# TDDE01 – Machine Learning Individual Laboration Report 2

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#### 1 Assignment 1

In this assignment we were tasked with implementing feature selection using the k-fold cross validation and linear regression algorithms. The result of this can be found in appendix: A - Code assignment 1.

The feature selection algorithm iterates through all possible combinations of *features* from the predictor variables and apply the *k-fold* cross validation on all of them. The feature combination with the lowest *sum of squared error* gets picked as the best combination.

$$\mathbf{A} = \{c_1, c_2, ..., c_n\}, 1 \le n \le ncol(X) \tag{1}$$

A in the equation above represents all the permutations of column index for the response variable. Each of theses combinations are sent to the k-fold cross validation where the data is split into K parts of equal size. The K are then iterated through and each data subset is used as testing data with the other K-1 sets as training data. The k-fold cross validation uses the linear regression with ordinary least squares estimator. The produced predictions are then error checked with the sum of squared error

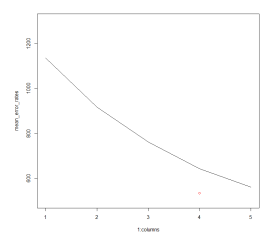
$$\sum_{i=1}^{n} (\hat{y}_i - y_i), n = nrow(y)$$
(2)

for fold K. After all the K folds for a given combination of features are calculated, the mean of all fold errors are calculated and returned as the final error value for the feature combination.

$$\frac{1}{K} \sum_{k=1}^{K} \left( \sum_{i=1}^{n} (\hat{y}_i - y_i) \right) \tag{3}$$

where K = |folds| and n = |y|.

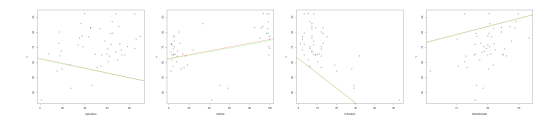
In the figure below the mean error for a given number of selected features can be shown.



Figur 1: The mean error rate for different number of feature combinations.

The red dot indicates the best feature combination, i.e. the one with the lowest error score. For this data set it's {Agriculture, Education, Catholic, Infant.Mortality}. In general terms we can observe that the more features that are selected, the lower the error rate becomes.

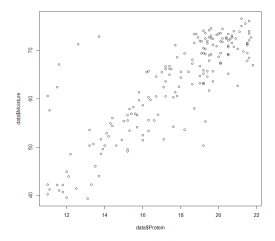
The following graphs show the linear regression result for each of the different features that was returned by the best feature subset function described above.



The models aren't perfect but manges to hit fairly close to what humans would recognize as the best linear function for each data set.

### 2 Assignment 2

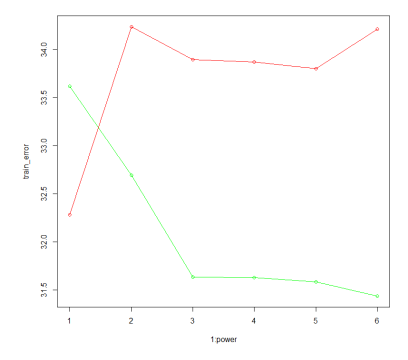
In this assignment we are tasked with analyzing the data set found in the file **tecator.xlsx**. The data is grouped into the columns: {Sample, Channel1, ..., Channel100, Moisur The first task is to plot the Moisture and Protein and determine if a linear model would be a good approximation for the data. The plot can be seen in the figure below.



Figur 2: The *Moisture* plotted by *Protein*.

A linear approximation seems to be a good fit for the data.

Next we examine if a polynomial approximation of the data up to the nth power and using mean squared error (MSE) to examine the fitness of the polynomial model. The data set is split into two sets of equal size where one of the sets are the training set and the other the validation set.



Figur 3: The training and validation error rate for the polynomial estimation given the MSE.

The **green line** is the error rate for the training data and the **red line** is the error rate for the validation set over the different power-levels. We can observe in the graph that the fitting of the training data decreases when the power-level increases but the error rate for the validation data increases with about the same magnitude. This can be explained with the fact that, as observed in the plot of the full data set above, the data would follow a linear model and the polynomial approximation leads to over fitting of the training set.

 ${f T.B.C}$  The next part involves performing variable selection on the full data set with the step variable selection function. We get 64 coefficients form the step function.

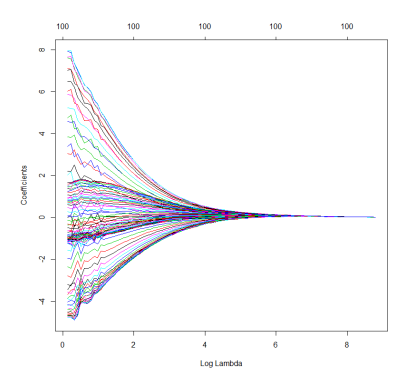
The next part involves fitting a ridge regression model to the data set.

$$\min \sum_{i=1}^{n} V(\hat{x}_{i} \cdot w, \hat{y}_{i}) + \lambda \|w\|_{2}^{2}$$
(4)

The equations results in the ridge estimation to intersect the linear regression

estimation in an edge point. This property will result in not many coefficients being zeroed although many will be very small.

