Lab Report: Computer lab 1 block 1

Group_A10

2024-11-25

Assignment 1: Handwritten Digit Recognition with K-Nearest Neighbors

Training Misclassification Error: 0.0429095
 Test Misclassification Error: 0.0324607

Confusion Matrix: Test Data

Predicted										
Actual	0	1	2	3	4	5	6	7	8	9
0	97	0	0	0	0	0	0	0	0	0
1	0	97	0	0	0	0	0	0	0	0
2	0	0	98	0	1	0	0	0	0	0
3	0	0	1	99	0	2	0	1	0	0
4	0	0	0	0	88	0	0	2	0	0
5	0	0	0	1	1	82	0	1	0	2
6	0	1	0	0	0	0	95	0	0	0
7	0	0	0	0	1	0	0	83	0	1
8	0	9	0	0	0	0	0	0	94	1
9	0	0	0	3	1	1	0	1	0	91

Confusion Matrix: Training Data

Predicted										
Actual	0	1	2	3	4	5	6	7	8	9
0	196	0	0	0	0	0	0	0	0	0
1	0	175	8	0	0	0	0	0	1	4
2	1	0	185	0	0	0	0	1	0	0
3	0	0	0	179	0	1	0	3	1	0
4	0	0	0	0	189	0	2	4	2	4
5	1	0	0	0	0	188	0	1	0	7
6	1	1	0	0	0	1	178	0	0	0
7	0	1	1	1	0	0	0	198	0	0
8	0	9	0	3	0	1	2	0	165	0
9	1	2	0	5	1	1	0	5	5	176

Summary of Prediction Quality

Key Results:

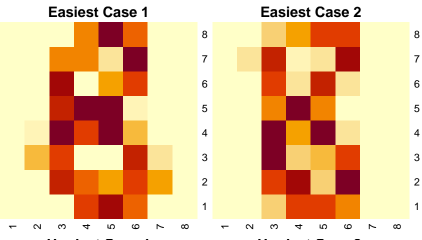
- Training Error: 4.36%
- Test Error: 4.62%
- Strong generalization across datasets.

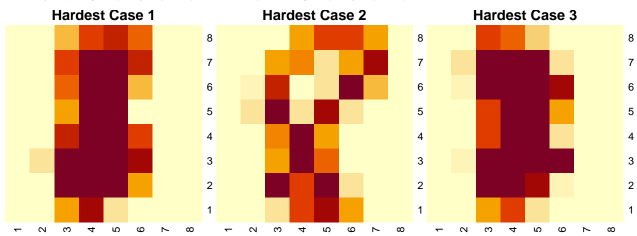
Class-Specific Observations:

• Best Performance: Digit 0 (perfect classification).

- Strong Performance: Digits 1, 2, 6, 7 (minimal errors).
- Challenging Cases:
 - Digits 3, 4, 5: Overlap with similar digits.
 - Digits 8, 9: High confusion with 1 and 3.

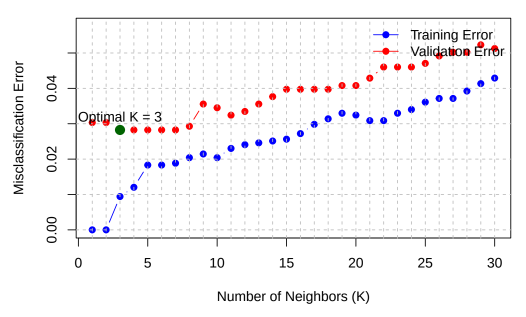
Visualizations of Easiest and Hardest Cases for Digit '8'





The hardest 3 cases are visually ambiguous and challenging to identify as "8," likely due to overlapping features or incomplete digit representation. In contrast, the easiest 2 cases are visually distinct and clearly recognizable as "8," making them straightforward to classify.

Training and Validation Errors vs. K in KNN



• Model Complexity:

- As K increases, the model becomes less complex because it considers more neighbors for classification. This effectively smooths the decision boundaries.
- For small K (e.g., K = 1), the model is highly complex as it relies only on the nearest neighbor, leading to potential overfitting.

• Effect on Errors:

- Training Error:

- * For small K, the training error is very low (almost 0) due to overfitting.
- * As K increases, the training error gradually rises because the model becomes less sensitive to individual training points.

- Validation Error:

- * Validation error initially decreases as K increases, reducing overfitting.
- * Beyond a certain point, validation error starts increasing due to underfitting when the model becomes overly smoothed.

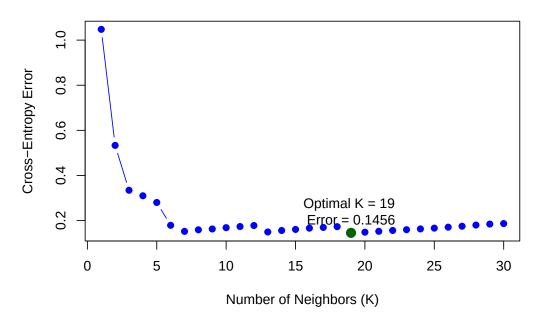
Optimal K According to the Plot

- From the plot, the **optimal** K corresponds to the minimum validation error.
- Optimal K=3 in this case, as the validation error is lowest at this point.

Comparison of Test, Training, and Validation Errors

The test misclassification error for K=3 is 0.0325, which is slightly higher than the validation error (0.0282) and significantly higher than the training error (0.0094). This indicates that the model generalizes well to unseen data while maintaining a low error on the training data.

Validation Cross-Entropy Error vs. K



Optimal K Value and Suitability of Cross-Entropy Error

- Optimal K:
 - From the graph, the **optimal** K is **19**, as the validation cross-entropy error is minimized at this point (**Error** = **0.1456**).
- Why Cross-Entropy is More Suitable:
 - Cross-entropy considers the confidence of predictions by penalizing incorrect or low-probability predictions more heavily.
 - Unlike misclassification error, which treats all incorrect predictions equally, cross-entropy error differentiates between predictions with varying levels of certainty.
 - In a multinomial distribution (as in this case with multiple classes), cross-entropy provides a
 probabilistic perspective, which is essential for understanding model confidence and calibration,
 making it more informative for fine-tuning models.

