machine learning(732A99) lab1

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Assignment 1

Loading The Libraries

Loading Input files

```
spam_data <- read.xlsx("spambase.xlsx", sheetName = "spambase_data")
spam_data$Spam <- as.factor(spam_data$Spam)

tecator_data <- read.xlsx("tecator.xlsx", sheetName = "data")
tecator_data <- tecator_data[,2:NCOL(tecator_data)] # removing sample column</pre>
```

1.1 Import the data into R and divide it into training and test sets (50%/50%) by using the following code

```
set.seed(12345)

n = NROW(spam_data)
id = sample(1:n, floor(n*0.5))
train = spam_data[id,]
test = spam_data[-id,]
```

1.2 Use logistic regression (functions glm(), predict()) to classify the training and test data by the classification principles

Manual Feature Selection

```
best_model <- glm(formula = Spam ~., family = binomial, data = train)
summ(best_model)

## MODEL INFO:
## Observations: 1370

## Dependent Variable: Spam
## Type: Generalized linear model
## Family: binomial
## Link function: logit

##
## MODEL FIT:
## <U+0001D6D8>²(48) = 768.27, p = 0.00
## Pseudo-R² (Cragg-Uhler) = 0.60
## Pseudo-R² (McFadden) = 0.45
## AIC = 1026.54, BIC = 1282.45
```

```
##
## Standard errors: MLE
                           S.E. z val. p
##
                  Est.
                           0.20 7.50 0.00 ***
## (Intercept)
                  1.51
## Word1
                 -0.72
                           0.50 -1.43 0.15
## Word2
                  0.04
                           0.30
                                 0.13 0.89
## Word3
                 -0.35
                           0.18 -1.92 0.06
                                  1.23 0.22
## Word4
                  0.14
                           0.11
## Word5
                  0.12
                           0.14
                                  0.86 0.39
## Word6
                           0.42
                                 0.70 0.49
                  0.29
## Word7
                 -0.29
                           0.33
                                 -0.88 0.38
                                 -0.29 0.77
## Word8
                 -0.10
                           0.35
## Word9
                 -0.11
                           0.41
                                 -0.270.79
                                 -0.19 0.85
## Word10
                 -0.03
                           0.17
## Word11
                 -0.61
                           0.68
                                 -0.90 0.37
## Word12
                  0.16
                           0.11
                                  1.50 0.13
## Word13
                  0.78
                           0.36
                                  2.19 0.03
## Word14
                 -0.48
                           0.30
                                 -1.60 0.11
## Word15
                 -0.13
                           0.39
                                 -0.34 0.74
## Word16
                  0.32
                           0.23
                                  1.36 0.17
## Word17
                 -0.09
                           0.28
                                 -0.330.74
## Word18
                  0.02
                           0.23
                                 0.10 0.92
                                  0.01 0.99
## Word19
                  0.00
                           0.06
## Word20
                  0.02
                           0.32
                                  0.05 0.96
## Word21
                 -0.03
                                -0.26 0.79
                           0.11
## Word22
                 -0.48
                           0.32 -1.49 0.14
## Word23
                  0.25
                           0.34
                                 0.74 0.46
## Word24
                 -0.25
                           0.63
                                 -0.40 0.69
## Word25
                 -0.08
                           0.06
                                -1.340.18
## Word26
                  0.00
                           0.14
                                 0.03 0.98
## Word27
                 -0.22
                           0.11
                                 -2.08 0.04
## Word28
                  0.13
                           0.19
                                 0.70 0.48
                                 -0.89 0.37
## Word29
                 -0.08
                           0.09
## Word30
                 -1.82
                           0.62
                                 -2.93 0.00
## Word31
                 -4.69
                           1.85
                                 -2.53 0.01
## Word32
               -119.45 15134.18
                                 -0.01 0.99
## Word33
                 -2.90
                           0.68
                                 -4.27 0.00 ***
## Word34
                 -3.71
                           4.35 -0.85 0.39
## Word35
                 -7.03
                           2.00
                                 -3.52 0.00 ***
                                -4.40 0.00 ***
## Word36
                 -1.68
                           0.38
## Word37
                 -0.86
                           0.22
                                 -3.95 0.00 ***
## Word38
                 -0.60
                           1.28
                                 -0.470.64
## Word39
                 -1.88
                           0.43
                                 -4.38 0.00 ***
                           0.34
                                  0.22 0.83
## Word40
                  0.07
## Word41
               -332.55 16559.52
                                 -0.02 0.98
                           1.30
                                 -4.11 0.00 ***
## Word42
                 -5.35
                                 -3.53 0.00 ***
## Word43
                 -2.59
                           0.74
## Word44
                 -2.93
                           0.66
                                -4.44 0.00 ***
## Word45
                 -1.14
                           0.17
                                 -6.79 0.00 ***
                                 -6.35 0.00 ***
## Word46
                 -3.29
                           0.52
## Word47
                 -3.74
                           2.03
                                 -1.84 0.07
                           1.47 -2.98 0.00 **
## Word48
                 -4.39
```

