# $report\_lab1$

# Yanjie Lyu, Yi Yang, Qingxuan Cui

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# Contents

T	Sta	Statement of Contribution							
2	Assignment 1 2.1 Comment on the quality of predictions for different digits and on the overall prediction quality 2.2 cases of digit "8"								
3	$\mathbf{Ass}$	Assignment 2							
	3.1 3.2 3.3 3.4 3.5	Compute a linear regression from the training data	3 4 4 4						
4	Assignment 3								
	4.1	Step1	4						
	4.2	Step2	4						
	4.3	Step3	5						
	4.4	Step4	5						
	4.5	Step5	5						
5	$\mathbf{Ass}$	ignment 4. Theory	6						
	5.1	Importance to consider various probability thresholds in classification problems	6						
	5.2	Methods of collecting correct values of the labels for the supervised learning problems	6						
	5.3	Express the cost fucntion of the linear regression in the matrix form	6						
6	Apı	pendix	7						
	6.1	Code for assignment 1	7						
	6.2	Code for assignment 2	9						
	6.3	Code for assignment 3	12						
	6.4	Figures	14						

### 1 Statement of Contribution

The contributions are distributed as follows:

Yanjie Lyu: Worked on Assignment 1 and Question 1 from Assignment 4.

Yi Yang: Worked on Assignment 2 and Question 3 from Assignment 4.

Qingxuan Cui: Worked on Assignment 3 and Question 2 from Assignment 4.

After completing their respective assignments (including code writing and analysis), all results were shared and thoroughly discussed among the three members. the group report was created based on this discussion.

# 2 Assignment 1

# 2.1 Comment on the quality of predictions for different digits and on the overall prediction quality

According to the test confusion matrix and misclassification rate (see Figure 4 and 5), the model performs best on number "0", but there is a high rate on number "5", "8", and "9".

The overall misclassification rates of training data and test data are 0.04238619 and 0.05857741, which is acceptable.

Overall Performance: The model performs well with low misclassification rates and generalizes effectively across training and test sets.

#### 2.2 cases of digit "8"

Accroding to Figure 6, index 209 is the easiest one to be recognized, and in Figure 7, index 1663 can barely make out the shape of eight.

#### 2.3 Fit a K-nearest neighbor classifiers with misclassification errors

## the optimal value of K is: 3

1.As the value of K increases, the model becomes less sensitive, i.e. the model complexity decreases. Smaller K values result in a more complex model (sensitive to noise), while larger K values make the model smoother (potentially resulting in underfitting).

2. The training error rate curve increases with the increase of K. With the increase of K, the validation error rate curve first decreases, reaches a lowest point (the best value k), and then may slightly increase or become stable.

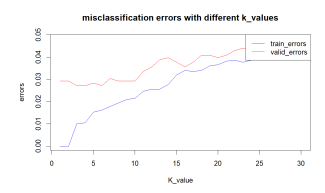


Figure 1: A.1.1: Misclassification Error

#### 2.3.1 test error compared with training and valid error

```
## the test error is: 0.03242678
## the training error is: 0.009942439
## the validation error is: 0.02722513
```

The classification error rate of the model on training, verification and test sets is low, and the difference is not large, indicating that the model has consistent performance on different data sets.

### 2.4 Fit K-nearest neighbor classifiers with cross-entropy

## the optimal value of K is: 8

1. For multiple classification problems, cross entropy can reflect the degree of confidence of the model in predicting the correct category.

2. The misclassification rate only considers correct or incorrect classifications, but the cross-entropy is more sensitive to the probabilistic output of the model and therefore reflects the performance of the model in more detail. If a model misclassifies, but the probability of prediction is very close to the correct answer, the increase in cross entropy loss will be smaller than the misclassification rate, reflecting a more nuanced error situation.

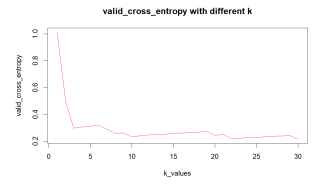


Figure 2: A.1.2: Validation Cross Entropy

# 3 Assignment 2

### 3.1 Compute a linear regression from the training data

Using 'lm()'to fit a linear regression model without intercept, we got the summary of the model as follows:

Figure 3: A.2.1: Summary

#### 3.2 Estimate traing and test MSE

The MSE for training and test data show as follows:

```
## MSE for the training data is : 0.8785431
```

## MSE for the test data is : 0.9354477

### 3.3 Commenting on Significant Variable Contributions to the Model

The coefficientJitter.Abs.,Shimmer.APQ5,Shimmer.APQ11,NHR,HNR,DFA,PPE is highly significant, as their P-value are less than 0.001,which suggests that they have a strong and statistically reliable impact on the response variable. The coefficient Shimmer is statistically significant,with a P-value < 0.01,indicating that Shimmer has a meaningful and reliable impact on the response variable.

