

AI BASED DIABETES PREDICTION SYSTEM



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Al Based Diabetes Prediction System

Problem Definition

An Al-based diabetes prediction system is a valuable tool for identifying individuals at risk of developing diabetes or for helping manage the condition in those who already have it. Such a system typically relies on machine learning algorithms and data analysis to make predictions. Here's an overview of the steps involved in developing and deploying such a system to use:

Design Thinking

- . Data collection
- . Data Pre-processing
- . Exploratory Data Analytics
- . Feature Engineering
- . Model Evaluation
- . Deployment Selection

Data Collection:

Gather a comprehensive dataset containing medical records, patient demographics, lifestyle factors (e.g., diet, physical activity), family history, and clinical measurements (e.g., blood glucose levels, BMI) of individuals.

Data Preprocessing:

Clean and preprocess the dataset, handling missing values, outliers, and ensuring data consistency. Transform categorical variables into numerical representations if necessary.

Exploratory Data Analysis (EDA):

Conduct EDA to understand data distributions, relationships between features, and identify potential risk factors associated with diabetes. Visualization and statistical analysis will aid in feature selection.

Feature Engineering:

Create relevant features that can enhance the predictive capabilities of the model. This may involve deriving new features from existing data, considering interactions between variables, and selecting the most informative features.

Predictive Modelling with Logistic Regression:

Utilize logistic regression as the primary predictive modelling technique. Train the model on historical data to predict the probability of an individual developing diabetes based on selected features.

Model Evaluation:

Evaluate the logistic regression model using appropriate metrics, such as accuracy, precision, recall, F1-score, and the area under the Receiver Operating Characteristic (ROC-AUC) curve. Perform cross-validation to assess model generalization.

Deployment and Maintenance:

Determine the deployment strategy, such as integrating the model into electronic health records (EHR) systems, mobile applications, or web platforms.

Regularly update the model with new patient data to ensure its accuracy and relevance in predicting diabetes risk.

Implement data security measures and adhere to healthcare data regulations to protect patient privacy (e.g., HIPAA compliance).

Ethical Considerations:

Adhere to ethical standards and guidelines for healthcare AI, ensuring responsible handling of patient data and obtaining informed consent.

Implementation Procedure

Introduction

The goal of this document is to outline the steps for transforming the design of the Albased diabetes prediction system, as previously defined, into an innovative solution.

Building a complete Al-based diabetes prediction system involves multiple steps, including obtaining the dataset, preprocessing the data, and building a predictive model. Below, I'll provide a step-by-step guide with code examples for each stage of the project.

Prerequisite step:

As you very well know, there are a quite a few programming languages (like Python, R) and tools that can be used to machine learning projects like this. But the most popular one is python and I going to use Python to do this project.

Initially, I will setup the project environment, here are the steps I will follow:

- Download Miniconda (an environment management system for installing and maintaining software packages), we use this to ease the process of installing necessary libraries for our project.
- . Then I will create a repository and create a conda environment along with the necessary dependencies **Pandas, NumPy, Matplotlib and Scikit.**

conda create --prefix ./env pandas numpy matplotlib scikit-learn

. Activate the environment and install Jupyter notebook – an IDE to run our project.

conda activate c:\Users\tvaru\Desktop\sample_project_ 1\env

In **Jupyter notebook**, I will create a new python file and began the coding process.

conda install jupyter

Step 1: Obtaining the Dataset

The first step in any machine learning project is to obtain a dataset to work with. There are many online resources available to find the datasets suitable for our project. One such popular platform that hosts datasets is **Kaggle**. Since the goal of our project is to predict Diabetes, I search for diabetes patient's dataset, which is readily available as were already many such similar projects.

- 1. Go to the Kaggle dataset page you mentioned: Diabetes Data Set.
- 2. Click the "Download" button to get the dataset files.
- 3. Unzip the downloaded files to a directory on your local machine into your project repository.

This dataset has the following attributes: Pregnancies, Glucose, Blood Pressure, Skin Thickness, Insulin, BMI, Diabetes Pedigree Function and Age, It also contains the Outcome label which tells whether the patient has diabetes or not. Since there is a class label present by default, this dataset can be studied using Classification based algorithms.

Step 2: Data Preprocessing

Now that we have the dataset, we need to prepare it for analysis and modeling. This involves cleaning and organizing the data. Now, since I took this dataset from Kaggle, it is already fully pre-processed (i.e.) there are no missing values, all features have numerical values.

```
# Import necessary libraries
import pandas as pd
import numpy as np

# Load the dataset
data = pd.read_csv('diabetes_data.csv')

# Check the first few rows of the dataset
print(data.head())

# Check for missing values
print(data.isnull().sum())

# Split the data into features (X) and target variable (y)
X = data.drop('diabetes', axis=1)
```

```
y = data['diabetes']
```

In this step, we load the dataset, inspect the first few rows, and check for missing values. Since the data set contains the target class label, I can go ahead with classification or regression based approaches to build the model.

