Assignment 2

Andrea Di Dio, Chenghan Song, Jiacheng Lu — Group 22

Exercise 1

a)

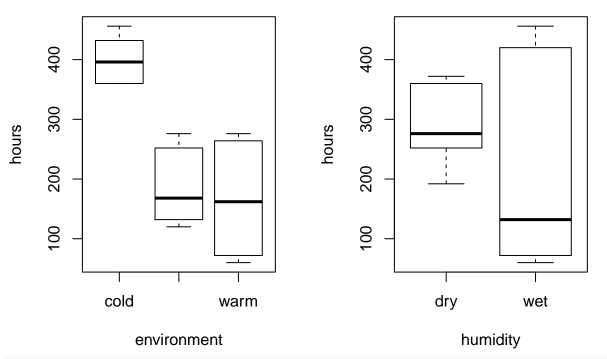
We assign combine levels (i,j) of the factors to a random set of N units

```
I=3; J=2; N=3
rbind(rep(1:I,each=N*J),rep(1:J,N*I),sample(1:(N*I*J)))
        [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10] [,11] [,12] [,13] [,14]
## [1,]
                                                      2
                            1
                                      1
                                            2
                                                 2
                                                             2
                      1
                                 1
                                                             2
                            2
                                                                          2
                                                                                      2
## [2,]
           1
                 2
                      1
                                      2
                                            1
                                                 2
                                                      1
                                                                   1
                                                                                1
                                 1
                                13
                                      4
                                           12
                                                 6
                                                      8
                                                                         11
                                                                                7
                                                                                      2
## [3,]
          14
                15
                      9
                          17
                                                            10
                                                                  18
        [,15] [,16] [,17] [,18]
## [1,]
            3
                   3
                         3
## [2,]
            1
                   2
                         1
                                2
## [3,]
             3
                   1
                        16
b)
breadata=read.table(file="bread.txt",header=TRUE);par(mfrow=c(1,2))
```

```
breadata=read.table(file="bread.txt",header=TRUE);par(mfrow=c(1,2))
boxplot(hours~environment,data=breadata, main="box plot of hours-environment")
boxplot(hours~humidity, data=breadata,main="box plot of hours-humidity")
```

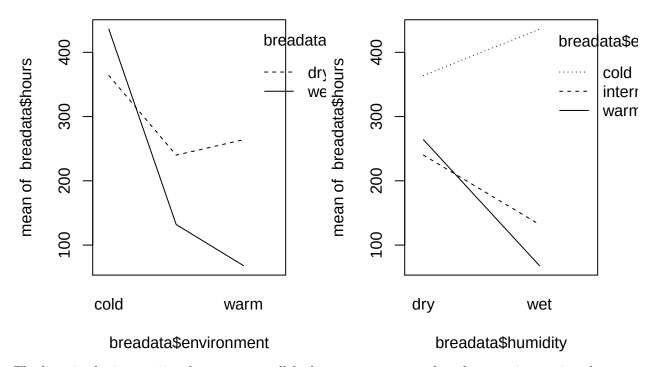
box plot of hours-environment

box plot of hours-humidity



interaction.plot(breadata\$environment,breadata\$humidity,breadata\$hours, main="interaction plot(humidity
interaction.plot(breadata\$humidity,breadata\$environment,breadata\$hours, main="interaction plot(environment)

interaction plot(humidity fixed) interaction plot(environment fixed)



The lines in the interaction plots are unparallel, thus we can assume that there are interactions between temperature and humidity.

c)

```
breadaov=lm(hours~environment*humidity, data = breadata); anova(breadaov)
```

```
## Analysis of Variance Table
##
## Response: hours
##
                       Df Sum Sq Mean Sq F value Pr(>F)
## environment
                         2 201904
                                  100952
                                            233.7 2.5e-10 ***
                            26912
                                             62.3 4.3e-06 ***
## humidity
                                    26912
                         1
## environment:humidity
                        2
                            55984
                                    27992
                                             64.8 3.7e-07 ***
## Residuals
                        12
                             5184
                                      432
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

 $H_0:\alpha_i=0$ for all i: The temperature does not have a main effect on the time of decay

 H_0 : $\beta_i = 0$ for all j: The humidity does not have a main effect on the time of decay

 H_0 : $\gamma_{ij} = 0$ for all (i, j): Temperature and humidity do not have interaction effects on the time of decay

The p-value for testing $H_0:\alpha_i=0$ for all i is 2.461e-10, hence H_0 is rejected; for $H_0:\beta_j=0$ for all j is 4.316e-06, hence H_0 is rejected; for $H_0:\gamma_{ij}=0$ for all (i,j) is 3.705e-07, hence H_0 is rejected. So both temperature and humidity have main effects and there are also interactions between two factors.

summary(breadaov)

