

FACULTY OF COMPUTER SCIENCE & INFORMATION TECHNOLOGY

WQD 7004 - Programming for Data Science

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Group Project

Project Title: Prediction of Thyroid Hormone Levels and Diagnosis of Thyroid Disorders

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Contents

Introduction		
Project Objectives	9	
Project Questions	9	
Step 1: Preliminaries	4	
Step 2: Data Preprocessing	6	
Step 3: Exploratory Data Analysis (EDA)	11	
Step 4: Model Training	14	
Regression	14	
Classification	15	
Step 5: Model Evaluation	17	
Regression	17	
Classification	19	
Step 6: Interpretation of Results	21	
Result Interretation - Regression	21	
Result Interpretaion - Classification	21	
References		

Introduction

Thyroid disorders, including hypothyroidism and hyperthyrodism, are prevalent endocrine conditions that can significantly impact metabolic health. Globally, the incidence of thyroid dysfunction is increasing, with studies indicating that thyroid disorders affect a substantial portion of the population [1]. Thyroid-stimulating hormone (TSH), produced by the pituitary gland, plays a pivotal role in regulating thyroid function and is a primary marker for assessing thyroid health [2].

In this project, we are utilizing the Hyperthyroid dataset, originally sourced from the UCI Machine Learning Repository and made available via Kaggle [3], to develop **predictive models** with two primary objectives. First, we are constructing a regression model to **estimate TSH levels** based on **clinical and demographic features**. Second, we are developing a **binary classification model** to determine whether an individual exhibits signs of thyroid dysfunction. By integrating both regression and classification, this projects aims to **enhance early diagnosis** and **support effective clinical decision-making** for thyroid health management.

Project Objectives

- 1. To build a regression model that estimates thyroid-stimulating hormone (TSH) levels based on clinical and demographic features.
- 2. To develop a binary classification model that distinguishes between individuals with thyroid dysfunction and those with normal thyroid function.
- 3. To evaluate the effectiveness of both models in providing accurate predictions that can aid in medical decision-making and early intervention.

Project Questions

- 1. Can we accurately predict an individual's TSH (thyroid-stimulating hormone) level based on clinical and demographic data?
- 2. Can we determine whether an individual has thyroid dysfunction (e.g., hypothyroidism) or is in a normal state using available features in the dataset?

Step 1: Preliminaries

In this preliminary step, we set up the working environment by loading the necessary R packages that will be used throughout the project and importing the thyroid dataset.

