# Assignment 2: Group 45

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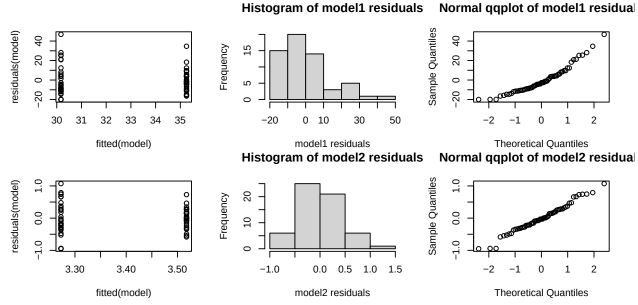
03 March 2023

Note: we made a function checkNorm() which prints a histogram, qqplot, and p-value from the shapiro-wilk normality test. And we made a function printPval() which simply prints a given p-value to 3 significant figures. We utilize both functions throughout this assignment.

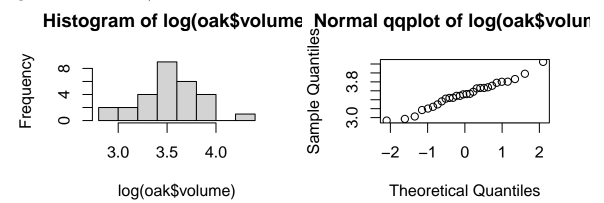
#### Exercise 1: Trees

1 a)

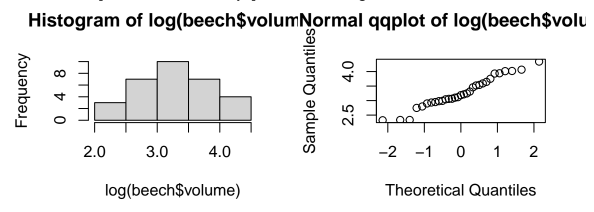
```
trees = read.table("treeVolume.txt", header=T)
trees$type = factor(trees$type)
par(mfrow=c(1,3))
model1 = lm(volume~type, data=trees)
pval = checkAnovaNorm(model1, "model1")
## [1] "Shapiro-Wilk normality p-value for model1 residuals: 0.001"
model2 = lm(log(volume)~type, data=trees)
pval = checkAnovaNorm(model2, "model2")
## [1] "Shapiro-Wilk normality p-value for model2 residuals: 0.551"
print("model coefficients:"); summary(model2)$coefficients
## [1] "model coefficients:"
##
               Estimate Std. Error t value Pr(>|t|)
                  3.273
                                     41.84 1.75e-44
## (Intercept)
                            0.0782
## typeoak
                  0.245
                            0.1135
                                      2.16 3.52e-02
res = anova(model2)
sprintf("ANOVA p-value for type = %.3f", res["type", "Pr(>F)"])
## [1] "ANOVA p-value for type = 0.035"
```



We tested two models, and looked at the residuals of each for normality. The results graphs suggests that using log(volume) in model2 is preferred as the resulting residuals are normally distributed (whereas this wasn't the case in model1). Additionally, the ANOVA analysis of model2 suggests that we can reject the  $H_0$  that type is not a significant predictor of (the log of) volume (as the p-value 0.035 < 0.05).



## [1] "Shapiro-Wilk normality p-value for log(oak\$volume): 0.902"



## [1] "Shapiro-Wilk normality p-value for log(beech\$volume): 0.377"

```
## [1] "oak mean volume = 35.250, beech mean volume = 30.171"
```

We can split the data into two samples of tree volume based on the tree types. Checking normality, we see that the log transformed volumes of each tree type are normal (whereas the volumes themselves aren't as in 1a).

We can indeed compare the means of the transformed samples using a t-test to determine whether, based on this data, there is a significant difference in mean volume between the two tree types.

```
new_oak = data.frame(type="oak"); new_beech = data.frame(type = "beech")
pred1 = exp(predict(model2, new_oak)); pred2 = exp(predict(model2, new_beech))
sprintf("predicted volumes: oak = %.3f, beech = %.3f", pred1, pred2)
```

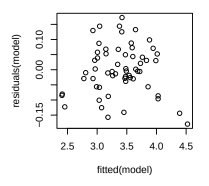
# ## [1] "predicted volumes: oak = 33.706, beech = 26.383"

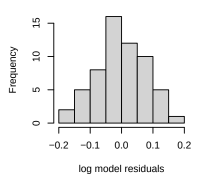
Above we use model2 to predict the log volume for each tree type, and use exp() to convert it to actual volume prediction.

