AI BASED DIABETES PREDICTION SYSTEM

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AI _Phase 1 Submission

PROJECT: Creating an AI model predicting diabetes risk using medical data





OBJECTIVE:

The objective of this project is to develop a machine learning model that accurately predicts the diabetes risk using the given medical data.

PHASE 1: Problem Defining and Design Thinking

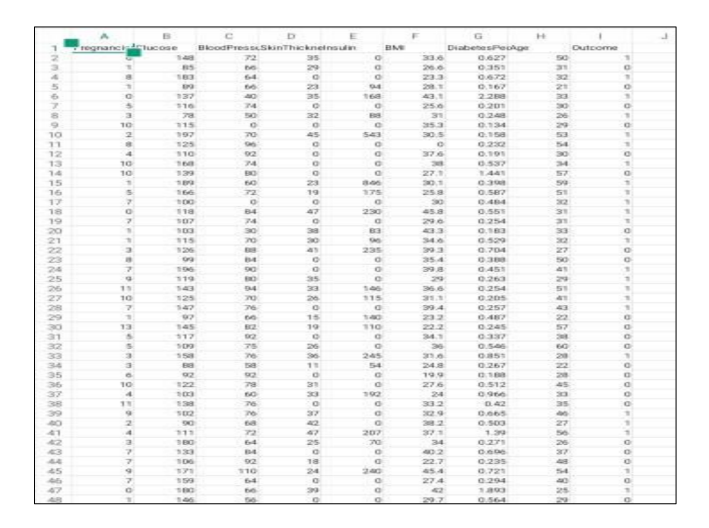
1.DATA COLLECTION

There are a variety of sources of data that can be used for an AI-based diabetes prediction system, including: Medical records: This data can provide information

about a patient's medical history, including demographics, lab results, medications, and diagnoses.

- Wearable devices: Wearable devices can track a variety of health metrics, such as blood sugar levels, heart rate, and activity levels.
- Patient surveys: Patient surveys can collect information about a patient's lifestyle, symptoms, and risk factors for diabetes.

It is important to collect data from a diverse population in order to train a model that is generalizable. The data should also be representative of the population that the model will be used to predict diabetes.



DATASETLINK:

https://www.kaggle.com/code/shobiarosalinmarriex/diabetesprediction-1e119d/edit

2.DATA PREPROCESSING

Once the data has been collected, it needs to be preprocessed to make it suitable for machine learning. This may involve the following steps:

- Cleaning the data: This involves removing errors and inconsistencies from the data. For example, you may need to remove missing values, correct typos, and standardize the format of the data.
- Scaling the data: This involves normalizing the values of different features so that they are on the same scale. This can be done using a variety of scaling techniques, such as min-max scaling and standard scaling.
- Encoding categorical features: Categorical features, such as gender and race, need to be encoded into numerical values before they can be used by machine learning algorithms. This can be done using a variety of encoding techniques, such as one-hot encoding and label encoding.

Once the data has been preprocessed, it is ready to be used to train a machine learning model to predict diabetes.

• Here are some additional tips for data collection and data preprocessing for an Al-based diabetes prediction system:

Use a large and diverse dataset. The more data you have, the better your model will be able to learn to predict diabetes. Additionally, it is important to use a dataset that is representative of the population that you are trying to predict diabetes for.

- Remove outliers. Outliers can skew the results of your model. It is important to remove outliers from the data before training the model.
- Use a variety of data preprocessing techniques. There is no one-size-fits-all approach to data preprocessing. It is important to use a variety of techniques to clean, scale, and encode the data in a way that is appropriate for your machine learning algorithm.

PYTHON PROGRAM:

INPUT AND OUTPUT:

import num p y as np # linear algebra

import pandas as pd # data processing, CSV file I/O (e g. pd. read_ csv)

import mat plot lib.py plot as p l t

import seaborn as s ns

In [2]:

dataset=pd.read_csv("/kaggle/input/diabetes-data-set/diabetes.csv")

In [3]:

dataset. Head ()

Out[3]: Pregnanc ies	Glucose	Blood Pressur	e	Skin Thickness	Insulin	BMI	Diabetes Pedigree Function	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

4]:

(768, 9)

In [5]:

#check if null value is present Dataset .is null(). Values .any()

Out[5]: False In [6]:

dataset.info()