# 3.4 Using RidgeOptto compute optimal parameters when lambda=1, 100 and 1000

The training and test MSE for different lambda and the degrees of freedom shows as follws:

```
## Training data MSE when lambda = 1 : 0.8786271
```

## Test data MSE when lambda =1: 0.9349969

## Degree of freedom when lambda = 1: 13.86074

## Training data MSE when lambda = 100 : 0.8844104

## Test data MSE when lambda = 100 : 0.9323316

## Degree of freedom when lambda = 100 : 9.924887

## Training data MSE when lambda = 1000 : 0.9211216

## Test data MSE when lambda = 1000 : 0.9539482

## Degree of freedom when lambda = 1000 : 5.643925

#### 3.5 Comenting on which is the most appropriate penalty parameter

Among the selected penalty parameters, lambda = 100 is the most appropriate .It provides the lowest test data MSE(0.9323316). Besides, the degree of freedom for lambda = 100 is 9.924887, which shows a balanced model complexity that avoids both overfitting and underfitting.

# 4 Assignment 3

## 4.1 Step1

# 4.1.1 Make a scatterplot showing a Plasma glucose concentration on Age where observations are colored by Diabetes levels

The plot is shown in Figure 8.

#### 4.1.2 Debate whether Diabetes is easy to classify by a standard logistic regression model

From the scatter plot between the given 2 features, that there is no any apparent boundary to classify the green points and red points representing whether being healthy or having diabetes. Based on that, I think it is not easy.

#### 4.2 Step2

#### 4.2.1 Report the probabilistic equation of the estimated model

The equation is:

$$P(y|X,\beta) = \sigma(X\beta) = \frac{1}{1 + exp(-X\beta)}$$

where:  $\sigma()$  represents the sigmod function

X is input data

 $\beta$  is coefficients

y is predicted output

#### 4.2.2 Comment on the quality of the classification by using these results

Based on accuracy of the prediction, besides the comparison between plots, it is obvious that the quality is not good enough to classify them well.

- ## [1] "The accuracy can be considered as quality of classification: 0.734028683181226"
- ## [1] "Misclassification error is: 0.265971316818774"

### 4.2.3 Make the scatter plot

The plot is shown in Figure 9.

### 4.3 Step3

4.3.1 Use the model estimated in step 2 to report the equation of the decision boundary between the two classes

```
## Decision Boundary: -5.897858 + 0.0355825 * x1 + 0.02450157 * x2 = y bar
```

#### 4.3.2 Add a curve showing this boundary to the scatter plot in step 2

The plot with the curve is shown in Figure 10.

#### 4.3.3 Comment whether the decision boundary seems to catch the data distribution well

The decision boundary is represented by a linear function, while the data distribution is more complicated than the data which can be caught by linear function.

#### 4.4 Step4

#### 4.4.1 Make same kind of plots as in step 2 but use thresholds r=0.2 and r=0.8

The plot with threhold = 0.2 and threhold = 0.8 are shown in Figure 11 and Figure 12, repectively.

# 4.4.2 By using these plots, comment on what happens with the prediction when r value changes

With the increasing value of r, less sample would be predicted as positive.

### 4.5 Step5

4.5.1 Create a scatterplot of the same kind as in step 2 for this model and compute the training misclassification rate

The plot is shown in Figure 13

## The misclassification error of the present model is: 0.2464146

# 4.5.2 What can you say about the quality of this model compared to the previous logistic regression model

After adding the non-linear elements into features, the performance of the model improved a little, but either cannot classify well.

- ## The misclassification error of the present model is: 0.2464146
- ## The misclassification error of the previous model is: 0.2659713

# 4.5.3 How have the basis expansion trick affected the shape of the decision boundary and the prediction accuracy

The decision boundary generated from original features is a linear function, while the present decision boundary in the shape of exponential function plot. In the meanwhile, it does improve the accuracy but slightly.

