# 732A99\_lab2\_block1\_A2

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A:
Assignment 3
<ol> <li>Reorder your data with respect to the increase of MET and plot EX versus MET. Discuss what kind of model can be appropriate here. Use the reordered data in steps 2-5.</li> <li>Use package tree and fit a regression tree model with target EX and feature MET in which the</li> </ol>
number of the leaves is selected by cross-validation, use the entire data set and set minimum number of observations in a leaf equal to 8 (setting minsize in tree.control). Report the selected tree. Plot the original and the fitted data and histogram of residuals. Comment on the distribution of the residuals and the quality of the fit.
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whether results of the regression model in step 2 seem to be reliable
4.Compute and plot the $95%$ confidence and prediction bands the regression tree model from step
2 (fit a regression tree with the same settings and the same number of leaves as in step 2 to the resampled data) by using a parametric bootstrap, assume Normal distribution with mean as labels in the tree leaves, while varience is residual varience. Consider the width of the
confidence band and comment whether results of the regression model in step 2 seem to be
reliable. Does it look like only $5\%$ of data are outside the prediction band? Should it be?
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more appropriate here
Assignment 4

1.	Conduct a standard PCA by using the feature space and provide a plot explaining how much	
	variation is explained by each feature. Does the plot show how many PC should be extracted?	
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#### ${\bf Contributions}$

During the lab, Anubhav focused on assignment 2, Lennart focused on assignment 3 and Thijs focused on assignment 4. All codes and analysis was indepedently done and is also reflected in the individual reports.

#### Loading The Libraries

#### Assignment 2

2.1 Import the data to R and divide into training/validation/test as 50/25/25: use data partitioning code specified in Lecture 1e.

```
set.seed(12345)

data <- read.xlsx("creditscoring.xls", sheetName = "credit")
# changing class of dependent variable good_bad
data$good_bad = as.factor(data$good_bad)
# dividing data into train (50%), validation (25%) and test set (25%)
n = dim(data)[1]
set.seed(12345)
id = sample(1:n, floor(n*0.5))
train = data[id,]
id1 = setdiff(1:n, id)
set.seed(12345)
id2 = sample(id1, floor(n*0.25))
valid = data[id2,]
id3 = setdiff(id1,id2)
test = data[id3,]</pre>
```

2.2 Fit a decision tree to the training data by using the following measures of impurity: a. Deviance b. Gini index and report the misclassification rates for the training and test data. Choose the measure providing the better results for the following steps.

```
set.seed(12345)

fitDecisionTree = function(testData, splitMethod) {
    # fitting decision tree
    decisionTreeFit = tree(formula = good_bad ~ ., data = train, split = splitMethod)
    # getting prediction results
    yFit = predict(object = decisionTreeFit, newdata = testData, type = "class")
    # printing misclassification rate
    pasteO("Misclassification rate: ", mean(yFit != as.character(testData$good_bad)))
```

```
}
fitDecisionTree(testData = train, splitMethod = "deviance")
## [1] "Misclassification rate: 0.212"
fitDecisionTree(testData = test, splitMethod = "deviance")
## [1] "Misclassification rate: 0.268"
fitDecisionTree(testData = train, splitMethod = "gini")
## [1] "Misclassification rate: 0.236"
fitDecisionTree(testData = test, splitMethod = "gini")
## [1] "Misclassification rate: 0.36"
# fitting tree with measure providing better results ("deviance")
unprunedTree = tree(formula = good_bad ~ ., data = train, split = "deviance")
# Create a decision tree model
credit_tree_deviance <- tree(good_bad~., data=train, split = c("deviance"))</pre>
credit_tree_gini <- tree(good_bad~., data=train, split = c("gini"))</pre>
# predicting on the test dataset to get the misclassification rate.
predict_tree_deviance <- predict(credit_tree_deviance, newdata = test, type = "class")</pre>
predict_tree_gini <- predict(credit_tree_gini, newdata = test, type = "class")</pre>
conf_tree_deviance <- table(test$good_bad, predict_tree_deviance)</pre>
names(dimnames(conf_tree_deviance)) <- c("Actual Test", "P2redicted Test")</pre>
caret::confusionMatrix(conf_tree_deviance)
## Confusion Matrix and Statistics
##
              P2redicted Test
##
## Actual Test bad good
                28
##
          bad
                     48
          good 19 155
##
##
##
                  Accuracy: 0.732
                    95% CI: (0.6725, 0.7859)
##
##
       No Information Rate: 0.812
       P-Value [Acc > NIR] : 0.9992613
##
##
##
                     Kappa: 0.2904
   Mcnemar's Test P-Value: 0.0006245
##
##
               Sensitivity: 0.5957
##
##
               Specificity: 0.7635
            Pos Pred Value: 0.3684
##
            Neg Pred Value: 0.8908
##
                Prevalence: 0.1880
##
##
            Detection Rate: 0.1120
```

```
##
      Detection Prevalence: 0.3040
##
         Balanced Accuracy: 0.6796
##
##
          'Positive' Class : bad
conf_tree_gini <- table(test$good_bad, predict_tree_gini)</pre>
names(dimnames(conf_tree_gini)) <- c("Actual Test", "Predicted Test")</pre>
caret::confusionMatrix(conf tree gini)
  Confusion Matrix and Statistics
##
##
              Predicted Test
## Actual Test bad good
##
          bad
                17
                     59
##
          good 36
                   138
##
##
                  Accuracy: 0.62
                    95% CI: (0.5567, 0.6804)
##
##
       No Information Rate: 0.788
       P-Value [Acc > NIR] : 1.000
##
##
##
                     Kappa: 0.0184
    Mcnemar's Test P-Value : 0.024
##
##
##
               Sensitivity: 0.3208
##
               Specificity: 0.7005
##
            Pos Pred Value: 0.2237
            Neg Pred Value: 0.7931
##
##
                Prevalence: 0.2120
##
            Detection Rate: 0.0680
##
      Detection Prevalence: 0.3040
##
         Balanced Accuracy: 0.5106
##
##
          'Positive' Class : bad
##
```

Analysis: On the Training dataset model with 'deviance' had a misclassification rate of 21.05% while the model with 'gini' split had the misclassification rate of 23.68%.

For the test dataset we see that the model with 'deviance' type of split has a accuracy of 73.2% or misclassification rate of 26.8%, we see that to predict 'good' the accuracy is 89.08% but for predicting bad its just 36.84%. Thus our our model is heavily baised towards predicting cases as 'good'.

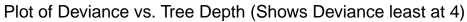
For the test dataset we see that the model with 'gini' type of split has a accuracy of 63.6% or misclassifiaction rate of 36.4%, we also see that to predict 'good' the accuracy is 81.03% but for predicting bad its just 18.97%. Thus our model is heavily baised towards predicting cases as 'good' even more than the model which uses 'deviance' to split variable.

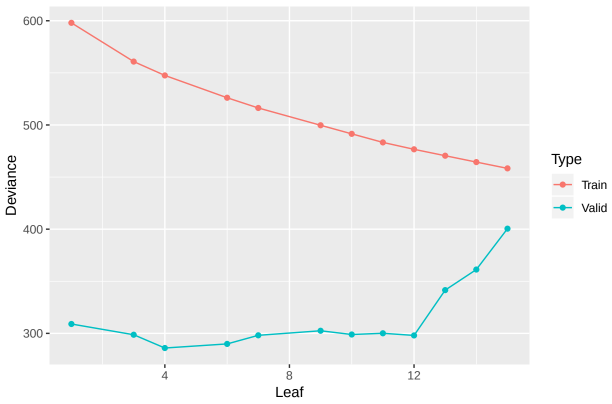
Both our models would lead to many bad loan applicants to be given loans which is never a good thing, however among the model the one using 'deviance' mode for split is better by 9.6%.

Thus we will select model using 'deviance' for further model building.

3. Use training and validation sets to choose the optimal tree depth. Present the graphs of the dependence of deviances for the training and the validation data on the number of leaves. Report the optimal tree, report it's depth and the variables used by the tree. Interpret the information provided by the tree structure. Estimate the misclassification rate for the test data.

