## Statistical Models: Homework 4

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### Question 1

In this problem, you practice working with predictor variables that are discrete. Consider the Boston dataset in the package MASS. Take as response the median property value.

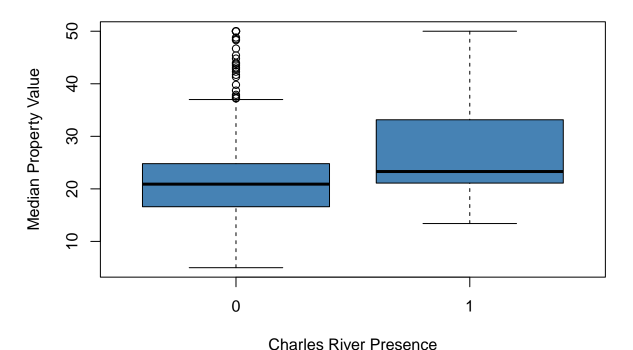
```
library(MASS)
library(ggplot2)
library(quantreg)
## Loading required package: SparseM
##
## Attaching package: 'SparseM'
## The following object is masked from 'package:base':
##
##
       backsolve
head(Boston)
##
        crim zn indus chas
                                     rm
                                         age
                                                dis rad tax ptratio
                                                                      black 1stat
## 1 0.00632 18
                 2.31
                         0 0.538 6.575 65.2 4.0900
                                                      1 296
                                                                15.3 396.90
                                                                             4.98
## 2 0.02731
                 7.07
                         0 0.469 6.421 78.9 4.9671
                                                      2 242
                                                                17.8 396.90
                                                                             9.14
                                                      2 242
## 3 0.02729
              0
                7.07
                         0 0.469 7.185 61.1 4.9671
                                                                17.8 392.83
                                                                             4.03
## 4 0.03237
                2.18
                         0 0.458 6.998 45.8 6.0622
                                                      3 222
                                                                18.7 394.63
## 5 0.06905
                                                      3 222
              0 2.18
                         0 0.458 7.147 54.2 6.0622
                                                                18.7 396.90
                                                                             5.33
## 6 0.02985
              0 2.18
                         0 0.458 6.430 58.7 6.0622
                                                      3 222
                                                                18.7 394.12 5.21
##
     medv
## 1 24.0
## 2 21.6
## 3 34.7
## 4 33.4
## 5 36.2
## 6 28.7
```

#### 1(a) For chas variable(Charles river)

Look at side-by-side boxplots for medv where the groups are defined by chas. Comment on what you observe. In particular, compare the different groups visually. Then fit a model explaining medv as a function of chas. Output an ANOVA table. What is the F-test testing? Is the result consistent with the boxplots?

#### **Creating Boxplot**

# Median Property Value(Medv) grouped by Charles River(chas)



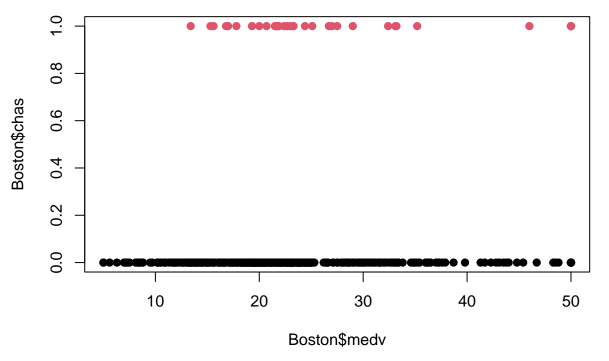
#### Inference:

- For chas = 1, the medy values lie between the 75 th and 25th quantile whereas for chas = 0, there are many values after the 75th quantile.
- The median values for both 0 and 1 category can be seen very closer to each other, approximately ranging near 20 to 25.
- We can see that median property values tend to be higher for properties near the Charles River compared to properties far from the Charles River
- This suggests that proximity to the Charles River has a positive effect on median property values.

```
#Comparing different groups visually table(Boston$chas)
```

```
## 0 1
## 471 35
```

plot(Boston\$medv,Boston\$chas,pch = 19, col=as.factor(Boston\$chas))



#### Inference:

- We can also visually see from the table and graph above that there are more data points for chas =0 than chas =1
- We can also observe that for chas = 0 the cluster values are more spread which can also be seen from the boxplot.

