Diabetes Patients Predictive Analysis

Project Overview:

This project focuses on predictive analysis for diabetes diagnosis using a dataset originally sourced from the National Institute of Diabetes and Digestive and Kidney Diseases. The dataset contains various medical and demographic variables for a group of Pima Indian heritage females who are at least 21 years old.

The primary objective of this project is to develop a predictive model that can diagnostically predict whether a patient has diabetes based on the provided diagnostic measurements and demographic information.

Dataset Description:

Features	Description
Pregnancies	The number of pregnancies a patient has had.
Glucose	Plasma glucose concentration, an indicator of blood sugar levels.
BloodPressure	Diastolic blood pressure.
SkinThickness	Skinfold thickness, which may be related to body composition.
Insulin	2-Hour serum insulin level.
BMI	A measure of body weight and height, indicating body fat.
DiabetesPedigreeFunction	A measure of the diabetes heredity risk based on family history.
Age	The age of the patient in years.
Outcome	The target variable indicating whether the patient has diabetes (1 for positive, 0 for negative).

Importing Important Libraries For This Project

```
import pandas as pd
         import numpy as np
        import matplotlib.pyplot as plt
        import seaborn as sns
         import plotly.express as px
        import math
         #Spliting Data into Train and Test:
         from sklearn.model_selection import train_test_split
         #For Feature Scaling:
        from sklearn.preprocessing import StandardScaler
         #Support Vector Machine:
         from sklearn.svm import SVC
         #Logistic Regression:
        \textbf{from} \  \, \textbf{sklearn.linear\_model import} \  \, \textbf{LogisticRegression}
         #Evaluation:
         from sklearn.model selection import cross val score
         from sklearn.metrics import accuracy_score
         #For ignoring warnings:
         import warnings
        warnings.filterwarnings('ignore')
In [3]: #Import Dataset
        data = pd.read csv("G:/data analyst/DIABETES PATIENTS/diabetes.csv")
In [4]: #Copy the dataset
        df = data.copy()
```

Data Exploration

In [5]:	df	df.head()										
Out[5]:		Pregnancies	Glucose	BloodPressure	SkinThickness	Insulin	вмі	DiabetesPedigreeFunction	Age	Outcome		
	0	6	148	72	35	0	33.6	0.627	50	1		
	1	1	85	66	29	0	26.6	0.351	31	0		
	2	8	183	64	0	0	23.3	0.672	32	1		
	3	1	89	66	23	94	28.1	0.167	21	0		
	4	0	137	40	35	168	43.1	2.288	33	1		

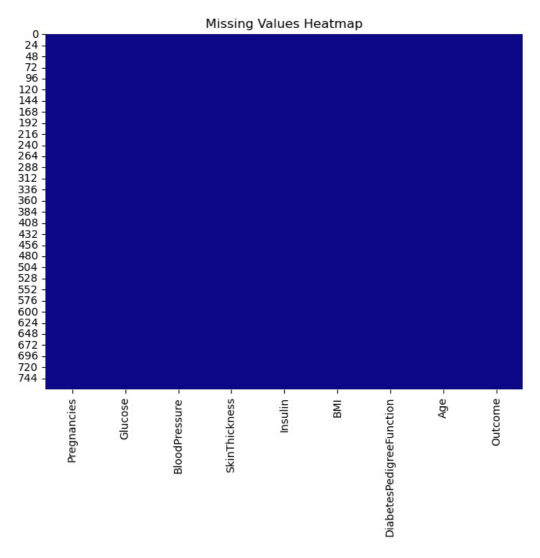
Out[6]:		Pregnancies	Glucose	BloodPressure	SkinThickness	Insulin	ВМІ	DiabetesPedigreeFunction	Age	Outcome
	763	10	101	76	48	180	32.9	0.171	63	0
	764	2	122	70	27	0	36.8	0.340	27	0
	765	5	121	72	23	112	26.2	0.245	30	0
	766	1	126	60	0	0	30.1	0.349	47	1
	767	1	93	70	31	0	30.4	0.315	23	0