This includes loading essential libraries for data manipulation and visualization, and reading the dataset into a data frame. We are also previewing the first few rows to confirm that the data has been successfully imported and to understand its structure before proceeding with data preprocessing.

```
# Load required libraries

# Data Manipulation and Visualization
library(tidyverse)

# Plotting
library(ggplot2)

# Data Wrangling
library(dplyr)

# Correlation Plots
library(corrplot)
```

```
# Load thyroid data
thyroid_data_df <- read.csv("../data/hypothyroid.csv")</pre>
# Preview the dataset
head(thyroid_data_df)
     age sex on.thyroxine query.on.thyroxine on.antithyroid.medication sick
## 1
           F
                         f
                                              f
                                                                          f
                                                                               f
## 2
      23
           F
                         f
                                              f
                                                                          f
                                                                               f
## 3
                         f
                                              f
                                                                          f
                                                                               f
      46
           М
           F
                                              f
## 4
      70
                                                                          f
                                                                               f
                          t
```

```
## 5
      70
           F
                          f
                                              f
                                                                           f
                                                                                f
## 6
           F
     pregnant thyroid.surgery I131.treatment query.hypothyroid query.hyperthyroid
##
## 1
             f
                              f
                                              f
                                                                  f
## 2
             f
                              f
                                              f
                                                                  f
                                                                                       f
## 3
             f
                              f
                                              f
                                                                  f
                                                                                       f
## 4
             f
                              f
                                              f
                                                                  f
                                                                                       f
## 5
             f
                              f
                                               f
                                                                  f
                                                                                       f
## 6
             f
                              f
                                               f
                                                                  f
                                                                                       f
     lithium goitre tumor hypopituitary psych TSH.measured TSH T3.measured T3
                                                              t 1.3
## 1
                   f
                          f
                                         f
                                                f
                                                                                t 2.5
## 2
           f
                   f
                                                              t 4.1
                          f
                                         f
                                                f
## 3
           f
                   f
                          f
                                         f
                                                f
                                                              t 0.98
                                                                                f
                                                                                     ?
                          f
## 4
           f
                   f
                                         f
                                                f
                                                              t 0.16
                                                                                t 1.9
## 5
           f
                                                f
                   f
                                                              t 0.72
                                                                                t 1.2
## 6
           f
                   f
                          f
                                         f
                                                f
                                                              t 0.03
     TT4.measured TT4 T4U.measured T4U FTI.measured FTI TBG.measured TBG
##
## 1
                 t 125
                                    t 1.14
                                                       t 109
## 2
                 t 102
                                                            ?
                                                                          f
                                                                              ?
                                    f
                                                       f
                                                                              ?
## 3
                 t 109
                                    t 0.91
                                                       t 120
                                                                          f
## 4
                 t 175
                                    f
                                         ?
                                                       f
                                                            ?
                                                                          f
                                                                              ?
## 5
                 t 61
                                    t 0.87
                                                          70
                                                                          f
                                                                              ?
                 t 183
                                    t 1.3
                                                                          f
                                                                              ?
## 6
                                                       t 141
     referral.source binaryClass
## 1
                 SVHC
## 2
                other
## 3
                other
                                 P
## 4
                other
                                 Р
                  SVI
## 5
                                 Ρ
## 6
                other
                                 Р
```

Step 2: Data Preprocessing

In this data preprocessing step, we clean and prepare the thyroid dataset for analysis.

This includes handling missing values by replacing "?" with NA, removing duplicate and irrelevant columns, converting character columns to numeric types, and filtering out outliers (e.g., age >= 100).

We are also imputing missing numeric values using the column mean and encode categorical variables (e.g., sex, binaryClass) into numerical format to prepare the data for model training.

The thyroid dataset has 3772 rows and 30 columns.

```
# List all column names in the dataset
for (idx in seq_along(thyroid_data_df)) {
  cat(idx, ":", colnames(thyroid_data_df)[idx], "\n")
}
```

```
## 1 : age
## 2 : sex
## 3 : on.thyroxine
## 4 : query.on.thyroxine
## 5 : on.antithyroid.medication
## 6 : sick
## 7 : pregnant
## 8 : thyroid.surgery
## 9 : I131.treatment
## 10 : query.hypothyroid
## 11 : query.hyperthyroid
## 12 : lithium
## 13 : goitre
## 14 : tumor
## 15 : hypopituitary
## 16 : psych
## 17 : TSH.measured
## 18 : TSH
## 19 : T3.measured
## 20 : T3
## 21 : TT4.measured
## 22 : TT4
## 23 : T4U.measured
## 24 : T4U
## 25 : FTI.measured
## 26 : FTI
## 27 : TBG.measured
```