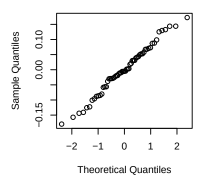
#### 1 b)

```
par(mfrow = c(1, 3))
model = lm(log(volume)~type*diameter+height, data=trees)
pval = checkAnovaNorm(model, "log model")
```

## Histogram of log model residual: Normal applot of log model residu







## [1] "Shapiro-Wilk normality p-value for log model residuals: 0.788"

```
res = drop1(model, test="F")
sprintf("drop1 p-value for type:diameter = %.3f", res["type:diameter", "Pr(>F)"])
```

#### ## [1] "drop1 p-value for type:diameter = 0.212"

We built a linear model that added an interaction term between diameter and type, the p-value 0.212 > 0.05 for this term suggests there's insufficient evidence to reject the  $H_0$  (that the influence of diameter on volume is the same for both tree types).

```
model = lm(volume~type*height+diameter, data=trees)
res = drop1(model, test="F")
sprintf("drop1 p-value for type:diameter = %.3f", res["type:height", "Pr(>F)"])
```

## [1] "drop1 p-value for type:diameter = 0.176"

Now running another linear model that includes an interaction term between height and type instead, the p-value 0.176 > 0.05 for this term suggests there's insufficient evidence to reject the  $H_0$  (that the influence of height on volume is the same for both tree types).

So based on the results from our two models above, there's insufficient evidence to suggest that the influences of diameter and height aren't similar for both tree types.

#### 1 c)

We construct a linear model to investigate how diameter, height and type influence volume.

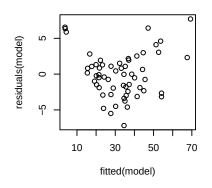
```
model = lm(volume~diameter+height+type, data=trees)
par(mfrow = c(1, 3)); pval = checkAnovaNorm(model, "model")
## [1] "Shapiro-Wilk normality p-value for model residuals: 0.524"
print("model coefficients:"); print(summary(model)$coefficients)
## [1] "model coefficients:"
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                -63.781
                             5.5129
                                     -11.57 2.33e-16
                  4.698
## diameter
                             0.1645
                                      28.56 1.14e-34
## height
                  0.417
                             0.0752
                                       5.55 8.42e-07
## typeoak
                                      -1.49 1.43e-01
                 -1.305
                             0.8779
drop1(model, test="F")
## Single term deletions
##
## Model:
## volume ~ diameter + height + type
            Df Sum of Sq RSS AIC F value Pr(>F)
##
## <none>
                           578 143
## diameter
                    8577 9155 304
                                    815.61 < 2e-16 ***
## height
             1
                     324
                           903 167
                                     30.82 8.4e-07 ***
```

# 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 Histogram of model residuals Normal qq

0.14

2.21

#### Normal applot of model residual:



1

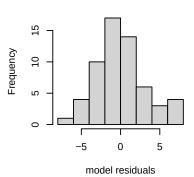
## type

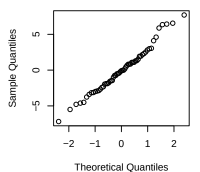
## Signif. codes:

## ---

23

602 143



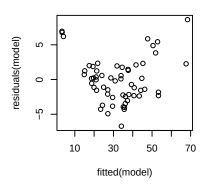


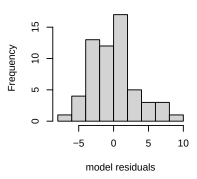
We built a linear model (where the normality assumptions hold). Based on the drop1 p-values, type is not a significant predictor for volume (p-value 0.14 > 0.05), while height and diameter are significant (p-values less than 0.05). Diameter and height are both positively correlated with the volume, with diameter having the largest contribution (coefficient) of the two.

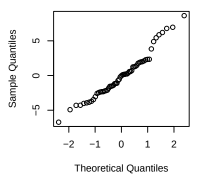
```
# build better model where type isn't considered
modelC = lm(volume~height+diameter, data=trees)
par(mfrow = c(1, 3)); pval = checkAnovaNorm(modelC, "model")
```

#### Histogram of model residuals

#### Normal applot of model residuals







## [1] "Shapiro-Wilk normality p-value for model residuals: 0.089"

```
avgTree = data.frame(height=mean(trees$height), diameter=mean(trees$diameter))
pred = predict(modelC, avgTree)
sprintf("predicted volume of average tree = %.3f", pred)
```

## [1] "predicted volume of average tree = 32.581"

```
#mean(trees$volume) # this also gives the same result as expected

r2 = summary(modelC)$r.squared; ar2 = summary(modelC)$adj.r.squared
sprintf("modelC: R^2 = %.3f, Adj. R^2 = %.3f", r2, ar2)
```

```
## [1] "modelC: R^2 = 0.949, Adj. R^2 = 0.947"
```

We omitted type from our model as it was shown above to be insignificant, we also observe that the normality assumptions hold. Using the resulting model, the volume of a tree with the average height and diameter is predicted to be 32.581.

## 1 d)

We propose to transform the data to create a new column that contains the volume of a (theoretical) cylinder based on the tree's diameter and height. (Note we omit tree type from the model as we found it to not be a significant predictor above).

```
# create predictor as cylinderical volume
trees$cylinder = pi * (trees$diameter / 2)^2 * trees$height
modelD = lm(volume~cylinder, data=trees)
print("model coefficients:"); summary(modelD)$coefficients
```

```
## [1] "model coefficients:"

## Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.37942  7.63e-01 -0.497 6.21e-01
## cylinder  0.00273  5.82e-05  46.913 3.09e-47

r2 = summary(modelD)$r.squared; ar2 = summary(modelD)$adj.r.squared
sprintf("model: R^2 = %.3f, Adj. R^2 = %.3f", r2, ar2)
```

```
## [1] "model: R^2 = 0.975, Adj. R^2 = 0.974"
```

After constructing a linear model for predicting the actual tree volume from our proposed cylindrical estimator, we see that the cylinder variable is a significant predictor of volume (p < 0.05). The adjusted  $R^2$  values (and the regular  $R^2$  values) for this model are both greater than that of the model in part c), so this model appears to be superior to using just the provided height and diameter variables in the model.