Assignment 2: Linear regression and ridge regression

• **Training MSE**: 6.0799301

• Test MSE: 6.5498889

The linear regression model shows that age, sex, total_UPDRS, Jitter.Abs., Shimmer.APQ5, and Shimmer.APQ11 are significant predictors of motor_UPDRS based on their low p-values (< 0.05). The total_UPDRS variable has the strongest influence, while others contribute moderately. The training MSE and test MSE indicate the model's accuracy on respective datasets.

Optimal theta parameters for different lambda values

- lambda = 1
- $\begin{array}{llll} \bullet & \textbf{Optimized Theta (coefficients)::} & 21.1858461, \; -0.1147378, \; -0.2160224, \; 0.3566923, \; -0.008651, \\ 7.7114936, \; 0.4944555, \; -0.5970692, \; -0.1654761, \; 0.1721167, \; -0.1563581, \; 0.491552, \; 0.129306, \; -0.0841912, \\ -1.3061248, \; 0.9164303, \; -0.0856044, \; 0.1199405, \; 0.03486, \; -0.2063699, \; -0.0694022, \; 0.4154516 \\ \end{array}$
- lambda = 100

- Optimized Theta (coefficients):: 21.189791, 0.4885039, 0.5076479, -0.1029531, 0.1492808, 2.935959, 0.0308058, -0.0974992, 0.0059612, 0.0123203, 0.0060051, 0.0234413, 0.0460907, -0.030851, -0.0016557, 0.1461861, -0.0308543, -0.0230367, -0.1611598, 0.0867156, -0.3214618, 0.2400944
- lambda = 1000
- Optimized Theta (coefficients):: 21.1858461, -0.1147378, -0.2160224, 0.3566923, -0.008651, 7.7114936, 0.4944555, -0.5970692, -0.1654761, 0.1721167, -0.1563581, 0.491552, 0.129306, -0.0841912, -1.3061248, 0.9164303, -0.0856044, 0.1199405, 0.03486, -0.2063699, -0.0694022, 0.4154516

Table 1: Ridge Regression Results for Different Lambda Values

Lambda	Train_MSE	Test_MSE	Degrees_of_Freedom	Optimized_Sigma
1	6.085829	6.56323	19.84347	2.467403
100	25.886877	26.94831	15.59068	5.088249
1000	61.361718	62.80625	10.04924	7.833606

From the table above, the choice of the penalty parameter λ significantly affects the model's performance and complexity. Here's the detailed analysis:

- lambda = 1:
 - Achieves the lowest **Train MSE** (6.085893) and **Test MSE** (6.560836).
 - Maintains a **Degrees of Freedom (DF)** of 19.84347, meaning more predictors contribute effectively to the model.
 - Indicates a balance between model complexity and generalization.
- lambda = 100:
 - Results in higher Train MSE (25.88616) and Test MSE (26.94751) compared to lambda = 1.
 - Reduces **DF** to 15.59068, indicating a stronger penalty and increased regularization.
 - May lead to underfitting as the model becomes overly simplified.
- lambda = 1000:
 - Has the highest **Train MSE** (61.36093) and **Test MSE** (62.8066), demonstrating significant underfitting.
 - Reduces **DF** further to 10.04924, suggesting excessive shrinkage of predictor coefficients.
 - Results in a simpler but less accurate model.

Based on the results and analysis:

- lambda = 1 is the most appropriate penalty parameter. It minimizes both Train MSE and Test MSE, while maintaining a high **Degrees of Freedom**, ensuring that the model is neither over-regularized nor under-regularized.
- As **lambda** increases, the model becomes overly regularized, leading to underfitting (higher MSE and lower DF).

Assignment 3

In this assignment we will try to predict diabetes using age and plasma glucose concentration, using logistic regression.

3.1

Below is scatterplot of the data.

```
set.seed(12345)
library(ggplot2)
df <- read.csv('data/pima-indians-diabetes.csv')</pre>
```

```
colnames(df) <- c('pregnant',</pre>
                  'plasma',
                  'blood_pressure',
                  'triceps',
                  'insulin',
                  'bmi',
                  'diabetes_pedigree',
                  'age',
                  'diabetes')
df$diabetes <- df$diabetes == 1
df |> ggplot() +
       geom_point(aes(x = age, y = plasma, color = diabetes)) +
       labs(x = "Age", y = "Plasma glucose concentration", color = "Has diabetes?")
   200
Plasma glucose concentration
   150 -
                                                                                    Has diabetes?
   100 -
                                                                                         FALSE
                                                                                         TRUE
    50 -
```

The success of doing logistic regression is predicated on being able to draw a good linear decision boundary. We see that there is chunk of false cases that can be easily separated, but then the remaining positive cases are mixed with some negative cases. So it should be possible to model with linear regression, but the accuracy might be affected by the mix of negative cases in the region that will be identified as the positive cases.

