DATA621Assignment2

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Load the Data

Download the classification output data set (attached in Blackboard to the assignment)

classification_df <- read.csv("https://raw.githubusercontent.com/uzmabb182/Data_621/refs/heads/main/Ass
head(classification_df)</pre>

##		pregnant	glucose	diastolic	${\tt skinfold}$	${\tt insulin}$	bmi	pedigree	age	class
##	1	7	124	70	33	215	25.5	0.161	37	0
##	2	2	122	76	27	200	35.9	0.483	26	0
##	3	3	107	62	13	48	22.9	0.678	23	1
##	4	1	91	64	24	0	29.2	0.192	21	0
##	5	4	83	86	19	0	29.3	0.317	34	0
##	6	1	100	74	12	46	19.5	0.149	28	0
##		scored.cl	ass scor	red.probabi	llity					
##	1		0	0.3284	15226					
##	2		0	0.2731	19044					
##	3		0	0.1096	6039					
##	4		0	0.0559	99835					
##	5		0	0.1004	19072					
##	6		0	0.0551	15460					

Confusion Matrix

The data set has three key columns we will use:

- class: the actual class for the observation
- scored.class: the predicted class for the observation (based on a threshold of 0.5)
- scored.probability: the predicted probability of success for the observation

Use the table() function to get the raw confusion matrix for this scored dataset. Make sure you understand the output. In particular, do the rows represent the actual or predicted class? The columns?

Confusion Matrix: The table() function in R will give you the raw confusion matrix. In this table:

- The rows generally represent the actual class
- The columns generally represent the predicted class

For example: Predicted Positive Predicted Negative Actual Positive True Positive (TP) False Negative (FN) Actual Negative False Positive (FP) True Negative (TN)

```
confusion_matrix <- table(classification_df$class, classification_df$scored.class)
confusion_matrix</pre>
```

This can be interpreted as:

- True Negatives (TN): 119 These are the instances where the actual class was 0, and the predicted class was also 0
- False Positives (FP): 5 These are the instances where the actual class was 0, but the predicted class was 1
- False Negatives (FN): 30 These are the instances where the actual class was 1, but the predicted class was 0
- True Positives (TP): 27 These are the instances where the actual class was 1, and the predicted class was also 1

Accuracy

Write a function that takes the data set as a dataframe, with actual and predicted classifications identified, and returns the accuracy of the predictions.

Accuracy: Represents the proportion of correct predictions. For instance, if accuracy is 90%, it means the model correctly classified 90% of the observations as either success or failure.

```
Formula Accuracy = (TP + TN) / (TP + FP + TN + FN)
```

```
calculate_accuracy <- function(data) {
  confusion_matrix <- table(data$class, data$scored.class)
  TP <- confusion_matrix[2, 2]
  TN <- confusion_matrix[1, 1]
  FP <- confusion_matrix[1, 2]
  FN <- confusion_matrix[2, 1]
  accuracy <- (TP + TN) / (TP + FP + TN + FN)
  return(accuracy)
}</pre>
```

Classification Error

Write a function that takes the data set as a dataframe, with actual and predicted classifications identified, and returns the classification error rate of the predictions.

Error Rate: Complements accuracy and indicates the proportion of incorrect predictions. If it's 10%, it tells us the model misclassified 10% of observations.

Formula Classification Error Rate == (FP + FN) / (TP + FP + TN + FN)

```
calculate_error_rate <- function(data) {
  confusion_matrix <- table(data$class, data$scored.class)
  FP <- confusion_matrix[1, 2]
  FN <- confusion_matrix[2, 1]
  TP <- confusion_matrix[2, 2]
  TN <- confusion_matrix[1, 1]
  error_rate <- (FP + FN) / (TP + FP + TN + FN)
  return(error_rate)
}</pre>
```

Precision

Write a function that takes the data set as a dataframe, with actual and predicted classifications identified, and returns the precision of the predictions.