#### Fitting model

```
#Fitting a linear model
model_chas = lm(medv ~ chas, data = Boston)
summary(model_chas)
##
## Call:
## lm(formula = medv ~ chas, data = Boston)
##
## Residuals:
##
                                3Q
       Min
                1Q
                    Median
                                       Max
##
   -17.094 -5.894
                    -1.417
                             2.856
                                    27.906
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 22.0938
                            0.4176
                                    52.902 < 2e-16 ***
## chas
                 6.3462
                            1.5880
                                     3.996 7.39e-05 ***
##
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

```
##
## Residual standard error: 9.064 on 504 degrees of freedom
## Multiple R-squared: 0.03072, Adjusted R-squared: 0.02879
## F-statistic: 15.97 on 1 and 504 DF, p-value: 7.391e-05
```

#### Analysis of Variance (ANOVA)

1

## Residuals 504 41404

## chas

```
anova(model_chas)$F

## [1] 15.97151 NA

anova(model_chas)

## Analysis of Variance Table
##
## Response: medv
## Df Sum Sq Mean Sq F value Pr(>F)
```

Inference: What is the F-test testing? Is the result consistent with the boxplots?

## Signif. codes: 0 '\*\*\* 0.001 '\*\* 0.01 '\* 0.05 '.' 0.1 ' 1

82.15

1312 1312.08 15.972 7.391e-05 \*\*\*

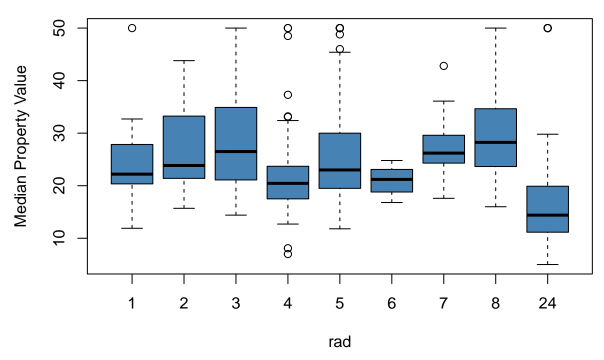
- F-test testing procedure that compares the variability of two or more sets of data
- For F test value =1 the variances are equal which is Null hypothesis and if it is not equal to 1 then there is variability
- We can also see that the F value here is 15.97 which denotes significant variability
- Null hypothesis for this case is: that the coefficients for the chas variable are equal to zero. This means that the chas variable has no effect on the median property value (medv). The null hypothesis assumes that the chas variable is not a significant predictor of medv. The alternative hypothesis is that the coefficients for the chas variable are not equal to zero, meaning that the chas variable does have an effect on medv and is a significant predictor.
- Since the p value if smaler than the significance level (0.05), the null hypothesis fails and we conclude that the **chas** variable is significantly related to **medv** meaning that the **chas** variable does have an effect on **medv** and is a significant predictor.
- This result is also consistent with what we can see from the boxplots that Charles river has impact that proximity to the Charles River has a positive effect on median property values.

#### 1(b) For Radial highway variable

Repeat with rad in place of chas.

#### Creating Boxplot

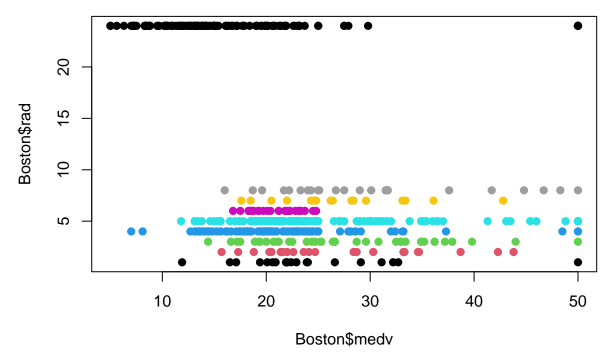
## Median Property Value(Medv) grouped by rad



#### Inference:

- Since, the boxes are of different heights, we can say that the median property value is not the same across all levels of rad.
- We can also observe that for higher value of rad (24), the median is less as compared to the closer rad values(1-8), that is for higher rad value (24) we can see that medv values is less. This means that the accessibility to radial highways generally lowers the median property values.

```
#Visually seeing the difference
table(Boston$rad)
##
##
         2
             3
                  4
                      5
                          6
                              7
                                  8
                                     24
     1
            38 110 115
    20
        24
                         26
                             17
                                 24 132
plot(Boston$medv,Boston$rad,pch = 19, col=as.factor(Boston$rad))
```



#### Inference:

• We can see that the data values for rad from 1-8 are quite well spread but for rad =24 are sparsely spread out with one data point above 40.

### Fitting the model