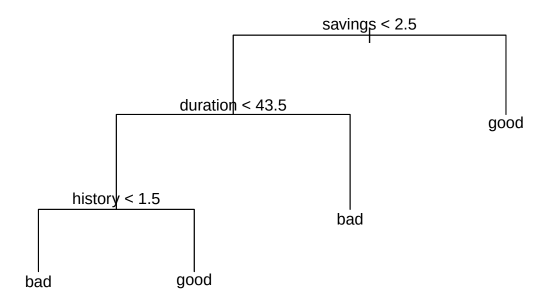
```
set.seed(12345)
credit_tree <- tree(good_bad~., data=train, split = c("deviance"))</pre>
credit tree purned train <- prune.tree(credit tree, method = c("deviance"))</pre>
credit_tree_purned_valid <- prune.tree(credit_tree, newdata = valid ,method = c("deviance"))</pre>
result_train <- cbind(credit_tree_purned_train$size,</pre>
                       credit_tree_purned_train$dev, "Train")
result_valid <- cbind(credit_tree_purned_valid$size,</pre>
                       credit tree purned valid$dev, "Valid")
result <- as.data.frame(rbind(result_valid, result_train))</pre>
colnames(result) <- c("Leaf", "Deviance", "Type")</pre>
result$Leaf <- as.numeric(as.character(result$Leaf))</pre>
result$Deviance <- as.numeric(as.character(result$Deviance))</pre>
# plot of deviance vs. number of leafs
ggplot(data = result, aes(x = Leaf, y = Deviance, colour = Type)) +
  geom_point() + geom_line() +
  ggtitle("Plot of Deviance vs. Tree Depth (Shows Deviance least at 4)")
```





```
# prune the tree to the required depth
credit_tree_sniped <- prune.tree(credit_tree, best=4)

plot(credit_tree_sniped)
text(credit_tree_sniped)</pre>
```



```
# misclassification rate for best pruned tree
result_prune_test <- predict(credit_tree_sniped, newdata = test, type = "class")
conf_prune_tree_test <- table(test$good_bad, result_prune_test)</pre>
names(dimnames(conf_prune_tree_test)) <- c("Actual Test", "Predicted Test")</pre>
caret::confusionMatrix(conf_prune_tree_test)
## Confusion Matrix and Statistics
##
##
              Predicted Test
## Actual Test bad good
                18
##
          bad
##
          good
                 6 168
##
##
                  Accuracy: 0.744
##
                    95% CI: (0.6852, 0.7969)
##
       No Information Rate: 0.904
       P-Value [Acc > NIR] : 1
##
##
##
                     Kappa: 0.2507
   Mcnemar's Test P-Value : 0.00000000183
##
##
##
               Sensitivity: 0.7500
##
               Specificity: 0.7434
##
            Pos Pred Value: 0.2368
```

```
## Neg Pred Value : 0.9655
## Prevalence : 0.0960
## Detection Rate : 0.0720
## Detection Prevalence : 0.3040
## Balanced Accuracy : 0.7467
##
## 'Positive' Class : bad
##
```

Analysis: Choosing optimal depth tree we get that '4' as the best depth. The variables used in the best tree are-duration, history, savings.

From the tree structure we can see that the following variables are best to split on, 'savings' < 2.5 then 'duration' < 43.5 and 'history' < 1.5.

The accuracy on the model trained on 'train' dataset is accuracy 73.2% and misclassification 26.8% while on the 'test' dataset accuracy is 74.4%, thus the misclassification rate is 25.6%. We see that model predicts 'good' applicants very well (accuracy of 96.55%) while it classifies 'bad' applicant way badly (accuracy is 23.68%).

Thus this model would be very bad for the business and would likely to run the business bankrupt.

4. Use training data to perform classification using Naïve Bayes and report the confusion matrices and misclassification rates for the training and for the test data. Compare the results with those from step 3.

```
#Fitting the Naive Bayes model
credit_naive_model = naiveBayes(good_bad ~., data=train)
#Prediction on the dataset
predict_naive_train = predict(credit_naive_model, newdata=train, type = "class")
predict_naive_test = predict(credit_naive_model, newdata=test, type = "class")
conf_naive_train <- table(train$good_bad, predict_naive_train)</pre>
names(dimnames(conf_naive_train)) <- c("Actual Train", "Predicted Train")</pre>
caret::confusionMatrix(conf_naive_train)
## Confusion Matrix and Statistics
##
##
               Predicted Train
##
  Actual Train bad good
##
           bad
                 95
                      52
##
           good 98
                     255
##
##
                  Accuracy: 0.7
##
                    95% CI: (0.6577, 0.7399)
##
       No Information Rate: 0.614
       P-Value [Acc > NIR] : 0.00003655
##
##
##
                     Kappa: 0.3378
   Mcnemar's Test P-Value: 0.0002386
##
##
##
               Sensitivity: 0.4922
##
               Specificity: 0.8306
            Pos Pred Value: 0.6463
##
```

```
##
            Neg Pred Value: 0.7224
##
                Prevalence: 0.3860
            Detection Rate: 0.1900
##
##
      Detection Prevalence: 0.2940
##
         Balanced Accuracy: 0.6614
##
          'Positive' Class : bad
##
##
conf_naive_test <- table(test$good_bad, predict_naive_test)</pre>
names(dimnames(conf_naive_test)) <- c("Actual Test", "Predicted Test")</pre>
caret::confusionMatrix(conf_naive_test)
## Confusion Matrix and Statistics
##
##
              Predicted Test
  Actual Test bad good
##
          bad
                46
                     30
##
          good
                49
                    125
##
##
                  Accuracy: 0.684
                    95% CI : (0.6224, 0.7411)
##
##
       No Information Rate: 0.62
##
       P-Value [Acc > NIR] : 0.02075
##
##
                      Kappa: 0.3024
    Mcnemar's Test P-Value: 0.04285
##
##
##
               Sensitivity: 0.4842
##
               Specificity: 0.8065
##
            Pos Pred Value: 0.6053
##
            Neg Pred Value: 0.7184
##
                Prevalence: 0.3800
##
            Detection Rate: 0.1840
      Detection Prevalence: 0.3040
##
##
         Balanced Accuracy: 0.6453
##
##
          'Positive' Class : bad
##
```

#### Analysis:

For the train dataset using NaiveBayes method we get accuracy 70% or misclassification of 30%, here we also notice that the accuracy of class 'bad' is 64.63% while for class 'good' is 72.24%, thus the model is more balanced in predicting, thus its still baised in predict one class over the other.

For the test dataset using NaiveBayes method we get accuracy 68.4% or misclassification of 31.6%, here we also notice that the accuracy of class 'bad' is 60.53% while for class 'good' is 71.84%, thus the model is almost the same compared to train.

Compared to step3, we see that for the 'train' dataset the optimal tree has accuracy of 73.2% while it is 74.4% on the 'test' dataset. For the NaiveBayes model, accuracy on the 'train' dataset is 70% and while it is 68.4% on the 'test' dataset.

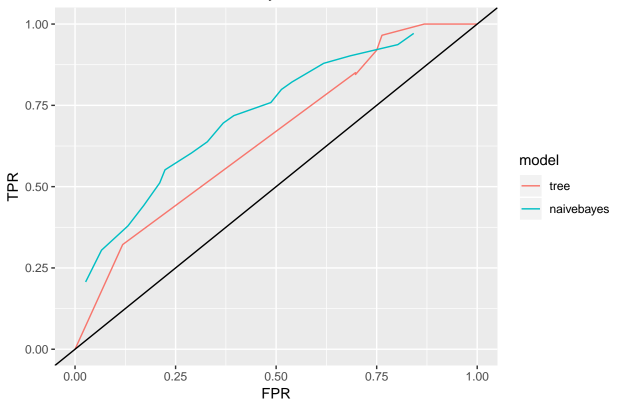
Accuracy is only part of the story what we see is better here is that this model classifies 'bad' customers better better for both train and test dataset than decision tree (60+% for both train and test for naive compared 36.84% train, 23.68% test for decision tree).

Thus the model is better to be used for the business than the one in the step3, the risk of providing loans to bad applicant is lesser than the previous model but its still not good enough!

5. Use the optimal tree and the Naïve Bayes model to classify the test data by using the following principle: where prob(Y|'good')=A, where A=0.05,0.10,....0.95. Compute the TPR and FPR values for the two models and plot the corresponding ROC curves. Conclusion?

```
set.seed(12345)
credit tree <- tree(good bad~., data=train, split = c("deviance"))</pre>
credit_naive_model = naiveBayes(good_bad ~., data=train)
# prune the tree to the required depth
credit_tree_sniped <- prune.tree(credit_tree, best=7)</pre>
# predicting class, getting probability
predict_prune_test_prob <- predict(credit_tree_sniped, newdata = test)</pre>
predict_naive_test_prob <- predict(credit_naive_model, newdata=test, type = "raw")</pre>
# data mugging
probability_data_naive <- as.data.frame(cbind(predict_naive_test_prob,</pre>
                                                 as.character(test$good_bad), "naivebayes"))
probability_data_tree <- as.data.frame(cbind(predict_prune_test_prob,</pre>
                                                as.character(test$good bad), "tree"))
probability_data_combined <- rbind(probability_data_tree, probability_data_naive)</pre>
colnames(probability_data_combined) <- c("prob_bad", "prob_good",</pre>
                                            "actual_test_class", "model")
# final dataset
probability_data_combined$prob_good <- as.numeric(as.character(probability_data_combined$prob_good))</pre>
# changing the threshold and printing the probability
tree_list <- NULL</pre>
naive list <- NULL
final <- NULL
for(threshold in seq(from = 0.05, to = 0.95, by = 0.05)){
  probability_data_combined$predicted_class <- ifelse(probability_data_combined$prob_good > threshold,
  df2 <- probability_data_combined[,c("model", "actual_test_class", "predicted_class")]</pre>
  df2$threshold <- threshold
  df2\statch <- ifelse(df2\statch actual_test_class == df2\statch predicted_class, 1, 0)
  final <- rbind(df2, final)</pre>
# Creating the FRP and TRP for each model and threshold
final$temp <- 1</pre>
final_summary <- final %>%
group_by(model, threshold) %>%
```

#### ROC curve for the Naive Bayes vs. Tree Model



Analysis: We find that 'naivebayes' model is better than 'tree' model for almost across varying threshold values, except the higher values of threshold (0.75+). Thus the decision of using NaiveBayes model here is further reinforced using the ROC curve.