```
##
## Call:
## lm(formula = hours ~ environment * humidity, data = breadata)
##
## Residuals:
##
      Min
              10 Median
                            30
                                  Max
      -48
              -7
                                   36
##
                      0
                            11
##
## Coefficients:
##
                                       Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                                              30.33 1.0e-12 ***
                                             364
                                                         12
## environmentintermediate
                                            -124
                                                         17
                                                              -7.31
                                                                     9.4e-06 ***
## environmentwarm
                                            -100
                                                         17
                                                              -5.89
                                                                     7.3e-05 ***
## humiditywet
                                              72
                                                         17
                                                               4.24
                                                                      0.0011 **
## environmentintermediate:humiditywet
                                            -180
                                                         24
                                                              -7.50
                                                                     7.2e-06 ***
## environmentwarm:humiditywet
                                                         24
                                                             -11.17
                                                                     1.1e-07 ***
                                            -268
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 20.8 on 12 degrees of freedom
## Multiple R-squared: 0.982, Adjusted R-squared: 0.975
## F-statistic: 132 on 5 and 12 DF, p-value: 4.68e-10
```

 H_0 for the interaction of intermediate environment and wet humidity: there is no interaction effect of intermediate environment and wet humidity. The p-value is 7.23e-06, therefore we reject H_0 and conclude that there is a significant interaction between intermediate temperature and wet humidity.

 H_0 for the interaction of warm environment and wet humidity: there is no interaction effect of warm environment and wet humidity. The p-value is 1.07e-07, therefore we reject H_0 and conclude that there is a significant interaction between warm temperature and wet humidity.

d)

This is not a good question. Because the interaction effects between two factors are significant. We can not compare the influence of the first and second factor when there are interactions between them.

e) par(mfrow=c(1,2));qqnorm(residuals(breadaov), main="QQ-plot residuals"); plot(fitted(breadaov),residual

residuals VS fitted values QQ-plot residuals 0 0 20 20 0 residuals(breadaov) Sample Quantiles 0 00 0 0 0 0 0 0 0 0 0 -20 2 **-1** 0 1 200 300 100 400 Theoretical Quantiles fitted(breadaov)

From the QQ-Plot we can see that the residuals are not normally distributed.

The spread in the residuals seems to be bigger for some certain fitted values. Due to the spread in the residuals should not change systematically with any variable, in particular not with the fitted values, there might be some outliers.

Exercise 2

a)

For the randomized block design, we perform the experiment with 5 blocks (B) using 3 interfaces (I) and have one replicate per treatment level per block (N).

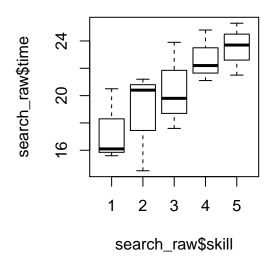
```
search_raw <- read.delim("./search.txt", header = TRUE, sep = "")
I <- 3; B <- 5; N <- 1;
for(i in 1:B) print(sample(1:(N * I)))

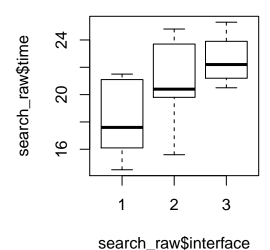
## [1] 2 3 1
## [1] 3 1 2
## [1] 1 3 2
## [1] 2 3 1</pre>
```

b) par(mfrow = c(1,2)) boxplot(search_raw\$time ~ search_raw\$skill, main = "time VS skill Boxplot") boxplot(search_raw\$time ~ search_raw\$interface, main = "time VS interface Boxplot")

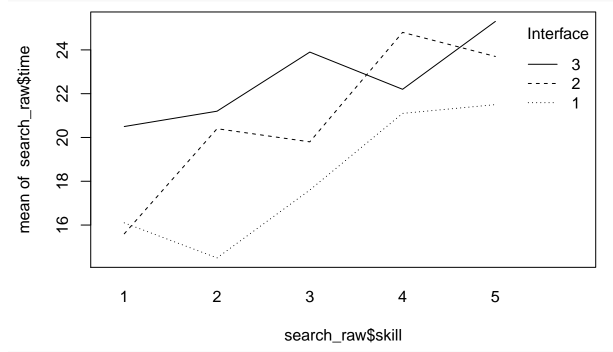
time VS skill Boxplot

time VS interface Boxplot

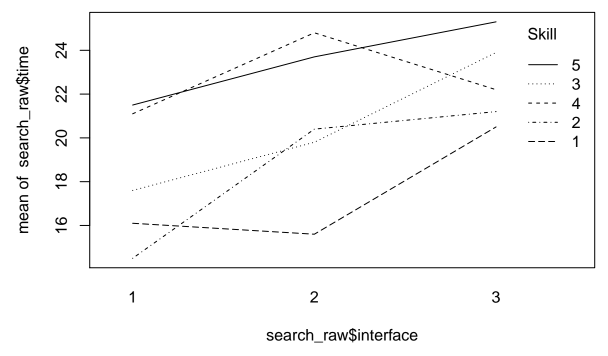




interaction.plot(search_raw\$skill, search_raw\$interface, search_raw\$time, trace.label = "Interface")



interaction.plot(search_raw\$interface, search_raw\$skill, search_raw\$time, trace.label = "Skill")



We have used two different graphical summaries in order to investigate the interaction between the interfaces and skill of the students on the time taken to complete the task. The boxplots show that both factors (skill and interface) affect the dependent variable (time). The lower the value of the skill factor (and hence the higher the competence of the student) result in lower times to complete the task. Also the interface factor seems to result in a difference of time taken to complete the task. This shows that there exists a dependence of the dependent variable on the factors of interest. The interaction plots show that the lines plotted are not parallel to one another, implying a significant interaction between the skill and the interface used to complete the task.

c)

 H_0 : The search time is the same for all the interfaces (There is no interaction between the factors). Given that our H_0 says that there is no interactions, we test using the additive model:

```
search_raw$skill <- as.factor(search_raw$skill)</pre>
search_raw$interface <- as.factor(search_raw$interface)</pre>
search_aov <- lm(search_raw$time ~ search_raw$skill + search_raw$interface, data = search_raw)
print(anova(search_aov), signif.stars = F)
## Analysis of Variance Table
##
## Response: search_raw$time
##
                         Df Sum Sq Mean Sq F value Pr(>F)
## search_raw$skill
                              80.1
                                      20.01
                                               6.21 0.014
                          2
                              50.5
                                      25.23
                                               7.82 0.013
## search_raw$interface
## Residuals
                              25.8
                                       3.23
```

The p-value $0.013 < \alpha$ meaning that we can safely reject H_0 concluding that the interfaces used affect the search time.

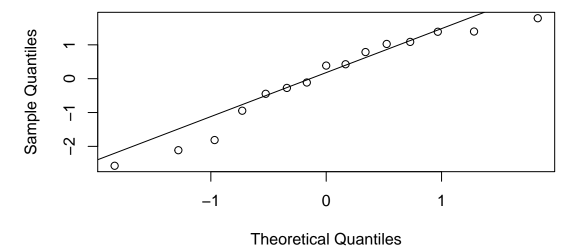
```
contrasts(search_raw$interface) <- contr.sum
contrasts(search_raw$skill) <- contr.sum
search_aov2 <- lm(search_raw$time ~ search_raw$interface + search_raw$skill)
summary(search_aov2)</pre>
```