```
#Fitting the model
model_rad = lm(medv ~ rad, data = Boston)
summary(model_rad)
##
## Call:
## lm(formula = medv ~ rad, data = Boston)
##
## Residuals:
##
                1Q Median
                                3Q
       Min
                                       Max
                    -1.967
                                    33.292
##
  -17.770 -5.199
                             3.321
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 26.38213
                           0.56176
                                    46.964
                                              <2e-16 ***
## rad
               -0.40310
                           0.04349
                                    -9.269
                                              <2e-16 ***
##
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Signif. codes:
## Residual standard error: 8.509 on 504 degrees of freedom
## Multiple R-squared: 0.1456, Adjusted R-squared: 0.1439
## F-statistic: 85.91 on 1 and 504 DF, \, p-value: < 2.2e-16
```

#### Analysis of Variance (ANOVA)

```
anova(model_rad)$F
## [1] 85.91428
                     NA
anova(model_rad)
## Analysis of Variance Table
##
## Response: medv
##
              Df Sum Sq Mean Sq F value
                        6221.1 85.914 < 2.2e-16 ***
## rad
                   6221
## Residuals 504
                 36495
                          72.4
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

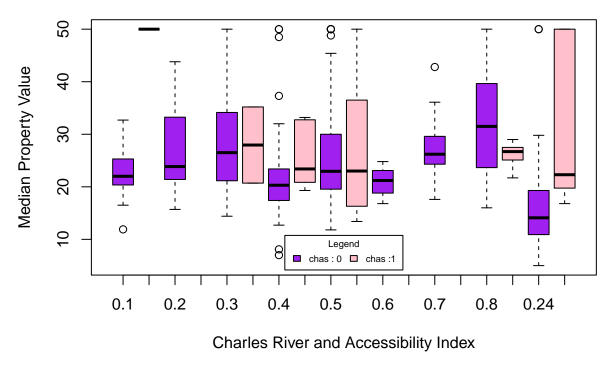
Inference: What is the F-test testing? Is the result consistent with the boxplots?

- F-test testing procedure that compares the variability of two or more sets of data
- For F test value =1 the variances are equal which is Null hypothesis and if it is not equal to 1 then there is variability
- We can also see that the F value here is 85.91 which denotes significant variability
- Null hypothesis for this case is: that the coefficients for the rad variable are equal to zero. This means that the rad variable has no effect on the median property value (medv). The null hypothesis assumes that the rad variable is not a significant predictor of medv. The alternative hypothesis is that the coefficients for the rad variable are not equal to zero, meaning that the rad variable does have an effect on medv and is a significant predictor.
- Since the p value if smaller than the significance level (0.05), the null hypothesis fails and we conclude that the rad variable is significantly related to medv meaning that the rad variable does have an effect on medv and is a significant predictor.
- This result is also consistent with what we can see from the boxplots that less accessibility to highways has an effect on median property values

#### 1(c) Boxplot

Produce a nice boxplot display of medv where the groups are defined by chas and rad jointly. Comment on what you observe. Then look at an interaction plot. Then fit a model explaining medv as a function of chas and rad with interactions. Output an ANOVA table. What are the different F-tests testing? Compare with the previous F-test as appropriate. Are the results of these tests consistent with the plots you just looked at?

# Median Property Value by Charles River and Accessibility Index



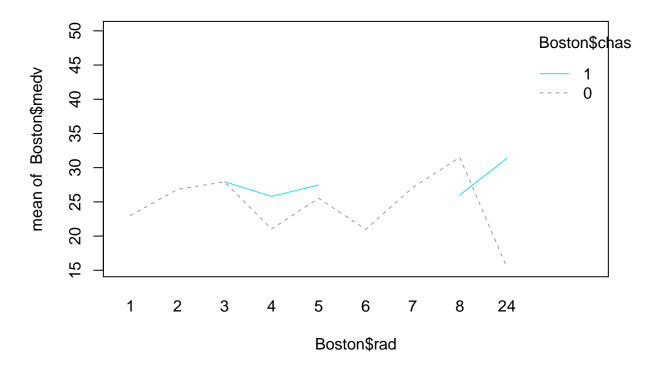
#### Inference:

- We can see that the grouping is done in 2 forms for chas where 1 and 0 indicate pink and purple boxes along with 9 values of rad for each, hence it creates 18 plots
- Another observation is that for some chas groups values(1) the rad values are missing.

### Interaction plot

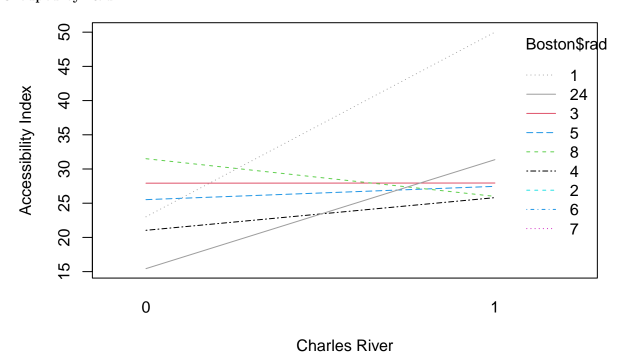
```
interaction.plot(Boston$rad, Boston$chas, Boston$medv, col = Boston$medv)
```

#### Grouped by Chas



Inference: We can see the variable trend for different values of rad for each of 0 and 1 groups chas.

