Diabetes Diagnosis Project

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1. Introduction

Diabetes is a very common disease that affects many people around the world. Its incidence is increasing at an alarming rate every year. The medical professionals need some reliable predictive systems for diagnosing diabetes. Machine learning is one of the applied technologies and methods in medical diagnosis.

This report describes the process of building a machine learning model for this project, including the analysis of the dataset and necessary pre-processing, the use of different methods and algorithms for model building, and a comparison of the models and their predictive accuracy.

We will use the Pima Indians Diabetes Database, acquired from Kaggle. The dataset consists of several medical predictor (independent) variables and one target (dependent) variable, Outcome. Independent variables include the number of pregnancies the patient has had, their BMI, insulin level, age, and so on.

The goal of this diabetes diagnosis project is to build a machine learning model to predict as accurately as possible whether a patient in a dataset has diabetes? And we try to make the accuracy of the model as high as possible, and then select the model with the highest prediction accuracy as the final model.

2. Importing the libraries and dataset

```
library(tidyverse)
library(caret)
library(data.table)
library(ggplot2)
library(matrixStats)
library(corrplot)
library(factoextra)
library(randomForest)
library(GGally)
library(gam)
```

The dataset we use is the Pima Indians Diabetes Database, downloaded from Kaggle:

https://www.kaggle.com/datasets/uciml/pima-indians-diabetes-database/download/

```
# importing the dataset
urlfile = "https://raw.githubusercontent.com/dapingz/Diabetes-Project/main/diabetes.csv"
# loading the dataset as data frame
diab_dat <- as.data.frame(read_csv(url(urlfile)))</pre>
```

We now display the first 10 rows of the dataset in the table below:

```
knitr::kable(head(diab_dat, 10))
```

Pregnancies	Glucose	BloodPressure Skir	nThickness 1	Insulin	BMI	DiabetesPedigreeFunction	A ge	Outcome
6	148	72	35	0	33.6	0.627	50	1
1	85	66	29	0	26.6	0.351	31	0
8	183	64	0	0	23.3	0.672	32	1
1	89	66	23	94	28.1	0.167	21	0
0	137	40	35	168	43.1	2.288	33	1
5	116	74	0	0	25.6	0.201	30	0
3	78	50	32	88	31.0	0.248	26	1
10	115	0	0	0	35.3	0.134	29	0
2	197	70	45	543	30.5	0.158	53	1
8	125	96	0	0	0.0	0.232	54	1

The dataset has 768 rows and 9 columns. Each row contains a different patient and their personal details, such as BMI, insulin levels, age, pregnancy, and diabetes status.

```
str(diab_dat)
```

```
## 'data.frame':
                   768 obs. of 9 variables:
##
   $ Pregnancies
                             : num 6 1 8 1 0 5 3 10 2 8 ...
  $ Glucose
                             : num 148 85 183 89 137 116 78 115 197 125 ...
##
  $ BloodPressure
                             : num 72 66 64 66 40 74 50 0 70 96 ...
## $ SkinThickness
                             : num 35 29 0 23 35 0 32 0 45 0 ...
## $ Insulin
                             : num 0 0 0 94 168 0 88 0 543 0 ...
## $ BMI
                             : num 33.6 26.6 23.3 28.1 43.1 25.6 31 35.3 30.5 0 ...
## $ DiabetesPedigreeFunction: num 0.627 0.351 0.672 0.167 2.288 ...
                             : num 50 31 32 21 33 30 26 29 53 54 ...
## $ Age
## $ Outcome
                             : num 1 0 1 0 1 0 1 0 1 1 ...
```

We also look at how many people have diabetes and how many do not. and the proportion of "no" and "yes"

```
table(diab_dat$Outcome)
```

```
prop.table(table(diab_dat$Outcome))
```