# Exercise 2: Expenditure on criminal activities

2 a)

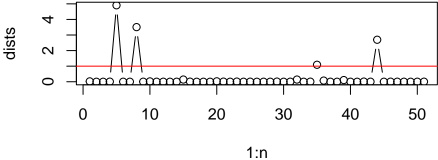
##

```
crimes = read.table("expensescrime.txt", header=T)
pairs(crimes[,-1])
                   200
                                          40000
                                                                  15000
    expend
                  bad
                             crime
00009
                                        lawvers
                                                    employ
                                                                  pop
   0
      3000
                           3000 6000
                                                   0
                                                      60000
crimes$state = factor(crimes$state)
model = lm(expend~crime+bad+lawyers+employ+pop, data=crimes)
summary(model)$coefficients; anova(model)
```

Estimate Std. Error t value Pr(>|t|)

```
## (Intercept) -299.1341
                          1.40e+02
                                     -2.14 0.03817
## crime
                 0.0324
                          2.81e-02
                                      1.15 0.25534
## bad
                -2.8319
                          1.24e+00
                                     -2.28 0.02719
## lawyers
                 0.0232
                          8.04e-03
                                      2.89 0.00592
## employ
                 0.0230
                          7.46e-03
                                      3.08 0.00354
                 0.0779
                          3.51e-02
                                      2.22 0.03184
## pop
## Analysis of Variance Table
## Response: expend
##
            Df
                 Sum Sq Mean Sq F value Pr(>F)
## crime
               7888219 7888219 155.03 3.5e-16 ***
## bad
             1 41265535 41265535 811.00 < 2e-16 ***
             1 17237521 17237521 338.77 < 2e-16 ***
## lawyers
## employ
             1
               1590235 1590235
                                   31.25 1.3e-06 ***
                                    4.91
                                           0.032 *
## pop
             1
                 249704
                          249704
## Residuals 45 2289716
                           50883
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
n = length(crimes[,1])
dists = cooks.distance(model)
plot(1:n, dists, type="b", main="Cook's Distance by Dataset Index")
abline(1, 0, col = 'red') # plot y=1 for reference
```

# **Cook's Distance by Dataset Index**



```
# cook's points:
print("Influence points:"); crimes[dists > 1,]
## [1] "Influence points:"
##
      state expend
                     bad crime lawyers employ
                                                 pop
## 5
              6539 336.2 6518
                                 82001 118149 27663
         CA
## 8
         DC
               435 23.3 8339
                                 28399
                                         7925
                                                 622
## 35
              5220 176.7 5589
         NY
                                 72575 111518 17825
## 44
         TX
              2313 370.1
                          6569
                                 39028 65488 16789
# investigating collinearity:
cor(crimes[,-1])
```

```
##
           expend
                    bad crime lawyers employ
## expend
          1.000 0.834 0.334
                                0.968
                                       0.977 0.953
## bad
            0.834 1.000 0.373
                                0.832 0.871 0.920
## crime
            0.334 0.373 1.000
                                       0.311 0.275
                                0.375
## lawyers 0.968 0.832 0.375 1.000
                                       0.966 0.934
## employ
                                       1.000 0.971
            0.977 0.871 0.311
                                0.966
## pop
            0.953 0.920 0.275 0.934 0.971 1.000
# using 0.8 as a threshold to help with visiblility:
res = cor(crimes[,-1])
res[res >= 0.8] = T; res[res <= 0.8] = F; res
           expend bad crime lawyers employ pop
## expend
                1
                    1
                          0
                                  1
## bad
                1
                    1
                          0
                                  1
                                          1
                                              1
## crime
                0
                    0
                          1
                                  0
                                         0
                                              0
## lawvers
                1
                    1
                          0
                                  1
                                         1
                                              1
## employ
                    1
                          0
                                  1
                                         1
                                              1
                                  1
## pop
                          0
```

Based on the correlation coefficients, it appears that all the explanatory variables are correlated with each other, except for crime which has no correlation with any of the other variables (its highest correlation coefficient is 0.375). The other variables all have a correlation coefficient of at least 0.832 between each other.