Step 3: Data Exploration (Optional)

Data exploration helps us understand the dataset better. We can create visualizations to gain insights into the data. We can visualize the data using the graphing library – **matplotlib**. I use the pairplot to understand the relationship between variables.

```
import matplotlib.pyplot asplt
import seaborn as sns

# Pairplot to visualize relationships between variables
sns.pairplot(data, hue='diabetes', diag_kind='kde')
plt.show()
```

Step 4: Data Splitting

The necessary step is to split the dataset into training and testing sets to train and evaluate our model. I use **Scikit's train_test_split** method to achieve this, the code for which as follows.

```
from sklearn.model_selection import train_test_split

# Split the data into training and testing sets (80% train, 20% test)

X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_state=42)
```

Step 5: Model Building (Logistic Regression)

Now, we'll build a logistic regression model to predict diabetes. As previously mentioned in the report, I am going to use Logistic Regression to build the model since

it is best suitable for datasets with class label, and also the class label has binary values, which in our case is true, since we have 0 or 1 for the Outcome.

The **Logistic Regression** method is found Scikit's linear_model sub module. After building and fitting the model to the training dataset, we predict the result of the testing dataset. And I evaluate the prediction using measures like accuracy, confusion matrix and classification report which standard metrics of classification based models.

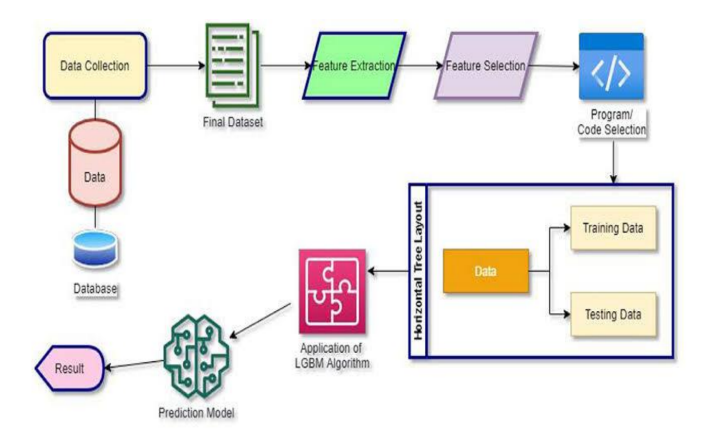
```
from sklearn.preprocessing import StandardScaler
from sklearn.linear_model import LogisticRegression
from sklearn.metrics import accuracy_score, classification_report,
confusion matrix
# Data preprocessing: Standardize features
scaler = StandardScaler()
X_{train} = scaler.fit_transform(X_train)
X_{test} = scaler.transform(X_{test})
# Create and train the logistic regression model
model = LogisticRegression()
model.fit(X_train,y_train)
# Make predictions on the test set
y_pred = model.predict(X_test)
# Evaluate the model
accuracy = accuracy_score(y_test, y_pred)
confusion = confusion_matrix(y_test, y_pred)
report = classification_report(y_test, y_pred)
print(f'Accuracy: {accuracy}')
```

print(f'Confusion Matrix:\n{confusion}')
print(f'Classification Report:\n{report}')

Step 6: Model Deployment (Optional)

To deploy the model, you would need to integrate it into a healthcare system, application, or web platform, which involves additional coding and considerations beyond the scope of this explanation.

Diabetes Graphical representation:



Data collection and Preprocessing:

Gather a dataset containing relevant information about individuals, including features such as age, preganacies, BMI, insulin, blood pressure, and glucose levels. Datasets like the Diabetes Database can be useful.

Data splitting:

Split the dataset into training and testing sets to evaluate your model's performance.

Model Training:

Train the selected model on the training data using appropriate algorithms.

Model selection:

Choose an appropriate machine learning or deep learning model for diabetes prediction. Common models include logistic regression, decision trees or random forests.

Evaluation Performance:

Evaluate the model's evaluation in given diabetes database using

Load the Dataset:

Load your dataset into a Pandas DataFrame. You can typically find kaggle Diabetes datasets in CSV format, but you can adapt this code to other formats as needed

To predict database using Diabetes

```
[1]: import pandas as pd
    import numpy as np
    import matplotlib.pyplot as plt
    import seaborn as sns
[2]: dataset = pd.read_csv('Documents/PS/diabetes.csv')
[4]: dataset.head()
                                                                   BMI
[4]:
       Pregnancies
                   Glucose BloodPressure SkinThickness
                                                          Insulin
                 6
                                       72
                                                               0 33.6
                       148
                                                      35
    1
                 1
                         85
                                      66
                                                      29
                                                               0 26.6
    2
                 8
                                       64
                                                     0
                                                              0 23.3
                        183
                 1
                                       66
                                                      23
                                                             94 28.1
                        89
                        137
                                       40
                                                      35
                                                             168 43.1
       DiabetesPedigreeFunction Age Outcome
    0
                         0.627 50
    1
                         0.351
                                 31
    2
                         0.672 32
                                           1
    3
                                           0
                         0.167
                                 21
                         2.288
                                 33
[5]: dataset.shape
[5]: (768, 9)
[6]: dataset.info()
   <class 'pandas.core.frame.DataFrame'>
   RangeIndex: 768 entries, 0 to 767
   Data columns (total 9 columns):
        Column
                                  Non-Null Count Dtype
                                  768 non-null
       Pregnancies
                                                 int64
                                 768 non-null
       Glucose
                                                int.64
                                 768 non-null int64
       BloodPressure
    3 SkinThickness
                                 768 non-null
                                                int64
```

```
4
   Insulin
                            768 non-null
                                           int64
                            768 non-null
                                           float64
5
   BMI
6
   DiabetesPedigreeFunction 768 non-null
                                           float64
                            768 non-null
                                           int64
   Age
                            768 non-null
                                           int64
8
   Outcome
```

