Machine Learning Lab1

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Assignment 1. Handwritten digit recognition with Knearest neighbors.

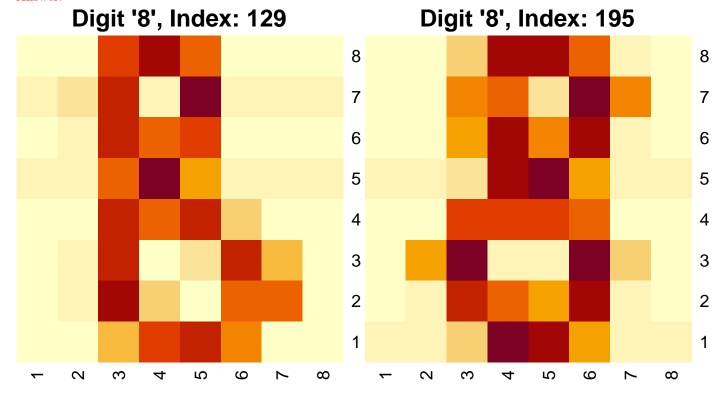
```
Answer:
2.
## Confusion Matrix for Training Data:
##
##
                  2
                                            8
                                                9
         0
                      3
                               5
                           0
                               0
                                                0
##
     0 202
              0
                  0
##
     1
         0 179
                 11
                      0
                           0
                               0
                                                3
##
                190
                      0
                           0
                               0
                                                0
##
     3
         0
              0
                  0 185
                           0
                                                1
                      0 159
         0
                  0
                           0 171
                                    0
##
              0
                      1
                               0 190
##
         0
                      0
                           0
##
     7
         0
              3
                  0
                      0
                           0
                               0
                                    0 178
##
             10
                           2
##
                                            3 183
  Confusion Matrix for Test Data:
##
        0
##
     0 82
           0
        0 90
##
               0 85
                    89
##
     6
               0
     7
##
                            0
                              99
##
        0
           7
               0
                  0
                     0
                        0
                            0
                               0 84
                     0
## Misclassification errors for the training data are: 0.04500262
## Misclassification errors for the test data are: 0.0585162
## The accuracy of prediction of for digit 0 is: 0.9761905
```

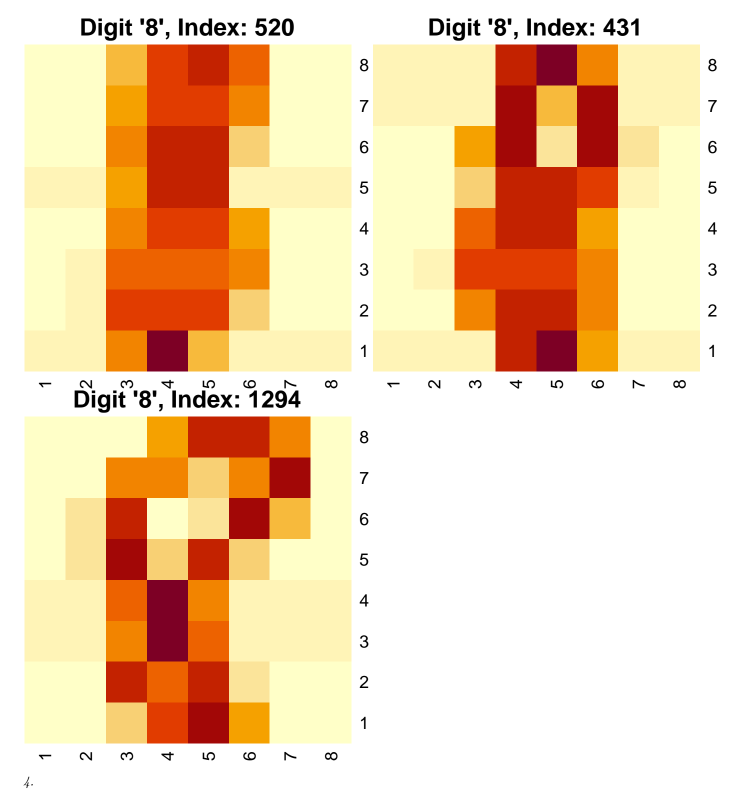
1.

```
## The accuracy of prediction of for digit 1 is: 0.9473684
## The accuracy of prediction of for digit 2 is:
                                                  0.9583333
## The accuracy of prediction of for digit 3 is:
## The accuracy of prediction of for digit 4 is:
                                                  0.872549
## The accuracy of prediction of for digit 5
                                             is:
## The accuracy of prediction of for digit 6
                                             is:
## The accuracy of prediction of for digit
                                              is:
                                                  0.980198
## The accuracy of prediction of for digit 8
                                                  0.9230769
                                             is:
## The accuracy of prediction of for digit 9 is:
## The overall accuracy of prediction is: 0.9414838
```

3.