Age

60

80

40

3.2

0 -

20

We will model as the following. We will have a linear predictor $z = k1 + k2 \cdot x_2 + k_3 \cdot x_1$ where x_1 is plasma glucose concentration and x_2 is age. Then the random variable Y which is 1 if one has diabetes and 0 otherwise will have the distribution $p(Y = 1|x_1, x_2) = \frac{e^z}{1+e^z}$.

```
m = glm(diabetes ~ age + plasma, data=df, family="binomial")
df$predicted_prob <- predict(m, newdata = df, type = "response")</pre>
classification_threshold <- 0.5</pre>
df$prediction <- df$predicted_prob > classification_threshold
missclassification_rate <- sum(df$diabetes != df$prediction) / nrow(df)</pre>
missclassification rate
## [1] 0.2659713
df |> ggplot() +
       geom_point(aes(x = age, y = plasma, color = prediction)) +
       labs(x = "Age", y = "Plasma glucose concentration", color = "Predicted to have diabetes?")
   200 -
Plasma glucose concentration
   150 -
                                                                   Predicted to have diabetes?
   100 -
                                                                        FALSE
                                                                        TRUE
    50 -
     0 -
                         40
       20
                                          60
                                                           80
                                 Age
```

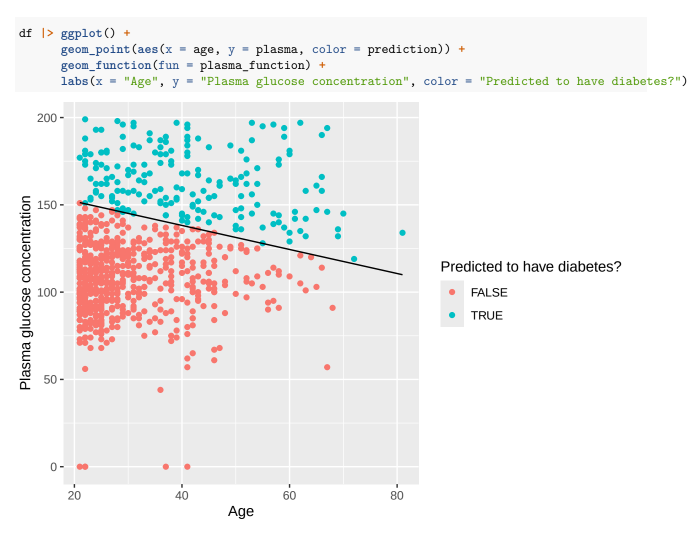
3.3

Since r = 0.5 we have that the decision boundary can be solved from the equation $\frac{e^{k_0 + k_1 \cdot x_2 + k_2 \cdot x_1}}{1 + e^{k_0 + k_1 \cdot x_2 + k_2 \cdot x_1}} = 0.5$

Which is equivalent to $k_0 + k_1 \cdot x_2 + k_2 \cdot x_1 = 0$

```
# 0 = k1 + k2 * age + k3 * plasma
k1 <- summary(m)$coefficients[1]
k2 <- summary(m)$coefficients[2]
k3 <- summary(m)$coefficients[3]

# plasma = (-k1 -k2 * age) / k3
plasma_function <- function(x)(-k1 -k2 * x) / k3</pre>
```



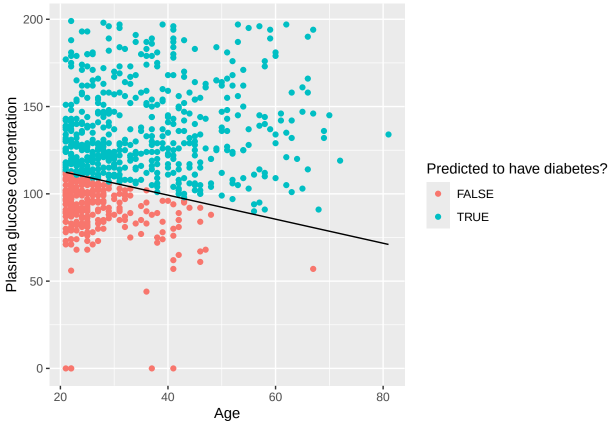
We see that it as we commented in the beginning of this assignment that it divides into a region where the negative cases are, but the remaining region for the positive cases isn't as good.

3.4

By changing r we can have our predictor be less or more sensitive. If we have a low value then it will do more positive guesses and with a high it will be reserved to guess a case as positive. We illustrate it with 0.2 and 0.8.

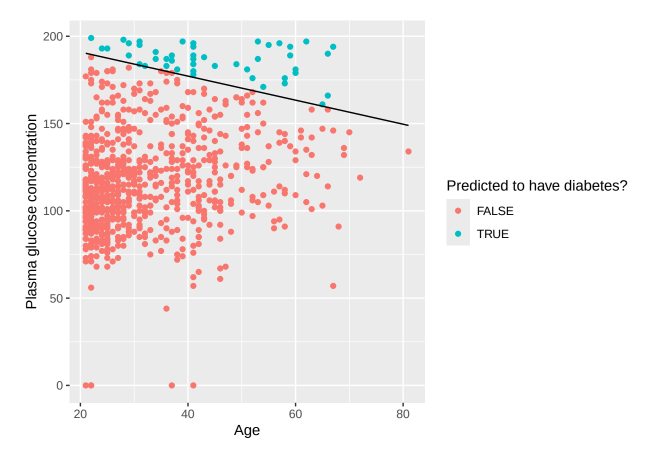
```
p <- 0.2
df$prediction02 <- df$predicted_prob > p
k4 <- log (p / (1-p))
plasma_function <- function(x)( k4 -k1 -k2 * x) / k3

df |> ggplot() +
    geom_point(aes(x = age, y = plasma, color = prediction02)) +
    geom_function(fun = plasma_function) +
    labs(x = "Age", y = "Plasma glucose concentration", color = "Predicted to have diabetes?")
```



```
p <- 0.8
df$prediction08 <- df$predicted_prob > p
k4 <- log (p / (1-p))
plasma_function <- function(x)( k4 -k1 -k2 * x) / k3

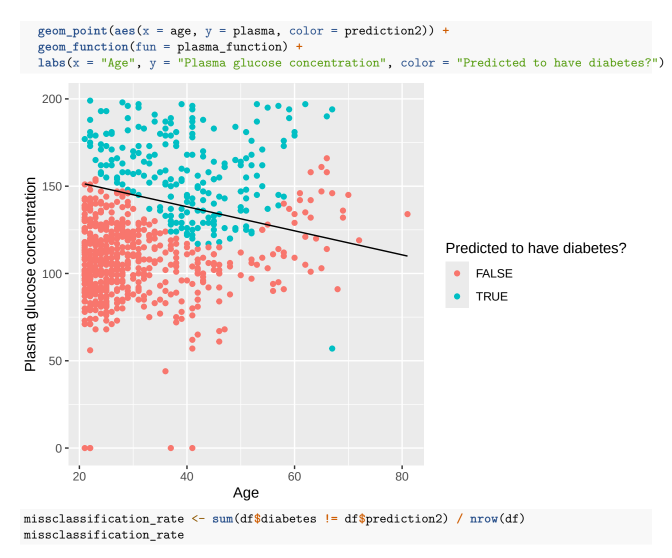
df |> ggplot() +
    geom_point(aes(x = age, y = plasma, color = prediction08)) +
    geom_function(fun = plasma_function) +
    labs(x = "Age", y = "Plasma glucose concentration", color = "Predicted to have diabetes?")
```



