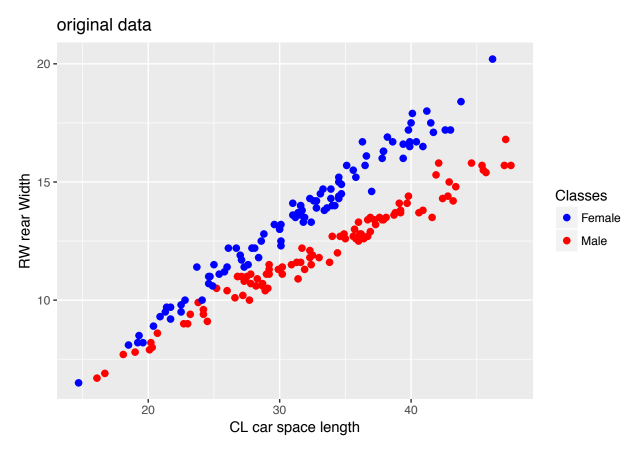
# Group8\_lab2

Group 8

November 29, 2017

## Assignment 1 LDA and Logistic Regression

Part 1



Yes , it is easy to classify the data by linear discriminant function because from the plot it can be observed that the two classes have cooperative distribution having equal covariance.

### Part 2

Discriminant function coefficients for male

**##** [1] 2.5658514 -0.2138144 -12.5634175

Discriminant function coefficients for Female

**##** [1] 8.248698 -2.161318 -22.428769

### **Decision boundary Equation**

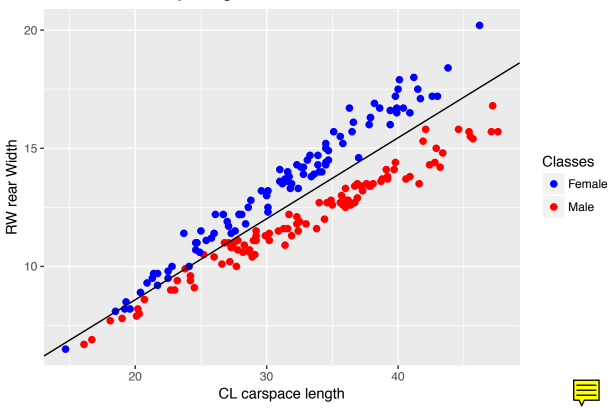
$$RW*W_{1R} + CL*W_{1C} + W_{01} = RW*W_{2R} + CL*W_{2C} + W_{02}$$

#### Final equation in term or RW as a Function of CL

$$\begin{split} RW &= \frac{(CL(W_{2C} - W_{1C}) + W_{02} - W_{02})}{W_{1R} - W_{2R}}\\ intercept &= \frac{CL(W_{2C} - W_{1C})}{W_{1R} - W_{2R}}\\ slope &= \frac{W_{02} - W_{02}}{W_{1R} - W_{2R}} \end{split}$$

#### Part 3

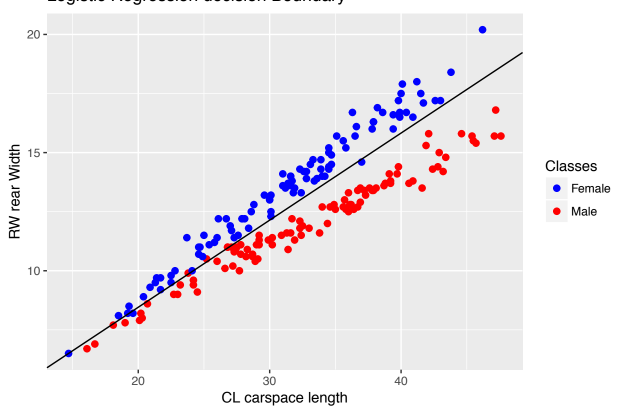
### Descion Boundary using LDA



It is evident from the plot that decision boundary is separating the data between the two classes classifying points belonging to each class on the respective side of the boundary. There are some points which are misclassified but since there is not a lot of misclassified data that is misclassification rate is low therefore it can be assumed that the quality of fit is good.

