$assignment_2$

2024-10-16

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Problem 1. Regression

```
data <- read.csv("qsar_aquatic_toxicity.csv", sep = ";", header = FALSE)

# Since the raw data does not have column names, we will assign them manually
names(data) <- c(
    "TPSA",
    "SAacc",
    "H050",
    "MLOGP",
    "RDCHI",
    "GATS1p",
    "nN",
    "C040",
    "LC50"
)</pre>
head(data)
```

```
##
      TPSA
            SAacc H050 MLOGP RDCHI GATS1p nN C040 LC50
## 1
      0.00
            0.000 0 2.419 1.225 0.667 0
                                             0 3.740
## 2
     0.00 0.000
                  0 2.638 1.401 0.632 0
                                             0 4.330
## 3
     9.23 11.000 0 5.799 2.930 0.486 0
                                             0 7.019
     9.23 11.000 0 5.453 2.887 0.495 0
                                             0 6.723
      9.23 11.000
## 5
                    0 4.068 2.758 0.695 0
                                             0 5.979
## 6 215.34 327.629
                    3 0.189 4.677 1.333 0
                                             4 6.064
```

a. Data splitting

Dimension of Test Set: 182x9

We split the data into a training and a test set, with approximately 2/3 and 1/3 of the observations, respectively.

```
# Use 2/3 of dataset as training set and remaining 1/3 as testing set
set.seed(123)
sample <- sample.split(data$LC50, SplitRatio = 2/3)
train <- subset(data, sample == TRUE)
test <- subset(data, sample == FALSE)</pre>
```

```
cat("Dimension of Training Set:", paste(dim(train), collapse = "x"), "\nDimension of Test Set:", paste(
## Dimension of Training Set: 364x9
```

(i) Original Model

First, we will fit a linear regression model on the training data using all the predictors.

```
# To make sure we use the same split in (i) and (ii)
train_i = train
test_i = test
```

The initial linear regression model shows significant predictors including TPSA, SAacc, MLOGP, RDCHI, GATS1p, and nN based on p-values less than 0.05. However, H050 and C040 do not appear to have a significant impact.

```
# Fit linear regression model on training data
model <- lm(LC50 ~ ., data=train_i)</pre>
summary(model)
##
## Call:
## lm(formula = LC50 ~ ., data = train_i)
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
## -2.8548 -0.8166 -0.1830 0.6771
                                   4.8867
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
                          0.312580
                                    8.411 1.00e-15 ***
## (Intercept) 2.629264
## TPSA
               0.027092
                          0.003336
                                    8.121 7.74e-15 ***
                          0.002652 -6.017 4.42e-09 ***
## SAacc
              -0.015959
## H050
              -0.003879
                          0.076369 -0.051 0.959522
## MLOGP
               0.400783
                          0.081760
                                    4.902 1.45e-06 ***
## RDCHI
               0.654990
                          0.177787
                                     3.684 0.000265 ***
              -0.589994
                          0.195299 -3.021 0.002702 **
## GATS1p
              -0.199466
                          0.059602 -3.347 0.000906 ***
## nN
## C040
              -0.046002
                          0.091165 -0.505 0.614156
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.219 on 355 degrees of freedom
## Multiple R-squared: 0.5024, Adjusted R-squared: 0.4912
## F-statistic: 44.8 on 8 and 355 DF, p-value: < 2.2e-16
```

We will predict the LC50 values on the training and test datasets to evaluate the model using

```
• Mean Squared Error (MSE): MSE = \sum_{i=1}^{D} (x_i - y_i)^2
• Root Mean Squared Error (RMSE): RMSE = \sqrt{MSE}
• R-squared: R^2 = 1 - \frac{\sum_{i=1}^{D} (x_i - y_i)^2}{\sum_{i=1}^{D} (x_i - \bar{x}_i)^2}
```

```
# Predict on training and test datasets
pred_train <- predict(model, newdata=train_i)
pred_test <- predict(model, newdata=test_i)</pre>
```

```
# Adding predictions columns to the datasets
train_i$predicted_LC50 <- pred_train
test_i$predicted_LC50 <- pred_test</pre>
```

We have the following result for the model on the training and test set as follows. We see that The training and test set metrics are reasonably close, with R-squared values indicating that approximately 50% of the variance is explained by the model in the training set and around 43% in the test set, suggesting the model generalizes fairly well.

```
cat(paste0(
  "Training Metrics:\n",
  "MSE (Train): ", mse_train, "\n",
  "RMSE (Train): ", rmse_train, "\n",
  "R-squared (Train): ", r2_train, "\n\n",
 "Test Metrics:\n",
  "MSE (Test): ", mse_test, "\n",
  "RMSE (Test): ", rmse_test, "\n",
  "R-squared (Test): ", r2_test, "\n"
## Training Metrics:
## MSE (Train): 1.44990640082018
## RMSE (Train): 1.20412059230801
## R-squared (Train): 0.502397645581479
##
## Test Metrics:
## MSE (Test): 1.40224882922927
## RMSE (Test): 1.18416587910194
## R-squared (Test): 0.433587696937759
```

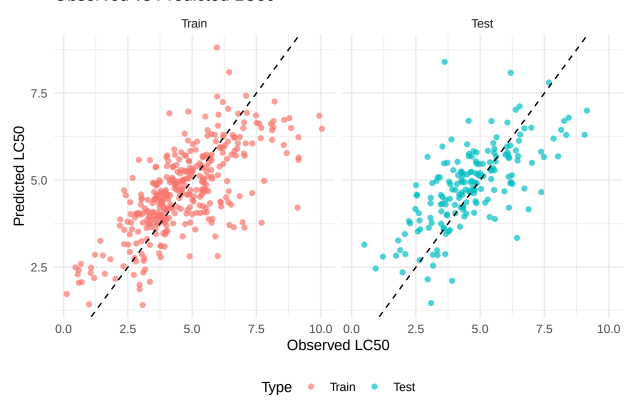
Plotting the observed vs predicted LC50 values for the training and test sets, we can see that the model generally performs well, with most points falling close to the dashed line (y=x) indicating perfect predictions.

```
# Combine data for plotting
train_i$Type <- 'Train'
test_i$Type <- 'Test'
combined_data <- rbind(train_i, test_i)

combined_data$Type <- factor(combined_data$Type, levels = c('Train', 'Test'))</pre>
```

```
# Plotting observed vs predicted LC50 values
ggplot(combined_data, aes(x = LC50, y = predicted_LC50, color = Type)) +
  geom_point(alpha = 0.7) +
  geom_abline(intercept = 0, slope = 1, linetype = "dashed") +
  labs(title = "Observed vs Predicted LC50", x = "Observed LC50", y = "Predicted LC50") +
  theme_minimal() +
  facet_wrap(~Type) +
  theme(legend.position = "bottom")
```

Observed vs Predicted LC50



(ii). Dummy encoding

We can see that in our data set, there are 3 count variables (H050, nN, C040) that represent the number of specific atoms in the chemical compounds. We will transform these variables using a 0/1 dummy encoding, where 0 represents the absence of the specific atom, and 1 represents the presence of the specific atoms. In this case, I suppose that the model will perform a litter bit worse than the original model because we lose some information by transforming the count variables into binary variables. On the other hand, it may help to reduce overfitting because it simplifies the model.

```
# To make sure we use the same split in (i) and (ii)
train_ii = train
test_ii = test
```

Transform 3 count variables (H050, nN, C040) into 0/1 in train and test datasets

```
train_ii$H050 <- ifelse(train_ii$H050 > 0, 1, 0)
train_ii$nN <- ifelse(train_ii$nN > 0, 1, 0)
train_ii$C040 <- ifelse(train_ii$C040 > 0, 1, 0)

test_ii$H050 <- ifelse(test_ii$H050 > 0, 1, 0)
test_ii$nN <- ifelse(test_ii$nN > 0, 1, 0)
test_ii$C040 <- ifelse(test_ii$C040 > 0, 1, 0)
```

head(train ii)

```
##
       TPSA
              SAacc H050 MLOGP RDCHI GATS1p nN C040 LC50
## 1
       0.00
              0.000
                       0 2.419 1.225 0.667 0
                                                  0 3.740
## 3
       9.23 11.000
                       0 5.799 2.930
                                      0.486 0
                                                  0 7.019
## 6
    215.34 327.629
                       1 0.189 4.677
                                      1.333 0
                                                  1 6.064
## 7
       9.23 11.000
                       0 2.723 2.321
                                      1.165
                                            0
                                                  0 7.337
## 9
       0.00
              0.000
                       0 2.067 1.800 1.250 0
                                                  0 3.941
## 10
       0.00
              0.000
                       0 2.746 1.667 1.400 0
                                                  0 3.809
```

After transforming the count variables into binary variables, we will fit a linear regression model on the training data using all the predictors.

```
# Fit linear regression model on transformed training data
model_transform_dummy <- lm(LC50 ~ ., data = train_ii)
summary(model_transform_dummy)</pre>
```

```
##
## Call:
## lm(formula = LC50 ~ ., data = train_ii)
## Residuals:
##
      Min
              1Q Median
                             3Q
                                    Max
## -3.0873 -0.8306 -0.1303 0.6571 5.0526
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.693545 0.315949
                                 8.525 4.44e-16 ***
## TPSA
              0.023267 0.003365
                                 6.914 2.20e-11 ***
## SAacc
             0.161558 -0.559 0.57684
## H050
             -0.090233
## MLOGP
              0.436885
                        0.082632
                                 5.287 2.18e-07 ***
                                 3.070 0.00231 **
## RDCHI
              0.553158
                        0.180181
## GATS1p
             -0.539057
                        0.190296 -2.833 0.00488 **
## nN
              0.018072
                        0.156479
                                  0.115 0.90812
## C040
             -0.124928
                        0.169094 -0.739 0.46051
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.239 on 355 degrees of freedom
## Multiple R-squared: 0.4864, Adjusted R-squared: 0.4749
## F-statistic: 42.03 on 8 and 355 DF, p-value: < 2.2e-16
```

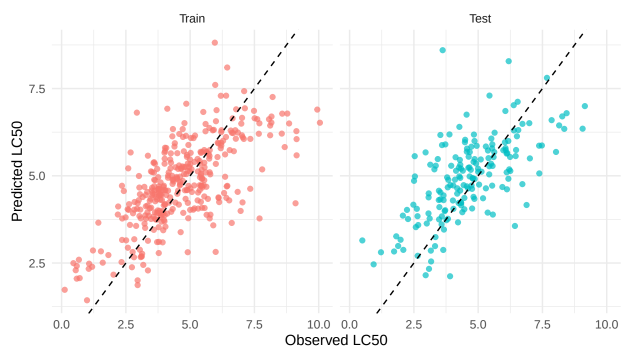
```
# Predict on training and test datasets
pred_train_transform_dummy <- predict(model, newdata=train_ii)
pred_test_transform_dummy <- predict(model, newdata=test_ii)

# Adding predictions columns to the datasets
train_ii$predicted_LC50 <- pred_train_transform_dummy
test_ii$predicted_LC50 <- pred_test_transform_dummy</pre>
```

Based on the results, we see that R^2 values dropped from 43% in the original model to around 38% after the transformation. For MSE and RMSE in test set, it increased slightly from 1.18 to 1.53 and 1.18 to 1.23, respectively.

```
# Evaluate model: calculate MSE, RMSE, and R-squared for training and test sets
mse_train_transform_dummy <- mean((train_ii$LC50 - train_ii$predicted_LC50)^2)</pre>
rmse_train_transform_dummy <- sqrt(mse_train_transform_dummy)</pre>
r2_train_transform_dummy <- 1 - (sum((train_ii$LC50 - train_ii$predicted_LC50)^2) / sum((train_ii$LC50
mse_test_transform_dummy <- mean((test_ii$LC50 - test_ii$predicted_LC50)^2)</pre>
rmse_test_transform_dummy <- sqrt(mse_test_transform_dummy)</pre>
r2_test_transform_dummy <- 1 - (sum((test_ii$LC50 - test_ii$predicted_LC50)^2) / sum((test_ii$LC50 - me
cat(paste0(
  "Training Metrics:\n",
  "MSE (Train): ", mse_train_transform_dummy, "\n",
  "RMSE (Train): ", rmse_train_transform_dummy, "\n",
  "R-squared (Train): ", r2_train_transform_dummy, "\n\n",
  "Test Metrics:\n",
 "MSE (Test): ", mse_test_transform_dummy, "\n",
  "RMSE (Test): ", rmse_test_transform_dummy, "\n",
  "R-squared (Test): ", r2_test_transform_dummy, "\n"
))
## Training Metrics:
## MSE (Train): 1.53935201233042
## RMSE (Train): 1.24070625545711
## R-squared (Train): 0.471700252387877
## Test Metrics:
## MSE (Test): 1.53043849004967
## RMSE (Test): 1.23710892408457
## R-squared (Test): 0.381807870490008
# Combine data for plotting
train_ii$Type <- 'Train'</pre>
test_ii$Type <- 'Test'</pre>
combined_data <- rbind(train_ii, test_ii)</pre>
combined_data$Type <- factor(combined_data$Type, levels = c('Train', 'Test'))</pre>
# Plotting observed vs predicted LC50 values
ggplot(combined_data, aes(x = LC50, y = predicted_LC50, color = Type)) +
```

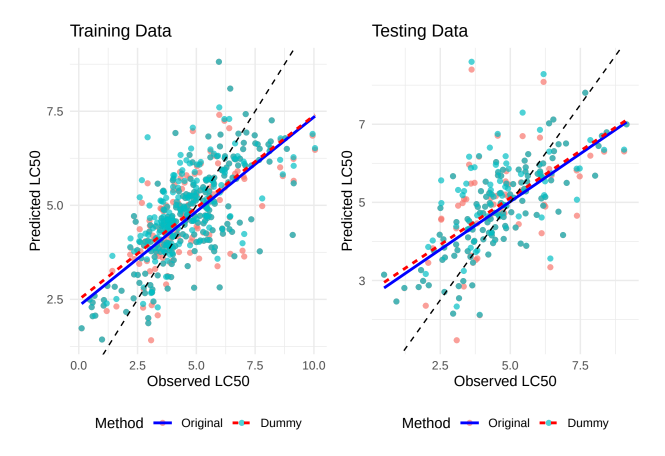
Dummy Encoding: Observed vs Predicted LC50



