays
- Very **fast** operations (written in C under the hood)
- Supports **vectorized operations** (no need for loops)
- Tools for working with matrices, Fourier transforms, and statistics

#### What is Pandas?

**Pandas** is a powerful **open-source Python library** used for **data analysis and data manipulation**. It is built on top of **NumPy** and makes working with **structured data** (like tables or spreadsheets) easy and efficient.

#### **Core Data Structures in Pandas**

- 1. **Series** 1D labeled array (like a single column)
- 2. DataFrame 2D table with rows and columns (like an Excel sheet)

#### **Key Features of Pandas**

Feature	Description		
DataFrame Support	Handles labeled 2D data like SQL tables or Excel files		
Data Cleaning	Handle missing data, filter, replace, or fill values		
Data Aggregation	Grouping, summarizing, counting, mean, etc.		
Data Analysis	Perform statistical operations and data exploration		
File Handling	Easily read/write from CSV, Excel, JSON, SQL, etc.		
Feature	Description		
Time Series Handling	Built-in support for date/time functions and indexing		

**Data Visualization** Integrates well with libraries like Matplotlib and Seaborn for plotting **Fast Operations** Built on top of NumPy, optimized for performance

# What is Matplotlib?

**Matplotlib** is a Python library used for **data visualization**. It lets you create **line charts, bar charts, scatter plots**, and many other types of graphs.

#### **Key Features of Matplotlib**

Feature Description

Plotting Variety Line, bar, scatter, pie, histogram, etc. Customization Control over colors, labels, legends, etc. Subplots

Support Create multiple plots in one figure

Exporting Save plots as PNG, PDF, SVG, etc. Annotating Graphs

Add labels, arrows, and custom styling

#### What is Seaborn?

**Seaborn** is a Python visualization library built **on top of Matplotlib**. It provides a **high-level interface** for making beautiful and informative statistical graphics.

### **Key Features of Seaborn**

Feature Description

Easy Plotting One-liner plots like histograms and box plots
Statistical Graphs Built-in support for correlation and regression

Beautiful Themes Pre-set styles and color palettes Works with Pandas Supports DataFrames directly Complex Plots Made

Easy Heatmaps, pairplots, violin plots, etc.

# What is SciPy?

**SciPy** (Scientific Python) is a Python library used for **scientific and technical computing**. It builds on NumPy and adds advanced features.

# **Key Features of SciPy**

Feature Description

Advanced Math Integration, differentiation, optimization Statistics

Probability distributions, statistical tests Linear Algebra Matrix

operations, eigenvalues, etc.

Signal & Image Processing Tools for filtering, image analysis, etc.

Performance Fast calculations built in C/Fortran

# What is Scikit-learn (sklearn)?

**Scikit-learn** is a powerful **machine learning library** in Python. It provides simple tools for **classification**, **regression**, **clustering**, and more.

## **Key Features of Scikit-learn**

Description

Machine Learning Classification, regression, clustering Model Tools Training, testing, model evaluation

Preprocessing Feature scaling, encoding, normalization

Model Selection Cross-validation, hyperparameter tuning

Pipeline Support Combine multiple steps into one workflow

# 1. Write a Python program to calculate the Mean, Median, and Standard Deviation of a dataset using a NumPy array.

```
[1]: import numpy as np
[14]: marks = [23,23,12,45,50,22,21,45,34,36]
      print(type(marks))
      print(marks)
      marks = np.array(marks)
      print(marks)
      print(type(marks))
      <class 'list'>
      [23, 23, 12, 45, 50, 22, 21, 45, 34, 36]
      [23 23 12 45 50 22 21 45 34 36]
      <class 'numpy.ndarray'>
[8]: avg_marks = np.mean(marks)
      middle_marks = np.median(marks)
      stdv_marks = np.std(marks)
[12]: print(f"Marks:\nAverage: {avg_marks} \nMiddle_marks: {middle_marks} \nDeviation_marks: {stdv_marks}")
      Marks:
      Average: 31.1
      Middle_marks: 28.5
      Deviation_marks: 12.070211265756702
[]:
```

2)Write a Python program to read a CSV file, clean the data by handling missing values, and perform basic data analysis using the Pandas library.