Prediction for probability greater than 50% and 90%

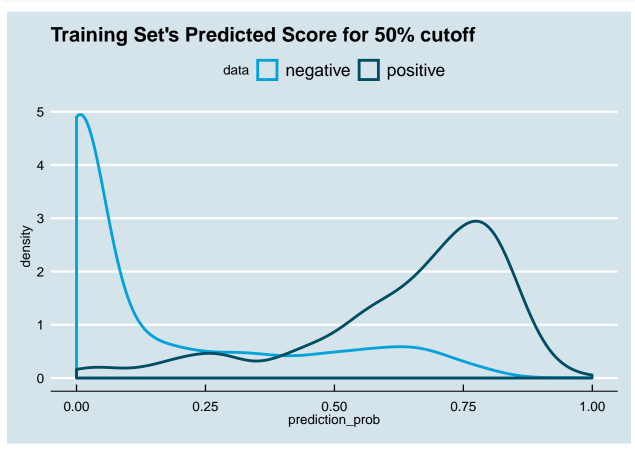
```
# prediction
train$prediction_prob <- predict(best_model, newdata = train, type = "response")
test$prediction_prob <- predict(best_model, newdata = test , type = "response")

train$prediction_class_50 <- ifelse(train$prediction_prob > 0.50, 1, 0)
test$prediction_class_50 <- ifelse(test$prediction_prob > 0.50, 1, 0)

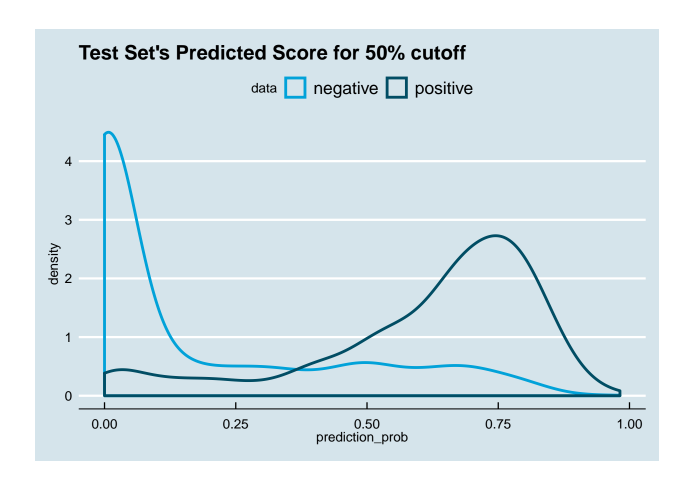
train$prediction_class_90 <- ifelse(train$prediction_prob > 0.90, 1, 0)
test$prediction_class_90 <- ifelse(test$prediction_prob > 0.90, 1, 0)
```

Assessing the Model

```
# plots
ggplot(train, aes(prediction_prob, color = Spam)) +
geom_density(size = 1) + ggtitle("Training Set's Predicted Score for 50% cutoff") +
scale_color_economist(name = "data", labels = c("negative", "positive")) +
theme_economist()
```



```
ggplot(test, aes(prediction_prob, color = Spam)) +
geom_density(size = 1) + ggtitle("Test Set's Predicted Score for 50% cutoff") +
scale_color_economist(name = "data", labels = c("negative", "positive")) +
theme_economist()
```



1.2 Assessing the Fit on train dataset for 50%

```
#confusion table
conf_train <- table(train$Spam, train$prediction_class_50)</pre>
names(dimnames(conf_train)) <- c("Actual Train", "Predicted Train")</pre>
confusionMatrix(conf_train)
## Confusion Matrix and Statistics
##
##
               Predicted Train
##
  Actual Train
                 0
              0 803 142
##
              1 81 344
##
##
                  Accuracy : 0.8372
##
                    95% CI: (0.8166, 0.8564)
##
##
       No Information Rate: 0.6453
##
       P-Value [Acc > NIR] : < 0.0000000000000022
##
                     Kappa : 0.6341
##
##
   Mcnemar's Test P-Value: 0.00005872
##
##
               Sensitivity: 0.9084
               Specificity: 0.7078
##
##
            Pos Pred Value: 0.8497
```

```
##
            Neg Pred Value: 0.8094
##
                Prevalence: 0.6453
##
            Detection Rate: 0.5861
##
      Detection Prevalence: 0.6898
##
         Balanced Accuracy: 0.8081
##
##
          'Positive' Class: 0
##
conf_test <- table(test$Spam, test$prediction_class_50)</pre>
names(dimnames(conf_test)) <- c("Actual Test", "Predicted Test")</pre>
confusionMatrix(conf_test)
## Confusion Matrix and Statistics
##
##
              Predicted Test
## Actual Test
                 0
                    1
##
             0 791 146
##
             1 97 336
##
##
                  Accuracy : 0.8226
                    95% CI: (0.8014, 0.8425)
##
##
       No Information Rate: 0.6482
##
       P-Value [Acc > NIR] : < 0.0000000000000022
##
##
                     Kappa: 0.6018
    Mcnemar's Test P-Value: 0.002076
##
##
##
               Sensitivity: 0.8908
##
               Specificity: 0.6971
##
            Pos Pred Value: 0.8442
##
            Neg Pred Value: 0.7760
##
                Prevalence: 0.6482
##
            Detection Rate: 0.5774
      Detection Prevalence: 0.6839
##
##
         Balanced Accuracy: 0.7939
##
##
          'Positive' Class: 0
##
```

Analysis: Distribution of the prediction score grouped by known outcome given that our model's final objective is to classify new instances into one of two categories (spam vs. non-spam). We will want the model to give high scores to positive instances (1: spam) and low scores (0: not spam) otherwise. Ideally you want the distribution of scores to be separated, with the score of the negative instances to be on the left and the score of the positive instance to be on the right.

From the confusion matrix it is apparent that Accuracy on train and test dataset when cutoff=50% is about $\sim 84\%$ for train and $\sim 82\%$ for test, thus the misclassification rate is ~ 16 and $\sim 18\%$ for the train and test dataset.

1.3 Assessing the Fit on train dataset for 90%

```
#confusion table
conf_train1 <- table(train$Spam, train$prediction_class_90)</pre>
```

```
names(dimnames(conf_train1)) <- c("Actual Train", "Predicted Train")</pre>
conf_train1
##
                Predicted Train
## Actual Train
                   0
                        1
##
               0 944
                        1
##
               1 419
                        6
conf_test1 <- table(test$Spam, test$prediction_class_90)</pre>
names(dimnames(conf_test1)) <- c("Actual Test", "Predicted Test")</pre>
conf_test1
               Predicted Test
##
## Actual Test
                  0
                      1
##
              0 936
                      1
              1 427
##
```

Analysis: Strange, the model almost only predicts one class!! We know that the prediction of a logistic regression model is a probability, thus in order to use it as a classifier, we'll have to choose a cutoff value, or threshold (cutoff). Where scores above this value will classified as positive, those below as negative. Lets us find this optimum value.