28 : TBG

29 : referral.source
30 : binaryClass

Statistical summary for each column in the dataset summary(thyroid_data_df)

```
##
                                            on.thyroxine
                                                                 query.on.thyroxine
        age
                            sex
##
    Length: 3772
                        Length: 3772
                                            Length: 3772
                                                                 Length: 3772
##
    Class : character
                        Class : character
                                            Class : character
                                                                 Class : character
    Mode :character
                        Mode : character
                                            Mode :character
                                                                 Mode : character
##
    on.antithyroid.medication
                                    sick
                                                      pregnant
  Length:3772
##
                               Length: 3772
                                                    Length: 3772
   Class : character
##
                               Class :character
                                                    Class : character
   Mode :character
                               Mode :character
##
                                                    Mode :character
   thyroid.surgery
                        I131.treatment
                                            query.hypothyroid query.hyperthyroid
##
    Length: 3772
                        Length: 3772
                                            Length: 3772
                                                                 Length: 3772
   Class : character
                        Class : character
                                            Class : character
                                                                 Class : character
##
   Mode :character
                        Mode :character
                                            Mode :character
                                                                Mode :character
##
      lithium
                           goitre
##
                                               tumor
                                                                hypopituitary
    Length: 3772
                        Length: 3772
                                            Length: 3772
                                                                Length: 3772
##
    Class : character
                        Class : character
                                            Class : character
                                                                 Class : character
    Mode :character
                        Mode : character
                                            Mode : character
                                                                Mode :character
##
                        TSH.measured
                                                                T3.measured
##
       psych
                                                TSH
##
    Length: 3772
                        Length: 3772
                                            Length: 3772
                                                                Length: 3772
    Class : character
                        Class : character
                                            Class : character
                                                                 Class : character
##
##
    Mode :character
                        Mode : character
                                            Mode : character
                                                                 Mode : character
                        TT4.measured
                                                TT4
                                                                 T4U.measured
##
         Т3
    Length: 3772
                        Length: 3772
                                            Length: 3772
                                                                Length: 3772
##
##
    Class : character
                        Class : character
                                            Class : character
                                                                 Class : character
    Mode : character
                        Mode : character
                                            Mode : character
                                                                 Mode : character
##
##
        T4U
                        FTI.measured
                                                FTI
                                                                 TBG.measured
    Length: 3772
                        Length: 3772
                                            Length: 3772
                                                                Length: 3772
##
##
    Class : character
                        Class : character
                                            Class : character
                                                                 Class : character
##
    Mode :character
                        Mode : character
                                            Mode : character
                                                                 Mode : character
##
        TBG
                        referral.source
                                            binaryClass
##
    Length: 3772
                        Length: 3772
                                            Length: 3772
   Class : character
                        Class : character
                                            Class : character
##
    Mode :character
                        Mode :character
                                            Mode : character
```