# Cleaning of data

# Method1: Droping all the null value

```
[8]: med_data_final = medical_data.dropna()

[9]: med_data_final

[9]: Patient_ID Name Age Gender Diagnosis Blood_Pressure Heart_Rate Cholesterol
```

# Method1: Droping all the null value

[8]: med\_data\_final = medical\_data.dropna()

[9]: med\_data\_final

[9]: Patient\_ID Name Age Gender Diagnosis Blood\_Pressure Heart\_Rate Cholesterol 0 101 Alice 29.0 F Diabetes 120/80 78.0 190.0 102 Bob 45.0 85.0 M Hypertension 140/90 220.0

# Method2: Replacing with the Central tendencies

[10]: medical_dat	ta
-------------------	----

[10]: **Patient ID** Diagnosis **Blood Pressure Heart Rate Cholesterol** Name Age Gender 0 F 101 Alice 29.0 Diabetes 120/80 78.0 190.0 1 45.0 Hypertension 85.0 220.0 102 Bob 140/90 2 Charlie 90.0 103 34.0 Μ Asthma 110/70 NaN 3 104 David NaN Diabetes 0.88 240.0 Μ 135/85 4 105 41.0 **Heart Disease** 125/80 210.0 Eva NaN NaN 5 106 Frank 50.0 Μ NaN 145/95 0.08 200.0 6 107 Hypertension 230.0 NaN 38.0 NaN 76.0

```
[35]:
         Patient_ID
                      Name Age Gender
                                                        Blood_Pressure Heart_Rate Cholesterol
                                              Diagnosis
                101
                       Alice 29.0
                                        F
                                               Diabetes
                                                                               78.0
                                                                                          190.0
      0
                                                                 120/80
      1
                102
                        Bob 45.0
                                           Hypertension
                                                                 140/90
                                                                               85.0
                                                                                          220.0
      2
                     Charlie 34.0
                                                                                          205.0
                103
                                        Μ
                                                Asthma
                                                                 110/70
                                                                               90.0
      3
                104
                      David 39.8
                                        Μ
                                               Diabetes
                                                                 135/85
                                                                               0.88
                                                                                          240.0
      4
                105
                        Eva 41.0
                                        M
                                           Heart Disease
                                                                 125/80
                                                                               82.5
                                                                                          210.0
       5
                106
                       Frank 50.0
                                               Diabetes
                                                                 145/95
                                                                               0.08
                                                                                          200.0
                                        Μ
      7
                108 Hannah 39.8
                                        F
                                               Diabetes
                                                                 130/85
                                                                               72.0
                                                                                          180.0
[70]: #funciton to check is data is cleaned or not
      def Check_clean(data):
           d = dict(data.isnull().sum())
           null_values = []
           for key,value in d.items():
               if value != 0:
                   null_values.append((key,value))
           return 'Cleaned' if data.isnull().sum().sum() == 0 else null_value
      Check_clean(data)
[71]:
```

[71]: 'Cleaned'