6. Repeat Naïve Bayes classification as it was in step 4 but use the following loss matrix (good loss 1, bad loss 10) and report the confusion matrix for the training and test data. Compare the results with the results from step 4 and discuss how the rates has changed and why.

```
set.seed(12345)

credit_naive_model = naiveBayes(good_bad ~., data=train)

# predicting class, getting probability
```

```
predict_naive_train_prob <- predict(credit_naive_model, newdata=train, type = "raw")</pre>
predict_naive_test_prob <- predict(credit_naive_model, newdata=test, type = "raw")</pre>
train <- cbind(predict_naive_train_prob, train)</pre>
test <- cbind(predict_naive_test_prob, test)</pre>
# class based on the loss matrix
train$predicted_class <- ifelse(train$good > 10*train$bad, "good", "bad")
test$predicted_class <- ifelse(test$good > 10*test$bad, "good", "bad")
# confusion matrix
conf_naive_train <- table(train$good_bad, train$predicted_class)</pre>
names(dimnames(conf naive train)) <- c("Actual Train", "Predicted Train")</pre>
caret::confusionMatrix(conf_naive_train)
## Confusion Matrix and Statistics
##
##
               Predicted Train
## Actual Train bad good
##
           bad 137
           good 263
                       90
##
##
##
                  Accuracy: 0.454
##
                    95% CI: (0.4097, 0.4988)
##
       No Information Rate: 0.8
       P-Value [Acc > NIR] : 1
##
##
##
                      Kappa: 0.1244
   Mcnemar's Test P-Value : <0.0000000000000002
##
##
##
               Sensitivity: 0.3425
##
               Specificity: 0.9000
##
            Pos Pred Value: 0.9320
##
            Neg Pred Value: 0.2550
##
                Prevalence: 0.8000
##
            Detection Rate: 0.2740
      Detection Prevalence: 0.2940
##
##
         Balanced Accuracy: 0.6213
##
##
          'Positive' Class : bad
conf_naive_test <- table(test$good_bad, test$predicted_class)</pre>
names(dimnames(conf_naive_test)) <- c("Actual Test", "Predicted Test")</pre>
caret::confusionMatrix(conf_naive_test)
## Confusion Matrix and Statistics
##
##
              Predicted Test
## Actual Test bad good
##
          bad
               71
                      5
##
          good 122
##
##
                  Accuracy: 0.492
```

```
##
                    95% CI: (0.4284, 0.5557)
##
       No Information Rate: 0.772
       P-Value [Acc > NIR] : 1
##
##
##
                     Kappa: 0.1626
   Mcnemar's Test P-Value : <0.0000000000000002
##
##
##
               Sensitivity: 0.3679
##
               Specificity: 0.9123
##
            Pos Pred Value: 0.9342
##
            Neg Pred Value: 0.2989
                Prevalence: 0.7720
##
##
            Detection Rate: 0.2840
##
      Detection Prevalence: 0.3040
##
         Balanced Accuracy: 0.6401
##
##
          'Positive' Class : bad
##
```

Analysis: We see a major drop in accuracy for both train and test cases for using the new loss matrix, the accuracy for train is 45.4% and for test cases its 49.2%, this was 70% and 68.4% respectively.

The biggest difference is in recognizing the bad customers, where the accuracy is 93.20% and 93.42% for the train and test cases respectively. This implies our model identifies 'bad' customers with a very high accuracy while it suffers to detect 'good' customers (accuracy of 25% and 29%). This is actually good for business because its many times more important to identify the bad customers instead of good customer (minimize risk).

The change in the accuracy is due to loss matrix where we have ensured that a customer is considered 'good' customer only if the probability of them being a 'good' customer is 10 times greater than them being a 'bad' customer.

# Assignment 3

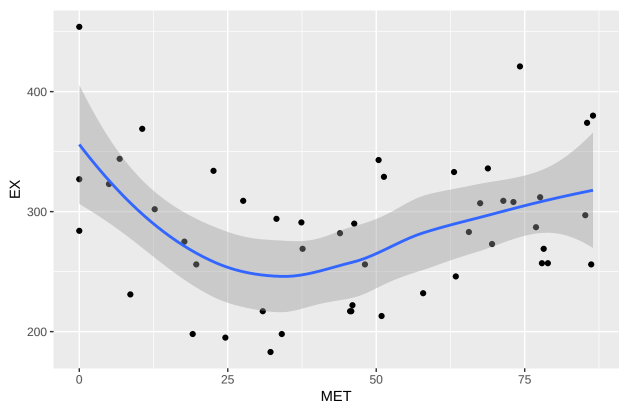
1. Reorder your data with respect to the increase of MET and plot EX versus MET. Discuss what kind of model can be appropriate here. Use the reordered data in steps 2-5.

```
rm(list=ls())
set.seed(12345)
state_data <- read.csv2("state.csv")

state_data <- state_data %>% arrange(MET)

ggplot(data = state_data, aes(x=MET, y = EX)) +
    geom_point() +
    geom_smooth(method = 'loess') +
    ggtitle("Plot of MET vs. EX")
```

#### Plot of MET vs. EX

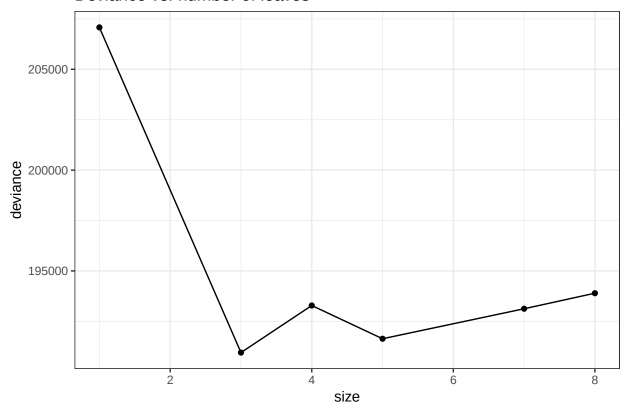


Analysis: As evident from the graph the best model, linear regression will not be a good fit, even the trend is non linear. Piece wise linear model(spline) might be a good one, thus regression per group/cluster will be a good approach.

2. Use package tree and fit a regression tree model with target EX and feature MET in which the number of the leaves is selected by cross-validation, use the entire data set and set minimum number of observations in a leaf equal to 8 (setting minsize in tree.control). Report the selected tree. Plot the original and the fitted data and histogram of residuals. Comment on the distribution of the residuals and the quality of the fit.

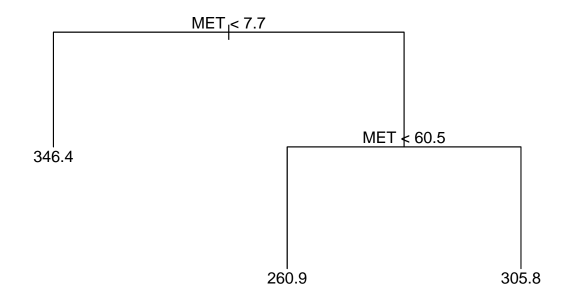
```
geom_line() +
    theme_bw() +
ggtitle("Deviance vs. number of leaves")
```

### Deviance vs. number of leaves



```
# The best size is either 3 or 4

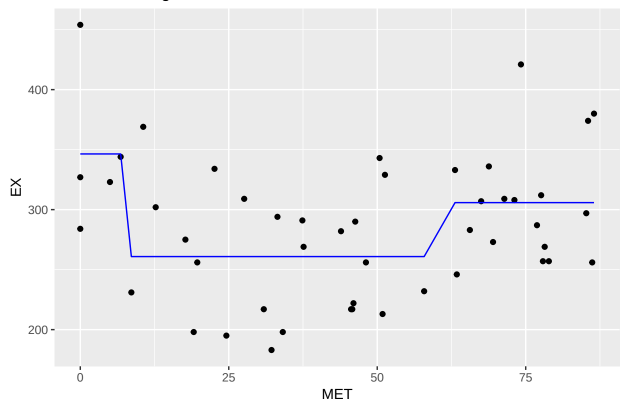
# puring the tree for leaf size of 3
state_cv_tree_purned <- prune.tree(state_tree_regression, best = 3)
plot(state_cv_tree_purned, main="Pruned Tree for the given dataset")
text(state_cv_tree_purned)</pre>
```



