# Factor Target Gene Prediction through Multi-Omics Datasets"

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# Introduction

#### Aim:

The aim of this project is to apply classification techniques to predict novel transcription factor target genes, focusing on Sox2 and Nanog during embryonic stem cell (ESC) differentiation.

## Background:

Transcriptional regulation is a fundamental ess in all living organisms, driven by transcription factors that control mRNA expression. These transcriptional networks are crucial in development, lineage specification, and cell fate decisions during early embryonic development (Theunissen and Jaenisch, 2017). Recent advances in omics technologies allow for the profiling of genome-wide transcriptional and epigenetic events, providing a deeper understanding of these networks.

In this project, we utilize high-temporal-resolution multi-omics data of ESC differentiation (Yang et al., 2019) to predict novel substrates of Sox2 and Nanog—two key transcription factors involved in maintaining pluripotency and guiding cell differentiation.

**Dataset Overview:** - **Transcriptome:** Time-course mRNA profiles during ESC differentiation. - **Proteome:** Time-course protein expression profiles during ESC differentiation. - **Epigenome:** Time-course ESC differentiation epigonme profiles of 6 histone marks.

We will develop and validate a classification model to predict novel transcription factor target genes, focusing on Sox2 and Nanog, using the provided multi-omics datasets.

# Load Required Libraries and Data

We start by loading the necessary R packages and the dataset Final\_Project\_ESC.RData, which contains the transcriptome, proteome, and epigenome data, along with a subset of known Sox2/Nanog target genes.

```
# Load necessary packages and data
load("Final_Project_ESC.RData", verbose = TRUE)
```

```
## Loading objects:
     Transcriptome
##
##
     Proteome
##
     H3K4me3
##
     H3K27me3
##
     PolII
##
     H3K4me1
##
     H3K27ac
     H3K9me2
##
##
     cMyc_target_genes
##
     cMyc_target_genes_subset
##
     OSN_target_genes
##
     OSN_target_genes_subset
```

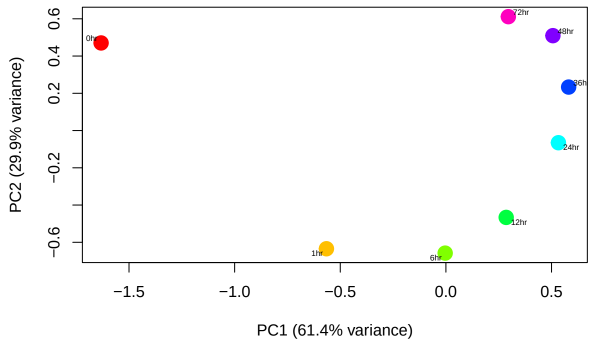
```
suppressPackageStartupMessages({
    library(e1071)
    library(ggplot2)
    library(ROCR)
    library(dplyr)
    library(tibble)
    library(reshape2)
    library(kernlab)
    library(caret)
    library(randomForest)
    library(adabag)
    library(gbm)
    library(xgboost)
    library(pROC)
    library(doParallel)
    library(calibrate)
})
```

# Describe and explore the Data set details

Before beginning data analysis it is important understand and investigate the data. The goal of this report is to predict to predict novel transcription factor target genes from multi-omics data. For each of our datasets lets look at the structure of the data and perform PCA Analysis as means of identifying trends in the dataset.