Part 4

Logistic Regression decision Boundary



#### Decision boundary equation for Logistic Regression

$$RW(W_{RW}) + CL(W_{CL}) + W_{01} = 0.5$$

### In terms of RW

$$\begin{split} RW &= \frac{-CL*W_{CL}}{W_{RW}} - \frac{W_{01}}{W_{RW}} + \frac{0.5}{W_{RW}}\\ intercept &= \frac{-CL(W_{CL})}{W_{RW}} \ slope = -\frac{W_{01}}{W_{RW}} \end{split}$$

#### Comaprision

Apparently there are more misclassified points for female in LDA while on applying logistic regression, the female points are classified better than that of male. One more thing which can be observed from the plot is that for LDA, there are very few points which lie on the decision boundary while for Logistic Regression, there are comparatively more points which lie on the decision boundary that its which are neither classified as male nor as female.

### Assignment 2. Analysis of Credit Scoring

#### Part 2

Misclassification for Training Data

#### Missclassification Rate of Deviance

```
##
## devi_yfit bad good
                   20
##
       bad
             61
       good 86 333
## [1] 0.212
Missclassification rate Gini
##
## gini_yfit bad good
              68
##
       bad
                   36
       good 79 317
##
```

#### Misclssification for Test Data

#### Missclassification Rate of Deviance

```
##
## devi_yfit bad good
## bad 24 17
## good 54 155
## [1] 0.284
```

## [1] 0.23

#### Missclassification Rate of Gini

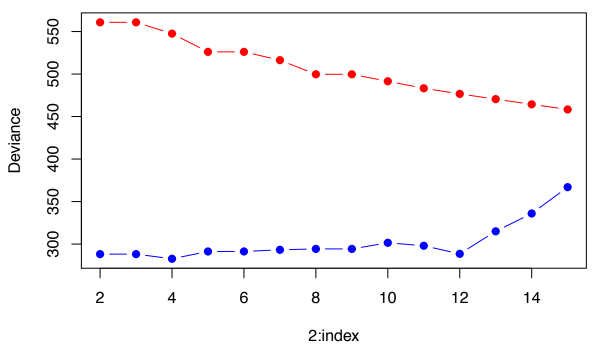
```
## ## gini_yfit bad good
## bad 23 30
## good 55 142
## [1] 0.34
```



Since the misclassification rate for both training and test data for Gini is less therefore it is a better measure.

### Part 3

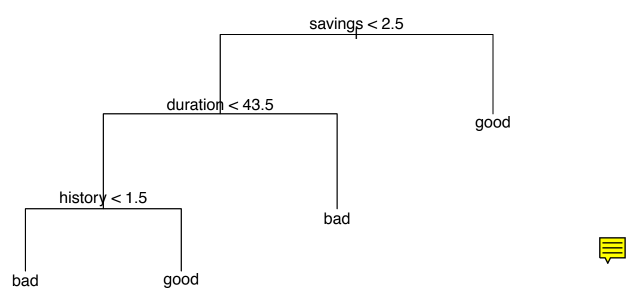
## **Dependence of Deviance**



```
##
## Classification tree:
## snip.tree(tree = tree_deviance, nodes = c(5L, 3L, 9L))
## Variables actually used in tree construction:
## [1] "savings" "duration" "history"
## Number of terminal nodes: 4
## Residual mean deviance: 1.117 = 547.5 / 490
## Misclassification error rate: 0.251 = 124 / 494
```

Optimal depth of tree is 4 and the variables selected by the tree are savings, duration and history

```
## [1] "Confusion Matrix"
         yfit
##
##
          bad good
##
     bad
           19
                59
     good
            6 166
## [1] "Misclassification rate"
## [1] 0.26
```



According to the selected tree, if savings is greater than 2.5 then its good else bad. Moreover, if the duration of savings is less than 43.5 months then its good else bad and if history of savings is maintained for less more than 1.5 then its good.