<class 'pandas. core. frame. Data Frame'>

Range Index: 768 entries, 0 to 767 Data columns (total 9 columns): # Column Non-Null Count D type

--- -----

- 0 Pregnancies 768 non-null int64
- 1 Glucose 768 non-null int64
- 2 Blood Pressure 768 non-null int64
- 3 Skin Thickness 768 non-null int64
- 4 Insulin 768 non-null int64
- 5 BMI 768 non-null float64
- 6 Diabetes Pedigree Function 768 non-null float64
- 7 Age 768 non-null int64
- 8 Outcome 768 non-null int64

D types: float64(2), int64(7)

memory usage: 54.1 KB

In [7]:

dataset. describe()

Pregnanc ies	Glucose	Blood Pressur	Skin e Thickr	Insu ness	in BM		abetes edigree	Age	Outcome
						Fι	ınction		
count	768.000 000	768.000 000	768.000 000	768.000 000	768.000 000	768.000 000	768.000 000	768.000 000	768.000 000

Pregnanc	Glucose	Blood	Skin	Insul	in BMI	l Di	abetes	Age	Outcome
ies		Pressur	e Thickr	ness		Pe	digree		
						Fu	nction		
mean	3.84505	120.894	69.1054	20.5364	79.7994	31.9925	0.47187	33.2408	0.34895
	2	531	69	58	79	78	6	85	8
std	3.36957	31.9726	19.3558	15.9522	115.244	7.88416	0.33132	11.7602	0.47695
	8	18	07	18	002	0	9	32	1
min	0.00000	0.00000	0.00000	0.00000	0.00000	0.00000	0.07800	21.0000	0.00000
	0	0	0	0	0	0	0	00	0
25%	1.00000	99.0000	62.0000	0.00000	0.00000	27.3000	0.24375	24.0000	0.00000
	0	00	00	0	0	00	0	00	0
50%	3.00000	117.000	72.0000	23.0000	30.5000	32.0000	0.37250	29.0000	0.00000
	0	000	00	00	00	00	0	00	0
75%	6.00000	140.250	80.0000	32.0000	127.250	36.6000	0.62625	41.0000	1.00000
	0	000	00	00	000	00	0	00	0
max	17.0000	199.000	122.000	99.0000	846.000	67.1000	2.42000	81.0000	1.00000
	00	000	000	00	000	00	0	00	0

Dataset is null().sum()

Out[8]: Pregnancies 0

Glucose 0

Blood Pressure 0

Skin Thickness 0

Insulin 0

BMI 0

Diabetes Pedigree Function 0

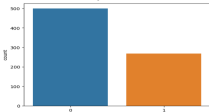
Age 0 Outcome 0 D type: int64

In [9]:

#data visualization

S ns. Count plot(x = 'Outcome', data = dataset)

<Axes: x label='Outcome', y label='count>

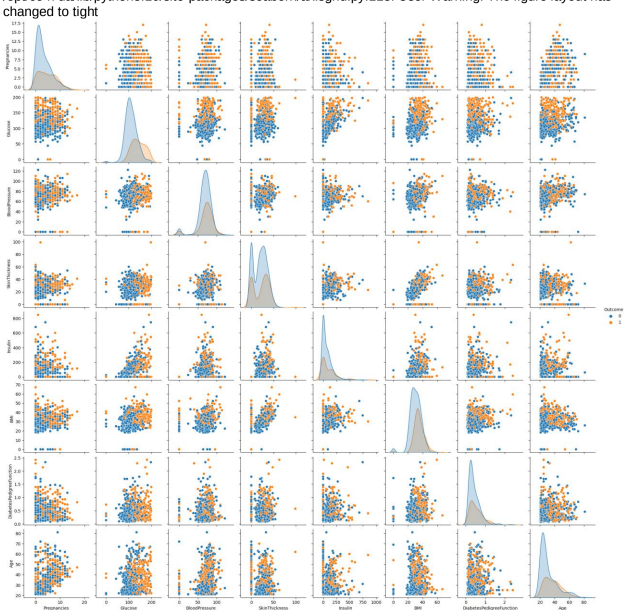


In [10]: # Pair plot

S ns. pair plot(data = dataset, hue = 'Outcome')

P It. show()

/opt/co n da/lib/python3.10/site-packages/seaborn/axisgrid.py:118: User Warning: The figure layout has



self._ figure. tight_ layout(*a r g s, **k w a r g s)