Type • Train • Test

```
# Prepare combined data
train_combined <- train_i[, c("LC50", "predicted_LC50")]
train_combined$Method <- 'Original'
train_combined$Type <- 'Train'
train_ii_combined <- train_ii[, c("LC50", "predicted_LC50")]
train_ii_combined$Method <- 'Dummy'
train_ii_combined$Type <- 'Train'
train_combined_all <- rbind(train_combined, train_ii_combined)
test_combined <- test_i[, c("LC50", "predicted_LC50")]
test_combined$Method <- 'Original'
test_combined$Type <- 'Test'
test_ii_combined$Method <- 'Dummy'
test_ii_combined$Method <- 'Dummy'
test_ii_combined$Type <- 'Test'
test_ii_combined$Type <- 'Test'
test_combined$Type <- 'Test'</pre>
```

```
# Convert 'Method' and 'Type' to factors
train_combined_all$Method <- factor(train_combined_all$Method, levels = c('Original', 'Dummy'))</pre>
test_combined_all$Method <- factor(test_combined_all$Method, levels = c('Original', 'Dummy'))</pre>
# Function to draw regression lines
add_regression_lines <- function(df, original_model, dummy_model) {</pre>
ggplot(df, aes(x = LC50, y = predicted_LC50, color = Method)) +
geom_point(alpha = 0.7) +
geom smooth(method = "lm", formula = y ~ x, se = FALSE,
aes(linetype = Method),
data = df[df$Method == 'Original', ],
color = 'blue') +
geom_smooth(method = "lm", formula = y ~ x, se = FALSE,
aes(linetype = Method),
data = df[df$Method == 'Dummy', ],
color = 'red') +
geom_abline(intercept = 0, slope = 1, linetype = "dashed") +
labs(x = "Observed LC50", y = "Predicted LC50", title = df$Type[1]) +
theme_minimal() +
theme(legend.position = "bottom")
}
# Plot training data with both regression lines
train_plot <- add_regression_lines(train_combined_all, model, model_transform_dummy)</pre>
train_plot <- train_plot + labs(title = "Training Data")</pre>
# Plot testing data with both regression lines
test_plot <- add_regression_lines(test_combined_all, model, model_transform_dummy)</pre>
test plot <- test plot + labs(title = "Testing Data")</pre>
# Display plots side by side
grid.arrange(train_plot, test_plot, ncol = 2)
```



In initial conclusion in one time spliting, the original model (without dummy encoding) provides a better fit to both the training and testing data, as evidenced by its closer alignment with the ideal prediction line and lower dispersion in the test data. This is likely because it retains the continuous information in the count variables, which adds more nuance to the model's predictions. So, in part b, we will draw a more reliable conclusion by repeating the procedure 200 times and comparing the average test errors.

b. Repeating the procedure 200 times

Procedure

- Randomly spiting training vs test set (2/3 vs 1/3).
- Fit the models with 2 options (i) Original model and (ii) Dummy encoding.
- Record the test errors (MSE/RMSE/ R^2).

```
# Initialize vectors to store test errors
mse_test_errors_i <- numeric(200)
rmse_test_errors_i <- numeric(200)
r2_test_errors_i <- numeric(200)
mse_test_errors_ii <- numeric(200)
rmse_test_errors_ii <- numeric(200)
r2_test_errors_ii <- numeric(200)
# Repeat the procedure 200 times
set.seed(2)</pre>
```

```
for (i in 1:200) {
  # Split the data
  sample <- sample.split(data$LC50, SplitRatio = 2/3)</pre>
  train <- subset(data, sample == TRUE)</pre>
  test <- subset(data, sample == FALSE)</pre>
  # Option (i): Original model
  model <- lm(LC50 ~ ., data=train)</pre>
  pred_test_i <- predict(model, newdata=test)</pre>
  mse_test_i <- mean((test$LC50 - pred_test_i)^2)</pre>
  rmse_test_i <- sqrt(mse_test_i)</pre>
   r2\_test\_i \leftarrow 1 - (sum((test\$LC50 - pred\_test\_i)^2) / sum((test\$LC50 - mean(test\$LC50))^2)) 
  # Option (ii): Dummy encoding
  train$H050 <- ifelse(train$H050 > 0, 1, 0)
  train$nN <- ifelse(train$nN > 0, 1, 0)
  train$C040 <- ifelse(train$C040 > 0, 1, 0)
  test$H050 <- ifelse(test$H050 > 0, 1, 0)
  test$nN <- ifelse(test$nN > 0, 1, 0)
  test$C040 \leftarrow ifelse(test$C040 > 0, 1, 0)
  model_ii <- lm(LC50 ~ ., data = train)</pre>
  pred_test_ii <- predict(model_ii, newdata = test)</pre>
  mse_test_ii <- mean((test$LC50 - pred_test_ii)^2)</pre>
  rmse_test_ii <- sqrt(mse_test_ii)</pre>
  r2_{test_{ii}} \leftarrow 1 - (sum((test_{LC50} - pred_{test_{ii}})^2) / sum((test_{LC50} - mean(test_{LC50}))^2))
  # Record the test errors
  mse_test_errors_i[i] <- mse_test_i</pre>
  rmse_test_errors_i[i] <- rmse_test_i</pre>
  r2_test_errors_i[i] <- r2_test_i
  mse_test_errors_ii[i] <- mse_test_ii</pre>
  rmse_test_errors_ii[i] <- rmse_test_ii</pre>
  r2_test_errors_ii[i] <- r2_test_ii
```

There are a few key reasons for repeating the procedure 200 times:

- To reduce the influence of random data splits
- To provide a more reliable estimate of the model's performance

```
# Calculate and print average test errors
average_test_error_i <- mean(mse_test_errors_i)
average_rmse_error_i <- mean(rmse_test_errors_i)
average_r2_error_i <- mean(r2_test_errors_i)
average_test_error_ii <- mean(mse_test_errors_ii)
average_rmse_error_ii <- mean(rmse_test_errors_ii)
average_r2_error_ii <- mean(r2_test_errors_ii)</pre>
```

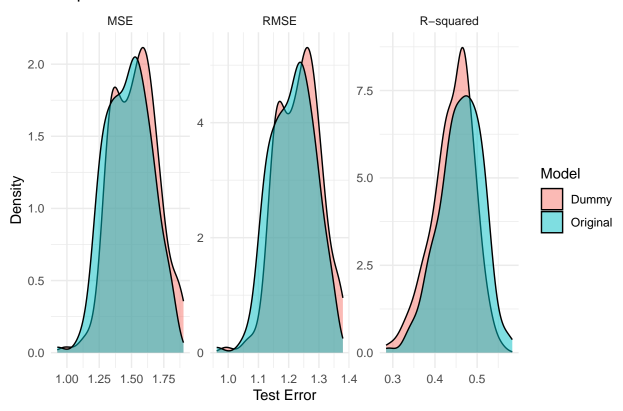
```
cat(paste0(
  "Average Test Errors (Original Model):\n",
  "MSE: ", average_test_error_i, "\n",
  "RMSE: ", average_rmse_error_i, "\n",
  "R-squared: ", average_r2_error_i, "\n\n",
  "Average Test Errors (Dummy Model):\n",
  "MSE: ", average test error ii, "\n",
  "RMSE: ", average rmse error ii, "\n",
  "R-squared: ", average_r2_error_ii, "\n"
## Average Test Errors (Original Model):
## MSE: 1.47708146772242
## RMSE: 1.21330895603276
## R-squared: 0.460485255063669
##
## Average Test Errors (Dummy Model):
## MSE: 1.52752950559007
## RMSE: 1.23398478875802
## R-squared: 0.442128799570138
```

The original model consistently achieves lower MSE and RMSE than the dummy-encoded model, as indicated by the density distributions. The peak of the distribution for the original model is shifted left compared to the dummy model, meaning the original model typically has smaller test errors. The dummy-encoded model shows slightly larger and more spread-out test errors, indicating poorer performance.

```
# Create data frames for plotting
errors_df_mse <- data.frame(</pre>
 Error = c(mse_test_errors_i, mse_test_errors_ii),
 Metric = 'MSE',
  Model = factor(rep(c("Original", "Dummy"), each = 200))
errors_df_rmse <- data.frame(</pre>
  Error = c(rmse_test_errors_i, rmse_test_errors_ii),
 Metric = 'RMSE',
  Model = factor(rep(c("Original", "Dummy"), each = 200))
errors_df_r2 <- data.frame(</pre>
  Error = c(r2_test_errors_i, r2_test_errors_ii),
 Metric = 'R-squared',
 Model = factor(rep(c("Original", "Dummy"), each = 200))
)
errors_df <- rbind(errors_df_mse, errors_df_rmse, errors_df_r2)</pre>
# Ensure the 'Metric' factor has the correct level order
errors_df$Metric <- factor(errors_df$Metric, levels = c('MSE', 'RMSE', 'R-squared'))
# Plot the empirical distributions of the test errors
ggplot(errors_df, aes(x = Error, fill = Model)) +
  geom_density(alpha = 0.5) +
 facet_wrap(~ Metric, scales = "free") +
```

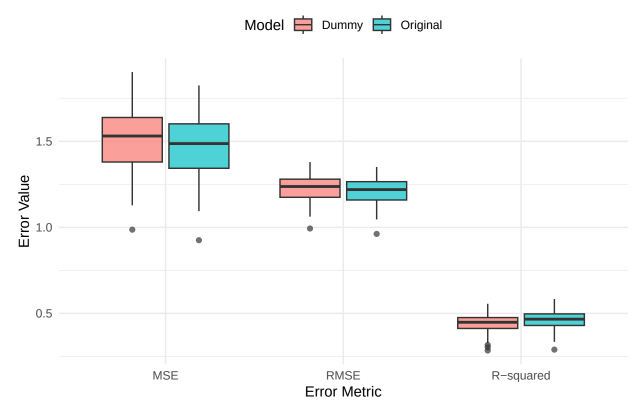
```
labs(title = "Empirical Distributions of Test Errors", x = "Test Error", y = "Density") +
theme_minimal()
```

Empirical Distributions of Test Errors



```
# Plot the empirical distributions of the test errors using boxplots
ggplot(errors_df, aes(x = Metric, y = Error, fill = Model)) +
  geom_boxplot(alpha = 0.7) +
  labs(title = "Boxplots of Test Errors", x = "Error Metric", y = "Error Value") +
  theme_minimal() +
  theme(legend.position = "top")
```

Boxplots of Test Errors



In conclusion, the higher test errors and greater variability of the dummy-encoded model (option ii) occur because it sacrifices important information present in the original continuous variables. Repeating the process confirms that the original model (option i) is generally superior.

c. Variable selection procedures

```
# Split the data into training (2/3) and test (1/3) sets
set.seed(123)
sample <- sample.split(data$LC50, SplitRatio = 2/3)
train <- subset(data, sample == TRUE)
test <- subset(data, sample == FALSE)

# Set up full and null model
full.model <- lm(LC50 ~ ., data = train)
null.model <- lm(LC50 ~ 1, data = train)

# Set up target and number of variables
y <- train$LC50
num_vars <- ncol(train) - 1 # exclude the response variable column</pre>
```

Forward Selection

Forward Selection is a stepwise regression method that starts with an empty model and adds predictors one by one based on a criterion (e.g., AIC, BIC) until no more predictors can be added.

- Key variables that were consistently selected include MLOGP, TPSA, SAacc, nN, RDCHI, and GATS1p.
- Both AIC and BIC agreed on the significance of these variables. However, BIC, being stricter, might lead to simpler models in other datasets, but in this case, the results remained the same across both criteria.

```
# With AIC
model.forward.aic <- stepAIC(null.model, scope = list(lower = null.model, upper = full.model), direction
summary(model.forward.aic)
##
## Call:
## lm(formula = LC50 ~ MLOGP + TPSA + SAacc + nN + RDCHI + GATS1p,
##
       data = train)
##
## Residuals:
##
      Min
                                30
                1Q Median
                                       Max
## -2.8194 -0.8018 -0.1737 0.6654
                                   4.8981
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.653540
                          0.285743
                                     9.286 < 2e-16 ***
## MLOGP
                           0.078544
                                    5.144 4.44e-07 ***
               0.404067
## TPSA
               0.027138
                           0.003284
                                    8.265 2.78e-15 ***
## SAacc
               -0.016185
                           0.002177 -7.435 7.84e-13 ***
               -0.201305
                           0.058114 -3.464 0.000597 ***
## RDCHI
               0.639082
                           0.174662
                                    3.659 0.000291 ***
              -0.589921
                           0.183821 -3.209 0.001452 **
## GATS1p
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.216 on 357 degrees of freedom
## Multiple R-squared: 0.502, Adjusted R-squared: 0.4937
## F-statistic: 59.98 on 6 and 357 DF, p-value: < 2.2e-16
# With BIC
# If we set it to k = log(n), the function considers the BIC.
model.forward.bic <- stepAIC(null.model, scope = list(lower = null.model, upper = full.model), direction
summary(model.forward.bic)
##
## Call:
## lm(formula = LC50 ~ MLOGP + TPSA + SAacc + nN + RDCHI + GATS1p,
##
       data = train)
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
## -2.8194 -0.8018 -0.1737 0.6654 4.8981
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
```

```
## (Intercept) 2.653540
                          0.285743
                                     9.286 < 2e-16 ***
## MLOGP
                                     5.144 4.44e-07 ***
               0.404067
                          0.078544
               0.027138
                          0.003284
                                     8.265 2.78e-15 ***
## TPSA
                                    -7.435 7.84e-13 ***
## SAacc
               -0.016185
                          0.002177
## nN
               -0.201305
                          0.058114
                                    -3.464 0.000597 ***
## RDCHI
               0.639082
                          0.174662
                                     3.659 0.000291 ***
## GATS1p
              -0.589921
                          0.183821 -3.209 0.001452 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.216 on 357 degrees of freedom
## Multiple R-squared: 0.502, Adjusted R-squared:
## F-statistic: 59.98 on 6 and 357 DF, p-value: < 2.2e-16
```

Backward Elimination

Backward Elimination is a stepwise regression method that starts with the full model and removes predictors one by one based on a criterion (e.g., AIC, BIC) until no more predictors can be removed.

- Similar to forward selection, backward elimination with both AIC and BIC resulted in a model that includes MLOGP, TPSA, SAacc, nN, RDCHI, and GATS1p.
- The consistency between forward and backward selection indicates that these predictors are strong, regardless of the method or criterion (AIC vs. BIC) used.

```
# With AIC
model.backward.aic <- stepAIC(full.model, direction = 'backward', trace = FALSE)
summary(model.backward.aic)</pre>
```

```
##
## Call:
## lm(formula = LC50 ~ TPSA + SAacc + MLOGP + RDCHI + GATS1p + nN,
##
       data = train)
##
## Residuals:
##
      Min
                1Q Median
                                30
                                       Max
## -2.8194 -0.8018 -0.1737 0.6654
                                   4.8981
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.653540
                           0.285743
                                      9.286 < 2e-16 ***
## TPSA
                0.027138
                           0.003284
                                      8.265 2.78e-15 ***
## SAacc
                           0.002177 -7.435 7.84e-13 ***
               -0.016185
## MLOGP
                0.404067
                           0.078544
                                      5.144 4.44e-07 ***
## RDCHI
                0.639082
                           0.174662
                                      3.659 0.000291 ***
               -0.589921
                           0.183821
                                     -3.209 0.001452 **
## GATS1p
                           0.058114
                                    -3.464 0.000597 ***
## nN
               -0.201305
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.216 on 357 degrees of freedom
## Multiple R-squared: 0.502, Adjusted R-squared: 0.4937
## F-statistic: 59.98 on 6 and 357 DF, p-value: < 2.2e-16
```

```
model.backward.bic <- stepAIC(full.model, direction = 'backward', k = log(nrow(train)), trace = FALSE)</pre>
summary(model.backward.bic)
##
## Call:
## lm(formula = LC50 ~ TPSA + SAacc + MLOGP + RDCHI + GATS1p + nN,
##
       data = train)
##
## Residuals:
##
       Min
                1Q Median
                                ЗQ
                                       Max
## -2.8194 -0.8018 -0.1737
                           0.6654
                                    4.8981
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.653540
                           0.285743
                                      9.286 < 2e-16 ***
## TPSA
                0.027138
                           0.003284
                                      8.265 2.78e-15 ***
## SAacc
               -0.016185
                           0.002177 -7.435 7.84e-13 ***
## MLOGP
                0.404067
                           0.078544
                                     5.144 4.44e-07 ***
## RDCHI
                0.639082
                           0.174662
                                     3.659 0.000291 ***
## GATS1p
               -0.589921
                           0.183821 -3.209 0.001452 **
               -0.201305
                           0.058114 -3.464 0.000597 ***
## nN
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.216 on 357 degrees of freedom
## Multiple R-squared: 0.502, Adjusted R-squared: 0.4937
## F-statistic: 59.98 on 6 and 357 DF, p-value: < 2.2e-16
```

Stepwise Selection

##

##

Min

1Q Median

-2.8194 -0.8018 -0.1737 0.6654

3Q

Stepwise Selection is a combination of forward and backward selection, where predictors are added or removed based on a criterion (e.g., AIC, BIC) until no more changes can be made.