### Grouped by Rad



Inference: We can see the lines for rad of different groups of chas for each of 0 and 1 groups

#### Fitting model and ANOVA

```
model_combined= lm(medv ~ chas * rad, data = Boston)
summary(model_combined)
##
## Call:
## lm(formula = medv ~ chas * rad, data = Boston)
##
## Residuals:
##
                1Q Median
      Min
                                ЗQ
                                       Max
## -17.527 -5.127 -1.796
                             3.548
                                    34.216
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 26.2762
                            0.5662 46.409 < 2e-16 ***
## chas
                0.7775
                            2.2042
                                     0.353 0.72445
                -0.4372
## rad
                            0.0437 -10.005 < 2e-16 ***
## chas:rad
                0.5860
                            0.1777
                                     3.297 0.00105 **
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## Residual standard error: 8.287 on 502 degrees of freedom
## Multiple R-squared: 0.1929, Adjusted R-squared: 0.188
## F-statistic: 39.98 on 3 and 502 DF, p-value: < 2.2e-16
anova(model_combined)
## Analysis of Variance Table
##
## Response: medv
##
              Df Sum Sq Mean Sq F value
                                           Pr(>F)
                  1312 1312.1 19.104 1.505e-05 ***
## chas
              1
```

#### Inference:

## rad

## chas:rad

F tests are used for different purposes that is for:

6179

746

746.5

68.7

## Signif. codes: 0 '\*\*\* 0.001 '\*\* 0.01 '\* 0.05 '.' 0.1 ' 1

1

1

## Residuals 502 34478

• testing equality of variance to test hypothesis of equality of two population variances

6179.4 89.972 < 2.2e-16 \*\*\*

10.869 0.001047 \*\*

- testing equality of several means to test for equality of several means is carried out by the technique called ANOVA
- for testing significance of regression is used to test the significance of the regression model

Comparision to previous F test:

As seen before, the chas and rad F test values were 15.97 and 85.91 but here we can see an increase in F test values to be 19.10 and 89.97 for degree of freedom 1

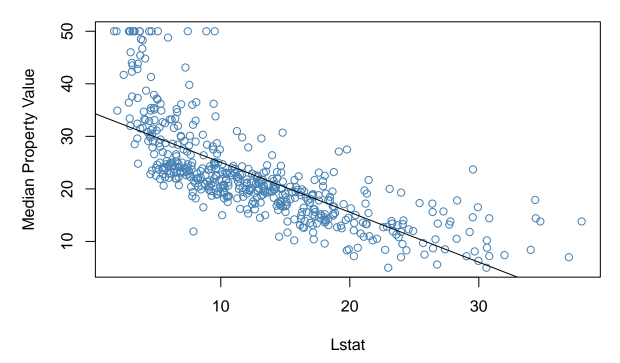
We can also see the combined effect of chas and rad where the F test values is 10.86 for degree of freedom 1. We can also see a decrease in significance of p values.

The observed values don't seem very consistent with the plots

#### 1(d) Checking for chas

It makes sense that median property value decreases with the percentage of lower status population lstat, and this is indeed what is observed here. Does the rate of decrease depend on whether the area borders the Charles River? Produce a plot that helps answer that question.

# Median Property Value(Medv) vs Lstat

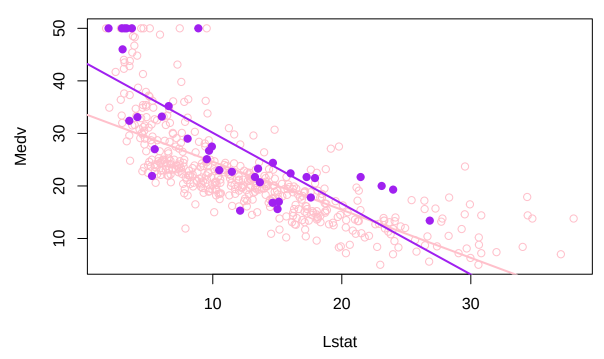


We can observes that as the the value of lstat decreases, the values of medv increases. It is almost inversely proportional.

```
ind = (Boston$chas==0)
plot(Boston$medv[ind] ~ Boston$lstat[ind], data = Boston,
    main = "Plot medv vs lstat",col="pink",
    xlab="Lstat", ylab = "Medv")
fit = lm(Boston$medv[ind] ~ Boston$lstat[ind], data = Boston)
```

```
abline(fit, col="pink", lwd=2)
ind = (Boston$chas==1)
points(Boston$medv[ind] ~ Boston$lstat[ind], data = Boston, col="purple", pch=19)
fit = lm(Boston$medv[ind] ~ Boston$lstat[ind], data = Boston)
abline(fit, col="purple", lwd=2)
```

### Plot medv vs Istat



#### Inference:

• We can see from the interaction plot that the houses near the Charles river (chas =1) that is the purple dots has a higher median property value as compared to the ones that are far away from the Charles river(chas =0) that is the pink dots.

