Diabetes Risk prediction via SEMMA w/ Regularized Logistic Regression

Kevin Acosta* M.S. in Data & Information Sciences (UTEP)

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1 Data Import

data <- read.csv("/Users/kevinacosta/Desktop/Stats ML 2/Comp Proj 01/diabetes_data_upload.csv"
dim(data)</pre>

[1] 520 17

summary(data)

##	Age	Gender	Polyuria	Polydipsia
##	Min. :16.00	Length:520	Length:520	Length:520
##	1st Qu.:39.00	Class :character	Class :character	Class :character
##	Median :47.50	Mode :character	Mode :character	Mode :character
##	Mean :48.03			
##	3rd Qu.:57.00			
##	Max. :90.00			
##	sudden.weight.loss weakness		Polyphagia	Genital.thrush
##	Length:520	Length: 520	Length: 520	Length:520
##	Max. :90.00 sudden.weight.l		V. 0	

^{*}kmacosta 2@miners.utep.edu

```
Class : character
                        Class : character
                                            Class :character
                                                                Class : character
    Mode :character
                        Mode :character
                                                  :character
##
                                            Mode
                                                                Mode :character
##
##
##
    visual.blurring
                                            Irritability
                                                                delayed.healing
##
                          Itching
    Length:520
                        Length:520
                                            Length: 520
                                                                Length:520
    Class : character
                        Class : character
                                            Class :character
                                                                Class : character
    Mode :character
                        Mode :character
                                            Mode :character
                                                                Mode :character
##
##
##
##
   partial.paresis
                        muscle.stiffness
                                              Alopecia
##
                                                                   Obesity
    Length: 520
                        Length: 520
                                            Length: 520
                                                                Length: 520
##
    Class : character
                        Class : character
                                            Class : character
                                                                Class : character
    Mode :character
                        Mode :character
                                            Mode :character
                                                                Mode : character
##
##
##
##
       class
    Length:520
##
    Class : character
    Mode :character
##
##
##
head(data) [c(1,2,3),]
##
     Age Gender Polyuria Polydipsia sudden.weight.loss weakness Polyphagia
      40
                                                               Yes
## 1
           Male
                       No
                                  Yes
                                                       No
## 2
     58
           Male
                       No
                                   No
                                                       No
                                                               Yes
                                                                            No
## 3
           Male
                      Yes
                                   No
                                                       No
                                                               Yes
                                                                           Yes
##
     Genital.thrush visual.blurring Itching Irritability delayed.healing
## 1
                 No
                                   No
                                          Yes
                                                                         Yes
                                                         No
## 2
                 No
                                  Yes
                                           No
                                                         No
                                                                          No
## 3
                                   No
                                          Yes
                 No
                                                         No
                                                                         Yes
     partial.paresis muscle.stiffness Alopecia Obesity
##
## 1
                                             Yes
                  No
                                    Yes
                                                      Yes Positive
## 2
                  Yes
                                             Yes
                                                       No Positive
## 3
                  No
                                    Yes
                                             Yes
                                                       No Positive
```

2 Exploratory Data Analysis

```
table(data$class) # simple frequency table
##
## Negative Positive
```

```
##
        200
                 320
for (i in 1:(ncol(data)-1)) {
  cat("Column:", colnames(data)[i],"--> ")
 x \leftarrow data[,i]
  cat("Number of unique values:", length(unique(x, incomparables=TRUE)),"\n")
  # checking for missing values per column
  col_count <- sum(is.na(x))</pre>
  if(col_count>0) {
    cat("Missing Values:", col_count, "\n")
  }
  else {
    cat("\t\tNONE Missing for this column\n")
}
## Column: Age --> Number of unique values: 51
        NONE Missing for this column
## Column: Gender --> Number of unique values: 2
        NONE Missing for this column
## Column: Polyuria --> Number of unique values: 2
        NONE Missing for this column
## Column: Polydipsia --> Number of unique values: 2
        NONE Missing for this column
##
## Column: sudden.weight.loss --> Number of unique values: 2
        NONE Missing for this column
## Column: weakness --> Number of unique values: 2
##
        NONE Missing for this column
## Column: Polyphagia --> Number of unique values: 2
##
        NONE Missing for this column
## Column: Genital.thrush --> Number of unique values: 2
        NONE Missing for this column
## Column: visual.blurring --> Number of unique values: 2
        NONE Missing for this column
## Column: Itching --> Number of unique values: 2
        NONE Missing for this column
## Column: Irritability --> Number of unique values: 2
##
        NONE Missing for this column
## Column: delayed.healing --> Number of unique values: 2
        NONE Missing for this column
## Column: partial.paresis --> Number of unique values: 2
##
        NONE Missing for this column
## Column: muscle.stiffness --> Number of unique values: 2
        NONE Missing for this column
## Column: Alopecia --> Number of unique values: 2
        NONE Missing for this column
```

```
## Column: Obesity --> Number of unique values: 2
## NONE Missing for this column
```