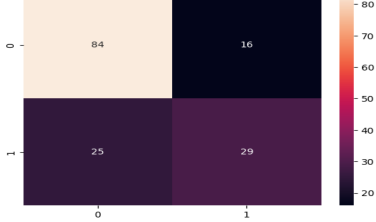
Heatmap S ns. heat map(data set .c o r r(), a n not = True) P lt. show()

```
Pregnancies - 1 0.13 0.14 -0.082-0.074 0.018-0.034
           0.082 0.057 0.21 1 0.44 0.39 0.18 -0.11 0.075
                 0.33 0.089 0.44 1 0.2 0.19 0.042 0.13
              34 0.14 0.041 0.18 0.19 0.14 1 0.034 0.17
             54 0.26 0.24 -0.11 -0.042 0.036 0.034 1
```

```
In [12]:
# Replacing zero values with Na N
Data set  new = dataset
dataset_new[["Glucose", "Blood Pressure", "Skin Thickness", "Insulin", "BMI"]] = dataset_new[["Glucose",
"Blood Pressure", "Skin Thickness", "Insulin", "BMI"]].replace(0, np.NaN)
# Count of Na N
dataset_ new. Is null(). sum()
Out[13]:
Pregnancies 0
Glucose 5
Blood Pressure 35
Skin Thickness 227
Insulin 374
BMI 11
Diabetes Pedigree Function 0
Age 0
Outcome 0
D type: int64
In [14]:
# Replacing Na N with mean values
dataset new["Glucose"].fill n a(dataset new["Glucose"].mean(), in place = True)
dataset_new["BloodPressure"].fillna(dataset_new["BloodPressure"].mean(), in place = True)
dataset_new["SkinThickness"].fillna(dataset_new["SkinThickness"].mean(), in place = True)
dataset_ new["Insulin"].fill n a(dataset_ new["Insulin"].mean(), in place = True)
dataset_ new["BMI"] .fill n a (dataset_ new["BMI"].mean(), in place = True)
In [15]:
dataset_ new is null (). Sum ()
Out[15]:
Pregnancies 0
Glucose 0
Blood Pressure 0
Skin Thickness 0
Insulin 0
BMI<sub>0</sub>
Diabetes Pedigree Function 0
Age 0
Outcome 0
D type: int64
```

In [16]: #Logistic regression y = dataset_ new['Outcome'] X = dataset_ new. drop('Outcome', axis=1) In [17]:

```
# Splitting X and Y
from s k learn. 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'])
In [18]:
from s k learn. linear model import Logistic Regression
model = Logistic Regression()
model. fit(X_ train, Y_ train)
y_ predict = model. predict(X_ test)
/opt/conda/lib/python3.10/site-packages/sklearn/linear_model/_logistic.py:458: Convergence
Warning: I b f g s failed to converge (status=1):
STOP: TOTAL NO. of ITERATIONS REACHED LIMIT.
Increase the number of iterations (max_ I ter) 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-regression
n Iter i = check optImIz result(
In [19]:
v predict
Out[19]:
array([1, 0, 0, 0, 0, 0, 0, 1, 0, 1, 0, 0, 0, 0, 0, 0, 1, 0, 1, 0, 0, 1,
0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 0, 1, 1, 0, 0, 0, 1, 1, 0, 0, 0, 0, 0
0, 0, 0, 0, 1, 0, 0, 1, 0, 0, 0, 1, 0, 1, 0, 1, 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, 0, 1, 1, 0, 0, 0, 1, 1, 1, 1, 0, 0, 0, 0, 0, 1, 0, 1, 0, 1, 1,
0, 1, 0, 0, 0, 0, 0, 1, 0, 1, 0, 0, 1, 0, 1, 1, 1, 1, 0, 0, 0, 1,
0, 1, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 1, 0, 0, 0, 1, 0]
In [20]:
# Confusion matrix
from s k learn . metrics import confusion_ matrix
cm = confusion matrix (Y test, y predict)
cm
Out[20]:
array([[84, 16],
[25, 29]])
In [21]:
# Heatmap of Confusion matrix
S ns .heat map(pd .Data Frame(cm), anno t=True)
Out[21]:
                                                                        80
                                                                        70
                                                   16
```