# Question 2

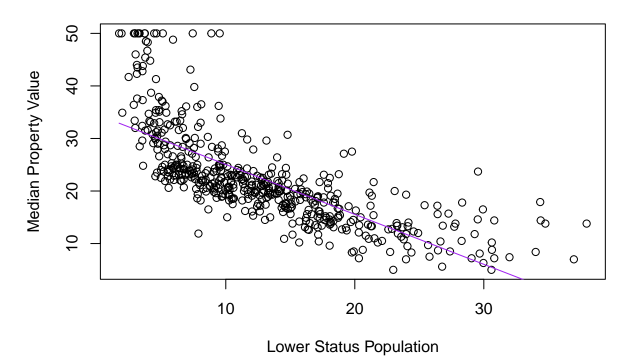
Consider the same dataset and turn to the problem of fitting a polynomial model explaining med as a function of lstat.

#### Fit a polynomial model of degree 1

First we will fit a model of degree of freedom = 1

```
ylab = "Median Property Value")
lines(sort(Boston$lstat), predict(fit,
    newdata = data.frame(lstat = sort(Boston$lstat))),
    col = "purple")
```

### Median Property Value vs. Lower Status Population(DEGREE 1)

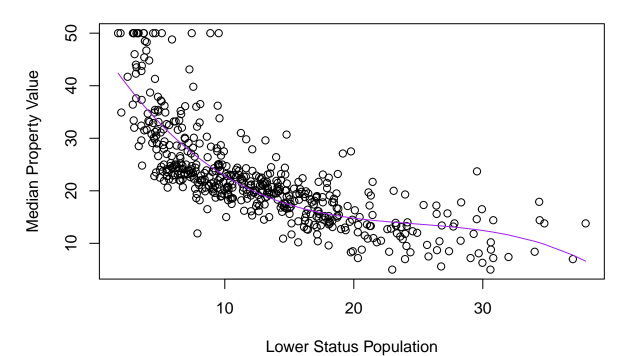


### #summary(fit)

2a Fit a polynomial model of degree 3 by least squares.

Fit a polynomial model of degree 3 by least squares.

### Median Property Value vs. Lower Status Population (DEGREE 3)



#### summary(fit)

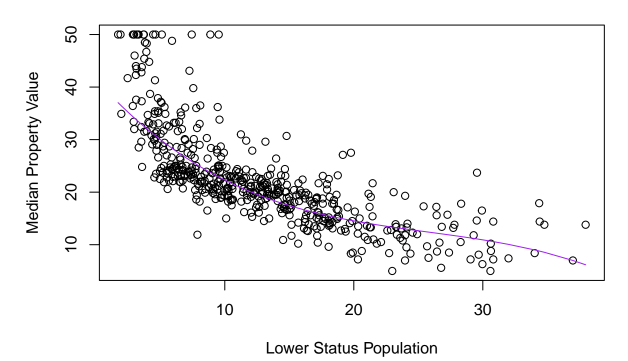
```
##
## lm(formula = medv ~ poly(lstat, 3), data = Boston)
##
## Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                           Max
## -14.5441 -3.7122 -0.5145
                               2.4846
                                       26.4153
## Coefficients:
                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                0.2399 93.937 < 2e-16 ***
                    22.5328
## poly(lstat, 3)1 -152.4595
                                5.3958 -28.255 < 2e-16 ***
## poly(lstat, 3)2
                                5.3958 11.903 < 2e-16 ***
                    64.2272
## poly(lstat, 3)3 -27.0511
                                5.3958 -5.013 7.43e-07 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 5.396 on 502 degrees of freedom
## Multiple R-squared: 0.6578, Adjusted R-squared: 0.6558
## F-statistic: 321.7 on 3 and 502 DF, p-value: < 2.2e-16
```

Similar plot with different library:

```
#Fit a polynomial model of degree 3 by regression qualtile library - rq
fit <- rq(medv ~ poly(lstat, 3), data = Boston, tau = 0.5)
plot(Boston$lstat, Boston$medv,</pre>
```

```
main = "Median Property Value vs. Lower Status Population (DEGREE 3)",
    xlab = "Lower Status Population",
    ylab = "Median Property Value")
lines(sort(Boston$lstat),
    predict(fit, newdata = data.frame(lstat = sort(Boston$lstat))),
    col = "purple")
```

# Median Property Value vs. Lower Status Population (DEGREE 3)



# #summary(fit)

### Inference:

As we can see from the above plots of degree of polynomial = 1 and degree of polynomial = 3, we can see a better fit in degree of polynomial = 3 rather than degree of polynomial = 1.