```
##
## Call:
## lm(formula = search_raw$time ~ search_raw$interface + search_raw$skill)
##
## Residuals:
              1Q Median
##
      Min
                            3Q
                                  Max
  -2.573 -0.697 0.387
                        1.057
                                1.787
##
## Coefficients:
##
                         Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                           20.547
                                       0.464
                                                44.31
                                                       7.4e-11 ***
                           -2.387
## search_raw$interface1
                                       0.656
                                                -3.64
                                                        0.0066 **
## search_raw$interface2
                            0.313
                                       0.656
                                                 0.48
                                                        0.6456
## search_raw$skill1
                                                        0.0095 **
                           -3.147
                                       0.927
                                                -3.39
## search_raw$skill2
                           -1.847
                                                -1.99
                                                        0.0816 .
                                       0.927
## search_raw$skill3
                           -0.113
                                       0.927
                                                -0.12
                                                        0.9057
                                                 2.32
                                                        0.0488 *
## search_raw$skill4
                            2.153
                                       0.927
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.8 on 8 degrees of freedom
## Multiple R-squared: 0.835, Adjusted R-squared:
## F-statistic: 6.74 on 6 and 8 DF, p-value: 0.0084
```

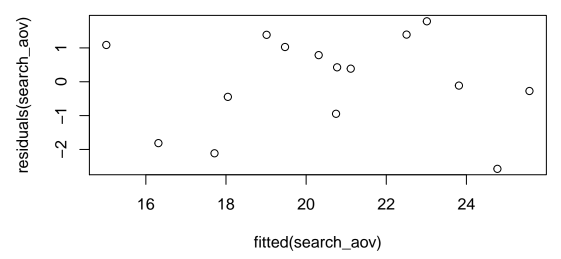
To estimate the time taken by a student with skill level 3 to complete the search task using interface 2, we should sum the intercept term with the coefficients of skill3 and interface 2.20.5467 + (-0.1133) + 0.3133 = 20.7467.

```
d)
options(digits = 6)
qqnorm(residuals(search_aov));qqline(residuals(search_aov))
```

Normal Q-Q Plot



```
plot(fitted(search_aov), residuals(search_aov))
```



print(shapiro.test(residuals(search_aov))) # fail to reject HO, might be normal

```
##
## Shapiro-Wilk normality test
##
## data: residuals(search_aov)
## W = 0.9309, p-value = 0.282
```

The QQ plot of the residuals for the data shows that the data might seem to be normally distributed and this is also confirmed by the **Shapiro-Wilk test** which shows a p-value of 0.282. The plot which shows the fitted values VS the residuals show that there is no systematic change in the residuals based on the fitted values as the points are well-spread out, suggesting that the two populations have equal variances.

e)

 H_0 : There is no effect in the interface used on the search time.

```
options(digits = 5)
friedman.test(search_raw$time, search_raw$interface, search_raw$skill)

##
## Friedman rank sum test
##
## data: search_raw$time, search_raw$interface and search_raw$skill
## Friedman chi-squared = 6.4, df = 2, p-value = 0.041
```

The non-parametric Friedman test gives a p-value of $0.041 < \alpha$ meaning that we can safely reject H_0 meaning that the choice in the interface used for the search engine has a statistically significant effect on the search time.

f)

 H_0 : The interface used does not affect the search time.

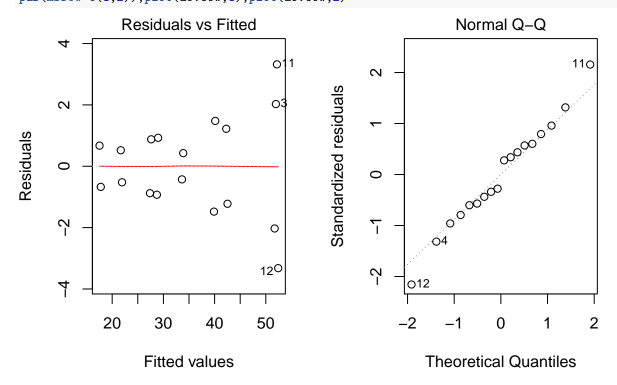
```
search_aov_one_way <- lm(search_raw$time ~ search_raw$interface, data = search_raw)
print(anova(search_aov_one_way), signif.stars = F)
## Analysis of Variance Table</pre>
```

```
## search_raw$interface 2 50.5 25.23 2.86 0.096 ## Residuals 12 105.9 8.82
```

The resulting p-value is $0.096 > \alpha$ meaning that we fail to reject H_0 and hence could be possible that the interfaces have no effect on the search time. However, given that the testing in part b) suggest that there is interaction between the skill and the interface factors, this test is not useful and wrong. In order for this test to be valid, we must assume no interaction between the two factors on the dependent variable, but this is not the case meaning that this test is not statistically meaningful.

Exercise 3