# 2 b)

```
evalModel = function(model, name) {
 r2 = summary(model)$r.squared; ar2 = summary(model)$adj.r.squared
 pVal = summary(model)$coefficients[name, "Pr(>|t|)"]
  cat(sprintf("trying var '%s'\t\tPr(>|t|) = \%.3f, model R^2 = \%.3f\n", name, pVal, r2))
}
doStepUp = function() {
  cat("\n****round1 (building model with 1 var)****\n")
  evalModel(lm(expend~bad, data=crimes), name="bad")
  evalModel(lm(expend~crime, data=crimes), name="crime")
  evalModel(lm(expend~lawyers, data=crimes), name="lawyers")
  evalModel(lm(expend~employ, data=crimes), name="employ")
  evalModel(lm(expend~pop, data=crimes), name="pop")
  # employ has highest adj. R^2 (0.954) and is significant
  cat("\n***round2 (building on model with employ)***\n")
  evalModel(lm(expend~employ+bad, data=crimes), name="bad")
  evalModel(lm(expend~employ+crime, data=crimes), name="crime")
  evalModel(lm(expend~employ+lawyers, data=crimes), name="lawyers")
  evalModel(lm(expend~employ+pop, data=crimes), name="pop")
  cat("\n***round3 (building on model with employ+laywers)****\n")
```

```
evalModel(lm(expend~employ+lawyers+bad, data=crimes), name="bad")
evalModel(lm(expend~employ+lawyers+crime, data=crimes), name="crime")
evalModel(lm(expend~employ+lawyers+pop, data=crimes), name="pop")
}
doStepUp()
```

```
##
## ****round1 (building model with 1 var)****
## trying var 'bad'
                        Pr(>|t|) = 0.000, model R^2 = 0.696
## trying var 'crime'
                            Pr(>|t|) = 0.016, model R^2 = 0.112
                            Pr(>|t|) = 0.000, model R^2 = 0.937
## trying var 'lawyers'
                            Pr(>|t|) = 0.000, model R^2 = 0.954
## trying var 'employ'
## trying var 'pop'
                        Pr(>|t|) = 0.000, model R^2 = 0.907
##
## ****round2 (building on model with employ)****
## trying var 'bad'
                        Pr(>|t|) = 0.279, model R^2 = 0.955
## trying var 'crime'
                            Pr(>|t|) = 0.289, model R^2 = 0.955
                            Pr(>|t|) = 0.001, model R^2 = 0.963
## trying var 'lawyers'
## trying var 'pop'
                        Pr(>|t|) = 0.555, model R<sup>2</sup> = 0.954
##
## ****round3 (building on model with employ+laywers)****
## trying var 'bad'
                        Pr(>|t|) = 0.345, model R^2 = 0.964
                            Pr(>|t|) = 0.896, model R^2 = 0.963
## trying var 'crime'
## trying var 'pop'
                        Pr(>|t|) = 0.399, model R^2 = 0.964
```

In the 1st round of the "step up" method we found "employ" to lead to the model with the largest  $R^2$ , while still being statistically significant. In the 2nd round (building up from the model with "employ"), "lawyers" was found to lead to the largest increase in  $R^2$  while still being statistically significant (p < 0.05), so we add it to the model. The 3rd round of the "step up" method showed that no further variables could be added while being statistically significant. So our final model is lm(expend~employ+lawyers, data=crimes).

#### 2 c)

Taking the resulting model from part 2b, we found the normality assumptions weren't met, so we adjusted the model by taking the square root of expend. The residuals then indicated normality assumptions were met, however the p-value for lawyers was no longer significant.

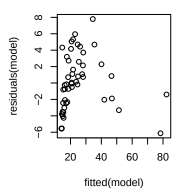
So we tried the best model from round 1 of step up which was lm(expend~employ, data=crimes) (noting that this model was comporable as adding laywers had only increased the  $R^2$  by 0.963 - 0.954 = 0.009). Finally, after checking the normality assumptions of this model, we again saw they weren't met, and so settled on the final model of lm(sqrt(expend)~employ, data=crimes) for which the assumptions are now met (the qqplot of residuals below is approximately linear now).

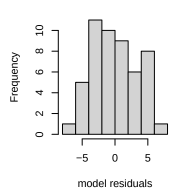
```
# this model meets normality assumptions, but p-value for lawyers is 0.44 > 0.05
#model = lm(sqrt(expend)~employ+lawyers, data=crimes)

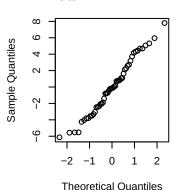
model = lm(sqrt(expend)~employ, data=crimes)
summary(model)$coefficients
```

```
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1.30e+01 6.27e-01 20.8 6.27e-26
## employ 5.86e-04 1.96e-05 29.9 3.73e-33
par(mfrow = c(1, 3)); pval = checkAnovaNorm(model, "model")
```

#### Histogram of model residua Normal applot of model residu







## [1] "Shapiro-Wilk normality p-value for model residuals: 0.485"

```
state = data.frame(bad=50, crime=5000, lawyers=5000, employ=5000, pop=5000)
pred = predict(model, state, interval="prediction")^2
sprintf("predicted expenditure CI ="); pred
```

```
## [1] "predicted expenditure CI ="
## fit lwr upr
## 1 254 80.3 526
```

The predicted interval [80.3, 526] can't be further improved in this case, as both numbers are > 0 and the process of finding a solid model already improved the accuracy of this CI. Arguably, you could consider removing some influence points to further "improve" this CI (but this is risky).