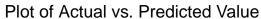
```
# Original vs. Fitted values
compare_data <- predict(state_cv_tree_purned, newdata = state_data)
compare_data <- cbind(compare_data, state_data$EX, state_data$MET)
compare_data <- as.data.frame(compare_data)
colnames(compare_data) <- c("predicted_value", "EX", "MET")
compare_data$residual <- compare_data$EX - compare_data$predicted_value

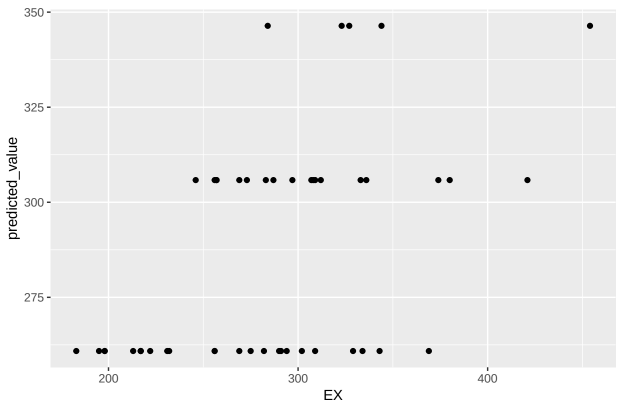
# plots
ggplot(data=compare_data, aes(x = MET, y = EX)) +
    geom_point(aes(x = MET, y=EX)) +
    geom_line(aes(x = MET, y=predicted_value), colour="blue") +
    ggtitle("EX value along with Predicted Value")</pre>
```

# EX value along with Predicted Value



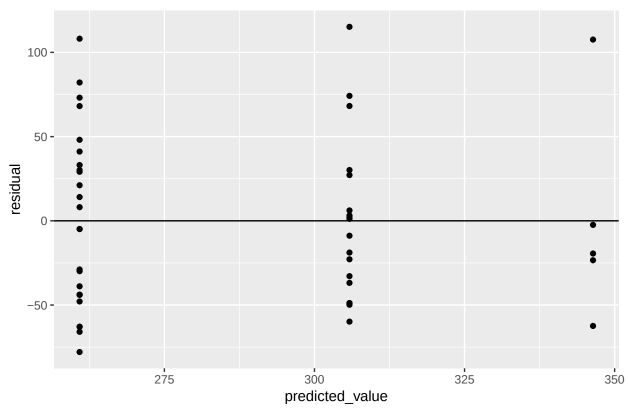
```
ggplot(compare_data, aes(x = EX, y = predicted_value)) +
geom_point() +
ggtitle("Plot of Actual vs. Predicted Value")
```



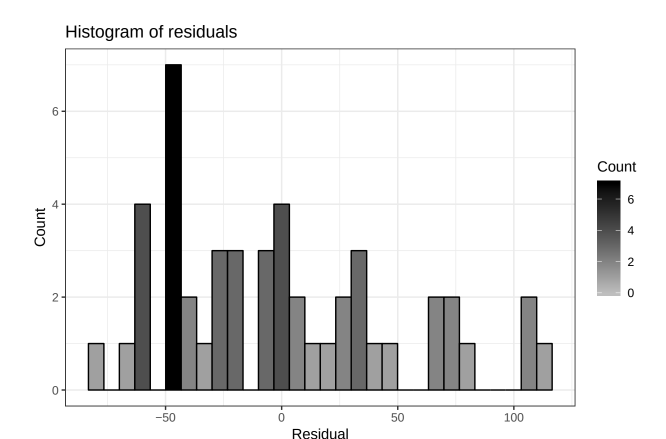


```
ggplot(compare_data, aes(x = predicted_value, y = residual)) +
geom_point() + geom_abline(slope=0, intercept=0) +
ggtitle("Plot of Predicted Value vs. Residual")
```

## Plot of Predicted Value vs. Residual



```
# Residuals
ggplot(data = compare_data, aes(residual)) +
geom_histogram(col="black",aes(fill=..count..),bins = 30) +
scale_fill_gradient("Count", low = "grey", high = "black") +
theme_bw() +
ggtitle("Histogram of residuals") +
labs(x="Residual", y="Count")
```



#### Analysis:

The predicted vs. Actual provides us the insight that for lower values of variable, our model is over predicting (actual value  $\sim 200$ ) while the predicted value is  $\sim 250$ . While for larger values ( $\sim 400$ ) our model is under predicting ( $\sim 330$ ). At around the mean value ( $\sim 300$ ) the predicted values are both under and over predicted thus no bias. Thus for values that are away from the mean our model is baised towards over/under predicted while values close to mean our model is not biased.

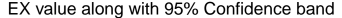
From the plot of Predicted vs. Residual values we can see that error appears random, neither is large bias/concentration of the error towards any value expect at lower/higher values(more points on one side of the line)

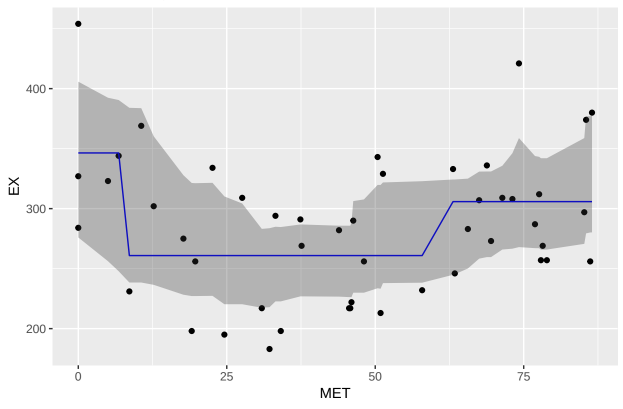
From the histogram we can see that the histogram has higher values on the left of zero, and a longer tail on the right. Thus from the above three points we can see that there is scope of improvement in the model especially in the extreme values of the predicted values.

3. Compute and plot the 95% confidence bands for the regression tree model from step 2 (fit a regression tree with the same settings and the same number of leaves as in step 2 to the resampled data) by using a non-parametric bootstrap. Comment whether the band is smooth or bumpy and try to explain why. Consider the width of the confidence band and comment whether results of the regression model in step 2 seem to be reliable.

set.seed(12345)

```
# computing bootstrap samples
bootstrap <- function(data, indices){</pre>
  data <- state_data[indices,]</pre>
 model <- tree(data = data,</pre>
       EX~MET,
       control = tree.control(nobs=NROW(data),
                               minsize = 8))
  model_purned <- prune.tree(model, best = 3)</pre>
 final_fit_boot <- predict(model_purned, newdata = state_data)</pre>
 return(final_fit_boot)
}
res <- boot(state_data, bootstrap, R=1000) #make bootstrap</pre>
## Warning in prune.tree(model, best = 3): best is bigger than tree size
e <- envelope(res, level = 0.95)
state_tree_regression <- tree(data = state_data, EX~MET,</pre>
                               control = tree.control(nobs=NROW(state_data),
                                                        minsize = 8))
# puring the tree for leaf size of 3
state_cv_tree_purned <- prune.tree(state_tree_regression, best = 3)</pre>
predict_for_ci <- predict(state_cv_tree_purned, state_data)</pre>
data_for_ci <- cbind(upper_bound = e$point[1,],</pre>
                      lower_bound = e$point[2,],
                      EX = state_data$EX,
                      MET = state data$MET,
                      predicted_value = predict_for_ci) %>% as.data.frame()
#plot cofidence bands
ggplot(data=data_for_ci, aes(x = MET, y = EX)) +
 geom_point(aes(x = MET,y=EX)) +
  geom_line(aes(x = MET, y=predicted_value), colour="blue") +
 geom_ribbon(aes(x = MET, ymin=lower_bound, ymax=upper_bound),alpha = 0.3) +
  ggtitle("EX value along with 95% Confidence band")
```





#### Analysis:

The confidence bands certainly appear to be bumpy and not smooth, the confidence bands are bumpy because the predicted values shows large flucations. From the width of the confidence band we can assume that our model is not a good one. Ideally we want the confidence band to be narrow thus a wider band suggests we need further tuning to the model.