# Transcriptome

```
head(Transcriptome)
##
                  0hr
                              1hr
                                          6hr
                                                     12hr
                                                                 24hr
                                                           0.71864514
## GNAI3 8.881784e-16
                      0.29131114
                                  0.618947976
                                               0.48337689
## CDC45 0.00000e+00
                       0.25062323
                                  0.199000752
                                               0.38800303
                                                           0.47163966
         0.000000e+00 0.38475910
                                  0.471216601
## H19
                                               0.55341685
                                                           1.87279375
## SCML2 8.881784e-16 0.64099671
                                  0.917100751
                                               0.73622935
                                                           0.74708761
## NARF -8.881784e-16 -0.40736349 -1.186942510 -0.69986049 -0.15754727
## CAV2
         5.551115e-17 0.01608049 0.002154149 0.07278802 0.06737809
##
               36hr
                           48hr
                                       72hr
## GNAI3 0.90833508 0.63408090 0.81560693
## CDC45 0.20338260 -0.03095025 -0.55159826
         2.77535428 4.17558703 4.34079023
## H19
## SCML2 0.71202112 -0.20856228 -1.12588475
## NARF
        -0.73012533 -0.11611985 -0.01534405
## CAV2
         dim(Transcriptome)
## [1] 19788
colnames(Transcriptome)
## [1] "0hr" "1hr" "6hr" "12hr" "24hr" "36hr" "48hr" "72hr"
# PCA analysis on the correlation matrix of the transcriptome data
cor.mat <- cor(Transcriptome)</pre>
pca.mat <- prcomp(cor.mat)</pre>
```

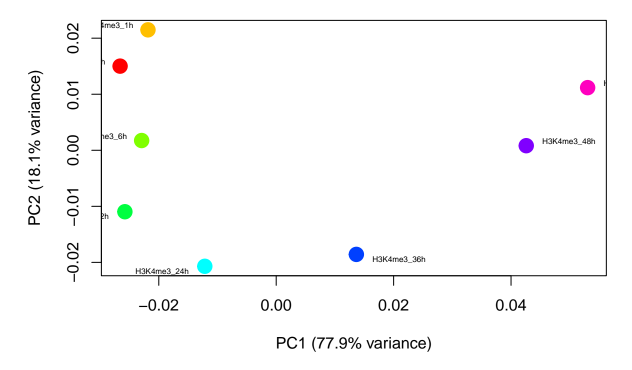


## Proteome

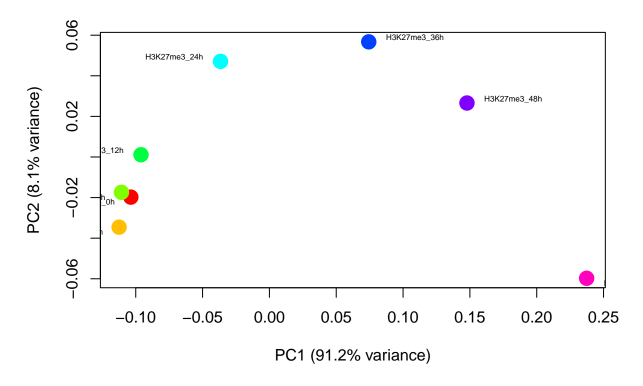
```
cor.proteome <- cor(Proteome)</pre>
pca.proteome <- prcomp(cor.proteome)</pre>
summary(pca.proteome)$importance
##
                                 PC1
                                            PC2
                                                      PC3
                                                                 PC4
                                                                             PC5
## Standard deviation
                           0.5233315 0.2915203 0.1624012 0.1053089 0.07579296
## Proportion of Variance 0.6763300 0.2098700 0.0651300 0.0273900 0.01419000
## Cumulative Proportion 0.6763300 0.8862000 0.9513300 0.9787100 0.99290000
##
                                  PC6
## Standard deviation
                           0.05362587 2.715731e-18
## Proportion of Variance 0.00710000 0.000000e+00
## Cumulative Proportion 1.00000000 1.000000e+00
# Using the previous correlation matrix and PCA results
cor.proteome <- cor(Proteome)</pre>
pca.proteome <- prcomp(cor.proteome)</pre>
```

```
# Get group labels and colors
grp <- rownames(pca.proteome$x) # Set groups according to your data</pre>
grp.col <- rainbow(nrow(pca.proteome$x))</pre>
names(grp.col) <- rownames(pca.proteome$x)</pre>
# Plot the PCA
plot(pca.proteome$x[,1], pca.proteome$x[,2], col=grp.col[grp], pch=19, cex=2,
     xlab=paste0("PC1 (", round(summary(pca.proteome)$importance[2,1]*100,1), "% variance)"),
     ylab=paste0("PC2 (", round(summary(pca.proteome)$importance[2,2]*100,1), "% variance)"))
      9.4
PC2 (21% variance)
      0.2
      0.0
            -0.8
                       -0.6
                                  -0.4
                                             -0.2
                                                                    0.2
                                                                               0.4
                                                         0.0
                                      PC1 (67.6% variance)
```