- The stepwise selection, which combines both forward and backward methods, also identified the same set of variables: MLOGP, TPSA, SAacc, nN, RDCHI, and GATS1p.
- There is no significant difference between the AIC and BIC results in this specific case.

Max

4.8981

```
# With AIC
model.stepwise.aic <- stepAIC(null.model, scope = list(lower = null.model, upper = full.model), directi
summary(model.stepwise.aic)

##
## Call:
## lm(formula = LC50 ~ MLOGP + TPSA + SAacc + nN + RDCHI + GATS1p,
## data = train)
##
## Residuals:</pre>
```

```
## Coefficients:
            Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 2.653540 0.285743 9.286 < 2e-16 ***
## MLOGP
            0.404067 0.078544 5.144 4.44e-07 ***
            0.027138 0.003284
## TPSA
                             8.265 2.78e-15 ***
            ## SAacc
            ## nN
            0.639082 0.174662
                             3.659 0.000291 ***
## RDCHI
## GATS1p
            -0.589921 0.183821 -3.209 0.001452 **
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.216 on 357 degrees of freedom
## Multiple R-squared: 0.502, Adjusted R-squared: 0.4937
## F-statistic: 59.98 on 6 and 357 DF, p-value: < 2.2e-16
# With BIC
model.stepwise.bic <- stepAIC(null.model, scope = list(lower = null.model, upper = full.model), directi
summary(model.stepwise.bic)
##
## lm(formula = LC50 ~ MLOGP + TPSA + SAacc + nN + RDCHI + GATS1p,
##
     data = train)
##
## Residuals:
     Min
             1Q Median
                          ЗQ
## -2.8194 -0.8018 -0.1737 0.6654 4.8981
## Coefficients:
            Estimate Std. Error t value Pr(>|t|)
## (Intercept) 2.653540 0.285743 9.286 < 2e-16 ***
## MLOGP
            0.404067 0.078544 5.144 4.44e-07 ***
            ## TPSA
## SAacc
            ## nN
           ## RDCHI
            0.639082 0.174662 3.659 0.000291 ***
## GATS1p
            ## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.216 on 357 degrees of freedom
## Multiple R-squared: 0.502, Adjusted R-squared: 0.4937
## F-statistic: 59.98 on 6 and 357 DF, p-value: < 2.2e-16
Model Comparison
```

```
# Predict on the test set using all models
test$pred_backward_aic <- predict(model.backward.aic, newdata = test)</pre>
test$pred_forward_aic <- predict(model.forward.aic, newdata = test)</pre>
test$pred stepwise aic <- predict(model.stepwise.aic, newdata = test)
test$pred_backward_bic <- predict(model.backward.bic, newdata = test)</pre>
```

```
test$pred_forward_bic <- predict(model.forward.bic, newdata = test)</pre>
test$pred_stepwise_bic <- predict(model.stepwise.bic, newdata = test)</pre>
# Calculate MSE, RMSE, and R-squared for each model
mse <- function(actual, predicted) mean((actual - predicted)^2)</pre>
rmse <- function(actual, predicted) sqrt(mse(actual, predicted))</pre>
r2 <- function(actual, predicted) 1 - (sum((actual - predicted)^2) / sum((actual - mean(actual))^2))
metrics <- data.frame(</pre>
  Model = c("Backward AIC", "Forward AIC", "Stepwise AIC", "Backward BIC", "Forward BIC", "Stepwise BIC
  MSE = c(
    mse(test$LC50, test$pred_backward_aic),
    mse(test$LC50, test$pred_forward_aic),
    mse(test$LC50, test$pred_stepwise_aic),
    mse(test$LC50, test$pred_backward_bic),
    mse(test$LC50, test$pred_forward_bic),
    mse(test$LC50, test$pred_stepwise_bic)
  ),
  RMSE = c(
    rmse(test$LC50, test$pred_backward_aic),
    rmse(test$LC50, test$pred_forward_aic),
    rmse(test$LC50, test$pred_stepwise_aic),
    rmse(test$LC50, test$pred_backward_bic),
    rmse(test$LC50, test$pred_forward_bic),
    rmse(test$LC50, test$pred stepwise bic)
  ),
  R2 = c(
    r2(test$LC50, test$pred_backward_aic),
    r2(test$LC50, test$pred_forward_aic),
    r2(test$LC50, test$pred_stepwise_aic),
    r2(test$LC50, test$pred_backward_bic),
    r2(test$LC50, test$pred_forward_bic),
    r2(test$LC50, test$pred_stepwise_bic)
  )
print(metrics)
##
            Model
                        MSE
                                RMSE
                                            R.2
## 1 Backward AIC 1.398176 1.182445 0.4352328
## 2 Forward AIC 1.398176 1.182445 0.4352328
## 3 Stepwise AIC 1.398176 1.182445 0.4352328
```

In conclusion, the variable selection procedures (forward, backward, and stepwise) consistently identified the same set of predictors: MLOGP, TPSA, SAacc, nN, RDCHI, and GATS1p.

d. Ridge Regression

4 Backward BIC 1.398176 1.182445 0.4352328 ## 5 Forward BIC 1.398176 1.182445 0.4352328 ## 6 Stepwise BIC 1.398176 1.182445 0.4352328

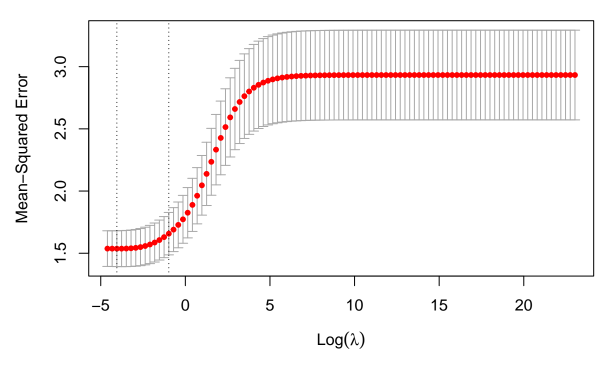
```
# Split the data into training (2/3) and test (1/3) sets
set.seed(123)
sample <- sample.split(data$LC50, SplitRatio = 2/3)
train <- subset(data, sample == TRUE)
test <- subset(data, sample == FALSE)

# Set up the training and test data
x_train <- as.matrix(train[, -9])
y_train <- train$LC50
x_test <- as.matrix(test[, -9])
y_test <- test$LC50</pre>
```

Cross Validation

```
# Reference: https://bookdown.org/ssjackson300/Machine-Learning-Lecture-Notes/choosing-lambda.html
# Define a grid of lambda values
lambda_grid <- 10^seq(10, -2, length = 100)
# Perform cross-validation for ridge regression
cv_ridge <- cv.glmnet(x_train,</pre>
                      y_train,
                      alpha = 0,
                      lambda = lambda_grid,
                       standardize = TRUE
best lambda cv <- cv ridge$lambda.min
print(paste("Best Lambda from Cross-Validation:", best_lambda_cv))
## [1] "Best Lambda from Cross-Validation: 0.0174752840000768"
# Predict and evaluate on test data
ridge_pred_cv <- predict(cv_ridge, s = best_lambda_cv, newx = x_test)</pre>
mse_cv <- mean((ridge_pred_cv - y_test)^2)</pre>
rmse cv <- sqrt(mse cv)</pre>
r2_cv <- 1 - (sum((ridge_pred_cv - y_test)^2) / sum((y_test - mean(y_test))^2))
cat(paste0(
 "MSE: ", mse_cv, "\n",
  "RMSE (Test): ", rmse_cv, "\n",
  "R-squared (Test): ", r2_cv, "\n"
))
## MSE: 1.39984196703498
## RMSE (Test): 1.18314917361885
## R-squared (Test): 0.434559904102569
plot(cv_ridge)
```





Bootstrap Procedure

```
# Reference: https://pages.stat.wisc.edu/~kdlevin/teaching/Fall2022/STAT340/lecs/L13_bootstrap.html
# Define ridge regression function for bootstrap
ridge_bootstrap <- function(data, lambda, B = 100) {</pre>
  n <- nrow(data) # number of observations</pre>
  boot_mses <- numeric(B)</pre>
  for (i in 1:B) {
    resample_indices <- sample(1:n, n, replace = TRUE)</pre>
    # resampled_data <- fin_pairs[resample_indices,] fin_pairs = [X, Y]</pre>
    resampled_data <- data[resample_indices, ]</pre>
    x_boot <- as.matrix(resampled_data[, -9])</pre>
    y_boot <- resampled_data$LC50</pre>
    # Apply ridge regression and predict in this resampling data set
    ridge_model <- glmnet(x_boot, y_boot, alpha = 0, lambda = lambda, standardize = TRUE)</pre>
    boot_pred <- predict(ridge_model, s = lambda, newx = as.matrix(data[, -9]))</pre>
    boot_mses[i] <- mean((boot_pred - data$LC50)^2)</pre>
  }
```

```
return(mean(boot_mses))
}

# Perform bootstrap for ridge regression
set.seed(1)
boot_results <- sapply(lambda_grid, function(lambda) {
   ridge_bootstrap(train, lambda, B = 100)
})

# Find the optimal lambda
best_lambda_bootstrap <- lambda_grid[which.min(boot_results)]
print(paste("Best_Lambda_from_Bootstrap:", best_lambda_bootstrap))</pre>
```

[1] "Best Lambda from Bootstrap: 0.0132194114846603"

```
# Predict and evaluate on test data
ridge_pred_bootstrap <- predict(cv_ridge, s = best_lambda_bootstrap, newx = x_test)
mse_bootstrap <- mean((ridge_pred_bootstrap - y_test)^2)
rmse_bootstrap <- sqrt(mse_bootstrap)
r2_bootstrap <- 1 - (sum((ridge_pred_bootstrap - y_test)^2) / sum((y_test - mean(y_test))^2))
cat(paste0(
    "MSE: ", mse_bootstrap, "\n",
    "RMSE (Test): ", rmse_bootstrap, "\n",
    "R-squared (Test): ", r2_bootstrap, "\n"
))

## MSE: 1.39995494236231
## RMSE (Test): 1.18319691613962
## R-squared (Test): 0.434514269822825</pre>
```

Cross Validation vs Bootstrap Comparision

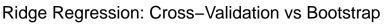
Using both cross-validation and bootstrap procedures, the optimal complexity parameter (lambda) was found to be approximately 0.017 using cross-validation and 0.013 using bootstrap. The performance of the ridge regression was very similar under both methods, with test set MSE around 1.40 and R-squared about 0.43, slightly better than the dummy encoded linear regression but on par with the original linear model.

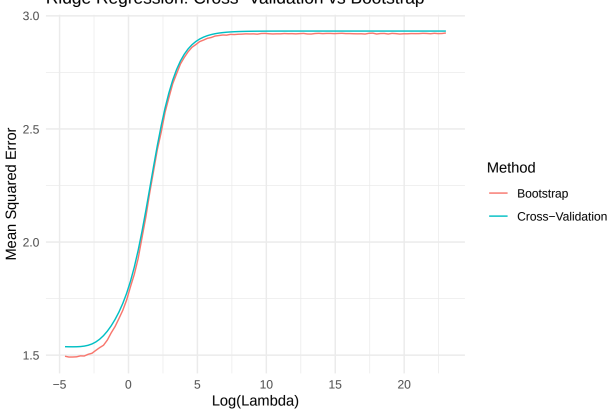
In the plot below, we observed that the both cross-validation and bootstrap should provide similar lambda values, but cross-validation typically has less variance in error estimates compared to bootstrap.

```
# Create comparison data frame
comparison_df <- data.frame(
   Lambda = rep(lambda_grid, 2),
   MSE = c(cv_ridge$cvm, boot_results),
   Method = rep(c("Cross-Validation", "Bootstrap"), each = length(lambda_grid))
)

# Plot the results
ggplot(comparison_df, aes(x = log(Lambda), y = MSE, color = Method)) +
   geom_line() +
   labs(title = "Ridge Regression: Cross-Validation vs Bootstrap",</pre>
```

```
x = "Log(Lambda)",
y = "Mean Squared Error") +
theme_minimal()
```



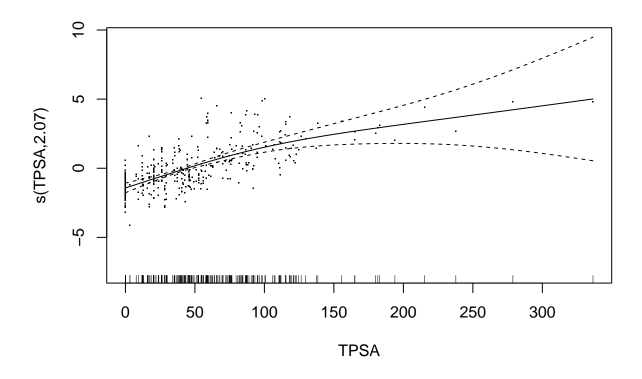


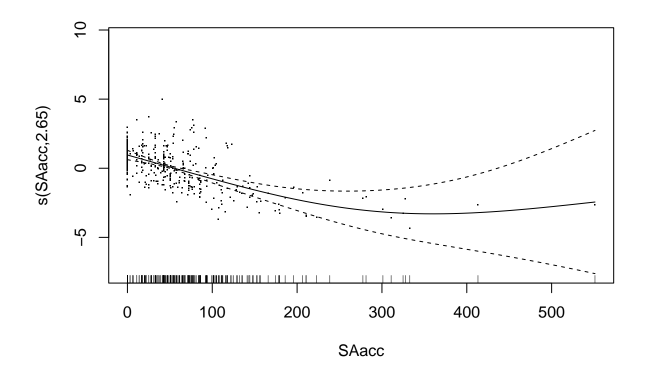
e. Generalised Additive Model (GAM)

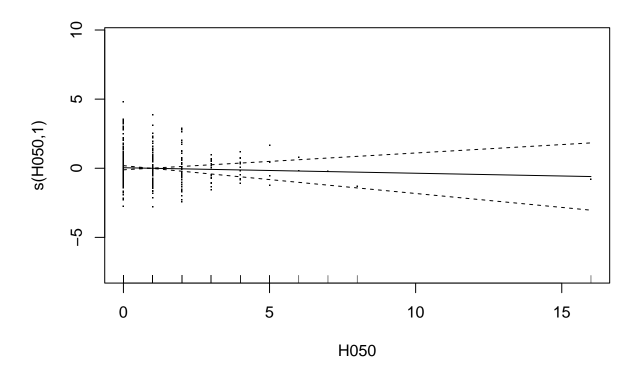
summary(train)