Precision: Focuses on the quality of positive predictions. For example, in a healthcare setting, a high precision rate means most patients predicted to have a condition actually do have it, reducing the risk of unnecessary interventions.

```
Formula Precision = TP / (TP + FP)
```

```
calculate_precision <- function(data) {
  confusion_matrix <- table(data$class, data$scored.class)
  TP <- confusion_matrix[2, 2]
  FP <- confusion_matrix[1, 2]
  precision <- TP / (TP + FP)
  return(precision)
}</pre>
```

Sensitivity (Recall / True Positive Rate)

Write a function that takes the data set as a dataframe, with actual and predicted classifications identified, and returns the sensitivity of the predictions. Sensitivity is also known as recall.

Sensitivity measures the ability of the model to correctly identify actual positives. This tells us how well the model is capturing positive cases. For example, if this is a medical dataset, sensitivity indicates the percentage of patients correctly identified as having the condition.

Formula Sensitivity = TP / (TP + FN)

```
calculate_sensitivity <- function(data) {
  confusion_matrix <- table(data$class, data$scored.class)
  TP <- confusion_matrix[2, 2]
  FN <- confusion_matrix[2, 1]
  sensitivity <- TP / (TP + FN)
  return(sensitivity)
}</pre>
```

Specificity (True Negative Rate)

Write a function that takes the data set as a dataframe, with actual and predicted classifications identified, and returns the specificity of the predictions.

Specificity measures the ability of the model to correctly identify actual negatives. In the context of the data, specificity shows how well the model avoids falsely predicting positives for negative cases.

Formula Specificity = TN / (TN + FP)

```
calculate_specificity <- function(data) {
  confusion_matrix <- table(data$class, data$scored.class)
  TN <- confusion_matrix[1, 1]
  FP <- confusion_matrix[1, 2]
  specificity <- TN / (TN + FP)
  return(specificity)
}</pre>
```

F1 Score

Write a function that takes the data set as a dataframe, with actual and predicted classifications identified, and returns the F1 score of the predictions

The F1 score is the harmonic mean of Precision and Sensitivity. It balances the trade-off between precision (quality of positive predictions) and sensitivity (coverage of actual positives). This metric is useful when there is an imbalance between positive and negative classes in the dataset.

Formula F1 Score = 2 * (precision * sensitivity) / (precision + sensitivity)

```
calculate_f1_score <- function(data) {
  confusion_matrix <- table(data$class, data$scored.class)
  TP <- confusion_matrix[2, 2]
  FP <- confusion_matrix[1, 2]
  FN <- confusion_matrix[2, 1]

  precision <- TP / (TP + FP)
  sensitivity <- TP / (TP + FN)

  f1_score <- 2 * (precision * sensitivity) / (precision + sensitivity)
  return(f1_score)
}</pre>
```

What are the bounds on the F1 score?

Before we move on, let's consider a question that was asked: What are the bounds on the F1 score? Show that the F1 score will always be between 0 and 1. (Hint: If 0 < a < 1 and 0 < b < 1 then ab < a.)

F1 Score is always between 0 and 1

- 1. Both Precision and Sensitivity are proportions, and thus range between 0 and 1
- 2. When multiplying two values between 0 and 1, the result is also between 0 and 1. For example, If 0 < a < 1 and 0 < b < 1 then a b < a and a b < b
- 3. Dividing a number between 0 and 1 (numerator) by another number greater than it (denominator) ensures the result remains between 0 and 1.

Thus, since Precision and Sensitivity are bounded by 0 and 1, their harmonic mean (F1 Score) must also be within this range.