Structure of the dataset

str(thyroid_data_df)

```
: chr "f" "f" "f" "f" ...
## $ query.on.thyroxine
## $ on.antithyroid.medication: chr "f" "f" "f" "f"
                                     "f" "f" "f" "f"
## $ sick
                               : chr
## $ pregnant
                               : chr
                                      "f" "f" "f" "f" ...
                                     "f" "f" "f" "f" ...
## $ thyroid.surgery
                              : chr
## $ I131.treatment
                                      "f" "f" "f" "f"
                              : chr
## $ query.hypothyroid
                                      "f" "f" "f" "f" ...
                              : chr
                               : chr
## $ query.hyperthyroid
                                      "f" "f" "f" "f" ...
                                     "f" "f" "f" "f" ...
## $ lithium
                               : chr
## $ goitre
                                      "f" "f" "f" "f" ...
                               : chr
                                      "f" "f" "f" "f" ...
## $ tumor
                               : chr
## $ hypopituitary
                                      "f" "f" "f" "f" ...
                               : chr
## $ psych
                              : chr
                                      "f" "f" "f" "f" ...
## $ TSH.measured
                               : chr
                                      "t" "t" "t" "t" ...
## $ TSH
                                      "1.3" "4.1" "0.98" "0.16" ...
                              : chr
## $ T3.measured
                               : chr
                                      "t" "t" "f" "t" ...
## $ T3
                                      "2.5" "2" "?" "1.9" ...
                              : chr
## $ TT4.measured
                              : chr
                                      "t" "t" "t" "t" ...
## $ TT4
                                      "125" "102" "109" "175" ...
                              : chr
                                      "t" "f" "t" "f" ...
## $ T4U.measured
                              : chr
## $ T4U
                                      "1.14" "?" "0.91" "?" ...
                               : chr
## $ FTI.measured
                              : chr
                                      "t" "f" "t" "f" ...
                              : chr
                                     "109" "?" "120" "?" ...
## $ FTI
## $ TBG.measured
                                      "f" "f" "f" "f" ...
                               : chr
                               : chr "?" "?" "?" "?" ...
## $ TBG
## $ referral.source
                               : chr
                                     "SVHC" "other" "other" "other" ...
## $ binaryClass
                               : chr "P" "P" "P" "P" ...
# Check for duplicate rows
duplicate_values <- sum(duplicated(thyroid_data_df))</pre>
cat("\nDuplicate values in the dataset: \n")
print(duplicate_values)
# Remove duplicate rows
thyroid_data_df <- thyroid_data_df[!duplicated(thyroid_data_df), ]</pre>
cat("\nDuplicate values after removal: \n")
print(sum(duplicated(thyroid_data_df)))
thyroid_data_df
# Loop through all columns and print value counts
for (colname in names(thyroid_data_df)) {
  cat("\nColumn:", colname, "\n")
  print(table(thyroid_data_df[[colname]], useNA = "ifany"))
}
```

```
# Removing irrelevant columns
thyroid_data_df$TBG <- NULL</pre>
thyroid_data_df$referral.source <- NULL</pre>
for (idx in seq along(thyroid data df)) {
  cat(idx, ":", colnames(thyroid_data_df)[idx], "\n")
# Replace data points '?' with NA
thyroid_data_df[thyroid_data_df == "?"] <- NA</pre>
na counts <- data.frame(</pre>
  Column = names(thyroid_data_df),
 Missing_Values = colSums(is.na(thyroid_data_df)),
  row.names = NULL
print(na_counts)
# Data encoding
thyroid_data_df$binaryClass <- ifelse(thyroid_data_df$binaryClass == "P", 0,
                                 ifelse(thyroid_data_df$binaryClass == "N", 1, NA))
thyroid_data_df$sex <- ifelse(thyroid_data_df$sex == "F", 1,</pre>
                        ifelse(thyroid_data_df$sex == "M", 0, NA))
thyroid_data_df[thyroid_data_df == "t"] <- 1</pre>
thyroid_data_df[thyroid_data_df == "f"] <- 0</pre>
thyroid_data_df
str(thyroid_data_df)
# Converting character columns to numeric
char_cols <- sapply(thyroid_data_df, is.character)</pre>
thyroid_data_df[char_cols] <- lapply(thyroid_data_df[char_cols],</pre>
                                       function(x) as.numeric(as.character(x)))
str(thyroid_data_df)
# Replace data points '?' with NA
thyroid_data_df[thyroid_data_df == "?"] <- NA</pre>
na_counts <- data.frame(</pre>
  Column = names(thyroid_data_df),
  Missing_Values = colSums(is.na(thyroid_data_df)),
  row.names = NULL
)
```

```
print(na_counts)
# Impute missing numeric values using the column mean
thyroid_data_df$age[is.na(thyroid_data_df$age)] <- mean(thyroid_data_df$age,
                                                          na.rm = TRUE)
thyroid_data_df$sex[is.na(thyroid_data_df$sex)] <- mean(thyroid_data_df$sex,</pre>
                                                          na.rm = TRUE)
# Impute missing numeric values using the column mean
cols_to_impute <- c("TSH", "T3", "TT4", "T4U", "FTI")</pre>
# Mean imputation
thyroid_data_df <- thyroid_data_df %>%
  mutate(across(all_of(cols_to_impute), ~ ifelse(is.na(.), mean(., na.rm = TRUE), .)))
# Replace data points '?' with NA
thyroid_data_df[thyroid_data_df == "?"] <- NA</pre>
na_counts <- data.frame(</pre>
  Column = names(thyroid_data_df),
 Missing_Values = colSums(is.na(thyroid_data_df)),
 row.names = NULL
print(na_counts)
# Filtering out outliers
thyroid_data_df <- subset(thyroid_data_df, age < 100)</pre>
# Preview cleaned dataset
thyroid_data_df
# Preview standardised structured of the dataset
str(thyroid_data_df)
```