Number of Rows and Columns in Dataset

```
In [7]: df.shape
         print("Total Number of Rows in Dataset :",data.shape[0])
print("Total Number of Columns in Dataset:",data.shape[1])
         Total Number of Rows in Dataset : 768
         Total Number of Columns in Dataset: 9
In [8]: df.info()
         <class 'pandas.core.frame.DataFrame'>
         RangeIndex: 768 entries, 0 to 767
        Data columns (total 9 columns):
                                         Non-Null Count Dtype
         # Column
             Pregnancies
                                         768 non-null
         0
                                                          int64
                                                        int64
         1
              Glucose
                                         768 non-null
              BloodPressure
                                         768 non-null
                                                          int64
         3
              SkinThickness
                                         768 non-null
                                                          int64
          4
              Insulin
                                         768 non-null
                                                          int64
         5
              BMI
                                         768 non-null
                                                          float64
             DiabetesPedigreeFunction 768 non-null
                                                          float64
                                          768 non-null
              Age
                                                          int64
             Outcome
                                                          int64
         8
                                          768 non-null
         dtypes: float64(2), int64(7)
         memory usage: 54.1 KB
```

Heatmap to Check Missing Values in Dataset

```
In [9]: 
plt.figure(figsize=(8, 6))
sns.heatmap(df.isnull(), cmap='plasma', cbar=False)
plt.title('Missing Values Heatmap')
plt.show()
```



We have verified that the dataset is free of any missing or null values.

Overall Statistics About The Dataset

10]:	df.describe()												
10]:		Pregnancies	Glucose	BloodPressure	SkinThickness	Insulin	ВМІ	DiabetesPedigreeFunction	Age	Outcome			
	count	768.000000	768.000000	768.000000	768.000000	768.000000	768.000000	768.000000	768.000000	768.000000			
	mean	3.845052	120.894531	69.105469	20.536458	79.799479	31.992578	0.471876	33.240885	0.348958			
	std	3.369578	31.972618	19.355807	15.952218	115.244002	7.884160	0.331329	11.760232	0.476951			
	min	0.000000	0.000000	0.000000	0.000000	0.000000	0.000000	0.078000	21.000000	0.000000			
	25%	1.000000	99.000000	62.000000	0.000000	0.000000	27.300000	0.243750	24.000000	0.000000			
	50%	3.000000	117.000000	72.000000	23.000000	30.500000	32.000000	0.372500	29.000000	0.000000			
	75%	6.000000	140.250000	80.000000	32.000000	127.250000	36.600000	0.626250	41.000000	1.000000			
	max	17.000000	199.000000	122.000000	99.000000	846.000000	67.100000	2.420000	81.000000	1.000000			

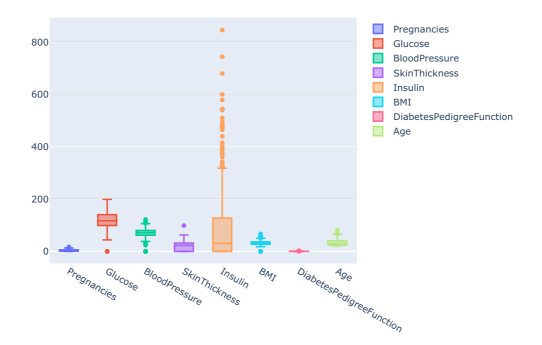
Data Summary Report

Features	Description
Pregnancies	Most data falls within the range of 1 to 6 pregnancies.
Glucose	Glucose levels are typically distributed between 99 to 140.
BloodPressure	Most values fall within the 62 to 80 range.
SkinThickness	A significant portion of the data has values between 0 and 32.
Insulin	A substantial portion of the data has low insulin values.
BMI	Most values fall in the range of 27.3 to 36.6.
DiabetesPedigreeFunction	The majority of values are below 0.626.
Age	Most of the data represents individuals between the ages of 24 and 41.
Outcome	The dataset appears to be imbalanced, with a lower number of positive outcomes (diabetes) compared to negative outcomes (no diabetes).

Checking Outliers

```
In [11]: import plotly.graph_objs as go
         # Create a list to store the box plot traces
         box traces = []
         # Iterate through each column and create a box plot
         for column in df.columns:
             if column != 'Outcome': # Exclude 'Outcome' if it's the target variable
                 trace = go.Box(y=df[column], name=column)
                 box_traces.append(trace)
         # Create a layout
         layout = go.Layout(title='Box Plots for Dataset Columns')
         # Create a figure and add the traces and layout
         fig = go.Figure(data=box traces, layout=layout)
         # Show the figure
         fig.show()
```

Box Plots for Dataset Columns



Outliers have been identified in our dataset.