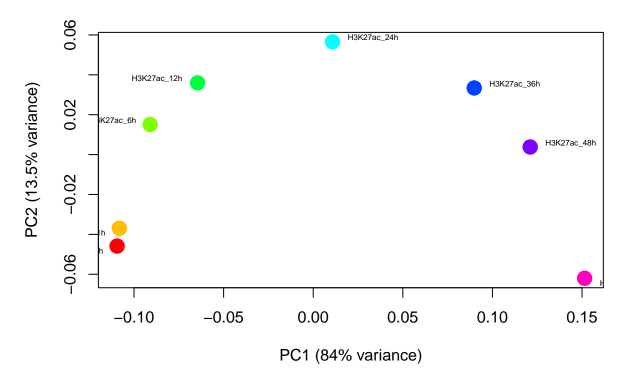
## h3k4me3



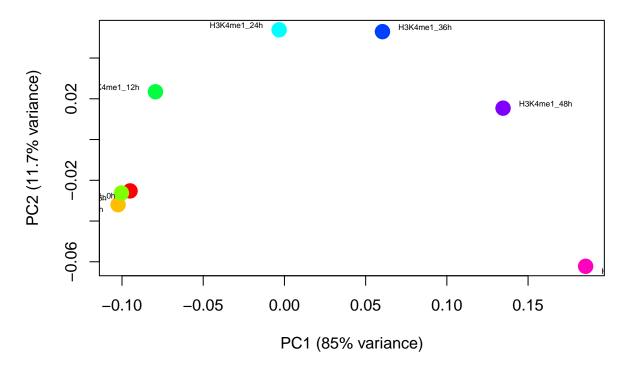
## H3K27me3



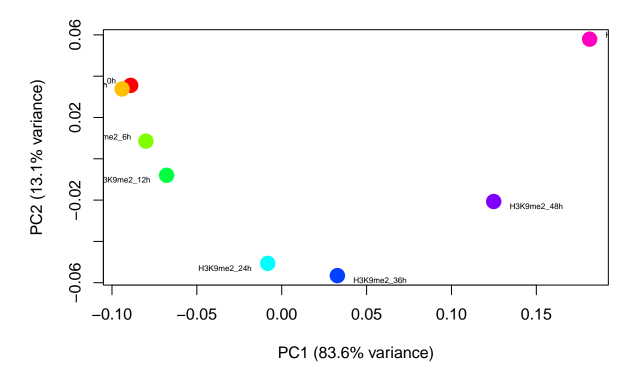
## H3K27ac



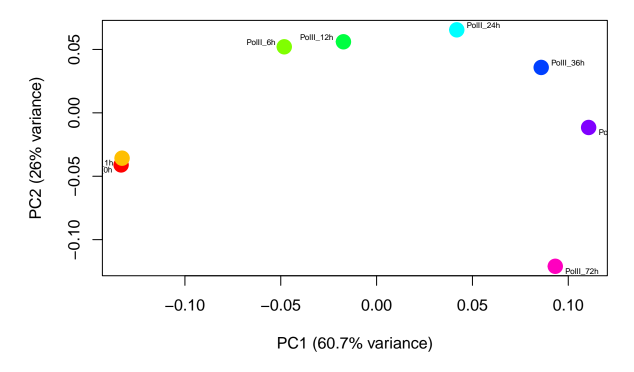
## H3K4me1



## H3K9me2



## PolII



## Classification

## Filter and Combine Datasets

In order to properly model and predicting novel transcription factors target genes we join the 8 datasets together and perform our calculations on the larger dataset.