Function that generates an ROC curve

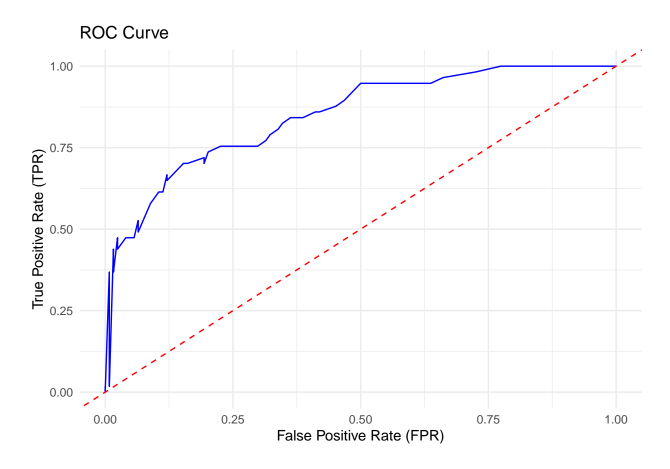
Write a function that generates an ROC curve from a data set with a true classification column (class in our example) and a probability column (scored.probability in our example). Your function should return a list that includes the plot of the ROC curve and a vector that contains the calculated area under the curve (AUC). Note that I recommend using a sequence of thresholds ranging from 0 to 1 at 0.01 intervals.

```
generate_roc <- function(data) {</pre>
  true class <- data$class
  predicted_probabilities <- data$scored.probability</pre>
  thresholds \leftarrow seq(0, 1, by = 0.01)
  TPR <- numeric(length(thresholds))</pre>
  FPR <- numeric(length(thresholds))</pre>
  for (i in seq_along(thresholds)) {
    threshold <- thresholds[i]</pre>
    predicted_class <- ifelse(predicted_probabilities >= threshold, 1, 0)
    TP <- sum(predicted_class == 1 & true_class == 1)</pre>
    TN <- sum(predicted class == 0 & true class == 0)
    FP <- sum(predicted_class == 1 & true_class == 0)</pre>
    FN <- sum(predicted_class == 0 & true_class == 1)
    TPR[i] <- TP / (TP + FN) # Sensitivity</pre>
    FPR[i] <- FP / (FP + TN) # 1 - Specificity</pre>
  auc <- sum(diff(FPR) * (TPR[-1] + TPR[-length(TPR)]) / 2)</pre>
  auc <- abs(auc)
  roc_data <- data.frame(FPR = FPR, TPR = TPR)</pre>
  roc_plot <- ggplot(roc_data, aes(x = FPR, y = TPR)) +</pre>
    geom_line(color = "blue") +
    geom abline(slope = 1, intercept = 0, linetype = "dashed", color = "red") +
    labs(title = "ROC Curve", x = "False Positive Rate (FPR)",
         y = "True Positive Rate (TPR)") +
         theme_minimal()
  return(list(plot = roc_plot, auc = auc))
```

Use your created R functions and the provided classification output data set to produce all of the classification metrics discussed above

Consider the functions confusionMatrix, sensitivity, and specificity

```
print(paste("Confusion Matrix: "))
## [1] "Confusion Matrix: "
confusion_matrix
##
##
##
     0 119
     1 30
           27
print(paste("Accuracy: ", calculate_accuracy(classification_df)))
## [1] "Accuracy: 0.806629834254144"
print(paste("Error rate: ", calculate_error_rate(classification_df)))
## [1] "Error rate: 0.193370165745856"
print(paste("Precision: ", calculate_precision(classification_df)))
## [1] "Precision: 0.84375"
print(paste("Sensitivity: ", calculate_sensitivity(classification_df)))
## [1] "Sensitivity: 0.473684210526316"
print(paste("Specifivity: ", calculate_specificity(classification_df)))
## [1] "Specifivity: 0.959677419354839"
print(paste("Classification: ", calculate_f1_score(classification_df)))
## [1] "Classification: 0.606741573033708"
roc_gen <- generate_roc(classification_df)</pre>
print(roc_gen$plot)
```



print(paste("AUC:", roc_gen\$auc))

[1] "AUC: 0.848896434634975"

Investigate the caret package

Investigate the caret package. In particular, consider the functions confusionMatrix, sensitivity, and specificity. Apply the functions to the data set. How do the results compare with your own functions?