Managing Outliers in Dataset

```
In [12]:
             #Create a function to handle Outliers
             def remove outliers(data, column name):
                  Q1 = data[column name].quantile(0.25)
                  Q3 = data[column_name].quantile(0.75)
                  IQR = Q3 - Q1
                  upper_limit = Q3 + 1.5 * IQR
                   lower_limit = Q1 - 1.5 * IQR
                  data[column name] = data[column name].clip(lower=lower limit, upper=upper limit)
                   return data
In [13]: #Handle outliers using "remove_outliers" function
             df = remove_outliers(df, 'Pregnancies')
            df = remove_outliers(df, 'Fregnancies')
df = remove_outliers(df, 'Glucose')
df = remove_outliers(df, 'BloodPressure')
df = remove_outliers(df, 'SkinThickness')
df = remove_outliers(df, 'Insulin')
df = remove_outliers(df, 'BMI')
```

In [14]: import plotly graph obis as go

```
# Create a list to store the box plot traces
box_traces = []

for column in df.columns:
    if column != 'Outcome': # Exclude 'Outcome' if it's the target variable
        trace = go.Box(y=df[column], name=column)
        box_traces.append(trace)

# Create a layout
layout = go.Layout(title='Box Plots for Dataset Columns')

# Create a figure and add the traces and layout
fig = go.Figure(data=box_traces, layout=layout)

# Show the figure
fig.show()
```

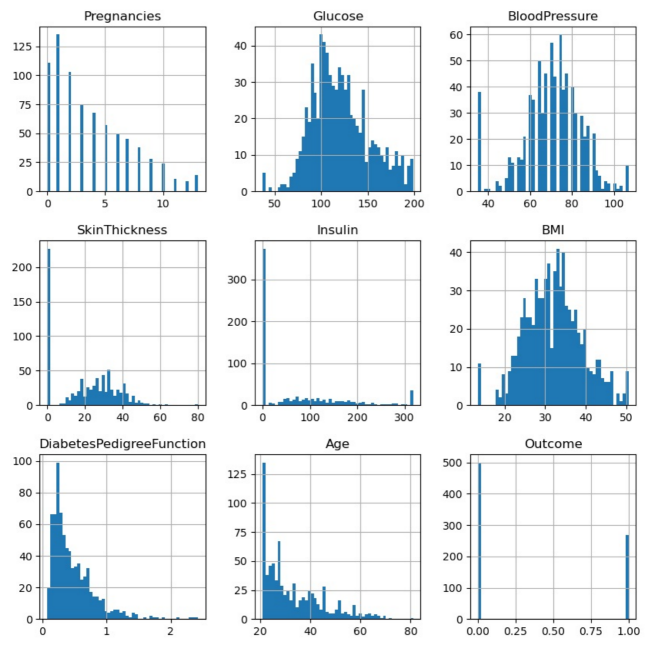
After handling outliers , the datatype of some columns has changed to float. We also need to convert them back to int32.

```
In [15]: df['Pregnancies']=round(df['Pregnancies'].astype('int32'))
         df['Glucose']=round(df['Glucose'].astype('int32'))
df['Insulin']=round(df['Insulin'].astype('int32'))
In [16]: df.info()
         <class 'pandas.core.frame.DataFrame'>
         RangeIndex: 768 entries, 0 to 767
         Data columns (total 9 columns):
          # Column
                                         Non-Null Count Dtype
          0 Pregnancies
                                                          int32
                                          768 non-null
          1
             Glucose
                                         768 non-null int32
              BloodPressure
                                          768 non-null
                                                          int64
                                         768 non-null int64
          3
             SkinThickness
          4
              Insulin
                                          768 non-null
                                                         int32
          5
              BMI
                                          768 non-null
                                                          float64
             DiabetesPedigreeFunction 768 non-null
          6
                                                         float64
          7
                                          768 non-null
              Age
                                                          int64
             Outcome
          8
                                          768 non-null
                                                          int64
         dtypes: float64(2), int32(3), int64(4)
         memory usage: 45.1 KB
```

Visualizing the Dataset for Better Understanding:

```
fig, ax = plt.subplots(figsize=(10, 10))
df.hist(bins=50, ax=ax)

# Show the plot
plt.show()
```



Observations:

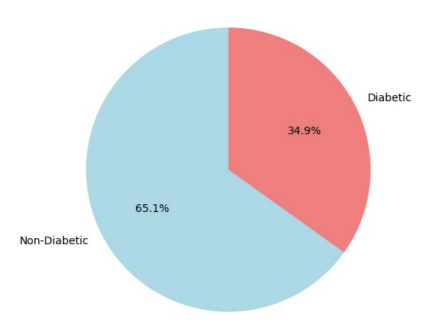
Distributions are mostly skewed to the right

Small peaks at higher values for glucose, blood pressure, skin thickness, insulin, BMI, and diabetes pedigree function

