

# AWS Machine Learning Engineer Nanodegree

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## 1 Definition

## 1.1 Project Overview

Electronic Health records or Electronic Medical Records data is the data being collected when we see a doctor, pick up a prescription at the pharmacy, or even from a visit to the dentist.

This data is used for a variety of use-cases. From personalizing healthcare to discovering novel drugs and treatments to helping providers diagnose patients better and reduce medical errors.

Diabetes mellitus, or simply diabetes, is a leading non-communicable disease (NCD) globally, almost doubling in cases since 1980. It is a chronic illness that develops either when the pancreas are not able to generate sufficient insulin or when the body does not utilize the insulin produced effectively. There is no cure for this disease. Diabetes is thought to result from a combination of genetic and environmental factors. Several risk factors that are attributed to diabetes include ethnicity, family history of diabetes, age, excess weight, unhealthy diet, physical inactivity, and smoking. In addition to this, the absence of early detection of diabetes has been known to contribute to the development of other chronic diseases such as kidney disease.

#### 1.2 Dataset

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 datasets consists of several medical predictor variables and one target variable, Outcome. Predictor variables includes the number of pregnancies the patient has had, their BMI, insulin level, age, and so on<sup>1</sup>.

#### 1.3 Problem Statement

Can you build a machine learning model to accurately predict whether or not the patients in the dataset have diabetes or not?

#### 1.4 Metrics

#### 1.4.1 AUC

AUC is used for binary classification, multiclass classification, and ranking problems. AUC measures the proportion of correctly ordered objects and the capability of the model to distinguish between the classes.

The AUC has an important statistical property: the AUC of a classifier is equivalent to the probability that the classifier will rank a randomly chosen positive instance higher than a randomly chosen negative instance.<sup>2</sup>

AUC is the Area Under the ROC Curve. The best AUC = 1 for a model that ranks all the objects right (all objects with class 1 are assigned higher probabilities then objects of class 0). AUC for the 'bad' classifier which is working as random guessing is equal to  $0.5.^2$ 

The ROC curve shows the model's ability to distinguishing between classes.

The model which randomly assigns a class to object is a 'bad' classifier and has a diagonal ROC curve. The better is the classifier, the higher is the ROC curve. The ROC curve is plotted with TPR, True Positive Rate, on the y-axis against the FPR, False Positive Rate, on the x-axis. The curve also could be interpreted in terms of Sensitivity and Specificity of the model with Sensitivity on the y-axis and (1-Specificity) on the x-axis.

Building and visualizing the ROC curve could be used to measure classification algorithm performance with different probability boundaries and select the probability boundary required to achieve the specified false-positive or false-negative rate.<sup>2</sup>

## 2 Analysis

Data was stored in AWS S3 cloud service for training and evaluation of the model.

```
train_file = "train.csv"
train.to_csv(train_file, index=False)
train_s3_path = session.upload_data(train_file, key_prefix="{}/data".format(prefix))

test_file = "test.csv"
test.to_csv(test_file, index=False)
test_s3_path = session.upload_data(test_file, key_prefix="{}/data".format(prefix))

X_test_file = "X_test.csv"
X_test_file = "X_test.csv"
X_test.to_csv(X_test_file, index=False)
X_test_s3_path = session.upload_data(X_test_file, key_prefix="{}/data".format(prefix))
```

### 2.1 Data Exploration

DataFrame.describe() method generates descriptive statistics that summarize the central tendency, dispersion and shape of a dataset's distribution, excluding NaN values. This method tells us a lot of things about a dataset. One important thing is that the describe() method deals only with numeric values. It doesn't work with any categorical values. So if there are any categorical values in a column the describe() method will ignore it and display summary for the other columns unless parameter include="all" is passed.

- count tells us the number of NoN-empty rows in a feature.
- mean tells us the mean value of that feature.
- std tells us the Standard Deviation Value of that feature.
- min tells us the minimum value of that feature.
- 25%, 50%, and 75% are the percentile/quartile of each features. This quartile information helps us to detect Outliers.
- max tells us the maximum value of that feature.