#### Part 4

```
##
##
  nav_test bad good
##
       bad
             47
                   49
##
       good
             31
                 123
##
##
  nav_train bad good
              95
                    98
##
        bad
##
              52
                  255
        good
## Misclassification train data value Using Naive Bayes is: 0.3
## Misclassification test data value Using Naive Bayes is: 0.32
```

The misclassification rate calculated by Naive Bayes in step 4 for test data is 0.32 which is greater than that of misclassification calculated by the decision tree in step 3 that is 0.26

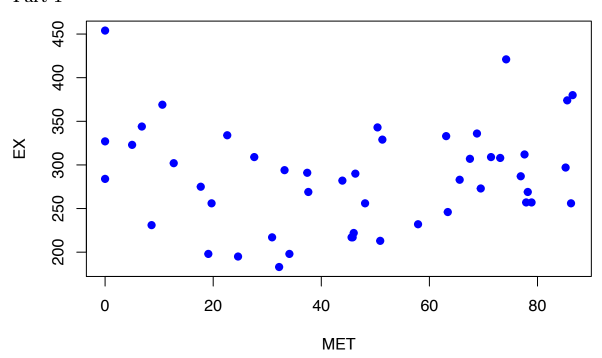
```
##
          nav_train
##
           FALSE TRUE
##
     bad
             137
                     10
             263
                    90
##
     good
   [1] 0.546
##
##
          nav_test
           FALSE TRUE
##
##
     bad
              70
                      8
             131
                     41
##
     good
## [1] 0.556
```

The misclassification rate calculated using the given loss function is 0.546 for training data and 0.556 for test data which is much greater than the one calculated by Naive Bayes without using loss function that is

0.3 for training data and 0.32 for test data. The increase in misclassification is due to the introduction of loss function giving a threshold for the true and false values.

## Assignment 3 Uncertainty estimation

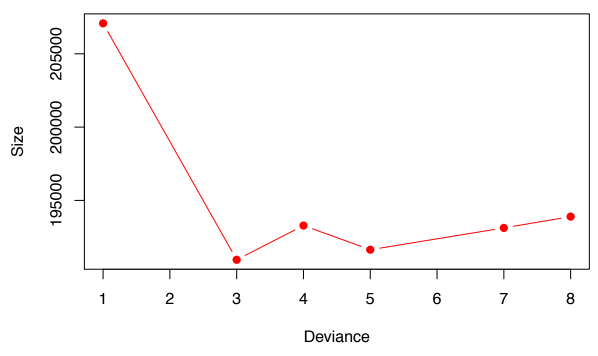
Part 1



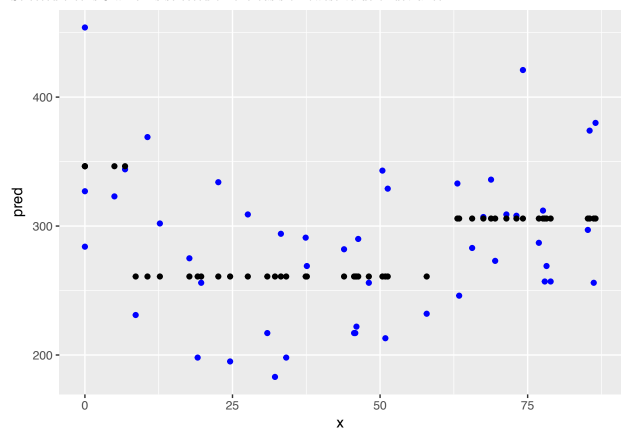
From the plot it can be observed that the data is scattered that is variance is high thus for the this type of data, decision trees would be the appropriate method