Bimodal distribution for outcome variable (diabetes vs. no diabetes)

```
In [58]: # Count the occurrences of each outcome value
  outcome_counts = df['Outcome'].value_counts()

# Create a pie chart
  plt.figure(figsize=(6, 6))
  plt.pie(outcome_counts, labels=['Non-Diabetic', 'Diabetic'], autopct='%1.1f%%', startangle=90, colors=['lightbl
  plt.title('Distribution of Outcomes')
  plt.show()
```



Converting Numerical Features into Categorical Features for Data Clarity:

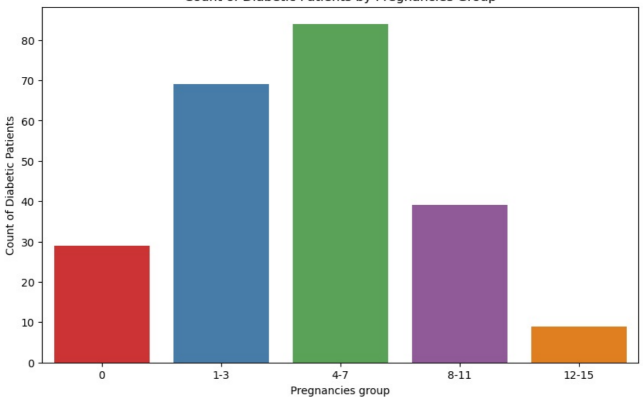
Create bins for the 'Pregnancies' column

```
In [53]:
    pregnancies_bins = [0, 1, 4, 8, 12, 16]
    pregnancies_labels = ['0', '1-3', '4-7', '8-11', '12-15']
    df['PregnanciesGroup'] = pd.cut(df['Pregnancies'], bins=pregnancies_bins, labels=pregnancies_labels)

# Filter the dataset to include only records with 'Outcome' equal to 1 (Diabetic patients)
    diabetic_df = df[df['Outcome'] == 1]

# Create a bar chart for Diabetic patients with 'PregnanciesGroup' as the x-axis
    plt.figure(figsize=(10, 6))
    sns.countplot(data=diabetic_df, x='PregnanciesGroup', order=pregnancies_labels, palette="Set")
    plt.xlabel('Pregnancies group')
    plt.ylabel('Count of Diabetic Patients')
    plt.title('Count of Diabetic Patients by Pregnancies Group')
    plt.show()
```





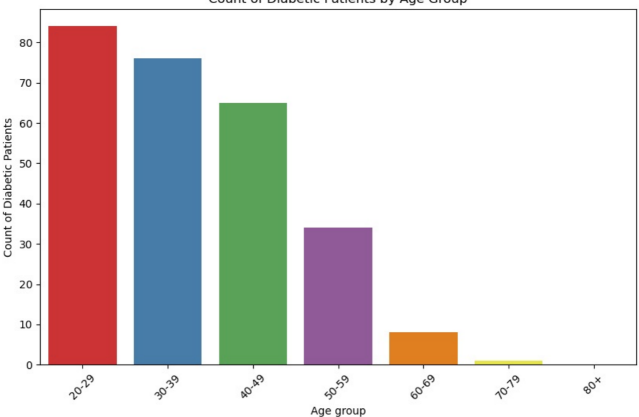
Create age groups based on the 'Age' column

```
In [21]: bins = [20, 30, 40, 50, 60, 70, 80, 200]
labels = ['20-29', '30-39', '40-49', '50-59', '60-69', '70-79', '80+']
df['AgeGroup'] = pd.cut(df['Age'], bins=bins, labels=labels, right=False)

In [52]: # Filter the dataset to include only records with 'Outcome' equal to 1 (Diabetic patients)
diabetic_df = df[df['Outcome'] == 1]

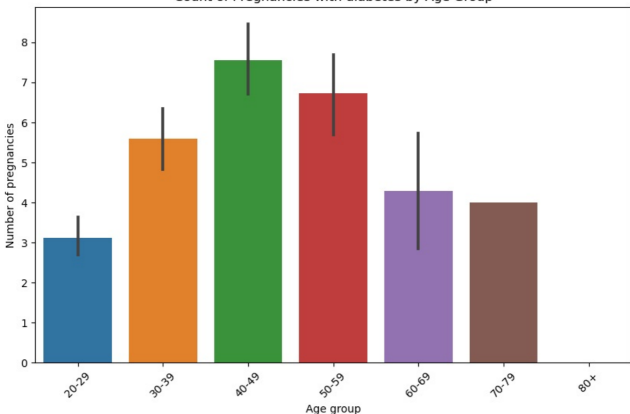
# Create a bar chart for Diabetic patients with age groups
plt.figure(figsize=(10, 6))
sns.countplot(data=diabetic_df, x='AgeGroup', order=labels, palette="Set1")
plt.xlabel('Age group')
plt.ylabel('Count of Diabetic Patients')
plt.title('Count of Diabetic Patients by Age Group')
plt.xticks(rotation=45)
plt.show()
```