3. Data Pre-processing & Data Explorations

3.1 Convert the target variable type

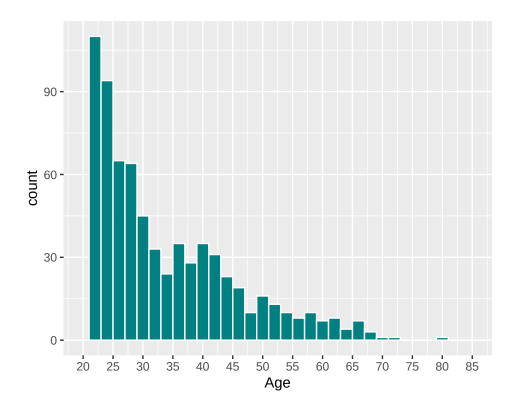
We convert the Outcome to factor, while changing '1' and '0' to 'yes' and 'no' to be more meaningful.

```
diab_dat$Outcome <- as.factor(ifelse(diab_dat$Outcome == "1", "Yes", "No"))</pre>
```

3.2 Age Analysis and grouping ages

Have a look at the Age variable. As can be seen from the distribution in the figure below, the age distribution is uneven, and there are too many young people in the data set.

```
ggplot(aes(x = Age), data = diab_dat) +
geom_histogram(binwidth = 2 , color = "white", fill = "#008080") +
scale_x_continuous(limits = c(20,85), breaks = seq(20,85,5))
```



Let's divide age into different age groups

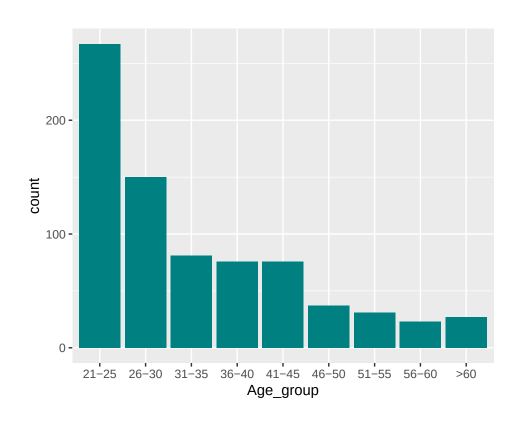
```
ifelse((diab_dat2$Age>35) & (diab_dat2$Age<=40), "36-40",
    ifelse((diab_dat2$Age>40) & (diab_dat2$Age<=45), "41-45",
    ifelse((diab_dat2$Age>45) & (diab_dat2$Age<=50), "46-50",
    ifelse((diab_dat2$Age>50) & (diab_dat2$Age<=55), "51-55",
        ifelse((diab_dat2$Age>55) & (diab_dat2$Age<=60), "56-60",
        ">60"))))))))))
```

```
# Convert Age_group to factor
diab_dat2$Age_group <- factor(diab_dat2$Age_group,
    levels=c('<21','21-25','26-30','31-35','36-40','41-45','46-50','51-55','56-60','>60'))

# Barplot by Age_group
```

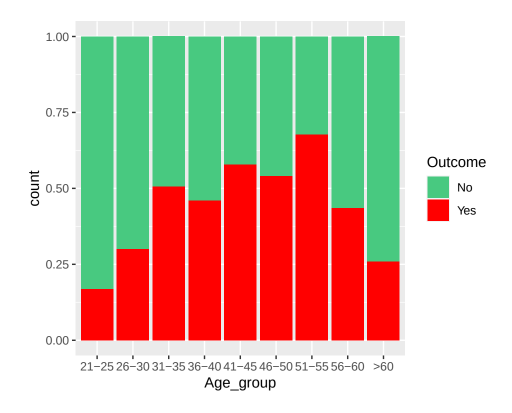
ggplot(aes(x = Age_group), data = diab_dat2) +

geom_bar(fill='#008080')



From the plot above we can see that people aged 21-25 are the most in the dataset. Next, we have a look at the portion of the people who have been diagnosed diabetes by age group

```
ggplot(diab_dat2, aes(x = Age_group, fill = Outcome)) +
  geom_bar(position = "fill") +
  scale_fill_manual(values = c("No"="#48C980", "Yes"="red"))
```



The plot above shows that the 40-50 age group has the highest rate of diabetes diagnoses, followed by the 30-35 and 35-40 age groups.