To ensure consistency across datasets, we filter each dataset to include only the common genes present in all omics layers. We then combine these filtered datasets into a single data frame for further analysis.

```
# Ensure all data has the same set of genes
genes <- intersect(rownames(Transcriptome), rownames(Proteome))</pre>
genes <- intersect(genes, rownames(H3K4me3))</pre>
genes <- intersect(genes, rownames(H3K27me3))</pre>
genes <- intersect(genes, rownames(H3K27ac))</pre>
genes <- intersect(genes, rownames(H3K4me1))</pre>
genes <- intersect(genes, rownames(H3K9me2))</pre>
genes <- intersect(genes, rownames(PolII))</pre>
# Filter each dataset for the common genes
Transcriptome_filter <- Transcriptome[genes, ]</pre>
Proteome_filter <- Proteome[genes, ]</pre>
H3K4me3_filter <- H3K4me3[genes, ]</pre>
H3K27me3_filter <- H3K27me3[genes, ]</pre>
H3K27ac_filter <- H3K27ac[genes, ]</pre>
H3K4me1_filter <- H3K4me1[genes, ]</pre>
H3K9me2_filter <- H3K9me2[genes, ]</pre>
PolII_filter <- PolII[genes, ]</pre>
# Confirm that all datasets share the same gene
identical(rownames(Transcriptome_filter), rownames(H3K4me3_filter))
```

## [1] TRUE

```
identical(rownames(Proteome_filter), rownames(H3K4me3_filter))
## [1] TRUE
identical(rownames(H3K27ac_filter), rownames(H3K4me3_filter))
## [1] TRUE
identical(rownames(H3K4me1_filter), rownames(H3K4me3_filter))
## [1] TRUE
identical(rownames(H3K9me2_filter), rownames(H3K4me3_filter))
## [1] TRUE
identical(rownames(PolII_filter), rownames(H3K4me3_filter))
## [1] TRUE
# Rename columns to avoid conflicts
colnames(Transcriptome_filter) <- paste("T_", colnames(Transcriptome_filter), sep = "")</pre>
colnames(Proteome_filter) <- paste("P_", colnames(Proteome_filter), sep = "")</pre>
colnames(H3K4me3_filter) <- paste("H3K4me3_", colnames(H3K4me3_filter), sep = "")</pre>
colnames(H3K27me3_filter) <- paste("H3K27me3_", colnames(H3K27me3_filter), sep = "")</pre>
colnames(H3K27ac_filter) <- paste("H3K27ac_", colnames(H3K27ac_filter), sep = "")</pre>
colnames(H3K4me1_filter) <- paste("H3K4me1_", colnames(H3K4me1_filter), sep = "")</pre>
colnames(H3K9me2_filter) <- paste("H3K9me2_", colnames(H3K9me2_filter), sep = "")</pre>
colnames(PolII_filter) <- paste("PolII_", colnames(PolII_filter), sep = "")</pre>
# Combine the datasets
combined data <- cbind(</pre>
  Transcriptome_filter,
 Proteome_filter,
 H3K4me3_filter,
 H3K27me3 filter,
 H3K27ac_filter,
  H3K4me1_filter,
  H3K9me2_filter,
  PolII_filter
# Add the labels
label <- ifelse(genes %in% OSN_target_genes_subset, "OSN", "Other")</pre>
combined_data <- data.frame(combined_data)</pre>
combined_data$label <- factor(label)</pre>
# Number of genes which are known to be targets for Sox2 and Nanog
length(OSN_target_genes_subset)
## [1] 100
```

We have 100 known target genes for OSN, and as seen below the we have 95 genes that have been identified as novel Sox2/Nanog targets on our combined filtered dataset.

```
# Check the initial label distribution
print(table(combined_data$label))
```