Count of Diabetic Patients by Age Group



```
In [51]: new_df = df[(df['Outcome'] == 1) & (df['Pregnancies'] > 0)]
# Create a bar chart with 'Outcome' as hue
plt.figure(figsize=(10, 6))
ax = sns.barplot(data=new_df, x='AgeGroup', y='Pregnancies')
plt.xlabel('Age group')
plt.ylabel('Number of pregnancies')
plt.title('Count of Pregnancies with diabetes by Age Group')
plt.xticks(rotation=45)
plt.show()
```

Count of Pregnancies with diabetes by Age Group



Define the bins and labels for 'BloodPressure'

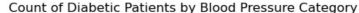
```
In [24]: # Define the bins and labels for 'BloodPressure'
blood_pressure_bins = [0, 80, 89, 99, 119, 1000] # Adjust the boundaries as needed
blood_pressure_labels = ['Low', 'Normal', 'Prehypertension', 'Stage 1 hypertension', 'Stage 2 hypertension']

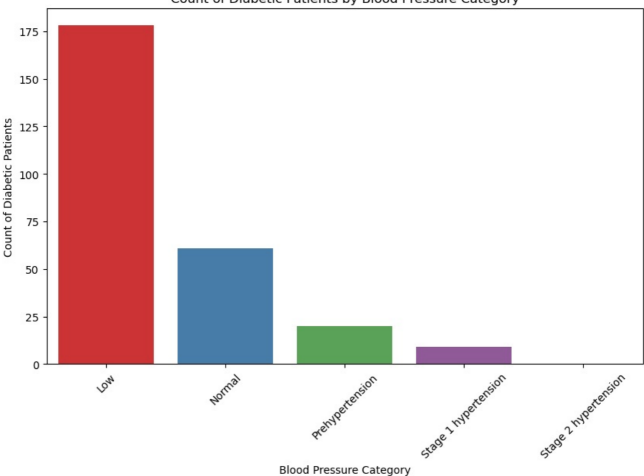
# Create a new column 'BloodPressureCategory' based on the bins and labels
df['BloodPressureCategory'] = pd.cut(df['BloodPressure'], bins=blood_pressure_bins, labels=blood_pressure_label
df.head()
```

Out[24]:		Pregnancies	Glucose	BloodPressure	SkinThickness	Insulin	BMI	DiabetesPedigreeFunction	Age	Outcome	PregnanciesGroup	AgeGrou
	0	6	148	72	35	0	33.6	0.627	50	1	4-7	50-
	1	1	85	66	29	0	26.6	0.351	31	0	0	30-0
	2	8	183	64	0	0	23.3	0.672	32	1	4-7	30-0
	3	1	89	66	23	94	28.1	0.167	21	0	0	20-2
	4	0	137	40	35	168	43.1	2.288	33	1	NaN	30-(

```
In [50]: # Filter the dataset to include only records with 'Outcome' equal to 1 (Diabetic patients)
diabetic_df = df[df['Outcome'] == 1]

# Create a bar chart for Diabetic patients with 'BloodPressureCategory' as the x-axis
plt.figure(figsize=(10, 6))
sns.countplot(data=diabetic_df, x='BloodPressureCategory', order=blood_pressure_labels, palette="Set1")
plt.xlabel('Blood Pressure Category')
plt.ylabel('Count of Diabetic Patients')
plt.title('Count of Diabetic Patients by Blood Pressure Category')
plt.xticks(rotation=45)
plt.show()
```





Define the bins and labels for 'SkinThickness'