dtypes: float64(2), int64(7)
memory usage: 54.1 KB

[7]: dataset.describe().T

	count	mean	std	min	25%	\
Pregnancies	768.0	3.845052	3.369578	0.000	1.00000	
Glucose	768.0	120.894531	31.972618	0.000	99.00000	
BloodPressure	768.0	69.105469	19.355807	0.000	62.00000	
SkinThickness	768.0	20.536458	15.952218	0.000	0.00000	
Insulin	768.0	79.799479	115.244002	0.000	0.00000	
BMI	768.0	31.992578	7.884160	0.000	27.30000	
DiabetesPedigreeFunction	768.0	0.471876	0.331329	0.078	0.24375	
Age	768.0	33.240885	11.760232	21.000	24.00000	
Outcome	768.0	0.348958	0.476951	0.000	0.00000	
	50)% 75	% max			
Pregnancies	3.000	6.0000	17.00			
Glucose	117.000	00 140.2500	0 199.00			
BloodPressure	72.000	80.0000	0 122.00			
SkinThickness	23.000	32.0000	0 99.00			
Insulin	30.500	00 127.2500	0 846.00			
BMI	32.000	36.6000	0 67.10			
	Glucose BloodPressure SkinThickness Insulin BMI DiabetesPedigreeFunction Age Outcome Pregnancies Glucose BloodPressure SkinThickness Insulin	Pregnancies 768.0 Glucose 768.0 BloodPressure 768.0 SkinThickness 768.0 Insulin 768.0 BMI 768.0 DiabetesPedigreeFunction 768.0 Age 768.0 Outcome 768.0 Pregnancies 3.000 Glucose 117.000 BloodPressure 72.000 SkinThickness 23.000 Insulin 30.500	Pregnancies 768.0 3.845052 Glucose 768.0 120.894531 BloodPressure 768.0 69.105469 SkinThickness 768.0 20.536458 Insulin 768.0 79.799479 BMI 768.0 31.992578 DiabetesPedigreeFunction 768.0 0.471876 Age 768.0 33.240885 Outcome 768.0 0.348958 Pregnancies 3.0000 6.00000 Glucose 117.0000 140.2500 BloodPressure 72.0000 80.0000 SkinThickness 23.0000 32.0000 Insulin 30.5000 127.2500	Pregnancies 768.0 3.845052 3.369578 Glucose 768.0 120.894531 31.972618 BloodPressure 768.0 69.105469 19.355807 SkinThickness 768.0 20.536458 15.952218 Insulin 768.0 79.799479 115.244002 BMI 768.0 31.992578 7.884160 DiabetesPedigreeFunction 768.0 0.471876 0.331329 Age 768.0 33.240885 11.760232 Outcome 768.0 0.348958 0.476951 ***Pregnancies* 3.0000 6.00000 17.00 Glucose 117.0000 140.25000 199.00 BloodPressure 72.0000 80.00000 122.00 SkinThickness 23.0000 32.00000 99.00 Insulin 30.5000 127.25000 846.00	Pregnancies 768.0 3.845052 3.369578 0.000 Glucose 768.0 120.894531 31.972618 0.000 BloodPressure 768.0 69.105469 19.355807 0.000 SkinThickness 768.0 20.536458 15.952218 0.000 Insulin 768.0 79.799479 115.244002 0.000 BMI 768.0 31.992578 7.884160 0.000 DiabetesPedigreeFunction 768.0 0.471876 0.331329 0.078 Age 768.0 33.240885 11.760232 21.000 Outcome 768.0 0.348958 0.476951 0.000 Fregnancies 3.0000 6.00000 17.00 17.00 140.25000 199.00 199.00 BloodPressure 72.0000 80.00000 122.00 199	Pregnancies 768.0 3.845052 3.369578 0.000 1.00000 Glucose 768.0 120.894531 31.972618 0.000 99.00000 BloodPressure 768.0 69.105469 19.355807 0.000 62.00000 SkinThickness 768.0 20.536458 15.952218 0.000 0.00000 Insulin 768.0 79.799479 115.244002 0.000 0.00000 BMI 768.0 31.992578 7.884160 0.000 27.30000 DiabetesPedigreeFunction 768.0 0.471876 0.331329 0.078 0.24375 Age 768.0 33.240885 11.760232 21.000 24.00000 Outcome 768.0 0.348958 0.476951 0.000 0.00000 Pregnancies 3.0000 6.00000 17.00 17.00 17.00 17.00 17.00 18.00 17.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 18.00 1

0.3725

29.0000

0.0000

0.62625

41.00000

1.00000

2.42

1.00

81.00

[8]: dataset.isnull().sum()

