Predicting the risk of diabetes

STATS/CSE 780 Course Project

Pao Zhu Vivian Hsu (Student Number: 400547994)

2023-12-12

Abstract

Introduction

Diabetes is a chronic disease that occurs when the body cannot effectively produce or use insulin to regulate sugar levels in the blood. According to the World Health Organization, 422 million people have diabetes worldwide and 1.5 million deaths that occur every year are directly linked to the disease (World Health Organization, n.d.). Due to its high prevalence and burden, finding ways to predict diabetes has become critical in improving population health.

With today's technological capabilities and abundance of data, machine learning has become the forefront of diabetes prediction. In literature, a large variety of techniques, such as naive Bayes, decision trees, and neural networks have been studied to find the most accurate method of prediction (Kumar & Velide, 2014; Rabina & Chopra, 2016).

One notable example is Islam et al.'s study on diabetes prediction (2020). In this study, the authors collected data from patients in Sylhet Diabetes Hospital in Sylhet, Bangladesh. Naive Bayes, logistic regression, and random forest techniques were applied to the data to predict the risk of diabetes. The study concluded that a random forest model had the greatest accuracy (Islam et al., 2020).

In our study, we also utilize machine learning techniques to predict the risk of diabetes. In particular, we focus on decision tree and neural network methods. The data we used was downloaded from an open source website called Kaggle (Larxel, 2023) and is the same data used in Islam et al.'s study (2020).

Methods

Data Exploration

The data set consists of 520 observations and 17 attributes. Before any analysis was done, a transformation was applied to the data to ensure that all categorical variables were expressed with binary indicators to ease the analysis. The response variable is a binary attribute called class that indicates whether the patient has a positive or negative risk for diabetes. The remaining attributes describe the patient and if they experience common symptoms related to the disease, such as weakness, itching, and obesity. A full list of the attributes and their meanings are outlined in Supp. Table 1.

There are no missing values in this data set since missing data was already addressed by Islam et al. after data collection (2020). To verify this, we performed a check for nulls and blanks as summarized in Supp. Table 2.

A correlation plot was created to check if there are any strong correlations between the attributes. We define a strong correlation as those with a correlation coefficient of 0.7 or larger. Based on Figure 1, all values are lower than 0.7 so there is no evidence of strong correlations.

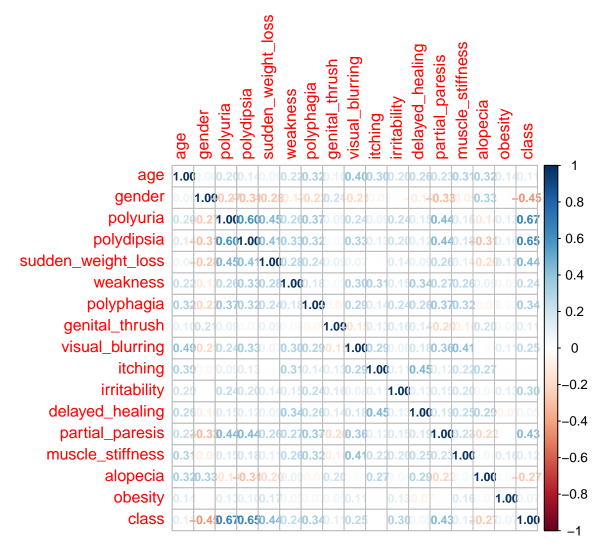


Figure 1: Correlation plot

Next, each of the individual attributes were explored. Figure 2 below shows a box plot of patient ages. The median age in the population is 47. There are a few outliers that exist outside of the interquartile range. These values will be capped at 79, the upper bound of the range.

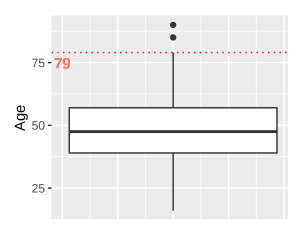


Figure 2: Boxplot of age

We also plotted the 16 categorical variables in bar charts as illustrated in Supp. Figure 3. There appears to be a somewhat even split between the predictor values for polyuria and itching, while the remaining predictors are not evenly split. This is especially important for the class variable because there are about 200 observations with a negative diabetes risk and about 300 observations with a positive risk. This suggests that if the model does not perform well, a resampling method such as cross-validation or bootstrapping may help improve the model.

Decision Tree

The first machine learning method will be a decision tree. This method was selected because Islam et al.'s study found that a decision tree model was most accurate (2020) and so this study will aim to reproduce such results. The index on the subtrees α will be tuned using cross-validation. Pruning will be applied to reduce the cost complexity of the tree, and random forest ensembling will be used to improve model performance.