Choosing the best cutoff for test

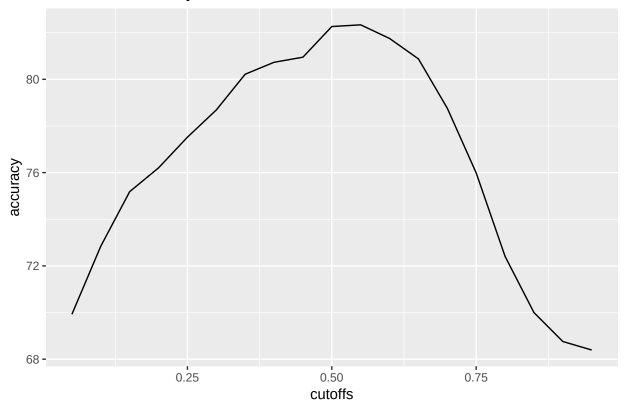
```
cutoffs <- seq(from = 0.05, to = 0.95, by = 0.05)
accuracy <- NULL

for (i in seq_along(cutoffs)){
    prediction <- ifelse(test$prediction_prob >= cutoffs[i], 1, 0) #Predicting for cut-off
    accuracy <- c(accuracy,length(which(test$Spam == prediction))/length(prediction)*100)}

cutoff_data <- as.data.frame(cbind(cutoffs, accuracy))

ggplot(data = cutoff_data, aes(x = cutoffs, y = accuracy)) +
    geom_line() +
    ggtitle("Cutoff vs. Accuracy for Test Dataset")</pre>
```

Cutoff vs. Accuracy for Test Dataset



Analysis: Our small detour suggests that the cutoff value of $\sim 60\%$ was the best for our purpose and going higher than this leads to worse results, at 70% and above the accuracy drastically reduces which is what we see when we make cutoff as 90%.

From the confusion matrix it is evident that the model becomes a trivial model (predicts all cases as one class) and thus the prediction is worse than just tossing a coin. This should be the absoutely the worst case that we should avoid.

The missclassication rate is about 31% for both the training dataset and test dataset.

1.4 Use standard classifier kknn() with K=30 from package kknn, report the the misclassification rates for the training and test data and compare the results with step 1.2.

```
knn_model30 <- train.kknn(Spam ~., data = train, kmax = 30)

train$knn_prediction_class <- predict(knn_model30, train)
test$knn_prediction_class <- predict(knn_model30, test)

conf_train2 <- table(train$Spam, train$knn_prediction_class)
names(dimnames(conf_train2)) <- c("Actual Train", "Predicted Train")
confusionMatrix(conf_train2)

## Confusion Matrix and Statistics
##
## Predicted Train</pre>
```

```
## Actual Train 0 1
##
             0 845 100
             1 91 334
##
##
##
                 Accuracy : 0.8606
##
                   95% CI: (0.8411, 0.8785)
##
      No Information Rate: 0.6832
      ##
##
##
                    Kappa: 0.6761
##
   Mcnemar's Test P-Value : 0.5627
##
              Sensitivity: 0.9028
##
##
              Specificity: 0.7696
           Pos Pred Value: 0.8942
##
##
           Neg Pred Value: 0.7859
##
               Prevalence: 0.6832
##
           Detection Rate: 0.6168
##
     Detection Prevalence: 0.6898
##
        Balanced Accuracy: 0.8362
##
##
         'Positive' Class : 0
##
conf_test2 <- table(test$Spam, test$knn_prediction_class)</pre>
names(dimnames(conf_test2)) <- c("Actual Test", "Predicted Test")</pre>
confusionMatrix(conf_test2)
## Confusion Matrix and Statistics
##
##
             Predicted Test
               0 1
## Actual Test
            0 801 136
            1 139 294
##
##
##
                 Accuracy : 0.7993
                   95% CI : (0.7771, 0.8202)
##
      No Information Rate: 0.6861
##
##
      ##
##
                    Kappa: 0.5348
##
   Mcnemar's Test P-Value : 0.904
##
##
              Sensitivity: 0.8521
              Specificity: 0.6837
##
##
           Pos Pred Value: 0.8549
##
           Neg Pred Value: 0.6790
##
               Prevalence: 0.6861
##
           Detection Rate: 0.5847
##
     Detection Prevalence: 0.6839
##
        Balanced Accuracy: 0.7679
##
##
         'Positive' Class: 0
##
```

Analysis: Using KKNN with K = 30, we increased our training accuracy to 86%, thus misclassification is 14%, however for the test dataset misclassification rate is about ~20%.

Thus compared to using logisite model the misclassification error for the training dataset decreased by 2% to 14%, while for the test dataset the misclassification error increased by 2% to 20%.

1.5 Repeat step 4 for K=1 and compare the results with step 4. What effect does the decrease of K lead to and why?

```
knn_model1 <- train.kknn(Spam ~., data = train, kmax = 1)</pre>
train$knn_prediction_class <- predict(knn_model1, train)</pre>
test$knn_prediction_class <- predict(knn_model1, test)</pre>
conf_train2 <- table(train$Spam, train$knn_prediction_class)</pre>
names(dimnames(conf_train2)) <- c("Actual Train", "Predicted Train")</pre>
confusionMatrix(conf_train2)
## Confusion Matrix and Statistics
##
##
               Predicted Train
## Actual Train
                  Ω
                       1
              0 945
                       0
##
##
              1
                   0 425
##
##
                   Accuracy: 1
                     95% CI : (0.9973, 1)
##
##
       No Information Rate: 0.6898
       P-Value [Acc > NIR] : < 0.0000000000000022
##
##
##
                      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: 0.6898
            Detection Rate: 0.6898
##
##
      Detection Prevalence: 0.6898
##
         Balanced Accuracy: 1.0000
##
          'Positive' Class: 0
##
conf_test2 <- table(test$Spam, test$knn_prediction_class)</pre>
names(dimnames(conf_test2)) <- c("Actual Test", "Predicted Test")</pre>
confusionMatrix(conf_test2)
## Confusion Matrix and Statistics
##
##
              Predicted Test
## Actual Test 0 1
             0 750 187
##
```

```
##
             1 150 283
##
##
                  Accuracy: 0.754
                    95% CI : (0.7303, 0.7766)
##
##
       No Information Rate: 0.6569
       P-Value [Acc > NIR] : 0.00000000000004691
##
##
##
                     Kappa: 0.4438
##
   Mcnemar's Test P-Value: 0.04987
##
##
               Sensitivity: 0.8333
               Specificity: 0.6021
##
##
            Pos Pred Value: 0.8004
##
            Neg Pred Value: 0.6536
##
                Prevalence: 0.6569
##
            Detection Rate: 0.5474
##
      Detection Prevalence: 0.6839
##
         Balanced Accuracy: 0.7177
##
##
          'Positive' Class: 0
##
```

Analysis: Using KKNN with K = 1, we increased our training accuracy to 100%, thus misclassification is 0%, however for the test dataset accuracy is ~75% thus misclassification rate is about ~25%, thus we improved on the training accuracy but did bad on the test case, thus this is an example of overfitting leading to more varience

Explaination: The KKNN works in the following way, An object is classified by a majority vote of its neighbors, with the object being assigned to the class most common among its k nearest neighbors. If k = 1, then the object is simply assigned to the class of that single nearest neighbor. Thus K=1, makes the separation boundary to be very complex and locally optimised (lots of local clusters), while as K goes higher, the decision boundary becomes more linear/simple.

