# AI BASED DIABETES PREDICTION SYSTEM

## **PHASE 3 SUBMISSION**

## **DATA LOADING & PREPROCESSING**

SUBMITTED BY

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### INTRODUCTION:

Loading and preprocessing data are crucial steps in building a diabetic prediction system. The accuracy and reliability of predictions heavily depend on the quality of data and how well it's prepared for analysis. In this phase we are going to start develop the diabetes prediction system by loading and preprocessing the dataset

#### DATA SOURCE:

https://www.kaggle.com/datasets/mathchi/diabetes-data-set/

### OBJECTIVE OF DATA SOURCE:

This dataset is originally from the National Institute of Diabetes and Digestive and Kidney Diseases. The objective of the dataset is to diagnostically predict whether or not a patient has diabetes, based on certain diagnostic measurements included in the dataset.

Several constraints were placed on the selection of these instances from a larger database. In particular, all patients here are females at least 21 years old of Pima Indian heritage.

#### The Pima Indian Diabetes data set consists of

- → Pregnancies: Number of times pregnant
- → Glucose: Plasma glucose concentration a 2 hours in an oral glucose tolerance test
- → Blood Pressure: Diastolic blood pressure (mm Hg)
- → Skin Thickness: Triceps skin fold thickness (mm)
- → Insulin: 2-Hour serum insulin (mu U/ml)
- $\rightarrow$  BMI: Body mass index (weight in kg/(height in m)^2)
- → Diabetes Pedigree Function: Diabetes pedigree function
- → Age: Age (years)
- → Outcome: Class variable (0 or 1) 268 of 768 are 1, the others are 0

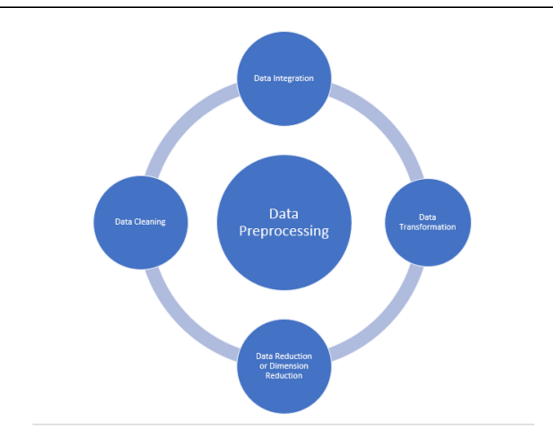
### WHAT IS DATA PREPROCESSING?

Data preprocessing is the process of transforming raw data into an understandable format. It is also an important step in data mining as we cannot work with raw data. The quality of the data should be checked before applying machine learning or data mining algorithms.

### Major Tasks in Data Preprocessing

There are 4 major tasks in data preprocessing –

- 1. Data cleaning,
- 2. Data integration,
- 3. Data reduction
- 4. Data transformation.



## O DATA CLEANING

Data cleaning is the process of removing incorrect data, incomplete data, and inaccurate data from the datasets, and it also replaces the missing values

### Data Integration

The process of combining multiple sources into a single dataset. The Data integration process is one of the main components of data management.

### DATA REDUCTION

This process helps in the reduction of the volume of the data, which makes the analysis easier yet produces the same or almost the same result. This reduction also helps to reduce storage space. Some of the data reduction techniques are dimensionality reduction, numerosity reduction, and data compression.

### Data Transformation

The change made in the format or the structure of the data is called data transformation. This step can be simple or complex based on the requirements. There are some methods for data transformation.

**Smoothing:** With the help of algorithms, we can remove noise from the dataset, which helps in knowing the important features of the dataset. By smoothing, we can find even a simple change that helps in prediction.

**Aggregation:** In this method, the data is stored and presented in the form of a summary. The data set, which is from multiple sources, is integrated into with data analysis description. This is an important step since the accuracy of the data depends on the quantity and quality of the data. When the quality and the quantity of the data are good, the results are more relevant.