```
a)
cow=read.table(file="cow.txt",header=TRUE)
cow$id=factor(cow$id); cow$per=factor(cow$per)
aovcow=lm(milk~treatment+id,data=cow); anova(aovcow)
## Analysis of Variance Table
##
##
  Response: milk
##
              {\tt Df \; Sum \; Sq \; Mean \; Sq \; F \; value}
                                           Pr(>F)
                             0.3
                                     0.05
                                              0.83
##
   treatment
##
               8
                    2467
                           308.4
                                    57.74 2.8e-06
   id
               8
                             5.3
## Residuals
                      43
##
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
par(mfrow=c(1,2));plot(aovcow,1);plot(aovcow,2)
```



 H_0 : the type of feedingstuffs does not influence milk production

The p-value for testing H_0 is 0.8281, thus we fail to reject H_0 , the type of feedingstuffs does not have an effect on milk production. From the plot above, the residuals do not seem to deviate significantly from normal.

However, an ordinary "mixed effects" model is not suitable in this case where the assumption of "exchangebility" may fail. Cows may be happy with or bored at feedingstuff, then a block design is invalid.

b)

```
cowlm=lm(milk~treatment+per+id,data=cow); anova(cowlm);summary(cowlm)
## Analysis of Variance Table
##
## Response: milk
            Df Sum Sq Mean Sq F value
                    0
                          0.3
                                 0.11
                                        0.751
## treatment
             1
                   25
                          25.4
                                 10.25
                                        0.015 *
## per
                         308.4 124.48 7.5e-07 ***
## id
             8
                  2467
## Residuals 7
                   17
                           2.5
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Call:
## lm(formula = milk ~ treatment + per + id, data = cow)
##
## Residuals:
##
     Min
             1Q Median
                            3Q
                                  Max
## -2.260 -0.438 0.000 0.438
                               2.260
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
                30.300
                            1.244
                                     24.35 5.0e-08 ***
## (Intercept)
                                     -0.68 0.51654
## treatmentB
                -0.510
                            0.747
## per2
                -2.390
                            0.747
                                     -3.20 0.01505 *
                23.000
## id2
                             1.574
                                    14.61 1.7e-06 ***
## id3
                11.150
                             1.574
                                     7.08 0.00020 ***
                                    -0.86 0.41948
## id4
                -1.350
                            1.574
## id5
                -7.050
                            1.574
                                     -4.48 0.00287 **
## id6
                23.450
                             1.574
                                     14.90 1.5e-06 ***
## id7
                            1.574
                                     8.61 5.7e-05 ***
                13.550
## id8
                 4.900
                            1.574
                                     3.11 0.01701 *
## id9
               -11.200
                             1.574
                                     -7.12 0.00019 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.57 on 7 degrees of freedom
## Multiple R-squared: 0.993, Adjusted R-squared: 0.983
## F-statistic: 101 on 10 and 7 DF, p-value: 1.35e-06
library("lme4")
## Loading required package: Matrix
cowlmer=lmer(milk~treatment+order+per+(1|id),REML=FALSE, data = cow);summary(cowlmer)
## Linear mixed model fit by maximum likelihood ['lmerMod']
## Formula: milk ~ treatment + order + per + (1 | id)
##
      Data: cow
##
```

```
##
        AIC
                 BIC
                       logLik deviance df.resid
##
      119.3
               124.7
                         -53.7
                                  107.3
                                               12
##
##
  Scaled residuals:
##
       Min
                1Q Median
                                 3Q
                                        Max
   -1.5311 -0.3710 0.0269 0.2675
                                    1.7249
##
##
## Random effects:
##
    Groups
             Name
                          Variance Std.Dev.
##
    id
             (Intercept) 133.15
                                   11.54
    Residual
                            1.93
                                    1.39
## Number of obs: 18, groups: id, 9
##
## Fixed effects:
               Estimate Std. Error t value
##
## (Intercept)
                 38.500
                              5.811
                                       6.63
                                      -0.77
                 -0.510
                              0.658
## treatmentB
## orderBA
                 -3.470
                              7.768
                                      -0.45
                 -2.390
                              0.658
                                      -3.63
##
  per2
##
## Correlation of Fixed Effects:
              (Intr) trtmnB ordrBA
##
## treatmentB -0.063
              -0.743 0.000
## orderBA
## per2
              -0.063 0.111 0.000
cowlmer1 = lmer(milk~order+per+(1|id),REML=FALSE, data = cow)
anova(cowlmer, cowlmer1)
## Data: cow
## Models:
## cowlmer1: milk ~ order + per + (1 | id)
## cowlmer: milk ~ treatment + order + per + (1 | id)
##
            Df AIC BIC logLik deviance Chisq Chi Df Pr(>Chisq)
## cowlmer1 5 118 122 -53.9
                                    108
## cowlmer
             6 119 125 -53.7
                                    107
                                         0.58
                                                    1
                                                            0.45
```

 H_0 : the type of feedingstuffs does not influence milk production

In fixed effects analysis, the p-value for testing H_0 is 1.349e-06, thus H_0 is rejected, the type of feedingstuffs influences milk production

In mixed effects analysis, the p-value for testing H_0 is 0.446, thus we accept H_0 , the type of feedingstuffs does not influence milk production

The estimated treatment and period effects under fixed effects of mixed effects analysis are identical to those in fixed effects analysis.

```
c)
```

```
attach(cow)
t.test(milk[treatment=="A"],milk[treatment=="B"],paired=TRUE)

##
## Paired t-test
##
## data: milk[treatment == "A"] and milk[treatment == "B"]
```

```
## t = 0.224, df = 8, p-value = 0.83
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -2.2679 2.7568
## sample estimates:
## mean of the differences
## 0.24444
```

The paired-sample t-test produces a invalid test for a difference in milk production, because repeated measures may not be exchangeable because of time effect: Cows may mature or get older and learning effect: Cows may be happy with or bored at feedingstuff.

The p-value for treatment is identical to the one of the previous "fixed effects" obtained in a) (the order of the treatments was ignored). They are compatible because in the case of two repeated measurements, t^2 value for paired t-test is identical to the F value of the repeated measures ANOVA.

Exercise 4