##

```
## OSN Other
## 95 8085
```

## **Data Splitting and Balancing**

The dataset is split into training (90%) and testing (10%) sets. The label column is reassigned to the test set to ensure that it is included correctly.

```
# Split the dataset into training (90%) and testing (10%) sets
set.seed(123)
train_index <- createDataPartition(combined_data$label, p = 0.9, list = FALSE)</pre>
train data <- combined data[train index, ]
test_data <- combined_data[-train_index, ]</pre>
# Reassign the label column to test_data
test_data$label <- combined_data[-train_index, "label"]</pre>
# Check the distribution of labels in the training and test sets
print("Training set label distribution:")
## [1] "Training set label distribution:"
print(table(train_data$label))
##
##
     OSN Other
      86 7277
print("Test set label distribution:")
## [1] "Test set label distribution:"
print(table(test_data$label))
##
##
     OSN Other
##
       9
           808
```

#### Balancing the Training Data

Since the dataset is imbalanced, downsampling is used to create a balanced training dataset. This ensures that both classes (OSN and Other) are equally represented, improving the robustness of the classification model.

##

```
## OSN Other
## 86 86

# Final check of training dataset dimensions
print(dim(downsampled_train_data))
## [1] 172 64
```

## **Model Training**

Train SVM and Random Forest Models We train two machine learning models, SVM (with the radial kernel) and Random Forest, using the balanced training dataset.

```
# Set a consistent seed
set.seed(123)

# Train an SVM model on the downsampled training data with radial basis function kernel
svm_model <- svm(label ~ ., data = downsampled_train_data, kernel = "radial", probability = TRUE)

# Train a Random Forest model on the downsampled training data
set.seed(123)
rf_model <- randomForest(label ~ ., data = downsampled_train_data, ntree = 1000)</pre>
```

## Bagging w/ Bagged Trees

```
bagged_trees <- train(
  label ~ .,
  data = downsampled_train_data,
  method = "treebag",
  trControl = trainControl(method = "cv", number = 5),
  tuneLength = 5
)
print(bagged_trees)

## Bagged CART</pre>
```

```
##
## 172 samples
## 63 predictor
    2 classes: 'OSN', 'Other'
##
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 137, 138, 138, 137, 138
## Resampling results:
##
##
    Accuracy
                Kappa
##
    0.7447059 0.4888148
```

#### GBM w/ Hyperparam Tuning and xgboost (using parallel processing)

```
tune_grid_xgb <- expand.grid(
  nrounds = c(100, 200),
  max_depth = c(3, 6),
  eta = c(0.1, 0.3),
  gamma = c(0, 0.1),</pre>
```

```
colsample_bytree = c(0.5, 1),
  min_child_weight = c(1, 10),
  subsample = c(0.5, 1)
)

cl <- makeCluster(detectCores())
registerDoParallel(cl)

train_control <- trainControl(
  method = "cv",
  number = 3,
  savePredictions = "final",
  verboseIter = TRUE,
  allowParallel = TRUE
)</pre>
```

xgb\_model\_tuned <- train( label  $\sim$  ., data = downsampled\_train\_data, method = "xgbTree", tr-Control = train\_control, tuneGrid = tune\_grid\_xgb, metric = "Accuracy" # Performance Metric ) print(xgb\_model\_tuned)

## Printing the best model's details

print(xgb\_model\_tuned\$bestTune)

## Plotting model performance

plot(xgb\_model\_tuned):