The result can be observed in the following plot. We can see as the lambda increases, the coefficients converges to zero. In total we get 64 coefficients from the ridge estimation.



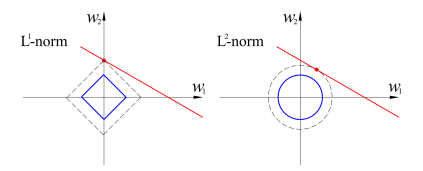
Figur 4: Size of the coefficients over the log of the penalty factor lambda.

We can observe that the number of coefficients stay fairly high over all the lambdas but the penalty factor causes the size of the coefficients to decrease.

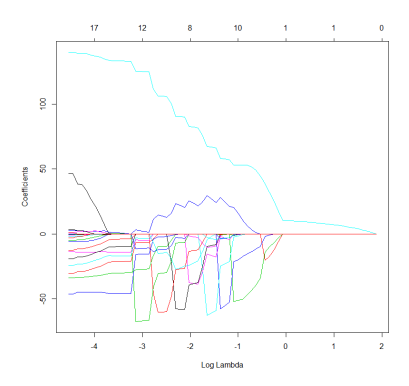
The lasso regression model results in fewer coefficients and less more non continuous coefficient functions over the lambdas.

$$\min \sum_{i=1}^{n} V(\hat{x}_i \cdot w, \hat{y}_i) + \lambda \|w\|$$
 (5)

The equation is similar to the ridge estimation with the big difference being that the estimation bound has a different shape where coefficients can be zeroed. This can be observed in the figure below.



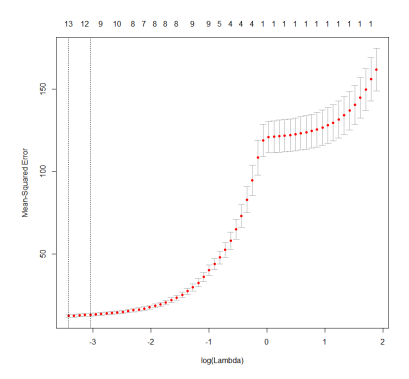
Figur 5: Shape of the Ridge and LASSO estimation bounds respectively, credit to Rezamohammadighazi for original image.



Figur 6: Size of the coefficients over the log of the penalty factor lambda.

As observed with the ridge regression the coefficients decrease over log lambda but the number of coefficients have drastically decreased and many coefficient functions jumps erratic in magnitude. This is caused by the shape of the lasso estimation where model intersection in the corners will cause one of the coefficients to be equal to zero.

The final part involves plotting to cross validation score using the lasso model, the result can be observed in the graph below. The variance increase with the lambda and the validation score.



Figur 7: Mean squared error over the log lambdas with variance included.

In comparison with the step aic model the cross validation using lasso and k-fold resulted in much less coefficients.