# 3.) Create a Python program to plot a scatter plot, histogram, and boxplot using Matplotlib and Seaborn.

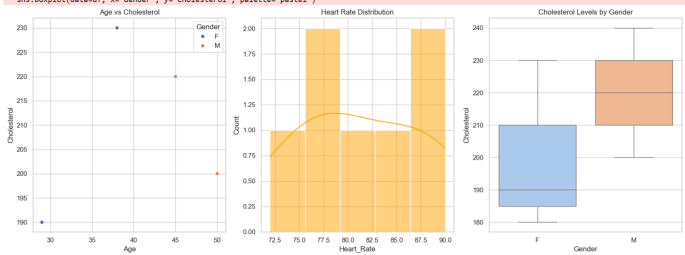
```
[5]: import numpy as np
      import pandas as pd
      import matplotlib.pyplot as plt
      import seaborn as sns
[6]: data = pd.read_csv("medical.csv")
[8]: df = pd.DataFrame(data)
[11]: def plotshow(data):
          # Set the plotting style
          sns.set(style="whitegrid")
          plt.figure(figsize=(16, 6))
          # Scatter Plot: Age vs Cholesterol
          plt.subplot(1, 3, 1)
          sns.scatterplot(data=df, x='Age', y='Cholesterol', hue='Gender')
          plt.title("Age vs Cholesterol")
          # Histogram: Heart Rate distribution
          plt.subplot(1, 3, 2)
          sns.histplot(df['Heart Rate'].dropna(), kde=True, bins=5, color='orange')
          plt.title("Heart Rate Distribution")
          # Boxplot: Cholesterol by Gender
          plt.subplot(1, 3, 3)
          sns.boxplot(data=df, x='Gender', y='Cholesterol', palette='pastel')
          plt.title("Cholesterol Levels by Gender")
          plt.tight_layout()
          plt.show()
```

#### [12]: plotshow(data)

C:\Users\Keshav Barawal\AppData\Local\Temp\ipykernel\_2124\3806559445.py:18: FutureWarning:

Passing `palette` without assigning `hue` is deprecated and will be removed in v0.14.0. Assign the `x` variable to `hue` and set `legend=False` for the same effect.

sns.boxplot(data=df, x='Gender', y='Cholesterol', palette='pastel')



4) Write Python Programme to implement linear regression.

```
[1]: import pandas as pd
      from sklearn.linear_model import LinearRegression
      from sklearn.model_selection import train_test_split
      from sklearn.metrics import mean_squared_error, r2_score
      import matplotlib.pyplot as plt
      import seaborn as sns
[2]: data = pd.read_csv("medical.csv")
[6]: df = pd.DataFrame(data)
[7]: # Drop rows with missing Age or Cholesterol for regression
      df_clean = df[['Age', 'Cholesterol']].dropna()
[8]: # Split into features and target
      X = df_clean[['Age']] # feature
      y = df_clean['Cholesterol'] # target
[9]:
      # Split into train and test sets
      X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_state=42)
[10]: # Linear Regression Model
      model = LinearRegression()
      model.fit(X_train, y_train)
```

```
[11]: # Predictions
      y_pred = model.predict(X_test)
[12]: # Evaluation
      print(f"Mean Squared Error: {mean_squared_error(y_test, y_pred):.2f}")
       print(f"R2 Score: {r2_score(y_test, y_pred):.2f}")
       Mean Squared Error: 113.78
       R<sup>2</sup> Score: nan
      C:\MY_PC\4thsem\Datamininglab\dm_env\Lib\site-packages\sklearn\metrics\_regression.py:1266: Undef
       less than two samples.
         warnings.warn(msg, UndefinedMetricWarning)
[13]: # Plotting
      plt.figure(figsize=(8, 6))
      sns.scatterplot(x='Age', y='Cholesterol', data=df_clean, color='blue', label='Actual')
      plt.plot(X_test, y_pred, color='red', label='Regression Line')
      plt.title('Linear Regression: Age vs Cholesterol')
      plt.xlabel('Age')
      plt.ylabel('Cholesterol')
      plt.legend()
      plt.grid(True)
      plt.show()
```