Part 2

## **Deviance Vs Size of Tree**

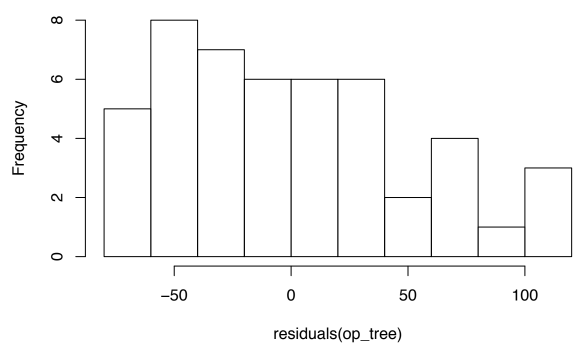


Selected tree is 3 which is selected on the basis of lowest value of deviance





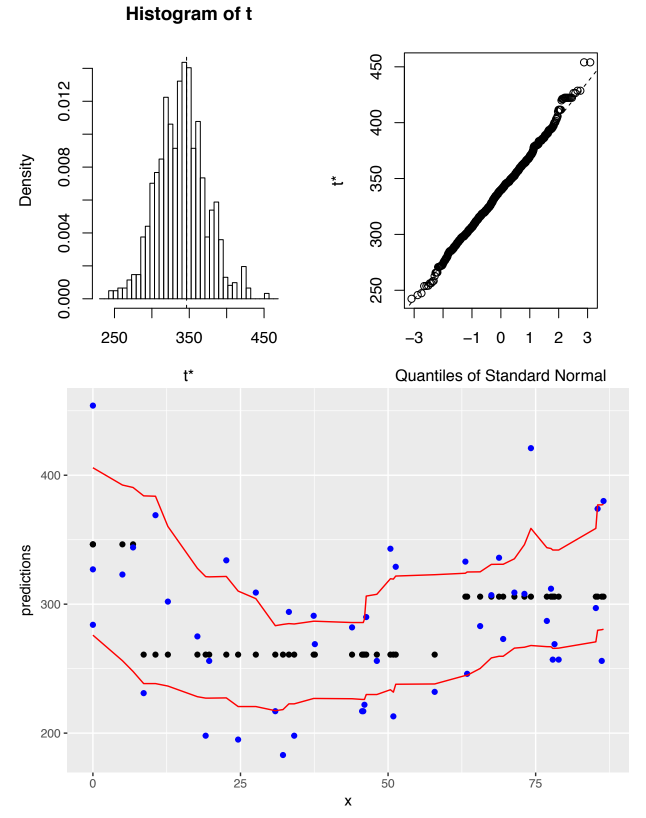
## Histogram of residuals(op\_tree)



Residuals seems to be marginally distributed across the data. According to the histogram, residuals represents that fitting of data can be improved that is the residuals can be reduced if better fitting is applied.



Part 3

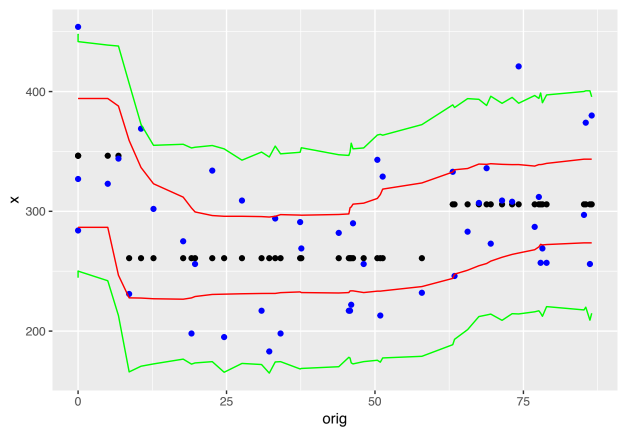


Confidence band is the bootstrap percentile interval which is combination of different confidence intervals

computed for different replicates of bootstrap.

The confidence band for the parametric bootstrap is bumpy due to the impact of bias on bootstrap.

Considering the width of confidence interval, the result of regression model computed in part 2 appears to be reliable as it appears to lie within the confidence band.



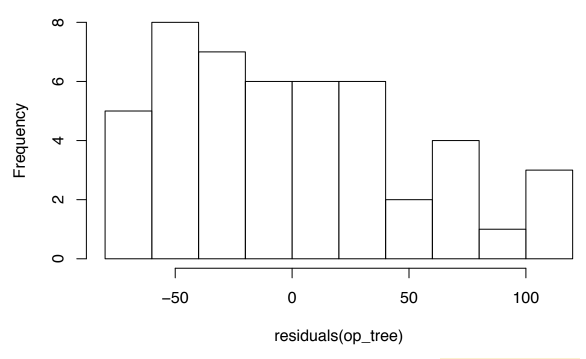
The red line in the above plot represents the confidence band. The confidence band seems to enclose the fitted data from step 2 and thus the regression model in step 2 appears to be reliable.

The green lines represent the prediction band. It can be observed that the prediction band encloses almost all the points and only 5% or less data resides outside the prediction band. If the prediction band would enclose all points, it will indicate that there might be overfitting.