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
              0
                    1
            0 119
                   30
##
##
            1
              5 27
##
                  Accuracy : 0.8066
##
                    95% CI: (0.7415, 0.8615)
##
       No Information Rate: 0.6851
##
       P-Value [Acc > NIR] : 0.0001712
##
##
##
                     Kappa: 0.4916
##
##
   Mcnemar's Test P-Value: 4.976e-05
##
##
               Sensitivity: 0.4737
##
               Specificity: 0.9597
            Pos Pred Value: 0.8438
##
##
            Neg Pred Value: 0.7987
##
                Prevalence: 0.3149
##
            Detection Rate: 0.1492
      Detection Prevalence: 0.1768
##
         Balanced Accuracy: 0.7167
##
##
          'Positive' Class : 1
##
##
```

The results from using the caret function compared to our own function are nearly identical.

Investigate the pROC package

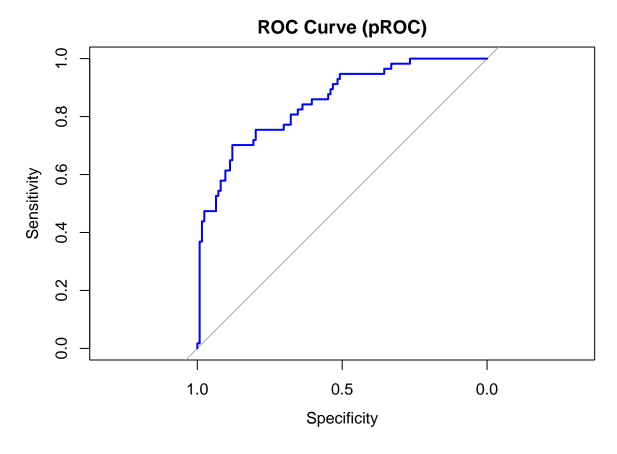
Investigate the pROC package. Use it to generate an ROC curve for the data set. How do the results compare with your own functions?

```
roc_result <- roc(classification_df$class, classification_df$scored.probability)

## Setting levels: control = 0, case = 1

## Setting direction: controls < cases

plot(roc_result, col = "blue", main = "ROC Curve (pROC)")</pre>
```



```
auc_value <- auc(roc_result)
print(paste("AUC (pROC):", auc_value))</pre>
```

```
## [1] "AUC (pROC): 0.850311262026033"
```

The curve produced by pROC is nearly identical to the one generated by the manual function, and the AUC values were nearly the same. While writing the function by hand was valuable for learning the underlying mechanics of ROC analysis, using pROC is clearly more efficient and practical for real-world work.

Fit Logistic Regression Model

```
classification_df$class <- as.factor(classification_df$class)
model <- glm(class ~ ., data = classification_df, family = binomial)
classification_df$predicted_prob <- predict(model, classification_df, type = "response")
classification_df$predicted_class <- ifelse(classification_df$predicted_prob > 0.5, 1, 0)
```

Generate and Plot the ROC Curve

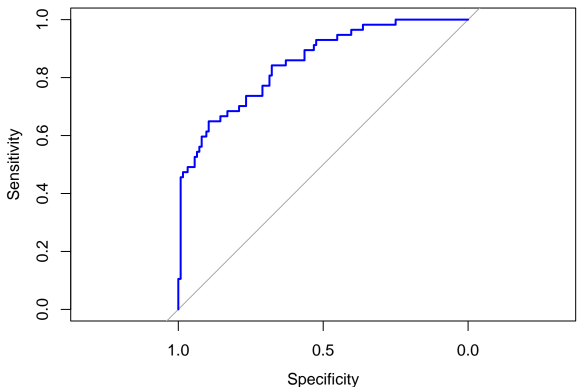
```
roc_curve <- roc(classification_df$class, classification_df$predicted_prob)