3.5

We can attempt to improve our model by adding new features that are functions of the already existing features. We will add the following features: $z_1 = x_1^4, z_2 = x_1^3 x_2, z_3 = x_1^2 x_2^2, z_4 = x_1 x_2^3, z_5 = z_2^4$.

```
df$z1 <- df$plasma ^ 4
df$z2 <- df$plasma ^ 3 * df$age
df$z3 <- df$plasma ^ 2 * df$age ^ 2</pre>
df$z4 <- df$plasma * df$age ^ 3</pre>
df$z5 <- df$age ^ 4
m2 = glm(diabetes \sim age + plasma + z1 + z2 + z3 + z4 + z5,
          data=df, family="binomial")
df$predicted_prob2 <- predict(m2, newdata = df, type = "response")</pre>
classification_threshold <- 0.5</pre>
df$prediction2 <- df$predicted_prob2 > classification_threshold
\# 0 = k1 + k2 * age + k3 * plasma
k1 <- summary(m)$coefficients[1]</pre>
k2 <- summary(m)$coefficients[2]</pre>
k3 <- summary(m)$coefficients[3]
\# plasma = (-k1 - k2 * age) / k3
plasma_function \leftarrow function(x)(-k1 -k2 * x) / k3
df |> ggplot() +
```



[1] 0.2464146

And we can see we get a better missclassification rate. It might seem like the decission boundary hasn't changed drastically, but now we must consider that we haven't graphed out all the variables being considered, thus we have decision plan in a higher dimensional space, and it just happens that the intersection with the age-plasma plane has similar line.

Assignment 4

It can be important to choose different probability thresholds since sometimes one might prioritize identifying all the positive cases more than avoiding false positives. An example is when detecting diseases, as the consequences for giving a false positive is less severe than missing out on identifying a true case (page 50).

The cost function associated to linear regression can be derived using MLE which is included in equation 3.21 on page 44

On page 14 it mentions that collecting values for supervised problems may be done by having an expert manually label data, or the data might already have recorded the variable we want to predict. # Appendix