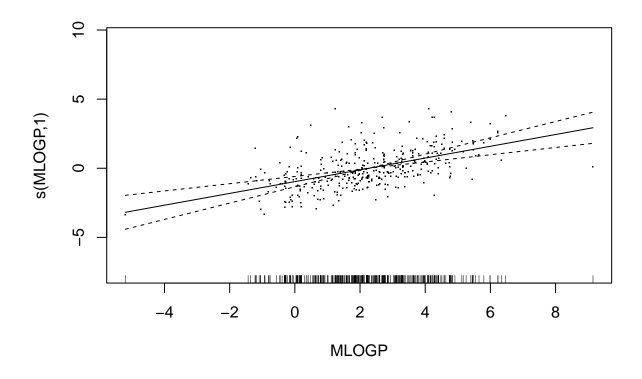
##	TPSA	SAacc	Н050	MLOGP
##	Min. : 0.00	Min. : 0.00	Min. : 0.0000	Min. :-5.199
##	1st Qu.: 16.05	1st Qu.: 13.13	1st Qu.: 0.0000	1st Qu.: 1.139
##	Median : 40.46	Median : 42.92	Median : 0.0000	Median : 2.226
##	Mean : 48.02	Mean : 58.75	Mean : 0.9313	Mean : 2.273
##	3rd Qu.: 70.14	3rd Qu.: 78.20	3rd Qu.: 1.0000	3rd Qu.: 3.455
##	Max. :336.43	Max. :551.10	Max. :16.0000	Max. : 9.148
##	RDCHI	GATS1p	\mathtt{nN}	C040
##	Min. :1.000	Min. :0.2880	Min. : 0.000	Min. : 0.0000
##	1st Qu.:1.946	1st Qu.:0.7578	1st Qu.: 0.000	1st Qu.: 0.0000
##	Median :2.329	Median :1.0485	Median : 1.000	Median : 0.0000
##	Mean :2.469	Mean :1.0682	Mean : 1.025	Mean : 0.3654
##	3rd Qu.:2.913	3rd Qu.:1.2902	3rd Qu.: 2.000	3rd Qu.: 0.0000
##	Max. :6.439	Max. :2.3530	Max. :11.000	Max. :11.0000
##	LC50			

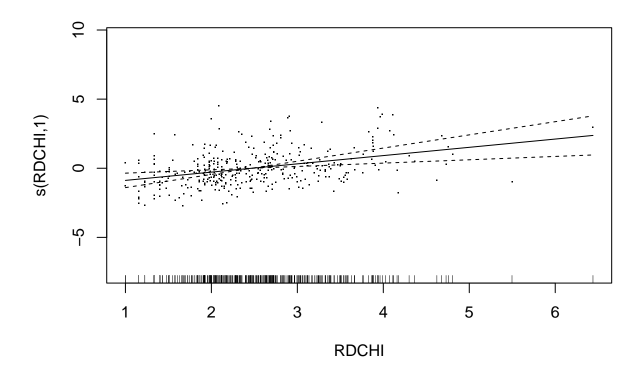
```
## Min. : 0.122
## 1st Qu.: 3.603
## Median : 4.516
## Mean
                      : 4.666
## 3rd Qu.: 5.637
## Max. :10.047
Lower Complexity (k = 4)
# Fit GAM with smoothing splines (lower complexity)
gam_model_1 \leftarrow gam(LC50 \sim s(TPSA, k = 4) + s(SAacc, k = 4) + s(HO50, k = 
                                                  s(MLOGP, k = 4) + s(RDCHI, k = 4) + s(GATS1p, k = 4) +
                                                  s(nN, k = 4) + s(CO40, k = 4), data = train)
# Summarize models
summary(gam_model_1)
## Family: gaussian
## Link function: identity
## Formula:
## LC50 ~ s(TPSA, k = 4) + s(SAacc, k = 4) + s(HO50, k = 4) + s(MLOGP,
                k = 4) + s(RDCHI, k = 4) + s(GATS1p, k = 4) + s(nN, k = 4) +
                s(C040, k = 4)
##
##
## Parametric coefficients:
                                 Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.66605
                                                        0.06325 73.77 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Approximate significance of smooth terms:
##
                                                                       F p-value
                                  edf Ref.df
## s(TPSA)
                             2.070 2.398 29.935 < 2e-16 ***
## s(SAacc) 2.653 2.839 13.121 1.57e-07 ***
                            1.000 1.000 0.243 0.622164
## s(H050)
## s(MLOGP) 1.000 1.000 26.936 6.42e-07 ***
## s(RDCHI) 1.000 1.000 11.284 0.000867 ***
## s(GATS1p) 1.000 1.000 8.847 0.003138 **
                          1.000 1.000 8.832 0.003164 **
## s(nN)
## s(CO40)
                             1.000 1.000 0.216 0.642396
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
##
## R-sq.(adj) = 0.502 Deviance explained = 51.6\%
## GCV = 1.5049 Scale est. = 1.4564
                                                                                           n = 364
\# reference: https://stackoverflow.com/questions/67077306/plotting-output-of-gam-model
p_obj <- plot(gam_model_1, residuals = TRUE)</pre>
```

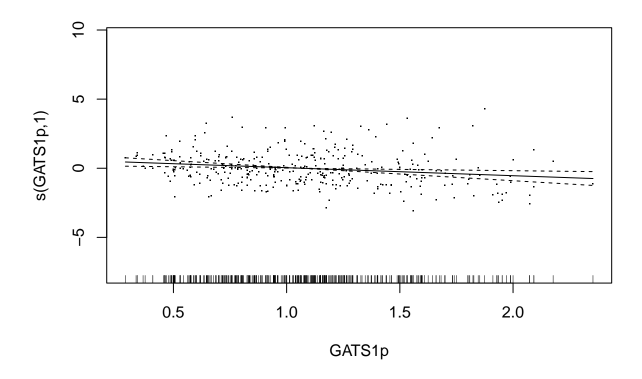


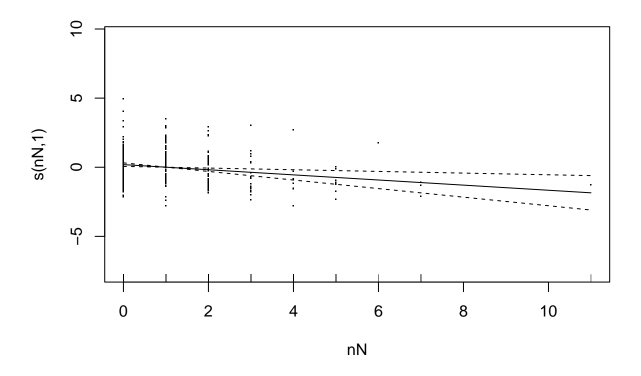


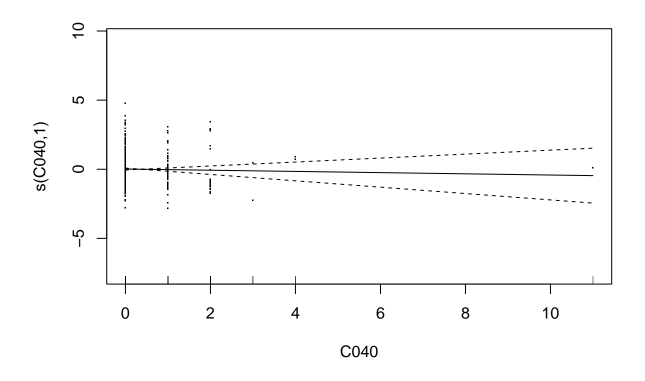


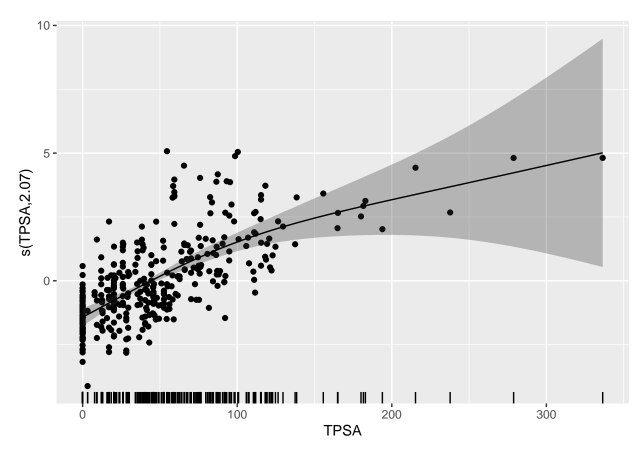












```
gam_pred_1 <- predict(gam_model_1, newdata = test)
gam_mse_1 <- mean((gam_pred_1 - y_test)^2)
gam_rmse_1 <- sqrt(gam_mse_1)
gam_r2_1 <- 1 - (sum((gam_pred_1 - y_test)^2) / sum((y_test - mean(y_test))^2))

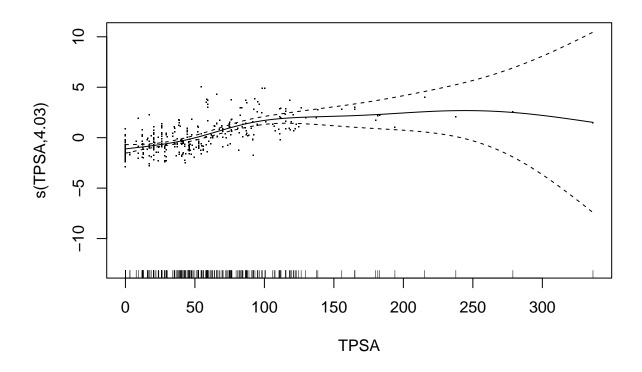
cat(paste0(
    "MSE: ", gam_mse_1, "\n",
    "RMSE (Test): ", gam_rmse_1, "\n",
    "R-squared (Test): ", gam_r2_1, "\n"
))

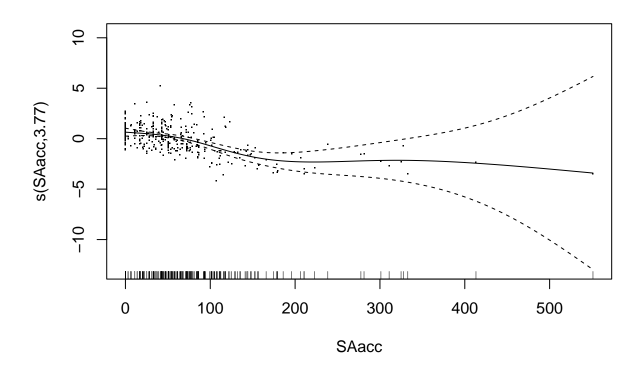
## MSE: 1.40582163542459
## RMSE (Test): 1.18567349444296</pre>
```

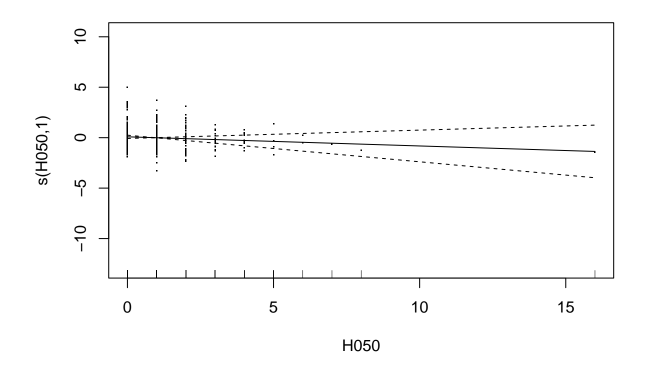
Higher Complexity (k = 6)

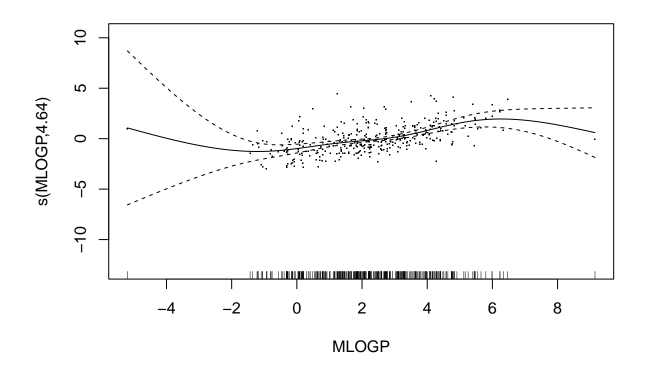
R-squared (Test): 0.432144528404967

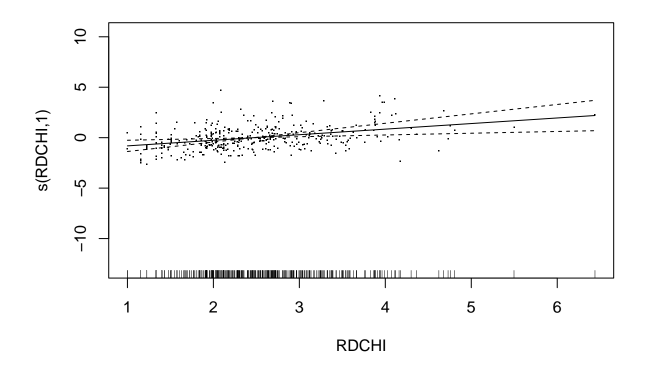
```
##
## Family: gaussian
## Link function: identity
##
## Formula:
## LC50 ~ s(TPSA, k = 6) + s(SAacc, k = 6) + s(HO50, k = 6) + s(MLOGP,
      k = 6) + s(RDCHI, k = 6) + s(GATS1p, k = 6) + s(nN, k = 6) +
      s(C040, k = 6)
##
##
## Parametric coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.66605
                         0.06113 76.33 <2e-16 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Approximate significance of smooth terms:
##
              edf Ref.df
                              F p-value
## s(TPSA)
            4.026 4.425 18.481 < 2e-16 ***
## s(SAacc) 3.775 4.260 9.769 5.29e-07 ***
            1.000 1.000 1.095 0.29606
## s(H050)
## s(MLOGP) 4.643 4.865 7.208 2.61e-06 ***
## s(RDCHI) 1.000 1.000 8.520 0.00375 **
## s(GATS1p) 1.871 2.363 3.643 0.01718 *
## s(nN)
           3.721 4.295 3.602 0.00821 **
## s(CO40)
           1.000 1.000 2.129 0.14545
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) = 0.534 Deviance explained = 56.1\%
## GCV = 1.4478 Scale est. = 1.3601
p_obj_2 <- plot(gam_model_2, residuals = TRUE)</pre>
```