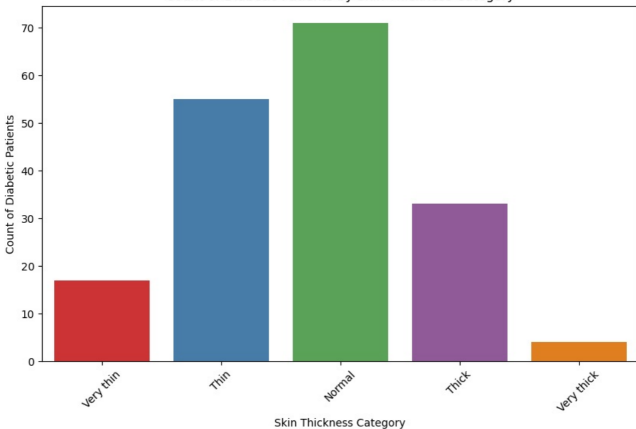
```
In [49]:
    skin_thickness_bins = [0, 20, 30, 40, 50, 100]
    skin_thickness_labels = ['Very thin', 'Thin', 'Normal', 'Thick', 'Very thick']

# Create a new column 'SkinThicknessCategory' based on the bins and labels
    df['SkinThicknessCategory'] = pd.cut(df['SkinThickness'], bins=skin_thickness_bins, labels=skin_thickness_label

# Filter the dataset to include only records with 'Outcome' equal to 1 (Diabetic patients)
    diabetic_df = df[df['Outcome'] == 1]

# Create a bar chart for Diabetic patients with 'SkinThicknessCategory' as the x-axis
    plt.figure(figsize=(10, 6))
    sns.countplot(data=diabetic_df, x='SkinThicknessCategory', order=skin_thickness_labels, palette="Set1")
    plt.xlabel('Skin Thickness Category')
    plt.xlabel('Skin Thickness Category')
    plt.ylabel('Count of Diabetic Patients')
    plt.title('Count of Diabetic Patients by Skin Thickness Category')
    plt.xticks(rotation=45)
    plt.show()
```

Count of Diabetic Patients by Skin Thickness Category



Define the custom bins and labels for 'BMI'

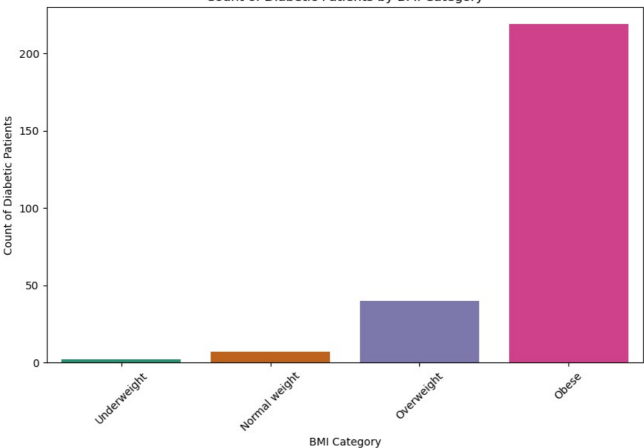
```
In [46]: # Define the custom bins and labels for 'BMI'
    bmi_bins = [0, 18.5, 24.9, 29.9, 1000]
    bmi_labels = ['Underweight', 'Normal weight', 'Overweight', 'Obese']

# Create a new column 'BMICategory' based on the custom bins and labels
    df['BMICategory'] = pd.cut(df['BMI'], bins=bmi_bins, labels=bmi_labels)

# Filter the dataset to include only records with 'Outcome' equal to 1 (Diabetic patients)
    diabetic_df = df[df['Outcome'] == 1]

# Create a bar chart for Diabetic patients with 'BMICategory' as the x-axis
    plt.figure(figsize=(10, 6))
    sns.countplot(data=diabetic_df, x='BMICategory', order=bmi_labels, palette="Dark2")
    plt.xlabel('BMI Category')
    plt.ylabel('Count of Diabetic Patients')
    plt.title('Count of Diabetic Patients by BMI Category')
    plt.xticks(rotation=45)
    plt.show()
```

Count of Diabetic Patients by BMI Category



Define the custom bins and labels for 'Insulin'

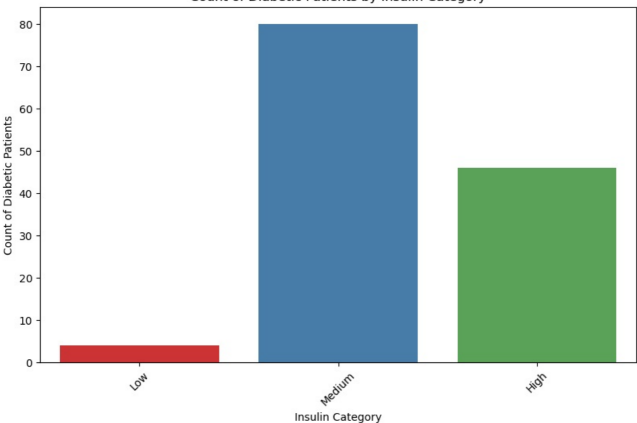
```
insulin_bins = [0, 50, 200, 10000]
insulin_labels = ['Low', 'Medium', 'High']

# Create a new column 'InsulinCategory' based on the custom bins and labels
df['InsulinCategory'] = pd.cut(df['Insulin'], bins=insulin_bins, labels=insulin_labels)

# Filter the dataset to include only records with 'Outcome' equal to 1 (Diabetic patients)
diabetic_df = df[df['Outcome'] == 1]

# Create a bar chart for Diabetic patients with 'InsulinCategory' as the x-axis
plt.figure(figsize=(10, 6))
sns.countplot(data=diabetic_df, x='InsulinCategory', order=insulin_labels, palette="Set1")
plt.xlabel('Insulin Category')
plt.ylabel('Count of Diabetic Patients')
plt.title('Count of Diabetic Patients by Insulin Category')
plt.xticks(rotation=45)
plt.show()
```