Code for Asssignment 1

```
#Load and Preprocess Data
# Load dataset
file_path <- "data/optdigits.csv"</pre>
optdigits_data <- read.csv(file_path)</pre>
colnames(optdigits_data) <- c(paste0("pixel_", 1:64), "label")</pre>
optdigits_data$label <- as.factor(optdigits_data$label)</pre>
# Split into training and test sets
set.seed(123)
n <- nrow(optdigits_data)</pre>
# Generate indices
train_indices <- sample(1:n, size = 0.5 * n) # 50% for training
remaining_indices <- setdiff(1:n, train_indices) # Remaining 50%
test_indices <- sample(remaining_indices, size = 0.25 * n) # 25% for testing
validation_indices <- setdiff(remaining_indices, test_indices) # Remaining 25% for validation
# Subset the data
train data <- optdigits data[train indices, ]</pre>
test_data <- optdigits_data[test_indices, ]</pre>
validation_data <- optdigits_data[validation_indices, ]</pre>
# Train the KNN classifier
knn_model <- kknn(label ~ ., train = train_data, test = train_data, k = 30, kernel = "rectangular")
# Predictions and performance
train_predictions <- fitted(knn_model)</pre>
conf_matrix_train <- table(Actual = train_data$label, Predicted = train_predictions)</pre>
print("Confusion Matrix (Training):")
print(conf_matrix_train)
# Predict on test data
test_predictions <- fitted(kknn(label ~ ., train = train_data, test = test_data, k = 30, kernel = "rect
conf_matrix_test <- table(Actual = test_data$label, Predicted = test_predictions)</pre>
print("Confusion Matrix (Test):")
print(conf_matrix_test)
# Misclassification error for training data
train_total <- sum(conf_matrix_train) # Total instances in training data
train_correct <- sum(diag(conf_matrix_train)) # Correctly classified instances (sum of diagonal)
train_misclassification_error <- 1 - train_correct / train_total</pre>
cat("Training Misclassification Error:", train_misclassification_error, "\n")
# Misclassification error for test data
test_total <- sum(conf_matrix_test) # Total instances in training data
test correct <- sum(diag(conf matrix test)) # Correctly classified instances (sum of diagonal)
test_misclassification_error <- 1 - test_correct / test_total</pre>
cat("Test Misclassification Error:", test_misclassification_error, "\n")
```

```
# Get predicted probabilities for training data
train_probabilities <- knn_model$prob # Matrix of probabilities</pre>
train_actual <- train_data$label # Actual labels for training data
# Extract rows corresponding to digit "8" in the training data
digit_8_indices <- which(train_actual == "8")</pre>
digit_8_probs <- train_probabilities[digit_8_indices, ] # Probabilities for digit "8"</pre>
digit_8_predicted_probs <- digit_8_probs[, "8"]</pre>
easiest_indices <- digit_8_indices[order(digit_8_predicted_probs, decreasing = TRUE)[1:2]]</pre>
hardest_indices <- digit_8_indices[order(digit_8_predicted_probs, decreasing = FALSE)[1:3]]
# Extract feature data (pixels) for easiest and hardest cases
easiest_features <- train_data[easiest_indices, 1:64] # Features for easiest cases
hardest_features <- train_data[hardest_indices, 1:64] # Features for hardest cases
# Reshape into 8x8 matrices
# Reshape and store as numeric matrices
easiest_matrices <- lapply(1:nrow(easiest_features), function(i) {</pre>
  matrix(as.numeric(easiest_features[i, ]), nrow = 8, byrow = TRUE)
})
hardest_matrices <- lapply(1:nrow(hardest_features), function(i) {</pre>
  matrix(as.numeric(hardest_features[i, ]), nrow = 8, byrow = TRUE)
})
# Visualize easiest cases
heatmap(easiest_matrices[[1]], Colv = NA, Rowv = NA, scale = "none", main = paste("Easiest Case", 1), x
heatmap(easiest_matrices[[2]], Colv = NA, Rowv = NA, scale = "none", main = paste("Easiest Case", 2), x
# Visualize hardest cases
heatmap(hardest_matrices[[1]], Colv = NA, Rowv = NA, scale = "none", main = paste("Hardest Case", 1), x
heatmap(hardest_matrices[[2]], Colv = NA, Rowv = NA, scale = "none", main = paste("Hardest Case", 2), x
heatmap(hardest_matrices[[3]], Colv = NA, Rowv = NA, scale = "none", main = paste("Hardest Case", 3), x
# Initialize vectors to store errors
k_values <- 1:30 # Values of K to test
train_errors <- numeric(length(k_values)) # Training errors</pre>
validation_errors <- numeric(length(k_values)) # Validation errors</pre>
# Loop through each value of K
for (k in k_values) {
  # Train KNN model on training data
  knn_model <- kknn(label ~ ., train = train_data, test = train_data, k = k, kernel = "rectangular")
  train_predictions <- fitted(knn_model)</pre>
  # Calculate training misclassification error
  conf_matrix_train <- table(Actual = train_data$label, Predicted = train_predictions)</pre>
  train_correct <- sum(diag(conf_matrix_train))</pre>
  train_total <- sum(conf_matrix_train)</pre>
```

```
train_errors[k] <- 1 - train_correct / train_total</pre>
  # Validate on validation data
  validation_predictions <- fitted(kknn(label ~ ., train = train_data, test = validation_data, k = k, k</pre>
  conf_matrix_validation <- table(Actual = validation_data$label, Predicted = validation_predictions)</pre>
  validation_correct <- sum(diag(conf_matrix_validation))</pre>
  validation_total <- sum(conf_matrix_validation)</pre>
  validation errors[k] <- 1 - validation correct / validation total
}
#
# Plot training and validation errors
plot(k_values, train_errors, type = "b", col = "blue", pch = 19,
     ylim = c(0, max(c(train_errors, validation_errors)) * 1.1),
     xlab = "Number of Neighbors (K)",
     ylab = "Misclassification Error",
     main = "Training and Validation Errors vs. K in KNN")
lines(k_values, validation_errors, type = "b", col = "red", pch = 19)
# Add gridlines
abline(h = seq(0, max(c(train_errors, validation_errors)), by = 0.01), col = "gray", lty = 2)
abline(v = seq(1, 30, by = 1), col = "gray", lty = 2)
# Highlight optimal K
optimal_k <- which.min(validation_errors)</pre>
points(optimal_k, validation_errors[optimal_k], col = "darkgreen", pch = 19, cex = 1.5)
text(optimal_k, validation_errors[optimal_k], labels = paste("Optimal K =", optimal_k), pos = 3)
# Add legend
legend("topright", legend = c("Training Error", "Validation Error"),
       col = c("blue", "red"), pch = 19, lty = 1, bty = "n")
# Train the model with optimal K
optimal_k <- 3
knn_model_optimal <- kknn(label ~ ., train = train_data, test = test_data, k = optimal_k, kernel = "rec
# Predictions and confusion matrix
test_predictions <- fitted(knn_model_optimal)</pre>
conf_matrix_test <- table(Actual = test_data$label, Predicted = test_predictions)</pre>
# Calculate test misclassification error
test_total <- sum(conf_matrix_test)</pre>
test_correct <- sum(diag(conf_matrix_test))</pre>
test_misclassification_error <- 1 - test_correct / test_total</pre>
cat("Test Misclassification Error for K =", optimal_k, ":", test_misclassification_error, "\n")
#Plot cross-entropy errors for validation data
plot(k_values, cross_entropy_errors, type = "b", col = "blue", pch = 19,
     xlab = "Number of Neighbors (K)", ylab = "Cross-Entropy Error",
```