Assignment 2 Feature selection by cross-validation in a linear model

2.1 Implement an R function that performs feature selection (best subset selection) in linear regression by using k-fold cross-validation without using any specialized function like lm() (use only basic R functions)

```
subset_function <- function(X,Y,N){

# X = swiss[,1:5]
# Y = swiss[,6:6]
# N = 5

df <- cbind(X,Y)
temp <- NULL
final <- NULL

for(i in 1:NCOL(X)){
    combs <- as.data.frame(gtools::combinations(NCOL(X), r=i, v=colnames(X), repeats.allowed=FALSE))
    combs <- tidyr::unite(combs, "formula", sep = ",")</pre>
```

```
temp <- rbind(combs, temp)</pre>
set.seed(12345)
df2 <- df[sample(nrow(df)),]</pre>
df2$k_fold <- sample(N, size = nrow(df), replace = TRUE)</pre>
result <- NULL
for (j in 1:NROW(temp))
  for(i in 1:N){
train = df2[df2$k_fold != i,]
test = df2[df2$k_fold == i,]
vec <- temp[j,]</pre>
train_trimmed = lapply(strsplit(as.character(vec), ","), function(x) train[x])[[1]]
test_trimmed = lapply(strsplit(as.character(vec), ","), function(x) test[x])[[1]]
y_train = train[,c("Y"), drop = FALSE]
y_test = test[,c("Y"), drop = FALSE]
train_trimmed = as.matrix(train_trimmed)
test_trimmed = as.matrix(test_trimmed)
y_test = as.matrix(y_test)
y_train = as.matrix(y_train)
t_train = as.matrix(t(train_trimmed))
t_test = as.matrix(t(test_trimmed))
betas = solve(t_train %*% train_trimmed) %*% (t_train %*% y_train)
train_trimmed = as.data.frame(train_trimmed)
test_trimmed = as.data.frame(test_trimmed)
train_trimmed$type = "train"
test_trimmed$type = "test"
final <- rbind(train_trimmed, test_trimmed)</pre>
y_hat_val = as.matrix(final[,1:(ncol(final)-1)]) %*% betas
mse = (Y - y_hat_val)^2
data <- cbind(i, vec, mse, type = final$type)</pre>
result <- rbind(data, result)</pre>
}
}
result <- as.data.frame(result)
colnames(result) <- c("kfold", "variables", "mse", "type")</pre>
```

```
result$mse <- as.numeric(result$mse)
result$no_variables <- nchar(as.character(result$variables)) - nchar(gsub('\\,', "", result$variables))
variable_performance <- result %>% group_by(kfold, no_variables) %>%
    summarise(MSE = mean(mse, na.rm = TRUE))

myplot <- ggplot(data = variable_performance, aes(x = no_variables, y = MSE, color=kfold)) +
geom_line() + ggtitle("Plot of MSE vs. Number of variables by folds")

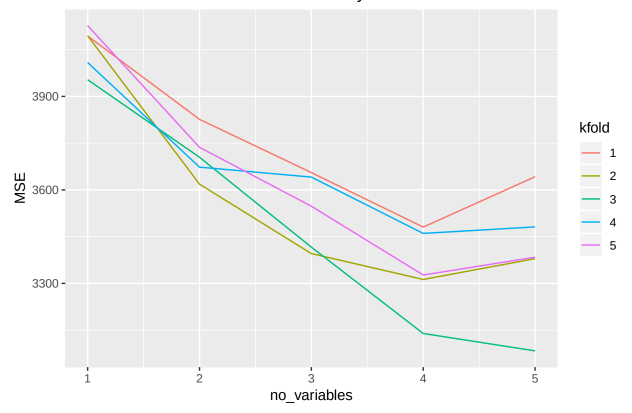
myplot2 <- ggplot(data = result, aes(x = variables, y = mse, color=kfold)) +
geom_bar(stat="identity") + ggtitle("Plot of RMSE vs. Features by folds") + coord_flip()

return(list(myplot, myplot2))
}</pre>
```

2.2 Test your function on data set swiss available in the standard R repository:

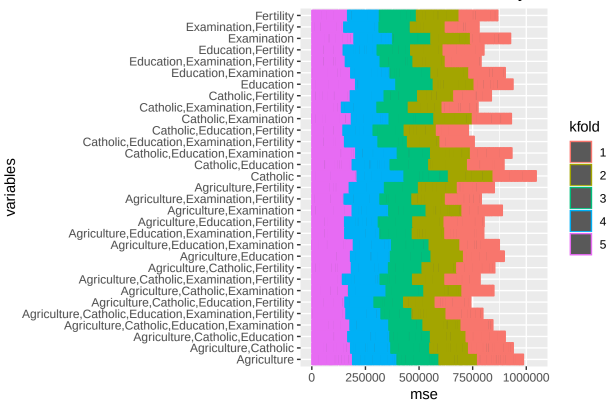
```
subset_function(X = swiss[,1:5], Y = swiss[,6], N = 5)
## [[1]]
```

Plot of MSE vs. Number of variables by folds



[[2]]

Plot of RMSE vs. Features by folds



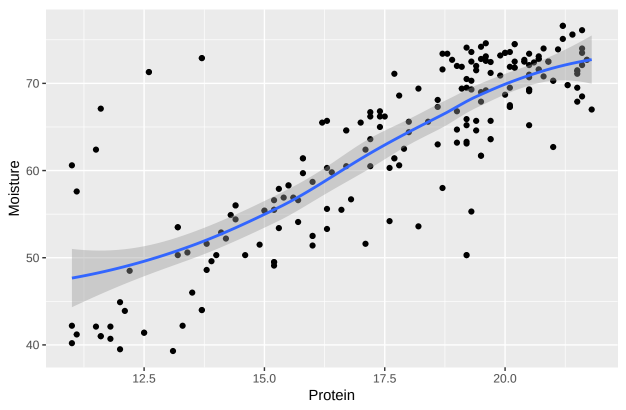
Analysis:

Assignment 3 Linear regression and regularization

3.1 Import data to R and create a plot of Moisture versus Protein. Do you think that these data are described well by linear model.

```
ggplot(data = tecator_data, aes(x = Protein, y = Moisture)) +
  geom_point() +
  geom_smooth( method = 'loess') +
  ggtitle("Plot of Moisture vs. Protein")
```

Plot of Moisture vs. Protein



Analysis: The data seems fairly linear in nature however there are many outliers. As we can see that data is fairly distributed around the line drawn (above and below) thus there is little bias.