Define the custom bins and labels for 'Glucose'

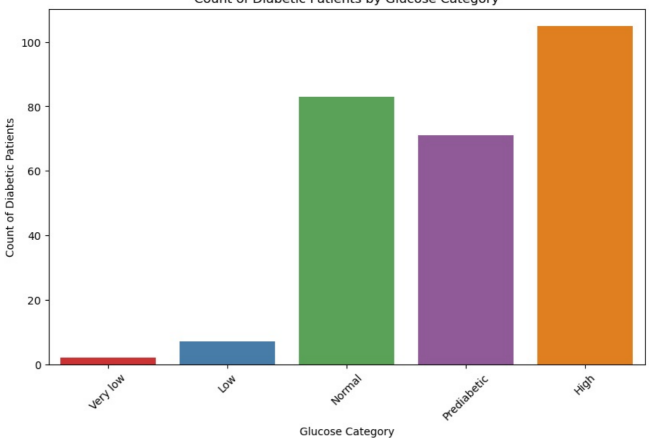
```
glucose_bins = [0, 75, 90, 125, 150, 1000] # Adjust the boundaries as needed
glucose_labels = ['Very low', 'Low', 'Normal', 'Prediabetic', 'High']

# Create a new column 'GlucoseCategory' based on the custom bins and labels
df['GlucoseCategory'] = pd.cut(df['Glucose'], bins=glucose_bins, labels=glucose_labels)

# Filter the dataset to include only records with 'Outcome' equal to 1 (Diabetic patients)
diabetic_df = df[df['Outcome'] == 1]

# Create a bar chart for Diabetic patients with 'GlucoseCategory' as the x-axis
plt.figure(figsize=(10, 6))
sns.countplot(data=diabetic_df, x='GlucoseCategory', order=glucose_labels, palette="Set1")
plt.xlabel('Glucose Category')
plt.ylabel('Gount of Diabetic Patients')
plt.title('Count of Diabetic Patients by Glucose Category')
plt.xticks(rotation=45)
plt.show()
```





Data Wrangling:

Seprate Independent Variable(X) and Dependent Variable(y)

```
df.columns
In [30]:
        Out[30]:
              dtype='object')
In [31]: #Independent Variables
         X = df[['Pregnancies', 'Glucose', 'BloodPressure', 'SkinThickness', 'Insulin', 'BMI', 'DiabetesPedigreeFunction'] \\
         #Target variable
         y = df['Outcome']
In [32]: X.head()
           Pregnancies Glucose BloodPressure SkinThickness Insulin BMI DiabetesPedigreeFunction Age
Out[32]:
                                                       0 33.6
                         85
                                     66
                                                29
                                                       0 26.6
                                                                            0.351
                                                                                  31
         2
                   8
                        183
                                     64
                                                 0
                                                       0 23.3
                                                                            0.672
                                                                                   32
                                                 23
                                                          28.1
                                                                            0.167
                                                                                   21
                        137
                                     40
                                                 35
                                                      168 43.1
                                                                            2.288
                                                                                   33
In [33]: y.head()
Out[33]:
             0
         2
             1
         3
             0
         Name: Outcome, dtype: int64
```

Split the data into training and testing sets

Feature Scaling

Train a Modal

SVM:

```
In [37]: sv_model = SVC(C= 0.1,kernel='linear',random_state=15)
sv_model.fit(X_train,y_train)

#Prediction on Traing Data
sv_pred_train = sv_model.predict(X_train)
#Prediction on Test Data
sv_pred_test = sv_model.predict(X_test)

#Evaluation
SVM_Train_Accuracy = accuracy_score(y_train,sv_pred_train)*100
SVM_Test_Accuracy = accuracy_score(y_test,sv_pred_test)*100
SVM_CV = cross_val_score(sv_model,X_test,y_test,cv=5,scoring="accuracy").mean()*100

print(f"Train Accuracy: {SVM_Train_Accuracy:.2f}%")
print(f"Test Accuracy: {SVM_Test_Accuracy:.2f}%")
print(f"cross Validataion Score: {SVM_CV:.2f}%")
Train Accuracy: 76.55%
```