Code for Asssignment 2

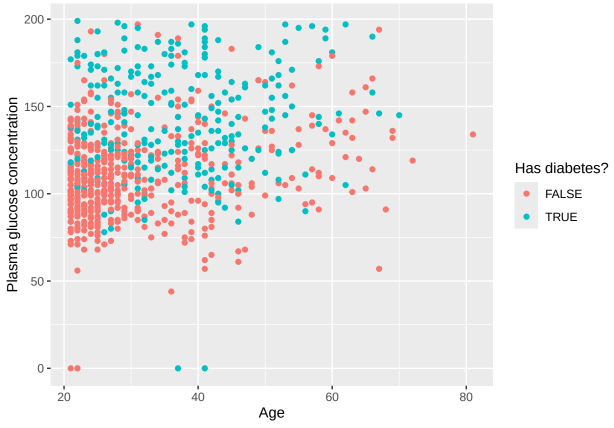
```
scale_features <- function(data) {</pre>
  data_scaled <- data
  data_scaled[ , -which(names(data) == "motor_UPDRS")] <- scale(data[ , -which(names(data) == "motor_UPDRS")]</pre>
  return(data_scaled)
# Define the Loglikelihood function
Loglikelihood <- function(theta, sigma, X, y) {
  n \leftarrow nrow(X)
  y_pred <- X %*% theta
  residual_sum_squares <- sum((y - y_pred)^2)</pre>
  log_likelihood <- - (n / 2) * log(2 * pi) - (n / 2) * log(sigma^2) -
    (1 / (2 * sigma^2)) * residual_sum_squares
  return(log_likelihood)
}
# Define Ridge function
Ridge <- function(theta, sigma, X, y, lambda) {</pre>
  log_likelihood <- Loglikelihood(theta, sigma, X, y)</pre>
  ridge_penalty <- lambda * sum(theta[-1]^2) # Exclude intercept</pre>
  return(log_likelihood - ridge_penalty)
}
# Define Ridge optimization function
RidgeOpt <- function(X, y, lambda) {</pre>
  initial_theta <- rep(0, ncol(X)) # Initialize theta to zeros</pre>
  initial_sigma <- 1
                                      # Initial quess for sigma
  initial_values <- c(initial_theta, initial_sigma)</pre>
  objective_function <- function(params) {</pre>
    theta <- params[1:(length(params) - 1)]</pre>
    sigma <- params[length(params)]</pre>
    return(-Ridge(theta, sigma, X, y, lambda))
  }
```

```
# Minimize the negative Ridge log-likelihood
  result <- optim(par = initial_values, fn = objective_function, method = "BFGS")
  optimized theta <- result$par[1:(length(result$par) - 1)]
  optimized_sigma <- result$par[length(result$par)]</pre>
  return(list(theta = optimized_theta, sigma = optimized_sigma, value = result$value))
# Function to calculate degrees of freedom for Ridge regression
DF <- function(X, lambda) {</pre>
 p <- ncol(X)
  I <- diag(p)</pre>
  XtX \leftarrow t(X) \% X
  ridge_matrix <- solve(XtX + lambda * I)</pre>
  H <- X %*% ridge_matrix %*% t(X)</pre>
  return(sum(diag(H)))
#Load and Preprocess Data
# Load dataset
file_path <- "data/parkinsons.csv"</pre>
parkinson_data <- read.csv(file_path)</pre>
# Split into training and test sets
set.seed(123)
n <- nrow(parkinson_data)</pre>
train_indices <- sample(1:n, size = 0.6 * n)</pre>
train_data <- parkinson_data[train_indices, ]</pre>
test_data <- parkinson_data[-train_indices, ]</pre>
# Scale features
train_data_scaled <- scale_features(train_data)</pre>
test_data_scaled <- scale_features(test_data)</pre>
#Train Linear Model
# Fit linear model
linear_model <- lm(motor_UPDRS ~ ., data = train_data_scaled)</pre>
# Predictions
train_predictions <- predict(linear_model, train_data_scaled)</pre>
test_predictions <- predict(linear_model, test_data_scaled)</pre>
# Calculate MSE
train_mse <- mean((train_data_scaled$motor_UPDRS - train_predictions)^2)</pre>
test_mse <- mean((test_data_scaled$motor_UPDRS - test_predictions)^2)</pre>
# Display MSE
cat("Train MSE:", train_mse, "\n")
cat("Test MSE:", test_mse, "\n")
```

```
#Ridge Regression
# Prepare data for Ridge regression
X <- as.matrix(cbind(1, train data scaled[, -which(names(train data scaled) == "motor UPDRS")]))
y <- train data scaled$motor UPDRS
# Initial log-likelihood
theta_initial <- coef(linear_model)</pre>
sigma initial <- summary(linear model)$sigma</pre>
log_likelihood_value <- Loglikelihood(theta_initial, sigma_initial, X, y)</pre>
cat("Log-Likelihood (Linear Model):", log_likelihood_value, "\n")
# Ridge optimization with lambda= 0.001
lambda <- 0.001
ridge_optimization_result <- RidgeOpt(X, y, lambda)</pre>
cat("Optimized Theta (coefficients):\n", ridge_optimization_result$theta, "\n")
cat("Optimized Sigma (residual standard error): \n", ridge_optimization_result$sigma, "\n")
cat("Minimized Negative Log-Likelihood:\n", ridge_optimization_result$value, "\n")
#Degrees of Freedom Analysis
# Degrees of Freedom for Ridge regression
lambda <- 10
degrees of freedom <- DF(X, lambda)
\#cat("Degrees of Freedom for Ridge Regression (lambda = ", lambda, "):", degrees_of_freedom, "\n")
# Compute predictions and MSE
predict_ridge <- function(X, y, theta) {</pre>
 y_pred <- X %*% theta</pre>
 mse <- mean((y - y_pred)^2)</pre>
 return(list(predictions = y_pred, mse = mse))
}
# Perform Ridge regression and compute metrics for given lambda values
#Display results
for (lambda in names(results)) {
result <- results[[lambda]]
cat("Lambda =", result$lambda, "\n")
cat("Train MSE:", result$train_mse, "\n")
cat("Test MSE:", result$test_mse, "\n")
cat("Degrees of Freedom:", result$degrees_of_freedom, "\n")
cat("Optimized Sigma:", result$optimized_sigma, "\n")
cat("\n")
}
```