**Discretization:** The continuous data here is split into intervals. Discretization reduces the data size. For example, rather than specifying the class time, we can set an interval like (3 pm-5 pm, or 6 pm-8 pm).

**Normalization:** It is the method of scaling the data so that it can be represented in a smaller range. Example ranging from -1.0 to 1.0.

### • IMPLEMENTATION:

# 1. import libraries

```
#import libraries
import numpy as np
import pandas as pd
import matplotlib.pyplot as plt
import seaborn as sns
import warnings
warnings.filterwarnings('ignore')
```

# 2. Load the dataset

```
#Reading the dataset
df = pd.read_csv("/kaggle/input/diabetes-predictionsys/diabetes.csv")
df.head()
    Pregnancies Glucose BloodPressure SkinThickness Insulin
                                                                 BMI
 0
                     148
                                                                33.6
              6
                                                   29
 1
                                                             0
              1
                     85
                                    66
                                                                26.6
 2
              8
                     183
                                    64
                                                    0
                                                             0 23.3
 3
              1
                                                   23
                     89
                                    66
                                                            94 28.1
 4
              0
                     137
                                    40
                                                   35
                                                           168 43.1
    DiabetesPedigreeFunction Age Outcome
 0
                      0.627
                              50
 1
                      0.351
                              31
                                        0
 2
                      0.672
                              32
                                        1
 3
                       0.167
                              21
                                        0
 4
                       2.288
                              33
                                        1
```

# 3. Analysis the dataset

```
#columns available in our dataset
df.columns
 Index(['Pregnancies', 'Glucose', 'BloodPressure', 'SkinThickness', 'Insulin',
        'BMI', 'DiabetesPedigreeFunction', 'Age', 'Outcome'],
       dtype='object')
#Information about the data
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
                              768 non-null
                                              int64
    Glucose
  1
                              768 non-null
                                              int64
  2
     BloodPressure
                              768 non-null
                                              int64
                                             int64
  3
     SkinThickness
                               768 non-null
  4
    Insulin
                               768 non-null
                                             int64
                               768 non-null
                                              float64
     DiabetesPedigreeFunction 768 non-null
                                              float64
  7
     Age
                               768 non-null
                                             int64
     Outcome
                               768 non-null
                                              int64
 dtypes: float64(2), int64(7)
 memory usage: 54.1 KB
```

```
#more about the dataset
df.describe().T
                                                                       25%
                           count
                                                             min
                                        mean
                                                     std
                                                           0.000
 Pregnancies
                           768.0
                                    3.845052
                                                3.369578
                                                                   1.00000
 Glucose
                                                           0.000
                           768.0 120.894531
                                               31.972618
                                                                  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
                           768.0
 BMI
                                   31.992578
                                                7.884160
                                                           0.000
                                                                  27.30000
 DiabetesPedigreeFunction
                           768.0
                                                           0.078
                                    0.471876
                                                0.331329
                                                                   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
                             3.0000
                                       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
 BMI
                                                 67.10
                            32.0000
                                     36.60000
 DiabetesPedigreeFunction
                             0.3725
                                       0.62625
                                                  2.42
```

41.00000

1.00000

81.00

1.00

# 4. Clean the data

Outcome

Clean the data after analyse the null, impossible value and duplicated value

29.0000

0.0000

```
#analyse the null and duplicated values
print(f'Duplicated rows are: \n { df.duplicated().sum()} \n \n \n Null
values per column are: \n {df.isnull().sum()}\n \n Zero values per
column are: \n {(df == 0).sum()} , \n \n \n \data types of each column is:
\n {df.dtypes} ')
 Duplicated rows are:
  Null values per column are:
 Pregnancies
                             0
 Glucose
                            0
 BloodPressure
                            0
 SkinThickness
                            0
 Insulin
                            0
                            0
 DiabetesPedigreeFunction
                            0
                            0
 Age
 Outcome
                            0
```