## The accuracy of the present model: 0.7535854

## The accuracy of the previous model: 0.7340287

## 5 Assignment 4. Theory

## 5.1 Importance to consider various probability thresholds in classification problems.

In classification problems, r=0.5 may not always the most important aspect of classifier. Many classification problems are asymmetric or imbalanced. For example, it can be important not to falsely predict the negative class than to falsely to predict the positive class in a medical diagnosis application. Besides, the medical problem could be imbalanced if the disorder is very rare. Adjusting the threshold r is essential to achieve performance based on different context(page 47-48).

# 5.2 Methods of collecting correct values of the labels for the supervised learning problems.

The vaules of label can be:

manually assigned by domain experts or possibly some auxiliary examination

calculated via specialised experiments and theories

obtained from the historical records(page 4-6)

#### 5.3 Express the cost function of the linear regression in the matrix form.

The cost function for the linear regression in the matrix form is defined as:

$$J(\theta) = \frac{1}{n} \sum_{i=1}^n \left( \hat{y}(x_i; \theta) - y_i \right)^2 = \frac{1}{n} \|\hat{y} - y\|_2^2 = \frac{1}{n} \|X\theta - y\|_2^2 = \frac{1}{n} \|\epsilon\|_2^2$$

where:  $\hat{y} = X\theta$  is the vector of the predicted values

y is the vector of true values

 $\epsilon = y - \hat{y}$  is the residuals

 $\|\cdot\|$  denotes the Euclidean norm

This cost function is also referred to as the least squares cost. (page 40,formula(3.11))