4.Compute and plot the 95% confidence and prediction bands the regression tree model from step 2 (fit a regression tree with the same settings and the same number of leaves as in step 2 to the resampled data) by using a parametric bootstrap, assume Normal distribution with mean as labels in the tree leaves, while varience is residual varience. Consider the width of the confidence band and comment whether results of the regression model in step 2 seem to be reliable. Does it look like only 5% of data are outside the prediction band? Should it be?

```
set.seed(12345)
mle=prune.tree(state_tree_regression, best = 3)
#data2 = state_data

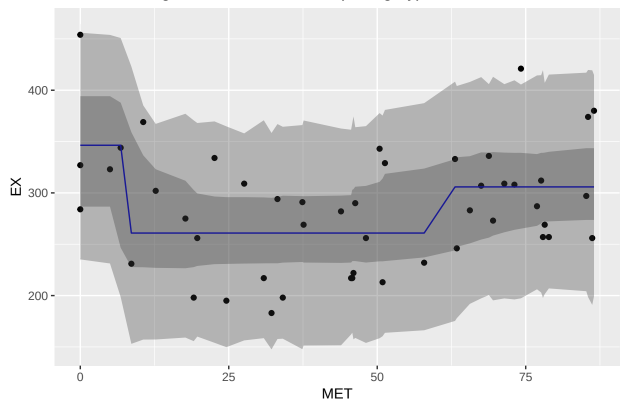
rng=function(data, mle) {
  data1=data.frame(EX=data$EX, MET=data$MET)
  n=length(data$EX)
```

```
pred <- predict(mle, newdata = data1)</pre>
  residual <- data1$EX - pred
#generate new Price
 data1$EX=rnorm(n, pred, sd(residual))
  return(data1)
# computing bootstrap samples
conf.fun <- function(data){</pre>
 model <- tree(data = data,</pre>
       EX~MET,
       control = tree.control(nobs=NROW(data),
                                minsize = 8))
  model_purned <- prune.tree(model, best = 3)</pre>
 final_fit_boot <- predict(model_purned, newdata = state_data)</pre>
  return(final_fit_boot)
}
# computing bootstrap samples
pred.fun <- function(data){</pre>
 model <- tree(data = data,</pre>
       EX~MET,
       control = tree.control(nobs=NROW(data),
                               minsize = 8))
 model_purned <- prune.tree(model, best = 3)</pre>
 final_fit_boot <- predict(model_purned, newdata = state_data)</pre>
 final_fit <- rnorm(n = length(final_fit_boot), mean = final_fit_boot, sd=sd(residuals(mle)))</pre>
 return(final_fit)
}
conf_para = boot(state_data, statistic=conf.fun, R=1000, mle=mle, ran.gen=rng, sim="parametric")
pred_para = boot(state_data, statistic=pred.fun, R=1000, mle=mle, ran.gen=rng, sim="parametric")
e1 <- envelope(conf para, level = 0.95)
e2 <- envelope(pred_para, level = 0.95)
## Warning in envelope(pred_para, level = 0.95): unable to achieve requested
## overall error rate
# puring the tree for leaf size of 3
state_cv_tree_purned <- prune.tree(state_tree_regression, best = 3)</pre>
predict_for_ci <- predict(state_cv_tree_purned, state_data)</pre>
data_for_ci_para <- cbind(upper_bound = e1$point[1,],</pre>
                      lower_bound = e1$point[2,],
                      upper_bound_pred = e2$point[1,],
                      lower_bound_pred = e2$point[2,],
                      EX = state_data$EX,
                      MET = state_data$MET,
```

```
predicted_value = predict_for_ci) %>% as.data.frame()

ggplot(data=data_for_ci_para, aes(x = MET, y = EX)) +
  geom_point(aes(x = MET, y=EX)) +
  geom_line(aes(x = MET, y=predicted_value), colour="blue") +
  geom_ribbon(aes(x = MET, ymin=lower_bound, ymax=upper_bound), alpha = 0.3) +
  geom_ribbon(aes(x = MET, ymin=lower_bound_pred, ymax=upper_bound_pred), alpha = 0.3) +
  ggtitle("EX value along with 95% Confidence(dark grey) and Prediction band")
```

### EX value along with 95% Confidence(dark grey) and Prediction band



#### Analysis:

From the plot we see that the width of the 'confidence band' (darker grey) is narrower compared to non-parametric. We also notice that most of the data is at in the 'confidence band', thus I believe that our regression model does make sense.

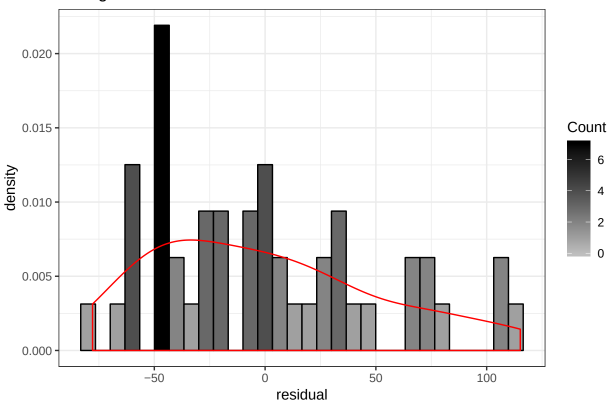
We can defintely see that most of the data is under the 'prediction band' more than 95%, thus we can reasonably expect that less than 5% of the data is outside of the 'prediction band'. Based on our assumption of data is normally distributed it does makes sense for prediction band to contain 95% of the data.

# 5. Consider the histogram of residuals from step 2 and suggest what kind of bootstrap is actually more appropriate here.

```
set.seed(12345)
# Residuals
```

```
ggplot(data = compare_data, aes(residual)) +
geom_histogram(col="black",aes(fill=..count.., y = ..density..),bins = 30) +
scale_fill_gradient("Count", low = "grey", high = "black") +
geom_density(colour = "red") +
theme_bw() +
ggtitle("Histogram of residuals")
```

#### Histogram of residuals



#### Analysis:

Judging from the density plot we can say that the density resembles that of a 'Beta Distribution' thus a known distribution is fit better using parametric bootstrap.

# Assignment 4

1.Conduct a standard PCA by using the feature space and provide a plot explaining how much variation is explained by each feature. Does the plot show how many PC should be extracted? Select the minimal number of components explaining at least 99% of the total variance. Provide also a plot of the scores in the coordinates (PC1, PC2). Are there unusual diesel fuels according to this plot?

```
rm(list=ls())
set.seed(12345)
```

Table 1: Contribution of PCA axis towards varience explaination

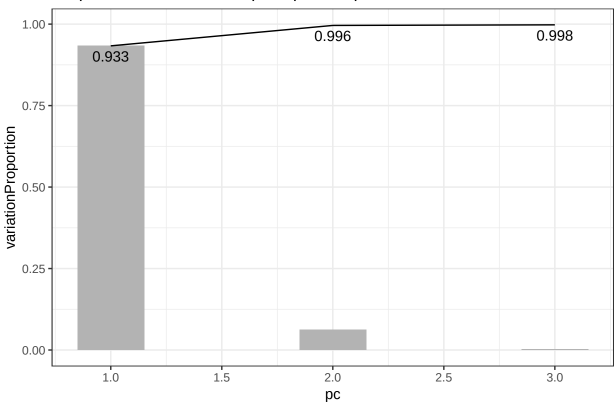
	PC1	PC2	PC3	PC4	PC5
Standard deviation	0.122062	0.0316205	0.0054353	0.0040107	0.0033031
Proportion of Variance	0.933320	0.0626300	0.0018500	0.0010100	0.0006800
Cumulative Proportion	0.933320	0.9959600	0.9978100	0.9988200	0.9995000

```
eigenvalues = pca_result$sdev^2

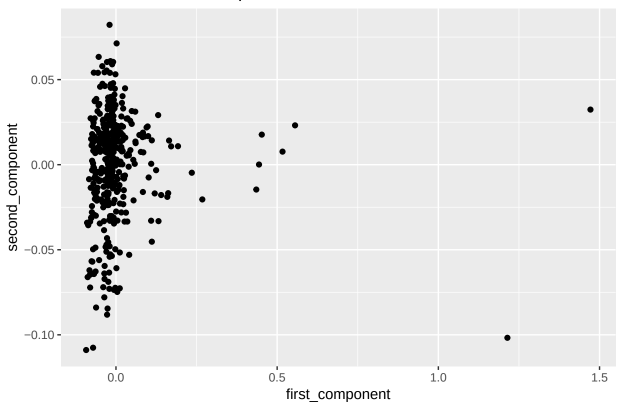
# plotting proportion of variation for principal components
plotData = as.data.frame(cbind(pc = 1:3,
    variationProportion = eigenvalues[1:3]/sum(eigenvalues),
    cummulative = cumsum(eigenvalues[1:3]/sum(eigenvalues))))

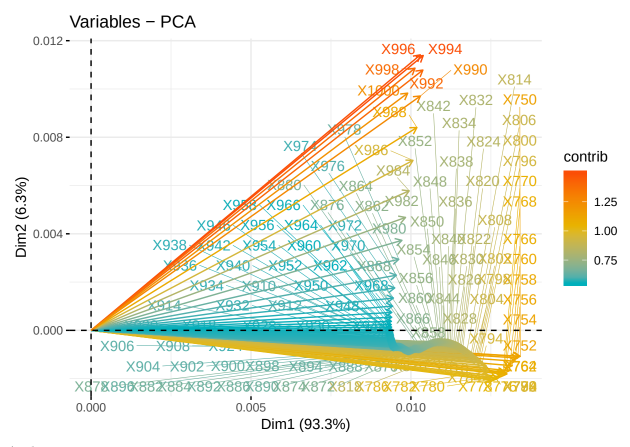
ggplot(data = plotData) +
geom_col(aes(x = pc, y = variationProportion), width = 0.3, fill = "grey70") +
geom_line(data = plotData,
    aes(x = pc, y = cummulative)) +
geom_text(aes(x = pc, y = cummulative, label = round(cummulative, 3)), size = 4,
position = "identity", vjust = 1.5) +
theme_bw() +
ggtitle("Proportion of variation for principal components")
```

## Proportion of variation for principal components



# Score Plot of PCA components





#### Analysis:

From the plot of PCA component vs. Viscoity, we can see that 2 components are needed (second component of PCA is needed), this is due to the fact the most of the data is vertically spread, removing this dimension would make it impossible to differentiate the types of diesel.