50



Age

30

35

5) Write a python programme to implement logistic regression.

```
[1]: import pandas as pd
     from sklearn.linear_model import LogisticRegression
     from sklearn.model_selection import train_test_split
     from sklearn.metrics import classification_report, confusion_matrix
     import seaborn as sns
     import matplotlib.pyplot as plt
[2]: data = pd.read_csv("medical.csv")
[3]: df = pd.DataFrame(data)
[4]: # Create target column: 1 if Diagnosis is Diabetes, else 0
     df['Diabetes'] = df['Diagnosis'].apply(lambda x: 1 if x == 'Diabetes' else 0)
[5]:
     # Keep only rows where we have no missing values for Age and Cholesterol
     df_clean = df[['Age', 'Cholesterol', 'Diabetes']].dropna()
[6]:
     # Features and target
     X = df_clean[['Age', 'Cholesterol']]
     y = df_clean['Diabetes']
[7]: # Split into train and test sets
     X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.3, random_state=42)
```

```
[8]: # Logistic Regression model
                     logreg = LogisticRegression()
                     logreg.fit(X_train, y_train)
    [8]: v LogisticRegression
                    LogisticRegression()
    [9]: # Predict
                     y_pred = logreg.predict(X_test)
 [10]: # Evaluation
                      print("Confusion Matrix:\n", confusion_matrix(y_test, y_pred))
                     print("\nClassification Report:\n", classification_report(y_test, y_pred))
                     Confusion Matrix:
                        [[2]]
                     Classification Report:
                                                                  precision
                                                                                                      recall f1-score support
                                                                             1.00
                                                                                                          1.00
                                                                                                                                        1.00
                                                                                                                                                                               2
                                                                                                                                        1.00
                                                                                                                                                                              2
                                 accuracy
                                                                             1.00
                                                                                                           1.00
                                                                                                                                        1.00
                              macro avg
                     weighted avg
                                                                                                           1.00
                                                                             1.00
                                                                                                                                        1.00
[11]:
                 # Optional: Heatmap of confusion matrix
                 plt.figure(figsize=(5, 4))
                 sns.heatmap(confusion_matrix(y_test, y_pred), annot=True, fmt='d', cmap='Blues')
                 plt.title("Confusion Matrix")
                 plt.xlabel("Predicted")
                 plt.ylabel("Actual")
                 plt.show()
                 C:\MY_PC\4thsem\Datamininglab\dm_env\Lib\site-packages\sklearn\metrics\_classification.py:407: UserWarning: A single like the control of the 
                 _pred'. For the confusion matrix to have the correct shape, use the 'labels' parameter to pass all known labels.
                   warnings.warn(
                                                                   Confusion Matrix
                                                                                                                                                                     - 2.20
                                                                                                                                                                     - 2.15
                                                                                                                                                                     - 2.10
                                                                                                                                                                     - 2.05
                                                                                                                                                                     - 2.00
                                                                                                                                                                    - 1.95
                                                                                                                                                                    - 1.90
```

- 1.85

- 1 20

6) Write python programme to implement decision tree.

```
[1]: import pandas as pd
     from sklearn.tree import DecisionTreeClassifier, plot_tree
     from sklearn.model_selection import train_test_split
     from sklearn.metrics import classification_report, accuracy_score
     import matplotlib.pyplot as plt
     import seaborn as sns
[2]: data = pd.read_csv("medical.csv")
[3]: df = pd.DataFrame(data)
[4]: # Target variable: 1 for Diabetes, 0 otherwise
     df['Diabetes'] = df['Diagnosis'].apply(lambda x: 1 if x == 'Diabetes' else 0)
[5]: # Clean data: drop rows with missing values in features
     df_clean = df[['Age', 'Cholesterol', 'Heart_Rate', 'Diabetes']].dropna()
[6]: # Features and target
     X = df_clean[['Age', 'Cholesterol', 'Heart_Rate']]
     y = df_clean['Diabetes']
[7]: # Train-test split
     X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.3, random_state=1)
```

```
[8]: # Create and train decision tree model
      clf = DecisionTreeClassifier(criterion='entropy', random_state=1)
      clf.fit(X_train, y_train)
 [8]:
                      DecisionTreeClassifier
      DecisionTreeClassifier(criterion='entropy', random_state=1)
 [9]: # Predict and evaluate
      y_pred = clf.predict(X_test)
[10]:
      print("Accuracy Score:", accuracy_score(y_test, y_pred))
      print("\nClassification Report:\n", classification_report(y_test, y_pred))
      Accuracy Score: 1.0
      Classification Report:
                    precision recall f1-score
                                                support
                        1.00
                                 1.00
                                          1.00
          accuracy
                                          1.00
                                                      2
                                          1.00
         macro avg
                        1.00
                                 1.00
                                                      2
                                          1.00
                      1.00
                                 1.00
      weighted avg
[11]: # Plot the decision tree
     plt.figure(figsize=(10, 6))
     plot_tree(clf, feature_names=['Age', 'Cholesterol', 'Heart_Rate'], class_names=['No Diabetes', 'Diabetes'], filled=True)
     plt.title("Decision Tree - Predicting Diabetes")
     plt.show()
                                 Decision Tree - Predicting Diabetes
                                    Age \leq 37.0
                                   entropy = 1.0
                                    samples = 2
                                   value = [1, 1]
                             class = No Diabetes
```