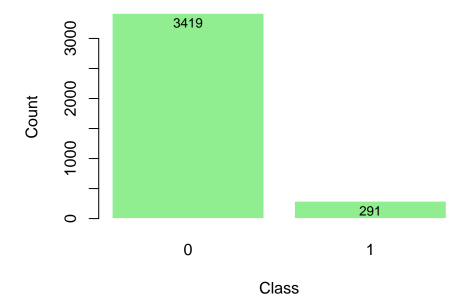
Step 3: Exploratory Data Analysis (EDA)

In this EDA step, we explore the thyroid dataset to uncover patterns, trends, and relationships between variables.

We begin by examining the distribution of individual features, such as binary class labels, using a bar plot. Relationships between numerical features (e.g., Age vs TSH levels) are explored using a scatter plot.

Lastly, a correlation matrix is generated for key thyroid-related features (e.g., T3, TT4, TSH) to visualize interdependencies and identify potential predictor variables for TSH levels.

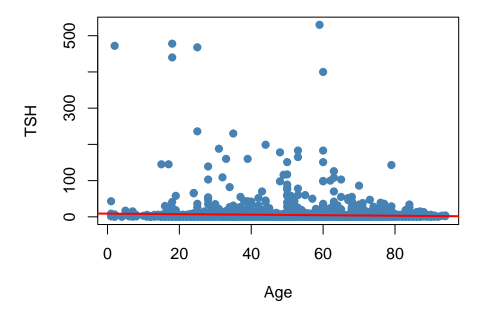
Distribution of Binary Class (Thyroid vs Normal)



```
# Scatterplot between Age and TSH levels
plot(thyroid_data_df$age, thyroid_data_df$TSH,
    main = "Scatterplot of Age vs TSH",
    xlab = "Age", ylab = "TSH",
    pch = 19, col = "steelblue")

# Add a linear regression line to observe trend
abline(lm(TSH ~ age, data = thyroid_data_df), col = "red", lwd = 2)
```

Scatterplot of Age vs TSH



```
correlation <- cor(thyroid_data_df$age, thyroid_data_df$TSH, use = "complete.obs")
cat("The correlation between Age and TSH is: ", round(correlation, 3), "\n")</pre>
```

The correlation between Age and TSH is: -0.059

```
# Heatmap of Thyroid-Related Features
selected_cols <- c("age", "sex", "FTI", "T3", "TT4", "TSH", "binaryClass")
subset_data <- thyroid_data_df[, selected_cols]

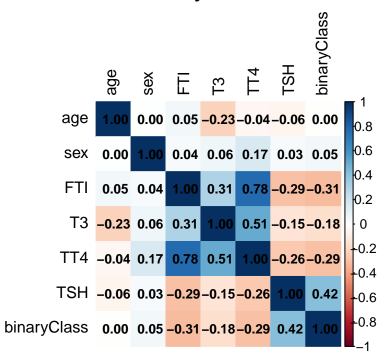
# Compute correlation matrix
cor_matrix <- cor(subset_data, use = "complete.obs")

library(corrplot)

corrplot(cor_matrix,</pre>
```

```
method = "color",
tl.col = "black",
addCoef.col = "black",
number.cex = 0.8,
title = "Correlation Matrix of Thyroid-Related features",
mar = c(0, 0, 2, 0))
```

Correlation Matrix of Thyroid-Related features



Step 4: Model Training

Regression

In this step, we build and evaluate predictive models to answer our regression research question. The goal is to assess whether applying machine learning can effectively predict TSH

We train three different regression models:

- 1. Random Forest a robust ensemble model that captures non-linear relationships.
- 2. Linear Regression a simple baseline model assuming linear relationships.
- 3. XGBoost a high-performance gradient boosting model

```
# Data preparation
rf_thyroid_data <- thyroid_data_df[!is.na(thyroid_data_df$TSH), ]</pre>
# Remove all columns that contains 'measured'
rf_thyroid_data <- rf_thyroid_data[, !grepl("measured", names(rf_thyroid_data))]</pre>
# Remove the 'binaryClass' column
rf_thyroid_data <- subset(rf_thyroid_data, select = -binaryClass)</pre>
rf_thyroid_data
# Splitting the dataset into training and test sets
set.seed(42)
sample_index <- sample(1:nrow(rf_thyroid_data), 0.7 * nrow(rf_thyroid_data))</pre>
thyroid_regress_train_data <- rf_thyroid_data[sample_index, ]</pre>
thyroid_regress_test_data <- rf_thyroid_data[-sample_index, ]</pre>
# Random Forest - Training
library(randomForest)
rf_model <- randomForest(TSH ~ ., data = thyroid_regress_train_data,</pre>
                          ntree = 100, importance = TRUE)
rf_preds_regress_train <- predict(rf_model, thyroid_regress_train_data)</pre>
rf_preds_regress_test <- predict(rf_model, thyroid_regress_test_data)
# Linear Regression - Training
lm_model <- lm(TSH ~ ., data = thyroid_regress_train_data)</pre>
lm_preds_regress_train <- predict(lm_model, thyroid_regress_train_data)</pre>
lm_preds_regress_test <- predict(lm_model, thyroid_regress_test_data)</pre>
# XGBoost - Training
library(xgboost)
```

Classification

In this step, we build a classification model to answer our classification research question. The goal is to predict whether an individual has thyroid dysfunction (binaryClass) based on clinical and demographic features.

We use the **Random Forest Classifier** to train the model and evaluate its performance using metrics such as accuracy, sensitivity, specificity, and confusion matrix.

To address the class imbalance in the dataset, **SMOTE** is applied to the training set, ensuring that the model is **not biased toward the majority class**.

```
# Data preparation
thyroid_data_df$binaryClass <- as.factor(thyroid_data_df$binaryClass)

# Remove all columns that contains 'measured'
clf_thyroid_data <- thyroid_data_df[, !grepl("measured", names(thyroid_data_df))]

# Remove TSH as its also the target variable
clf_thyroid_data <- clf_thyroid_data[, setdiff(names(clf_thyroid_data), "TSH")]

clf_thyroid_data

# Splitting the dataset into training and test sets
set.seed(42)
sample_index <- sample(1:nrow(clf_thyroid_data), 0.7 * nrow(clf_thyroid_data))

thyroid_classify_train_data <- clf_thyroid_data[sample_index, ]
thyroid_classify_test_data <- clf_thyroid_data[-sample_index, ]</pre>
```

Step 5: Model Evaluation

Regression

In this step, each model were trained on a subset of the dataset (training set) and evaluated on unseen data (test set) to assess generalizability.

Key evaluation metrics include:

- R-squared (R2) proportion of variance in TSH explained by the model.
- Root Mean Squared Error (RMSE) average prediction error magnitude.

Finally, we use SHAP (Shapley Additive Explanations) to interpret the predictions of the best-performing model, identifying which features most influence TSH levels.

```
library(Metrics)
r2_score <- function(actual, predicted) {</pre>
  cor(actual, predicted)^2
}
results_df <- data.frame(</pre>
  Model = c("Random Forest", "Linear Regression", "XGBoost"),
  R_squared_Train = round(c(
   r2_score(thyroid_regress_train_data$TSH, rf_preds_regress_train),
   r2_score(thyroid_regress_train_data$TSH, lm_preds_regress_train),
   r2_score(thyroid_regress_train_data$TSH, xgb_preds_regress_train)
  ), 3),
  R squared Test = round(c(
   r2_score(thyroid_regress_test_data$TSH, rf_preds_regress_test),
   r2_score(thyroid_regress_test_data$TSH, lm_preds_regress_test),
   r2_score(thyroid_regress_test_data$TSH, xgb_preds_regress_test)
 ), 3),
  RMSE = round(c(
   rmse(thyroid_regress_test_data$TSH, rf_preds_regress_test),
   rmse(thyroid_regress_test_data$TSH, lm_preds_regress_test),
   rmse(thyroid_regress_test_data$TSH, xgb_preds_regress_test)
  ), 3)
)
```

```
## Model R_squared_Train R_squared_Test RMSE
## 1 Random Forest 0.784 0.598 17.110
## 2 Linear Regression 0.112 0.185 23.150
## 3 XGBoost 0.997 0.592 17.135
```