The minimum number of components that account for 99% of the total varience is 2, mainly PC1 and PC2.

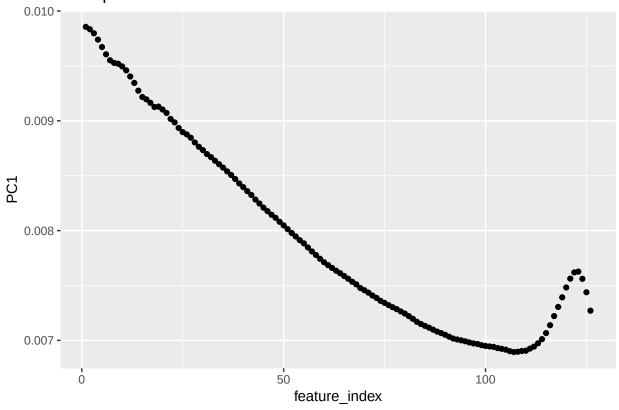
From the plot we can also see that there are evidently some diesel that are outliers (diesel with first\_component > 0.5 and second component~4).

# 2.Make trace plots of the loadings of the components selected in step 1. Is there any principle component that is explained by mainly a few original features?

```
# creating extra columns
aload <- abs(pca_result$rotation[,1:2])
components <- sweep(aload, 2, colSums(aload), "/")
components <- as.data.frame(components)
components$feature_name <- rownames(components)
components$feature_index <- 1:nrow(components)

ggplot(data = components, aes(x = feature_index, y = PC1)) +
    geom_point() +
    ggtitle("Traceplot of feature index vs. PC1")</pre>
```





```
ggplot(data = components, aes(x = feature_index, y = PC2)) +
geom_point() +
ggtitle("Traceplot of feature index vs. PC2")
```

### Traceplot of feature index vs. PC2

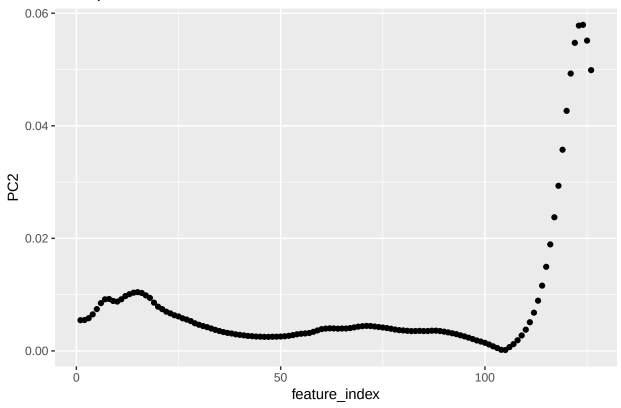


Table 2: Contribution of Features towards the Principle Components

	PC1	PC2	$feature\_name$	feature_index
X750	0.0098560	0.0054568	X750	1
X752	0.0098341	0.0054945	X752	2
X754	0.0097973	0.0058118	X754	3
X756	0.0097397	0.0064942	X756	4
X758	0.0096720	0.0074338	X758	5
X760	0.0096057	0.0084896	X760	6
X762	0.0095511	0.0091662	X762	7
X764	0.0095275	0.0092284	X764	8
X766	0.0095199	0.0088733	X766	9
X768	0.0094956	0.0087674	X768	10

#### Analysis:

While for PC1, the loadings of the variables do not differ too much (maximum loading  $\sim 0.11$ , minimum loading  $\sim 0.075$ ), for PC the loadings differ extremely. This principal component is explained mainly by the variables with a high index (on the right of the plot). Here, the maximum loading ( $\sim 0.35$ ) is much higher than for most of the other variables.

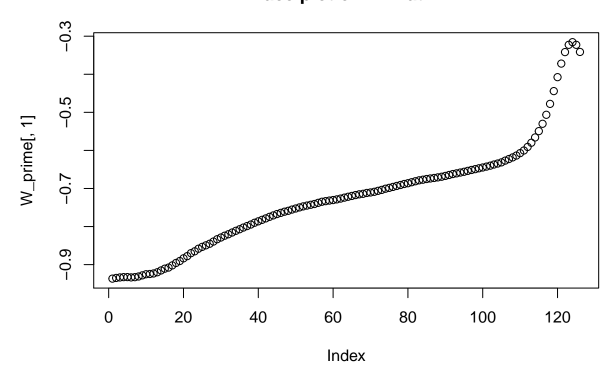
From these two plots is evident that there are no few features (<5) which form the core essence of the PCA components, thus the PCA components is made up of many (20+) features at least and they cannot be limited

to few original features.

3. Perform Independent Component Analysis with the number of components selected in step 1 (set seed 12345). Check the documentation for the fastICA method in R and do the following: a.Compute W(prime) = K.W and present the columns of W(prime) in the form of trace plots. Compare with the trace plots in step2 and make conclusion. What kind of measure is represented by the matrix W(prime.), b. Make a plot of the scores of the first two latent features and compare it with the score plot from step 1.

```
set.seed(12345)
\# X \rightarrow pre-processed data matrix
# K -> pre-whitening matrix that projects data onto the first n.compprincipal components.
# W -> estimated un-mixing matrix (see definition in details)
# A -> estimated mixing matrix
# S -> estimated source matrix
X <- as.matrix(pca_data)</pre>
ICA_extraction <- fastICA(X, 2, alg.typ = "parallel", fun = "logcosh", alpha = 1,</pre>
method = "R", row.norm = FALSE, maxit = 200,
tol = 0.0001, verbose = TRUE)
## Centering
## Whitening
## Symmetric FastICA using logcosh approx. to neg-entropy function
## Iteration 1 tol = 0.01930239
## Iteration 2 tol = 0.01303959
## Iteration 3 tol = 0.002393582
## Iteration 4 tol = 0.0006708454
## Iteration 5 tol = 0.0001661602
## Iteration 6 tol = 0.00003521604
W_prime <- ICA_extraction$K %*% ICA_extraction$W
#trace plots
plot(W_prime[,1], main = "Trace plot of W` Matrix")
```

# Trace plot of W` Matrix



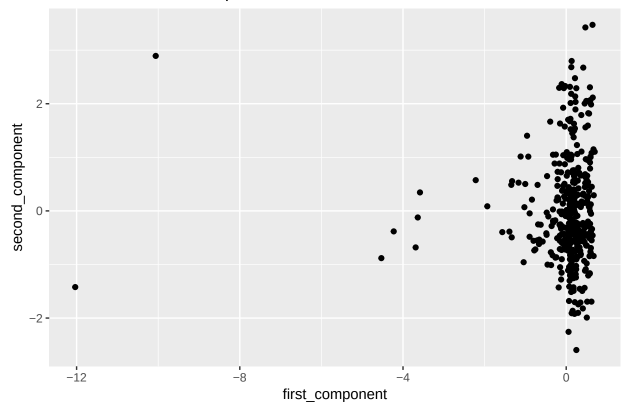
```
#trace plots
plot(W_prime[,2], main = "Trace plot of W` Matrix")
```

# Trace plot of W` Matrix



```
# pca components and the viscocity
ICA_result_data = cbind(first_component = ICA_extraction$S[,1],
                        second_component = ICA_extraction$S[,2],
                        Viscosity = NIR_data$Viscosity) %>% as.data.frame()
# plotting the data variation
ggplot(data = ICA_result_data, aes(x = first_component, y = second_component)) +
  geom_point() + ggtitle("Score Plot for ICA components")
```

#### Score Plot for ICA components



#### Analysis:

Comparing with the trace plots from step 2, we see that ICA trace plots are almost a (flipped using horizontal axis) flipped image of the PCA trace plots.

The W'(multipled matrix) represents the loadings of the ICA components, which are very similar to the PCA loadings.

Comparing the Score plots here we find that the plot is again a flipped version of the score plot from PCA (flipped along vertical axis).