3.2 Multiple Models of varying degree.

$$M_{i} = \sum_{i=0}^{p} X^{i} Protein * \beta i + \epsilon$$

$$\epsilon \sim N(0, \sigma^{2})$$

$$\epsilon = M_{i} - \sum_{i=0}^{p} X^{i} Protein * \beta i$$

$$M_{i} \sim N\left(\sum_{i=0}^{p} X^{i} Protein * \beta i, \sigma_{M}^{2}\right)$$
or

$$P\left(M_i|X_{Protein}, \vec{\beta}\right) = N\left(\sum_{i=0}^{p} X^i Protein * \beta i, \sigma_M^2\right)$$

Where,

 σ_M^2 : variance of Moisture

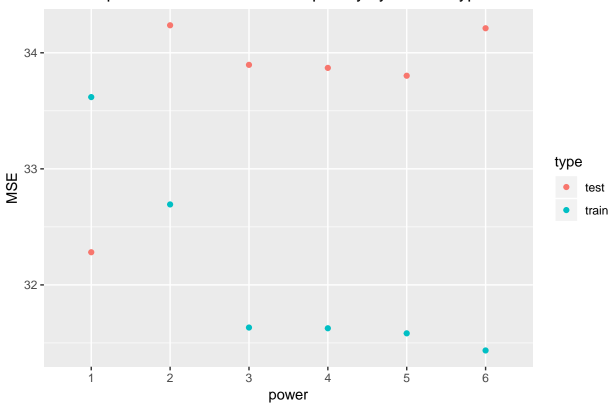
p: degree of the polynomial

3.3 Validation of the Model

```
final_data <- tecator_data
magic_function <- function(df, N)</pre>
df2 <- df
for(i in 2:N)
  df2[paste("Protein_",i,"_power", sep="")] <- (df2$Protein)^i</pre>
df2 <- df2[c("Protein_2_power", "Protein_3_power",</pre>
              "Protein_4_power", "Protein_5_power",
              "Protein_6_power")]
df <- cbind(df,df2)</pre>
return(df)
}
final_data <- magic_function(final_data, 6)</pre>
set.seed(12345)
n = NROW(final_data)
id = sample(1:n, floor(n*0.5))
train = final_data[id,]
test = final_data[-id,]
# model building
M_1 <- lm(data = train, Moisture~Protein)</pre>
M_2 <- lm(data = train, Moisture~Protein+Protein_2_power)</pre>
M_3 <- lm(data = train, Moisture~Protein+Protein_2_power+Protein_3_power)
M_4 <- lm(data = train, Moisture~Protein+Protein_2_power+Protein_3_power+
             Protein_4_power)
M_5 <- lm(data = train, Moisture~Protein+Protein_2_power+Protein_3_power+
             Protein_4_power+Protein_5_power)
M_6 <- lm(data = train, Moisture~Protein+Protein_2_power+Protein_3_power+
            Protein_4_power+Protein_5_power+Protein_6_power)
train$type <- "train"</pre>
test$type <- "test"</pre>
final_data <- rbind(test, train)</pre>
# predicting new values
M_1_predicted <- predict(M_1, newdata = final_data)</pre>
M_2_predicted <- predict(M_2, newdata = final_data)</pre>
```

```
M_3_predicted <- predict(M_3, newdata = final_data)</pre>
M_4_predicted <- predict(M_4, newdata = final_data)</pre>
M_5_predicted <- predict(M_5, newdata = final_data)</pre>
M_6_predicted <- predict(M_6, newdata = final_data)</pre>
# calculating the MSE
final_data$M_1_error <- (final_data$Moisture - M_1_predicted)^2</pre>
final_data$M_2_error <- (final_data$Moisture - M_2_predicted)^2</pre>
final_data$M_3_error <- (final_data$Moisture - M_3_predicted)^2</pre>
final_data$M_4_error <- (final_data$Moisture - M_4_predicted)^2</pre>
final_data$M_5_error <- (final_data$Moisture - M_5_predicted)^2</pre>
final_data$M_6_error <- (final_data$Moisture - M_6_predicted)^2</pre>
# Chainning like Chainsaw
final_error_data <- final_data %>% select(type, M_1_error, M_2_error, M_3_error,
                                            M_4_error, M_5_error, M_6_error) %>%
  gather(variable, value, -type) %>%
  separate(variable, c("model", "power", "error"), "_") %>%
  group_by(type, power) %>%
  summarise(MSE = mean(value, na.rm=TRUE))
ggplot(final_error_data, aes(x = power, y = MSE, color=type)) + geom_point() +
  ggtitle("Mean squared error vs. model complexitiy by dataset type")
```

Mean squared error vs. model complexitiy by dataset type



Analysis: As evident from the plot above, we see that as we increase the model complexitiy (higher powers of the 'protein'), the training error reduces however the model becomes too biased towards the training set

(overfits) and misses the test datasets prediction by larger margins in higher powers.

The best model is M1, that is Moisture~Protein as evident from the least test error (MSE).

The above is a classical case of bias-varience trade-off, which is as follows, as one makes the model fit the trainning dataset better the model becomes more biased and its ability to handle variation to new dataset decreases(varience), thus one should also maintain a good trade off between these two.