<Axes: >

```
from s k l ear . metrics import accuracy_ score
In [23]:
accuracy =accuracy_ score(Y_ test, y_ predict)
accuracy
Out[23]:
0.7337662337662337
In [24]:
#Example: Let's check whether the person have diabetes or not using some random values
y_predict = model.predict([[1,148,72,35,79.799,33.6,0.627,50]])
print (y_ predict)
if y_ predict==1:
print("Diabetic")
else:
print (" Non Diabetic ")
[1]
Diabetic
/opt /co n da /lib/python3.10/site-packages/s k learn/base.py:439: User Warning: X does not have
valid feature names, but Logistic Regression was fitted with feature names
Warnings warn (
In[]:
```

3.FEATURE SELECTION

We will select relevant features that can impact diabetes risk prediction.

Feature selection is the process of identifying and selecting the most relevant and informative features from a dataset for use in a machine learning model. This is an important step in developing an Al-based diabetes prediction system, as it can help to improve the accuracy and efficiency of the model.

There are a number of different feature selection algorithms that can be used, and the best algorithm to use will depend on the specific dataset and machine learning model being used. Some common feature selection algorithms include:

- Filter methods: These methods rank features based on a statistical measure of their importance, such as information gain or correlation.
- Wrapper methods: These methods evaluate features by training and testing a machine learning model on different subsets of features.
- Embedded methods: These methods incorporate feature selection into the machine learning process itself.

4.MODEL SELECTION

Model selection is the process of choosing the best machine learning model for a given dataset and task. In the context of AI-based diabetes prediction, model

selection involves evaluating the performance of different machine learning models on a held-out test set.

Once the most relevant features have been selected, they can be used to train a machine learning model to predict diabetes. There are a number of different machine learning models that can be used for diabetes prediction, such as:

- Logistic regression: This is a simple but effective model for binary classification tasks, such as predicting whether or not someone has diabetes.
- Support vector machines (SVMs): SVMs are a powerful machine learning model that can be used for both classification and regression tasks.
- Decision trees: Decision trees are a type of machine learning model that can be used to create rules for predicting diabetes.
- Random forests: Random forests are an ensemble machine learning model that combines the predictions of multiple decision trees to improve accuracy.

5.EVALUATION:

Some common metrics used to evaluate the performance of machine learning models for diabetes prediction include:

- Accuracy: This metric measures the proportion of predictions that are correct.
- Sensitivity: This metric measures the proportion of people with diabetes who are correctly predicted to have diabetes.
- Specificity: This metric measures the proportion of people without diabetes who are correctly predicted not to have diabetes.
- AUC-ROC: This metric measures the area under the receiver operating characteristic (ROC) curve, which is a plot of sensitivity against 1 specificity.

The best machine learning model for AI-based diabetes prediction will depend on the specific dataset and the desired trade-off between sensitivity and specificity.

6.ITERATIVE IMPROVEMENT:

We will fine tune the model parameters and explore techniques like feature engineering to enhance prediction accuracy.

ADDITIONAL CONSIDERATION:

In addition to the tasks listed above, there are a few other things to keep in mind when collecting and preprocessing data for an AI-based diabetes prediction system:

- a)Data privacy and security: It is important to ensure that the data is collected and stored in a secure manner, and that the privacy of the patients is protected.
- b)Data bias: It is important to be aware of the potential for bias in the data. For example, if the data is collected from a single hospital or clinic, it may be biased towards patients with certain characteristics.

Data imbalance: Diabetes is a relatively rare disease, so it is common for datasets of patients with and without diabetes to be imbalanced. This means that there are many more patients without diabetes than with diabetes. Imbalanced datasets can make it difficult for machine learning models to learn to predict diabetes accurately.

There are a variety of ways to handle imbalanced datasets, such as oversampling the minority class (patients with diabetes) or under sampling the majority class (Patients without diabetes).

By following these guidelines, you can collect and preprocess data for an AI-based diabetes prediction system that is high quality and suitable for machine learning.

CONCLUSION:

It is also important to note that AI-based diabetes prediction systems should be carefully evaluated and validated before being used in clinical practice. This is because these systems can have a significant impact on people's lives, and it is important to ensure that they are accurate and reliable. In Phase 1, we have established a clear understanding of our goal: to predict diabetes of given data using machine language. We have outlined a structured approach that includes data collection, data preprocessing, feature selection, model selection, evaluation and iterative improvement. This sets the stage for our project's successful execution in subsequent phases.