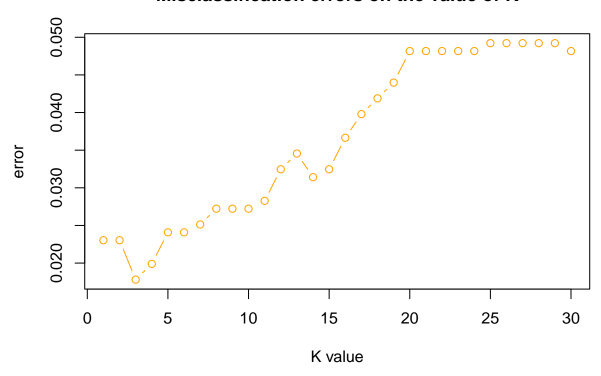
Answer:





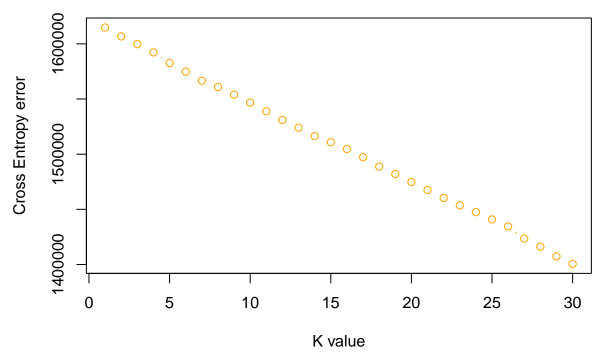
Answer:

Misclassification errors on the value of K



5. Answer:

Cross Entropy errors on the value of K



The optimal k is: 30

Assignment 2. Linear regression and ridge regression

```
1.
Answer:
2.
Answer:
## The training data's MSE is: 0.8785431
## The training data's MSE is: 0.9191113
##
## Call:
## lm(formula = motor_UPDRS ~ ., data = scaled_train_data)
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -3.0255 -0.7363 -0.1087 0.7333 2.1960
##
## Coefficients:
##
                  Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                 6.563e-15 1.583e-02 0.000 1.000000
## Jitter...
                1.869e-01 1.496e-01
                                      1.250 0.211496
## Jitter.Abs.
                -1.696e-01 4.081e-02 -4.156 3.32e-05 ***
                -5.270e+00 1.884e+01 -0.280 0.779688
## Jitter.RAP
## Jitter.PPQ5
              -7.457e-02 8.778e-02 -0.850 0.395659
## Jitter.DDP
              5.250e+00 1.884e+01 0.279 0.780541
                5.924e-01 2.060e-01 2.876 0.004055 **
## Shimmer
## Shimmer.dB.
                -1.727e-01 1.393e-01 -1.239 0.215380
## Shimmer.APQ3 3.207e+01 7.717e+01 0.416 0.677738
## Shimmer.APQ5 -3.875e-01 1.138e-01 -3.405 0.000669 ***
## Shimmer.APQ11 3.055e-01 6.124e-02
                                      4.989 6.37e-07 ***
## Shimmer.DDA -3.239e+01 7.717e+01 -0.420 0.674739
## NHR
                -1.854e-01 4.557e-02 -4.068 4.85e-05 ***
## HNR
                -2.385e-01 3.640e-02 -6.553 6.45e-11 ***
                4.068e-03 2.267e-02
## RPDE
                                      0.179 0.857576
## DFA
                -2.803e-01 2.014e-02 -13.919 < 2e-16 ***
## PPE
                2.265e-01 3.289e-02
                                      6.886 6.75e-12 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.9396 on 3508 degrees of freedom
## Multiple R-squared: 0.1212, Adjusted R-squared: 0.1172
## F-statistic: 30.24 on 16 and 3508 DF, p-value: < 2.2e-16
3.
Answer:
4.
Answer:
## Ridge Training MSE: 7.323041e+28 1.357493e+29 4.687817e+28
## Ridge Test MSE: 7.299556e+28 1.356537e+29 4.692501e+28
## Degrees of Freedom: 13.86074 9.924887 5.643925
```