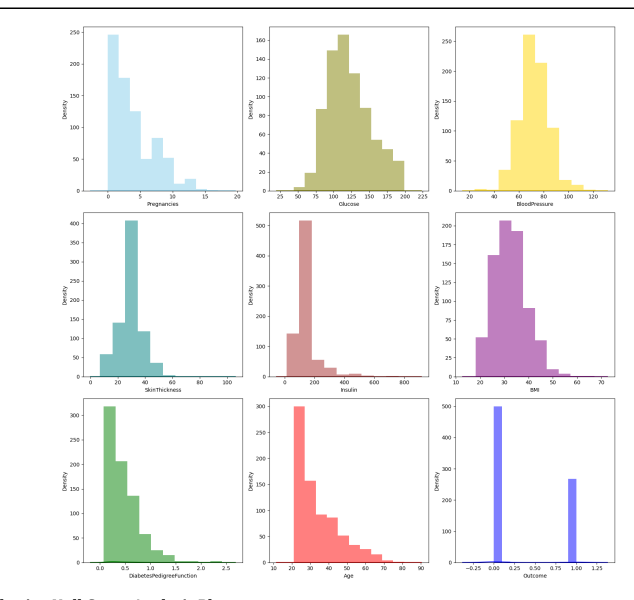
```
dtype: int64
    Zero values per column are:
    Pregnancies
                               111
   Glucose
                                5
                               35
   BloodPressure
   SkinThickness
                              227
   Insulin
                              374
   BMI
                               11
   DiabetesPedigreeFunction
                                0
   Age
                                0
                              500
   Outcome
   dtype: int64,
    data types of each column is:
    Pregnancies
                                 int64
   Glucose
                                int64
   BloodPressure
                                int64
   SkinThickness
                                int64
   Insulin
                                int64
   BMI
                              float64
   DiabetesPedigreeFunction
                              float64
                                int64
   Age
   Outcome
                                int64
   dtype: object
  #analyse the impossible values
  impossible_values = (df["Glucose"] == 0) | (df["BloodPressure"] == 0) |
  (df["SkinThickness"] == 0) | (df["Insulin"] == 0) | (df["BMI"] == 0)
  impossible_values.sum()
   376
Replace the null value with median value
  #replace the null value with median value
  lst=['Glucose', 'BloodPressure', 'SkinThickness', 'Insulin', 'BMI']
  for i in 1st:
      df[i].replace(0, np.nan, inplace=True)
      df[i].fillna(df[i].median(), inplace=True)
  df.head()
      Pregnancies Glucose BloodPressure SkinThickness Insulin
                                                                   BMI
   0
                     148.0
                                    72.0
                                                   35.0
                                                           125.0 33.6
                6
                                                           125.0 26.6
   1
                1
                                                   29.0
                     85.0
                                     66.0
   2
                8
                     183.0
                                     64.0
                                                   29.0
                                                           125.0 23.3
   3
                1
                     89.0
                                     66.0
                                                   23.0
                                                            94.0 28.1
                                                           168.0 43.1
                0
                     137.0
                                     40.0
                                                   35.0
```

	DiabetesPedigreeFunction	Age	Outcome
0	0.627	50	1
1	0.351	31	0
2	0.672	32	1
3	0.167	21	0
4	2.288	33	1