#### 2 d)

# library(glmnet)

```
par(mfrow=c(1,2))
set.seed(42) # ensuring results don't change each time its run
x = as.matrix(crimes[,-1]) # remove states column
x = x[,-1] # remove expenditure
y = crimes[,2]

train = sample(1:nrow(x),0.67*nrow(x))
x.train = x[train,]; y.train = y[train]
x.test = x[-train,]; y.test = y[-train]

lm.model = lm(expend~bad+crime+lawyers+employ+pop,data=crimes, subset=train)
y.predict.lm = predict(lm.model,newdata=crimes[-train,])
```

```
mse.lm = mean((y.test-y.predict.lm)^2)
sprintf("linear model mse = %.3f", mse.lm)
## [1] "linear model mse = 54935.728"
lasso.model = glmnet(x.train,y.train,alpha=1)
lasso.cv = cv.glmnet(x.train,y.train,alpha=1,type.measure="mse",nfolds=5)
plot(lasso.model,label=T,xvar="lambda"); plot(lasso.cv)
                                    Mean-Squared Error
         5
               5
                    3
                          3
                                                     5
                                                        3
                                                            3
                                          2000000
Coefficients
     -3.0
         0
               2
                    4
                          6
                                              0
                                                    2
                                                         4
                                                               6
                                                     Log(\lambda)
              Log Lambda
lambda.min = lasso.cv$lambda.min; lambda.1se=lasso.cv$lambda.1se;
sprintf("lambda.min = %.3f, lambda.1se = %.3f", lambda.min, lambda.1se)
## [1] "lambda.min = 29.891, lambda.1se = 305.942"
# lasso min (minimizing R^2)
coef(lasso.model,s=lasso.cv$lambda.min)
## 6 x 1 sparse Matrix of class "dgCMatrix"
##
## (Intercept) -132.2017
## bad
## crime
## lawyers
                   0.0261
## employ
                   0.0132
                   0.0703
## pop
# 1 SE (minimizing number of vars, while remaining within 1 std. err.)
coef(lasso.model,s=lasso.cv$lambda.1se)
## 6 x 1 sparse Matrix of class "dgCMatrix"
## (Intercept) 130.4846
## bad
## crime
## lawyers
                  0.0171
## employ
                  0.0132
## pop
                  0.0449
```

```
lasso.pred1 = predict(lasso.model,s=lambda.min,newx=x.test)
lasso.pred2 = predict(lasso.model,s=lambda.1se,newx=as.matrix(x.test))
mse1.lasso = mean((y.test-lasso.pred1)^2)
mse2.lasso = mean((y.test-lasso.pred2)^2)
sprintf("mse1 = %.3f, mse2 = %.3f", mse1.lasso, mse2.lasso)
```

```
## [1] "mse1 = 45429.278, mse2 = 169706.874"
```

As we can see from the lambdas calculated by the model above, for the minimum error, the relevant variables are laywers, employ, and crime. To obtain a reduced model that is within one standard error of the minimum, we can take into account only these relevant variables.

Considering the scale differences of the relevant variables, the coefficient of pop is not only the largest, but it also has the largest contribution for predictions (as the pop variable will always be larger than employ and lawyer for any given state by a considerable margin).

As a disclaimer, the results from the lasso method will always vary with random chance, however this interpretation holds for our resulting model.

The coefficients of the other two relevant variables (lawyers and pop) are much smaller due to their collinearity with employ.

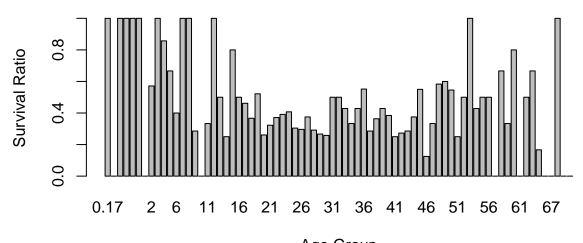
#### Exercise 3: Titanic

3 a)

```
titanic = read.table("titanic.txt", header = T)
titanic$PClass = factor(titanic$PClass)
titanic$Sex = factor(titanic$Sex)
# titanic$Survived = factor(titanic$Survived)
library(plyr)
# round age to nearest 10 years
# (https://stackoverflow.com/a/6466894)
titanic$Ager = round_any(titanic$Age, 20)
titanic$Ager = factor(titanic$Ager)
titanic$Age2 = titanic$Age^2
print("total number of individuals for each combination of class and gender:")
## [1] "total number of individuals for each combination of class and gender:"
tot = xtabs(~PClass + Sex, data = titanic)
tot
##
         Sex
## PClass female male
             143 179
##
      1st
##
      2nd
             107 173
##
      3rd
             212 499
```

```
print("Survival rate by class and gender combinations:")
## [1] "Survival rate by class and gender combinations:"
totc = xtabs(Survived ~ PClass + Sex, data = titanic)
round(totc/tot, 2)
##
         Sex
## PClass female male
            0.94 0.33
##
      1st
##
      2nd
            0.88 0.14
##
            0.38 0.12
      3rd
# plot(titanic$Age, y=titanic$Survived, xlim=c(0, 100))
totage = xtabs(~Age, data = titanic)
barplot(xtabs(Survived ~ Age, data = titanic)/totage, main = "% Survival by Age Group",
    xlab = "Age Group", ylab = "Survival Ratio")
```