3.3 Checking missing values

First check if there is any null values in the dataset

[1] 0

There are no null values in the dataset, but we see that there are some zero values in the dataset.

knitr::kable(head(diab_dat))

Pregnancies	Glucose	BloodPressure Skir	Thickness I	nsulin	BMI	DiabetesPedigreeFunction	A ge	Outcome
6	148	72	35	0	33.6	0.627	50	Yes
1	85	66	29	0	26.6	0.351	31	No
8	183	64	0	0	23.3	0.672	32	Yes
1	89	66	23	94	28.1	0.167	21	No
0	137	40	35	168	43.1	2.288	33	Yes
5	116	74	0	0	25.6	0.201	30	No

A value of zero does not make sense and thus indicates missing value. We will replace these zeros with the median of each column.

```
# step1: replace all zeros with NAs
diab_dat[diab_dat == 0] <- NA

# step2: replace all "NA"s with the median of each column
diab_dat <- diab_dat %>%
  mutate_if(is.numeric, function(x) ifelse(is.na(x), median(x, na.rm = T), x))
```

Now showing the dataset again, we can see that all columns containing "0" are now replaced with median.

```
knitr::kable(head(diab_dat))
```

Pregnancies	Glucose	BloodPressure Skin	Thickness I	nsulin	BMI	${\bf Diabetes Pedigree Functio \textbf{A}}$	ge	Outcome
6	148	72	35	125	33.6	0.627	50	Yes
1	85	66	29	125	26.6	0.351	31	No
8	183	64	29	125	23.3	0.672	32	Yes
1	89	66	23	94	28.1	0.167	21	No
4	137	40	35	168	43.1	2.288	33	Yes
5	116	74	29	125	25.6	0.201	30	No

3.4 Splitting the dataset into independent and dependent datasets

We use row 1 to 8 as independent dataset and covert it to a matrix and use the Outcome as dependent dataset

```
diab_x <- as.matrix(diab_dat[,1:8])
diab_y <- diab_dat$Outcome</pre>
```

For independent dataset: there are 768 samples and 8 predictors in the dataset (features matrix).

```
dim(diab_x)[1]
```

[1] 768

```
dim(diab_x)[2]
```

[1] 8

For dependent dataset: there 768 outcomes in the dataset (target variable).

```
length(diab_y)
```

[1] 768

Check the proportions of No and Yes samples in the dataset.

A. See how many patients are in each category (No or Yes).

```
summary(diab_y)
```

```
## No Yes
## 500 268
```

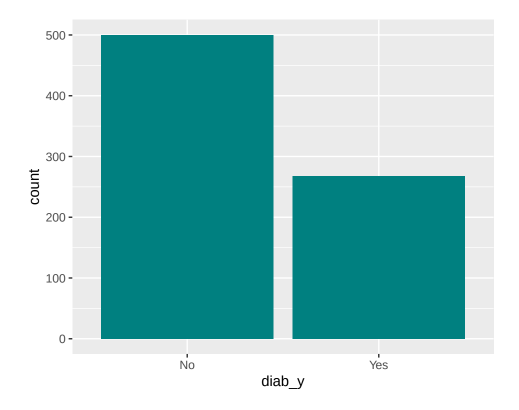
B. See what the proportion of No and Yes in a table.

```
prop.table(table(diab_y))
```

```
## diab_y
## No Yes
## 0.6510417 0.3489583
```

C. See the proportion of No and Yes on the bar plot

```
data.frame(diab_y) %>%
  ggplot(aes(diab_y)) +
  geom_bar(fill='#008080')
```