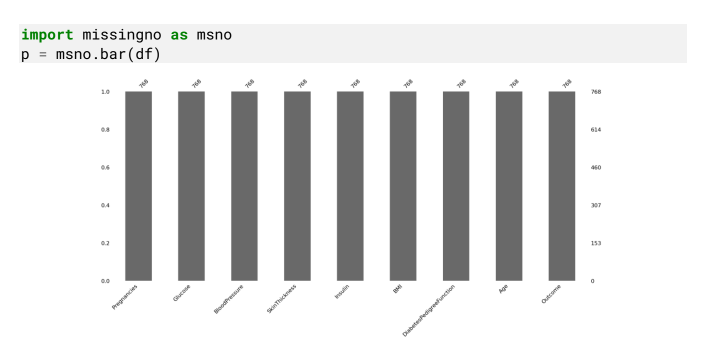
# 5. Visualize the dataset

### After clean the data visualize the data distribution by plotting it as histogram

```
lst1=['Pregnancies','Glucose','BloodPressure','SkinThickness','Insulin','BM
I','DiabetesPedigreeFunction','Age','Outcome']
lst_col=['skyblue','olive','gold','teal','brown','purple','green','red','bl
ue']
f, axes = plt.subplots(3, 3, figsize=(15, 15), sharex=False) # Set up the
matplotlib figure
axes = axes.flatten() # Plot a simple histogram with binsize determined
automatically
for ax,k,m in zip(axes,lst1,lst_col):
    ax.hist(df[k], color=m, bins=10, alpha=0.5)
    sns.distplot(df[k], color=m, ax=ax)
plt.tight_layout()
```



## **Plotting Null Count Analysis Plot**



Inference: Now in the above graph also we can clearly see that there are no null values in the dataset.

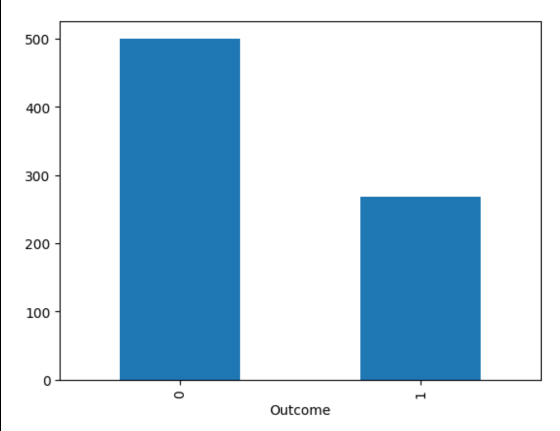
### check that how well our outcome column is balanced

dataset is completely imbalanced in fact the number of patients who are diabetic is half of the patients who are non-diabetic

```
color_wheel = {1: "#0392cf", 2: "#7bc043"}
colors = df["Outcome"].map(lambda x: color_wheel.get(x + 1))
print(df.Outcome.value_counts())
p=df.Outcome.value_counts().plot(kind="bar")
```

Outcome 0 500 1 268

Name: count, dtype: int64

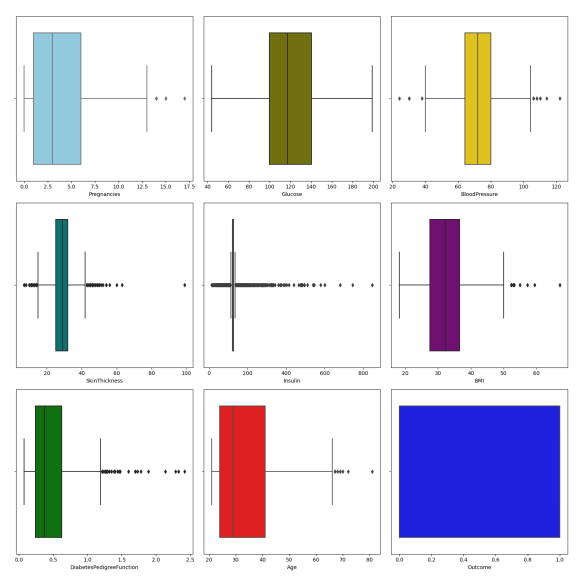


Interference:dataset is completely imbalanced in fact the number of patients who are diabetic is half of the patients who are non-diabetic

# Visualization of Outliers

```
# Generate a box plot for each feature
lst1=['Pregnancies','Glucose','BloodPressure','SkinThickness','Insulin','BMI',
'DiabetesPedigreeFunction','Age','Outcome']
lst_col=['skyblue','olive','gold','teal','brown','purple','green','red','blue'
```

```
f, axes = plt.subplots(3, 3, figsize=(15, 15), sharex=False) # Set up the
matplotlib figure
axes = axes.flatten() # Plot a simple histogram with binsize determined
automatically
for ax,k,m in zip(axes,lst1,lst_col):
    sns.boxplot(data=df, x=k, color=m, ax=ax)
plt.tight_layout()
plt.show()
```