3.4 Perform variable selection of a linear model in which Fat is response and Channel1:Channel100 are predicted by using stepAIC.

```
min.model1 = lm(Fat ~ 1, data=tecator_data[,-1])
biggest1 <- formula(lm(Fat ~., data=tecator_data[,-1]))</pre>
step.model1 <- stepAIC(min.model1, direction ='forward', scope=biggest1, trace = FALSE)</pre>
summ(step.model1)
## MODEL INFO:
## Observations: 215
## Dependent Variable: Fat
## Type: OLS linear regression
##
## MODEL FIT:
## F(29,185) = 4775.35, p = 0.00
## R^2 = 1.00
## Adj. R^2 = 1.00
##
## Standard errors: OLS
##
                   Est.
                            S.E. t val.
                            1.59 58.86 0.00 ***
## (Intercept)
                  93.46
## Moisture
                  -1.03
                            0.02 -54.25 0.00 ***
## Protein
                  -0.64
                            0.06 -10.91 0.00 ***
## Channel100
                  66.56
                           48.18
                                   1.38 0.17
## Channel41
                         826.92
                                  -3.95 0.00 ***
               -3268.11
## Channel7
                 -64.03
                           20.80
                                  -3.08 0.00
## Channel48
               -2022.46 254.46
                                  -7.95 0.00 ***
                4934.22 1124.96
## Channel42
                                   4.39 0.00 ***
## Channel50
                1239.52 236.09
                                   5.25 0.00 ***
## Channel45
                4796.22 783.38
                                   6.12 0.00 ***
## Channel66
                2435.79 1169.85
                                   2.08 0.04
## Channel56
                2373.00 540.06
                                   4.39 0.00 ***
## Channel90
                -258.27
                         247.22
                                  -1.040.30
## Channel60
                -264.27
                         708.11
                                  -0.37 0.71
## Channel70
                  14.25
                         327.12
                                   0.04 0.97
## Channel67
               -2015.92
                         543.74
                                  -3.71 0.00 ***
## Channel59
                 635.71
                         996.31
                                   0.64 0.52
## Channel65
                -941.61 1009.23
                                  -0.93 0.35
## Channel58
                1054.24
                         927.95
                                   1.14 0.26
## Channel44
               -5733.84 1079.19
                                  -5.31 0.00 ***
## Channel18
                 299.80
                          88.43
                                   3.39 0.00 ***
## Channel78
                2371.11
                         361.25
                                   6.56 0.00 ***
## Channel84
                -428.99
                         338.35
                                  -1.27 0.21
## Channel62
                3062.97
                        769.59
                                   3.98 0.00 ***
## Channel53
                -804.39
                         203.44
                                  -3.95 0.00 ***
```

```
## Channel75
               -1461.42 402.26
                                 -3.63 0.00 ***
## Channel57
               -3266.79
                         876.71
                                 -3.73 0.00 ***
               -2844.66
## Channel63
                         906.40
                                 -3.14 0.00
## Channel24
                -308.71
                          97.87
                                 -3.15 0.00
## Channel37
                 401.64
                         151.76
                                  2.65 0.01
```

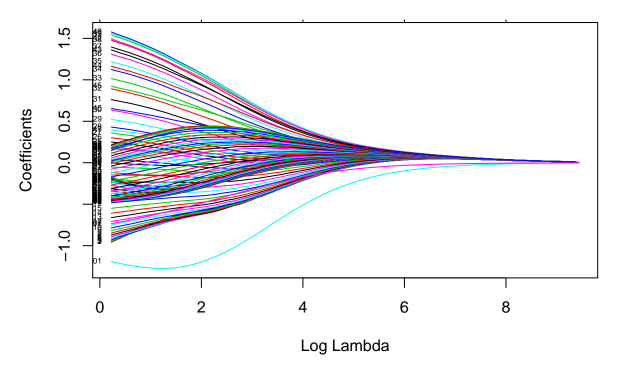
Analysis: 29 variables were choose out of 107. Even among these there are many which have very low p values thus statistically it is a practice to remove variables which are above 0.0005 p values, thus the true variables may not even include these many.

3.5 Fit a Ridge regression model with the same predictor and response

```
y <- tecator_data %>% select(Fat) %>% data.matrix()
x <- tecator_data %>% select(-c(Fat)) %>% data.matrix()

ridge_fit <- glmnet(x, y, alpha = 0, family = "gaussian")
plot(ridge_fit, xvar = "lambda", label = TRUE, main = "Plot showing shrinkage of coefficents with rise</pre>
```

Plot showing shrinkage of coefficents with rise in log of lambda



```
## Chnage of coefficent with respect to lambda
result <- NULL
for(i in 1:100){
temp <- t(coef(ridge_fit, i)) %>% as.matrix()
temp <- cbind(temp, lambda = i)
result <- rbind(temp, result)
}
result <- result %>% as.data.frame() %>% arrange(lambda)
```

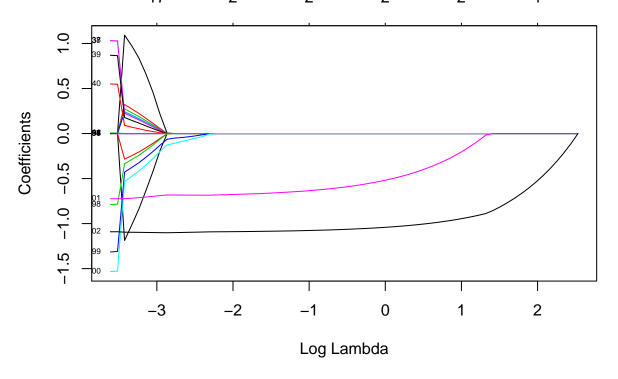
Table 1: Coefficent of Ridge Regression vs. Lambda

Channel1	Channel2	Channel3	Channel4	Protein	lambda
-0.9508076	-0.9588835	-0.9467108	-0.9309440	-1.190667	1
-0.8093030	-0.8337643	-0.8358230	-0.8310237	-1.247494	2
-0.7177628	-0.7473382	-0.7582023	-0.7577328	-1.272872	3
-0.6601701	-0.6865170	-0.7020438	-0.7064153	-1.270796	4
-0.6272862	-0.6550612	-0.6719332	-0.6770183	-1.255186	5
-0.5999460	-0.6206878	-0.6390387	-0.6491696	-1.233227	6
-0.5800181	-0.5980050	-0.6158099	-0.6275854	-1.207645	7
-0.5644019	-0.5799424	-0.5965167	-0.6089792	-1.180696	8
-0.5508957	-0.5643019	-0.5792399	-0.5915380	-1.153425	9
-0.5374226	-0.5490345	-0.5622751	-0.5739280	-1.126430	10