After seeing the first frequency table, we can confirm this does resemble an unbalanced classification problem favoring 'positive' labels with 200 negative cases for Diabetes and 320 positive cases. Using a for loop I checked for every column/feature excluding our response variable and confirmed that there are no missing values found.

3 Variable Screening

```
#library(tidyverse)
response <- data$class
p_values <- numeric(ncol(data) - 1)</pre>
data2 <- data
data2[, 1:(ncol(data2) - 1)] <- lapply(data2[, 1:(ncol(data2) - 1)], as.factor)
for (i in 1:(ncol(data2) - 1)) {
  attrib <- data2[, i]
  if (is.factor(attrib)){ # categorical attribute --> chi-squared test
    cat_attr <- chisq.test(table(attrib, response))</pre>
    p_values[i] <- cat_attr$p.value</pre>
  }
  else{ # continuous attribute --> t-test
    cont_attr <- t.test(attrib ~ response)</pre>
    p_values[i] <- cont_attr$p.value</pre>
  }
  cat("Attribute: ", colnames(data2)[i], "\n\t\tP-value: ", p_values[i], "\n")
}
## Warning in chisq.test(table(attrib, response)): Chi-squared approximation may
## be incorrect
## Attribute: Age
##
        P-value: 2.654685e-11
## Attribute: Gender
##
        P-value: 3.289704e-24
## Attribute: Polyuria
        P-value: 1.740912e-51
##
## Attribute: Polydipsia
        P-value: 6.18701e-49
##
## Attribute: sudden.weight.loss
        P-value:
                  5.969166e-23
##
## Attribute: weakness
##
        P-value: 4.869843e-08
## Attribute: Polyphagia
        P-value: 1.165158e-14
##
```

```
## Attribute: Genital.thrush
        P-value: 0.0160979
## Attribute: visual.blurring
        P-value: 1.701504e-08
##
## Attribute: Itching
        P-value:
                  0.8297484
##
## Attribute: Irritability
        P-value: 1.771483e-11
## Attribute: delayed.healing
        P-value: 0.3266599
## Attribute: partial.paresis
        P-value: 1.565289e-22
##
## Attribute: muscle.stiffness
        P-value: 0.006939096
##
## Attribute: Alopecia
        P-value: 1.909279e-09
## Attribute: Obesity
##
        P-value: 0.127108
# identifying the statistically significant attributes
significant_attrs <- which(p_values < 0.25 & !is.na(p_values))</pre>
significant_names <- colnames(data2)[significant_attrs]</pre>
print("Significant attributes based on level alpha 0.25:\n")
## [1] "Significant attributes based on level alpha 0.25:\n"
print(significant_names)
    [1] "Age"
                             "Gender"
##
                                                   "Polyuria"
##
   [4] "Polydipsia"
                             "sudden.weight.loss" "weakness"
## [7] "Polyphagia"
                             "Genital.thrush"
                                                   "visual.blurring"
## [10] "Irritability"
                             "partial.paresis"
                                                   "muscle.stiffness"
```

4 Data Partition

[13] "Alopecia"

Partition the data into two parts, the training data D1 and the test data D2, with a ratio of 2:1.

"Obesity"

```
data$Itching <- NULL
data$delayed.healing <- NULL

n_samples <- nrow(data)
# 2/3 for train set below
n_train <- round(n_samples * 0.67) - 1
train <- sample(n_samples, n_train)
train_data <- data[train,]
test_data <- data[-train,]</pre>
```

```
dim(train_data)
## [1] 347 15
dim(test_data)
## [1] 173 15
```