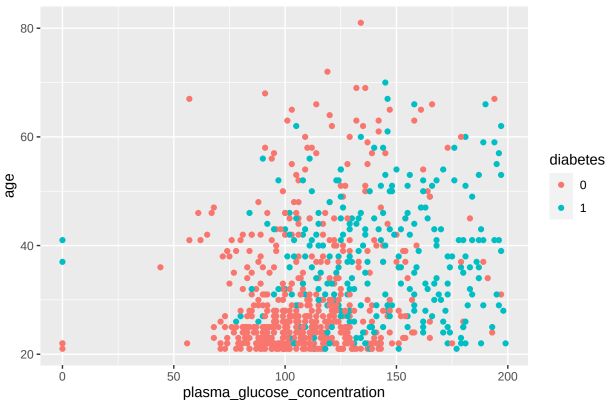
Assignment 3. Logistic regression and basis function expansion

1.

Answer:

```
## Loading required package: lattice
##
## Attaching package: 'caret'
## The following object is masked from 'package:kknn':
##
## contr.dummy
```

Scatter Plot of Plasma Glucose Concentration on Age



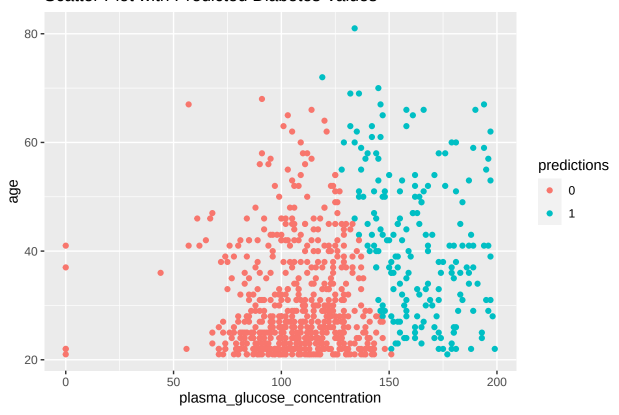
2.

Answer:

```
##
## Call:
  glm(formula = diabetes ~ plasma_glucose_concentration + age,
       family = binomial, data = pima_data)
##
## Deviance Residuals:
       Min
                 1Q
                      Median
                                   3Q
                                           Max
## -2.3367 -0.7775 -0.5087
                               0.8367
                                        3.1630
## Coefficients:
                                 Estimate Std. Error z value Pr(>|z|)
```

```
## (Intercept)
                               -5.912449
                                           0.462620 -12.78 < 2e-16 ***
## plasma_glucose_concentration  0.035644
                                           0.003290
                                                      10.83 < 2e-16 ***
                                0.024778
                                           0.007374
                                                       3.36 0.000778 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 993.48 on 767
                                     degrees of freedom
## Residual deviance: 797.36 on 765 degrees of freedom
## AIC: 803.36
##
## Number of Fisher Scoring iterations: 4
## [1] "Probability(Diabetes=1) = 1 / (1 + exp(-(-5.91244906318139 + 0.0356440425816302*x1 + 0.02477835)
## Misclassification Error: 0.2630208
```

Scatter Plot with Predicted Diabetes Values

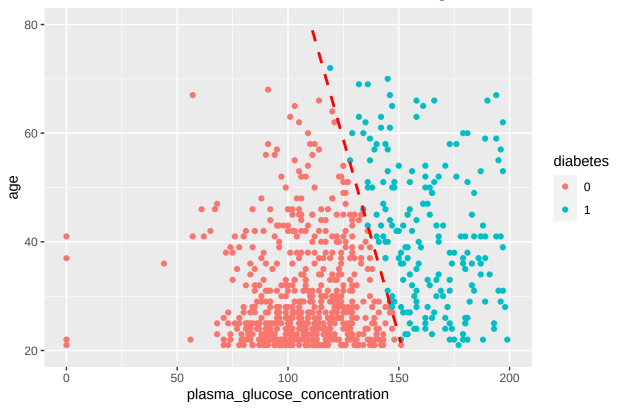


3.