3.5 Feature Scaling and Splitting Datasets

```
# Scaling all predictor variables
x_centered <- sweep(diab_x, 2, colMeans(diab_x))
x_scaled <- sweep(x_centered, 2, colSds(diab_x), FUN = "/")
# After scaling, the standard deviation is 1 for all columns
colSds(x_scaled)</pre>
```

[1] 1 1 1 1 1 1 1 1

summary(x scaled)

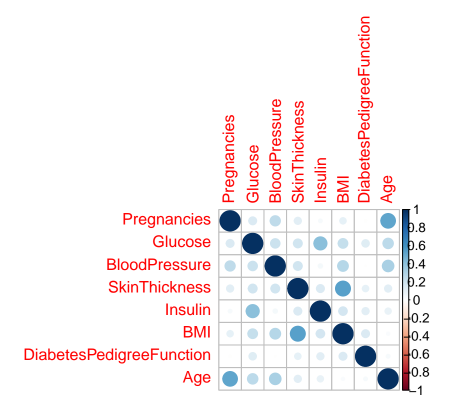
```
BloodPressure
                                                        SkinThickness
##
    Pregnancies
                       Glucose
          :-1.1485
                    Min. :-2.5513
                                     Min. :-4.00001
                                                              :-2.51479
   Min.
                                                        Min.
   1st Qu.:-0.8130
                                                        1st Qu.:-0.46729
                    1st Qu.:-0.7197
##
                                      1st Qu.:-0.69331
## Median :-0.1420
                    Median :-0.1530
                                     Median :-0.03197
                                                        Median :-0.01229
  Mean : 0.0000
##
                    Mean : 0.0000
                                      Mean : 0.00000
                                                        Mean : 0.00000
                    3rd Qu.: 0.6109
##
   3rd Qu.: 0.5291
                                      3rd Qu.: 0.62937
                                                        3rd Qu.: 0.32896
## Max. : 4.2197
                    Max. : 2.5410
                                           : 4.10141
                                                              : 7.95020
                                     Max.
                                                        Max.
      Insulin
                         BMI
##
                                      DiabetesPedigreeFunction
## Min.
         :-1.4664
                    Min. :-2.07343
                                     Min.
                                             :-1.1888
##
  1st Qu.:-0.2219
                    1st Qu.:-0.72074
                                      1st Qu.:-0.6885
## Median :-0.1814
                    Median :-0.02258
                                      Median :-0.2999
##
  Mean
         : 0.0000
                    Mean : 0.00000
                                      Mean : 0.0000
   3rd Qu.:-0.1554
                    3rd Qu.: 0.60286
                                      3rd Qu.: 0.4659
  Max. : 8.1651
                    Max. : 5.03911
##
                                      Max. : 5.8797
##
        Age
##
         :-1.0409
  Min.
  1st Qu.:-0.7858
## Median :-0.3606
## Mean : 0.0000
## 3rd Qu.: 0.6598
## Max. : 4.0611
```

3.6 Correlation Analysis

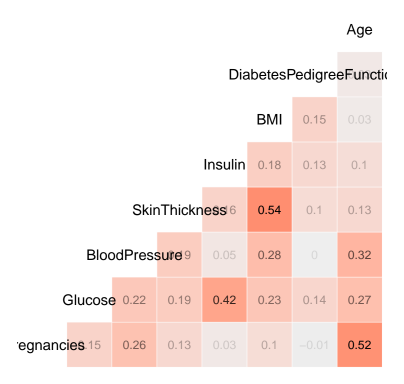
We now perform correlation analysis between independent variables. We observe the correlations through two plots.