Code for Assignment 3

```
set.seed(12345)
library(ggplot2)
df <- read.csv('data/pima-indians-diabetes.csv')</pre>
```



df\$diabetes

TRUE FALSE TRUE FALSE TRUE TRUE FALSE ## [1] FALSE TRUE FALSE TRUE FALSE ## TRUE TRUE TRUE TRUE FALSE TRUE FALSE FALSE TRUE [25] TRUE FALSE FALSE FALSE TRUE FALSE FALSE FALSE FALSE ## TRUE TRUE TRUE FALSE FALSE FALSE TRUE FALSE TRUE FALSE FALSE ## [37] TRUE [49] FALSE FALSE FALSE TRUE FALSE FA ## TRUE FALSE FALSE FALSE [61] TRUE FALSE FALSE TRUE FALSE TRUE FALSE TRUE FALSE FALSE FALSE FALSE TRUE ## [73] FALSE FALSE FALSE FALSE ## [85] FALSE FALSE FALSE TRUE FALSE FALSE FALSE TRUE FALSE FALSE [97] FALSE FALSE TRUE TRUE FALSE FALSE FALSE FALSE FALSE FALSE FALSE ## [109] TRUE TRUE TRUE FALSE FALSE TRUE TRUE TRUE FALSE FALSE TRUE ## [121] FALSE FALSE TRUE TRUE FALSE FALSE TRUE TRUE TRUE TRUE ## [133] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE TRUE FALSE ## [145] FALSE FALSE FALSE FALSE FALSE FALSE TRUE FALSE TRUE TRUE FALSE ## [157] FALSE FALSE TRUE FALSE FALSE FALSE FALSE TRUE TRUE FALSE FALSE FALSE ## [169] FALSE TRUE TRUE FALSE FALSE TRUE FALSE TRUE FALSE TRUE FALSE ## [181] FALSE FALSE FALSE TRUE TRUE TRUE TRUE TRUE FALSE FALSE TRUE ## [193] TRUE FALSE TRUE FALSE TRUE TRUE TRUE FALSE FALSE FALSE FALSE ## [205] FALSE TRUE TRUE FALSE TRUE FALSE FALSE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE FALSE FALSE FALSE FALSE TRUE FALSE ## [217] FALSE ## [229] FALSE TRUE TRUE FALSE FALSE TRUE TRUE TRUE TRUE TRUE FALSE FALSE ## [241] FALSE TRUE TRUE FALSE TRUE FALSE FALSE FALSE FALSE FALSE FALSE ## [253] FALSE TRUE TRUE FALSE FALSE TRUE FALSE TRUE FALSE TRUE ## [265] FALSE TRUE FALSE FALSE TRUE TRUE FALSE FALSE FALSE FALSE TRUE ## [277] FALSE FALSE TRUE FALSE FALSE TRUE TRUE FALSE FALSE TRUE FALSE ## [289] FALSE FALSE TRUE TRUE TRUE FALSE FALSE TRUE FALSE TRUE FALSE TRUE ## [301] TRUE FALSE TRUE FALSE TRUE FALSE TRUE TRUE FALSE FALSE TRUE ## [313] FALSE TRUE FALSE TRUE FALSE TRUE FALSE TRUE TRUE TRUE TRUE FALSE ## [325] FALSE TRUE FALSE TRUE FALSE FALSE FALSE FALSE FALSE FALSE FALSE ## [337] TRUE TRUE TRUE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE ## [349] TRUE FALSE FALSE FALSE FALSE TRUE TRUE TRUE TRUE TRUE TRUE ## [361] FALSE FALSE TRUE FALSE FALSE TRUE FALSE FALSE TRUE TRUE FALSE FALSE ## [373] FALSE FALSE TRUE FALSE FALSE TRUE FALSE FALSE FALSE FALSE FALSE ## [385] FALSE TRUE TRUE TRUE FALSE FALSE TRUE FALSE FALSE TRUE FALSE FALSE ## [397] TRUE FALSE TRUE TRUE FALSE TRUE FALSE TRUE FALSE TRUE ## [409] TRUE FALSE FALSE FALSE TRUE TRUE FALSE TRUE FALSE TRUE FALSE ## [421] FALSE FALSE FALSE TRUE TRUE FALSE TRUE FALSE TRUE FALSE FALSE ## [433] FALSE FALSE TRUE FALSE FALSE FALSE TRUE FALSE TRUE TRUE TRUE ## [445] TRUE FALSE FALSE TRUE FALSE FALSE TRUE FALSE FALSE TRUE FALSE ## [457] FALSE TRUE FALSE FALSE FALSE FALSE FALSE FALSE FALSE TRUE ## [469] FALSE FALSE FALSE FALSE FALSE FALSE FALSE TRUE FALSE FALSE TRUE ## [481] FALSE FALSE FALSE TRUE TRUE FALSE FALSE FALSE FALSE FALSE FALSE ## [493] TRUE FALSE FALSE FALSE TRUE FALSE FALSE FALSE TRUE FALSE FALSE ## [505] FALSE TRUE FALSE FALSE TRUE FALSE FALSE FALSE TRUE TRUE ## [517] FALSE FALSE FALSE FALSE FALSE TRUE FALSE FALSE FALSE FALSE FALSE ## [529] FALSE FALSE FALSE FALSE FALSE TRUE FALSE FALSE TRUE TRUE ## [541] TRUE TRUE FALSE FALSE TRUE TRUE FALSE FALSE FALSE FALSE FALSE ## [553] FALSE FALSE FALSE FALSE FALSE FALSE TRUE TRUE FALSE FALSE FALSE [565] FALSE FALSE FALSE TRUE FALSE FALSE FALSE FALSE FALSE FALSE [577] TRUE FALSE TRUE TRUE FALSE FALSE FALSE TRUE FALSE TRUE FALSE TRUE [589] FALSE TRUE FALSE TRUE FALSE TRUE FALSE FALSE TRUE FALSE FALSE ## [601] FALSE FALSE TRUE TRUE FALSE TRUE FALSE FALSE FALSE TRUE TRUE ## [613] FALSE TRUE FALSE FALSE TRUE TRUE FALSE FALSE FALSE FALSE ## [625] FALSE FALSE FALSE FALSE TRUE FALSE FALSE FALSE TRUE FALSE ## [637] FALSE TRUE FALSE FALSE TRUE FALSE FALSE FALSE TRUE TRUE TRUE ## [649] FALSE FALSE FALSE FALSE FALSE TRUE FALSE FALSE TRUE FALSE [661] TRUE TRUE TRUE TRUE FALSE TRUE TRUE FALSE FALSE FALSE FALSE [673] FALSE FALSE TRUE TRUE FALSE TRUE FALSE TRUE FALSE TRUE FALSE ## [685] FALSE FALSE FALSE TRUE FALSE TRUE FALSE TRUE FALSE TRUE TRUE ## [697] FALSE FALSE FALSE TRUE TRUE FALSE FALSE TRUE FALSE TRUE ## [709] TRUE FALSE FALSE TRUE FALSE FALSE TRUE TRUE FALSE FALSE TRUE FALSE ## [721] FALSE TRUE FALSE FALSE FALSE FALSE FALSE FALSE TRUE TRUE TRUE ## [733] FALSE FALSE FALSE FALSE FALSE TRUE TRUE FALSE FALSE TRUE FALSE ## [745] FALSE TRUE FALSE TRUE TRUE TRUE FALSE FALSE TRUE TRUE TRUE FALSE

```
## [757] TRUE FALSE TRUE FALSE TRUE FALSE FALSE FALSE TRUE FALSE
m = glm(diabetes ~ age + plasma, data=df, family="binomial")
df$predicted_prob <- predict(m, newdata = df, type = "response")</pre>
classification_threshold <- 0.5</pre>
df$prediction <- df$predicted_prob > classification_threshold
missclassification rate <- sum(df$diabetes != df$prediction) / nrow(df)
\# 0 = k1 + k2 * age + k3 * plasma
k1 <- summary(m)$coefficients[1]</pre>
k2 <- summary(m)$coefficients[2]</pre>
k3 <- summary(m)$coefficients[3]
\# plasma = (-k1 - k2 * age) / k3
plasma_function <- function(x)(-k1 -k2 * x) / k3</pre>
df |> ggplot() +
      geom_point(aes(x = age, y = plasma, color = prediction)) +
      geom_function(fun = plasma_function) +
      labs(x = "Age", y = "Plasma glucose concentration", color = "Predicted to have diabetes?")
   200
Plasma glucose concentration
   150
                                                                  Predicted to have diabetes?
   100 -
                                                                      FALSE
                                                                      TRUE
    50 -
     0 -
       20
                        40
                                         60
                                                          80
                                Age
p < -0.2
df$prediction02 <- df$predicted_prob > p
k4 \leftarrow log (p / (1-p))
plasma_function <- function(x)( k4 -k1 -k2 * x) / k3</pre>
```