Answer:

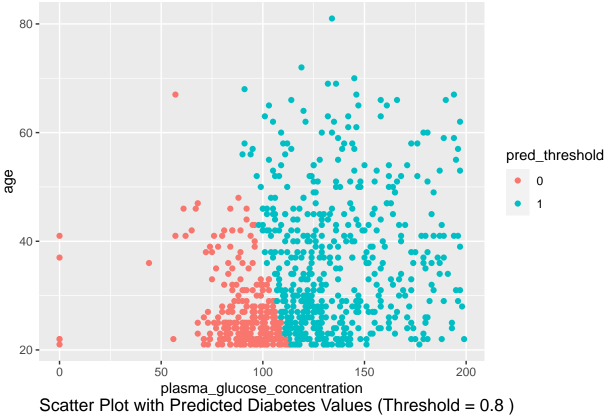
Warning: Removed 1 rows containing missing values (`geom_point()`).
Warning: Removed 454 rows containing missing values (`geom_line()`).

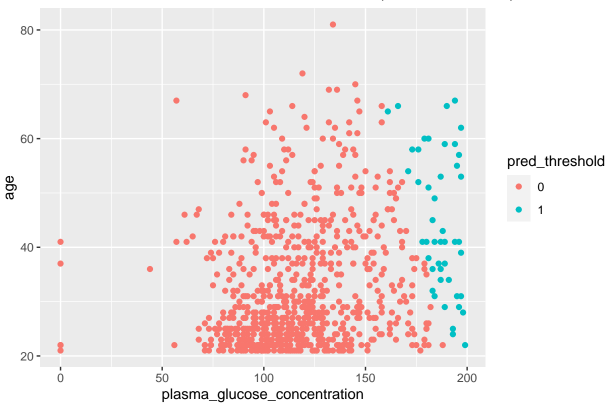
Scatter Plot of Plasma Glucose Concentration on Age with Decision Bounda



4. Answer:

Scatter Plot with Predicted Diabetes Values (Threshold = 0.2)

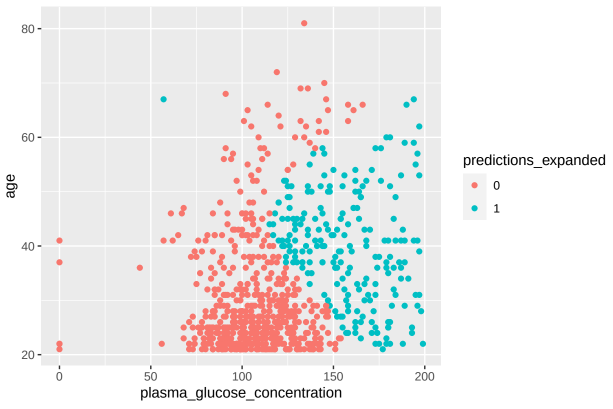




5.

Answer:

Scatter Plot with Predicted Diabetes Values (Expanded Features)



Misclassification Error (Expanded Features): 0.2447917

Appendix:

knearest.R

```
# load necessary libraries
library(ggplot2)
library(kknn)
# import data set
optdigits_data <- read.csv('optdigits.csv', header = FALSE)</pre>
colnames(optdigits_data) <- c(paste0("a",1:64),"digit")</pre>
optdigits_data$digit <- as.factor(optdigits_data$digit)</pre>
#head(optdigits_data, 5)
n=dim(optdigits_data)[1]
set.seed(12345)
id=sample(1:n, floor(n*0.5))
train_data=optdigits_data[id,]
id1=setdiff(1:n, id)
id2=sample(id1, floor(n*0.25))
valid_data=optdigits_data[id2,]
id3=setdiff(id1,id2)
test_data=optdigits_data[id3,]
```

```
# 30-nearest neighbor classification
k_fit_train <- kknn(formula = digit ~ ., train_data, train_data, k = 30, kernel = "rectangular")</pre>
k_fit_test <- kknn(formula = digit ~ ., train_data, test_data, k = 30, kernel = "rectangular")
# Confusion matrices for the training and test data
train_confusion <- table(train_data$digit, fitted(k_fit_train))</pre>
cat("Confusion Matrix for Training Data:\n")
print(train confusion)
test_confusion <- table(test_data$digit, fitted(k_fit_test))</pre>
cat("Confusion Matrix for Test Data:\n")
print(test_confusion)
# Misclassification errors for the training and test data
train_error <- 1 - sum(diag(train_confusion)) / sum(train_confusion)</pre>
cat("Misclassification errors for the training data are: ", train_error, "\n")
test_error <- 1 - sum(diag(test_confusion)) / sum(test_confusion)</pre>
cat("Misclassification errors for the test data are: ", test_error, "\n")
# the quality of predictions for different digits
for ( i in 1:nrow(test confusion)) {
 digit_accuracy <- test_confusion[i,i] / sum(test_confusion[i,])</pre>
  cat("The accuracy of prediction of for digit ", i-1, " is: ", digit_accuracy, "\n")
overall_accuracy <- sum(diag(test_confusion)) / sum(test_confusion)</pre>
cat("The overall accuracy of prediction is:", overall_accuracy, "\n")
# Get probabilities of class "8"
probabilities <- k_fit_train$prob[,"8"]</pre>
# Get indices of training data for class "8"
indices_8 <- which(train_data$digit == "8")</pre>
# Get probabilities for class "8"
probabilities 8 <- probabilities[indices 8]</pre>
# Find 2 easiest (highest probability) and 3 hardest (lowest probability) to classify cases
easiest_indices <- indices_8[order(probabilities_8, decreasing = TRUE)[1:2]]</pre>
hardest_indices <- indices_8[order(probabilities_8)[1:3]]
# Reshape features as 8x8 matrix and visualize
for (index in c(easiest_indices, hardest_indices)) {
 digit_8 <- matrix(as.numeric(train_data[index, 1:64]), nrow = 8, byrow = TRUE)
 heatmap(digit_8, Colv = NA, Rowv = NA, main = paste("Digit '8', Index:", index))
# Fit KNN for different K values and plot errors
errors <- data.frame()</pre>
for (k in 1:30) {
 fit <- kknn(digit ~ ., train_data, valid_data, k = k, kernel = "rectangular")</pre>
```

```
pred <- fit$fitted.values</pre>
  confusion_matrix <- table(valid_data$digit, pred)</pre>
  error <- 1 - sum(diag(confusion_matrix)) / sum(confusion_matrix)</pre>
  errors <- rbind(errors, data.frame(K = k, Error = error))</pre>
\# Plot the misclassification errors on the value of K
plot(errors$K, errors$Error, type = "b", col='orange', main="Misclassification errors on the value of K
# Initialize a data frame to store the results
ce_errors <- data.frame(k_value = integer(), cross_entropy_error = numeric())</pre>
for (k in 1:30) {
  # Fit K-nearest neighbor classifier
  fit <- kknn(digit ~ ., train_data, valid_data, k = k, kernel = "rectangular")</pre>
  # Get predicted probabilities
  predicted_probabilities <- fit$prob</pre>
  # Compute cross-entropy error
  #actual_probabilities <- ifelse(valid_data$digit == "8", 1, 0)</pre>
  #cross_entropy <- -sum(actual_probabilities * log(predicted_probabilities + 1e-15))</pre>
  cross_entropy <- -sum(as.numeric(valid_data$digit) * log(predicted_probabilities + 1e-15))</pre>
  ce_errors <- rbind(ce_errors, data.frame(k_value = k, cross_entropy_error = cross_entropy))</pre>
}
# Plot the results
plot(ce_errors$k_value, ce_errors$cross_entropy_error, type = "b", col='orange', main="Cross Entropy er
\# Find the optimal K