3.6.1 corrplot

```
corr_mat <- cor(diab_x)
corrplot(corr_mat)</pre>
```



3.6.2 plot correlation matrix



The output of ggcorr shows that pregnancy and age have the highest correlation coefficient at 0.6, followed by SkinThickness and BMI at 0.5, and glucose and insulin at 0.4. The remaining variables do not have strong correlations. Since the highest correlation coefficient among these variables is only 0.6, and we only have 8 predictors, we will keep all predictors.

4. Splitting Datasets into Training Set and Test Set

Split the dataset into a training set (80%) and test set (20%)

```
set.seed(1, sample.kind = "Rounding") # if using R 3.6 or later
test_index <- createDataPartition(diab_y, times = 1, p = 0.2, list = FALSE)
test_x <- x_scaled[test_index,]
test_y <- diab_y[test_index]
train_x <- x_scaled[-test_index,]
train_y <- diab_y[-test_index]</pre>
```

Below we can see that by calculating the proportion of No and Yes in the training set and the proportion in the test set respectively, both datasets have very similar proportions.

```
# check the proportion in training set
prop.table(table(train_y))

## train_y
## No Yes
## 0.6514658 0.3485342
```

```
# check the proportion in test set
prop.table(table(test_y))

## test_y
## No Yes
## 0.6493506 0.3506494
```

5. Building the Models

We will use some different methods to build 8 models and see which model has the highest prediction accuracy.

Model 1: Logistic Regression Model

Model 2: rpart Model

Model 3: LDA Model

Model 4: QDA Model

Model 5: Loess Model

```
# set.seed(5)
set.seed(5, sample.kind = "Rounding") # simulate R 3.5

# train a loess model on the training set
train_loess <- train(train_x, train_y, method = "gamLoess")

# generate predictions on the test set
loess_preds <- predict(train_loess, test_x)

# calculate the accuracy, sensitivity, specificity, and prevalence of the model on test set
cm_loess <- confusionMatrix(data = loess_preds, reference = test_y)</pre>
```

Model 6: K-Nearest Neighbours Model

```
# train a KNN model on the training set, and find the best value in the model
set.seed(7,sample.kind = "Rounding")
train_knn <- train(train_x, train_y,</pre>
                   method = "knn",
                   tuneGrid = data.frame(k = seq(3, 21, 2))
# determine the value of k
train_knn$bestTune # the best value is 10 or 21
##
       k
## 10 21
# use the final value in the model to generate predictions on the test set
knn_preds <- predict(train_knn, test_x)</pre>
# calculate the accuracy, sensitivity, specificity, and prevalence of the model on test set
cm_knn <- confusionMatrix(data = knn_preds, reference = test_y)</pre>
result_knn <- c(cm_knn$byClass[c("Sensitivity", "Specificity",</pre>
                                  "Prevalence")], cm_knn$overall["Accuracy"])
result_knn
## Sensitivity Specificity Prevalence
                                           Accuracy
   0.9300000 0.5740741
                             0.6493506
                                        0.8051948
```

Model 7: Random Forest model

mtry ## 1 3

```
# find the most important variable in the random forest model
varImp(train_rf)
## rf variable importance
##
##
                            Importance
## Glucose
                                100.000
## BMI
                                 34.970
## Age
                                28.771
## Pregnancies
                                 22.484
## Insulin
                                11.927
## DiabetesPedigreeFunction
                                10.178
## SkinThickness
                                 2.177
## BloodPressure
                                  0.000
# use the best value of mtry in the model to generate predictions on the test set
rf_preds <- predict(train_rf, test_x)</pre>
# calculate the accuracy, sensitivity, specificity, and prevalence of the model on test set
cm_rf <- confusionMatrix(data = rf_preds, reference = test_y)</pre>
result_rf <- c(cm_rf$byClass[c("Sensitivity", "Specificity",</pre>
                                "Prevalence")], cm_rf$overall["Accuracy"])
result_rf
## Sensitivity Specificity Prevalence
                                           Accuracy
    0.8600000 0.6481481 0.6493506 0.7857143
```