Neural Network

The second method will be a neural network. This method was selected since Islam et al.'s study has not explored a model using neural networks (2020) and so there is a possibility that a neural network may be more accurate than a tree-based model. The number of hidden layers and the units per layer will be tuned through trial and error. Based on James et al.'s book on statistical

learning (2021), the units per layer will be set to some large value and overfitting will be controlled with ridge regularization. The strength of the regularization λ will be tuned at each layer.

Results

After the two models are built, they will each be assessed using misclassification rate, accuracy, specificity, and sensitivity. A comparison of these four measurements will reveal which model is a stronger fit to predict diabetes. By nature, decision trees are easier to interpret and reproducible compared to neural networks. Thus, these qualities will also be considered in its comparison.

Supplementary Materials

Tables and Figures

Table 1: Description of attributes

Attribute	Values
Age	In years
Gender	1 = Male, 0 = Female
Polyuria	1 = Yes, 0 = No
Polydipsia	1 = Yes, 0 = No
Sudden weight loss	1 = Yes, 0 = No
Weakness	1 = Yes, 0 = No
Polyphagia	1 = Yes, 0 = No
Genital thrush	1 = Yes, 0 = No
Visual blurring	1 = Yes, 0 = No
Itching	1 = Yes, 0 = No
Irritability	1 = Yes, 0 = No
Delayed healing	1 = Yes, 0 = No
Partial paresis	1 = Yes, 0 = No
Muscle stiffness	1 = Yes, 0 = No
Alopecia	1 = Yes, 0 = No
Obesity	1 = Yes, 0 = No
Class	1 = Positive risk, 0 = Negative risk