LogisticRegression

Test Accuracy: 76.62%

cross Validataion Score: 74.69%

```
In [38]: logistic = LogisticRegression(C=100,penalty='l1',solver='liblinear',random_state=16)
logistic.fit(X_train,y_train)

#Prediction on Traing Data
log_pred_train = logistic.predict(X_train)
#Prediction on Test Data
log_pred_test = logistic.predict(X_test)

log_Train_Accuracy = accuracy_score(y_train,log_pred_train)*100
log_Test_Accuracy = accuracy_score(y_test,log_pred_test)*100
Log_CV = cross_val_score(logistic,X_test,y_test,cv=5,scoring="accuracy").mean()*100

print(f"Train Accuracy: {log_Train_Accuracy:.2f}%")
print(f"Test Accuracy: {log_Test_Accuracy:.2f}%")
print(f"cross Validataion Score: {Log_CV:.2f}%")
Train Accuracy: 77.20%
```

Conclusion

Test Accuracy: 77.27%

cross Validataion Score: 75.94%

After evaluating the performance metrics of the models, specifically Support Vector Machine and Logistic Regression, and keeping in mind the objective of maximizing the accuracy in predicting Diabetic Patients, the **Logistic Regression** model stands out as the most suitable choice.

Developing a Prediction System

```
input_data = (2,174,88,37,120,44.5,0.646,24)

# changing the input_data to numpy array
input_data_as_numpy_array = np.asarray(input_data)

# reshape the array as we are predicting for one instance
input_data_reshaped = input_data_as_numpy_array.reshape(1,-1)

# standardize the input data
std_data = scaler.transform(input_data_reshaped)

prediction = logistic.predict(std_data)
print(prediction)

if (prediction[0] == 0):
    print('The person is not diabetic')
else:
    print('The person is diabetic')

[1]
```

Graphical User Interface

The person is diabetic

```
In [40]:
         import tkinter as tk
         from tkinter import Label, Entry, Button
         # Function to make predictions
         def predict():
             input_data = [
                 pregnancies_entry.get(),
                 glucose entry.get(),
                 blood_pressure_entry.get(),
                 skin_thickness_entry.get(),
                 insulin entry get(),
                 bmi_entry.get(),
                 pedigree_function_entry.get(),
                 age_entry.get()
             1
             # Convert input_data to a NumPy array and standardize it
             input data as numpy array = np.asarray(input data, dtype=float).reshape(1, -1)
             std input data = scaler.transform(input_data_as_numpy_array)
             # Make the prediction using sv_model
             prediction = logistic.predict(std input data)
             # Interpret the prediction
             result = 'diabetic' if prediction[0] == 1 else 'not diabetic'
             result_label.config(text=f'Prediction: {result}')
         # Create the main window
         root = tk.Tk()
         root.title("Diabetes Prediction")
         # Create labels and entry fields for input features
         Label(root, text="Pregnancies").grid(row=0, column=0)
         pregnancies_entry = Entry(root)
         pregnancies entry.grid(row=0, column=1)
         Label(root, text="Glucose").grid(row=1, column=0)
         glucose_entry = Entry(root)
         glucose_entry.grid(row=1, column=1)
         Label(root, text="Blood Pressure").grid(row=2, column=0)
         blood pressure entry = Entry(root)
         blood pressure entry.grid(row=2, column=1)
         Label(root, text="Skin Thickness").grid(row=3, column=0)
         skin thickness entry = Entry(root)
         skin_thickness_entry.grid(row=3, column=1)
         Label(root, text="Insulin").grid(row=4, column=0)
         insulin entry = Entry(root)
         insulin_entry.grid(row=4, column=1)
         Label(root, text="BMI").grid(row=5, column=0)
         bmi_entry = Entry(root)
```

```
bmi_entry.grid(row=5, column=1)
Label(root, text="Diabetes Pedigree Function").grid(row=6, column=0)
pedigree_function_entry = Entry(root)
pedigree_function_entry.grid(row=6, column=1)
Label(root, text="Age").grid(row=7, column=0)
age_entry = Entry(root)
age_entry.grid(row=7, column=1)

# Create a button to make predictions
predict_button = Button(root, text="Predict", command=predict)
predict_button.grid(row=8, columnspan=2)

# Create a label to display predictions
result_label = Label(root, text="Prediction: ")
result_label.grid(row=9, columnspan=2)

# Start the main loop
root.mainloop()
```