```
a)
nausea_raw <- read.delim("./nauseatable.txt", header = TRUE, sep = "")
med_vec <- c();nausea_vec <- c()

for(i in 1:3) { med_name <- rownames(nausea_raw)[i]
    for(j in 1:2) { nausea <- colnames(nausea_raw)[j]
        for(k in 1:nausea_raw[,j][i]) { med_vec <- append(med_vec, med_name)
            if(j == 1) { nausea_vec <- append(nausea_vec, "no")}
            else if(j == 2){ nausea_vec <- append(nausea_vec, "yes")}
    }
}
nausea_frame <- data.frame(medicin = med_vec, naus = nausea_vec)
xtabs(~nausea_frame$medicin + nausea_frame$naus)</pre>
```

nausea_frame\$naus
nausea_frame\$medicin no yes
Chlorpromazine 100 52
Pentobarbital(100mg) 32 35
Pentobarbital(150mg) 48 37

b)

 H_0 : There is no difference between the treatment with different medicines (populations are the same).

The p-value resulting from the permutation test $0.056 > \alpha$ meaning that we fail to reject H_0 and thus could mean that the two medicines perform equally well to treat nausea. However, it must be said that in a permutation test, the p-value can change depending on the randomly generated samples meaning that a p-value which is just above the significance level, might be lower than the significance level on a different run.

c)

```
H_0: There is no difference between the treatment with different medicines (populations are the same).
```

```
p_val_chi <- chisq.test(xtabs(~nausea_frame$medicin + nausea_frame$naus))[[3]]</pre>
```

The p-value for the χ^2 -test for contingency tables is 0.036 which is lower than that computed in part b) and is, in fact less than true significance value α meaning that we can safely reject H_0 suggesting that the different medicines are not equally as good in treating nausea in patients. This could be due to the χ^2 T-values sampled in the permutation test being slightly different from the true χ -distribution.

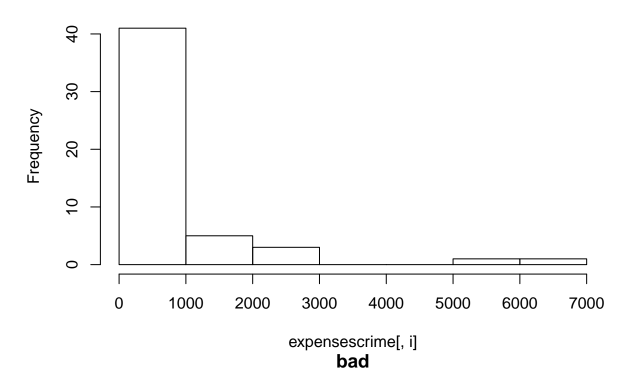
Exercise 5

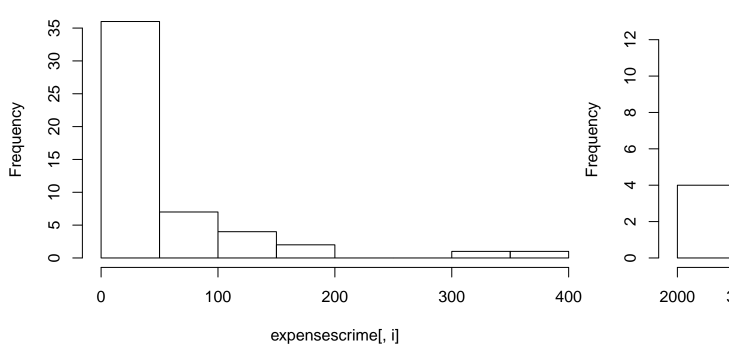
The data in expensescrime.txt were obtained to determine factors related to state expenditures on criminal activities (courts, police, etc.) The variables are: state (indicating the state in the USA), expend (state expenditures on criminal activities in \$1000), bad (crime rate per 100000), crime (number of persons under criminal supervision), lawyers (number of lawyers in the state), employ (number of persons employed in the state) and pop (population of the state in 1000). In the regression analysis, take expend as response variable and bad, crime, lawyers, employ and pop as explanatory variables.

a) Make some graphical summaries of the data. Investigate the problem of potential and influence points, and the problem of collinearity.

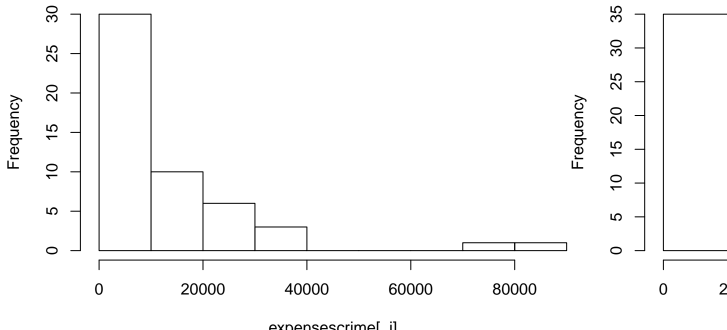
```
expensescrime=read.table(file="expensescrime.txt",header=TRUE)
for (i in c(2,3,4,5,6,7)) hist(expensescrime[,i],main=names(expensescrime[i]))
```

expend

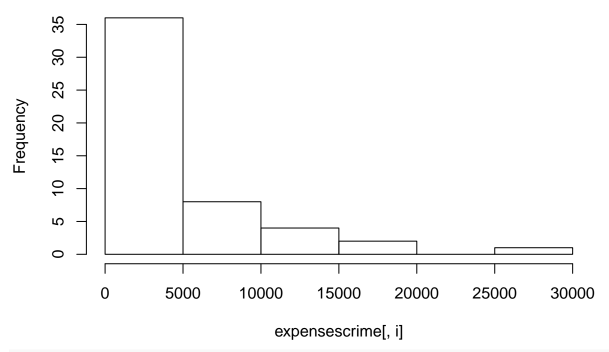




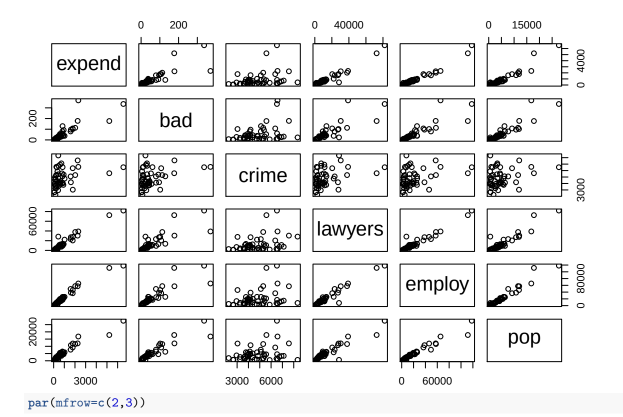




 $\begin{array}{c} \text{expensescrime[, i]} \\ \textbf{pop} \end{array}$



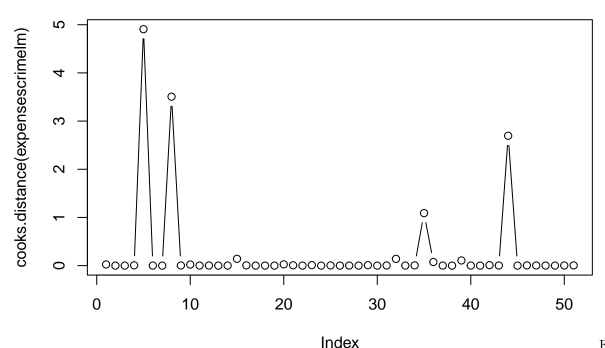
plot(expensescrime[,c(2,3,4,5,6,7)])



From the plot we can find potential points for each variable. For bad:336.2(CA) & 370.1(TX); For lawyers:82001(CA)&72575(NY); For employ: 118149(CA) & 111518(NY); For pop: 27663(CA).