3.6 Fit a Lasso regression model with the same predictor and response

```
lasso_fit <- glmnet(x, y, alpha = 1, family = "gaussian")
plot(lasso_fit, xvar = "lambda", label = TRUE, main = "Plot showing shrinkage of coefficents with rise</pre>
```

Plot showing shrinkage of coefficents with rise in log of lambda



Apendix

```
knitr::opts_chunk$set(echo = TRUE)
if (!require("pacman")) install.packages("pacman")
pacman::p_load(xlsx, glmnet, MASS, jtools, huxtable, ggplot2,
               ggthemes, gridExtra, ROCR, broom, caret, e1071,
               kknn, tidyr, dplyr, reshape2, glmnet)
options("jtools-digits" = 2, scipen = 999)
spam_data <- read.xlsx("spambase.xlsx", sheetName = "spambase_data")</pre>
spam_data$Spam <- as.factor(spam_data$Spam)</pre>
tecator_data <- read.xlsx("tecator.xlsx", sheetName = "data")</pre>
tecator_data <- tecator_data[,2:NCOL(tecator_data)] # removing sample column</pre>
set.seed(12345)
n = NROW(spam_data)
id = sample(1:n, floor(n*0.5))
train = spam_data[id,]
test = spam_data[-id,]
best_model <- glm(formula = Spam ~., family = binomial, data = train)</pre>
summ(best_model)
```

```
# prediction
train$prediction_prob <- predict(best_model, newdata = train, type = "response")</pre>
test$prediction prob <- predict(best model, newdata = test , type = "response")</pre>
train*prediction_class_50 <- ifelse(train*prediction_prob > 0.50, 1, 0)
test$prediction_class_50 <- ifelse(test$prediction_prob > 0.50, 1, 0)
train$prediction_class_90 <- ifelse(train$prediction_prob > 0.90, 1, 0)
test$prediction class 90 <- ifelse(test$prediction prob > 0.90, 1, 0)
# plots
ggplot(train, aes(prediction_prob, color = Spam)) +
geom_density(size = 1) + ggtitle("Training Set's Predicted Score for 50% cutoff") +
  scale_color_economist(name = "data", labels = c("negative", "positive")) +
  theme_economist()
ggplot(test, aes(prediction_prob, color = Spam)) +
geom_density(size = 1) + ggtitle("Test Set's Predicted Score for 50% cutoff") +
  scale_color_economist(name = "data", labels = c("negative", "positive")) +
 theme economist()
#confusion table
conf_train <- table(train$Spam, train$prediction_class_50)</pre>
names(dimnames(conf_train)) <- c("Actual Train", "Predicted Train")</pre>
confusionMatrix(conf_train)
conf_test <- table(test$Spam, test$prediction_class_50)</pre>
names(dimnames(conf_test)) <- c("Actual Test", "Predicted Test")</pre>
confusionMatrix(conf_test)
#confusion table
conf_train1 <- table(train$Spam, train$prediction_class_90)</pre>
names(dimnames(conf_train1)) <- c("Actual Train", "Predicted Train")</pre>
conf_train1
conf_test1 <- table(test$Spam, test$prediction_class_90)</pre>
names(dimnames(conf_test1)) <- c("Actual Test", "Predicted Test")</pre>
conf test1
cutoffs \leftarrow seq(from = 0.05, to = 0.95, by = 0.05)
accuracy <- NULL
for (i in seq_along(cutoffs)){
    prediction <- ifelse(test$prediction_prob >= cutoffs[i], 1, 0) #Predicting for cut-off
    accuracy <- c(accuracy,length(which(test$Spam == prediction))/length(prediction)*100)}</pre>
cutoff_data <- as.data.frame(cbind(cutoffs, accuracy))</pre>
ggplot(data = cutoff_data, aes(x = cutoffs, y = accuracy)) +
  geom line() +
  ggtitle("Cutoff vs. Accuracy for Test Dataset")
```

```
knn_model30 <- train.kknn(Spam ~., data = train, kmax = 30)
train$knn_prediction_class <- predict(knn_model30, train)</pre>
test$knn_prediction_class <- predict(knn_model30, test)</pre>
conf_train2 <- table(train$Spam, train$knn_prediction_class)</pre>
names(dimnames(conf_train2)) <- c("Actual Train", "Predicted Train")</pre>
confusionMatrix(conf train2)
conf_test2 <- table(test$Spam, test$knn_prediction_class)</pre>
names(dimnames(conf_test2)) <- c("Actual Test", "Predicted Test")</pre>
confusionMatrix(conf_test2)
knn_model1 <- train.kknn(Spam ~., data = train, kmax = 1)
train$knn_prediction_class <- predict(knn_model1, train)</pre>
test$knn_prediction_class <- predict(knn_model1, test)</pre>
conf_train2 <- table(train$Spam, train$knn_prediction_class)</pre>
names(dimnames(conf_train2)) <- c("Actual Train", "Predicted Train")</pre>
confusionMatrix(conf_train2)
conf_test2 <- table(test$Spam, test$knn_prediction_class)</pre>
names(dimnames(conf_test2)) <- c("Actual Test", "Predicted Test")</pre>
confusionMatrix(conf_test2)
subset_function <- function(X,Y,N){</pre>
\# X = swiss[,1:5]
# Y = swiss[,6:6]
#N = 5
df <- cbind(X,Y)</pre>
temp <- NULL
final <- NULL</pre>
for(i in 1:NCOL(X)){
combs <- as.data.frame(gtools::combinations(NCOL(X), r=i, v=colnames(X), repeats.allowed=FALSE))</pre>
combs <- tidyr::unite(combs, "formula", sep = ",")</pre>
temp <- rbind(combs, temp)</pre>
}
set.seed(12345)
df2 <- df[sample(nrow(df)),]</pre>
df2$k_fold <- sample(N, size = nrow(df), replace = TRUE)</pre>
result <- NULL
for (j in 1:NROW(temp))
  for(i in 1:N){
train = df2[df2$k_fold != i,]
test = df2[df2$k_fold == i,]
```

```
vec <- temp[j,]</pre>
train_trimmed = lapply(strsplit(as.character(vec), ","), function(x) train[x])[[1]]
test_trimmed = lapply(strsplit(as.character(vec), ","), function(x) test[x])[[1]]
y_train = train[,c("Y"), drop = FALSE]
y_test = test[,c("Y"), drop = FALSE]
train trimmed = as.matrix(train trimmed)
test trimmed = as.matrix(test trimmed)