The figure 1 below shows the summary of the describe function and also the information about the dataset

The results of this summary of describe function. There are some value of below listed columns have zero minimum, the value of zero does indicates missing value. Following columns or variables have an invalid zero value:

- Glucose
- BloodPressure
- SkinThickness
- Insulin
- BMI

## 2.2 Exploratory Visualization

#### 2.2.1 Data Distribution

A left-skewed distribution has a long left tail. Left-skewed distributions are also called negatively-skewed distributions. That's because there is a long tail in the negative direction on the number line. The mean is also to the left of the peak.

Figure 2 below shows data distribution.

Aright-skewed distribution has a long right tail. Right-skewed distributions are also called positive-skew distributions. That's because there is a long tail in the positive direction on the number line. The mean is also to the right of the peak.

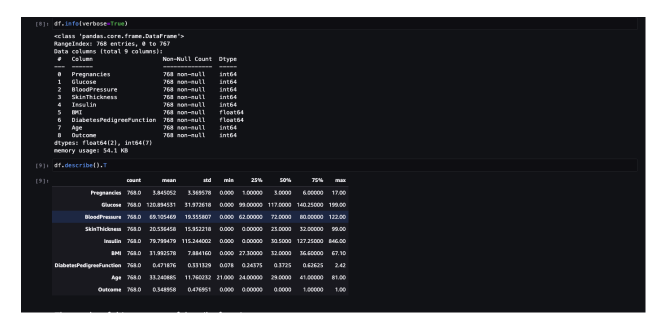


Figure 1: The describe function results

This data is mostly left skewed for example the pregrancy plot, insulin, age and diabetes pedigree function.

#### 2.2.2 Correlation visualisation

Heatmap is good method to visualize correlation between features. This heatmap helps to know the following pairs had a positive correlation coefficient between them as compared to other parameters.

- Pregnancies and age
- Insulin and Skin thickness
- BMI and Skin thickness
- Insulin and Glucose

Glucose and BMI values are related the most. This indicates the two parameters need special attention.

The heatmap figure 3 shows correlation coeficient

#### 2.2.3 True Diabetes Distribution

A violin plot is a hybrid of a box plot and a kernel density plot, which shows peaks in the data. It is used to visualize the distribution of numerical data. Unlike a box plot that can only show summary statistics, violin plots depict summary statistics and the density of each variable.<sup>3</sup>

To understand data we have to plot data using violin visualization. This plotting shows where the outcome of diabetes was 1. This shows the distribution of the diabetes resulting from different factors. This gives us clear picture that Glucose, BMI and Insulin had the most effect on the outcome value. This also determines on the data division for model training and model testing. The median age is 35 of those having diabetes according the violin plot with the higher probability. Figure 4 shows the violin plots where the outcome is 1

#### 2.2.4 False Diabetes Distribution

To understand data we have to plot data using violin visualization. This plotting shows where the outcome of diabetes was 0. Figure 5 shows the violin plots where the outcome is 0