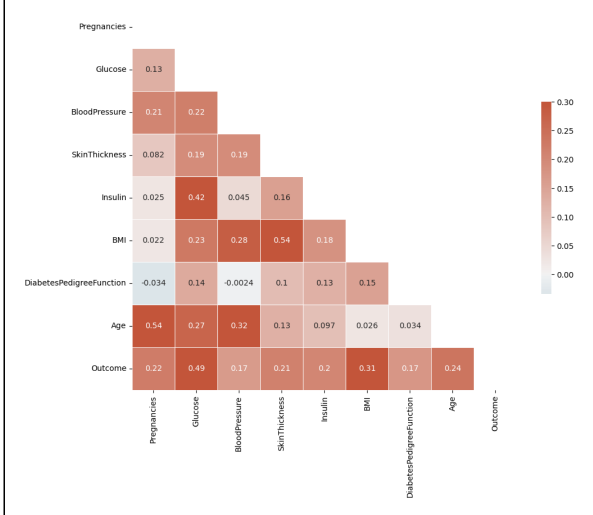


# Check the Relationship Between Variables

Then we wanna check the relationships between the different variables (columns). This can provide insights into which variables are strongly or weakly associated with each other.

```
corr = df.corr() # Compute the correlation matrix
mask = np.triu(np.ones_like(corr, dtype=bool)) # Generate a mask for the
```

```
upper triangle
f, ax = plt.subplots(figsize=(11, 9)) # Set up the matplotlib figure
cmap = sns.diverging_palette(230, 20, as_cmap=True) # Generate a custom
diverging colormap
sns.heatmap(corr, mask=mask, cmap=cmap, vmax=0.3, center=0, square=True,
linewidths=.5, cbar_kws={"shrink": .5}, annot=True) # Draw the heatmap with
the mask and correct aspect ratio
plt.tight_layout()
```



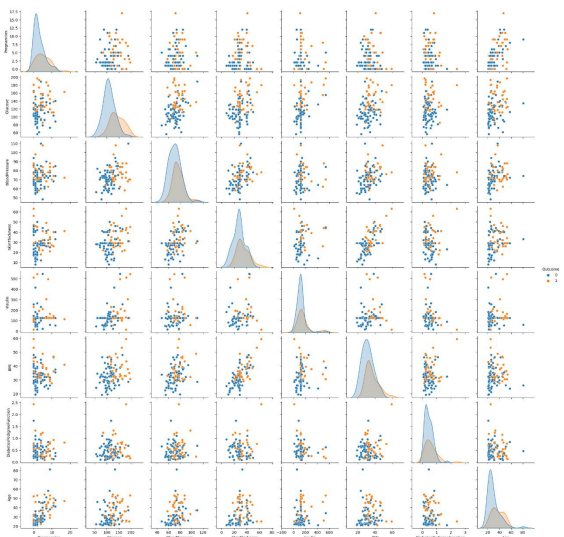
Interference: The correlation coefficient values range from -1 to 1. If the correlation coefficient is close to 1, it means that there is a strong positive correlation between the two variables. When it is close to -1, the variables have a strong negative correlation. Glucose, Age and BMI are moderately correlated with Outcome. Pregnancies and Age show a strong correlation.

# Pairwise Variable Relationship

This can be very helpful to understand how the variables interact with each other and identify any potential patterns or trends in the data.

```
# We'll use a sample of the data to make the pairplot faster to generate
df_sample = df.sample(100, random_state=1)
```





# 6.Split the data

we need to split the data into a training set and a test set. This allows us to evaluate how well our model generalizes to unseen data. We'll use 80% of the data for training and 20% for testing.

```
from sklearn.model_selection import train_test_split
X = df.drop("Outcome", axis=1)
y = df["Outcome"]
X_train, X_test, y_train, y_test = train_test_split(X, y, test_size=0.2, random_state=42)# Split the data into a training set and a test set
X_train.shape, X_test.shape
```