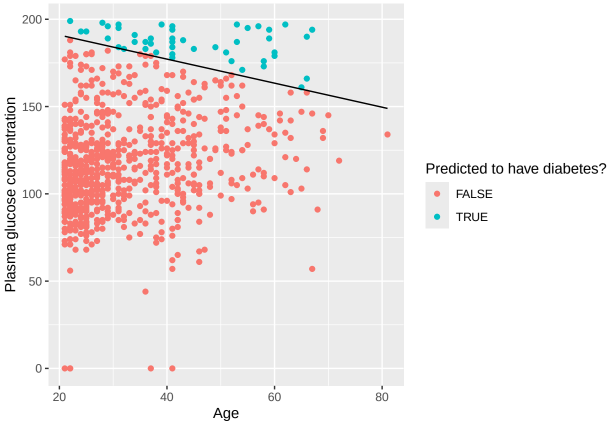
```
df |> ggplot() +
      geom_point(aes(x = age, y = plasma, color = prediction02)) +
      geom_function(fun = plasma_function) +
      labs(x = "Age", y = "Plasma glucose concentration", color = "Predicted to have diabetes?")
   200 -
Plasma glucose concentration
   150 -
                                                                   Predicted to have diabetes?
   100 -
                                                                       FALSE
                                                                       TRUE
    50 -
     0 -
       20
                         40
                                         60
                                                           80
                                 Age
p < -0.8
df$prediction08 <- df$predicted_prob > p
k4 < -\log (p / (1-p))
plasma_function <- function(x)( k4 - k1 - k2 * x) / k3
```

labs(x = "Age", y = "Plasma glucose concentration", color = "Predicted to have diabetes?")

geom_point(aes(x = age, y = plasma, color = prediction08)) +

geom_function(fun = plasma_function) +

df |> ggplot() +



```
df$z1 <- df$plasma ^ 4
df$z2 <- df$plasma ^ 3 * df$age
df$z3 <- df$plasma ^ 2 * df$age ^ 2</pre>
df$z4 <- df$plasma * df$age ^ 3</pre>
df$z5 <- df$age ^ 4
m2 = glm(diabetes \sim age + plasma + z1 + z2 + z3 + z4 + z5,
          data=df, family="binomial")
df$predicted_prob2 <- predict(m2, newdata = df, type = "response")</pre>
classification_threshold <- 0.5</pre>
df$prediction2 <- df$predicted_prob2 > classification_threshold
\# 0 = k1 + k2 * age + k3 * plasma
k1 <- summary(m)$coefficients[1]</pre>
k2 <- summary(m)$coefficients[2]</pre>
k3 <- summary(m)$coefficients[3]
\# plasma = (-k1 - k2 * age) / k3
plasma_function <- function(x)(-k1 -k2 * x) / k3</pre>
df |> ggplot() +
      geom_point(aes(x = age, y = plasma, color = prediction)) +
      geom_function(fun = plasma_function) +
      labs(x = "Age", y = "Plasma glucose concentration", color = "Predicted to have diabetes?")
```