# % Survival by Age Group



# Age Group

```
model = glm(Survived ~ Age + Sex + PClass, data = titanic, family = binomial)
summary(model)
##
## Call:
## glm(formula = Survived ~ Age + Sex + PClass, family = binomial,
##
       data = titanic)
##
## Deviance Residuals:
      Min
               1Q Median
                               3Q
                                      Max
## -2.723 -0.707 -0.392
                            0.649
                                     2.529
##
## Coefficients:
```

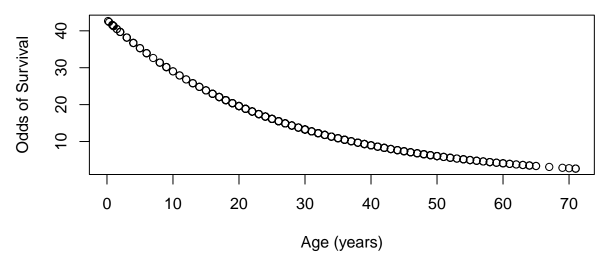
Estimate Std. Error z value Pr(>|z|)

# build logistic regression model

##

```
## (Intercept)
                            0.39757
                3.75966
                                       9.46
                                             < 2e-16 ***
## Age
               -0.03918
                            0.00762
                                      -5.14
                                             2.7e-07 ***
## Sexmale
               -2.63136
                            0.20151
                                     -13.06
                                             < 2e-16 ***
## PClass2nd
               -1.29196
                                      -4.97
                            0.26008
                                             6.8e-07 ***
## PClass3rd
               -2.52142
                            0.27666
                                      -9.11
                                             < 2e-16 ***
## Signif. codes:
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 1025.57
                                on 755
                                        degrees of freedom
## Residual deviance:
                       695.14
                                on 751
                                        degrees of freedom
     (557 observations deleted due to missingness)
##
## AIC: 705.1
##
## Number of Fisher Scoring iterations: 5
plot(titanic$Age, exp(3.759662 + titanic$Age * -0.039177), ylab = "Odds of Survival",
    xlab = "Age (years)", main = "Survival Odds of a First Class Female by Age")
```

# Survival Odds of a First Class Female by Age



In the model above, the intercept can be interpreted as a female who traveled in 1st class. We can see that other classes (namely 2nd and 3rd) lower the odds of survival by the negative coefficients, where 3rd class has a larger impact than 2nd. Similarly, an increase in age has a negative impact on survival odds, however the impact is low because of the parabolic shape of the graph shown above. How we interpret this is that the fact that very young and very old people have the highest survival rates are balancing each other out to form a coefficient that is close to 0, which inaccurately represents the actual relationship between age and survival, because it is clear that it does have an effect. Finally the odds of survival of a male are also significantly lower than that of a female, as shown by the negative coefficient.

```
3 b)
```

```
ageclass_model = glm(Survived~Age*PClass+Sex,data=titanic, family=binomial)
summary(ageclass_model)
##
## Call:
## glm(formula = Survived ~ Age * PClass + Sex, family = binomial,
       data = titanic)
##
## Deviance Residuals:
     Min
##
              1Q Median
                              3Q
                                     Max
## -2.721 -0.695 -0.399
                           0.630
                                   2.356
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                 3.7534
                             0.5405
                                      6.94 3.8e-12 ***
## Age
                 -0.0380
                             0.0117
                                      -3.23 0.0012 **
## PClass2nd
                 -0.5762
                             0.6637
                                      -0.87 0.3853
## PClass3rd
                                      -4.75 2.0e-06 ***
                 -3.0180
                             0.6351
## Sexmale
                             0.2058 -13.07 < 2e-16 ***
                 -2.6893
                             0.0187 -1.38 0.1663
## Age:PClass2nd -0.0258
## Age:PClass3rd
                  0.0211
                             0.0182
                                       1.16
                                              0.2450
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 1025.57 on 755
                                      degrees of freedom
## Residual deviance: 689.64 on 749 degrees of freedom
     (557 observations deleted due to missingness)
## AIC: 703.6
##
## Number of Fisher Scoring iterations: 5
agesex_model = glm(Survived~Age*Sex+PClass,data=titanic, family=binomial)
summary(agesex_model)
##
## Call:
## glm(formula = Survived ~ Age * Sex + PClass, family = binomial,
      data = titanic)
##
##
## Deviance Residuals:
##
     Min
              1Q Median
                              3Q
                                     Max
## -2.435 -0.656 -0.353
                          0.696
                                   2.728
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
```

```
6.30 3.0e-10 ***
## (Intercept) 2.75656
                          0.43764
                                              0.83
## Age
               0.00244
                          0.01141
                                    0.21
## Sexmale
             -0.50819
                          0.44251
                                   -1.15
                                              0.25
## PClass2nd -1.54337
                                   -5.37 7.8e-08 ***
                          0.28736
## PClass3rd -2.65398
                          0.29142 -9.11 < 2e-16 ***
## Age:Sexmale -0.07559
                                    -5.04 4.7e-07 ***
                          0.01501
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 1025.57 on 755 degrees of freedom
## Residual deviance: 667.08 on 750 degrees of freedom
     (557 observations deleted due to missingness)
## AIC: 679.1
##
## Number of Fisher Scoring iterations: 5
male1 = data.frame(Age=55, Sex = "male", PClass = "1st")
male2 = data.frame(Age=55, Sex = "male", PClass = "2nd")
male3 = data.frame(Age=55, Sex = "male", PClass = "3rd")
female1 = data.frame(Age=55, Sex = "female", PClass = "1st")
female2 = data.frame(Age=55, Sex = "female", PClass = "2nd")
female3 = data.frame(Age=55, Sex = "female", PClass = "3rd")
predict(model,female1, type="response")
##
      1
## 0.833
predict(model,female2, type="response")
##
       1
## 0.578
predict(model,female3, type="response")
##
       1
## 0.286
predict(model,male1, type="response")
##
## 0.264
predict(model,male2, type="response")
##
## 0.0896
predict(model,male3, type="response")
```

```
## 1
## 0.028
```