# 7. Data Transformation

## Transform data as needed, for example, scaling

## a)Before Scaling

```
df.head()
```

```
Pregnancies
                Glucose BloodPressure SkinThickness Insulin
                                                                  BMI
0
             6
                  148.0
                                  72.0
                                                  35.0
                                                          125.0
                                                                33.6
1
             1
                   85.0
                                  66.0
                                                  29.0
                                                          125.0
                                                                26.6
2
             8
                  183.0
                                  64.0
                                                  29.0
                                                          125.0
                                                                 23.3
             1
3
                                                  23.0
                   89.0
                                  66.0
                                                           94.0 28.1
4
             0
                  137.0
                                  40.0
                                                  35.0
                                                          168.0 43.1
   DiabetesPedigreeFunction Age Outcome
0
                      0.627
                              50
                      0.351
                                        0
1
                              31
2
                      0.672
                              32
                                        1
3
                              21
                                         0
                      0.167
4
                      2.288
                              33
```

### b) After Scaling

### Scaled it by trained model

```
from sklearn.preprocessing import StandardScaler
scaler = StandardScaler() # Initialize the scaler
scaler.fit(X_train) # Fit the scaler to the training data
X_train_scaled = scaler.transform(X_train)
X_test_scaled = scaler.transform(X_test)
pd.DataFrame(X_train_scaled, columns=['Pregnancies',
'Glucose', 'BloodPressure', 'SkinThickness', 'Insulin', 'BMI',
'DiabetesPedigreeFunction', 'Age']).head()
    Pregnancies
                 Glucose BloodPressure SkinThickness
                                                        Insulin
                                                                      BMI
 0
      -0.526397 -1.256881
                              -0.018995
                                             0.034298 -0.175620 -0.007450
 1
      1.588046 -0.326051
                               0.808174
                                            -0.560583 -0.175620 -0.599092
 2
      -0.828460 0.571536
                                            -1.155463 -0.652193 -0.526941
                              -2.169636
 3
      -1.130523 1.302903
                              -1.838768
                                             0.034298 -0.175620 -1.508200
 4
      0.681856 0.405316
                              0.642740
                                             0.986106 2.604392 1.998360
    DiabetesPedigreeFunction
                                  Age
 0
                  -0.490735 -1.035940
 1
                   2.415030 1.487101
 2
                   0.549161 -0.948939
```

```
3 -0.639291 2.792122
4 -0.686829 1.139095
```

### Scaled it by test model

```
X_test_scaled = scaler.transform(X_test)
pd.DataFrame(X_test_scaled, columns=['Pregnancies',
'Glucose', 'BloodPressure', 'SkinThickness', 'Insulin', 'BMI',
'DiabetesPedigreeFunction', 'Age']).head()
                  Glucose BloodPressure SkinThickness
    Pregnancies
                                                        Insulin
                                                                      BMI
                                                                           \
 0
       0.681856 -0.791466
                              -1.177033
                                              0.510202 0.561935 0.237865
 1
      -0.526397 -0.326051
                               0.229156
                                              0.391226 -0.175620 0.483180
                              -0.680731
0.642740
1.469910
      -0.526397 -0.459026
 2
                                              0.034298 -0.175620 -0.223904
 3
       1.285983 -0.492270
                                            0.034298 -0.175620 -1.118582
       0.983919 0.471804
                                              0.034298 -0.175620 -0.353777
    DiabetesPedigreeFunction
                                  Age
 0
                  -0.116372 0.878091
 1
                   -0.954231 -1.035940
 2
                  -0.924520 -1.035940
 3
                   1.149329 0.095078
 4
                   -0.770021 1.487101
```

### CONCLUSION:

Loading and preprocessing the dataset are crucial steps in building a prediction system. The use of Pandas for data loading and scikit-learn for preprocessing ensures efficiency and consistency in handling the data

These steps set the foundation for subsequent stages, such as model training and evaluation, in the development of a diabetes prediction system.