### 3 Appendix: A - Code assignment 1

Listing 1: Code for assignment 1

library(readxl)

```
# folding indexes, returns start indexes for each
   fold
indexes <- function(n,k){</pre>
  s = floor(n/k)
  indexes = matrix(1,k,2)
  for(i in 1:k){
    indexes[i,1] = (i-1) * s
    indexes[i,2] = i * s -1
  }
  indexes = indexes + 1
  indexes[k,2] = n
  return(indexes)
}
# Deprecated because of reasons
binary_permutations <- function(n){</pre>
  indexes = matrix(0,2^n-1,n)
  for(i in 1:2<sup>n-1</sup>){
    indexes[i,] = as.numeric(intToBits(i))[1:n]
  }
  return(indexes)
}
k_fold <- function(X,Y,K){</pre>
  n = nrow(X)
  fold_errors = matrix(0,K,1)
  fold_weights = matrix(0,K,(ncol(X)+1))
  indexes = indexes(n,K)
  for( i in 1:K){
    test_fold_indexes = indexes[i,1]:indexes[i,2]
    train_fold_indexes = (1:n)[-test_fold_indexes]
    test_y = Y[test_fold_indexes]
    test_x = X[test_fold_indexes,]
    train_x = X[train_fold_indexes,]
    train_y = Y[train_fold_indexes]
    result = linear_regression(as.matrix(train_x),
       train_y, as.matrix(test_x), test_y)
    fold_errors[i,] = result$err
    fold_weights[i,] = result$param
  }
```

```
return (list(weights=colMeans(fold_weights), err=
    fold_errors))
best_subset <- function(X,Y,K){</pre>
 n = nrow(X)
  columns = ncol(X)
  print(columns)
  errors = matrix(0,columns,2^columns)
  mean_error_rates = matrix(0,columns,1)
  best_error = Inf
  best_indexes = c()
  best_weights = c()
  for(n_features in 1:columns){
    binary_permutations = binary_permutations(ncol(X
    binary_permutations = t(combn(1:columns,n_
       features))
    n_combinations = nrow(binary_permutations)
    combination_errors = matrix(Inf,n_combinations
       , 1)
    for(combination in 1:n_combinations){
      current_features = binary_permutations[
         combination,]
      filtered_x = as.matrix(X[current_features])
      k_fold = k_fold(filtered_x,Y,K)
      combination_errors[combination,] = mean(k_fold
         $err)
      best_combination_index = which.min(combination
         _errors)
      if(combination_errors[best_combination_index,]
          < best_error){
        best_weights = k_fold$weights
        best_indexes = binary_permutations[
           combination,]
        best_error = combination_errors[best_
           combination_index,]
```

```
}
    }
    mean_error_rates[n_features,] = mean(combination
      _errors)
  plot(1:columns, mean_error_rates, type = "1", ylim
    = c(500,1300)
  points(length(best_indexes), best_error, col="Red")
  print(best_error)
  return(list(indexes=best_indexes, weights=best_
    weights, err=best_error))
}
 # Linear regression between two samples, one as x-
   values and one as y-values
 linear_regression <- function(x_train,y_train,x_</pre>
   test,y_test){
    x_train = cbind(matrix(1,nrow(x_train),1),x_
       train)
    x_test = cbind(matrix(1,nrow(x_test),1),x_test)
    w = solve(t(x_train) %*% x_train) %*% t(x_train)
       %*% y_train
    w = as.vector(w)
    pY = x_test %*% w
    errors = sum((y_test - pY)^2)
   return(list(param=w, pred=pY, err=errors))
 }
data = swiss
set.seed(12345)
s = sample(1:nrow(data),nrow(data))
data = data[s,]
# ========
#fold_indexes(nrow(data),5)
X = data[,2:ncol(data)]
Y = as.matrix(data$Fertility)
best_features = best_subset(X,Y,5)
```

## 4 Appendix: B - Code assignment 2

Listing 2: Code for assignment2

```
library (MASS)
library(readxl)
library(Matrix)
library(glmnet)
data = read_excel("tecator.xlsx")
plot(data$Protein,data$Moisture)
# answer: yes
# task 2
# Find a probablist model explaining Mi. where M is
   a polynomial model of the protein up to power i
# Why is it important to use mean squared error whne
    fitting model?
# We approximate the model as w0 + w1x^1 + w2x^2 \dots
# Use MSE instead of SSE because MSE is normalized
set.seed(12345)
n = nrow(data)
```

```
train_indexes = sample(1:n,floor(n*0.5))
train_data = data[train_indexes,]
test_data = data[-train_indexes,]
poly = 6
train_error = matrix(0,poly,1)
test_error = matrix(0,poly,1)
for(i in 1:poly) {
 model = lm(Moisture ~ poly(Protein,i), data=train_
  train_predictions = predict(model,train_data)
  test_predictions = predict(model,test_data)
 train_error[i,] = mean((train_data$Moisture -
    train_predictions)^2)
  test_error[i,] = mean((test_data$Moisture - test_
    predictions)^2)
}
vlim = c(min(rbind(train_error, test_error)), max(
  rbind(train_error, test_error)))
plot(1:poly,train_error, col="Green", ylim=ylim)
lines(1:poly,train_error, col="Green")
points(1:poly,test_error,col="Red")
lines(1:poly,test_error, col="Red")
# Observation of the plot indicates that as we
   increase the polynomial level the function
  becomes more fitted for the training data and
  results in increasingly worse fit for the test
  data
model = lm(Fat ~ . - Protein - Moisture - Sample,
  data=data)
steps = stepAIC(model,direction="both", trace=FALSE)
coeff_aics = steps$coefficients
n_coeff_aics = length(coeff_aics)
print(n_coeff_aics)
# 64 columns were selected
data_y = data\$Fat
```

```
data = data[,-which(colnames(data) == "Sample")]
data = data[,-which(colnames(data) == "Fat")]
data = data[,-which(colnames(data) == "Protein")]
data = as.matrix(data[,-which(colnames(data) == "
  Moisture")])
ridge = glmnet(x=data, y=data_y, alpha=0)
plot(ridge, xvar="lambda")
# task 6
# Do the same stuff but with LASSO, compare with
lasso = glmnet(x=data, y=data_y, alpha=1)
plot(lasso, xvar="lambda")
# task 7
    CV to find the optimal LASSO, report the optimal
   lambda and how many variables were chosen by the
   model '
    make conculsions
   show a plot of CV scores in comparasion to
   lambda
lasso_cv = cv.glmnet(data,data_y, alpha=1)
lasso_cv_lambda = min(lasso_cv$lambda)
n_lass_cv = sum(coef(lasso_cv) != 0)
print(lasso_cv_lambda)
print(n_lass_cv)
plot(lasso_cv)
# task 8
# compare result from 4 and 7
```