Given the results of the two models produced here, as well as the one from 3a), we will select the model that considers the interaction between sex and age, given that the model found that the interaction was significant (the model in 3a) found that sex and age by themselves were also significant). Furthermore, the model that considered the interaction between age and class found that the interaction was not significant (P > 0.05 for both 2nd class and 3rd class).

The estimates for the probability of survival of each of the combinations of factors is as shown below:

#### 3 c)

We propose to do k-fold cross-validation, using 10% of the data in each fold, and calculating the average prediction accuracy per combination of factor levels. We would use a slightly modified version of the model from 3b), namely glm(Survived~Ager\*Sex+PClass,data=titanic, family=binomial), where the difference would be that the age would be split into several groups (represented as the variable Ager.

To be able to predict for the odds resulting from the model, we would use a threshold of 1, where if the odds were above 1, the subject would be predicted as survived and not survived otherwise. (Because when the odds are  $\geq 1$ , there's a  $\geq 50\%$  chance of survival).

# 3 d)

Based on the (3x2) table in part 3a, every unique combination of classes in this dataset has > 5 individuals, so a chi-squared test is applicable.

To investigate the effect of the factors PClass and Sex separately, we'll perform two separate chisquare tests which each test the significance of a single factor.

```
# build contigency table
cont1 = as.matrix(xtabs(~PClass, data=titanic)); cont1
##
       [,1]
## 1st
        322
## 2nd
        280
## 3rd 711
chisq.test(cont1)
##
##
    Chi-squared test for given probabilities
##
## data: cont1
## X-squared = 258, df = 2, p-value <2e-16
cont2 = as.matrix(xtabs(~Sex, data=titanic)); cont2
##
          [,1]
## female
           462
## male
           851
```

#### chisq.test(cont2)

```
##
## Chi-squared test for given probabilities
##
## data: cont2
## X-squared = 115, df = 1, p-value <2e-16</pre>
```

Both chisquare tests yield p-values < 0.05 so we reject the null hypotheses that Sex and PClass aren't significantly explanatory variables for predicting survival.

# 3 e)

Comparing the linear regression approach vs the chisquare approach, the linear regression approach has the advantage of being able to make predictions on a given individual's chance of survival (which chisquare has no ability to do), and the linear regression approach can give p-values for each variable.

Both methods can also give insights into the significance of explanatory variables.

# Exercise 4: Military Coups

#### 4 a)

```
coups = read.table("coups.txt", header=T)
coups$pollib=factor(coups$pollib)
coupsglm = glm(miltcoup~oligarchy+pollib+parties+pctvote+popn+size+numelec+numregim, family=po
summary(coupsglm)
##
## Call:
## glm(formula = miltcoup ~ oligarchy + pollib + parties + pctvote +
      popn + size + numelec + numregim, family = poisson, data = coups)
##
##
## Deviance Residuals:
##
     Min
              1Q Median
                              3Q
                                     Max
## -1.508 -0.953 -0.310
                           0.486
                                   1.646
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -0.233427
                          0.997611
                                     -0.23
                                             0.8150
## oligarchy
              0.072566
                          0.035346 2.05
                                             0.0401 *
## pollib1
                          0.655811
              -1.103244
                                     -1.68
                                             0.0925
## pollib2
              -1.690306
                          0.676650 - 2.50
                                             0.0125 *
## parties
               0.031221
                          0.011166
                                      2.80
                                             0.0052 **
## pctvote
               0.015441
                          0.010103
                                      1.53
                                             0.1264
## popn
              0.010959
                          0.007149
                                      1.53
                                             0.1253
## size
                                     -0.99
              -0.000265
                          0.000269
                                             0.3244
```

```
## numelec
              -0.029619
                          0.069625
                                             0.6705
                                     -0.43
## numregim
               0.210943
                          0.233933
                                      0.90
                                             0.3672
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 65.945 on 35 degrees of freedom
## Residual deviance: 28.249 on 26 degrees of freedom
## ATC: 113.1
##
## Number of Fisher Scoring iterations: 5
```

As can be seen by the results of the Poisson model above, it is seen that the variables oligarchy, pollib, and parties are seen as significant in predicting the number of successful military coups from independence to 1989.