# 6 Appendix

## 6.1 Code for assignment 1

```
library(kknn)
data <- read.csv("LAB1data/optdigits.csv", header = TRUE)</pre>
data$X0.26 <- as.factor(data$X0.26)</pre>
n <- nrow(data)</pre>
set.seed(12345)
id \leftarrow sample(1:n,floor(n*0.5))
train <- data[id,]</pre>
valid_test <- data[-id,]</pre>
id_2 <- sample(1:nrow(valid_test),floor(nrow(valid_test)*0.5))</pre>
valid <- valid_test[id_2,]</pre>
test <- valid_test[-id_2,]</pre>
train_model <- kknn(X0.26~.,train,train,k=30,kernel = "rectangular")</pre>
train_predictions <- fitted(train_model)</pre>
test_model <- kknn(X0.26~.,train,test,k=30,kernel = "rectangular")</pre>
test_predictions <- fitted(test_model)</pre>
train_confusion_matrix <- table(Predicted = train_predictions, Actual = train$X0.26)</pre>
test_confusion_matrix <- table(Predicted = test_predictions, Actual = test$X0.26)
misclassification_train <- 1 - sum(diag(train_confusion_matrix)) / sum(train_confusion_matrix)
misclassification_test <- 1 - sum(diag(test_confusion_matrix)) / sum(test_confusion_matrix)
train_prob <- train_model$prob</pre>
index_eight <- which(train$X0.26 == "8")</pre>
prob_eight <- train_prob[index_eight, "8"]</pre>
ordered_indices <- order(prob_eight, decreasing = TRUE)</pre>
easiest_indices <- index_eight[ordered_indices[1:2]]</pre>
n <- length(ordered indices)</pre>
hardest_indices <- index_eight[ordered_indices[(n-2):n]]
feature_columns <- setdiff(names(train), "X0.26")</pre>
easiest_features_1 <- as.numeric(train[easiest_indices[1], feature_columns])</pre>
easiest_features_2 <- as.numeric(train[easiest_indices[2], feature_columns])</pre>
hardest_features_1 <- as.numeric(train[hardest_indices[1], feature_columns])
hardest_features_2 <- as.numeric(train[hardest_indices[2], feature_columns])
hardest_features_3 <- as.numeric(train[hardest_indices[3], feature_columns])
digit_matrix_easy_1 <- matrix(easiest_features_1, nrow = 8, ncol = 8, byrow = TRUE)</pre>
digit_matrix_easy_2 <- matrix(easiest_features_2, nrow = 8, ncol = 8, byrow = TRUE)</pre>
digit_matrix_hard_1 <- matrix(hardest_features_1, nrow = 8, ncol = 8, byrow = TRUE)</pre>
digit_matrix_hard_2 <- matrix(hardest_features_2, nrow = 8, ncol = 8, byrow = TRUE)</pre>
digit_matrix_hard_3 <- matrix(hardest_features_3, nrow = 8, ncol = 8, byrow = TRUE)</pre>
```

```
heatmap(digit_matrix_easy_1, Colv = NA, Rowv = NA,
        main = paste("easiest_indices:",easiest_indices[1] ))
heatmap(digit_matrix_easy_2, Colv = NA, Rowv = NA,
        main = paste("easiest indices:",easiest indices[2] ))
heatmap(digit_matrix_hard_1, Colv = NA, Rowv = NA,
        main = paste("hardest_indices:",hardest_indices[1] ))
heatmap(digit_matrix_hard_2, Colv = NA, Rowv = NA,
        main = paste("hardest indices:",hardest indices[2] ))
heatmap(digit_matrix_hard_3, Colv = NA, Rowv = NA,
        main = paste("hardest_indices:",hardest_indices[3] ))
k_values <- c(1:30)
train_errors <- numeric(length(k_values))</pre>
valid_errors <- numeric(length(k_values))</pre>
for (i in k_values) {
  model <- kknn(X0.26 ~ ., train = train, test = train, k = i, kernel = "rectangular")</pre>
  pred_train <- fitted(model)</pre>
  train_cm <- table(Predicted = pred_train, Actual = train$X0.26)</pre>
  train_errors[i] <- 1 - sum(diag(train_cm)) / sum(train_cm)</pre>
  model_valid <- kknn(X0.26 ~ ., train = train, test = valid, k = i, kernel = "rectangular")</pre>
  pred valid <- fitted(model valid)</pre>
 valid_cm <- table(Predicted = pred_valid, Actual = valid$X0.26)</pre>
  valid_errors[i] <- 1 - sum(diag(valid_cm)) / sum(valid_cm)</pre>
plot(k_values, train_errors, type = "1", col = "blue", ylim = c(0, max(c(train_errors, valid_errors))),
     xlab = "K_value", ylab = "errors", main = "errors with different k_values")
lines(k_values, valid_errors, type = "l", col = "red")
legend("topright", legend = c("train_errors", "valid_errors"), col = c("blue", "red"), lty = 1)
optimal_K <- which.min(valid_errors)</pre>
final_model <- kknn(X0.26 ~ ., train = train, test = test, k = optimal_K, kernel = "rectangular")</pre>
pred_test <- fitted(final_model)</pre>
test_cm <- table(Predicted = pred_test, Actual = test$X0.26)</pre>
test error <- 1 - sum(diag(test cm)) / sum(test cm)
valid_cross_entropy <- numeric(length(k_values))</pre>
class_levels <- levels(train$X0.26)</pre>
for (i in k_values) {
  model <- kknn(X0.26 ~ ., train = train, test = valid, k = i, kernel = "rectangular")</pre>
  prob_matrix <- model$prob</pre>
  epsilon <- 1e-15
  prob_matrix <- pmax(prob_matrix, epsilon)</pre>
  prob_matrix <- pmin(prob_matrix, 1 - epsilon)</pre>
  actual_classes <- as.numeric(valid$X0.26)</pre>
  N <- nrow(valid)</pre>
  cross_entropy <- 0</pre>
```