Part 5

### Histogram of residuals(op\_tree)



On the basis of histogram plot from step 2, it can be suggested that the **parametric bootstrap** would be more appropriate model since for the given amount of data, non-parametric bootstrap might gives underfitted values. Moreover, the parametric bootstrap is more reliable if distribution is pre-determined regardless of the size of data.

### **APPENDIX**



```
## Assignment 1

library(ggplot2)
data<- read.csv2("australian-crabs.csv" ,sep = ",",dec=".")
p <- ggplot(data, aes(x=CL, y=RW)) + geom_point(aes(color=sex), size=2) +
    scale_color_manual (values = c('blue', 'red')) +
    labs(x="CL carspace length", y="RW rear Width", colour="Classes") +
    ggtitle("original data")

X<- data.frame(RW=data$RW , CL=data$CL )
Y <- data$sex

#1.2
library(MASS)
disc_fun=function(label, S)
{
    X1=X[Y==label,]
    mean_v <- c(mean(X1$RW) ,mean(X1$CL))
    covaiance_mat_inverse <- solve(S)</pre>
```

```
prior_prob <- nrow(X1) / nrow(X)</pre>
  w1 <- covaiance_mat_inverse %*% mean v
  b1 <- ((-1/2) %*% t(mean_v) %*% covaiance_mat_inverse %*% mean_v) + log(prior_prob)
  w1<- as.vector(w1)
 return(c(w1[1], w1[2], b1[1,1]))
X1=X[Y=="Male",]
X2=X[Y=="Female",]
S=cov(X1)*dim(X1)[1]+cov(X2)*dim(X2)[1]
S=S/dim(X)[1]
#discriminant function coefficients
res1=disc_fun("Male",S)
res2=disc_fun("Female",S)
#1.2
#decision boundary coefficients 'res'
res <- c( -(res1[1]-res2[1]) , (res2[2]-res1[2]), (res2[3]-res1[3]))
# classification
d=res[1]*X[,1]+res[2]*X[,2]+res[3]
Yfit=(d>0)
plot(X[,1], X[,2], col=Yfit+1, xlab="CL", ylab="RW")
#slope and intercept
slope <- (res[2] / res[1]) * -1
intercept \leftarrow res[3] /res[1] * -1
#1.3
#plot decision boundary
X<- cbind(X,sex=Y)</pre>
p <- ggplot(X, aes(x=CL, y=RW)) + geom_point(aes(color=sex), size=2) +</pre>
  scale_color_manual (values = c('blue', 'red')) +
  labs(x="CL carspace length", y="RW rear Width", colour="Classes") +
  geom_abline(slope = slope, intercept = intercept) +
  ggtitle("Descion Boundary LDA")
#1.4
logistic_fit <- glm(sex ~ RW + CL, data= X,</pre>
                    family='binomial') #glm fitting
r.unlist <- as.numeric(unlist(logistic_fit$coefficients))</pre>
r.unlist <- data.frame(intercept=r[1] ,</pre>
                        RW=r[2],
                        CL=r[3]) #convert data into datafreame
glm_intercept <- (-r$intercept/r$RW) #+ (0.5/r$RW)</pre>
glm_slope <- (-r^{CL} / r^{RW} ) #+ (0.5) / ^{RW}
p <- ggplot(X, aes(x=CL, y=RW)) + geom_point(aes(color=sex), size=2 ) +</pre>
  scale_color_manual (values = c('blue', 'red')) +
  labs(x="CL carspace length", y="RW rear Width", colour="Classes") +
  geom_abline(slope = glm_slope,
```

```
intercept = glm_intercept) +
    ggtitle("Logistic Regression
            decision Boundary")
  ## Assignment 2
library(readxl)
library(tree)
library(e1071)
# Importing Data - 2.1
data <- read_excel("creditscoring.xls")</pre>
data <- as.data.frame(data)</pre>
data$good_bad <- as.factor(data$good_bad)</pre>
# Dividing Data into three Train(50%) Test(25%) Validation(25%)
n = nrow(data)
set.seed(12345)
n=dim(data)[1]
# 50% Training Data
id=sample(1:n, floor(n*0.5))
train=data[id,]
# 25% validation & testing Data
Sub id = data[-id,]
m = dim(Sub_id)[1]
part1 = sample(1:m, floor(m*0.5))
validation = Sub_id[part1,]
testing = Sub_id[-part1,]
# Step 2
# Fitting data using Deviance and gini
tree_deviance = tree(as.factor(good_bad) ~ ., data = train, split = "deviance")