```
expensescrimelm=lm(expend~bad+crime+lawyers+employ+pop,data=expensescrime)
expendbadlm=lm(expend~bad,data=expensescrime)
expendcrimelm=lm(expend~crime,data=expensescrime)
expendlawyerslm=lm(expend~lawyers,data=expensescrime)
expendemploylm=lm(expend~employ,data=expensescrime)
expendpoplm=lm(expend~pop,data=expensescrime)
round(cooks.distance(expensescrimelm),2)
##
               3
                        5
                             6
                                 7
                                      8
                                           9
                                               10
                                                    11
                                                        12
                                                             13
                                                                  14
                                                                      15
                                                                           16
## 0.02 0.00 0.00 0.01 4.91 0.00 0.00 3.51 0.00 0.02 0.00 0.00 0.00 0.00 0.14 0.01
##
    17
         18
              19
                  20
                       21
                            22
                                 23
                                     24
                                          25
                                               26
                                                    27
                                                        28
                                                             29
                                                                  30
                                                                      31
37
                                 39
                                     40
##
         34
              35
                  36
                            38
                                          41
                                               42
                                                    43
                                                        44
                                                             45
                                                                  46
                                                                       47
## 0.00 0.00 1.09 0.07 0.00 0.00 0.11 0.00 0.00 0.01 0.00 2.70 0.00 0.00 0.00 0.00
##
    49
         50
              51
## 0.00 0.00 0.00
plot(cooks.distance(expensescrimelm), type="b", main="Cook's distances")
```

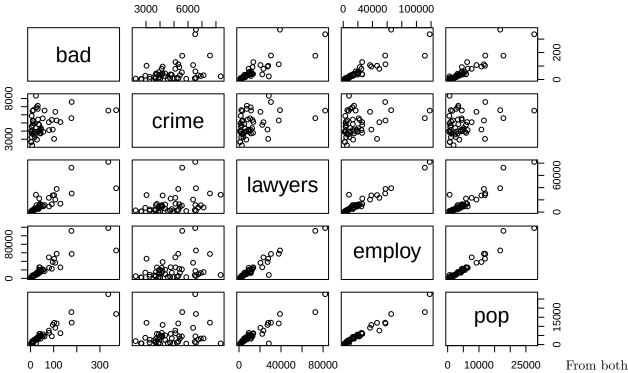
Cook's distances



fluence points, we check the Cook's distance, from the plot we can see that there are four. No.5(CA) & No.8(DC) & No.35(NY) & No.44(TX).

```
library(car)
```

```
## Loading required package: carData
## Registered S3 methods overwritten by 'car':
##
     method
                                      from
##
     influence.merMod
                                      lme4
##
     cooks.distance.influence.merMod lme4
     dfbeta.influence.merMod
##
                                      lme4
     dfbetas.influence.merMod
                                      lme4
vif(expensescrimelm)
##
       bad
             crime lawyers
                             employ
                                        pop
##
      8.36
                     16.97
                              33.59
              1.49
                                      32.94
round(cor(expensescrime[,c(3,4,5,6,7)]),2)
##
            bad crime lawyers employ pop
## bad
           1.00
                 0.37
                          0.83
                                 0.87 0.92
## crime
           0.37
                 1.00
                          0.38
                                 0.31 0.28
## lawyers 0.83
                 0.38
                          1.00
                                 0.97 0.93
## employ 0.87
                 0.31
                          0.97
                                 1.00 0.97
           0.92
                 0.28
                          0.93
                                 0.97 1.00
pairs(expensescrime[,c(3,4,5,6,7)])
```



the scatter plot and pairwise linear correlation and the vif, bad and pop (correlation value=0.92), lawyers and employ (correlation value=0.97), lawyers and pop (correlation value=0.93), employ and pop (correlation value=0.97), so crime and lawyers and employ and pop are collinear.

b) Fit a linear regression model to the data. Use both the step-up and the step-down method to find the best model. If step-up and step-down yield two different models, choose one and motivate your choice.

First, in the step-up method, we starts with fitting all p possible simple linear regression models: $Y_n = \beta_0 + \beta 1 X_{nj} + e_n$ To save pages, Only parts of summary are shown.