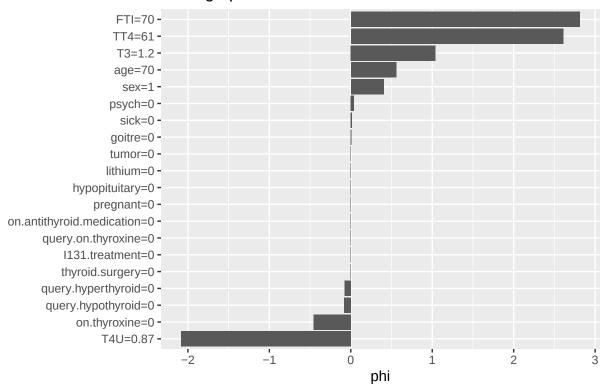
```
library(iml)
```

```
# Remove TSH from features
X <- thyroid_regress_test_data[, setdiff(names(thyroid_regress_test_data), "TSH")]
y <- thyroid_regress_test_data$TSH

# Create iml Predictor object
predictor <- Predictor$new(
    model = rf_model,
    data = X,
    y = y
)

# Choose a single observation to interpret
shap <- Shapley$new(predictor, x.interest = X[1, ])
plot(shap)</pre>
```

Actual prediction: 10.00 Average prediction: 5.06



Classification

In this step, the classification model selected was trained on a subset of the dataset (training set) and evaluated on unseen data (test set) to assess its ability to generalize to new cases.

Key evaluation metrics include:

- Accuracy the overall proportion of correct predictions.
- Sensitivity (Recall) the model's ability to correctly identify individuals with thyroid dysfunction.
- Specificity the ability to correctly identify normal individuals.
- Balanced Accuracy average sensitivity and specificity, useful for imbalanced data.

To enhance interpretability, we also used SHAP (Shapley Additive Explanations) to understand how individual features contribute to the classification decisions. This helps identify the most influential clinical and demographic features in classifying thyroid dysfunction.

```
library(caret)

# Confusion matrix
conf_mat <- confusionMatrix(rf_preds_classify_test, thyroid_classify_test_data$binaryClass)
print(conf_mat)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
               0
                    1
##
            0 972
                   59
##
            1 45
##
                  Accuracy: 0.9066
##
                    95% CI: (0.8879, 0.923)
##
       No Information Rate: 0.9137
##
##
       P-Value [Acc > NIR] : 0.8187
##
                     Kappa: 0.3653
##
##
##
    Mcnemar's Test P-Value: 0.2024
##
               Sensitivity: 0.9558
##
##
               Specificity: 0.3854
##
            Pos Pred Value: 0.9428
##
            Neg Pred Value: 0.4512
                Prevalence: 0.9137
##
##
            Detection Rate: 0.8733
##
      Detection Prevalence: 0.9263
         Balanced Accuracy: 0.6706
##
##
##
          'Positive' Class : 0
##
```

```
cat("Test Accuracy:", round(mean(rf_preds_classify_test == thyroid_classify_test_data$binaryClass), 3), "\n")
```