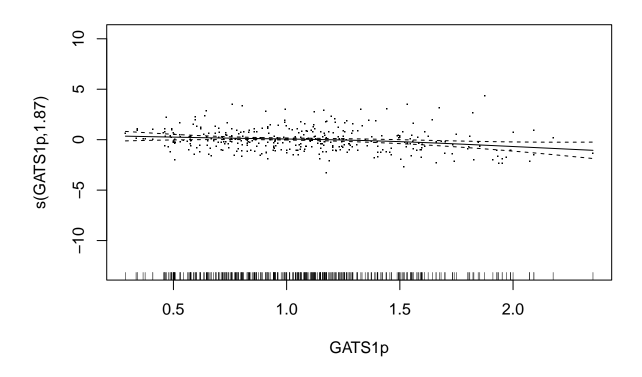


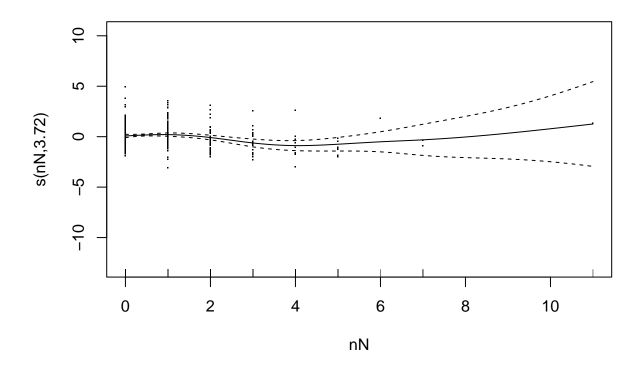


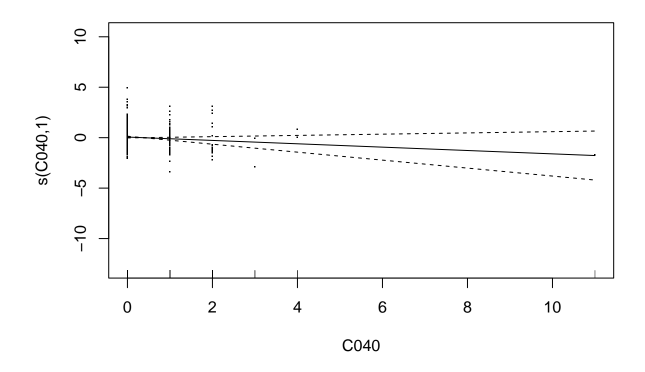


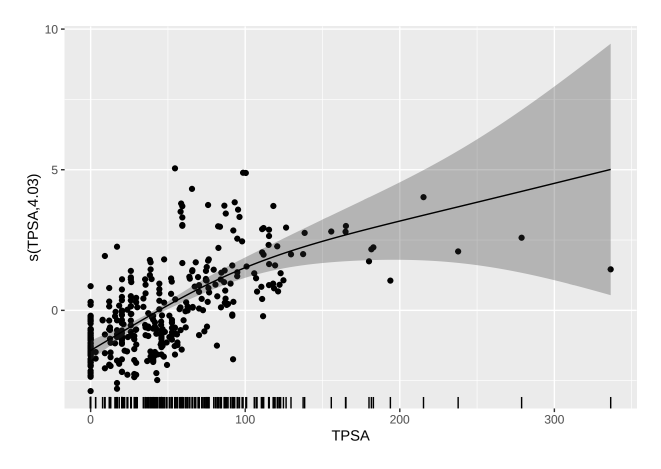












```
gam_pred_2 <- predict(gam_model_2, newdata = test)
gam_mse_2 <- mean((gam_pred_2 - y_test)^2)
gam_rmse_2 <- sqrt(gam_mse_2)
gam_r2_2 <- 1 - (sum((gam_pred_2 - y_test)^2) / sum((y_test - mean(y_test))^2))

cat(paste0(
    "MSE: ", gam_mse_2, "\n",
    "RMSE (Test): ", gam_rmse_2, "\n",
    "R-squared (Test): ", gam_r2_2, "\n"
))</pre>
```

```
## MSE: 1.3585368150292
## RMSE (Test): 1.16556287476446
## R-squared (Test): 0.451244351105306
```

In conclusion, fitting GAM models with different levels of smoothing complexity (k=4 and k=6) showed that increasing complexity slightly improved the fit, with R^2 improving from 0.43 to 0.45 and RMSE decreasing from 1.19 to 1.17. However, the benefit of increasing complexity is relatively minor, suggesting that moderate smoothing (k=4) suffices for this problem without introducing too much overfitting.

f. Regression Tree with Cost-Complexity Pruning

The complexity parameter (CP) table below shows the relative error, cross-validated error, and standard deviation for each tree size, helping to determine the best trade-off between bias and variance. The final

pruned tree has about 26 splits, meaning that it captures sufficient detail to provide accurate predictions without being overly complex.

```
# Fit a regression tree model
tree_model <- rpart(LC50 ~ ., data = train, method = "anova", control = rpart.control(cp = 0.001))
printcp(tree_model) # Display the cost complexity pruning table
##
## Regression tree:
## rpart(formula = LC50 ~ ., data = train, method = "anova", control = rpart.control(cp = 0.001))
## Variables actually used in tree construction:
## [1] CO40
              GATS1p H050
                            MLOGP RDCHI SAacc TPSA
##
## Root node error: 1060.6/364 = 2.9138
##
## n= 364
##
##
             CP nsplit rel error xerror
                                             xstd
## 1
     0.2198608
                     0
                         1.00000 1.00582 0.083377
     0.1015513
                         0.78014 0.85037 0.073595
## 2
                     1
     0.0470255
## 3
                     2
                        0.67859 0.73790 0.061890
## 4 0.0384187
                     3
                        0.63156 0.67470 0.059262
## 5
     0.0282925
                     6
                         0.51631 0.66571 0.059021
## 6
     0.0225187
                     7
                         0.48801 0.66047 0.058312
## 7
     0.0132815
                     8
                         0.46550 0.65154 0.056470
                    10
                         0.43893 0.60906 0.054135
## 8 0.0109658
## 9 0.0094706
                    11
                         0.42797 0.60140 0.053811
## 10 0.0086901
                    14
                         0.39955 0.59657 0.052692
## 11 0.0066615
                    15
                         0.39086 0.61671 0.053956
## 12 0.0063800
                    16
                         0.38420 0.63467 0.055231
## 13 0.0039482
                         0.37782 0.63634 0.058081
                    17
## 14 0.0038386
                    18
                         0.37387 0.62956 0.057522
## 15 0.0031083
                    19
                         0.37004 0.63425 0.057628
## 16 0.0027054
                    20
                         0.36693 0.63142 0.057532
## 17 0.0021546
                    21
                         0.36422 0.63076 0.058458
## 18 0.0019759
                    22
                         0.36207 0.62731 0.058474
## 19 0.0014224
                    23
                         0.36009 0.62838 0.058471
## 20 0.0014058
                         0.35867 0.63024 0.058550
                    24
## 21 0.0010000
                         0.35586 0.62934 0.058560
                    26
# Prune the tree
optimal cp <- tree model cptable [which.min(tree model cptable [, "xerror"]), "CP"]
pruned_tree <- prune(tree_model, cp = optimal_cp)</pre>
```

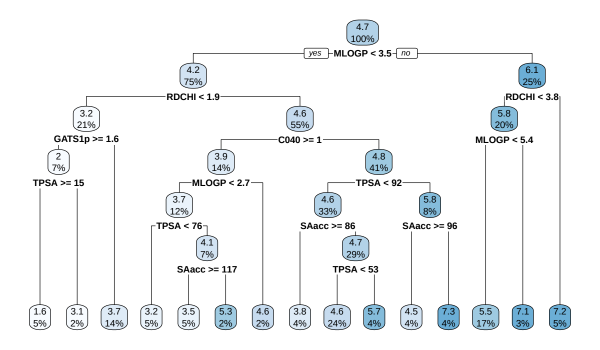
The root node splits based on MLOGP (lipophilicity), which is the most important predictor for determining the LC50 value. The tree continues to split based on other descriptors, such as:

- RDCHI (topological index), which also be the first split in the tree
- C040 (the number of certain carbon atoms)
- TPSA (polar surface area)

The terminal nodes represent the predicted LC50 values for the corresponding groups of observations.

```
# Visualize the tree
rpart.plot(pruned_tree, main = "Pruned Regression Tree")
```

Pruned Regression Tree



```
# Predict and evaluate on test data
tree_pred <- predict(pruned_tree, newdata = test)
tree_mse <- mean((tree_pred - y_test)^2)
tree_rmse <- sqrt(tree_mse)
tree_r2 <- 1 - (sum((tree_pred - y_test)^2) / sum((y_test - mean(y_test))^2))

cat(paste0(
   "MSE: ", tree_mse, "\n",
   "RMSE (Test): ", tree_rmse, "\n",
   "R-squared (Test): ", tree_r2, "\n"
))</pre>
```

MSE: 1.55285972999018 ## RMSE (Test): 1.24613792574906 ## R-squared (Test): 0.372751228125619

However, the performance of the tree, with an R-squared of around 0.37 on the test set, suggests that the tree structure, while interpretable, may not capture the relationships in the data as effectively as other methods.

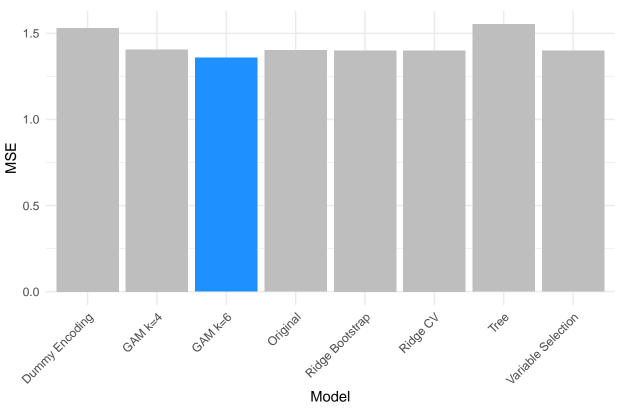
g. Compare all the models implemented

[TBD]

- Linear Regression (Original)
- Linear Regression model by Dummy Encoding Method (Dummy Encoding)
- Ridge Regression with Cross Validation (Ridge CV)
- Ridge Regression with Bootstrapping ("Ridge Bootstrap")
- Generalized Additive Model with lower complexity (GAM k=4)
- Generalized Additive Model with higher complexity (GAM k=6)
- Regression Tree with Cost-Complexity Pruning (Tree)

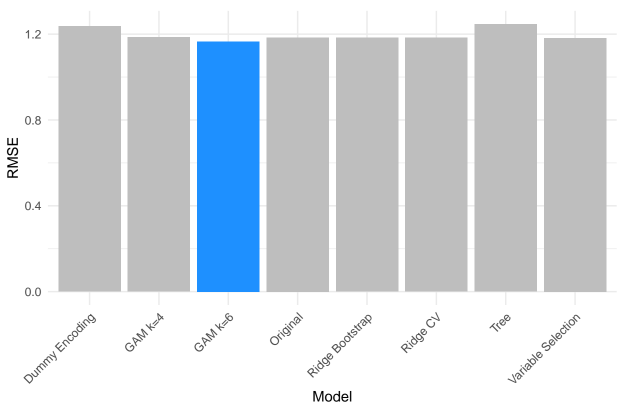
```
# Create a comprehensive metrics data frame
all_models_metrics <- data.frame(</pre>
  Model = c("Original", "Dummy Encoding", "Variable Selection", "Ridge CV", "Ridge Bootstrap", "GAM k=4
  MSE = c(mse_test, mse_test_transform_dummy, metrics$MSE[metrics$Model == "Forward AIC"], mse_cv, mse_
          gam_mse_2, tree_mse),
  RMSE = c(rmse_test, rmse_test_transform_dummy, metrics$RMSE[metrics$Model == "Forward AIC"], rmse_cv,
           gam_rmse_2, tree_rmse),
  R2 = c(r2_test, r2_test_transform_dummy, metrics$R2[metrics$Model == "Forward AIC"], r2_cv, r2_bootst
         gam_r2_2, tree_r2)
# Print the all models metrics
print(all_models_metrics)
##
                  Model
                             MSE
                                     RMSE
                                                 R.2
## 1
               Original 1.402249 1.184166 0.4335877
## 2
         Dummy Encoding 1.530438 1.237109 0.3818079
## 3 Variable Selection 1.398176 1.182445 0.4352328
## 4
               Ridge CV 1.399842 1.183149 0.4345599
## 5
        Ridge Bootstrap 1.399955 1.183197 0.4345143
                GAM k=4 1.405822 1.185673 0.4321445
## 6
## 7
                GAM k=6 1.358537 1.165563 0.4512444
                   Tree 1.552860 1.246138 0.3727512
## 8
## Visualization of Model Comparisons
# Identify the model with the minimum MSE, RMSE and maximum R-squared
best_mse_model <- all_models_metrics$Model[which.min(all_models_metrics$MSE)]
best_rmse_model <- all_models_metrics$Model[which.min(all_models_metrics$RMSE)]
best_r2_model <- all_models_metrics$Model[which.max(all_models_metrics$R2)]
# Plot MSE across selected models
ggplot(all_models_metrics, aes(x = Model, y = MSE, fill = Model == best_mse_model)) +
  geom_bar(stat = "identity") +
  scale_fill_manual(values = c("gray", "dodgerblue"), guide = "none") +
  theme_minimal() +
  labs(title = "MSE Across Selected Models", x = "Model", y = "MSE", fill = "Best Model") +
  theme(axis.text.x = element_text(angle = 45, hjust = 1))
```



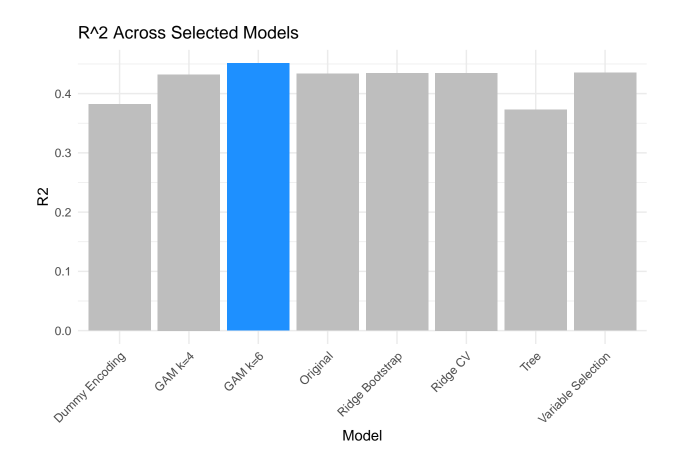


```
# Plot RMSE across selected models
ggplot(all_models_metrics, aes(x = Model, y = RMSE, fill = Model == best_mse_model)) +
  geom_bar(stat = "identity") +
  scale_fill_manual(values = c("gray", "dodgerblue"), guide = "none") +
  theme_minimal() +
  labs(title = "RMSE Across Selected Models", x = "Model", y = "RMSE", fill = "Best Model") +
  theme(axis.text.x = element_text(angle = 45, hjust = 1))
```





```
# Plot RMSE across selected models
ggplot(all_models_metrics, aes(x = Model, y = R2, fill = Model == best_mse_model)) +
   geom_bar(stat = "identity") +
   scale_fill_manual(values = c("gray", "dodgerblue"), guide = "none") +
   theme_minimal() +
   labs(title = "R^2 Across Selected Models", x = "Model", y = "R2", fill = "Best Model") +
   theme(axis.text.x = element_text(angle = 45, hjust = 1))
```