Table 2: No missing data

Nulls	Blank
0	(

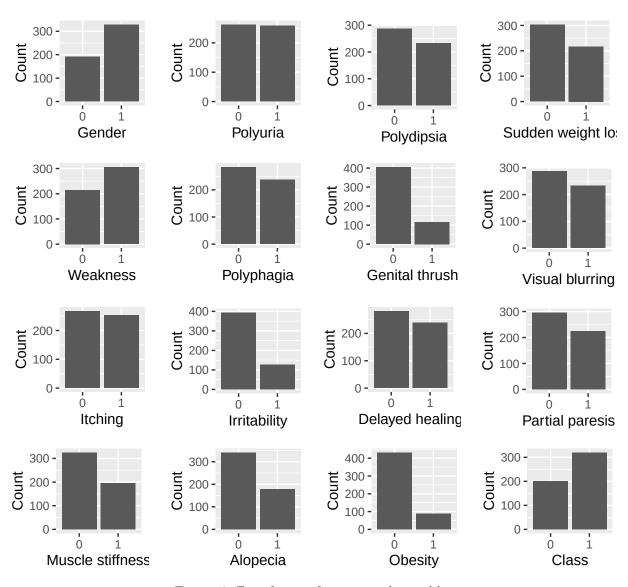


Figure 3: Bar charts of categorical variables

Code

```
library(knitr)
library(tidyverse)
library(corrplot)

# ---- DATA CLEANSING ----- #
diabetes_raw <- read.csv("diabetes_data.csv", sep=";")</pre>
```

```
diabetes_int <- diabetes_raw %>%
 mutate(gender = as.integer(ifelse(gender=="Male",1,
                            ifelse(gender=="Female",0,
                            NA))))
diabetes <- diabetes raw %>%
 mutate(gender = as.factor(ifelse(gender=="Male",1,
                            ifelse(gender=="Female",0,
         polyuria = as.factor(polyuria),
         polydipsia = as.factor(polydipsia),
         sudden_weight_loss = as.factor(sudden_weight_loss),
         weakness = as.factor(weakness),
         polyphagia = as.factor(polyphagia),
         genital_thrush = as.factor(genital_thrush),
         visual blurring = as.factor(visual blurring),
         itching = as.factor(itching),
         irritability = as.factor(irritability),
         delayed healing = as.factor(delayed healing),
         partial_paresis = as.factor(partial_paresis),
         muscle_stiffness = as.factor(muscle_stiffness),
         alopecia = as.factor(alopecia),
         obesity = as.factor(obesity),
         class = as.factor(class))
# ---- DATA EXPLORATION ---- #
# Data description
attribute <- c("Age", "Gender", "Polyuria", "Polydipsia",</pre>
               "Sudden weight loss", "Weakness", "Polyphagia",
               "Genital thrush", "Visual blurring", "Itching",
               "Irritability", "Delayed healing", "Partial paresis",
               "Muscle stiffness", "Alopecia", "Obesity", "Class")
```

```
values <- c("In years", "1 = Male, 0 = Female", "1 = Yes, 0 = No", "1 = Yes, 0 = No",
            "1 = Yes, 0 = No", "1 = Yes, 0 = No", "1 = Yes, 0 = No", "1 = Yes, 0 = No",
            "1 = Yes, 0 = No", "1 = Yes, 0 = No", "1 = Yes, 0 = No", "1 = Yes, 0 = No",
            "1 = Yes, 0 = No", "1 = Yes, 0 = No", "1 = Yes, 0 = No", "1 = Yes, 0 = No",
            "1 = Positive risk, 0 = Negative risk")
data_summary <- data.frame(Attribute=attribute, Values=values)</pre>
kable(data_summary)
# Check for missing data
nulls <- sapply(diabetes,</pre>
                function(col){ifelse(is.na(sum(col == "")), 0, sum(col == ""))})
blanks <- sapply(diabetes,</pre>
                 function(col){ifelse(is.na(sum(col == "")), 0, sum(col == ""))})
kable(data.frame(Nulls=sum(nulls), Blanks=sum(blanks)))
# Boxplot of continuous variable
bplot <- ggplot(diabetes, aes(y = age)) + geom_boxplot() + labs(x="",y="Age") +</pre>
  theme(axis.text.x=element blank(), axis.ticks.x=element blank())
bplot_a1 <- as.integer(unlist(ggplot_build(bplot)$data)["ymax"])</pre>
bplot + geom hline(yintercept = bplot a1, linetype="dotted", color="red") +
  geom_text(aes(-0.4,bplot_a1,label = bplot_a1, vjust = 1.5, color="red"),
            show.legend = FALSE)
# Cap outliers
diabetes_no_outliers <- diabetes %>% mutate(age = ifelse(age > bplot_a1, bplot_a1, age))
# Barplots for each categorical variable
ggplot(diabetes, aes(x = gender)) + geom bar() + labs(y = "Count", x = "Gender")
ggplot(diabetes, aes(x = polyuria)) + geom_bar() + labs(y = "Count", x = "Polyuria")
ggplot(diabetes, aes(x = polydipsia)) + geom_bar() +
  labs(y = "Count", x = "Polydipsia")
```

```
ggplot(diabetes, aes(x = sudden_weight_loss)) + geom_bar() +
  labs(y = "Count", x = "Sudden weight loss")
ggplot(diabetes, aes(x = weakness)) + geom_bar() + labs(y = "Count", x = "Weakness")
ggplot(diabetes, aes(x = polyphagia)) + geom_bar() +
  labs(y = "Count", x = "Polyphagia")
ggplot(diabetes, aes(x = genital_thrush)) + geom_bar() +
  labs(y = "Count", x = "Genital thrush")
ggplot(diabetes, aes(x = visual_blurring)) + geom_bar() +
  labs(y = "Count", x = "Visual blurring")
ggplot(diabetes, aes(x = itching)) + geom_bar() + labs(y = "Count", x = "Itching")
ggplot(diabetes, aes(x = irritability)) + geom_bar() +
  labs(y = "Count", x = "Irritability")
ggplot(diabetes, aes(x = delayed_healing)) + geom_bar() +
  labs(y = "Count", x = "Delayed healing")
ggplot(diabetes, aes(x = partial_paresis)) + geom_bar() +
  labs(y = "Count", x = "Partial paresis")
ggplot(diabetes, aes(x = muscle_stiffness)) + geom_bar() +
  labs(y = "Count", x = "Muscle stiffness")
ggplot(diabetes, aes(x = alopecia)) + geom_bar() + labs(y = "Count", x = "Alopecia")
ggplot(diabetes, aes(x = obesity)) + geom_bar() + labs(y = "Count", x = "Obesity")
ggplot(diabetes, aes(x = class)) + geom bar() + labs(y = "Count", x = "Class")
# Correlation plot
corr_matrix <- cor(diabetes_int)</pre>
corrplot(round(corr_matrix,2), method = "number", number.cex=0.75)
```

References

- Islam, M. M. F., Ferdousi, R., Rahman, S., & Bushra, H. Y. (2020). Likelihood prediction of diabetes at early stage using data mining techniques. In M. Gupta, D. Konar, S. Bhattacharyya, & S. Biswas (Eds.), Computer vision and machine intelligence in medical image analysis (pp. 113–125). Springer Singapore. https://doi.org/10.1007/978-981-13-8798-2 12
- James, G., Witten, D., Hastie, T., & Tibshirani, R. (2021). Deep learning. In An introduction to statistical learning with applications in r 2nd edition (pp. 403–458). Springer.
- Kumar, V., & Velide, L. (2014). A data mining approach for prediction and treatment of diabetes disease.
- Larxel. (2023). Early classification of diabetes. https://www.kaggle.com/datasets/andrewmvd/heart-failure-clinical-data/data
- R Core Team. (2023). R: A language and environment for statistical computing. R Foundation for Statistical Computing. https://www.R-project.org/
- Rabina, & Chopra, Er. A. (2016). Diabetes prediction by supervised and unsupervised learning with feature selection.
- World Health Organization. (n.d.). *Diabetes*. https://www.who.int/health-topics/diabetes#tab=tab_1