entropy = 0.0 samples = 1 value = [0, 1] class = Diabetes entropy = 0.0 samples = 1 value = [1, 0] class = No Diabetes 7) Write python programme to implement Ensemble Techniques.

```
[1]: import pandas as pd
      from sklearn.ensemble import RandomForestClassifier, GradientBoostingClassifier, VotingClassifier
      from sklearn.model_selection import train_test_split
      from sklearn.metrics import accuracy_score, classification_report
      from sklearn.preprocessing import LabelEncoder
      import numpy as np
[2]: data = pd.read_csv("medical.csv")
[15]: df = pd.DataFrame(data)
[16]: # Create target column: 1 if Diagnosis is 'Diabetes', else 0
      df['Diabetes'] = df['Diagnosis'].apply(lambda x: 1 if x == 'Diabetes' else 0)
[17]: # Drop rows with missing values in necessary columns
      df_clean = df[['Age', 'Gender', 'Cholesterol', 'Heart_Rate', 'Diabetes']].dropna()
[14]: # Encode Gender
      le = LabelEncoder()
      df_clean['Gender'] = le.fit_transform(df_clean['Gender']) # F=0, M=1
[13]: # Features and Target
      X = df_clean[['Age', 'Gender', 'Cholesterol', 'Heart_Rate']]
      y = df_clean['Diabetes']
[12]: # Split dataset
      X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.3, random_state=42)
```

```
[11]: # 1. Random Forest
      rf_model = RandomForestClassifier(n_estimators=100, random_state=42)
[19]: # 2. Gradient Boosting
      gb_model = GradientBoostingClassifier(n_estimators=100, random_state=42)
[21]: # 3. Voting Classifier (Hard Voting)
      voting_clf = VotingClassifier(estimators=[
          ('rf', rf_model),
          ('gb', gb_model)
      ], voting='hard')
[22]: # Train all models
      rf_model.fit(X_train, y_train)
      gb_model.fit(X_train, y_train)
      voting_clf.fit(X_train, y_train)
[22]: 🕨
                                    VotingClassifier
                       rf
                                                             gb
           RandomForestClassifier
                                             GradientBoostingClassifier
[23]: # Predictions
      rf_pred = rf_model.predict(X_test)
      gb_pred = gb_model.predict(X_test)
      voting_pred = voting_clf.predict(X_test)
```

```
# Results

print("Random Forest Accuracy:", accuracy_score(y_test, rf_pred))

print("Gradient Boosting Accuracy:", accuracy_score(y_test, gb_pred))

print("Voting Classifier Accuracy:", accuracy_score(y_test, voting_pred))

Random Forest Accuracy: 0.5

Gradient Boosting Accuracy: 0.5

Voting Classifier Accuracy: 0.5
```

[25]: print("\nVoting Classifier Classification Report:\n", classification\_report(y\_test, voting\_pred))

Voting Classifier Classification Report:

	precision	recall	f1-score	support
0	1.00	0.50	0.67	2
1	0.00	0.00	0.00	0
accuracy			0.50	2
macro avg	0.50	0.25	0.33	2
weighted avg	1.00	0.50	0.67	2