#### 6.2 Code for assignment 2

```
parkinsons <- read.csv("LAB1data/parkinsons.csv",header=TRUE)</pre>
parkinsons <- parkinsons[,!names(parkinsons)%in%c("subject.","age","sex","test_time","total_UPDRS")]
library(caret)
#data scaling
set.seed(12345)
n <- nrow(parkinsons)</pre>
id <- sample(1:n,floor(n*0.6))</pre>
train <- parkinsons[id,]</pre>
test <- parkinsons[-id,]</pre>
library(caret)
scaler <- preProcess(train)</pre>
trainS <- predict(scaler,train)</pre>
testS <- predict(scaler,test)</pre>
#fit the model
fit <- lm(motor UPDRS~.-1,data=trainS)</pre>
summary(fit)
#using model to predict training and test data
predictions_train <- predict(fit,newdata=trainS)</pre>
predictions_test <- predict(fit, newdata = testS)</pre>
#compute MSE for trainning and test data
MSE_trainning <- mean((trainS$motor_UPDRS-predictions_train)^2)</pre>
MSE_test <- mean((testS$motor_UPDRS-predictions_test)^2)</pre>
cat("MSE for the training data is :",MSE_trainning,"\n")
cat("MSE for the test data is :",MSE_test,"\n")
#The coefficient`Jitter.Abs.`,`Shimmer.APQ5`,`Shimmer.APQ11`,`NHR`,`HNR`,`DFA`,`PPE`
#is highly significant, as their P-value < 0.001, which suggests that they have a strong
#and statistically reliable impact on the response variable.
```

```
#The coefficient for `Shimmer.dB.` is statistically significant, as its P-value < 0.01.
#This suggests that `Shimmer.dB.` has a meaningful and reliable impact on the response variable.
#log_likelihood
Loglikelihood <- function(theta,sigma,X,Y){</pre>
  n <- length(Y)
  predicted Y <- X%*%theta
  squared_error <- (Y-predicted_Y)^2</pre>
  log_likelihood <- -0.5*n*log(2*pi*sigma^2)-0.5/sigma^2*sum(squared_error)
  return(log_likelihood)
}
#compute the loglikelihood for the fitting model
 #extract theta
theta <- coef(fit)
length(theta)
#extract sigma
residual_train <- trainS$motor_UPDRS-predictions_train</pre>
sigma <- sqrt(mean(residual_train^2))</pre>
#extract X as feature matrix for the training data
X_train <- as.matrix(trainS[,-1])</pre>
#extract Y
Y_train <- as.numeric(trainS$motor_UPDRS)</pre>
 #compute loglikelihood for the training data
Loglikelihood(theta,sigma,X_train,Y_train)
predictions_train
#Ridge
Ridge <- function(theta,sigma,lambda,X,Y){</pre>
  penalty <- lambda*sum(theta^2) #penalty</pre>
  loglikelihood <- Loglikelihood(theta,sigma,X,Y)</pre>
  ridge_loglikelihood <- -loglikelihood+penalty</pre>
  return(ridge_loglikelihood)
}
#RidgeOpt
RidgeOpt <- function(lambda,X,Y){</pre>
  #object function
  objfun <- function(params){</pre>
    #extract theta and sigma
    theta <- params[1:ncol(X)]</pre>
    sigma <- params[ncol(X)+1]</pre>
    #ensure sigma is non-negative
    if(sigma<=0)</pre>
      return(Inf)
    #compute ridge-loglikelihood
```

```
RidgeLikelihood <- Ridge(theta, sigma, lambda, X, Y)</pre>
    return(RidgeLikelihood)
  #initialize theta and sigma
  initialized_theta <- rep(0,ncol(X)) #theta to be 0</pre>
  initialized_sigma <- sd(Y)</pre>
  #combine theta with sigma
  initialized_params <- c(initialized_theta,initialized_sigma)</pre>
  #using optim function to optimize
  result <- optim(par=initialized_params,</pre>
                   fn=objfun,
                   method = "BFGS",
                   control = list(maxit=1000))
   if(result$convergence!=0){
     warning("Optimization did not converge.")
  optimezed_params <- result$par</pre>
  return(list(theta=optimezed_params[1:ncol(X)],
               sigma=optimezed_params[ncol(X)+1],
               value=-result$value,
               convergence=result$convergence))
}
#degree of freedom
DF <- function(lambda, X){</pre>
  XtX \leftarrow t(X)%*%X
                                                    #t(X)*x
                                                    \#t(X)*x+lambda*I
  XtX_lambda <- XtX+lambda*diag(ncol(X))</pre>
  Hat_matrix <- X%*%solve(XtX_lambda)%*%t(X)</pre>
  df <- sum(diag(Hat_matrix))</pre>
                                                              #trace of the hat matrix
  return(df)
}
X_test <- as.matrix(testS[,-1])</pre>
Y_test <- as.numeric(testS$motor_UPDRS)</pre>
#result and predicted values for training and testing data and MSE value
\#lambda=1
result1 <- RidgeOpt(1, X_train, Y_train)</pre>
theta1 <- result1$theta
sigma <- result1$sigma
predicted_1_Y_train <- X_train%*%theta1</pre>
predicted_1_Y_test <- X_test%*%theta1</pre>
MSE_1_trian <- mean((Y_train-predicted_1_Y_train)^2)</pre>
MSE_1_test <- mean((Y_test-predicted_1_Y_test)^2)</pre>
df_1 \leftarrow DF(1,X_{train})
cat("Training data MSE when lambda=1:",MSE_1_trian,"\n")
cat("Test data MSE when lambda=1:", MSE_1_test, "\n")
cat("Degree of freedom when lambda=1:",df_1,"\n")
```

```
\#lambda=100
result2 <- RidgeOpt(100, X_train, Y_train)</pre>
theta2 <- result2$theta
sigma2 <- result2$sigma</pre>
predicted_100_Y_train <- X_train%*%theta2</pre>
predicted_100_Y_test <- X_test%*%theta2</pre>
MSE_100_trian <- mean((Y_train-predicted_100_Y_train)^2)</pre>
MSE 100 test <- mean((Y test-predicted 100 Y test)^2)</pre>
df 100 <- DF(100, X train)
cat("Training data MSE when lambda=100:", MSE_100_trian,"\n")
cat("Test data MSE when lambda=100:", MSE_100_test, "\n")
cat("Degree of freedom when lambda=100:",df_100,"\n")
\#lambda=1000
result3 <- RidgeOpt(1000, X_train, Y_train)
theta3 <- result3$theta
sigma3 <- result3\sigma
predicted_1000_Y_train <- X_train%*%theta3</pre>
predicted_1000_Y_test <- X_test%*%theta3</pre>
MSE_1000_trian <- mean((Y_train-predicted_1000_Y_train)^2)</pre>
MSE 1000 test <- mean((Y test-predicted 1000 Y test)^2)</pre>
df_1000 <- DF(1000,X_train)</pre>
cat("Training data MSE when lambda=1000:", MSE 1000 trian,"\n")
cat("Test data MSE when lambda=1000:",MSE_1000_test,"\n")
cat("Degree of freedom when lambda=1000:",df_1000,"\n")
```