This is because degree of polynomial = 3 model adjusts more to the current data

For degree of polynomial = 1 we can see high bias whereas for degree of polynomial = 3 there is much more optimal fit.

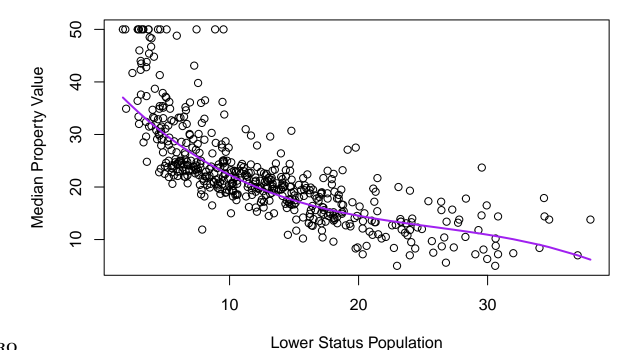
2b Repeat with each robust method covered in the lecture notes/slides.

### Robust methods

```
fit_rq <- rq(medv ~ poly(lstat, 3), data = Boston, tau = 0.5)
plot(Boston$lstat, Boston$medv,
    main = "Median Property Value vs. Lower Status Population(RQ)",
    xlab = "Lower Status Population",</pre>
```

```
ylab = "Median Property Value")
lines(sort(Boston$lstat),
    predict(fit_rq, newdata = data.frame(lstat = sort(Boston$lstat))),
    col = "purple", lwd = 2)
```

### Median Property Value vs. Lower Status Population(RQ)



 $\mathbf{R}\mathbf{Q}$ 

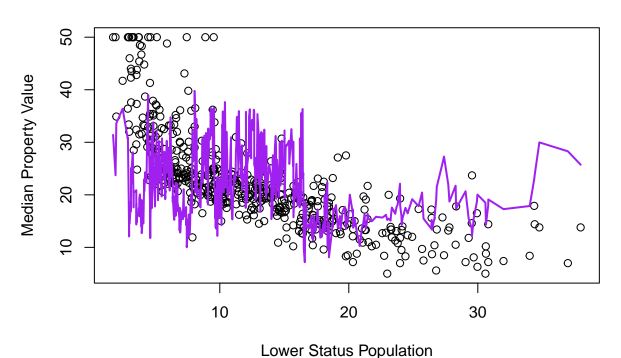
```
m.huber = rlm(Boston$medv ~ poly(Boston$lstat, 3), psi = psi.huber)
summary(m.huber)
```

#### **Huber's M-estimation:**

```
##
## Call: rlm(formula = Boston$medv ~ poly(Boston$lstat, 3), psi = psi.huber)
## Residuals:
##
        Min
                  1Q
                       Median
                                             Max
## -13.8373 -3.0157 -0.2219
                                2.7329
                                        26.8409
##
## Coefficients:
##
                          Value
                                    Std. Error t value
## (Intercept)
                            21.9774
                                        0.2037
                                                 107.8773
## poly(Boston$1stat, 3)1 -143.0378
                                        4.5827
                                                 -31.2126
## poly(Boston$1stat, 3)2
                            55.9111
                                                  12.2005
                                        4.5827
## poly(Boston$1stat, 3)3 -21.7848
                                                  -4.7537
                                        4.5827
## Residual standard error: 4.323 on 502 degrees of freedom
```

```
plot(Boston$lstat, Boston$medv,
    main = "Median Property Value vs. Lower Status Population(Huber)",
    xlab = "Lower Status Population",
    ylab = "Median Property Value")
lines(sort(Boston$lstat),
    predict(m.huber, newdata = data.frame(lstat = sort(Boston$lstat))),
    col = "purple", lwd = 2)
```