tree_gini = tree(as.factor(good_bad) ~ ., data = train, split = "gini")
# Prediction
## Misclassification for training data
devi_yfit = predict(tree_deviance, newdata = train,type="class")
gini_yfit = predict(tree_gini, newdata = train,type="class")
plot(tree_deviance)
plot(tree_gini)
devi_table = table(devi_yfit,train$good_bad)
gini_table = table(gini_yfit,train$good_bad)
devi_table
```

```
# Missclassification rate Deviance
missclass_devi <- 1-sum(diag(devi_table))/sum(devi_table)</pre>
missclass devi
gini_table
# Missclassification rate Gini
missclass_gini <- 1-sum(diag(gini_table))/sum(gini_table)</pre>
missclass_gini
## Misclssification for test data:
devi_yfit = predict(tree_deviance, newdata = testing,type="class")
gini_yfit = predict(tree_gini, newdata = testing,type="class")
plot(tree_deviance)
plot(tree_gini)
devi_table = table(devi_yfit,testing$good_bad)
gini_table = table(gini_yfit,testing$good_bad)
devi_table
# Missclassification rate Deviance
missclass_devi <- 1-sum(diag(devi_table))/sum(devi_table)</pre>
missclass_devi
gini table
# Missclassification rate Gini
missclass_gini <- 1-sum(diag(gini_table))/sum(gini_table)</pre>
missclass_gini
### Step 3
index = summary(tree_deviance)[4]$size
trainScore = rep(0,index)
testScore = rep(0,index)
# Graph training and validation
for(i in 2:index) {
  prunedTree=prune.tree(tree_deviance,best=i)
  pred=predict(prunedTree, newdata=validation,type="tree")
  trainScore[i] = deviance(prunedTree)
  testScore[i] = deviance(pred)
}
plot(2:index,trainScore[2:index], col="Red",type = "b", main = "Dependence of Deviance",
     ylim=c(min(testScore[2:index]),max(trainScore)), pch=19, cex=1, ylab="Deviance")
points(2:index,testScore[2:index],col="Blue",type="b", pch=19, cex=1)
# misclassification rate for test data
missclass_test_t = prune.tree(tree_deviance, best = 4)
summary(missclass_test_t)
```

```
yfit = predict(missclass_test_t, newdata = testing, type="class")
valid_ = table(testing$good_bad,yfit)
print("Confusion Matrix")
mc <- 1-sum(diag(valid_))/sum(valid_)</pre>
print("Misclassification rate")
plot(missclass test t)
text(missclass_test_t)
### Step 4
# Na?ve Bayes 2.4
naye = naiveBayes(good_bad ~., data=train)
nav_test = predict(naye, newdata = testing[,-ncol(testing)], type = "class")
nav_train = predict(naye,newdata = train[,-ncol(train)])
# Confusion Matrix Using Naive Bayes
nv_tbl_test = table(testing$good_bad,nav_test)
print(nv tbl test)
nv_tbl_train <- table(train$good_bad,nav_train)</pre>
print(nv_tbl_train)
# Missclassification train data value Using Naive Bayes
mc_nav_train <- 1-sum(diag(nv_tbl_train))/sum(nv_tbl_train)</pre>
cat("Misclassification train data value Using Naive Bayes is:",mc_nav_train)
# Missclassification test data value Using Naive Bayes
mc_nav_test <- 1-sum(diag(nv_tbl_test))/sum(nv_tbl_test)</pre>
cat("Misclassification test data value Using Naive Bayes is:",mc_nav_test)
### Step 5
# Naive Bayes With loss matrix 2.5
naye = naiveBayes(good_bad ~ ., data = train)
# Predicting using Naive
nav_test = predict(naye, testing[,-ncol(testing)] , type="raw")
nav_train = predict(naye, train[,-ncol(train)] , type="raw")
# applying loss matrix if greater then 10 True else False
nav_test = (nav_test[, 2] / nav_test[, 1]) > 10
nav_train = (nav_train[, 2] / nav_train[, 1]) > 10
# confusion matrix for train & test
naive_table = table(testing$good_bad,nav_test)