Problem 2. Classification

```
library(mlbench)
data("PimaIndiansDiabetes2")
data <- PimaIndiansDiabetes2</pre>
head(data)
    pregnant glucose pressure triceps insulin mass pedigree age diabetes
##
           6
                  148
                            72
                                    35
                                            NA 33.6
                                                       0.627 50
                  85
                                    29
                                            NA 26.6
## 2
            1
                            66
                                                       0.351 31
                                                                       neg
## 3
            8
                  183
                            64
                                    NA
                                            NA 23.3
                                                       0.672 32
                                                                       pos
## 4
            1
                   89
                            66
                                    23
                                            94 28.1
                                                       0.167 21
                                                                       neg
## 5
            0
                  137
                            40
                                    35
                                           168 43.1
                                                       2.288 33
                                                                       pos
## 6
            5
                            74
                                            NA 25.6
                  116
                                    NA
                                                       0.201 30
                                                                       neg
# Checking missing value
sapply(data, function(x) sum(is.na(x)))
## pregnant glucose pressure triceps insulin
                                                    mass pedigree
                                                                        age
##
                   5
                           35
                                   227
                                            374
                                                      11
                                                                          0
## diabetes
##
```

```
# Remove rows with missing values
data <- na.omit(data)</pre>
head(data)
##
      pregnant glucose pressure triceps insulin mass pedigree age diabetes
                              66
## 4
                     89
                                       23
                                               94 28.1
                                                           0.167
                                                                  21
             1
                                                                           neg
## 5
             0
                    137
                                       35
                                              168 43.1
                                                           2.288
                              40
                                                                  33
                                                                           pos
## 7
             3
                                       32
                    78
                              50
                                               88 31.0
                                                           0.248
                                                                  26
                                                                           pos
             2
                                              543 30.5
## 9
                    197
                              70
                                       45
                                                           0.158
                                                                  53
                                                                           pos
## 14
                    189
                                       23
                                              846 30.1
                                                           0.398
             1
                              60
                                                                  59
                                                                           pos
## 15
                    166
                              72
                                       19
                                              175 25.8
                                                           0.587 51
                                                                           pos
summary(data)
                         glucose
                                                            triceps
##
       pregnant
                                          pressure
          : 0.000
                                             : 24.00
                                                                : 7.00
   Min.
                      Min.
                             : 56.0
                                       Min.
                                                        Min.
##
    1st Qu.: 1.000
                      1st Qu.: 99.0
                                       1st Qu.: 62.00
                                                         1st Qu.:21.00
##
   Median : 2.000
                      Median :119.0
                                       Median : 70.00
                                                        Median :29.00
##
   Mean
           : 3.301
                             :122.6
                                       Mean
                                             : 70.66
                                                         Mean
                                                                :29.15
                      Mean
    3rd Qu.: 5.000
                                       3rd Qu.: 78.00
                      3rd Qu.:143.0
                                                         3rd Qu.:37.00
##
    Max.
           :17.000
                             :198.0
                                       Max.
                                              :110.00
                                                                :63.00
                      Max.
                                                         Max.
##
       insulin
                           mass
                                          pedigree
                                                              age
                                                                          diabetes
##
   Min.
           : 14.00
                      Min.
                             :18.20
                                              :0.0850
                                                         Min.
                                                                :21.00
                                                                          neg:262
   1st Qu.: 76.75
                      1st Qu.:28.40
                                       1st Qu.:0.2697
                                                         1st Qu.:23.00
                                                                          pos:130
## Median :125.50
                      Median :33.20
                                       Median :0.4495
                                                         Median :27.00
## Mean
           :156.06
                                              :0.5230
                             :33.09
                                       Mean
                                                         Mean
                                                                :30.86
                      Mean
## 3rd Qu.:190.00
                      3rd Qu.:37.10
                                       3rd Qu.:0.6870
                                                         3rd Qu.:36.00
   Max.
           :846.00
                      Max.
                             :67.10
                                       Max.
                                              :2.4200
                                                         Max.
                                                                :81.00
# Checking how balance is with the dependent variable
```

```
# Checking how balance is with the dependent variable
prop.table(table(data$diabetes))
```

```
## neg pos
## 0.6683673 0.3316327
```

Randomly split the dataset into a training set (approximately 2/3 of the sample size) and a test set, such that the class distributions (i.e. the empirical distribution of diabetes) is similar in the two sets.

```
set.seed(123)

sample <- sample.split(data$diabetes, SplitRatio = 2/3)
train <- subset(data, sample == TRUE)
test <- subset(data, sample == FALSE)</pre>
```

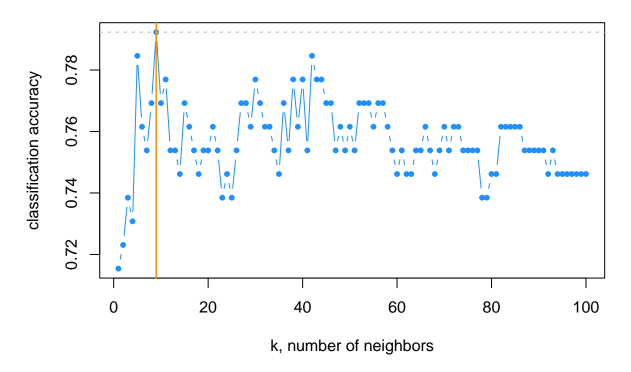
```
# Class distribution in the training set
prop.table(train$diabetes))
```

```
## neg pos
## 0.6679389 0.3320611
```

```
# Class distribution in the testing set
prop.table(table(test$diabetes))
##
##
         neg
## 0.6692308 0.3307692
cat("Dimension of Training Set:", paste(dim(train), collapse = "x"), "\nDimension of Test Set:", paste(
## Dimension of Training Set: 262x9
## Dimension of Test Set: 130x9
a. k-NN classifier
library(class)
library(caret)
## Loading required package: lattice
##
## Attaching package: 'lattice'
## The following object is masked from 'package:boot':
##
       melanoma
library(FNN)
## Warning: package 'FNN' was built under R version 4.3.3
## Attaching package: 'FNN'
## The following objects are masked from 'package:class':
##
       knn, knn.cv
##
X_train <- train[, -ncol(train)]</pre>
y_train <- train$diabetes</pre>
X_test <- test[, -ncol(test)]</pre>
y_test <- test$diabetes</pre>
```

```
accuracy = function(actual, predicted) {
  mean(actual == predicted)
}
set.seed(42)
k_to_try = 1:100
acc_k = rep(0, length(k_to_try))
# Loop over values of k
for (i in seq_along(k_to_try)) {
  pred <- knn(</pre>
   train = scale(X_train),
   test = scale(X_test),
   cl = y_train,
   k = k_{to_{try}[i]}
  acc_k[i] <- accuracy(y_test, pred)</pre>
ex_seq = seq(from = 1, to = 100, by = 5)
seq_along(ex_seq)
## [1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
ex_storage = rep(x = 0, times = length(ex_seq))
for(i in seq_along(ex_seq)) {
  ex storage[i] = mean(rnorm(n = 10, mean = ex seq[i], sd = 1))
ex_storage
## [1] 1.547297 5.836543 10.821920 15.636096 20.979785 26.018394 31.539077
## [8] 35.782125 41.251106 45.912805 51.229248 55.801428 60.205034 66.210242
## [15] 70.797793 75.754132 81.101802 86.093609 90.743936 96.187940
# Reference: https://daviddalpiaz.github.io/r4sl/k-nearest-neighbors.html
# plot accuracy vs choice of k
plot(acc_k, type = "b", col = "dodgerblue", cex = 1, pch = 20,
     xlab = "k, number of neighbors", ylab = "classification accuracy",
     main = "Accuracy vs Neighbors")
# add lines indicating k with best accuracy
abline(v = which(acc_k == max(acc_k)), col = "darkorange", lwd = 1.5)
# add line for max accuracy seen
abline(h = max(acc_k), col = "grey", lty = 2)
# add line for prevalence in test set
abline(h = mean(y_test == "No"), col = "grey", lty = 2)
```

Accuracy vs Neighbors



```
max(acc_k)
## [1] 0.7923077

max(which(acc_k == max(acc_k)))

## [1] 9

# Get K-NN result using best k
best_k <- max(which(acc_k == max(acc_k)))
set.seed(42)
knn_pred <- knn(train = scale(X_train), test = scale(X_test), cl = y_train, k = best_k)
knn_accuracy <- accuracy(y_test, knn_pred)

# Convert K-NN predicted labels to probabilities for ROC
knn_pred_prob <- ifelse(knn_pred == "pos", 1, 0)

# Calculate ROC curve for K-NN
roc_curve_knn <- roc(as.numeric(y_test == "pos"), knn_pred_prob)

## Setting levels: control = 0, case = 1</pre>
```

```
## Setting direction: controls < cases
```

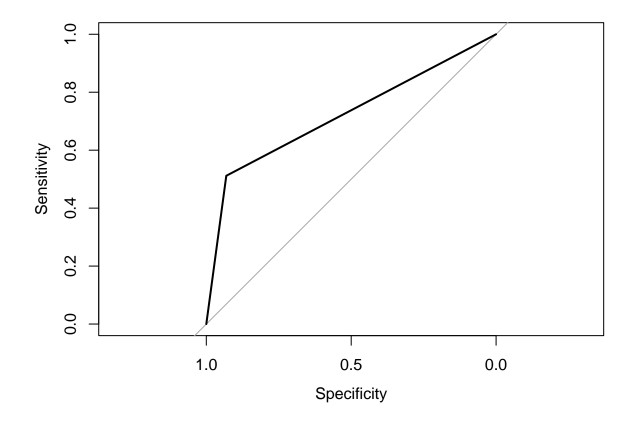
```
# Calculate and print AUC for K-NN
auc_knn <- auc(roc_curve_knn)
cat("k-NN - AUC:", auc_knn, "\n")

## k-NN - AUC: 0.7213312

cat("Accuracy:", knn_accuracy, "\n")

## Accuracy: 0.7923077

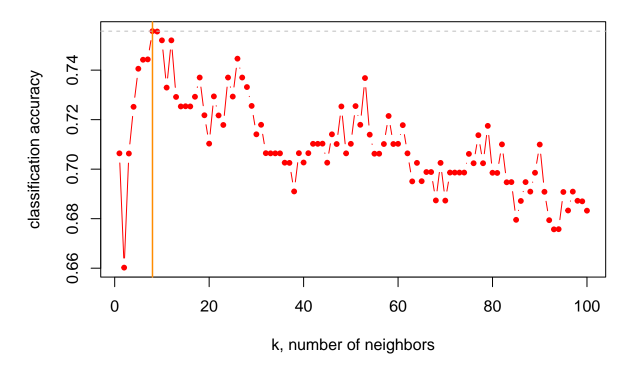
# Plot ROC
plot(roc_curve_knn)</pre>
```



Using 5-fold

```
trControl = train_control,
                   tuneGrid = expand.grid(k = k_to_try))
## plot(train_knn)
cv_results <- train_knn$results</pre>
head(cv_results)
                     Kappa AccuracySD
    k Accuracy
                                        KappaSD
## 1 1 0.7063861 0.3512222 0.05274457 0.1132618
## 2 2 0.6602322 0.2180471 0.05024751 0.1199193
## 3 3 0.7063135 0.3304328 0.05489854 0.1044619
## 4 4 0.7251814 0.3627845 0.05461723 0.1114380
## 5 5 0.7405660 0.3922546 0.07858248 0.1721106
## 6 6 0.7441945 0.3957179 0.08260038 0.2122957
# Plot the 5-fold CV accuracy vs choice of k
plot(cv_results$k, cv_results$Accuracy, type = "b", col = "red", cex = 1, pch = 20,
     xlab = "k, number of neighbors", ylab = "classification accuracy",
     main = "5-fold CV Accuracy vs Neighbors")
#legend("bottomright", legend = "5-fold CV Accuracy", col = "red", pch = 19, lty = 1)
# Add lines indicating k with best accuracy
abline(v = cv_results\$k[which.max(cv_results\$Accuracy)], col = "darkorange", lwd = 1.5)
# Add line for max accuracy seen
abline(h = max(cv_results$Accuracy), col = "grey", lty = 2)
```

5-fold CV Accuracy vs Neighbors



```
max(cv_results$Accuracy)

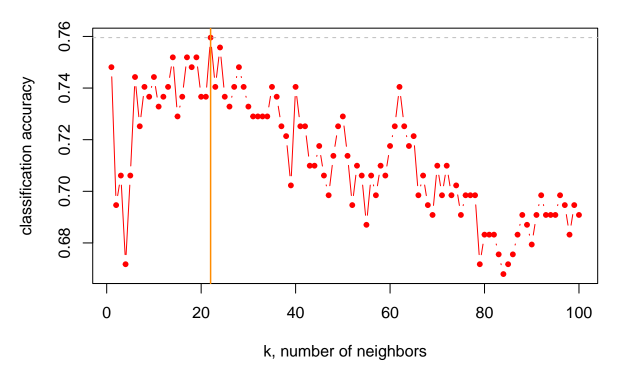
## [1] 0.7557329

cv_results$k[which.max(cv_results$Accuracy)]

## [1] 8
```

Using leave-one-out cross-validation

LOOCV Accuracy vs Neighbors



```
max(cv_results_loocv$Accuracy)
```

[1] 0.759542

k Accuracy

1 1 0.7480916 0.4353817

Kappa

```
cv_results_loocv$k[which.max(cv_results_loocv$Accuracy)]
## [1] 22
```

b. Generalized Additive Model (GAM)

```
# Fit a GAM with automatic smoothness selection
gam_model <- gam(
    diabetes ~ s(glucose) + s(pressure) + s(insulin) + s(mass) + s(pedigree) + s(age) + s(pregnant),
    data = train,
    family = binomial(link = 'logit'),
    select = TRUE,
    method= "REML")</pre>
```

In the summary(model) output, the Approximate significance of smooth terms table shows an estimated degrees of freedom (edf) and Chi square score (Chi.sq) close to zero, with a p-value > 0.05:

```
summary(gam_model)
```

```
##
## Family: binomial
## Link function: logit
##
## Formula:
## diabetes ~ s(glucose) + s(pressure) + s(insulin) + s(mass) +
##
      s(pedigree) + s(age) + s(pregnant)
##
## Parametric coefficients:
              Estimate Std. Error z value Pr(>|z|)
##
                           0.1918 -5.895 3.75e-09 ***
## (Intercept) -1.1305
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Approximate significance of smooth terms:
                    edf Ref.df Chi.sq p-value
## s(glucose) 9.761e-01
                             9 36.884 < 2e-16 ***
## s(pressure) 1.297e-05
                             9 0.000 0.832820
## s(insulin) 5.976e-06
                           9 0.000 0.697822
## s(mass)
              1.889e+00
                            9 10.598 0.001791 **
## s(pedigree) 8.059e-01
                             9 3.887 0.027410 *
## s(age)
              2.127e+00
                             9 12.199 0.000871 ***
## s(pregnant) 1.798e+00
                             9 4.837 0.047140 *
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## R-sq.(adj) = 0.421
                        Deviance explained = 39.2%
## -REML = 114.77 Scale est. = 1
```

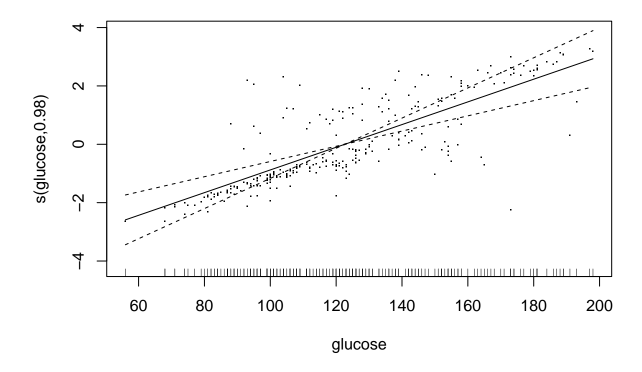
Reference: https://osf.io/wgc4f/wiki/mgcv:%20model%20selection/