Age

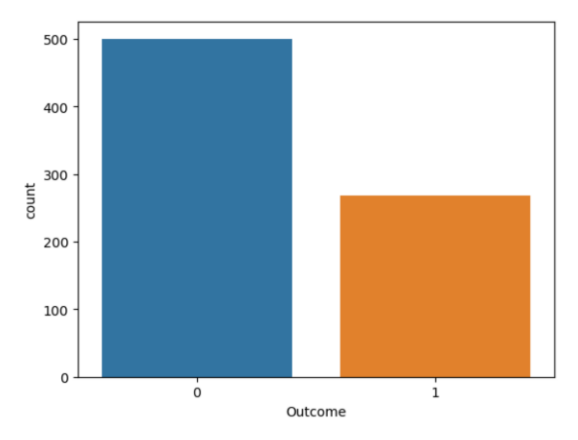
Outcome

[8]: Pregnancies 0 Glucose 0 BloodPressure 0 SkinThickness 0 Insulin 0 BMI 0 DiabetesPedigreeFunction Age 0 Outcome 0 dtype: int64

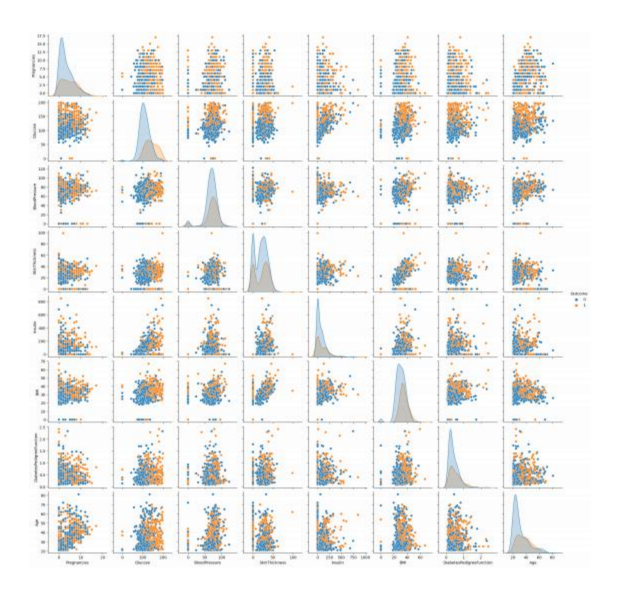
DiabetesPedigreeFunction

[9]: sns.countplot(x = 'Outcome', data = dataset)

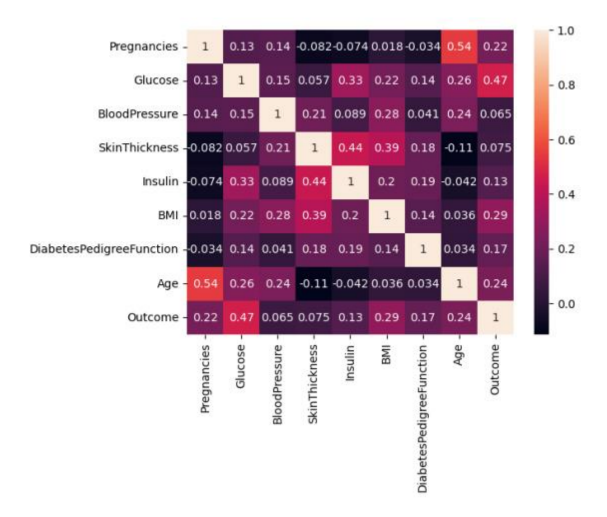
[9]: <Axes: xlabel='Outcome', ylabel='count'>



```
[12]: # Pairplot
    sns.pairplot(data = dataset, hue = 'Outcome')
    plt.show()
```