naive_table_train = table(train$good_bad,nav_train)
\# missclasification for train \& test
naive_table_train
1-sum(diag(naive_table_train))/sum(naive_table_train)
```

```
naive_table
1-sum(diag(naive_table))/sum(naive_table)
# Assignment 3
library(tree)
# 3.1 Data import, reorder and Plot
set.seed(12345)
data = read.csv2("State.csv", header = TRUE)
data = data[order(data$MET),]
plot(EX ~ MET, data = data, pch = 19, cex = 1,col="blue")
# 3.2
set.seed(12345)
control_parameter = tree.control(nobs = nrow(data),minsize = 8)
fit_tree = tree(formula = EX ~ MET,data = data,control = control_parameter)
leave_fit = cv.tree(fit_tree)
plot(leave_fit$size, leave_fit$dev, main = "Deviance Vs Size of Tree" ,
     xlab="Deviance", ylab = "Size", type="b",col="red", pch= 19,cex=1)
op_tree = prune.tree(fit_tree,best = leave_fit\$size[which.min(leave_fit\$dev)])
fitted_val = predict(op_tree, newdata=data)
df = data.frame(x = data$MET, pred = fitted_val, ex_val = data$EX)
fit_original_plot = ggplot(df, aes(x, pred, ex_val)) +
  geom_point(aes(x,ex_val), colour = "blue") +
  geom_point(aes(x, pred))
fit_original_plot
hist(residuals(op_tree))
# 3.3 Non-Paramatric Bootstrap
library(boot)
f_np = function(data,index){
  sample = data[index,]
 Ctrl = tree.control(nrow(sample), minsize = 8)
 fit = tree( EX ~ MET, data=sample, control = Ctrl)
 optimal_tree = prune.tree(fit, best= leave_fit$size[which.min(leave_fit$dev)])
 return(predict(optimal_tree, newdata=data))
np_bs = boot(data, statistic = f_np, R=1000)
conf_bound = envelope(np_bs,level=0.95)
```

```
predictions = predict(op_tree,data)
plot(np_bs)
fig_data = data.frame(orig = data$EX, x=data$MET, pred=predictions,
                      upper=conf_bound$point[1,], lower=conf_bound$point[2,])
fig = ggplot(fig_data, aes(x,predictions,upper,lower))
p = fig + geom_point(aes(x, pred)) +
  geom_point(aes(x, orig),colour="blue") +
  geom_line(aes(x,upper),colour="red") +
  geom_line(aes(x,lower),colour="red")
р
# 3.4 Paramatric Bootstrap
set.seed(12345)
parama_conf = function(data){
  controll = tree.control(nrow(data), minsize = 8)
  fit = tree( EX ~ MET, data=data, control = controll)
  op_tree = prune.tree(fit, best=leave_fit\$size[which.min(leave_fit\$dev)])
  return(predict(op_tree, newdata=data))
param_predict = function(data){
  controll = tree.control(nrow(data), minsize = 8)
  fit = tree( EX ~ MET, data=data, control = controll)
  op_tree = prune.tree(fit, best=leave_fit\$size[which.min(leave_fit\$dev)])
  predictions = predict(op_tree, newdata=data)
  return(rnorm(nrow(data), predictions, sd(resid(fit))))
rnd = function(data, model){
  sample = data.frame(MET=data$MET, EX=data$EX)
  sample$EX = rnorm(nrow(data), predict(model,newdata=data),sd(resid(model)))
  return(sample)
}
set.seed(12345)
param_boot_conf = boot(data, statistic = parama_conf, R=1000, mle = op_tree,
                       ran.gen = rnd, sim = "parametric")
confidence_bound_param = envelope(param_boot_conf, level=0.95)
param_boot_predict = boot(data, statistic = param_predict, R=1000, mle = op_tree, ran.gen = rnd, sim =
prediction_bound_param = envelope(param_boot_predict, level=0.95)
plot(param_boot_conf)
plot(param_boot_predict)
predictions = predict(op_tree,data)
fig_data = data.frame(orig = data$EX, x=data$MET, pred=predictions,
                      upper_c=confidence_bound_param$point[1,],
                      lower_c=confidence_bound_param$point[2,],
```