```
summary(lm(expend~bad,data=expensescrime))[8]

## $r.squared
## [1] 0.696

summary(lm(expend~crime,data=expensescrime))[8]

## $r.squared
## [1] 0.112

summary(lm(expend~lawyers,data=expensescrime))[8]

## $r.squared
## [1] 0.937

summary(lm(expend~employ,data=expensescrime))[8]

## $r.squared
## [1] 0.954

summary(lm(expend~pop,data=expensescrime))[8]
```

\$r.squared

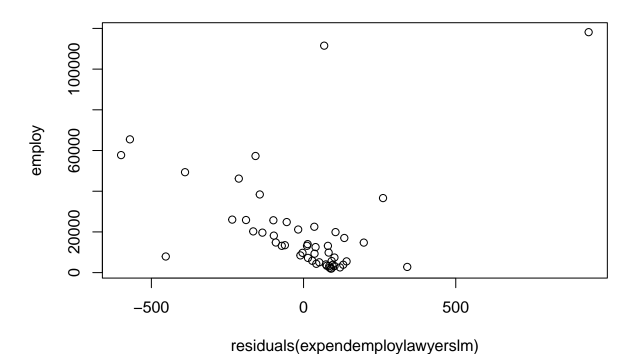
```
## [1] 0.907
The employ yields the highest R^2 increase.
summary(lm(expend~employ+bad,data=expensescrime))[8]
## $r.squared
## [1] 0.955
summary(lm(expend~employ+crime,data=expensescrime))[8]
## $r.squared
## [1] 0.955
summary(lm(expend~employ+lawyers,data=expensescrime))[8]
## $r.squared
## [1] 0.963
summary(lm(expend~employ+pop,data=expensescrime))[8]
## $r.squared
## [1] 0.954
Adding bad or crime or pop yields insignificant explanatory varibles. Therefore stop. The resulting model of
the step-up method is expend = -110.7 + 0.02971 * employ + 0.02686 * lawyers.
summary(lm(expend~bad+crime+lawyers+employ+pop,data=expensescrime))[8]
## $r.squared
## [1] 0.968
summary(lm(expend~bad+lawyers+employ+pop,data=expensescrime))[8]
## $r.squared
## [1] 0.967
summary(lm(expend~bad+lawyers+employ,data=expensescrime))[8]
## $r.squared
## [1] 0.964
summary(lm(expend~lawyers+employ,data=expensescrime))[8]
## $r.squared
## [1] 0.963
In the step-down method, we remove variables whose p-value is larger than 0.05, we stop after remove bad
because remaining variables are significant. the model is the same as the model of step-up model.
```

c) Check the model assumptions by using relevant diagnostic tools.

The model is expend = -110.7 + 0.02971 * employ + 0.02686 * lawyers.

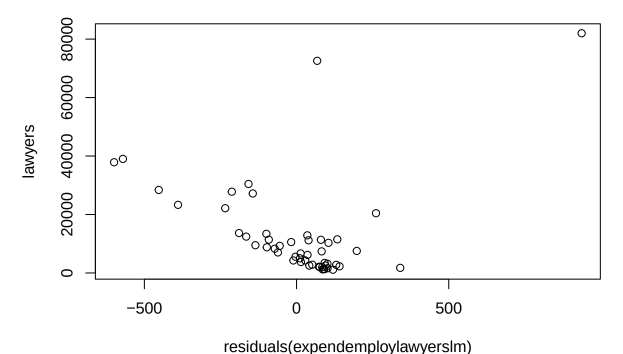
```
employ=expensescrime$employ
expend=expensescrime$expend
lawyers=expensescrime$lawyers
expendemploylawyerslm=lm(expend~employ+lawyers)
plot(residuals(expendemploylawyerslm),employ,main="plot of residuals against employ")
```

plot of residuals against employ



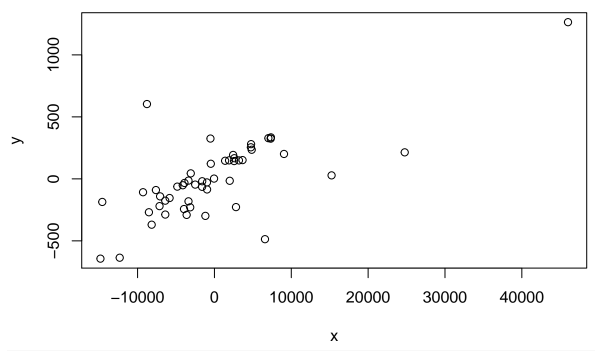
plot(residuals(expendemploylawyerslm),lawyers,main="plot of residuals against lawyers")

plot of residuals against lawyers



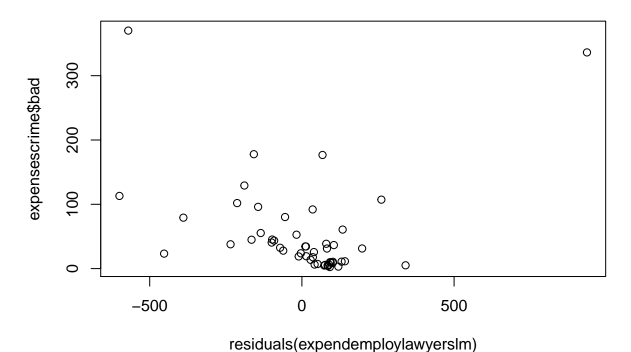
x=residuals(lm(employ+lawyers~expensescrime\$bad+expensescrime\$crime+expensescrime\$pop))
y=residuals(lm(expend~expensescrime\$bad+expensescrime\$crime+expensescrime\$pop))
plot(x,y,main="Added variable plot for employ+lawyers")