# Appendix

```
set.seed(12345)
options("jtools-digits" = 2, scipen = 999)
# colours (colour blind friendly)
cbPalette <- c("#999999", "#E69F00", "#56B4E9", "#009E73", "#F0E442", "#0072B2",
               "#D55E00", "#CC79A7")
set.seed(12345)
data <- read.xlsx("creditscoring.xls", sheetName = "credit")</pre>
# changing class of dependent variable good_bad
data$good_bad = as.factor(data$good_bad)
# dividing data into train (50%), validation (25%) and test set (25%)
n = dim(data)[1]
set.seed(12345)
id = sample(1:n, floor(n*0.5))
train = data[id,]
id1 = setdiff(1:n, id)
set.seed(12345)
id2 = sample(id1, floor(n*0.25))
valid = data[id2,]
id3 = setdiff(id1,id2)
test = data[id3,]
set.seed(12345)
fitDecisionTree = function(testData, splitMethod) {
# fitting decision tree
decisionTreeFit = tree(formula = good_bad ~ ., data = train, split = splitMethod)
# getting prediction results
yFit = predict(object = decisionTreeFit, newdata = testData, type = "class")
# printing misclassification rate
paste0("Misclassification rate: ", mean(yFit != as.character(testData$good_bad)))
}
fitDecisionTree(testData = train, splitMethod = "deviance")
fitDecisionTree(testData = test, splitMethod = "deviance")
fitDecisionTree(testData = train, splitMethod = "gini")
fitDecisionTree(testData = test, splitMethod = "gini")
# fitting tree with measure providing better results ("deviance")
unprunedTree = tree(formula = good_bad ~ ., data = train, split = "deviance")
# Create a decision tree model
credit_tree_deviance <- tree(good_bad~., data=train, split = c("deviance"))</pre>
credit_tree_gini <- tree(good_bad~., data=train, split = c("gini"))</pre>
# predicting on the test dataset to get the misclassification rate.
predict_tree_deviance <- predict(credit_tree_deviance, newdata = test, type = "class")</pre>
predict_tree_gini <- predict(credit_tree_gini, newdata = test, type = "class")</pre>
```

```
conf_tree_deviance <- table(test$good_bad, predict_tree_deviance)</pre>
names(dimnames(conf_tree_deviance)) <- c("Actual Test", "P2redicted Test")</pre>
caret::confusionMatrix(conf_tree_deviance)
conf_tree_gini <- table(test$good_bad, predict_tree_gini)</pre>
names(dimnames(conf_tree_gini)) <- c("Actual Test", "Predicted Test")</pre>
caret::confusionMatrix(conf_tree_gini)
set.seed(12345)
credit_tree <- tree(good_bad~., data=train, split = c("deviance"))</pre>
credit_tree_purned_train <- prune.tree(credit_tree, method = c("deviance"))</pre>
credit_tree_purned_valid <- prune.tree(credit_tree, newdata = valid ,method = c("deviance"))</pre>
result_train <- cbind(credit_tree_purned_train$size,
                       credit_tree_purned_train$dev, "Train")
result_valid <- cbind(credit_tree_purned_valid$size,</pre>
                       credit_tree_purned_valid$dev, "Valid")
result <- as.data.frame(rbind(result_valid, result_train))</pre>
colnames(result) <- c("Leaf", "Deviance", "Type")</pre>
result$Leaf <- as.numeric(as.character(result$Leaf))</pre>
result$Deviance <- as.numeric(as.character(result$Deviance))</pre>
# plot of deviance vs. number of leafs
ggplot(data = result, aes(x = Leaf, y = Deviance, colour = Type)) +
  geom_point() + geom_line() +
  ggtitle("Plot of Deviance vs. Tree Depth (Shows Deviance least at 4)")
# prune the tree to the required depth
credit_tree_sniped <- prune.tree(credit_tree, best=4)</pre>
plot(credit_tree_sniped)
text(credit_tree_sniped)
# misclassification rate for best pruned tree
result_prune_test <- predict(credit_tree_sniped, newdata = test, type = "class")
conf_prune_tree_test <- table(test$good_bad, result_prune_test)</pre>
names(dimnames(conf_prune_tree_test)) <- c("Actual Test", "Predicted Test")</pre>
caret::confusionMatrix(conf prune tree test)
#Fitting the Naive Bayes model
credit_naive_model = naiveBayes(good_bad ~., data=train)
#Prediction on the dataset
predict_naive_train = predict(credit_naive_model, newdata=train, type = "class")
predict_naive_test = predict(credit_naive_model, newdata=test, type = "class")
conf_naive_train <- table(train$good_bad, predict_naive_train)</pre>
names(dimnames(conf_naive_train)) <- c("Actual Train", "Predicted Train")</pre>
caret::confusionMatrix(conf_naive_train)
```

```
conf_naive_test <- table(test$good_bad, predict_naive_test)</pre>
names(dimnames(conf_naive_test)) <- c("Actual Test", "Predicted Test")</pre>
caret::confusionMatrix(conf naive test)
set.seed(12345)
credit_tree <- tree(good_bad~., data=train, split = c("deviance"))</pre>
credit_naive_model = naiveBayes(good_bad ~., data=train)
# prune the tree to the required depth
credit_tree_sniped <- prune.tree(credit_tree, best=7)</pre>
# predicting class, getting probability
predict_prune_test_prob <- predict(credit_tree_sniped, newdata = test)</pre>
predict_naive_test_prob <- predict(credit_naive_model, newdata=test, type = "raw")</pre>
# data mugging
probability_data_naive <- as.data.frame(cbind(predict_naive_test_prob,</pre>
                                                as.character(test$good_bad), "naivebayes"))
probability_data_tree <- as.data.frame(cbind(predict_prune_test_prob,</pre>
                                               as.character(test$good_bad), "tree"))
probability_data_combined <- rbind(probability_data_tree, probability_data_naive)</pre>
colnames(probability_data_combined) <- c("prob_bad", "prob_good",</pre>
                                           "actual_test_class", "model")
# final dataset
probability_data_combined$prob_good <- as.numeric(as.character(probability_data_combined$prob_good))</pre>
# changing the threshold and printing the probability
tree_list <- NULL</pre>
naive_list <- NULL</pre>
final <- NULL
for(threshold in seq(from = 0.05, to = 0.95, by = 0.05)){
  probability_data_combined$predicted_class <- ifelse(probability_data_combined$prob_good > threshold,
 df2 <- probability_data_combined[,c("model", "actual_test_class", "predicted_class")]</pre>
  df2$threshold <- threshold
 df2\statch <- ifelse(df2\statch actual_test_class == df2\statch predicted_class, 1, 0)
 final <- rbind(df2, final)</pre>
}
# Creating the FRP and TRP for each model and threshold
final$temp <- 1
final_summary <- final %>%
group_by(model, threshold) %>%
summarise(total_positive = sum(temp[actual_test_class == "good"]),
          total_negative = sum(temp[actual_test_class == "bad"]),
          correct_positive = sum(temp[actual_test_class == "good" & predicted_class == "good"]),
          false_positive = sum(temp[actual_test_class == "bad" & predicted_class == "good"])) %>%
    mutate(TPR = correct_positive/total_positive, FPR = false_positive/total_negative) %>%
  select(model, threshold, TPR, FPR)
```

```
ggplot(data = final_summary, aes(x = FPR, y=TPR)) + geom_line(aes(colour = model)) +
  geom_abline(intercept = 0, slope = 1) +
  ggtitle("ROC curve for the Naive Bayes vs. Tree Model")
set.seed(12345)
credit naive model = naiveBayes(good bad ~., data=train)
# predicting class, getting probability
predict_naive_train_prob <- predict(credit_naive_model, newdata=train, type = "raw")</pre>
predict_naive_test_prob <- predict(credit_naive_model, newdata=test, type = "raw")</pre>
train <- cbind(predict_naive_train_prob, train)</pre>
test <- cbind(predict_naive_test_prob, test)</pre>
# class based on the loss matrix
train$predicted_class <- ifelse(train$good > 10*train$bad, "good", "bad")
test$predicted_class <- ifelse(test$good > 10*test$bad, "good", "bad")
# confusion matrix
conf_naive_train <- table(train$good_bad, train$predicted_class)</pre>
names(dimnames(conf naive train)) <- c("Actual Train", "Predicted Train")</pre>
caret::confusionMatrix(conf_naive_train)
conf_naive_test <- table(test$good_bad, test$predicted_class)</pre>
names(dimnames(conf_naive_test)) <- c("Actual Test", "Predicted Test")</pre>
caret::confusionMatrix(conf_naive_test)
rm(list=ls())
set.seed(12345)
state_data <- read.csv2("state.csv")</pre>
state_data <- state_data %>% arrange(MET)
ggplot(data = state_data, aes(x=MET, y = EX)) +
 geom point() +
 geom smooth(method = 'loess') +
 ggtitle("Plot of MET vs. EX")
set.seed(12345)
state_tree_regression <- tree(data = state_data, EX~MET,</pre>
                               control = tree.control(nobs=NROW(state_data),
                                                       minsize = 8))
state_cv_tree <- cv.tree(state_tree_regression, FUN = prune.tree)</pre>
temp <- cbind(size = state_cv_tree$size, deviance = state_cv_tree$dev) %>% as.data.frame()
ggplot(data = temp, aes(x = size, y = deviance)) +
         geom_point() +
         geom_line() +
```

```
theme_bw() +
ggtitle("Deviance vs. number of leaves")
# The best size is either 3 or 4
# puring the tree for leaf size of 3
state_cv_tree_purned <- prune.tree(state_tree_regression, best = 3)</pre>