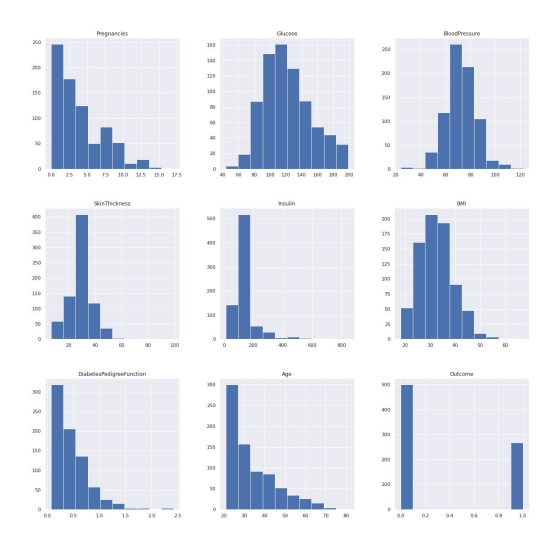


Figure 2: Data Distribution

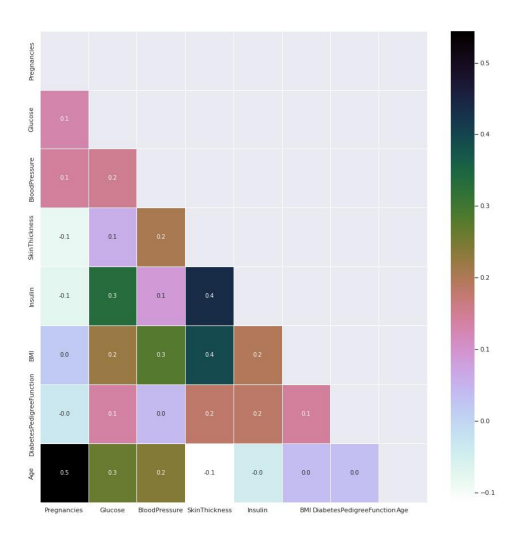


Figure 3: Heat map Correlation Coeffi

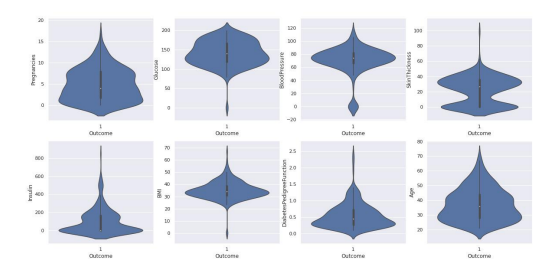


Figure 4: True Diabetes

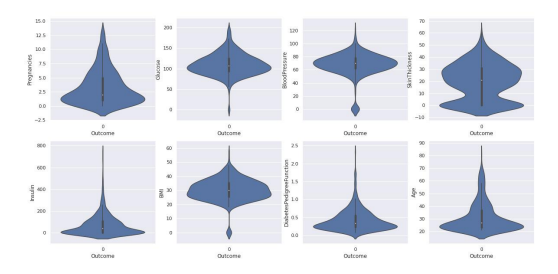


Figure 5: False Diabetes

## 2.3 Algorithms and Techniques

AutoGluon automates machine learning tasks enabling you to easily achieve strong predictive performance in your applications. With just a few lines of code, you can train and deploy high-accuracy machine learning and deep learning models on image, text, time series, and tabular data.

AutoGluon enables easy-to-use and easy-to-extend AutoML with a focus on automated stack ensembling, deep learning, and real-world applications spanning image, text, and tabular data.<sup>4</sup>

## 3 Methodology

### 3.1 Data Preprocessing

Data was diveded into training set and testing set. The training set was 70 percent of the whole data while the testing set was 30 percent of the data.

```
train = df.sample(frac=0.7, random_state=42)
test = df.drop(train.index)

label = "Outcome"
y_test = test[label]
X_test = test.drop(columns=[label])
```

## 3.2 Implementation

The platfrom for implimentation was AWS sagemaker studio. The data was stored in AWS S3 after it was devided for training set and testing set. The procedure for that was to create an S3 bucket using a script.

```
import os
   import sys
   import boto3
   import sagemaker
   from time import sleep
   from collections import Counter
   import numpy as np
   import pandas as pd
   from autogluon.tabular import TabularPredictor
   from sagemaker import get_execution_role, local, Model, utils, s3
   from sagemaker.estimator import Estimator
   from sagemaker.predictor import Predictor
   from sagemaker.serializers import CSVSerializer
   from sagemaker.deserializers import StringDeserializer
   from sklearn.metrics import accuracy_score, classification_report
   from IPython.core.display import display, HTML
   from IPython.core.interactiveshell import InteractiveShell
18
   import matplotlib.pyplot as plt
19
   import seaborn as sns
20
   sns.set()
21
   import warnings
   warnings.filterwarnings('ignore')
23
   import missingno as msno
24
   %matplotlib inline
25
   session = sagemaker. Session()
   bucket = session.default_bucket()
   prefix = "sagemaker/autogluon-tabular"
   region = session.boto_region_name
  role = get_execution_role()
```

```
client = session.boto_session.client("sts", region_name=region, endpoint_url=utils.
sts_regional_endpoint(region))
```