y_test = as.matrix(y_test)
y_train = as.matrix(y_train)
t_train = as.matrix(t(train_trimmed))
t_test = as.matrix(t(test_trimmed))
betas = solve(t_train %*% train_trimmed) %*% (t_train %*% y_train)
train_trimmed = as.data.frame(train_trimmed)
test_trimmed = as.data.frame(test_trimmed)
train trimmed$type = "train"
test_trimmed$type = "test"
final <- rbind(train_trimmed, test_trimmed)</pre>
y_hat_val = as.matrix(final[,1:(ncol(final)-1)]) %*% betas
mse = (Y - y_hat_val)^2
data <- cbind(i, vec, mse, type = final$type)</pre>
result <- rbind(data, result)</pre>
}
}
result <- as.data.frame(result)</pre>
colnames(result) <- c("kfold", "variables", "mse", "type")</pre>
result$mse <- as.numeric(result$mse)</pre>
result$no_variables <- nchar(as.character(result$variables)) - nchar(gsub('\\,', "", result$variables))
variable_performance <- result %>% group_by(kfold, no_variables) %>%
  summarise(MSE = mean(mse, na.rm = TRUE))
myplot <- ggplot(data = variable_performance, aes(x = no_variables, y = MSE, color=kfold)) +
geom_line() + ggtitle("Plot of MSE vs. Number of variables by folds")
myplot2 <- ggplot(data = result, aes(x = variables, y = mse, color=kfold)) +</pre>
geom_bar(stat="identity") + ggtitle("Plot of RMSE vs. Features by folds") + coord_flip()
return(list(myplot, myplot2))
subset_function(X = swiss[,1:5], Y = swiss[,6], N = 5)
```

```
ggplot(data = tecator_data, aes(x = Protein, y = Moisture)) +
  geom_point() +
  geom_smooth( method = 'loess') +
  ggtitle("Plot of Moisture vs. Protein")
final_data <- tecator_data</pre>
magic function <- function(df, N)</pre>
df2 <- df
for(i in 2:N)
  df2[paste("Protein_",i,"_power", sep="")] <- (df2$Protein)^i
df2 <- df2[c("Protein_2_power", "Protein_3_power",</pre>
              "Protein_4_power", "Protein_5_power",
              "Protein_6_power")]
df <- cbind(df,df2)</pre>
return(df)
}
final_data <- magic_function(final_data, 6)</pre>
set.seed(12345)
n = NROW(final data)
id = sample(1:n, floor(n*0.5))
train = final_data[id,]
test = final_data[-id,]
# model building
M_1 <- lm(data = train, Moisture~Protein)</pre>
M_2 <- lm(data = train, Moisture~Protein+Protein_2_power)</pre>
M_3 <- lm(data = train, Moisture~Protein+Protein_2_power+Protein_3_power)
M_4 <- lm(data = train, Moisture~Protein+Protein_2_power+Protein_3_power+
             Protein_4_power)
M_5 <- lm(data = train, Moisture~Protein+Protein_2_power+Protein_3_power+
             Protein_4_power+Protein_5_power)
M_6 <- lm(data = train, Moisture~Protein+Protein_2_power+Protein_3_power+
            Protein_4_power+Protein_5_power+Protein_6_power)
train$type <- "train"</pre>
test$type <- "test"</pre>
final_data <- rbind(test, train)</pre>
# predicting new values
M_1_predicted <- predict(M_1, newdata = final_data)</pre>
M_2_predicted <- predict(M_2, newdata = final_data)</pre>
M_3_predicted <- predict(M_3, newdata = final_data)</pre>
M_4_predicted <- predict(M_4, newdata = final_data)</pre>
M_5_predicted <- predict(M_5, newdata = final_data)</pre>
```

```
M_6_predicted <- predict(M_6, newdata = final_data)</pre>
# calculating the MSE
final_data$M_1_error <- (final_data$Moisture - M_1_predicted)^2</pre>
final_data$M_2_error <- (final_data$Moisture - M_2_predicted)^2</pre>
final_data$M_3_error <- (final_data$Moisture - M_3_predicted)^2</pre>
final_data$M_4_error <- (final_data$Moisture - M_4_predicted)^2</pre>
final_data$M_5_error <- (final_data$Moisture - M_5_predicted)^2</pre>
final_data$M_6_error <- (final_data$Moisture - M_6_predicted)^2</pre>
# Chainning like Chainsaw
final_error_data <- final_data %>% select(type, M_1_error, M_2_error, M_3_error,
                                           M_4_error, M_5_error, M_6_error) %>%
  gather(variable, value, -type) %>%
  separate(variable, c("model", "power", "error"), "_") %>%
  group_by(type, power) %>%
  summarise(MSE = mean(value, na.rm=TRUE))
ggplot(final_error_data, aes(x = power, y = MSE, color=type)) + geom_point() +
  ggtitle("Mean squared error vs. model complexitiy by dataset type")
min.model1 = lm(Fat ~ 1, data=tecator_data[,-1])
biggest1 <- formula(lm(Fat ~., data=tecator_data[,-1]))</pre>
step.model1 <- stepAIC(min.model1, direction ='forward', scope=biggest1, trace = FALSE)</pre>
summ(step.model1)
y <- tecator data %>% select(Fat) %>% data.matrix()
x <- tecator_data %>% select(-c(Fat)) %>% data.matrix()
ridge_fit <- glmnet(x, y, alpha = 0, family = "gaussian")</pre>
plot(ridge_fit, xvar = "lambda", label = TRUE, main = "Plot showing shrinkage of coefficents with rise
## Chnage of coefficent with respect to lambda
result <- NULL
for(i in 1:100){
temp <- t(coef(ridge_fit, i)) %>% as.matrix()
temp <- cbind(temp, lambda = i)</pre>
result <- rbind(temp, result)</pre>
result <- result %>% as.data.frame() %>% arrange(lambda)
table_cofe <- head(result,10) %>% select(Channel1, Channel2,
                                           Channel3, Channel4,
                                           Protein, lambda)
knitr::kable(table_cofe, caption = "Coefficent of Ridge Regression vs. Lambda")
lasso_fit <- glmnet(x, y, alpha = 1, family = "gaussian")</pre>
plot(lasso_fit, xvar = "lambda", label = TRUE, main = "Plot showing shrinkage of coefficents with rise
```