## 6.3 Code for assignment 3

```
data = read.csv("LAB1data/pima-indians-diabetes.csv")
colnames(data) = list("pegnant_times",
                      "glucose_level",
                      "blood_pressure",
                      "skin_thickness",
                      "serum_insulin",
                      "boby_mass",
                      "diabetes_pedigree_func",
                      "age",
                      "diabetes")
library(lattice)
library(caret)
library(ggplot2)
# plot 1
ggplot(data, aes(age, glucose_level, color = as.factor(diabetes))) +
  geom_point(size = 2) +
  labs(title = "Scatter Plot between Age and Glucose Level",
       x = "Age",
       v = "Glucose Level") +
  scale_color_manual(values = c("0" = "green", "1" = "red")) +
  theme_minimal()
# model 1
```

```
model_data = as.data.frame(x = cbind(data$glucose_level, data$age, data$diabetes))
colnames(model_data) = list("x1", "x2", "label")
model = train(as.factor(label) ~ .,
              data = model data,
              method = "glm",
              family = "binomial")
output_prob = predict(model, model_data, type = "prob")[,2]
output_label = ifelse(output_prob >= 0.5, 1, 0)
loss_model1 = mean((as.numeric(output_label) - as.numeric(model_data$label)) ** 2)
cat("The misclassification error is:", loss_model1,"\n")
acc_model1 = mean(output_label == model_data$label)
cat("The accuracy can be considered as quality of classification:", acc_model1)
# plot 2
ggplot(cbind(model_data,output_label), aes(x2, x1, color = as.factor(output_label))) +
  geom_point(size = 2) +
  labs(title = "Scatter Plot between Age and Glucose Level",
      x = "Age",
       y = "Glucose Level") +
  scale_color_manual(values = c("0" = "green", "1" = "red")) +
 theme_minimal()
coeff = coef(model$finalModel)
intercept = coeff[1]
param_x1 = coeff[2]
param_x2 = coeff[3]
cat("probabilistic equation:", intercept, "+", param_x1, "* x1 +", param_x2, "* x2 = y bar")
x2_range = seq(min(model_data$x2), max(model_data$x2), length.out = dim(model_data))
x1_bound = -(intercept + param_x2 * x2_range) / param_x1
# plot 3
ggplot(cbind(model_data,output_label), aes(x2, x1)) +
  geom_point(aes(color = as.factor(output_label)), size = 2) +
  geom line(aes(x = x2 range, y = x1 bound), color = "black", linetype = "dashed")+
  labs(title = "Scatter Plot between Age and Glucose Level",
      x = "Age",
      y = "Glucose Level") +
  scale_color_manual(values = c("0" = "green", "1" = "red")) +
 theme minimal()
r = c(0.2, 0.8)
output_label = ifelse(output_prob >= r[1], 1, 0)
# plot 4
ggplot(cbind(model_data,output_label), aes(x2, x1, color = as.factor(output_label))) +
  geom_point(size = 2) +
  labs(title = "Scatter Plot between Age and Glucose Level",
      x = "Age",
       y = "Glucose Level") +
  scale_color_manual(values = c("0" = "green", "1" = "red")) +
```

```
theme_minimal()
output_label = ifelse(output_prob >= r[2], 1, 0)
ggplot(cbind(model_data,output_label), aes(x2, x1, color = as.factor(output_label))) +
  geom_point(size = 2) +
  labs(title = "Scatter Plot between Age and Glucose Level",
       x = "Age",
       y = "Glucose Level") +
  scale_color_manual(values = c("0" = "green", "1" = "red")) +
  theme minimal()
label = model_data$label
model data$label = NULL
for (i in 0:4){
 feature_name = paste0("z", as.character(i+1))
  model_data[[feature_name]] = (model_data$x1 ** (4 - i)) * (model_data$x2 ** i)
}
model_data$label = label
head(model_data)
# model 2
model = train(as.factor(label) ~ .,
              data = model_data,
              method = "glm",
              family = "binomial")
output_prob = predict(model, model_data, type = "prob")[,2]
output_label = ifelse(output_prob >= 0.5, 1, 0)
loss_model2 = mean((as.numeric(output_label) - as.numeric(model_data$label)) ** 2)
acc_model2 = mean(output_label == model_data$label)
cat("The misclassification error is:", loss_model2, "\n")
cat("The accuracy can be considered as quality of classification:", acc_model2)
db_points = model_data[abs(output_prob - 0.5) < 0.05,]</pre>
# plot 5
ggplot() +
  geom_point(data = cbind(model_data, output_label), aes(x = x2, y = x1, color = as.factor(output_label)
  labs(title = "Scatter Plot between Age and Glucose Level",
       x = "Age",
       y = "Glucose Level") +
  scale_color_manual(values = c("0" = "green", "1" = "red")) +
  geom_smooth(data = db_points, aes(x = x2, y = x1), method = "loess", color = "black", linetype = "dash
 theme minimal()
```