# Median Property Value vs. Lower Status Population(Huber)



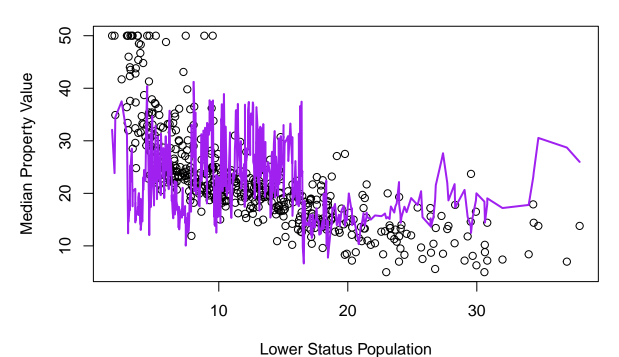
m.hampel = rlm(Boston\$medv ~ poly(Boston\$lstat, 3), psi = psi.hampel)
summary(m.hampel)

#### Hampel's M-estimation:

```
##
## Call: rlm(formula = Boston$medv ~ poly(Boston$1stat, 3), psi = psi.hampel)
## Residuals:
##
       Min
                  1Q
                      Median
## -14.0819 -3.3423 -0.3086
                                2.6869
                                        26.7536
## Coefficients:
                          Value
                                    Std. Error t value
                                       0.2161
## (Intercept)
                            22.2032
                                                102.7330
## poly(Boston$1stat, 3)1 -147.0125
                                       4.8616
                                                -30.2395
## poly(Boston$1stat, 3)2
                            60.7791
                                       4.8616
                                                12.5019
## poly(Boston$1stat, 3)3 -25.7454
                                       4.8616
                                                 -5.2957
```

```
##
## Residual standard error: 4.547 on 502 degrees of freedom
```

# Median Property Value vs. Lower Status Population(Hampel)



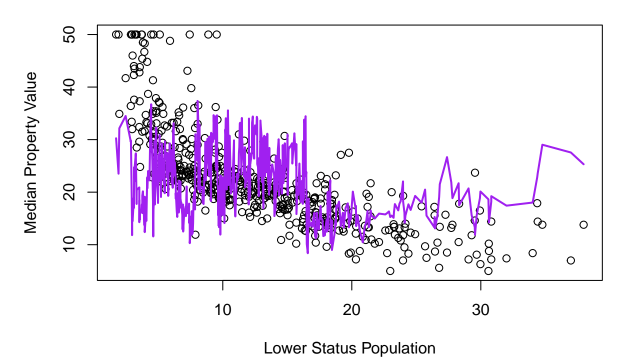
```
m.tukey = rlm(Boston$medv ~ poly(Boston$lstat, 3), psi = psi.bisquare)
summary(m.tukey)
```

#### Tukey's M-estimation:

```
##
## Call: rlm(formula = Boston$medv ~ poly(Boston$lstat, 3), psi = psi.bisquare)
## Residuals:
                       Median
                                            Max
                  1Q
## -13.4084 -2.7037 -0.1377
                                2.9575 27.0035
## Coefficients:
                          Value
                                    Std. Error t value
##
## (Intercept)
                            21.6329
                                       0.1962
                                               110.2426
```

```
## poly(Boston$1stat, 3)1 -134.8470
                                       4.4141
                                                -30.5493
## poly(Boston$1stat, 3)2 48.3681
                                       4.4141
                                                 10.9577
## poly(Boston$1stat, 3)3 -15.4625
                                                 -3.5030
                                       4.4141
##
\#\# Residual standard error: 4.172 on 502 degrees of freedom
plot(Boston$lstat, Boston$medv,
     main ="Median Property Value vs. Lower Status Population(Tukey)",
     xlab = "Lower Status Population",
     ylab = "Median Property Value")
lines(sort(Boston$1stat),
      predict(m.tukey, newdata = data.frame(lstat = sort(Boston$lstat))),
      col = "purple", lwd = 2)
```

## Median Property Value vs. Lower Status Population(Tukey)



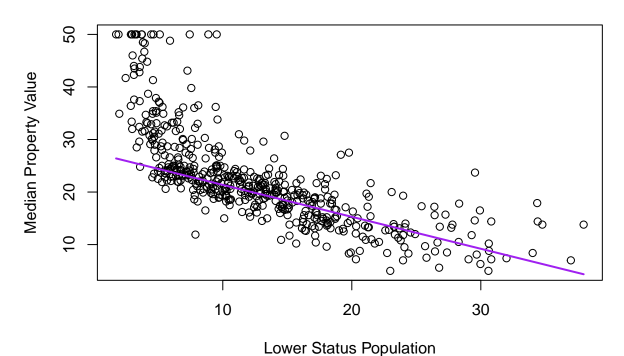
```
# high breakdown point methods
fit.lms = lmsreg(medv ~ lstat, data = Boston)
summary(fit.lms)
```