5 Logistic Regression Modeling

```
library(glmnet)

## Loading required package: Matrix

## Loaded glmnet 4.1-8

y <- as.factor(train_data$class)

X <- model.matrix(class ~ ., data=train_data)[,-1] #

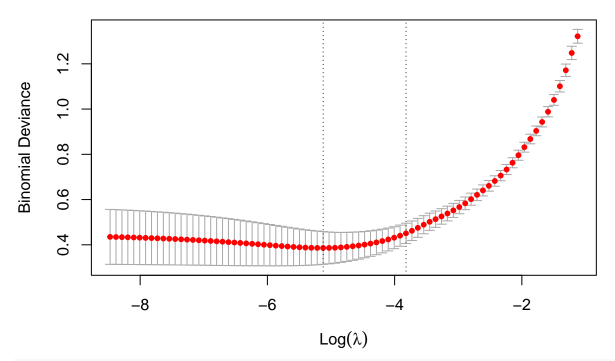
# fitting a lasso logistic regression with cv to help select the best lambda value.

cv_lasso_fit <- cv.glmnet(x=X,y=y, nfolds=10, family="binomial", alpha=1)

best_lambda <- cv_lasso_fit$lambda.min

plot(cv_lasso_fit)</pre>
```

14 14 14 14 14 13 11 11 9 9 7 6 4 3 2 2



here we're simply fitting again but to include the best lambda & afterwards
inspect the coefficients
best_lasso <- glmnet(x=X, y=y, family="binomial", alpha=1, lambda = best_lambda)
cat("Best lambda:", best_lambda)</pre>

Best lambda: 0.005954938

```
print("Final lasso model coefficients:")
## [1] "Final lasso model coefficients:"
print(coef(best_lasso))
## 15 x 1 sparse Matrix of class "dgCMatrix"
##
                                   s0
                           2.57905167
## (Intercept)
## Age
                          -0.05492271
## GenderMale
                         -3.42280679
## PolyuriaYes
                           2.99271946
## PolydipsiaYes
                          3.47185990
## sudden.weight.lossYes 0.66408553
## weaknessYes
## PolyphagiaYes
## Genital.thrushYes
                           1.07112130
## visual.blurringYes
## IrritabilityYes
                           1.83898204
## partial.paresisYes
                          1.60617446
## muscle.stiffnessYes
                         -0.24020360
## AlopeciaYes
                         -0.01279120
## ObesityYes
                         -0.32272431
```

The model coefficient results presented above can vary from run to run, but as long as they have a value not zero, we would list them as significant to the model.

In general and on average, the most relevant coefficients are: 'PolydipsiaYes'- (a lot of thirst is indicated with the highest positive coefficient), 'PolyuriaYes'- (excessive urination is also indicated by a large coefficient, so its another strong predictor of diabetes) 'GenderMale'- (so just being male significantly decreases the odds of getting a diabetes diagnosis because the coefficient is negative!).

6 Model Assessment/Deployment

```
library(pROC)

## Type 'citation("pROC")' for a citation.

##

## Attaching package: 'pROC'

## The following objects are masked from 'package:stats':

##

## cov, smooth, var

y_test <- as.factor(test_data$class)

x_test <- model.matrix(class ~ ., data=test_data)[,-1]

# predict probability for a positive classification/diagnosis
y_pred <- predict(best_lasso, newx=x_test, type="response")</pre>
```

```
# generate ROC curve and get AUC
roc_curve <- roc(y_test, y_pred)

## Setting levels: control = Negative, case = Positive

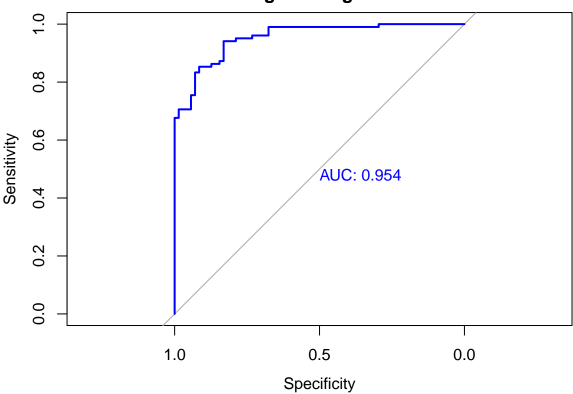
## Warning in roc.default(y_test, y_pred): Deprecated use a matrix as predictor.

## Unexpected results may be produced, please pass a numeric vector.

## Setting direction: controls < cases
auc_value <- auc(roc_curve)

plot(roc_curve, col = "blue", main = "ROC Curve: Logistic Regression w/LASSO", print.auc = TRUC</pre>
```

ROC Curve: Logistic Regression w/LASSO



```
print(paste("AUC/C-Statistic:", round(auc_value, 4)))
```

[1] "AUC/C-Statistic: 0.9542"

Overall the Logistic Regression Classifier's performance is very good at predicting what is considered the true positives in classifying patients with diabetes when they do in fact have it.