optimal_k <- ce_errors$k_value[which.min(ce_errors$cross_entropy_error)]
cat("The optimal k is:", optimal_k, "\n")
linear ridge.R
# import data set
parkinson_data <- read.csv('parkinsons.csv', header = TRUE)</pre>
#head(parkinson_data, 5)
# Split the data into training and test sets
parkinson_data <- subset(parkinson_data, select = -c(1:4,6))</pre>
n=dim(parkinson_data)[1]
set.seed(12345)
id=sample(1:n, floor(n*0.6))
train_data=parkinson_data[id,]
test_data=parkinson_data[-id,]
# Scale the data
\#scaled\_train\_data \leftarrow as.data.frame(cbind(scale(subset(train\_data, select = -c(motor\_UPDRS))), motor\_UPDRS)))
\#scaled\_test\_data < -as.data.frame(cbind(scale(subset(test\_data, select = -c(motor\_UPDRS))), motor\_UPDR
scaled_train_data <- as.data.frame(scale(train_data))</pre>
scaled_test_data <- as.data.frame(scale(test_data))</pre>
# linear regression model
```

```
lm_model <- lm(motor_UPDRS ~ ., data = scaled_train_data)</pre>
 # Training and test MSE
train_pred <- predict(lm_model, newdata = scaled_train_data)</pre>
train_mse <- mean((scaled_train_data$motor_UPDRS - train_pred)^2)</pre>
cat("The training data's MSE is: ", train_mse)
test pred <- predict(lm model, newdata = scaled test data)</pre>
test mse <- mean((scaled test data$motor UPDRS - test pred)^2)
cat("The training data's MSE is: ", test_mse)
# Significant variables
summary(lm_model)
 # Log-likelihood function
log_likelihood_fun <- function(theta, sigma, y, X) {</pre>
       n <- length(y)</pre>
       \log_{1/2} - \frac{1}{2 \cdot \sin^2} \times \frac{1}{2 \cdot \sin^2} - \frac{1}{2 \cdot \sin^2} \times \frac{
       return(log_likelihood)
}
 # Ridge log-likelihood function
ridge_log_Likelihood_fun <- function(theta, sigma, lambda, y, X) {</pre>
       log_like <- log_likelihood_fun(theta, sigma, y, X)</pre>
       ridge_penalty <- lambda * sum(theta^2)</pre>
       return(-log_like + ridge_penalty)
 # Ridge log-likelihood optimization function
ridge_log_likelihood_opt <- function(lambda, y, X) {</pre>
       start <- c(rep(0, ncol(X)), sd(y)) # combine theta and sigma into one numeric vector
       theta_length <- ncol(X)</pre>
       # Define a function for the optim() call
       fn_to_optim <- function(params) {</pre>
              theta <- params[1:theta_length]</pre>
              sigma <- params[theta_length + 1]</pre>
              return(-ridge_log_Likelihood_fun(theta, sigma, lambda, y, X))
       }
       # Call optim()
       opt_res <- optim(start, fn = fn_to_optim, method = "BFGS")</pre>
       # Return theta and sigma separately
       return(list(theta = opt_res$par[1:theta_length], sigma = opt_res$par[theta_length + 1]))
 # Degrees of freedom function
df_fun <- function(lambda, X) {</pre>
      H <- solve(t(X) %*% X + lambda * diag(ncol(X))) %*% t(X) %*% X</pre>
       return(sum(diag(H)))
}
```

```
y <- scaled_train_data$motor_UPDRS</pre>
X <- as.matrix(subset(scaled_train_data, select = -c(motor_UPDRS)))</pre>
# Ridge optimization for different lambdas
lambdas \leftarrow c(1, 100, 1000)
opt_params <- lapply(lambdas, function(lambda) ridge_log_likelihood_opt(lambda, y, X))</pre>
# Predictions and MSE for different lambdas
ridge_train_preds <- lapply(1:length(opt_params), function(i) {</pre>
  theta <- opt_params[[i]]$theta</pre>
  as.matrix(subset(scaled_train_data, select = -c(motor_UPDRS))) %*% theta
})
#train_mses <- colMeans((scaled_train_data$motor_UPDRS - ridge_train_preds)^2)</pre>
ridge_train_mse <- sapply(ridge_train_preds, function(pred) mean((pred - scaled_train_data$motor_UPDRS)
cat("Ridge Training MSE:", ridge_train_mse, "\n")
ridge_test_preds <- lapply(1:length(opt_params), function(i) {</pre>
  theta <- opt_params[[i]]$theta
  as.matrix(subset(scaled_test_data, select = -c(motor_UPDRS))) %*% theta
})
ridge_test_mse <- sapply(ridge_test_preds, function(pred) mean((pred - scaled_test_data$motor_UPDRS)^2)
cat("Ridge Test MSE:", ridge_test_mse, "\n")
# Degrees of freedom for different lambdas
degrees_of_freedom <- sapply(lambdas, function(lambda) df_fun(lambda, X))</pre>
cat("Degrees of Freedom:", degrees_of_freedom, "\n")