8) Write a python program to implement clustering.

```
[1]: import pandas as pd
    from sklearn.cluster import KMeans
    from sklearn.preprocessing import StandardScaler
    import matplotlib.pyplot as plt
    import seaborn as sns

[2]: data = pd.read_csv("medical.csv")

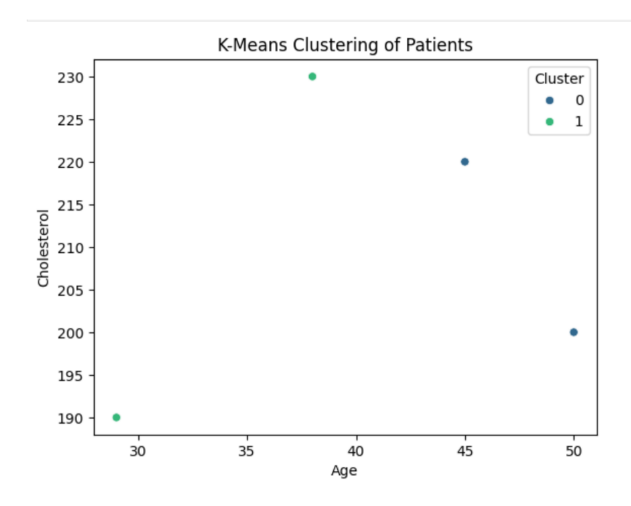
[3]: df = pd.DataFrame(data)

[4]: # Select features for clustering and handle missing values
    df_cluster = df[['Age', 'Cholesterol', 'Heart_Rate']].dropna()

[5]: # Standardize the features
    scaler = StandardScaler()
    scaled_features = scaler.fit_transform(df_cluster)

[7]: # Apply KMeans clustering
    kmeans = KMeans(n_clusters=2, random_state=42)
    df_cluster['Cluster'] = kmeans.fit_predict(scaled_features)
```

```
[8]: # Print clustered data
     print("Clustered Data:\n")
     print(df_cluster)
     Clustered Data:
         Age Cholesterol Heart_Rate Cluster
       29.0
                    190.0
                                 78.0
     1 45.0
                    220.0
                                  85.0
                                              0
     5 50.0
                    200.0
                                 80.0
                                             0
     6 38.0
                    230.0
                                 76.0
[9]: # Optional: Visualize the clusters
     sns.scatterplot(x='Age', y='Cholesterol', hue='Cluster', data=df_cluster, palette='v
     plt.title("K-Means Clustering of Patients")
     plt.xlabel("Age")
     plt.ylabel("Cholesterol")
     plt.show()
```



```
[1]: import pandas as pd
     from sklearn.model_selection import train_test_split
     from sklearn.preprocessing import StandardScaler
     from sklearn.svm import SVC
     from sklearn.metrics import classification report, confusion matrix
[2]: data = pd.read_csv("medical.csv")
[3]: df = pd.DataFrame(data)
[5]: # Create binary labels: 1 if Diagnosis is Diabetes, else 0
     df['Target'] = df['Diagnosis'].apply(lambda x: 1 if x == 'Diabetes' else 0)
[6]: # Select features and target, dropping rows with missing values
     features = df[['Age', 'Cholesterol', 'Heart_Rate']]
     df_model = pd.concat([features, df['Target']], axis=1).dropna()
[7]: X = df_model[['Age', 'Cholesterol', 'Heart_Rate']]
     y = df_model['Target']
[8]: # Split dataset
     X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.3, random_state=42)
[9]: # Standardize the features
     scaler = StandardScaler()
     X_train_scaled = scaler.fit_transform(X_train)
     X test scaled = scaler.transform(X test)
```

```
[10]: # Train SVM classifier
      svm_model = SVC(kernel='linear') # You can try 'rbf', 'poly' too
      svm_model.fit(X_train_scaled, y_train)
[10]:
            SVC
     SVC(kernel='linear')
[11]: # Predict
      y_pred = svm_model.predict(X_test_scaled)
[12]: # Evaluation
      print("Confusion Matrix:\n", confusion_matrix(y_test, y_pred))
      print("\nClassification Report:\n", classification_report(y_test, y_pred))
      Confusion Matrix:
       [[2]]
      Classification Report:
                    precision recall f1-score support
                0
                        1.00
                                 1.00
                                           1.00
                                                       2
                                           1.00
                                                       2
          accuracy
                        1.00
                                  1.00
                                           1.00
                                                       2
         macro avg
                                                       2
      weighted avg
                        1.00
                                 1.00
                                           1.00
```