## Setting levels: control = 0, case = 1

## Setting direction: controls < cases

plot(roc_curve, col = "blue", main = "ROC Curve for Logistic Regression")</pre>
```



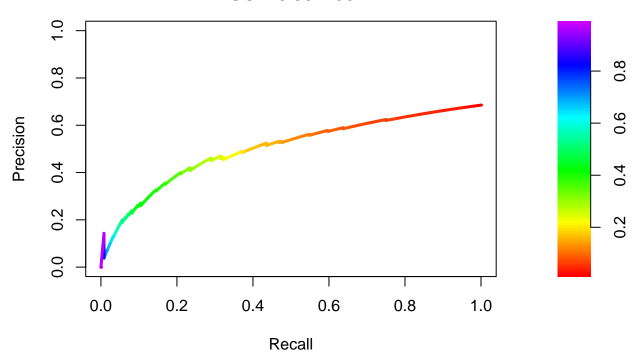


```
auc_value <- auc(roc_curve)
print(paste("AUC Score:", auc_value))</pre>
```

[1] "AUC Score: 0.851018675721562"

Generate and Plot the Precision-Recall Curve

Precision–Recall Curve AUC = 0.5014507



Generate a Confusion Matrix Heatmap

```
generate_confusion_matrix_heatmap <- function(data) {
  confusion_matrix <- table(data$class, data$scored.class)
  library(reshape2)
  confusion_df <- melt(confusion_matrix)
  ggplot(confusion_df, aes(x = Var2, y = Var1, fill = value)) +
    geom_tile() +
    geom_text(aes(label = value), color = "white") +
    scale_fill_gradient(low = "blue", high = "red") +
    labs(title = "Confusion Matrix Heatmap", x = "Predicted", y = "Actual") +
    theme_minimal()</pre>
```

```
}
generate_confusion_matrix_heatmap(classification_df)
##
## Attaching package: 'reshape2'
## The following object is masked from 'package:tidyr':
##
##
       smiths
        Confusion Matrix Heatmap
    1.5
    1.0
                          30
                                                           27
                                                                                  value
                                                                                       90
    0.5
                                                                                       60
```

Histogram of Predicted Probabilities

119

0.0

0.0

-0.5

-0.5

```
ggplot(classification_df, aes(x = predicted_prob, fill = as.factor(class))) +
  geom_histogram(binwidth = 0.1, alpha = 0.7, position = "identity") +
  scale_fill_manual(values = c("red", "green"), labels = c("Negative", "Positive")) +
  labs(title = "Histogram of Predicted Probabilities", x = "Predicted Probability", fill = "Class")
```

0.5

Predicted

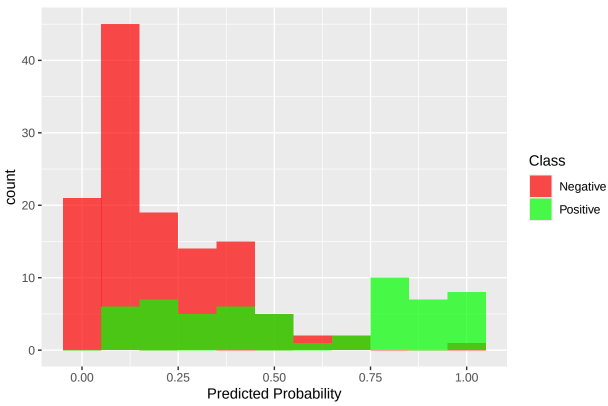
5

1.0

30

1.5





Summary:

ROC Curve (AUC Score) Measures how well the model separates classes. Precision-Recall Curve Evaluates performance on imbalanced datasets. Confusion Matrix Heatmap Visualizes classification errors and correct predictions. Histogram of predicted probabilities This gives insights into how confident the model is about its predictions.