#### 4 b)

```
# prints just the top 3 least significant coefficients of a
# model:
evalModel = function(model, name) {
    cat(sprintf("\n%s:\n", name))
    res = summary(model)$coefficients
    print(head(res[order(res[, 4], decreasing = T), ], n = 3))
}
coupsglm1 = glm(miltcoup ~ oligarchy + pollib + parties + pctvote +
    popn + size + numelec + numregim, family = poisson, data = coups)
evalModel(coupsglm1, "coupsglm1")
##
## coupsglm1:
               Estimate Std. Error z value Pr(>|z|)
                            0.9976 - 0.234
## (Intercept)
                -0.2334
                                               0.815
## numelec
                -0.0296
                            0.0696
                                    -0.425
                                               0.671
## numregim
                 0.2109
                            0.2339
                                     0.902
                                               0.367
# numelec has highest p-value (0.6705) so we remove it
coupsglm2 = glm(miltcoup ~ oligarchy + pollib + parties + pctvote +
    popn + size + numregim, family = poisson, data = coups)
evalModel(coupsglm2, "coupsglm2")
##
## coupsglm2:
                Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -0.457746
                           0.860234
                                     -0.532
                                                0.595
## numregim
                0.180442
                           0.224117
                                       0.805
                                                0.421
```

```
-0.000269
                          0.000269 -1.000
                                               0.317
## size
# numregim has highest p-value so we remove it
coupsglm3 = glm(miltcoup ~ oligarchy + pollib + parties + pctvote +
   popn + size, family = poisson, data = coups)
evalModel(coupsglm3, "coupsglm3")
##
## coupsglm3:
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) 0.041976 0.577410 0.0727
                                               0.942
## size
              -0.000258
                          0.000266 -0.9688
                                               0.333
               0.007165
                          0.005684 1.2604
                                               0.208
## popn
# size has highest p-value so we remove it
coupsglm4 = glm(miltcoup ~ oligarchy + pollib + parties + pctvote +
   popn, family = poisson, data = coups)
evalModel(coupsglm4, "coupsglm4")
##
## coupsglm4:
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.23143
                           0.52889 -0.438
                                             0.662
                                     1.032
                                              0.302
## popn
               0.00566
                           0.00548
              -0.68359
## pollib1
                           0.49582 - 1.379
                                              0.168
# popn has highest p-value so we remove it
coupsglm5 = glm(miltcoup ~ oligarchy + pollib + parties + pctvote,
    family = poisson, data = coups)
evalModel(coupsglm5, "coupsglm5")
##
## coupsglm5:
              Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -0.1165
                          0.51375 -0.227
                                              0.821
               -0.6208
                                              0.203
## pollib1
                           0.48753 - 1.273
                0.0121
                          0.00907
                                   1.329
                                             0.184
## pctvote
# pctvote has highest p-value so we remove it
coupsglm6 = glm(miltcoup ~ oligarchy + pollib + parties, family = poisson,
    data = coups)
evalModel(coupsglm6, "coupsglm6")
##
## coupsglm6:
              Estimate Std. Error z value Pr(>|z|)
##
                                            0.6407
## (Intercept)
                 0.208
                        0.446
                                    0.467
```

The number of explanatory variables was reduced following the step-down approach, and in doing so the variables were removed in the following order: numelec, numregim,  $size\ popn$ , pctvote. This left the model with only relevant explanatory variables (significant as P < 0.05). These variables were the same as those seen as relevant in the previous model in 4a), namely oligarchy, pollib, and parties.

## 4 c)

## [1] "country2 prediction = 1.772"
## [1] "country3 prediction = 0.956"

```
country1 = data.frame(oligarchy = mean(coups$oligarchy),
   pollib = factor(0), parties = mean(coups$parties))
country2 = data.frame(oligarchy = mean(coups$oligarchy),
   pollib = factor(1), parties = mean(coups$parties))
country3 = data.frame(oligarchy = mean(coups$oligarchy),
   pollib = factor(2), parties = mean(coups$parties))

printResults = function() {
   print(sprintf("country1 prediction = %.3f", predict(coupsglm6, country1, type="response")))
   print(sprintf("country2 prediction = %.3f", predict(coupsglm6, country2, type="response")))
   print(sprintf("country3 prediction = %.3f", predict(coupsglm6, country3, type="response")))
}
printResults()

## [1] "country1 prediction = 2.908"
```

Using the model from 4b), we found that the coefficient of *pollib* was significant and negative. This entails that with an increase in the value of the political liberalization (i.e. with an increase in civil rights), there would be a decrease in the predicted number of successful military coups. The predictions above confirm this.