plot(state cv tree purned, main="Pruned Tree for the given dataset")
text(state_cv_tree_purned)
# Original vs. Fitted values
compare_data <- predict(state_cv_tree_purned, newdata = state_data)</pre>
compare data <- cbind(compare data, state data$EX, state data$MET)</pre>
compare_data <- as.data.frame(compare_data)</pre>
colnames(compare_data) <- c("predicted_value", "EX", "MET")</pre>
compare_data$residual <- compare_data$EX - compare_data$predicted_value
# plots
ggplot(data=compare_data, aes(x = MET, y = EX)) +
  geom_point(aes(x = MET,y=EX)) +
  geom_line(aes(x = MET, y=predicted_value), colour="blue") +
  ggtitle("EX value along with Predicted Value")
ggplot(compare_data, aes(x = EX, y = predicted_value)) +
  geom point() +
  ggtitle("Plot of Actual vs. Predicted Value")
ggplot(compare_data, aes(x = predicted_value, y = residual)) +
  geom_point() + geom_abline(slope=0, intercept=0) +
  ggtitle("Plot of Predicted Value vs. Residual")
# Residuals
ggplot(data = compare_data, aes(residual)) +
geom_histogram(col="black",aes(fill=..count..),bins = 30) +
scale_fill_gradient("Count", low = "grey", high = "black") +
theme_bw() +
ggtitle("Histogram of residuals") +
labs(x="Residual", y="Count")
set.seed(12345)
# computing bootstrap samples
bootstrap <- function(data, indices){</pre>
  data <- state_data[indices,]</pre>
 model <- tree(data = data,</pre>
       EX~MET,
       control = tree.control(nobs=NROW(data),
                               minsize = 8))
  model_purned <- prune.tree(model, best = 3)</pre>
  final_fit_boot <- predict(model_purned, newdata = state_data)</pre>
```

```
return(final_fit_boot)
}
res <- boot(state_data, bootstrap, R=1000) #make bootstrap
e <- envelope(res, level = 0.95)
state_tree_regression <- tree(data = state_data, EX~MET,</pre>
                               control = tree.control(nobs=NROW(state_data),
                                                       minsize = 8))
# puring the tree for leaf size of 3
state_cv_tree_purned <- prune.tree(state_tree_regression, best = 3)</pre>
predict_for_ci <- predict(state_cv_tree_purned, state_data)</pre>
data for ci <- cbind(upper bound = e$point[1,],
                      lower_bound = e$point[2,],
                      EX = state_data$EX,
                      MET = state_data$MET,
                      predicted_value = predict_for_ci) %>% as.data.frame()
#plot cofidence bands
ggplot(data=data_for_ci, aes(x = MET, y = EX)) +
  geom_point(aes(x = MET,y=EX)) +
  geom_line(aes(x = MET, y=predicted_value), colour="blue") +
  geom_ribbon(aes(x = MET, ymin=lower_bound, ymax=upper_bound),alpha = 0.3) +
  ggtitle("EX value along with 95% Confidence band")
set.seed(12345)
mle=prune.tree(state_tree_regression, best = 3)
#data2 = state_data
rng=function(data, mle) {
  data1=data.frame(EX=data$EX, MET=data$MET)
  n=length(data$EX)
  pred <- predict(mle, newdata = data1)</pre>
  residual <- data1$EX - pred
#generate new Price
  data1$EX=rnorm(n, pred, sd(residual))
  return(data1)
}
# computing bootstrap samples
conf.fun <- function(data){</pre>
  model <- tree(data = data,</pre>
       EX~MET.
       control = tree.control(nobs=NROW(data),
                               minsize = 8))
  model_purned <- prune.tree(model, best = 3)</pre>
  final_fit_boot <- predict(model_purned, newdata = state_data)</pre>
  return(final_fit_boot)
```

```
# computing bootstrap samples
pred.fun <- function(data){</pre>
  model <- tree(data = data,
       EX~MET,
       control = tree.control(nobs=NROW(data),
                              minsize = 8))
 model_purned <- prune.tree(model, best = 3)</pre>
 final fit boot <- predict(model purned, newdata = state data)</pre>
 final_fit <- rnorm(n = length(final_fit_boot), mean = final_fit_boot, sd=sd(residuals(mle)))</pre>
 return(final fit)
}
conf_para = boot(state_data, statistic=conf.fun, R=1000, mle=mle, ran.gen=rng, sim="parametric")
pred_para = boot(state_data, statistic=pred.fun, R=1000, mle=mle, ran.gen=rng, sim="parametric")
e1 <- envelope(conf_para, level = 0.95)
e2 <- envelope(pred_para, level = 0.95)
# puring the tree for leaf size of 3
state_cv_tree_purned <- prune.tree(state_tree_regression, best = 3)</pre>
predict_for_ci <- predict(state_cv_tree_purned, state_data)</pre>
data_for_ci_para <- cbind(upper_bound = e1$point[1,],</pre>
                     lower_bound = e1$point[2,],
                     upper_bound_pred = e2$point[1,],
                     lower_bound_pred = e2$point[2,],
                     EX = state_data$EX,
                     MET = state_data$MET,
                     predicted_value = predict_for_ci) %>% as.data.frame()
ggplot(data=data_for_ci_para, aes(x = MET, y = EX)) +
  geom_point(aes(x = MET,y=EX)) +
  geom_line(aes(x = MET, y=predicted_value), colour="blue") +
  geom_ribbon(aes(x = MET, ymin=lower_bound, ymax=upper_bound),alpha = 0.3) +
  geom ribbon(aes(x = MET, ymin=lower bound pred, ymax=upper bound pred), alpha = 0.3) +
  ggtitle("EX value along with 95% Confidence(dark grey) and Prediction band")
set.seed(12345)
# Residuals
ggplot(data = compare_data, aes(residual)) +
geom_histogram(col="black",aes(fill=..count.., y = ..density..),bins = 30) +
scale_fill_gradient("Count", low = "grey", high = "black") +
geom_density(colour = "red") +
theme_bw() +
ggtitle("Histogram of residuals")
rm(list=ls())
set.seed(12345)
```

```
NIR_data <- read.csv2("NIRSpectra.csv")</pre>
pca_data = select(NIR_data,-c(Viscosity))
pca_result = prcomp(pca_data)
contribution <- summary(pca_result)$importance</pre>
knitr::kable(contribution[,1:5],
             caption = "Contribution of PCA axis towards varience explaination")
eigenvalues = pca_result$sdev^2
# plotting proportion of variation for principal components
plotData = as.data.frame(cbind(pc = 1:3,
variationProportion = eigenvalues[1:3]/sum(eigenvalues),
cummulative = cumsum(eigenvalues[1:3]/sum(eigenvalues))))
ggplot(data = plotData) +
geom_col(aes(x = pc, y = variationProportion), width = 0.3, fill = "grey70") +
geom_line(data = plotData,
aes(x = pc, y = cummulative)) +
geom_text(aes(x = pc, y = cummulative, label = round(cummulative, 3)), size = 4,
position = "identity", vjust = 1.5) +
theme bw() +
ggtitle("Proportion of variation for principal components")
# pca components and the viscocity
pca_result_data = cbind(first_component = pca_result$x[,1],
                                 second_component = pca_result$x[,2]) %>% as.data.frame()
# plotting the data variation and the viscocity
ggplot(data = pca_result_data, aes(x = first_component, y = second_component)) +
  geom_point() + ggtitle("Score Plot of PCA components")
# showing the score of PCA component
factoextra::fviz_pca_var(pca_result,
             col.var = "contrib", # Color by contributions to the PC
             gradient.cols = c("#00AFBB", "#E7B800", "#FC4E07"),
             repel = TRUE  # Avoid text overlapping
set.seed(12345)
# creating extra columns
aload <- abs(pca_result$rotation[,1:2])</pre>
components <- sweep(aload, 2, colSums(aload), "/")</pre>
components <- as.data.frame(components)</pre>
components$feature_name <- rownames(components)</pre>
components$feature_index <- 1:nrow(components)</pre>
ggplot(data = components, aes(x = feature_index, y = PC1)) +
  geom_point() +
  ggtitle("Traceplot of feature index vs. PC1")
```

```
ggplot(data = components, aes(x = feature_index, y = PC2)) +
  geom_point() +
  ggtitle("Traceplot of feature index vs. PC2")
knitr::kable(components[1:10,],
             caption = "Contribution of Features towards the Principle Components")
set.seed(12345)
# X -> pre-processed data matrix
# K -> pre-whitening matrix that projects data onto the first n.compprincipal components.
\#\ \mathbb{W}\ 	ext{->}\ estimated\ un-mixing\ matrix} (see definition in details)
# A -> estimated mixing matrix
\# S \rightarrow estimated source matrix
X <- as.matrix(pca_data)</pre>
ICA_extraction <- fastICA(X, 2, alg.typ = "parallel", fun = "logcosh", alpha = 1,</pre>
method = "R", row.norm = FALSE, maxit = 200,
tol = 0.0001, verbose = TRUE)
W_prime <- ICA_extraction$K %*% ICA_extraction$W
#trace plots
plot(W_prime[,1], main = "Trace plot of W` Matrix")
#trace plots
plot(W_prime[,2], main = "Trace plot of W` Matrix")
# pca components and the viscocity
ICA_result_data = cbind(first_component = ICA_extraction$S[,1],
                         second_component = ICA_extraction$S[,2],
                         Viscosity = NIR_data$Viscosity) %>% as.data.frame()
# plotting the data variation
ggplot(data = ICA_result_data, aes(x = first_component, y = second_component)) +
 geom_point() + ggtitle("Score Plot for ICA components")
```