[13]: # Heatmap sns.heatmap(dataset.corr(), annot = True) plt.show()



```
[14]: # Replacing zero values with NaN
     dataset_new = dataset
     dataset_new[["Glucose", "BloodPressure", "SkinThickness", "Insulin", "BMI"]] = _
       -dataset_new[["Glucose", "BloodPressure", "SkinThickness", "Insulin", "BMI"]].
       replace(0, np. NaN)
[15]: # Count of NaN
     dataset_new.isnull().sum()
[15]: Pregnancies
                                    0
     Glucose
                                    5
     BloodPressure
                                   35
     SkinThickness
                                  227
     Insulin
                                  374
     BMI
                                   11
     DiabetesPedigreeFunction
                                    0
                                    0
     Age
```

```
dtype: int64
[16]: # Replacing NaN with mean values
      dataset_new["Glucose"] . fillna(dataset_new["Glucose"] . mean(), inplace = True)
      dataset_new["BloodPressure"] . fillna(dataset_new["BloodPressure"] . mean(), _
       inplace = True)
      dataset_new["SkinThickness"] . fillna(dataset_new["SkinThickness"] . mean(), _
       inplace = True)
      dataset_new["Insulin"] . fillna(dataset_new["Insulin"] . mean(), inplace = True)
      dataset new["BMI"] . fillna(dataset new["BMI"] . mean(), inplace = True)
[17]: dataset_new.isnull().sum()
[17]: Pregnancies
                                   0
      Glucose
                                   0
      BloodPressure
                                   0
                                   0
      SkinThickness
      Insulin
                                   0
      BMI
      DiabetesPedigreeFunction
                                   0
      Age
      Outcome
                                   0
      dtype: int64
[18]: y = dataset_new['Outcome']
      X = dataset_new.drop('Outcome', axis=1)
[19]: # Splitting X and Y
      from sklearn.model selection import train_test_split
      X_train, X_test, Y_train, Y_test = train_test_split(X, y, test_size = 0.20, __
       random_state = 42, stratify = dataset_new['Outcome'] )
[20]: from sklearn.linear model import LogisticRegression
      model = LogisticRegression()
      model.fit(X_train, Y_train)
      y_predict = model.predict(X_test)
     C: \text{Users}\text{CSE}_BAY4\text{\text{anaconda}}\text{alib}\text{\text{site}}-
     packages\sklearn\linear_model\left\_logistic.py:460: ConvergenceWarning: lbfgs failed
     to converge (status=1):
     STOP: TOTAL NO. of ITERATIONS REACHED LIMIT.
     Increase the number of iterations (max_iter) or scale the data as shown in:
         https://scikit-learn.org/stable/modules/preprocessing.html
     Please also refer to the documentation for alternative solver options:
         https://scikit-learn.org/stable/modules/linear_model.html#logistic-
```

0

Outcome

```
regression
      n iter i = check optimize result(
[21]: y predict
[21]: array([1, 0, 0, 0, 0, 0, 1, 0, 1, 0, 0, 0, 0, 0, 0, 1, 0, 1, 0, 1,
            0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1, 1, 0, 0, 0, 1, 1, 0, 0, 0,
            0, 0, 0, 0, 1, 0, 0, 1, 0, 0, 1, 0, 1, 0, 1, 0, 1, 0, 0, 1, 0, 0,
            1, 0, 0, 1, 0, 0, 0, 1, 0, 1, 0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0,
            0, 0, 1, 1, 0, 0, 0, 1, 0, 1, 1, 0, 0, 0, 0, 0, 1, 0, 1, 0, 1, 1,
            0, 1, 0, 0, 0, 0, 0, 1, 0, 1, 0, 1, 0, 1, 1, 1, 1, 0, 0, 0,
            0, 1, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 1, 0, 0, 0, 1, 0],
           dtype=int64)
[22]: # Confusion matrix
     from sklearn.metrics import confusion matrix
     cm = confusion_matrix(Y_test, y_predict)
     cm
[22]: array([[85, 15],
            [25, 29]], dtype=int64)
[23]: # Heatmap of Confusion matrix
     sns.heatmap(pd.DataFrame(cm), annot=True)
[23]: <Axes: >
```

Descision Tree:

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

[]: DecisionTreeClassifier()

Making prediction:

Logistic Regression:

Model Evaluation for Logistic Regression:

Train Score and Test Score

Train Accuracy of Logistic Regression: 77.36156351791531
Accuracy (Test) Score of Logistic Regression: 77.272727272727
Accuracy Score of Logistic Regression: 77.272727272727

```
[]: # For Decesion Tree:
print("Train Accuracy of Decesion Tree: ", dt.score(X_train, y_train)*100)
print("Accuracy (Test) Score of Decesion Tree: ", dt.score(X_test, y_test)*100)
print("Accuracy Score of Decesion Tree: ", accuracy_score(y_test, dt_pred)*100)
```

Train Accuracy of Decesion Tree: 100.0

Accuracy (Test) Score of Decesion Tree: 80.51948051948052

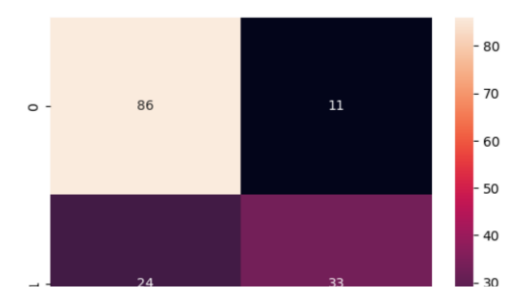
Accuracy Score of Decesion Tree: 80.51948051948052

Confusion Matrix

. Confusion Matrix of "Logistic Regression"

[]: sns.heatmap(confusion_matrix(y_test, lr_pred), annot=True, fmt="d")

[]: <Axes: >



```
[]: TN = cm[0, 0]
    FP = cm[0, 1]
    FN = cm[1, 0]
    TP = cm[1, 1]
[ ]: TN, FP, FN, TP
[ ]: (86, 11, 24, 33)
[]: from sklearn.metrics import classification_report, confusion_matrix
     from sklearn.metrics import accuracy_score, roc_auc_score, roc_curve
    cm = confusion_matrix(y_test, lr_pred)
    print('TN - True Negative {}'.format(cm[0,0]))
    print('FP - False Positive {}'.format(cm[0, 1]))
    print('FN - False Negative {}'.format(cm[1,0]))
    print('TP - True Positive {}'.format(cm[1, 1]))
    print('Accuracy Rate: {}'.format(np.divide(np.sum([cm[0, 0], cm[1, 1]]), np.
      -sum(cm))*100)
    print('Misclassification Rate: {}'.format(np.divide(np.sum([cm[0,1], cm[1,0]]), _
      -np. sum(cm))*100)
    TN - True Negative 86
    FP - False Positive 11
    FN - False Negative 24
    TP - True Positive 33
    Accuracy Rate: 77.272727272727
    Misclassification Rate: 22.7272727272727
[ ]: 77.27272727272727+22.727272727272727
[ ]: 100.0
```