#### 3.2.1 Training

The model was trained using AUTOML autogluon<sup>4</sup>

## 4 Results

The results of Autogluon training can be seen in the summary below.

```
predictor.fit summary(show plot=True)
   *** Summary of fit() ***
   Estimated performance of each model:
                                       fit_time pred_time_val_marginal
   model score_val pred_time_val
       fit_time_marginal stack_level
                                       can infer fit order
   0
         WeightedEnsemble_L2
                                0.849699
                                               0.910417
                                                          242.822065
      0.000679
                                               2
                          0.457114
                                                       True
           LightGBMXT_BAG_L1
                                0.848956
                                                0.067625
                                                           41.802209
   1
      0.067625
                         41.802209
                                                       True
      NeuralNetFastAI_BAG_L1
                                0.831127
                                                0.364630
                                                          104.587116
      0.364630
                        104.587116
                                               1
                                                       True
   3
       NeuralNetTorch\_BAG\_L1
                                0.830445
                                                0.477483
                                                           95.975625
10
      0.477483
                                                                      3
                         95.975625
                                                       True
   Number of models trained: 4
   Types of models trained:
12
   { 'StackerEnsembleModel_TabularNeuralNetTorch', 'WeightedEnsembleModel', '
13
       StackerEnsembleModel_NNFastAiTabular', 'StackerEnsembleModel_LGB'}
   Bagging used: True (with 5 folds)
   Multi-layer stack-ensembling used: False
   Feature Metadata (Processed):
   (raw dtype, special dtypes):
   ('float', []) : 2 | ['BMI', 'DiabetesPedigreeFunction']
18
                 : 6 | ['Pregnancies', 'Glucose', 'BloodPressure', 'SkinThickness', '
   ('int', [])
19
       Insulin', ...]
   Plot summary of models saved to file: AutogluonModels/ag-20230310_110459/
      SummaryOfModels.html
   *** End of fit() summary ***
```

### 4.1 Model Evaluation

The model was evaluated by using predctor.evaluate command which is a built in command from the autogluon model. The result of the execution is shown in the script below.

```
predictor.evaluate(test_s3_path)
```

```
Loaded data from: s3://sagemaker-us-east-1-495962688195/sagemaker/autogluon-tabular/
       data/test.csv | Columns = 9 / 9 | Rows = 230 \rightarrow 230
   Evaluation: roc_auc on test data: 0.8274792522424343
   Evaluations on test data:
       "roc auc": 0.8274792522424343,
       "accuracy": 0.7478260869565218,
       "balanced accuracy": 0.6872327940313522,
       "mcc": 0.41186364034362855,
       "f1": 0.5735294117647058,
11
       "precision": 0.6842105263157895,
       "recall": 0.4936708860759494
14
   { 'roc auc': 0.8274792522424343,
     accuracy ': 0.7478260869565218,
    'balanced accuracy': 0.6872327940313522,
17
    'mcc': 0.41186364034362855,
18
    'f1': 0.5735294117647058,
    'precision': 0.6842105263157895,
20
    'recall': 0.4936708860759494}
```

- The roc-auc is 82 percent there is diabetes. In machine learning, we use ROC Curves to analyze the predictive power of a classifier: they provide a visual way to observe how changes in our model's classification thresholds affect our model's performance.
- Accuracy 75 percent shows that there is diabetes
- Balanced Accuracy 69 percent there is diabetes
- F1 score is 0.57 showing the good performance of the model. The F1-score is a great way to compare the performance of multiple classifiers

## 5 Conclusion

AutoML frameworks offer an enticing alternative. For the novice, they remove many of the barriers of deploying high performance ML models. For the expert, they offer the potential of implementing best ML practices only once (including strategies for model selection, ensembling, hyperparameter tuning, feature engineering, data preprocessing, data splitting, etc.), and then being able to repeatedly deploy them. This allows experts to scale their knowledge to many problems without the need for frequent manual intervention

## References

- 1. Aamna. HW1 machine learning for EHR. (2023).
- 2. Volodkevich, A. AUC and its implementation in CatBoost, medium. Towards data science.
- 3. Carron, J. Violin plots 101: Visualizing distribution and probability density.
- 4. Autogluon. AutoML for text, image, time series, and tabular data AutoGluon documentation 0.7.0 documentation (2023).