logistic.R
library(ggplot2)
library(caret)
# import data set
pima_data <- read.csv('pima-indians-diabetes.csv', header = FALSE)</pre>
colnames(pima_data) <- c("num_of_pregnant", "plasma_glucose_concentration", "blood_pressure",</pre>
                                            "skinfold_thickness", "serum_insulin", "bmi",
                                            "diabetes_predigree", "age", "diabetes")
#head(pima_data, 5)
# Scatterplot
ggplot(pima_data, aes(x=plasma_glucose_concentration, y=age, color=as.factor(diabetes))) +
  geom_point() +
  labs(color="diabetes", title='Scatter Plot of Plasma Glucose Concentration on Age')
# Logistic Regression
logsitic_model <- glm(diabetes ~ plasma_glucose_concentration + age, data=pima_data, family=binomial)
summary(logsitic_model)
# Predict probabilities
probabilities <- predict(logsitic_model, type="response")</pre>
print(paste0("Probability(Diabetes=1) = 1 / (1 + exp(-(",coef(logsitic_model)[1]," + ", coef(logsitic_m
```

```
# Classify observations
predictions <- ifelse(probabilities >= 0.5, 1, 0)
# Compute misclassification error
mis_error <- mean(predictions != pima_data$diabetes)</pre>
cat("Misclassification Error:", mis_error, "\n")
# Scatter plot with predicted values
ggplot(data.frame(pima_data, predictions), aes(x=plasma_glucose_concentration, y=age, color=as.factor(p
   geom_point() +
   labs(color="predictions",title='Scatter Plot with Predicted Diabetes Values')
# Decision boundary equation
# Decision boundary is where the logistic function equals 0.5
\# 0 = intercept + coef1*x1 + coef2*x2
# x2 = -(intercept + coef1*x1) / coef2
\# decision\_boundary <--(coef(logsitic\_model)[1] + coef(logsitic\_model)[2]*pima\_data\$age) / coef(logsitic\_model)[2] + pima\_data\$age) / coef(logsitic\_model)[2] + pima\_data§age) / coef(logsitic\_mode
decision_boundary_x1 <- pima_data$plasma_glucose_concentration</pre>
decision_boundary_x2 <- -(coef(logsitic_model)[1] + coef(logsitic_model)[2]*decision_boundary_x1) / coe
# Add decision boundary to scatter plot
ggplot(data.frame(pima_data, predictions), aes(x=plasma_glucose_concentration, y=age, color=as.factor(p
   geom_point() +
   geom_line(aes(x=decision_boundary_x1, y=decision_boundary_x2), color='red', linetype='dashed', linewi
   labs(color="diabetes", title='Scatter Plot of Plasma Glucose Concentration on Age with Decision Bound
# Predictions with different thresholds
thresholds \leftarrow c(0.2, 0.8)
for ( threshold in thresholds) {
   pred_threshold <- ifelse(predict(logsitic_model, newdata=pima_data, type='response') >= threshold, 1,
   # Scatter plot with predicted values and threshold
   scatter_plot <- ggplot(data.frame(pima_data, pred_threshold), aes(x=plasma_glucose_concentration, y=a
       geom point() +
       labs(color="pred_threshold", title=paste('Scatter Plot with Predicted Diabetes Values (Threshold ='
   print(scatter_plot)
# Create new features
pima_data$z1 <- pima_data$plasma_glucose_concentration^4</pre>
pima_data$z2 <- pima_data$plasma_glucose_concentration^3 * pima_data$age
pima_data$z3 <- pima_data$plasma_glucose_concentration^2 * pima_data$age^2
pima_data$z4 <- pima_data$plasma_glucose_concentration * pima_data$age^3</pre>
pima_data$z5 <- pima_data$age^4</pre>
# Logistic Regression with new features
model_expanded <- glm(diabetes ~ plasma_glucose_concentration + age + z1 + z2 + z3 + z4 + z5, data=pima
```

```
# Predict probabilities and classify observations
probabilities_expanded <- predict(model_expanded, type="response")
predictions_expanded <- ifelse(probabilities_expanded >= 0.5, 1, 0)

# Scatter plot for the model with expanded features
ggplot(data.frame(pima_data, predictions_expanded), aes(x=plasma_glucose_concentration, y=age, color=as
    geom_point() +
    labs(color="predictions_expanded",title='Scatter Plot with Predicted Diabetes Values (Expanded Featur

# Compute misclassification error
mis_error_expanded <- mean(predictions_expanded != pima_data$diabetes)
cat("Misclassification Error (Expanded Features):", mis_error_expanded, "\n")</pre>
```