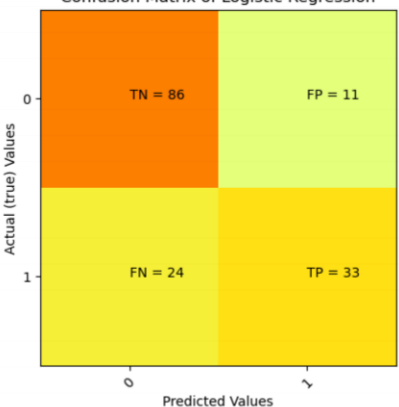
```
import matplotlib.pyplot as plt
import numpy as np

plt.clf()
plt.imshow(cm, interpolation='nearest', cmap=plt.cm.Wistia)
classNames = ['0', '1']
plt.title('Confusion Matrix of Logistic Regression')
plt.ylabel('Actual (true) Values')
plt.xlabel('Predicted Values')
tick_marks = np.arange(len(classNames))
plt.xticks(tick_marks, classNames, rotation=45)
plt.yticks(tick_marks, classNames)
```

```
s = [['TN', 'FP'], ['FN', 'TP']]
for i in range(2):
    for j in range(2):
        plt.text(j, i, str(s[i][j]) + " = " + str(cm[i][j]))

plt.show()
```

Confusion Matrix of Logistic Regression



86 11

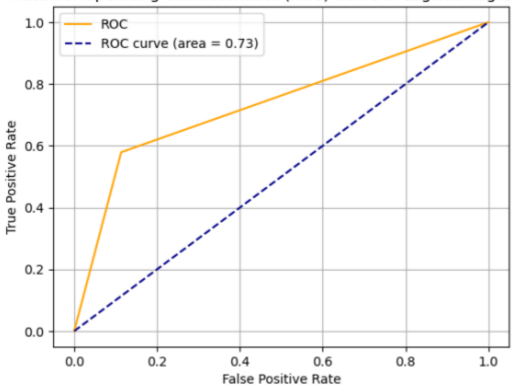
24 33 57

97

```
print('F1_Score of Macro: ', f1_score(y_test, Ir_pred)*100)
   F1 Score of Macro: 65.34653465346535
[ ]: print("Micro Average f1 Score is: ", f1_score(y_test, Ir_pred, _
      average= 'micro')*100)
    print("Macro Average f1 Score is: ", f1_score(y_test, Ir_pred, _
      -average= 'macro')*100)
    print("Weighted Average f1 Score is: ", f1_score(y_test, Ir_pred, _
      average='weighted')*100)
    print("f1 Score on Non Weighted score is: ", f1_score(y_test, Ir_pred, _
      average=None) *100)
   Micro Average f1 Score is: 77.272727272727
   Macro Average f1 Score is: 74.21916104653944
   Weighted Average f1 Score is: 76.52373933045479
   f1 Score on Non Weighted score is: [83.09178744 65.34653465]
    Classification Report of Logistic Regression:
    from sklearn.metrics import classification_report
print('Classification Report of Logistic Regression: \n', _
     -classification_report(y_test, lr_pred, digits=4))
   Classification Report of Logistic Regression:
                  precision
                                recall f1-score
                                                   support
                     0.7818
                               0.8866
               0
                                         0.8309
                                                       97
               1
                     0.7500
                               0.5789
                                         0.6535
                                                       57
                                         0.7727
                                                      154
        accuracy
                                         0.7422
                                                      154
                     0.7659
                               0.7328
      macro avg
                     0.7700
                               0.7727
                                         0.7652
                                                      154
   weighted avg
   ROC Curve& ROC AUC
    auc= roc_auc_score(y_test, lr_pred)
    print("ROC AUC SCORE of logistic Regression is ", auc)
   ROC AUC SCORE of logistic Regression is 0.7327726532826913
[]: from sklearn.metrics import roc_curve, auc
     import matplotlib.pyplot as plt
    fpr, tpr, thresholds = roc_curve(y_test, lr_pred)
```

[]: from sklearn.metrics import f1_score

Receiver Operating Characteristics (ROC) Curve of Logistic Regression



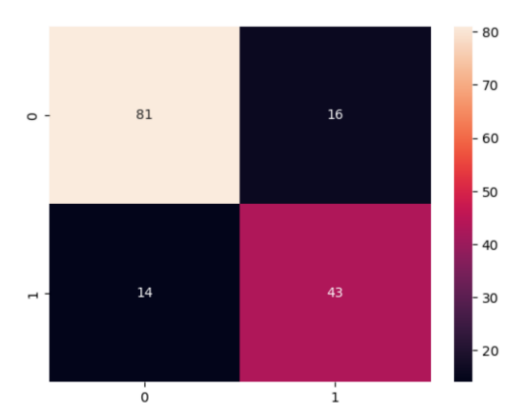
Confusion Matrix:

• Confusion matrix of "Decision Tree"

```
[]: from sklearn.metrics import classification_report, confusion_matrix

cm = confusion_matrix(y_test, dt_pred)

cm
```



```
[]: TN =cm[0, 0]
FP =cm[0,1]
FN = cm[1,0]
TP = cm[1,1]

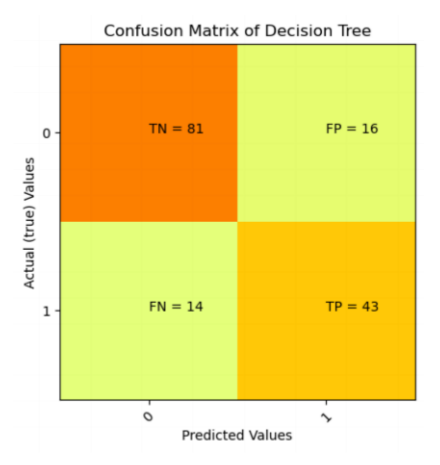
[]: TN, FP, FN, TP

[]: (81, 16, 14, 43)
```

```
[]: from sklearn.metrics import classification_report, confusion_matrix
    from sklearn.metrics import accuracy_score, roc_auc_score, roc_curve
    cm = confusion_matrix(y_test, dt_pred)

print('TN - True Negative {}'.format(cm[0,0]))
print('FP - False Positive {}'.format(cm[0,1]))
```

```
[ ]: import matplotlib.pyplot as plt
    import numpy as np
    plt.clf()
    plt.imshow(cm, interpolation='nearest', cmap=plt.cm.Wistia)
    classNames = ['0', '1']
    plt.title('Confusion Matrix of Decision Tree')
    plt.ylabel('Actual (true) Values')
    plt.xlabel('Predicted Values')
    tick_marks = np.arange(len(classNames))
    plt.xticks(tick_marks, classNames, rotation=45)
    plt.yticks(tick_marks, classNames)
    s = [['TN', 'FP'], ['FN', 'TP']]
    for i in range(2):
        for j in range(2):
            plt.text(j, i, str(s[i][j]) + " = " + str(cm[i][j]))
    plt.show()
```



Precision:

```
[ ]: Precision score:

precision_score = TP/float(TP+FP)*100
print('Precision Score: {0:0.4f}'.format(precision_score))
```

Precision Score: 72.8814

```
dt_pred, average=None) * 100)
   Precision Score is: 72.88135593220339
   Micro Average Precision Score is: 80.51948051948052
   Macro Average Precision Score is: 79.07225691347011
   Weighted Average Precision Score is: 80.68028314237056
   Precision Score on Non Weighted score is: [85.26315789 72.88135593]
   Recall:
[ ]: recall_score = TP/ float(TP+FN)*100
    print('recall_score', recall_score)
    recall score 75.43859649122807
[]: from sklearn.metrics import recall_score
    print('Recall or Sensitivity_Score: ', recall_score(y_test, dt_pred)*100)
   Recall or Sensitivity_Score: 75.43859649122807
[ ]: print("recall Score is: ", recall_score(y_test, dt_pred)*100)
    print("Micro Average recall Score is: ", recall_score(y_test, dt_pred, _
      -average='micro')*100)
    print("Macro Average recall Score is: ", recall_score(y_test, dt_pred, _
      average= 'macro')*100)
    print("Weighted Average recall Score is: ", recall_score(y_test, dt_pred, _
      -average='weighted')*100)
    print("recall Score on Non Weighted score is: ", recall_score(y_test, dt_pred, _
      average=None) *100)
   recall Score is: 75.43859649122807
   Micro Average recall Score is: 80.51948051948052
   Macro Average recall Score is: 79.47187556520167
   Weighted Average recall Score is: 80.51948051948052
   recall Score on Non Weighted score is: [83.50515464 75.43859649]
   FPR
[ ]: FPR = FP / float(FP + TN) * 100
    print('False Positive Rate: {:.4f}'.format(FPR))
   False Positive Rate: 16,4948
```

print ("Precision Score on Non Weighted score is:", precision_score(y_test, _

Specificity:

accuracy

macro avg weighted avg

[]: specificity = TN / (TN+FP) *100

```
print('Specificity : {0:0.4f}'.format(specificity))
    Specificity: 83.5052
[ ]: from sklearn.metrics import f1_score
    print('F1_Score of Macro: ', f1_score(y_test, dt_pred)*100)
   F1 Score of Macro: 74, 13793103448276
[ ]: print("Micro Average f1 Score is: ", f1_score(y_test, dt_pred, _
      -average='micro')*100)
    print("Macro Average f1 Score is: ", f1_score(y_test, dt_pred, _
      -average= 'macro')*100)
    print("Weighted Average f1 Score is: ", f1_score(y_test, dt_pred, _
      -average='weighted')*100)
    print("f1 Score on Non Weighted score is: ", f1_score(y_test, dt_pred, _
      average=None) *100)
   Micro Average f1 Score is: 80.51948051948051
   Macro Average f1 Score is: 79.25646551724138
   Weighted Average f1 Score is: 80.58595499328258
   f1 Score on Non Weighted score is: [84.375
                                                   74. 13793103]
    Classification Report of Decision Tree:
[ ]: from sklearn.metrics import classification_report
    print('Classification Report of Decision Tree: \n', _
      -classification_report(y_test, dt_pred, digits=4))
   Classification Report of Decision Tree:
                   precision
                                recall f1-score
                                                   support
                     0.8526
                               0.8351
                                         0.8438
                                                       97
               0
               1
                     0.7288
                               0.7544
                                         0.7414
                                                       57
```

0.8052

0.7926

0.8059

0.7947

0.8052

0.7907

0.8068

154

154

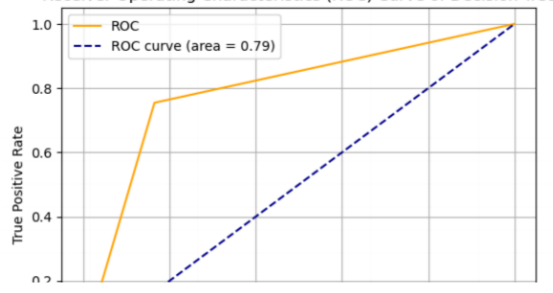
154

ROC Curve& ROC AUC

```
[ ]: auc= roc_auc_score(y_test, dt_pred)
print("ROC AUC SCORE of Decision Treeis ", auc)
```

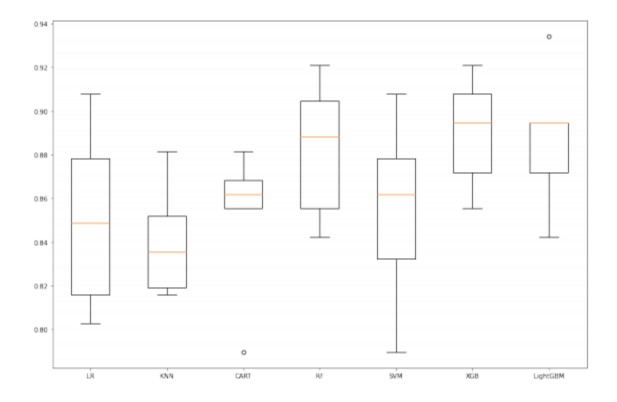
ROC AUC SCORE of Decision Tree is 0.7947187556520168

Receiver Operating Characteristics (ROC) Curve of Decision Tree



Base Models

```
[]: models = []
    models.append(('LR', LogisticRegression(random state = 12345)))
    models.append(('KNN', KNeighborsClassifier()))
    models.append(('CART', DecisionTreeClassifier(random state = 12345)))
    models.append(('RF', RandomForestClassifier(random state = 12345)))
    models.append(('SVM', SVC(gamma='auto', random state = 12345)))
    models.append(('XGB', GradientBoostingClassifier(random state = 12345)))
    models.append(("LightGBM", LGBMClassifier(random state = 12345)))
    results = []
    names = []
[ ]: for name, model in models:
            kfold = KFold(n splits = 10, random state = 12345)
            cv_results = cross_val_score(model, X, y, cv = 10, scoring= "accuracy")
            results.append(cv results)
            names.append(name)
            msg = "%s: %f (%f)" % (name, cv results.mean(), cv results.std())
            print (msg)
    fig = plt.figure(figsize=(15,10))
    fig.suptitle('Algorithm Comparison')
    ax = fig.add subplot(111)
    plt.boxplot(results)
    ax.set xticklabels(names)
    plt.show()
   LR: 0.848684 (0.036866)
   KNN: 0.840789 (0.023866)
   CART: 0.857895 (0.024826)
   RF: 0.881579 (0.026316)
   SVM: 0.853947 (0.036488)
   XGB: 0.890789 (0.020427)
   LightGBM: 0.885526 (0.024298)
```



5 Reporting

The aim of this study was to create classification models for the diabetes data set and to predict whether a person is sick by establishing models and to obtain maximum validation scores in the established models. The work done is as follows:

- 1) Diabetes Data Set read.
- 2) With Exploratory Data Analysis; The data set's structural data were checked. The types of variables in the dataset were examined. Size information of the dataset was accessed. The 0 values in the data set are missing values. Primarily these 0 values were replaced with NaN values. Descriptive statistics of the data set were examined.
- 3) Data Preprocessing section; df for: The NaN values missing observations were filled with the median values of whether each variable was sick or not. The outliers were determined by LOF and dropped. The X variables were standardized with the rubost method..
- 4) During Model Building; Logistic Regression, KNN, SVM, CART, Random Forests, XGBoost, LightGBM like using machine learning models Cross Validation Score were calculated. Later Random Forests, XGBoost, LightGBM hyperparameter optimizations optimized to increase Cross Validation value.
- 5) Result; The model created as a result of XGBoost hyperparameter optimization became the