#### 6.4 Figures

Actual													
Predicted	0	1	2	3	4	5	6	7	8	9			
0	88	0	0	0	1	0	0	0	0	0			
1	0	85	0	0	0	0	2	1	4	1			
2	0	3	105	0	0	0	0	0	0	0			
3	0	0	0	88	0	3	0	0	1	2			
4	0	0	0	0	99	0	0	0	1	0			
5	0	0	0	1	0	80	0	1	0	0			
6	0	0	0	1	0	1	79	0	0	0			
7	0	0	1	3	4	0	0	88	0	6			
8	0	0	3	2	1	1	0	0	93	0			
9	0	1	1	0	2	6	0	1	1	95			

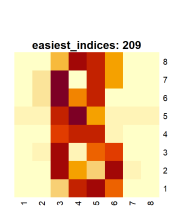
Figure 4: A.1.3: Test Confusion Matrix

Misclassification rates:

Training data: 0.0424 | Test data: 0.0586

Misclassification of 0: 0.0000 Misclassification of 1: 0.0449 Misclassification of 2: 0.0455 Misclassification of 3: 0.0737 Misclassification of 4: 0.0748 Misclassification of 5: 0.1209 Misclassification of 6: 0.0247 Misclassification of 7: 0.0330 Misclassification of 8: 0.0700 Misclassification of 9: 0.0865