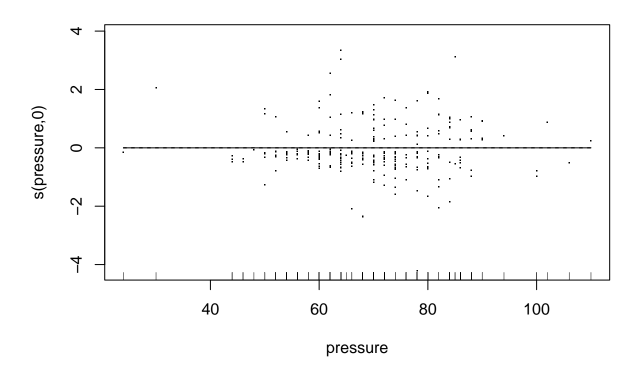
[TBD] You can use these statistics to detect a shrunk term. It is usually sufficient to check for p-values > 0.05:

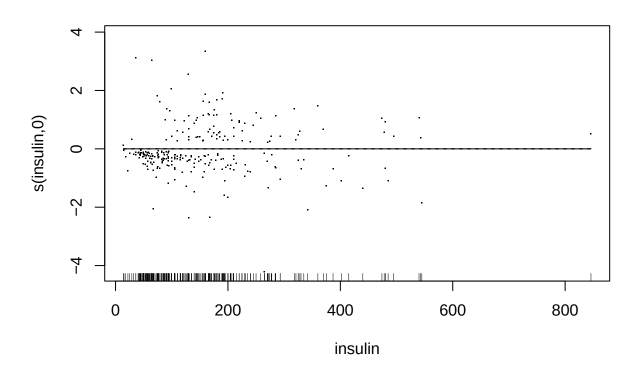
```
summary(gam_model)$s.pv
```

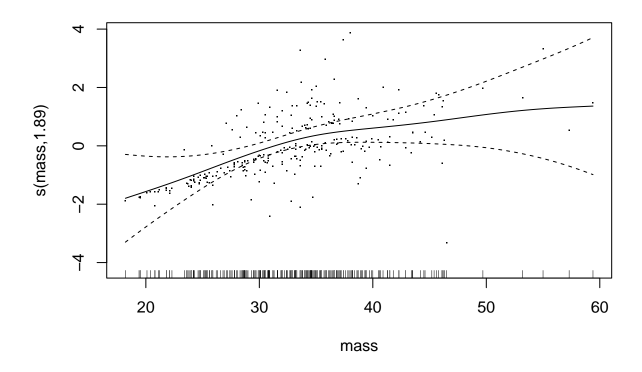
We can see that the glucose and pedigree is a shrunk term

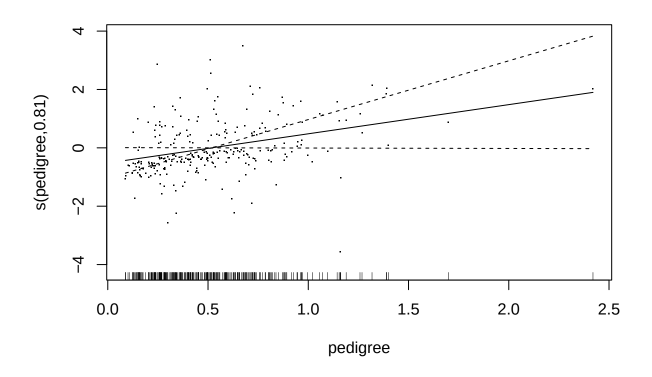
```
p_obj <- plot(gam_model, residuals = TRUE)</pre>
```

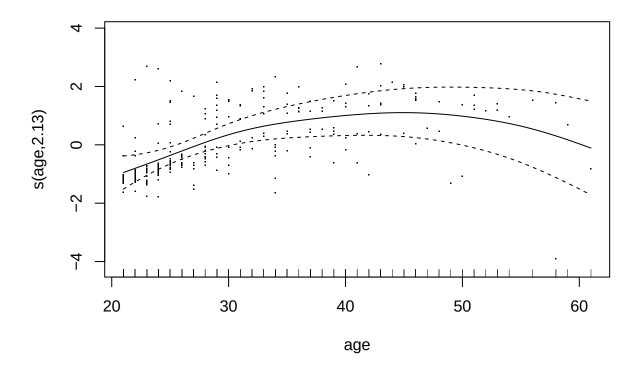


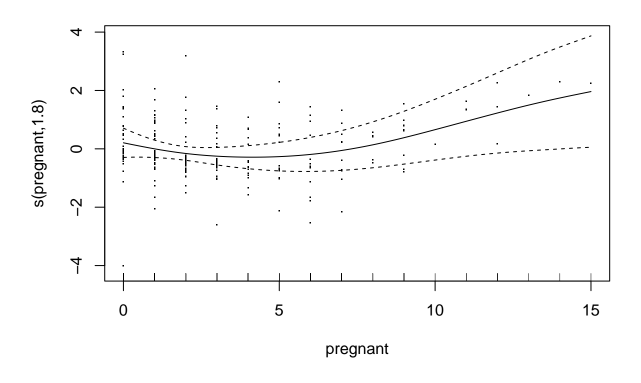


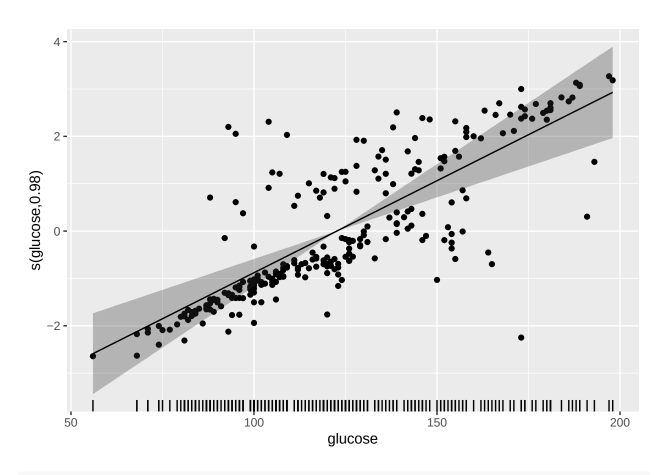












```
# Predictions and probabilities for GAM
gam_pred_prob <- predict(gam_model, newdata = test, type = "response")

# Convert probabilities to binary predictions for confusion matrix
gam_pred_class <- ifelse(gam_pred_prob > 0.5, "pos", "neg")

# Calculate accuracy for GAM
gam_accuracy <- mean(gam_pred_class == test$diabetes)
cat("GAM - Test Accuracy:", gam_accuracy, "\n")

## GAM - Test Accuracy: 0.7769231

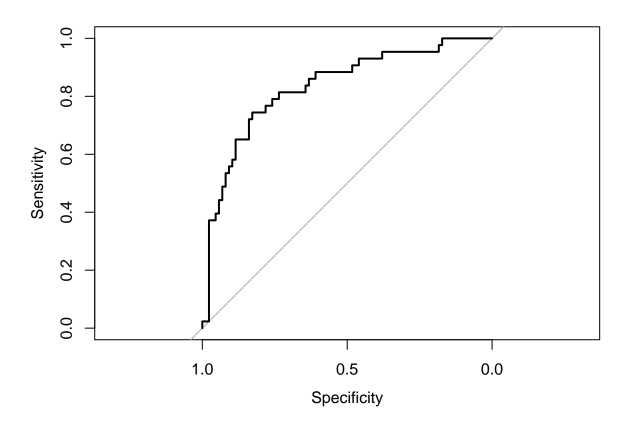
# Calculate ROC curve for GAM
roc_curve_gam <- roc(test$diabetes, gam_pred_prob)

## Setting levels: control = neg, case = pos

## Setting direction: controls < cases

# Calculate and print AUC for GAM
auc_gam <- auc(roc_curve_gam)
cat("GAM - AUC:", auc_gam, "\n")</pre>
```

```
# Plot ROC
plot(roc_curve_gam)
```

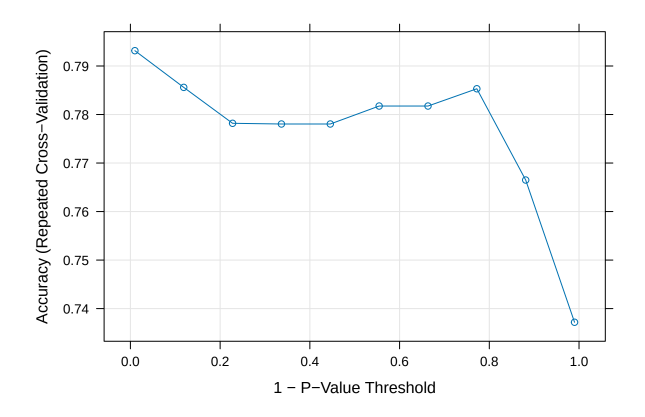


c. Tree-based methods

Setting up the k-fold cross validation k=10 cross-validation folds. Reference: https://quantdev.ssri.psu.edu/sites/qdev/files/09_EnsembleMethods_2017_1127.html

(i) Classification tree

```
## Conditional Inference Tree
##
## 262 samples
   8 predictor
##
##
    2 classes: 'neg', 'pos'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 1 times)
## Summary of sample sizes: 235, 236, 235, 236, 235, 237, ...
## Resampling results across tuning parameters:
##
##
     mincriterion Accuracy
                              Kappa
##
    0.0100000
                  0.7931567 0.5294069
##
    0.1188889
                  0.7855954 0.4963222
##
    0.2277778
                  0.7781880 0.4816163
##
     0.3366667
                  0.7780456 0.4835965
##
    0.4455556
                  0.7780456 0.4835965
##
     0.5544444
                  0.7817493 0.4930560
##
    0.6633333
                  0.7817493 0.4930560
##
     0.7722222
                  0.7853105 0.4941199
##
    0.8811111
                  0.7664957 0.4422783
##
     0.9900000
                  0.7371852 0.3930795
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mincriterion = 0.01.
```



plot(train.tree\$finalModel)

```
glucose
                                     p < 0.001
                     2
                                                            21
                              ≤ 131
                                                  > 131
                    age
                                                          glucose
                 p < 0.001
                                                           = 0.088
                                12
         3
               ≤ 28
      pedigree
                               insulin
                                                     pregnan
                                                              triceps
      മ = 0.112
                                                      p = 0.0
                                                             p = 0.718
                               = 0.036
     4 4
                                            23
                             13
                                                                    33
          > 0
  pressure
               triceps
                          glucose
                                  insulin
                                           mass
                                                                   age
              പ്പ= 0.254
                          p = 0.2 \sqrt{p} = 0.42
  p = 0.347
                                           = 0.398
                                                                 p = 0.398
             8
                        14
                                               <del>2</del>5م
                              120
                                              glucose
          glucose
                        age
                                             p = 0.859
                       = 0.948
          p = 0.198
     > 78
                                   ≤ > 191
                                                                   ≤ > 37
                                                 pressure
               119
                         > 32
                                                 p = 0.671
                                                     > 70
```

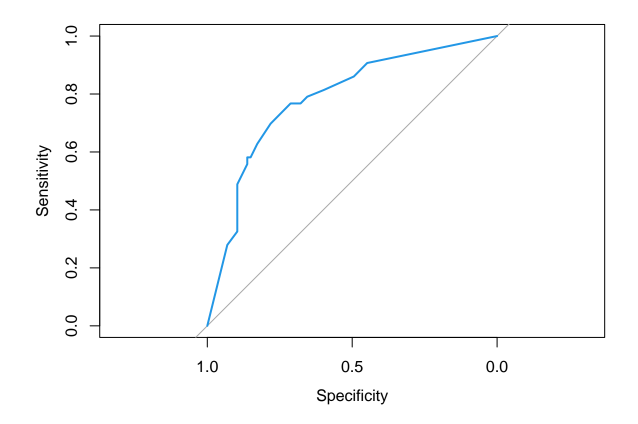
```
# obtaining class predictions for training
tree.classTrain <- predict(train.tree, type="raw")

# Check accuracy and confusion matrix for training set
confusionMatrix(train$diabetes, tree.classTrain)</pre>
### Confusion Matrix and Statistics
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction neg pos
          neg 164 11
##
          pos 24 63
##
##
##
                  Accuracy: 0.8664
                    95% CI: (0.8191, 0.9052)
##
##
       No Information Rate: 0.7176
       P-Value [Acc > NIR] : 7.327e-09
##
##
##
                     Kappa: 0.6871
##
##
   Mcnemar's Test P-Value: 0.04252
##
##
               Sensitivity: 0.8723
               Specificity: 0.8514
##
##
            Pos Pred Value: 0.9371
            Neg Pred Value: 0.7241
##
```

```
Prevalence: 0.7176
##
##
            Detection Rate: 0.6260
##
      Detection Prevalence: 0.6679
         Balanced Accuracy: 0.8618
##
##
##
          'Positive' Class : neg
##
Accuracy of training set is 86.64%
# Obtaining class predictions for test set
tree.classTest <- predict(train.tree,</pre>
                            newdata = test,
                            type="raw")
# Check accuracy and confusion matrix for training set
confusionMatrix(test$diabetes, tree.classTest)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction neg pos
          neg 74 13
##
##
          pos 18 25
##
##
                  Accuracy : 0.7615
##
                    95% CI: (0.6789, 0.8319)
       No Information Rate: 0.7077
##
       P-Value [Acc > NIR] : 0.1034
##
##
##
                     Kappa: 0.4451
##
    Mcnemar's Test P-Value: 0.4725
##
##
               Sensitivity: 0.8043
##
##
               Specificity: 0.6579
##
            Pos Pred Value: 0.8506
##
            Neg Pred Value: 0.5814
                Prevalence: 0.7077
##
##
            Detection Rate: 0.5692
##
      Detection Prevalence: 0.6692
##
         Balanced Accuracy: 0.7311
##
##
          'Positive' Class : neg
##
tree_accuracy <- mean(tree.classTest == test$diabetes)</pre>
cat("Accuracy of Classification Tree:", tree_accuracy, "\n")
```

Accuracy of Classification Tree: 0.7615385



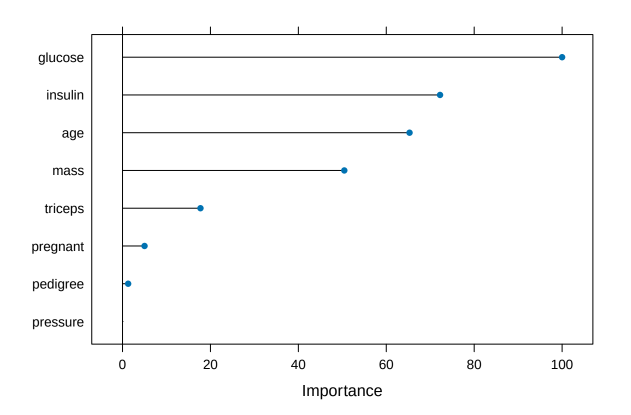
#calculate the area under curve
auc(rocCurve.tree)