Added variable plot for employ+lawyers



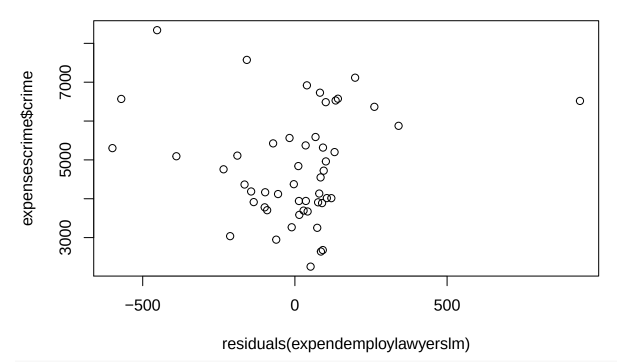
plot(residuals(expendemploylawyerslm),expensescrime\$bad,main="plot of residuals against bad")

plot of residuals against bad



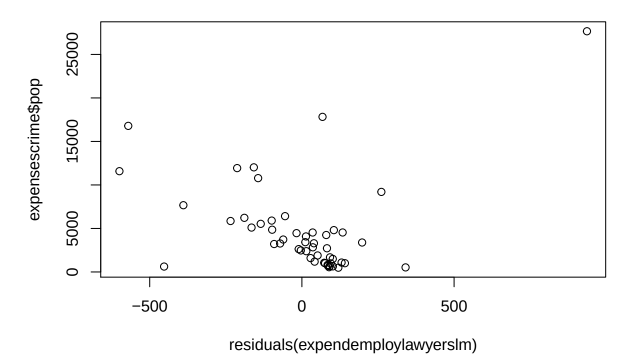
plot(residuals(expendemploylawyerslm),expensescrime\$crime,main="plot of residuals against crime")

plot of residuals against crime



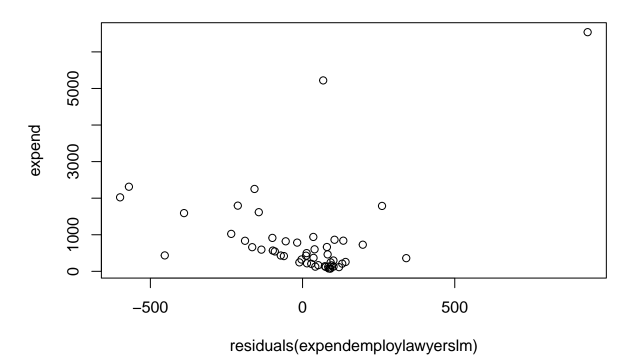
plot(residuals(expendemploylawyerslm),expensescrime\$pop,main="plot of residuals against pop")

plot of residuals against pop

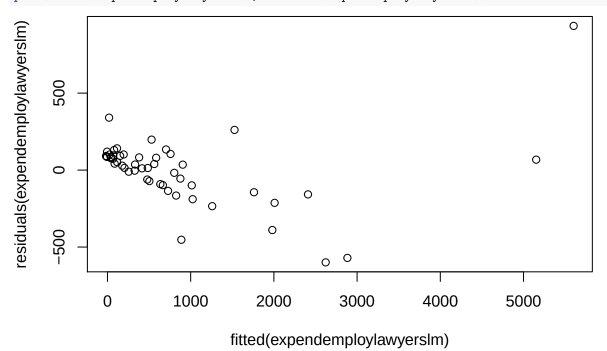


plot(residuals(expendemploylawyerslm),expend,main="plot of residuals against expend")

plot of residuals against expend

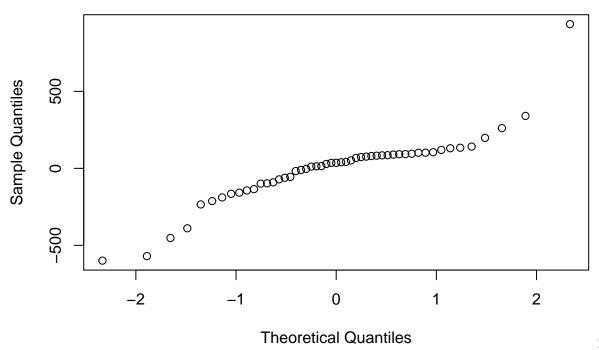


plot(fitted(expendemploylawyerslm),residuals(expendemploylawyerslm))



qqnorm(residuals(expendemploylawyerslm))

Normal Q-Q Plot



the above plots we can see that spread of the residuals against employ and the residuals against lawyers are alike. The normal Q-Q plot of residuals doesn't show normal distribution. JHowever, plot residuals against Y shows some pattern of decrease. From a), we already know that there is a problem of collinearity between lawyers and employ. Compare with other models, expend~employ has less variables and only a slightly lower value of R-squared. The model is expend = -116.7 + 0.046811 * employ.

```
vif(expendemploylawyerslm)
##
    employ lawyers
      14.8
summary(lm(expend~lawyers,data=expensescrime))[8]
## $r.squared
## [1] 0.937
summary(lm(expend~employ,data=expensescrime))
##
## Call:
  lm(formula = expend ~ employ, data = expensescrime)
##
##
  Residuals:
##
      Min
              1Q Median
                            3Q
                                  Max
##
  -636.0 -84.3
                   47.6
                         108.0 1124.7
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -1.17e+02
                           4.71e+01
                                       -2.48
                                                0.017 *
##
  employ
                4.68e-02
                           1.47e-03
                                       31.87
                                               <2e-16 ***
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
##
```

Residual standard error: 257 on 49 degrees of freedom
Multiple R-squared: 0.954, Adjusted R-squared: 0.953
F-statistic: 1.02e+03 on 1 and 49 DF, p-value: <2e-16</pre>