Figure 5: A.1.4: Misclassification Rate



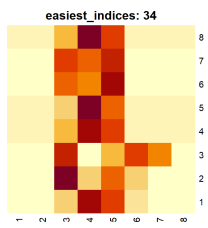


Figure 6: A.1.5: Heatmap of Digit 8\_easiest

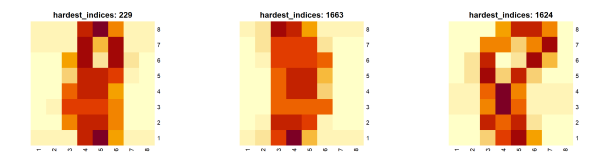


Figure 7: A.1.6: Heatmap of Digit 8\_hardest

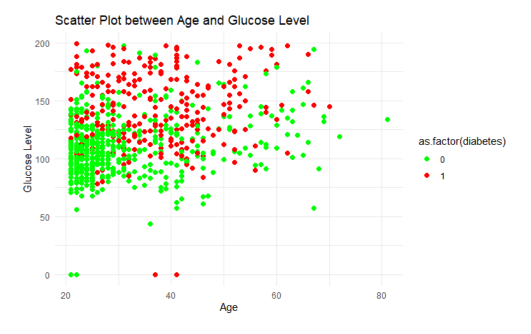


Figure 8: A.3.1: Raw Data Distribution

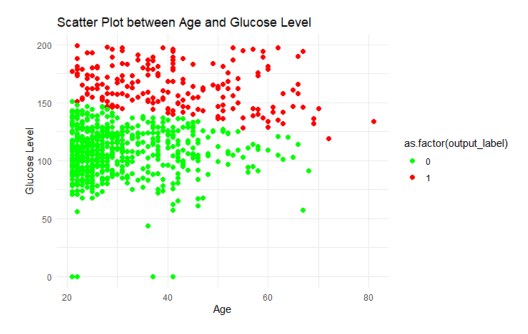


Figure 9: A.3.2: Prediction Outcome

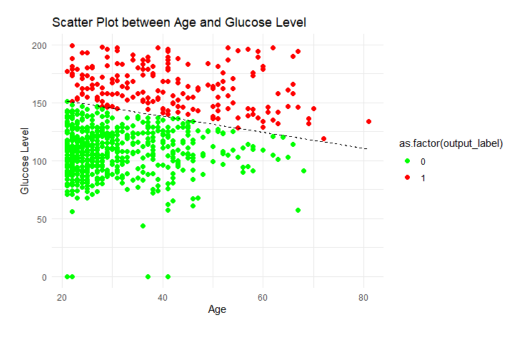


Figure 10: A.3.3: Prediction Outcome with Decision Boundary

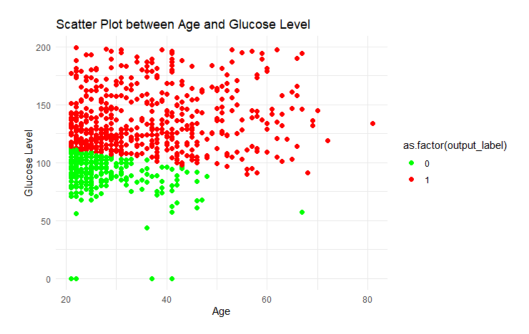


Figure 11: A.3.4.1: Prediction Outcome with Threhold = 0.2

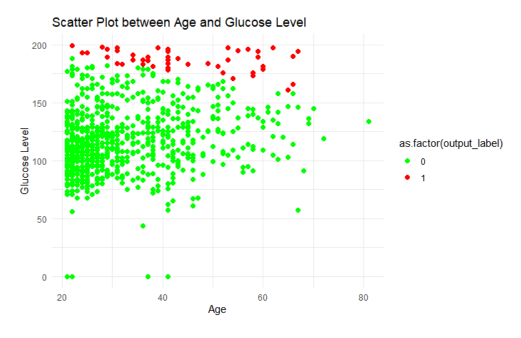


Figure 12: A.3.4.2: Prediction Outcome with Threhold = 0.8

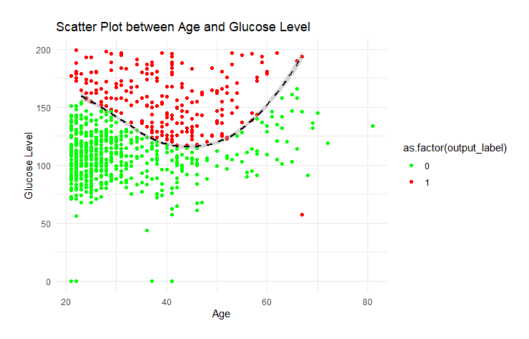


Figure 13: A.3.5: Prediction Outcome based Computed Features with Decision Boundary