#### Model Evaluation

Predictions and Confusion Matrix We use the trained models to make predictions on the test set and evaluate their performance using confusion matrices.

```
# Ensure column names in the test set match the training data
colnames(test_data) <- colnames(downsampled_train_data) [1:(ncol(downsampled_train_data) - 1)] # Exclud
# Check if the label column is present and correctly populated
if ("label" %in% colnames(combined_data) && length(combined_data[-train_index, "label"]) == nrow(test_d
    # Assign the label to test_data
    test_data$label <- combined_data[-train_index, "label"]</pre>
} else {
    stop("The label vector is empty or has a different length than expected. Check the data preparation
# Ensure the label is a factor with the correct levels
test_data$label <- factor(test_data$label, levels = c("OSN", "Other"))</pre>
# SVM Predictions on the test set
svm_test_pred <- predict(svm_model, newdata = test_data[, -ncol(test_data)], probability = TRUE)</pre>
svm_test_prob <- attr(svm_test_pred, "probabilities")[, "OSN"]</pre>
# Random Forest Predictions on the test set
rf_test_pred <- predict(rf_model, newdata = test_data[, -ncol(test_data)], type = "prob")[, "OSN"]
# SVM Confusion Matrix on the test set
```

```
svm_test_conf_matrix <- table(Predicted = ifelse(svm_test_prob > 0.5, "OSN", "Other"), Actual = test_da
print("SVM Test Confusion Matrix:")
## [1] "SVM Test Confusion Matrix:"
print(svm_test_conf_matrix)
##
            Actual
## Predicted OSN Other
##
       OSN
               7
                   194
##
       Other
               2
# Random Forest Confusion Matrix on the test set
rf_test_conf_matrix <- table(Predicted = ifelse(rf_test_pred > 0.5, "OSN", "Other"), Actual = test_data
print("Random Forest Test Confusion Matrix:")
## [1] "Random Forest Test Confusion Matrix:"
print(rf_test_conf_matrix)
            Actual
## Predicted OSN Other
##
       OSN
               8
                   204
##
       Other
               1
                   604
```

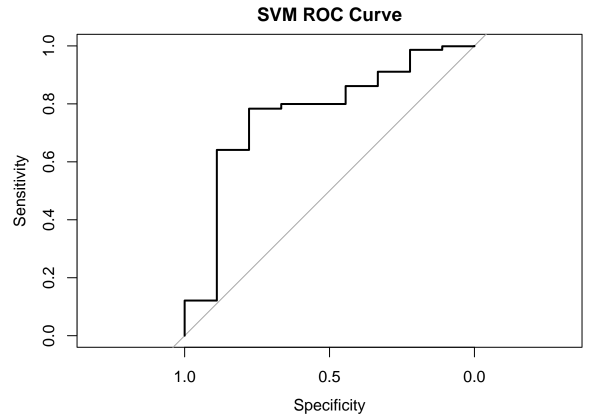
#### **ROC Curve and AUC**

To further assess model performance, we plot the ROC curves and calculate the AUC for both the SVM and Random Forest models.

```
# Evaluate the models on the test set (AUC and ROC)
svm_test_roc <- roc(test_data$label, svm_test_prob)

## Setting levels: control = OSN, case = Other

## Setting direction: controls > cases
plot(svm_test_roc, main = "SVM ROC Curve")
```



```
print(paste("Final SVM Test AUC:", auc(svm_test_roc)))

## [1] "Final SVM Test AUC: 0.766914191419142"

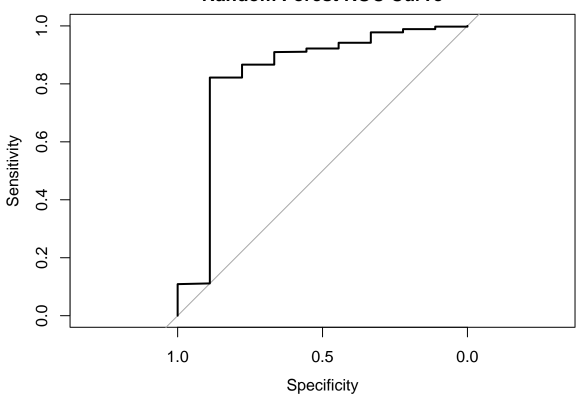
rf_test_roc <- roc(test_data$label, rf_test_pred)

## Setting levels: control = OSN, case = Other

## Setting direction: controls > cases

plot(rf_test_roc, main = "Random Forest ROC Curve")
```