Area under the curve: 0.7848

(ii) Ensemble of bagged trees

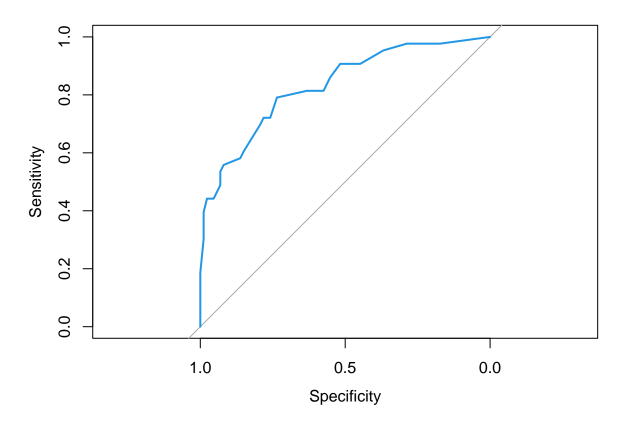
```
## Bagged CART
##
## 262 samples
    8 predictor
##
##
     2 classes: 'neg', 'pos'
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 1 times)
## Summary of sample sizes: 235, 237, 235, 236, 236, ...
## Resampling results:
##
##
     Accuracy
                Kappa
     0.7817721 0.4906951
##
```

plot(varImp(train.bagg))



```
#obtaining class predictions
bagg.classTrain <- predict(train.bagg, type="raw")</pre>
confusionMatrix(train$diabetes, bagg.classTrain)
## Confusion Matrix and Statistics
##
             Reference
##
## Prediction neg pos
##
          neg 175 0
              0 87
          pos
##
##
                  Accuracy: 1
##
                    95% CI : (0.986, 1)
##
##
       No Information Rate: 0.6679
##
       P-Value [Acc > NIR] : < 2.2e-16
##
##
                     Kappa: 1
##
##
   Mcnemar's Test P-Value : NA
##
##
               Sensitivity: 1.0000
##
               Specificity: 1.0000
##
            Pos Pred Value : 1.0000
##
            Neg Pred Value: 1.0000
##
                Prevalence: 0.6679
            Detection Rate: 0.6679
##
##
      Detection Prevalence: 0.6679
##
         Balanced Accuracy: 1.0000
##
##
          'Positive' Class : neg
##
bagg.classTest <- predict(train.bagg,</pre>
                           newdata = test,
                           type="raw")
confusionMatrix(test$diabetes, bagg.classTest)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction neg pos
          neg 74 13
##
          pos 17 26
##
##
##
                  Accuracy : 0.7692
##
                    95% CI : (0.6872, 0.8386)
##
       No Information Rate: 0.7
##
       P-Value [Acc > NIR] : 0.04933
##
##
                     Kappa: 0.4662
```

```
##
## Mcnemar's Test P-Value: 0.58388
##
##
               Sensitivity: 0.8132
##
               Specificity: 0.6667
##
            Pos Pred Value: 0.8506
##
            Neg Pred Value: 0.6047
                Prevalence : 0.7000
##
##
            Detection Rate: 0.5692
##
      Detection Prevalence : 0.6692
##
         Balanced Accuracy: 0.7399
##
##
          'Positive' Class : neg
##
bagg_accuracy <- mean(bagg.classTest == test$diabetes)</pre>
cat("Accuracy of Classification Tree:", bagg_accuracy, "\n")
## Accuracy of Classification Tree: 0.7692308
#Obtaining predicted probabilities for Test data
bagg.probs=predict(train.bagg,
                 newdata=test,
                 type="prob")
#Calculate ROC curve
rocCurve.bagg <- roc(test$diabetes, bagg.probs[,"neg"])</pre>
## Setting levels: control = neg, case = pos
## Setting direction: controls > cases
#plot the ROC curve
plot(rocCurve.bagg,col=c(4))
```



```
#calculate the area under curve (bigger is better)
auc(rocCurve.bagg)
```

Area under the curve: 0.8311

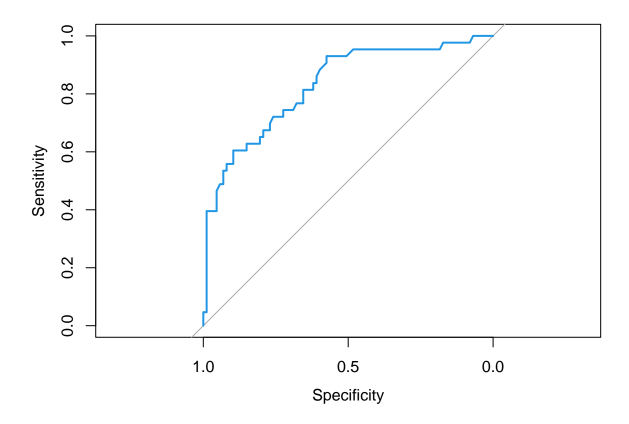
(iii) Random Forest

```
## Random Forest
##
## 262 samples
## 8 predictor
## 2 classes: 'neg', 'pos'
##
## No pre-processing
```

```
## Resampling: Cross-Validated (10 fold, repeated 1 times)
## Summary of sample sizes: 235, 235, 236, 236, 236, ...
## Resampling results across tuning parameters:
##
##
    mtry Accuracy
                      Kappa
##
           0.7938917 0.5193219
     2
##
           0.8014416 0.5417510
           0.7898917 0.5141065
##
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 5.
#obtaining class predictions
rf.classTrain <- predict(train.rf, type="raw")</pre>
confusionMatrix(train$diabetes, rf.classTrain)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction neg pos
          neg 175
##
##
          pos
              0 87
##
##
                  Accuracy: 1
                    95% CI: (0.986, 1)
##
##
       No Information Rate: 0.6679
##
       P-Value [Acc > NIR] : < 2.2e-16
##
##
                     Kappa: 1
##
##
   Mcnemar's Test P-Value : NA
##
##
               Sensitivity: 1.0000
##
               Specificity: 1.0000
##
            Pos Pred Value: 1.0000
##
            Neg Pred Value: 1.0000
##
                Prevalence: 0.6679
##
            Detection Rate: 0.6679
##
      Detection Prevalence: 0.6679
##
         Balanced Accuracy: 1.0000
##
##
          'Positive' Class : neg
##
rf.classTest <- predict(train.rf,</pre>
                         newdata = test,
                          type="raw")
confusionMatrix(test$diabetes, rf.classTest)
## Confusion Matrix and Statistics
```

##

```
##
             Reference
## Prediction neg pos
##
          neg 73 14
          pos 16 27
##
##
##
                  Accuracy : 0.7692
##
                    95% CI: (0.6872, 0.8386)
       No Information Rate: 0.6846
##
##
       P-Value [Acc > NIR] : 0.02154
##
##
                     Kappa: 0.4725
##
    Mcnemar's Test P-Value: 0.85513
##
##
##
               Sensitivity: 0.8202
##
               Specificity: 0.6585
##
            Pos Pred Value: 0.8391
##
            Neg Pred Value: 0.6279
##
                Prevalence: 0.6846
            Detection Rate: 0.5615
##
##
      Detection Prevalence: 0.6692
##
         Balanced Accuracy: 0.7394
##
##
          'Positive' Class : neg
##
rf_accuracy <- mean(bagg.classTest == test$diabetes)</pre>
cat("Accuracy of Classification Tree:", rf_accuracy, "\n")
## Accuracy of Classification Tree: 0.7692308
#Obtaining predicted probabilities for Test data
rf.probs=predict(train.rf,
                 newdata=test,
                 type="prob")
#Calculate ROC curve
rocCurve.rf <- roc(test$diabetes, rf.probs[,"neg"])</pre>
## Setting levels: control = neg, case = pos
## Setting direction: controls > cases
#plot the ROC curve
plot(rocCurve.rf,col=c(4))
```



```
#calculate the area under curve (bigger is better)
auc(rocCurve.rf)
```

Area under the curve: 0.8292

d. Neural Network

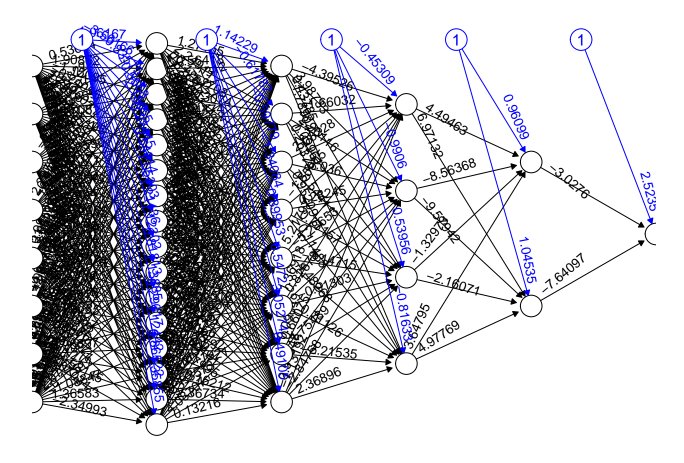
Reference:

 $\bullet \ \, https://www.datacamp.com/tutorial/neural-network-models-r$

library(tidyverse)

```
----- tidyverse 2.0.0 --
## -- Attaching core tidyverse packages --
## v dplyr
                                     2.1.5
               1.1.4
                         v readr
## v forcats
               1.0.0
                         v stringr
                                     1.5.1
## v lubridate 1.9.3
                         v tibble
                                     3.2.1
                         v tidyr
## v purrr
               1.0.2
                                     1.3.1
## -- Conflicts ----
                                                      ----- tidyverse_conflicts() --
## x stringr::boundary() masks strucchange::boundary()
## x dplyr::collapse()
                         masks nlme::collapse()
## x dplyr::combine()
                         masks gridExtra::combine()
## x tidyr::expand()
                         masks Matrix::expand()
```

```
## x dplyr::filter()
                          masks stats::filter()
## x dplyr::lag()
                          masks stats::lag()
## x purrr::lift()
                          masks caret::lift()
                          masks Matrix::pack()
## x tidyr::pack()
## x dplyr::select()
                           masks MASS::select()
## x tidyr::unpack()
                          masks Matrix::unpack()
## x dplyr::where()
                          masks party::where()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
library(neuralnet)
##
## Attaching package: 'neuralnet'
## The following object is masked from 'package:dplyr':
##
##
       compute
# Extract features and labels
X_train_nn <- as.data.frame(train[, -ncol(train)])</pre>
y_train_nn <- as.numeric(train$diabetes == "pos") # Convert to binary 0/1
X_test_nn <- as.data.frame(test[, -ncol(test)])</pre>
y_test_nn <- as.numeric(test$diabetes == "pos")</pre>
# Scale the features
X_train_nn_scaled <- scale(X_train_nn)</pre>
X_test_nn_scaled <- scale(X_test_nn)</pre>
# Combine scaled features and labels for training
train_combined <- cbind(X_train_nn_scaled, diabetes = y_train_nn)</pre>
nn_model = neuralnet(
    diabetes~.,
    data=train_combined,
    hidden=c(16,8,4,2),
    linear.output = FALSE
)
plot(nn_model,rep = "best")
```



```
# Predict probabilities on test set
nn_pred <- compute(nn_model, X_test_nn_scaled)</pre>
nn_pred_prob <- nn_pred$net.result</pre>
# Convert probabilities to binary predictions
nn_pred_class <- ifelse(nn_pred_prob > 0.5, 1, 0)
# Calculate and print confusion matrix
conf_matrix <- table(Predicted = nn_pred_class, Actual = as.numeric(test$diabetes == "pos"))</pre>
print(conf_matrix)
            Actual
##
## Predicted 0 1
           0 72 16
##
           1 15 27
##
# Calculate accuracy
check <- as.numeric(test$diabetes == "pos") == nn_pred_class</pre>
nn_accuracy <- (sum(check) / nrow(test))</pre>
print(paste("Neural Network Test Accuracy:", nn_accuracy))
```

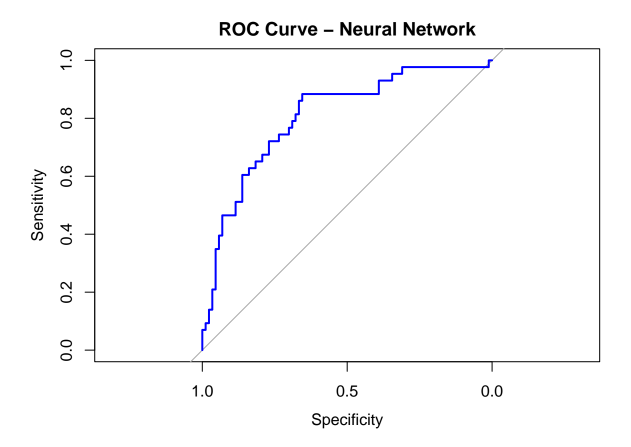
[1] "Neural Network Test Accuracy: 0.761538461538461"

```
# Calculate ROC curve
roc_curve_nn <- roc(test$diabetes, as.numeric(nn_pred_prob))

## Setting levels: control = neg, case = pos

## Setting direction: controls < cases

# Plot the ROC curve
plot(roc_curve_nn, col = "blue", main = "ROC Curve - Neural Network")</pre>
```



```
# Calculate and print AUC
auc_nn <- auc(roc_curve_nn)
cat("Neural Network - AUC:", auc_nn, "\n")</pre>
```

Neural Network - AUC: 0.8067362

e. Compare all models

- KNN
- Tree-based Method
 - Classification Tree

- Ensemble of bagged trees
- Random Forest
- Neural Network

```
# Create a summary table for accuracy and AUC
accuracy_results <- data.frame(</pre>
 Model = c("KNN", "GAM", "Classification Tree", "Bagged Trees", "Random Forest", "Neural Network"),
 Accuracy = c(knn_accuracy, gam_accuracy, tree_accuracy, bagg_accuracy, rf_accuracy, nn_accuracy),
 AUC = c(auc_knn, auc_gam, auc(rocCurve.tree), auc(rocCurve.bagg), auc(rocCurve.rf), auc_nn)
# Print the summary table
print(accuracy_results)
##
                  Model Accuracy
## 1
                     KNN 0.7923077 0.7213312
## 2
                     GAM 0.7769231 0.8334670
## 3 Classification Tree 0.7615385 0.7848169
## 4
           Bagged Trees 0.7692308 0.8310612
## 5
          Random Forest 0.7692308 0.8291901
         Neural Network 0.7615385 0.8067362
## 6
# Plot ROC curves for ALL models
plot(rocCurve.tree, col = 4, main = "ROC Curve Comparison", lwd = 2)
lines(rocCurve.bagg, col = 6, lwd = 2) # Add ROC curve for bagged trees
lines(rocCurve.rf, col = 1, lwd = 2) # Add ROC curve for random forest
lines(roc_curve_nn, col = "blue", lwd = 2) # Add ROC curve for neural network
lines(roc_curve_knn, col = "green", lwd = 2) # Add ROC curve for k-NN
lines(roc_curve_gam, col = "purple", lwd = 2) # Add ROC curve for GAM
# Add a legend to the plot
legend("bottomright", legend = c("Classification Tree", "Bagged Trees", "Random Forest", "Neural Networ
```

col = c(4, 6, 1, "blue", "green", "purple"), lwd = 2, bty = "n")