### High breakdown point methods - Least Median of Squares (LMS)

##		Length	Class	Mode
##	crit	1	-none-	numeric
##	sing	1	-none-	${\tt character}$
##	coefficients	2	-none-	numeric
##	bestone	2	-none-	numeric

```
## fitted.values 506
                        -none-
                                    numeric
## residuals
                 506
                                    numeric
                        -none-
## scale
                        -none-
                                    numeric
## terms
                   3
                        terms
                                    call
## call
                   4
                        -none-
                                    call
## xlevels
                   0
                        -none-
                                    list
## model
                        data.frame list
plot(Boston$lstat, Boston$medv,
     main = "Median Property Value vs. Lower Status Population(LMS)",
     xlab = "Lower Status Population", ylab = "Median Property Value")
lines(sort(Boston$1stat),
      predict(fit.lms,
      newdata = data.frame(lstat = sort(Boston$lstat))),
      col = "purple", lwd = 2)
```

### Median Property Value vs. Lower Status Population(LMS)



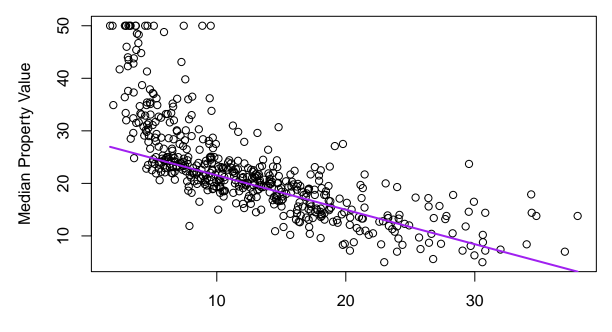
```
fit.lts = ltsreg(medv ~ lstat, data = Boston)
summary(fit.lts)
```

#### High breakdown point methods - Least Trimmed Squares (LTS)

```
## Length Class Mode
## crit 1 -none- numeric
## sing 1 -none- character
## coefficients 2 -none- numeric
```

```
## bestone
                        -none-
                                    numeric
## fitted.values 506
                                    numeric
                        -none-
## residuals
                 506
                        -none-
                                    numeric
## scale
                   2
                                    numeric
                        -none-
## terms
                   3
                        terms
                                    call
## call
                   4
                                    call
                        -none-
## xlevels
                   0
                         -none-
                                    list
## model
                        data.frame list
plot(Boston$lstat, Boston$medv,
     main = "Median Property Value vs. Lower Status Population(LTS)",
     xlab = "Lower Status Population", ylab = "Median Property Value")
lines(sort(Boston$1stat),
      predict(fit.lts,
      newdata = data.frame(lstat = sort(Boston$lstat))),
      col = "purple", lwd = 2)
```

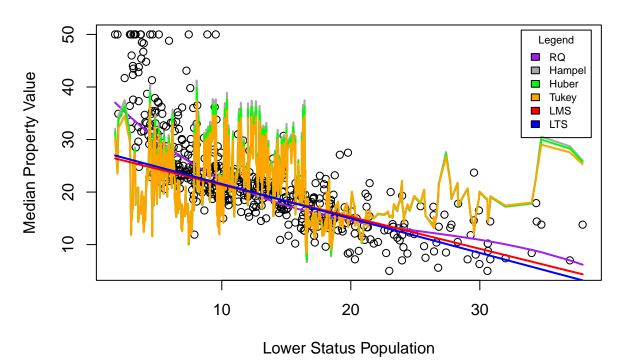
### Median Property Value vs. Lower Status Population(LTS)



**Lower Status Population** 

```
lines(sort(Boston$lstat), predict(m.tukey, newdata = data.frame(lstat = sort(Boston$lstat))),
        col = "orange", lwd = 2)
lines(sort(Boston$lstat), predict(fit.lms, newdata = data.frame(lstat = sort(Boston$lstat))),
        col = "red", lwd = 2)
lines(sort(Boston$lstat), predict(fit.lts, newdata = data.frame(lstat = sort(Boston$lstat))),
        col = "blue", lwd = 2)
legend("topright", inset = 0.02,
        title = "Legend", c("RQ", "Hampel", "Huber", "Tukey", "LMS", "LTS"),
        fill = c("purple", "darkgrey", "green", "orange", "red", "blue"),
        horiz = FALSE, cex = 0.7)
```

# 2c Produce a scatterplot and overlay all these fits with different colors and a legend. Scatterplot and Overlay for robust methods



#### Team Contributions:

Both the team members Sourabh Prakash and Priyanshi Shah have contributed equally to the homework by discussing the key points and logic together and doing pair programming. For the implementation part question 1 a,b,c was contributed by Priyanshi Shah and question 1d and 2a,b,c by Sourabh Prakash. The conclusions were written together for all the parts of question 1 and 2