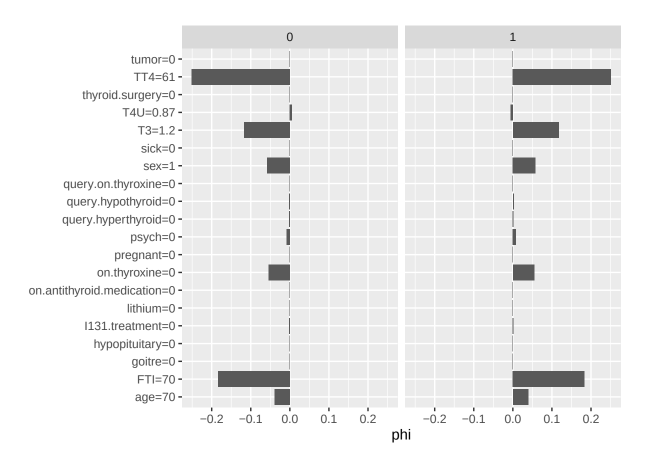
Test Accuracy: 0.907

```
library(iml)

# Remove TSH from features
X <- thyroid_classify_test_data[, setdiff(names(thyroid_classify_test_data), "binaryClass")]
y <- thyroid_classify_test_data$binaryClass

# Create iml Predictor object
predictor <- Predictor$new(
    model = rf_clf,
    data = X,
    y = y,
    type = "prob"
)

# Choose a single observation to interpret
shap <- Shapley$new(predictor, x.interest = X[1, ])
plot(shap)</pre>
```



Step 6: Interpretation of Results

Result Interretation - Regression

Key Takeaways:

Model Summary:

- Random Forest and XGBoost both performed reasonably well on test data (R2 approximately 0.60), explaining about 60% of the variance in TSH levels.
- Linear Regression performed poorly, indicating a non-linear relationship between predictors and TSH.
- XGBoost has the best fit on training data, but a similar test performance to Random Forest, suggesting
 possible overfitting.

Feature Summary:

- The top contributors to TSH prediction include FTI, TT4, T3, age, and sex.
- Features like T4U and on.thyroxine negatively influenced the prediction for this individual.
- This matches clinical expectations hormone levels and demographics strong indicators of thyroid health.

Overall, it is **feasible to predict an individual's TSH level** using machine learning models trained on clinical and demographic data. The models demonstrate **good predictive performance** and **feature interpretability**, making them promising for **clinical use** and **early detection** of thyroid issues.

Result Interpreation - Classification

Key Takeaways:

Model Summary:

- A Random Forest Classifier was used to predict binaryClass (0 = Positive Thyroid, 1 = Negative Thyroid)
- The model achieved a test accuracy of 90.7%, which reflects strong overall performance.
- However, due to class imbalance, there is a trade-off between sensitivity and specificity:
- Sensitivity (Recall for Thyroid cases): 0.9558 -> Excellent at identifying individuals with thyroid dysfunction.
- Specificity (Recall for Normal cases): 0.3854 -> Poor at identifying truly normal individuals.
- Balanced Accuracy: 0.6706 -> Moderate overall performance when accounting for imbalance.
- Kappa: 0.3653 -> Indicates fair agreement between predicted and actual classes.

Feature Summary:

- ${f FTI},\,{f TT4},\,{f T3}$ Hormone levels relevant to thy roid function.
- on.thyroxine Medication use is a strong signal for thyroid issues
- ${\bf age,\,sex}$ Demographic features also contribute meaningfully

Overall, it is feasible to classify individuals as having thyroid dysfunction or being in a normal state using the features in the dataset. The Random Forest model achieved high overall accuracy and strong sensitivity for detecting thyroid conditions.

However, the **low specificity** suggests **room for improvement** in detecting truly negative thyroid individuals. To further improve the model:

- Experiment with other classifiers with threshold tuning (e.g., XGBoost, Logistic Regression).
- Perform threshold optimization to better balance sensitivity and specificity.

References

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- 2. MedlinePlus. (2023). TSH (Thyroid-stimulating hormone) test. U.S. National Library of Medicine. https://medlineplus.gov/lab-tests/tsh-thyroid-stimulating-hormone-test/
- 3. Yasserhessein. (2022). Thyroid Disease Data Set. Kaggle. Retrieved from https://www.kaggle.com/datasets/yasserhessein/thyroid-disease-data-set

file.rename("src/thyroid_health.pdf", "output/thyroid_health_report.pdf")