# **Random Forest ROC Curve**



```
print(paste("Final Random Forest Test AUC:", auc(rf_test_roc)))

## [1] "Final Random Forest Test AUC: 0.8373899889989"

# Evaluate SVM model on the test set (AUC and ROC)

svm_test_roc <- roc(test_data$label, svm_test_prob)

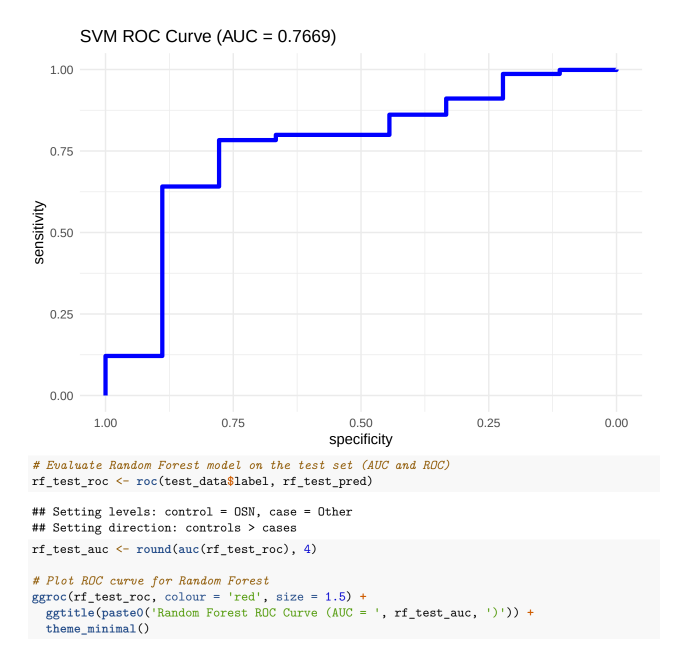
## Setting levels: control = OSN, case = Other

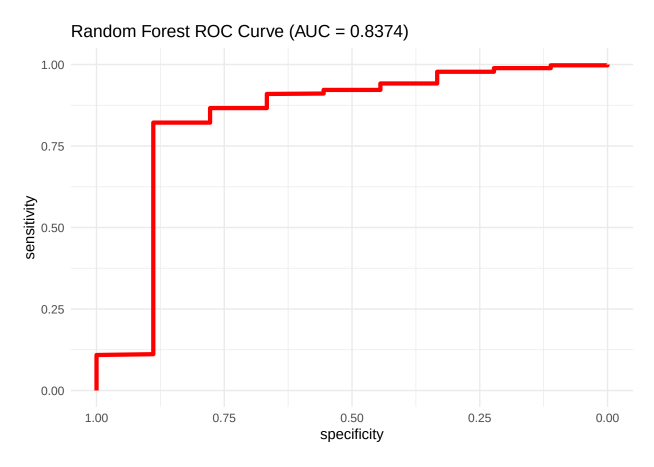
## Setting direction: controls > cases

svm_test_auc <- round(auc(svm_test_roc), 4)

# Plot ROC curve for SVM

ggroc(svm_test_roc, colour = 'blue', size = 1.5) +
    ggtitle(pasteO('SVM ROC Curve (AUC = ', svm_test_auc, ')')) +
    theme_minimal()</pre>
```





## **Future Directions:**

## Model Improvement:

Further hyperparameter tuning using automated methods like grid search or random search could refine the model's accuracy. Additionally, exploring ensemble methods that combine predictions from several models might yield better results.

#### **Data Expansion:**

Incorporating additional omics layers, such as metabolomics or additional transcription factor binding profiles, could help to improve the model's predictive power and generalisability.

## Integration with Clinical Data:

Linking omics profiles with clinical outcomes could also be explored to improve the translational impact of the research.