Model 8: Ensemble Model

```
# Creating an ensemble model that combine all the models trained before
ensemble <- cbind(glm = glm_preds=="No", lda = lda_preds=="No",
                  qda = qda_preds=="No", loess = loess_preds=="No",
                  rf = rf_preds=="No", knn = knn_preds=="No",
                  rpart = rpart_preds=="No")
# generate predictions
ensemble_preds <- ifelse(rowMeans(ensemble) > 0.5, "No", "Yes")
# calculate the accuracy, sensitivity, specificity, and prevalence of the model on test set
cm_ensemble <- confusionMatrix(data = factor(ensemble_preds), reference = test_y)</pre>
result\_ensmble <- c(cm\_ensemble\$byClass[c("Sensitivity","Specificity", "Prevalence")],\\
                    cm_ensemble$overall["Accuracy"])
result_ensmble
## Sensitivity Specificity Prevalence
                                          Accuracy
    0.8800000 0.5555556 0.6493506
                                       0.7662338
```

Creating an ensemble list that shows the accuracy of these models.

```
# create a list of accuracy by model names
model names <- c("Logistic regression", "rpart", "LDA", "QDA", "Loess",</pre>
                 "K nearest neighbors", "Random forest", "Ensemble")
models_accuracy <- c(cm_glm$overall["Accuracy"], cm_rpart$overall["Accuracy"],</pre>
                      cm_lda$overall["Accuracy"], cm_qda$overall["Accuracy"],
                      cm_loess$overall["Accuracy"], cm_knn$overall["Accuracy"];
                      cm_rf$overall["Accuracy"], cm_ensemble$overall["Accuracy"])
data.frame(Model = model_names, Accuracy = models_accuracy)
##
                   Model Accuracy
## 1 Logistic regression 0.7597403
## 2
                   rpart 0.7792208
## 3
                     LDA 0.7467532
## 4
                     QDA 0.7337662
## 5
                   Loess 0.7727273
## 6 K nearest neighbors 0.8051948
## 7
           Random forest 0.7857143
## 8
                Ensemble 0.7662338
# select a best prediction model according to highest accuracy
best_model <- subset(models_accuracy, models_accuracy==max(models_accuracy))</pre>
best model
## Accuracy
## 0.8051948
```

Model Comparisons

Compare the accuracy of the models, and how they perform in terms of sensitivity, specificity.

```
##
            Sensitivity Specificity Prevalence Accuracy
## result_glm
                0.89 0.5185185 0.6493506 0.7597403
## result_rpart
                0.92
                     ## result_lda
                0.87
## result qda
                ## result loess
                0.89
                     0.5555556  0.6493506  0.7727273
## result_knn
                0.93
                     0.5740741 0.6493506 0.8051948
## result_rf
                0.86
                     0.6481481 0.6493506 0.7857143
                 ## result_ensmble
```

We can see that the best model is KNN model because it has highest accuracy, it also has better performance in Sensitivity, Specificity on the same Prevalence.

6. Conclusion

We used a few different methods to build some models and calculated and compared their accuracy. We recommend the KNN model as the best model because it has the highest prediction accuracy. In the future we can try to use other methods to train the models to get better accuracy; We may also divide the Age variable into different "Age Group" and find some different methods to improve the accuracy. I actually tried to add "Age Group" as one of the predictor, and trained the model used LDA and QDA methods and better accuracy by creating a "Age Group" variable.

References

- 1. HarvardX PH125.8x Data Science: Machine Learning: 7.1 Final Assessment: Breast Cancer Prediction Project
- 2. Kaggle: Pima Indians Diabetes Database, Dataset Notebooks
- 3. Kaggle: MIRI CHO, [Classification] Diabetes or Not (with basic 12ML)
- 4. Kaggle: LEARNING-MONK, PIMA Indian Diabetes Logistic Regression with R
- 5. ggcorr: https://briatte.github.io/ggcorr